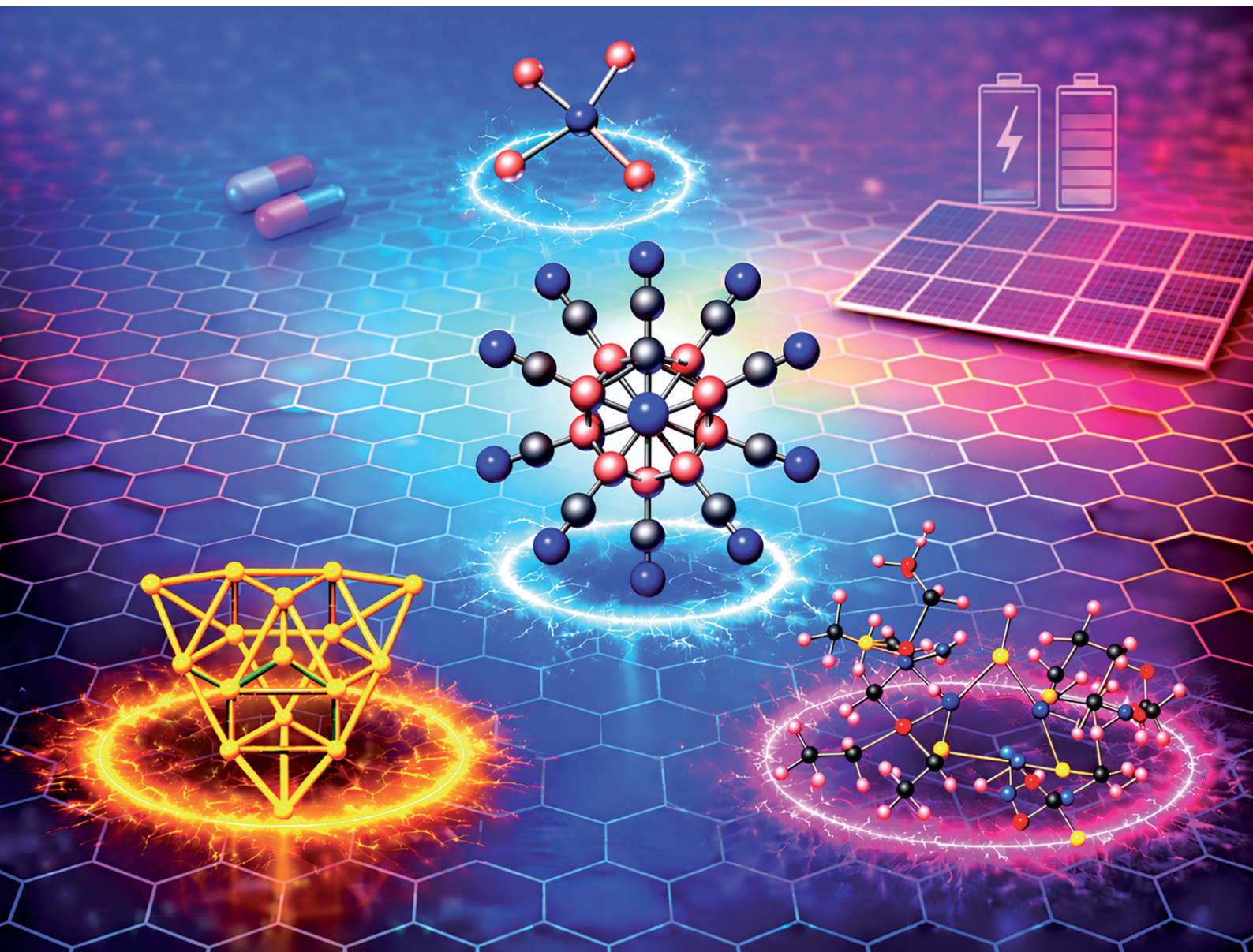


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## Cluster-based materials: design and applications

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Nanoclusters constitute an interdisciplinary field bridging chemistry, physics, biology, medicine, and materials science. Its broad appeal is fueled not only by the expectation that one can gain a fundamental understanding of how the structure and properties of matter evolve one atom at a time, but also, how they could be designed, with specific size and composition, to mimic the chemistry of atoms in the periodic table. Referred to as superatoms, these clusters can function as modular building units for a new generation of cluster-assembled materials, provided they retain their structural identity upon integration into extended architectures. C<sub>60</sub> fullerene is an example of such a cluster. Clusters, protected by ligands or supported on a substrate, can also form cluster-assembled materials with unique properties. In contrast, atoms in an existing crystal can be replaced by corresponding superatoms to form “cluster-based materials”. While several perspectives and review articles have been published to highlight the unique structure- and composition-specific properties of isolated clusters and cluster-assembled materials, it is during the past few years that attention has been focused on cluster-based materials. This review focuses on the recent developments in the design, synthesis, and applications of clusters with uncommon properties and highlights their potential as building blocks of cluster-based materials. The topics include isolated clusters enabling unusual reactions, their role in the design and synthesis of materials such as super-electrides, solid state electrolytes with fast ionic conductivity and stable interfaces, moisture resistant hybrid perovskite solar cells, thermoelectric materials with high figure of merit, and a new class of single-superatom catalysts. Also outlined are the challenges and opportunities going forward.

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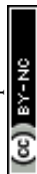
*systems as well as behaviors of ion transport and kinetics at the interfaces. He has over 60 peer-reviewed publications in the related fields.*



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*theoretical prediction of stable ferromagnetism in CrI<sub>3</sub> monolayer, later confirmed experimentally, initiating the field of 2D magnetism. He was elected an IAAM Fellow in 2023 and was recognized among the top 0.1% Highly Cited Scientists in 2025.*



## I. Introduction

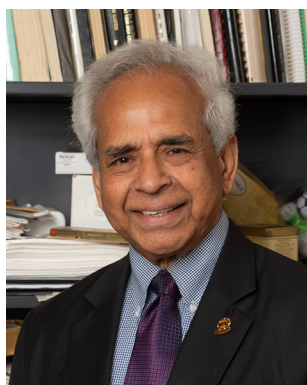
The periodic table of elements is the foundation of all materials, however, its scope is constrained by the finite number of naturally occurring elements, many of which are rare or pose environmental and toxicity concerns. The Alchemists' dream to replace gold with base metals never materialized because the chemistry of gold is pre-determined by its atomic orbitals that cannot be altered. Fortunately, advancement in new experimental and theoretical techniques over the past few decades have enabled researchers to manipulate and control properties of matter at the nanoscale by tailoring their size, shape, and composition and producing materials with unique properties. However, at this length scale, properties are not influenced by the addition or removal of a single atom.

Atomic clusters, defined as an aggregate of atoms with finite size and composition, bridge not only this gap but also their properties can be changed by the addition or removal of a single atom or even a single electron.<sup>1–15</sup> With developments in experimental techniques, one can now produce clusters of most elements with varying size and composition in the gas phase.<sup>16–22</sup> Soft-landing of mass-isolated clusters on substrates<sup>23–25</sup> as well as those assisted with ligands<sup>26–34</sup> or guided by molecular templates<sup>35–38</sup> have made it possible to synthesize cluster-assembled materials with unique properties. These experimental techniques combined with high-throughput calculations based on the density functional theory have enabled researchers to study the structure and properties of homo- and hetero-atomic clusters containing many atoms.<sup>39–42</sup> The early motivation in studying clusters as a function of size was to understand how the properties of matter evolve, one atom at a time.<sup>43–45</sup> While some understanding has been achieved in this regard, the answer to “when a cluster becomes a crystal” remains largely unanswered. This is because different properties evolve differently, making it difficult to find a unique answer. For example, a recent study showed that five

H<sub>2</sub>O molecules can make a water drop,<sup>46,47</sup> but five gold atoms do not make an inert gold. However, past studies have revealed many unusual size- and composition-specific properties of clusters. These include clusters mimicking the properties of atoms,<sup>5</sup> otherwise nonmagnetic elements becoming magnetic,<sup>48</sup> chemically inert elements becoming reactive,<sup>49</sup> atoms achieving oxidation states higher than nature intended,<sup>50</sup> clusters with like charges attracting,<sup>51</sup> noble gas atoms reacting under ambient conditions,<sup>52</sup> *etc.* These findings have fueled much interest in cluster science.<sup>53–56</sup>

One of the unique properties of clusters this review focuses on is clusters as “superatoms” and their role in the design and synthesis of cluster-based materials. The concept of superatoms originated from the pioneering work of Khanna and Jena,<sup>57,58</sup> who demonstrated that atomic clusters with specific size and composition can emulate the chemical properties of elements in the periodic table. This idea was motivated by two key experiments: (1) the observation of magic numbers in sodium clusters by Walter Knight and coworkers who explained their unusual stability using the jellium model to be due to electronic shell closure (*e.g.*, 1S, 1P, 1D).<sup>59</sup> (2) The observation by Will Castleman and co-workers<sup>60</sup> that the Al<sub>13</sub><sup>−</sup> cluster is highly stable and chemically inert towards oxygen due to atomic and electronic shell closure. Building on these findings, Jena and Khanna proposed that nearly spherical clusters with delocalized electrons filling superatomic orbitals according to an Aufbau principle could be designed to mimic atomic behavior. They further suggested that such clusters could serve as building blocks for cluster-assembled materials with uncommon properties, provided their structural and electronic integrity is retained upon assembly. Studies of superatoms have increased over the past 30 years as evidenced by the growing number of publications and their unusual properties (see Fig. 1).

The superatom concept has since expanded to include superalkalis and superhalogens, first proposed by Gutsev and Boldyrev in 1980's.<sup>61,62</sup> Superalkalis (*e.g.*, Li<sub>2</sub>F) have ionization potentials lower than that of alkali metals, while superhalogens (*e.g.*, LiF<sub>2</sub>) have electron affinities exceeding those of halogens. Although these clusters are not spherical and do not exhibit superatomic orbitals, they do mimic the chemistry of alkali and halogen atoms, respectively. Some ions such as BH<sub>4</sub><sup>−</sup>, CN<sup>−</sup>, CB<sub>11</sub>H<sub>12</sub><sup>−</sup>, traditionally known in chemistry as polyatomic ions or complex ions, also exhibit superhalogen properties and have been used to substitute halogens, yielding materials with uncommon properties. In the past few years, numerous superhalogens and superalkalis have been designed using the octet rule,<sup>63,64</sup> 18-electron rule,<sup>65,66</sup> 32-electron rule,<sup>67,68</sup> Hückel's rule,<sup>69,70</sup> and Wade's rule.<sup>71–74</sup> Some of these have been synthesized, forming novel materials. In addition, core-shell structures such as ligated metal chalcogenides<sup>75–78</sup> and thiolated noble metal clusters<sup>26–34,79–83</sup> have been classified as superatoms as they are nearly spherical and do exhibit superatomic orbitals, even though they may not mimic the chemistry of a particular atom in the periodic table. The bonding in these clusters has been characterized as concentric bonding in

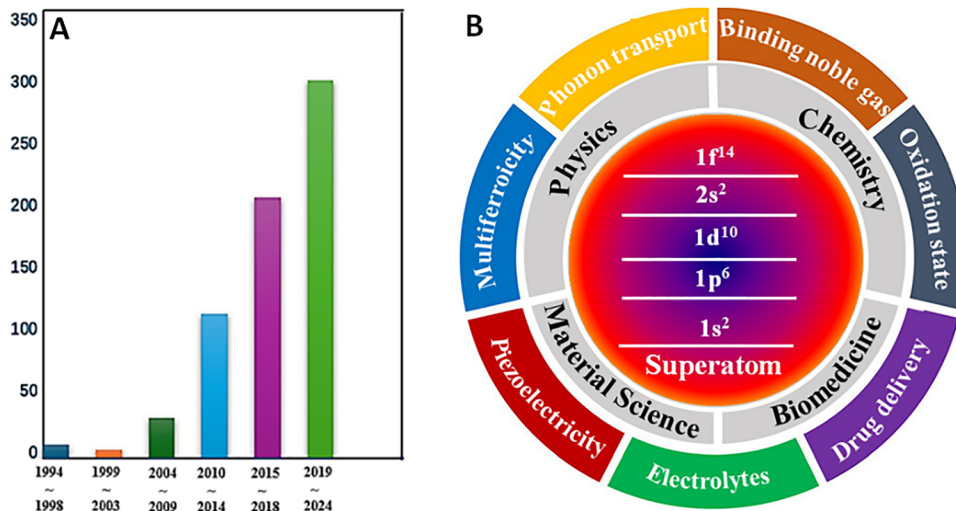


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**Fig. 1** (A) Increasing number of papers containing the key word of superatoms either in the title or in the abstract published the past 30 years. (B) Works on isolated clusters and cluster-assembled materials bridging physics, chemistry, materials science, biology, and medicine, and enriching our understanding of the physical and chemical properties of matter at the nano-scale.

superatoms.<sup>84</sup> This refers to a unique bonding pattern where atoms within a cluster arrange in nested spherical layers, resembling the electronic shell structures of noble gas atoms. In addition to the ligated clusters, some other examples of concentric bonding in metallic clusters include Au<sub>13</sub>, Au<sub>18</sub>, Pd<sub>6</sub>O<sub>4</sub>, and a cage-like boron cluster such as B<sub>12</sub>. Inspired by the quark model, a grand unified model for ligated Au clusters has also been proposed that accounts for the structure of 71 such clusters.<sup>85</sup> Regardless of whether the above species are termed superatoms, polyatomic ions, or complex ions, their fundamental role as building blocks for new materials remains unchanged. To quote William Shakespeare, “a rose by any other name would smell as sweet”. The ligated clusters have rich chemistry and have been used to synthesize materials with unique properties.

In Table 1, we summarize the key reasons why superatomic clusters, instead of atoms whose chemistry they mimic, can serve as transformative building blocks for constructing novel materials. However, atomic clusters produced in the gas phase under inert condition are typically unstable once removed from

that environment. Upon exposure to ambient conditions, they tend to react with surrounding gases and/or coalesce with neighboring clusters, thereby losing their individual identity, structure, and properties. To retain their identity, mass-isolated clusters could be soft-landed on substrates,<sup>23–25</sup> protected by ligands,<sup>26–34</sup> or guided by molecular templates.<sup>35–38</sup> However, in these cases, interaction of clusters with the substrates or ligands must be considered which can also be used for further functionalization of the material properties. Another way to use clusters as a building block is to replace an atom in an existing material with a corresponding superatomic cluster. We refer to such materials as “cluster-based materials,” which forms the focus of this review. Given that several recent reviews have already comprehensively addressed core metal clusters protected by complex ligands (including metal chalcogenide clusters, Au/Ag nanoclusters, and mass-isolated clusters on substrates), we do not discuss these topics further here due to space constraints.<sup>26–34,75–83,86–104</sup>

Although several review articles and perspectives on superatoms have been published,<sup>53–56,105</sup> this review focuses primarily

**Table 1** Key advantages of superatomic clusters, instead of atoms, serving as the building block of materials

| Atoms as building blocks  | Superatoms as building blocks   |
|---|---|
| Number of atoms is fixed.   | Number of superatoms is unlimited.  |
| Properties of atoms are governed by their atomic orbitals which cannot be changed.        | Molecular orbitals of superatoms mimicking the chemistry of an atom can be changed by varying their size and composition.   |
| Ionization potentials of alkali atoms and electron affinities of halogen atoms are fixed. | Superalkalis are characterized by ionization potentials lower than those of alkali metals, and superhalogens display electron affinities surpassing those of halogens. Such extreme electronic properties are not fixed; rather, they can be engineered by tailoring the cluster's size and chemical composition. |
| Most atoms cannot accommodate more than one extra electron.                               | Superatoms can accommodate multiple electrons.  |
| In atom-assembled solids, the phonons are due to the vibration of atoms.                  | In cluster-based solids, phonons are due to intra-cluster as well as inter-cluster vibrations, thus fundamentally altering electron–phonon coupling.  |
| Lattice constants and lattice symmetries are fixed.                                       | Lattice constants and symmetries can be varied with size and composition.   |
| Energy bands are formed by overlap of atomic orbitals.                                    | Energy bands are formed by overlap of superatomic orbitals.   |



Table 2 Electron counting rules for designing superatoms: principles, conditions, and examples<sup>ab</sup>

| Rule             | Principle  | Conditions  | Best for   | Properties   | Examples  | Notes   |
|------------------|--|---|--|--|---|---|
| Jellium rule     | Stability arises from closed-shell electronic configurations in a spherical potential well: 2, 8, 18, 20, 34, 40, 58, 92... electrons.   | <ul style="list-style-type: none"> <li>Delocalized valence electrons</li> <li>Nearly spherical geometry</li> <li>Uniform positive charge background</li> </ul>                  | Alkali, noble metal clusters (free or ligand-protected)        | <ul style="list-style-type: none"> <li>Ionization energies peak at magic numbers</li> <li>Delocalized metallic bonding</li> </ul>  | Na <sub>8</sub> , Na <sub>20</sub> , Mg <sub>40</sub> , Al <sub>13</sub> <sup>-</sup> , Au <sub>25</sub> (SR) <sub>18</sub> <sup>-</sup> , Ag <sub>14</sub> (SR) <sub>30</sub> , Au <sub>102</sub> (SR) <sub>44</sub> , Au <sub>144</sub> (SR) <sub>60</sub> , SiAl <sub>14</sub> (C <sub>5</sub> Me <sub>5</sub> ) <sub>6</sub>  | The count of 'free' or delocalized electrons: $n = NV_n - YV_X + sV_L - z$ , where $N$ is the number of metal atoms (M); $S$ is the number of charge-donating ligands (L); $Y$ is the number of charge-withdrawing ligands (X); $Z$ is the overall charge; $V_M$ , $V_X$ and $V_L$ are the valence numbers of the metal and the ligands. <sup>c</sup> |
| Wade's rule      | Polyhedral skeletal structures require specific numbers of bonding electron pairs (NBEP) based on cluster geometry ( <i>closo</i> , <i>nido</i> , <i>arachno</i> , <i>hypho</i> ). | <ul style="list-style-type: none"> <li>Polyhedral or cage-like clusters</li> <li>Mostly applies to boranes, metallo boranes, and ligand-stabilized metal clusters</li> </ul>    | Boranes, carboranes, metallo-boranes ligand-protected clusters | <ul style="list-style-type: none"> <li>NBEP = <math>n + 1</math>: <i>closo</i> (complete polyhedron)</li> <li>NBEP = <math>n + 2</math>: <i>nido</i> (one missing vertex from <i>closo</i>)</li> <li>NBEP = <math>n + 3</math>: <i>arachno</i> (two missing vertices)</li> <li>NBEP = <math>n + 4</math>: <i>hypho</i> (three missing vertices)</li> </ul> | B <sub>6</sub> H <sub>6</sub> <sup>2-</sup> , C <sub>2</sub> B <sub>10</sub> H <sub>12</sub> , Fe <sub>3</sub> (CO) <sub>12</sub> , Au <sub>25</sub> (SR) <sub>18</sub> <sup>-</sup> , Ag <sub>25</sub> (SR) <sub>18</sub> <sup>-</sup> Al <sub>2</sub> C <sup>-</sup> , [Pd <sub>6</sub> (CO) <sub>12</sub> ] <sup>2+</sup> , Al <sub>4</sub> H <sub>6</sub> <sup>6-</sup> , Al <sub>4</sub> H <sub>4</sub> <sup>2-</sup> , Sn <sub>12</sub> <sup>+</sup> , Pb <sub>12</sub> <sup>2-</sup> , AlPb <sub>12</sub> <sup>+</sup> | <ul style="list-style-type: none"> <li>divide remaining electrons by 2 to get skeletal electron pairs.</li> <li>Match to NBEP formula to determine geometry type.</li> </ul>  |
| Octet rule       | Main-group atoms are most stable with 8 valence electrons (full s and p orbitals).   | <ul style="list-style-type: none"> <li>Small main-group clusters</li> <li>Atoms with accessible s and p orbitals</li> <li>No significant d-orbital participation</li> </ul>     | Main-group anions, superhalogens, molecular superclusters      | Superhalogens and superalkalis   | ZrO, BO <sub>2</sub> <sup>-</sup> , AlO <sub>2</sub> <sup>-</sup> , BH <sub>4</sub> <sup>-</sup> , AlF <sub>4</sub> <sup>-</sup> , BF <sub>4</sub> <sup>-</sup> , MnO <sub>4</sub> <sup>-</sup>   | Octet-rule superatoms often resemble covalent or ionic molecular systems with localized bonding.  |
| 18-Electron rule | Maximum stability occurs when a transition metal atom or cluster has 18 valence electrons (full s, p, d shells).   | <ul style="list-style-type: none"> <li>Transition metal center</li> <li>Ligand field strong enough to define orbital splitting</li> <li>No strong electron repulsion</li> </ul> | Organometallic complexes, transition metal-centered superatoms | Transition metal based 18-e superatom can be magnetic such as Mn@Si <sub>12</sub> and Cr@Si <sub>12</sub>  | Fe(CO) <sub>5</sub> , Ni(CO) <sub>4</sub> , Mo(CO) <sub>6</sub> , V(CO) <sub>6</sub> , Ru <sub>3</sub> (CO) <sub>12</sub>   | Total valence electrons = metal valence electrons + ligand electrons ± charge.  |
| 32-Electron rule | 32 valence electrons occupying cluster orbitals with spherical harmonics character, the cluster is stable with a closed-shell electronic configuration                             | <ul style="list-style-type: none"> <li>Spherical or near-spherical symmetry.</li> <li>Electrons are delocalized over the whole cluster</li> </ul>                               | 5f-Element encapsulated cages                                  | Can be used to mimic xenon or a heavy element analog.  | Pu@Pb <sub>12</sub> , Ge@Sn <sub>12</sub> <sup>-</sup> , Pu@C <sub>24</sub> An@C <sub>28</sub> (An = Th, Pa, U, Pu <sup>4+</sup> ), CBe <sub>8</sub> H <sub>12</sub> , U@B <sub>40</sub>  | <ul style="list-style-type: none"> <li>The 5f orbitals in actinides are chemically accessible, radially extended, and capable of covalent bonding, making them uniquely well-suited for stabilizing 32-electron clusters.</li> <li>Both the central atom's electrons and ligand/cage contributions are included in the counting.</li> </ul>           |



Table 2 (continued)

| Rule          | Principle   | Conditions   | Best for   | Properties   | Examples  | Notes   |
|---------------|---|--|--|--|---|---|
| Hückel's rule | Aromatic stability in planar, cyclic systems with $(4n + 2)$ $\pi$ -electrons   | <ul style="list-style-type: none"> <li>Planar, cyclic geometry</li> <li>Conjugated <math>\pi</math>-system</li> <li>Electron delocalization over ring</li> </ul> | $\pi$ -Conjugated rings, metal Delocalized 2D aromaticity rings, small planar clusters |  | Benzene, $C_6H_4^{2-}$ , $B_6^{2-}$ , $Al_4^{2-}$ , $LiAl_4$ , $Na_3^+$ , $Be_3^-$ , $Cu_4^{2+}$ , $Ag_4^+$ | Some B clusters are aromatic with $4n$ electrons due to multi-center bonding.       |
| Hirsch's rule | A 3D superatom is aromatic containing $2(n + 1)^2$ delocalized $\pi$ -electrons | Spherical symmetry, delocalized $\pi$ -electrons   | Spherical delocalization with correct number of $\pi$ -electrons                       | Spherical electron cloud distributes electronic charge evenly across the structure to minimize repulsion | $C_{20}^{2+}$ , $C_{60}^{10+}$ , $[Sn_{12}]^{2-}$ , $Au_{13}^{5+}$  | Often applies to charged species or zintl ions where $\pi$ -electrons can be tuned. |

<sup>a</sup> No single rule universally applies; a hybrid approach sometimes is needed due to complexity of bonding and compositions (transition metals, main-group elements, and ligands). <sup>b</sup> Design rules are guidelines not absolutes, real superatoms can defy them due to factors such as geometry, ligand effects, relativistic stabilization, and electronic correlation. <sup>c</sup> Ref. 79.

on the developments over the past 5–10 years and illustrates how superatoms can promote unusual reactions enabling noble gas atoms to form chemical bonds at room temperature, and lead to unconventional electrides, solid state electrolytes with fast ionic conductivity and stable interfaces, moisture resistant hybrid perovskite solar cells, thermoelectric materials with high figure of merit, and superior catalysts. Also discussed are challenges and opportunities moving forward.

In Section II, we briefly outline the design rules and synthesis techniques of superatoms. Their unusual chemical and physical properties are described in Sections III and IV, respectively. Section V deals with the applications of cluster-based materials in energy production and storage, focusing on light absorbers in solar cells and solid-state electrolytes in metal-ion batteries. Superatoms with unusual ferroelectric, piezoelectric and thermoelectric properties are discussed in Section VI. A summary of our conclusions and the outlook focusing on challenges and opportunities are presented in Section VII.

## II. Design rules and synthesis of superatoms and cluster-based materials

In this section, we discuss briefly how the superatomic clusters and cluster-based materials are theoretically designed and experimentally synthesized. In Table 2, we summarize the various electron counting rules used to design superatoms. The design rule that was initially used exclusively for simple metals is the jellium rule, originally used to describe the magic numbers in Na clusters.<sup>59</sup> A good example of this design is the  $Al_{13}$  cluster.<sup>60</sup> With 39 valence electrons,  $Al_{13}$  requires only one electron to complete its 1S 1P 1D 2S 1F 2P electronic shells. This is similar to halogens, which also require one electron to complete their outermost np shell. Hence,  $Al_{13}$  behaves like a halogen atom. Indeed, the electron affinity of  $Al_{13}$  cluster, namely, 3.6 eV, is very close to that of the chlorine atom. Later, the octet,<sup>63,64</sup> 18-electron,<sup>65,66</sup> and 32-electron rules<sup>67,68</sup> have been used to design superatoms composed of low-atomic-weight atoms, transition metal atoms, and rare-earth atoms, respectively. Similarly, Wade's rule<sup>71–74</sup> and Hückel's and Hirsch's rule<sup>69,70,106</sup> have been used for boranes and aromatic molecules, respectively.<sup>107,108</sup> Several superatoms mimicking the chemistry of halogens, chalcogens, pnictogens, and alkalis, as well as transition metals, have been designed and will be discussed in the following sections. Unlike in atoms, multiple electrons counting rules can be simultaneously used to create superatoms with uncommon properties. An example of such a system is  $B_{12}(CN)_{12}$ , where both the Wade's rule and the octet rule are simultaneously used to create an extraordinarily stable dianion. This will be discussed at length in the forthcoming sections.

In Table 3 we summarize the different experimental techniques used for the synthesis of superatoms. In the gas phase, superatoms can be formed by using gas evaporation, laser vaporization, or sputtering techniques.<sup>16–25</sup> Metal-

**Table 3** Synthesis of superatomic clusters. In the table, SPhMe<sub>2</sub> = 2,4-dimethylbenzenethiolate, PET<sub>3</sub> = triethylphosphine, SR = thiolate, Cp\* = pentamethylcyclopentadienyl, and PET = phenylethanethiol

| Methods   | Features   | Examples <sup>16–38,109–121</sup>  |
|---|--|--|
| Gas-phase synthesis of isolated clusters        | Providing high precision in producing atomically precise superatomic clusters but being impractical for large-scale production.  | Al <sub>13</sub> , Na <sub>8</sub> , Na <sub>20</sub> , Al <sub>13</sub> <sup>–</sup> , Au <sub>20</sub> , W@Au <sub>12</sub> , Nb <sub>11</sub> O <sub>15</sub> <sup>–</sup>  |
| Supported clusters aided by gas phase synthesis |  | M@Si <sub>16</sub> (M = Sc, Ti, V, Zr, Ta, Hf, Lu, W)  |
| Photo-induced fusion reaction in solution       | Stabilizing atomically precise superatomic clusters by ligands and fusion, but this technique is valid for specific elements and sizes.  | M@Au <sub>12</sub> (M = Pd, Pt)  |
| Solution-based synthesis of ligated clusters    |  | Au <sub>25</sub> (SR) <sub>18</sub> , PdAu <sub>24</sub> (SR) <sub>18</sub> , PtAu <sub>24</sub> (SR) <sub>18</sub> , PtAg <sub>42</sub> , Co <sub>6</sub> Se <sub>8</sub> (-PET <sub>3</sub> ) <sub>6</sub> , Co <sub>6</sub> S <sub>8</sub> (PET <sub>3</sub> ) <sub>6</sub> , PtAg <sub>18</sub> , MAG <sub>24</sub> (SR) <sub>18</sub> (M = Ni, Pd, Pt), [IrHAg <sub>24</sub> (SPhMe <sub>2</sub> ) <sub>18</sub> ] <sup>2–</sup> , [OsH <sub>2</sub> Ag <sub>24</sub> (SPhMe <sub>2</sub> ) <sub>18</sub> ] <sup>2–</sup> , [RuH <sub>2</sub> Ag <sub>24</sub> (SPhMe <sub>2</sub> ) <sub>18</sub> ] <sup>2–</sup> , [Au <sub>6</sub> Al <sub>6</sub> ](Cp*) <sub>6</sub> , [HAu <sub>7</sub> Al <sub>6</sub> ](Cp*) <sub>6</sub> , Al <sub>13</sub> <sup>–</sup> , PtBr <sub>4</sub> , FeSn <sub>12</sub> , Au <sub>25</sub> PET <sub>18</sub> |
| Molecular template-guided synthesis             | Employing molecular templates to direct the assembly of atoms into well-defined superatomic structures with precise atomic control, but limited heavily by the availability and design of suitable molecular templates |  |

encapsulated clusters such as M@Si<sub>16</sub> (M = Sc, Ti, V, Zr, Ta, Hf, Lu, W) generated in the gas phase can be soft landed on substrates such as C<sub>60</sub>, preserving the cage structure of 16 Si atoms uniformly distributed around the central metal atom.<sup>24</sup> Substrates can also be used to further tune the electronic and chemical properties of the deposited superatoms through cluster-surface interaction. For instance, a p-type organic substrate of hexa-*tert*-butyl-hexa-*peri*-hexabenzocoronene (HBHBC, C<sub>66</sub>H<sub>66</sub>) substrates is employed for the deposition of halogen-like superatom Lu@Si<sub>16</sub>, where the substrate can donate one electron to the superatom to close its electron shell.<sup>24</sup> Similarly, a W@Si<sub>16</sub> superatom, when deposited on an n-type substrate C<sub>60</sub>, is charge positive, while it is negatively charged with enhanced oxidation reactivity when deposited on a p-type HBHBC substrate.<sup>109</sup> In addition, ligated superatomic M<sub>2</sub>@Au<sub>17</sub> (M = Pd, Pt) clusters have been synthesized *via* a novel photoinduced fusion of M@Au<sub>12</sub> superatoms triggered by photo-irradiation under visible light.<sup>110</sup> The unprecedented M<sub>2</sub>@Au<sub>17</sub> superatom has a structure of two M@Au<sub>12</sub> cores sharing Au<sub>5</sub> facets in the neutral form and exhibit an electronic configuration like that of the N<sub>2</sub> molecule.<sup>110</sup>

Ligand-protected nanoclusters with Au/Ag/Cu/Al metal cores constitute a large family of superatoms, where the metal core is electronically stabilized by ligands against aggregation in solution or in solid state. A well-known example is [Au<sub>25</sub>(SR)<sub>18</sub>]<sup>–</sup> (SR = thiolate), which exhibits an icosahedral Au<sub>13</sub> core protected by six Au<sub>2</sub>(SR)<sub>3</sub> ligands. There are many ligated metalloid gold clusters with different core sizes, such as Au<sub>13</sub>, Au<sub>20</sub>, Au<sub>23</sub>, Au<sub>24</sub>, Au<sub>25</sub>, Au<sub>26</sub>, Au<sub>30</sub>, Au<sub>36</sub>, Au<sub>38</sub>, Au<sub>55</sub>, Au<sub>60</sub>, Au<sub>99</sub>, Au<sub>102</sub>, Au<sub>103</sub>, Au<sub>133</sub>, and Au<sub>144</sub>.<sup>111</sup> Among these, larger clusters can be converted to smaller ones by thermochemical etching with exogenous thiols, while smaller-to-larger conversion can be realized using fusion reactions in solutions.<sup>93</sup> In general, three methods are used to synthesize ligated Au clusters, including reduction of gold precursors, conversion between different-sized clusters, and metallic Au evaporation.<sup>82</sup> Other superatomic clusters can also be prepared by doping heterometals into the existing ligand-protected Au and Ag clusters. For example, a set of diphosphine-protected MAu<sub>12</sub> (M = Ru, Rh, Ir) clusters have

been synthesized by doping d-shell metals in the Au<sub>13</sub> cluster,<sup>112</sup> with new synthetic methods to make atomically precise open sites by controlled removal of ligands.<sup>30</sup> Similarly, d-metal doped Ag clusters have been synthesized, including MAG<sub>24</sub>(SR)<sub>18</sub> (M = Ni, Pd, Pt), [RhH@Ag<sub>24</sub>(SPhMe<sub>2</sub>)<sub>18</sub>]<sup>2–</sup>, [IrHAg<sub>24</sub>(SPhMe<sub>2</sub>)<sub>18</sub>]<sup>2–</sup>, [OsH<sub>2</sub>Ag<sub>24</sub>(SPhMe<sub>2</sub>)<sub>18</sub>]<sup>2–</sup>, and [RuH<sub>2</sub>Ag<sub>24</sub>(SPhMe<sub>2</sub>)<sub>18</sub>]<sup>2–</sup> (SPhMe<sub>2</sub> = 2,4-dimethylbenzenethiolate).<sup>113–115</sup> Most recently, new hydrocarbon ligated heteronuclear gold clusters with p-metals (*e.g.*, Al), such as [Au<sub>6</sub>Al<sub>6</sub>](Cp\*)<sub>6</sub> and [HAu<sub>7</sub>Al<sub>6</sub>](Cp\*)<sub>6</sub>, have also been synthesized by using various phosphines as additives on the reduction of the Au precursors.<sup>116</sup>

Another large family of superatoms is based on ligated metal chalcogenide clusters, M<sub>6</sub>E<sub>8</sub> (M = metal, E = chalcogen), where the metal atoms are arranged on an octahedron and are contained in a cube of chalcogen atoms. Each metal atom in these clusters is coordinated by ligands L (*e.g.*, phosphines), forming M<sub>6</sub>E<sub>8</sub>L<sub>6</sub>. These superatoms are typically synthesized by combining a precursor of a metal source and phosphine chalcogenide in solutions.<sup>77</sup> By mixing such clusters of similar size, shape, and chemical properties, *e.g.*, Co<sub>6</sub>Se<sub>8</sub>(PET<sub>3</sub>)<sub>6</sub> and Cr<sub>6</sub>Te<sub>8</sub>(PET<sub>3</sub>)<sub>6</sub> (PET<sub>3</sub> = triethylphosphine), solid solutions at the cluster level and the corresponding [cluster]<sub>2</sub>[C<sub>60</sub>]<sub>2</sub> crystals can be synthesized with novel properties.<sup>28</sup> Controlling the ligand coordination spheres of the metal chalcogenide clusters in the solution phase can lead to novel cluster fusion and materials, allowing rational control over the synthetic process beyond traditional photolytic or thermal methods.<sup>27</sup> For example, carbene and multi-carbene adducts have been used to selectively modify the ligands of [Co<sub>6</sub>Se<sub>8</sub>L<sub>6</sub>] (L = PET<sub>3</sub>, CO, C(H)SiMe<sub>3</sub>) clusters to tailor the cluster fusion reactivity and the shape of the cluster core, as well as create site-specific functionalized dimers and unprecedented cyanide/pyridine adducts that lead to novel solids.<sup>27</sup>

Atom-by-atom substitution in the metal core of the metal chalcogenide clusters has also proven to be a viable strategy for the design of new superatoms with tailored electronic, magnetic, and optical properties. For example, the stability, magnetic, and optical properties of cobalt sulfide superatom



$[\text{Co}_6\text{S}_8\text{L}_6]^+$  ligated with triethylphosphine ( $\text{L} = \text{PEt}_3$ ) can be controlled by substituting Fe or Ni for Co in the core.<sup>78</sup> Interestingly, it is found most recently that, by using  $\text{NiCl}_2$  as a precursor in the solution-phase synthesis, nickel sulfide clusters with a  $\text{Ni}_3\text{S}_3$  core (with three Ni atoms arranged in a planar triangle and three bridging S atoms residing between Ni) are preferentially formed, instead of having the regular metal chalcogenide  $\text{Ni}_6\text{S}_8$  core. The subsequently protonated  $\text{Ni}_3\text{S}_3\text{H}(\text{PEt}_3)_5$  exhibits strong ferromagnetic ordering and coercivity.<sup>117</sup>

Solution-phase synthesis of superatomic cluster using dendrimer templates have become another attractive method, since the size of the generated clusters can be controlled in the state of the precursors.<sup>38,118–121</sup> Compared to the conventional methods, such a template-guided method can support short-time synthesis, circumventing special conditions, such as low temperature or complex reactions. For example, the thiol-protected  $\text{Au}_{25}$ ,  $\text{PdAu}_{24}$ , and  $\text{PtAu}_{24}$  superatoms can be facily synthesized using  $\text{AuCl}_3$  and a phenylazomethine dendrimer template with a tetraphenyl methane core part.<sup>38</sup> Most recently, the  $\text{FeSn}_{12}$  superatomic cluster that had only been synthesized in the gas phase before was synthesized in solution using  $\text{FeCl}_3$  and  $\text{SnCl}_2$  on the dendrimer templates.<sup>37</sup>

### III. Unusual chemical properties of superatoms

The chemistry of atoms is governed primarily by the valence electrons which are fixed. On the other hand, the chemistry of the superatoms is governed by their molecular orbitals which can be tailored by their size and composition, hence enabling superatoms to exhibit rich chemistry. We demonstrate this by focusing on how a multiply charged superatom can be stable while the atom whose chemistry it mimics cannot hold more than one excess electron. Such superatoms can usher unusual reactions by making noble gas atoms to form chemical bonds at room temperature. Similarly, superalkalis can be used as building blocks of super electrides and a single superatom

composed of earth-abundant materials can outperform single-atom catalysts such Pd.

#### III.1 Stabilizing multiply charged anions

Although multiply charged anions, protected by charge compensating cations, can be stable in matrices or in solvents, isolated atoms cannot hold more than one electron. On the contrary, an isolated superatom can accommodate more than one electron if its binding energy is larger than the repulsive energy between the additional electrons. An intriguing question always has been: what is the smallest superatom that can be stable as a dianion? A classic example of a stable dianion is dodecaborane,  $\text{B}_{12}\text{H}_{12}^{2-}$ , whose second electron is bound by 0.9 eV, although  $\text{C}_{60}^{2-}$ , which is larger in size, turns out to be unstable. Further, the second electron affinity of  $\text{B}_{12}\text{H}_{12}^{2-}$  can be systematically increased by replacing H with halogen atoms<sup>122</sup> and can reach as high as 5.28 eV when H is replaced by CN.<sup>52,123</sup> This highlights the importance of the electronic structure of the parent cluster as well as those of the ligands, suggesting that it may be possible to increase the stability of a multiply charged cluster by tailoring its size and composition. For example, it has been shown theoretically that by replacing a B atom by Be in  $\text{B}_{12}(\text{CN})_{12}^{2-}$  cluster, the resulting  $\text{BeB}_{11}(\text{CN})_{12}^{3-}$  trianion can be stabilized.<sup>124</sup> This is consistent with the zintl-Klemm pseudo-element concept, given that  $\text{Be}^-$  is isoelectronic with B.<sup>125</sup>

Fang and Jena established a general model to design clusters that can be stable while accommodating as many as 5 extra electrons.<sup>126,127</sup> They started with the underlying principle that in stabilizing a multiply charged anionic cluster in the gas phase, the excess electrons must occupy a series of bound states within a 3D potential well, whose depth can be approximated by the total energy required to reduce the highest negatively charged state to neutrality—that is, the cumulative sum of the electron affinities ( $\text{EA}_n$ ) associated with each added electron, while the width of the potential well can be measured by the radius of a cluster defined as half of its longest dimension, per Fig. 2A. The general trend is that a more negative state

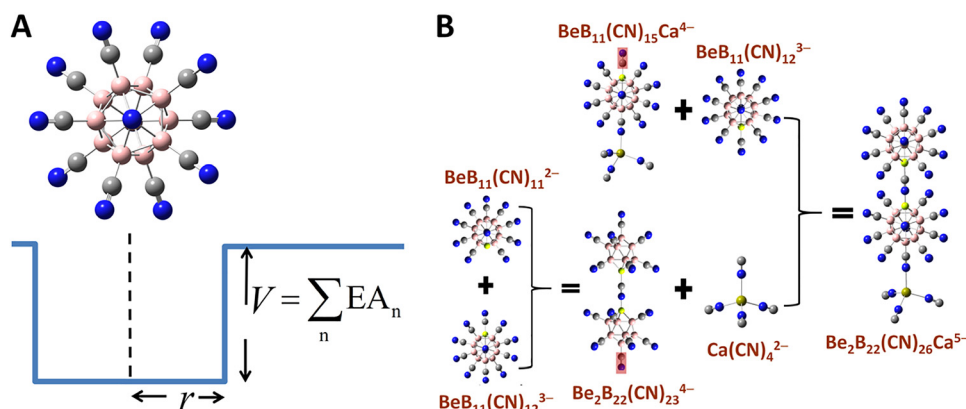


Fig. 2 Systematic approach to design stable multiply charged clusters in the gas phase. (A) The general potential well model of a stable cluster with its dimension corresponding to the radius ( $R$ ) and the sum of electron affinities ( $\text{EA}_n$ ) of the cluster.<sup>126</sup> (B) Stable tetra- and penta-anions created by assembling lesser-charged clusters based on the general model, involving removal of  $\text{CN}^-$  ligands in some cases (highlighted in red shades).<sup>126</sup>



requires the cluster to be in a deeper and larger “well” to hold more bound electrons. In practice, the “well depth” as well as the “well width” of a cluster can be manipulated by tailoring the core and terminal groups attached to the periphery of the cluster. Thus, stable clusters carrying multiple charges (beyond “−2”) can be formed by assembling lesser-charged clusters and terminal groups each of which already satisfies a closed shell rule with high EA (at the corresponding charge state) and large size. For example, dodecaborane cluster  $B_{12}H_{12}^{2-}$ , stabilized at −2 charge state, can be readily functionalized and reach a variety of charge states and electron seizing powers by changing one of the core atoms as well as the ligands, *e.g.*, B by Be and H by CN which can be accomplished by experiments.<sup>52</sup>

In Table 4, we list the calculated electron affinities of clusters as successive electrons are added. Note that among all the atoms, halogens have the highest electron affinity, but superhalogens such as  $CN^-$ ,  $SCN^-$ ,  $BH_4^-$ , and  $BF_4^-$  (these are often referred to as polyatomic ions) have electron affinities even higher than that of the halogen atoms. However, none of these superhalogens are stable as di-anions such as  $M(CN)_4^{2-}$  ( $M = Mg, Ca, Sr, Ba$ ).<sup>128</sup> Replacing H by CN in  $B_{12}H_{12}^{2-}$  increases the second electron affinity substantially, namely to 5.28 eV, in  $B_{12}(CN)_{12}^{2-}$  dianion. This prediction has been experimentally confirmed with measured second electron affinity at 5.5 eV.<sup>52</sup> Replacing one B atom by Be in  $B_{12}(CN)_{12}^{2-}$  makes it possible for  $BeB_{11}(CN)_{12}^{3-}$  to be stable with the calculated third electron affinity of 2.65 eV. This trianion is a combination of monoanion (superhalogen)  $CN^-$  and a stable dianion  $BeB_{11}(CN)_{11}^{2-}$ . According to the generalized theory given above,<sup>126</sup> these stable dianions and trianions can be used to assemble tetra- and penta-anions that are stable at “−4” and “−5” charge states, respectively. Combining dianions  $BeB_{11}(CN)_{11}^{2-}$  and  $M(CN)_4^{2-}$  ( $M = Ca, Sr, Ba$ ) leads to a set of stable tetra-anions  $BeB_{11}(CN)_{11}M(CN)_4^{4-}$  ( $M = Ca, Sr, Ba$ ) with  $EA_4$  (the fourth electron affinity) being 0.79, 0.20 and 0.25 eV, respectively. Another stable tetra-anion is  $Be_2B_{22}(CN)_{23}^{4-}$ , with

$EA_4$  as large as 1.48 eV by attaching  $BeB_{11}(CN)_{11}^{2-}$  to  $BeB_{11}(CN)_{12}^{3-}$ . A stable penta-anion  $Be_2B_{22}(CN)_{26}Ca^{5-}$  with  $EA_5$  of 30 meV and a vertical detachment energy of 180 meV is created by combining two  $BeB_{11}(CN)_{11}^{2-}$  with one  $Ca(CN)_4^{2-}$ , as shown in Fig. 2B. These results are based on density functional theory and fully optimized clusters and have predictive capability as evidenced by the good agreement between theory and experiment for  $B_{12}(CN)_{12}^{2-}$ .

### III.2 Super-electrophiles enabling Argon and $N_2$ to form chemical bonds at room temperature

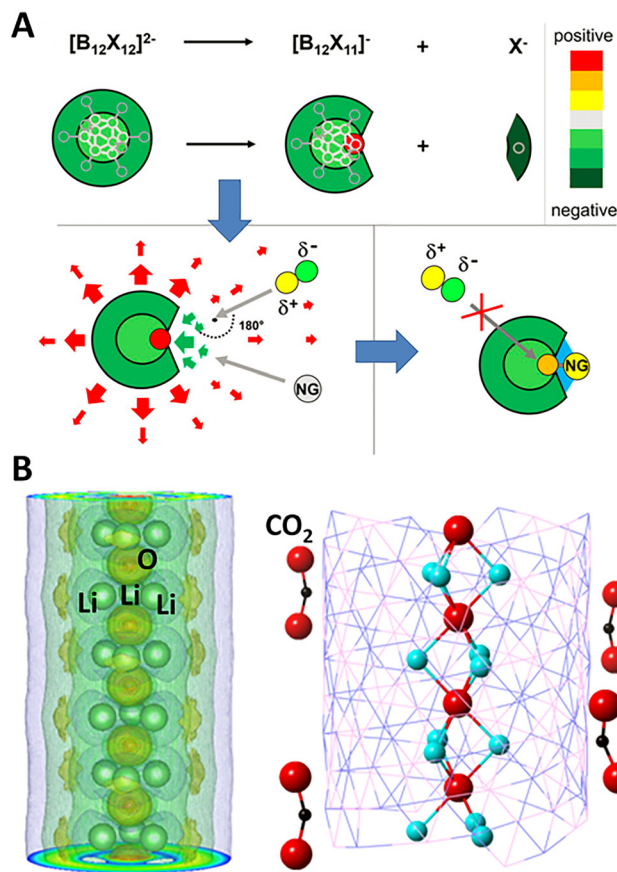
Here, we discuss how very stable multi-anionic clusters can be used to fundamentally change the chemistry of noble gas atoms, making them form chemical bonds at room temperature. Note that noble gas atoms are chemically inert due to the octet shell closure rule.<sup>63,64</sup> They are characterized with high ionization potential and zero to negative electron affinity. Ways to make noble gas atoms to form chemical bonds has been a topic of interest for more than a century. Pauling first suggested that Kr and Xe, due to their large size, can be polarized and form chemical bonds.<sup>129,130</sup> Thirty years later, this was experimentally confirmed by Bartlett.<sup>131</sup> Since then, numerous unsuccessful efforts have been made to see if Ne and Ar can form chemical bonds at room temperature.<sup>132–139</sup>

The prediction of highly stable  $B_{12}(CN)_{12}^{2-}$  (ref. 123) and its experimental discovery<sup>52</sup> changed this scenario. Recent studies have shown that superatomic electrophiles can exert enough oxidative power to even bind the noble gases. When one of the ligands in  $B_{12}(CN)_{12}^{2-}$  is removed, the B atom without a ligand in  $B_{12}(CN)_{11}^-$  remains positively charged and becomes super electrophilic. In addition,  $B_{12}(CN)_{11}^-$  with the extra electron spread over the 11 CN ligands and the positive charge on the “naked” B atom, possesses a dipole moment. The superatomic electrophile becomes reactive, capable of binding an Ar atom chemically at room temperature, as shown in Fig. 3A.

Table 4 Ionic radii (IR) and electron affinities ( $EA_n$ , in eV) at different charge states of stable anions in gas phase<sup>126</sup>

| Anion species  | IR (Å)    | $EA_1$    | $EA_2$    | $EA_3$    | $EA_4$    | $EA_5$ |
|--|-----------|-----------|-----------|-----------|-----------|--------|
| $Cl^-$   | 1.81      | 3.62      | —         | —         | —         | —      |
| $Br^-$   | 1.96      | 3.36      | —         | —         | —         | —      |
| $I^-$  | 2.20      | 3.06      | —         | —         | —         | —      |
| $CN^-$   | 1.40      | 4.10      | —         | —         | —         | —      |
| $SCN^-$  | 2.50      | 3.50      | —         | —         | —         | —      |
| $SeCN^-$   | 2.70      | 3.39      | —         | —         | —         | —      |
| $BH_4^-$   | 2.24      | 3.42      | —         | —         | —         | —      |
| $BF_4^-$   | 2.75      | 7.33      | —         | —         | —         | —      |
| $BCl_4^-$  | 3.69      | 5.01      | —         | —         | —         | —      |
| $B_{12}(CN)_{11}^-$                                  | 5.86      | 8.49      | —         | —         | —         | —      |
| $B_{12}H_{12}^{2-}$                                  | 3.91      | 4.57      | 0.86      | —         | —         | —      |
| $B_{12}(CN)_{12}^{2-}$                               | 5.88      | 8.56      | 5.28      | —         | —         | —      |
| $B_{12}(SCN)_{12}^{2-}$                              | 7.79      | 5.65      | 3.28      | —         | —         | —      |
| $BeB_{11}(CN)_{11}^{2-}$                             | 5.89      | 8.10      | 4.62      | —         | —         | —      |
| $M(CN)_4^{2-}$ ( $M = Ca, Sr, Ba$ )                  | 3.77–4.07 | 7.21–7.08 | 2.62–2.79 | —         | —         | —      |
| $BeB_{11}(CN)_{12}^{3-}$                             | 6.11      | 8.44      | 4.76      | 2.65      | —         | —      |
| $BeB_{23}(CN)_{22}^{3-}$                             | 10.15     | 8.86      | 6.38      | 3.25      | —         | —      |
| $BeB_{11}(CN)_{11}M(CN)_4^{4-}$ ( $M = Ca, Sr, Ba$ ) | 7.03–7.39 | 7.63–4.07 | 5.20–5.15 | 1.77–2.44 | 0.79–0.20 | —      |
| $Be_2B_{22}(CN)_{23}^{4-}$                           | 10.15     | 8.86      | 5.98      | 3.54      | 1.48      | —      |
| $Be_2B_{22}(CN)_{26}Ca^{5-}$                         | 11.66     | 8.37      | 5.98      | 4.25      | 1.88      | 0.18   |





**Fig. 3** Superatoms as super-electrophiles and electrides. (A) Schematics showing the mechanism of binding noble gas atom by electrophilic clusters  $B_{12}X_{11}^{-}$  ( $X = \text{halogen, CN}$ ). Near the positively charged binding site (green arrows), the electric field is strong and attractive; at greater distances, it becomes repulsive (red arrows) owing to the overall negative charge. This leads to the Ng atom being attracted and bond to the positive binding site. Adapted with permission from ref. 52. Copyright 2019 *Proc. Natl. Acad. Sci. U. S. A.* (B) The 1D chain structure of  $Li_3O^{+}\cdot e^{-}$  electride, where the anionic electrons are shown by the yellow shells around the chain on the  $(Li_3)$  planes. The formed nanostructure of  $Li_3O^{+}\cdot e^{-}$  electride grown in a boron-nitride nanotube can simultaneously activate (bind) multiple  $CO_2$  molecules. Adapted with permission from ref. 149. Copyright 2018 The Royal Society of Chemistry.

Consequently, argon loses its nobility.<sup>49,52,140</sup> The binding energy of Ar to  $B_{12}(\text{CN})_{11}^{-}$  is 0.61 eV.

In another experiment, it was shown that  $B_{12}\text{Br}_{11}^{-}$ , with one Br removed from  $B_{12}\text{Br}_{12}^{2-}$ , can bind inert molecular nitrogen ( $N_2$ ) strongly to form  $B_{12}\text{Br}_{11}\text{N}_2^{-}$  at room temperature.<sup>141</sup> More pronounced binding energies to the noble gas atoms can be achieved by incorporating highly electronegative terminal groups or/and replacing a B atom by Be in the boron cage of the clusters. According to the theoretical calculations, super-electrophilic  $\text{BeB}_{11}(\text{CN})_{11}^{2-}$  dianion and  $\text{BeB}_{11}(\text{CN})_{10}^{-}$  monoanion can be formed by subsequently removing  $\text{CN}^{-}$  terminal groups from a stable trianion  $\text{BeB}_{11}(\text{CN})_{12}^{3-}$ . Both electrophiles can then interact strongly with noble gas atoms ( $\text{Ng} = \text{He-Xe}$ ) at the exposed Be and B sites, forming  $[\text{NgBeB}_{11}(\text{CN})_{11}]^{2-}$  and  $[(\text{Ng})_2\text{BeB}_{11}(\text{CN})_{10}]^{-}$  compounds with the highest binding

energies to Ne and Ar among the series being 0.15 and 0.38 eV, respectively.<sup>142</sup> Even higher binding energies of 0.40 and 0.59 eV to Ne and Ar, respectively, can be achieved in trianions  $[\text{BeB}_{23}(\text{CN})_{22}\text{Ng}]^{3-}$ ,<sup>143</sup> where the superatomic electrophile  $\text{BeB}_{23}(\text{CN})_{22}^{3-}$  is obtained by removing one  $\text{CN}^{-}$  from the stable tetra-anion  $\text{BeB}_{23}(\text{CN})_{23}^{4-}$ .<sup>126</sup> Further computational studies show that more than one noble gas atoms can be bound simultaneously in  $[B_{12}Y_{10}-2\text{Ng}]^0$  ( $Y = \text{CN}^{-}$  and  $\text{BO}^{-}$ ) with the highest binding energies of 0.25 and 0.84 eV per Ne and Ar, respectively.<sup>144</sup>

### III.3 Superalkalis as super-electrides

Electrides are ionic compounds and are used as catalysts and reducing agents.<sup>145</sup> Here, anion components are electrons (instead of atoms) localized in cavities in a periodic lattice.<sup>146</sup> Given the extraordinarily low ionization potential, superalkalis can be used to form a new class of super-electrides. The non-nucleus-bound electrons exhibit high mobility and low work function, which can serve as powerful catalytic agents.<sup>147,148</sup> Fang *et al.* have computationally shown that  $Li_3O$  super-alkali can form a one-dimensional electride chain with every unit cell containing two  $Li_3O^{+}\cdot e^{-}$  units.<sup>149</sup> The designed electride and its constructed nanostructures exhibit extremely low electronic work function of 2 eV and can, therefore, readily bind to and activate inert molecules such as  $CO_2$ , as shown in Fig. 3B. Interestingly, the loosely bound anionic (p-type) electrons in this electride even carries a magnetic moment which is found to be  $0.21\mu_B/Li_3O$ .<sup>149</sup> Other superalkalis including  $Cs_3O$  and  $Li_4N$  can also form bulk electrides in 0-, 2-, and 3-dimensions.<sup>150,151</sup> Most recently,  $2[\text{CaCl}]^{+}\cdot 2e^{-}$  is theoretically designed to be a 0-D inorganic electride in which  $\text{CaCl}$  is a superalkali.<sup>152</sup> The interstitial electrons of this electride form a tetragonal lattice between the atomic layers, leading to a highly anisotropic Dirac fermion at the X and M points, protected by symmetries. This electride also shows low work function of 3.4 eV which can effectively facilitate  $N_2$  cracking by electron donation.<sup>152</sup> In another computational design, superalkali  $\text{Ca}_4\text{N}_2$  can form a 1D electride  $[\text{Ca}_4\text{N}_2]^{2+}\cdot 2e^{-}$  with the anionic electrons forming 1D chains confined in the surface channels formed by Ca atoms.<sup>153</sup> It is found that these anionic species are highly mobile, leading to exceptionally high conductance of the material at room temperature.

### III.4 Superatoms as super-catalysts

Catalysts play a critical role in the chemical industry as they are used in nearly 90% of all manufactured goods and are responsible for about 35% of the world GDP. Unfortunately, catalysts are usually composed of expensive metals like Pd, Pt, Rh, Ru, and Ir.<sup>154,155</sup> Naturally, there is considerable interest in either reducing the amount of the metal used or finding earth-abundant metals to replace the expensive ones. As much of the chemical activity occurs on the surface, nanoparticles with large surface to volume ratio have been the catalysts of choice, smaller the particle greater is its reactivity (Fig. 4A).<sup>156</sup> However, nanoparticles still have non-active atoms, and their size variation causes unwanted reactions. Consequently, in the past



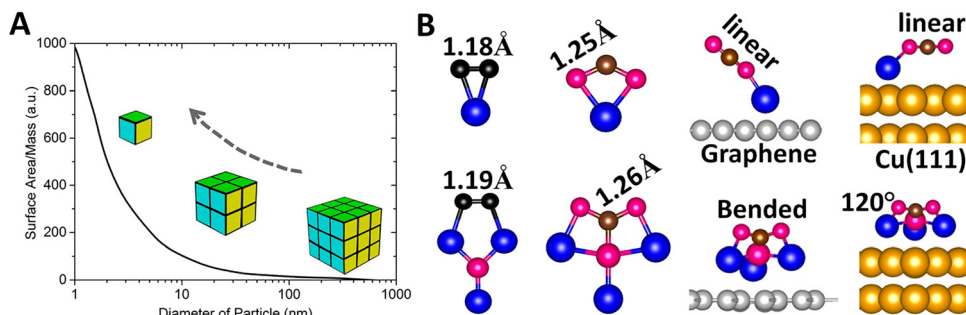


Fig. 4 Superatoms as catalysts. (A) Schematics showing surface-mass ratio with the catalytic power increasing with the reduction of the particle size. Adapted with permission from ref. 156. Copyright 2007 AIP Publishing. (B) Interactions between chemically inert gas molecules (e.g.,  $\text{N}_2$  and  $\text{CO}_2$ ) and superatom  $\text{Li}_3\text{O}$  compared to that of atom Li, with N in black, O in red, C in brown, Li in blue, graphene in grey, and Cu in gold. Adapted with permission from ref. 182. Copyright 2024 Wiley-VCH.

decade, attention has been focused on single-atom catalysts,<sup>157–162</sup> where single metal atoms, the ultimate nanoparticle, are dispersed on a substrate. This too has limitations; single-atom catalysts only offer a single active site that may not be sufficient for reactions requiring the co-adsorption of multiple reactants.<sup>162–166</sup> While dual- and triple-atom catalysts (DAC and TAC) have been suggested as a solution, there is no unique way to choose their composition.

Superatoms offer an alternative. Their sizes can be very small with almost all the atoms lying on the surface and participating in chemical reactions. While several studies have demonstrated the benefits of using atomic clusters as catalysts, they deal with clusters in the gas phase.<sup>167–181</sup> However, for practical applications, clusters need to be deposited on substrates which raises further concerns: (1) once deposited on a substrate, clusters would interact with the substrate atoms. This not only may change the structure and properties of the clusters but may also affect the substrate topology. Consequently, the reaction of molecules with clusters in the gas phase may be very different from those when supported on a substrate. (2) Secondly, the results may depend upon the adopted substrate itself.

Recently, Kilic and Jena<sup>182</sup> addressed these issues by focusing on the reaction of small molecules such as  $\text{H}_2$ ,  $\text{O}_2$ ,  $\text{N}_2$ ,  $\text{CO}$ , and  $\text{CO}_2$  with Li and  $\text{Li}_3\text{O}$  in the gas phase as well as when supported on graphene, Au(111) and Cu(111) substrates. Using calculations based on density functional theory, they showed that while the reactions of the above molecules with Li and  $\text{Li}_3\text{O}$  in the gas phase are rather similar,  $\text{Li}_3\text{O}$  binds and activates the above molecules more strongly than the Li atom (Fig. 4B). When deposited on the above substrates, the geometry of  $\text{Li}_3\text{O}$  and that of the substrates are barely changed. As shown in Fig. 4B, the  $\text{CO}_2$  molecule maintains its linear geometry when interacting with the Li atom on the substrate, but undergoes significant bending when it interacts with the  $\text{Li}_3\text{O}$  superatom, with the bond angle  $\angle \text{OCO}$  reducing to  $120^\circ$  on the Cu(111) substrate. This is due to the low ionization potential of  $\text{Li}_3\text{O}$ , allowing it to readily donate an electron to the  $\text{CO}_2$  molecule and lead to increased binding energy and activation. Although neither Li nor  $\text{Li}_3\text{O}$  are traditional

catalysts, the above study demonstrates that a single-superatom catalyst can offer advantages over a single-atom catalyst, since they can provide multiple active sites with a facile electron transfer. As superatoms provide a new paradigm in catalyst design, identifying superatoms that can mimic the chemistry of commonly used catalysts, such as Pd, would be important.

In a photoelectron spectroscopy experiment on negatively charged ions, Castleman and coworkers compared the electronic structure of Pd atom with that of ZrO dimer.<sup>183</sup> Considering the electronic configurations of Pd ( $[\text{Kr}]4d^{10}$ ), Zr ( $[\text{Kr}]4d^25s^2$ ), and O ( $[\text{He}]2s^22p^4$ ), one finds that the ZrO dimer is isoelectronic with the Pd atom. The good agreement in the electronic structures of these two species led the authors to suggest that ZrO could be a replacement catalyst for Pd. While early work of the interaction of gas molecules with isolated  $\text{Pd}_n$  and  $\text{ZrO}_n$  clusters casted doubt on this possibility,<sup>178</sup> we should note that an understanding of the reactions of gas molecules with clusters in the gas phase may not be sufficient to ascertain their suitability as good catalysts, because in practical applications the catalysts need to be supported on a substrate. Recently, Kilic and Jena studied the hydrogen evolution reaction (HER),  $\text{CO}_2$  reduction reaction ( $\text{CO}_2\text{RR}$ ), and ammonia production by supporting ZrO dimer and Pd atom on graphene, Cu(111), Au(111) surfaces.<sup>182,184,185</sup>

First, it is important to ensure that the structure and properties of isolated clusters remain unaffected by their interaction with the substrate and that gas molecules bind and get activated by the supported clusters. Kilic and Jena confirmed that the geometry of ZrO remains unchanged when the dimer is supported on the above substrates and that the ZrO dimer is even more strongly bound to the substrate compared to the Pd atom, as shown in Fig. 5.<sup>184</sup>

In Fig. 6, we present the interaction  $\text{CO}_2$  with Pd atom and ZrO dimer supported on graphene, Cu(111) and Au(111) surfaces.<sup>184</sup> The first step in a catalytic process is the binding and subsequent activation of the  $\text{CO}_2$  molecule. Note that  $\text{CO}_2$  is a linear molecule and very stable because of the covalent bond between C and O. Its activation requires charge transfer to the molecule so that not only the CO bond is elongated, but



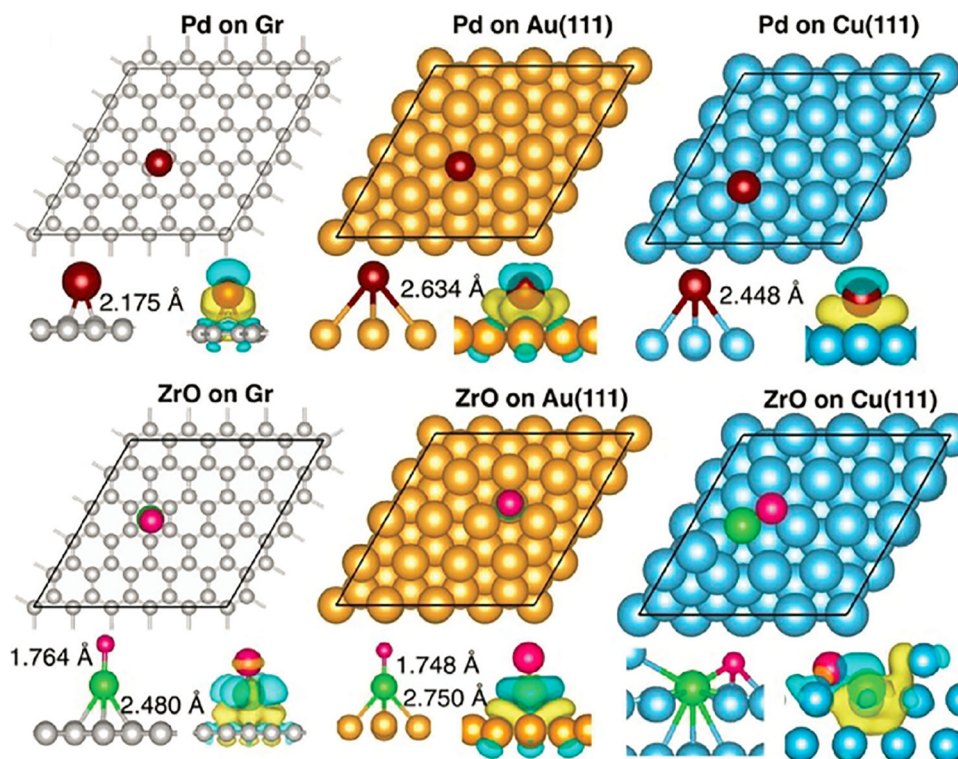


Fig. 5 Optimized positions of Pd and ZrO supported on the graphene, Au(111), and Cu(111) surfaces, with C, Au, Cu, Pd, Zr, and O in gray, yellow, cyan, violet, green, and red, respectively. The corresponding charge density difference is shown in the lower panel, with yellow and cyan iso-surfaces representing charge accumulation and depletion, respectively. Adapted with permission from ref. 184. Copyright 2025 Wiley-VCH.

also, the O–C–O bond angle is decreased from  $180^\circ$ . For  $\text{CO}_2$  adsorbed on Pd (ZrO) doped graphene, Au, and Cu surfaces, the calculated C–O and C–O\* bond lengths—where O\* represents the oxygen atom bound to Pd or Zr—are 1.199/1.241 Å (1.215/1.329 Å), 1.174/1.183 Å (1.207/1.293 Å), and 1.229/1.236 Å (1.217/1.312 Å), respectively. The corresponding  $\angle\text{OCO}$  bond angles are  $151^\circ$  ( $132^\circ$ ),  $179^\circ$  ( $131^\circ$ ),  $142^\circ$  ( $130^\circ$ ). This clearly shows that ZrO can activate  $\text{CO}_2$  more strongly than the Pd atom, irrespective of the substrate used. The calculated  $\text{CO}_2$  reduction reaction and reaction pathways also shows that ZrO outperforms Pd as a catalyst, establishing the single-supercatalysts are superior to single-atom catalysts. Investigations of  $\text{NH}_3$  synthesis further reveal that graphene-supported single-supercatalysts (TiO, ZrO, and WC) outperform their single-atom counterparts (Ni, Pd, and Pt) in both stability and catalytic performance toward the electrochemical nitrogen reduction reaction.

#### IV. Unusual physical properties of superatomic crystals – phonon transport and thermal conductivity

The transformative power of atomic clusters lies in their ability to serve as well-defined building blocks for the design of novel materials. By treating clusters as “superatoms” with tunable electronic structures, one can effectively extend and enrich the

conventional periodic table, enabling access to chemical compositions and functionalities unattainable with individual elements alone. The principal classes of clusters employed as building units in experimentally synthesized assemblies include: (1) fullerenes and their analogues, such as  $\text{C}_{60}$ ,  $\text{K}_3\text{C}_{60}$ , and endohedral fullerenes (e.g.,  $\text{M@Si}_{16}$  and  $\text{M@Ge}_{16}$ );<sup>25,186–188</sup> (2) boranes and carboranes, including  $\text{B}_{12}\text{H}_{12}^{2-}$  and  $\text{C}_2\text{B}_{10}\text{H}_{12}$ ;<sup>189,190</sup> (3) metal chalcogenide clusters, such as  $\text{M}_6\text{X}_8$  (M = Cr, Co, Re; X = S, Se, Te);<sup>28,77,191,192</sup> (4) ligand-protected metal clusters,<sup>193–198</sup> including  $\text{Au}_{25}$ ,  $\text{AgAu}_{24}$ ,  $\text{Au}_1\text{Ag}_{22}$ ,  $\text{Pt}_1\text{Ag}_{18}$ ,  $\text{Ag}_{14}$ , and  $\text{Pd}_3$ .

As summarized in Table 1, materials assembled from cluster building blocks often exhibit properties markedly distinct from those of their atom-based counterparts, owing to their unique electronic shells, size-dependent effects, and inter-cluster coupling. In this section, we focus specifically on their phonon properties and the unique lattice dynamical behaviors that emerge from cluster-based architectures.

Phonons are collective excitations of vibrational modes in solids; they carry energy and momentum and can exhibit wave-particle duality. The ability to manipulate phonons holds significant implications for tailoring intrinsic properties of materials. However, phonons are bosons and do not follow the conventional polarity regulation used for fermions like electrons. Superatomic crystals (SACs)<sup>186,199</sup> provide a unique platform for phonon manipulation. In conventional materials, phonons arise from atomic vibrations. In contrast, SACs, where superatomic clusters serve as the building blocks, exhibit two



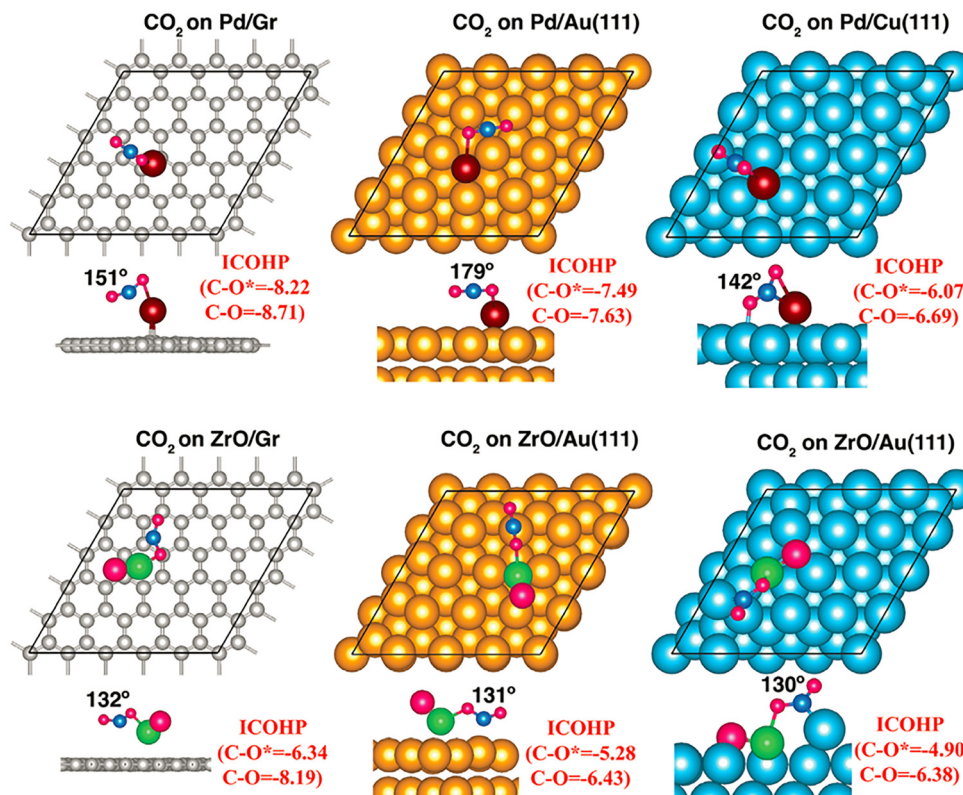


Fig. 6 The top and lateral views of CO<sub>2</sub> adsorbed on Pd single atom and ZrO speratom supported on the graphene, Au(111), and Cu(111) surfaces, with C (graphene), Au, Cu, Pd, Zr, and O in gray, yellow, cyan, violet, green, and red, respectively. For the CO<sub>2</sub> molecule, C is in blue and O in red. Adapted with permission from ref. 184. Copyright 2025 Wiley-VCH.

distinct vibrational modes: (1) intra-cluster vibrations of atoms within a superatomic unit, and (2) inter-cluster vibrations of the clusters themselves. These combined vibrational modes can alter the phonon spectra and influence phonon scattering and coherence, leading to properties distinct from those of conventional atomic crystals as described in Table 5. In addition, the interactions between phonons and electrons are also different as demonstrated in K<sub>3</sub>C<sub>60</sub>. Neither K nor C<sub>60</sub> is superconducting, but K<sub>3</sub>C<sub>60</sub> is where the C<sub>60</sub> fullerenes composed of 60 carbon atoms play a fundamental role.<sup>200</sup>

Coupling between intra-superatom and inter-superatom vibrations can effectively suppress phonon mobility, while

enhancing higher-order phonon scattering and phonon coherence. These factors contribute to intriguing properties such as ultralow lattice thermal conductivity and remarkable thermal rectification. Experimentally, Ong *et al.*<sup>201</sup> conducted the first measurement of thermal transport in unary SACs of Co<sub>6</sub>X<sub>8</sub>(-PET<sub>3</sub>)<sub>6</sub> [X = S, Se, Te; PET<sub>3</sub> = triethylphosphine, Et = (CH<sub>2</sub>CH<sub>3</sub>)<sub>3</sub>] and the binary SACs of [Co<sub>6</sub>X<sub>8</sub>(PET<sub>3</sub>)<sub>6</sub>][C<sub>60</sub>]<sub>2</sub>. Their results indicate that heat conduction in SACs is mediated by collective inter-superatom phonons, with mean free paths determined by the inter-superatom separation and coupling strength. Upon transitioning from an orientationally disordered to an ordered structure, phonon coherence is enhanced, resulting in longer

Table 5 Key differences in phonon transport and thermal conductivity between superatomic crystals and conventional atomic crystals

| Feature                            | Superatomic crystals  | Conventional atomic crystals                                |
|------------------------------------|---|---|
| Building blocks                    | Large, complex superatoms   | Individual atoms  |
| Unit cell size                     | Large   | Small (atomic scale)  |
| Phonon dispersion                  | Flattened bands with many localized modes                               | Well-defined dispersive phonon bands                        |
| Phonon group velocity              | Low (flat dispersion)   | High (steep dispersion)                                     |
| Phonon scattering                  | Strong 3- and 4-phonon scattering caused by structural complexity       | Lower scattering  |
| Phonon coherence                   | Weak in most cases  | Strong in most cases  |
| Dominant phonon modes              | Localized intra-superatom modes and low-frequency inter-superatom modes | Acoustic and optical phonons                                |
| Anharmonicity                      | High  | Low for most systems  |
| Phonon mean free path (MFP)        | Short   | Long  |
| Thermal conductivity ( $\kappa$ )  | Ultralow  | Moderate to high (depending on materials)                   |
| Temperature dependence of $\kappa$ | Weak or plateau-like  | Strong ( $\kappa \propto 1/T$ at high $T$ for most systems) |



mean free paths and increased thermal conductivity at lower temperatures. Additionally, they found that the hierarchical bonding features expand the phonon space by combining localized 0D modes with the delocalized 2D and 3D modes. This phenomenon is observed in a SAC consisting of  $\text{Re}_6\text{Se}_8$  as the superatomic building block, forming a layered van der Waals lattice.<sup>202</sup> Theoretically, it was also shown that the rotational dynamics and orientational ordering of the superatoms significantly impact the lattice thermal conductivity.<sup>203</sup>

Moreover, the bonding hierarchy in SACs introduce significant four-phonon anharmonicity. For instance, in a superatom-based superionic conductor  $\text{Na}_3\text{OBH}_4$ , strong four-phonon scattering extends the phonon scattering phase space by 33% at 300 K, leading to a 24% reduction in lattice thermal conductivity.<sup>204</sup> A similar phenomenon is observed in calculations of supersalt  $\text{NaNO}_3$  monolayer,<sup>205</sup> which is composed of superhalogen  $\text{NO}_3$ , significantly reducing the lattice thermal conductivity from  $4.13$  to  $1.65 \text{ W m}^{-1} \text{ K}^{-1}$  at 300 K. Moving from atom-superatom hybrid systems to a solely superatom-based supersalt  $\text{PH}_4\text{AlBr}_4$  composed of superalkali  $\text{PH}_4$  and superhalogen  $\text{AlBr}_4$ , Du *et al.*<sup>206</sup> showed theoretically that the structural and bonding hierarchies are significantly enhanced with a much more complex geometric configuration. Notably, the phonon mean-free paths of several modes are comparable to the interatomic distances. This, known as the Ioffe–Regel limit, results in strong phonon coherence. Consequently, the lattice thermal conductivity is controlled by a two-channel mechanism, incorporating both phonon scattering and phonon coherence.

These unusual phonon transport characteristics make SACs promising for applications in phononic devices. One example is the thermal rectification (TR), which is similar to the diode-like effect observed in electronics. TR occurs in structures that transfer heat in a nonreciprocal manner, driven by the interplay between structural gradient and lattice nonlinearity. Recent computational studies have shown that the superatom-based TR devices exhibit two notable features that distinguish them from atom-based systems with atomically flat surfaces:<sup>207,208</sup> (1) the large size and non-spherical shape of superatoms not only make their assembled structures porous but also induce significantly high roughness on both internal and external surfaces. This enhances the tuning of phonon transport, resulting in unconventional thermal rectification behavior. (2) SACs-based fractals have high surface-to-volume ratio and surface roughness that changes exponentially with the fractal levels. This leads to highly nonlinear temperature variations with spatial positions, inducing nonlinear heat transfer; and causing the thermal conductivity to exhibit high asymmetry, which is sensitive to the direction of the heat current.

## V. Polyatomic ions as building blocks of energy materials

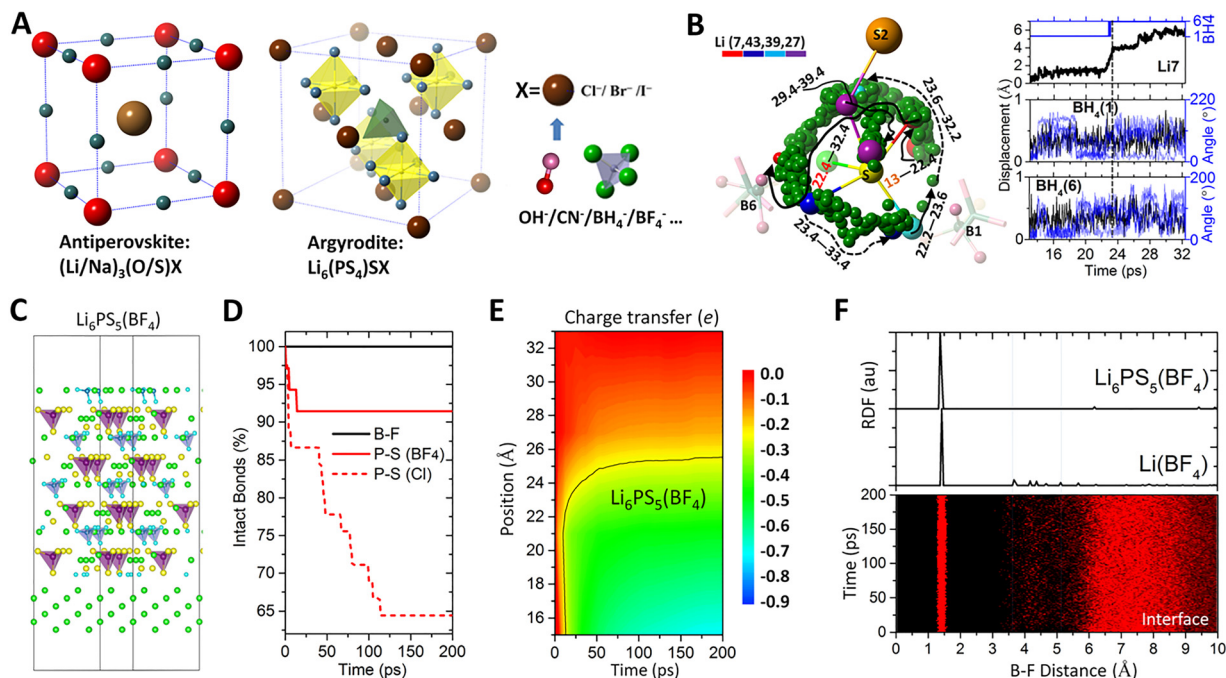
Polyatomic ions such as  $\text{BH}_4^-$  hold the key to overcome some of the key challenges in developing materials for energy

production and storage. As mentioned before, these ions not only mimic the chemistry of halogens, but also their electron affinities are larger than those of halogens. In this context, they can be regarded as superatoms. Each being a group of atoms (cluster) stabilized at a certain charge state; polyatomic ions can provide a much wider materials design palette than employing individual atoms. Instead of limited number of elements in each group of the periodic table, many more polyatomic species are available due to mix-and-match of elements in their compositions. Compared to the spherical shape and periodic bonding characteristics of atomic ions, polyatomic ions carry various chemical compositions, symmetries (shapes), sizes, charge distributions among the composite atoms, and electron affinities. Accordingly, polyatomic ions serving as dopants or substitutes in materials can bring additional degrees of freedom to tailor and optimize the bulk and interface properties. Significant breakthroughs in application and understanding of polyatomic ions as building blocks of energy materials have been achieved, especially in the development of solid-state electrolytes for batteries and hybrid halide perovskites for solar cells. These studies are highlighted by synergy between theory and experiment.

### V.1 Solid-state electrolytes

Solid-state electrolytes hold the promise to achieve safe, cost-effective, and high energy electrochemical systems that are essential for next-generation long-duration energy storage. Two major challenges remain in the development of solid electrolytes – to achieve adequate ionic conductivity at room temperature on par with those of liquid electrolytes and to have stable interface against metal anode for high energy density. However, solid electrolytes with high ionic conductivities are often characterized by weak anion–cation interactions and high formation energies, and, therefore, exhibit poor interfacial stability against the metal anode. Recent theoretical studies have shown that substituting halogen (a ubiquitous composite in solid electrolytes) by selected polyatomic ions (*e.g.*,  $\text{BH}_4^-$ ,  $\text{BF}_4^-$ ,  $\text{CN}^-$  *etc.*) can simultaneously enhance the ionic conductivity and stabilize the interfacial stability of solid electrolytes. Doping polyatomic ions to the halogen sites was first proposed for antiperovskite solid electrolytes due to their large structural tolerance.<sup>209,210</sup> The strategy was later extended to the argyrodite family due to their well-known high ionic conductivity but poor interfacial stability against Li metal anode.<sup>211–213</sup> Once incorporated into the structure of a solid electrolyte, the polyatomic ions can introduce motional dynamics into the host structure, with an extraordinarily large length scale that is 1–2 orders of magnitude higher than the regular thermal vibrational amplitude ( $0.01$ – $0.1 \text{ \AA}$ ) of individual atoms, as shown in Fig. 7A and B. The translational and rotational dynamics of the non-spherical polyatomic ions effectively alter the surrounding energy landscape, which can significantly lower the diffusion barrier and enhance the ionic diffusion in the electrolyte.<sup>209,210</sup> Additional channel spaces for ionic diffusions can be formed around smaller halogen ions when they are co-existing with relatively large polyatomic ions in a





**Fig. 7** Solid state electrolytes based on superatomic ions. (A) Design of solid electrolytes by substituting halogen for selected polyatomic ions. (B) Diffusion trajectory (from 22.4 to 39.4 ps) of  $\text{Li}^+$  (indexed by 7, 43, 39 and 27 in the simulation cell) displacements (up to 6 Å) coupled with translational displacements ( $\sim 0.5$  Å) and rotations ( $\sim 200^\circ$ ) of  $\text{BH}_4^-$  clusters (B1 and B6). Adapted with permission from ref. 211. Copyright 2022 Springer Nature. (C) An interface model between argyrodite solid electrolyte  $\text{Li}_6\text{PS}_5(\text{BF}_4)$  based on polyatomic  $\text{BF}_4^-$  (violet tetrahedra) and Li metal (in green). (D) Explicit interface modelling showing that incorporating  $\text{BF}_4^-$  can stabilize the P–S bonds in  $\text{Li}_6\text{PS}_5(\text{BF}_4)$ , in contrast to the case of  $\text{Li}_6\text{PS}_5\text{Cl}$  based on Cl. (E) Presence of  $\text{BF}_4^-$  can inhibit electron transfer from Li metal into the solid electrolyte across the interface. (F) Radial distribution function analysis at the  $\text{Li}_6\text{PS}_5(\text{BF}_4)$ –Li interface suggests the formation of  $\text{Li}(\text{BF}_4)$  interphase which is electronically insulating and ionically conductive. Adapted with permission from ref. 212. Copyright 2025 American Physical Society.

structure.<sup>210</sup> Upon interfacial decomposition of the polyatom-based solid electrolyte, the polyatomic ions can form interphases that are ionically conductive yet electronically insulating, which can effectively stabilize the interface against metal anode,<sup>212</sup> as shown in Fig. 7C–F.

Inspired by these early theoretical studies, recent experiments<sup>214–219</sup> have substituted the halogen site with polyatomic ions including  $\text{BF}_4^-$  and  $\text{BH}_4^-$  in the antiperovskite solid electrolytes  $\text{M}_3\text{YX}$  and  $\text{M}_{3-x}\text{OH}_x\text{X}$  ( $\text{M}$  = alkali metal;  $\text{Y}$  = chalcogen;  $\text{X}$  = halogen). A mechanochemical synthesis is typically used, with ball milling powder mixing of metal borohydride/tetrafluoroborate, metal halides, and metal oxides.<sup>214,215</sup> It is found that, among the doping series  $\text{Li}_2\text{OHCl}_{1-x}(\text{BF}_4)_x$  ( $x = 0.02, 0.05, 0.1$ ),  $\text{Li}_2\text{OHCl}_{0.98}(\text{BF}_4)_{0.02}$  exhibits the lowest activation energy and the highest ionic conductivity which is about 2 times higher than that of  $\text{Li}_2\text{OHCl}$ . The polyatom-doped system also exhibits much enhanced stability against Li metal anode.<sup>214</sup> The doping concentration of the polyatomic ion is crucial to achieve improved properties. High concentration of the polyatomic dopants can be detrimental to the ionic conductivity. This is evidenced by another recent experiment, where  $\text{Na}_3\text{O}(\text{BH}_4)$  with 100% substitution of Br in antiperovskite  $\text{Na}_3\text{OBr}$  by  $\text{BH}_4^-$  exhibits even lower ionic conductivity than that of  $\text{Na}_3\text{OBr}$ .<sup>215</sup>

The polyatomic doping strategy has also been applied to the argyrodite solid electrolytes  $\text{Li}_6\text{PS}_5\text{X}$  ( $\text{X}$  = halogen) in recent

experiments. With the halogen replaced by polyatomic  $\text{CN}^-$ ,  $\text{Li}_6\text{PS}_5\text{CN}$  is synthesized using low-temperature solution-based processing.<sup>216</sup> It is found that  $\text{Li}_6\text{PS}_5\text{CN}$  exhibits lower activation barrier than that of  $\text{Li}_6\text{PS}_5\text{Br}$  which is the lowest among the halide-argyrodite family.<sup>216</sup> In another experimental work,  $\text{Li}_6\text{PS}_5\text{Cl}_{0.9}(\text{BH}_4)_{0.1}$  is mechanochemically synthesized with ball milling.<sup>217</sup> It is found that the partial substitution of  $\text{Cl}^-$  by  $\text{BH}_4^-$  greatly improves the ionic conductivity to  $2.83 \text{ mS cm}^{-1}$  which is twice as high as that of  $\text{Li}_6\text{PS}_5\text{Cl}$ . The experiment also shows that the  $\text{Li}/\text{Li}_6\text{PS}_5\text{Cl}_{0.9}(\text{BH}_4)_{0.1}/\text{Li}$  symmetric cell can stably run for more than 400 h, with a critical current density of up to  $3.5 \text{ mA cm}^{-2}$ .<sup>217</sup> Doping high concentration of  $\text{BH}_4^-$  in argyrodite may lead to mixed phases, as shown by the recently synthesized thiophosphate solid electrolytes  $(1-x)\text{Li}_3\text{PS}_4 \cdot 2x\text{LiBH}_4$  using a simple milling method without heat treatment.<sup>218</sup> The ionic conductivity increases, and the activation energy decreases with increasing  $\text{BH}_4^-$  up to  $x = 0.54$  (with the highest room-temperature ionic conductivity of  $11 \text{ mS cm}^{-1}$ ), beyond which the trends are reversed. The resulting solid electrolyte also exhibits superior cycle and rate performances in a Li–In/NCM all-solid-state cell.<sup>218</sup> Further experiments find that even higher ionic conductivities can be achieved in the argyrodite by simultaneously substituting halogen and  $\text{BH}_4^-$  for  $\text{S}^{2-}$  using room-temperature ball milling.<sup>218</sup> Among the resulting series,  $\text{Li}_{7-a-b}\text{PS}_{6-a-b}(\text{BH}_4)_a\text{X}_b$  ( $\text{X} = \text{Cl}, \text{Br}, \text{I}; a + b \leq 1.8$ ),  $\text{Li}_{5.35}\text{PS}_{4.35}(\text{BH}_4)_{1.15}\text{Cl}_{0.5}$  exhibits the highest



ionic conductivity of  $26.1 \text{ mS cm}^{-1}$  at room temperature. The corresponding symmetric cell maintains stable lithium plating/stripping over 2000 hours at  $1 \text{ mA cm}^{-2}$  and delivers a high critical current density of  $2.1 \text{ mA cm}^{-2}$ ,<sup>219</sup> underscoring its excellent interfacial stability. A most recent experiment shows that  $\text{LiBH}_4$  doping into  $\text{Li}_6\text{PS}_5\text{Cl}$  creates to a tri-layer solid electrolyte interphase of  $\text{Li}_3\text{P}/\text{LiBH}_4/\text{Li}_2\text{S}$  that blocks electrons while facilitating ion diffusion.<sup>220</sup> This leads to greatly enhanced electrochemical stability, with the symmetric cells achieving a critical current density (CCD) of  $7.3 \text{ mA cm}^{-2}$  which is almost three times that of the baseline without doping.<sup>220</sup>

The working mechanism of lowering activation energy and improving ionic conductivity demonstrated in the above experiments relies on the interplay between the doped anion cluster and the diffusing metal ions. The dynamical reorientation of the polyatomic anions in the solid electrolytes at room temperature are verified by experimental characterizations using neutron diffraction or high-resolution synchrotron powder X-ray diffraction.<sup>216,221</sup> Modeling studies and data analysis are necessary to uncover the possible correlation between the dynamics of the anion clusters and the diffusion metal ions. Most recent theoretical studies explain the coupling between the dynamics of the doped polyatomic ions and the metal ion diffusion.<sup>211,222,223</sup> Using light mono-anion clusters ( $\text{BH}_4^-$ ,  $\text{CN}^-$ ,  $\text{SH}^-$ , *etc.*) working in the argyrodite as a model system, it is found that the dynamics of the anion cluster inside the structure can be categorized into two types. One is the so-called “responsive dynamics”, where, due to their exceptionally large motional degrees of freedom in the structure, the polyatom can effectively accommodate the motion of a passing metal ion by rotation and translation. This can lower the migration barrier and the activation energy, and, therefore, enhance the ionic diffusion. The other is called “active dynamics”, where the polyatomic anions are rotating and translating under thermal excitations. This type of dynamics may inhibit rather than facilitate the diffusion of a passing ion, depending on the specific migration path at that moment. Increasing the concentration of polyatomic dopants will increase both types of dynamics at the same time. Therefore, the polyatomic concentration that can enable the optimal properties of a system can only be achieved when the “responsive dynamics” is maximized against the detrimental effect of the “active dynamics”.<sup>211</sup> Such cluster dynamics has been extended most recently to heavy polyanion clusters (*e.g.*,  $\text{PS}_4^{3-}$ ) in the sulfide electrolytes.<sup>222</sup> Thus, the coupling between the dynamics of the polyatomic ions and the diffusing ion is beyond the so-called “paddle-wheel mechanism” that has always been assumed, where a rotating cluster ‘drags’ or ‘pulls’ its nearby metal ions. Realizing the difference between the “responsive dynamics” and the “active dynamics” is crucial to accurately understand the effect of the doped anion cluster on the ionic diffusion. Had the clusters’ dynamics always facilitate the ionic diffusion as depicted in the early “paddle-wheel mechanism”, ionic conduction would increase with elevated doping level of the cluster. This contradicts the observation in the above-mentioned experiments.

## V.2 Hybrid perovskite solar cells

In another field of interest in energy materials, halide hybrid perovskite (*e.g.*,  $\text{MAPbX}_3$ ,  $\text{MA} = \text{CH}_3\text{NH}_3$ ,  $\text{X} = \text{halogen}$ ) solar cells have attracted considerable attention due to their low-cost solution-based synthesis and decent power conversion efficiency (PCE). One of the main challenges for the commercialization of these materials is their instability against moisture and heat. Recent studies have found that the polyatomic substitution for the halogen inside the hybrid perovskites has great potential to overcome this challenge. Early theoretical studies showed that partial replacement of halogen by selected polyatomic ions (*e.g.*,  $\text{BH}_4^-$ ,  $\text{SCN}^-$ , *etc.*) can greatly enhance the stability of the hybrid perovskites.<sup>224,225</sup> Recent experimental works in this field have been summarized in two comprehensive reviews.<sup>226,227</sup> Here, we focus on the underlying mechanism of the stabilization of hybrid perovskites by incorporating polyatomic ions, which has not been adequately emphasized in the published reviews.

Based on the concept of superatoms, one can view a hybrid perovskite as a “super” alkali halide. The organic cation  $\text{MA}^+$  behaves as a superalkali with an ionization potential lower than that of Li, while the inorganic framework anion  $[\text{MX}_3]^-$  ( $\text{M} = \text{Pb}$ ,  $\text{Sn}$ ,  $\text{Ge}$ ;  $\text{X} = \text{halogen}$ ) functions as a superhalogen with electron affinity greater than that of Cl. This pronounced ionic character enhances Coulombic attraction toward a polarized water molecule, which becomes confined between  $\text{MA}^+$  and  $[\text{PbX}_3]^-$ . Water then mediates degradation by promoting  $\text{I}^-$  removal from  $[\text{PbI}_3]^-$ ; the released  $\text{I}^-$  combines with  $\text{H}^+$  from the organic cation to generate HI and  $\text{PbI}_2$ ,<sup>225</sup> as shown in Fig. 8A.

Partial replacement of the halogen (X) in  $[\text{PbX}_3]^-$  by selected polyatomic ions (*e.g.*,  $\text{BH}_4^-$ ,  $\text{SCN}^-$ ,  $\text{SeCN}^-$  *etc.*) can break the above degradation process. Given that the polyatomic cluster contains multiple atoms, the ‘-1’ charge of the whole cluster is unevenly distributed among the constitute atoms (*e.g.*, S/Se, C, and N in  $\text{SCN}^-/\text{SeCN}^-$ , or B and H in  $\text{BH}_4^-$ ). It has been proved that such multi-centered charge distribution inside a polyatomic ion can effectively reduce the Coulombic attraction to water, so that the water molecule is unlikely to be trapped between the superalkali  $\text{MA}^+$  and superhalogen  $[\text{PbX}_3]^-$  in the first place. Meanwhile, such charge distribution enhances the bonding between Pb and X within  $[\text{PbX}_3]^-$  so that  $\text{X}^-$  is unlikely to be dislodged from the group to react with  $\text{H}^+$  of the organic cation,<sup>224</sup> as shown in Fig. 8B.

Optimal bandgaps and band edges are crucial to achieve high PCE in the hybrid perovskites. For the regular alkali halide ionic compounds AX ( $\text{A} = \text{alkali}$ ,  $\text{X} = \text{halogen}$ ), the bandgap decreases from  $\text{X} = \text{Cl}$  to  $\text{I}$  with increasing size of the halogen (X). Similarly, the bandgaps of hybrid perovskites viewed as “super” alkali halides also depend on the size of the doped superhalogen. Theoretical studies find that the polyatomic anions  $\text{BH}_4^-$ ,  $\text{SCN}^-$  and  $\text{SeCN}^-$  all have ionic radii similar or smaller than that of  $\text{I}^-$ .<sup>225</sup> Therefore, the resulting  $\text{MAPbI}_{2.75}\text{Y}_{0.25}$  ( $\text{Y} = \text{BH}_4$ ,  $\text{SCN}$ ,  $\text{SeCN}$ ) systems exhibit the optimal bandgaps for solar cell as that of  $\text{MAPbI}_3$ . Besides the proper bandgap, the doped anion clusters can simultaneously achieve



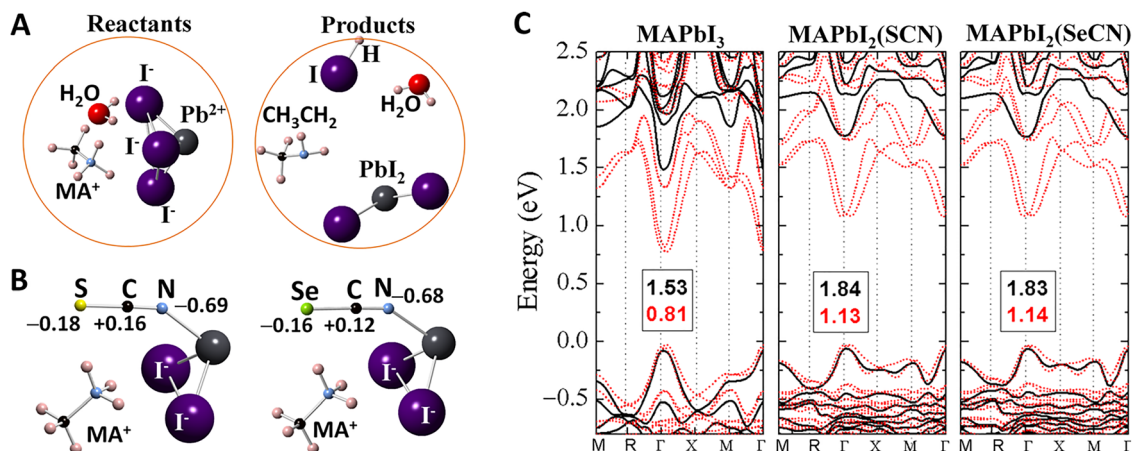


Fig. 8 Doping polyatomic ions in hybrid perovskites to improve properties. (A) Mechanism of the hybrid perovskite catalyzed by water at the molecular level. (B) Partial replacement of halogen by SCN<sup>-</sup> or SeCN<sup>-</sup> with the linear configuration. The charge distribution (values in e) along these polyatomic ions entails reduced attraction with water molecule and enhanced bonding with Pb in the functional group. (C) Calculated band gaps (1.13 and 1.14 eV) and band edges (Rashba splitting highlighted in red) of the polyatom-doped hybrid perovskites compared to those of MAPbI<sub>3</sub>. Adapted with permission from ref. 224. Copyright 2017 American Chemical Society.

decent carrier mobility and the large Rashba splitting at the band edges, which are essential to reach high PCE in the hybrid perovskites,<sup>224</sup> as shown in Fig. 8C.

Indeed, most recent experiments confirm the above theory. It has been demonstrated that the stability of the hybrid perovskite against humidity, thermal, and ultraviolet illumination are significantly enhanced by incorporating BH<sub>4</sub><sup>-</sup>. According to the experiment, the polyatomic dopant not only effectively inhibits the evaporation and decomposition of the methylamine group (MA<sup>+</sup> = CH<sub>3</sub>NH<sub>3</sub><sup>+</sup>) in the resulting MAPbI<sub>3-x</sub>(BH<sub>4</sub>)<sub>x</sub>, but also increases the PCE from 18.43 to 21.10%, accompanied by increased short-circuit current density and open-circuit voltage with near-zero current-voltage hysteresis.<sup>228</sup> In another telling experiment, it is demonstrated that adding appropriate alkali metal halide KSeCN into perovskite can result in MAPbI<sub>3-x</sub>(SeCN)<sub>x</sub> which exhibits significantly improved stability, PCE of 18.41%, hysteresis index of 1.5%, and efficiency of 85.3% after 500 h.<sup>229</sup> More beneficiary effects of the polyatomic anion additives due to various mechanisms have been reported in recent experiments.<sup>230-232</sup> High-quality perovskite CsPbBr<sub>3</sub> nanocrystals featuring ultrahigh stability are obtained by incorporating polyatomic BF<sub>4</sub><sup>-</sup> anions due to an *in situ* surface passivation.<sup>233</sup> In another experiment, MABF<sub>4</sub> serving as the BF<sub>4</sub><sup>-</sup> additive can effectively improve the photovoltaic performance of the formamidine (FA) lead iodide (FAPbI<sub>3</sub>) perovskite solar cell by facilitating the perovskite crystal growth and enhancing the heterointerfaces between the FAPbI<sub>3</sub> layer and carrier-transport materials, while retaining the favorable narrow bandgap of FAPbI<sub>3</sub>.<sup>234</sup>

## VI. Superatom-based ferroelectric, piezoelectric and thermoelectric materials

A common characteristic of ferroelectric, piezoelectric and thermoelectric materials is that their properties are all related

to electric polarization which can be enhanced in superatom-based structures due to the following factors: (1) the lack of spherical symmetry in superatoms can lead to anisotropic polarization; (2) the larger size of superatoms compared to atoms results in electron distribution over a broader phase space, making them more easily polarizable; (3) superatoms such as superhalogens and superalkalis have higher electron affinities and lower ionization potentials, respectively, facilitating charge transfer and thereby enhancing polarization.

The early experimental study of ferroelectricity in the gas-phase Nb clusters was conducted by de Heer and colleagues in 2003.<sup>235</sup> They observed ferroelectric-like state in cryogenically cooled neutral Nb<sub>*n*</sub> clusters (*n* = 2–150), with the transition temperature decreasing as cluster size increased. Additionally, pronounced even-odd alternations were noted for *n* > 38. Later, using molecular beam Stern-Gerlach deflection measurement, they found that the odd-*n* clusters deflected due to a single unpaired spin, and spin uncoupling coincided with the transition to ferroelectric state.<sup>236</sup> Coexistence of ferroelectricity and ferromagnetism was also observed in rhodium clusters,<sup>237</sup> where the transition temperatures exhibited size-dependent variations. Similar behavior was also predicted in Ta clusters.<sup>238</sup> The ferroelectric state of free clusters can be further tuned through doping, as demonstrated in Nb clusters doped with Al, Au, Mn, Fe, and Co.<sup>239</sup> These findings open avenues for designing new “multiferroic” materials composed of clusters. An intriguing question arises from these observations: as clusters exist in an intermediate phase between atoms and bulk matter, how many atoms are required to exhibit traditional ferroelectric effect? A recent experimental study determined this threshold to be approximately 5000 atoms in zero-dimensional systems.<sup>240</sup> These advances have strongly motivated research on unconventional ferroelectricity in cluster-based materials.

Using an unbiased structure search combined with DFT calculations, Gao *et al.*<sup>241</sup> theoretically proposed a family of



stable three-dimensional ionic supersalts,  $\text{PnH}_4\text{MX}_4$  ( $\text{Pn} = \text{N}, \text{P}$ ;  $\text{M} = \text{B}, \text{Al}, \text{Fe}$ ;  $\text{X} = \text{Cl}, \text{Br}$ ), constructed from superalkali  $\text{PnH}_4^+$  and superhalogen  $\text{MX}_4^-$  building blocks. The exothermic nature of the reactions  $\text{MPn} + 4\text{HX} \rightarrow \text{PnH}_4\text{MX}_4$  or  $\text{PnH}_4\text{X} + \text{MX}_3 \rightarrow \text{PnH}_4\text{MX}_4$  suggests the potential of efficient synthesis. These supersalts adopt distorted zinc blende (ZB) structures characterized by covalent-like directional bonding induced by the anisotropy induced by the aspherical geometry of the superatoms, as shown in Fig. 9A. Such a unique structural configuration enables ferroelectricity with significant ion displacement, as well as ferroelasticity with ultra-large reversible strain. Notably, the ionic bonding features reduce the switching barriers associated with large deformations, very different from conventional ferroelectric materials which are brittle in nature due to covalent bonding.  $\text{PH}_4\text{FeBr}_4$  is particularly remarkable for its triferroic character, combining ferroelectricity, ferroelasticity, and antiferromagnetism within a single phase, where the spin configuration can be reversibly tuned by ferroelastic or  $90^\circ$  ferroelectric switching. Moreover, a supersalt crystal built exclusively from  $\text{SbCl}_4$  superhalogens is stabilized through valence disproportionation of Sb into  $\text{Sb}^{5+}$  and  $\text{Sb}^{3+}$  species.<sup>242</sup> The coexistence of mixed valence and lone-pair activity generates an unprecedented form of ferroelectricity featuring multiple coupled polarization modes.

This discovery of a purely superhalogen-based supersalt crystal and multimode ferroelectricity underscores the unique role of superatoms in designing novel functional materials.

Experimentally, Ai *et al.*<sup>243</sup> successfully observed ferroelectricity in a fullerene adduct formed by  $\text{C}_{60}$  and  $\text{S}_8$  clusters. The combination of  $\text{C}_{60}$ , the most symmetric molecule with  $I_h$  symmetry, and the highly symmetric  $\text{S}_8$  cluster with  $D_{4d}$  symmetry resulted in the formation of a polar  $\text{C}_{60}\text{S}_8$  adduct with  $C_{2v}$  point group at room temperature.  $\text{C}_{60}\text{S}_8$  undergoes polar-to-polar ferroelectric phase transition, the resulting ferroelectricity was confirmed through the observation of a ferroelectric hysteresis loop and ferroelectric domain switching. This experimental demonstration of ferroelectricity in 3D cluster assemblies is expected to inspire further research and exploration in this field.

For 2D assemblies, Huang *et al.*<sup>244</sup> used *ab initio* molecular dynamics simulations and Monte Carlo methods to propose a 2D magnetic honeycomb lattice material  $\text{TM}_2(\text{C}_{60})_3$  ( $\text{TM} = \text{Mo}$ ,

and W) consisted in exohedral metallofullerene clusters  $\text{TM}(\text{C}_{60})_3$ , where the transition metal atoms deviate from the central plane, leading to a spontaneous electric dipole moment perpendicular to the plane. The energy barrier between the two dipolar states (up and down) was found to be 0.50 and 0.49 eV, respectively, low enough to allow switching between these states under an external electric field (gate voltage). Consequently, this system qualifies as a ferroelectric material with out-of-plane polarization.

For 1D assemblies, Zhao *et al.*<sup>245</sup> computationally designed magnetic fullerene-based 1D chains of  $\text{U}_2\text{C}@\text{C}_{80}\text{-M}$  ( $\text{M} = \text{Cr}, \text{Mn}, \text{Mo}, \text{and Ru}$ ), which exhibit enhanced spontaneous polarization ( $\text{Ps}$ ) compared to isolated clusters. This enhancement ranges from two to four times and is attributed to the interaction between fullerene and metal atoms. The calculated  $\text{Ps}$  values (in  $\text{pC m}^{-1}$ ) for Cr, Mn, Mo, and Ru are 22.77, 34.60, 27.34, and 20.34, respectively, and the energy barrier  $E_B$ 's for polarization switching (in  $\text{meV atom}^{-1}$ ) are 7.98, 17.14, 3.45, and 2.74, respectively.

Compared to the advancements in cluster-based ferroelectricity, much less attention has been given to cluster-based piezoelectric materials. Recently, Wang *et al.*<sup>246</sup> synthesized a superhalogen  $\text{GaCl}_4$  based piezoelectric material  $[(\text{CH}_3)_3\text{NCH}_2\text{Cl}][\text{GaCl}_4]$ , which exhibits an exceptionally high piezoelectric coefficient ( $d_{33} = 226 \text{ pC N}^{-1}$ ) and a remarkable piezoelectric voltage coefficient ( $g_{33} = 1318 \times 10^{-3} \text{ V m N}^{-1}$ ), along with outstanding mechanical durability and stability.

Unlike ferroelectricity and piezoelectricity, significant experimental works have been dedicated to synthesizing cluster-based thermoelectric materials as summarized in ref. 230. However, the maximum figure of merit ( $ZT$ ) values reported so far remain below 1.0, leading to low conversion efficiency. To address this limitation, Li *et al.*<sup>247</sup> studied the recently synthesized superatomic  $\text{Re}_6\text{Se}_8\text{I}_2$  monolayer,<sup>192</sup> depicted in Fig. 9B. Their results reveal a clear phonon–electron decoupling mechanism: the heavy and anharmonic Re–I bonds strongly suppress lattice heat transport, reducing the thermal conductivity to  $1.20 \text{ W m}^{-1} \text{ K}^{-1}$  at 300 K, while the stiff and harmonic Re–Se framework sustains high carrier mobility, producing a TE power factor of  $4344 \text{ } \mu\text{W m}^{-1} \text{ K}^{-2}$  along the  $b$  axis under n-type doping—approximately an order of magnitude greater than that of comparable cluster-based systems.

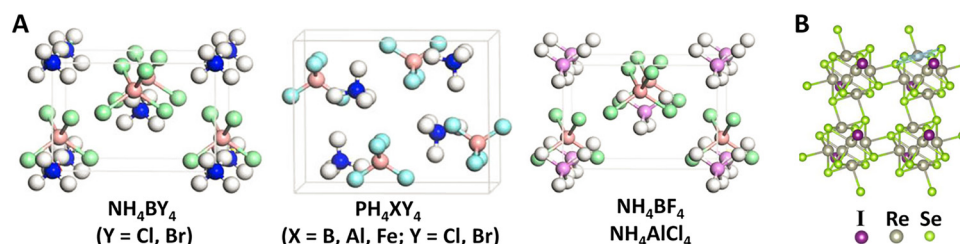


Fig. 9 Functional materials formed by superatomic ions. (A) Typical crystal structures of multiferroic supersalts featured by non-centrosymmetric superatomic ions  $\text{NH}_4^+$  or  $\text{PH}_4^+$  interacting with  $\text{XY}_4^-$  ( $\text{X} = \text{B}, \text{Al}, \text{Fe}$ ;  $\text{Y} = \text{Cl}, \text{Br}$ ). Adapted with permission from ref. 241. Copyright 2021 Springer Nature. (B) Monolayer structure of thermoelectric material  $\text{Re}_6\text{Se}_8\text{I}_2$  composed of superatomic cluster  $\text{Re}_6\text{Se}_8^{2+}$  and two  $\text{I}^-$  in unit cell. Adapted with permission from ref. 247. Copyright 2022 American Physical Society.



Table 6 Summary on superatom-based ferroelectricity, piezoelectricity and thermoelectricity

| Property                                     | Ferroelectricity  | Piezoelectricity   | Thermoelectricity  |
|--|---|--|--|
| Mechanism                                    | Reversible spontaneous electric polarization  | Polarization induced by mechanical stress  | Electrical current generated from a temperature gradient   |
| Required symmetry                            | Non-centrosymmetric with switchable dipoles   | Non-centrosymmetric, polar axis  | No strict symmetry requirement   |
| Tunable features <i>via</i> superatom design | Electric dipole moment; switching field strength  | Piezoelectric coefficient; mechanical compliance; anisotropy of response   | Seebeck coefficient; electrical conductivity; phonon scattering for low thermal conductivity   |
| Role of superatoms                           | <p>(1) The internal dipole moments of superatoms enable strong and switchable polarization due to asymmetric charge distribution within superatoms.</p> <p>(2) The high polarizability of superatoms enhances dielectric constant and strengthens electric field response.</p> <p>(3) The structural flexibility of superatoms allows low-energy switching between polar states.</p> <p>(4) The anisotropic properties of superatoms enable directional polarization switching and control.</p> | <p>(1) The internal dipole moments of superatoms provide built-in polarization that responds to mechanical strain.</p> <p>(2) The high polarizability of superatoms improves electromechanical conversion efficiency.</p> <p>(3) The structural flexibility enhances strain-to-polarization coupling <i>via</i> deformable bonds.</p> <p>(4) The tunable electronic structure of superatoms allows engineered piezoelectric coefficients <i>via</i> electronic deformation.</p> <p>(5) The anisotropic properties of superatoms enhance the directional sensitivity to mechanical stimuli.</p> | <p>(1) The internal dipoles of superatoms can affect carrier distribution.</p> <p>(2) The high polarizability of superatoms modulates carrier behavior and contributes to thermopower.</p> <p>(3) The tunable electronic structure of superatoms boosts Seebeck coefficient and electrical conductivity through density-of-states tuning.</p> <p>(4) The anisotropic properties of superatoms support high power factor along optimal conduction directions.</p> |
| Performance metric                           | Polarization, coercive field  | Piezoelectric coefficient, coupling factor   | Figure of merit  |

Consequently, superatomic  $\text{Re}_6\text{Se}_8\text{I}_2$  exhibits outstanding thermoelectric performance, with  $ZT$  values reaching 1.20 at 500 K and 1.43 at 900 K. As summarized in Table 6, the distinct merits of superatom-based ferroelectricity, piezoelectricity, and thermoelectricity underscore the immense potential of clusters in designing novel functional materials.

## VII. Conclusion and outlook

In summary, we have reviewed the recent progress in elucidating the unique properties of superatomic clusters across disciplines and their potential as building blocks of novel materials. For example, boron-based superatomic clusters with unprecedented stability as multiply charged ions can enable noble gas atoms to react at room temperature. Distinctive properties of the superatoms, such as the non-spherical symmetry, variable chemical compositions, internal charge distribution, as well as extraordinary electronic structure and interaction, can render novel properties of the host materials and provide new understanding of the structure–property relationship. For example, a supersalt can be formed by only using a superhalogen anion such as  $\text{SbCl}_4$  as the building block. Superalkalis can be used to create a new class of electrides with enhanced catalytic properties while single-superatom catalysts can outperform single-atom catalysts. Superatoms can be used as building blocks of novel thermoelectric, piezoelectric, and multiferroic materials as well as materials for energy production and storage. Substitution of regular atomic ions by selected superatomic ions in energy materials can enhance both transport properties and stability of the bulk and interface. These

achievements show that the future of this field is bright and, with new design and synthesis paradigm, superatoms with unique functionalities can unravel new science and applications.

There are challenges as well as opportunities. While considerable work over the past two decades have showcased numerous superhalogens and superalkalis, similar progress has not been made in the design and synthesis of superatoms that mimic the chemistry of transition metal, noble metal, and rare-earth metal atoms. This is important because of the importance of these materials in technology such as catalysis and magnetism. For example, significant amount of work has been done on single atom-catalysts yet work on single-superatom catalysts is in its infancy. Work using ligands to manipulate the properties of core metal clusters is important for their applications in catalysis, magnetism, biomedical applications, and optical materials.

In the field of superatom-based materials, although many recent experiments have confirmed and supported the theoretical designs and mechanisms, many more systematic syntheses and characterizations are needed to further narrow the gap between the prototype modeling and experimental realization. In practice, doping superatomic ions into materials is often limited by the metastability and the creation of impurity phases. Besides optimizing the known synthetic routes, other non-equilibrium synthetic pathways are worth exploring to kinetically stabilize the metastable phases, such as laser deposition and ablation,<sup>248–251</sup> high-temperature quenching,<sup>252</sup> and high-energy ball milling.<sup>253</sup> Beyond conventional techniques to characterize structural features, such as the X-ray diffraction



and neutron scattering, optical spectroscopy, electron microscopy, and solid-state NMR, probing the dynamics of the superatomic ions inside a material would require development of characterization techniques with improved spatial and temporal resolutions. Meanwhile, new computational tools and machine-learning methods<sup>254</sup> are needed to expedite property-oriented materials discovery based on different superatoms, as well as to enable accurate evaluation of phase stability. Large-scale molecular dynamics simulations are also needed to best capture the compositional and defect complexity possibly introduced by superatomic dopants in the bulk and interface.

Different from the conventional atom-based crystals, superatomic crystals have complex geometries with large unit cells and manifold orientations, resulting in strong phonon scattering (including higher order phonon scattering) and phonon coherence. To accurately calculate the phonon transport in superatomic crystals, machine-learning-based potentials are highly desirable. Such potentials are efficient for simulation but challenging for fitting, which is also true for simulating superatom-based thermoelectric materials involving both phonons and electrons. Accurate electronic band structure is also needed for calculating the figure of merit to measure the thermoelectric performance.

In any field of science, a common question to ask is about its future. For cluster science in general and superatoms in particular, the answer that comes to our mind is same as the one Walt Disney gave when asked about the future of Disney World. His answer, that Disney World will survive as long as there is human imagination, may very well apply to cluster science.

## Author contributions

P. J., H. F., and Q. S. proposed the topic of the Review. All authors wrote and discussed the manuscript.

## Conflicts of interest

The authors declare no competing interests.

## Data availability

The data underlying this study are available in the published article.

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