The successful discovery and isolation of graphene in 2004, and the subsequent synthesis of layered semiconductors and heterostructures beyond graphene have led to the exploding field of two-dimensional (2D) materials that explore their growth, new atomic-scale physics, and potential device applications. This review aims to provide an overview of theoretical, computational, and machine learning methods and tools at multiple length and time scales, and discuss how they can be utilized to assist/guide the design and synthesis of 2D materials beyond graphene. We focus on three methods at different length and time scales as follows: (i) nanoscale atomistic simulations including density functional theory (DFT) calculations and molecular dynamics simulations employing empirical and reactive interatomic potentials; (ii) mesoscale methods such as phase-field method; and (iii) macroscale continuum approaches by coupling thermal and chemical transport equations. We discuss how machine learning can be combined with computation and experiments to understand the correlations between structures and properties of 2D materials, and to guide the discovery of new 2D materials. We will also provide an outlook for the applications of computational approaches to 2D materials synthesis and growth in general.

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Thermodynamic stability. The thermodynamic stability of a 2D material requires its formation energy to be negative. The formation energies of various 2D materials have been efficiently calculated by DFT in combination with high-throughput screening platforms\(^{14}\). The success of DFT in predicting energies relies on the accuracy of well-tempered approximate exchange-correlation functionals. Standard functionals, e.g., the Perdew-Burke-Ernzerhof parametrization of the generalized gradient approximation exchange-correlation functional (GGA-PBE), accompanied by appropriate corrections\(^{15}\), describe well the formation and atomization energies. However, they are limited by intrinsic delocalization errors when bonds are stretched, resulting in underestimated reaction barriers\(^{16}\). A more realistic criterion for a stable 2D material is a low “above-hull” energy\(^{17}\). Exfoliation energies within 0.2 eV/atom are suggested as a general rule of thumb for the stability in 2D form. A slightly stricter criterion to have low surface energy (<20 meV/Å) is suggested to rule out potential high-surface-energy but multi-atom-thick sheets.

Kinetically stabilized 2D Materials. As-grown products may be metastable structures rather than ground-state polytypes\(^{18}\). The well-known phase-stability competition between the 2H and 1T\(^{\prime}\) types in transition metal dichalcogenides (TMDs) has been studied extensively using DFT. Several possible mechanisms for the 1T\(^{\prime}\)-phase stabilization in the disulfide, including growth-related factors such as the presence of point defects and residual strain, have been elucidated through DFT calculations\(^{19}\). Phase-stability analysis for the entire group IV metal dichalcogenide family\(^{20}\) revealed that the 1T\(^{\prime}\) phases are generally more stable for ditellurides. In addition to pure compounds, mixing/segmentation behavior and order–disorder transitions in alloys have also been discussed\(^{21}\). Within the W/Mo + S/Se/Te combinations, mixing either chalcogen or metals is thermodynamically favorable\(^{22}\), which can be experimentally quantified using the Warren-Cowley order parameters\(^{23}\). For TMDs, theory predicts that at finite temperatures, the formation of random alloys is favorable. However, kinetically stabilized atomically thin strips of alternating W and Mo in a sulfide alloy have been reported\(^{24}\). Interlayer/substrate interactions. Accurate description of vdW interactions becomes critical when interlayer sheet–sheet or sheet–substrate interactions are considered, e.g., for identifying the growth orientation. As a non-local effect, vdW forces are not correctly described by DFT with semi-local exchange-correlation functionals and are often corrected by adding pairwise interatomic terms from empirical fittings\(^{25}\), based on charge densities\(^{26}\), or by introducing non-local exchange-correlation functionals, i.e., van der Waals Density Functional\(^\ast\). Thorough testing of these methods has been examined in ref.\(^{28}\). One common technical problem in this computational approach is that substrates are modeled in a slab geometry where the one surface not in contact with the 2D sheet may host surface states, unless it is passivated. To suppress artificial charge transfer that these may induce, capping using pseudohydrogen\(^{29}\) with fractional charge is often performed. With the advent of new software platforms automating the generation of solid surfaces in combination with stacked 2D sheets and adsorption geometries (e.g., MPInterfaces\(^{30}\)), high-throughput screening of possible substrates may lead to a systematic approach to substrate engineering.

Precursor chemistry and kinetics. Transient intermediate states are commonly calculated using transition state theory and the nudged elastic band (NEB) method\(^{31}\) by calculating bond dissociation energies or corresponding activation barriers\(^{32}\). One recently attempted approach to capture precursor reaction kinetics is constrained molecular dynamics (MD)\(^{33}\), where slowly varying coordination constraints are enforced on reacting species to map out free energy barriers, as implemented in VASP\(^{34}\). It is different from NEB calculations in that (1) it calculates free energies at finite temperatures; (2) the final state of the reaction need not to be known; and (3) it has a better numerical stability\(^{35}\). Constrained MD was used to study a sulfur precursor, S\(_2\), reacting with a MoO\(_3\) surface\(^{36}\), concluding that MoO\(_3\) surface vacancies favor the sulfurization process both kinetically and thermodynamically. The downside of constrained MD is that a reaction coordinate is chosen a priori, which may bias the system towards unnatural products with incorrect reaction mechanisms\(^{33}\) and higher reaction barriers\(^{37}\).

Other methods for sampling of rare events such as nucleation of a new structure include umbrella sampling, transition interface sampling, and metadynamics. The umbrella sampling technique was introduced by Torrie and Valleau\(^{37}\), to improve the sampling of systems with energy landscapes containing high energy barriers. The weighted histogram analysis method\(^{38}\) can be used to analyze a series of umbrella sampling methods. In the transition interface sampling\(^{39}\), the transition region is divided into subregions of intermediate states. The rate constant of a reaction in this method is the multiplication of transition probabilities between different intermediate states. The metadynamics technique was introduced in 2002 and is usually used within an atomistic modeling framework\(^{40}\). Selected collective variables of the system not only evolve with time but also periodically leave behind positive Gaussian potentials that are added to the original

### Table 1. Examples of applications of DFT to 2D materials, the DFT-based first-principles methodology, and corresponding experimental observables.

| Aspect of growth | First-principles framework | Experimental observable and impact |
|------------------|----------------------------|-----------------------------------|
| Thermodynamic stability | Thermochemistry, high-throughput screening | Successful synthesis |
| Interlayer interaction | Dispersion forces, commensurate supercell construction | Orientation control |
| Precursor chemistry and kinetics | Transition state theory/NEB, thermochemistry, constrained molecular dynamics | Growth rate, residual gas analysis |
| Growth front advancement | Kinetic Monte Carlo, edge energetics | Morphology and growth rate, microscopy |
| Defects | Formation energies | Defect population statistics |

**ATOMISTIC COMPUTATIONAL METHODS**

First-principles calculations

This section discusses the computational approaches based on the density functional theory (DFT). The primary application of DFT in modeling 2D materials is to determine the relative thermodynamic stabilities of different crystal structures of a 2D material by computing their chemical potentials or to identify kinetic pathways by analyzing the energetics of potential transient structures from one stable equilibrium structure to another as thermodynamic conditions change. Examples of applications of DFT to understanding 2D materials and their growth, as well as the corresponding DFT-based first-principles methodologies and the corresponding experimental observables for validation are summarized in Table 1.
potential energy of the system, which effectively push the system out of a local minimum and into a neighboring energy well. Informally, metadynamics resembles “filling the free energy wells with computational sand.”

**Growth front advancement.** Early studies focused on calculating the surplus energies of edges in the high-symmetry directions and then applying the thermodynamic Wulff construction. One difficulty in calculating the edge energies is the polar and non-centrosymmetric nature of materials, where the usual ribbon calculation geometry makes the energies of unlike edges inseparable. The usual workaround is by constructing triangular flakes exposing a single type of edge. Another “energy density method” was introduced in ref. and a recently proposed method aimed at finding a general method suitable for high-throughput calculations introduces capping groups to passivate a surface.

One major criticism of further employing thermodynamic Wulff constructions is that growth is by definition out of equilibrium. Therefore, edges that dominate over others should not be the energetically favorable ones, but the slowest growing ones. The step-flow approach was formulated in ref. for graphene growth and was further developed in ref. for polar materials, establishing the use of kinetic Wulff construction. A similar approach involves DFT calculations and fitting to experimental grain morphologies, to construct a kinetic Monte Carlo (KMC) model.

**Defects.** The formation of defects may occur within thermodynamic equilibrium such as thermally generated point defects or follow from growth imperfections such as dislocations and grain boundaries (GBs), or reflect the finite size of crystals such as edges and surfaces. Defects can also be deliberately introduced using methods such as electron or ion irradiation and chemical treatment. The calculation of defect formation energies follows a well-established procedure detailed in ref. 48. 2D materials also host lattice-specific defect types, such as the Stone-Wales defect in graphene. Single-atom vacancies are another type of defects in 2D materials. A general strategy to ensure correct convergence behavior for charged defects in 2D systems is presented in ref. 49. Defect formation energies in MoS2 have been comprehensively studied in ref. 50; possible strategies of introducing extrinsic dopants have been examined in ref. 51. Defect complexes are also frequently studied to identify likely combinations between simple intrinsic defects and external contamination, devise possible defect-pairing strategies to neutralize harmful defects, and investigate their influence on the growth behavior of 2D materials on a different 2D sheet.5,53. Multiple vacancies may exist in 2D materials such as double vacancies in graphene, resulting in (i) two pentagons and one octagon—V2 (5-8-5) defect; or (ii) three pentagons and three heptagons—V2 (555-777) defect; or (iii) four pentagons, a hexagon, and four heptagons—V2 (555-6-777) defect. The formation of defects with an even number of vacancies in graphene is energetically favorable due to the lack of any dangling bond, whereas a large number of vacancies may bend and wrap the 2D material. Table 2 lists the typical values of formation and migration energies of these point defects from atomistic calculations.

Dislocations are line defects in 2D materials that can be of in-plane edge or out-of-plane screw type. The former consists of pentagon–heptagon (5–7) pairs and the latter makes the structure to become 3D. In general, dislocations in 2D materials have high gliding energy barriers and are immobile at room temperature. They hinder the plastic deformation of 2D materials. Dislocation structures are energetically favorable when the straight GB line filling the free energy wells out of equilibrium.

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**Empirical potentials.** Empirical potentials are faster and more computationally efficient, but less accurate than reactive potentials. Common empirical potentials for modeling 2D materials are reviewed below.

**Lennard-Jones potential:** In 1924, Lennard-Jones (LJ) published the 12–6 pairwise potential to describe the vdW interaction between two neutral particles (i.e., rare gases)

$V_{\text{LJ}} = 4\epsilon \left( \frac{1}{r^{12}} - \frac{1}{r^6} \right)$

where $r$ is the distance between two interacting atoms, $\epsilon$ is the potential well depth and $\sigma$ is the distance at which the potential is zero. The $r^{-12}$ term captures the strong repulsion occurring when two atoms get closer. The repulsive effect originates from the Pauli exclusion principle, which penalizes the overlap of electron orbitals. The $r^{-6}$ term describes the vdW interaction, which is due to the coupling between the instantaneous polar charges induced in two neutral molecules and is significantly weaker than the electron overlap effects.

The popularity of the LJ potential is due to its simplicity and it has been widely applied to simulate the interlayer interactions in atomically layered materials. While being used to capture vdW interactions, a limitation of the pairwise interaction is the inability to capture frictional processes, such as the sliding of atomic layers.

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**Table 2.** Typical values of formation energies and migration energy barriers of point defects in 2D materials from atomistic calculations.

| Type of 2D materials | Type of point defects | Formation energy | Migration energy barriers |
|----------------------|-----------------------|------------------|--------------------------|
| Graphene             | Stone-Wales          | $\approx 5 \text{ eV}$ | $\approx 1.5 \text{ eV}$ |
| SV$^{61}$            | $\approx 7.5 \text{ eV}$ | $\approx 7 \text{ eV}$ |
| V$_2$ (5-8-5)$^{56}$ | $\approx 8 \text{ eV}$ | $\approx 7 \text{ eV}$ |
| V$_2$ (555-777)$^{56}$ | $\approx 7 \text{ eV}$ | $\approx 7 \text{ eV}$ |
| V$_2$ (5555-6-777)$^{56}$ | $7 - 8 \text{ eV}$ | $\approx 7 \text{ eV}$ |
| Phosphorene          | SV$^{95}$            | $\approx 1.65 \text{ eV}$ | $\approx 0.4 \text{ eV}$ |
| MoS$_2$              | V$_5^{59,60}$         | $1.22 - 2.25 \text{ eV}$ | $\approx 2.27 \text{ eV}$ |
Different corrections have been proposed to overcome this shortcoming. The two potential parameters and can be determined by using one energy quantity like the binding energy and a structural quantity like the density. The LJ parameters for most elements can be determined systematically, as demonstrated by Rappe et al. The LJ parameters for typical 2D materials and substrates in Table 3 are listed in Table 4 for various materials via analytic expressions, including many 2D materials and substrates.

The FF potential includes both pair (two-body) and three-body interactions, where V2 describes the pairwise bond-stretching energy and V3 captures the bending energy associated with an angle with initial value θ0.

\[ V_{ij} = A \exp\left(\frac{r_{ij} - r_{ij}^0}{\sigma}\right) \]

where r_{ij} is the distance between atoms i and j, and r_{ij}^0 is the equilibrium distance. The FF potential is additive, and the energies in the FF model are additive.

For the bond-stretching energy, Vb is the bond-angle-bending energy, Vc is the torsional energy for a bond, and Ve is the inversion energy among four atoms. The Keating model is the simplest model including only the harmonic terms. Parameters of the potential are usually determined by fitting to experimental data, and the volume and accuracy of the fitted data set determine the accuracy of the FF model.

The FF potential has two significant features: (i) the number of interaction terms and (ii) the force-constant parameters and the equilibrium structural parameters (bond lengths and angles). MM3 and COMPASS use all of the interaction terms and can simulate large deformations and nonlinear phenomena. In contrast, the Keating model is the simplest model including only the harmonic terms. Parameters of the potential are usually fitted to available data and, thus, the volume and accuracy of the fitted data set determine the accuracy of the FF model. Several generations of FF models have been developed as more data become available, including four versions of the molecular mechanics FF: MM3, MM2, MM3, and MM4. The UFF provides the FF parameters for all the elements in the periodic table.

As the energies in the FF model are additive, this potential has been widely used to calculate physical or mechanical properties for various materials via analytic expressions, including many 2D materials. For example, an FF model, with bond stretching and angle-bending terms, has been used to derive the phonon dispersion of graphene layers. The bond-stretching and angle-bending terms were utilized to extract the bending properties of graphene. A mapping between the FF model and the elastic beam model has been developed, which is used to derive analytic expressions for Young’s modulus and Poisson’s ratio of graphene and carbon nanotubes.
Reactive interatomic potentials. The reactive-potential approach has been widely used to explore the synthesis and properties of different materials. The reactive potentials can capture the bond breaking and formation during the classical simulations of 2D materials. They are also computationally less expensive than first-principles methods such as DFT calculations and ab-initio MD. Typical reactive potentials that have been frequently considered are Tersoff Bond Order (Tersoff) \(^{89}\), Adaptive Interatomic Reactive Empirical Bond Order (REBO) \(^{90}\), Adaptive Intermolecular Reactive Empirical Bond Order (AIReBO) \(^{91}\), Charge Optimized Many-Body (COMB) \(^{92}\), and Reactive Force Field (ReaxFF) \(^{93}\). These different reactive potentials are reviewed in detail in ref. \(^{94}\). Performances of several reactive and non-reactive potentials in graphene-based materials are compared in ref. \(^{95}\). These potentials allow the realization of the thermodynamic stability of graphene-based materials under different conditions, environmental chemistry, substrate, interlayer interactions, and structural defects. However, only a few of them have been employed in the study of computational synthesis of other 2D materials to reveal thermodynamic stability, simulate the process of the growth of layers, model the top-down synthesis of 2D materials such as exfoliation techniques \(^{38}\), and evaluate the properties of 2D nanostructures. This is due to several challenges including the following: (1) the long time required to develop a reactive potential, (2) the non-transferability of potential parameters from one material to another, and (3) the lack of universal functional forms for the reactive potentials.

The Tersoff potentials are mainly used to explore various structures and properties of 2D hexagonal boron nitride (h-BN) materials \(^{96}\). The REBO and AIReBO potentials are used to simulate the CVD process and growth of 2D amorphous carbon on substrates, mechanical properties of MoS\(_2\), and even study the thermal stability of C\(_{60}\) 2D nanostructures \(^{97}\). The COMB3 potentials have been used to simulate the experimental CVD deposition and growth of graphene and metal on substrates \(^{98}\). The ReaxFF potentials have been used to explore the structures of several 2D materials under intercalation, study the defects and groups on the surfaces of MXene materials, simulate CVD growth of MoS\(_2\) layers, and explore the synthesis of 2D polymeric materials in experiments \(^{99}\). In general, COMB3 and ReaxFF can further explore 2D materials, as their functional forms can describe heterogeneous systems. It remains crucial to find accelerated methods to develop reactive potentials with high accuracy and transferability for 2D materials.

Tersoff potential: The Tersoff potential was first published in 1986 \(^{100}\) and further developed in 1988 \(^{101}\). The second version of the Tersoff potential has the following form,

\[
E = \sum_i E_i = \frac{1}{2} \sum_{i \neq j} V_{ij},
\]

(9)

\[
V_{ij} = f_c(r_{ij})[a_{ij}f_b(r_{ij}) + b_{ij}f_a(r_{ij})];
\]

\[
f_c(r) = \begin{cases} 
1 & r < R - D \\
\frac{1}{2} - \frac{1}{2} \sin \left[ \pi (r - R)/D \right] & R - D < r < R + D \\
0 & r > R + D
\end{cases}
\]

(11)

A distinct feature of the Tersoff potential is the absence of an explicit three-body interaction term. Instead, the concept of bond order is introduced, in which the strength of a given bond depends strongly on its local environment. Bond order is defined through a set of bond, over-coordination penalty, under-coordination stability, lone pair, valence, torsion, non-bonded vDW interactions, and Coulomb energies, respectively. The bond-order parameter determines the bonded interactions between the

\[
E_{\text{system}} = E_{\text{bond}} + E_{\text{over}} + E_{\text{under}} + E_p + E_{\text{val}} + E_{\text{tor}} + E_{\text{vdW}} + E_{\text{Coulomb}}.
\]

(14)

where \(f_b(r_{ij})\) and \(f_a(r_{ij})\) are the repulsive and attractive components, respectively. The cutoff function is:

\[
f_c(r) = \begin{cases} 
1 & r < R - D \\
\frac{1}{2} - \frac{1}{2} \sin \left[ \pi (r - R)/D \right] & R - D < r < R + D \\
0 & r > R + D
\end{cases}
\]

(11)

\[
g(\theta) = 1 + \frac{c^2}{d^2} - \frac{c^2}{d^2 + (h - \cos \theta)^2}.
\]

(12)

\[
a_0 = (1 + \alpha n_0)^{1/2n};
\]

(13)

\[
\eta_0 = \sum_{\text{configurations}} f_c(r_{ij})a_0^{1/(n-\tau)}
\]

The 13 unknown parameters are typically determined by fitting to data from first-principles calculations or experiments.

The Tersoff potential was used for modeling of graphene, which gives a better description of the phonon spectrum \(^{102}\). It was also used to model germanene \(^{103}\) and silicene \(^{103}\). By introducing one scaling parameter for the quantity \(b_{ij}\) with atoms \(i\) and \(j\) of different elements \(^{104}\), the Tersoff potential was generalized to describe the interaction for multicomponent systems including C-Si and Si-Ge \(^{104}\), and boron nitride \(^{105}\). Using the Tersoff potential, the thermal conductivity of silicene was calculated to be 61.7 and 68.5 W mK\(^{-1}\) for the armchair and zigzag configurations, which is reduced when it is doped with heavier silicon isotopes \(^{106}\). This potential has also been used for modeling the misfit dislocations in 2D materials, where a critical thickness was found for the formation of an interface misfit dislocation \(^{107}\).

ReaxFF potential: The ReaxFF \(^{93}\) is a bond-order-based force field, which allows formation and breaking of bonds. It has a general form of

\[
E_{\text{system}} = E_{\text{bond}} + E_{\text{over}} + E_{\text{under}} + E_p + E_{\text{val}} + E_{\text{tor}} + E_{\text{vdW}} + E_{\text{Coulomb}}.
\]

(14)

where the energy terms on the right-hand side in the order of appearance are bond, over-coordination penalty, under-coordination stability, lone pair, valence, torsion, non-bonded vDW interactions, and Coulomb energies, respectively. The bond-order parameter determines the bonded interactions between the
where $BO_\sigma$, $BO_\pi$, and $BO_{\pi\pi}$ are the partial contributions of associated $\sigma$, $\pi$, and $\pi\pi$ bonds between the $i$ and $j$ atoms; $r_{ij}$ is the distance between the $i$ and $j$ atoms; $r_\sigma$, $r_\pi$, and $r_{\pi\pi}$ are the associated bond radii; and $p_{bo}$ are empirical constants that will be determined from ab-initio simulations or experiments. The energy terms in Eq. (14) are then formulated as a function of the bond-order parameter (see ref. 91 for details).

ReaxFF studies on 2D materials were initiated from studies of various aspects of graphene\textsuperscript{109} and inspired investigations for a number of other 2D materials, which are elaborated below.

Graphene—Effect of the environment’s chemistry on properties and responses of graphene has been investigated using ReaxFF. For example, effect of oxygen on the mechanical response of graphene\textsuperscript{109}, effect of the addition of fluorine and hydrogen\textsuperscript{110} on the structural quality of graphene, as well as the Ni-catalyzed growth process of single-layer graphene\textsuperscript{111}, are investigated. The growth mechanism involves atomic deposition and segregation of C atoms on the Ni substrate, forming dome-like carbon nanotube cap and polygonal ring patterns via atoms→chains→graphene-like layer formation (Fig. 1a). The ReaxFF has also been applied to study reactive events at the graphene–water interface (Fig. 1b)\textsuperscript{112} and selective desalination via the bare and hydrogenated graphene nanopore\textsuperscript{113}.

Silicene and phosphorene—Silicene and black phosphorene (BP) are the silicon and phosphorus analogs of graphene, respectively. A favorable route for silicene formation has been proposed by employing the graphene bilayer as a template (Fig. 1c)\textsuperscript{114}. It was found that vacancy defects reduce the thermal stability of silicene, where the critical temperature reduces by more than 30%\textsuperscript{115}. A P/H parameter set has been developed for monolayer BP\textsuperscript{114} to examine the mechanical, thermal, and chemical stability of both pristine and defective BP.

Molybdenum disulfide—The ReaxFF potential for 2D MoS\textsubscript{2} systems was introduced in ref. 6. This potential could precisely calculate the formation energy of ripplocations in MoS\textsubscript{2} (Fig. 1d)\textsuperscript{116}, which result from the slippage of the upper layer against the lower layer to relax the strain without breaking covalent bonds. This potential was also trained over the vacancy formation energies, diffusion barriers, bending rigidity, and kinetics. It has...
also been applied to design S vacancy defects on MoS$_2$ surface\textsuperscript{117}. Reactive MD simulations have also been used to model the synthesis of MoS$_2$ layers via sulfidation of MoO$_2$ surfaces\textsuperscript{7}. These simulations identified a three-step reaction pathway as follows: (i) evolution of O$_2$ and reduction of MoO$_2$ surface; (ii) S$_2$-assisted reduction and formation of SO/3SO$_2$; and (iii) sulfidation of the surface via Mo-S bond formation. The ReaxFF MD simulations have also been employed within a multiscale framework to capture thermal properties of MoS$_2$\textsuperscript{118}. The predicted thermal conductivity of MoS$_2$ structures with infinite length (37 ± 3 W/mK) falls within the range of experimental results (34.5–110 W/mK)\textsuperscript{19}.

MXenes—MD simulations with ReaxFF of MXenes revealed that intercalating with potassium cations drastically improves water stability and homogeneity of MXenes, and reduce the self-diffusion coefficient of water by two orders of magnitude\textsuperscript{120}. The dynamical response of MXene with different surface terminations to intercalating ions was studied with DFT calculations and ReaxFF MD simulations\textsuperscript{121}. Figure 1e compares the interlayer distance of Ti$_3$C$_2$(OH)$_2$ MXene bilayer with and without K$^+$ ions. Influence of metal ions intercalation on vibrational properties of water molecules trapped between MXene layers was studied using ReaxFF MD simulations\textsuperscript{122}. Widening of the interlayer gap may enable the penetration of molecular reactants such as urea, which decomposes readily and leads to the intercalation of ammonium cations\textsuperscript{123}. ReaxFF simulations were performed for defect formation and homoeipitaxial growth of TIC layer on Ti3C2 MXene structures\textsuperscript{124}.

**REBO/AIREBO potentials**: The REBO potential ($E^{\text{REBO}}_{ij}$) is a combination of attractive ($V^A_{ij}$) and repulsive ($V^R_{ij}$) interactions with certain ratio ($b_{ij}$), $E^{\text{REBO}}_{ij} = V^A_{ij} + b_{ij} V^R_{ij}$. The bond-order interaction ratio, $b_{ij}$, is dependent on the local coordination in the atomic environment. The REBO potential can be used to study the growth of 2D amorphous carbon, mechanical properties of MoS$_2$, and even thermal stabilities of C$_60$ 2D nanostructures using atomic coordination, internal stress, and mass density analysis\textsuperscript{126}. The self-potential phenomenon during CVD, i.e., penetration of the substrate bulk carbon by carbon atoms, can be explained by this REBO potential. REBO potential is also suitable for modeling and analysis of TMD materials such as MoS$_2$. The predicted thermal conductivity of MoS$_2$ structures with interlayer gap may enable the penetration of molecular reactants such as urea, which decomposes readily and leads to the intercalation of ammonium cations\textsuperscript{123}. ReaxFF simulations were performed for defect formation and homoeipitaxial growth of TIC layer on Ti3C2 MXene structures\textsuperscript{124}.

The AIREBO potential is the modification over the REBO potential, which also considers the torsional ($E^{\text{TORSION}}_{kijl}$) and LJ ($E^{\text{LJ}}_{ij}$) interactions. The integration of pairwise interactions in the AIREBO potential can be represented by the following equation

$$E = \frac{1}{2} \sum_{i < j} \sum_{k} \left[ E^{\text{REBO}}_{ij} + E^{\text{LJ}}_{ij} + \sum_{k=1}^{\text{NN}} E^{\text{TORSION}}_{kijl} \right].$$  \hspace{1cm} (23)

The AIREBO potential can describe the breaking, formation, and hybridization of covalent bonds. The REBO and AIREBO potentials have been used to study the growth of 2D amorphous carbon, mechanical properties of MoS$_2$, and even thermal stabilities of C$_60$ 2D nanostructures using atomic coordination, internal stress, and mass density analysis\textsuperscript{126}. The self-potential phenomenon during CVD, i.e., penetration of the substrate bulk carbon by carbon atoms, can be explained by this REBO potential. REBO potential is also suitable for modeling and analysis of TMD materials such as MoS$_2$. The predicted thermal conductivity of MoS$_2$ structures with interlayer gap may enable the penetration of molecular reactants such as urea, which decomposes readily and leads to the intercalation of ammonium cations\textsuperscript{123}. ReaxFF simulations were performed for defect formation and homoeipitaxial growth of TIC layer on Ti3C2 MXene structures\textsuperscript{124}.

The total potential energy consists of electrostatic energy ($U_{\text{el}}[q, r]$), short-range interactions ($U_{\text{short}}[q, r]$), long-range vdW interaction ($U_{\text{vdW}}[q, r]$), and the correction terms. Here, $q$ and $r$ represent the charge and coordinate array of the elements. The electrostatic energies ($U_{\text{el}}[q, r]$) can be represented as

$$U_{\text{el}}[q, r] = U_{\text{off}}[q, r] + U_{\text{short}}[q, r] + U_{\text{vdW}}[q, r] + U_{\text{corr}}[q, r].$$  \hspace{1cm} (24)

The self-interaction energy ($U_{\text{off}}[q, r]$) consists of field-effect and ionization energy, or the affinity energy. Charge-to-charge interaction ($U_{\text{short}}[q, q])$ is associated with the charge density distribution functions. Nuclear-charge interaction ($U_{\text{vdW}}[q, r]$) depends on the coulombic interaction between charge density distributions in the system. Polar interaction energy ($U_{\text{corr}}[q, r]$) depends on the charge interactions and dipole distributions in response to the external electric field. The charge-dependent short-range interaction in Eq. (24) is associated with bond distance functions of the atom and the associated charge functions. This energy term can be expressed by

$$U_{\text{short}}[q, r] = \sum_{i < j} V_{b_{ij}} \sum_{k} \left[ F_i(r) |q_i| |q_j| - b_{ij} F_i(r) |q_i| |q_j| \right].$$  \hspace{1cm} (25)

Here, $V_{b_{ij}}$ is the bond energy that is associated with pairwise attraction ($V^A_{ij}(q_i, q_j, q_k)$) and pairwise repulsion energy ($V^R_{ij}(q_i, q_j, q_k)$). They exponentially decrease with the interatomic distance $r_{ij}$. $F_i(r)$ is the cutoff function.

The long-range interaction energy is defined by the vdW interaction, which can be expressed by the LJ interaction formula\textsuperscript{130}

$$U_{\text{vdW}}[q, r] = \frac{1}{2} \sum_{i < j} \sum_{k} \left[ \left( \frac{\sigma_{vdW}}{r_{ij}} \right)^{12} - \left( \frac{\sigma_{vdW}}{r_{ij}} \right)^{6} \right].$$  \hspace{1cm} (27)
Here, $c^{GW}$ and $c^{DW}$ are the equilibrium distance and interaction energy, respectively; $r_i$ is the cutoff radii where the interaction assumes zero.

The correction factor in Eq. (24) adjusts the energy penalties associated with specified angles. It can be calculated by a sixth-order Legendre polynomial and bond bending term.

$$U'(\{q\}, \{r\}) = \sum_{i=1}^{N} \sum_{j=1}^{N} \sum_{k=1}^{N} K_{ij}^{\theta} \left( \cos(\theta_{ijk}) + K_{ij}^{\theta} \left( \cos(\theta_{ijk}) - \cos(\theta_{ijk}^{\theta}) \right) \right)^2$$

Here, $K_{ij}^{\theta}$ indicates the specific bond angles and $\theta_{ijk}$ is the bond angle.

The COMB3 potential has been introduced to study the CVD process for various 2D materials. The charge-optimized COMB3 potentials have been used to simulate the experimental CVD processes for various 2D materials. The charge-optimized COMB3 potential has been used to simulate the experimental CVD process for various 2D materials. The charge-optimized COMB3 potential has also been used to investigate the effect of surface hydroxylation of amorphous SiO$_2$ substrate on heat conduction of supported graphene.

**MESOSCALE SIMULATIONS**

Phase-field model

At mesoscale, a simple and effective model for 2D materials growth is the classical Burton-Cabrera-Frank (BCF) theory, where the island growth is realized by the deposition of adatoms from the supersaturated gas atmosphere and the incorporation of diffusing atoms at the step edge. The direct numerical implementation of the BCF model requires tracking the evolving step edges and applying the boundary conditions due to the sharp-interface nature of the model; in contrast, phase-field diffuse-interface models avoid the explicit tracking of the interfaces.

The BCF-based phase-field models have been developed since the 1990s for a series of problems related to crystal growth, including collective step motions in a 1D step train, spiral surface growth during thin-film epitaxy, and step flow under different kinetic regimes. More recently, the combined effects of edge diffusion, the Ehrlich-Schoebel barrier, deposition, and desorption for epitaxial growth have been investigated in 2D.

Based on these studies, Meca et al. applied the model to the growth of 2D materials with anisotropic diffusion on substrates, using the epitaxial graphene growth on copper foil during CVD as an example, with experimental validations. The typical BCF-based phase-field model for 2D materials growth uses a smoothly varying order parameter $\phi$ to distinguish the phases, e.g., $\phi = 1$ in the 2D island, $\phi = -1$ in the substrate, and $-1 < \phi < 1$ at step edges. The free energy functional $\mathcal{F}$ of the system has the general form of:

$$\mathcal{F} = \int_V \left( f(\phi) + \lambda u g(\phi) + \frac{1}{2} W(\theta)^2 |\nabla \phi|^2 \right) dV$$

where the first term $f(\phi)$ is a double-well (or multi-well, e.g., see refs. 136, 141) function with global minima at $\phi = 1$ and $\phi = -1$; the second term describes the coupling with the reduced saturation field $u$, with $g(\phi)$ an interpolation function and $\lambda$ the coupling coefficient; the third term is the gradient energy due to the inhomogeneous distribution of $\phi$ at interfaces, where $W(\theta)$ is the angle-dependent interface thickness with $\theta = \arctan(\phi_\parallel/\phi_\perp)$. $u$ can be directly related to the adatom concentration $c$ through $u = \Omega (c - c_0)$. $c_0$ is the equilibrium concentration for a flat interface and $\Omega$ is the atomic area of the solid phase. The evolution of $\phi$ is governed by

$$\frac{\partial u}{\partial t} = D \nabla^2 u + F_d - \frac{u}{\tau_s} - \frac{1}{2} \frac{\partial \phi}{\partial t}$$

where $D$ is the diffusion coefficient, $F_d$ is the effective deposition rate, and $\tau_s$ is the characteristic time scale for atom desorption. The phase-field equations (Eqs. (30) and (31)) can recover the sharp-interface BCF model in the limit of $W(\theta) \to 0$.

The applications of the phase-field models in 2D materials growth can be categorized into the following three aspects. (1) Reproducing and explaining experimentally observed growth morphologies. Examples include the reconstruction of the spiral growth of SnSe$_2$; the verification of impurity-induced bilayered graphene growth; and the explanation to island shape change from quasi-hexagons to triangles and dendrite-like morphologies. (2) Investigating the effect of experimental parameters on growth morphologies. Taking the CVD growth of graphene as an example, the effects of flux of carbon species, deposition rates, and equilibrium saturation on graphene island morphologies have been investigated using phase-field simulations, which could provide useful guidance to the experimental control of the growth qualities. (3) Further development of the model to include the effect of additional physical/chemical processes during growth. In addition to the deposition, edge diffusion, and edge anisotropies, phase-field simulations have been extended to model multi-island interactions, tilt GB topology of 2D materials on substrates with topological cones, and to include the chemical reaction kinetics in the gas phase using a microkinetic model. These works improve the fidelity and applicable range of the phase-field model for 2D materials growth.

Nevertheless, a primary obstacle to applying phase-field models for computational synthesis of 2D materials is the lack of connection between the model parameters and the experimentally controllable CVD processing parameters, which is largely due to the difference in size scales of interest. Most of the existing phase-field simulations only focus on the growth of a few 2D islands (up to micrometer scale), which can hardly represent the effect of the macroscopic CVD parameters. One possible solution is to enable the “multiscale” modeling approach, integrating the FEM (see the section on “Macroscopic models”) to account for the effect of macroscopic CVD parameters on transport phenomena and the mesoscale phase-field model for 2D crystal growth within selected zones of the whole substrate. In particular, the steady-state velocity, concentration, and temperature profiles from FEM can be incorporated in the phase-field model, as it has been proposed and applied by Momeni et al. (see Fig. 2). The incorporation of chemical reaction kinetics in the phase-field model can improve the accuracy of the predictions and expand its applicability. Phase-field models have also been applied to phase changes in 2D materials. Notable examples include the multi-domain microstructure during H-T’ structure transformation in MoTe$_2$ and the domain switching behaviors under external stimuli. A fast Fourier transform algorithm has been implemented to study the domain switching during bending of MoTe$_2$ (see Fig. 3).
Phase-field-crystal model

Different from the continuous mesoscale phase-field approach, the phase-field-crystal (PFC) approach\(^{150}\) describes the thermodynamics and dynamics of phase transformations through an atomically varying order parameter related to the atomic density field. With the application of the classical dynamic DFT, PFC can capture the atomistic-scale morphology and evolution dynamics, within diffusional time scales, which is usually challenging for conventional atomistic methods. Recently, PFC has been actively applied to investigate the defect formation and the atomic structure of interfaces, GBs, and triple junctions during the growth of 2D materials\(^{151}\). The unique GB structure and collective domain dynamics in binary 2D materials, e.g., h-BN, was investigated (see Fig. 4\(^{152}\)).

Compared with the continuous phase-field approach, the advantage of PFC lies in resolving atomistic-scale interface structures and growth dynamics of 2D islands; however, due to the limitation of simulation size, interface dynamics under the effect of realistic CVD parameters are difficult to realize. Future directions of PFC simulations for 2D materials growth include the consideration of atomistic-scale island-substrate interactions and the integration with the larger-scale phase-field or FEM simulations.

Kinetic Monte Carlo

KMC is another mesoscale technique for predicting the growth morphology and kinetic mechanisms of 2D materials, where all possible kinetic events are listed in an event catalog, and their stochastic sequence are randomly selected based on their activation energies\(^{153}\). Activation energies of the related kinetic processes are usually obtained from atomistic-scale calculations. KMC identifies the governing kinetic pathways and simulates the kinetic process. Details of its implementation can be found in ref. \(^{154}\).

KMC has been extensively applied to simulate the growth of graphene, e.g., the evolution of vacancy complexes and formation of vacancy chains\(^{155}\), the formation and kinetic effect of multi-member carbon ring complexes\(^{156}\), etching of graphene GBs due to oxygen migration and reaction\(^{157}\), and GB evolution following the Stone-Wales mechanism\(^{158}\). Growth of nanoscale graphene islands

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Fig. 2 A coupled macroscale/phase-field simulation and experimental comparison for MoS\(_2\). a Initial precursor concentration distribution from FEM. b Phase-field simulation results for MoS\(_2\) morphology on substrate at \(t = 0.25\) h, the enlarged images of i–iii boxes are shown in d–f. c Experimental results for deposition of MoS\(_2\) after deposition for \(t = 0.25\) h, the enlarged images of 1–3 boxes are shown in g, h. Reprinted from ref. 8 under Creative Commons Attribution 4.0 International License. The scale bar in a, b shows 400 \(\mu\)m and the one in d–f shows 400 \(\mu\)m.
**Fig. 3** Variant selection and transformation morphologies in MoTe$_2$ under an applied in-plane strain $\varepsilon_{ii}$. 

- **a** Phase diagram in $\{\varepsilon_{xx}, \varepsilon_{yy}\}$ space. Colored (hatched) regions correspond to one (two) phase regions. Solid lines (analytic) and filled data points (simulations) indicate the strain at which H and a given T’ variant have equal energies. Dashed lines and open symbols similarly represent two-phase coexistence bounds. ($\blacksquare$, $\triangle$) and ($\bullet$, ○) correspond to simulations with and without bending, respectively, and blue circles (blue solid circle and blue open circle) represent DFT results at 0.

- **b–d** Images of the microstructures with increasing time/strain from left to right for strain paths b–d in phase diagram a. Strain values $\{\varepsilon_{xx}, \varepsilon_{yy}\}$ are given below each panel. 

- **e** Corresponding maximum principal stress (GPa) maps for microstructures in d. All simulations are for 50 $\mu$m $\times$ 50 $\mu$m simulations. Adapted with permission from ref. 149. Copyright (2017) American Chemical Society.

**Fig. 4** PFC simulations of the grain growth and coalescence in h-BN. 

- **a–f** Grain coalescence and inversion domain dynamics.

- **c** Atomic site number $N$ vs. $t$, showing two regimes $N = -0.065t + 3.045 \times 10^4$ and $N = -0.085t + 3.612 \times 10^4$ via fitting.

- **d, e** Domain shrinking in the white boxed region of b. f Transient of three merging heart-shaped defects before their annihilation.

- **g–i** Time evolution of collective atomic displacements in the yellow boxed corner of d. Adapted figure with permission from ref. 152. Copyright (2017) by the American Physical Society.
from carbon monomer nuclei or pre-existing growth fronts has been captured, showing catalytic growth behaviors, anisotropic morphological patterns, temperature- and deposition-flux-rate-dependent size and shapes, inhomogeneous and nonlinear growth kinetics due to lattice mismatch, and geometry-determined growth mechanisms. The "coastline" graphene morphology during sublimation and the step-flow growth of epitaxial graphene have also been reported. Based on KMC simulations for graphene growth on Cu (111) with and without hydrogen, the growth protocol was designed for bilayer growth and N-doped graphene growth.

Regarding TMDs, the KMC simulations for the growth of WSe2 monolayer on graphene have been used to develop the phase diagram of domain morphologies as a function of flux and precursor stoichiometry. It was found that the fast kink nucleation and propagation, rather than edge attachment and diffusion, could lead to ultrafast growth of monolayer WSe2. KMC simulations also guided CVD growth of large-scale WSe2 grains by controlling the three-stage adsorption-diffusion-attachment mechanisms. KMC model is also a key component in a more generalized mechanistic model for growth morphology predictions of 2D materials.

In summary, the KMC model can be a useful tool for investigating the kinetic pathways and morphologies during the growth of 2D materials. However, the probability-based nature of KMC makes it most suitable for cases where atomic fluctuations are high, i.e., the atomistic and nanoscale morphology and kinetics of 2D materials. For larger-scale simulations, due to the significantly increased system size and the disparate rates of different KMC events, a full-KMC is computationally expensive. As a compromise, multiscale KMC has been developed, simplifications should also be made to account for the key events that are most relevant to the large-scale kinetics. Under such situations, the phase-field approach could be a more efficient option. Meanwhile, the KMC model can provide the governing kinetic mechanisms for phase-field simulations to improve the validity and accuracy of the simulation.

**MACROSCALE MODELS**

Although the growth of 2D materials occurs at the nano- and mesoscale, it is controlled by physics and parameters that have a macroscopic nature, e.g., heat and mass transfer, furnace configuration, and gas-phase reactions. Thus, having a thorough understanding of the macroscale physics and processes is essential for controlling the growth of 2D materials and their synthesis by design. We can classify the macroscale models of the growth chamber into four groups: (i) experiment-based models, where rate equations and their constants used to describe the growth are determined from experiments; (ii) analytical models, where the governing equations are simplified and solved analytically; (iii) adaptive models, where a set of experiments is used to train the model; and (iv) multiphysics models, where the coupled system of governing equations at different length and temporal scales are solved numerically. Among these methods, the last group of models have key advantages, providing a profound understanding to the growth process, flexibility to apply to different growth conditions, and the ability to optimize the process.

A practical macroscale model of the growth chamber should capture the critical governing physics, e.g., the heat and mass transports and chemical reactions. Setting up these models requires several key information that can be obtained from lower scale simulations or experiments. Identifying the gas-phase reactions, we may approximate some of the reaction parameters using classical theories, e.g., the collision rate to estimate the chemisorption rate of species or the group contribution methods to determine the diffusion coefficient. The other approximation is for low concentrations of reactive species, where the change in pressure and heat of reaction can be neglected. In the latter case, we may decouple the fluid and heat transfer of gaseous materials from the mass transfer and kinetics. In contrast, for high concentrations of reactive species in the gas phase, such as in metalorganic chemical vapor deposition (MOCVD), the coupled system of equations must be solved. A summary of main equations in macroscale models are presented below.

**Gas flow**

The Navier-Stokes equation governs the flow rates in the growth chamber,

\[
\rho \frac{\partial \mathbf{u}}{\partial t} + \rho (\mathbf{u} \cdot \nabla) \mathbf{u} = \nabla \cdot \left( -\rho I + \mu \left( \nabla \mathbf{u} + (\nabla \mathbf{u})^T \right) \right) + \mathbf{F}; \quad \rho \nabla \cdot \mathbf{u} = 0.
\]

where \( \mathbf{u} \) is the velocity field, \( \rho \) is pressure, \( I \) is the unit matrix, \( \mu \) is the dynamic viscosity, and \( \mathbf{F} \) is the volumetric applied force, i.e.,
weight. Furthermore, the density $\rho$ is a function of temperature and precursor concentration. In general, the concentration of precursor in the gas phase is low and $\rho$ can be assumed to be only a function of temperature.

Heat transfer
The two main heat transfer mechanisms in the growth chamber are convection and conduction that respectively are

$$Q = \rho C_p \frac{\partial T}{\partial t} + \rho C_p a \cdot \nabla T + \nabla \cdot q; \quad q = -k \nabla T,$$

where $Q$ is the heat source, $C_p$ is the heat capacity at constant pressure, $T$ is the temperature, $q$ is the heat flux vector, and $k$ is the thermal conductivity coefficient.

Mass transfer
Commonly, the concentration of precursor materials in the growth chamber is negligible. Thus, the equation governing the mass transfer can be formulated with the flow-assisted diffusion of dilute species

$$R = \frac{\partial c}{\partial t} + \nabla \cdot (-D \nabla c) + \mathbf{u} \cdot \nabla c; \quad N = -D \nabla c + \mathbf{uc},$$

where $R$ is the source term for precursor, $c$ is the precursor concentration, $D$ is the diffusion coefficient, and $N$ is the flux of the precursor. Equations (32)–(34) must be solved for a set of initial and boundary conditions, as shown in Table 6.

Chemical reaction
Consider a set of $j$ reactions involving $i$ species of the form $aA + bB + \ldots = xX + yY + \ldots$ with forward and reverse reaction rate constants $k_f$ and $k_r$, respectively. These reaction rates depend on the temperature via an Arrhenius expression, e.g.,

$$k_f = \tilde{k}(T/T_\text{ref})^n \exp\left(-\frac{E_f}{RT}\right).$$

Here, $\tilde{k}$ denotes the frequency factor, $n$ the temperature exponent, $E_f$ the activation energy, and $R_g$ the gas constant. According to the mass action law, the reaction rates $r_j$ can be described as

$$r_j = k_f \prod_{i \in \text{react}} c_i^{n_i} - k_r \prod_{j \in \text{prod}} c_j^{n_j},$$

where $c_i$ is the concentration of $i$-th species and $v_j$ is the stoichiometric coefficient (negative for reactants and positive for products). The rate equations provide the information regarding the concentration of different species, which in combination with mass and heat balance equations form the complete system of equations.

Macroscale models provide insight into the growth mechanisms and flow regimes, and play a crucial role in determining proper furnace design and growth parameters. Figure 6 shows the application of a macroscopic model of the flow and concentration diffusion in a vertical furnace with a side inlet to study the effect of the gap between the susceptor and substrate. It shows that the presence of the gap results in the formation of eddy currents close to the substrate edge, which distorts precursor concentration and hinders the formation of 2D materials that is consistent with the experimental observations. These models have been successfully used in determining the mechanisms governing the growth of 2D materials. For example, it was revealed that the concentration gradient of the precursor is a critical factor in determining the growth morphology of 2D materials, where a low concentration gradient along the substrate leads to the formation of multigrain films. In contrast, moderate planar concentration gradients lead to isolated islands of varying morphologies, and an out-of-plane gradient leads to the formation of standing 2D materials. Furthermore, macroscale models can be combined with lower scale models to form a multiscale platform with high fidelity, e.g., ref. 9.

### MATERIALS GENOME APPROACHES

Data-driven methods have been employed to satisfy the urgent need for theoretical development of 2D materials. The improvement of computational power has resulted in extensive collections of data and scientifically organized databases, which can be used by methods such as data mining and machine learning, to accelerate the computational design of new 2D materials.

#### Database of 2D materials

Compared with the databases of 3D bulk materials such as the Crystallographic Open Database, Inorganic Crystal Structure Database, and the Cambridge Crystallographic Data Centre, the database focused on 2D materials is still under development. Several 2D materials databases have been attempted primarily based on a high-throughput approach coupled with first-principles and quantum theory to address this shortcoming. Besides, data-mining methods have also been applied to screen those bulk materials databases and identify possible 2D materials from 3D structures. These approaches can provide various information about 2D materials that have been or not been synthesized in experiments. For example, using a criterion to identify 2D materials, a database has been developed on the basis of experimental and DFT data. This database provides information including exfoliation energy, bandgap, work function, and elastic-constant values of various 2D materials, and also predicted hundreds of new 2D materials for the experimental synthesis. DFT and many-body perturbation theory have been used to generate the Computational 2D Materials Database (C2DB). The C2DB contains structural, thermodynamic, elastic, electronic, magnetic, and optical information of about two thousand 2D materials in more than 30 crystal structures. A few hundred new monolayer structures...
that were previously unknown, but potentially synthesizable were also identified. A data-mining method has also been used to screen the bulk materials from the open database and identified a few hundred 2D weakly bonded solid materials. The band gaps and point groups that determine piezoelectric and nonlinear optical properties in these materials were also provided. A search in the database of bulk materials was performed and DFT calculations were used to explore these materials for possible exfoliation, where more than one thousand materials have been identified. Information including electronic, vibrational, magnetic and topological properties of promising materials was also provided. Similarly, the crystal database was screened for layered motifs, and hundreds of stable layered materials were found that can be considered as candidates for the formation of 2D monolayers via exfoliation.

Machine learning method

Machine learning models trained using 2D material databases can be parameterized, validated, and therefore used for predictive exploration of novel 2D materials with desired properties. Machine learning methods can also aid the understanding of the complex correlations between structures and properties in 2D materials. A combination of machine learning with MD simulations and in-situ high-resolution transmission electron microscopy to explore TMDs has been performed, where machine learning provided information for structural optimization and evolution of defects to help understand the structural transition in 2D TMDs. A practical method to explore hybrid 2D materials was developed by coupling machine learning with DFT calculations. The structural and electronic properties of different hybrid 2D materials were provided and various parameters for vdW heterostructures were screened. A machine learning model with force-filed-inspired descriptors in material screening for complex systems has been introduced and used to discover exfoliated 2D-layered materials. An artificial neural network for titanium dioxide systems was trained based on the a DFT calculated database, where a novel quasi 2D titanium dioxide structure was revealed. Similarly, new 2D materials with high magnetic moments were found using a machine learning model trained by first-principles data. Machine learning methods have also been used to aid the development of force fields for classical simulations of materials. For instance, a force field for classical simulations of stanene was developed using a machine learning method trained by data sets from ab-initio results to calculate the mechanical and thermal properties of stanene. 2D materials without bulk layered counterparts are also being discovered, using genetic algorithms or particle swarm optimization, both using energy-based merit criteria and additional biases towards 2D sheets.

OUTLOOK

This overview summarizes the state-of-the-art of modeling efforts on the growth of 2D-layered materials and outlines the advantages and limitations of computational models and methods at different length and time scales. The emphasis is on modeling, understanding, and predicting the thermodynamics and kinetics of 2D materials growth. The eventual goal is to develop computational tools and techniques to realize the “synthesis by design” of 2D materials.

An ideal computational model should not only capture accurately the correct physics to allow robust designs but also be computationally efficient to be employed for active control of the growth process. The complex nature of the growth of 2D materials requires applications of a range of models, which operate at multiple length and temporal scales to capture the full range of the growth phenomena from macroscale flow and concentration profiles in the furnace to atomistic reactions leading to formation and growth of 2D materials at mesoscale. Modeling the full spectrum of mechanisms involved in the growth of 2D materials is challenging due to the limitations associated with each of the computational models at different length and temporal scales.

A high-fidelity prediction of 2D materials growth often requires an integration of methods applicable across different scales, ranging from electronic and atomistic, to mesoscale phenomenological, and large-scale continuum models. For example, chemical potentials of different phases and the migration energies of atomic species in different structural and chemical environment from electronic and atomistic models can be employed to parametrize mesoscale phase-field models for predicting the morphology and thus characteristics of as-grown 2D materials with the boundary conditions specified by the information obtained from large-scale continuum models such as temperature distribution in the growth chamber as presented in ref. . Furthermore, technical obstacles to in-situ experimental monitoring of growth hinder the validation of computational predictions. The current mathematical and numerical models are computationally expensive and commonly do not capture simultaneously all the physics involved in a growth process of 2D materials, limiting their widespread adoption in active control of 2D materials growth. One promising way to circumvent this challenge is to use machine learning models trained using experimental and
computational databases of the 2D materials structure and growth conditions such as the ones available at the 2D Crystal Consortium—Materials Innovation Platform. Finally, the models and approaches applied in understanding the growth of 2D materials can be adapted to the growth of other materials that use the same synthesis techniques, e.g., CVD growth of thin films.

As a future perspective, we can envision two main strategies for the development of computational models and their applications to the design and synthesis of 2D materials, which are long-term challenging goals of the field. First is the open-loop design approach, where the growth chamber design and synthesis conditions are determined by performing a series of simulations mimicking the thermophysical conditions and reaction and growth kinetics that would be expected in experiments. In this approach, the models must have high fidelity and a comprehensive sensitivity analysis should be performed to obtain a robust design. The second approach is the closed-loop design that the developed models will be utilized to determine the states of the system to be passed to a controller which adjusts the growth conditions in real-time. In this approach, the computational efficiency of the models is key to the success of the design.

**DATA AVAILABILITY**

All data that was obtained during this project are available from the authors.

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