Metallene-related materials for electrocatalysis and energy conversion

Minghao Xie, Sishuang Tang, Bowen Zhang and Guihua Yu

As a member of graphene analogs, metallenes are a class of two-dimensional materials with atomic thickness and well-controlled surface atomic arrangement made of metals or alloys. When utilized as catalysts, metallenes exhibit distinctive physicochemical properties endowed from the under-coordinated metal atoms on the surface, making them highly competitive candidates for energy-related electrocatalysis and energy conversion systems. Significantly, their catalytic activity can be precisely tuned through the chemical modification of their surface and subsurface atoms for efficient catalyst engineering. This minireview summarizes the recent progress in the synthesis and characterization of metallenes, together with their use as electrocatalysts toward reactions for energy conversion. In the Synthesis section, we pay particular attention to the strategies designed to tune their exposed facets, composition, and surface strain, as well as the porosity/cavity, defects, and crystallinity on the surface. We then discuss the electrocatalytic properties of metallenes in terms of oxygen reduction, hydrogen evolution, alcohol and acid oxidation, carbon dioxide reduction, and nitrogen reduction reaction, with a small extension regarding photocatalysis. At the end, we offer perspectives on the challenges and opportunities with respect to the synthesis, characterization, modeling, and application of metallenes.

1. Introduction

Materials with an ultrathin two-dimensional (2D) structure exhibit unique and unprecedented chemical, electronic, and catalytic properties owing to the distinctive electron confinement featured in two dimensions which is unachievable either in other classes of nanomaterials or their bulk counterparts. Since the exfoliation of the first 2D nanocrystal, graphene, the family of 2D materials has experienced a big boom. As a remarkable group of 2D materials, graphene analogs characterized by atomic layers are essentially important for surface-related applications due to their ultrahigh surface-to-volume ratio and thus maximized atomic utilization efficiency (AUE). To date, the reported graphene analogs are essentially divided into three categories (Fig. 1A):

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Minireview

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(1) metal-free analogs such as carbon nitride, boron nitride, phosphorene, etc.; (2) transition metal compounds such as carbides (also known as MXenes), chalcogenide, hydroxide, etc.; (3) metallic materials such as monometallenes, bimetallenes, and trimetallenes, etc. Featured with laminar structures stacked by van der Waals forces, the first two categories can be synthesized via top-down exfoliation or bottom-up growth down to single atomic layers.8 In contrast, due to the robust and isotropic nature of metallic bonds, the majority of metal materials are characterized by closely packed crystal structures, making their atomically thin morphology thermodynamically unfavorable.9 As such, the preparation of mettalenes remains a significant challenge, hindering their generation for potential applications in the past few decades.

As a newcomer of 2D materials, mettalenes have garnered ever-increasing attention in recent years owing to their unique superiors including the maximized exposure of active metal sites and high conductivity for catalysis, electronics, and energy-related applications.10,11 Particularly, the precisely tunable electronic structure, modifiable surface state, and high carrier mobility make mettalenes competitive candidates for energy conversion devices. In essence, mettalenes refer to a class of atomically thin 2D materials composed of metals and alloys. Fig. 1B instinctively demonstrates an overview of the classification of mettalenes from the perspective of compositions and the metal elements reported so far. Mettalenes composed of noble metals like Pd,12,13 Ir,14 and Rh15,16 which are commonly used in electrocatalysis for energy conversion delivered outstanding electrocatalytic performance for practical applications. In addition, several strategies were developed to boost their performance to a new limit, including atomic doping, lattice engineering, surface modification, etc.10 On top of that, the recent success of PdMo bimetallene with four atomic layers significantly proved the superior performance of mettalenes in electrocatalysis of oxygen reduction, triggering substantial studies of mettalenes as electrocatalysts for energy-related applications.17

In this minireview, we summarize some of the recent advancements and achievements in the fabrication, characterization, and electrocatalytic applications of mettalenes. We begin with a brief introduction to general approaches to the synthesis and characterization of mettalenes. We pay particular attention to the design principles of mettalenes from the viewpoint of catalyst engineering with high performance. We then discuss the electrocatalytic properties of mettalenes and showcase a set of energy-related reactions including oxygen reduction, hydrogen evolution, alcohol and acid oxidation, carbon dioxide reduction, and nitrogen reduction reaction, with extended discussion regarding photocatalysis. Finally, on the basis of the present progress, we highlight some potential prospects on the challenges and outlooks with respect to synthesis, characterization, modeling, and stability of mettalenes for realistic energy application.

2. Definition, synthesis, and characterization of mettalenes

The definition of mettalle is adopted from that of graphene. In principle, a well-defined mettalle refers to a single atomic layer composed of metal atoms. Similar to graphene, the monolayer structure is highly flexible which can feature atomic surface flatness when deposited on a substrate or surface curvature when it is freestanding due to the high surface free energy. The use of “ene” for metal is also due to the comparable electronic properties between mettalenes and graphene when the thickness of metal is reduced to a monolayer.8 In reality, the high energy barrier to overcome and the thermodynamic instability of metal monolayers limits the synthesis approaches for obtaining this type of nanostructure either on a
substrate or in an isolated state. In this Minireview, the definition of metallenes generally relates to atomically thin 2D metals with a thickness of around 1 nm (corresponding to atomic layers) with rare exceptions above 2 nm. Different from conventional 2D metal nanoplates typically with flat surface morphology, metallenes feature a lower thickness and much larger lateral size, which makes them favorable to form highly curved morphology in most cases.

2.1. Synthesis of metallene: bottom-up, top-down, and topotactic metallization

The controlled synthesis of metallene enables the alteration of thickness, composition, and crystal structure to vary their electrochemical properties and thus catalytic performance. Thanks to the efforts of many groups, a number of approaches have been developed for the synthesis of 2D metal nanomaterials with controlled thickness, composition, and crystal structure. The typical synthesis methods for metallene are summarized in Table 1, which can be generally categorized into three major strategies including bottom-up wet-chemical methods, top-down exfoliation methods, and topotactic metallization approaches.

In the bottom-up methods, the major concern for the generation of metallene is the introduction of anisotropic growth of metal at the nanoscale. According to the classic LaMer model, the anisotropic nucleation of metal atoms for the birth of 2D nuclei is critical to induce the subsequent production of an ultrathin nanostructure of metal. To confine the thickness of the 2D nanostructure, the growth perpendicular to the basal planes of the nanostructure needs to be passive, which can be achieved by well-selected ligands preferentially attached to the basal planes to make this ultrathin nanostructure thermodynamically favorable. As demonstrated in Fig. 2A and B, the ligands used for anisotropic confinement...
can be inorganic molecules such as CO or organic surfactants to produce ultrathin Pd metallenes.\textsuperscript{10,20,21} Similar to ligand-confined growth, the anisotropy can also be induced by laminar structures in which the growth of metals is well-confined within the interlayer space of the layered structures. As shown in Fig. 2C, the PdCo single atomic layers can be obtained by a space-confined growth of PdCo monolayers in the angstrom-sized interlamination of layered crystalline clay mineral montmorillonite.\textsuperscript{28} In addition to the confined growth approaches, the 2D nanostructure can be thermodynamically favored by template-mediated anisotropy. Starting from a template with an atomically flat 2D surface, metal atoms can be preferentially deposited on the surface of the template for the generation of ultrathin metallic layers. As an intuitively selected template, graphene has been used for the preparation of metallenes. As reported by Zhang and coworkers, Au metallene with an unconventional hexagonal close packed (hcp) crystal structure can be obtained by depositing Au atoms on graphene.\textsuperscript{31} It is worth noting that the thermodynamically unfavored metallene can also be stabilized in the pores of graphene, by which the free-standing Fe monolayer suspended in graphene pores was gained.\textsuperscript{30} Besides graphene, metallic templates are also widely used for the epitaxial growth of metal atoms by atomic layer-by-layer deposition. By using this strategy, a variety of metal atoms can be deposited on the surface of templates in a well-controlled way, which paves the way for the formation of metallene with higher complexity in terms of composition. For example, Guo and coworkers reported metallene composed of high-entropy alloys \textit{via} galvanic replacement reaction and co-reduction of various metal ions on the surface of an Ag template followed by selective etching of the template.\textsuperscript{32} Beyond the aforementioned methods, metallene can also be synthesized \textit{via} other unique bottom-up approaches including pulsed laser deposition, pyrolysis, electrodeposition, and photoreduction, \textit{etc.}\textsuperscript{11}

It is well known that laminar materials such as graphite, transition metal dichalcogenides, and layered double hydroxides feature strong in-plane covalent bonds and weak interlayer van der Waals interactions, providing possibility to exfoliate them into mono- or few layers.\textsuperscript{47–49} Likewise, metallenes composed of elements with layered structures (such as Ga, Sb, and Bi) can also be straightforwardly obtained from their bulk counterparts \textit{via} various exfoliation techniques.\textsuperscript{11} Among them, liquid-phase exfoliation is widely used due to the high convenience and effectiveness of this method. As demonstrated in Fig. 2D, free-standing layers of Bi can be obtained by ion intercalation followed by exfoliation driven by the power of ultrasonication.\textsuperscript{50} In addition to liquid-phase exfoliation, 2D materials such as graphene can be fabricated by mechanical exfoliation from their bulk counterparts, which has been utilized in the synthesis of Sb metallenes. Using a mechanical exfoliation realized by pregrinding-induced shear force followed by ultrasonic exfoliation, Sb plates and Sb metallenes with mono- and few layers thickness can be subsequently

<table>
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<tr>
<th>Synthesis approaches</th>
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<tr>
<td>Bottom-up Ligand-confined growth</td>
<td>Pd</td>
<td>5 layers</td>
<td>CO\textsubscript{2}RR</td>
<td>20</td>
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<td></td>
<td></td>
<td>3–8 layers</td>
<td>ORR</td>
<td>13</td>
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<tr>
<td></td>
<td></td>
<td>0.8 nm</td>
<td>Glycerol oxidation</td>
<td>21</td>
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<td></td>
<td></td>
<td>~1.0 nm</td>
<td>ORR</td>
<td>12</td>
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<tr>
<td></td>
<td>Rh</td>
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<td>Hydrogenation</td>
<td>15</td>
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<tr>
<td></td>
<td></td>
<td>~1.0 nm</td>
<td>Hydrogenation</td>
<td>16</td>
</tr>
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<td></td>
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<td>0.84 nm</td>
<td>CO\textsubscript{2}RR</td>
<td>22</td>
</tr>
<tr>
<td></td>
<td>RuRh</td>
<td>0.83 nm</td>
<td>HER</td>
<td>23</td>
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<tr>
<td></td>
<td>PdMo</td>
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<td>ORR</td>
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<td>PdIr</td>
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<td>24</td>
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<td></td>
<td>PdFe\textsubscript{2}</td>
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<td>NRR</td>
<td>25</td>
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<td>RhPd-H</td>
<td>~7 layers</td>
<td>HER</td>
<td>26</td>
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<td>28</td>
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<td></td>
<td>Pt</td>
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<td>29</td>
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<td>Template-directed growth Fe</td>
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<td>—</td>
<td>30</td>
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<tr>
<td>Au</td>
<td>~2.4 nm</td>
<td>—</td>
<td>31</td>
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<td>HEA</td>
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<td>ORR</td>
<td>32</td>
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<td>Top-down Liquid-phase exfoliation Bi</td>
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<td>33</td>
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<td>Sb</td>
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<td></td>
<td>1.5 nm</td>
<td>HER</td>
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<td>1.3 nm</td>
<td>CO\textsubscript{2}RR</td>
<td>36</td>
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<tr>
<td>Mechanical exfoliation Co</td>
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<td>Perovskite solar cells</td>
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<td>Solid-melt exfoliation Sb</td>
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<tr>
<td>Electron beam dealloying Ga</td>
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<td>—</td>
<td>39</td>
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<td>Electron beam dealloying Au</td>
<td>Monolayer</td>
<td>—</td>
<td>39</td>
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<td>Topotactic metallization Chemical metallization Bi</td>
<td>0.65 nm</td>
<td>CO\textsubscript{2}RR</td>
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<td>Rh</td>
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<td>NRR</td>
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<td>NiFe</td>
<td>0.65 nm</td>
<td>NRR</td>
<td>42</td>
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<td>CO\textsubscript{2}RR</td>
<td>43</td>
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<tr>
<td></td>
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<td>CO\textsubscript{2}RR</td>
<td>44</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1.21–1.55 nm</td>
<td>CO\textsubscript{2}RR</td>
<td>45</td>
<td></td>
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<tr>
<td></td>
<td>1.02 nm</td>
<td>CO\textsubscript{2}RR</td>
<td>46</td>
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produced. Furthermore, some other top-down approaches are also utilized in the synthesis of metallasenes, including electrochemical exfoliation, solid-melt exfoliation, hot-pressing exfoliation, and dealloying. For instance, Ajayan and coworkers successfully prepared Ga metallasenes from bulk Ga by solid-melt exfoliation. The low melting point (\( T_m \approx 302.85 \) K) of bulk Ga provides the possibility to strip a few layers of Ga from liquid Ga on different substrates (Fig. 2F). In another typical example, a free-standing Au monolayer framed in bulk crystals can be obtained by in situ dealloying of a bulk Au–Ag alloy by electron beam in a transmission electron microscope (TEM) (Fig. 2G).

The special morphology and composition of metallasenes make topotactic metallization of the layered metal compounds another important synthetic method. Specifically, ultrathin layers composed of pure metals can be obtained by the extraction of the in-plane/interlayered non-metallic groups from the laminar metal compounds under reductive conditions without compromising the original layered structures. For instance, as shown in Fig. 2H, Ru monolayer can be prepared via topotactic metallization of layered RuO\(_2\) under a H\(_2\) atmosphere. Moreover, non-noble metallasenes composed of Co or Ni have been successfully synthesized from their hydroxide precursors under reductive conditions. The reductive conditions can be achieved by chemical or electrochemical methods, which have been applied in the synthesis of diverse Bi metallasenes with different morphology and thickness.
2.2. Characterization of mettallene at the nanoscale: morphology, thickness, and composition

The atomic thin nature of mettallenes requires special characterization techniques to confirm their morphology, thickness, and composition. In terms of the morphology characterization of metals, the TEM technique is usually used to show the phase and mass contrast generated by the presence of metal atoms. However, the thermodynamically unstable metallic layers often suffer from stability issues under an electron beam, leading to the morphology and/or phase evolution from thermodynamically unfavorable states to stable states. One typical example of the morphology change of mettallene is observed in the TEM characterization of Rh mettallene. As shown in Fig. 3A, an obvious increased porosity of Rh mettallene can be induced after beam irradiation for 28 s, indicating the high sensitivity of the mettallenes to external energy due to the atomic thin nature of Rh mettallene. In another example, an electron beam-induced hcp to face-centered cubic (fcc) phase transformation of Au mettallene was observed after irradiation for ~20 s. Therefore, TEM characterization of ultrathin mettallenes requires special attention with consideration of the morphology change under different exposure times under an electron beam.

The subnanometer thickness of mettallenes call for techniques with high sensitivity to the surface properties of materials. By flatly depositing mettallenes on a substrate, their thickness can be directly reflected by the change of the height of the substrate surface, which can be detected by scanning probe microscopy such as atomic force microscopy (AFM). As shown in Fig. 3C and D1, partially oxidized Co mettallene and PdMo mettallenes with four atomic layers and a thickness below 9 angstroms can be clearly revealed by AFM. Moreover, the thickness of mettallene can also be directly illustrated by their lateral view using atomic resolution scanning transmission electron microscopy (STEM)/TEM. Compared with AFM, the use of high resolution STEM/TEM not only confirms the thickness of mettallenes but also clearly indicates the number of atomic layers and the corresponding atomic arrangement perpendicular to the basal planes of mettallenes. As shown in Fig. 3E and F1, the lateral STEM and TEM images of Co and Rh mettallenes provide evidence for the construction of the corresponding models showing the crystal structures and atomic arrangements in the mettallenes.

Besides morphology and thickness, other significant characteristics of mettallenes are their surface properties and composition, which essentially determine their physicochemical properties. In general, the surface properties of mettallenes relate to the coordination number of surface atoms which is highly dependent on surface atomic arrangement. Similar to the lateral view of mettallene by atomic resolution STEM, the projection of mettallene perpendicular to the basal plane can clearly indicate the atomic arrangement over the surface, which can be used for the analysis of lattice spacings along different crystallographic orientations. As demonstrated in Fig. 3G and H, the hexagonal arrangement of surface atoms was clearly observed for PdMo-C and RhPd-H mettallenes. In mettallenes composed of alloys, the divergence of mass contrast of surface atoms caused by different atomic numbers of involved metals can also be reflected in the STEM image, which can be analyzed by the corresponding integrated intensity profile. As shown in Fig. 3I and J1, the STEM image of pentamery mettallene demonstrate square arrangement of surface atoms with distinguishable integrated pixel intensities over different regions. To quantitatively confirm the composition of mettallene especially those consisting of solid solution alloys, typical metallic detection techniques are required, such as inductively coupled plasma mass spectrometry (ICP-MS) and inductively coupled plasma atomic emission spectrometry (ICP-AES). The distribution of different metals over the mettallene can be revealed by energy dispersive X-ray spectroscopy (EDS). For example, Fig. 3J and J1 show the EDS elemental mapping of an individual mettallene, clearly demonstrating the uniform distribution of Rh and Pd over the entire RhPd-H mettallene. When a monometallene is doped with a trace amount of a secondary metal atoms, single atom alloy mettallene can be acquired with an atomically distributed secondary metal on the surface. In this case, the detection of a single atom metal becomes much trickier, and requires characterization approaches that are usually used in the analysis of single atom alloys, such as topographic atom imaging analysis, X-ray absorption near-edge structure (XANES), and extended X-ray absorption fine structure (EXAFS). As a typical example, the presence of single Fe atoms on the surface of PdFe single atom alloy mettallene was evidently illustrated by a well-performed 3D topographic atom imaging analysis (Fig. 3K). Taken together, the characterization of mettallene is considerably complicated, and requires more attention relative to other metal-based nanomaterials.

3. Design of mettallenes for electrocatalysis

The diversity of synthetic approaches provides the possibility to enable modification of the physicochemical properties of mettallenes for electrocatalytic applications. According to the Sabatier principle, the activity of a heterogeneous catalyst is highly dependent on the interaction between its surface atoms and the intermediate(s) involved in the rate-determining step of the catalytic reaction. Fundamentally, the above interaction originates from the formation and occupancy of bonding and antibonding orbitals between the intermediate(s) and the catalytic site. According to this principle, the adsorption strength of intermediate species should be at a mediate value, neither too weak nor too strong, for the catalyst with an optimal activity. On the basis of the d-band theory, the change of electronic (d-band) structure can intrinsically alter the coupling strength between the adsorbate and the d-band of transition metals and thus result in the variation of their electrocatalytic activity. The electronic structure of metals is essentially determined by surface atomic arrangement, elemental composition, and lattice strain. In addition, the coordination number (CN) of active sites can also intrinsically alter the adsorption and dissociation energy of reactant species on a metal surface,
Fig. 3 Characterization of mettallenes at the nanoscale. (A and B) Morphology characterization. (A) TEM images of an individual Rh mettallen after electron beam exposure for 28 s (scale bar, 20 nm). Reproduced from ref. 15 with permission from Springer Nature. Copyright 2014. (B) Scheme showing the electron beam-induced phase transformation of an Au mettallen. (B1) TEM image of a porous Au mettallen after electron beam irradiation for about 20 s (scale bar, 50 nm). Reproduced from ref. 31 with permission from Springer Nature. Copyright 2011. (C–F1) Thickness characterization. (C) AFM image and (C1) corresponding height profiles of Co mettallen. Reproduced from ref. 22 with permission from Springer Nature. Copyright 2016. (D) AFM image and (D1) corresponding height profiles of PdMo bimettallen. Reproduced from ref. 17 with permission from Springer Nature. Copyright 2019. (E) lateral
varying the catalytic performance of electrocatalysts made of metals. In light of the above understanding, two major routes are adopted for the controlled synthesis of metallenes with respect to the design principles of catalyst engineering. One is modulation of electronic structure, including tuning the surface facet (facet effect), surface composition (ligand effect), and surface strain (strain effect). The other route is engineering the coordination environment of surface atoms. These general strategies are demonstrated in Fig. 4 and will be discussed in the following section.

### 3.1. Surface facet control

Since the electronic structure of transition metals is essentially determined by the atomic arrangement, designing their metallenes with well-controlled surface facets is an effective strategy to improve their electrocatalytic activity. Fig. 5A clearly illustrates the three types of low-index facets, (111), (100), and (110), which are typically taken by the transition metal with an fcc structure. The difference in atomic arrangement not only changes the packing density of atoms but also varies the number of broken bonds per surface atoms, therefore leading to different specific surface free energy as well as catalytic activity of these facets.

Generally, the exposed facets of metal nanocrystals synthesized by wet chemical methods can be controlled by using a well-selected capping agent which can be preferentially adsorbed on the desired planes to reduce their surface free energy and thus passivate their growth rate for exposure. In terms of metallenes, similar strategies can be adopted to control their surface atomic arrangement via ligand-assisted methods. For example, Liu and coworkers discovered that

**Fig. 4** Schematic illustration of the design principle for optimizing electrocatalytic properties of metallenes. The optimization strategies are categorized into two major routes: (1) modulation of electronic structure via different routes including facet effect, ligand effect, and strain effect. (2) Engineering the coordination environment via inducing surface porosity/cavity, defects, or amorphization.
selective adsorption of functional groups of COOH, pyridyl, and quaternary ammonium on different planes of Pd could induce the anisotropic growth of ultrathin Pd enclosed by \{100\}, \{110\}, and \{111\} facets, respectively (Fig. 5B–D). The length of the alkyl chain of the ligand needs to be controlled long enough to form stable lamellar mesophases for the confined growth of Pd. This tunability of surface atomic arrangement provides an opportunity to evaluate facet-dependent catalytic properties of Pd for various energy-related reactions.

3.2. Surface and subsurface composition control

Since the d-band structures varies over different transition metals, they intrinsically feature different activities on a specific catalytic reaction. For example, platinum group metals (PGMs) have been discovered as the prime catalysts for formic acid decomposition due to mediate values of heat for the formation of metal formate \((\Delta H_f)\) on their surfaces and thus an optimal net reaction rate balanced by the rate of adsorption and desorption. Therefore, the synthesis of mettallenes made of various single metals is important to study the 2D morphological effects of metals on electrocatalysis. Thanks to the synthetic capabilities developed over recent decades, a number of monometallenes, including Co, Rh, Pd, Ir, and Au, have been synthesized with well-controlled size, thickness, and surface facets (Fig. 6A–A5). The successful synthesis of monometallenes composed of diverse metals enables further

development of bi-, tri-, and even high-entropy metallenes with improved electrocatalytic activity and selectivity.

As demonstrated in Fig. 2, the inclusion of a secondary metal element into the lattice of a transition metal can change the surrounding atoms of the surface catalytic sites and thus modify the electronic structure of the electrocatalyst, also known as the ligand effect.63 In terms of metallenes, atomic doping of the secondary metal can be controlled at either
top surface or subsurface, and both can intrinsically change the catalytic activity and selectivity of metallenes due to electron transfers between the primary and secondary metal elements. It is worth noting that the metallene structure features maximized surface area with a high surface free energy which is unfavorable for the stabilization of dopant. Therefore, sublayer doping of the secondary metals are usually adopted in the synthesis of bimetallenes, which prevents the leaching of dopant during electrocatalysis.17,64 Based on the requirements of compositions of alloys for various electrocatalytic reactions, a variety of bimetallenes and trimetallenes, including PdCu, PdMo, PdFe, PdPt, and PtPdNi, have been reported with remarkable catalytic properties (Fig. 6B–B5).17,27,64–66

When the number of metal elements increases above five in an alloy with an equant stoichiometry, the mix configuration entropy (S) of the system will pass a critical value (1.5R) which defines the concept of high-entropy alloys (HEAs). After the successful carbothermal shock synthesis of HEA nanoparticles and utilization in catalysis by the Hu group, increasing studies were conducted for the synthesis of HEA at the nanoscale with different structures for electrocatalysis.67–69 However, achieving 2D anisotropic growth of HEA remains a grand challenge owing to the distinct nucleation and growth kinetics of the composed metals and the intrinsic thermodynamic instability of 2D HEA. Very recently, Guo and coworkers proposed a general wet chemical method for generating up to an octonary HEA phase with a subnanometer ribbon structure using Ag nanowire as a template.32 As demonstrated in Fig. 6C, the production of this HEA 2D structure includes three major steps: (1) heterogeneous nucleation induced by the galvanic replacement reaction between metal ions and Ag template, (2) co-reduction of various metal ions on the Ag template, (3) removal of the Ag template by dealloying. The well-designed seed-mediated growth provided the possibility of incorporating diverse transition metals with catalytic properties into the 2D structure for the study of the effect of high-entropy alloys on electrocatalysis.

3.3. Surface strain control

Regarding the strain effect, the lattice distortion plays an essential role in shifting the d-band center of transition metals, changing the binding energy of an adsorbate on the surface of the electrocatalyst. Specifically, an expanded lattice reduces the overlap of d electrons between adjacent metal atoms, resulting in a narrowed bandwidth and the upshift of d states toward the Fermi level (EF) for the retention of d-band occupancy.70 According to the d-band center model, the upshifted d-band also lifts the antibonding state between the adsorbate and the d-band, decreasing the occupancy of the antibonding state and thus increasing the binding energy of the adsorbate.55 This fundamental correlation between surface strain and binding strength means that strain engineering is commonly used in designing electrocatalysts. In general, the lattice distortion of transition metals can be induced through four major routes: (1) epitaxial growth on a substrate with different lattice constants, (2) defect-induced lattice distortion such as vacancy, dislocation, and grain boundary, (3) the incorporation of secondary metal atoms with a different atomic radius in the lattice points, and (4) the intercalation of nonmetal atoms with small atomic radii at the interstitial sites of the lattice. With regard to metallenes, all of the routes have been reported for strain engineering. Specifically, the first route regards core-shell metal crystals with 2D morphology and a thickness of a few nanometers.9–71 The second route regards porous metallenes with a modified coordination environment of surface atoms which will be discussed in detail in Section 2.4. As discussed in Section 2.2, the third route results in bimetallenes and trimetallenes, in which the strain effect is coupled with the ligand effect for the engineering of electrocatalysts. Very recently, our group used the fourth route to synthesize Pd metallenes with controlled lattice strain.72 As demonstrated in Fig. 7A1–A4, the lattice expansion achieved by intercalations can be clearly demonstrated by an atomic resolution scanning transmission electron microscope image and X-ray diffraction pattern. A variety of nonmetal atoms, including H, C, and N, were inserted into the Pd lattice as the interstitial atoms, which offered a capability to study the difference of strain and ligand effect induced by various nonmetal elements.

The special 2D morphology of metallenes enables more approaches to strain engineering. The simplest way is downsampling the thickness, leading to the loss of nearby atoms and thus the reduced surface electron density. As a result, the attractive forces dominate on the surface and reduce the interatomic distance to equilibrate charge distributions. In contrast to bulk counterparts, the compressive strain caused by the contraction of surface atoms exhibit manifest effects on metallenes with a maximized ratio of surface atoms. A systematic study by the Greeley group offers a quantitative correlation between the lattice strain and the thickness of Pd metallenes.13 In this study, Pd metallenes of varying thickness were created by simply altering the CO source and the temperature environment involved in the synthesis. As shown in Fig. 7B1–B8, Pd metallenes with a compressive strain of 0.3, 1.2, and 1.5% were achieved on 8, 5, and 3 layers of Pd, respectively, demonstrating an inversely proportional correlation between intrinsic strain and thickness of Pd metallenes. In addition to the changed thickness, the curvature of metallenes caused by the minimization of surface free energy can also induce lattice distortion, leading to tensile strains generated on the surface of metallenes.17 The intrinsic tensile strains featured by highly curved metallenes have been proved to be favorable in electrocatalysis for specific reactions, which will be discussed in Section 3.1.17

3.4. Engineering of coordination environment

In addition to modulating the electronic structure of metallenes, another considerable way to regulate the adsorption behavior of adsorbates is adjusting the coordination environment of active sites. The essential descriptor of the coordination environment is CN which refers to the number of bonds formed between the atoms of interest and the neighboring atoms. For perfect crystalline materials, the CN of a surface atom is primarily determined by the lattice structure and the type of surface facet. Regarding metallenes with specific types of surface facet, inducing porosity on the surface is a specific
A way to modify the CN of active sites. Specifically, atoms situated at the edge of pores compose under coordinated surface sites with lower CN, greatly contributing to the catalytic activity of reactions that demand strong reaction species to active site bindings relative to saturated atoms located on the middle of the surface facet. In this regard, a variety of metallenes with a highly porous structure were reported for electrocatalysis (Fig. 8A–A3). Additionally, the presence of defects in imperfect crystals can also change the CN of atoms located around the defects (Fig. 8B). Therefore, altering the defect density on the surface of metallenes has also been utilized to improve the catalytic performance. As reported by Wang and coworkers, Pd metallenes with abundant Pd vacancies on the surface could provide active sites for reactant adsorption and accelerate charge transfer for the catalytic reaction (Fig. 8B1–B3).

Apart from the first-nearest neighbor atoms, the coordination environment can also be affected by the extended scale of atoms. As a remarkable example, the Bandarenka group proposed the concept of “generalized” CN, which involved the effect from the adjacent atoms of first-nearest neighbor atoms on CN of active sites, to predict the electrocatalytic activity of Pt-based catalysts. The study concluded that Pt atoms on a concaved (111) surface with CN larger than 7.5 featured an optimal binding strength towards the adsorbates, delivering a greater catalytic performance relative to Pt atoms on a pristine (111) surface with CN of 7.5. On the basis of this generalized coordination number theory, Guo and coworkers reported local coordination regulated Pd metallenes with atomic scale surface cavities and systematically demonstrated the relationship between the CN and catalytic activity of metallenes, which will be discussed in detail in Section 3.1. Another important way to change the coordination environment of active sites is amorphization of metallenes by ligand exchange or intercalation of nonmetal atoms. As shown in Fig. 9A–A1, partially amorphous Pd metallene intercalated with S atoms can be obtained from crystalline Pd metallene via solvothermal sulphuration. Besides, Zhang and coworkers reported ligand-exchange-induced amorphization of Pd, in which an order to disorder phase transition of fcc Pd was achieved by a thiol molecule (1,3,4-thiadiazole-2,5-dithiol, C₇H₇N₂S₂), namely bis-muthiol I, under ambient conditions without destroying their original morphologies (Fig. 9B–B5). The reduced crystallinity caused by amorphization not only increases the distance between metal atoms but also changes the coordination numbers of active sites on the surface, resulting in a large alteration of the electronic structure of metallenes for improved electrocatalytic performance.
4. Electrocatalysis and photocatalytic energy conversion

Given the intriguing structural and electronic properties and the aforementioned design strategies, 2D metallenes, especially those made of noble metals, have attracted increasing attention in a wide range of electrocatalysis and devices for energy conversion, including water splitting, CO₂ reduction, fuel cells, and metal-air batteries. Here we summarize the recent progress in utilizing metallenes as electrocatalysts for diverse catalytic reactions, including the oxygen reduction reaction (ORR), hydrogen evolution reaction (HER), alcohol and acid...
oxidation, CO₂ reduction (CO₂RR), and nitrogen reduction reaction (NRR), as well as photocatalysts for energy conversion.

4.1. Oxygen reduction reaction

As the cathodic reaction in fuel cells, the ORR has sluggish kinetics and limits the commercialization of the device.⁷⁶–⁷⁸ Although state-of-the-art Pt-based catalysts can essentially accelerate the ORR kinetics, the high price of Pt greatly increases the cost of fuel cells and hinders the extensive application of this class of catalyst.⁷⁹–⁸¹ Therefore, exploring catalysts with considerable activity relative to that of Pt is a considerable strategy for the commercialization of fuel cells. Recently, 2D Pd-based catalysts were identified as an alternative to Pt. With a structure of maximized AUE, Pd metallenes with atomic thickness feature abundant active sites on the surface for a superior ORR mass activity. As discussed in Section 2, the ORR activity of Pd metallenes can be further improved by dopants, thickness control, and modification of the coordination environment of surface atoms. For example, the PdMo metallenes reported by the Guo group delivered a mass activity of 16.37 A mg⁻¹ at 0.9 V versus the reversible hydrogen electrode (RHE) in alkaline electrolytes, which was 78 and 327 times higher than those of commercial Pt/C and Pd/C catalysts, respectively.¹⁷ The enhanced ORR performance could be attributed to the significant ligand effect induced by the doping of Mo atoms into Pd metallene with intrinsic surface tensile strains, in which charge transfer downshifted the d-band center by 0.26 eV relative to the pure Pd metallene and moved the binding energy of oxygen (ΔE₀) toward the optimal value for the ORR (Fig. 10A–A³).

The ORR performance of Pd metallene can also be finely tuned by altering their thickness. As demonstrated by the Wang group, the Pd metallene with a thickness of five monolayers delivered a higher ORR activity relative to those with three and eight monolayers in both acidic and alkaline electrolytes (Fig. 10B–B³).¹³ The optimal performance of five monolayer metallene arises from the optimal compressive surface strain, generating appropriate binding between the oxygen species and the surface atoms. Regarding the engineering of coordination environment, Guo and coworkers reported Pd metallenes with...
a series of average CN of surface atoms to systematically study the CN-dependent ORR activity of Pd.\textsuperscript{74} As shown in Fig. 10C–C3, the Pd metallene with an average CN of about 7.83 exhibited the highest activity and half-wave potential owing to the elevated CN and moderately downshifted d-band center.

4.2. Hydrogen evolution reaction

As a key half reaction for hydrogen production via water splitting, the HER is usually catalyzed by noble metals such as Pt, Pd, Ir, and Rh. Taking advantage of the 2D morphology, a variety of noble metallenes have been utilized in the HER with outstanding performance. For instance, Cheng and coworkers demonstrated that defective and porous Ir metallenes delivered a superior HER performance with an overpotential of 14 mV at a current density of 10 mA cm\textsuperscript{–2}, a Tafel slope of 31.2 mV dec\textsuperscript{–1}, and a double-layer capacitance of 4.30 mF cm\textsuperscript{–2}.\textsuperscript{14} The HER activity of Ir metallene surpassed that of Pt/C due to the large specific surface area, high conductivity, and efficient mass transfer benefitting from the ultrathin 2D morphology. In addition, the partially hydroxylated surface achieved by polyallylamine (PAH) modification featured a modulated electronic structure and efficient proton carriers, which is favorable to the HER in acidic media. As discussed in Section 2.1, the exposed facet on the surface of metallenes can also alter their HER performance. The facet-dependent HER activity was investigated by Liu and coworkers, comparing the HER activity of Pd metallenes with (100), (111), and (110) facets.\textsuperscript{60} Due to the optimal balance between hydrogen adsorption and desorption on Pd(100) planes, the Pd metallene with a surface facet of (100) planes showed the lowest overpotentials at current densities of 10, 50, and 100 mA cm\textsuperscript{–2} in an acidic electrolyte relative to those exposed with (111) and (110) planes (Fig. 11A and A1). Moreover, the anisotropic 2D morphology could largely retard the electrochemical dissolution and ripening process of Pd metallene, realizing excellent HER durability with a slight degradation over 20 h (Fig. 11A2 and A3).

The HER activity of metallenes can be further improved \textit{via} inducing a strain and ligand effect achieved by the
incorporation of either metallic or nonmetallic elements. For example, bimetallenes such as RuRh and IrRh metallenes have been demonstrated as outstanding electrocatalysts towards the HER.\textsuperscript{82,83} In addition to forming alloys, intercalation of light atoms, defined as being within the third-row elements, has also been proved an effective way to modify the electronic structure of metallenes.\textsuperscript{84–88} Different from the secondary metal atoms located at the lattice point of metallenes, the much smaller atomic radii of light atoms enable them to permeate into the interstitial sites of the lattice, inducing lattice enlargement and charge transfer between the host and guest atoms. This strain and ligand effect caused by light atoms enhances the catalytic activity and selectivity of metallenes.\textsuperscript{84} The RhPd-H bimetallene also delivered an outstanding HER durability due to the ultrathin 2D sheet-like morphology with a faster electron and mass transfer rate on the open double-sided surfaces.\textsuperscript{26} The amorphization of Pd metallene can also enhance the HER activity. As reported by our group, the sulphuration-induced Pd amorphization resulted in the greater HER performance of Pd–S metallene relative to those of the crystalline counterparts (Fig. 11C–C\textsubscript{3}).\textsuperscript{72} This improvement can be ascribed to the well-modified electronic structure caused by the ligand effect and amorphous structure. The elongated Pd–Pd bond length and increased d-band downshift in amorphous structure weakened the hydrogen bonding and thus improved the HER activity. The disordered surface and presence of S atoms also altered the coordination environment of surface Pd atoms, potentially providing abundant active sites on Pd–S metallene for the HER.

### 4.3. Alcohol and acid oxidation reaction

Chemical fuels, such as formic acid, ethanol, and ethylene glycol, are commonly used in the anodic reaction in direct liquid fuel cells.\textsuperscript{77,89} The oxidation reactions of these chemicals
can be greatly promoted when using Pt- and Pd-based electrocatalysts.\textsuperscript{90–93} Owing to the largely exposed surface area, abundant active sites, and excellent poisoning resistance, Pd-based metallenes have been widely used in these oxidation reactions.\textsuperscript{65,66,94,95} Moreover, the electrocatalytic activity can be further improved when alloying Pd with other metal or nonmetal elements to modify the electronic structure and crystal phase. By alloying Pd with Cu, Zhang and coworkers reported ultrathin PdCu bimetallenes with a superior mass activity of 1.66 A mg\textsuperscript{-1} Pd toward the formic acid oxidation reaction (FAOR), which was about 14 times higher than that of commercial Pd black (Fig. 12A–A\textsubscript{3}).\textsuperscript{65} The enhancement in FAOR activity could be attributed to the 2D morphology, the ligand effect between Pd and Cu, and the maximized exposure of surface active sites by the post-treatment with ethylenediamine (EN).

Alloying Pd with another transition metal and basic metals such as Zn, Cd, and Pb can not only alter the electronic structure but also result in phase transition of Pd.\textsuperscript{94,96} For example, Zhang and coworkers reported the successful synthesis of PdZn and PdCd bimetallenes with a face-centered tetragonal (fct) structure.\textsuperscript{94} The PdZn bimetallene displayed an improved mass activity of 2.73 A mg\textsuperscript{-1} Pd toward the ethanol oxidation reaction (EOR), which was about 1.74 and 2.97 fold higher than those of Pd metallene and commercial Pd black, respectively (Fig. 12B–B\textsubscript{3}). Apart from the 2D nature with plenty of undercoordinated active sites on the surface, the improved EOR performance of PdZn bimetallene might also result from the unique electronic and crystal structure caused by the incorporation of Zn into the Pd lattice, together with the polycrystalline structure of the metallene with extra active sites for the EOR. When alloying Pd with p-block nonmetal elements, the unconventional p–d hybridization interaction between the d orbital of Pd and p orbital of nonmetal elements can bring remarkable improvement of electrocatalytic properties of Pd.\textsuperscript{97} As reported by the Zhang group, Pd metallene doped with Se atoms (Pd–PdSe) could exhibit 6.6 and 2.6 fold enhancement in mass activity for the ethylene glycol oxidation reaction (EGOR) compared with those of Pd metallene and commercial Pd/C (Fig. 12C–C\textsubscript{2}).\textsuperscript{95} Both experimental results and theoretical calculations revealed the existence of strong p–d orbital hybridization and tensile strain in Pd–PdSe bimetallene, which could reduce the activation energy of C–C bond cleavage and accelerate CO\textsuperscript{*} oxidation and thus boost the complete oxidation of EG (Fig. 12C\textsubscript{3}).
4.4. Carbon dioxide reduction reaction

Converting CO$_2$ into chemical fuels, such as methane, methanol, formate, ethylene, and ethanol, is one of the most considerable strategies for addressing the global energy crisis. To date, both noble (Pd, Ag, Au) and non-noble metals (Co, Cu, Zn, Bi) have been involved in the catalysis of the CO$_2$RR. Remarkably, 2D non-noble metal-based electrocatalysts have attracted increasing research interest because of the maximized AUE and higher cost-effectiveness relative to noble metals. For instance, Xie and coworkers prepared partially oxidized Co metallene with four-atomic-layer thickness for highly selective electroreduction of CO$_2$ to formate (Fig. 13A). As demonstrated in Fig. 13A$_1$ and A$_2$, a stable current density of 10.59 mA cm$^{-2}$ and a high faradaic efficiency (FE) of 90.1% could be achieved at an overpotential of 0.24 V versus RHE, which were substantially remained over 40 h. The study suggests that regulation of the oxidation state of 2D metallenes plays a key role in improving the catalytic properties of the CO$_2$RR, offering a design principle for the engineering of active sites at an atomic scale. In another example, ultrathin bismuthene with an average thickness of 1.28–1.45 nm was prepared by Xu and coworkers. When used as an CO$_2$RR catalyst, a nearly 100% FE for formate and a partial current density up to 72 mA cm$^{-2}$ were obtained in a wide potential window over 0.35 V (Fig. 13B). To further evaluate the feasibility of bismuthene in energy-related devices, the catalyst was also applied in a self-designed flow cell and a bipolar membrane separated fuel cell (Fig. 13B$_2$). The bismuthene could deliver a steady current density of 100 and 200 mA cm$^{-2}$ at $-0.57$ and $-0.75$ V versus RHE, respectively, with FE close to 100%, meeting the requirements for commercial application. Meanwhile, a new pathway with the direct involvement of HCO$_3^-$ in the CO$_2$RR process for formate production was proposed based on in situ attenuated total reflection-infrared (ATR-IR) spectroscopy analysis combined with theoretical calculation (Fig. 13B$_3$).

4.5. Nitrogen reduction reaction

Production of ammonia by direct reduction of N$_2$ under ambient conditions becomes a promising alternative way relative to...
the conventional Haber–Bosch process in the industrial production of fertilizer. However, the high bond energy of the N≡N triple bond and the competing HER lead to an unsatisfactory faradaic efficiency (FE) of NRR for NH₃ production. Besides, the involved dissociative and associative mechanisms make the NRR process complicated and less understandable. Therefore, well-designed electrocatalysts are required to perform the experimental and theoretical studies and thus overcome these issues associated with the NRR. Among the catalysts reported, Pd-based nanomaterials have recently emerged as an excellent catalyst for the NRR with high activity and selectivity for NH₃ production. The NRR performance can be fundamentally improved by facilitating the adsorption and activation of N₂ on the surface of Pd, which can be realized by the intercalation of lattice hydrogen atoms as well as engineering lattice vacancies of Pd. As such, the synthesis of Pd-H metallene with surface vacancies serves as a promising strategy for the generation of a highly efficient NRR catalyst. As reported by Wang and coworkers, electrochemical conversion of N₂ to NH₃ can be performed and catalyzed by Pd-H metallene with Pd vacancies in a three-electrode H-cell under ambient conditions (Fig. 14A). The yield rate and FE of NH₃ at different applied potentials is shown in Fig. 14A, which can be well-maintained within 10 cycles (Fig. 14A). The NRR mechanism on the surface...
of Pd-H metallene was studied by in situ Raman spectroscopy, indicating an association pathway of the NRR as demonstrated in Fig. 14A3 and A4. Doping Pd with another NRR active material such as Fe, Co, and Cu is another effective way to improve the intrinsic activity and selectivity of Pd metallene. As shown in Fig. 14B and B1, theoretical simulations were conducted to demonstrate the enhanced N₂ adsorption preference to the Pd-coordinated isolated Fe atoms on the surface of PdFe₁ single atom alloy metallene.²⁵ The electron-deficient Fe atoms can act as a Lewis acid site to attract the lone pair electrons of Lewis base N₂ with a hole-rich region, suggesting the activated N₂ on PdFe₁ metallene. As shown in Fig. 14B₂, the PdFe₁ metallene delivered the highest NH₃ yield of 111.9 µg h⁻¹ mg⁻¹ and FE of 37.8% at −0.2 V, indicating the exceptional NRR activity and selectivity. The PdFe₁ metallene also exhibited excellent electrochemical stability with a steady...

**Fig. 15** Utilization of metallenes in photocatalysis. (A) NH₃ yield rates under UV-Vis or visible light irradiation over antimonene. (A₁) NH₃ yield rate of antimonene against recycling cycle. (B–B₂) Transient photocurrent responses of bismuthene for the CO₂RR at 0.4, 1.0, and 0.6 V versus RHE. (A and A₁) Reproduced from ref. 103 with permission from Elsevier. Copyright 2020. (B–B₂) Reproduced from ref. 104 with permission from American Chemical Society. Copyright 2021.
current density in the chronoamperometry test for 100 h continuous NRR electrolysis and no apparent degradation in NH₃ yields and FEs during the cycling tests (Fig. 14B₃).

4.6. Photocatalysis

Beyond the electrochemical reactions mentioned above, some metallenes also performed well in photocatalytic reactions. For example, Sun and coworkers reported the defect engineering of two-dimensional oxidized Sb metallene to activate intrinsically inert Sb for efficient visible light-driven NRR to produce NH₃.¹⁰³ Compared with bulk Sb and Sb₂O₃, the Sb metallene delivered substantially enhanced N₂ photofixation with remarkable NH₃ yield rates, likely resulting from the plentiful surface catalytic sites associated with atomic thickness and a high density of active surface defects to greatly facilitate charge carrier migration and separation (Fig. 15A). The defective Sb metallene maintained good stability and thus reusability without apparent decays in activity after five cycling tests (Fig. 15A₁). In another example of photocatalytic applications, Bi metallene served as an electron mediator in a 2D Z-scheme photocatalyst for the CO₂RR.¹⁰⁴ In the photoelectrochemical experiment, high cathode photocurrent densities were observed for the photocatalyst, indicating an enhanced charge utilization efficiency in either the oxidation or the reduction reaction (Fig. 15B and B₁). Moreover, no significant photocurrent change was observed within 2000 s at a constant bias of 0.6 V versus RHE, confirming the satisfactory photoelectrochemical stability of the catalyst (Fig. 15B₂).

5. Conclusions and perspectives

Taking advantage of the rapid advancement of synthetic techniques at the nanoscale, remarkable development has been made in the synthesis of metallenes. This minireview summarizes the recent progress in this domain, highlighting the synthesis, characterization, and design principles of metallenes with respect to electrocatalysis for energy conversion. Here we discuss some challenges and future research directions to motivate more significant works in this field (Fig. 16).

5.1. Precision synthesis control

Despite the successful synthesis of metallenes with different variations in terms of thickness, composition, lattice constants, and porosity, precise control over their synthesis at an atomic scale is still challenging due to the difficulty of high-level manipulation with the consideration of anisotropic growth of metals for maintaining the 2D morphology with atomic thickness. This precision in terms of synthesis control includes but is not limited to: (1) downscaling thickness to a new limit (sub-nanometer or even monolayer thickness) with a freestanding and stable structure, (2) generation of anisotropic strain over the surface, (3) precise elemental distribution in multielement metallenes, and (4) new crystal phases such as body-centered cubic (bcc) or orthorhombic structures with atomic thickness. In general, all the parameters associated with metallenes need to be precisely controlled and optimized in order to meet the requirements of electrocatalysis for various energy conversions.
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5.2. Characterization methods

Although great efforts have been made in the controlled synthesis of numerous metallenes by the bottom-up growth, the formation mechanism of these unique 2D metals still needs to be fully understood using both experimental tools and theoretical simulations. Given this, high-level characterization techniques are required for the in-depth investigation of the formation process from nucleation to anisotropic growth of metallenes. Thanks to the advancement of in situ characterization tools including in situ TEM, XRD, Raman, and XAS, different formation mechanisms will be revealed for the guidance and prediction of synthesis. Moreover, the delicate nature of metallenes typically leads to difficulties during characterization of the atomic structures by using conventional tools. For example, the high sensitivity of metallenes to the electron beams make them susceptible to falling apart when observed by TEM. This electron beam instability of metallenes needs to be taken into consideration especially when characterizing metallenes composed of solution alloys. In particular, to reveal the elemental distribution over the metallenes, their ultrathin nature requires an elongated time to collect a sufficient EDS signal, which leads to potentially aggravated electron beam-induced structural and even chemical deviation. This issue can be potentially addressed by using cryo-electron microscopy. In addition, the thermodynamic unfavorable structure is more susceptible to go through structural changes during their application in electrochemical and practical applications. Therefore, it is essentially important to develop new in situ characterization tools for observing these possible evolutions in different applications of metallenes.

5.3. Scalable production

One challenge for most energy-related applications is to produce metallenes in large quantities without the compromise of control over thickness, composition, surface facets, and porosity. In the bottom-up synthesis by wet-chemical approaches, a typical strategy for increasing the productivity is to increase the volume of the reaction mixture. For example, by increasing the reaction volume from 5 mL to over 200 mL, the yield of PdMo metallene can also be increased by 40 times with a tight control over various parameters. However, the intrinsic thermal and compositional inhomogeneity over a large volume makes it highly challenging to obtain metallenes with high quality when the reaction volume is further scaled up. Therefore, other alternative approaches need to be developed to ensure the anisotropic growth of metallenes. For example, continuous-flow reactors have been used to address the inhomogeneity issue associated with the reaction volume by dividing the reaction mixture into droplets, plugs, or flow with a millimeter scale diameter for maintaining the homogeneity of the reaction solution. This new technique is able to achieve well-controlled reaction parameters for the scalable generation of nanomaterials without compromising their quality.

5.4. Physical modeling

Physical modeling is fundamental for new structures to reveal their importance in various applications. However, relevant studies on metallene-related materials are still in the initial stage. As demonstrated in Fig. 1B, the elements involved in metallenes in the periodic table achieved by either experiments or theoretical calculations are still limited. So far, the expansion of theoretical modelling of 2D metal materials with atomic thickness is important to make this class of materials become more attractive to different communities for potential applications including electronics, optics, catalysis, and biomedicines.

5.5. Stability and realistic device applications

Although progress in the synthesis and utilization of metallenes as electrocatalysts has recently been made, a number of challenges still need to be addressed before this class of nanomaterials can be used in real applications. Indeed, many of the electrocatalytic applications were merely proof-of-concept and performed in a laboratory. The cost, processability, and scale-up feasibility of metallenes all need to be taken into account for applications in realistic devices. Additionally, most of the metallenes might lack long-term stability and contamination resistance in these practical applications, which can be potentially addressed by oriented assembly of this class of materials to make them robust and functional in realistic devices. Therefore, there is still a long way to go before this unique class of 2D metals can be used in real-life energy conversion, especially when the operation conditions are harsh.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

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Materials Horizons


