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## Structure, stability, reactivity and bonding in noble gas compounds

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Noble gases (Ngs) are recognized as the least reactive elements due to their fully filled valence electronic configuration. Their reluctance to engage in chemical bond formation necessitates extreme conditions such as low temperatures, high pressures, and reagents with high reactivity. In this Perspective, we discuss our endeavours in the theoretical prediction of viable Ng complexes, emphasizing the pursuit of synthesizing them under nearly ambient conditions. Our research encompasses various bonding categories of Ng complexes and our primary aim is to comprehend the bonding mechanisms within these complexes, utilizing state-of-the-art theoretical tools such as natural bond orbital, energy decomposition, and electron density analyses. These complex types manifest distinct bonding scenarios. In the non-insertion type, the donor–acceptor interaction strength hinges on the polarizing ability of the binding atom, drawing the electron density of the Ng towards itself. In certain instances, especially with heavier Ng elements, this interaction reaches a magnitude where it can be considered a covalent bond. Conversely, in most insertion cases, the Ng prefers to share electrons to form a covalent bond on one side while interacting electrostatically on the other side. In rare cases, both bonds may be portrayed as electron-shared covalent bonds. Furthermore, a host cage serves as an excellent platform to explore the limits of achieving Ng–Ng bonds (even for helium), under high pressure.

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### Introduction

In the grand tapestry of chemical elements, noble gases (Ngs) have long occupied a unique and seemingly immutable position. Their full valence electron shells, a testament to the octet



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rule, were once thought to render them chemically inert and isolated from interatomic bonding. This perception, preserved in textbooks and reinforced by decades of seemingly inert behaviour, stood as a cornerstone of chemical understanding. However, high-pressure synthesis techniques and sophisticated theoretical calculations have cracked this facade. In terms of electronic configurations, they are characterized by fully occupied valence orbitals,  $ns^2np^6$  for all except helium, which has  $1s^2$ . These electronic configurations offer exceptional stability and impede their proclivity to readily engage in chemical bonding with other elements, thereby designating them as “inert gases.” The pronounced elevation of their first ionization potentials (IPs), showcasing a gradual decrease from helium to radon,<sup>1</sup> further reinforces their reputation for chemical inertness. Nonetheless, this decrease in IP signifies that the outermost electrons of the heavier noble gases, notably xenon, become progressively more vulnerable to external influences, paving the way for the plausible formation of noble gas compounds.

Before the successful synthesis of noble gas compounds, several prominent chemists theorized the possibility of their formation based on atomic properties and periodic trends. In 1916, Kossel predicted the existence of xenon and krypton fluorides based on their loosely bound outermost electrons and fluorine's high electronegativity.<sup>2</sup> Leveraging ionic radii data, Pauling predicted the potential formation of xenon fluorides ( $XeF_6$  or  $XeF_8$ ) and xenic acid ( $H_4XeO_6$ ).<sup>3</sup> In 1962, Bartlett's intuition regarding the resemblance between the first IPs of  $O_2$  and Xe yielded fruitful results. While undertaking a reaction between dioxygen and  $PtF_6$  to form  $O_2^+PtF_6^-$ ,<sup>4</sup> he made the said realization and immediately aimed to form an  $Xe^+PtF_6^-$  complex by reacting Xe and  $PtF_6$ .<sup>5</sup> This reaction resulted in the precipitation of an orange-yellow solid. This landmark discovery shattered the longstanding notion of noble gas inertness, marking the inception of the “Noble Gas Chemistry” era. Initially presumed to be  $Xe^+PtF_6^-$ , the formula of this first Ng compound underwent revision upon X-ray powder diffraction (XRPD) analysis. The actual product unveiled itself as  $XeF^+Pt_2F_{11}^-$ , a more intricate species than Bartlett's envisioned simple adduct.<sup>6</sup> The reaction mechanism, as delineated by Christie's quantum-chemical calculations, implicated  $F^-$  ion catalysis, yielding a mixture comprising  $XeF^+PtF_6^-$ ,  $PtF_5$ , and the ultimate product,  $XeF^+Pt_2F_{11}^-$ .<sup>7,8</sup>

In the subsequent years, a surge of investigative endeavours resulted in the synthesis and characterization of numerous xenon compounds, encompassing  $XeF_2$ ,  $XeF_4$ ,  $XeF_6$ ,  $XeOF_4$ , and  $XeO_3$ . Krypton, too, participated in this chemical exploration with the introduction of  $KrF_2$ .<sup>9,10</sup> Xenon, distinguished by its loosely bound outer electrons, emerged as the focal point of scrutiny. A multitude of scholarly publications detailed the existence of species such as  $XeF_{2-6}$ ,  $XeOF_4$ ,  $XeO_3$ , and others.<sup>11-16</sup> Even the radioactive radon found recognition, forming compounds like  $RnF_2$  and  $[RnF][Sb_2F_{11}]$ .<sup>17</sup> The beginning of the 21st century marked another watershed moment: the isolation of the first argon compound,  $HArF$ , accomplished by Räsänen's research group within a low-temperature

matrix.<sup>18,19</sup> Subsequent to this achievement, the identification of various weak neon complexes, including  $NeAuF$  and  $(NeAr)-Be_2O_2$ , expanded the horizons of Ng reactivity.<sup>20-23</sup> In the year 1925, the  $HeH^+$  cation was observed by Hogness and Lunn.<sup>24</sup> Later, Dong *et al.* demonstrated that, under high pressures, even helium can engage in chemical bonding, resulting in the formation of the  $Na_2He$  compound.<sup>25</sup>

Presently, it is established that all Ng elements exhibit chemical reactivity, albeit contingent on specific conditions. The trajectory from inertness to reactivity persists, with ongoing research continuously expanding the frontiers of noble gas chemistry and revealing its diverse applications in domains such as anaesthesia, nuclear waste storage, and materials science.

## Bonding pattern

The bonding in compounds involving Ng elements can be categorized into three primary types, namely, (a) non-insertion  $NgAB$ , (b) insertion  $ANgB$ , and (c)  $Ng_n@cage$  complexes.

Ng adducts, where the binding of the Ng atom occurs at an exposed end of a molecule (say,  $AB$ ), constitute the non-insertion type of compounds. The interaction between Ng and A stems from the ability of the A atom to induce polarization in the electron cloud of the outermost orbital of the Ng, leading to the formation of an attractive donor-acceptor bond. This polarization is contingent upon the polarizing power of A, influenced by its size and charge. Smaller, highly charged A atoms result in more pronounced polarization and, consequently, stronger Ng-A interactions. Additionally, the difference in electronegativity between A and the counterion, B, plays a pivotal role. A substantial difference induces a dipole in the  $AB$  molecule, intensifying the polarization of the Ng atom. It is to be noted that there is a heightened effectiveness in inducing polarization by ionic counterions ( $B^-$ ) compared to that by the neutral ones. Finally, in neutral  $AB$  molecules with minimal electronegativity differences, dispersion forces contribute to bonding, albeit to a lesser degree. This intricate interplay of factors underscores the multifaceted nature of Ng-A interactions, rendering these adducts a compelling focal point in contemporary chemistry research.

For the insertion complex, the Ng atom inserts itself within the A-B bond to form an  $ANgB$  complex. The formation of this complex inherently disrupts the pre-existing A-B bond, presenting a thermodynamic challenge. While the collective interactions of A-Ng and Ng-B do contribute, they often fall short of fully compensating for the lost A-B stability. In contrast, the stability of  $NgAB$  primarily hinges on the thermodynamics of the Ng-A bond. Vigorous donor-acceptor interactions between the Ng atom and strongly polarizing centers in A contribute to their heightened stability. The insertion complex,  $ANgB$  species, tends to be metastable, and its existence is frequently contingent on kinetic factors. In-depth thermochemical analyses illuminate this kinetic reliance. The consideration of

extensive dissociation pathways is imperative, with two predominant routes prevailing (others are mostly endergonic),  $\text{ANgB} \rightarrow \text{Ng} + \text{AB}$  (highly exergonic) and  $\text{ANgB} \rightarrow \text{A} + \text{Ng} + \text{B}$  (occasionally weakly exergonic). Evaluating the stability of  $\text{ANgB}$  entails assessing the activation energy barriers of these dissociation pathways.

Beyond classical  $\text{NgAB}$  and  $\text{ANgB}$  complexes, a captivating domain unfolds where noble gas chemistry relies on the influences generated by the restricted environment facilitated by specific cages. Instances include  $\text{Ng}_2$  dimers enclosed within various hollow cages, such as fullerene, borospherene,  $\text{C}_{20}\text{H}_{20}$ ,  $\text{B}_{12}\text{N}_{12}$ ,  $\text{B}_{16}\text{N}_{16}$ , octa acids, cucurbit[ $n$ ]uril, clathrate hydrates, and carbon nanotubes doped with BN. Our research group has extensively investigated these systems over the years. In this context, the confinement itself narrates the structure, bonding, and reactivity of the entrapped  $\text{Ng}_2$  guests. We systematically analyze them along with their confined movements within the cages, comparing them to their unconfined counterparts.

## Computational details

Designing Ng compounds requires careful computational optimization. Selecting the right method is crucial, aiming for minimal errors in predicting geometry, electronic properties, and energy. For highly accurate calculations, especially on small systems, *ab initio* methods like CCSD(T) with advanced basis sets are ideal. However, for larger systems, computationally expensive CCSD(T) becomes impractical, necessitating DFT-based methods. Benchmark studies suggest specific DFT levels like MPW1B95/6-311+G(2df,2pd), BMK/aug-cc-pVTZ, DSD-BLYP/aug-cc-pVTZ, and B2GP-PLYP/aug-cc-pVTZ levels, which offer excellent accuracy in calculating Ng bond energies. Notably, MP2 with aug-cc-pVDZ shows less accuracy. Relativistic effects of heavier elements like Xe and Rn are often handled with effective core potentials or ZORA approximations. Interestingly, Ng insertion compounds are particularly sensitive to the chosen level of theory. Structures stable at DFT or MP2 levels can sometimes dissociate at the higher-level CCSD(T), making these high-level calculations essential for predicting truly stable Ng-inserted molecules. Conversely, non-insertion type systems are less sensitive, with only bond dissociation energies changing slightly across different levels of theory. It is highly unlikely for a stable species at one level to become unstable at another in these cases.

The natural population analysis (NPA)<sup>26</sup> scheme is used in NBO<sup>27</sup> to determine atomic charges ( $q$ ), followed by the computation of Wiberg bond indices (WBI)<sup>28</sup> to assess the formation of potential bonds between atoms. Utilizing second-order perturbation allows for a qualitative understanding of interactions between filled Lewis-type NBOs ( $i$ , acting as donors) and empty non-Lewis NBOs ( $j$ , acting as acceptors). The stabilization energy of delocalization from  $i$  to  $j$  is given by eqn (1).

$$E(2) = \Delta E_{ij} = q_i \frac{F(i,j)^2}{\varepsilon_j - \varepsilon_i} \quad (1)$$

where  $q_i$  and  $\varepsilon_i$ ,  $\varepsilon_j$  are the occupancy of the donor orbital and the orbital energies, respectively and  $F(i, j)$  is the off-diagonal NBO

Fock matrix element. Enhancing the interaction between the donor and the acceptor within NBO involves the removal of the Fock matrix element followed by re-diagonalization.<sup>29,30</sup> The calculations are carried out using the Gaussian program.<sup>31</sup>

The bonding nature is further elucidated through electron density (ED,  $\rho(r)$ ) topological analysis using Bader's quantum theory of atoms-in-molecules (QTAIM)<sup>32</sup> method in the Multiwfn software.<sup>33</sup> This analysis involves utilizing  $\nabla\rho(r)$  and  $\nabla^2\rho(r)$  to delineate atomic regions and interatomic bond paths. Specifically, the bond path is identified as the one with the maximum gradient of the ED starting and ending in nuclei passing through the bond critical point (BCP). Covalent bonding is typically indicated by high  $\rho(r_c)$  and negative  $\nabla^2\rho(r_c)$  values, and non-covalent bonding by low values of  $\rho(r_c)$  coupled with positive  $\nabla^2\rho(r_c)$ . In the case of heavy atoms, a negative total electron energy density ( $H(r_c)$ ) alongside  $G(r_c)/\rho(r_c)$  approximately equal to 1 suggests partial covalency ( $G(r_c)$  denotes the local kinetic energy density).

Energy decomposition analysis (EDA)<sup>34</sup> is conducted in conjunction with the natural orbitals for the chemical valence (NOCV)<sup>35</sup> method, employing the ADF 2013.01 program<sup>36</sup> package, to provide further insights into the bonding scenario. The interaction between the prepared fragments is quantified as the intrinsic interaction energy ( $\Delta E_{\text{int}}$ ), consisting of four distinct energy components: electrostatic energy ( $\Delta E_{\text{elstat}}$ ), orbital interaction energy ( $\Delta E_{\text{orb}}$ ), dispersion interaction energy ( $\Delta E_{\text{disp}}$ ), and Pauli's repulsion energy ( $\Delta E_{\text{Pauli}}$ ). While the first three terms contribute to the total attractive interaction energy, the fourth one represents a repulsive force.

$$\Delta E_{\text{int}} = \Delta E_{\text{elstat}} + \Delta E_{\text{orb}} + \Delta E_{\text{disp}} + \Delta E_{\text{Pauli}} \quad (2)$$

$\Delta E_{\text{elstat}}$  reflects the quasi-classical electrostatic interaction between the unperturbed EDs of the interacting fragments, while  $\Delta E_{\text{Pauli}}$  quantifies the repulsive energy when it transforms into a wavefunction superposition. The resultant wavefunction has an energy penalty due to the Pauli's exclusion principle.  $\Delta E_{\text{orb}}$  encompasses the energy stabilization gained through electron sharing, polarization, and charge transfer between fragments. In the NOCV methodology,  $\Delta E_{\text{orb}}$  is further delineated as the summation of pairwise orbital interaction energies ( $\Delta E_{\text{orb}}^k$ ) associated with the pairwise charge contributions ( $\Delta\rho^k(r)$ ) (eqn (3)).

$$\Delta E_{\text{orb}} = \sum_k \Delta E_{\text{orb}}^k \quad (3)$$

Finally,  $\Delta E_{\text{disp}}$  considers the long-range interactions attributed to the instantaneous polarization multipoles induced within the molecules. A few other important studies regarding the bonding interactions within Ng complexes are reported in the literature.<sup>37–40</sup>

## Discussion

### Non-insertion complexes

Inspired by Pauzat *et al.*'s findings<sup>41–43</sup> on triangular  $\text{H}_3^+$  clusters interacting with noble gases, our research team has

computationally demonstrated that  $\text{H}_3^+$  can form remarkably strong bonds with a maximum of three Ng atoms.<sup>44</sup> Notably, the first bond between H and Ng in  $\text{H}_3(\text{Ng})^+$  displayed significantly higher strength compared to larger clusters. This stability was linked to delocalization of the cationic charge, as revealed by conceptual DFT-based descriptors. Interestingly,  $\text{Li}_3^+$  offered a contrasting picture. While each Li center can bind with an Ng atom forming  $\text{Li}_3\text{Ng}_3^+$ , the bonds of Ng with Li are notably weaker than that with H atoms observed in the  $\text{H}_3^+$  species. Comparing it to previously reported  $\text{NgLiH}$  and  $\text{NgLiF}$ , the Li–Ng bond strength turns out to be similar.<sup>45–49</sup> Motivated by the intriguing interaction of Ng in such species and Li's potential for hydrogen storage, a comprehensive study explored Ng interactions with star-shaped  $\text{C}_5\text{Li}_7^{+50}$  and  $\text{O}_2\text{Li}_5^{+51}$  clusters to check how far Ng atoms can replace  $\text{H}_2$  molecules.<sup>52,53</sup> The former can bind up to seven helium atoms, with other Ng atoms (Ne–Xe) reaching a maximum of 12. Similarly,  $\text{O}_2\text{Li}_5^+$  hosted up to seven Ng atoms (He–Xe). Further comparative analyses unveiled that, in contrast to helium and neon, which exhibit slightly weaker interactions with  $\text{C}_5\text{Li}_7^+$  and  $\text{O}_2\text{Li}_5^+$  clusters compared to hydrogen molecules, heavier noble gases display higher bond strength.<sup>47,48</sup> This intriguing phenomenon extends beyond specific Li-decorated clusters. Our examination of various Li-decorated and super-alkali clusters consistently reaffirmed that clusters with highly positively charged Li centers tend to exhibit an inclination for bonding with noble gases.<sup>52,54</sup> Applying an external electric field further strengthened the Li–Ng bonds.

The higher ionic potential of Be compared to Li affords it a notably stronger affinity for these elusive Ng elements. As early as 1988, Frenking *et al.* theoretically predicted the potential formation of  $\text{NgBeO}$  complexes,<sup>55</sup> which materialized into experimental reality in 1994 when Andrews *et al.* employed the pulsed-laser matrix isolation technique to detect these Ar, Kr and Xe analogues.<sup>56</sup> Motivated by these revelations, we have explored some additional viable Ng–Be complex structures, *viz.*,  $\text{CN}_3\text{Be}_3^+$ , featuring two distinct types of beryllium centers. While each ring-connected beryllium can accommodate a single Ng atom, the lone beryllium outside the ring can interact with two, resulting in a total of four Ng partners. Our computational analyses suggest that the  $\text{NgCN}_3\text{Be}_3^+$  cluster may feasibly exist at room temperature for lighter noble gases (He–Ar), while colder temperatures may be required for the heavier ones (Kr–Rn). Significantly, the formation of a nearly half-bond between Be and heavier noble gases (Kr–Rn) is indicated by high Wiberg Bond Index (WBI) values. Apart from  $\text{BeO}$ , the entire class of  $\text{NgBeX}$  compounds, where  $\text{X} = \text{O}, \text{S}, \text{Se}, \text{Te}$ , and their positively charged counterparts are explored, where the latter forms exhibited a heightened affinity for Ng atoms compared to their neutral counterparts when considering a specific X atom. Moreover, within a given noble gas, the bond strength with Be diminished as we progressed from O to Te. Apart from these, the global minimum (GM) energy structures of  $\text{Be}_2\text{N}_2$ ,  $\text{Be}_3\text{N}_2$ , and  $\text{BeSiN}_2$  species were explored.<sup>57</sup> Markedly positive Be centers in these structures serve as attractive binding partners for the Ng atoms.  $\text{Be}_2\text{N}_2$  accommodates two Ng atoms,  $\text{Be}_3\text{N}_2$  hosts three, and  $\text{BeSiN}_2$  binds one, all through

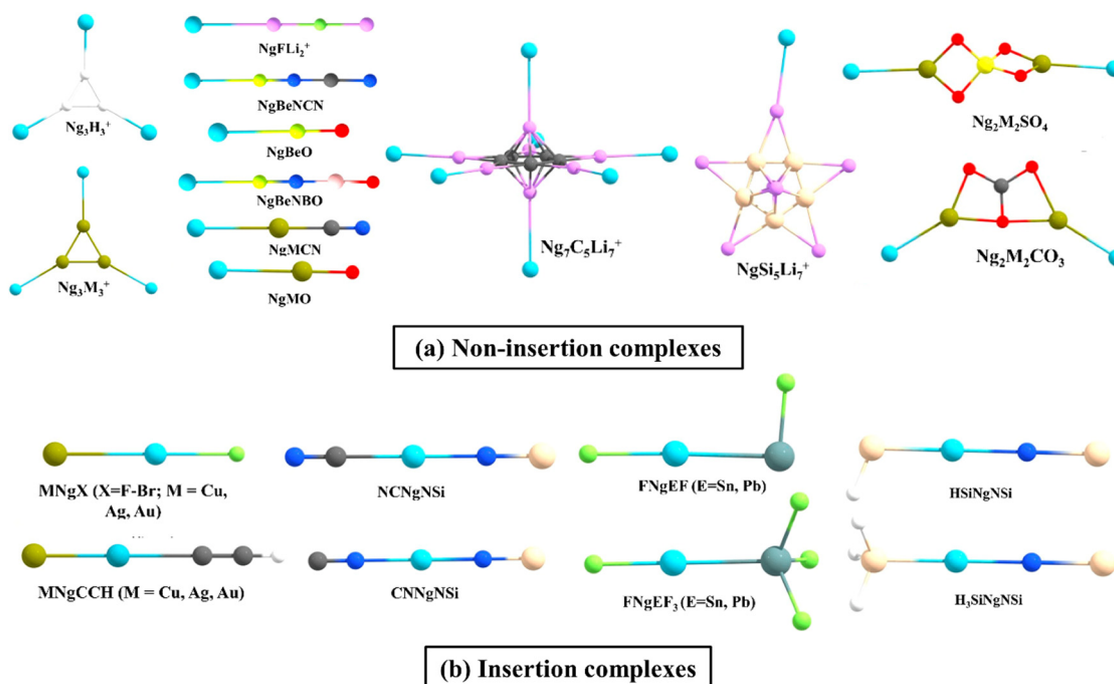


Fig. 1 A few examples of the (a) non-insertion NgAB and (b) insertion ANgB type complexes [reproduced from ref. 88 with permission from MDPI, © 2019].

their Be centers. Notably, the presence of Ng can alter the energy rankings of these isomers. For instance, the linear (GM) structure of BeSiN<sub>2</sub> has Be in the center, compared to the next higher energy isomer with Be situated terminally (Fig. 1a). This energy positioning changes in the presence of heavier Ngs (Ar–Rn), as the second isomer optimally positions the Be atom terminally (compared to the central position of Be in the former) for a more robust interaction with the Ng atoms.

Attaching a Lewis acid, like BH<sub>3</sub>, to the oxygen center of BeO, as proposed by Grandinetti *et al.* enhances the interaction with Ng atoms. Similarly, substituting the H atom in BeNH with various groups, as observed by the same authors, proved effective in augmenting its Ng binding capabilities.<sup>58</sup> Nevertheless, a trade-off was evident, as these modified systems did not represent the most stable configurations. Identifying two Be-containing molecules, BeNCN and BeNBO, proved pivotal as they achieved both exceptional stability (as both are GM structures) and robust Ng binding (highest and second-highest among reported neutral Be systems). These molecules, formed by replacing the H atom in BeNH with CN and BO, respectively, strategically position the Be atom terminally for optimal interaction with the Ng partners.<sup>59</sup> In the pursuit of ideal Be-based salts for strong Ng–Be bonds, our investigation delved into the Ng-binding capacity of various BeX systems, where X = CO<sub>3</sub>, SO<sub>4</sub>, CrO<sub>4</sub>, and HPO<sub>4</sub>.<sup>60,61</sup> A comparative analysis unveiled that NgBeSO<sub>4</sub> forms the most potent bonds with Ngs in contrast to NgBeCO<sub>3</sub> and NgBeO (with the exception of He–Be).

The interactions between Ngs and 1-tris(pyrazolyl)-borate salts of Be and Mg are also studied.<sup>62</sup> Be demonstrates superior affinity for Ngs compared to Mg because of the former's smaller size and higher charge density, enabling a stronger polarizing force on the Ng electron cloud. Moving beyond the Be–Mg duo, we explored half-sandwich complexes where Ngs (He–Rn) reside atop metal cations (M = Be–Ba) with a cyclopentadienyl ring (Cp).<sup>63</sup> These NgMCP<sup>+</sup> complexes exhibited a hierarchy of bond strengths where, for a specific Ng, the interaction with Be was the most robust, followed by Mg, Ca, Sr, and Ba. Interestingly, for a fixed metal, the bond strength increased with the Ng atomic mass, from He to Rn. To understand the underlying mechanism of this interaction, we analyzed deformation densities related to the orbital contributors. The primary interaction was characterized by Ng p<sub>σ</sub> orbitals donating σ electron density to the BeCp<sup>+</sup> complex, followed by the π-donation from Ng p<sub>π</sub> orbitals to BeCp<sup>+</sup>.

We expanded our scope beyond beryllium and magnesium to explore the binding abilities of Ngs with group 14 elements as well. The EX<sub>3</sub><sup>+</sup> system, where E = elements of group 14 and X = H, F, Cl, Br, exhibited substantial yet comparatively lower attraction towards noble gases than boron in B<sub>3</sub><sup>+</sup>.<sup>64,65</sup> Notably, EH<sub>3</sub><sup>+</sup>, for a given noble gas, showed the order C > Si > Ge in terms of binding strength. The substitution of a hydrogen atom in EH<sub>3</sub><sup>+</sup> with X (F, Cl, Br) introduced a complex interplay of two opposing factors: the inductive (–I) effect of X enhanced the Lewis acidity of E, while the π-backdonation from X to E had a negative effect on its ability to bind Ngs. Notably, while SiF<sub>3</sub><sup>+</sup> and GeF<sub>3</sub><sup>+</sup> exhibited higher affinity for Ngs compared to their

EH<sub>3</sub><sup>+</sup> counterparts; due to the highly efficient F → C π-backdonation in CF<sub>3</sub><sup>+</sup>, its Ng binding ability significantly diminished compared to CH<sub>3</sub><sup>+</sup>. Further findings revealed that EH<sub>3</sub><sup>+</sup> (E = Si, Ge) and EF<sub>3</sub><sup>+</sup> (E = Si, Ge, Sn, Pb) can effectively bind two Ng atoms simultaneously, and their strong Lewis acidity extends to the Lewis base CO.<sup>66</sup>

Among the transition metal (TM) compounds, particularly superhalogens MF<sub>3</sub> (M = Ru–Ag and Os–Au in the second and third row TM series, respectively), RuF<sub>3</sub> and AuF<sub>3</sub>, exhibited the highest and the second highest affinity for binding with Xe, respectively.<sup>67</sup> Subsequent studies revealed that the coinage metals (Cu, Ag, and Au) demonstrated very high Ng-binding abilities, generally surpassing those of other TMs. The σ-aromatic coinage metal clusters M<sub>3</sub><sup>+</sup> and their M<sub>3</sub>Ng<sub>3</sub><sup>+68</sup> complexes exhibited a bond dissociation energy range of 2.2–19.0 kcal mol<sup>–1</sup> for the Ng–M bonds, following the order, Au > Cu > Ag. Unlike the orbital-dominated bonding observed in the main group elements, these Ng–M bonds derived nearly equal strength from both coulombic and covalent contributions. The crucial orbital factor stabilizing the Ng–M bonds originated from electron donation from a filled p<sub>Ng</sub> to the LUMO<sub>M3</sub><sup>+</sup>. The strong interactions between noble gases and coinage metals (also known as noble metals) uncovered within compounds like NgMNO<sub>3</sub>, NgCu(NO<sub>3</sub>)<sub>2</sub>, NgMSO<sub>4</sub>, Ng<sub>2</sub>M<sub>2</sub>SO<sub>4</sub>, NgCuCO<sub>3</sub>, Ng<sub>2</sub>M<sub>2</sub>CO<sub>3</sub>, NgMCN, NgMO, and [Ng–M(bipy)]<sup>+</sup> complexes<sup>69–74</sup> revealed a partial covalent interaction in the Ng–M bonds. For [Ng–M(bipy)]<sup>+</sup> complexes, finding a suitable counterion like [SbF<sub>6</sub>]<sup>–</sup> that stabilizes the complex while maintaining the strength of the Ng–M bond becomes crucial for successful synthesis.

### Insertion complexes

Continuing our investigation into noble gas interactions, we explore compounds, inspired by the works of Merino *et al.*,<sup>75,76</sup> which provided insights into HNgY molecules (Y = CCH, CN, NC, F, Cl, Br, I and Ng = Xe, Rn), containing electron-sharing H–Ng bonds as well as ionic Ng–Y interactions. These systems can be conceptualized as Ng<sup>+</sup> interacting with [H · Y]<sup>–</sup>, forming a polar covalent bond. Despite their metastable nature, they exhibit significant kinetic barriers against dissociation, preventing immediate breakdown into Ng and the parent molecule.

We subsequently examined compounds like H<sub>3</sub>SiNgNSi and HSiNgNSi (Ng = Xe, Rn).<sup>77</sup> In both the cases, the free energy change associated with dissociation pathways leading to free Ng and the parent compound is negative, implying thermodynamic stability. While the 2B path is exergonic, the 3B dissociation channel appears endergonic for the Rn analogues, and the Xe variant exhibits a marginally exergonic process, although calculations at lower temperatures suggest endergonicity. Negative ∇<sup>2</sup>ρ(r<sub>c</sub>) and H(r<sub>c</sub>) values, coupled with a high ELF at the Si–Ng bond critical point, reveal its covalent nature. In contrast, the Ng–N bond displays ionic characteristics. (H<sub>3</sub>SiNg)<sup>+</sup>(NSi)<sup>–</sup> and (HSiNg)<sup>+</sup>(NSi)<sup>–</sup> bonding representations turn out to be the most accurate. Energy decomposition analysis (EDA)<sup>78,79</sup> further corroborates these findings, with ΔE<sub>orb</sub> as

the dominant contributor to the Si–Ng bond and  $\Delta E_{\text{elstat}}$  playing a significant role in the Ng–N interaction. Computed activation energy barriers suggest that  $\text{H}_3\text{SiNgNSi}$  possesses sufficient kinetic stability for detection at 250–300 K, whereas  $\text{HSiNgNSi}$  requires a relatively cold environment (150–200 K).

Expanding our exploration of these unique Ng compounds, we reported the first set featuring E–Ng covalent bonds (E = Sn, Pb and Ng = Kr, Xe, Rn) in  $\text{FNgEF}_3$  and  $\text{FNgEF}$  compounds.<sup>80</sup> These systems also exhibit metastability, with only the Ng release path being highly exergonic with substantial energy barriers ensuring their protection. Natural bond orbital (NBO) analysis,<sup>81</sup>  $H(r_c)$  values,<sup>82</sup> and EDA calculations collectively indicate a covalent character in the Ng–E bonds and ionic in the Ng–F bonds. The significant role of superhalogens in stabilizing Ng insertion compounds was demonstrated in  $\text{HNgY}$  (Y =  $\text{BO}_2$ ,  $\text{BF}_4$ ) molecules by their enhanced stability compared to the halogen analogues.<sup>83</sup> Building on this insight, we explored the efficacy of  $\text{BeF}_3$ , a potent superhalogen, to stabilize  $\text{HNgBeF}_3$  (Ng = Ar–Rn) complexes.<sup>84</sup> Interestingly, the 2B dissociation path ( $\text{HNgBeF}_3 \rightarrow \text{Ng} + \text{HBeF}_3$ ) is exergonic for all Ng variants, indicating thermodynamic stability. In contrast, the 3B path ( $\text{HNgBeF}_3 \rightarrow \text{H} + \text{Ng} + \text{BeF}_3$ ) and an additional identified route ( $\text{HNgBeF}_3 \rightarrow \text{Ng} + \text{HF} + \text{BeF}_2$ ) are endergonic, with the latter revealing a two-step process upon further scrutiny. Notably, Xe and Rn analogues might exhibit half-lives of up to  $10^2$  seconds at 100 K, further supporting their viability. Thorough bonding analysis reveals the covalent nature of H–Ng bonds and the ionic character of Ng–F interactions, presenting the most accurate representation of the molecule as  $(\text{HNg})^+(\text{BeF}_3)^-$ .

For a unique class of compounds,  $\text{NCNgNSi}$  (Ng = Kr–Rn), representing the first observed instance of the C–Ng–N unit with covalent bonds on either side of Ng, a deviation from the typical  $\text{XNgY}$  insertion molecules represented as  $\text{X}^+(\text{NgY})^-$  was explored.<sup>85</sup> The bonds of Ng with both C and N in  $\text{NCNgNSi}$  exhibit electron-sharing characteristics. All dissociation channels, except for  $\text{NCNgNSi} \rightarrow \text{Ng} + \text{CNSiN}$ , are endergonic for Ng = Xe and Rn. Remarkably, the 3B dissociation of  $\text{NCKrNSi}$  has low exergonicity at room temperature, but lower temperatures will render it endergonic again. A gradual increase in free energy barriers of the 2B dissociation pathway from Kr to Rn (25.2 to 39.3 kcal mol<sup>−1</sup>) suggests that  $\text{NCNgNSi}$  (Ng = Xe, Rn) systems are viable candidates for detection under ambient conditions, while the Kr analogue might require colder temperatures. Covalent C–Ng and Ng–N bonds are confirmed by substantial WBI (>0.5),  $H(r_c) < 0$ , and EDA, providing compelling evidence for this bonding picture. Additional analyses using AdNDP reinforce this description by identifying a delocalized 3c–2e  $\pi$ -bonding associated with the C–Ng–N unit and a completely delocalized 5c–2e  $\pi$ -bond. These findings solidify the unique bonding characteristics of this fascinating molecule. Intriguingly, Ng dissociation from  $\text{NCNgNSi}$  leads to  $\text{CNSiN}$  (the higher energy isomer), instead of the more stable  $\text{NCNSi}$ . However, the presence of Ngs significantly lowers the barrier for this transformation, facilitating the detection of the less stable isomer and emphasizing the role of Ngs in

manipulating the energy landscape of these systems. More recently, another Ng insertion complex with the C–Ng–N motif,  $\text{HCCNgNSi}$  (Ng = Kr, Xe, Rn), has been studied.<sup>86</sup> This Ng insertion within the parent compound  $\text{HCCNSi}$  compound helped realize the other isomer,  $\text{HCCSiN}$ , which was experimentally undetected. The most probable explanation behind this is that the Ng insertion within the C–N bond provided enough space between the HCC and NSi for the latter to rotate to SiN and form the  $\text{HCCNgSiN}$  complex, which dissociates to form Ng and  $\text{HCCSiN}$ . A DFT-based investigation was also performed on the stability and bonding of the novel complex,  $\text{XNgOPO}(\text{OH})_2$  (X = F, Cl, Br; Ng = Kr, Xe, Rn).<sup>87</sup>

### Confined complexes

The intriguing domain of noble gas chemistry takes a compelling turn when confinement factors into the equation. Fullerenes, particularly  $\text{C}_{60}$ , stand out as versatile tools to investigate the impact of confinement on these traditionally inert elements. The encapsulation of two noble gas atoms within the  $\text{C}_{60}$  cage ( $\text{Ng}_2@C_{60}$ ) provides an exceptional platform for examining bond formation within spatial constraints. Krapp and Frenking's ground-breaking discovery<sup>89</sup> revealed that confinement within  $\text{C}_{60}$  substantially reduces the bond distance in  $\text{Xe}_2$  compared to the free molecule, surpassing even that of free  $\text{Xe}_2^{2+}$ . Bonding analysis attributes this phenomenon to the potent steric pressure applied by the cage, bringing the Ng atoms into closer proximity, which encourages orbital overlap, leading to negative  $H(r_c)$  values at the BCP for Ar, Kr, and Xe. The increasingly negative values with heavier elements suggest a more pronounced covalent character in the bond. A noteworthy conclusion emerged by further comparing encapsulated  $\text{Ng}_2$  reactivity with their free counterparts: confinement induces a true chemical bond between Ar–Xe, while for smaller He and Ne, the large  $\text{C}_{60}$  cavity allows for distant positions, minimizing Pauli repulsion and favoring weaker van der Waals interactions. To assess the kinetic stability of  $\text{Ng}_2@C_{60}$ , we have employed *ab initio* molecular dynamics, unveiling a remarkable precessional movement of the encapsulated pair, behaving as a unified entity.<sup>90</sup> This synchronized movement provides additional support for the formation of a bond under confinement. Interestingly, the extent of precession reduces with increasing Ng atom size.

In another interesting observation, encapsulating the  $\text{Xe}_2$  dimer necessitated a significant change in the structure of the host cage,  $\text{C}_{60}$ . To accommodate the larger guest, the cage deviated from the isolated pentagonal rule (IPR),<sup>91</sup> adopting a geometry with two adjacent pentalene units.<sup>92</sup> Driven by the pursuit of  $\text{He}_2$  bonding, our investigation ventured into even smaller cages than  $\text{C}_{60}$ .<sup>93</sup> By trapping two helium atoms within the  $\text{C}_{20}\text{H}_{20}$  cage, we achieved a He–He internuclear separation of 1.265 Å, which is less than half of the distance observed in the free  $\text{He}_2$  dimer. Despite this remarkable proximity, detailed analysis revealed very low charge transfer, a practically zero WBI, and molecular orbital plots indicative of a closed-shell interaction. This study underscores that a short internuclear distance alone does not guarantee a chemical bond; compelling

evidence of significant electron sharing and orbital hybridization is necessary.<sup>94</sup> Undeterred, our pursuit of a chemically bound He<sub>2</sub> unit continued within B<sub>12</sub>N<sub>12</sub> and B<sub>16</sub>N<sub>16</sub> cages.<sup>95</sup> Although these heteroatomic cages exhibited slightly improved charge transfer from He to the cage compared to C<sub>20</sub>H<sub>20</sub>, they remained thermochemically unstable. Nonetheless, *ab initio* simulations suggested their kinetic stability. More excitingly, He<sub>2</sub>@B<sub>12</sub>N<sub>12</sub> exhibited negative  $H(r_c)$  values at the He–He BCP, hinting at some degree of orbital involvement. Furthermore, EDA calculations revealed that 40.9% of the total attraction between the He atoms constitutes the  $\Delta E_{orb}$  term, supporting the observed orbital contribution.

The adaptability of Ng interactions extends beyond confined fullerenes, encompassing a diverse array of host molecules. Two systems, namely, clathrate hydrates and cucurbit[*n*]uril (CB[*n*]) frameworks, both traditional and HF-doped, demonstrate their potential as hosts for Ng atoms.<sup>96–99</sup> A comprehensive study involving 5<sup>12</sup>, HF5<sup>12</sup> (doped), 5<sup>12</sup>6<sup>8</sup>, and HF5<sup>12</sup>6<sup>8</sup> cages as hosts and He, Ne, and Ar as guests unveiled the stabilizing effect of HF doping. While 5<sup>12</sup> and HF5<sup>12</sup> cages can accommodate one Ng atom each, the larger HF5<sup>12</sup>6<sup>8</sup> can house up to 10 He, 6 Ne, or 6 Ar atoms. Crucially, all interactions within these complexes manifest as purely non-covalent. The CB[*n*] macrocycle, notably, CB[6] accommodates up to three Ne atoms, while only two Ar or Kr atoms fit within its confines. This varying capacity arises from the larger sizes of Ar and Kr, causing distortion of the cage walls due to repulsive forces. Interestingly, the distances between the encapsulated Ng atoms fall below the sum of their van der Waals radii, indicating close proximity. However, electron density analysis and EDA calculations unveil a closed-shell, dispersion-dominated interaction between the Ng atoms within the CB[6] cage, revealing the non-covalent nature of the association. MD simulations further affirm the stability of these confined species at 77 K. Similar investigations have been conducted with octa acids,<sup>100</sup> highlighting the diversity of potential host molecules.

The domain of Ng interactions transcends mere confinement, with the electronic charge distribution of the host emerging as a pivotal factor in stabilizing these enigmatic elements. Our scrutiny of BN-doped carbon nanotubes (BNCNTs) serves as an exemplification of this phenomenon.<sup>101</sup> Helium dimers confined within BNCNTs exhibit an unexpectedly compact He–He bond distance of 1.824 Å, notably smaller than that observed in pristine CNTs (2.596 Å). This substantial disparity underscores the heightened polarization of He atoms within the BNCNT cavity compared to their behavior in CNTs. Analogous trends manifest in other Ng–Ng distances within Ng<sub>*n*</sub>@BNCNT systems (Ng = He, Ar, Kr; *n* = 3 for He, and 2 for Ar and Kr), with AIM analysis providing additional clarity on the nature of these interactions. Helium pairs display closed-shell interactions, while Ar–Ar and Kr–Kr interactions within the BNCNT cavity assume a partial covalent character.

Borospherene (B<sub>40</sub>) emerges as another interesting host for encapsulating Ng atoms. Utilizing DFT methods, we explored systems like Ng<sub>*n*</sub>@B<sub>40</sub> (*n* = 1 and 2 for He–Kr, and only 1 for Xe and Rn).<sup>102,103</sup> While lacking thermodynamic stability

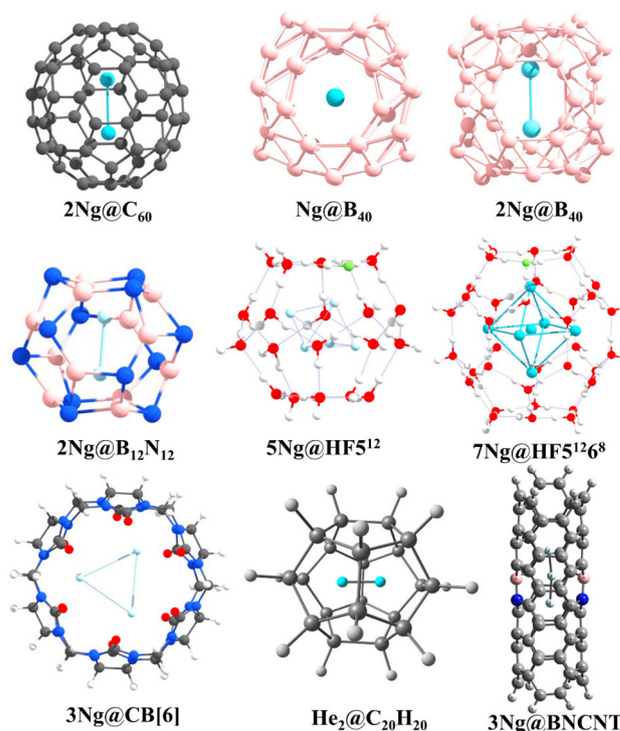


Fig. 2 A few examples of the Ng-encapsulated systems [reproduced from ref. 88 with permission from MDPI, © 2019].

concerning dissociation, these complexes exhibit remarkable kinetic stability due to elevated free energy barriers ranging from 84.7 to 206.3 kcal mol<sup>−1</sup> for the Ng@B<sub>40</sub> complexes. The increasing size of the Ng atom induces greater distortion in the encapsulated B<sub>40</sub> system, optimizing Ng<sub>2</sub> dimer formation up to the Kr analogue within the B<sub>40</sub> cavity. Notably, the Ng–Ng bonds in Ar<sub>2</sub>@B<sub>40</sub> and Kr<sub>2</sub>@B<sub>40</sub> acquire partial covalent character inside the cage, emphasizing the subtle influence of the host. The inherently fluxional nature of the B<sub>40</sub> cage, involving continuous interconversion between the hexagonal and heptagonal boron rings, has some effect on the Ng encapsulation.<sup>104–109</sup> The corresponding free energy barrier increases upon encapsulation, delineating the altered kinetic landscape. Moreover, the presence of Xe within the B<sub>40</sub> cage augments its complexation ability with [Fe(η<sup>5</sup>-C<sub>5</sub>Me<sub>5</sub>)]<sup>+</sup> compared to the free cage. This insightful observation underscores the potential for leveraging encapsulated Ng atoms to tailor the reactivity and functional properties of host molecules (Fig. 2).

## Summary

This Perspective discusses our research contributions concerning the prediction of novel noble gas (Ng) complexes and the associated bonding phenomena. Our investigations have spanned mainly three distinct categories of Ng compounds, namely, the non-insertion (NgAB), insertion (ANgB), and the cage complexes encapsulating Ng. Here, A and B can denote a single atom or a group of atoms. The thermochemical stability

of NgAB hinges on the strength of the interaction, allowing stability at specific temperatures. Conversely, ANgB compounds have an exergonic dissociation channel,  $\text{ANgB} \rightarrow \text{Ng} + \text{AB}$ , but are kinetically stable. ANgB molecules exhibit sensitivity to the level of theory employed. Instances have arisen where a system considered energetically stable at the DFT level may undergo dissociation while optimization at the coupled cluster [CCSD(T)] level. Consequently, we strongly recommend the latter for studying ANgB molecules. While bond dissociation energy may vary with changes in the level of theory for the other types of Ng compounds, stability remains relatively consistent. Analyzing the bonding scenario entails utilizing a comprehensive theoretical toolkit, which includes NBO, AIM, EDA, and AdNDP analyses. In NgAB systems, the extent of interaction between the donor and the acceptor is contingent on the polarizing ability of the A center, potentially leading to covalent bonds, especially for heavier Ng elements. Within ANgB complexes, Ng often shares electrons to form covalent bonds with A while engaging in electrostatic interactions with B, portraying itself as  $[\text{ANg}]^+\text{B}^-$ . Uncommon instances, exemplified by  $\text{NCNgNSi}$ , exhibit bonds on either side of Ng as electron-shared covalent bonds. Cage hosts serve as excellent entities to explore the limits of achieving Ng–Ng bonding, even for helium, under the condition of high pressure. The constraining influence exerted by the small  $\text{B}_{12}\text{N}_{12}$  cage can induce some degree of covalency between two He atoms in  $\text{He}_2@ \text{B}_{12}\text{N}_{12}$ . Our investigations into BN-doped nanotubes and fluxional borospherene contribute compelling chapters to the ongoing narrative of noble gas interactions. These studies not only unveil the nuanced interplay between host electronics and Ng atom stabilization, but also illuminate how confinement induces intriguing alterations in behaviour and reactivity. As we delve deeper into this captivating realm, exciting possibilities for manipulating and harnessing the unique properties of noble gas systems beckon on the horizon.

## Conflicts of interest

The authors declare that they have no conflict of interest regarding the publication of this article, financial, and/or otherwise.

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