2D materials and van der Waals heterostructures

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BACKGROUND: Materials by design is an appealing idea that is very hard to realize in practice. Combining the best of different ingredients in one ultimate material is a task for which we currently have no general solution. However, we do have some successful examples to draw upon: Composite materials and III-V heterostructures have revolutionized many aspects of our lives. Still, we need a general strategy to solve the problem of mixing and matching crystals with different properties, creating combinations with predetermined attributes and functionalities.

ADVANCES: Two-dimensional (2D) materials offer a platform that allows creation of heterostructures with a variety of properties. One-atom-thick crystals now comprise a large family of these materials, collectively covering a very broad range of properties. The first material to be included was graphene, a zero-overlap semimetal. The family of 2D crystals has grown to includes metals (e.g., NbSe2), semiconductors (e.g., MoS2), and insulators [e.g., hexagonal boron nitride (hBN)]. Many of these materials are stable at ambient conditions, and we have come up with strategies for handling those that are not. Surprisingly, the properties of such 2D materials are often very different from those of their 3D counterparts. Furthermore, even the study of familiar phenomena (like superconductivity or ferromagnetism) in the 2D case, where there is no long-range order, raises many thought-provoking questions. A plethora of opportunities appear when we start to combine several 2D crystals in one vertical stack. Held together by van der Waals forces (the same forces that hold layered materials together), such heterostructures allow a far greater number of combinations than any traditional growth method. As the family of 2D crystals is expanding day by day, so too is the complexity of the heterostructures that could be created with atomic precision.

When stacking different crystals together, the synergetic effects become very important. In the first-order approximation, charge redistribution might occur between the neighboring (and even more distant) crystals in the stack. Neighboring crystals can also induce structural changes in each other. Furthermore, such changes can be controlled by adjusting the relative orientation between the individual elements.

Such heterostructures have already led to the observation of numerous exciting physical phenomena. Thus, spectrum reconstruction in graphene interacting with hBN allowed several groups to study the Hofstadter butterfly effect and topological currents in such a system. The possibility of positioning crystals in very close (but controlled) proximity to one another allows for the study of tunneling and drag effects. The use of semiconducting monolayers leads to the creation of optically active heterostructures.

The extended range of functionalities of such heterostructures yields a range of possible applications. Now the highest-mobility graphene transistors are achieved by encapsulating graphene with hBN. Photovoltaic and light-emitting devices have been demonstrated by combining optically active semiconducting layers and graphene as transparent electrodes.

OUTLOOK: Currently, most 2D heterostructures are composed by direct stacking of individual monolayer flakes of different materials. Although this method allows ultimate flexibility, it is slow and cumbersome. Thus, techniques involving transfer of large-area crystals grown by chemical vapor deposition (CVD), direct growth of heterostructures by CVD or physical epitaxy, or one-step growth in solution are being developed. Currently, we are at the same level as we were with graphene 10 years ago: plenty of interesting science and unclear prospects for mass production. Given the fast progress of graphene technology over the past few years, we can expect similar advances in the production of the heterostructures, making the science and applications more achievable.

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Production of van der Waals heterostructures. Owing to a large number of 2D crystals available today, many functional van der Waals heterostructures can be created. What started with mechanically assembled stacks (top) has now evolved to large-scale growth by CVD or physical epitaxy (bottom).
The physics of two-dimensional (2D) materials and heterostructures based on such crystals has been developing extremely fast. With these new materials, truly 2D physics has begun to appear (for instance, the absence of long-range order, 2D excitons, commensurate-incommensurate transition, etc.). Novel heterostructure devices—such as tunneling transistors, resonant tunneling diodes, and light-emitting diodes—are also starting to emerge. Composed from individual 2D crystals, such devices use the properties of those materials to create functionalities that are not accessible in other heterostructures. Here we review the properties of novel 2D crystals and examine how their properties are used in new heterostructure devices.

Among the tools for band-structure engineering in van der Waals heterostructures are the relative alignment between the neighboring crystals, surface reconstruction, charge transfer, and proximity effects (when one material can borrow the property of another by contact via quantum tunneling or by Coulomb interactions). Thus, a moiré structure for graphene on hexagonal boron nitride (hBN) leads to the formation of secondary Dirac points and gap opening in the electronic spectrum, and spin-orbit interaction can be enhanced in graphene by neighboring transition metal dichalcogenides (TMDCs) (11, 12).

Here we provide a review of 2D materials, analyzing the physics that can be observed in such crystals. We discuss how these properties are put to use in new heterostructure devices.

**Transition metal dichalcogenides**

Transition metal dichalcogenides, with the formula MX\(_2\) (where M is a transition metal and X is a chalcogen), offer a broad range of electronic properties, from insulating or semiconducting (e.g., Ti, Hf, Zr, Mo, and W dichalcogenides) to metallic or semimetallic (V, Nb, and Ta dichalcogenides). The different electronic behavior arises from the progressive filling of the nonbonding d bands by the transition metal electrons. The evolution of the electronic density of states (DOS) is shown in Fig. 1 (adapted from (13–17)) for the most stable phase of each of the dichalcogenides.

All TMDCs have a hexagonal structure, with each monolayer comprising three stacked layers (X-M-X). The two most common polytypes of the monolayers are trigonal prismatic (e.g., MoS\(_2\) and WS\(_2\)) and octahedral (e.g., TiS\(_2\)); these terms refer to the coordination of the transition metal atom. Inversion symmetry is broken in the former, giving rise to piezoelectricity and having important consequences for the electronic structure. In addition, many of the tellurides, TeS\(_2\), ReS\(_2\), and other dichalcogenides adopt lower-symmetry structures in which the metal atom is displaced away from the center of the coordination unit.

**Metallic TMDCs**

As shown in Fig. 1, the DOS of metallic TMDCs has two main properties: (i) The Fermi level of the undoped material is always crossing a band with d-orbital character; implying that the electrons move mostly in the metal layers, and (ii) the DOS at the Fermi level is usually quite high, which hints at a common explanation for the phase transitions observed in these materials (18).

The interest in these materials comes from the existence of CDWs and superconductivity in their phase diagrams (19). Whereas the CDW phase has clear insulating tendency (opening a gap and suppressing the DOS at the Fermi level), the superconducting phase needs finite DOS to exist, resulting in a direct competition between the two many-body states. This competition leads to a complex phase diagram with the presence of inhomogeneous electronic and structural patterns, which have been observed in electron microscopy and neutron scattering in the 3D parent compound. Measurements of specific heat and magnetic susceptibility in 3D samples have shown partial gaps of the Fermi surface. In some cases (e.g., TaS\(_2\)), the CDW transition leads to the decoupling of the unit cells along the axis perpendicular to the planes, with an enormous increase in transverse resistivity.

These unusual properties of metal TMDCs have been the subject of intense theoretical debate, but no consensus has been reached. The mechanism for the CDW transition does not fit standard weak-coupling mean field theories such as Fermi surface nesting or transitions induced by van Hove singularities. Many angle-resolved photoemission experiments have been performed in 3D samples with contradictory results (20). The existence of several Fermi surface sheets and the partial gapping of the Fermi surface make the theoretical interpretation of the experimental data quite difficult. Furthermore, the coexistence of CDWs and superconductivity (clearly seen in local probes) (21) indicates that many-body effects play a very important role in these materials.

Critical information can be obtained from transport data in these materials, when transport measurements are performed in conjunction with the application of electric and magnetic fields. External electric field changes the Fermi energy and the carrier concentration in the 2D material, without the need for chemical doping (which was the case in 3D materials and which introduces appreciable disorder).

In a recent experiment on 1T-TiS\(_2\), a 2D film was encapsulated by hBN and subjected to transverse electric and magnetic fields (22). By applying an external electric field to change the carrier density, it was possible to tune the CDW transition temperature from 170 to 40 K and, concomitantly, the superconducting transition temperature from 0 to 3 K. Controlling the transition temperatures using an electric field allows the critical exponents for the phase transition to be determined.
with high accuracy. Moreover, applying an external transverse magnetic field at the same time reveals novel physical phenomena associated with periodic motion of the Cooper pairs in the superconducting phase. Such behavior seems to be tied up with the formation of discommensurations between different CDW domains—namely, the electronic system broke down in perfectly ordered superconducting and CDW domains.

Phase transitions in 2D materials
Electrons in a solid are characterized by several quantum numbers that include charge and spin. Due to electron-electron or electron-ion interactions, electrons can organize themselves in phases characterized by an order parameter that is associated with these degrees of freedom. In a CDW state, as in the case of TMDCs, the order parameter is the local electron density \( \rho(r) \), where \( r \) is the position vector, which orders with a well-defined periodicity. This periodicity implies that the Fourier transform of the density, \( \rho(Q) \), where \( Q \) is the so-called ordering wave vector of the CDW, acquires a finite expectation value. For a CDW, the expectation value of \( \rho(Q) \) is the order parameter, which is zero in the disordered (or normal) phase and finite in the ordered phase. The transition between these phases can be driven by external forces such as electric, mechanical, and thermal.

Two-dimensional systems play a particular role in the physics of phase transitions. For a system with a continuous order parameter, it is not possible to have true long-range order in less than three dimensions at any finite temperature \( T \), implying that even minute thermal fluctuations can destroy order \( (23) \). In two dimensions, long-range order is possible only at strictly zero temperature. At \( T = 0 \), it is also possible for a system to be disordered if one varies an external parameter such as pressure or electric field, \( E \) (Fig. 2). The point at which a system becomes ordered at \( T = 0 \) is called the quantum critical point, and the transitions are called quantum phase transitions. In this case, it is not thermal motion that drives the system from order to disorder but quantum fluctuations. In this type of transition, the scale at which order is created is characterized by a correlation length \( \xi \), which diverges at the phase transition as

\[
\xi(E) \sim 1/|E - E_\text{c}|^\nu
\]

where \( E_\text{c} \) is the critical field and \( \nu \) is the critical exponent. Fluctuations of the order parameter at different points in space decay exponentially with \( \xi \). Variations in length scales lead to fluctuations in energy scales as well. In a second-order phase transition, the characteristic energy scale, \( \Delta \), associated with the particular order (that is, the energy gap in the system) vanishes at the phase transition with another dynamical exponent, \( \nu \), as

\[
\Delta(E) \sim 1/\xi^\nu - |E - E_\text{c}|^{\nu\nu}
\]

The simplest theory for understanding the effect of critical fluctuations close to a phase transition assumes that the order parameter (SDW, CDW, etc.) couples locally with the relevant degree of freedom (spin, charge, etc.). The resistivity is then given by the standard de Gennes–Friedel formula, in which the electron mean free path scales with the differential scattering cross section of the order parameter fluctuations.

In a classical phase transition, the behavior is driven by thermal fluctuations. The resistivity has the same kind of singularity as internal energy, implying that the critical behavior of the derivative of the resistivity is the same as the specific heat at the phase transition. This indicates that in a classical phase transition the critical behavior is marked by an inflection point in the resistivity at \( T_\text{c} \).

Even though, for a 2D system, long-range order is not possible at any finite temperature, the

Fig. 1. Electronic properties of different classes of 2D materials. The Fermi level is set to the zero of the energy scale. The DOS is given in states per electron volt per cell.
system can undergo a transition to quasi-long-range order (KT transition) with the presence of vortex-antivortex pairs (24). In this case, the order parameter correlation length obeys the exponential dependence with temperature $T$

$$\xi(T) \sim a \exp(b/|T - T_{KT}|^{1/2})$$

where $a$ and $b$ are constants and $T_{KT}(E)$ is the KT transition temperature, which is a function of the external tuning parameter $E$. The resistivity scales with some power of the inverse correlation length and hence is supposed to have an exponential dependence with temperature.

**Semiconducting group-VIB dichalcogenides**

Because of the charge confinement and reduced dielectric screening, the optical properties of semiconducting 2D materials are dominated by excitonic effects. The optical spectra of MoS$_2$, one of the most studied TMDCs, is characterized by three main transitions, named the A, B, and C peaks. The A exciton is the lowest energy corresponding to the fundamental optical gap of the material. The corresponding exciton binding energy is ~1 eV, according to theory. The B exciton also corresponds to a transition at the K point but for opposite spin. The C peak is of a different nature, as it has contributions from excitons from a large, annular-shaped region of the $k$-space with nearly identical transition energies.

In nearly neutral monolayer samples, other quasi-particles have been observed, including positively and negatively charged excitons (i.e., trions) and bie excitons (25–27). The large trion binding energies (20 to 30 meV) have no parallel in traditional semiconductors and allow for these quasi-particles to be observed even at room temperature.

The series of Rydberg exciton states above the 1s (A) exciton of WS$_2$ reveals an exciton series that deviates considerably from the hydrogen model (28, 29). Not only do the 1s, 2s, 3s, ... $n$s states have a closer spacing for small $n$, reflecting a weaker screening at short range (~$\log r$, where $r$ is the electron-hole separation) (28), they also have an entirely different dependence on the angular momentum. Ab initio GW calculations show that the states in the same shell but with higher angular momentum are at lower energy levels—that is, 3d, 3p, and 3s are in order of increasing energy.

From the technologic point of view, however, the most relevant transitions are those close to the fundamental gap at $K(K')$ points of the Brillouin zone, which can be used for manipulating quantum information stored as spin and momentum (valley index) of individual electrons, holes, or excitons. The selection rule for optical transitions is valley-dependent, with the $K(K')$ valley coupling exclusively to right (left) circularly polarized light. Thus, the valley index, or pseudospin, can be controlled coherently by using polarized light. Because the two valleys have non-zero and symmetrical Berry curvature, in the presence of in-plane electric field they give rise to Hall currents with sign depending on the valley index, an effect known as the valley Hall effect. The orbital magnetic moment is also valley-dependent, which allows for coupling with magnetic fields (30–36).

Quantum dots of TMDCs inherit the valley properties of the monolayer and therefore are appealing for valleytronics due to the possibility of controlling spin and valley states of single confined electrons or holes—for example, via interaction with propagating single photons. Quantum dots can be created by growing finite islands on a monolayer substrate or by applying confinement potentials using patterned electrodes.

**Phosphorene and group-IV monochalcogenides**

Phosphorene, a monolayer of black phosphorus, is a monoelemental 2D material. Monolayer, few-layer, and bulk black phosphorus are all semiconducting materials, with a direct or nearly direct band gap (4). Additionally, phosphorene has a very high mobility that can reach 1000 cm$^2$/V⋅s for devices of ~10-nm thickness at room temperature (37). This exceeds the carrier mobility of TMDCs. According to theoretical predictions, the phonon-limited hole mobilities can reach 10,000 to 25,000 cm$^2$/V⋅s for the monolayer (zigzag direction) (38).

Both the optical and transport properties of phosphorene are highly anisotropic, as a consequence of this material’s orthorhombic, wavelike structure. Optical selection rules dictate that the absorption threshold is lower for linear polarized light along the armchair direction than along the perpendicular direction. Optical conductivity and Raman spectra are also anisotropic, providing a fast way to determine phosphorene’s lattice orientation. In addition to its optical and electronic properties, fundamental research in phosphorene has unraveled a growing number of physical phenomena, including superconductivity, high thermoelectric figure of merit (39), birrefringence, and colossal ultraviolet (UV) absorption.

The group-IV monochalcogenides SnS$_2$, GeS$_2$, SnSe, and GeSe are isoelectronic with phosphorene and share its orthorhombic structure, but the two atoms types break the inversion symmetry of the monolayer. As a consequence, they feature spin-orbit splitting (19 to 86 meV) (40) and piezoelectricity with large coupling between deformation and polarization change in plane (with piezoelectric coefficients $e_{31}$ ranging from $7 \times 10^{-10}$ to $23 \times 10^{-10}$ C/m, largely exceeding those of MoS$_2$ and hBN (41)).

SnS$_2$, SnSe, and GeSe are semiconductors, with gap energies covering part of the infrared and visible range for different numbers of layers (40). Even though the indirect band gap (in most cases) makes these materials less attractive for optical applications, the existence of two pairs of twofold degenerate valence and conduction band valleys, each placed on one principal axis of the Brillouin zone, makes them suitable for valleytronics applications. In this case, the symmetry is orthorhombic and, thus, the valley manipulation processes are different from those for TMDCs. Valley pairs can be selected using linear rather than circularly polarized light. Furthermore, there is no valley Hall effect, so the transverse valley current under an electric field is a second-order effect. Group-IV monochalcogenides are more stable against oxidation than phosphorene, can be grown by chemical vapor deposition (CVD), and have been recently exfoliated down to their bilayers.

**Gallium and indium monochalcogenides**

GaX and InX (where X is a chalcogen, like S, Se, or Te) are additional members of the family of hexagonal 2D materials. In this case, each layer can be viewed as a double layer of metal M = Ga, intercalated between two layers of chalcogen (X-M-M-X). The band structure of monolayers of such materials is rather unusual, having a “Mexican hat” dispersion at the top of the valence band, leading to a high DOS (42, 43) (bulk materials are most probably direct band-gap semiconductors). Thus, these materials have high and fast photosensitivity (44, 45) and large second-harmonic generation and have attracted attention mostly due to their optical properties. If the Fermi level is close to this singularity in p-doped materials, a ferromagnetic instability arises (46).

**Hexagonal boron nitride**

Layers of hBN consist of hexagonal rings of alternating B and N atoms, with strong covalent sp$^3$ bonds and a lattice constant nearly identical to that of graphite. hBN is very resistant both to mechanical manipulation and chemical interactions and also has a large band gap in the UV range. For these reasons, hBN is a material of choice as an encapsulating layer or substrate for 2D stacked devices, providing an atomically smooth surface free of dangling bonds and charge traps. hBN substrates leave the band structure of graphene near the Dirac point virtually unperturbed (if crystallographic orientations of the two crystals are misaligned) and dramatically improve the mobility of graphene devices (47, 48).
Oxide layers and other insulators

Many oxides have layered structures and can therefore be seen as a source for new 2D materials. These include lead oxide and lead salts [PbO, Pb₂O₃SO₄, Na₂PbO₂, etc.], phosphorus oxides and phosphates, molybdenum and vanadium oxides, and other transition metal oxides. In these materials, the layers are often connected by weak covalent bonds, oxygen bridges, or intercalating elements and are normally nonstoichiometric (due to the presence of oxygen vacancies). Further, layered oxides are normally polycrystalline, and mechanical exfoliation methods are usually limited to those available in higher-quality crystals. For the chemical means of production of such monolayers, intercalation with bulky guest species (such as tetrabutylammonium ions) has been used. Some of these layered oxides have been studied due to their importance as battery cathode materials (e.g., MoO₃, V₂O₅, and other Mo and V oxides), superconductors (e.g., copper and cobalt layered oxides) (49, 50), passivating layers (phosphorus oxide) (51, 52), and other areas of technological interest. Layered oxides allow for alloying, combination of different layers, and intercalation of ions and molecules; the possibilities of materials design are immense.

Among the most studied 2D insulators are hybrid perovskites, which are noteworthy for their high optical absorption coefficient within the solar spectrum and strong luminescence. Thin-film perovskite-based solar cells have emerged with a 20% power conversion efficiency, a notable value for a new technology (53). A hybrid perovskite is formed by layers of a metal halide intercalated with layers of organic chains. The high solar cell efficiency is thought to be greatly attributable to the confinement of excitons to the layers. Few-layer hybrid perovskites have been isolated by mechanical exfoliation and found to be stable in air in a time scale of minutes.

Novel van der Waals heterostructures

Two-dimensional crystals can be assembled into heterostructures (54), where the monolayers are held together by van der Waals forces. Considering that a large number of 2D crystals is currently available, it should be possible to create a substantial variety of heterostructures. However, the assembly technique currently in use (micromechanical stacking), allows only certain combinations of the interfaces. At the same time, an alternative technique, which potentially allows mass production of such structures (i.e., sequential growth of monolayers) comes with its own limitations and is presently in its infancy. Nevertheless, a large variety of novel experiments and prototypes have already been carried out with van der Waals heterostructures, which indicates that these materials are versatile and practical tools for future experiments and applications.

Assembly techniques

Currently, the most versatile technique for heterostructure assembly is direct mechanical assembly. This technique flourished starting in 2010 with Dean et al.’s work, which demonstrated the very high performance of graphene devices placed on an hBN substrate (47).

The technique used in the early works is based on preparing a flake of 2D crystal (Fig. 3, A to F) on a sacrificial membrane, aligning and placing it on another flake, and then removing the membrane. The process is then repeated to deposit further layers. Although the crystals are exposed to sacrificial membrane and solvents, which can contaminate the interface, annealing allows one to remove the contaminating and achieve very high interface quality (55), reaching high mobility (~10⁴ cm²/V-s) in graphene devices prepared this way.
A substantially cleaner method (dubbed the “pick-and-lift” method) is based on strong van der Waals interactions that exist between the crystals. When the membrane with a 2D crystal on it is brought into contact with another 2D crystal, it is not dissolved but rather is lifted up (Fig. 3, G to O); then there is a chance that the second crystal will stick to the first and will be lifted together with it. The process can be repeated several times. This technique results in clean interfaces over large areas and yet higher electron mobility (56). Further advances could be achieved by transferring the whole process into a glovebox with a controllable atmosphere.

1D contacts

The later method (Fig. 3, G to O) has one particular disadvantage: Having a completely assembled stack would prohibit one to make contacts to the inner layer. Luckily, it has been demonstrated that one can achieve various profiles of the edges of such a stack by reactive plasma etching. Thus, it is possible to etch the edge of the stack in such a way that the desired layer becomes exposed and can be contacted by metal evaporation (56) (Fig. 4A). The contact resistance for graphene can be as low as 35 ohm-μm.

Self-cleansing mechanism

The transmission electron microscopy (TEM) studies (55) demonstrate that interfaces can be atomically flat and free of any contamination [Fig. 4, B to D; adopted from (55)]. The reason for such behavior is the so-called “self-cleansing” mechanism (57). If the affinity between the two 2D crystals is larger than that of graphene, which leads to the formation of a moiré pattern (5).

It has been demonstrated that the most favorable configuration for graphene on hBN is when boron atoms lay on top of one of the sublattices in graphite and nitrogen is situated at the center of the hexagon (58). Then, by stretching itself to match the interatomic spacing of hBN, graphene tries to increase the area where the favorable configuration is achieved. Owing to the high Young modulus of graphene, such perfect stacking cannot be achieved across the whole interface (unless the hBN can contract, as the loss in elastic energy would not be compensated by the gain in the van der Waals interaction). Thus, such stretching of graphene can only be local, and the stretched regions would be separated by areas where the graphene lattice is not commensurate with hBN (Fig. 4, K and L).

This effect has been observed for graphene on hBN when the crystallographic orientations of the two crystals are practically aligned (10). In this case, the large regions of the moiré pattern where the two crystals are commensurate are separated by areas where the graphene lattice is relaxed. This effect disappears when the graphene is misoriented with respect to hBN. Such commensurate-incommensurate transition happens at a critical angle, which is given by the crystal mismatch (10).

Stacks of several other 2D crystals—including MoS₂ and MoSe₂ (59), MoS₂ and WS₂ (60), fluorographene and MoS₂ (61), and many others—have been investigated for electronic properties (62) and possible surface reconstruction. Thus, layer-breathing phonon modes have been observed by means of Raman spectroscopy for MoSe₂/MoS₂ heterobilayers (63). However, because the lattice-constant mismatch for those pairs is usually above 2%, the surface reconstruction would be hard to observe. It has been experimentally detected for silicene on MoS₂, where vertical buckling of silicene allows perfect stacking between the two crystals (64).

Spectrum reconstruction for graphene on hBN

Moiré patterns in graphene on hBN provide periodic scattering potential for electrons. This leads to the reconstruction of the electronic spectrum in graphene at the wave vectors determined by the...
periodicity of the moiré structure, as has been observed in scanning tunneling microscopy (5) and, later, in transport (6–8) and capacitance (9) measurements. Secondary Dirac points appear in the electronic spectrum, in both the valence and conduction bands (Fig. 4M; adapted from (85)). The energy range where the spectrum is reconstructed is given by the strength of the van der Waals interaction between graphene and hBN and is estimated to be on the order of 50 meV. Furthermore, the surface reconstruction leads to the strong asymmetry between the sublattices in graphene, which opens a gap in the graphene spectrum.

**Capacitively coupled van der Waals heterostructures**

Conceptually, the simplest devices based on van der Waals heterostructures are those for capacitance measurements. hBN is an ideal insulator that can sustain large electric fields (0.5 V per layer and above), allowing the preparation of capacitors with a very thin dielectric. The use of a thin dielectric ensures a large contribution of the quantum capacitance, which is directly proportional to the DOS in the electrode, making capacitance measurements a viable tool to study both single-particle and interaction phenomena in 2D materials. A number of systems have been investigated so far, including quantum capacitance in graphene (66), various sandwiches of graphene with TMDCs (57), and black phosphorus (67).

Capacitive coupling between two graphene layers through a thin layer of hBN can also lead to a number of noteworthy phenomena. This method allows for very-high-quality Coulomb drag devices, where two graphene layers, separated galvanically, interact through Coulomb forces between the charge carriers in the two layers (68). Because it is an atomically flat crystal with a very large gap in the electronic spectrum, hBN allows very thin barriers (on the order of a few nanometers) before any tunneling kicks in, bringing the two graphene layers closer than the characteristic distance between electrons in each of the layers (10 nm for a characteristic density of $10^{22}$ cm$^{-2}$). This opens the new regime of effective zero-layer separation in Coulomb drag experiments.

**Tunneling devices**

Graphene can be combined with semiconductor and insulating 2D crystals to create a tunnel junction (89). The use of hBN as a tunneling barrier is particularly attractive due to its large band gap (~6 eV), low number of impurity states within the barrier, and high breakdown field. Because the position of the Fermi energy and the DOS in graphene can be varied by external gate, the same applies for the tunneling current, which allows such structures to be used as field-effect tunneling transistors (FETTs) (70).

The architecture of FETTs enables tunneling spectroscopy to probe DOS in graphene, as well as to observe impurity- and phonon-assisted tunneling (71). Elastic tunneling through impurities gives peaks in $dI/dV_c$ ($d$-current, $V_c$ bias voltage); peak positions depend on both bias and gate voltages (Fig. 5C). On the other hand, inelastic phonon-assisted tunneling is characterized by a set of plateaus in $dI/dV_c$ independent of gate voltage (71) [more pronounced in $d^2I/dV_c^2$ (Fig. 5B)]. When the bias voltage is large enough to emit a phonon ($eV_F = h
\nu_{\text{ph}}$, where $e$ is the electron charge, $h$ is Planck’s constant $h$ divided by $2\pi$, and $\nu_{\text{ph}}$ is the frequency of the emitted phonon), an additional channel opens for electron tunneling, which increases transmission probability and, hence, tunnel conductance. Tunneling through impurities and with the phonon emission is especially visible if the crystallographic lattices of the two graphene electrodes are strongly misoriented with respect to each other, which prohibits direct electron tunneling because it is impossible to fulfill the momentum conservation requirements.

If the crystallographic lattices of the two graphene electrodes are aligned, momentum conservation for tunneling electrons can be achieved without impurity or phonon scattering. Rotational misalignment of the two graphene crystals corresponds to a relative rotation of the two graphene Brillouin zones in the reciprocal space. If the misalignment is small enough (~2°), then the momentum difference between the electronic states in the top and bottom graphene layers can be compensated electrostatically by applying bias and gate voltages (72), leading to the resonant tunneling and observation of the negative differential resistance (72) (Fig. 5D). A sharp negative differential resistance feature allows one to build a tunable radio-frequency oscillator with the potential to reach subterahertz frequencies.

The highest on-off ratio for FETTs can be achieved if the changes in the Fermi energy in graphene are comparable with the gap in the tunneling barrier—the situation achieved if hBN is replaced with WS$_2$ (on-off ratio of 10$^3$) (73) or MoS$_2$ (on-off ratio of 10$^5$ to 10$^6$, probably because of the presence of impurity bands) (70). In addition to logic applications, tunneling in van der Waals heterostructures was exploited for memory devices (74) with a floating gate, logic circuits (75), radio-frequency oscillators (72), and resonant tunneling diodes (76).

**Interaction with light**

Optoelectronic devices based on graphene (77) as well as other 2D materials (78) have been studied intensively. However, graphene photodetectors typically have low responsivity, which is a consequence of low adsorption cross-section. Such issues are eliminated when other 2D materials are used for such purposes. Thus, TMDCs (78), GaN (79), InSe (80), black phosphorus (81), and other materials (82) have been used as photodetectors (83) in photodiode or photoconductor regimes. The advantages of using such materials are the large DOS (which guarantees large optical adsorption), the materials’ flexibility, and the possibility of local gating, which allows one to control the spectral response in such devices.

**Van der Waals heterostructures for photovoltaic applications**

Still, even larger opportunities open up when such materials are combined. Combinations of graphene (as a channel material) and TMDCs (as light-sensitive material, where trapped charges are controlled by illumination) allow creation of simple and efficient phototransistors (85).

Combining materials with different work functions can lead to photoexcited electrons and holes accumulated in different layers, giving rise to indirect excitons (e.g., as has been observed for the pairs MoS$_2$/WS$_2$ (86) and MoS$_2$/WSe$_2$ (87) (Fig. 5, E and F)). Such excitons typically have long lifetimes, and their binding energy could be tuned by controlling the distance between the semiconductor layers.

If p- and n-doped materials are used in such devices, then atomically sharp p-n junctions can be created (88, 89). Such devices are extremely efficient in carrier separation, so they demonstrate very high quantum efficiency (for instance, GaTe/MoS$_2$ devices had external quantum efficiencies of >60% (88)). Furthermore, their performance can be tuned externally by gate voltage, as has been demonstrated for black phosphorus/MoS$_2$ heterostructures (90).

Even more efficient photovoltaic devices can be created by combining thin layers of TMDCs (91) or metal chalcogenides (92) with graphene. By sandwiching the photosensitive material between graphene electrodes, one can achieve very efficient photocarrier extraction from the device into graphene electrodes (which typically form good ohmic contacts with the TMDCs and serve as a transparent electrode as well). Because these structures are typically symmetric (Fig. 5G), one needs to create an electric field inside the TMDC to produce efficient carrier separation by bias, external gating (because the electric field is not fully screened by graphene due to its low DOS), or different doping of the two graphene layers.

**Light-emitting diodes**

The p-n junctions described above can be operated in the regime of electrical injection of the charge carriers, which leads to electron-hole recombination and light emission (89). However, such arrangement is limited by the requirements of synthesizing p- and n-type materials, which have not yet been demonstrated for all 2D crystals. Furthermore, the resistance of the junction is comparable to the resistances of the p and n electrodes, which makes it hard to control the current distribution.

A more straightforward arrangement is the carrier injection from highly conductive transparent electrodes directly into the 2D material in a vertical structure. Such a scheme, however, requires careful control of the dwell time of the injected electrons and holes in the semiconductor crystal, because photoemission is a slow process in comparison with the characteristic time required to penetrate the junction between graphene and the semiconductor. The dwell time can be controlled by introducing additional tunnel barriers (5, 51). Thus, two to three layers of hBN have...
been used (93) to increase the time electrons and holes spend inside the monolayer TMDC, allowing their radiative recombination. Devices based on WSe₂ are particularly efficient: Their quantum efficiency increases with increasing temperature and injection current, reaching 20% at room temperature (94). One can increase the quantum efficiency of such structures by placing several layers of TMDCs in series (93) (Fig. 5).

**Plasmonic devices**

Plasmons in graphene attract a lot of attention because it is possible to tune their frequency by changing the carrier concentration and, thus, the plasmonic frequency (95). Simultaneously, plasmonic and phonon-polaritonic properties have been studied in other 2D materials. For instance, hBN has polar dielectric properties, so it supports surface phonon polaritons with very low optical losses (96).

A number of new polaritonic effects can be seen in van der Waals heterostructures. Encapsulation of graphene with hBN allows one to eliminate the scattering of graphene plasmons with impurities, increasing their inverse damping ratio by a factor of 5 in comparison with bare graphene (97).

By sandwiching several graphene layers separated by hBN spacers, one can hybridize plasmonic modes in such multilayers, which can be further controlled with external gate voltage (98).

In such heterostructures, it is possible to enter a regime where the plasmon polaritons in graphene and the phonon polaritons in hBN coexist (Fig. 5, J to L; adapted from (99)). Strong coupling between the two leads to formation of the new collective modes: plasmon-phonon polaritons (100). Both the amplitude and the wavelength of the new collective modes can be controlled by gating graphene.

In aligned graphene/hBN heterostructures, the formation of the moiré pattern provides further modification of the graphene plasmonic spectrum. Zone folding results in the formation of the secondary Dirac points (5, 8, 85) (Fig. 4M), which allows a new type of vertical optical transition. Such transitions are immediately reflected in the modified damping factor, which exhibits a maximum at such Fermi energies (101). It has also

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**Fig. 5.** Electronic and optoelectronic applications of van der Waals heterostructures. (A to D) Tunneling in graphene/hBN/graphene tunnel transistors. (A) Schematic representation of a graphene tunneling device. Graphene electrodes are shown in dark purple, and the hBN tunneling barrier is light blue. The electrodes can be aligned with respect to each other. (B) $dI/dV$ map of phonon-assisted tunneling. Color scale: yellow to red corresponds to 0 to $3.8 \times 10^{-6}$ ohm$^{-1}$ V$^{-1}$. (C) $dI/dV$ map of resonant tunneling due to the presence of impurities in the hBN tunnel layer. Color scale: yellow to red corresponds to 0 to $7 \times 10^{-8}$ ohm$^{-1}$. (D) $dI/dV$ map of resonant tunneling with momentum conservation due to crystallographic alignment of two graphene electrodes. Blue coloring denotes the range of voltages where the negative differential conductivity is observed. Color scale: blue to white to red corresponds to $-6 \times 10^{-6}$ to 0 to $6 \times 10^{-8}$ ohm$^{-1}$. (E and F) Indirect excitons in a MoS₂/ WSe₂ heterostructure. Photoexcited electrons from WSe₂ are accumulated in MoS₂. Photoexcited holes from MoS₂ are accumulated in WSe₂. (G) TMDCs (large blue and white spheres) sandwiched between two graphene electrodes (small light-blue spheres) for photovoltaic applications. Photocarriers generated in TMDCs are separated by external gating (not shown). The structure can be encapsulated in hBN (purple and yellow spheres). (H and I) Vertical light-emitting diode heterostructures. hBN barriers increase the dwell time of the electron and hole in the TMDC, allowing their radiative recombination. Multiple quantum wells, formed by different materials, can be used in such structures (I). Coloring of the atoms is the same as in (F) and (G). (J to L) Polaritonic dispersions of graphene, hBN, and a graphene/hBN heterostructure. $q$, polariton momentum; $\omega$, polariton frequency; $r_p$, the imaginary part of the reflectivity.
Assembling van der Waals heterostructures in liquid and from liquid-phase-exfoliated 2D materials

A very powerful method of preparing graphene, which can also be extended to other materials, is liquid-phase exfoliation (102). Ink formulation based on such suspensions led to the development of graphene-based printed electronics (103). However, many applications would strongly benefit from properties beyond the capabilities of graphene inks. Thus, high thermal conductivity combined with dielectric property can be delivered by hBN, and optoelectronic capabilities can be delivered by inks of 2D semiconductors.

The ability to print combinations of such materials opens the door for low-cost fabrication of various devices (104). Planar (105) and vertical (106) photovoltaic devices based on TMDCs, as well as planar (107) and tunneling transistors (106) based on graphene and hBN, have recently been demonstrated.

By solution synthesis of 2D crystals or by controlling the charge on individual flakes in suspensions, heterostructures can be formed directly in the liquid phase (108) and can be used for energy applications. For instance, MoSe2/graphene structures have been used for Li-ion battery applications (109). Similar heterostructures have also been used for catalytic applications (110).

Growing van der Waals heterostructures

Direct growth methods such as CVD are promising techniques for scalable manufacturing of van der Waals heterostructures (111). Such techniques can be grouped as follows: (i) sequential CVD growth of 2D crystals on top of mechanically transferred or grown 2D materials, (ii) direct growth of TMDC heterostructures by vapor-solid reactions, and (iii) van der Waals epitaxial growth of 2D crystals on top of mechanically transferred or grown 2D mate-

terials, (ii) direct growth of TMDC heterostructures by vapor-solid reactions, and (iii) van der Waals epitaxy. State-of-the-art CVD, direct growth, and van der Waals epitaxy methods have already enabled the growth of many vertical heterostructures, such as graphene/hBN (112–116), MoS2/graphene (117–120), GaSe/graphene (121), MoS2/hBN (122, 123), WSe2/hBN (124), MoTe2/MoS2 (125), WS2/MoS2 (126), VSe2/GeSe2 (127), MoSe2/Bi2Se3 (128, 129), MoSe2/HfSe2 (129), MoS2/WSe2/graphene, and WSe2/MoS2/graphene (76).

In situ CVD growth of encapsulated graphene in a hBN/graphene/hBN heterostructure was an important achievement because it demonstrated the scalability of high-mobility graphene-based field-effect transistors (116). Also, some of the TMDC heterostructures can be grown directly in a single-step process: A WS2/MoS2 heterobilayer was grown on a SiO2/Si substrate at 850°C from precursors (W, S, MoO3) placed in the growth tube [Fig. 6A; adapted from (126)]. Because of the difference in the growth rates of MoS2 and WS2, the formation of a Mo2WxSi1–x alloy is suppressed. A clean interface enabled a band alignment of

been predicted that new plasmonic modes with carrier density dependence characteristic of parabolic electronic bands should appear in the vicinity of the van Hove singularities of the reconstructed spectrum (100) (Fig. 4M).

\[ H = \sum_k \epsilon_k | \phi_k \rangle \langle \phi_k | \]

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the two constituent layers, which led to the observation of indirect excitons in the WS2/MoS2 heterostructure.

Van der Waals epitaxy

Van der Waals epitaxy was introduced more than 30 years ago with the growth of a NbSe2 mono-layer on a cleaved face of MoS2 bulk crystal (330). This work also led to the successful growth of monolayer MoSe2 on SnS2 (331), as well as growth of a two-component heterostructure of monolayer NbSe2/trilayer MoSe2 on HBN (332). To grow graphene on HBN, Yang et al. used plasma to break down precursor methane molecules (49), after which growth occurred at 500°C over the course of 2 to 3 hours on HBN crystals mechanically exfoliated on a SiO2/Si substrate. Van der Waals interactions during epitaxial growth defined the preferential growth directions so that graphene crystals were aligned to the HBN substrate [Fig. 6, E to G; adapted from (48)].

Mechanically exfoliated HBN has also served as a substrate for CVD-based van der Waals epitaxy of a rotationally commensurate MoS2/HBN heterostructure (122). Another example of van der Waals epitaxy is the growth of high-quality wafer-scale MoSe2/BlSi2 heterostructures on the low-cost dielectric substrate AlN/Si in ultrahigh vacuum conditions (129).

Graphene is also a good substrate for van der Waals epitaxy: Grown WS2/graphene heterostructures show an atomically sharp interface and nearly perfectly crystalline orientation between graphene and WS2, despite a large (22%) lattice mismatch (333). Few-layer MoS2 and HBN structures were also grown using epitaxial graphene as a growth substrate (117, 134). Recently, monolayers of WSe2 and MoS2 were grown on free-standing CVD graphene (135). TMD crystal structures were also explored as substrates for epitaxy when a MoTe2 monolayer was grown on a bulk MoS2 substrate (125).

Finally, van der Waals epitaxy can be repeated several times to grow complex multicomponent heterostructures, such as atomically thin resonant tunneling diodes based on MoS2/WSe2/graphene and WSe2/MoS2/graphene [Fig. 6, H to I; adapted from (70)]. To this end, an epitaxial graphene trilayer was used as a substrate to grow monolayers of either MoS2 (at 750°C) or WSe2 (at 850°C) via powder vaporization or metal-organic CVD processes. Subsequently, a second TMD layer (WSe2 or MoS2) was grown on top of the initially grown heterostructure.

Lateral heterostructures

Lateral heterostructures can also be grown by a variety of methods. Thus, CVD-grown graphene was lithographically patterned and etched away, and HBN was grown via CVD, forming lateral 1D heterojunctions [Fig. 6B; adapted from (136)]. Beyond graphene and HBN, lateral heterostructures based on 2D TMDs can be disruptive for integrated optoelectronic devices. Although direct growth favors TMD alloys because of a similar chemistry and a small lattice mismatch between different TMDs (137), two-step epitaxial growth of a MoS2/WSe2 lateral heterostructure was recently demonstrated [Fig. 6, C and D; adapted from (338)]. To avoid alloying during growth, two separate temperature regimes were used (338). The atomically sharp WSe2/MoS2 heterojunction has a depletion width of >300 nm due to the potential difference between the MoS2 and WSe2 regions.

Lateral heterostructures of MoS2/WS2 and WSe2/MoSe2 were grown directly by controlling the growth temperature at ~650°C (126). Growth at relatively low temperatures was facilitated by either reducing tellurium into the CVD process (296) or using perylene-based growth promoters (339). The use of growth-promoting perylene-based aromatic molecules was recently extended to stitch together largely dissimilar 2D materials.

Conclusion

The family of 2D crystals is continuously growing, both in terms of variety and number of materials, and it looks like this process is only beginning. Almost every new 2D material possesses unusual physical properties. The 2D physics (e.g., KT transitions) in such materials is just starting to emerge. Still, we argue that the most interesting phenomena can be realized in van der Waals heterostructures, which now can be mechanically assembled or grown by a variety of techniques. Among the unsolved problems is the control of surface reconstruction, charge transfers, and built-in electric fields in such heterostructures. The standard band diagrams with quasi-electric fields are not a useful concept in 2D heterostructures; therefore, a new framework must be developed.

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