Ligand exchange reactions on thiolate-protected gold nanoclusters

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As a versatile post-synthesis modification method, ligand exchange reaction exhibits great potential to extend the space of accessible nanoclusters. In this review, we summarized this process for thiolate-protected gold nanoclusters. In order to better understand this reaction we will first provide the necessary background on the synthesis and structure of various gold clusters, such as Au_{25}(SR)_{18}, Au_{38}(SR)_{24}, and Au_{102}(SR)_{44}. The previous investigations illustrated that ligand exchange is enabled by the chemical properties and flexible gold–sulfur interface of nanoclusters. It is generally believed that ligand exchange follows a SN2-like mechanism, which is supported both by experiments and calculations. More interesting, several studies show that ligand exchange takes place at preferred sites, i.e. thiolate groups –SR, on the ligand shell of nanoclusters. With the help of ligand exchange reactions many functionalities could be imparted to gold nanoclusters including the introduced of chirality to achiral nanoclusters, size transformation and phase transfer of nanoclusters, and the addition of fluorescence or biological labels. Ligand exchange was also used to amplify the enantiomeric excess of an intrinsically chiral cluster. Ligand exchange reaction accelerates the prosperity of the nanocluster field, and also extends the diversity of precise nanoclusters.

1. Introduction

During the past decade, the interest in monolayer-protected metal clusters has drastically increased. The progress in the field has been reported in review articles focussing for example on precise synthesis,\textsuperscript{1–3} metal doping,\textsuperscript{4–7} and applications of gold nanoclusters.\textsuperscript{8,9} In general, the chemical properties of nanomaterials largely depend on their surface properties. As a flexible surface chemistry method, ligand exchange reaction (LER) is therefore an important tool for the development of nanomaterials. For instance, Dinkel and co-workers summarized the findings obtained by \textit{in situ} spectroscopic measurements to probe the ligand exchange at the surface of colloidal gold and silver nanoparticles. Specifically, the results obtained by the inherently surface sensitive technique second-harmonic scattering (SHS) was discussed.\textsuperscript{10} In addition, Xu and co-workers concentrated on exchange reactions on metal–organic...
frameworks (MOFs). More focussing on metal clusters, Zhu and co-worker published a review on transformations of precise nanoclusters induced by ligand exchange. They focused on the size conversion of gold and silver nanoclusters during ligand exchange reaction. However, a comprehensive summary focussing on the ligand exchange reaction of thiolate-protected gold clusters is still rare.

Recently, many new findings have emerged on the mechanism and peculiarities of the ligand exchange reactions on thiolate-protected clusters, including ligand exchange between clusters, ligand exchange including chiral thiols, and ligand exchange induced amplification of enantiomeric excess. These new findings are of great interest and further expanded the knowledge in the field of the ligand exchange reaction on thiolate-protected gold nanoclusters. Considering the recent findings in the field and its implications for applications an attempt to summarize to current knowledge seems appropriate.

The ligand exchange reaction is an important post-synthesis method for the modification and further functionalization of gold nanoclusters. In this review, we specifically address the ligand exchange reactions (LERs) on thiolate-protected gold nanoclusters. The main points to be addressed are illustrated in Scheme 1. We mainly focus on atomically precise gold nanoclusters with known crystal structure, such as Au_{25}(SR)_{18}, Au_{38}(SR)_{24}, and Au_{102}(SR)_{44}, which allows to obtain molecular level insight into the process. The clusters contain diverse surface motifs, e.g. monomeric and dimeric Au–sulfur staples and the distinct surface thiolates show different reactivity towards ligand exchange. It is also emerging from recent experiments and calculations that the surface region of these clusters, i.e. the Au–sulfur interface, is far from being static. Furthermore, not only free thiols in solution can exchange with thiols on the cluster, but thiols can exchange among the clusters without addition of free thiols. In addition, LERs also takes place between free thiols and supported clusters. Ligand exchange can lead to important modifications of the cluster properties. For example, LERs can induce chirality on achiral nanoclusters and also lead to the enhancement of enantiomeric excess of chiral clusters. LERs can promote size transfer of clusters, and phase transfer between organic and aqueous phase. In general, ligand exchange reaction is an important process that largely extends the diversity of precise nanoclusters and their potential for applications. In the following, after an introduction to monolayer-protected gold clusters, we summarize the current knowledge on LERs with special focus on the more recent findings.

2. Precise monolayer-protected gold nanoclusters

2.1 Synthesis of precise gold nanoclusters

Metal nanoclusters are composed of tens to hundreds of atoms and bridge the gap between single atoms or molecules and nanoparticles. Different from the plasmon excitation in nanoparticles, which is a collective excitation of conduction electrons, metal nanoclusters are characterized by strong quantum confinement effect and molecular-like properties. Nowadays, metal nanoclusters include several sub-classes, such as thiolate-protected silver nanoclusters, phosphine-protected gold nanoclusters, thiolate-protected gold nanoclusters and so on. Among them, thiolate-protected gold nanoclusters are the most studied due to their relatively long history and unique properties. In the 1990s, triggered by the work of Brust and Schiffrin, several groups started the research on gold nanoclusters and obtained a series of ultrasmall species. Typically in these syntheses mixtures of different cluster sizes were obtained. At this size every atom counts and minor changes in composition can have large effects on electronic and optical properties of the nanoclusters. Therefore, the precise control of the composition and purity is critically important.

In order to obtain pure nanoclusters, Whetten and co-workers tried to separate cluster mixtures by polyacrylamide gel electrophoresis (PAGE) and solvent-extraction, however, it is challenging to obtain pure cluster samples in this way. Murray and co-workers prepared the Au_{25} cluster by a two-phase protocol or the conversion of Au_{11} to Au_{25}, but mixtures of clusters of different size were obtained. After purification by solvent-extraction and other methods, clusters of different size were obtained with very low yield. Later, Tsukuda and co-workers applied PAGE, recycling size exclusion chromatograph and solvent-extraction to isolate the precise nanoclusters in a time-consuming strategy. After about ten years of struggling to obtain pure cluster samples in high yield, there was a strong need for efficient methods towards synthesis of precise gold nanoclusters.

The first breakthrough was reported for Au_{25}, a prototypical gold cluster. In 2008 Zhu and co-workers reported a high yield synthesis of Au_{25}(SCH_{2}CH_{2}Ph)_{18} by kinetic control with a two-phase method. Later on, by using THF as solvent, Wu and co-workers achieved a big improvement of the synthesis of Au_{25} nanoclusters using a one-pot method. Importantly, a “size-
focusing” process was identified in the growth of Au25 nanoclusters, which accounts for the formation of atomically monodisperse Au25 nanoclusters from the one-pot reaction.

Similar to the success in the synthesis of monodisperse Au25, a facile solution synthetic method for obtaining monodisperse Au38(S–C12H25)24 was developed by exploring a two phase ligand exchange process in which glutathione-capped Aun clusters are utilized as the starting material. Here a LER was carried out at elevated temperature (e.g. 80 °C) to obtain high purity Au38(S–C12H25)24 clusters, still at relatively low yield. After that, Qian and co-workers modified the protocol by replacing methanol by acetone in the first step (synthesis of Au38(SG)18) and by using an excess of phenylethylthiol (PhC2H4SH) as the etching ligand for the ligand exchange reaction. Using these modifications they could increase the synthesis yield of Au38 to 25% (Au atom). This work illustrated that the formation of Au38(SG)18 nanoclusters resulted from the conversion from larger clusters in the thiol etching induced growth process in a remarkable “size-focusing” process. The above results revealed that size-focusing is a very efficient methodology for synthesizing atomically precise gold nanoclusters. The underlying principle can be illustrated as shown in Fig. 1. Initially several nanoclusters with different sizes and vastly different stability coexist in the reaction mixture. The most stable nanocluster within the initially controlled size distribution will be sieved out. After the success in obtaining Au25 and Au144 nanoclusters using the size focusing method, a series of Au38(SR)m nanoclusters were synthesized in high yield and on a large scale, using this methodology, such as Au32,34 Au40,35 Au99,36 Au144 (ref. 37) and Au33.38 As the size-focusing involves core-etching from larger clusters to smaller clusters, the size (distribution) of the initial thiolate-protected gold cluster is critical. In addition, the high yield product may still coexist with some by-products, making additional purification steps such as size exclusion chromatography (SEC) necessary. Knappe and co-workers used SEC to separate Au38 and Au144 clusters, which differ by only two gold atoms. This example demonstrates the power of this separation technique.

The development of synthesis methods resulted in several monodisperse nanoclusters including Au25(SR)18, Au38(SR)24, Au99(SR)42,36 and Au144(SR)60.37 Such clusters have already become excellent candidates for applications in catalysis,36,40,41 as optical active materials,42,43 as chemical sensors,44,45 and for biological applications.46,47 The availability of numerous atomically precise thiolate-protected gold clusters allowed one to study the evolution of properties as a function of size. Compared with the bulk metal, which has continuous energy band structure, ultrasmall nanoclusters possess size-dependent electronic structure, a HOMO–LUMO gap and molecule-like properties. The HOMO–LUMO gap of Au25(SR)18 was found to be 1.3 eV,48 which is bigger than the value of 1.0 eV found for Au38(SR)24.33 This implies that the larger clusters have a smaller energy gap, as expected. Having many additional clusters at hand, the relationship between HOMO–LUMO gap and size of the nanoclusters became clearer. There is a general trend of shrinking gap with increasing size. Plots of the HOMO–LUMO gap Es against n1/3 or against ln(n) (n: number of gold atoms) revealed linear relationship and two groups of clusters.7 The two sets of clusters have difference structure and atomic packing, which will be addressed in the next subsection.

2.2 Structure of precise monolayer-protected gold nanoclusters

In order to understand the fundamental science of nanoclusters, it is necessary to determine their structures. X-ray single-crystal diffraction is the most straightforward and important technique for the structure determination of nanoclusters. Among the most important milestones in the research history of structural characterization of nanoclusters goes back to 2007, when Kornberg and co-workers published the first total X-ray structure of a thiolate-protected gold cluster, containing 102 gold atoms and 44 p-mercaptobenzoic acid (p-MBA) ligands.49 The structure of Au102(p-MBA)44 was unexpected and did not match the view emerging from the “cluster of clusters” growth mechanism proposed for cluster chemistry.49 As shown in Fig. 2a, the gold core has a decahedron structure and is surrounded by a gold thiolate layer characterized by “staple” motifs. In these structures, one gold atom is bound to two sulfur atoms. Similar structures were previously predicted theoretically as gold-thiolate rings.50 The Au102(p-MBA)44 cluster contains 19 short and two long motif SR(AuSR)x (x = 1, 2). In addition, the cluster is chiral because of the geometry of the equatorial atoms (Fig. 2b). The stability of the cluster structure has several origins. The cluster has 58 free electrons and thus has a closed electronic shell within a superatom model.51 The thiolate monolayer contributes as well to the stability of the cluster and within this layer additional stability stems from the π–π stacking between phenyl rings of the p-MBA molecules.

Owing to the development of precise synthesis of nanoclusters and X-ray single-crystal diffraction, structure determination of gold nanoclusters ushered in a period of vigorous development after the first breakthrough of Au102(p-MBA)44.48 The experimental crystal structure of ultrasmall gold nanocluster Au25(SR)18 was reported separately by Jin’s and Murray’s groups.52,53 Grönbeck and co-workers also predicted the same

Fig. 1 Schematic of the “size-focusing” process. Adapted with permission from ref. 2. Copyright 2010 American Chemical Society.
Au\(_{25}\)(SR)\(_{18}\) structure by DFT calculations. X-ray crystallographic analysis shows that the Au\(_{25}\) cluster has an Au\(_{13}\) icosahedral core (Fig. 3) surrounded by a monolayer containing 12 gold atoms and 18 thiolates (Fig. 3). The crystal structure reveals three types of gold atoms ((i) one central gold atom (ii) 12 gold atoms as the vertices of the icosahedron, and (iii) 12 gold atoms forming staple) and two types of SR groups (terminal SR groups which link to the gold core, and central SR groups which link to two staple gold atoms). The high symmetry structure shown above was obtained from the anion Au\(_{25}\)(SR)\(_{18}\), which is an 8 electron superatom with S\(^2\)P\(^6\) electronic structure, but a similar structure was found for neutral Au\(_{25}\)(SR)\(_{18}\). \(^{52}\)

Compared with Au\(_{25}\)(SR)\(_{18}\), the crystal structure of Au\(_{38}\)(SR)\(_{24}\) is slightly more complex. \(^{55}\) As illustrated in Fig. 4A, the Au\(_{38}\)(SR)\(_{24}\) cluster has a face-fused biicosahedral Au\(_{23}\) core covered by six dimeric (\(\text{SR}-\text{Au-SR-Au-SR}\)) and three monomeric staples (\(\text{SR}-\text{Au-SR}\)). A similar structure has been predicted by Zeng \(^{56}\) just slightly differing in the arrangement of the dimeric staples on the icosahedral Au\(_{13}\) unit. The monomeric staples were found for Au\(_{38}\)(SR)\(_{24}\) but not for Au\(_{25}\)(SR)\(_{18}\) which is just protected by six dimeric staples. This observation led to the prediction that the smaller size clusters need longer staples for the protection considering the large curvature for small gold core. \(^{57}\)

Similar to the case of Au\(_{102}\)(SR)\(_{44}\) and Au\(_{38}\)(SR)\(_{24}\), the structure of Au\(_{28}\)(TBBT)\(_{20}\) (where TBBT = 4-tert-butylbenzenethiolate) is also chiral (Fig. 4B). \(^{58,55,58}\) The cluster has an Au\(_{20}\) chiral kernel protected by four dimeric “staples” and eight bridging thiolates, and the chirality of the structure arises due to the rotational arrangement of the four dimeric staples and the arrangement of eight bridging thiolates. Later, Jin’s group

**Fig. 2** X-ray crystal structure of the Au\(_{102}\)(p-MBA)\(_{44}\) crystal. (A) Electron density map and atomic structure (Au atoms depicted as yellow spheres, and p-MBA shown as framework and with small spheres [S in cyan, C in gray, and O in red]). (B) Top view of the two enantiomeric clusters. Colour scheme as in (A), only the sulfur atoms of p-MBA are shown. Adapted with permission from ref. 48. Copyright 2007 American Association for the Advancement of Science.

**Fig. 3** Crystal structure of the Au\(_{25}\)(SR)\(_{18}\) cluster. (A) The icosahedral Au\(_{13}\) core; (B) the Au\(_{13}\) core plus the exterior 12 Au atoms; (C) whole Au\(_{25}\) cluster protected by 18 thiolate ligands (just S is shown for clarity, magenta, Au; yellow, S). Adapted with permission from ref. 52. Copyright 2008 American Chemical Society.

**Fig. 4** Crystal structures of (A) Au\(_{38}\)(PET)\(_{24}\), adapted with permission from ref. 55. Copyright 2010 American Chemical Society. (B) Au\(_{28}\)(3,4-DBTBT)\(_{20}\), adapted with permission from ref. 58. Copyright 2013 American Chemical Society. (C) Au\(_{28}\)(S-c-C\(_6\)H\(_{11}\)S)\(_{20}\), adapted with permission from ref. 59. Copyright 2016 American Chemical Society. (D) Au\(_{28}\)(PPH\(_3\)\(_{10}\)C\(_6\)H\(_7\)Ph)\(_{20}\), adapted with permission from ref. 60. Copyright 2012 American Chemical Society. (E) Au\(_{24}\)(SCH\(_2\)Ph-tBu)\(_{20}\), adapted with permission from ref. 61. Copyright 2014 Royal Society of Chemistry. (F) Au\(_{24}\)(SeC\(_6\)H\(_5\))\(_{20}\), adapted with permission from ref. 62. Copyright 2014 American Chemical Society. The ligands are partially omitted for clarity.
reported an isomer of Au_{124}(TBBT)_{20} obtained after the ligand exchange with excess SH—C_{6}H_{11} ligand. The obtained Au_{124}(S—C_{6}H_{11})_{20} has two trimeric staples arranged on the surface as shown in Fig. 4C.\textsuperscript{39} The above results revealed that, for a given number of gold atoms, the structure of a cluster may be modified by the ligand. This was also observed for Au_{114} clusters.\textsuperscript{60—62} Au_{24}L_{20} clusters have different structures depending on the ligands (L): (i) Mixed ligand shell composed of phosphine and thiolate (Fig. 4D),\textsuperscript{68} (ii) thiolate (Fig. 4E)\textsuperscript{68} and (iii) selenolate-capped Au_{24} clusters (Fig. 4F).\textsuperscript{68} This confirms that the structure of gold clusters is affected by the protecting ligand on the surface.

Apart from the clusters we mentioned above, the crystal structure of some other clusters have also been reported including Au_{18}(SR)_{14},\textsuperscript{63} Au_{30}(TBBT)_{16},\textsuperscript{64} Au_{32}(S—tBu)_{15},\textsuperscript{65} Au_{36}(SR)_{24},\textsuperscript{66} Au_{23}(SR)_{44},\textsuperscript{67} Au_{38}(p-MBT)_{50},\textsuperscript{68} and Au_{144}(SCH_{2}Ph)_{60}.\textsuperscript{69} Among those clusters, Au_{144}(SCH_{2}Ph)_{60} has attracted a lot of attention. The structure of Au_{144}(SCH_{2}Ph)_{60} was predicted by Olga Lopez-Acevedo and co-workers in 2009,\textsuperscript{70} but the experimental structure was published almost one decade later, in 2018, by Zhikun Wu and co-workers.\textsuperscript{69} The Au_{144}(SCH_{2}Ph)_{60} has a Au_{12}icosahedral inner core covered by two Au shells resulting in a Au_{114} core. 30 monomeric staple motifs are distributed on the surface of the Au_{114} core in a highly ordered pattern. Because of the arrangement of the 30 monomeric staples, the Au_{144}(SCH_{2}Ph)_{60} structure is chiral. With more and more crystal structures of gold clusters determined, the influence of size and structure on the HOMO–LUMO gap can be established. Atom packing plays an important role for \(E_g\), as mentioned above, and the FCC (face-centered cubic) structures have much higher \(E_g\) energies than the icosahedral ones of comparable sizes.\textsuperscript{77}

### 2.3 Flexibility of the gold–sulfur interface of gold nanoclusters

The formation of a gold–sulfur bond is the driving force for the anchoring of thiol ligands on gold surfaces, nanoparticles or clusters,\textsuperscript{71} leading to the formation of self-assembled monolayers (SAM).\textsuperscript{72} From the available structural investigations it emerges that the staple-type binding motif is the preferred structure element of the gold–sulfur interface, as first evidenced by the crystal structure determination of Au_{102}(SR)_{46}.\textsuperscript{48} supported by calculations.\textsuperscript{27} After analysis of various nanocluster structures such as Au_{23}(SR)_{18}, Au_{38}(SR)_{24} and Au_{36}(SR)_{20}, the bridging motif (Fig. 5A), monomeric staple (Fig. 5B) and dimeric staple motif (Fig. 5C) are the most common surface structure elements. Actually the tetrameric staple motif also has been evidenced for Au_{34}(SCH_{2}Ph—Bu)_{20}.\textsuperscript{41} Also other staple motifs are expected in order to adapt to the different curvatures of nanoclusters.

The properties of gold–sulfur interface and its dynamics determine the further functionalization of thiolate-protected clusters and their applications. However, due to the experimental difficulties in probing the dynamic properties of the interface, this field was barren for a long time. Since the structure of the Au_{38}(SR)_{24} cluster has been solved it became a good candidate for studying the gold–sulfur interface.\textsuperscript{71} The cluster is intrinsically chiral due to the chiral arrangement of the dimeric staples at the poles of the cluster. Dolamic and co-workers reported the separation of the enantiomers of Au_{38}(SCH_{2}CH_{2}Ph)_{24} (ref. 73) by chiral high-performance liquid chromatography. Using enantiopure Au_{38}(SR)_{24} clusters Knoppe and co-workers showed that the cluster can racemize (Fig. 6, top).\textsuperscript{73} The reaction was started with A-Au_{38}(SCH_{2}CH_{2}Ph)_{24} and C-Au_{38}(SCH_{2}CH_{2}Ph)_{24} (A: anticlockwise, C: clockwise), respectively, and the evolution of the clusters was followed by circular dichroism spectroscopy. From this investigation, the racemization of the enantiopure Au_{38}(SR)_{24} clusters takes place at modest temperatures without significant decomposition. Furthermore, from the temperature-dependent kinetic data, the activation barrier for the rearrangement within the thiolate-gold interface of the Au_{38}(SR)_{24} clusters is about 28 kcal mol\textsuperscript{–1}, which is lower than the gold–sulfur bond energy (about 50 kcal mol\textsuperscript{–1}). The relatively low activation barrier indicated that the racemization process of Au_{38}(SR)_{24} proceeds without complete Au–S bond

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**Fig. 5** Schematic drawing of the bridging thiolate (A), monomeric Au–SR—Au staple motif (B) and dimeric Au–SR—Au—SR—Au staple motif. Yellow: Au (0) atom; orange: Au(l) atom; green: sulphur atom.

**Fig. 6** Schematic drawing of the racemization reaction of Au_{38}(SR)_{24} cluster. (Top) In this scheme (view along the long axis of the cluster), the left-handed cluster (left, A-Au_{38}(SR)_{24}) is converted into the right-handed enantiomer (right, C-Au_{38}(SR)_{24}) and vice versa. Two enantiomers are separated by an activation barrier of ca. 28 kcal mol\textsuperscript{–1} as determined from experiments. Adapted with permission from ref. 74. Copyright 2012 American Chemical Society. (Bottom) Density functional theory computational simulation of the chiral inversion of Au_{38}(SR)_{24} clusters through rotation of the three gold core atoms at the poles of the cluster. Adapted with permission from ref. 75. Copyright 2019 American Chemical Society.
breaking and evidenced the flexibility of the gold-thiolate interface. In addition, using density functional theory (DFT) computations, Håkkinen and co-workers proposed a mechanism for this inversion of the Au–S framework of Au_{38}(SR)_{24} as shown in Fig. 6 (bottom). In their model, the racemization proceeds via a rotational reconstruction of the metal core without any Au–S bonds being broken.

The flexibility of the gold–sulfur interface is affected by many factors, such as Pd doping, which is normally used for increasing the stability of the cluster and enhance its reactivity in catalytic and ligand exchange reactions. Barrabés and co-workers reported that the racemization process takes place at significantly lower temperature after doping the Au_{38}(SR)_{24} cluster with two Pd atoms compared with the parent cluster. This case indicated that the Pd doping of the cluster renders the Au–S interface more flexible. In contrast, after introducing of a rigid dithiolate, 1,1’-binaphthyl-2,2’-dithiol (BINAS), into the ligand shell of the Au_{38}(SR)_{24} cluster, the racemization drastically slows down. For example, the racemization of Au_{38}(2-PET)_{24}(BINAS) at 70 °C is about 27 times slower compared to the parent cluster, meaning that the introduction of the dithiol reduces the flexibility of the Au–S interface. Interestingly, the vibrational spectrum of the Au–S interface is also drastically influenced by introducing one BINAS dithiol into the ligand shell of the cluster, possibly related to the fact that the dithiol bridges two staples. Similarly, the racemization of the Au_{40}(SR)_{24} cluster was reported to take place at higher temperature compared with Au_{38}(SR)_{24} but with a similar activation barrier, further confirming the flexibility of gold–sulfur interface.

The racemization described above is not the only dynamic process observed for the Au_{38}(SR)_{24} cluster. In fact, ligands can migrate on the cluster surface between different symmetry unique sites. It has been reported that HPLC can be used to separate and isolate one specific regioisomer after ligand exchange between Au_{38}(2-PET)_{24} clusters and enantiopure chiral [2.2]-paracyclophane-4-thiol 1 (PCP-4-SH). The isolated species is stable at room temperature, however, the adsorbed thiolate migrates between the different symmetry-unique sites at 80 °C as followed by HPLC. The mechanism underlying this observation is not yet clear: it might involve the exchange of two ligands between different sites on one cluster or the exchange of ligands between clusters (see below).

The chemical properties and flexibility of the gold–sulfur interface, as discussed above, form the basis for the post-synthesis modification and further functionalization of the gold nanoclusters. Among them, ligand exchange reaction is extraordinarily sparkly and we will now focus on this topic in the following sections.

3. Ligand exchange reactions

3.1 Mechanism of ligand exchange reactions

Depending on the flexibility and the reactivity of the gold–sulfur interface, the thiolates within the ligand shell can be displaced by other thiols in the solution, which is named place-exchange reactions, or more commonly ligand exchange reactions (LERs).
Table 1  Summary of the examples of ligand exchange reactions mentioned in each section. The order in the table follows the order of discussion in the text.

| Section | Topic              | Reaction                                                                 | Reference |
|---------|--------------------|--------------------------------------------------------------------------|-----------|
| 3.1     | Ligand exchange    | [Au25(SC12H25)18] + dihydrolipoic acid (DHLA) → Au25(SG)18 + Au25(S-Adm)38PA2 | 87        |
|         | with free ligand   | [Au25(SC12H25)18] + dihydrolipoic acid (DHLA) → Au25(SG)18 + Au25(S-Adm)38PA2 | 87        |
| 4.1     | Ligand exchange    | [Au25(SC12H25)18] + dihydrolipoic acid (DHLA) → Au25(SG)18 + Au25(S-Adm)38PA2 | 87        |
| 4.2     | Ligand exchange    | [Au25(SC12H25)18] + dihydrolipoic acid (DHLA) → Au25(SG)18 + Au25(S-Adm)38PA2 | 87        |
| 4.3     | Ligand exchange    | [Au25(SC12H25)18] + dihydrolipoic acid (DHLA) → Au25(SG)18 + Au25(S-Adm)38PA2 | 87        |
| 4.4     | Addition of        | [Au25(SC12H25)18] + dihydrolipoic acid (DHLA) → Au25(SG)18 + Au25(S-Adm)38PA2 | 87        |
|         | fluorescence       | [Au25(SC12H25)18] + dihydrolipoic acid (DHLA) → Au25(SG)18 + Au25(S-Adm)38PA2 | 87        |

Independent of the nature of the thiolate ligands and the presence of the protein. But the presence of positive residues at the protein C-terminal tail is critical for forming attractive intermolecular interactions with the carboxylate groups of the SG ligands, facilitating the adsorption of the protein cysteine on the gold cluster surface.

Because of the fast dynamics and complexity of the ligand exchange it is difficult to disclose the microscopic details of the S<sub>N</sub>2-like process by experimental alone. However, with the assistance of computational method such as DFT calculations, some light could be shed on the S<sub>N</sub>2-like mechanism of initial ligand exchange. Interestingly, recently Wu and co-workers proposed an unimolecular nucleophilic substitution (S<sub>N</sub>1)-like mechanism (in addition to a S<sub>N</sub>2-like process), based on single crystal structure analysis of ligand exchanged clusters. The crystal structures showed that part of the sulfur atoms retained their configuration upon ligand exchange, which seems not possible for a S<sub>N</sub>2-like process. This argument only holds if the configuration at the sulfur atom is retained in solution, which was not addressed.
Apart from the possibility for exchange of thiols via free ligand, as discussed above, there is a second mechanism for thiol exchange, which takes place through intercluster collisions. In 2013, Yoshiki Niihori and co-workers observed the intercluster LERs between Au_{25}(SC_{10}H_{21})_{18} and Au_{25}(SC_{12}H_{25})_{18} when they studied the influence of Pd atom doping of thiolate-protected Au_{25} nanoclusters on ligand exchange reactivity. In that case the authors assumed that this exchange resulted from the detachment of ligand or gold–ligand species from the cluster.

Later, Salassa and co-workers carried out the intercluster LERs between Au_{25}(SBut)_{18} and Au_{25}(2-PET)_{18}, and experimentally showed that the process is fast. As shown in Fig. 9A, 15 min after mixing the two clusters at room temperature peaks from clusters with mixed ligand layer already appear in the mass spectrum. In addition, the interclusters ligand exchange seems to take place without release of thiol or thiol–gold complex into the solution, as illustrated in Fig. 9B. In this experiment Au_{25}(SBut)_{18} was put inside a dialysis membrane and thus the two clusters were physically separated, whereas low molecular weight species could still penetrate and pass the membrane. The MALDI mass spectra showed that the clusters did not undergo any exchange of thiols.

Recently, we confirmed that bidentate thiol ligands can also exchange between clusters by observing the mass peak belonging to Au_{25}(2-PET)_{16}(R-BINAS)_{x} (BINAS = 1,1'-binaphthyl-2,2'-dithiol) after mixing Au_{138}(2-PET)_{24-x}(R-BINAS)_{x} (x corresponds to the number of BINAS ligands, which was about 2 on average in this experiment) and Au_{138}(2-PET)_{18} clusters. Compared to the exchange of monothiols higher temperatures are needed for the process to take place, 70 °C in this case.
However, compared with the free thiol exchange, the mechanism of intercluster ligand exchange is still rather unexplored and further investigations are needed.

Over the past decades, since the first report of ligand exchange reactions for gold nanoclusters, this reaction has been widely used to introduce new ligands to the parent nanoclusters,77,88,90–92 and to add functionality or chemical properties to clusters as a post-modification method.85,94 This methodology strongly extends the possibilities to modify surface properties of gold nanoclusters, normally without changing the metal core.94 Understanding the mechanism of the ligand exchange reaction may help us engineer the surface chemistry of nanomaterials in order to build multifunctional nano-platforms.

3.2 Site-selective ligand exchange reactions

Several reports show that during the ligand exchange reaction, many sites on the ligand shell exchange slowly or not at all, whereas others are comparatively reactive.85,94,95 These observations revealed that the ligand shell is heterogeneous and offers a diversity of ligand binding sites and the exchange reaction occurs preferentially at selected ones.95 Before going into detail, it is essential to discuss the diversity sulfur groups in the ligand shell which represent the reactive site.

Since the dynamic exchange process were mostly investigated with Au25(SR)18 and Au38(SR)24,88,89 which are easily prepared and relatively stable, we focus some more on their crystal structure as illustrated in Fig. 10. The Au25(SR)18 cluster (Fig. 10 left structure) has an icosahedral Au13 core surrounded by six SR-Au-SR-Au-SR staple units as we mentioned before.77,89,92,97–99 Because of the symmetry of Au25(SR)18, six dimeric staples share the same chemical environment, but the SR groups in one staple can be divided into central –SR group, which is bound to the two gold atoms in the staple, and terminal –SR groups, which are linked to the gold core.100 In contrast, Au38(SR)24 consists of a bicuboctahedral Au23 core (Fig. 10 right structure) and is covered not only by six dimeric units but also by three monomeric units. The structure of Au130(SR)4 is elongated, with the three monomeric staples at the equator and the six dimeric staples at the two poles.101 The dimeric staples can be divided into central –SR group and terminal –SR groups. However, the two terminal –SR groups in the dimeric staples of Au130(SR)4 have different chemical environment.88,102 Overall the 24 thiolate ligands are divided into four groups of symmetry unique sites with different chemical environment, which can give rise to preferential exchange sites during LERs.88,92

From recent experimental and computational studies, it became clear that LERs between clusters and free monothiolates in solution start preferentially at the terminal SR groups of Au25(SR)18 and Au38(SR)24 clusters,92,100,102 via an associative SP2-like mechanism.98 This tendency was experimentally demonstrated by Yoshiki Niihori and co-workers.99 In this case the isomer distributions of the product after ligand exchange between Au25(Pd(SC2H4Ph)18 and C12H25SH were determined by high resolution high-performance liquid chromatography. The quantitative evaluation of the expected coordination isomers and the products obtained by the reactions, showed that the exchange reaction starts at the thiolate which is bound to the core site. This also holds for exchange with other chalcogenate ligands such as HSePh or HTePh.99 Niihori and co-workers also found that Pd doping of the Au25(SR)18 cluster drastically increased the ligand exchange rate. Pd doping reduces the number of valance electrons of the metal core, which facilitates the attack by the incoming ligand. Furthermore Pd doping induces the distortion of the cluster geometry.99

In addition, the site preference of LERs can be investigated by computational methods. Aikens and co-workers employed density functional theory (DFT) to examine the ligand exchange on model Au25(SH)18 and Au38(SH)24 clusters with an incoming methanethiol.100,102 They calculated the intermediates and transition states, and predicted the barrier heights and reaction energies for this ligand exchange process. In Fig. 10, the different possible ligand substitution sites are marked on the crystal structure of Au25(SR)18 (Fig. 10 left) and Au38(SR)24 (Fig. 10 right) nanoclusters. The former cluster offers three different sites, whereas the latter has eight. The energies of the intermediates, transition state and products corresponding to ligand exchange at different sites determined by DFT calculation are summarized in Table 2. The results for Au25 indicated that the most favourable ligand exchange process is at site B, and the site C has highest energy barrier, which resulted in ligand exchange at the central SR group. This tendency was consistent with the one for Au38(SR)24 as is evident from the calculated energies listed at Table 2. LERs at site C had higher energy barriers compared with other sites, meaning that the reaction between the staple gold atom and the sulfur atom of the central-SR units will proceed at a slower rate. Here, we should mention that each sulfur atom is participating in two bonds, for example Au25 A&B, Au38 C&D, and Au38 E&F, which represent different reactive sites, however, those two sites will lead to the same product. Above predictions from computational studies on Au25(SH)18 and Au38(SR)24 indicated that the
exchange is preferred at the terminal SR groups, which is congruous with the experimental results.\(^{88,92}\) However, this might not always be the case as shown by Zhu and co-workers. After the ligand exchange reaction between Au\(_{25}\)Cd(SCH\(_2\)Ph)\(_{18}\) and SCH\(_2\)Ph-Bu Au\(_{25}\)Cd(SCH\(_2\)Ph-Bu)\(_2\)S(SCH\(_2\)Ph)\(_3\) has been obtained.\(^{90}\) The integrated area of the peaks extracted from \(^1\)H NMR spectra of this species illustrated that the remaining parent ligand SCH\(_2\)Ph was equally distributed among the terminal and central sulfur atoms, in contrast with the results discussed above. This result may be caused by several reasons, such as steric hindrance within the ligand shell or the inevitable migration of the ligand on the cluster surface at 80 °C. In addition, Pengo and co-workers also showed by NMR spectroscopy that the distribution of incoming ligand on the central and terminal sites depends on the properties of the ligand.\(^{98}\)

Apart from the monothiolate ligands, nanoclusters are also reactive towards dithiol ligands. Amala Dass and co-workers systematic studied LERs with aliphatic dithiol ligands of various chain length, HS-(CH\(_2\))^\(_n\)-SH.\(^{104}\) They documented that C\(_1\) and C\(_2\) prefer interstaple coupling, and C\(_3\) and C\(_4\) are good candidates for intrastaple binding, whereas the length of C\(_2\) ligand is not enough for the bidentate binding. Aromatic dithiols have also been used for the LERs. However, they are sterically more demanding and more rigid compared to aliphatic thiols. One example is 1,1′-binaphthalene-2,2′-dithiol (BINAS), which is a chiral rigid dithiol ligand (Fig. 11 left), used to introduce chirality to achiral clusters.\(^{77,105}\) Several studies provide insight into the LERs on BINAS with Au\(_{25}\) and Au\(_{18}\) clusters,\(^{77,103,105-108}\) and from the experimental and computational results it emerges that the bidentate ligand connects two neighboring staples by interstaple coupling.\(^{103,105,108}\) Sels and co-workers reported the isolation of different exchange products and isomers of clusters containing one or two BINAS adsorbed PdAu\(_{24}\)(2-PET)\(_{18}\).\(^{106}\) The investigated structure of the PdAu\(_{24}\)(SR)\(_{18}\)(S-BINAS)\(_{1}\) cluster is shown in Fig. 11, which indicates the two different sulfur environments of the cluster and marks some bond distances.\(^{103}\) As displayed in Fig. 11, there are several possible binding sites for BINAS at the ligand shell, illustrated with curved lines. However, considering the distances between the two S-atoms in undistorted BINAS, 4.1 Å, the interstaple binding site between a S\(_{\text{apex}}\) and a S\(_{\text{core}}\) sulfur atom has the appropriate distance, 4.05 Å. Based on X-ray absorption spectroscopy (XAS), the authors also showed that the BINAS interstaple binding mode does not perturb the cluster structure.\(^{103}\)

Compared with the monothiolate ligand exchange it may be easier to isolate isomers of clusters with mixed ligand shell when using dithiolate ligands. A better understanding of the preferred reactive sites of the LERs will enable the rational design of mixed ligand shell clusters.

### 3.3 Comparison of ligand exchange reactions of clusters in solution and on supports

LERs is a process that takes advantage of the dynamic nature of the thiol-gold surface,\(^{77}\) and in most cases it is performed with clusters in solution. As a further development, Truttmann and co-workers recently reported the first successful ligand exchange reaction on supported (immobilized) Au\(_{11}\) nanoclusters.\(^{109}\) In this case, Au\(_{11}\)(PPh\(_3\))^\(_3\)-Cl\(_3\) clusters were deposited on Al\(_2\)O\(_3\), and then the dropcast sample was exposed to a solution of thiol ligand. In such a procedure cluster–ligand combinations are possible that are not accessible through normal LERs due to solvent incompatibility. Indeed, in the study mentioned above the hydrophilic thiol ligand, 1-gluta-thione, and the hydrophobic 2-PET (2-phenylethanethiol) ligand were used. According to previous reports, when performing ligand exchange with Au\(_{11}\)(PPh\(_3\))^\(_3\)-Cl\(_3\) and GSH in solution, the Au\(_{11}\) clusters grow to form Au\(_{25}\).\(^{106,110}\) Here, in contrast, no cluster growth has been observed as indicated from MALDI-MS after ligand exchange reaction on supported clusters. As evidenced by IR (PM-IRRAS, ATR-IR) and NMR spectroscopy, partial ligand exchange took place and the halide ligands of the parent nanocluster were primarily exchanged with thiolates.

In contrast to the ligand exchange in solution, the reaction on the supported cluster did not induce a change of the core size of the nanoclusters. However, due to the hindered
accessibility of the ligand on the support side of the cluster, the number of exchanged ligands was relatively low compared to typical solution phase ligand exchange reactions. In addition, similar to the functionalization in solution, fluorescence could be introduced to supported clusters by reacting them with fluorescein probe, which also provides new insight for use of the clusters in biological applications.  

4. Functionalization of gold nanoclusters induced by LERs

4.1 Chirality induced by LERs

Chirality is a geometric property of objects, which widely exists in nature from molecules over proteins to even larger structures. During the past decade, with the use of X-ray single-crystal diffraction, it has emerged that chirality is a ubiquitous property for gold nanoclusters. Because of the potential applications in sensing, catalysis, molecular recognition and so on, chiral gold nanoclusters have attracted a lot of interest. The reported chiral gold nanoclusters can be categorized into three types: (i) chiral Au–S framework with achiral ligands, (ii) achiral Au–S framework with chiral ligands, and (iii) a combination of the two (chiral Au–S framework and chiral ligand). The type I chiral nanoclusters are sometimes also called intrinsically chiral nanoclusters, and some examples are Au25(SR)18, Au28(SR)20, Au38(SR)24, Au102(SR)44 and Au133(SR)52, in which all the different R groups are achiral. The chirality of type II nanoclusters is due to the ligands SR. Fundamentally, such chiral clusters can be prepared by direct synthesis using a chiral thiol. However, resulting from the solubility and steric effect of different ligands, direct synthesis of nanoclusters with some ligands was unsuccessful. Alternatively, ligand exchange is a good method to incorporate chirality or other functionality onto the gold nanoclusters.

By using ligand exchange on achiral Au25(SR)18 with chiral thiols (SR*), a series of chiral Au25(SR)18-S(R*)x clusters with chiroptical activity have become available. For example, Si and co-workers reported the ligand exchange on [Au25(PET)18][TOA] in the following named as Au25BINAS. As illustrated in Fig. 12, the CD (left panel) and UV-Vis absorption (right panel) spectra of Au25 clusters were recorded before (a) and after exchange with R/S-BINAS (b/c). As expected, the Au25 cluster is optically inactive before ligand exchange. Once the cluster exchange with BINAS, intense bands at 305, 352, and 390 nm are observed in the CD spectra with opposite sign for clusters covered by the two enantiomers of BINAS. The CD signals refer to the chiral ligand but the spectra are different compared with the one of free BINAS. Furthermore, Fig. 12 reveals that the absorption spectrum of the cluster became less defined after exchange with BINAS. Apart from Au25 clusters, similar exchange with R/S-BINAS also takes place on Au133 and Au40 clusters. In addition, as we mentioned before, the introduction of BINAS ligand on Au38 clusters stabilized the structure against inversion (enantiomerization), probably due to the reduced flexibility of the gold–sulfur interface. Also, different from the LERs with monothiols, exchange with dithiol ligand reduces the number of isomers. Due to the reduction of the reaction rate after the first exchange, it is also relatively easy to obtain species with just one ligand exchanged.

Compared with the direct synthesis, the ligand exchange of nanoclusters may lead to a heterogeneous system with a distribution of the number of exchanged ligands when the exchange is incomplete. In order to obtain completely exchanged nanoclusters, large excess of free incoming ligand with respect to the clusters and/or repetition of the exchange reaction are necessary, with purification of the intermediate cluster after each step.

The optical activity is usually considerably stronger for the type I chiral clusters, i.e., clusters with intrinsic chirality in their Au–S framework, compared to type II chiral clusters, i.e., achiral Au–S framework with chiral ligands. The absorption spectrum of a given cluster is roughly ligand independent. In contrast,
Au$_{25}$ (SR) shows circular dichroism (CD) spectra of Au$_{25}$ and Au$_{38}$ clusters stabilized with three different chiral R* groups. All these chiral Au$_{25}$ (SR*)$_{18}$ nanoclusters reveal the typical UV/Vis absorption spectrum of Au$_{25}$, but their CD signals show differences depending on the R* groups (Fig. 13 left panel). A similar phenomenon was also observed for Au$_{38}$ (SR*)$_{24}$ nanoclusters with different chiral thiolate ligands (Fig. 13 right panel). This ligand-dependence provides a possibility to engineer the optical characteristics of a given gold nanocluster.

In addition, the chiral ligand R-BINAS was also used to amplify the enantiomeric excess of Au$_{38}$ (SR*)$_{24}$ at 70 °C. In a mixture containing the two enantiomers of the Au$_{38}$ cluster and the corresponding clusters containing one R-BINAS in their ligand shell (diastereomers), the fraction of anticlockwise (A) clusters increases with time at the expense of clockwise (C) clusters (Fig. 14B). For the experiment shown in Fig. 14B the initial sample was prepared with high fraction of C-Au$_{38}$. The sample was then heated to 70 °C and the HPL chromatograms were recorded as a function of time. The fractions of different species were determined from the HPL chromatograph and plotted as function as time (Fig. 14B), showing the increase of A-Au$_{38}$. The proposed mechanism of the process is shown in Fig. 14A. This dynamic inversion is due to the diastereoselective intercluster exchange of R-BINAS between chiral clusters and the fast racemization of Au$_{38}$ (2-PET)$_{24}$. This example shows that the dynamic nature of these clusters can be used as a benefit. Overall, the ligand exchange reaction with chiral ligands will continue to add new impetus to the field of chiral nanomaterials and its applications.

4.2 LERs induced size transformation of clusters

LERs is a process that takes advantage of the dynamic nature of the thiol-gold interface, and in most cases it does not cause any change in size or structure of the cluster, only replacing one capping ligand by another one. During the past decade, due to the development of precise synthesis methods and structure determination by X-ray single-crystal diffraction numerous new nanoclusters were reported. LERs became an important new methodology for controlling the size and structure of nanoclusters, and the process was named ligand-exchange-induced size/structure transformation (LEIST for short), which was first proposed by the Jin’s group.

Up to now, most of the reported LEIST work focuses on PPh$_3$-stabilized and thiolate-protected Au nanoclusters. However, the driving forces for LEIST is quite different for these two kinds of Au nanoclusters. First, for LEIST of PPh$_3$-stabilized Au nanoclusters, exchanging the phosphine ligands by thiolate ligands changes the surface bonding completely (from Au–P to Au–S). The different coordination modes, Au–P vs. Au–S, lead to different surface motifs and drives the structural transformation of the cluster. Second, for LEIST of thiolate-protected Au nanoclusters, the new thiolates would not significantly alter the gold–ligand coordination; however, due to the distinctly different physical–chemical properties (e.g., size, rigidity, bulkiness, interactions etc.) of different thiolate ligands, the original structures may not be the most stable for the ligand-exchanged nanoclusters, and thus the structural transformation takes place. In the following we mainly focus on the LEIST on thiolate-protected Au nanoclusters. Depending on the size or structural change of the thiolate-protected Au nanoclusters resulting from the LEIST, this process has been classified into three groups.

(1) Transformation between structural isomers without size change. For example, Au$_{28}$ (SR)$_{20}$ nanocluster reversibly changes its structure upon ligand exchange between R = c-C$_6$H$_{11}$ and R = Ph$_2$Bu at elevated temperatures (e.g., 80 °C) as shown in Fig. 15. The structures of these two nanoclusters are remarkable different. Au$_{28}$ (SPhBu)$_{20}$ contains a FCC Au$_{20}$ kernel capped by four Au$_4$(SR)$_3$ staple motifs and eight bridging thiolates, whereas Au$_{28}$ (S–c-C$_6$H$_{11}$)$_{20}$ adopts a more lose structure with a FCC Au$_{20}$ kernel plus two Au$_4$(SR)$_3$, two Au$_4$(SR)$_4$, and eight bridging SR staple-like structures. DFT calculations revealed that the origin of reversible isomerization lay in the thiolate ligand’s carbon tail structure, which was found to dictate the
isomer's stability.\textsuperscript{29} In LEIST the isomerization of clusters is rare.

(II) Transformations from a smaller to a larger nanocluster.\textsuperscript{26,28,42,119,125–127} An interesting example is the transformation of Au\textsubscript{25}(PET)\textsubscript{18} into Au\textsubscript{28}(TBBT)\textsubscript{20}.\textsuperscript{28} Au\textsubscript{25}(PET)\textsubscript{18} is probably the most studied thiolate-protected nanocluster, due to its prototypical character. At 80 °C and in large excess of TBBT the Au\textsubscript{25}(PET)\textsubscript{18} cluster is transformed into Au\textsubscript{28}(TBBT)\textsubscript{20}. It is worth to mention that the resulting Au\textsubscript{28}(TBBT)\textsubscript{20} cluster is chiral, and the origin of chirality is primarily rooted in the rotating arrangement of the four dimeric staples as well as the arrangement of the bridging thiolates as we mentioned before. Moreover, the pair of enantiomers of Au\textsubscript{28}(TBBT)\textsubscript{20} can be separated by chiral-HPLC.\textsuperscript{28} Another example is the transformation of Au\textsubscript{11} to Au\textsubscript{25} upon ligand exchange with GSH.\textsuperscript{26,110}

(III) Transformations from a larger to a smaller nanocluster.\textsuperscript{27,28,120,123–125,128,129} For instance, highly stable Au\textsubscript{114}(SC\textsubscript{2}H\textsubscript{4}Ph)\textsubscript{60} reacted with thiophenol, HSPH, to form a different 99 atom cluster species Au\textsubscript{99}(SPh\textsubscript{X})\textsubscript{42}.\textsuperscript{129} In addition, Zeng and co-worker reported that, by LEIST, the very stable and widely investigated Au\textsubscript{44}(SC\textsubscript{2}H\textsubscript{4}Ph)\textsubscript{24} cluster was transformed to Au\textsubscript{44}(SPh\textsubscript{X}Bu)\textsubscript{24}, as shown in Fig. 16A.\textsuperscript{129} To unravel details of the intriguing one-size-to-another size transformation, Zeng carried out time-dependent mass spectrometry and optical spectroscopy analyses (Fig. 16B) and found a remarkable disproportionation in the transformation of rod-like biicosahedral Au\textsubscript{36}(SC\textsubscript{2}H\textsubscript{4}Ph)\textsubscript{24} to tetrahedral Au\textsubscript{36}(SPh\textsubscript{X}Bu)\textsubscript{24}. From the evolution of the mass spectra and corresponding UV curves, the reaction pathway can be roughly divided into four stages (Fig. 16C): (i) (0–5 min) ligand exchange reaction occurs between PET and TBBT; (ii) (10–15 min) ligand exchange reaction continues leading to TBBT-triggered structural distortion of Au\textsubscript{36} with an optical feature at 550 nm (Fig. 16D); (iii) (20–60 min) critical stage for the disproportionation of Au\textsubscript{36} to Au\textsubscript{38} and Au\textsubscript{40}, and (iv) size conversion of Au\textsubscript{40} to Au\textsubscript{36} evidenced by the decrease and complete disappearance of the Au\textsubscript{40} peak in time-dependent mass spectra. Recently, other LEIST, such as transformations from Au\textsubscript{36}(S–C\textsubscript{6}H\textsubscript{11})\textsubscript{25} to Au\textsubscript{36}(TBBT)\textsubscript{24} and from Au\textsubscript{326}(PET)\textsubscript{8} to Au\textsubscript{278}(SPh\textsubscript{X}Bu)\textsubscript{8} also revealed similar pathways but without disproportionation. In these cases the conversion through ligand exchange followed by the size focusing ultimately lead to size growth.\textsuperscript{127,129}

Apart from the irreversible transformation mentioned above, Amala Dass and co-workers presented the first reversible interconversion between two nanomolecules Au\textsubscript{36}(SPhX)\textsubscript{24}, (where X = –H or –Bu) and Au\textsubscript{34}(S–Bu)\textsubscript{24} as illustrated in Fig. 17.\textsuperscript{128,129} In this case, the gold core converted between bicuboctahedral Au\textsubscript{36} and 4-fused cuboctahedron Au\textsubscript{34} and the staple arrangement of these two cluster surfacers was also different. More interesting, the reversible conversion easily takes place under the same thermochemical conditions with different thiol ligands. Later, Wu and co-workers achieved the interconversion among Au\textsubscript{44}(TBBT)\textsubscript{28}, Au\textsubscript{44}(2,4-DMBT)\textsubscript{28}, and Au\textsubscript{43}(S–C\textsubscript{6}H\textsubscript{11})\textsubscript{25} nanoclusters by the ligand exchange,\textsuperscript{130,131} and they also investigated the thermostability of these nanoclusters. As monitored by time-dependent optical absorption, the Au\textsubscript{44}(2,4-DMBT)\textsubscript{28} cluster was less thermostable than Au\textsubscript{44}(TBBT)\textsubscript{28}, and much more stable than Au\textsubscript{44}(S–C\textsubscript{6}H\textsubscript{11})\textsubscript{25} at 80 °C.\textsuperscript{130} The interconversion not only offer many possibilities to obtain new nanoclusters but also leads to valuable insight into the inherent influence of the ligand on the composition and atomic structure of thiolate-protected gold clusters.

Focussing on the experimental conditions of the LEIST, mentioned above, it seems that the conditions necessary for LEIST to take place are large excess of incoming ligand, normally more than 100 times compared to the endogenous ligand, and elevated temperatures.\textsuperscript{134} In addition, it was argued that the incoming thiol must be significantly different from the endogenous thiol for the transformation to happen.\textsuperscript{61} However, we reported a new transformation of Au\textsubscript{32}(SR)\textsubscript{14} into Au\textsubscript{28}(SR)\textsubscript{21} by LER with a chiral ligand (R- or S-phenylpropane-1-thiol) at mild conditions, \textit{i.e.} at room temperature and with low thiol excess (with incoming to outgoing ligand molar ratio 2 : 1).\textsuperscript{135} In this case, the van der Waals interactions within the ligand shell and...
the “bulkiness” of the methyl are the main factors driving this process. In summary, LEIST provides new and exciting avenues to explore novel atom-precise nanoclusters and opens a new strategy to investigate the size growth of nanoclusters.

4.3 LERs induced organic/aqueous phase transfer of clusters

There are many types of ligands that can be used to stabilize nanoparticles or nanoclusters, as shown in Fig. 18. Depending on the solubility of the surface ligand, one can distinguish hydrophobic and hydrophilic clusters. However, for some (biological) applications, the hydrophobic clusters need to be dissolved in aqueous solutions. Also hydrophilic clusters sometimes need to made accessible to reactions in organic solvent. In this situation, phase transfer will be the first option to relocate the clusters to the desired phase.

In general, for phase transfer processes, reagents such as tetaoctylammonium bromide, tetramethylammonium bromide, and others have been successfully used. These chemical reagents form an additional molecular layer on the clusters and thus change their surface properties. This strategy was mainly used for the phase transfer from aqueous to organic phase. In addition to this strategy, ligand exchange has also been utilized as an important strategy for the phase transfer. For instance, when hydrophobic Au11(PPh3)7Cl3 reacted with GSH ligand in solution, the Au11 clusters grew to form water-soluble Au25 nanoclusters, which is a good evidence for ligand exchange. In addition, Zheng and co-workers reported the crystal structure of intermetallic nanocluster Au24Ag20(2-SPy)4(PA)20Cl2 and also investigated the phase transfer process after performing ligand exchange with mercaptosuccinic acid (MSA), leading to transfer of the Au24Ag20 cluster from DCM to water phase. Other reported cases of phase transfer induced by ligand exchange reactions, include ligands such as (phenylacetylene) PAH and Tiopronin. The latter ligand also introduced cancer therapy functions to the cluster.

It emerges from the cases mentioned above, that the phase transfer from aqueous to organic phase relies on phase transfer reagents or ligand modification, and in many cases the core size of the cluster is maintained. However, the inverse process, transfer from organic to aqueous phase, mostly relies on the ligand exchange reactions. Furthermore, the phosphine-stabilized clusters are more prone to be transferred to another phase after reaction with thiol ligands. For the choice of the ligand used for the phase transfer, three considerations have to be made: (i) the affinity to the metal core of new ligand compared with the original one; (ii) the solubility of the new ligand in the target solvent; and (iii) the capability of the ligand to maintain the core size of the cluster after phase transfer.

4.4 Florence induced by LERs

Fluorescent gold nanoclusters have been widely used for biological applications such as cell identification, to study interaction, differentiation, and tracking. Compared with the
more traditional QDs, fluorescent gold nanoclusters have decent quantum yield, excellent biocompatibility, good photostability, and lower cytotoxicity. The clusters had negligible influence on the cell viability at the considered dose. However, because of the synthesis method and the solubility of ligands, the diversity of the fluorescent gold nanoclusters has been limited.

LER is a very efficient strategy for preparing fluorescent gold nanoclusters. As one of the most intuitive cases, the non-fluorescent gold nanoclusters (AuNC@DDAB) can be converted to brightly red emitting nanoclusters (AuNC@DHLA) through an elegant ligand exchange reaction with dihydrolipoic acid (DHLA) as shown in Fig. 19. In this system, the DDAB-stabilized gold nanoparticles (AuNP@DDAB) are etched by the addition of Au precursors (HAuCl₄ or AuCl₃) to smaller nanoclusters (AuNC@DDAB), and the organic soluble and hydrophobic AuNC@DDAB become water soluble upon ligand exchange with dihydrolipoic acid (AuNC@DHLA). The AuNP@DDAB solution shows red color due to surface plasmon absorption, which is absent to AuNC@DDAB and AuNC@DHLA. More interesting, after ligand exchange with DHLA, the AuNC@DHLA solution shows strong red photoluminescence as shown in the Fig. 19. Here, the phase transfer and addition of fluorophore has been achieved together by the LERs.

In addition, Xu et al. reported ligand exchange by GSH and N-acetylcysteine (NAC) on AuNAC@Ag. The products showed maximum 20-fold fluorescence enhancement. During this ligand exchange process, silver ions and GSH have synergistic effects and the PL enhancement was found to be proportional to the concentration of GSH. This fluorescence enhancement was also used for selective imaging of intracellular glutathione. Besides dissolved clusters, the fluorescence can also be introduced to supported nanoclusters by using ligand exchange on the immobilized sample, as we mentioned above.

5. Conclusions

In conclusion, ligand exchange reaction plays an important role in the field of nanoclusters. As one of the post-synthesis methods for the modification and functionalization of the nanoclusters, the process is enabled by the flexibility of the gold–sulfur interface. Experimental and computational investigations show that LERs between clusters and free monothiols in solution starts preferably at the terminal SR groups of the staple motifs, which are linked to the gold core, via an associative S_N2-like mechanism. The process takes place in solution phase but also on immobilized Au nanoclusters. Understanding of the preferred reactive sites of the process will help us engineer precise clusters with mixed ligand shell. Compared with the monothiolate ligand exchange, diathiolate ligands may offer an easier way to obtain precise clusters with mixed ligand shell. Intercluster ligand exchange is also an important property of thiolate-protected metal clusters and has to be considered whenever clusters with different ligands are in the same solution. Despite the considerable effort made, the microscopic details of intercluster ligand exchange remain obscure.

Ligand exchange reactions are in general easy to perform and offer a great potential to introduce or amplify properties of the clusters, such as introduction of chirality to achiral nanoclusters and the amplification of optical activity and enantioselective, size transformation of the cluster, phase transfer, and the addition of fluorescent groups. Thus, ligand exchange reactions will continue to play an important role in the future and a better understanding of the process will further increase the potential of this method.

Conflicts of interest

There are no conflicts to declare.

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