**Abstract:** Metal nanoclusters (NCs), which consist of several, to about one hundred, metal atoms, have attracted much attention as functional nanomaterials for use in nanotechnology. Because of their fine particle size, metal NCs exhibit physical/chemical properties and functions different from those of the corresponding bulk metal. In recent years, many techniques to precisely synthesize metal NCs have been developed. However, to apply these metal NCs in devices and as next-generation materials, it is necessary to assemble metal NCs to a size that is easy to handle. Recently, multiple techniques have been developed to form one-, two-, and three-dimensional connected structures (CSs) of metal NCs through self-assembly. Further progress of these techniques will promote the development of nanomaterials that take advantage of the characteristics of metal NCs. This review summarizes previous research on the CSs of metal NCs. We hope that this review will allow readers to obtain a general understanding of the formation and functions of CSs and that the obtained knowledge will help to establish clear design guidelines for fabricating new CSs with desired functions in the future.

**Keywords:** metal cluster; one-dimensional connected structure; two-dimensional connected structure; three-dimensional connected structure; metal–organic framework; photoluminescence; electrical conductivity

1. **Introduction**

1.1. Metal Nanoclusters for Nanotechnology

Nanotechnology is technology to precisely manufacture small structures. In many countries, nanotechnology is being promoted as a national policy. Progress of nanotechnology allows information and functions to be integrated in smaller spaces, making it possible to manufacture devices with more functions at the same scale. In addition, since nanotechnology makes it possible to integrate the same function in a smaller volume than is the case for current devices, it is expected that devices will be downsized, which will increase their portability. This eliminates the need for the user to be dependent on the device location, which could solve the problems such as crowding and traffic jams and allow users to manage their time more effectively. In addition, the progress of nanotechnology has many advantages, such as saving resources and energy and decreasing waste and environmental damage [1].
Techniques to fabricate small materials can be roughly divided into two categories. One category is top-down methods, in which a desired structure is produced by decreasing the size of a larger substrate (Figure 1). Nanotechnology has been supported by the development of top-down methods. For example, increased functionality and miniaturization of electronic devices have been realized by the progress of top-down techniques. However, when a fine structure is manufactured by using tools (light, electron beam, scanning probe microscope, etc.), it is difficult to manufacture nanostructures with finer accuracy than that of the tools. Therefore, in recent years, bottom-up methods that assemble nanostructures from atoms and molecules have attracted attention (Figure 1) [1].

Figure 1. Fine processing techniques, including top-down (cyan) and bottom-up (red) methods. IC = integrated circuit.

Metal nanoclusters (NCs), which consist of several, up to about one hundred, metal atoms [1–10], are nanomaterials that can be synthesized by bottom-up methods. Metal NCs are not only small (<2 nm in size), but they also show physical/chemical properties and functions which are different from those of the corresponding bulk metals [11–35]. Furthermore, the physical/chemical properties and functions of metal NCs change considerably depending on the number of constituent atoms [36–65]. Therefore, if the number of constituent atoms of metal NCs is controlled, it is possible to produce various physical/chemical properties and functions by using only one type of metal element. If several types of elements can be used, it becomes possible to obtain more functionalities [66–82]. For these reasons, metal NCs have been attracting considerable attention as a central material in nanotechnology. In recent years, it has become possible to synthesize such metal NCs precisely at the atomic and molecular level by using thiolate (SR) [2,66], alkyne [59,83], phosphine [9,84–92], carbon monoxide [93–100], and dendrimers [7] as protective organic molecules. Investigation of the obtained precise metal NCs has revealed their geometrical structure (aggregation pattern of metal atoms) and the influences of miniaturization [1–65] and alloying [66–82] on the electronic structures and physical/chemical properties of metal NCs. In parallel, research on the applications of the optical properties and catalytic activity of metal NCs is being actively conducted [81,101–108].

1.2. Controlled Assembly of Metal Nanoclusters

As mentioned above, metal NCs show promise as constituent units of functional nanomaterials. Gold (Au) NCs have already been put to practical use in the fields of sensors, catalysts, and paints. On the other hand, at present, electronic devices are manufactured by top-down methods. Therefore, to replace the components of current devices with metal NCs, it is necessary to grow metal NCs to a size that allows their combination with structures manufactured by top-down methods. Moreover, in other applications, the small size of metal NCs often makes them difficult to handle. To realize nanodevices and next-generation materials with the advantageous characteristics of metal NCs, it is essential to establish techniques to assemble metal NCs to a size that makes them easy to handle.
To form a one-dimensional (1D) arrangement of metal NCs, templates [109,110] and host-guest interactions [111] are extremely effective. Two-dimensional (2D) and three-dimensional (3D) arrays of metal NCs can be fabricated by Langmuir–Blodgett [112] and alternate adsorption methods. There are many reports in which metal NCs are arranged in one, two, and three dimensions by using these methods. However, in the structures produced by these methods, the metal NCs are not regularly arranged in a strict sense. Since the conductivity (\( \sigma \)) of metal NCs changes exponentially with the length of the insulating organic ligands [113], the existence of a distribution in the distances between NCs (i.e., the length of the insulating part) is undesirable when applying the assembled metal NCs in electronic devices. Furthermore, although the Langmuir–Blodgett and alternate adsorption methods are suitable for arranging metal NCs in a wide size area, they are not suitable for arranging those in a tiny area.

Metal NCs are regularly arranged in single crystals. Therefore, it is possible to produce precise structures in which metal NCs are regularly connected in one, two, and three dimensions by crystallizing them while including an ingenious means to connect metal NCs. In fact, many crystals in which metal NCs are regularly linked by such a method have been reported in recent years. These structures contain strong bonds, such as Au–Au, Au–silver (Ag), Ag–oxygen (O), Ag–sulfur (S), Ag–chloride (Cl), Ag–nitrogen (N), cesium (Cs)–S, or hydrogen (H) bonds [114], and weak interactions, such as \( \pi \)–\( \pi \), anion–\( \pi \), cation···\( \pi \), aryl CH···Cl, and van der Waals interactions [115] (Figure 2). Similar to supramolecules, molecular assemblies [116], and metal–organic frameworks (MOFs) [117] that are self-assembled from metal ions and organic molecules, connected structures (CSs) of metal NCs can be formed by self-assembly during crystallization, using these bonds and interactions. Such structures, which could also be called “suprametal NC crystals”, exhibit physical/chemical properties different from those of an individual metal NC. Thus, the formation of CSs not only increases the size of the structure, but also enables the application of NCs in new fields.

![Figure 2](image-url). Representative methods for connecting metal NCs: (A) formation of metal–metal bond; (B) formation of Ag–O, Ag–S, and Ag–Cl bonds; (C) control of counterions; (D) introduction of linker molecules; and (E) use of inter-ligand interactions. In this review, 1D, 2D, and 3D CSs formed by inter-ligand interactions (E) are not introduced.

1.3. Contents of This Review

In recent years, it has become possible to control not only the geometrical structure of metal NCs, but also 1D, 2D, and 3D CSs of metal NCs. Further development of these techniques will lead to novel nanomaterials possessing the characteristics of metal NCs. Such development may enable a future in
which metal NCs are applied in devices. However, since these studies have been initiated in recent years, there have been few review articles focusing on 1D, 2D, and 3D CSs of metal NCs [118,119].

In this review, we summarize the existing research, with the purposes of understanding the current situation regarding these structures and giving perspective regarding clear design for producing new 1D, 2D, and 3D CSs with desired functions.

This review is structured as follows. Section 2 outlines the fabrication of 1D CSs consisting of metal NCs, their geometrical structures, and physical/chemical properties. Then, Sections 3 and 4 present research on 2D and 3D CSs, respectively. After summarizing this review article in Section 5, a brief future outlook is described in Section 6.

It should be noted that 1D, 2D, and 3D CSs of metal NCs can be formed by methods other than crystallization [120,121]; for example, hydrophilic Au NCs have been arranged in 1D and 3D form by the Xie’s group, although these NCs have not been crystallized. However, in this review, only the CSs of metal NCs in crystals are summarized, because our focus is on the regularly CSs in a strict sense. In addition, we described the synthesis methods only for the several examples. Thus, we recommend the readers who want to know the detail of synthesis methods for each example to refer to each original paper.

2. One-Dimensional Structures

The formation of 1D CSs composed of precise metal NCs is important from the viewpoint of the fabrication of controlled nanodevices by bottom-up methods. In this section, we introduce some typical examples of the construction of 1D CSs by the formation of metal–metal bonds (Figure 2A), formation of Ag–O bonds (Figure 2B), control of counterions (Figure 2C), and introduction of linker molecules (Figure 2D). The connection methods, NCs, linkers, year reported, and reference numbers of 1D CSs are summarized in Table 1. Chemical structures of some of the ligands used in these studies are shown in Scheme 1. The chemical structures of organic molecules used as linkers are illustrated in Scheme 2.
Scheme 2. Linker molecules used to connect metal NCs: (1) pyrazine, (2) pyridine, (3) p-methylpyridine, (4) bpy-NH₂, (5) bpy, (6) p-iah, (7) o-iah, (8) bpe, (9) dipyr idin-4-yl-diazene, (10) TPPA, (11) TPyP, (12) m-iah, (13) bpz-NH₂, (14) 1,4-bis(4-pyridyl)benzene, (15) CPPP, (16) tppe, and (17) mdppz.

2.1. Direct Connection via Metal–Metal Bonds

In 2014, Maran et al. [122] fabricated a 1D CS composed of an SR-protected Au25-atom NC ([Au25(SR)18]0). Since Au25(SR)18 NCs exhibit high stability among Au(n)(SR)m NCs, their geometrical/electronic structures and physical/chemical properties have been studied extensively [18,27,123–139]. However, because most of the studies were conducted on Au25(SR)18 in solution and there had been few studies on Au25(SR)18 in the solid phase, Maran’s group studied the behavior of Au25(SR)18 in the solid state.

In their study, butanethiolate (S-Bu, Scheme 1(1)) was used as the SR ligand. First, [Au25(S-Bu)18]− anion was synthesized by reducing the Au(I)-S-Bu complex by using sodium borohydride (NaBH4). Then, [Au25(S-Bu)18]− anion was oxidized into neutral [Au25(S-Bu)18]0 in open column packed with silica gel, and single crystals were grown by slow evaporation. Figure 3A(a) shows the geometrical structure of [Au25(S-Bu)18]0 obtained by single-crystal X-ray diffraction (SC-XRD). Each [Au25(S-Bu)18]0 has almost the same framework structure as that of [Au25(SR)18]0 protected by other SR ligands; e.g., phenylethanethiolate (PET, Scheme 1(2)) and ethanethiolate (S-Et, Scheme 1(3)) (Figure 3A(b)) [27,53]. However, Au–Au bonds formed between adjacent NCs in the [Au25(S-Bu)18]0 crystal, unlike the case for other [Au25(SR)18]0 crystals (Figure 3A(a),B(a)). This indicates that [Au25(S-Bu)18]0 is a suitable structural unit to form 1D CSs. The Au–Au distance between adjacent NCs of 3.15 Å was within the range of aurophilic interactions (2.9–3.5 Å) and shorter than the non-bonding Au–Au distance (3.80 Å) estimated from the van der Waals radius of Au. This result indicates that a 1D CS was formed in the crystal structure of [Au25(S-Bu)18]0 via Au–Au bonds. To form such a 1D CS, it was considered that
the repulsion between the ligands was suppressed and an attractive force between the ligands was induced because the adjacent NCs twisted and approached each other (twist-and-lock mechanism). It was suggested that 1D CSs did not form when S-Et and PET were used (Figure 3A(b),B(b)) because S-Et has a short alkyl group that leads to a weak attractive force between ligands and PET with a bulky functional group has large steric repulsion between ligands. In 2017, these researchers also succeeded in forming a 1D CS of [Au$_{25}$ (S-Pen)$_{18}$]$_0$ (S-Pen = pentanethiolate, Scheme 1(4)) [140]. In the same paper, they reported that the distance between NCs was shorter in this 1D CS than in the 1D CS of [Au$_{25}$ (S-Bu)$_{18}$]$_0$ (Figure 3C). In addition, in 2019, they formed 1D CSs of [Au$_{24}$Hg(S-Bu)$_{18}$]$_0$ (Hg = mercury) and [Au$_{24}$Cd(S-Bu)$_{18}$]$_0$ (Cd = cadmium), in which one Au of [Au$_{25}$ (S-Bu)$_{18}$]$_0$ was replaced with Hg or Cd [141].

Figure 3. (A,B) Crystal structures of (a) [Au$_{25}$ (S-Bu)$_{18}$]$_0$ and (b) [Au$_{25}$ (S-Et)$_{18}$]$_0$. In (B), R groups are omitted for clarity. (C) Crystal structure of [Au$_{25}$ (S-Pen)$_{18}$]$_0$. In (A–C), Au = yellow, S = red, C = light blue, and H = white. (D) Comparison of the continuous wave-electron paramagnetic resonance (EPR) spectra of solid (blue traces) and frozen toluene solution (red traces) for (a) [Au$_{25}$ (S-Bu)$_{18}$]$_0$ and (b) [Au$_{25}$ (S-Et)$_{18}$]$_0$ at −253 °C. The inset shows the same spectra with normalized peak intensity. The black curve corresponds to the EPR cavity signal, which is subtracted in the inset for clarity. All spectra were obtained by using the following parameters: microwave frequency = 9.733 GHz; microwave power = 150 µW; amplitude modulation = 1 G. Reproduced with permission from References [122,140]. Copyright 2014 American Chemical Society and 2017 American Chemical Society.
The same group also revealed that the 1D CSs had electronic structures and physical properties different from those of individual NCs. Since [Au25(S-Bu)18]0 has unpaired electrons, it exhibits paramagnetism in solution. Conversely, the 1D CS of [Au25(S-Bu)18]0 was non-magnetic (Figure 3D) [122]. This change was mainly ascribed to the formation of the 1D CS, which led to the close proximity of NCs, allowing the unpaired electrons of adjacent NCs to form electron pairs. Because of the formation of such electron pairs, the conduction band of the 1D CS was full and its valence band was empty, so the obtained 1D CS was predicted to have the properties of a semiconductor [122].

In 2020, we [142] conducted a detailed study on the factors responsible for the formation of 1D CSs via Au–Au bonds by using [Au4Pt2(SR)8]0 (Pt = platinum) as the NC. A similar NC, [Au4Pd2(PET)8]0 (Pd = palladium), was reported by Wu and colleagues in 2017 (Figure 4) [143]. Although it was not mentioned in their paper, [Au4Pd2(PET)8]0 formed a 1D CS in its crystal (Figure 4). Because this type of metal NC has a smaller metal core than that of [Au25(SR)18]0 described above (Figure 4), the distribution of the ligands in this type of NC should change depending on the ligand structure. Moreover, Au and Pt form a stronger bond than that between Au and Pd [144]. Therefore, it was expected that changing Pd to Pt would increase the stability of the NC [145], thereby expanding the variety of ligand functional group structures that can be used in 1D CSs. For these reasons, we chose [Au4Pt2(SR)8]0 as the building block of their 1D CS. The SR ligands shown in Scheme 1 (2), (5)–(8) were used. Because the functional group structures of these SR ligands differ greatly, it was expected that there would be different ligand–ligand interactions between the resulting NCs.

In the experiment, [Au4Pt2(SR)8]0 NCs with different SRs were precisely synthesized by reducing the metal–SR complex with NaBH4. Each [Au4Pt2(SR)8]0 NC was separated from by-products, using open column chromatography, and then single crystals were grown by vapor diffusion. The SC-XRD of the series of [Au4Pt2(SR)8]0 crystals revealed the following three points for [Au4Pt2(SR)8]0: (1) [Au4Pt2(SR)8]0 is a metal NC that can become a structural unit of 1D CSs via Au–Au bond formation (Figure 5A); (2) although all [Au4Pt2(SR)8]0 NCs have similar structures, the intra-cluster ligand interactions vary depending on the ligand structure. As a result, the distribution of the ligands in [Au4Pt2(SR)8]0 changes depending on the ligand structure; (3) the differences in the ligand distributions influence the inter-cluster ligand interactions, which in turn affect the formation of 1D CSs and change their structure (Figure 5B). These results demonstrate that we need to design intra-cluster ligand interactions, to produce 1D CSs with desired configurations. This study also explored the effects of 1D CS formation on the electronic structure of NCs. The results revealed that the formation of the 1D CS caused the band gap of the NCs to decrease (Figure 5C, D) [142].

Figure 4. Crystal unit cell of [Au4Pd2(PET)8]0. S = yellow, Au = red and orange, Pd = olive, C = gray. Reproduced with permission from Reference [143]. Copyright 2017 Wiley-VCH.
in the ligand layers (see Figure 6E) and in weak interactions in the ligand layers (see Section 2.3). It was interpreted that the conductivity and charge carrier mobility was increased by several orders of magnitude in their 1D CS consisting of Au clusters in the crystallographic axis is one-to-three orders of magnitude higher than the values reported for 1D CS consisting of Au\(^{10}\) crystals in the \(\approx 5\ \text{S/m}\) of these crystals in the \(\approx 1.3\ \text{eV}\) (Figure 6C). Field-effect transistors (FETs) fabricated with single crystals of 1D CS by density functional theory (DFT) calculations, which predicted that the single crystals of 1D CS have a band gap of about 1.3 eV (Figure 6C). Field-effect transistors (FETs) fabricated with single crystals of 1D CS (Figure 6D) showed highly anisotropic \(\approx 1800\)-fold conductivity in the direction of the polymer as compared to cross directions (Figure 6E), hole mobility of \(\approx 70\ \text{cm}^2/\text{Vs}\), and an ON/OFF ratio up to \(\approx 4000\). They noted that the conductivity (1.49 \times 10^{-5} \text{S/m}) of these crystals in the c-crystallographic axis is one-to-three orders of magnitude higher than the values reported for 1D CS consisting of Au\(^{21}\) clusters, where 1D CS was formed by modulating the weak interactions in the ligand layers (see Section 2.3). It was interpreted that the conductivity and charge carrier mobility was increased by several orders of magnitude in their 1D CS via direct linking.
of the metal NCs by the –Ag–Au–Ag– chains in the crystal. They described in this paper that this result holds promise for further design of functional cluster-based materials with highly anisotropic semiconducting properties.

![Figure 6](image_url)

**Figure 6.** (A) Structures of the cluster polymer (approximately orthogonal to the c-axis). (B) Au–Au distances in the distorted Au₆ hexagon and Ag–Ag distance in the “Ag–Au–Ag” unit of between alloy NCs. Au/Ag = golden and green, C = gray. All hydrogen atoms are omitted for clarity. (C) DFT-computed electronic density of states (DOS) of the cluster polymer crystal. Cluster model was used to build the periodic crystal, and the integration over the Brillouin zone was done in a 4 × 4 × 4 Monkhorst-Pack k-point mesh. The band gap is centered around zero. (D,E) Electrical transport properties of the cluster polymer crystals; (D) structure of the polymer crystal FET; (E) I–V plot of the polymer crystal along a-axis and c-axis, respectively, with the range of corresponding conductivity values shown in the inset. Reproduced with permission from Reference [146]. Copyright 2020 Springer-Nature.

### 2.2. Connection via Ag–O Bonds

When Ag NCs contain acetic acid ions (CH₃COO⁻), trifluoroacetic acid ions (CF₃COO⁻), or nitrate ions (NO₃⁻) in the ligand layer, it is possible to connect Ag NCs by forming Ag–O bonds. Su et al. [147] first reported the formation of such a 1D CS in 2014. In this study, the 1D CS was obtained by crystallization of the product which was obtained by the reaction between AgS₄Bu, (NH₄)₃[CrMo₃O₇H₆] (Cr = chromium, Mo = molybdenum), Ni(CH₃COO)₂ (Ni = nickel), AgCF₃COO, and AgBF₄. The each Ag NC had a chemical composition of Ag₂₀(CO₃)(S₄Bu)₁₀(CH₃COO)₆(DMF)₂ (CO₃²⁻ = carbonate anion; S₄Bu = tert-butylthiolate, Scheme 1(10)), DMF = N,N-dimethylformamide). The Ag₂₀(CO₃) core of the NC was formed by the aggregation of Ag around CO₃²⁻ (Figure 7A) as an anion template. In the crystal, the Ag NCs were connected in one dimension via two Ag–O–Ag bonds (Figure 7B). The obtained 1D CS was stable in both solid and solution states, had a bandgap of 3.22 eV, and exhibited reversible thermochromic emission.
Formation of 1D CSs based on a similar principle was also reported by Mak and co-workers in 2017 [148]. In their report, Ag$_{18}$(CO$_3$)$_3$(S$_2$Bu)$_{10}$(NO$_3$)$_6$(DMF)$_4$ was linked by the formation of Ag–O bonds (Figure 8). The Ag$_{18}$(CO$_3$)$_3$ core contained CO$_3^{2–}$ as an anion template at the center, like Ag$_{20}$(CO$_3$)$_3$. As described in Section 3.1, this group also succeeded in forming a 2D CS of Ag$_{20}$(CO$_3$)$_3$ by changing the SR structure.

Recently, Sun et al. [149] succeeded in the connection of Ag$_{44}$(V$_{10}$O$_{28}$)(S-Et)$_{20}$(PhSO$_3$)$_{18}$(H$_2$O)$_2$ (V$_{10}$O$_{28}^{6–}$, Scheme 1(11), PhSO$_3^{–}$ = benzenesulfonic acid ion). In this 1D CS, the Ag$_{44}$(V$_{10}$O$_{28}$) core was formed by using the polyoxometalate (POM) V$_{10}$O$_{28}^{6–}$ as an anion template (Figure 9A). This was the first report of the formation of a structure in which V$_{10}$O$_{28}^{6–}$ was covered with an Ag NC with SR as a ligand. This 1D CS assembled because two Ag–O bonds were formed between two PhSO$_3^{–}$ in the ligand layer and one Ag$_{44}$(V$_{10}$O$_{28}$)(S-Et)$_{20}$ NC (Figure 9B).

In the 1D CS of [Au$_7$Ag$_9$(dpdf)$_3$(CF$_3$COO)$_2$BF$_4$]$_n$ (dpdf = 1,1′-bis(diphenylphosphino)ferrocene, Scheme 1(12), BF$_4$ = tetrafluoroboric acid) reported by Wang et al. [150] in 2019, each NC was also connected via an Ag–O bond (Figure 10A), although this was not direct connection of metal NCs. In the above three NCs (i.e., Ag$_{20}$(CO$_3$)$_3$(S$_2$Bu)$_{10}$(CH$_3$COO)$_8$(DMF)$_2$, Ag$_{18}$(CO$_3$)$_3$(S$_2$Bu)$_{10}$(NO$_3$)$_6$(DMF)$_4$,
and Ag₄₄(V₁₀O₂₈)(S·Et)₂₀(PhSO₂)₁₈(H₂O)₂), the anion template was contained in the center, whereas Au₇Ag₈(dpff)₃(CF₃COO)₇ had an icosahedral metal core composed of Au₇Ag₈. Such an icosahedral core structure is often seen in metal NCs [27,53]. The 1D CS of [Au₇Ag₈(dpff)₃(CF₃COO)₇]BF₄劬 was synthesized in one pot. Probably, excess Ag binds to CF₃COO⁻ in the ligand layer during synthesis, resulting in the formation of a 1D CS composed of Au₇Ag₈(dpff)₃(CF₃COO)₇ NCs. The researchers also revealed that this 1D CS possessed a band gap of 2.18 eV (Figure 10B).

Figure 10. (A) View of the whole structure of [Au₇Ag₈(dpff)₃(CF₃COO)₇]BF₄劬 (anions and H atoms are omitted for clarity). Ag = green, Au = orange, O = red, P = purple, C = gray, Fe = blue. (B) Absorption spectrum of [Au₇Ag₈(dpff)₃(CF₃COO)₇]BF₄劬 in CH₂Cl₂ solution. Inset: absorption spectrum on the energy scale (eV) and photographs showing actual colors of [Au₇Ag₈(dpff)₃(CF₃COO)₇]BF₄劬 in CH₂Cl₂ and the crystalline state. Reproduced with permission from Reference [150]. Copyright 2019 Royal Society of Chemistry.

2.3. Control of Counterions

For certain metal NCs, the total number of valence electrons satisfies the number for the closed-shell electronic structure when it is a cation, so they are generated as a cation [151]. For example, [Au₂₁(S·C₅H₁₁)₁₂(dppm)]⁺ (S·C₅H₁₁ = cyclohexanethiolate, Scheme 1(13), and DPPM = bis(diphenylphosphinomethane), Scheme 1(14)) is synthesized as a cation. In 2018, Jin et al. [152] revealed that [Au₂₁(S·C₅H₁₁)₁₂(dpdm)]⁺ formed a 1D CS in a crystal by assembling as a pair with the counter anion and that the structure of the 1D CS changed depending on the counterion (Figure 11A,B).

In this study, [Au₂₁(S·C₅H₁₁)₁₂(dpdm)]⁺[AgCl₂]⁻ and [Au₂₁(S·C₅H₁₁)₁₂(dpdm)]⁺[Cl]⁻ were precisely synthesized and single crystals were grown. As shown in Figure 11B, a 1D CS was formed by the alternating connection of [Au₂₁(S·C₅H₁₁)₁₂(dpdm)]⁺ and [AgCl₂]⁻ in the [Au₂₁(S·C₅H₁₁)₁₂(dpdm)]⁺[AgCl₂]⁻ crystal. This 1D CS was considered to assemble via π-π, anion-π, and aryl-C·H···Cl interactions. The connection pattern of [Au₂₁(S·C₅H₁₁)₁₂(dpdm)]⁺ in the 1D CS changed slightly when the counterion was Cl⁻ rather than AgCl₂⁻. It was considered that the connection pattern of [Au₂₁(S·C₅H₁₁)₁₂(dpdm)]⁺ changed because the arrangement of phenyl groups in the NC was affected by the counterion (Figure 11A).

The obtained 1D CSs of [Au₂₁(S·C₅H₁₁)₁₂(dpdm)]⁺ had different electron transport properties depending on the counter anion. The 1D CS of [Au₂₁(S·C₅H₁₁)₁₂(dpdm)]⁺[AgCl₂]⁻ had a σ of only ~1.44 × 10⁻⁸ S/m, whereas that of [Au₂₁(S·C₅H₁₁)₁₂(dpdm)]⁺[Cl]⁻ was ~2.38 × 10⁻⁶ S/m (Figure 11C). Changing the counter anion from AgCl₂⁻ to Cl⁻ shortened the distance between NCs from 16.80 to 16.39 Å and formed an intra-cluster π-stacking structure that allowed electricity to flow easily (Figure 11B). These two reasons explained why σ of [Au₂₁(S·C₅H₁₁)₁₂(dpdm)]⁺[Cl]⁻ was two orders of magnitude higher than that of [Au₂₁(S·C₅H₁₁)₁₂(dpdm)]⁺[AgCl₂]⁻ (Figure 11D).

There are not only metal NCs synthesized as cations but also metal NCs synthesized as anions. Because the total number of valence electrons of [Ag₂₉(BDT)₁₂(PPh₃)₄]⁻ (BDT = 1,3-benzendithiolate, Scheme 1(15); PPh₃ = triphenylphosphine, Scheme 1(16)) satisfies the number for a closed-shell electronic structure as the anion, it is generated as the anion [153]. In 2019, Zhu et al. [154] reported that mixing this NC with Cs acetate in DMF induced Cs⁺ attachment to the NC and PPh₃ desorption from the NC, resulting in the formation of [Cs₃Ag₂₉(BDT)₁₂(DMF)₃]₀ (x = 5 or 6) and that the
obtained $[\text{Cs}_3\text{Ag}_{29}(\text{BDT})_{12}(\text{DMF})_3]^0$ formed a 1D CS in its crystal. Figure 12A shows the resulting 1D CS. In the crystal, $[\text{Cs}_3\text{Ag}_{29}(\text{BDT})_{12}(\text{DMF})_3]^0$ was connected by a series of bonds consisting of $-\text{Cs}^+ - \text{DMF} - \text{Cs}^+ - \text{S} - \text{Ag} - \text{Ag} - \text{S} -$. This 1D CS was considered to be formed because of the electrostatic attraction between $[\text{Ag}_{29}(\text{BDT})_{12}(\text{DMF})_3]^-$ and $\text{Cs}^+$, $\text{Cs} - \text{S}$ bond formation, and $\text{Cs} - \pi$ interactions (Figure 12B).

Figure 11. (A) Site-specific tailoring of the surface motifs and associated counterions of Au NCs. The two RS-Au-SR (R is cyclohexyl) surface motifs in Au$_{213}$ (precursor NC) were replaced by two DPPM motifs in Au$_{211}$. Each P atom was connected to two phenyl rings. Dashed lines indicate the motifs. (B) Packing of $[\text{Au}_{213}(\text{S-c-C}_6\text{H}_{11})_{12}(\text{DPPM})_{2}]^+[\text{AgCl}_2]^-$ and $[\text{Au}_{211}(\text{S-c-C}_6\text{H}_{11})_{12}(\text{DPPM})_{2}]^+[\text{Cl}]^-$ in their 1D assemblies. The orientation of Au NCs is modulated by the counterion. Au = magenta, Ag = gray, Cl = light green, S = yellow, P = orange, C = green. All H atoms are omitted for clarity. Yellow areas are the surface hooks connecting neighboring NCs. (C) Room-temperature conductivity of single crystals of $[\text{Au}_{213}(\text{S-c-C}_6\text{H}_{11})_{12}(\text{DPPM})_{2}]^+[\text{AgCl}_2]^-$ (green) and $[\text{Au}_{211}(\text{S-c-C}_6\text{H}_{11})_{12}(\text{DPPM})_{2}]^+[\text{Cl}]^-$ (red). (D) Schematic diagram of electron hopping in Au$_{211}$ NC assemblies. Different configurations of the interacting $\pi-\pi$ pairs result in tunneling barriers of different heights (white solid squares), thus changing the electron conductivity ($e^-$ represents an electron, $\sigma$ is the conductivity, $d$ is the interparticle distance, and $\beta$ is the tunneling decay constant). Reproduced with permission from Reference [152]. Copyright 2018 Springer-Nature.
Both \([\text{Ag}_{29}(\text{BDT})_{12}(\text{PPh}_3)_4]^-\) and \([\text{Cs}_3\text{Ag}_{29}(\text{BDT})_{12}(\text{DMF})_3]^0\) solutions showed similar absorption and photoluminescence (PL) spectra (Figure 12C). This indicates that Cs\(^+\) attachment and PPh\(_3\) desorption did not markedly change the electronic structure of the Ag\(_{29}(\text{BDT})_{12}\) NCs. In contrast, the absorption and PL spectra of \([\text{Ag}_{29}(\text{BDT})_{12}(\text{PPh}_3)_4]^-\) and \([\text{Cs}_3\text{Ag}_{29}(\text{BDT})_{12}(\text{DMF})_3]^0\) in the crystalline state were quite different (Figure 12D). The 1D CS of \([\text{Cs}_3\text{Ag}_{29}(\text{BDT})_{12}(\text{DMF})_3]^0\) was considered to show different optical behavior from that of the individual \([\text{Ag}_{29}(\text{BDT})_{12}(\text{PPh}_3)_4]^-\) because of the electronic interactions between adjacent NCs in the 1D CS of \([\text{Cs}_3\text{Ag}_{29}(\text{BDT})_{12}(\text{DMF})_3]^0\).

2.4. Introduction of Linker Molecules

In the examples described in Section 2.3, 1D CSs were formed by the counterion acting as a linker. When an organic molecule is used as the linker, the distance between NCs in a 1D CS can be freely controlled because the design of the structure of organic molecules is well understood. In fact, the geometry of an MOF is controlled by the design of the linker organic molecule [155]. In recent years, several 1D CSs of metal NCs with organic molecules as linkers have also been reported.

**Figure 12.** (A) 1D linear assembly of \([\text{Cs}_3\text{Ag}_{29}(\text{BDT})_{12}(\text{DMF})_3]^0\) in the crystal lattice. Ag = light blue/gray, Cs = dark purple, S = yellow and red, O = green. For clarity, all H, C, and N atoms, some Cs\(^+\), and DMF molecules are omitted. Each O atom represents a DMF molecule. (B) (a) Overall surface structure of \([\text{Cs}_3\text{Ag}_{29}(\text{BDT})_{12}(\text{DMF})_3]^0\) and (b) interactions between Ag\(_{29}(\text{BDT})_{12}\), Cs\(_1\), Cs\(_2\), and DMF. (C) Comparison of optical absorption and emission spectra of Ag\(_{29}(\text{BDT})_{12}(\text{PPh}_3)_4\) (black) and Cs\(_3\text{Ag}_{29}(\text{BDT})_{12}(\text{DMF})_3\) (red) NCs dissolved in DMF. (D) Comparison of optical absorption and emission spectra of Ag\(_{29}(\text{BDT})_{12}(\text{PPh}_3)_4\) (black) and Cs\(_3\text{Ag}_{29}(\text{BDT})_{12}(\text{DMF})_3\) (red) NCs in crystalline films. Reproduced with permission from Reference [154]. Copyright 2019 American Chemical Society.
slightly when the counterion was Cl\(^-\) rather than \([\text{AgCl}_2]^-\). It was considered that the connection of metal–metal bond (Figure 2A)

| Connection Mode | NC | Linker | Year | Ref. |
|-----------------|----|--------|------|------|
| Formation of Ag–Ag bond | \([\text{Au}_{25}(\text{S-Bu})_{18}]^3\) | - | 2014 | [122] |
| | \([\text{Au}_{25}(\text{S-Pen})_{18}]^3\) | - | 2017 | [140] |
| | \([\text{Au}_{22}\text{Hg}(\text{S-Bu})_{18}]^3\) | - | 2019 | [141] |
| | \([\text{Au}_{23}\text{Cd}(\text{S-Bu})_{18}]^3\) | - | 2019 | [141] |
| | \([\text{Au}_4\text{Pd}_2(\text{PET})_8]^3\) | - | 2017 | [143] |
| | \([\text{Au}_4\text{Pt}_2(\text{SCH}_2\text{PhCl})_8]^3\) | - | 2020 | [142] |
| | \([\text{Au}_4\text{Pt}_2(\text{PET})_8]^3\) | - | 2020 | [146] |

| Formation of Ag–O, Ag–S, Ag–Cl bond, etc. (Figure 2B) | \(\text{Ag}_{20}\text{CO}_2\text{(S-Bu)}_{18}\text{(CH}_3\text{COO)}_{3}\text{(DMF)}_2\) | - | 2014 | [147] |
| | \(\text{Ag}_{16}\text{CO}_2\text{(S-Bu)}_{10}\text{(NO}_2\text{)}_{3}\text{(DMF)}_4\) | - | 2017 | [148] |
| | \(\text{Au}_{4}\text{Cl}(\text{C}_6\text{H}_4\text{O}_3\text{)}_3\text{(Et)}_{12}\text{(H}_2\text{O)}_2\) | - | 2019 | [149] |
| | \(\text{Au}_{4}\text{Ag}3\text{H}(\text{dppf})_3\text{(CF}_3\text{COO)}_2\) | - | 2019 | [150] |

| Control of counter ion (Figure 2C) | \([\text{Au}_{25}(\text{S-c-C}_4\text{H}_12\text{)}_{12}\text{(DPPM)}_2]^3\) | \([\text{AgCl}_2]^3\) | 2018 | [152] |
| | \([\text{Au}_{25}(\text{S-c-C}_4\text{H}_12\text{)}_{12}\text{(DPPM)}_2]^3\) | \([\text{Cs}]^3\) | 2019 | [154] |

| Introduction of linker molecule (Figure 2D) | \(\text{Ag}_{14}(\text{DT-o-C})_8\) | pyrazine | 2018 | [156] |
| | \(\text{Ag}_{11}(\text{PhPO}_3\text{(S-Bu)}_{10}\text{(CF}_3\text{COO)}_2\text{(PhPO}_3\text{H})_4\) | bpy-NH$_2$ | 2019 | [157] |
| | \(\text{Ag}_{11}\text{Cl}(\text{S-Bu})_{10}\text{(CF}_3\text{COO)}_{3}\text{(S}\text{-Bu)}_{10}\text{(DMF)}_2\) | bpy | 2019 | [158] |
| | \(\text{Ag}_{10}\text{CF}_3\text{COO}(\text{S-Bu)}_{10}\text{(CH}_3\text{CN)}_2\) | p-iah | 2019 | [159] |
| | \(\text{Ag}_{10}\text{CF}_3\text{COO}(\text{S-Bu)}_{10}\text{(CH}_3\text{CN)}_2\) | e-iah | 2019 | [159] |
| | \(\text{Cd}_6\text{Au}_{24}(\text{S-Pt})_{10}\text{(DMF)}_2\text{(CH}_3\text{OH)}_2\) | bpe | 2020 | [160] |

\(a\) See Scheme 2.

For example, in 2018, Zang et al. [156] reported a 1D CS in which \(\text{Ag}_{14}(\text{DT-o-C})_{8}\) NCs (DT-o-C = 1,2-dithiolate-o-carborane, Scheme 1(17)) were linked by pyrazine (Scheme 2(1)). In this study, first, \(\text{Ag}_{14}(\text{DT-o-C})_{8}\) (pyridine/p-methylpyridine)$_8$ (Scheme 2(2),(3)) were identified as Ag NCs with high thermal stability that maintained their framework structure even at 150 °C or higher in air (Figure 13A). Then, they attempted to synthesize Ag NCs in which the pyridine or p-methylpyridine ligands of these Ag NCs were replaced by pyrazine. As a result, they obtained a 1D CS in which \(\text{Ag}_{14}(\text{DT-o-C})_{8}\) NCs were linked by pyrazine (Figure 13B). In the obtained structure, pyrazine was coordinated to each \(\text{Ag}_{14}(\text{DT-o-C})_{8}\) NC at a diagonal position, which caused the 1D CS to rotate in the clockwise direction with respect to the (001) axis. In this study, the researchers also succeeded in forming 2D and 3D CSs composed of \(\text{Ag}_{14}(\text{DT-o-C})_{8}\) NCs by changing the structure of the bipyridine ligand, as described later in Sections 3.2 and 4.3, respectively.

![Figure 13](image-url) (A) Structure of \(\text{Ag}_{14}(\text{DT-o-C})_{8}\) (pyridine/p-methylpyridine)$_8$. (B) 1D helix of \(\text{Ag}_{14}(\text{DT-o-C})_{8}\) NC. Ag = green and pink, S = yellow, C = gray, N = blue, carborane = turquoise. Reproduced with permission from Reference [156]. Copyright 2018 American Chemical Society.

Because N readily coordinates to Ag, bipyridine is often used to connect NCs containing Ag. In 2019, Zang and colleagues formed a 1D CS composed of \(\text{Ag}_{116}(\text{PhPO}_3\text{(S-Bu)}_{10}\text{(CF}_3\text{COO)}_2\text{(PhPO}_3\text{H})_4\) (PhPO$_3^{2-}$ = phenylphosphinic diion; PhPO$_3^{-}$ = phenylphosphinic acid ion) NC nodes with
bipyridine(3-amino-4,4′-bipyridine (bpy-NH₂, Scheme 2(4)) linkers [157]. In this experiment, NC synthesis, ligation, and crystallization were performed simultaneously in one pot (Figure 14A). The node Ag₁₈(PhPO₃)(S⁻Bu)₁₀ NCs contained PhPO₄²⁻ as an anion template in their center. In the crystal, adjacent Ag₁₈(PhPO₃)(S⁻Bu)₁₀ NCs were linked by two bpy-NH₂ to form a 1D CS (Figure 14B).

The obtained 1D CS was stable up to 110 °C in a nitrogen (N₂) atmosphere. When mechanical stimulation was applied to the 1D CS, its PL wavelength changed. When the 1D CS sample subjected to mechanical stimulation was recrystallized, its PL wavelength returned to the original value (Figure 14C). Thus, the PL of the 1D CS composed of Ag₁₈(PhPO₃)(S⁻Bu)₁₀ NCs exhibited reversible mechanochromism. Because this 1D CS emitted light at two wavelengths and its PL intensity ratio...
changed with temperature (thermochromism; Figure 14D), the authors suggested that this 1D CS could be applied as a thermometer.

In 2019, Bakr et al. [158] also reported the connection of Ag NCs by bipyridine. In this study, a 1D CS was synthesized in one pot (Figure 15A). \( \text{Ag}_{16}\text{Cl}(\text{S}^\text{-Bu})_6(\text{CF}_3\text{COO})_{5.67}(\text{NO})_{0.33}(\text{DMF})_2 \) was used as the node, and 4,4'-bipyridine (bpy, Scheme 2(5)) was used as the linker. In \( \text{Ag}_{15}\text{Cl}(\text{S}^\text{-Bu})_8(\text{CF}_3\text{COO})_{5.67}(\text{NO})_{0.33}(\text{DMF})_2 \), \( \text{Cl}^- \) acted as an anion template. The core of the \( \text{Ag}_{15}\text{Cl} \) NC had a geometry in which one Ag was lost from the \( \text{Ag}_{16}\text{Cl} \) core of the \( \text{Ag}_{16}\text{Cl}(\text{S}^\text{-Bu})_6(\text{CF}_3\text{COO})_{7}(\text{DMF})_4(\text{H}_2\text{O}) \) NC, which did not form a 1D CS, namely individual \( \text{Ag}_{16}\text{Cl}(\text{S}^\text{-Bu})_6(\text{CF}_3\text{COO})_{7}(\text{DMF})_4(\text{H}_2\text{O}) \) NCs (Figure 15B). The 1D CS with a ladder structure was formed by combining these \( \text{Ag}_{15}\text{Cl} \) NCs with three adjacent \( \text{Ag}_{15}\text{Cl} \) NCs via four bpy molecules (Figure 15C). It was found that the 1D CS displayed slightly higher thermal stability than that of the \( \text{Ag}_{16}\text{Cl} \) NCs (Figure 15D).

![Figure 15. (A) Synthesis of Ag NCs and NC-based frameworks. (B) Top views of the core structures of (a) \( \text{Ag}_{16}\text{Cl}(\text{S}^\text{-Bu})_6(\text{CF}_3\text{COO})_{7}(\text{DMF})_4(\text{H}_2\text{O}) \) and (b) \( \text{Ag}_{15}\text{Cl}(\text{S}^\text{-Bu})_8(\text{CF}_3\text{COO})_{5.67}(\text{NO})_{0.33}(\text{DMF})_2 \). (C) Crystal structure of the corresponding 1D CS. Free (co-crystallized) DMF molecules are not shown. The green semitransparent spheres in the Ag clusters are shown as a visual guide. H atoms were omitted for clarity. (D) Thermogravimetric analysis curves of NCs, 1D CS, and 2D CS (see Section 3.2). Reproduced with permission from Reference [158]. Copyright 2019 American Chemical Society.](image)

In the above three studies, the bipyridines had a rigid framework structure. In 2019, Cao et al. formed 1D CSs of Ag NCs by using pyridine derivatives \( p\text{-iah} = 4\)-pyridine carboxylic hydrazide, Scheme 2(6); \( o\text{-iah} = 2\)-carboxyhydrazide, Scheme 2(7)) that contained N in both the rigid pyridine framework and flexible substituents [159]. In this study, the 1D CSs were obtained by reacting \( \text{Ag}_{12}(\text{S}^\text{-Bu})_6(\text{CF}_3\text{COO})_6(\text{CH}_3\text{CN})_6(\text{CH}_3\text{CN} = \text{acetonitrile}) \) with the above-mentioned pyridine derivatives. The SC-XRD analysis of the products revealed that \( \text{Ag}_{10}(\text{CF}_3\text{COO})_4(\text{S}^\text{-Bu})_6(\text{CH}_3\text{CN})_2 \) and \( \text{Ag}_{10}(\text{CF}_3\text{COO})_4(\text{S}^\text{-Bu})_6(\text{CH}_3\text{CN})_2 \) were the nodes in the 1D CSs with \( p\text{-iah} \) and \( o\text{-iah} \), respectively (Figure 16A(a),B(a)). These 1D CSs containing \( p\text{-iah} \) and \( o\text{-iah} \) had cross-helical and parallel chain structures, respectively (Figure 16A(b),B(b)). The latter structure also contained hydrogen bonds \( (\text{N} - \text{H} \cdots \text{O}) \) between the parallel linker molecules. It is interesting that different 1D CSs formed depending on the position of N in the linker molecule (Scheme 2(6),(7)).
Both 1D CSs showed PL and that with o-iah as the linker exhibited weak green PL in highly polar solvents and strong yellow PL in solvents with low polarity. Based on these characteristics, the authors suggested that the 1D CS with o-iah could be used to measure the concentration of dichloromethane (CH$_2$Cl$_2$, Figure 16C) or trichloromethane (CHCl$_3$) in tetrachloromethane (CCl$_4$).

Bipyridines can also be used as linkers to form 1D CSs of Ag chalcogenide NCs. Very recently, Xu and co-workers reported the formation of a zigzag-type of 1D CS with Cd$_6$Ag$_4$(S-Ph)$_{16}$(DMF)$(\text{H}_2\text{O})$ (S-Ph = benzenethiolate, Scheme 18 and Figure 17A) as a node and trans-1,2-bis(4-pyridyl)ethylene (bpe, Scheme 28) as a linker (Figure 17B) [160]. This 1D CS was obtained by the reaction of Cd$_6$Ag$_4$(S-Ph)$_{16}$(DMF)$_3$(CH$_3$OH) (CH$_3$OH = methanol) with bpe. In the obtained 1D CS, N of bpe coordinated to Cd not Ag (Figure 17A). Such a coordination pattern has also been observed in 2D and 3D CSs composed of Cd$_6$Ag$_4$(S-Ph)$_{16}$ and bpe previously reported by Zhang et al. [161,162].
They compared the electronic structures of the resulting 1D CS and individual Cd₆Ag₄(S-Ph)₁₆(DMF)₃(CH₃OH) NCs. The results revealed that the band gap of the NCs was narrowed by the formation of the 1D CS (Figure 17C). In the 1D CS, the optical absorption onset was redshifted to the visible region. They used the 1D CS as a visible light (>420 nm)-responsive photocatalyst to decompose the organic dye Rhodamine B in water. The 1D CS exhibited higher photocatalytic activity toward Rhodamine B degradation than that of the Cd₆Ag₄(S-Ph)₁₆ NCs (Figure 17D) and high stability during the photocatalytic reaction.

3. Two-Dimensional Structures

To date, Ag NCs have been used as the building blocks in almost all 2D CSs. Like Au NCs, Ag NCs have unique electronic and optical properties [68,163–166] and are expected to be applied in various fields. However, Ag NCs are less stable than Au NCs against external stimuli, such as light and solvents. Therefore, studies have been actively conducted to improve the stability of Ag NCs by assembly of CSs and thereby improve their physical properties. In Sections 3.1 and 3.2 we focus on the connection of NCs by Ag–O bond formation (Figure 2B) and introducing linker molecules (Figure 2D), respectively. Table 2 summarizes the connection methods, NCs, linkers, reported years, and reference numbers of the relevant literature. Several of the ligands used in these studies are shown in Scheme 1. The organic molecules used as linkers are depicted in Scheme 2.

3.1. Connection via Ag–O Bonds

In 2017, Mak et al. [148] reported the formation of 2D CSs with Ag₂₀(CO₃)(S-iPr)₁₀(CF₃COO)₉(CF₃COOH)(CH₃OH)₂ (S-iPr = isopropylthiolate, Scheme 1(5)) or Ag₂₀(CO₃)(S-c-C₆H₁₁)₁₀(CF₃COO)₁₀(CF₃COOH)₂(H₂O)₂ as building blocks in their paper on the formation of 1D...
CSs. These NCs contained CO$_3^{2-}$ as an anion template at the center of their cores (Figure 18A,B). The structures of the SR in these two types of Ag$_{20}$(CO$_3$)(SR)$_{10}$ NCs were different (S-iPr vs. S-$c$-C$_8$H$_{11}$), which influenced the formation angle and bond distance between adjacent NCs in the 2D CSs. The O$^-$ of CF$_3$COO$^-$ and Ag of an adjacent NC were directly linked via an Ag$–$O bond with a length of 14 Å. In addition, isolated Ag was trapped between CF$_3$COO$^-$ of adjacent NCs, so the adjacent NCs were connected via an O$–$Ag–O bond with a length of 17 Å. The 2D CS consisting of Ag$_{20}$(CO$_3$)(S-iPr)$_{10}$(CF$_3$COO)$_6$(CF$_3$COOH)(CH$_3$OH)$_2$ showed dual emission at room temperature. Because both zero-dimensional [Ag$_{20}$(S-$i$-Bu)$_{10}$(CF$_3$COO)$_2$]Cl·(CF$_3$COO)$_7$·5CH$_3$OH NCs and the 1D CS of Ag$_{20}$(CO$_3$)(S-$i$-Bu)$_{10}$(CH$_3$COO)$_8$(DMF)$_2$ emit only single emission peaks, it was speculated that the formation of the 2D CS was related to the observed dual emission.

![Figure 18](image-url)

Figure 18. (A) The molecular building block in Ag$_{20}$(CO$_3$)(S-iPr)$_{10}$(CF$_3$COO)$_6$(CF$_3$COOH)(CH$_3$OH)$_2$ with its four linking sites, and condensation of blocks into a 4,4-net. (B) Similar condensation of molecular building blocks of Ag$_{20}$(CO$_3$)(S-$c$-C$_8$H$_{11}$)$_{10}$(CF$_3$COO)$_6$(CF$_3$COOH)$_2$(H$_2$O)$_2$. Note the difference between the ‘arms’ with lengths of 14 Å (orange) and 17 Å (cyan). Ag = blue and cyan, S = yellow, O = red, C = gray, F = green. Reproduced with permission from Reference [148].

In 2019, Sun and co-workers also reported the formation of a 2D CS, using Ag NCs [149]. In this study, a 2D CS consisting of Ag$_{46}$(V$_{10}$O$_{28}$)(S-Et)$_{25}$(PhSO$_3$)$_{15}$(CO$_3$)$_2$ was formed (Figure 19), using a different solvent from the case of Ag$_{44}$(V$_{10}$O$_{28}$)(S-Et)$_{20}$(PhSO$_3$)$_{18}$(H$_2$O)$_2$, which formed a 1D CS (Figure 9). These Ag NCs have the same total number of ligands (S-Et and PhSO$_3^-$) of 38 but different ratios of the ligand types. They considered that the ligand ratio affected the number of Ag atoms in the core and also the connection mode between adjacent NCs.
Xu et al. [166], also in 2019, reported 2D CSs of Ag$_{11}$Cl(N-L)$_8$(CF$_3$COO)$_2$·2CHCl$_3$ (Figure 20), Ag$_{11}$Cl(N-L)$_8$(NO$_3$)$_2$·2CHCl$_3$, and Ag$_{11}$Cl(N-L)$_8$(CF$_3$SO$_3$)$_2$·2CHCl$_3$ (N-L = 2-acetamido-5-methyl-1,3,4-thiadiazole, Scheme 1(19)), in which adjacent NCs are linked by Ag–O bonds. SR, alkyne, or phosphine ligands are generally used in metal NCs. In this study, their aim was to synthesize Ag NCs by using an N-donor ligand, which is not appropriate based on the hard/soft acid/base theory [167], and form corresponding CSs. The three kinds of 2D CSs obtained had similar frameworks regardless of the coordination ions (CF$_3$COO$^-$, NO$_3^-$, or CF$_3$SO$_3^-$), which means that the framework structure shown in Figure 20B is very rigid. It was found that Ag$_{11}$Cl(N-L)$_8$(CF$_3$COO)$_2$·2CHCl$_3$ showed dual-emission behavior and that its PL peaks had different optimal excitation wavelengths.

3.2. Introduction of Linker Molecules

As described below, in Section 4.3, in 2017, Zang et al. [168] reported the formation of a 3D CS in which Ag$_{12}$(S-4-Bu)$_8$(CF$_3$COO)$_4$ NCs were linked by bpy. This was a pioneering study on the formation of an MOF, using Ag NCs as nodes, and has greatly influenced subsequent studies. In 2018,
they synthesized a 2D CS consisting of Ag₁₂ NCs and bpy [169]. The core structure of the node was changed (isomerized) by dissolving an Ag₁₂(S⁻Bu)₆(CF₃COO)₆ NC MOF crystal in a mixed solvent consisting of N,N-dimethylethanamide (DMAC) and toluene, which changed the geometrical structure of the entire CS from 3D to 2D (Figure 21A,B). This result indicates that the solvent selection is important in the design of the structure of metal NCs and their CSs. In the 2D CS consisting of newly formed Ag₁₂(S⁻Bu)₆(CF₃COO)₆ NCs, each Ag₁₂(S⁻Bu)₆(CF₃COO)₆ NC was linked to six adjacent Ag₁₂(S⁻Bu)₆(CF₃COO)₆ NCs via linkers to produce a highly symmetric 2D CS (Figure 21C). The layers were separated by 7.23 Å, with weak interactions between them. It was also revealed that the reversible structural transformation between 3D and 2D CSs could be induced by appropriate solvent selection (Figure 21A,B).

![Figure 21](image_url)

**Figure 21.** Comparison of the Ag₁₂ core structures in (A) Ag₁₂(S⁻Bu)₆(CF₃COO)₆(bpy)₃ (Ag₁₂bpy-2) and (B) Ag₁₂(S⁻Bu)₆(CF₃COO)₆(bpy)₄ (Ag₁₂bpy). (C) (a) Perspective view of an Ag₁₂S₆ node with six pendant bpy linkers (ORTEP drawing at the 50% probability level). (b) Stacking of the 2D network structure of Ag₁₂(S⁻Bu)₆(CF₃COO)₆(bpy)₃ viewed along the crystallographic c-axis. Ag = green, S = yellow, C = gray, N = blue; CF₃COO⁻, tBu, and H atoms are omitted for clarity. (D) 3D-excitation emission matrix of Ag₁₂(S⁻Bu)₆(CF₃COO)₆(bpy)₃ measured at −190 °C. (E) Thermochromic images of the (a) exterior [001] surfaces and (b) exposed interior [010]/[100] planes of Ag₁₂(S⁻Bu)₆(CF₃COO)₆(bpy)₃/NH₂ (bpy:bpy-NH₂ = 20:1) solvated single crystals under ultraviolet light irradiation. Reproduced with permission from Reference [169]. Copyright 2018 Wiley-VCH.

Structural deformation of the CSs also induced changes in their electronic structure and PL properties. For example, the 3D CS only showed PL at a single wavelength, regardless of temperature and excitation wavelength, whereas the 2D CS exhibited PL of two colors (blue and red), depending on the excitation wavelength (Figure 21D). To enhance the blue emission of the 2D CS, the researchers introduced bpy-NH₂, which itself emits blue light, as a linker, to fabricate a 2D CS containing two types of linkers, bpy and bpy-NH₂. The intensity ratio of the red and blue PL signals depended on the mixing ratio of linker molecules. At the optimum linker mixing ratio, the PL intensity ratio of the red and blue peaks depended on temperature. Therefore, this 2D CS containing two types of linkers could be used as a temperature sensor (Figure 21E).

In 2018, the same group also reported the formation of a 2D CS consisting of Ag₁₄(DT-o-C)₆ NCs [156]. The 2D CS with Ag₁₄(DT-o-C)₆, as a structural unit (Figure 22A), was fabricated by changing the linker structure from that used to form the 1D CS (Section 2.4 and Figure 13) to
dipyridin-4-yl-diazene (Scheme 2(9)). The obtained 2D CS possessed a rhombic network structure with Ag₁₂₄(DT-o-C₆)(CH₃CN)₄ as nodes (Figure 22B).

Table 2. Connection modes, NCs, linkers, publication years, and references for 2D CS.

| Connection Mode | NC                                      | Linker            | Year  | Ref.   |
|-----------------|-----------------------------------------|-------------------|-------|--------|
| Formation of Ag−O, Ag−S, Ag−Cl bond, etc. (Figure 2B) | Ag₂₀(CO₃)₁₀(S-OH)₁₀(CF₃COO)₉(CF₃COOH)(CH₃OH)₂ | -      | 2017  | [148]  |
|                 | Ag₂₀(CO₃)₁₀(S-C₆)₁₀(CF₃COO)₁₀(CF₃COOH)(H₂O)₂ | -      | 2019  | [149]  |
|                 | Ag₄₆(VO₂)₁₂(S-Et)₁₂(P₃S₃C₆)₁₂(CO₃)     | -      | 2019  | [150]  |
|                 | Ag₂₁Cl(N-L₆)(CF₃COO)₂·2CHCl₃            | -      | 2019  | [166]  |
|                 | Ag₁₁Cl(N-L₆)(CF₃SO₂)₂·2CHCl₃            | -      | 2019  | [166]  |
| Introduction of linker molecule (Figure 2D) | Ag₁₂(S⁻²Bu)₁₂(CF₃COO)₉ | bpy²⁺        | 2018  | [169]  |
|                 | Ag₁₄(DT-o-C₆)                           | dipyridin-4-yl-diazene ²⁻ | 2018  | [156]  |
|                 | Ag₁₂(S⁻²Bu)₁₂(CF₃COO)₉                 | TPPA²⁺       | 2018  | [170]  |
|                 | Ag₁₂(S⁻²Bu)₁₂(CF₃COO)₉                 | TPyP²⁺       | 2018  | [171]  |
|                 | Ag₁₂(S⁻²Bu)₁₂(CF₃COO)₉(DMF)            | bpy²⁺        | 2019  | [158]  |
|                 | Ag₁₀(CF₃COO)₁₂(S⁻²Bu)₁₂(CH₃CN)₁₂      | m-iah²⁺     | 2019  | [159]  |
|                 | Ag₁₂(S⁻²Bu)₁₂(CF₃COO)₉(CH₃CN)₆           | bpz-NH₂²⁺ | 2019  | [172]  |

*See Scheme 2.

Figure 22. (A) Structural unit and (B) 2D CS of Ag₁₂₄(DT-o-C₆)(CH₃CN)₄(dipyridin-4-yl-diazene)₂.
Reproduced with permission from Reference [156]. Copyright 2018 American Chemical Society.

In 2018, Zang et al. [170] reported the formation of a 2D CS consisting of Ag₁₂(S⁻²Bu)₁₂(CF₃COO)₉ and tris(4-pyridylphenyl)-amine (TPPA; Scheme 2(10) and Figure 23A). This structure is interesting because the distance between the 2D layers can be changed. In this 2D CS, DMAC used as a solvent existed between layers immediately after the synthesis and the 2D layers overlapped, as shown in Figure 23B(a),(d). When the DMAC was partially removed from this structure, the overlap of the 2D layers changed, as illustrated in Figure 23B(b),(e). Furthermore, when this structure was immersed again in the mother liquor, the structure changed to that depicted in Figure 23B(c),(f). It was also found that the size of the crystal and its emission characteristics changed in accordance with the overlap manner in the 2D CS.

Recently, this group also formed a 2D CS with Ag₁₂(S⁻²Bu)₁₂(CF₃COO)₉ NCs as nodes by using 5,10,15,20-tetra(4-pyridyl)porphyrin (TPyP; Scheme 2(11)) as a linker (Figure 24A) [171]. TPyP has a photosensitizing effect. Therefore, the ability of the 2D CS to degrade the toxic substance 2-chloroethyl ethyl sulfide (CEES), also called mustard gas, was studied (Figure 24B). The obtained 2D CS showed higher photocatalytic activity than that of a reported MOF. This high photocatalytic activity was ascribed to the synergistic effect of Ag NCs and TPyP, promoting the production of singlet oxygen, which induced the degradation of CEES (Figure 24B). The 2D CS maintained its crystallinity after the photocatalytic reaction and was able to be used repeatedly. The authors pointed out that photocatalytic activity could be further increased by selecting appropriate Ag NCs and organic molecular linkers.

Two other types of 2D CSs were also reported in 2019. In their paper on 1D CS formation (Figure 15 and Section 2.4), Bakr et al. [158] also reported that a 2D CS with Ag₁₄₁(Cl(S⁻²Bu)₈(CF₃COO)₈(DMF) as nodes was formed by changing the concentration of bpy during synthesis (Figure 25A). This Ag₁₄₁ Cl core contained Cl⁻ at its center as an anion template. Compared with the Ag₁₅₁(Cl(S⁻²Bu)₈(CF₃COO)₈₆.₇(NO₃)₀₃.₃(DMF)₂
node of the 1D CS, the node of the 2D CS (Ag\textsubscript{12}Cl(S\textsuperscript{-1}Bu\textsubscript{6})(CF\textsubscript{3}COO)\textsubscript{2}(DMF)) had one less Ag atom. However, the frameworks of these NCs were similar to each other (Figure 25B). The 2D CS showed higher thermal stability than those of individual Ag NCs and the 1D CS (Figure 15D). Unlike individual Ag NCs and the 1D CS, the 2D CS emitted green light with a strong intensity, even at room temperature (Figure 25C). Based on the results of a DFT calculation, it was interpreted that the enhancement of PL intensity was caused by a linker-to-cluster charge transfer excitation. In addition, Gao et al. [159] recently formed a 2D CS (Figure 26) with Ag\textsubscript{10}(CF\textsubscript{3}COO)\textsubscript{6}(S\textsuperscript{-1}Bu\textsubscript{6})(CH\textsubscript{3}CN)\textsubscript{4} nodes, using 3-pyridine carboxylic hydrazide (m-iah, Scheme 2(12)) as a linker, which is the meta equivalent of p-iah and o-iah used to form 1D CSs (Figure 16).

Thus, 2D CS formation by using a linker leads to the assembly of a structure with high stability and quantum yield (QY). However, 2D CS formation decreases the solubility of NCs, which limits their processability and device applicability. Therefore, Zang et al. [172] recently established a method to polymerize 2D CS to overcome this problem and provide materials suitable for practical use (Figure 27A). In this study, Ag\textsubscript{12}(S\textsuperscript{-1}Bu\textsubscript{6})(CF\textsubscript{3}COO)\textsubscript{6} NCs reported in their previous work (Figure 21) [168] were used as nodes. Moreover, 1,4-bis (pyrid-4-yl)benzenamine (bpz-NH\textsubscript{2}, Scheme 2(13)) was used as the linker. The amino group of the bpz-NH\textsubscript{2} linker played an important role in polymerization. First, 2D CS crystals consisting of Ag\textsubscript{12}(S\textsuperscript{-1}Bu\textsubscript{6})(CF\textsubscript{3}COO)\textsubscript{6} and bpz-NH\textsubscript{2} were fabricated (Figure 27B). The crystal size was limited to about 200–300 nm by quenching the reaction within 1 min. A 2D CS film was obtained by reacting the crystals with methacrylic anhydride (MA). MA bound to the amino group of bpz-NH\textsubscript{2} (Figure 27C) and was then polymerized with acrylate monomers butyl methacrylate (BMA) and triethylene glycol dimethacrylate (TEGDMA), as shown in Figure 27D.

![Figure 23](image-url) **Figure 23.** (A) Perspective view of an Ag\textsubscript{12}S\textsubscript{6} subunit with six pendant TPPA linkers in each layer. (B) Stacking of the 2D network structure of Ag\textsubscript{12}(S\textsuperscript{-1}Bu\textsubscript{6})(CF\textsubscript{3}COO)\textsubscript{6} by TPPA. AA stacking viewed along the (a) c-axis and (d) b-axis, AB stacking viewed along the (b) c-axis and (e) a-axis, and ABC stacking viewed along the (c) c-axis and (f) a-axis. Reproduced with permission from Reference [170]. Copyright 2018 Royal Society of Chemistry.

![Figure 24](image-url) **Figure 24.** (A) Synthesis of Ag\textsubscript{12}(S\textsuperscript{-1}Bu\textsubscript{6})(CF\textsubscript{3}COO)\textsubscript{2}(TPyP). (B) Schematic illustration of the capture and photodetoxification of CEES by Ag\textsubscript{12}(S\textsuperscript{-1}Bu\textsubscript{6})(CF\textsubscript{3}COO)\textsubscript{6}(TPyP). Reproduced with permission from Reference [171]. Copyright 2019 American Chemical Society.
newly formed Ag$_{12}$(S-tBu)$_6$(CF$_3$COO)$_6$ NCs, each Ag$_{12}$(S-tBu)$_6$(CF$_3$COO)$_6$ NC was linked to six adjacent Ag NCs as nodes by using DMAC as a solvent and tris(4-diazene (N-tBu)CN) called mustard gas, was studied. The obtained 2D CS showed higher photocatalytic activity than that of a reported MOF. This high photocatalytic activity was ascribed to the synergistic effect of Ag NCs and TPyP acting as a linker, which is the photosensitizing effect. Therefore, the ability of the 2D CS to degrade the toxic substance 2 showed higher thermal stability than those of individual Ag NCs and the 1D CS (Figure 15D). Unlike individual Ag NCs and the 1D CS (Figure 13D), the layers existed between layers immediately after the synthesis and the 2D layers overlapped, as shown in Figure 23B(a). This structure is interesting because the distance between the 2D layers can be changed. In this 2D CS, DMAC used as a solvent again in the mother liquor, the structure changed to that described in Figure 23B(c) (Figure 26, reproduced with permission from Reference [159]). Copyright 2019 American Chemical Society.

Figure 25. (A) Structure of Ag$_{14}$Cl(S-tBu)$_8$(CF$_3$COO)$_5$(DMF)(bpy)$_2$. Free (co-crystallized) DMF molecules are not shown. (B) Top views of the core structures of (a) Ag$_{15}$Cl(S-tBu)$_8$(CF$_3$COO)$_5$(NO$_3$)$_{0.67}$(DMF)$_2$ and (b) Ag$_{14}$Cl(S-tBu)$_6$(CF$_3$COO)$_5$(DMF). The green semitransparent spheres in the Ag NCs are shown as a visual guide. H atoms have been omitted for clarity. (C) Steady-state PL and excitation spectra of 2D CS crystals measured at room temperature (~25 °C). Emission spectra were measured under 365 nm excitation. Reproduced with permission from Reference [158]. Copyright 2019 American Chemical Society.

Figure 26. (A) Structural unit and (B) spatial stacking diagram of Ag$_{10}$(CF$_3$COO)$_4$(S-tBu)$_6$(CH$_3$CN)$_4$(m-iah)$_4$. Reproduced with permission from Reference [159]. Copyright 2019 American Chemical Society.
Thus, 2D CS formation by using a linker leads to the assembly of a structure with high stability and quantum yield (QY). However, 2D CS formation decreases the solubility of NCs, which limits their processability and device applicability. Therefore, Zang et al. [172] recently established a method to polymerize 2D CS to overcome this problem and provide materials suitable for practical use (Figure 27A). In this study, Ag$_{12}$(S-tBu)$_6$(CF$_3$COO)$_6$(bpz-NH$_2$)$_3$ NCs reported in their previous work (Figure 21) [168] were used as nodes. Moreover, 1,4-bis(pyrid-4-yl)benzene (bpz-NH$_2$, Scheme 2(13)) was used as the linker. The amino group of the bpz-NH$_2$ linker played an important role in polymerization. First, 2D CS crystals consisting of Ag$_{12}$(S-tBu)$_6$(CF$_3$COO)$_6$ and bpz-NH$_2$ were fabricated (Figure 27B). The crystal size was limited to about 200–300 nm by quenching the reaction within 1 min. A 2D CS film was obtained by reacting the crystals with methacrylic anhydride (MA). MA bound to the amino group of bpz-NH$_2$ (Figure 27C) and was then polymerized with acrylate.

The resulting membrane exhibited PL with a QY of 14.8%, which was higher than that of the unpolymerized 2D CS crystals (Figure 28A(a)). The increased PL intensity was ascribed to the polymerization strengthening the structure of the 2D CS, which suppressed molecular vibrations and thus nonradiative decay. The membrane was stable in water regardless of pH (Figure 28A(b)). The researchers also attempted to use the membrane to sense the harmful substance nitrobenzene in solution. The results revealed that the membrane was able to detect nitrobenzene with a sensitivity of 3.19 ppb (Figure 28B). This membrane also displayed high reusability (Figure 28C). These results indicate that polymerizing 2D CSs is an effective approach to obtain Ag NCs suitable for applications.
In 2017, Mak et al. [148] formed a 3D CS consisting of Ag_{14}(S-iPr)_6(CF_3COO)_{11}(H_2O)_3(CH_3OH) NCs (Figure 29A). The NCs were connected via O–Ag–O bonds formed between CF_3COO^- and Ag ions (Figure 29B,C). Each Ag NC was connected to six other NCs, thereby forming a distorted octahedral-like coordination structure. It was speculated that this 3D CS formed by the assembly of NCs after NC generation.

## 4. Three-Dimensional Structures

Ag NCs are often used as nodes in 3D CSs. Because 3D CSs generally possess stronger frameworks than those of 2D CSs, the formation of 3D CSs is effective to enhance the stability of Ag NCs and thereby improve their physical properties. In 3D CS formation, the principles of NC assembly are similar to those in 1D and 2D CS formation, although the ligands used are often different. In Sections 4.1–4.3, we focus on the assembly of 3D CSs via the formation of Ag–O, Ag–S, or Ag–Cl bonds (Figure 2B), control of counterions (Figure 2C), and the introduction of linker molecules (Figure 2D), respectively. The metal NCs, connection modes, linker molecules, year reported, and reference numbers for these studies are summarized in Table 3. Several of the ligands used in 3D CSs are shown in Scheme 1. The organic molecules used as linkers are illustrated in Scheme 2.

### 4.1. Connection via Ag–O, Ag–S, or Ag–Cl Bonds

In 2017, Mak et al. [148] formed a 3D CS consisting of Ag_{14}(S-iPr)_6(CF_3COO)_{11}(H_2O)_3(CH_3OH) NCs (Figure 29A). The NCs were connected via O–Ag–O bonds formed between CF_3COO^- and Ag ions (Figure 29B,C). Each Ag NC was connected to six other NCs, thereby forming a distorted octahedral-like coordination structure. It was speculated that this 3D CS formed by the assembly of NCs after NC generation.

Figure 28. (A) (a) Photographs of the Ag_{12} clusters, nano-NH_2-Ag_{12}bpz, and an Ag_{12}bpz membrane under 365 nm ultraviolet light irradiation. (b) PXRD patterns of NH_2-Ag_{12}bpz and the Ag_{12}bpz membrane upon treatment with water, base, and acid for different periods. (B) Fluorescence spectra showing the response of the Ag_{12}bpz membrane to the incremental addition of a nitrobenzene solution. (C) Cycling test of the Ag_{12}bpz membrane upon exposure to nitrobenzene vapor. Reproduced with permission from Reference [172]. Copyright 2019 Royal Society of Chemistry.
In 2019, Sun and colleagues also produced a 3D CS, and theirs consisted of Ag$_{44}$(Mo$_6$O$_{19}$)(S-Et)$_{24}$(SCl)$_3$ NCs (Figure 30A) containing a POM as an anion template (Figure 30B); these NCs were reported in their paper on 1D CS (Figure 9) and 2D CS (Figure 19) formation [149]. In this 3D CS, Mo$_6$O$_{19}^{2−}$ was used as an anion template, which was different from the case of the 1D and 2D CSs, in which the POM V$_{10}$O$_{28}^{6−}$ was located in the center of the cluster. This was the first report in which Mo$_6$O$_{19}^{2−}$ was used as an anion template of Ag NCs. Mo$_6$O$_{19}^{2−}$ has octahedral symmetry, and thereby the outer Ag$_{44}$(S-Et)$_{24}$ layer also displayed high symmetry (Figure 30C). The Ag$_{44}$ shell had six quadrangles and 24 pentagonal faces. Ag at the vertices of these six quadrangles was connected with one S atom and four Cl atoms, leading to the formation of a 3D CS consisting of Ag$_{44}$(Mo$_6$O$_{19}$)(S-Et)$_{24}$(SCl)$_3$ (Figure 30D). The S and Cl atoms used in the connections were generated by the decomposition of S-Et and CH$_2$Cl$_2$ during the CS synthesis.

The 3D CS in which Ag NCs are linked by dithiocarb, reported in 2019 by Gao et al. [173], should also be included in this category. The researchers first synthesized an Ag$_{11}$S(C$_5$NS$_2$H$_{10}$)$_9$ precursor (C$_5$NS$_2$H$_{10}$ = diethyldithiocarbamate, Scheme 1(20)) [174]. The obtained precursor was reacted under high pressure, in an autoclave, to form a 3D CS with Ag$_{17}$(C$_5$NS$_2$H$_{10}$)$_{14}$ as a repeating unit. This structure consists of Ag$_9$ NCs bound to twelve C$_5$NS$_2$H$_{10}$ (Figure 31A) and Ag$_5$ NCs containing six C$_5$NS$_2$H$_{10}$ (Figure 31B). The 3D CS (Figure 31C) was formed by the sharing of S between the two types of NCs. In these structures, the Ag$_9$ NCs were the nodes for three-point bridges and the Ag$_5$ NCs were the nodes for four-point bridges.

### Table 3. Connection modes, NCs, linkers, publication years, and references for 3D CS.

| Connection Mode | NC or Metal Ion | Linker | Year | Ref. |
|-----------------|----------------|--------|------|------|
| Formation of Ag−O−, Ag−S, Ag−Cl bond, etc. (Figure 2B) | Ag$_{14}$(S-Pr)$_6$(CF$_3$COO)$_{11}$(H$_2$O)$_3$(CH$_3$OH)Ag$_3$ | - | 2017 | [148] |
| | Ag$_{44}$(Mo$_6$O$_{19}$)(S-Et)$_{24}$(SCl)$_3$ | - | 2019 | [149] |
| | Ag$_{17}$(C$_5$NS$_2$H$_{10}$)$_{14}$ | - | 2019 | [173] |
| Control of counter ion (Figure 2C) | [Au$_2$Ag$_{22}$(S-Adm)$_2$(Cl)$_{24}$]$_2^-$ | SbF$_6^-$ | 2020 | [175] |
| | [Au$_2$Ag$_{22}$(S-Adm)$_2$(Cl)$_{24}$]$_{12}^{2+}$ | SbF$_6^-$ | 2020 | [175] |
| Introduction of linker molecule (Figure 2D) | Ag$_{12}$(S$^2$Bu)$_6$(CF$_3$COO)$_4$ | bpy$^a$ | 2017 | [168] |
The researchers first synthesized an Ag$_{17}$ subunit (Figure 32A(h)). In the second 3D CS, [Au$_{24}$] CS possessed a diamondlike structure (Figure 32A(f)) consisting of interpenetrating clockwise and counterclockwise optical isomers, depending on the winding method of the oligomer (Figure 32A(b),(c)).

In the first 3D CS, [Au$_{14}$Ag$_{44}$] CSs were linked by dithiocarb, reported in 2019 by Gao et al. [173], should be included in this category. The researchers first synthesized an Ag$_{17}$ subunit (Figure 32A(h)). In the second 3D CS, [Au$_{24}$] CS possessed a diamondlike structure (Figure 32A(f)) consisting of interpenetrating clockwise and counterclockwise optical isomers, depending on the winding method of the oligomer (Figure 32A(b),(c)).

Very recently, Zhu’s group synthesized two types of 3D CSs in which [Au$_1$Ag$_{22}$] CSs were linked by dithiocarb, reported in 2019 by Gao et al. [173], should be included in this category. The researchers first synthesized an Ag$_{17}$ subunit (Figure 32A(h)). In the second 3D CS, [Au$_{14}$Ag$_{44}$] CSs were linked by dithiocarb, reported in 2019 by Gao et al. [173], should be included in this category.

Very recently, Zhu’s group synthesized two types of 3D CSs in which [Au$_1$Ag$_{22}$] CSs were linked by dithiocarb, reported in 2019 by Gao et al. [173], should be included in this category. The researchers first synthesized an Ag$_{17}$ subunit (Figure 32A(h)). In the second 3D CS, [Au$_{14}$Ag$_{44}$] CSs were linked by dithiocarb, reported in 2019 by Gao et al. [173], should be included in this category.

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Figure 30. (A) Structure of (a) Ag$_{14}$[Mo$_{6}$O$_{19}$](S-Et)$_{24}$[SCl$_4$]$_3$ and (b) the Mo$_6$O$_{19}^{2-}$ anion template. Tetragons (yellow) and pentagons (green) in an Ag$_{22}$ cage are shown. (B) Connections (cyan polyhedra) between Ag$_{14}$ subunits (highlighted in different colors) in the 3D framework. (C) Framework and (D) simplified primitive cubic topology with the Ag$_{14}$ subunit as a node (represented as red balls) of the Ag$_{22}$[Mo$_{6}$O$_{19}$](S-Et)$_{24}$[SCl$_4$]$_3$ 3D CS. Reproduced with permission from Reference [149]. Copyright 2019 Royal Society of Chemistry.

Figure 31. (A) Structures of the Ag$_9$ cluster, (B) Ag$_5$ subunits, and (C) 3D framework of Ag$_{17}$[C$_5$N$_2$H$_{10}$]$_{14}$. Reproduced with permission from Reference [173]. Copyright 2019 Royal Society of Chemistry.

4.2. Control of Counterions

Very recently, Zhu’s group synthesized two types of 3D CSs in which [Au$_1$Ag$_{22}$] CSs were linked by dithiocarb, reported in 2019 by Gao et al. [173], should be included in this category. The researchers first synthesized an Ag$_{17}$ subunit (Figure 32A(h)). In the second 3D CS, [Au$_1$Ag$_{22}$] CSs were linked by dithiocarb, reported in 2019 by Gao et al. [173], should be included in this category. The researchers first synthesized an Ag$_{17}$ subunit (Figure 32A(h)). In the second 3D CS, [Au$_1$Ag$_{22}$] CSs were linked by dithiocarb, reported in 2019 by Gao et al. [173], should be included in this category.

Very recently, Zhu’s group synthesized two types of 3D CSs in which [Au$_1$Ag$_{22}$] CSs were linked by dithiocarb, reported in 2019 by Gao et al. [173], should be included in this category. The researchers first synthesized an Ag$_{17}$ subunit (Figure 32A(h)). In the second 3D CS, [Au$_1$Ag$_{22}$] CSs were linked by dithiocarb, reported in 2019 by Gao et al. [173], should be included in this category.
3D CS was formed by connecting \([\text{Au}_1\text{Ag}_{22}(\text{S-Adm})_{12}]^{3+}\) to three \(\text{SbF}_6^-\) via \(\text{Ag}-\text{F}-\text{Ag}\) bonds (Figure 32B(a)–(c)). This structure only contained clockwise or counterclockwise optical isomers (Figure 32B(d)). As a result, this 3D CS had a larger pore diameter (15 Å, Figure 32B(e)) than that of the first 3D CS (6.2 Å, Figure 32A).

Figure 32. (A) Structure of the \(\text{Au}_1\text{Ag}_{22}\) superatom complex and interpenetrating 3D channel framework assembled from alloy NC monomers (for clarity, C and H atoms are omitted). (a) Icosahedral \(\text{Au}_1\text{Ag}_{12}\) core, (b) cage-like \(\text{Ag}_{10}(\text{SR})_{12}\) complex shell, (c) a pair of \(\text{Au}_1\text{Ag}_{22}\) isomers, (d) the connection of \(\text{SbF}_6^-\) and alloy NCs, (e) two alloy NCs connected by \(\text{SbF}_6^-\), (f) tetrahedral structure of NC monomers (the inset shows methane), (g) topology of the diamond-like structure, and (h) interconnected channels of \(\text{Au}_1\text{Ag}_{22}\) along the \(z\)-axis. The left- and right-handed enantiomers in (c), (g), and (h) are highlighted in pink and yellow, respectively. Atoms are denoted in conventional colors: \(\text{Au} = \text{gold, Ag in core and the Ag}\mu_2\text{ motif} = \text{pale blue, Ag in the Ag}\mu_3\text{ motif} = \text{green, S} = \text{red, F} = \text{light turquoise, Sb} = \text{purple.}

(B) Crystal and channel structure of left-handed chiral 3D channel framework (C and H atoms are omitted for clarity). (a) The connection of Ag and \(\text{SbF}_6^-\), (b) the connection of Ag and \(\text{SbF}_6^-\), (c) two alloy NCs linked by \(\text{SbF}_6^-\), (d) illustration of the hexagonal network structure, and (e) schematic of the large hexagonal channel structure. Note that the packing pattern of the right-handed chiral 3D channel framework was the same as that of the left-handed chiral 3D channel framework. Reproduced with permission from Reference [175]. Copyright 2020 Wiley-VCH.
Study of the physical and chemical properties of the 3D CSs revealed that both exhibited red PL in the presence of polar solvents such as CH₃OH, ethanol, and water, which disappeared when the solvent was evaporated (Figure 33A). This behavior indicates that the obtained 3D CSs can function as sensors for polar solvents. The 3D CS composed of only the right- or left-handed enantiomer exhibited circularly polarized luminescence (CPL) (Figure 33B).

Figure 33. (A) Luminescence switching response to protic solvents. Fluorescence of films coated with (a) the interpenetrating 3D CS (SCIF-1) and (b) the left-handed 3D CS (SCIF-2) before and after solvent evaporation. Inset images show solvatochromic photographs of the corresponding films excited with 365 nm ultraviolet light before (left) and after desolvation (middle), and fluorescence recovery after treatment with n-hexane containing 5% ethanol (right). (B) CPL spectra of SCIF-1, SCIF-2 (left-handed), and SCIF-2 (right-handed) single crystals and the superstructures of these three crystal samples. (a) CPL spectra of SCIF-1, SCIF-2 (left-handed), and SCIF-2 (right-handed). Insets show photographs of the corresponding crystals. (b) Crystal structure of the SCIF-1 framework. (c) Crystal structure of the SCIF-2 (left-handed framework). (d) Crystal structure of the SCIF-2 (right-handed) framework. Reproduced with permission from Reference [175]. Copyright 2020 Wiley-VCH.

4.3. Introduction of Linker Molecules

The formation of 3D CSs by using linker molecules is a technique often used to fabricate molecular assemblies and MOFs. When a 3D CS composed of metal NCs is formed by such a method, in addition to increasing the stability of the NCs, it is also expected to adsorb gas molecules within its pores and behave as a catalyst with high selectivity because of the narrow pores. Furthermore, because the metal NCs, which are used as nodes, have more diversity in terms of coordination direction than that of metal ions, metal NC-based MOFs may have different connection modes from those of normal MOFs formed by using metal ions as nodes, and thereby they construct novel framework structures. Thus, metal NC-based MOFs possess not only the characteristics of individual metal NCs and MOFs, but also the possibility to produce new functions through synergistic effects.

Zang’s group have been energetically researching 3D CSs with linkers, as well as the cases of 1D and 2D CSs. First, in 2017, Zang et al. [168] formed an Ag₁₂ NC-based MOF (Ag₁₂(S⁻⁻Bu₆)(CF₃COO)₄(bpy)₄)
in which Ag$_{12}$(S-Bu)$_6$(CF$_3$COO)$_4$ was bridged by bpy (Figure 34A). The obtained Ag$_{12}$ NC-based MOF possessed a bilayer structure (Figure 34B). The formation of such a 3D CS markedly improved the stability of the Ag$_{12}$ NCs. For example, a crystal of the individual Ag$_{12}$(S-Bu)$_6$(CF$_3$COO)$_4$(CH$_3$CN)$_6$ NCs discolored in just 30 min when left in the atmosphere. In contrast, the Ag$_{12}$ NC-based MOF showed almost no change in crystallinity, even when left in the air for one year (Figure 34C). The 3D CS also showed high stability during long-term gas adsorption and irradiation with visible light for several hours.

![Figure 34](image-url)

**Figure 34.** (A) Schematic representation of the ligand-exchange strategy used to obtain Ag$_{12}$(S-Bu)$_6$(CF$_3$COO)$_4$(bpy)$_4$ crystals (Method 1, giving low yield) and one-pot synthesis (Method 2, for gram-quantity production) under identical conditions. Interconnected channels of Ag$_{12}$(S-Bu)$_6$(CF$_3$COO)$_4$(bpy)$_4$ viewed along the $a$ and $b$ axes, where the yellow surface represents the pore surface. Ag = green, C = gray, O = red, S = yellow, F = turquoise. H atoms are omitted for clarity. Inset are photographs showing the changes of Ag$_{12}$(S-Bu)$_6$(CF$_3$COO)$_4$(CH$_3$CN)$_6$ and Ag$_{12}$(S-Bu)$_6$(CF$_3$COO)$_4$(bpy)$_4$ crystals under ambient conditions. (B) Schematic representation of the topology of Ag$_{12}$(S-Bu)$_6$(CF$_3$COO)$_4$(bpy)$_4$ along the $c$-axis. The square windows in one double layer are blocked by Ag-S cluster nodes of adjacent layers. (C) PXRD patterns of Ag$_{12}$(S-Bu)$_6$(CF$_3$COO)$_4$(bpy)$_4$ (in ascending order): simulated, experimental, gram-scale synthesis, after gas adsorption experiments ($O_2$, $N_2$), ethanol, after sensing experiments ($O_2$/$N_2$, $O_2$/vacuum, ethanol/air), after 10 h of visible-light irradiation under a Xe lamp equipped with a 420 nm cutoff filter, and the sample vial after one year under ambient conditions. Reproduced with permission from Reference [168]. Copyright 2017 Springer-Nature.
The formation of the 3D CS greatly changed the PL properties of the NCs. The individual Ag$_{12}$ NCs exhibited red PL with low QY. Conversely, the Ag$_{12}$ NC-based MOF exhibited green PL under vacuum, which was quenched by O$_2$ in the atmosphere (Figure 35A). The PL emission wavelength of the 3D CS under vacuum was independent of temperature and excitation wavelength, and its QY was 60 times higher than that of individual Ag$_{12}$ NCs. The authors ascribed this high QY to the efficient suppression of nonradiative decay in the 3D CS. Moreover, the fact that the PL of the 3D CS is quenched by O$_2$ means that it is highly sensitive to O$_2$. The 3D CS showed a fast response to O$_2$ in experiments in which the atmosphere was repeatedly switched between air and N$_2$. No such O$_2$ response was observed for the individual Ag$_{12}$ NCs. Based on these results, they suggested that the Ag$_{12}$ NC-based MOF can be applied as an O$_2$ sensor. In addition, the Ag$_{12}$ NC-based MOF was able to adsorb volatile organic compounds (VOCs) in its pores. The VOC-containing Ag$_{12}$ NC-based MOF exhibited different PL colors, depending on the kind of VOC (Figure 35B). This indicates that the Ag$_{12}$ NC-based MOF displays solvatochromism and therefore can be used for VOC detection.

In a paper on Ag$_{14}$(DT-o-C)$_6$ NCs published in 2018 (Figures 13 and 22), the same group reported that a 3D CS containing Ag$_{14}$(DT-o-C)$_6$ NCs formed when bpy was used as a linker (Figure 36A) [156]. The Ag$_{14}$ NC nodes in the 3D CS had a face-centered cubic structure like that of other Ag$_{14}$ NCs. This 3D CS was formed by connecting the eight vertices of each Ag$_{14}$ NC with bpy linker molecules. However, this 3D CS was not stable after solvent evaporation, similar to the case of the corresponding 1D CS (Figure 13) and 2D CS (Figure 22) [156]. Therefore, they synthesized a 3D CS with an interpenetrating framework by using 1,4-bis(d-pyridyl)benzene (Scheme 2(14)) as a linker, in order to decrease the pore size and form a 3D CS with a strong framework (Figure 36B). The obtained 3D CS showed high thermal stability, remaining intact up to 220 °C (Figure 36C), and possessed pores with a diameter of
about 1.12 nm (Figure 36D). This 3D CS showed optical absorption over a wide wavelength range and thermochromism (Figure 36E).

Figure 36. Three-dimensional CSs composed of (A) Ag$_{14}$(DT-o-C)$_6$ NCs and bpy (SCAM-3) and (B) Ag$_{14}$(DT-o-C)$_6$ NCs and 1,4-bis(4-pyridyl)benzene (SCAM-4). (C) Variable-temperature PXRD of SCAM-4 from 25 to 350 °C. (D) Different view of SCAM-4. (E) Evacuated SCAM-4 (excited at 380 nm) from −190 to 25 °C in air. Inset are corresponding photographs of SCAM-4 under 365 nm ultraviolet light irradiation. Reproduced with permission from Reference [156]. Copyright 2018 American Chemical Society.

Zang et al. [176] have also produced several other functional Ag NC-based MOFs. For example, in 2018, they reported the synthesis of a flexible Ag NC-based MOF. This structure consisted of 2D layers of Ag$_{10}$(S-^Bu)$_6$(CF$_3$COO)$_2$(PhPO$_3$H)$_2$ NCs linked via bpy, which were stacked through hydrogen bond (O-H···O) and C-H···O interactions, to form the 3D CS (Figure 37A,B). The 2D CS layers were thus linked by weak interactions, which facilitated the sliding of the layers, allowing the 3D CS to undergo structural deformation in response to guest organic molecules (Figure 37C). This Ag$_{10}$ NC-based MOF exhibited green PL in air. Upon inclusion of guest organic molecules, it exhibited PL with an emission color depending on the guest organic molecule (Figure 37D). As such, this Ag$_{10}$ NC-based MOF has potential as a sensor for distinguishing VOCs by its PL color. In 2019, they also formed a 3D CS consisting of Ag$_{12}$(S-^Bu)$_6$(CF$_3$COO)$_6$ NC nodes and 2,5-bis(4-cyanophenyl)-1,4-bis(4-(pyridine-4-yl)-phenyl)-1,4-dihydropyrrolo[3,2-b]pyrrole (CPPP, Scheme 2(15)) as a linker (Figure 38A) [177]. This was the first report in which a nitrile group (-CN) was used to link Ag NCs. The obtained 3D CS exhibited PL with a higher QY (61%) than that of CPPP in solution and solid states because the aggregation-induced quenching of CPPP was suppressed in the 3D CS (Figure 38B).
D-B-C e Ag NC was suppressed, which changed the excited-state dynamics of the 3D CS. The change in the emission behavior of the Ag NC-based MOF induced by DMAC was ascribed to this change of its excited-state emitting molecule. The PL wavelength of the 3D CS depended on the presence or absence of DMAC in its pores immediately after synthesis (Figure 40A). When the obtained 3D CS was exposed to the atmosphere, DMAC was removed, while maintaining the framework of the 3D CS.

In the above 3D CSs, the nodes consisted of only one kind of Ag NCs. Tang et al. [178] synthesized an Ag NC-based MOF that had two types of Ag NCs as nodes. This Ag NC-based MOF was composed of 1,1,2,2-tetrakis(4-(pyridin-4-yl)phenyl)ethene (tppe, Scheme 2(16)), Ag$_{10}$(S-$^t$Bu)$_6$(CF$_3$COO)$_2$(PhPO$_3$H)$_2$(bpy)$_2$ with complementary hydrogen bonding (O–H–O; the H–O distance is 1.750 Å) between interlayer −PO$_2$OH moieties. Illustration of reversible pore open/closed structural transformation induced by CH$_2$Cl$_2$, CHCl$_3$, and CCl$_4$ (represented as space-filling models) and switchable solvatochromism. (D) (a) Luminescence images of Ag$_{10}$(S-$^t$Bu)$_6$(CF$_3$COO)$_2$(PhPO$_3$H)$_2$(bpy)$_2$/solvent (guest free, CH$_2$Cl$_2$, CHCl$_3$, CCl$_4$, 1,4-dioxane, cyclohexane, DMAC, and acetone) combinations under 365 nm ultraviolet light irradiation. (b) Emission maxima of various Ag$_{10}$(S-$^t$Bu)$_6$(CF$_3$COO)$_2$(PhPO$_3$H)$_2$(bpy)$_2$/solvent combinations at room temperature. Reproduced with permission from Reference [176]. Copyright 2018 American Chemical Society.

Wang and colleagues reported the formation of a 3D CS, using metal NCs as linkers, before the use of metal NCs as nodes was developed [179]. In 2014, they succeeded in forming an NbO-type MOF by using Ag ions as nodes and [C(Au-mdppz)$_2$][(BF$_4$)$_2$] (mdppz = 2-(3-methylpyrazinyl)diphenylphosphine, Scheme 2(17)) NCs as linkers (Figure 41). [C(Au-mdppz)$_2$][(BF$_4$)$_2$] has a framework with C in the center.
(Figure 41A) and is luminescent. The 3D CS was formed by the outer N atom of mdppz (Scheme 2(17)) binding to an Ag ion (Figure 41B,C). The obtained 3D CS consisted of two interpenetrating frameworks (Figure 41D,E) with a 1D channel in the c-axis direction (Figure 41C). Because a luminescent NC was used as the linker, the obtained MOF also showed green PL. The 3D CS displayed a PL QY of 25.6%, which was much higher than that of the luminescent NCs (1.5%). This increase of QY was caused by the strengthening of the framework of the linker NCs upon MOF formation and the excited-state perturbation induced by the coordination of Ag ions.

Figure 38. (A) Structures of (a) Ag chalcogenolate cluster nodes, (b) cage in Ag₁₂(S⁻⁷Bu₆)(CF₃COO)₆(CPPP)₂ (Ag₁₂CPPP), and (c) distribution of the cages in Ag₁₂CPPP. All H atoms and guest solvent molecules are omitted for clarity. (B) Solid-state absorption (dashed lines) and emission (solid lines) spectra of Ag₁₂CPPP and CPPP at room temperature. Inset are photographs of the crystals of CPPP and Ag₁₂CPPP under 365 nm ultraviolet light irradiation. Reproduced with permission from Reference [177]. Copyright 2019 Wiley-VCH.

Figure 39. Structure of the Ag₈ cluster, Ag₁₂ cluster, and tppe ligand, and single net of the [Ag₁₂(S⁻⁷Bu₆)(CF₃COO)₆]₁₀[Ag₈(S⁻⁷Bu₄)(CF₃COO)₄]tppe₂(DMAC)₁₀ framework viewed along the c-axis. DMAC molecules are omitted for clarity. Reproduced with permission from Reference [178]. Copyright 2019 Wiley-VCH.
Figure 40. (A) Distribution of DMAC molecules in the [Ag_{12}(S-Bu)_6(CF_3COO)_6]_{0.5}[Ag_{8}(S-Bu)_4(CF_3COO)_4](tppe)_2(DMAC)_{10} framework. (B) (a) Gradual fluorescence changes of the same [Ag_{12}(S-Bu)_6(CF_3COO)_6]_{0.5}[Ag_{8}(S-Bu)_4(CF_3COO)_4](tppe)_2(DMAC)_{10} crystal under atmospheric exposure and (b) normalized fluorescence spectra of [Ag_{12}(S-Bu)_6(CF_3COO)_6]_{0.5}[Ag_{8}(S-Bu)_4(CF_3COO)_4](tppe)_2(DMAC)_{10}. (C) (a) Proposed fluorescence decay paths in [Ag_{12}(S-Bu)_6(CF_3COO)_6]_{0.5}[Ag_{8}(S-Bu)_4(CF_3COO)_4](tppe)_2(DMAC)_{10} (path a) and [Ag_{12}(S-Bu)_6(CF_3COO)_6]_{0.5}[Ag_{8}(S-Bu)_4(CF_3COO)_4](tppe)_2 (I) (path b) and (b) fluorescence decay profiles of [Ag_{12}(S-Bu)_6(CF_3COO)_6]_{0.5}[Ag_{8}(S-Bu)_4(CF_3COO)_4](tppe)_2 in DMAC, THF, toluene, and DMF (denoted as 1 ⊂ DMAC, 1 ⊂ THF, 1 ⊂ Toluene, and 1 ⊂ DMF, respectively). Reproduced with permission from Reference [178]. Copyright 2019 Wiley-VCH.
Figure 41. (A) Structure of $[\text{C}(\text{Au-mdppz})_6](\text{BF}_4)_2$. (B) Four extensions: Ag3, Ag3a, Ag3b, and Ag3c. Note that Ag2 and Ag2a are not involved in the structural extension. (C) Perspective view of the 3D CS along the $c$ direction. Au = orange, Ag = green, P = purple, N = blue, C = gray. (D) Two interpenetrating nets shown in different colors; anions and solvent molecules are omitted for clarity. (E) Schematic representation of NbO topology in the 3D CS. Reproduced with permission from Reference [179]. Copyright 2014 Wiley-VCH.

5. Summary

In this review, representative studies on the formation of 1D, 2D, and 3D CSs in which metal NCs were self-organized and regularly linked were summarized. From this summary, the following points became clear.

(1) **Methods.** The methods to connect metal NCs that have been reported to date can be roughly divided into the following five categories: (i) direct connection by formation of metal–metal bonds (Figure 2A); (ii) connection by Ag–O, Ag–S, or Ag–Cl bond formation (Figure 2B); (iii) connection by counterions (Figure 2C); (iv) connection by linker molecules (Figure 2D); and (v) connection by inter-ligand interactions (Figure 2E; not introduced in this review).

(2) **Diversity.** Among CSs produced by the above methods, there are many examples of the formation of 1D, 2D, and 3D CSs through the use of methods (ii) and (iv). An important point when constructing CSs by these methods is the design of the ligand of the NCs and linker,
respectively. It is presumed that the control of these species is relatively easy, which has led to the wider utilization of methods (ii) and (iv) than of the other methods. In particular, for method (iv), existing knowledge obtained in the study of normal MOFs can be considered.

(3) **Metal element.** To directly connect metal NCs, it is effective to use Au as a main element because it forms strong aurophilic interactions (intermetal interactions). In the connections involving metal–O or metal–Cl bonds, it is effective to use Ag as a main element because it readily bonds with O or Cl. Moreover, in the connections using bpy as a linker, Ag is attractive as the main element because of the high affinity of N and Ag.

(4) **Stability.** The formation of a CS generally improves the thermal stability of the component metal NCs regardless of the connection mode.

(5) **Electronic structure.** The formation of a CS often causes the band gap of the NC to narrow. This means that CS formation allows the use of a broader wavelength range of light, opening up the possibility of visible-light-driven photocatalysis by using CSs.

(6) **PL properties.** For 2D and 3D CS using linkers, CS formation often leads to an increase in PL emission intensity. When metal NCs are the PL source, there are many cases in which dual emission peaks appear upon connection with a linker. In addition, the PL color of a CS often changes depending on the kind of VOC trapped in its pores.

(7) **Electrical conduction.** The electron conductivity of CSs changes dramatically depending on the distance between each metal NC and the mode of connection; 1D CS formed by the direct connection via metal–metal bond shows the higher conductivity than 1D CS connected through counter ion.

(8) **Possible applications.** The reported CSs have potential applications in fields such as electronic devices, luminescent devices, gas and temperature sensing, and photocatalysis.

This review allowed us to obtain a common understanding of the CSs reported to date and their functions. We hope that the knowledge thus clarified will lead to clear design guidelines for developing new CSs with desired functions in the future.

### 6. Outlook

It is expected that the following studies will be conducted in the future, leading to new CSs.

(1) **Use of other metal elements.** At present, mostly Au and Ag are used as the metal element in CSs. This is largely related to the high stability of Au and Ag NCs. For Ag NCs, the good connectivity between Ag and linker molecules is also related to this fact. On the other hand, several syntheses of individual copper (Cu) NCs have been reported recently [10,180–183]. In addition, other metal ions are often used in normal MOFs with metal ions as nodes [153]. CS formation of NCs based on Cu or other metals may also lead to materials with high thermal stability. In the future, it is expected that many elements will be used in CSs, thereby realizing various functions and decreased cost of such materials.

(2) **Use of the alloying effect.** At present, there are few examples in which alloy NCs are connected to form CSs [142,143,146]. Mixing different elements leads to NCs with physical/chemical properties and functions that are different from those of monometal NCs. In fact, for individual metal NCs, many cases have been reported in which new physical properties/functions appeared because of mixing/synergistic effects [66,184–194]. The previous studies have established basic techniques for the formation of CSs consisting of Ag NCs. In the future, it is expected that more functional materials will be created by extending such CS formation techniques to Ag-based alloy NCs.

(3) **Connection of reported metal NCs.** Ag NC-based MOFs are interesting because they can be synthesized by a one-pot process. However, in CSs formed by such a method, metal NCs that are stable only in the CS are often found as nodes. For individual metal NCs, many NCs have already been synthesized with atomic precision [1–65]. In addition, much information has been
obtained on methods to generate novel functions in such NCs, including alloying [66–68]. In the future, it is expected that a method to more effectively utilize the reported metal NCs in CSs will be found. To achieve this, it may be necessary to establish new connection methods different from those described in this review (Figure 2).

(4) **Elucidation of electronic conductivity.** We believe that 1D CSs may be applied as nanodevices. However, at present, few experiments on the conductivity of 1D CSs have been reported [146,152]. In the future, it is expected that the conductivity of 1D CSs will be measured as a basic physical property. It is anticipated that the accumulation of such information will eventually lead to the production of nanodevices based on 1D CSs of metal NCs.

(5) **Exploration of other possible applications of connected structures.** Various applications, such as gas storage, gas separation, gas conversion, and reaction-selective catalysis, have been studied for normal MOFs with metal ions as nodes [117]. It has also been reported that, in the case of self-assembled complexes, a reaction different from that in the case of using an ordinary flask proceeds in the cage structure (i.e., the cage behaves as a nanoflask) [116]. In the future, it is expected that these possibilities will be investigated for metal NC-based MOFs and that their functions will be much higher than those of conventional MOFs and self-assembled complexes.

As mentioned in Section 1, the advance of bottom-up technology is essential for the further development of nanotechnology in the future. In previous research, multiple techniques to generate metal atoms on the molar scale in a solution and to self-organize them to form nanomaterials with the same number of constituent atoms and the same number of molecules have been studied; that is, precise synthesis techniques of metal NCs have been developed. However, to apply these metal NCs as devices and next-generation materials, developing techniques to assemble metal NCs to a size that is easy to handle is necessary (Figure 1). We hope that technologies that allow the self-organization of regularly arranged CSs composed of metal NCs will be further developed in the future. Ultimately, such nanotechnology is expected to enable resource conservation, energy conservation, decreased waste and environmental load, and the better use of time by human society.

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