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Antimonene: a tuneable post-graphene material for advanced applications in optoelectronics, catalysis, energy and biomedicine

Jose A. Carrasco, † Pau Congost-Escoin, † Mhamed Assebban and Gonzalo Abellán *

The post-graphene era is undoubtedly marked by two-dimensional (2D) materials such as quasi-van der Waals antimonene. This emerging material has a fascinating structure, exhibits a pronounced chemical reactivity (in contrast to graphene), possesses outstanding electronic properties and has been postulated for a plethora of applications. However, chemistry and physics of antimonene remain in their infancy, but fortunately recent discoveries have shed light on its unmatched allotropy and rich chemical reactivity offering a myriad of unprecedented possibilities in terms of fundamental studies and applications. Indeed, antimonene can be considered as one of the most appealing post-graphene 2D materials reported to date, since its structure, properties and applications can be chemically engineered from the ground up (both using top-down and bottom-up approaches), offering an unprecedented level of control in the realm of 2D materials. In this review, we provide an in-depth analysis of the recent advances in the synthesis, characterization and applications of antimonene. First, we start with a general introduction to antimonene, and then we focus on its general chemistry, physical properties, characterization and synthetic strategies. We then perform a comprehensive study on the allotropy, the phase transition mechanisms, the oxidation behaviour and chemical functionalization. From a technological point of view, we further discuss the applications recently reported for antimonene in the fields of optoelectronics, catalysis, energy storage, cancer therapy and sensing. Finally, important aspects such as new scalable methodologies or the promising perspectives in biomedicine are discussed, pinpointing antimonene as a cutting-edge material of broad interest for researchers working in chemistry, physics, materials science and biomedicine.

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1. Introduction

Since the isolation of phosphorene, the two-dimensional (2D)-pnictogen (P, As, Sb and Bi) science has emerged as one of the main topics in 2D materials.^{1–11} Indeed, 2D materials from group 15 of the periodic table offer a great variety of structures, properties and reactivities as a consequence of the progressive increase in the intensity of the interlayer bonds as one goes down the group, passing from purely van der Waals (vdW) materials such as phosphorene to materials with a marked covalent character that makes them quasi-vdW, as is the case with heavy pnictogens (Sb & Bi).^{12–15} The importance of these interactions in the physical and chemical properties of 2D-pnictogens is crucial and differentiates them from the rest of Xenes. This makes heavy 2D-pnictogens, and antimonene in particular, one of the most attractive materials in the field, as it opens the door to chemically engineer their properties from scratch.

Instituto de Ciencia Molecular (ICMol), Universidad de Valencia, Catedrático José Beltrán Martínez, 2, 46980 Paterna, Spain. E-mail: gonzalo.abellán@uv.es
† These authors contributed equally to this article.

This review provides a comprehensive analysis and updated study of antimonene, a material that since its prediction in 2015 and later isolation in 2016 has attracted increasing attention due to its unique electronic properties (thermoelectric, semiconducting, topological insulator, nonlinear optic, *etc.*) and its unmatched atomic structure and chemical flexibility, which allow its exotic (inter-allotrophic) phase engineering. Moreover, this review thoroughly analyses (>200 articles) the state-of-the-art of antimonene research, highlighting the recent progress in its experimental preparation with a strong emphasis on its properties, chemical reactivity, functionalization, and the potential applications reported to date. Despite the fact that there are a few reviews focused on pnictogens as a family of emerging materials, this work intends to address the antimonene state-of-the-art from in-depth and critical chemical points of view. To the best of our knowledge this approach has not been addressed to date. Indeed, in a post-graphene era, antimonene holds great promise for the development of fundamental science such as novel van der Waals (and beyond) heterostructures, molecular interface engineering, DNA sensors, high-performance (opto)electronic devices, modern



Na-ion batteries and many other scientific challenges to come. Scheme 1 summarizes the different synthetic approaches and main applications of this fascinating material, which will be detailed below, including a brief historical note about the origin of antimony and its properties.

1.1 History of antimony and general properties

Antimony (symbol Sb, from the Latin *stibium* and derived from the Greek word referring to the use of stibnite) has been known since ancient times (*ca.* 4000 BC) and has been used in the form



Jose A. Carrasco

ICMol (2019–2020) working on 2D-pnictogens, and at the Institute of Chemical Technology (ITQ) under the supervision of Dr. Antonio Leyva-Pérez, working in catalysis (CSIC-UPV, 2021). Now he is working in his true passion as a teacher of physics and chemistry, sharing his love for science to the younger generations.

Jose Alberto Carrasco obtained his degree in Chemistry from the University of Valencia (UV) in 2012. Afterwards, he received his Master's degree in Nanoscience and Nanotechnology in 2014. From 2014 to 2018 he pursued his PhD supervised by Prof. Eugenio Coronado and Dr. Gonzalo Abellán at the Institute of Molecular Science (ICMol, UV), focusing on inorganic two-dimensional materials. Thereafter, he worked as a postdoctoral fellow in the 2D-Chem group at

of cosmetics, medications, colouring agent for glass and casting pottery, to name a few. In this sense, ancient Egyptians used antimony in the form of stibnite (Sb_2S_3) as a rouge for lips. Similarly, ancient Romans used stibnite in a medical way or as black eye make-up.¹⁶ The name antimony comes from two different Greek words: *anti*, which means *not* and *monos*, whose meaning refers to *alone*. This compound noun is due to the rare appearance of isolated antimony in nature while is commonly found as combined with both metals and non-metals.



Pau Congost-Escoin

interactions of 2D materials in the biological environments and the mechanisms governing them.

Pau Congost-Escoin received his BSc in Biochemistry and Biomedical sciences from the University of Valencia (UV) in 2020, and a Master's degree in Nanoscience and Nanotechnology in 2021, when he started working on 2D materials. He is currently a PhD student in Nanoscience and Nanotechnology at ICMol (UV), under the supervision of Dr. Gonzalo Abellán, working on the biomedical applications of 2D pnictogens. His research interest lies in the



Mhamed Assebban

(ZMP, FAU) in Germany as a postdoctoral researcher in 2018, and then he moved to the Institute of Molecular Science (ICMol) at the University of Valencia in Spain under the supervision of Dr. Gonzalo Abellán. In 2020 he returned back to occupy the same position at the ZMP in Fürth where he focuses his research work on the study of novel 2D materials by tuning their physical and chemical properties to make them even more suitable for different applications, such as energy storage, catalysis and optoelectronics.

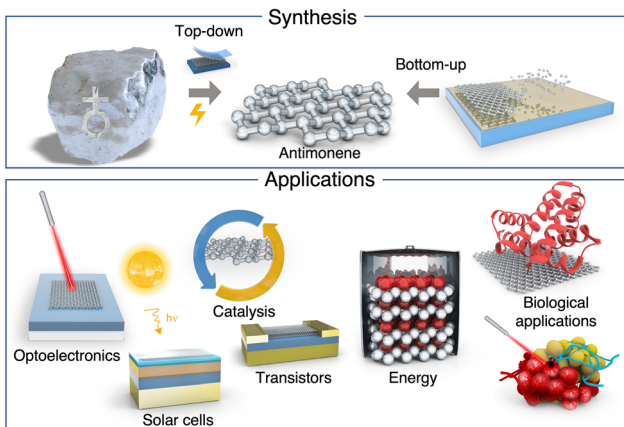
Mhamed Assebban obtained his BSc in Chemical engineering and materials science at the University of Abdelmalek Essaadi in 2009 and his Master's degree in 2011. He obtained his PhD in 2016 at the same university, in a collaborative research project with the University of Bielefeld in Germany, where he developed efficient heterogeneous catalysts for the removal of toxic atmospheric gases. He joined the Zentralinstitut für neue Materialien und Prozesstechnik



Gonzalo Abellán

Gonzalo Abellán obtained his PhD in nanoscience and nanotechnology in 2014 from the University of Valencia. Afterwards he gained a self-driven Marie Curie Fellowship and joined Prof. Andreas Hirsch's group at the Friedrich-Alexander-Universität, Erlangen-Nürnberg. In 2018 he returned to Spain as an Excellence Distinguished Researcher after getting a GenT-CIDEGENT contract (Generalitat Valenciana), the Ramón y Cajal fellowship, and the ERC Starting Grant, among others. Gonzalo Abellán's main research activities have been focused on three lines: 2D-pnictogens (group of P, As, Sb and Bi), layered hydroxides, and carbon nanoforms and related hybrid materials, with a special emphasis on energy-related applications. He is the leader of the 2D-Chem research group at the Institute of Molecular Science (ICMol, UV).





Scheme 1 Summary of antimonene synthesis and applications. The image above represents (top) the synthesis of antimonene from bulk antimony or via the assembly of antimony atoms on a substrate and (bottom) the applications of antimonene as a multifunctional material. From left to right: optoelectronic devices, solar cells, catalysis, field-effect transistors, energy storage, biomedical applications by binding molecules of interest and destroying carcinogenic cells.

The use of Sb was also described by medieval alchemists in the 15th century. The element intrigued the alchemists due to its similar properties to those of gold and the fact that it cannot be dissolved with aqua regia, hence being a good candidate for its transmutation into gold.¹⁷ During the 16th century, Paracelsus and his followers promoted the use of antimony and other metals as drugs. This statement was in stark contrast with the teachings of Galen (*ca.* 2nd century) who considered the use of metals as poisonous.¹⁸ In this context, the period between *ca.* 1560 and 1660 was coined as the antimony war due to the strong conflict between the Galenic medicine against the medical practices of Paracelsus.

Even though the original discoverer of Sb remains unknown, Nicolas Lémery, a French chemist, was the first one to scientifically study both antimony and its compounds, publishing his findings in 1707. Antimony compounds have been used for centuries in the treatment of schistosomiasis and leishmaniasis;¹⁹ however, one of the most recognisable aspects of antimony is its toxicity,²⁰ since it can even cause death by intoxication and its symptoms are disguised as general gastric disorders. The poisonous activity comes from its ability to be attached to particular enzymes because of its high affinity to the sulphur atoms of the enzyme's active site. Regarding the different species, one of the deadliest compounds of antimony is the gas stibine (SbH_3). Most antimony compounds such as stibine or oxides (Sb_2O_3 , Sb_2O_5 , *etc.*) can be absorbed from the respiratory tract due to the low particle size, therefore being retained in the organism for longer periods than larger particles.²¹

From the economic point of view and regarding the applications of antimony and its compounds, the element is a metalloid with an important impact. The actual annual production is at around 185 000 tonnes per year, with 85% coming from China. Additional producers can be found in countries like

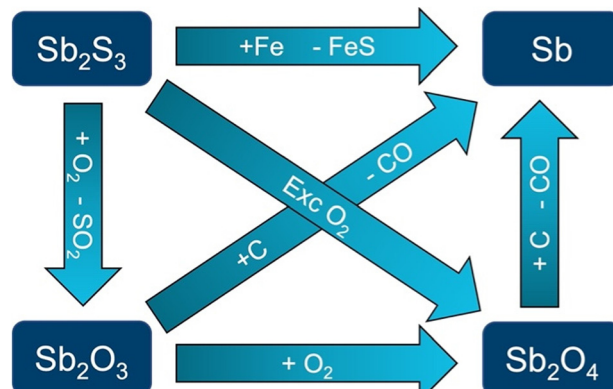


Fig. 1 Schematic representation of the production of elemental antimony from stibnite. Adapted from ref. 23 with permission from John Wiley & Sons, copyright 2011.

Russia, South Africa or Bolivia, among others. The main ores are in the form of stibnite and tetrahedrite, this one being a copper antimony sulfosalt mineral with formula: $(\text{Cu}, \text{Fe})_{12}\text{Sb}_4\text{S}_{13}$, which yields Sb as a by-product.¹⁷ Stibnite is easily separated from other minerals due to its low melting point of 546 °C. Afterwards, the roasting of stibnite produces Sb_2O_3 or Sb_2O_4 which can be both reduced to elemental Sb with coke. An alternative method to obtain Sb is to directly reduce the stibnite with scrap iron, followed by a purification procedure with sodium nitrate and sodium carbonate.^{22,23} Fig. 1 schematizes the production of elemental Sb from stibnite.

With respect to the main applications, antimony compounds can be used in different fields such as flame retardants (Sb_2O_3), catalysis (SbF_5), pyrotechnical articles ($\text{Sb}_2\text{S}_3 + \text{H}_2\text{S}$) or in electronics (alloyed with Ga and As).^{17,24} Related to electronic applications, Sb is mainly used in semiconductor devices such as diodes and infrared detectors. Alloying with Pb or other metals can enhance its hardness and strength. In this sense, a wide variety of alloys are described with Na, K, Ag, Au, Mg, Zn or Al, to name a few,²⁵ leading to additional applications like cable and bullet sheathing as well as its use in batteries in the form of a Sb–Pb alloy.²⁶

From the chemical point of view, Sb is considered a metalloid element, a member of the group 15 in the periodic table, *i.e.*, the family of pnictogens and with an electronic configuration of $[\text{Kr}] 4d^{10} 5s^2 5p^3$. Regarding the position of pnictogens in the periodic table, it is not unusual to come across some papers that refer to group VA instead of group 15. Here, we want to point out the fact that this old nomenclature based on roman numbers and capital letters A/B has been discarded by the IUPAC since 1990.²⁷ Therefore, the only correct labelling nowadays for the pnictogen group is 15. Sb can appear in two different forms: while the metallic one is bright, silvery, hard and brittle (and resembles Pb), the non-metallic one is a grey powder. Furthermore, it poorly conducts both electricity and temperature, whereas it shows good stability in dry air and is not affected by dilute acids or alkalis. Additionally, upon cooling, Sb and some of its alloys expand.²⁸ The most common



oxidation states are (+3) and (+5) as found in nature. Some typical compounds for the trivalent and pentavalent antimony are SbCl_3 and Sb_2O_5 , respectively. However, additional oxidation states of (-3) (stibine, SbH_3) and (0) (Sb metal) are also described,²⁹ as well as mixed oxidation state compounds.³⁰ The inorganic chemistry of Sb is extremely rich and mostly focused on both (+3) and (+5) oxidation states. In this context, there are several studies reporting antimony hydrides (SbH_3), compounds with elements of group 13 or 14 (GaSb, $\text{Sb}(\text{SiMe}_3)$), with other pnictogens such as N or P ($\text{Sb}(\text{NR}_2)$), antimony chalcogens (Sb_2O_3 , $\text{Sb}_2(\text{SO}_4)_3$, Sb_2S_3), antimony halides (SbCl_3 , SbF_5) or antimonides with Sb^{3-} such as Li_3Sb or Na_3Sb , to name a few.²³ Although it appears mainly in inorganic compounds, Sb also has a rich organometallic chemistry.³¹ After the synthesis of the first trialkyl antimony compounds in around 1850 – trimethyl- and triethylstibine – an incredible variety of organoantimony compounds have been reported to date, mainly in the oxidation states of (+3) and (+5). Trivalent Sb can be found in tertiary stibines, organoantimony(III) halides and pseudohalides, diorganoantimony cations and anions, or organoantimony hydrides, to name a few. On the other hand, regarding the pentavalent organoantimony compounds, one can find triorganoantimony(V) compounds, penta alkyls and aryls or organoantimony(V) chalcogen and nitrogen derivatives. Additionally, there are other organometallic compounds such as transition metal complexes with organoantimony ligands, low-coordination organoantimony derivatives and organometallic compounds with covalent single Sb–Sb bonds, including distibines, cyclotribines catena-stibines or polycycles.³²

The abundance of the element corresponds to *ca.* 0.00002% of the Earth's crust, and with respect to the existence of

isotopes, there are 37 known Sb isotopes, with masses that vary from 103 to 139 u. In this context, there are two naturally stable isotopes: Sb-121 (57%) and Sb-123 (43%). Some isotopes are radioactive and produced after fission experiments. Among these, Sb-125 is the longest-lived radioisotope, with a half-life of 2.76 years.³³ Additional elemental parameters of Sb can be found in Table 1.

Concerning the existence of allotropes, elemental bulk Sb has three known allotropes: explosive, black and gray.²² The most common form is the gray allotrope, with a rhombohedral structure like gray arsenic and a typical semi metallic behaviour. Black antimony is analogous to red phosphorus and explosive antimony usually evolve to the gray one after mechanical stress or on heating.³⁴ While other pnictogens such as P and As depict α and β layered stable phases, the heavy pnictogens Sb and Bi have their most stable phase as the β allotrope even though other allotropes are theoretically predicted and recently obtained under certain conditions, which will be detailed below.^{35,36} Fig. 2 summarizes the different layered crystal structures and stable phases.³⁴

1.2 2D materials: cutting-edge for the 21st century

After the rise of graphene in 2004,^{37–39} a myriad of new graphene-like two-dimensional (2D) materials have arisen.^{40–43} The number of documents by year related to the topic of 2D materials keeps growing and has reached a maximum of *ca.* 17 000 papers by mid-2022 (source: Scopus). Fig. 3 depicts the total number of papers related specifically to 2D-pnictogens. This trend is motivated for the outstanding properties of these materials towards new applications in the fields of

Table 1 Elemental parameters for antimony. Some data are extracted from ref. 51

Parameter	Value	Parameter	Value
Year of discovery	Known from <i>ca.</i> 4000 B.C., studied by Nicolas Lémery (1707)	Magnetic type	Diamagnetic
Alchemy symbol	⊕	Molar magnetic susceptibility ($\text{m}^3 \text{ mol}^{-1}$)	-1.327×10^{-9}
Atomic number	51	Electronic shell	[Kr] 4d ¹⁰ 5s ² 5p ³
Atomic mass (g mol ⁻¹)	121.75	Standard potential (V)	0.21 (Sb^{3+}/Sb) 0.60 ($\text{Sb}_2\text{O}_5/\text{SbO}^+$) -0.51 (Sb/SbH_3)
Density (g cm ⁻³)	6.684	Natural isotopes, abundance and radioactive stability	¹²¹ Sb (57.4%, stable) ¹²³ Sb (42.6%, stable)
Melting point (°C)	631	Abundance in Earth's crust	0.00002%
Boiling point (°C)	1587	Nuclear spin, I	¹²¹ Sb (+5/2) ¹²³ Sb (+7/2)
Heat of fusion (kJ mol ⁻¹)	19.7	Ionization energies (kJ mol ⁻¹)	
Heat of vaporization (kJ mol ⁻¹)	67	$\text{M} \rightarrow \text{M}^+$	833.7
Specific heat (J kg ⁻¹ K ⁻¹)	207	$\text{M}^+ \rightarrow \text{M}^{2+}$	1794
Thermal conductivity (W m ⁻¹ K ⁻¹)	24	$\text{M}^{2+} \rightarrow \text{M}^{3+}$	2443
Bulk modulus (GPa)	42 GPa	$\text{M}^{3+} \rightarrow \text{M}^{4+}$	4260
Shear modulus (GPa)	20	$\text{M}^{4+} \rightarrow \text{M}^{5+}$	5400
Young modulus (GPa)	55	Electron affinity (kJ mol ⁻¹)	101
Mohs hardness	3	$\text{M} \rightarrow \text{M}^-$	
Space group name	$R\bar{3}m$	Electronegativity, χ_P (Pauling scale)	2.05
Electrical type	Conductor	Atomic radius (Å)	1.82
Electrical conductivity (S m ⁻¹)	2.5×10^6	Single-bond covalent radius (Å)	1.41
Resistivity (m Ω)	4×10^{-7}	van der Waals radius (Å)	2.20
		Ionic radii (Å)	M^{3+} 0.62 M^{5+} 0.76



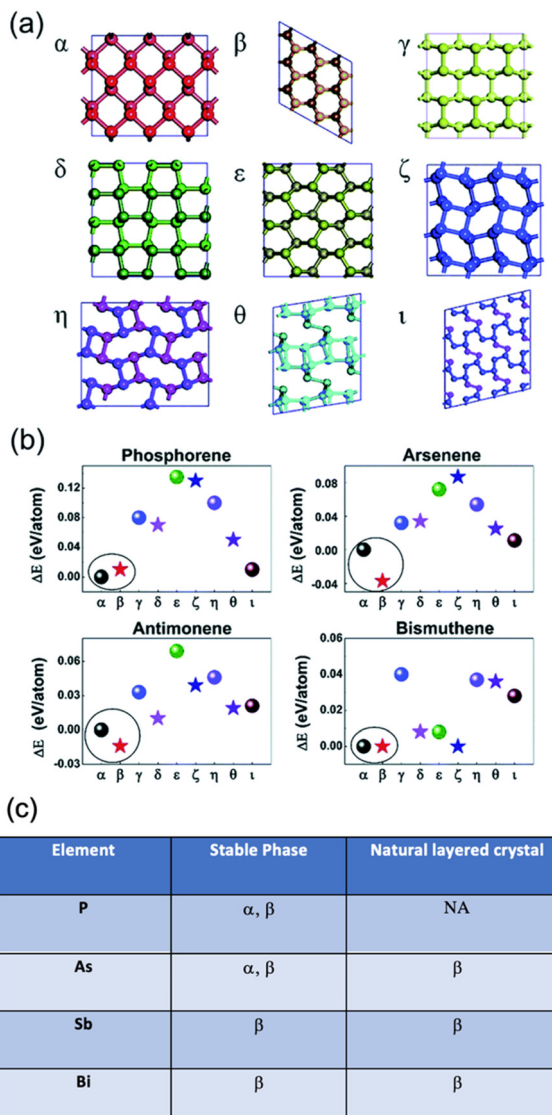


Fig. 2 (a) Typical honeycomb and non-honeycomb structures of 2D pnictogen allotropes. (b) Average binding energies of the different allotropes, highlighting the β phase as the most stable in average. (c) Table summarizing the stable phases of the different pnictogens. While α phase presents a parallel puckered layer with space group of $Cmca$, β phase exhibits parallel buckled layers with the $R\bar{3}m$ space group in a rhombohedral structure. Adapted from ref. 34 with permission from Royal Society of Chemistry, copyright 2018.

energy, electronics, catalysis, topological insulators, thermal management or anisotropic magnetization.⁴⁰

The wide variety of 2D materials that are described in the literature can be divided into three main groups: the layered van der Waals (vdW) solids, the 3D covalent/ionic solids, and the layered ionic solids. This classification is based on the main forces that ensemble the layered structure, *i.e.*, intermolecular van der Waals interactions or electrostatic forces. While in the first group one can find materials such as graphene, boron nitride or layered metal dichalcogenides,^{40,44} the second group is composed of covalent materials that can be exfoliated down

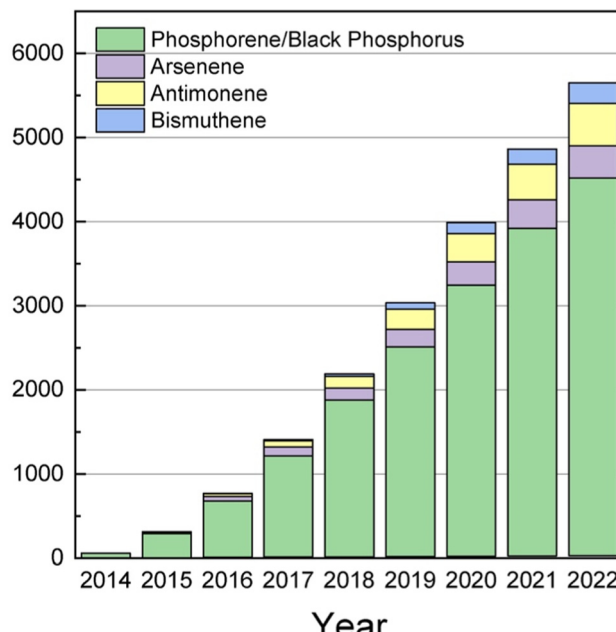


Fig. 3 Graphic showing the cumulative number of papers focused on 2D pnictogens. Data acquired from Scopus on October 2022.

to the 2D limit (*i.e.* germanene) or 3D ionic solids (like iron oxide) that can be exfoliated by controlling parameters such as the energy of cleavage planes or the stabilization energy of the nanolayers. Finally, the third group includes charged 2D layers that are held together, thanks to the presence of cations or anions in the interlayer space. This is the case of layered metal oxides, layered double hydroxides or cation-exchanged layers from Ruddlesden–Popper perovskite-type structures.^{45–50} Table 2 shows some of the reported 2D materials and Fig. 4 depicts how these can be assembled together to build van der Waals heterostructures.

The group of pnictogens (P, As, Sb and Bi) exhibits several allotropes with layered structures, both with rhombohedral and orthorhombic structures.³⁴ In this sense, true van der Waals forces held the layered assembly in the orthorhombic structure, while the interlayer interactions of the rhombohedral counterparts increase concomitantly with the atomic number, *i.e.*, the nature of the pnictogen.⁵³ The first one to be described was phosphorene in 2014, the 2D counterpart of layered black phosphorus, which exhibited great potential as a p-type semiconducting material (Fig. 5). The authors described a few-layer phosphorene field-effect transistor with a channel length of 1.0 μm at room temperature, which displayed a high on-current of 194 mA mm^{-1} together with a high hole field-effect mobility of 286 $\text{cm}^2 \text{V}^{-1} \text{s}^{-1}$ and an on/off ratio of up to 10^4 .⁵⁴ Phosphorene exhibits an orthorhombic crystalline form in which the P atoms are assembled composing hexagonal puckered layers that are sustained thanks to true van der Waals forces. It displays high reactivity as well as a rapid oxidation under ambient conditions leading to the formation of P_xO_y compounds which, in the presence of moisture, leads to the formation of phosphoric acid and related compounds.⁵⁵ Furthermore, it has a modulable



Table 2 Reported monolayers in the literature classified into different families according to its composition. Adapted from ref. 42 and 52

Carbon materials	Graphene	Hexagonal boron nitride (white graphene) and boron carbon nitride	Graphene oxide	Fluorographene
Xenes	Group 13: borophene, gallene	Group 14: graphene, silicene, germanene, stanene	Group 15: phosphorene, arsenene, antimonene, bismuthene	Group 16: selenene, tellurene
2D Chalcogenides	MoS ₂ , WS ₂ , MoSe ₂ , WSe ₂	Semiconducting dichalcogenides: MoTe ₂ , WTe ₂ , ZrS ₂ , ZrSe ₂ , etc.	Metallic dichalcogenides: NbSe ₂ , NbS ₂ , TaS ₂ , TiS ₂ , NiSe ₂ , etc.	Layered semiconductors: GaSe, GaTe, InSe, Bi ₂ Se ₃ , etc.
2D oxides and hydroxides	Micas, layered Cu oxides, etc.	MoO ₃ , WO ₃ , TiO ₂ , MnO ₂ , V ₂ O ₅ , CuO, RuO ₂ , etc.	Perovskite-type: LaNb ₂ O ₇ , (Ca,Sr) ₂ Nb ₃ O ₁₀ , Bi ₄ Tl ₃ O ₁₂ , Ca ₂ Ta ₂ TiO ₁₀ , etc.	Simple and double hydroxides: Ni(OH) ₂ , Eu(OH) ₂ , NiFe-LDH, CoAl-LDH, etc.

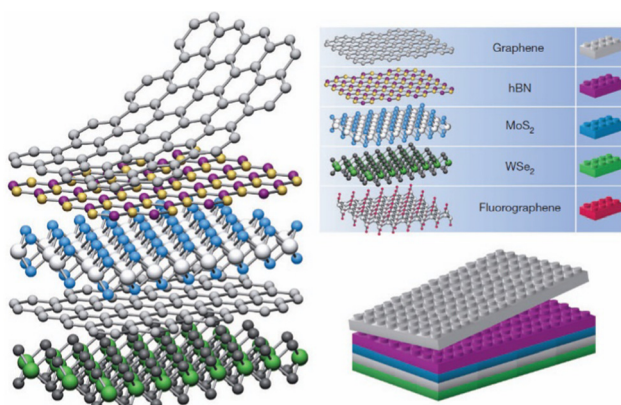


Fig. 4 Building van der Waals heterostructures by combining different layers of 2D materials. Adapted from ref. 42.

bandgap that ranges from 0.3 eV to 2.2 eV for bulk and monolayer black phosphorus, respectively.⁵⁶ Since then, phosphorene has been extensively studied, including its exfoliation,^{57–59} its covalent and noncovalent functionalization^{55,60,61} as well as its different applications.^{55,62–65} Analogous 2D pnictogens (arsenene, antimonene and bismuthene) have been also predicted and most of them have been experimentally obtained.^{53,66}

In this sense, it is important to point out that the -ene suffix in these materials (phosphorene, antimonene, bismuthene as well as others such as germanene or silicene) does not refer to a double bond between the individual atoms as in graphene. Nevertheless, these sorts of 2D materials composed of one single element are commonly referred to as Xenes by the scientific community.⁶⁷ The layered structure of As corresponds to rhombohedral gray arsenic, and exhibits a tunable bandgap depending on the number of stacked layers of the material, ranging from a predicted 0.47 to 2.49 eV for a bilayer and a monolayer arrangement, respectively.⁶⁸ Studies concerning arsenene are very scarce; nevertheless, there are some reports regarding the preparation of multilayer arsenene with a rhombohedral structure. For example, in the form of applying nitrogen plasma with indium arsenide, giving rise to the synthesis of multilayer arsenene nanoribbons on the surface of the InAs.⁶⁹ Regarding heavy pnictogens (Sb & Bi), the fact of descending through the group implies an increase in interlayer interactions, going from being purely vdW materials to having a

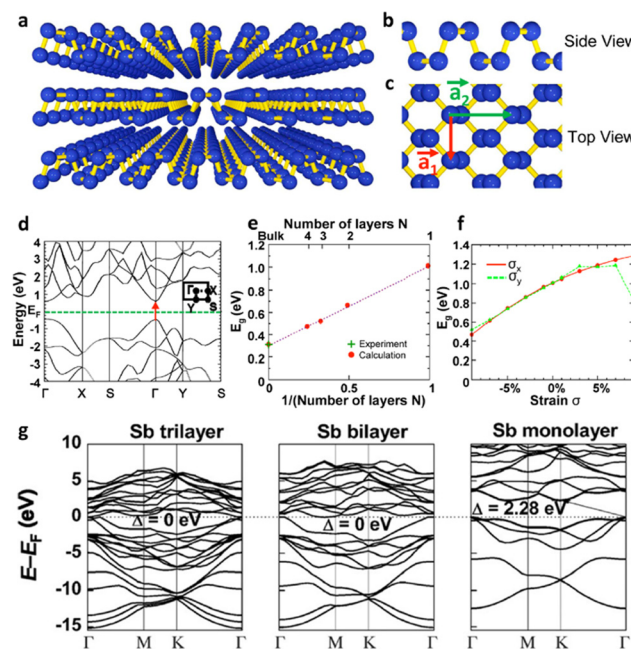


Fig. 5 Crystal structure and band structure of few-layer phosphorene. (a) Perspective side view of few-layer phosphorene. (b) and (c) Side and top views of few-layer phosphorene. (d) Theoretical band structure of a phosphorene monolayer calculated using DFT-HSE06. (e) and (f) DFT-HSE06 results for the dependence of the energy gap in few-layer phosphorene on (e) the number of layers and (f) the strain along the x- and y-directions in a monolayer system. (g) Representation of the electronic band structures of antimony trilayers, bilayers, and monolayers calculated using DFT-HSE06. Adapted from ref. 54 with permission from American Chemical Society, copyright 2014 and ref. 76 with permission from John Wiley & Sons, copyright 2015.

marked covalent character. Specifically, group 15 elements (without considering nitrogen) tend to favor the sp^3 hybridization, but in their $ns^2\ np^3$ valence configuration they tend to form layered structures showing in-plane covalent bonds together with weaker interlayer bonds, which are purely vdW in the case of phosphorene, but progressively incorporate interlayer orbital hybridization in heavier elements, thus forming quasi-vdW materials. This latter aspect is critical in the case of bismuth, thus precluding its micromechanical exfoliation. Moreover, the strong spin-orbit coupling characteristic of heavy pnictogens results in the fact that both known structures of bismuth (α and β forms) have been reported as topological



insulators, with predicted bandgaps of 0.18 and 0.23 eV for the bilayer α and β forms, respectively, that increase to 0.30 and 0.32 eV for the monolayer α and β -bismuthene, respectively.^{70,71} There are reports of the successful synthesis of few-layer bismuthene *via*, for example, liquid phase exfoliation giving rise to few-layer systems; however, the strong tendency to oxidation of this material usually leads to heavily oxidized systems with poor morphologies.^{72,73} Hence, recent studies on top-down bismuthene materials reveal promising results in energy storage (*i.e.* sodium- and potassium-ion batteries), electrocatalysis such as selectively catalyzing the electroreduction of CO₂ to HCOO⁻, or photonics.⁷³ Moreover, wet-chemical approaches have been reported involving thermal reduction or hot-injection reactions leading to 2D bismuth with different morphologies and degrees of oxidation.^{74,75}

Taking all this information into account, antimonene holds a privileged position in group 15 because despite being a heavy pnictogen, it still maintains a marked van der Waals character in its interlaminar interactions. This intermediate position allows us to exploit not only the physical applications of antimonene (*e.g.* optical or electronic) but also its pronounced chemical reactivity, of extreme importance in applications beyond electronics, such as biomedicine, sensing, energy or catalysis. In this review we will particularly focus on antimonene; we will give an overall view of the history and properties, later on we will thoroughly analyze the different synthetic methodologies and characterization techniques. We will also tackle the latest insights on chemical reactivity, including oxidation behavior and chemical functionalization, to finally address all potential applications reported so far for this brand-new material.

2. General physical and chemical properties of antimonene

2.1 History of antimonene: from the theoretical predictions to its experimental isolation

The story of antimonene begins in 2015 when Zhang *et al.* theoretically identified using density functional theory (DFT) the existence of novel 2D layered materials composed of Sb and As monolayers, which were labelled as antimonene and arsenene, respectively. These predictions anticipated that both materials would display good stability, high carrier mobility as well as a wide range of bandgaps according to the total number of layers.⁷⁶ That work showed that although Sb and As are mostly semi-metallic elements in bulk, they are both transformed into indirect semiconductors, highlighting a bandgap of 2.28 and 2.49 eV when thinned to the monolayer region (Fig. 5). In addition, antimonene and arsenene evolve from indirect semiconductors to direct bandgap semiconductors if a small biaxial strain is applied, exhibiting great potential for optoelectronic applications. In the modelling a buckled honeycomb structure was also predicted where each atom is bonded to three adjacent atoms of the same layer achieving octet stability. This arrangement is similar to those observed for

germanene and silicene and thus aids in the stabilization of the layered structure.⁷⁷ Fig. 5(g) presents the original electronic band structure of antimony trilayers, bilayers and monolayers (calculated at the HSE06 level of theory) as presented by Zhang *et al.*⁷⁶

After this initial study, numerous articles dealing with various theoretical aspects of antimonene appeared, pinpointing the interest of different properties of the material and postulating several potential applications.^{35,78-80} For instance, the studies by Karna and co-workers exhibited that free-standing α and β allotropes of antimonene are stable and semiconducting.³⁵ In this sense, while the α form has a puckered structure with two atomic sublayers, the β allotrope displays a buckled hexagonal lattice. Another interesting difference between both allotropes is the strong anisotropy of the α form which contrasts with the nearly isotropic mechanical properties of β -Sb. The possibility of the existence of additional allotropes (γ and δ) was also investigated, concluding in the structural instability of both of them compared with the α and β forms. The parameter chosen for analyzing the stability was the phonon dispersion curves, displaying no imaginary vibrating modes for the α and β allotropes, hence implying their stability as free-standing monolayers, a point additionally supported by using several exchange and correlational functional forms of DFT.³⁵ In contrast, γ and δ forms did exhibit imaginary vibrational modes, thus confirming their structural instability. This work also conveyed that a moderate tensile strain would induce a transformation from indirect to direct bandgap transition in antimonene, as well as a marked vibrational Raman bands that can be very useful in the characterization of monolayers. Fig. 6 presents, on the one hand, the structural configuration of antimonene allotropes, and, on the other hand, the predicted electronic properties of α and β allotropes. Interestingly, allotropic forms of antimonene have been experimentally investigated in the formation of van der Waals heterostructures, where it has been found that the rhombohedral β phase is the most stable,⁸¹⁻⁸³ but the orthorhombic α -phase can be metastable at the few-layer regime, showing a spontaneous transition from the orthorhombic to the rhombohedral bulk-like phase for thicknesses beyond 4 nm.⁸²

In 2016, Ares *et al.* were the first to obtain and experimentally isolate antimonene.⁸⁴ In this work, the micromechanical exfoliation of bulk crystals of Sb was carried out down to the single-layer regime, showing experimental evidence of the stability of antimonene under ambient conditions. Furthermore, the provided experimental data highlighting good stability and no degradation for months. Moreover, additional DFT simulations mimicked ambient conditions and confirmed the experimental results, predicting a bandgap of 1.2–1.3 eV corresponding to an interesting range of optoelectronics applications. Fig. 7 shows evidence of experimental antimonene flakes obtained by micromechanical exfoliation on SiO₂ substrates.

Transmission electron microscopy (TEM) of antimonene flakes confirmed a clear hexagonal periodicity as expected for the few-layer β -Sb phase. Regarding the feasibility of achieving a single-layer of antimonene, a monolayer terrace was observed



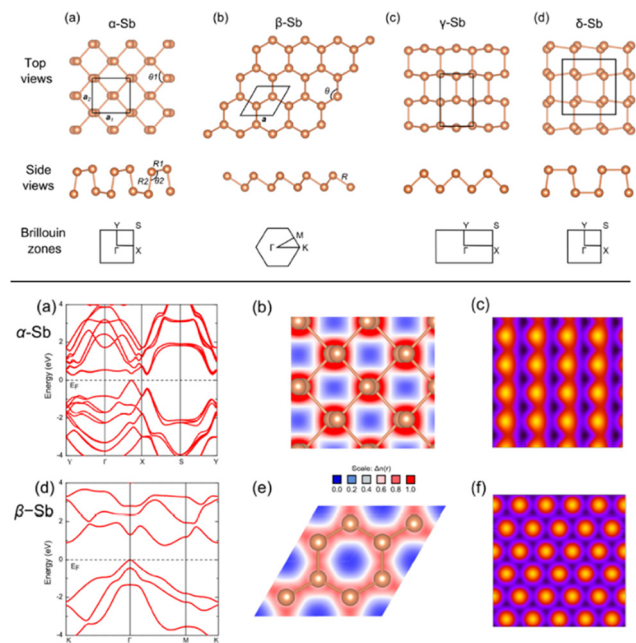


Fig. 6 Top: Structural configurations of antimonene allotropes: (a) α -Sb, (b) β -Sb, (c) γ -Sb, and (d) δ -Sb. Bottom: Electronic properties of (a)–(c) α -Sb and (d)–(f) β -Sb monolayers: (a), (d) band structure, (b), (e) charge density projected in the plane, and (c), (f) simulated scanning tunneling microscopy (STM) images. Adapted from ref. 35 with permission from American Chemical Society, copyright 2015.

with a height of *ca.* 0.9 nm that surely included additional water molecules. It is widely assumed that this layer of adsorbed

water molecules is always present under ambient conditions, with an average thickness of *ca.* 0.6 nm which is present between the deposited flakes of the material and the SiO_2 substrate.^{85,86} These pioneering studies paved the way for the development of the chemistry and physics of antimonene. Henceforth, hundreds of papers dealing with different aspects of antimonene have been published, corroborating the interest generated by this novel 2D material (Fig. 3).

2.2 Properties

Antimonene has been proposed as a good candidate for novel applications due to its outstanding properties. Nevertheless, one has to take into account that some of them have not been experimentally tested yet; therefore, the main properties of the material are a combination of the actual experimental and theoretical results as reviewed below:

2.2.1 Structural diversity. Antimonene presents many allotropic forms, creating a wide range of possibilities and tunability in terms of properties.⁸⁷ However, theoretical calculations have revealed that the only free-standing stable allotropes are the β and α allotropes. In this context, α and β allotropes of antimonene monolayers have been studied using DFT theory suggesting them as suitable materials for micro-electronic and optoelectronic nanodevices and solar cell applications (Fig. 6).⁸⁸

Nevertheless, the study of antimonene heterostructures revealed the plasticity of this novel material, the structure of which is critically dependent on its surroundings. This includes thickness driven phase transitions and substrate-layer stability dependence.^{82,83} This rich allotropic dynamism generates a

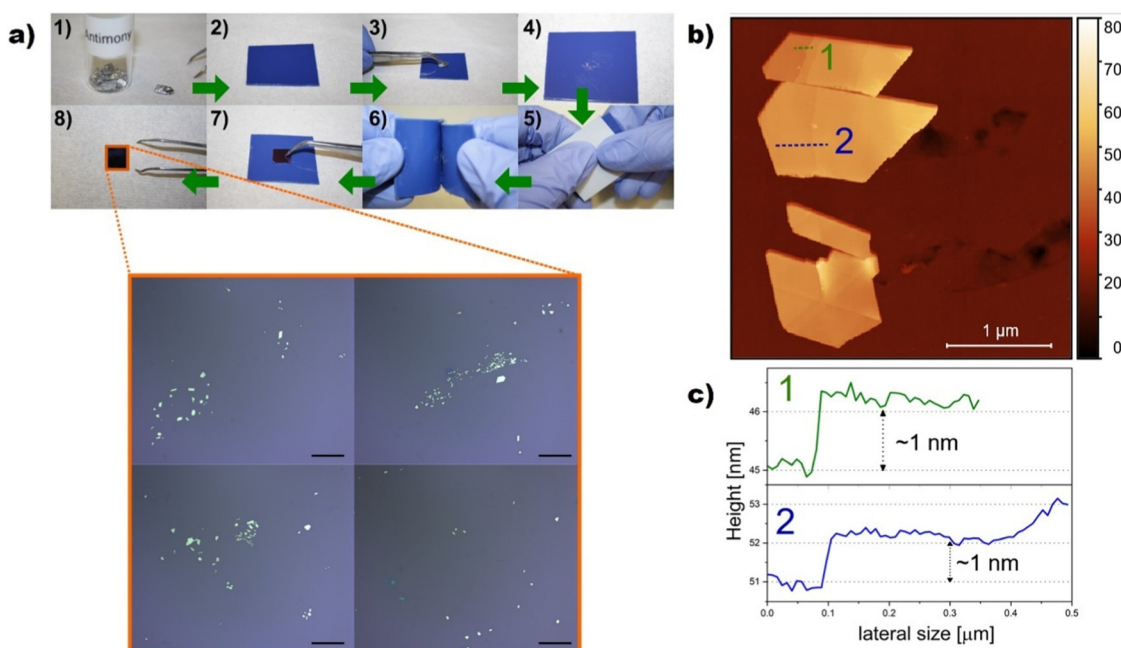


Fig. 7 (a) Step-by-step procedure (1–8) for the exfoliation of ANSs using micromechanical approach. Inset in (8) Optical micrographs showing exfoliated ANSs with different polygonal geometries. Scale bars 20 μm . (b) AFM image of typical ANSs and (c) height profiles along the dashed lines showing steps of approximately 1 nm, compatible with a single layer antimonene. Adapted from ref. 98 with permission from IOP Publishing, copyright 2020.



wide number of possible antimonene structures with unique properties that can be exploited specifically to meet the requirements of each application.

2.2.2 Mechanical properties. Concerning mechanical properties, and based on first-principles calculations, the tensile stress-strain of antimonene was investigated under uniaxial and equibiaxial strains. It was found that the ideal strength decreases with the increase of the atomic number, while the critical strain does not suffer strong deviations.⁸⁹ Furthermore, an increase of *ca.* 25% is observed in the Young's modulus of β -antimonene with respect to bulk Sb.⁹⁰ At the same time, the hexagonal lattice constant of the monolayer is significantly reduced (*ca.* 5%) from that in bulk, indicative of strong interlayer interactions. Regarding deformation, the integrity of the material is better preserved if the process takes place in the armchair direction. Applying uniaxial strain in both the armchair and zigzag directions leads to a direction-dependent trend in the electronic band structure. In this sense, it is observed that when the strain is applied in the zigzag direction, the nature of the gap is unaltered, yet if applied in the armchair direction it gives rise to an indirect–direct bandgap transition at a critical strain of 4%, motivated by a band switching mechanism.⁹⁰

2.2.3 Optical properties. From the optical properties point of view, the values of the refractive indices of α and β -phases have been investigated, revealing values of 2.3 and 1.5 at zero frequency and 3.6 and 2.0 in the UV region, respectively.⁹¹ Furthermore, one can expect this suitability towards new optical applications using various light emission, detection or modulation and manipulation functions, owing to the specific dielectric function, absorption, reflectivity, energy loss spectra and refractivity of the material. The high refractive indices found in the UV region of the electromagnetic spectrum allow its application as an inner layer in UV absorbers.

Moreover, with plasmon energies of *ca.* 9 eV, the combination of the dielectric function and electron energy loss spectroscopy (EELS) reveals sections of the electromagnetic spectrum where antimonene behaves as a metal.⁸⁸ Recent studies confirm these results and the applicability in optoelectronics.⁹² The authors remark the significant absorption from the visible to the ultraviolet region of α -antimonene that can be, thus, used as an adequate saturable absorber, while the β form displays an almost negligible absorption in the visible region.⁹²

2.2.4 Electric properties. As mentioned before, antimonene presents a semimetallic behaviour in the bulk, which is transformed into an indirect semiconductor in the monolayer region, with a bandgap of 2.28 eV.^{35,76} The possibility to modulate it into a direct bandgap by applying biaxial strain makes antimonene promising for optoelectronic applications.

Electrical transport in this material has also been recently studied, concluding that antimonene is expected to exhibit unusual transport properties due to its topological phase transition feature and quantum spin Hall edge states.⁹³ These intrinsically localized edge channels oriented to electrical transport can be controlled by applying an external electric field.⁹³

Furthermore, Ares *et al.* studied the morphological and electronic properties of micromechanically exfoliated antimonene, from single layer to few-layer regime.⁹⁴ They observed that the conductance of the antimonene sheet was constant with the number of layers, obtaining a carrier concentration similar to graphene under ambient conditions, and thus having a resistivity independent of the flake thickness ($1200 \Omega \text{ sq}^{-1}$) larger than that observed for graphene. The experimental measurements along with theoretical calculations suggest that the topologically protected surface states play a key role in antimonene electrical properties.

2.2.5 Thermal properties. The thermal properties of antimonene have been theoretically studied by Wang *et al.* using the Boltzmann transport equation, which revealed a low lattice thermal conductivity of $15.1 \text{ W m}^{-1} \text{ K}^{-1}$ at 300 K, thus demonstrating its potential applications for next-generation thermoelectric devices.^{95–97} This lattice thermal conductivity decreases with temperature, as expected for crystalline materials where phonon–phonon scattering rules the thermal conductivity in the high temperature range.⁹⁵ The low value at 300 K is related to the small group velocity, with low Debye temperature and large buckling height. Fig. 8(a) reflects the predicted lattice thermal conductivity of antimonene as a function of temperature. Experimentally, Fickert *et al.* studied the thermal and photothermal properties of antimonene nanosheets isolated on a SiO_2/Si substrate by micromechanical exfoliation.⁹⁸ Using a Raman spectroscopy-based thermometric measurement combined with numerical simulations of heat transfer, they estimated a thermal conductivity value of $90 \text{ W m}^{-1} \text{ K}^{-1}$ (Fig. 8(b) and (c)). A slight deviation of the estimated thermal

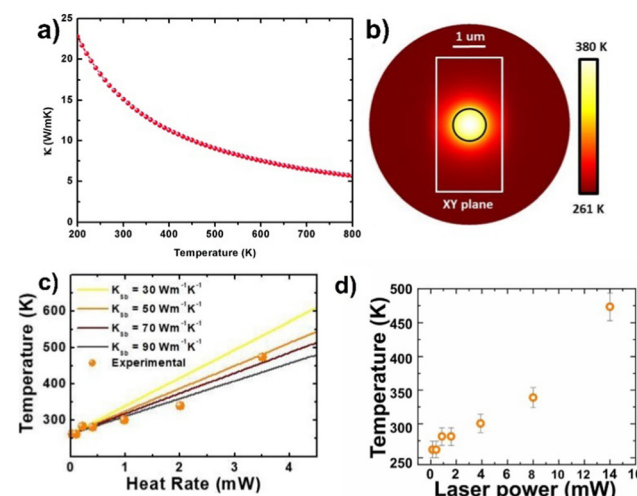


Fig. 8 (a) The lattice thermal conductivity of antimonene as a function of temperature. (b) Zenith view of the temperature mapping for a rectangular flake (white lines) considering a disk-like heat source (black circle). (c) Average temperature of the excitation spot as a function of the heat source rate for different thermal conductivities of the flake. (d) Temperature of the probed ANS as a function of laser power, estimated using both calculated power and temperature coefficients. Adapted from ref. 95 with permission from Royal Society of Chemistry, copyright 2016, and ref. 98 with permission from IOP Publishing, copyright 2020.



conductivity from the theoretically predicted values was attributed to reasons inherent to the method itself, such as the disregard of the convection loss and to the lack of accurate refractive index data of antimonene. Nevertheless, this estimated thermal conductivity remains the only experimental value reported so far, and further experimental investigations are still needed.

2.2.6 Magnetic behaviour. As far as magnetism is concerned, the reported magnetic nature of antimonene seems to be strongly influenced by the presence of vacancies, defects and doping. It is predicted that antimonene transits from a non-magnetic semiconductor to a magnetic half-metal as a consequence of vacancy inclusion.

Furthermore, the electronic and magnetic properties can be additionally modified by applying in-plane biaxial strains. When this strain is above 4%, the ferromagnetic half-metal nature will evolve into an antiferromagnetic semiconductor with no change in the total magnetic moment. Similarly, small compressive strains can reduce the magnetic response giving rise to an n-type doping semiconductive state at a strain of -4% . Regarding the amount of vacancy, a high number of these could even quench the induced magnetism.⁹⁹ These tunable magnetic response set antimonene as a good candidate towards the development of new low-dimensional spintronic devices.¹⁰⁰ With respect to the doping of the Sb layer with transition metals such as Cr, V, Ti or Fe, to name a few, the predicted influence on the magnetic behaviour diverges according to the chosen metal.¹⁰¹ For example, Cr-doped systems transform the magnetic semiconductor nature to a magnetic half-metal material after applying a biaxial tensile strain of 6%.

Also, the bandgap closing can be achieved under a biaxial strain between -4 to -6% , resulting in metallic features. The tensile strain also increases the local magnetic moments up to *ca.* 5.00 μ_B compared with the unstrained system.¹⁰² Fe-doped systems can lead to stable room temperature ferromagnetism, combined with a strong orbital hybridization (p-d) and a spin-orbit coupling effect which results in a significant spin splitting around the Fermi level. Introducing V atoms into an Fe-doped system changes the overall magnetic order from ferromagnetic to antiferromagnetic.¹⁰³ In this line, the structural and mechanical properties of antimonene monolayers doped with transition metals such as Sc, Ti, V, Cr, Fe, Co, Ni, Cu, and Zn have also been theoretically addressed *via* DFT calculations.¹⁰⁴ The study pointed out a reduction in the yield strain of the nanosheets under biaxial loading, resulting in the reduction of the Young's and bulk moduli. Overall, these results pinpoint strain and defect engineering as promising routes for the modulation of the magnetic behavior in antimonene.

2.2.7 Bulk chemical reactivity. The rich chemical reactivity of antimonene allows the formation of intermetallic phases, a source of novel materials with interesting physical and chemical properties, to be explored. In this sense, the combination of Sb with other elements giving rise to Zintl phases or alloys has been reported in the literature. First, Song *et al.* published the creation of the 2D layered Zintl phase by

dimensional manipulation of the crystal structure.¹⁰⁵ In that work, they established 2D polymorphism thanks to the discovery of a 2D layered structure in Zintl phases with a vast number of combinations with p-block metals. In this sense, it is expected that the 3D Zintl phases with sp^3 hybrid orbital bonding can transform to the sp^2 honeycomb 2D structure thanks to electron transfer processes.¹⁰⁶ Using this approach, a 2D layered structure of ZnSb is created after the successful modification of a 3D orthorhombic ZnSb. The strategy to obtain the resulting 2D Zintl phase consists of alloying with alkali metals such as Li, Na or K the 3D-ZnSb phase. Afterwards, the selective etching of the alkali metals can be carried out *via* chemical reaction in deionized (DI) water-incorporated solutions and electrochemical ion etching reaction in an alkali-based electrolyte. This methodology allows the use of phase transformations to create Zintl phases by manipulating the structural dimensionality broadening the scope of this sort of materials towards additional properties in 2D magnetism, ferroelectricity, thermoelectricity, and topological states.

Second, antimonene alloys have also been recently reported in different studies. For example, López-Marzo *et al.* demonstrated that 2D Sb_2Te_3 is suitable for optoelectronics and thermoelectric applications.¹⁰⁷ The antimony alloy was synthesized after putting the appropriate stoichiometric amount of Sb and Te in a quartz glass ampoule under high vacuum and posterior heating in the absence of air. Specifically, the few-layer and micrometer-sized Sb_2Te_3 was obtained after a glove box-free electrochemical exfoliation of the as-synthesized bulk material in Na_2SO_4 , which resulted in high yields and good stability. In this context, it is worth noting that the inherent oxidation of the alloy surface can be tuned by switching the voltage polarity during the exfoliation procedure. Antimonene alloys with arsenic can be found in different studies in the literature.^{108–110} 2D As_xSb_{1-x} (previously predicted by DFT^{109,110}) was experimentally grown *via* molecular beam epitaxy giving rise to a stable alloy under ambient conditions on Si, Ge or graphene substrates.¹⁰⁸ It was found that As composition can be tuned up to 15% using this co-deposition approach. The synthesis of new Sb alloys widens the scope of feasible applications towards new tunable devices. Further important reactivity aspects such as surface oxidation or functionalization will be thoroughly discussed in the following sections.

2.2.8 Electrochemical properties. Concerning electrochemical properties, antimonene exhibits a pronounced reactivity in both aqueous and organic media. Indeed, electrochemical sodiation/desodiation processes yielded a pronounced anisotropic volume change along the *a/b*-axis and partially reversible crystalline-phase transformations ($Sb \rightarrow NaSb \rightarrow Na_3Sb \rightarrow NaSb \rightarrow Sb$), leading to a high theoretical capacity of 660 mA h g^{-1} .⁷⁴ On the other hand, performance of antimonene as a hydrogen evolution reaction (HER) electrocatalyst seems to be somehow related to the irreversible formation of superficial antimony oxides (with different oxidation states) during the electrochemical procedure, as well as the thickness and the number of edges in the antimonene flakes, which contribute to the formation of electroactive sites (*vide infra*).^{111,112} This work



pinpoints the electrochemical importance of the antimonene surface oxidation species.

3. Synthetic approaches in the preparation of antimonene and its heterostructures

When producing 2D materials, including antimonene, it is crucial to have control over the thickness (*i.e.* number of layers), lateral dimensions, crystallinity or surface features in order to systematically study the properties of the material as well as point towards their specific applications. In broad strokes, one can divide the methods of synthesizing antimonene into two different groups: top-down and bottom-up approaches, as summarized in Fig. 9.^{113,114} Top-down refers to using exfoliation tools that are controlled by external forces like electrochemical potential, ultrasound or shear forces, to create nanoscale structures starting from bulk crystals and reducing them to the desired size, shape and features. In stark contrast, bottom-up approaches are based on using atomic or molecular components as building blocks for the direct self-assembly *via* specific reaction mechanisms, yielding 2D materials that can be prepared either free-standing or supported on surfaces.^{113,115,116} This strategy has allowed the preparation of the first antimonene-based heterostructures with other 2D materials like graphene, allowing the elucidation of the interface properties and several aspects related to the rich allotropic/polymorphic structural diversity. In the case of antimonene, both approaches have been reported in the literature and we will detail each of them below.

3.1 Top-down approaches

3.1.1 Micromechanical exfoliation (ME). This strategy is based on the paradigmatical “Scotch tape method” that was used to obtain graphene after successively peeling graphite flakes.^{38,85} It should be noted that the comparatively short

distance between the layers in bulk antimony and the higher strength of the interlayer interactions, makes it extremely difficult to strip-off thin antimony layers using the conventional Scotch-tape approach. This micromechanical exfoliation starting from a freshly cleaved crystal of β -antimony was used in the first isolation of antimonene by Zamora *et al.*⁸⁴ Fig. 7 depicts the resulting antimonene nanosheets obtained from millimetre-sized crystals of antimony. Initially, they observed that the direct transfer of the sub-millimetre flakes obtained from peeling the bulk Sb to a SiO_2 substrate led to a very low transfer yield. This problem was overcome by adding an initial transfer from the Scotch tape to a thin layer of a viscoelastic polymer adhered to a glass slide. As a result, a higher yield of deposited flakes was observed that were then transferred to the SiO_2 substrate. With this modification in the deposition method, the authors reported the presence of few-layer and single-layer antimonene in a more controlled way.⁸⁴

The as-obtained micrometric-flakes were then easily identified by means of optical microscopy. Then, AFM images depicted an average height of *ca.* 0.9 nm for a monolayer terrace at the bottom of a few-layer antimonene flake (Fig. 10). As previously commented, the 0.9 nm value takes into account the existence of water molecules trapped between the flake and the SiO_2 substrate.^{85,86} Nevertheless, statistics has to be taken into account when talking about AFM thicknesses. The mechanical stability of the monolayer terrace was also confirmed after folding it with no breakage by AFM-assisted nanomanipulation. Additional techniques such as high-resolution transmission electron microscopy (HRTEM) and X-ray energy dispersive spectroscopy (XEDS) in combination with high-resolution AFM confirmed the quality of the material and its rhombohedral structure, that is, corresponding to β -antimonene. Finally, the stability of the material under ambient conditions as a function of time (up to two months) was also studied. It was observed that not only is antimonene stable after that time but also is after the immersion of the sample in water, thus depicting the high overall stability of the material in the presence of moisture (the role exerted by surface oxidation will be discussed later, *vide infra*).⁸⁴

Recently, our group managed to introduce a deterministic method to easily isolate antimonene nanosheets (ANSs) through an adapted micromechanical exfoliation approach using a high-tack adhesive film.⁹⁸ Indeed, we have been able to isolate ANSs exhibiting well-defined polygonal shapes with smooth surfaces and sharp edges (Fig. 10(e) and (j)) in very high density over the SiO_2/Si substrate. Topographic analysis using AFM revealed ANSs in a few-layer regime with thicknesses down to 14 nm (Fig. 10(f) and (g)). Moreover, the Raman spectra of the thin flakes displayed a noticeable blue-shift due to the strong electron-lattice interaction resulting from the reduced stacking order, which falls perfectly in line with previous theoretical predictions (Fig. 10(h)).¹¹⁷ Furthermore, we have been able to demonstrate a perfect correlation using four different analytical techniques, namely optical microscopy, AFM, scanning Raman microscopy (SRM) and scanning electron microscopy (Fig. 10(e), (f), (i) and (j)), respectively. This

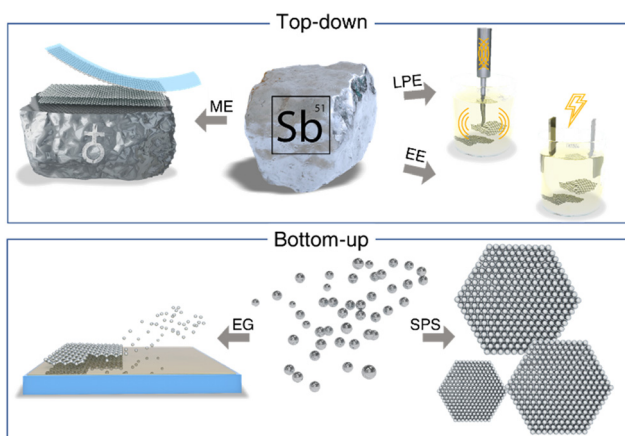


Fig. 9 Main synthetic approaches in the synthesis of antimonene classified in top-down and bottom-up approaches. Acronyms stand for mechanical exfoliation (ME), liquid phase exfoliation (LPE), electrochemical exfoliation (EE), epitaxial growth (EG) and solution-phase synthesis (SPS).



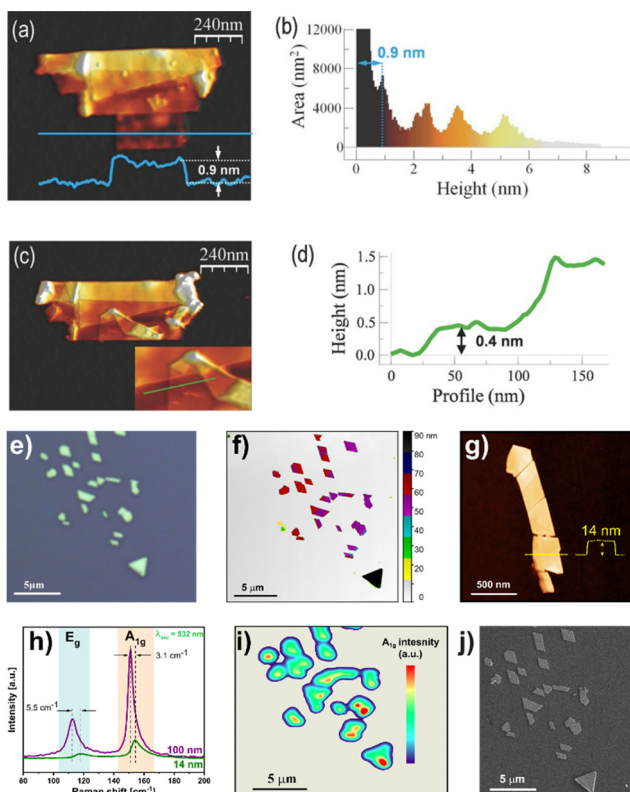


Fig. 10 AFM topography images of an antimonene flake with a monolayer terrace at the bottom. (a) AFM topography showing an antimonene flake with terraces of different heights. (b) Height histogram of the image in (a) pointing out the different thicknesses of the terraces. (c) Same flake as in (a) after a nanomanipulation process to confirm the stability of the material. The lower terrace of the flake was folded upward resulting in an origami structure with different folds. The inset corresponds to the area of the origami where the lowest step height is found. (d) Profile along the green line in the inset in (c). (e) Optical micrograph of a typical antimonene nanosheet isolated on SiO_2/Si substrate using micromechanical exfoliation. (f) False-colored AFM image of the same ANSs with incremental color code to highlight the terraces. Each color indicates a 10 nm step (g) AFM image of a few-layer antimonene nanosheet and height profile along the dashed yellow line. (h) Comparative Raman spectra of the pristine bulk (100 nm) and few-layer (14 nm) antimonene nanosheets, remarking the blue shift of the A_{1g} and E_g modes for the few-layer antimonene nanosheets. (i) Scanning Raman microscopy (SRM) map of the intensity of A_{1g} mode of the flakes in (e). (j) Scanning electron microscope (SEM) image of the same flakes in (e). Adapted from ref. 84 with permission from John Wiley & Sons, copyright 2016, and ref. 98 with permission from IOP Publishing, copyright 2020.

neat correlation allowed for the simultaneous and at the same time unambiguous identification of the morphology and the chemical structure of the probed sample. It is worth noting that such correlations are of critical importance in the characterization of 2D materials, and the reliance on individual measurements in distinct locations, even within the same sample, can be dangerously misleading, especially for new and not well understood systems such as the case of antimonene. It is commonplace to encounter washing solvent residues or impurities from the polymer used for delamination exhibiting flat morphologies, which might be erroneously identified as

antimonene flakes if solely measured by AFM without a direct correlation to other techniques, such as Raman for instance.

The main limitation of this micromechanical approach is that it yields a low quantity of exfoliated material, highlighting the need for alternative methods to develop antimonene on a large scale.

3.1.2 Liquid phase exfoliation (LPE). This top-down approach is widely known for its successful use towards the delamination of bulk materials to produce their 2D counterparts in large quantities.^{49,118–120} From van der Waals materials such as graphene to ionic solids like layered double hydroxides, LPE allows the preparation of colloidal suspensions of the desired material in a wide variety of solvents. The family of pnictogens was not foreign to this method since phosphorene has been successfully isolated yielding relatively concentrated samples.⁵⁹ As thoroughly discussed by Backes *et al.*,¹²⁰ LPE methods are based on the solubility parameters (such as the surface tension or the Hansen and Hildebrand parameters) theory. In this context, the stabilization of the nanomaterial is achieved when the solubility parameters of both solvent and solute match. Some of the most typical solvents that give stable dispersions are the following: *N*-methyl-2-pyrrolidone (NMP), *N*-cyclohexyl-2-pyrrolidone (CHP), dimethylformamide (DMF), dimethyl sulfoxide (DMSO) and isopropyl alcohol (IPA). The overall procedure consists of three stages: first, the interlayer attraction between the material's sheets is overcome by supplying energy (e.g. with ultrasonic waves). Then, the as-obtained nanosheets are stabilized with suitable solvents preventing the reaggregation of the material. Finally, a size-selection procedure is often required because of the polydispersity of the dispersions.¹²⁰ Regarding antimonene, Abellán, Zamora *et al.* reported in 2016 the first LPE of few-layer antimonene.¹²¹ In that work, they report a fast and simple method to obtain highly stable IPA/H₂O (4:1) suspensions of few-layer antimonene starting from bulk Sb crystals. Afterwards, the non-exfoliated material was removed by cascade centrifugation at 3000 rpm. To do this, the LPE process is carried out with tip sonication avoiding additional stabilizers such as surfactants. The high quality of the nanosheets was unveiled with AFM, scanning electron microscopy (SEM), Raman spectroscopy or EELS. AFM on SiO_2 substrates revealed step heights of 4 nm, related to the presence of few-layer material (Fig. 11). This value considers the overestimation because of residual solvent or additional contributions such as adhesion and capillary forces.^{59,122,123} The overall lateral dimensions were found to be in the 1–3 μm range and the atomic periodicity indicates a β -phase. Additionally, the obtained nanolayers were extremely stable under ambient conditions. Regarding the Raman characterization, the correlation of statistical Raman microscopy (SRM) with AFM in a polydisperse sample revealed antimonene flakes with thicknesses below 70 nm, corresponding to 17 layers, and barely showing any Raman signal. This result was similar to that observed for micromechanically exfoliated flakes.⁸⁴

After this initial work back in 2016, LPE is commonly being used as a methodology to obtain antimonene.^{112,124,125} Since



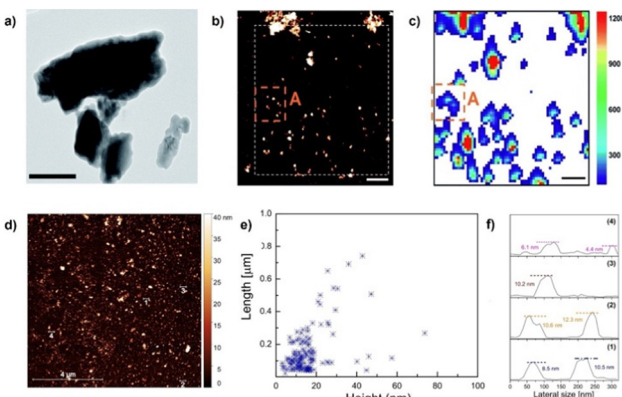


Fig. 11 (a) TEM image of FL antimonene nanolayers. (b) Representative topographic AFM image (scale bar 2 μm) of the exfoliated FL antimonene nanolayers. (c) Corresponding Raman $\text{A}_{1\text{g}}$ map (scale bar 2 μm) of the exfoliated FL antimonene nanolayers contained in the area dotted in white in (b). (d) AFM image of typical ANS showing polydisperse nanosheets exfoliated by ultrasonication. (e) Statistical dispersion of ANS lateral size as a function of thickness. (f) Thickness measured by AFM along the lines numbered in (d). Adapted from ref. 112 with permission from Royal Society of Chemistry, copyright 2019, and ref. 125 with permission from IOP Publishing, copyright 2020.

then, great effort has been made to maximize the concentration of the samples obtained by LPE, while maintaining an appropriate “dimensional anisotropy” (DA) ratio, which has been defined as the ratio between the height and length of the nanolayers, providing a good estimation of the morphology of the nanolayers (having the thin and large flakes a high DA ratio). We have reported a systematic study of the exfoliation process of antimony bulk crystals, with the aim to obtain stable FL antimonene suspensions with a high nanolayer concentration as well as a high DA ratio, by optimizing different parameters of the exfoliation procedure.¹¹² Specifically, the results show that the pre-processing of the material using wet-ball milling with 2-butanol strongly affects both the concentration and morphology of the suspension of FL antimonene. Moreover, the analysis of 28 different solvents previously selected based on Hansen parameters as well as other experimental observations yielded the highest concentration of FL antimonene for the mixture of NMP/H₂O in a (4 : 1) ratio, *ca.* 0.368 g L⁻¹ (*~*yield of 37 wt%), while the largest DA value was obtained using 2-butanol (*ca.* 27.6). The nanolayer heights were comprised between 2 and 8 nm and lateral dimensions lower than *ca.* 300 nm. Additionally, this work revealed that using tip sonication employing an ultrasonic wave amplitude of 100% yields better results in terms of DA than bath sonication. Interestingly, the use of 2-butanol also yielded a large number of edges and electroactive sites that strongly influence the electrochemical behavior of the exfoliated antimonene, forming also irreversible superficial antimony oxides.¹¹²

A particular case within the LPE is that of the antimonene quantum dots (AMQD). This type of material – of great interest in biomedical applications (*vide infra*) – which is characterized by lateral dimensions *<*8 nm, is obtained after subjecting the

bulk antimony samples to an iterative sonication and centrifugation process for long periods of time using high power and high amplitude values.^{126,127} More in detail, this procedure combines ultrasound tip sonication with ice bath sonication of bulk Sb in NMP or ethanol, followed by a low-speed centrifugation to remove the larger particles and a high-speed centrifugation to isolate the desired QD particles. Fig. 12 highlights the synthesis and characterization of AMQDs in EtOH according to ref. 127. The resulting solution displays a high yield in the synthesis of uniform and small particles with average size and thickness of *ca.* 3 and 2 nm, respectively, as detected by AFM, in good agreement with TEM images. The composition of the AMQDs and the presence of Sb was determined *via* XPS and Raman spectroscopy. Moreover, in order to improve their biocompatibility, a non-covalent functionalization with polyethylene glycol (PEG) moieties has been carried out.

The as-obtained PEG-coated AMQDs exhibited excellent stability in physiological experiments; no signs of toxicity in cells in the studied range of concentrations and excellent photothermal conversion efficiency upon NIR irradiation. Beyond the biomedical applications,¹²⁷ recent studies on AMQDs points to them as promising candidates in optoelectronic devices, and more specifically in quantum-dot-sensitized solar cells.^{128,129} In this sense, the work of Zhang *et al.*¹²⁹ presents liquid phase-synthesized AMQDs as an effective photoactive material in quantum-dot-sensitized solar cells. The AMQDs were prepared following the aforementioned protocol starting from bulk Sb crystals using a simple probe ultrasonic procedure followed by an ultrasound bath treatment. The temperature during the probe sonication was set under 25 °C to avoid possible oxidation or degradation processes. Afterwards, the dispersion was centrifuged to remove the residual Sb bulk particles and the resulting suspension was centrifuged at high speed (12 000 rpm) to collect the supernatant AMQDs, which were re-suspended in NMP. AFM

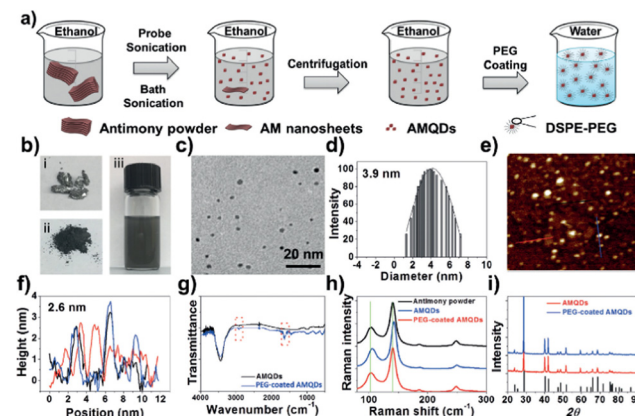


Fig. 12 (a) Fabrication of PEG-coated AMQDs. (b) Images of bulk antimony, antimony powder, and AMQD solution during the preparation process. (c) TEM image, (d) diameter distribution, (e) AFM image, and (f) thickness of the PEG-coated AMQDs. (g) FTIR spectrum, (h) Raman spectrum, and (i) XRD spectrum of AMQDs and PEG-coated AMQDs. Reproduced from ref. 127 with permission from John Wiley & Sons, copyright 2017.



revealed an average thickness of *ca.* 2.2 nm and the related Sb Raman bands A_{1g} and E_g modes confirmed the nature of the QDs. The combination of the AMQDs with strong light-matter interaction, moderate energy bandgap in the visible range and antioxidation properties allows a photoelectric conversion efficiency up to 3.07%, alongside the long-term stability of these quantum-dot-sensitized solar cells (more than 90% of the initial conversion efficiency after 1000 h).

3.1.3 Electrochemical exfoliation. Electrochemical exfoliation is considered as a low-cost, environmentally friendly and straightforward synthetic approach.¹³⁰ This method has been recently used for the synthesis of some 2D materials such as phosphorene or graphene, offering a new pathway for the large-scale synthesis of 2D materials.^{131,132} The first work showing the production of few-layer antimonene *via* electrochemical exfoliation was reported in 2017 (Fig. 13). This top-down approach uses a two-electrode electrochemical cell in combination with bulk Sb as the working electrode and a Pt wire as the counter electrode, both immersed in the electrolyte solution of Na_2SO_4 to avoid contamination from the conducting wire as well as the possibility of electrolysis. The exfoliation procedure was conducted with at a constant voltage of -6 V for 60 min. The resulting electrolyte solution containing the exfoliated antimonene flakes was centrifuged and the sediment was collected and washed, followed by the dispersion and storage of the antimonene suspension in NMP.¹²⁶ The work of Lu *et al.*

gives rise to both α -phase few-layer antimonene in combination with antimonene quantum dots presenting high quality and stability, characterizing the non-linear optical Kerr response at the visible band by means of spatial self-phase modulation (SSPM) measurement technique. In this sense, the few-layered system displays a larger nonlinear refractive index than that found for the quantum dot counterpart. This study also considers antimonene as a promising optical Kerr material at the short wavelength range, hence strengthens its use in optoelectronic applications.¹²⁶ Moreover, Li and co-workers proposed a mechanism to explain the cathodic exfoliation of 2D Sb nanosheets in 0.5 M Na_2SO_4 solution based on the Na^+ intercalation and interlayer expansion. Another approach consists of using square-wave voltage using bulk antimony as the anode and changing the voltage direction to perform exfoliation. This strategy has been explained on the basis of four different stages involving oxygen-containing free radicals, Na^+ intercalation into the gaps and holes formed by the radicals (and inducing an initial expansion of the layers), SO_4^{2-} intercalation after changing the voltage, and finally the exfoliation and dispersion induced by the gases formed during this drastic process (H_2 , O_2 and SO_2).¹⁰⁷

3.1.4 Pressurized alloy assisted exfoliation. One of the newest top-down approaches reported so far deals with alloy formation under high pressures generated during a conventional hydrothermal process. Indeed, the use of *n*-butyllithium creates a Li_3Sb alloy intermediate at the edge region of bulk antimony. Afterwards, a protonation process converts Li_3Sb to gaseous stibine (SbH_3) in a liquid solution, leading at $80\text{ }^\circ\text{C}$ to an appropriate buoyancy and subsequent endothermic opening of the galleries that facilitate LPE. The resulting material consists of antimonene nanosheets with lateral dimensions of $\approx 3\text{ }\mu\text{m}$ and a thickness of $<2\text{ nm}$.¹³³

The ability to intercalate alkali metals shown by Sb allows the formation of antimony intercalation compounds to be envisioned, in analogy to previously reported black phosphorus intercalation compounds (BPICs), which under appropriate solvents and under thorough exclusion of oxygen and water may lead to pnictogenide dispersions.^{61,134–136}

3.2 Bottom-up approaches

3.2.1 Epitaxial growth. Epitaxial growth is a set of techniques involving the deposition of crystalline layers on crystalline substrates that belong to the bottom-up approach and are conventionally used in order to obtain high-quality 2D materials.^{137,138} In addition, epitaxial synthesis allows for obtaining significant amounts of crystalline 2D materials, clearly surpassing micromechanical exfoliation. For antimonene, two different methods can be used: molecular beam epitaxy (MBE) and van der Waals epitaxy (vdWE).

The MBE approach was previously carried out by Lei *et al.* with the growth of single layer Bi (111) on Bi_2Te_3 (111) and Bi_2Te_3 (111) substrates with a small lattice mismatch.¹³⁹ Afterwards, Wu *et al.* were able to grow antimonene monolayers (thickness of *ca.* 0.28 nm as measured by the profile line across the edge of the antimonene film) on PdTe_2 substrates

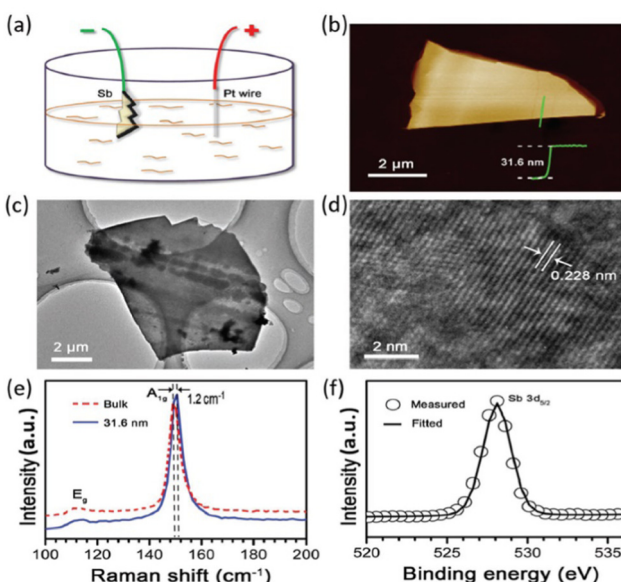


Fig. 13 Characterization of the prepared multilayer antimonene by electrochemical exfoliation. (a) Schematic illustration of the two-electrode system used for the procedure, depicting bulk Sb, Pt wire, and Na_2SO_4 aqueous solution as the working electrode, counter electrode, and electrolyte, respectively. (b) AFM image of a multilayer antimonene nanoflake obtained by electrochemical exfoliation. (c), (d) TEM and HRTEM images of the electrochemical exfoliated antimonene. (e) Raman spectra of the bulk antimony and multilayer antimonene shown in (b). (f) XPS spectrum showing the Sb 3d_{5/2} peak of the exfoliated multilayer antimonene. Reproduced from ref. 126 with permission from John Wiley & Sons, copyright 2017.



(Fig. 14).¹⁴⁰ In that work, both *in situ* low-energy electron diffraction (LEED) and scanning tunneling microscopy (STM) measurements confirmed the high quality of the epitaxially-grown antimonene with a honeycomb graphene-like structure. XPS results reported that the interaction between antimonene and the substrate is *via* weak van der Waals forces. The combined results of STM and XPS highlighted a good stability of the material against air exposure for 20 min.

Most recently, additional studies have used MBE to grow antimonene on Ge (111) and Ag (111) (Fig. 14) substrates, confirming the weak interactions between the monolayer and the substrate as well as the honeycomb structure.^{81,141}

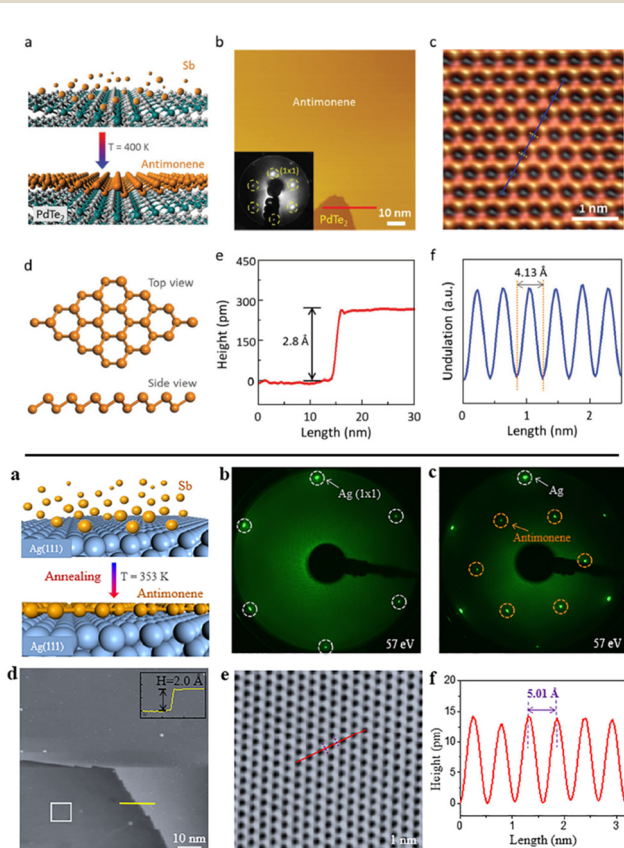


Fig. 14 Top: (a) Schematic of fabrication of an antimonene monolayer formed on the PdTe_2 substrate obtained by MBE approach. (b) STM topographic image of large antimonene island on PdTe_2 . (Inset: LEED pattern of antimonene on PdTe_2 .) (c) Atomic resolution STM image of monolayer antimonene showing the graphene-like honeycomb structure. (d) Top view and side view of the buckled conformation of the antimonene honeycomb. (e) Height profile along the red line in (b), showing that the apparent height of the antimonene island is 2.8 \AA . (f) Line profile corresponding to the blue line in (c), revealing the periodicity of the antimonene lattice ($4.13 \pm 0.02 \text{ \AA}$). Bottom: (a) Schematic of the MBE fabrication process. (b) LEED pattern of a clean $\text{Ag}(111)$ substrate. (c) LEED pattern of antimonene on $\text{Ag}(111)$. (d) Large scale STM image of monolayer antimonene on the $\text{Ag}(111)$ (inset: height profile along the yellow line at the terrace edge). (e) High-resolution STM image of antimonene depicted by the white square in (d). (f) Line profile corresponding to the red line in (e), revealing the periodicity of the antimonene lattice (5.01 \AA). Adapted from ref. 140 with permission from John Wiley & Sons, copyright 2016, and ref. 141 with permission from American Chemical Society, copyright 2018.

Moreover, these studies revealed a semi metallic behavior for a single layer of deposited antimonene. MBE has also been reported to obtain multilayer antimonene nanoribbons on sapphire substrates that can extend themselves to a length of few microns.³⁶ These ribbons were grown *via* *in situ* thermal annealing after the initial MBE deposition. Their formation mechanism, supported on SEM and AFM measurements, considers an adatom migration process at an elevated temperature in combination with strain relaxation and surface energy balance.

vdWE is based on the use of substrates with the absence of dangling bonds on their surface. Thus, the epitaxially grown layer of the layered material is connected to the substrate by means of weak van der Waals interactions in the absence of strong chemical bonds. Therefore, it is possible for the material to exhibit a different crystalline symmetry than that of the substrate. The first few-layer antimonene grown by vdWE was reported in 2016 by Ji *et al.*¹¹⁷ Fig. 15 highlights the antimonene layer synthesized on mica substrates by means of a van der Waals epitaxy.

In that work the high-quality synthesis of few-layer antimonene with a buckled hexagonal structure (β -phase) was reported as confirmed by HRTEM and Raman spectroscopy. This finding is consistent with the predicted as most stable allotrope of monolayered antimonene. In broad strokes, the synthesis is carried out in a two-zone tube furnace with separated temperature controls. The bulk Sb is placed in one of the regions and then heated to obtain Sb vapor. A transport flow of $\text{Ar} + \text{H}_2$ transfers the Sb vapor to the other region of the furnace, in which there is a substrate at a lower temperature than the first region, hence promoting the condensation of Sb. The high stability of the material and an average dimension ($\text{ca. } 4 \text{ nm}$, *i.e.*, few-layers) and thicknesses (*ca.* 4 nm, *i.e.*, few-layers) were analysed by optical microscopy, AFM, Raman spectroscopy and XPS. The

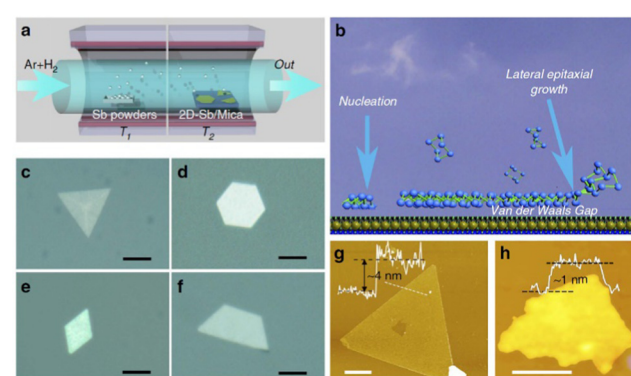


Fig. 15 Antimonene synthesized on mica substrates *via* vdW epitaxy. (a) Schematic illustration of the sample synthesis configurations. (b) Schematic diagram of the vdWE. (c)–(f) Optical images of typical antimonene polygons with triangular, hexagonal, rhombic, and trapezoidal shapes, respectively. The scale bar is 5 mm. (g) AFM image of a typical triangular antimonene sheet. The scale bar is 1 mm. (h) AFM image of a tiny antimonene sheet with a thickness of *ca.* 1 nm. The scale bar is 50 nm. Reproduced from ref. 117 with permission from Springer Nature, copyright 2016.



stability of the nanosheets was tested with an as-prepared and a 30 day aging sample, with no additional signature in the corresponding Raman spectra or X-ray energy-dispersive spectroscopy (XEDS). The good electrical conductivity of the synthesized antimonene (10^4 S m $^{-1}$) in combination with the wavelength independent high transparency in the visible light spectrum endows this material with potential applications as a flexible transparent electrode.¹¹⁷

Along this front, Hogan *et al.* reported the temperature-induced phase transition between α and β phases of antimonene in a van der Waals heterostructure on Bi₂Se₃.¹⁴² The Bi₂Se₃ surface was prepared *via* an ultra-high vacuum chamber, and the antimony was then sublimated and deposited at room temperature. Once antimony is deposited, it forms α -antimonene domains with different orientations with respect to the substrate. However, after mild annealing at 473 K, there is a growing of the β phase that overcomes α -antimonene, leading finally to a single domain of β -antimonene which exactly matches with the surface lattice structure of Bi₂Se₃. These results were also supported by DFT calculations, which highlight the lattice matching of the β in stark contrast with the α one.

Last but not least, recent studies using vdWE have demonstrated the experimental synthesis of antimonene-based vdW heterostructures, proving the stability of both the hexagonal β - and rectangular α -forms of antimonene, on top of the topological insulator α -bismuthene.¹⁴³ Moreover, these studies have allowed the elucidation of the rich allotrophic/polymorphic structural diversity of antimonene, revealing extremely interesting rotational vdW epitaxy in 2D Sb/graphene heterostructures. Furthermore, thanks to these novel heterostructures several aspects related to the dynamic oxidation and phase transition of antimonene have been corroborated. These interesting points will be further analyzed below.

3.2.2 Solution-phase synthesis. One of the most promising bottom-up approaches has been reported by Peng *et al.* using a solution-phase synthesis of few-layer antimonene *via* anisotropic growth.¹⁴⁴ This work addresses the fact that the actual synthetic methods for obtaining high-quality few-layer antimonene nanosheets (exfoliation and epitaxial growth) have hindered the large-scale production of this material. The authors propose the direct synthesis of few-layer and promoting their anisotropic growth in a colloidal solution, starting from SbCl₃, together with oleylamine and dodecanothiol. While oleylamine is employed as the reducing agent to obtain Sb(0), dodecanothiol is used to prevent the formation of oxides. In this synthesis the role of the chloride anions is crucial to induce the hexagonal final form. Fig. 16 depicts how the final morphology of the resulting antimony species can be tuned if some experimental parameters are modified during the synthesis. The same authors recently reported further insights into the formation of hexagonal antimonene as well as remarked on the role of the chloride salt (SbCl₃) as a key factor in the resulting morphology. Furthermore, they replace the use of dodecanothiol by alkyl phosphonic acids to avoid the presence of sulphur-side products.¹⁴⁵

Some of the advantages of the solution-phase methodologies rely on their simple procedures, together with the

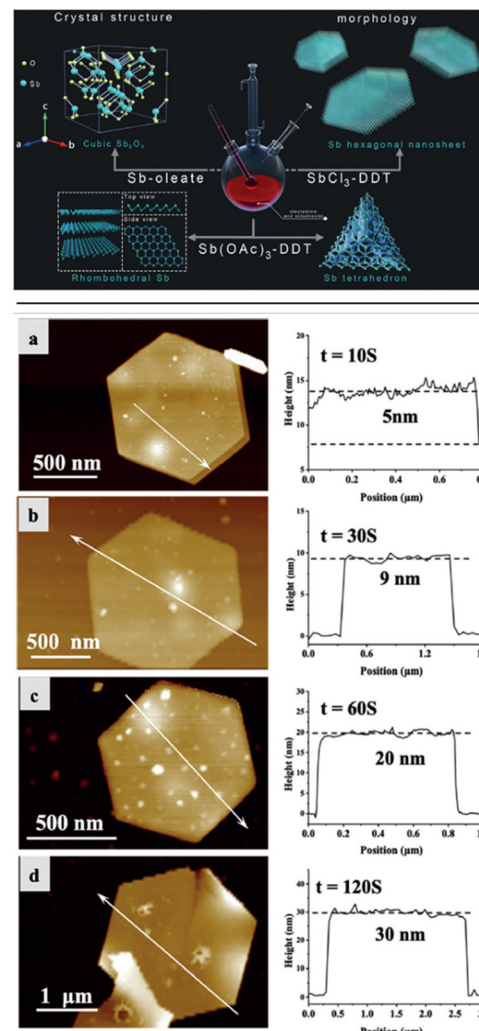


Fig. 16 Top: Illustration of the wet chemical solution-phase synthesis of antimony trioxide, antimony tetrahedral, and hexagonal antimonene nanosheets from Sb-oleate, Sb(OAc)₃-DDT, and SbCl₃-DDT as precursors, respectively. Bottom: AFM images and the corresponding height profiles of hexagonal antimonene nanosheets with tunable layer thicknesses obtained at different annealing times at 300 °C: (a) 10 s, (b) 30 s, (c) 60 s, and (d) 120 s. Reproduced from ref. 144 with permission from John Wiley & Sons, copyright 2019.

tunability of the material from the point of view of size, morphology and composition, as well as the monodisperse character of the obtained samples,^{146,147} opening the door to the large-scale synthesis of the desired material. Indeed, the authors modulated to some extent the average thickness of the nanosheets by tuning the annealing time. They characterized the nanosheets *via* Raman spectroscopy, TEM and XRD proving that the material consisted of β -antimonene. Furthermore, X-ray photoelectron spectroscopy (XPS) results confirmed the presence of chloride anions on the surface of the antimonene, as well as the coordination of dodecanothiol, thus suggesting surface functionalization.

Furthermore, the colloidal synthesis of antimonene has been very recently studied by Abellán, Zamora and co-workers, optimizing the different synthetic parameters to



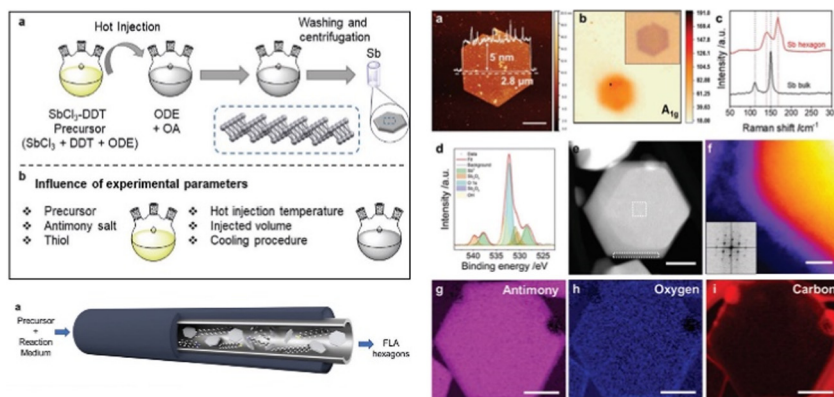


Fig. 17 Main approaches in solution-phase synthesis of antimonene hexagons. Above: Schematic representation of hot injection method and parameters influencing the outcome of the synthesis. Below: Illustration of the continuous-flow reactor. On the right, different characterization performed in the obtained antimonene hexagons from the optimized solution-phase synthesis of antimonene: (a) AFM, (b) Raman mapping and (c) Raman spectra, (d) XPS, (e) HAADF STEM and (f) Colorized HAADF STEM, inset showing the fast Fourier transformation. (g)–(i) Elemental compositional EELS maps. Scale bar (e)–(i) 100 nm. Adapted from ref. 148 with permission from John Wiley & Sons, copyright 2021.

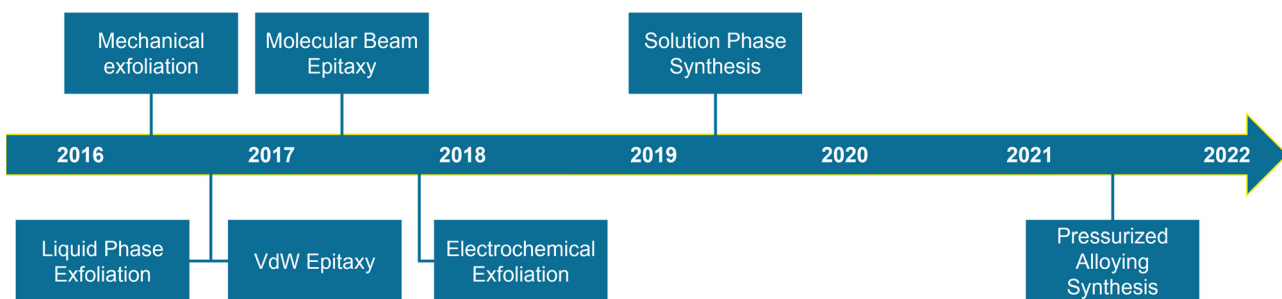


Fig. 18 Timeline highlighting the development of the different synthetic approaches to obtain antimonene.

obtain high-quality, ultrathin antimonene hexagons (Fig. 17).¹⁴⁸ By studying the effects of each synthetic parameter on the resulting antimonene hexagons, like the precursor solution and the amount of volume injected, the antimony salt used, the hot-injection temperature or the reaction time, the authors have been able to gain control over the particle morphology. This further encouraged them to develop a scale-up process, which is essential for the real application of antimonene. For this, they have developed a continuous-flow synthesis reactor that allows the large-scale production of excellent morphological and structural quality antimonene hexagons, producing up to 0.964 mg h^{-1} .

To sum up, Fig. 18 and Table 3 expose an historical timeline and general overview of the main synthetic approaches developed for antimonene, highlighting the main aspects associated with each of them.

4. Novel insights on the characterization and chemical reactivity of antimonene

Regarding the most typical characterization techniques to confirm the identity of antimonene, there are the following:

microscopy techniques, X-ray diffraction (XRD), Raman spectroscopy and XPS. Firstly, microscopy measurements (HRTEM, SEM, AFM or optical microscope, among others) are fast techniques to check the layered morphology of the material. As previously commented, optical contrast is reported as a feasible way to identify the quality of the layered system.¹⁴⁹ AFM is positioned as the best tool to study the thickness (*i.e.* the number of layers) of the deposited antimonene (Fig. 10). In this sense, it is mandatory to consider that the water molecules trapped between the antimonene and the substrate overestimate the measured thickness.⁸⁶ Taking this into account, a monolayer of antimonene exhibits an average height of *ca.* 0.9 nm, while a few-layered system is obtained in high yield with thicknesses in the 4 nm range and several μm in lateral dimensions.^{84,121} In order to unveil if a thin flake is a monolayer, one can fold the terrace *via* nanomanipulation with the AFM into an origami structure. As previously reported by Geim and Novoselov concerning graphene sheets, a single flake can be identified by measuring the step height of single folds.⁸⁵ As reported by Ares *et al.*,⁸⁴ the lowest step height was found to be 0.4 nm hence corresponding to a monolayer of antimonene. This origami folding was also performed days after the flake deposition on the substrate, thus confirming the mechanical stability of the material. Furthermore, high-resolution AFM



Table 3 Main synthetic approaches in the synthesis of antimonene and timeline with the number of documents

	Synthetic approach	Main aspects	Ref.
Top-down	Micromechanical exfoliation	<ul style="list-style-type: none"> • First isolation of few and single-layer antimonene • Millimeter-sized crystals • Easy identification (optical microscopy) • Low quantity of exfoliated material, time consuming, irregular morphology and lack of homogeneity in exfoliation. 	First time reported: ⁸⁴ Other works: ¹⁴⁹
	Liquid phase exfoliation	<ul style="list-style-type: none"> • Suspensions of the material in adequate solvents (NMP, DMF, and IPA...) • Based on solubility parameters • Size selection (cascade centrifugation) • Micrometer-sized crystals of few-layer antimonene (AFM) • Irregular morphology and polydispersity. 	First time reported: ¹²¹ Other works: ^{112,124,125,150}
	Electrochemical exfoliation	<ul style="list-style-type: none"> • Two-electrode cell • Electrolyte solution containing few-layer antimonene dispersion. • Size selection (cascade centrifugation) • Micrometer-sized crystals of few-layer antimonene (AFM) • Irregular morphology and polydispersity. 	First time reported: ¹²⁶ Other works: ^{107,151}
	Pressurized alloy assisted synthesis	<ul style="list-style-type: none"> • Uses pressure created from the protonation of the alloy, creating a buoyancy that exfoliates antimonene • Irregular morphology, wide size distribution, but very thin flakes (1–2 nm) • Molecular beam epitaxy (MBE) and van der Waals epitaxy (vdWE) 	First time reported: ¹³³
Bottom-up	Epitaxial growth	<ul style="list-style-type: none"> • Used to grow antimonene on substrates (mica, Ge(111), Ag(111)...) • Micrometer-sized crystals of single-layer antimonene (AFM) • Irregular morphology, low quantity if the material and polydispersity. 	First time reported: MBE, ¹⁴⁰ vdWE ¹¹⁷ Other works: ^{81,108,141,142,152–155}
	Solution-phase synthesis	<ul style="list-style-type: none"> • Based on the anisotropic growth of antimonene • Large-scale production of the material • Micrometer-sized crystals of few-layer well-defined hexagonal antimonene (AFM) • Polydispersity 	First time reported: ¹⁴⁴ Other works: ^{145,148}

allows the ability to study the atomic periodicity, and hence it is useful to relate it with the phase of antimonene. Due to the rippling effects caused by the conformation of antimonene to the underlying SiO_2 , it is very difficult to obtain well-resolved atomic force microscopy (AFM) images that exhibit the atomic periodicity. This effect is also visible for graphene on SiO_2 , therefore not ascribed to the antimonene itself.¹⁵⁶ Regarding AFM data, it is of utmost importance to correlate them with those of other techniques such as Raman spectroscopy to successfully identify the layered material (Fig. 10).¹⁵⁷ In the absence of this supporting evidence, one can easily mistake solvent impurities for “fake flakes” of the desired material.

Electron microscopy provides the morphology of the material and is revealed as a key technique in the search for a desired morphology, such as well-defined hexagonal samples in solution phase synthesis or the AMQDs.^{127,144} Even though TEM, HRTEM or SEM cannot be used to analyse the thickness of the sample, they can be used for measuring the average lateral dimension as well as taking wider areas of the deposited material throughout the whole substrate. In this context, aberration-corrected scanning transmission electron microscopy (STEM) combined with EELS can be used to investigate the local structure and chemistry of the flakes (Fig. 19–21). For example, for LPE samples, high-angle annular dark-field images taken at low voltages to prevent damaging the flakes highlight the atomic-resolution of the crystal structure, confirming a β -phase as well as the good crystallinity and the absence of defects in the studied antimonene flakes. The EEL

spectra depict the compositional maps according to absorption edges such as Sb M_{4,5}, O K or C K, pointing out some damage in the flake ends and the presence of elements such as C and O because of surface contamination and oxidation.¹²¹

Moutanabbir *et al.* have recently used scanning tunneling microscopy (STM) and STEM, together with state-of-the-art techniques, like *in situ* low-energy electron microscopy (LEEM) or X-ray photoemission electron microscopy (XPEEM) to observe the allotropic state of antimonene grown by MBE on top of graphene Ge heterostructures. These techniques are optimized for surface observations, as the low energy of the electrons gives high sensitivity to surface signals due to their strong interaction, with resolutions greater than 10 nm in some cases. The measurements revealed a stabilized α -phase (A17) of the antimonene in flakes thinner than 4 nm (Fig. 19(e)), which undergo a thickness-driven transition to the stable β -phase (A7) as the number of layers increase (Fig. 19(c)).⁸² This allotropic dynamism as function of thickness is intimately related to the electronic structure of antimonene.

Further studies about antimonene/graphene heterostructures have been carried out by Bayer *et al.*, where they studied the allotropic phases of antimonene grown by vdWE onto CVD graphene. They found the coexistence of two main morphologies: 2D rhombohedral β -antimonene, and a one-dimensional nanowire structure with rectangular basis.⁸³ Remarkably, both morphologies exhibit direct in-plane rotational vdW epitaxy with the graphene support. These results open the door for the scalable production of antimonene heterostructures by bottom-up approaches.



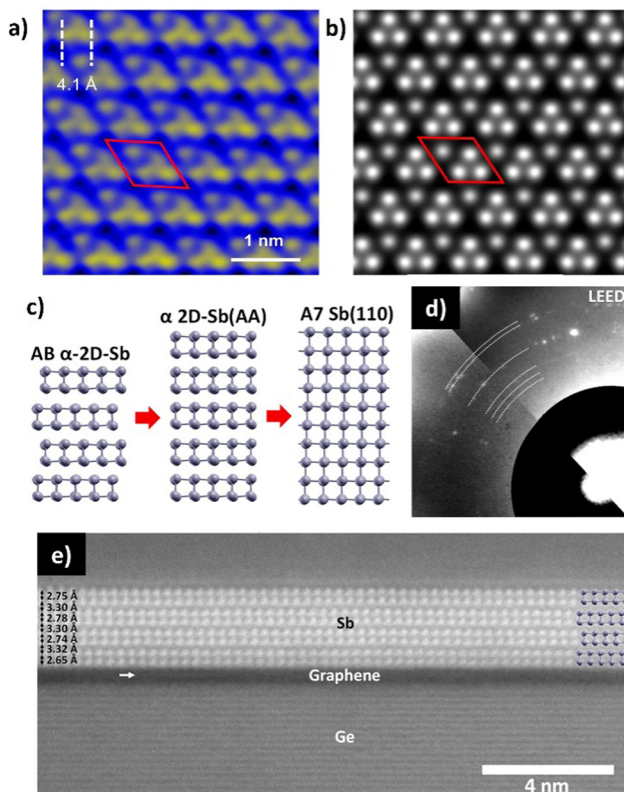


Fig. 19 (a) Atomically resolved STM image of an antimonene monolayer on SbAg₂ surface alloy on Ag(111). (b) The simulated STM in the constant height mode (~ 2 Å). (c) Schematic representation of the layer-dependent A17 (AB α -2D-Sb) transition to A7 in antimonene. (d) LEED pattern of 2D-Sb on graphene. Part of the Ewald sphere at 44 eV is shown in the bottom left and at 29 eV is shown in the top right. (e) STEM of a cross section of 4 bilayers A17 antimonene island on graphene. Adapted from ref. 82 and 152 with permission from American Chemical Society, copyright 2020.

As in other 2D materials prepared using top-down approaches, XRD is one of the most commonly used techniques because it can confirm the diffraction patterns of the resulting product allowing identification of the antimonene crystalline phase.^{112,144} In this context, bulk Sb and antimonene exhibit very similar XRD patterns. The main difference relies on the weaker intensity of the (003) and (006) peaks located at *ca.* 23.7° and 48.5°, respectively, of the antimonene compared with that of bulk Sb.¹⁵⁸ This effect is indicative of the reduction of the antimony crystals along the *c*-axis mainly.¹²⁴ The rhombohedral phase of Sb is indexed as PDF#035-0732.¹⁵⁹ Fig. 20(a) depicts XRD patterns of bulk Sb, Sb micro-crystals after ball-milling and antimonene.

Raman spectroscopy is a powerful technique widely used to characterize 2D materials.^{59,160} The Raman spectrum of bulk Sb when excited at 532 nm depicts two main phonon peaks: A_{1g} and E_g modes at 149.8 and 110 cm⁻¹, respectively, and related to the in-plane and out-of-plane vibrational modes.¹⁶¹ Few-layer antimonene displays a shift in both bands to a higher wavenumber region (also called blueshift). In this sense, it is reported that the correlation of AFM with SRM¹⁶² indicated that antimonene samples with a thickness below

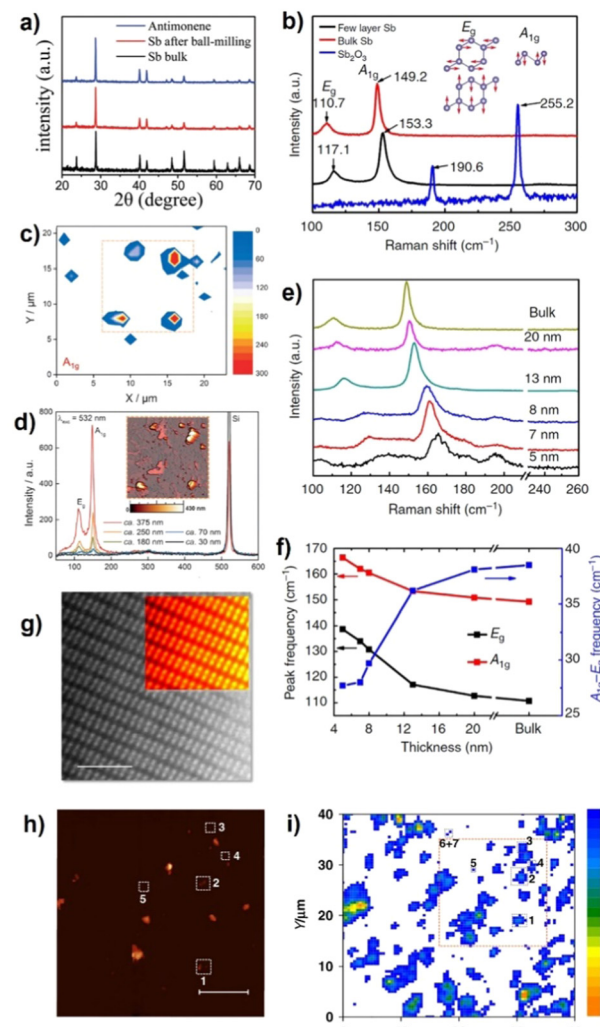


Fig. 20 (a) X-ray diffraction (XRD) patterns of bulk antimony crystals, antimony plates after ball-milling and antimonene. (b) Raman spectra of bulk antimony (β-phase), few-layer antimonene and antimony trioxide. Inset: Vibrational modes of β-phase antimonene. (c) A1g intensity Raman mapping of solvent-exfoliated flakes deposited on a SiO₂/Si substrate. (d) Single-point Raman spectra measured at different thicknesses according to the topographic AFM image (inset) of the same area studied in (c) (dashed lines). (e) Raman spectra of antimonene polygons with different thicknesses, from 5 nm to bulk. (f) A_{1g}, E₉ peak frequencies and energy difference of those two peaks plotted against sample thickness. (g) Atomic resolution HAADF image acquired on the edge of a free-standing portion of an antimonene flake obtained by LPE. The scale bar is 2 nm. (h) Representative AFM topography image (scale bar 5 μ m) of exfoliated antimonene onto SiO₂/Si substrates. (i) The corresponding Raman A_{1g} mapping of the same antimonene flakes in (h). Adapted from ref. 65 and 117 with permission from Springer Nature, copyright 2019 and 2016, ref. 121 with permission from John Wiley & Sons, copyright 2016, and ref. 158 with permission from the Royal Society of Chemistry, copyright 2020.

70 nm (*ca.* 17 layers) barely show any Raman signal (Fig. 20(b) and (c)).¹²¹ Nevertheless, taking single-point spectra at different points of the sample does evidence how the peak intensities decrease concomitantly with the thickness of antimonene (Fig. 20(d)).¹²¹ It is worth noting that these Raman results follow the same trends independently of the synthetic

approach to obtain antimonene. The work of Ji *et al.*¹¹⁷ clearly illustrates how the Raman peaks of bulk Sb evolve after obtaining its layered counterpart and also compares the fingerprint for a Sb_2O_3 species, which exhibits a completely different pattern (Fig. 20(b)). Indeed, Fig. 20(e) shows the Raman spectra of antimonene polygons synthesized by vdW epitaxy with thicknesses varying from 5 nm to bulk. Moreover, Fig. 19(f) shows the A_{1g} , E_g peak frequencies (left vertical axis) and the energy difference (right vertical axis) of those two peaks plotted against sample thickness, confirming the blueshift observed in LPE samples. These sorts of blueshifts have been also reported for other 2D materials and are probably attributed to the fact that lattice constant shrinks when the number of layers decreases.^{163,164} Last but not least, by performing the LPE under inert conditions using bmim- BF_4^- ionic liquid and argon-filled gloveboxes it is possible to obtain highly crystalline Sb nanosheets (Fig. 20(g)) that exhibit observable Raman signals with perfect AFM spatial correlation under environmental conditions down to the 5 nm limit, well below the previous LPE results using IPA/ H_2O mixtures (Fig. 20(h) and (i)).^{65,117} XPS analysis is an effective technique to study the oxidation state of the Sb atom as well as the surface oxidation state. In this sense, and due to the importance of this point, a thorough discussion is carried out in the following section.

4.1 Oxidation behaviour

As previously commented, it is reported that antimonene exhibits good structural integrity as well as high thermodynamic stability, even though theoretical studies predicted a high tendency for oxidation surpassing even that of phosphorene.⁶⁵ This is in good agreement with the recent experimental results in which O is always detected on the surface of the antimonene sheets.^{111,117,121} It is believed that the oxidation of antimonene leads to a surface oxidation layer that acts as a passivation coating, thus protecting the material from further structural decomposition.⁶⁰ This oxidation layer might potentially alter the final properties of the material, and eventually affect the associated applications.¹⁶⁵ Therefore, understanding the oxidation behaviour of antimonene from the fundamental viewpoint as well as from the experimental perspective is imperative for the future development of antimonene-based technologies. Recently, Assebban *et al.* published a seminal work comprehensively investigating the surface oxidation of liquid-phase exfoliated antimonene nanosheets (ANS).¹²⁵ First, they observed that the Raman spectra of the exfoliated ANSs featured, besides the typical vibrational modes at A_{1g} (149.8 cm^{-1}) and E_g (110 cm^{-1}), additional peaks at 190.5 and 254.6 cm^{-1} that were ascribed to Sb_2O_3 (Fig. 21).

Further STEM analysis combined with EELS elemental mappings has shown the formation of an amorphous layer rich in Sb and O, exclusively at the top surface and edges, which corroborated the observed partial oxidation caused by the sonochemical processing in IPA/ H_2O during exfoliation. The authors decided afterwards to submit exfoliated samples to thermal annealing at $210\text{ }^\circ\text{C}$ under ultra-high vacuum (UHV),

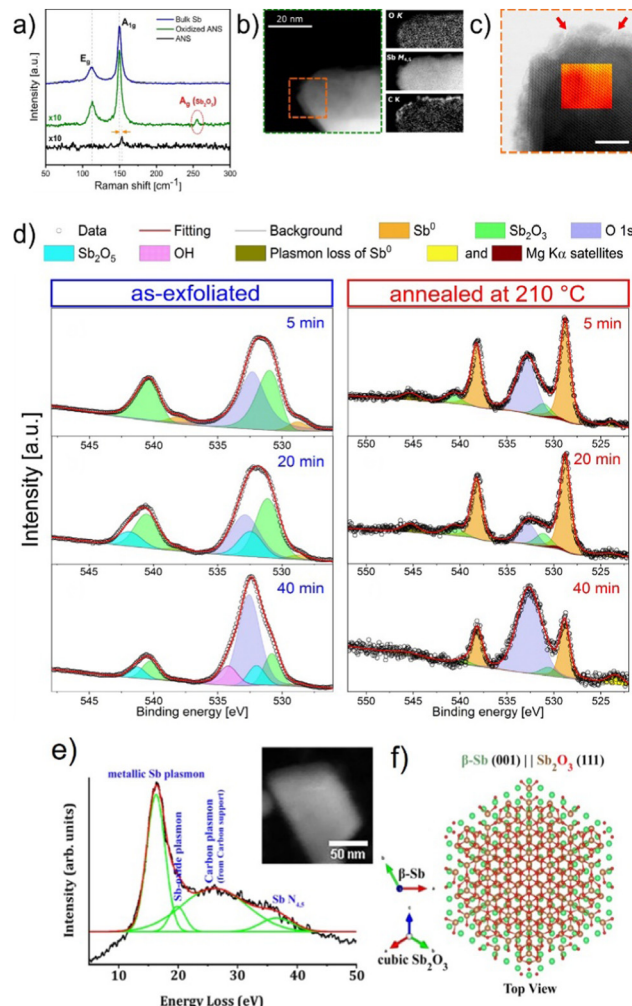


Fig. 21 (a) Raman spectra of bulk antimony, exfoliated ANS and oxidized-ANS exhibiting Sb_2O_3 Raman fingerprint at 254.6 cm^{-1} . (b) HAADF image of sub-nanometric ANS acquired at 80 kV and the corresponding elemental compositional maps derived from EELS. (c) High-magnification annular bright field (ABF) image near the edge of the nanosheet displayed in (b) (orange-dashed area) (scale bar 5 nm). (d) XPS line spectra in the Sb 3d region for antimonene nanosheet samples, as-exfoliated (left column) and after thermal annealing at $210\text{ }^\circ\text{C}$ under high vacuum (right column). (e) VEELS spectrum of the $\beta\text{-Sb}(001)$ crystal on suspended graphene (ADF STEM in the inset). Spectrum acquired after ~ 8 months ambient air exposure of the sample. (f) Atomic model of the suggested $\text{Sb}_2\text{O}_3(111)$ formed from ambient air exposure on $\beta\text{-Sb}(001)$ crystals. Adapted from ref. 83 with permission from Springer Nature, copyright 2021, and ref. 125 with permission from IOP Publishing, copyright 2020.

while tracking the surface chemistry using X-ray photoelectron spectroscopy (Fig. 21). They discovered that the thermal annealing resulted, to some extent, in the removal of previously observed surface oxide adlayer, as indicated by the *in situ* XPS evidence. More interestingly, by means of ultraviolet photoelectron spectroscopy (UPS), they demonstrated a change in the surface electronic properties, which upon annealing, switches from a metallic to an otherwise semiconducting material with an estimated band gap of approximately 1 eV , assuming the Fermi level is situated in the middle of the gap.



Bayer *et al.* also studied the oxidation behavior in PVD grown antimonene/graphene heterostructures by Valence EELS (VEELS), allowing any trace of surface oxidation to be precisely detected. VEELS revealed low levels of oxidation after 8 months of ambient exposure (Fig. 21(e)), and together with the ADF STEM FT patterns, revealed the presence of a sixfold structure that can be associated with cubic $\text{Sb}_2\text{O}_3(111)$. From these results, they suggest the formation of a thin crystalline layer of Sb_2O_3 on top of β -antimonene (Fig. 21(f)). These results again support the spontaneous formation of a 2D surface oxidation layer that passivates the structure of antimonene endowing the material with an outstanding environmental stability.^{83,166} This technique allows both spatial and chemical resolution of the surface of the flakes to be obtained with great precision. In particular, the XPS data of the flakes revealed the presence of an oxidation layer with a binding energy (BE; 530.9 eV) larger than the value expected for Sb_2O_3 (530 eV) and slightly higher than the value expected for Sb_2O_5 (530.8 eV). The O_2 peak also presented a higher BE value (532.61 eV) compared to most of the metallic oxides (531–528 eV), thus pointing to a superficial oxidized component present only on the outer surface and not presenting a conventional oxidation pattern. These results collected in 2017 by Abellán *et al.* already pointed to the spontaneous formation of antimonene oxides and suggested an explanation for the great stability of antimonene under environmental conditions.

Along this front, an experiment that has been key to understanding this behavior has been the one developed by Lloret *et al.* using highly concentrated bmim-BF₄ exfoliated

antimonene suspensions investigated by XPS. As can be observed in Fig. 22, next to the broad O 1s signal at 533 eV from the surface enriched ionic liquid contamination, the Sb 3d region between 525 and 545 eV of the FL-Sb_{sus} sample (Fig. 22 right, spectrum II) reveals very weak Sb 3d_{3/2,5/2} signals from Sb in oxidation state zero at 528.2 eV for the 3d_{5/2} level, along with minor contributions from Sb in the higher oxidation state at around 530.3 eV. Removing most of the excess IL by heating in ultra-high vacuum clearly showed Sb signals originating mostly from bulk antimony zero (spectrum III). Finally, exposing the heated FL-Sb_{sus} sample without the protecting IL medium for several hours in air led to a significant decrease in Sb(0) and concomitant increase of the oxidized Sb species (spectrum IV); these findings thus demonstrate the role of bmim-BF₄ in stabilizing the antimonene against oxidation, and more importantly, demonstrate the reactive nature of LPE antimonene with a pronounced oxophilicity. These findings are of uttermost importance as they first confirm the passivation of the antimonene surface by an oxide layer that shields it from further structural degradation, thus explaining the high stability of antimonene in comparison to its counterpart black phosphorus.¹⁶⁷ Second, they demonstrated a novel modification approach to tailor its properties by opening a bandgap. In the same context, and by means of first-principles calculations, Zhang *et al.* presented a new class of two-dimensional structures of antimonene oxide. These theoretical predictions are based on a monolayer antimonene with Sb=O double bonds perpendicular to the antimonene plane [type (I) structure], and depending on the oxygen content, tunable direct

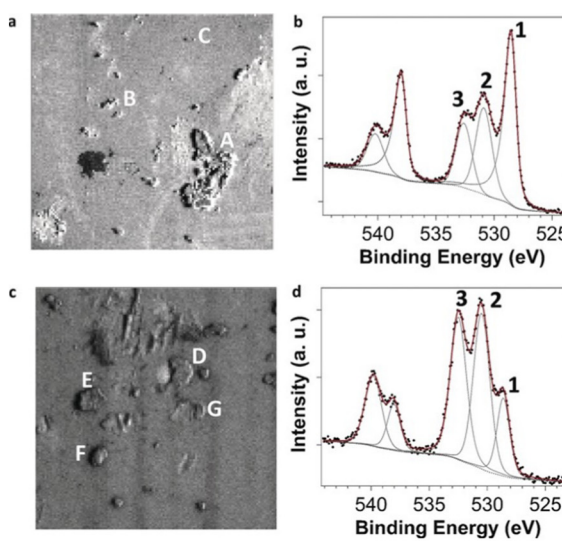
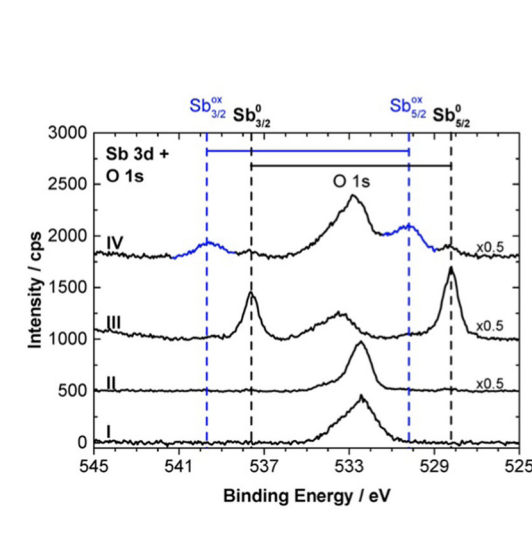


Fig. 22 Left: (a) Sb 3d image for the reference sample. Letters refer to the antimonene flakes studied. (b) Sb 3d peak for the reference sample, corresponding to point B in (a). Numbers identify the different components: 1 (Sb 3d_{5/2}), 2 (oxidized Sb 3d_{5/2}), and 3 (O 1s). Peaks at higher BEs are the Sb 3d_{3/2} components. (c) The Sb 3d image for the functionalized antimonene. (d) Sb 3d peak for the functionalized sample, corresponding to point E in (c). The peak identification and other details are as in (b). The size of images in (a) and (c) is $40 \times 40 \mu\text{m}$. Right: XPS Sb 3d and O 1s region of the neat bmim-BF₄ IL (I) showing oxygen signals from the IL surface contamination layer, of the highly-concentrated FL-Sb suspension (II) showing small signals of non-oxidized (Sb 3d_{5/2} at 528.2 eV) and minor contributions from oxidized (530.3 eV) antimony next to the oxygen contamination, after removal of most of the IL by heating in UHV (III), and after submitting the sample to environmental conditions for a day, showing a drastic decrease in Sb(0). Adapted from ref. 65 with permission from Springer Nature, copyright 2019, and ref. 165 with permission from John Wiley & Sons, copyright 2017.



bandgaps ranging from 0 to 2.28 eV have been calculated.¹⁶⁵ Later on, Wolff *et al.* realized that the fully oxidized antimonene monolayers presented by Zhang *et al.* are metastable and likely undergo a transition into a more stable configuration.¹⁶⁸ Based on density functional theory (DFT) calculations, they proposed a novel type of 2D antimony oxide single- and few-layer structures in which the oxygen atom is incorporated into the antimonene planes and is bound to at least two antimony atoms (Sb–O–Sb) [type (II) structure] (Fig. 23). They have also found out that, depending on stoichiometry and the type of bonding, the proposed 2D layers exhibit different structural stability and electronic properties, ranging from topological insulators to semiconductors with direct and indirect band gaps between 2.0 and 4.9 eV. More interestingly, they foresaw that semimetallic few-layer antimonene can naturally form heterostructures with semiconducting oxidized layers, and they have presented the corresponding vibrational modes, which agree with the experimental findings on liquid-phase exfoliated few-layer antimonene.¹²⁵

With this knowledge in mind, it is apparent that the surface oxidation of antimonene is in most cases inevitable, especially if exposed to ambient conditions. It becomes vital, now more than ever before, to reconsider the outcomes of such drastic surface changes on the measured physical properties of antimonene, which might eventually lead to the better understanding of the current divergence in the reported experimental findings. Furthermore, it might also explain the difficulties being encountered by different research groups in conducting some of what is considered basic measurements in the realm of 2D materials, such as electrical contacts and determination of the electronic band gap.

Through this section of the present review, we aim at stimulating the scientific community to revive the discussions related to this fundamental aspect of antimonene. For instance, Gibaja *et al.* observed that the ultrasonication-induced oxidation of liquid-phase exfoliated antimonene comparatively enhanced the electrocatalytic performance towards the hydrogen evolution reaction (HER) under acidic pH.¹¹² Likewise, Bat-Erdene *et al.* demonstrated that surface-oxidized few-layer antimonene nanosheets can be an efficient nitrogen reduction reaction (NRR) catalyst for the electrochemical synthesis of NH₃.¹⁶⁹ According to the same authors, the activation of the NN bond for the NRR process is facilitated by the oxidized species of antimonene as a result of the more favourable adsorption energy of N₂ caused by the increase in polarity, as compared to non-oxidized antimonene. Furthermore, Duo *et al.* proved that ultrathin antimonene nanoparticles (AMNPs) are effective radiosensitizers that can achieve efficient radiochemotherapy effects.¹⁷⁰ As the authors explained, X-ray irradiation induced reactive oxygen species (ROS) overproduction and accelerated the valence transition to toxic Sb₂O₃, eventually leading to cancer cell death. Finally, yet importantly, it turned out that the oxidation of antimonene is highly desired in many applications, as opposed to the detrimental oxidation of black phosphorus. Therefore, the development of a fast, effective, and reliable method to realize the expected antimonene oxide

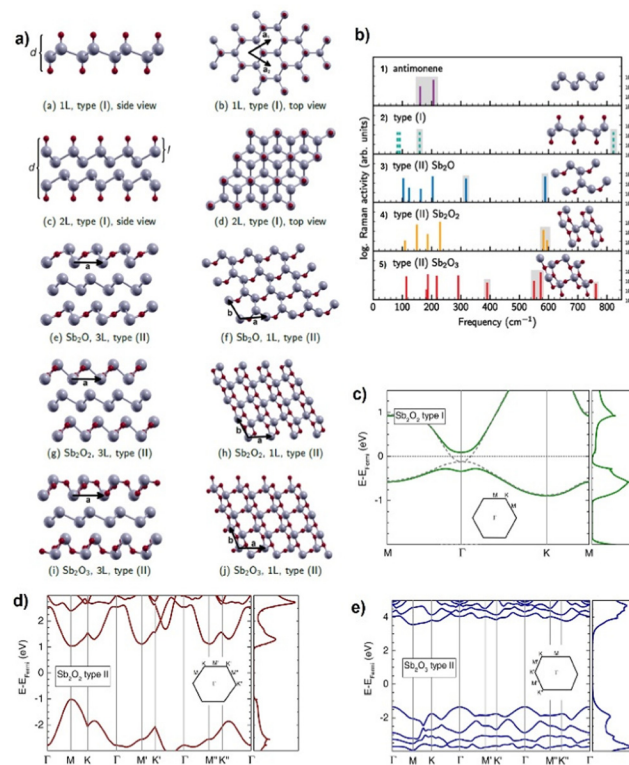


Fig. 23 (a) Type (I) antimonene oxide structures with one layer in side view (a) and top view (b), and two layers in (c) and (d). Type (II) antimonene oxide heterostructures with different stoichiometries of the oxidized layers: (e) and (f) Sb₂O; (g) and (h) Sb₂O₂; and (i) and (j) Sb₂O₃. Type (II) structures in the top view show the oxidized layer only. Oxygen (antimony) atoms are shown in red (gray). Labels a and b on the figures indicate the in-plane lattice vectors. (b) Calculated frequencies of Raman-active vibrational modes in (1) antimonene, (2) type (I), (3)–(5) type (II) antimonene oxide structures 1. (c)–(e) Electronic band structures and density of states calculated using the hybrid functional HSE12 and inclusion of spin-orbit interactions (SOI) for (c) type (I) Sb₂O₂, (d) type (II) Sb₂O₂, and (e) type (II) Sb₂O₃ monolayers. Gray dashed lines are the results without SOI and are almost congruent with the bands with SOI for (c) and (d). The zero of energy is set to the Fermi energy. Adapted from ref. 168 with permission from American Physical Society, copyright 2022.

systems is an urgent demand. Fickert *et al.* have recently introduced a proof of concept of such a method, by which they realized arbitrary heterostructures of Sb₂O₃/antimonene through a spatially resolved laser-induced oxidation of antimonene on the sub- μ m scale.⁹⁸ Although a fine control of the O distribution and concentration is still to be achieved, it provides an indisputable experimental evidence of its practical feasibility. In the same context, Han *et al.* have successfully synthesized two-dimensional Sb₂O₃ crystals down to the monolayer thickness using passivator-assisted vapour deposition (PAVD) (Fig. 24).¹⁷¹ The crystal structure and chemical composition have been confirmed by a set of analytical techniques, namely SAED, HAADF-STEM, EDX and EELS (Fig. 24(d)–(f) and (i)–(l)). Interestingly, the authors unravelled a reversible phase transition between α - and β -phases of Sb₂O₃, which could be controlled either by thermal annealing or by electron-beam irradiation. This remarkable phase transformation is accompanied



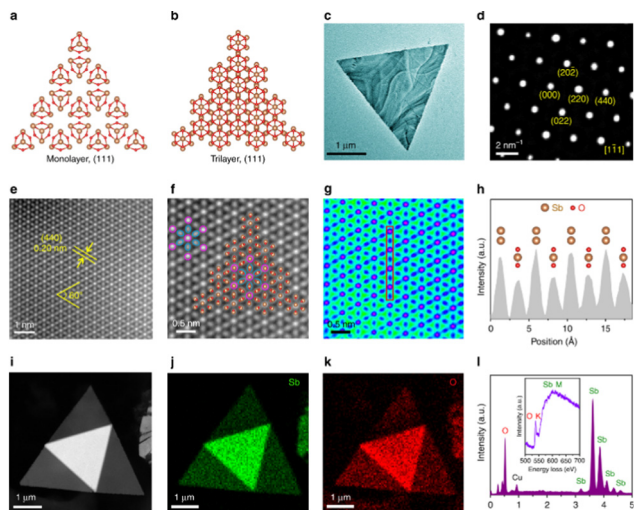


Fig. 24 Atomic structure of 2D Sb_2O_3 molecular crystals. (a), (b) Top-view structural models of monolayer and trilayer Sb_2O_3 flakes with the (111) plane. Brown balls, Sb atoms. Red balls, O atoms. (c) TEM image of a triangular Sb_2O_3 flake. (d) SAED pattern of the Sb_2O_3 flake. (e) Z-contrast atomic-level HAADF-STEM image of the Sb_2O_3 flake showing the perfect atomic lattice. (f) Enlarged HAADF image and the matched atomic ball model, white atoms are marked by yellow circles and gray atoms are marked by blue circles. (g) Scattered electron intensity color image for (f). (h) Intensity line profile along the red box in (g). (i)–(k) HAADF image of a stacked flake and the corresponding elemental maps for Sb and O. (l) EDX and EELS spectra of the Sb_2O_3 flake. Reproduced from ref. 171 with permission from Springer Nature, copyright 2019.

by changes in the electrical properties of the crystals which switches from the insulating α -phase of Sb_2O_3 to the semiconducting β -phase, thus demonstrating their potential applications in phase-change devices.

Last but not least, in contrast to predicted good semiconducting behaviour, experimental reports on antimonene have proved semimetal to semiconductor transitions, high carrier mobilities and strain-tunable indirect-to-direct bandgap transition, as well as the possibility of 2D and 3D topological insulator behaviour. However, due to the large allotropic/polymorphic structural diversity of antimonene, the preparation of structurally controlled ultrathin high-quality samples is very challenging. Indeed, from the physical point of view, the preparation of laterally large, defect-free monolayer antimonene films is the most important synthetic challenge. It is due to the lack of free-standing high-quality monolayer samples for device fabrication that some open questions regarding transport properties of antimonene remain unexplored. We believe that future synthetic efforts should pursue to tackle this challenge. Furthermore, once this hurdle has been overcome, the interesting topologically surface state physics of antimonene will allow its application in quantum computing or spintronic devices to be envisioned.

4.2 Chemical functionalization

Contrary to other 2D materials such as graphene or even other pnictogens like phosphorene,^{60,118,172} the chemical functionalization of antimonene is barely explored. Very scarce reports can be found in the literature regarding this topic.

After initial predictions concerning the effect of adatoms and physisorbed molecules on the overall properties of antimonene,^{96,173} the first experimental evidence of a noncovalent functionalization was reported by Abellán *et al.* in 2017.¹⁶⁶ In that work, the authors report the functionalization of antimonene with perylene diimide (PDI), resulting in a higher charge-transfer performance than that observed in black phosphorus (Fig. 25). The functionalization was carried out by drop-casting a THF solution of a tailor-made

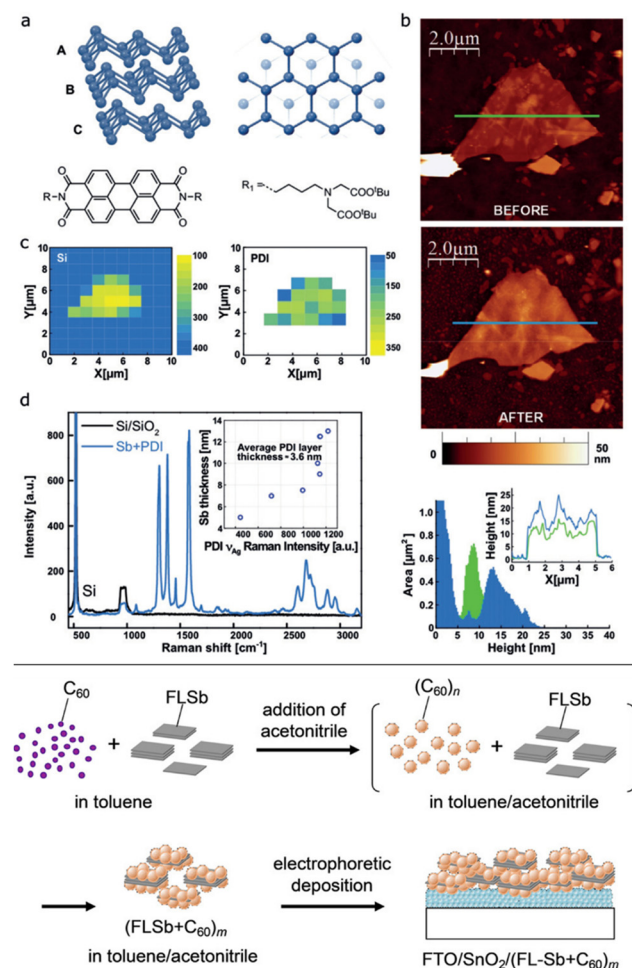


Fig. 25 Top: (a) Structure of β -antimonene (top panel) and the perylene bisimide (PDI) molecule (bottom panel). (b) AFM topographic images showing an antimonene flake of about 10 nm of thickness. Top: Flake as deposited. Middle: Same flake after the functionalization with PDI molecules. Bottom: Height histograms of the flake before (green) and after functionalization (blue), showing an average thickness increase of 4.1 nm. The average PDI coverage in all the studied flakes was 3.6 nm. The inset shows representative profiles corresponding to the lines in the images. (c) Scanning Raman microscopy (SRM) of the same flake. Left: Silicon intensity Raman map showing a decrease in the 521 cm^{-1} signal which clearly reveals the morphology of the flake. Right: Raman intensity mapping shows the exclusive self-assembly of the PDI on the antimonene flakes and not on the Si/SiO_2 substrate. (d) Mean Raman spectra (excitation at 532 nm) of the flake showing the PDI bands as a consequence of the quenching of its fluorescence. Bottom: Formation and electrophoretic depositions of FLSb- C_{60} composite clusters. Adapted from ref. 166 and 176 with permission from John Wiley & Sons, copyright 2017 and 2020.



ethylenediaminetetraacetic acid-PDI derivative on micromechanically exfoliated antimonene flakes on SiO_2/Si substrates. An increase in the average thickness of the material was observed after the functionalization due to the assembly of the PDI molecules on their surfaces. Most recently, Lloret *et al.* also pinpointed that noncovalent functionalization with PDI leads towards the efficient passivation of BP field effect transistors (FET), paving the way for the use of PDIs to preserve the electronic properties of 2D Xenes.¹⁷⁴

The scanning Raman microscopy of the flakes before and after PDI insertion highlights a significant quenching of fluorescence which allows the characterization of the PDI spectrum with excellent quality. The exclusive self-assembly of the PDI molecules on the top of antimonene flakes with no traces of them on the SiO_2/Si substrate is worth noting. Furthermore, there is a clear correlation between the thickness of the organic coverage and the quenching of fluorescence when the former is below 4.5 nm (*ca.* 10 monolayers). Nevertheless, for a thicker coverage, an increase of the background coverage takes place concomitantly with the decrease in PDI Raman intensities. XPS spectra confirm the grafting of the PDI molecules pointing out a shift in the functionalized sample and thus the charge-transfer effect. This effect is ascribed to the electron-withdrawing behaviour of PDI molecules as seen in other 2D materials.¹⁷⁵ Besides, scanning photoelectron microscopy combined with computational studies demonstrates the nature of the strong noncovalent interaction between PDI and antimonene indicating a significant charge transfer effect from the antimonene to the PDI of about twice as large as the observed for phosphorene.⁶⁰ Regarding the possibility to remove the PDI adlayer with temperature, a temperature-dependent Raman analysis was carried out. The results exhibited a complete degradation of the PDI coverage at 400 °C (which starts removing at 300 °C), while antimonene flakes remain stable up to 450 °C, where degradation occurs. In addition to the study of PDI noncovalent functionalization on micromechanically-exfoliated flakes, the work also tests the functionalization of antimonene dispersions prepared by LPE.¹⁶⁶ PDI grafting led to a fluorescence quenching of 80%, higher than that of black phosphorus (66%).⁶⁰ Finally, tetracyano-quinodimethane (TCNQ) was used as an additional grafting molecule (instead of PDI) to further confirm the charge-transfer process.¹⁶⁶

Another work related to noncovalent functionalization is the one reported by Imahori and co-workers with fullerene clusters.¹⁷⁶ In this approach, the resulting hybrid material was synthesized after combining few-layer antimonene with fullerene clusters in a mixture of a poor solvent such as acetonitrile with a good solvent like toluene as depicted in Fig. 25. This work is focused on the photochemical properties, exhibiting a rapid rise in transient conductivity for the hybrid, while no signal is observed in the single component. The reported results open the door to the fabrication of inorganic-organic antimonene hybrids of interest in optoelectronic devices.

In addition to the noncovalent functionalization previously reported with discrete molecules, Zamora and co-workers

recently reported the functionalization of few-layer antimonene with oligonucleotides for DNA sensing.¹⁷⁷ The work highlights the supramolecular interaction between the two materials supported theoretically and experimentally. Theoretical analysis confirmed that the single-stranded oligonucleotides adopt different conformational configurations after interaction with antimonene. Because of that, one of those possible arrangements allows the nucleotide to hybridize with its complementary nucleotide, which is the basis for the development of a DNA sensor.

A completely different approach was recently reported by Su *et al.* showing the synthesis of halogenated antimonene nanosheets composed of oxygen and halogen-decorated amorphous and crystalline domains.¹⁵¹ Halogens are common species used in the surface functionalization of 2D materials because of their high electronegativity, which allows them to bond most of the metallic or non-metallic atoms. In this sense, it is also reported that halogenated 2D materials exhibit enhanced properties if compared with the pristine samples. The halogenated antimonene was synthesized concomitantly with the electrochemical exfoliation of bulk Sb as detailed in Fig. 26. To carry out the halogenation, the electrolyte was an acetonitrile solution of 1-ethyl-3-methylimidazolium (EMIM) $[\text{BF}_4^-]$, [EMIM] $[\text{Cl}^-]$, [EMIM] $[\text{Br}^-]$ or [EMIM] $[\text{I}^-]$ as the source of F, Cl, Br or I atoms, respectively. It is expected that the halogenation process takes place *via* the formation of halogen radicals that intercalate themselves into the interlayer region of Sb crystals, thus weakening the van der Waals interactions between layers and hence expanding the interlayer space.

The XRD characterization of the halogenated sample shows no additional peaks after functionalization, discarding the presence of SbX species. Regarding Raman spectroscopy, the typical vibrational modes of antimonene are suppressed and become undetectable, in a similar way to what happens with fluorinated graphene.¹⁷⁸ All thicknesses were estimated with AFM giving rise to average values of 2.5 nm, ascribed to a few-layer system and taking into account the surface adsorption of O and halogen atoms. Interestingly, XPS also highlights the successful preparation of halogenated antimonene. All spectra evidence Sb-Sb bonds that indicate the intrinsic metallic antimonene, as well as Sb-O and Sb-X bonds. In this context, the degree of functionalization can be semiquantitatively estimated according to the chemical bonds concentration by dividing the overall peak area of Sb-O and Sb-X signals with the Sb $3d_{5/2}$ integral area. The results gave rise to a higher degree of functionalization for Cl and I (*ca.* 47%) followed by F (38%) and Br (25%).¹⁵¹ Concerning the application as photodetectors, the photoresponse performance under UV and visible light of halogenated antimonene was investigated. All samples depicted photocurrent on/off behaviour at 0 V, increasing the photocurrent signal under a fixed external applied voltage of 0.6 V. However, the overall performance of halogenated antimonene is lower than that of the pure material.

Finally, the covalent functionalization of antimonene has been recently reported by Xing *et al.* using 3-glycidoxypropyltrimethoxysilane achieving silane-functionalization.¹⁷⁹ Silane



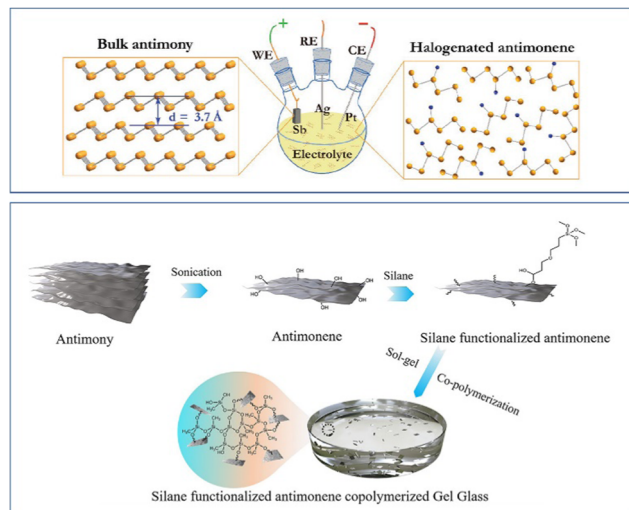


Fig. 26 Top, schematic illustration of the experimental setup for the electrochemical exfoliation and synchronous halogenation of antimonene in an ionic liquid-based electrolyte. Bottom, schematic illustration for the preparation of silane-functionalized antimonene nanosheets and their copolymerized gel glasses. Adapted from ref. 151 with permission from John Wiley & Sons, copyright 2019 and ref. 179 with permission from American Chemical Society, copyright 2021.

functionalization has been explored in 2D materials such as graphene or LDHs, among others.^{180,181} The functionalized antimonene material was obtained as a copolymerized gel glass after the silane hydrolysis and polycondensation reaction (Fig. 26). The resulting gel glass depicted outstanding nonlinear and optical performances, with a strong ultra-broad-band optical limiting performance focused at 532–2150 nm. This optical limiting feature comes as a combination of nonlinear absorption, nonlinear refraction, and nonlinear scattering. This work paves the way for using silane-functionalized antimonene and their copolymerized hybrids in different fields such as biology, energy or optoelectronics.

Rao *et al.* investigated the functionalization of antimonene using group 12 and 13 Lewis acids.¹⁸² They were able to obtain antimonene nanosheets with blue-shifted emission, depending on the Lewis acid used. The interaction of the lone pairs on Sb led to the formation of Lewis acid–base adducts that show a lattice distortion due to the charge transfer from Sb to the Lewis acid and steric repulsions. The functionalized antimonene nanosheets showed a red-shift in the vibrational modes which depended on the strength of the Lewis acid, being more prominent in the A_{1g} mode, showing that the effect in the out-of-plane lattice constant is greater than the in-plane. Together with the rehybridization of the band structure, that allowed the blue-shift of the antimonene nanosheets emission, they demonstrated a functionalization procedure that allowed for tunability and surface passivation.

Rao *et al.* also covalently functionalized antimonene and bismuthine nanosheets using *p*-nitrobenzene diazonium salts.¹⁸³ The study of the reaction mechanisms demonstrated the transfer of lone pairs of electrons from the metallic Sb to

the diazonium salts, forming a quaternary Sb⁺ species with organic moieties predominantly attached through Sb–C bonds. This functionalization led to lattice distortion and charge transfer in the antimonene nanosheets, reducing their bandgap from 2.23 to 2.18 eV, demonstrating the tunability of the optical properties and surface passivation.

5. Applications

As previously commented in this review, the practical applications and properties of antimonene are still emerging. However, there are different works that propose a broad range of feasible applications, including proof-of-concept devices and processes that can be classified into optoelectronic devices, catalysis and electrocatalysis, energy storage, and last but not least, in biomedical applications like cancer therapy or biosensing, which we believe holds great promise for the following years. Below we will detail each of them, showing their potential and future prospects.

5.1 Optoelectronic devices

Optoelectronic devices require materials that are both responsive to visible light as well as able to transfer carriers. In this sense, 2D conductors benefit from appropriate bandgaps and high mobilities.^{15,184} From the practical point of view, Wang *et al.* reported the first use of antimonene with the adequate bandgap as a hole transport layer (HTL) in a perovskite solar cell (Fig. 27).¹⁸⁵ The assembled device was built as an ITO/antimonene/perovskite/PCBM/Bphen/Al heterostructure (PCBM = [6,6]-phenyl-C₆₁-butyric acid methyl ester). Antimonene was obtained after pregrinding bulk Sb into Sb plates that were transformed into antimonene by means of LPE assisted by sonication. The final material was obtained after successive centrifugation at different speeds while studying the bandgap dependence on the thickness, which was tuned in the 0.8 to 1.44 eV range (probably related with different degrees of surface oxidation). The energy level of antimonene closely matches that of methylammonium lead iodide (MAPbI₃) perovskite, hence suggesting an adequate driving force for the hole extraction. This fact is also confirmed by the strongly quenched photoluminescence of the perovskite on top of the ITO/antimonene. This is indicative of the prevented radiative recombination due to photoinduced charge separation. The use of the material as an HTL gave rise to an enhancement of the hole extraction and photocurrent of *ca.* 30%, from 11.2 to 14.6 mA cm⁻² for the HTL-free (control sample) and the antimonene device, respectively. In addition, the external quantum efficiency (EQE) in the entirely visible light was enhanced from 45% to 55–60% for the control and antimonene device, respectively. Additionally, Wang *et al.* also reported the use of antimonene quantum dots (AMQDs) for assembling high-performance organic photovoltaics, enhancing the power conversion efficiency of the device *ca.* 25% higher than that compared with the reference device (Fig. 27).¹⁸⁶



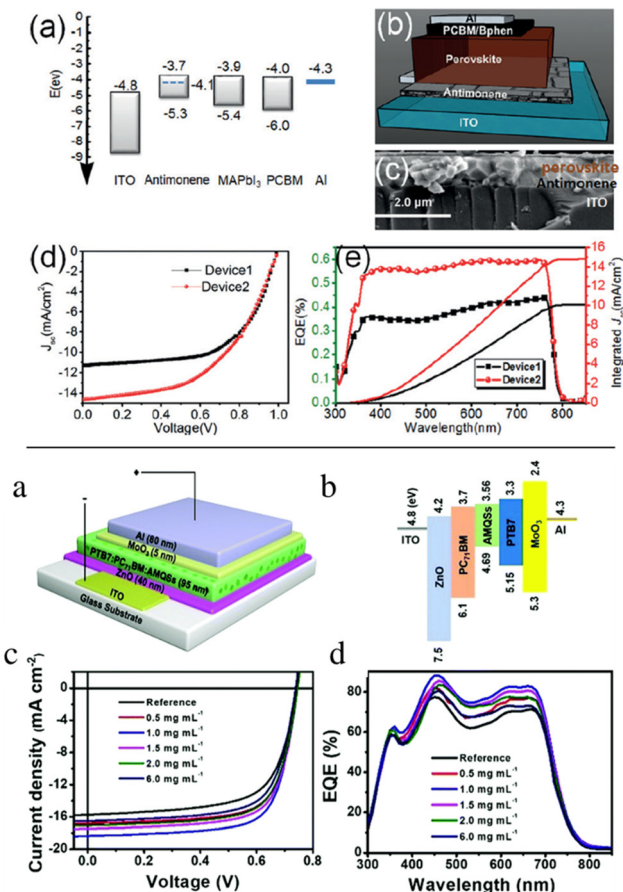


Fig. 27 Top: (a) Comparison of energy levels of each functional layer. The Fermi level of antimonene is represented by the dashed line. (b) Configuration of the antimonene-based device. (c) Cross-sectional SEM image of the studied device. (d) Current-density–voltage (J – V) curves of devices with different architectures. (e) External quantum efficiency (EQE) spectra together with EQE-data-based integrated short-circuit current densities (J_{sc}) for devices 1 (HTL-free) and 2 (ITO/antimonene/perovskite/PCBM/Bphen/Al). Bottom: Device structure and performance characterization of an antimonene organic photovoltaics device. (a) Device architecture including the AMQDs. (b) Band structure of the organic photovoltaics device. (c) Current density–voltage (J – V) curves and (d) EQE characteristics of the best reference device and the devices with different concentrations of AMQDs in their active layer. Adapted from ref. 185 with permission from John Wiley & Sons, copyright 2018, and ref. 186 with permission from the Royal Society of Chemistry, copyright 2018.

After the work of Wang *et al.*,¹⁸⁵ other authors have proposed additional works on the optoelectronic applications. Lu *et al.* reported α -phase few-layer antimonene and AMQDs with nonlinear optical Kerr response using spatial self-phase modulation (SSPM).¹²⁶ They reported a nonlinear refractive index of the few-layer antimonene of *ca.* 10^{-5} cm² W⁻¹, larger than that of AMQDs. Therefore, the study proposes antimonene as a new kind of promising optical Kerr material with enhanced stability, particularly at a short wavelength range. Song *et al.* also reported the study of the broadband nonlinear optical response of few-layer antimonene *via* open-aperture Z-scan laser measurement.¹⁸⁷ Their findings confirmed the use of a few-layer antimonene decorated microfiber as an optical

saturable absorber for ultrafast photonics operation and as a stable all-optical pulse threshold that can suppress the transmission noise, boost the signal-to-noise ratio (SNR), and reshape the deteriorated input signal.¹⁸⁷ That work was further expanded with an additional study in which they use the microfiber as an all-optical Kerr switcher with an extinction ratio up to 12 dB and a wavelength conversion of modulated high-speed signals at a frequency up to 18 GHz.¹⁸⁸ Both reports enlighten the applicability of antimonene-based devices in high-speed optical communication. Finally, the work of Zhang *et al.* demonstrated that semiconducting antimonene nanosheets (probably heavily oxidized ones) exhibited indirect bandgap properties with photoluminescence bandgap of *ca.* 2.33 eV and a lifetime of 4.3 ns. These nanosheets were also satisfactorily tested for the hole extraction layer in planar inverted perovskite solar cells, enhancing the device performance thanks to fast hole extraction and efficient hole transfer at the perovskite/hole transport layer interface.¹⁸⁹

Alongside optoelectronic applications, the preparation of electronic devices as field effect transistors (FETs) is also reported with antimonene. In this topic, Ji *et al.* reported the fabrication of thin film transistors on few-layer antimonene.¹¹⁷ The devices were directly prepared on mica substrates after the synthesis of β -antimonene using HfO₂ as a top gate dielectric. The contacts are made of Cr and Au in the form of 10 and 30 nm thick layers, respectively. Afterwards, a 15 nm thick layer of HfO₂ was grown on the top by means of atomic layer deposition, followed by a final 50 nm thick layer of Au as the top gate electrode. The electrical characterization was carried out with a semiconductor parameter analyser at room temperature, and three typical devices with antimonene thicknesses of 30, 40 and 50 nm were measured at zero gate bias. Regarding the thinnest device (30 nm), an electrical resistance of 600 Ω was measured, with an electrical conductivity of 1.6×10^4 S m⁻¹. This value is in good agreement with that expected for semimetals and consistent with the theoretical predictions. Furthermore, the antimonene devices on mica substrates exhibited high flexibility and no major changes in the conductivity were observed after bending 100 times. These results combined with the wavelength independent high transparency in the visible light range suggest antimonene as a good candidate in the elaboration of flexible transparent conducting electrodes.¹¹⁷

Further studies on preparing transistors are found in the works of Chang¹⁹⁰ and Sun *et al.*¹⁹¹ The first one is focused on novel antimonene tunneling field-effect transistors (TFETs) based on the lateral monolayer (semiconducting)/multilayer (metallic)/monolayer (semiconducting) heterostructure. The proposed antimonene TFETs consist of a semiconducting monolayer of the material in combination with a small metallic multilayer region between the source and the channel. The presence of a multilayer region introduces gapless metallic states that result in the enhancement of the tunneling probability, leading to a larger current. Using the *ab initio* electronic structure and quantum transport calculations for different antimonene TFETs, the simulations conclude that even a

1 nm scale nanostructured multilayer can boost the current and enable abrupt device switching.¹⁹⁰ Overall, antimonene heterostructure TFETs appear to surpass phosphorene ones because of the elimination of the tunneling barrier thanks to the locally constructed multilayer antimonene. Second, the work of Sun *et al.*¹⁹¹ is focused on the device performance of FETs composed of monolayer arsenene and antimonene in the sub-5 nm region using *ab initio* quantum transport simulations. They concluded that the sub-5 nm double gate monolayer assembly of arsenene and antimonene metal oxide semiconductor FETs successfully fulfills the low power requirements of the International Technology Roadmap for Semiconductors in 2028. The calculated performance outperforms that of a MoS₂ analogue in terms of on-current.¹⁹¹

In this context, it is noteworthy to note the work of Han *et al.* in the synthesis of Sb₂O₃ inorganic molecular crystals with thickness down to monolayer previously detailed in the section focused on the oxidation behaviour.¹⁷¹ 2D inorganic molecular crystals are still a challenge because of the difficulties in controlling both crystal phase and growth plane. The work reports the design of a passivator-assisted vapor deposition method that leads to 2D Sb₂O₃. The use of a passivator (InCl₃ or Se), prevents the heterophase growth and promotes the lateral growth in the high-energy crystal planes. The as-synthesized α -phase of Sb₂O₃ depicts a thermal-induced phase transition to a mixed-phase highlighting a thermal hysteresis loop. The mixed-phase is stable from 673 to 353 K. Additionally, the insulating α -phase Sb₂O₃ can evolve to a semiconducting β -phase under heat and electron-beam irradiation. This fact opens the door for new opportunities in potential molecular electronic devices based on Sb.

5.2 Electrocatalysis

One of the most appealing uses of 2D materials such as MoS₂ and WSe₂, as well as those from the pnictogen group, is their application as electrocatalysts for the electrochemical CO₂ reduction reaction (CO₂RR).^{192,193} In this context, Li *et al.* reported the production of few-layer antimonene nanosheets by cathodic exfoliation and tested them as a 2D electrocatalyst for the reduction of CO₂ to formate with high efficiency.¹⁹⁴ The cathodic exfoliation methodology has proven to be a simple, economic, efficient and environmentally friendly method for the large-scale synthesis of 2D materials.^{195,196} If compared with the bulk Sb electrode, the few-layer antimonene nanosheets electrode exhibited a lower onset potential and an enhanced catalytic current density. Furthermore, the electrode made with antimonene depicted a broad peak at *ca.* -1.06 V in the CO₂ saturated solution which is indicative of the CO₂ reduction process. The selectivity regarding H₂ and formate was dependent on the applied potential, pointing to a faradaic efficiency for the formate with a maximum at around 84% at -1.06 V. On the other hand, for the CO the faradaic efficiency was found to be almost constant independent of the applied potential. In addition, the smaller antimonene nanosheets gave rise to a polarization curve I_{mass} 3.6 times that corresponding to the larger flakes, confirming that the enhancement in the

electrocatalytic activity is related to the exposed edge sites after the exfoliation. Finally, the work of Li *et al.* also analysed an antimonene nanosheet-graphene composite, prepared by replacing the Pt anode with a graphite rod during the exfoliation procedure. The composite depicted a higher I_{mass} and selectivity towards formate at lower overpotentials, with a partial current density for formate 1.5 times higher than the pristine antimonene nanosheets. The enhanced activity of the composite is ascribed with both the exposure of large numbers of active sites as well as the strong electronic interaction between antimonene and graphene.¹⁹⁷

Most recently, Gibaja *et al.* reported the electrocatalytic properties of LPE-antimonene suspended in BuOH or NMP towards the hydrogen evolution reaction (HER).¹¹² In this study, two parameters were chosen to compare the electrochemical performance: the Tafel slope and the overpotential required at a current density of 10 mA cm⁻² (Fig. 28). The antimonene prepared in NMP showed a lower Tafel slope (145 mV dec⁻¹) than the antimonene prepared in BuOH, while the latter depicted a lower overpotential value (-0.7 V). The reduction in the onset potential could be ascribed to both the dimensions of the nanosheets and the oxidation degree. Cyclic voltammetry curves before and after the linear sweep voltammetry showed a more pronounced redox activity for the sample in BuOH. These results are indicative of a higher degree of exfoliation of the sample in BuOH than that prepared in NMP. The cyclic voltammetry curves also highlight the irreversible oxidation processes which indicate the oxidation of antimonene into novel surface Sb oxides with specific structural and physical properties. Overall, the higher HER performance of the BuOH sample can be explained based on its better exfoliation degree and anisotropic ratio, which yield a large number of edges and therefore electroactive sites. This HER performance is also linked to the irreversible formation of Sb oxides, an effect which is more pronounced in the BuOH sample than in the NMP one.¹¹²

5.3 Organic catalysis

Lewis acid catalysis is a chemical process that is mainly based on the ability of protons and/or cationic metal atoms to attract electron density from atoms of external molecules and activate them towards the attack of a third chemical entity. However, the use of non-metal species, beyond protons, is much scarcer, in particular of heteroatoms in the low-valence state. Pnictogens in the zero oxidation state (P, As, Sb, Bi) are rarely used in catalysis because of the difficulties in preparing stable materials from these elements. Nonetheless, Lloret *et al.* recently reported the first organic catalysis for antimonene and phosphorene taking advantage of the high degree of exfoliation and the protection against oxidation given by the ionic liquid used in the LPE process. The work focuses on the use of few-layered pnictogens as efficient catalysts for the alkylation of nucleophiles (alcohols, thiols and indoles) with alkyl carbonyl compounds (esters) in good yields and excellent selectivity (Fig. 29).⁶⁵ Regarding the reaction mechanism, it is observed that the pnictogen selectively adsorbs both the nucleophile and



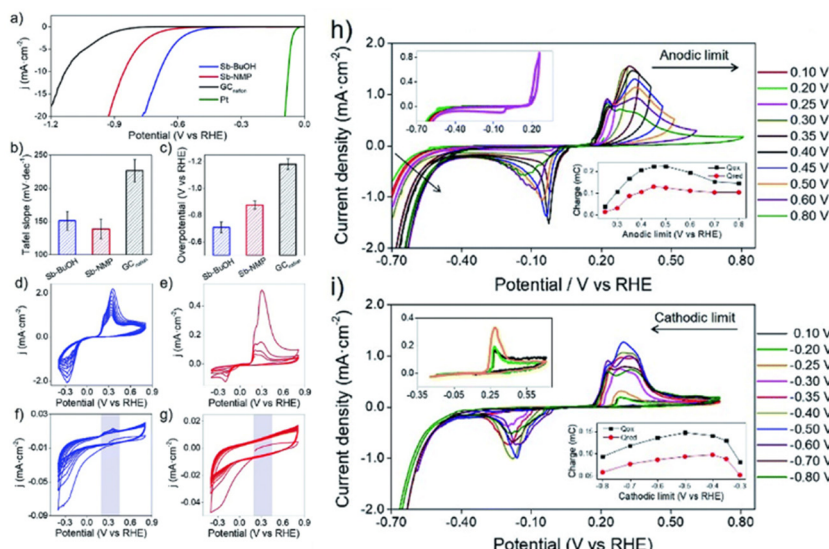


Fig. 28 (a) LSV curves of the different FL antimonene samples, Nafion modified glassy carbon blank and platinum blank. (b) Tafel slope values and (c) overpotentials required for 10 mA cm^{-2} calculated from LSV data. CV curves recorded after LSV measurements of (d) Sb-BuOH and (e) Sb-NMP. CV curves recorded before LSV measurements of the (f) Sb-BuOH and (g) Sb-NMP samples. (h) Voltammograms obtained between -0.9 V and different anodic potential limits measured using 50 mV s^{-1} . (i) Voltammograms obtained between 0.5 V and different cathodic potential limits measured using 50 mV s^{-1} . The insets show the calculated charges obtained when a positive current is passed (Q_{ox}) or negative current is passed (Q_{red}). Adapted from ref. 112 with permission from Royal Society of Chemistry, copyright 2018.

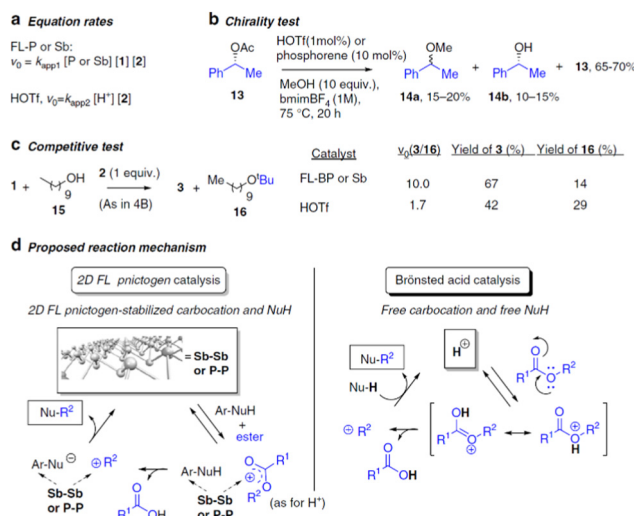


Fig. 29 Mechanistic studies. Experimental evidence and proposed mechanisms for few-layer black phosphorus (FL-Bp), FL-Sb, and superacid triflic acid (HOTf) catalyzed alkylation of nucleophiles with esters in bmimBF_4 . Reproduced from ref. 65 with permission from Springer Nature, copyright 2019.

ester on the surface thanks to the electronic stabilization of the few layers underneath promoting the coupling after formation and stabilization of alkyl carbocations on the lone electron pair network of P and Sb. Additionally, competitive tests between benzyl and decyl alcohols point to a one order of magnitude higher formation rate for the aromatic compound than for the alkyl one.

The higher polarizability of the antimonene results in a better performance than that found for the phosphorene. This mechanism enables Lewis acid catalysed processes under much milder conditions than with protons or metal cations, giving access to the alkylation of soft nucleophiles incompatible with the standard acid catalyzed conditions. This result is also in good agreement with the ability of antimonene to transfer electron charge to planar aromatic molecules, as shown in the noncovalent functionalization studies previously analyzed.¹⁶⁶

5.4 Energy storage

The use of antimonene in energy storage is focused on its application as an anode in Na storage thanks to its high theoretical capacity of 660 mA h g^{-1} for Na storage as well as a low discharge potential of *ca.* 0.5 V (*versus* Na^+/Na).¹⁹⁸ It is worth noting that Na-ion batteries (SIBs) have attracted increasing attention because of sodium abundance, affordability, ease and greenness of extraction procedure.¹⁹⁹ Nonetheless, the larger size of Na^+ compared to Li^+ , the slow insertion and extraction of ions, and large volume changes represent the main drawbacks for developing sodium-ion batteries (NaIBs). In this sense, 2D materials in general, and 2D-pnictogens in particular, are considered one of the best alternatives for electrode materials.

In fact, Gu *et al.* reported the synthesis of composite films made of antimonene nanosheets and graphene.¹²⁴ The synthesis of Sb nanosheets was carried out *via* LPE starting from Sb powder in an IPA solution with constant concentration of NaOH . The resulting composite films were prepared with

tunable densities, and it was observed that the volume change of metallic Sb can be successfully alleviated with the aid of the flexible graphene. As a consequence, the whole density of electrode films can be improved by harnessing the density of the antimonene nanosheets. The optimized composite film depicted a volumetric capacity of $1226 \text{ mA h cm}^{-3}$ in the first cycling. This result is higher than those reported for graphene films (80 mA h cm^{-3}) and Sb/C nanocomposites ($100\text{--}300 \text{ mA h cm}^{-3}$).¹²⁴ In addition to that, the composite exhibited high rate capabilities for Na storage at the highest applied current of 4.0 mA cm^{-2} , whose charge and discharge volumetric capacities were found to be initially 216 mA h cm^{-3} and 110 mA h cm^{-3} after 100 cycles.¹²⁴

Afterwards, Tian *et al.* discovered that the storage of Na in few-layer antimonene takes place through a sodiation/desodiation mechanism with a fast insertion of Na^+ cations and a posterior in-plane alloying reaction. This effect is produced thanks to the few-layer antimonene's small diffusion barrier of 0.14 V .⁷⁴ By *in situ* synchrotron XRD results they confirmed the reversible phase evolution of the antimonene ($\text{Sb} \rightleftharpoons \text{NaSb} \rightleftharpoons \text{Na}_3\text{Sb}$).²⁰⁰ One can clearly observe how the (003) and (110) planes of the few-layer antimonene peaks were shifted to smaller and larger angles during the sodiation and desodiation process, respectively. Besides, the authors reported an experimentally measured capacity value of 620 mA h g^{-1} , so close to the theoretically predicted value of 660 mA h g^{-1} , thus concluding that *ca.* 94% of Sb atoms participate in the process for nearly 150 cycles. The schematic illustration of the proposed discharge–charge mechanism for the Sb–C NIB is depicted in Fig. 30.

Finally, Zamora and co-workers have recently reported the use of antimonene in modified screen-printed electrodes (SPE) for supercapacitors.²⁰¹ They reported a significant improvement in the energy storage capabilities of a carbon electrode in both galvanostatic charging and cyclic voltammetry using $0.5 \text{ M H}_2\text{SO}_4$ as the electrolyte. Antimonene depicted an excellent performance with a specific capacitance of 1578 F g^{-1}

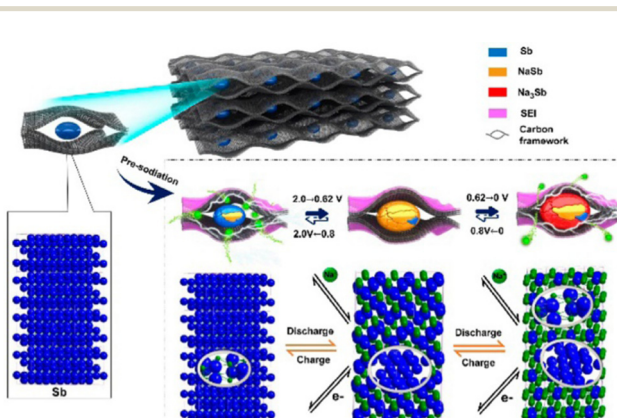


Fig. 30 Schematic illustration of the proposed discharge–charge mechanism for the Sb–C framework film anodes as the high-performance sodium battery with unusual reversible crystalline-phase transformation. Reproduced from ref. 200 with permission from American Chemical Society, copyright 2016.

measured at a current density of 14 A g^{-1} , hence being considered as a highly promising material for energy storage applications. In line with that, the system also exhibited a highly competitive energy and power densities of 20 mW h kg^{-1} and 4.8 kW kg^{-1} , respectively, in combination with good cycling capabilities. It is worth mentioning that the optimum capacitive performance has been achieved for quantities of 3.6 ng of antimonene/SPE.

5.5 Biomedical applications

5.5.1 Cancer therapy. Among the different techniques related to cancer therapy, photothermal therapy (PTT) is considered as one of the most promising strategies thanks to its high efficiency and minimal invasiveness.²⁰² Therefore, it is necessary to develop photothermal agents (PTAs) with both good biocompatibility and excellent photothermal conversion efficacies (PTCEs).

In this context, the outstanding thermophysical properties and chemical reactivity of antimonene make this material an excellent candidate for photothermal cancer therapy.⁹⁸

In this sense, Tao *et al.* reported the synthesis of novel PTAs using AMQDs coated with PEG *via* LPE.¹²⁷ The surface modification of AMQDs with PEG enhanced the biocompatibility and stability in the physiological medium (Fig. 31). The authors reported a clear dependence of the photothermal effect with the concentration, showing a PTCE of 45.5%, higher than that observed for other PTAs such as graphene, MoS_2 or black phosphorus.²⁰³ By measuring the *in vitro* cytotoxicity of the PEG-coated AMQDs all samples exhibited good biocompatibility. Nevertheless, it was observed that the cellular viability decreased when the AMQD concentration increased under near-infrared (NIR) radiation. This decrease was reported to be down to 10% of viable cells at a $200 \text{ }\mu\text{g mL}^{-1}$ concentration. Fig. 31 depicts a clear separation between the living and dead cells, coloured in green and red, respectively, thus confirming the killing of cancer cells using AMQDs as a PTA combined with NIR radiation. Additionally, *in vivo* studies were carried out in mice with MCF-7 tumours, confirming the successful treatment of NIR irradiation combined with intratumour injection of PEG-coated AMQDs, resulting in negligible or no regrowth of the tumour. In this line, Niu *et al.* also focused on the PTCEs of AMQDs presenting the first excited-state dynamics study of the PTT of antimonene. Interestingly, this work highlights that the enhanced PTT of AMQDs comes from their spontaneous partial oxidation. This feature induces additional band edge states that broaden the optical absorption range and strengthen the transition dipole moment. Henceforth, this study proposes the use of oxidative AMQDs in the biomedical field.²⁰⁴

Most recently, the same authors reported a photonic drug-delivery platform based on 2D PEGylated antimonene nanosheets.²⁰⁵ The main advantages of this platform are its excellent photothermal properties, high drug-loading capacity, spatiotemporally controlled drug release activated by NIR light and moderate acidic pH, deep tumour penetration by both extrinsic and intrinsic stimuli, excellent multimodal-imaging properties or substantial inhibition of tumour growth with no



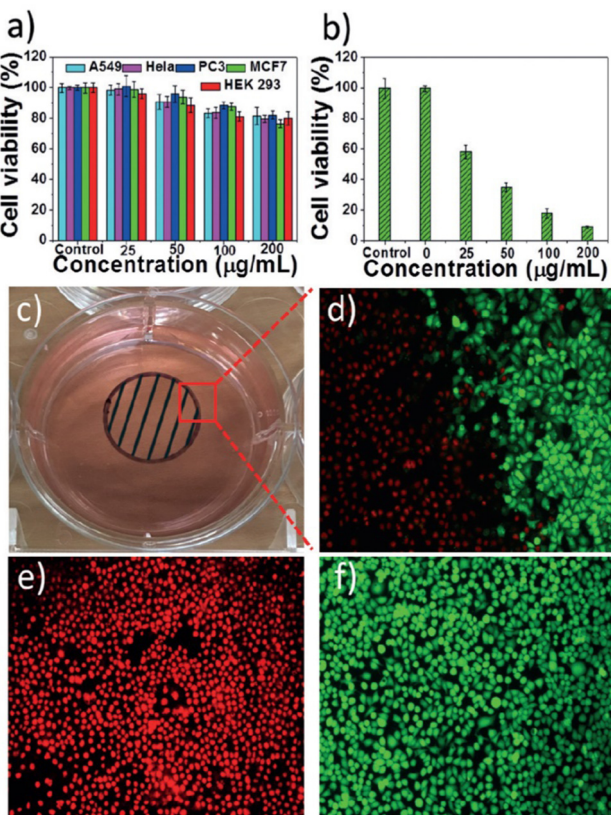


Fig. 31 (a) Cell viability after incubation with only PEG-coated AMQDs. (b) Cell viability of MCF-7 cells treated with PEG-coated AMQDs with NIR (808 nm, 1 W cm^{-2}) for 5 minutes. (c) A photo of the cell culture dish after incubation with PEG-coated AMQDs. The black circle with shadow shows the laser spot. (d)–(f) Confocal images of calcein antimонene (green, live cells) and propidium iodide (red, dead cells) co-stained MCF-7 cells after exposure to NIR irradiation (808 nm, 1 W cm^{-2}). The amplification of confocal images is $100\times$. Reproduced from ref. 127 with permission from John Wiley & Sons, copyright 2017.

apparent side effects and potential degradability, to name a few. Furthermore, the intracellular fate of the antimонene nanosheets was revealed for the first time, leading to a better understanding of the nano-bio interactions of this class of 2D materials. The work of Tao *et al.* represents the first study on antimонene-based photonic drug-delivery platforms, hence paving the way for the application of this pnicogen into cancer theranostics (Fig. 32).²⁰⁵

Additionally, Lu *et al.* also reported the successful use of antimонene for high-performance cancer theranostics.²⁰⁶ In this case, the authors propose the use of cell membrane camouflage alongside dimension optimization and size control to improve the stability of antimонene. The need for enhancing the stability of this material is crucial since its use as a photothermal therapy agent is limited by its rapid degradation in physiological medium. Therefore, the antimонene sheets cloaked with the cell membrane exhibited improved stability as well as increased photothermal efficacy and superior tumour targeting capacity. Furthermore, no notable side effects were observed after using the coated antimонene as a potent

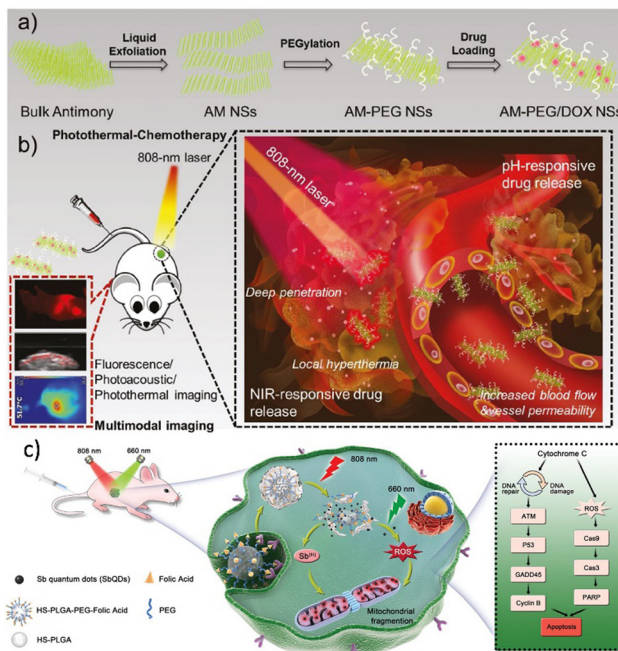


Fig. 32 Schematic illustration of (a) the preparation of 2D antimone-PEG-doxorubicin nanosheets and (b) their administration as photonic nanomedicines for cancer theranostics, (c) Nanopoxia treatment and methods of action. Adapted from ref. 205 and 206 with permission from John Wiley & Sons, copyright 2018 and 2020.

photodynamic/photothermal therapy agent as an anticancer treatment during *in vivo* studies. In conclusion, this strategy reveals as a safe and high-performance modality of cancer therapy.²⁰⁷

One of the current challenges that photothermal therapy faces is the activation of inherent defence mechanisms of tumorous cells due to the overexpression of heat shock proteins in response to the hyperthermia-based treatments. To prevent this thermoresistance, Wu *et al.* constructed a nanocatalyst consisting of calcium carbonate, glucose oxidase and AMQDs.²⁰⁸ When the nanocatalyst encounters the acidic tumour microenvironment, it rapidly degrades releasing the integrated glucose oxidase and the AMQDs. The enzymatic activity of glucose oxidase effectively depletes the cellular glucose levels, reducing energy supplies of the tumorous cells and down-regulating the expression of the heat shock proteins, increasing the therapeutic performance of the photonic hyperthermia treatment. This study reveals the potential of multimodal synergistic therapies for improved biomedical effects.

Interestingly, Qiu *et al.* recently presented a proof-of-concept of an hypoxia-based precision cancer therapy, which they called nanopoxia.²⁰⁹ Taking advantage of the hypoxia conditions usually found in tumours, they developed a nanoplatform based on AMQDs encapsulated by a thermosensitive polymer. Upon 808 nm laser irradiation, the SbQDs@HS-PLGA-PEG-FA outer shell is degraded, releasing the antimonene nanoparticles. In the hypoxic environment of tumours, Sb is



converted into the highly cytotoxic trivalent form Sb(III), while under the normoxia conditions of healthy tissues it is completely oxidized into the less toxic pentavalent form Sb(V), achieving tumour-specific targeting and delivery of Sb. Furthermore, the oxidation of Sb can be regulated and controlled using photothermia,^{98,204,209} allowing a multimodal approach by combining tumour-selective toxicity of Sb, ROS production and hyperthermia treatment.

Interestingly, semimetallic mesoporous Sb-based nanospheres have been reported recently by Chen *et al.*²⁰⁷ By controlling the amount of oxygen in the synthesis of the Sb nanospheres, they selectively etched the nanospheres to obtain mesoporous materials with tuneable pore size, presenting upstanding theragnostic properties as high PTCE (44%), high photoacoustic signal, accelerated photodegradability and high drug loading efficiency.

Finally, Song *et al.* investigated an effective strategy to synthesize Sb nanocrystals in ligand-guided growth strategy, observing a modulation effect on the localized surface plasmon resonance depending on the morphology on the obtained antimony nanopolyhedra.²¹⁰ This allowed them to increase the PTCE up to 62% when the excitation wavelength is well matched with the resonance frequency of the particles. This finding demonstrates the importance of morphology-controllable synthesis methods, allowing the fine tuning of the properties of the Sb-nanomaterials for maximizing their therapeutic potential. These results may serve as inspiration for future developments in antimonene chemistry.

5.5.2 Biosensing. Biosensing applications have been reported for elemental 2D materials such as graphene or phosphorene, taking advantage of the large surface area and the possibility of stabilizing different (macro)molecules by supramolecular interactions.^{211,212} Similarly, the first report showing the use of antimonene as a biosensor was reported by Mayorga-Martínez *et al.*²¹³ In that work, the authors optimized and implemented a highly sensitive and selective phenol enzymatic biosensor using exfoliated antimonene (among other pnictogens) as the sensing platform to enhance the electron transfer process. The resulting biosensor was used for phenol detection following the electroreduction of *o*-quinone to catechol (Fig. 33). The pnictogen nanosheets were obtained by shear-force exfoliation that underwent a down-sizing process alongside the delamination of the material. The phenol biosensor based on antimonene exhibited the best results among the tested pnictogens (phosphorene, arsenene, antimonene and bismuthene), with enhanced analytical performance from the point of view of sensitivity, selectivity, linearity and reproducibility. Furthermore, the boosted electrocatalytic reduction of phenol allows its detection in the presence of common interferents and real samples, with LOQ and LOD 10 times below the recommended limit.²¹³

Another study in the field of sensing can be found in the work of Xue *et al.*²¹⁴ reporting a surface plasmon resonance sensor which is based on 2D antimonene, with the aim of the label-free detection of clinically relevant bio-markers such as miRNA-21 and miRNA-155. By means of first-principles

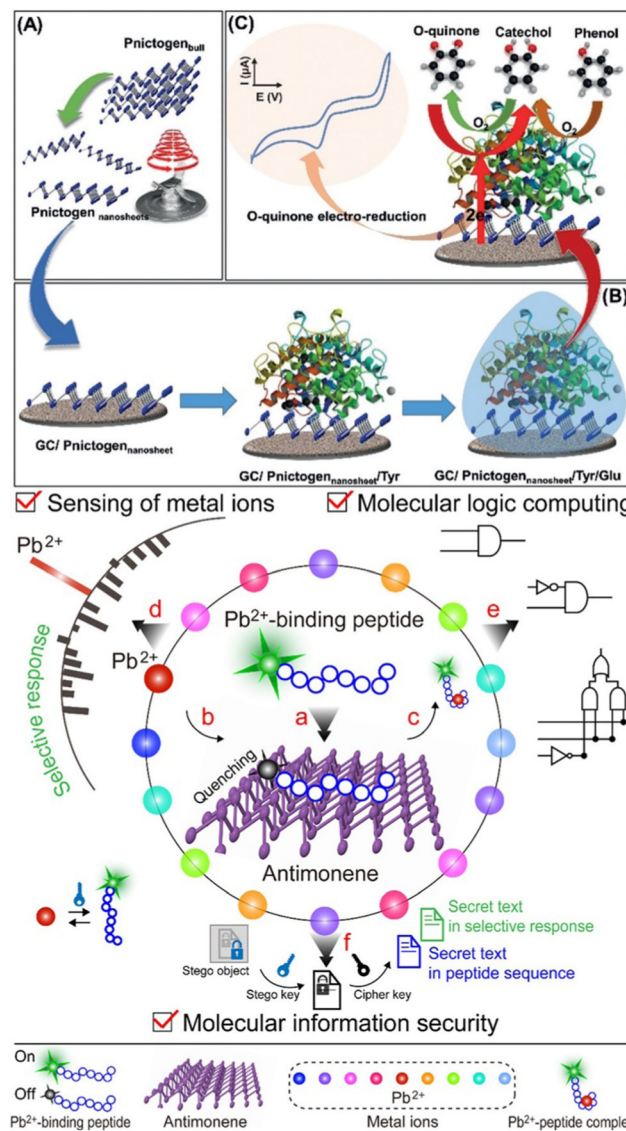


Fig. 33 Top (a) Shear exfoliated pnictogens using a kitchen blender. (b) Biosensor preparation using layer-by-layer drop-cast pnictogen nanosheets, tyrosinase (Tyr), and glutaraldehyde (Glu) onto a glassy carbon (GC) electrode. (c) Chemical mechanism of phenol detection by a biosensor based on exfoliated pnictogen and Tyr. Bottom, scheme of a peptide fluorescence sensing system using antimonene and its different applications in the detection of Pb^{2+} , molecular logic operations and crypto-steganography. Adapted from ref. 212 with permission from John Wiley & Sons, copyright 2019, and ref. 217 with permission from American Chemical Society, copyright 2022.

energetic calculations it is estimated that antimonene exhibits stronger interaction with single-stranded DNA than graphene because of the more delocalized 5s/5p orbitals in antimonene. With an enhanced LOD up to 10,000 times compared to existing miRNA sensors, antimonene reveals as a promising material in the development of these biosensors.²¹⁴

Taking advantage of this surface plasmon resonance, Singh *et al.* developed a nanolayered biosensing device for the detection of hemoglobin concentrations, by combining BaTiO_3 and antimonene.²¹⁵ In this device the antimonene is used as the



bio-recognition element, due to the large surface area, its hydrophilic nature and higher absorption energy. This study further confirms the usefulness of antimonene in the sensing field, especially for biomedical applications.

Recently, Najam-ul-Haq and co-workers have developed a cost-effective biosensor based on catalase immobilized in AMQDs with the aim of determining H_2O_2 . The key aspect in determining H_2O_2 is its role as a predictive biomarker of different types of cancers. The resulting material is used for the electrochemical sensing of H_2O_2 and tested in ovarian cancer serum pointing to a great stability and a quantitative determination of the analyte.²¹⁶

As a final point, antimonene has also been proposed as a platform for peptide-based sensing. In the study by Bu *et al.*,²¹⁷ antimonene has been used as a universal quencher of fluorescence for peptides used to sense the presence of metal ions by fluorescence. The authors claim that this platform can be used for molecular logic computing and as a nanosystem for cryptography (Fig. 33). This report provides a basic paradigm for the use of antimonene in molecular sensing platforms, opening the door to novel applications.

5.5.3. Perspectives on the biomedical applications of antimonene. The great potential exhibited by antimonene in biomedical applications deserves, in our opinion, a dedicated analysis, since we think that many of the most interesting applications of this material will arise in this field. Nanomaterials physicochemical properties, like size, shape, charge or surface chemistry are key factors influencing their performance in biological systems.^{218,219} Interactions with biological molecules and cells are intimately related to the surface properties of the material. Higher surface area will increase the number of interactions, and the surface chemistry will favour certain types of bindings with biomolecules, like protein corona formation,^{220,221} contact with cell membrane or with the extracellular matrix. Those interactions may affect at the same time the nanomaterial, impacting its stability, oxidation state, toxicity and even the (opto)electronic properties. This change of properties can be exploited from different perspectives, like in biosensing devices, as previously seen,^{213,214} or can be exploited to further enhance the therapeutic effect of a given nanodevice.²⁰⁹ The understanding of how these interactions govern the behaviour of antimonene in biological environments is very important for obtaining proper nanodevices with the desired function for a certain biomedical application. Unfortunately, there is a lack of systematic studies about the behaviour of cells in response to the properties of a nanomaterial, as well as the alteration of the properties of the nanomaterial upon the interaction with cellular environments (cellular media, extracellular matrix or intracellular environments.), despite being essential for the rational design of nanodevices.²²²

As previously discussed, antimonene shows a great potential in the biomedical field, with the first studies reporting the use of this material as PTA showing its great PTCE, together with an excellent biocompatibility of the PEG coated antimonene, both *in vitro* and *in vitro*.^{127,205,206} In terms of toxicity, the coated

antimonene platforms have been tested in different cell lines, revealing a good biocompatibility, which shows enhanced toxicity after NIR radiation exposure due to the partial oxidation of the material.²⁰⁴ Furthermore, the first investigations of the internalization mechanisms and intracellular fate of PEGylated antimonene nanosheets demonstrated that they enter the cell primarily through micropinocytosis and caveolin-dependent pathways, and follow the classic endocytic pathway through early and late endosomes to end up in lysosomes.²⁰⁵ Nevertheless, most of the studies reported so far use functionalized antimonene, being coated with PEG or camouflaged using tumorous cell membranes.^{127,204–206} Even though these studies revealed the biomedical potential of antimonene, the use of only functionalized antimonene hinders the understanding of the effects and behaviour of the material in cellular environments, which is of critical importance for the safe and efficient use of nanomaterials.

Getting the picture of what happens with antimonene when it interacts with the biological environments (cell culture media and/or blood plasma) up to the subcellular fate of the material, addressing the alterations that it may suffer, is not an easy task. From the moment that antimonene enters in contact with the cell culture media, protein–nanoparticle interactions form the protein corona, which might affect the antimonene chemical stability and undoubtedly affects the cellular response towards the nanomaterial.^{220,221} The surface modification plays a crucial role in its uptake and internalization, as it is intimately related to the size, shape and surface chemistry of the nanoparticles, affecting the relative contribution of the uptake route.^{218,219} This in turn affects the intracellular fate of antimonene, as each internalization mechanism has different trafficking pathways, ending in diverse subcellular compartments. It is important to keep in mind that cellular organelles and compartments have utterly different chemical environments, from neutral pH compartments like endosomes or caveosomes, high molecular density organelles as the nucleus, or even very acidic and very enzymatic active lysosomes, which could affect the chemical evolution of the nanomaterial, probably modifying the nanomaterial toxicity and response. The reliable assessment of the nanomaterial and cellular association appears as a key factor in understanding the role of novel nanomaterials as biomedical tools. Nevertheless, there is no single analytical methodology capable of providing all the required information, compelling a multimodal approach, a promising strategy that offers analytical and statistical evidence to be able to obtain the full picture of these complex processes.²²³

6. Conclusions and outlook

This review summarizes the most important aspects of antimonene highlighting why it is considered as one of the most promising 2D materials for energy storage, high-performance opto-electronic devices, DNA sensors, cancer therapy or novel vdW heterostructures, to name a few. From its properties to its



applications, this work thoroughly reviews the current knowledge on antimonene and provides an in-depth description of the material with the purpose of stimulating future research. In this sense, despite the tremendous efforts carried out during the last few years, a complete fundamental understanding of the structure–property relationships is still missing. Indeed, in contrast to graphene and other post-graphene 2D materials, experimental reports on antimonene have proved semimetal to semiconductor transitions, high carrier mobilities and strain-tunable indirect-to-direct bandgap transition, as well as the possibility of 2D and 3D topological insulator behaviour. However, due to the large allotropic/polymorphic structural diversity of antimonene, the preparation of structurally controlled ultrathin high-quality samples is very challenging. Indeed, from the physical point of view, the preparation of laterally large, defect-free monolayer antimonene samples is one of the most important synthetic challenges. This is due to the lack of this type of samples, that some open questions regarding electronic (and spintronic) properties of antimonene remain unexplored. We believe that future synthetic efforts should pursue to tackle this challenge. In any case, scientists can take advantage of the unprecedented rich allotropy of antimonene that can be engineered to some extent by playing with the synthetic strategy, substrates, morphology or thicknesses. In addition, it presents a series of phase transition mechanisms that imply great changes at the electronic level, and therefore in its interfacial properties. This dynamic behaviour holds great promise in the development of new heterostructures, both vdW and covalent, in which one can envision a novel epitaxy landscape of interest for twistronics. Another aspect that clearly differentiates antimonene from its peers (and specially from graphene) is its high chemical reactivity, leading to complex oxidation behaviour, which underlies the origin of antimonene's high environmental stability. This mouldable behaviour of antimonene that can be chemically engineered from the ground up distinguishes it from other 2D materials where little control can be exercised over the structure and properties.

To this end, we can foresee that the development of antimonene chemistry will allow in a short time to obtain a controlled functionalization and a fine adjustment of the properties of the material. This will enable the development of covalent and non-covalent heterostructures, hybrid materials, and functional devices that will expand the applications of antimonene to new and exciting horizons.

In terms of applications, the use of different forms of antimonene in optoelectronic devices holds great promise in field effect transistors, perovskite solar cells or non-linear optics. It is worth mentioning that in some cases the oxidized $\text{Sb}_x\text{O}_y/\text{Sb}$ heterostructures outperform pristine antimonene, emphasizing the importance of controlling the synthesis of the materials. Particularly, this aspect represents one of the main differences with lighter 2D-pnictogens as phosphorene, in which surface oxidation leads to a complete degradation of the flakes. The surface reactivity of antimonene has turned out to be key for the development of catalysts, both for boosting organic reactions or electrochemical processes. In fact, while in

organic catalysis it is important to preserve the antimonene surface intact in the zero oxidation state, in the electrocatalytic processes such as the hydrogen evolution reaction (HER) or the electrochemical CO_2 reduction reaction, surface oxidation plays a fundamental role. We believe that through a deep understanding of the reaction mechanisms, the support of theoretical calculations and the development of new hybrid materials, current limitations exhibited by these systems will be overcome. For energy storage applications, both supercapacitors and sodium ion batteries (NIBs) have been the applications that have received the most attention. For example, in the case of NIBs, it is necessary to improve the integration of antimonene with other nanocarbon materials such as graphene to improve the cyclability of batteries by minimizing the pulverization as a consequence of charge and discharge cycles, as well as the irreversible formation of intermetallic phases. One of the greatest challenges faced by this field of research is the development of synthetic methodologies that allow obtaining antimonene of high crystalline and morphological quality on a large-scale. Along this front, novel colloidal approaches are shown as a promising alternative for applications in electronics and beyond, complementing the most established vdW epitaxy processes.

Last but not least, access to monodisperse systems such as AMQDs or colloidal routes open the door to one of the most suggestive fields of application of antimonene: biomedicine. In this direction, to the best of our knowledge, there is a lack of systematic studies aiming to rationalize the chemical identity of bare antimonene (synthetic method, size, morphology and surface chemistry) on the biomedical performance of the material, hampering the comprehension of the mechanisms governing the cellular response towards antimonene and *vice versa*. It will be paramount to understand the mechanism controlling the uptake of antimonene nanomaterials and how they are influenced by its physicochemical properties, as well as depicting the toxic mechanism of antimonene, relating them to the chemical evolution that it may suffer during the interaction (e.g. oxidation). A methodical study of how the properties of antimonene affect its biological performance will allow a rational design of novel antimonene biomedical platforms, with control over their stability, toxicity and biodistribution, enabling more precise biomedical nanodevices based on this novel nanomaterial. Overall, we hope that this review article will inspire novel research regarding antimonene paving the way for the development of new exciting applications.

Conflicts of interest

There are no conflicts to declare.

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