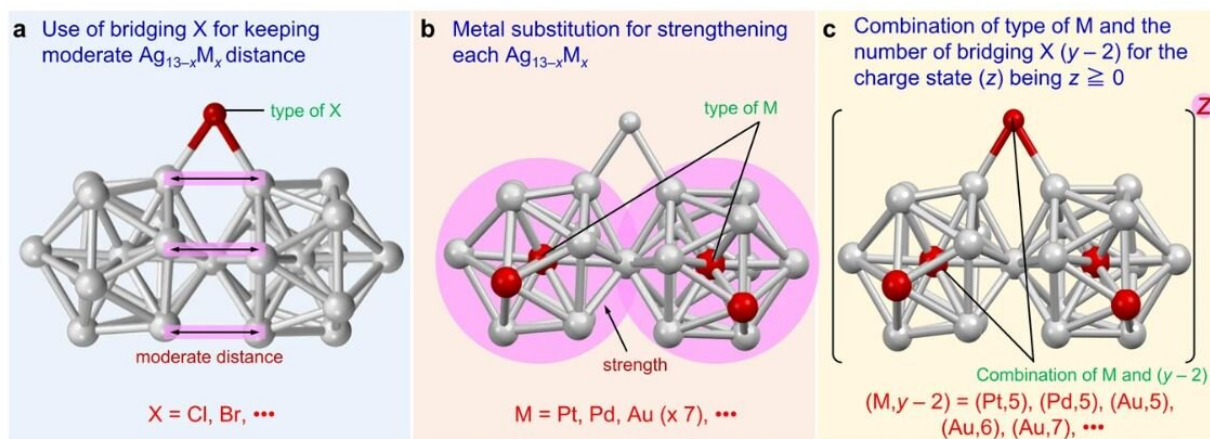


New study reveals design clues for silver-based superatomic molecules

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The formation of stable Ag-based nanoclusters require: (a) a bridging halogen that can maintain a moderate distance between the two $\text{Ag}_{13-x}\text{M}_x$ structures, (b) a metal atom for a stable icosahedral core and (c) a combination of heteroatoms and bridging halogens that results in 16 valence electrons. Credit: Yuichi Negishi, Tokyo University of Science

Superatomic molecules containing noble metal elements like gold and silver are studied for their potential in the synthesis of superatomic materials. However, the understanding of silver-based superatomic molecules has been limited. Addressing this gap, researchers from Japan studied two bimetallic superatomic molecules with silver as a main constituent to determine the key factors that enabled their formation. Their findings are expected to advance the development of novel

materials in the future.

In the past few decades, metal nanoclusters composed of noble metal elements such as [gold](#) (Au) and silver (Ag) have gained attention as superatoms for the synthesis of materials with [unique properties](#) and potential new applications. These superatoms (also known as "[artificial atoms](#)") typically consist of a cluster of a few to several hundred atoms and exhibit properties that are significantly different from their bulk, conventional counterparts. However, much like real atoms, the stability of these superatoms is determined by the formation of a closed-shell electron structure.

Ag-based superatoms are known for their superior properties and functions, including photoluminescence and selective catalytic activity, compared to those of Au-based superatoms. However, most of the research in this field has been primarily focused on Au-based superatomic molecules.

To overcome this research gap, researchers from Japan studied the formation of superatomic molecules composed of Ag and evaluated the factors involved in this formation. This study was published in the journal *Communications Chemistry* on March 28, 2023.

Speaking of the motivation behind studying Ag-based superatoms, Prof. Negishi says, "So far, we humans have created a variety of useful materials from the elements available to us on Earth. However, looking at a future with complex energy and environmental issues, the development of materials with new properties and functions is desired."

To this end, the researchers synthesized two di-superatomic molecules with bromine (Br) as the bridging ligand: $[\text{Ag}_{23}\text{Pt}_2(\text{PPh}_3)_{10}\text{Br}_7]^0$ and $[\text{Ag}_{23}\text{Pd}_2(\text{PPh}_3)_{10}\text{Br}_7]^0$ (PPh_3 = Triphenylphosphine). The former consisted of two icosahedral Ag_{12}Pt superatoms connected by vertex

sharing with platinum atoms (Pt) occupying the central position in each [superatom](#). In contrast, the other superatomic molecule consisted of two icosahedral Ag_{12}Pd structures with palladium (Pd) as the central atom.

The geometric/electronic structure and stability of these two nanoclusters was then analyzed and compared with $[\text{Ag}_{23}\text{Pt}_2(\text{PPh}_3)_{10}\text{Cl}_7]^0$ (1) and $[\text{Ag}_{23}\text{Pd}_2(\text{PPh}_3)_{10}\text{Cl}_7]^0$ (2)—two nanoclusters with geometrical similarity to the synthesized nanoclusters, consisting of chlorine (Cl) as the bridging atom.

On examining the geometric structures of the four nanoclusters, the researchers observed a twist between the two icosahedral structures containing Br as the bridging ligand. The researchers suggest that this twist stabilizes the nanocluster by shortening the distance between the two icosahedral structures.

Additionally, the larger Br atom was found to introduce steric hindrance in the molecule, causing both the PPh_3 molecule to be positioned further from the long axis of the metal nanocluster and, a change in the bond length of the Ag-P and Ag-Ag bonds. These findings indicate that although the type of bridging halogen slightly affects the geometric structures of the metal nanoclusters, it does not hinder their formation.

"The type of bridging halogen appears to have little effect on whether superatomic molecules can be formed or not, as long as the bridging halogen is large enough to maintain a moderate distance between the two Ag_{12}M structures," explains Prof. Negishi.

However, the stability of the nanocluster was largely dependent on the number of bridging halogens attached to it. Like atoms, stable metallic nanoclusters require a filled valence shell. In the case of the prepared nanoclusters—which had a total of 16 valence electrons—the researchers were only able to attach a maximum of five bridging halogens to

maintain the metal nanocluster in a stable neutral or cationic state.

The presence of Pd and Pt central atoms was found to be due to the formation of metallic nanoclusters. Substituting the central atom of Ag_{13} with Pt or Pd led to an increase in the average binding energy within the nanoclusters, making it favorable for the formation of superatomic molecules.

Overall, the researchers identified three key requirements for the formation and isolation of superatomic molecules consisting of two $\text{Ag}_{13-x}\text{M}_x$ structures connected by vertex sharing. These include the presence of a bridging halogen that can maintain an optimal distance between the two structures, a combination of heteroatoms and bridging halogens that results in 16 valence electrons, and the formation of an icosahedral core that is stronger than Ag_{13} .

In the words of Prof. Negishi, "These findings offer clear design guidelines for the creation of molecular devices with various properties and functions, and can potentially contribute to resolving pressing concerns regarding clean energy and the environment."

More information: Sayuri Miyajima et al, Key factors for connecting silver-based icosahedral superatoms by vertex sharing, *Communications Chemistry* (2023). [DOI: 10.1038/s42004-023-00854-0](https://doi.org/10.1038/s42004-023-00854-0)

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