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Elucidation of the electronic structures of thiolateprotected gold nanoclusters by electrochemical measurements

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Metal nanoclusters (NCs) with sizes of approximately 2 nm or less have different physical/chemical properties from those of the bulk metals owing to quantum size effects. Metal NCs, which can be size-controlled and heterometal doped at atomic accuracy, are expected to be the next generation of important materials, and new metal NCs are reported regularly. However, compared with conventional materials such as metal complexes and relatively large metal nanoparticles (>2 nm), these metal NCs are still underdeveloped in terms of evaluation and establishment of application methods. Electrochemical measurements are one of the most widely used methods for synthesis, application, and characterisation of metal NCs. This review summarizes the basic knowledge of the electrochemistry and experimental techniques, and provides examples of the reported electronic states of thiolate-protected gold NCs elucidated by electrochemical approaches. It is expected that this review will provide useful information for researchers starting to study metal NCs.

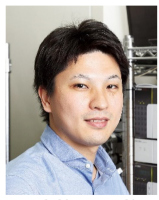
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Introduction

Metal nanoclusters (NCs) are a promising and widely studied group of next-generation nanomaterials. 1-7 Although there is no strict definition of a NC, when the size of a metal particle is below ~2 nm, it is often called a metal NC. Metal NCs are known to have different electronic/geometric structures from both the corresponding bulk metals and metal nanoparticles



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(NPs), and they thus have different physical properties and functions. Furthermore, because such properties and functions are dependent on the number of constituent atoms, creation of multiple functions with a single element is possible if the number of constituent atoms can be controlled in the size region of metal NCs. If multiple elements are used, 8,9 an even greater variety of functions can be created. Among these metal NCs, gold (Au) NCs protected by thiolate (SR) groups $(Au_n(SR)_m$, where n is the number of constituent Au atoms and m is the number of SR ligands) have been the most studied. This is because of their simple and easy synthesis under atmospheric conditions and their relatively high chemical and thermal stability. Furthermore, many $Au_n(SR)_m$ NCs can be analysed by single-crystal X-ray diffraction (SCXRD) to reveal their geometric structure and by density functional theory (DFT) calculations to reveal their electronic structure and origin of constructed orbitals. 10,11 Therefore, $Au_n(SR)_m$ NCs have attracted widespread attention as novel nanomaterials in environmental, energy, and biological fields. 12-17

Electrochemical methods play an important role in investigation of metal NCs with respect to elucidation of their electronic structures (Fig. 1). $^{18-20}$ In this review, we summarise previous reports using electrochemical approaches to study the electronic structures of $Au_n(SR)_m$ NCs. In section 2, we provide a brief history of synthesis of $Au_n(SR)_m$ NCs. In section 3, we discuss the principles of electrochemical measurement techniques and provide reported examples of the electronic structures of $Au_n(SR)_m$ NCs determined by electrochemical methods. In section 4, we summarise previous work and discuss the future outlook of this field. Studies on electrochemical catalysis $^{21-24}$ and synthesis of Au NCs $^{9,25-27}$ were excluded from the present review because they have been described in detail elsewhere.

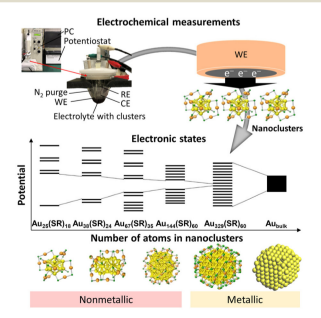


Fig. 1 Schematic of electrochemical measurements for elucidation of the electronic structure of metal clusters.

2 Thiolate-protected gold nanoclusters

The characteristics of the electronic structures of metal NCs are between those of metal complexes and metal NPs. Consequently, they have many similarities in their synthetic methods. As in general synthesis of metal NPs, metal NCs are obtained by mixing a solution containing organic ligands with metal salts as precursors with addition of a reducing agent. Such liquid-phase synthesis is called a bottom-up method. Since the late 1960s, efforts have been made to synthesise metal NCs consisting of Au, platinum (Pt), and palladium (Pd) with phosphine, halogen (X), and carbon monoxide (CO) as ligands.²⁸⁻³² Moreover, metal NCs comprising SR-protected Au, silver (Ag), copper (Cu), Pt, and Pd have also been reported. $Au_n(SR)_m$ NCs are extremely stable in air owing to the strong bonding between Au and sulfur (S). 33,34 Therefore, precise separation and synthesis of a single chemical composition of these $\operatorname{Au}_n(\operatorname{SR})_m$ NCs was attempted relatively early. ^{35,36} In general, $Au_n(SR)_m$ NCs are synthesised by (i) adding a metal salt (e.g., gold chloride) to a solvent, (ii) adding a thiol as a ligand, and (iii) mixing with a reducing agent (e.g., sodium borohydride). The byproducts and remaining SR complex are then washed off with solvent to obtain $Au_n(SR)_m$ NCs with a uniform number of constituent atoms. In addition to SR, a number of syntheses using phosphine, N-heterocyclic carbene, and alkynyl ligands have also been reported. 37,38 Moreover, some Au atoms in $Au_n(SR)_m$ NCs can be exchange by Ag, Cu, Pt, Pd, iridium (Ir), mercury (Hg), cadmium (Cd) and so on. For Au_n(SR)_m NCs, many of their geometric structures have been determined by SCXRD.³⁹ In conjunction with the development of these synthetic methods, the resulting $Au_n(SR)_m$ NCs has been used as luminescent materials, $^{40-42}$ sensors, 43 photosensitisers, 44-46 catalysts, 11,47-49 chemical electrocatalysts, 22,49-53 and photocatalysts. 49,54,55 This is because the unique electronic/geometric structures of $Au_n(SR)_m$ NCs give rise to properties and functions that are not observed in bulk metals and metal NPs. Research on metal NCs, which can be regarded as ultimate nanomaterials based on atomically precise control, has become increasingly active in recent years.

3 Electronic structures of thiolateprotected metal nanoclusters by electrochemical methods

In this section, we describe the basic experimental techniques of electrochemical measurements and describe a series of studies on the electronic structures of $\operatorname{Au}_n(\operatorname{SR})_m$ NCs among the studies of metal NCs using electrochemical measurement techniques.

3.1 Electrochemical methods for determination of the electronic structures of metal nanoclusters

There are two main parameters used in electrochemical measurements: the voltage (potential) and the current. The

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potential is the energy applied to the electrodes to cause a reaction (Fig. 2a), while the current indicates the reaction rate. By controlling these parameters, the causal relationship between them and their correlation with time can be investigated. In basic electrochemical measurements, three-electrode (working electrode (WE), reference electrode (RE), and counter electrode (CE)) or two-electrode (WE and CE) systems are used. The WE is the main electrode in the electrochemical reaction, and electrons are transferred and chemical reactions occur on its surface. The RE is used to control the potential of the WE and as a standard of the potential for the measurements. The CE is paired with the WE and serves to conduct unwanted current. These electrodes are immersed in an electrolyte containing the reactants, and the potential and current are controlled using a potentiostat to make electrochemical measurements. They can also be evaluated as a solid device by using a electrolyte instead of a solution Electrochemical measurement methods can be divided into several measurement methods depending on the settings of the potential and current (Fig. 2).

Among the electrochemical measurement methods, voltammetry is the most basic method for measuring the flowing current by continuously changing the potential of the WE. Linear sweep voltammetry (LSV) is a technique in which the potential is swept in one direction and the current is measured (Fig. 2b). The measurement method in which the potential is folded back and the applied potential cycle is swept repeatedly

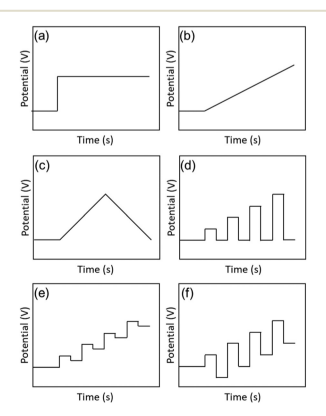


Fig. 2 Electrochemical methods: (a) chronoamperometry, (b) LSV, (c) CV, (d) NPV, (e) DPV, and (f) SWV.

in a specific potential range is called cyclic voltammetry (CV) (Fig. 2c). CV is widely used as a general purpose method because it provides a rough state of the electrochemical condition on the surface of the WE. Furthermore, it can provide information about (i) the ease of oxidation or reduction of the reactants dissolved in the electrolyte, (ii) the chemical reactions at the surface of the WE, and (iii) the species adsorbed on the surface of the WE. The direction and rate of sweep of the applied potential are important parameters, and their behaviour depends on the reversibility of the electrochemical reactions.

LSV and CV gradually change the applied voltage over a period of time and monitor the current during that time. In contrast, pulse voltammetry applies pulse-stepped potentials and measures the current. In LSV and CV, the reaction proceeds with the charging of the electrical double layer at the surface of the electrode, because the potential gradually changes with time. This makes it difficult to separate the nonfaradaic current (i.e., the capacitive current) from the faradaic current (i.e., the reaction current). However, in pulse voltammetry, the charging of the electric double layer immediately ends after the applied potential step, so that the faradaic current can be separated and extracted. Therefore, by giving a slightly different potential each time and plotting the current values at regular intervals, more detailed analysis can be performed using pulse voltammetry compared with LSV and CV. Depending on the shape of the pulse, the measurements can be classified as normal pulse voltammetry (NPV, Fig. 2d), differential pulse voltammetry (DPV, Fig. 2e), and square-wave voltammetry (SWV, Fig. 2f). SWV can significantly reduce the measurement time compared with normal pulse voltammetry and DPV because the step potential is gradually increased. In recent years, pulse voltammetry has been used to determine the electronic states of metal NCs, and it can be combined with optical absorption spectroscopy, luminescence measurements, and DFT calculations to investigate the electronic structures of metal NCs in detail.

3.2 Determination of the electronic structures of metal nanoclusters

The earliest study of voltammetry of small metal NPs was reported by Mulvaney and co-workers in 1997. 19,56 Initially, single-electron tunnelling microelectrode measurements using scanning tunnelling microscopy was the main focus. 57,58 In 1998, Whetten and co-workers⁵⁹ performed electrochemical analysis of $Au_n(SR)_m$ NCs, which were easily obtained by liquid-phase synthesis (Fig. 3). The electronic structures of $Au_n(SR)_m$ NCs with sizes of 8–38 kDa (1.1–1.9 nm in diameter) protected with an alkanethiolate (butanethiolate (SC₄H₉) or hexanethiolate (SC_6H_{13})) were evaluated by DPV measurement.

The double-layer capacitance of small NCs can be modelled by considering the NC to be a concentric sphere capacitor. The inner and outer spheres have different radii, which correspond to the metal-core radius (r) and the thickness of the SR ligand layer (d) in $Au_n(SR)_m$, respectively. Considering the dielectric

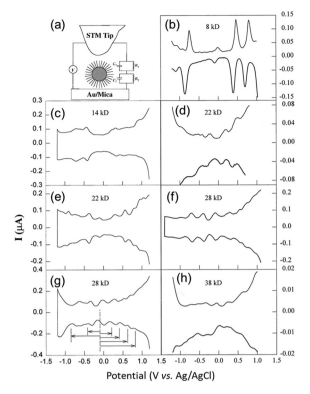


Fig. 3 (a). Schematic of the scanning tunnelling microscopy double tunnel-junction model. Differential pulse voltammograms of (b), (d), (f) and (h) hexanethiolate-, and (c), (e), and (g) butanethiolate-Au NCs as a function of the uniform core size in 0.05 M tetrabutylammonium perchlorate/toluene/acetonitrile (2:1 = v:v) at a 9.5×10^{-3} cm² Pt electrode (direct current potential scan at 10 mV s⁻¹ with a pulse amplitude of 50 mV). The sizes and concentrations were (b) 8 kDa and 0.30 mM, (c) 14 kDa and 0.086 mM, (d) 22 kDa and 0.10 mM, (e) 22 kDa and 0.032 mM, (f) 28 kDa and 0.10 mM, (g) 28 kDa and 0.10 mM, and (h) 38 kDa and 0.10 mM. Reproduced with permission from ref. 59. Copyright 1998, AAAS.

function of the SR ligand layer, the capacitance of the NC $(C_{\rm NC})$ is described by the following eqn (1):^{60,61}

$$C_{
m NC} = 4\pi \varepsilon \varepsilon_0 \frac{r}{d} (r+d)$$
 (1)

where ε is the effective dielectric constant of the SR layer and ε_0 is the permittivity of free space.

Assuming that C_{NC} does not vary with the charge of the metal core, a continuous series of charge steps (ΔV_c) occurs at regular intervals when the metal core catches and releases one electron:

$$V_{\rm c} = \frac{e}{C_{\rm NC}} \tag{2}$$

where ΔV_c is the critical voltage bias, $C_{\rm NC}$ is the capacitance, and e is the electron charge. Such constant-interval charging steps are actually observed in these $Au_n(SR)_m$ NCs. C_{NC} exhibits a minimum value close to the zero-charge potential of the charged interface. Au_n(SR)_m NCs also exhibit similar doublelayer capacitance regardless of whether they are dissolved in solution or deposited on a surface.⁶²

Based on eqn (2), the large $Au_n(SR)_m$ NCs (22-38 kDa) exhibited a Coulomb staircase-like response consistent with a metallic double-layer charge (Fig. 3e and f). In contrast, the small $Au_n(SR)_m$ NCs (8 and 14 kDa) exhibited trends like a molecule (Fig. 3b and c), including a large central gap (1.2 and 0.7 V vs. Ag/AgCl for the 8 kDa $Au_n(SR)_m$ NCs). Such moleculelike behaviour of the small $Au_n(SR)_m$ NCs is similar to that of Pt carbonyl NCs, 63 and it is consistent with the highest occupied molecular orbital (HOMO)-lowest unoccupied molecular orbital (LUMO) gaps obtained from the optical absorption spectra.

Studies then started to estimate the HOMO-LUMO (H-L) gaps of $Au_n(SR)_m$ NCs by electrochemical methods, such as DPV. $^{60,62,64-70}$ However, the electronic structures of $\mathrm{Au}_n(\mathrm{SR})_m$ NCs controlled with atomic precision were only revealed electrochemically from 2004. The development of methods for synthesis and isolation of $Au_n(SR)_m$ NCs and establishment of evaluation methods, such as the development of mass spectrometry instruments, significantly contributed to this breakthrough. In recent years, the electronic structures of a number of Au NCs have been clarified. Such information about the electronic structure can also be obtained by optical absorption spectroscopy and DFT calculations. The H-L gaps of $Au_n(SR)_m$ NCs determined by electrochemical methods are summarised in Table 1, along with information^{71,72} about the electronic structures obtained by other methods. In this review, the effects of the representative factors of the (i) size, (ii) ligand, (iii) introduction of other elements, and (iv) geometrical structure changes on the electronic structures of $Au_n(SR)_m$ NCs are introduced.

3.3 Size effect on the electronic structure of thiolateprotected gold nanoclusters

In 2004, Murray and co-workers⁷³ found that the electrochemically determined band gap of Au₃₈(PET)₂₄ (PET = 2-phenylethanethiolate), which was later modified to Au₂₅(PET)₁₈ (Fig. 4A) owing to developments in mass spectrometry, was consistent with the optically determined band gap. 74 Au₂₅(PET)₁₈ was dissolved in a mixture of toluene/acetonitrile or dichloromethane (CH₂Cl₂) containing 0.1 M tetrabutylammonium hexafluorophosphate (TBAPF₆) using 0.4 mm-diameter Pt for the WE, a Ag wire for the pseudo-RE (Ag QRE), and a Pt wire for the CE. The dissolved Au₂₅(PET)₁₈ was evaluated by DPV and CV (Fig. 4B). A resting potential of -0.1 V vs. Ag QRE was observed at 25 °C, with the first one-electron oxidation peak (O1) slightly on the positive side. Conversely, the first one-electron reduction peak (R1) was largely on the negative side, showing a large energy gap (1.62 V) (Fig. 4B and C). The potential intervals were calculated to be 0.73, 0.29, and 1.62 V for the O3-O2, O2-O1, and O1-R1 transitions, respectively. In this case, measurements at low temperature (-70 °C) contributed to the reversibility, especially in the high oxidation state. In general, the energy gap between the initial one-electron oxidation and one-electron reduction (O1-R1) is the H-L gap including the electron and hole charging energies. Therefore, to calculate the pseudo-H-L gap, this charging energy has to be subtracted.

Table 1 Experimental and theoretical HOMO-LUMO gaps ($E_{\rm q}$) of the thiolate-protected Au NCs mentioned in this review

Composition	Geometric structure	Conditions for the electrochemical measurements	$E_{\rm g}^{\rm e}$ [eV] a	$E_{\mathrm{g}}^{\circ}\left[\mathrm{eV}\right]^{b}$	$E_{\rm g}^{\rm c}$ [eV] c	$Ref.^d$
$Au_{18}(SC_6H_{11})_{14}$	Bi-O _h ^e	RT, DPV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂ ^f	1.91	1.7	_	75
$Au_{15}Ag_3(SC_6H_{11})_{14}$		RT, DPV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂	2.11	2.0	_	75
$[Au_{25}(SC_6H_5)_{18}]^-$	$I_{ m h}^{\ \ h}$	RT, DPV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂	1.16		_	113
$[Au_{25}(SC_6H_{13})_{18}]^-$		RT, SWV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂	1.32	1.3	_	88
		RT, DPV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂ : toluene	~1.65	_	_	108
		RT, DPV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂	1.25	_	_	113
		SWV, $0.1 \text{ M TBAPF}_6/\text{CH}_2\text{Cl}_2$	1.31	_	_	89
$Au_{25}(SC_nH_{2n+1})_{18} (n = 2-12)$		RT, DPV, $0.1 \text{ M TBAPF}_6/\text{CH}_2\text{Cl}_2$	1.29 - 1.31	_	_	99
$Au_{25}(MPS)_{18}^{g}$		RT, DPV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂ : toluene	1.39-1.66	_	_	108
[Au ₂₅ (PET) ₁₈] ⁻		RT, DPV, $0.1 \text{ M TBAPF}_6/\text{CH}_2\text{Cl}_2$	1.34	_	_	76
		RT, DPV, 0.1 M TBAPF ₆ /DMF	1.44	_	_	76
		RT, DPV, $0.1 \text{ M TBAPF}_6/\text{CH}_2\text{Cl}_2$	1.23	_	_	113
		273 K, DPV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂	1.30	1.17	_	116
$[Au_{25}(PET)_{18}]^{+}$		DPV, 0.1 M TBAP/benzene : acetonitrile ⁱ	1.54	_	_	80
$[Au_{24}Pt(SC_4H_9)_{18}]^0$		RT, DPV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂	0.38	_	_	112
$[Au_{24}Pt(SC_6H_{13})_{18}]^0$		SWV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂	0.34	_	_	110
$[Au_{24}Pd(SC_6H_{13})_{18}]^0$			0.32	_	_	110
[Au24Hg(PET)18]0		RT, DPV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂	1.20	_	_	113
			1.29	_	_	112
$[Au_{24}Hg(SC_4H_9)_{18}]^0$			1.28	_	_	
$[Au_{24}Cd(SC_4H_9)_{18}]^0$			1.41	1.37	_	
[Au24Cd(PET)18]0			1.46	1.41	_	
		273 K, DPV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂	1.29	1.22	_	116
$[Au_{24}Pt(PET)_{18}]^{2-}$			>1.80	1.40	_	
[Au ₂₃ CdPt(PET) ₁₈] ⁻			1.45	1.41	_	
$[Au_{25}(PPh_3)_{10}(SC_6H_{13})_5Cl_2]^{2+}$	$\operatorname{Bi-}I_{\operatorname{h}}^{j}$	RT, DPV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂	1.54	1.82	_	101
$[Au_{25}(PPh_3)_{10}(PET)_5Cl_2]^{2+}$		DPV, TBAPF ₆ /CH ₂ Cl ₂	1.57	~ 1.4	_	119
$[Au_{24}Pd(PPh_3)_{10}(PET)_5Cl_2]^+$			1.29	~1.2	_	
$Au_{36}(SPh)_{24}$	FCC^k	RT, DPV, 0.5 M BTPPATBF ₂₀ /CH ₂ Cl ₂ ¹	1.96	1.71	1.51-1.79	81
$[Au_{37}(PPh_3)_{10}(PET)_{10}Cl_2]^+$	Tri-I _h ^m	RT, DPV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂	0.77	0.83	_	82
$Au_{38}(PET)_{24}$	$I_{ m h}$	RT, DPV, $0.1 \text{ M TBAPF}_6/\text{CH}_2\text{Cl}_2$	1.00	0.92	_	83
$Au_{38}(SC_6H_{13})_{24}$		RT, SWV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂	0.99	0.9	_	88
$[Au_{38}(SC_6H_{13})_{24}]^0$		SWV, $0.1 \text{ M TBAPF}_6/\text{CH}_2\text{Cl}_2$	0.86	_	1.01	114
$[Au_{36}Pd_2(SC_6H_{13})_{24}]^0$		SWV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂	0.26	_	0.23	114
$[Au_{36}Pt_2(SC_6H_{13})_{24}]^{2-}$		SWV, $0.1 \text{ M TBAPF}_6/\text{CH}_2\text{Cl}_2$	0.95	_	1.10	114
$Au_{67}(PET)_{35}$	_	RT, DPV, 0.5 M TBAPF ₆ /THF	0.74	_	0.75	84
$Au_{67}(SC_6H_{13})_{35}$	_	RT, SWV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂	0.61	0.7	_	88
$Au_{102}(SC_6H_{13})_{44}$	${D_{ m h}}^o$	RT, SWV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂	0.18	<0.5	_	88
$Au_{103}S_2(S-Nap)_{41}^{n}$		RT, DPV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂	0.38	0.42	_	105
$Au_{130}(PET)_{50}$		RT, DPV, BTPPATBF ₂₀ /CH ₂ Cl ₂	0.21	_	_	85
$Au_{144}(SC_6H_{13})_{60}$	$I_{ m h}$	RT, SWV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂	0.15	<0.5	_	88
$Au_{333}(SC_6H_{13})_{79}$	_	RT, SWV, 0.1 M TBAPF ₆ /CH ₂ Cl ₂	0.04	< 0.5	_	88

 $[^]a$ Electrochemical HOMO-LUMO gap. b Optical HOMO-LUMO gap. c DFT-calculated HOMO-LUMO gap. d Reference number. e Bi- O_h : Bi-octahedral. f TBAPF₆: tetrabutylammonium hexafluorophosphate. g MPS: (3-mercaptopropyl)sulfonate. hI_h : icosahedral. i BAP: tetra-n-butylammonium perchlorate. j Bi- I_h : bi-icosahedral. k FCC: face-centered cubic. l BTPPATBF $_{20}$: bis(triphenyl phosphoranylidene)ammonium tetrakis(pentafluorophenyl). m Tri- I_h : tri-icosahedral. n S-Nap: 2-naphthalenethiolate. o D $_h$: decahedral. Room temperature (RT).

The charging energy was estimated to be 0.29 V, which corresponded to the O2–O1 transition. Because the energy gap between the initial one-electron oxidation and one-electron reduction in $\mathrm{CH_2Cl_2}$ was 1.62 V (O1–R1), a value of 1.33 eV was obtained by subtracting the charging energy of 0.29 V (O2–O1). The absorption edge wavelength (*i.e.*, the H–L gap) obtained from the optical absorption spectrum was 1.33 eV, which was in close agreement with the electrochemically determined H–L gap (Fig. 4C). Subsequently, several groups reported the H–L gaps of $\mathrm{Au_{18}(SC_6H_{11})_{14}}^{75}$ $[\mathrm{Au_{25}(SR)_{18}}]^{-,76-80}$ $\mathrm{Au_{36}(SPh)_{24}}$ (SPh = benzenethiolate), 81 $[\mathrm{Au_{37}(PPh_3)_{10}(PET)_{10}Cl_2}]^+$ (PPh₃ = triphenylphosphine), 82 $\mathrm{Au_{38}(SR)_{24}}^{,68,83}$ $\mathrm{Au_{67}(PET)_{35}}^{,84}$ $\mathrm{Au_{130}(PET)_{50}}^{,85}$ $\mathrm{Au_{130}(Dur-DT)_{29}(PET)_{22}}$ (Dur-DT = durene-dithiolate), 86 and $\mathrm{Au_{133}(TBBT)_{52}}$ (TBBT = tert-butylbenzenethiolate), 87 which were estimated by electrochemical measurements.

Lee and co-workers 88,89 used $Au_{25}(SC_6H_{13})_{18}$, $Au_{38}(SC_6H_{13})_{24}$, $Au_{67}(SC_6H_{13})_{35}$ $Au_{102}(SC_6H_{13})_{44}$ $Au_{144}(SC_6H_{13})_{60}$ $Au_{333}(SC_6H_{13})_{79}$ to obtain the size effect of the H-L gap. They performed SWV measurements of each $Au_n(SR)_m$ NC (Fig. 5A). The electrochemical H-L gaps were determined to be 1.32, 0.99, 0.61, 0.18, and 0.15 eV for $Au_{25}(SC_6H_{13})_{18}$, $Au_{38}(SC_6H_{13})_{24}$, $Au_{67}(SC_6H_{13})_{35}$, $Au_{102}(SC_6H_{13})_{44}$, and $Au_{144}(SC_6H_{13})_{60}$, respectively (Fig. 5B). A significantly size-dependent O1-R1 gap was observed for the small $Au_n(SR)_m$ NCs. The O1-R1 gaps of $Au_{102}(SC_6H_{13})_{44}$, $Au_{144}(SC_6H_{13})_{60}$, and $Au_{333}(SC_6H_{13})_{79}$ were considerably smaller than those of the relatively small $Au_n(SR)_m$ NCs (0.49, 0.39 and 0.22 V, respectively), and the O1 and R1 peaks were observed with equally spaced current peaks along with other peaks. This suggests that Au₁₀₂(SC₆H₁₃)₄₄, $Au_{144}(SC_6H_{13})_{60}$, and $Au_{333}(SC_6H_{13})_{79}$ work as quantum capaci-

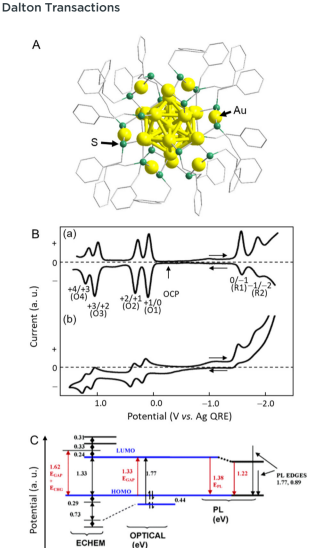


Fig. 4 (A) Geometric structure of [Au₂₅(PET)₁₈]⁻. (B) (a) 25 °C DPVs at 0.02 V s⁻¹, and (b) –70 °C cyclic voltammogram (0.1 V s⁻¹) of Au₂₅(PET)₁₈ in 0.1 M TBAPF₆ in degassed CH₂Cl₂ with a 0.4 mm-diameter Pt WE, a Ag-wire quasi-RE, and a Pt-wire CE. The arrows indicate the solution rest potentials, and * indicates the wave for incompletely removed O₂, which varied from experiment to experiment. (C) Energy level diagram for Au₂₅(PET)₁₈ based on the data in CH₂Cl₂ solution. Reproduced with permission from ref. 74. Copyright 2004, American Chemical Society.

tors that exhibit the charging characteristics of a quantum double layer. In other words, the double-layer capacitance of these $\mathrm{Au}_n(\mathrm{SR})_m$ NCs is sufficiently small that the potential spacing between successive single-electron changes of the charge on the NC core becomes experimentally observable. The optical absorption spectra of $\mathrm{Au}_{25}(\mathrm{SC}_6\mathrm{H}_{13})_{18}$, $\mathrm{Au}_{38}(\mathrm{SC}_6\mathrm{H}_{13})_{24}$, and $\mathrm{Au}_{67}(\mathrm{SC}_6\mathrm{H}_{13})_{35}$ showed absorption edges of 1.3, 0.9, and 0.7 eV, respectively. The absorption edges corresponded to the $\mathrm{Au}_n(\mathrm{SR})_m$ optical gaps for the electronic transitions, and the absorption edge decreased with increasing $\mathrm{Au}_n(\mathrm{SR})_m$ size. For relatively large $\mathrm{Au}_n(\mathrm{SR})_m$ NCs, such as $\mathrm{Au}_{102}(\mathrm{SC}_6\mathrm{H}_{13})_{44}$, $\mathrm{Au}_{144}(\mathrm{SC}_6\mathrm{H}_{13})_{60}$, and $\mathrm{Au}_{333}(\mathrm{SC}_6\mathrm{H}_{13})_{79}$, the optical energy gap

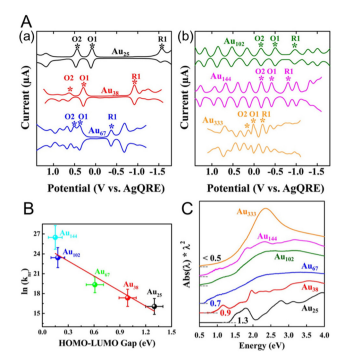


Fig. 5 (A) Square-wave voltammograms of (a) $Au_{25}(SC_6H_{13})_{18}$, $Au_{38}(SC_6H_{13})_{24}$, and $Au_{67}(SC_6H_{13})_{35}$ and (b) $Au_{102}(SC_6H_{13})_{44}$, $Au_{144}(SC_6H_{13})_{60}$, and $Au_{333}(SC_6H_{13})_{79}$ in CH_2Cl_2 containing 0.1 M TBAPF₆. (B) Plot of $ln(k_{nr})$ versus the H–L gap for gold clusters. The solid line is the best fit straight line for the data $(Au_{25}-Au_{144})$. (C) Ultravioletvisible–near-infrared absorption spectra of $Au_{25}(SC_6H_{13})_{18}$, $Au_{38}(SC_6H_{13})_{24}$, $Au_{67}(SC_6H_{13})_{35}$, $Au_{102}(SC_6H_{13})_{44}$, $Au_{144}(SC_6H_{13})_{60}$, and $Au_{333}(SC_6H_{13})_{79}$. Reproduced with permission from ref. 88. Copyright 2017, American Chemical Society.

was less than 0.5 eV and the absorption edge became less distinct. In this respect, the electrochemical method was able to calculate the H–L gap with higher accuracy than optical absorption spectroscopy. From the optical absorption spectrum, $Au_{333}(SC_6H_{13})_{79}$ showed a prominent band of surface plasmon resonance at approximately 2.3 eV. This indicates that electrons collectively oscillate within $Au_{333}(SC_6H_{13})_{79}$ in response to the incident optical electric field, indicating that it has metallic properties.

The electronic structure of $Au_{329}(PET)_{84}$, a relatively large $Au_n(SR)_m$ NC, was further analysed electrochemically by Dass and co-workers. They found that over a potential window of approximately 4 V, $Au_{329}(PET)_{84}$ exhibited 22 electronic states. In this study, the capacitance varied in proportion to the size of the $Au_n(SR)_m$ NC. The transition of the electronic state from molecular to metallic in $Au_n(SR)_m$ is still not completely understood. $Au_{246}(p\text{-MBT})_{80}$ (p-MBT = p-methylbenzenethiolate) is the largest non-metallic $Au_n(SR)_m$ NC and $Au_{279}(TBBT)_{84}$ is the smallest metallic $Au_n(SR)_m$ NC, as revealed by Jin and coworkers $^{91-94}$ by transient absorption spectroscopy. However, while $Au_{246}(p\text{-MBT})_{80}$ has an octahedral structure, $Au_{279}(TBBT)_{84}$ has a face-centred cubic (fcc) structure. The influences of the differences in the geometry and ligand effect

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have not been clarified, so further investigation is required.95,96

The small size region of $Au_n(SR)_m$ NCs was further investigated by Jin and co-workers 97 using Au₁₃₃(TBBT)₅₂ and $Au_{144}(BM)_{60}$ (BM = phenylmethanethiolate). They found that the charging energy decreases with the catch and release of $Au_n(SR)_m$ electrons in the more oxidised/reduced state (Fig. 6). This trend can be attributed to an increase in the capacitance of $Au_n(SR)_m$ due to an increase in the dielectric constant of the ligand monolayer by penetration or intercalation of electrolyte ions. Indeed, when the temperature during the electrochemical measurements of Au₁₃₃(TBBT)₅₂ was lowered to 195 K, the charging energy became independent of the redox state and showed a constant value (Fig. 6A and B). Interestingly, the charging energy was much less sensitive to temperature changes for Au₁₃₃(TBBT)₅₂ in the reduced state compared with oxidised Au₁₃₃(TBBT)₅₂. To access the core stabilised by the bulky tert-butyl terminal functional group, the tetrabutylammonium cation is considerably less accessible than the perchloride anion because of steric hinderance. The relatively large decrease in the charging energy of oxidised NCs in the low temperature measurements could be due to suppression of the movement of such sterically hindered molecules at low temperature. Conversely, the BM monolayer of Au₁₄₄(BM)₆₀ is more flexible owing to the absence of the bulky tert-butyl group between the S atom and the phenyl group. Therefore, Au₁₄₄(BM)₆₀ exhibits a smaller peak potential spacing owing to it having larger capacitance than Au₁₃₃(TBBT)₅₂ with roughly the same sized metal core (Fig. 6C and D).

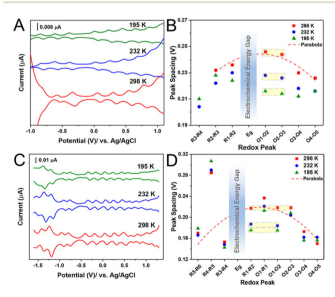


Fig. 6 Temperature dependence of electrochemical properties of Au₁₃₃(TBBT)₅₂ and Au₁₄₄(BM)₆₀. (A and C) The DPVs and (B and D) the electrochemical peak spacing of (A and B) Au₁₃₃(TBBT)₅₂ and (C and D) $Au_{144}(BM)_{60}$ at 298, 232, and 195 K. Peak spacing ΔVs in (B) and (C) analyzed from (A) and (C), respectively. Reproduced with permission from ref. 97. Copyright 2020, American Chemical Society.

3.4 Ligand effect on the electronic structure of thiolateprotected gold nanoclusters

Changes in the ligand from the S atom onwards have little effect on the electrochemically determined H-L gap in many cases, 74,90,97-101 but replacing the S atom with a different element causes a change in the H-L gap.

Negishi and co-workers102 investigated the ligand dependence using $[Au_{25}(PET)_{18-x}(SePh)_x]^-$ (SePh = benzeneselenolate) and $[Au_{25}(PET)_{18-x}(TePh)_x]^-$ (TePh = benzenetellurolate) by DPV measurements in 2016 (Fig. 7A). When the PET ligands were replaced by SePh or TePh, the redox potentials of R1 and O1 shifted towards the positive side (Fig. 7B). However, when the PET ligands were replaced by SePh, the redox potentials of O2 and O3 shifted to the negative side in the final step of ligand exchange (around x = 18) (Fig. 7B). Furthermore, the H-L gap narrowed as the number of exchanges increased. The relationships between the changes in the redox potentials of O2 and O3 and the number of exchanged ligands were not linear. This is presumably because of changes in the framework structure of the NC with increasing number of exchanged ligands, reflecting the difference in the atomic radii of S and Se. A similar trend was also reported by Zhu and co-workers 103 using $[Au_{25}(PET)_{18}]^-$ and $[Au_{25}(SePh)_{18}]^-$.

In 2019, Tsukuda and co-workers¹⁰⁴ reported the electronic structures of C=CR-protected Au NCs by electrochemical measurements. The electrochemically determined superatomic orbital 1P-1D gaps of $[Au_{25}(PET)_{18}]^-$ and $[Au_{25}(C \equiv CAr^F)_{18}]^-$

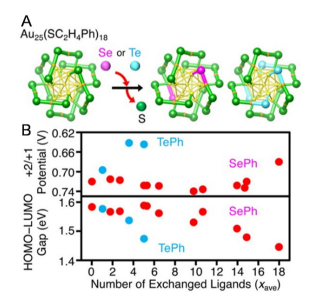


Fig. 7 (A) Framework structures of $[Au_{25}(PET)_{18-x}(SePh)_x]^-$ and [Au₂₅(PET)_{18-x}(TePh)_x] determined by SCXRD. (B) Relationships of the peak energy of O3 and the H-L gap with the average number of exchanged ligands (x_{ave}) for $[Au_{25}(PET)_{18}]^-$, $[Au_{25}(PET)_{18-x}(SePh)_x]^-$, and $[Au_{25}(PET)_{18-x}(TePh)_x]^-$. In the experiments, each cluster was dissolved in 0.1 M tetrabutylammonium perchlorate in CH₂Cl₂. DPV was performed at room temperature. Reproduced with permission from ref. 102. Copyright 2016, American Chemical Society.

(Ar^F = 3,5-(CF₃)₂C₆H₃) were 1.93 and 1.8 eV, respectively (Fig. 8A). A similar trend was observed when the superatomic orbital 1P–1D gaps were determined from the optical absorption spectra, with $[Au_{25}(C = CAr^F)_{18}]^-$ having a smaller 1P–1D gap than $[Au_{25}(PET)_{18}]^-$ (Fig. 8B). This can be attributed to the stronger cationic nature of Au in $[Au_{25}(C = CAr^F)_{18}]^-$ than in $[Au_{25}(PET)_{18}]^-$. The ligand dependence on the H–L gap owing to the change of the Au–S interface has been reported to be limited to relatively small $Au_n(SR)_m$ NCs. $^{105-107}$

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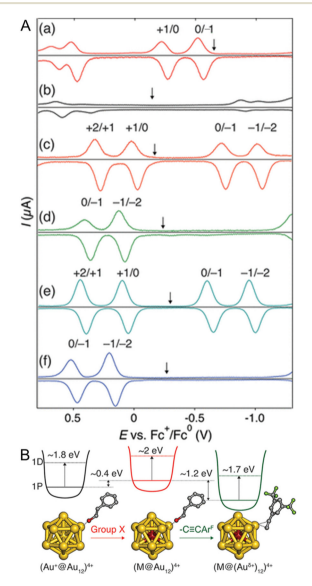
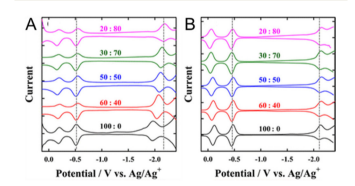


Fig. 8 (A) Differential pulse voltammograms of (a) $[Au_{25}(PET)_{18}]^-$, (b) $[Au_{25}(C = CAr^F)_{18}]^-$, (c) $[PdAu_{24}(PET)_{18}]^0$, (d) $[PdAu_{24}(C = CAr^F)_{18}]^{2-}$, (e) $[PtAu_{24}(PET)_{18}]^0$, and (f) $[PtAu_{24}(C = CAr^F)_{18}]^{2-}$ in a CH_2CI_2 solution of 0.1 M $TBAPF_6$ at room temperature. The down arrows indicate the corresponding open-circuit potentials. (B) Schematic energy diagram of M@Au $_{12}$ superatoms protected by thiolate or alkynyl groups. Reproduced with permission from ref. 104. Copyright 2019, American Chemical Society.

Solvent-ligand interactions also have an important effect on the electrochemically obtained H-L gap. Kwak and Lee¹⁰⁸ investigated the H-L gaps of water-soluble Au₂₅(MPS)₁₈ ((3mercaptopropyl)sulfonate) and [Au₂₅(SC₆H₁₃)₁₈] by optical electrochemical methods (Fig. 9A and B). Electrochemically, SWV measurements in CH2Cl2 containing 0.1 M TBAPF₆ were performed. For [Au₂₅(SC₆H₁₃)₁₈]⁻, a H-L gap of 1.65 V was obtained, whereas the H-L gap for Au₂₅(MPS)₁₈ was 1.39 V. The O1-O2 gap difference was not very large, suggesting that the dielectric environments were not significantly different in the NCs. Furthermore, the ratio of the CH₂Cl₂/toluene solvent mixture was varied, and SWV measurements were performed in these solvents (Fig. 9A and B). In $[Au_{25}(SC_6H_{13})_{18}]^-$, the H-L gap did not change when the ratio of the mixed solvent was changed (Fig. 9B). However, for Au₂₅(MPS)₁₈, the first reduction potential (R1) shifted in the negative potential direction as the proportion of toluene increased. As a result, the H-L gap shifted from 1.39 to 1.66 V, which was found to be in close agreement with [Au₂₅(SC₆H₁₃)₁₈] (Fig. 9A). This solvent dependence can be explained by the electrostatic effects of the MPS ligand and TOA+ (TOA = tetraoctylammonium): the SO₃ group of the MPS ligand produces an ionpair interaction with TOA+, resulting in depolarisation. Furthermore, depolarisation is expected to be influenced by the solvent polarity, which affects the strength of the ionpair interaction between the SO₃ group and TOA⁺. As a result, more positive charge is induced in the Au core by the polarised SO₃ group, which may result in a positive redox potential shift in Au₂₅(MPS)₁₈. Similarly, the charging energy is known to vary with the solvent, even when only hydrophobic ligands are used. These effects of the ligand and solvent interactions on the H-L gap and charging energies might be better understood when the details of how solvation occurs between the SR ligands and solvents



surrounding $Au_n(SR)_m$ NCs are further clarified.

Fig. 9 Square-wave voltammograms of (A) $Au_{25}(MPS)_{18}$ and (B) $Au_{25}(SC_6H_{13})_{18}$ in CH_2Cl_2/t oluene mixtures containing 0.1 M TBAPF₆. The CH_2Cl_2/t oluene compositions (v/v) are indicated on the voltammograms. The first oxidation and reduction potentials of $Au_{25}(SR)_{18}$ in 20:80 CH_2Cl_2/t oluene are indicated by dashed lines for comparison. Reproduced with permission from ref. 108. Copyright 2012, American Chemical Society.

3.5 Doping effect on the electronic structure of thiolate-

protected gold nanoclusters

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For Au₂₅(SR)₁₈, there are a number of different elements (Ag, Cd, Cu, Hg, Ir, Pd, and Pt) that can be substituted for the Au atoms, and these are the most widely studied doped $Au_n(SR)_m$ NCs. These $Au_{25-x}M_x(SR)_{18}$ alloys can be divided into three groups: (i) a group with Pt or Pd, (ii) a group with Ag or Cu, and (iii) a group with Hg or Cd as the heteroatom. For NCs in group (i), neutral $[Au_{24}M_1(SR)_{18}]^0$ (M = Pt or Pd) with a total valence electron number of six can be stably synthesized. 109,110 However, the NCs in group (ii) are stably synthesised as negative $[Au_{25-x}M_x(SR)_{18}]^-$ (M = Ag or Cu) NCs, and those in group (iii) are neutral $[Au_{25-x}M_x(SR)_{18}]^0$ (M = Hg or Cd) NCs with a total number of eight valence electrons. The difference in the total number of valence electrons in such Au_{25-x}M_x(SR)₁₈ NCs induces significant changes of the energy level in their HOMOs and LUMOs. Consequently, optical absorption of $Au_{25-x}M_x(SR)_{18}$ (M = Ag, Cd Cu, Hg, Pd or Pt) significantly differs between the six- and eight-electron systems. The sixelectron system shows a smaller H-L gap than the eight-electron system.

In 2012, Negishi and co-workers. 111 optically and electrochemically determined the electronic structures $[Au_{25-x}Cu_x(PET)_{18}]^-$ (x = 1-5) (Fig. 10). By DPV measurements, for [Au₂₅(PET)₁₈]-, the redox peaks corresponding to $[Au_{25}(PET)_{18}]^{-1/0}$ and $[Au_{25}(PET)_{18}]^{-2/-1}$ were observed at -282(O1) and -1942 mV (R1) vs. ferrocene (Fc) $^{+1/0}$, respectively (Fig. 10B). For $[Au_{25-x}Cu_x(PET)_{18}]^-$ (x = 1-5), the peaks were observed at more positive energies (-276 (O1) and -1896 (R1) mV vs. Fc^{+1/0}). This indicates that Cu doping decreases the HOMO and LUMO energies of [Au₂₅(PET)₁₈]⁻. The LUMO shift (46 mV) was larger than the HOMO shift (6 mV), indicating that Cu doping decreases the H-L gap. DFT calculations showed that Cu doped at the stable site does not significantly change the H-L gap, whereas Cu doped at the core surface site decreases the H-L gap.

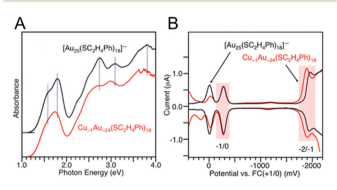


Fig. 10 (A) Optical absorption spectra of Cu_{~1}Au_{~24}(PET)₁₈ and [Au₂₅(PET)₁₈]⁻. The dotted lines indicate the main peak positions in the absorption spectrum of [Au₂₅(PET)₁₈]⁻. (B) DPV curves of $Cu_{\sim 1}Au_{\sim 24}(PET)_{18}$ and $[Au_{25}(PET)_{18}]^-$. The red shaded regions indicate the peaks derived from the redox potentials of -2/-1 and -1/0. Reproduced with permission from ref. 111. Copyright 2012, American Chemical Society.

In 2019, Maran and co-workers 112 revealed the electronic structures of Au₂₄M(SR)₁₈ (M = Au, Cd, Hg and Pt) by DPV measurements (Fig. 11A). Electrochemical measurements were performed using a glassy carbon microdisc electrode in CH₂Cl₂ containing 0.1 M TBAPF₆. The DPV curves of [Au₂₄Hg $(SC_4H_9)_{18}]^0$ and $[Au_{24}Cd(SC_4H_9)_{18}]^0$ were qualitatively similar: two successive one-electron oxidations (E_1° for O1 and E_2° for O2) were observed at 0.364 and 0.684 V vs. SCE for [Au24Hg $(SC_4H_9)_{18}]^0$ and at 0.332 and 0.636 V vs. SCE for $[Au_{24}Cd$ $(SC_4H_9)_{18}$ ⁰. On the timescale of the voltammetry experiments, both reactions were reversible processes. The first peaks were observed on the more positive side for $[Au_{24}Hg(SC_4H_9)_{18}]^0$ and $[Au_{24}Cd(SC_4H_9)_{18}]^0$ compared with those for $[Au_{25}(SC_4H_9)_{18}]^ (E_1^{\circ} = -0.188 \text{ and } E_2^{\circ} = 0.139 \text{ V} \text{ vs. SCE})$. A similar trend was observed using the PET ligand instead of the SC₄H₉ ligand. The first reduction peaks (R1) were observed at -1.23 and -1.39 V vs. SCE for $[Au_{24}Hg(SC_4H_9)_{18}]^0$ and $[Au_{24}Cd$ $(SC_4H_9)_{18}$ ⁰, respectively. These anion formation processes were chemically irreversible, in which case the potential can be detected reversibly by increasing the potential scanning rate. The H-L gap of [Au₂₄Cd(SC₄H₉)₁₈]⁰ was determined to be 1.41 eV, which was in good agreement with the H-L gap of 1.37 eV determined by optical absorption (Fig. 11B). Similarly, the H-L gap of $[Au_{24}Cd(PET)_{18}]^0$ was determined to be 1.46 eV, which was in good agreement with the H-L gap of 1.41 eV determined by optical absorption (Fig. 11B). Furthermore, a

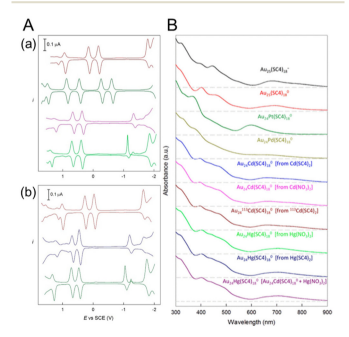


Fig. 11 (A) Comparison of the DPV curves of (from top to bottom) (a) $[\mathrm{Au}_{25}(\mathrm{SC}_4\mathrm{H}_9)_{18}]^-, \ [\mathrm{Au}_{24}\mathrm{Pt}(\mathrm{SC}_4\mathrm{H}_9)_{18}]^0, \ [\mathrm{Au}_{24}\mathrm{Cd}(\mathrm{SC}_4\mathrm{H}_9)_{18}]^0, \ \mathrm{and} \ [\mathrm{Au}_{24}\mathrm{Hg}]^0$ $(SC_4H_9)_{18}]^0$ and (b) $[Au_{25}(PET)_{18}]^-$, $[Au_{24}Cd(PET)_{18}]^0$, and $[Au_{24}Hg]^0$ $(PET)_{18}$]⁰ (glassy-carbon electrode, $CH_2Cl_2/0.1$ M TBAH, 25 °C). (B) Ultraviolet-visible absorption spectra of all of the SC₄H₉-protected samples (0.2 mM, 1 mm cuvette) in CH₂Cl₂. For better comparison, the curves have been shifted vertically. The dashed lines indicate the corresponding zero absorbance. Reproduced with permission from ref. 112. Copyright 2019, American Chemical Society.

decrease in the H-L gap was observed for Hg doping. The change of the H-L gap with Hg doping has also been reported by Yang, Wu and co-workers, 113 but the H-L gap difference was not as large as the differences in the Pt- and Pd-doped sixelectron systems.

In 2015, Lee and co-workers¹¹⁰ investigated the electronic structures of $[Au_{24}Pt(PET)_{18}]^0$ and $[Au_{24}Pd(PET)_{18}]^0$ (Fig. 12A). For $[Au_{24}Pd(PET)_{18}]^0$, the open-circuit potential was in the middle of the two redox peaks (O1 and R1) at -0.48 V vs. Fc^{+1/} o, indicating that it was neutral. Furthermore, the electrochemical gap between O1 and R1 was 0.75 V, giving a H-L gap of 0.32 V when the charge energy (O1-O2, 0.43 V) was considered. This was a dramatic decrease from the H-L gap of 1.32 eV for $[Au_{25}(PET)_{18}]^{-}$. $[Au_{24}Pt(PET)_{18}]^{0}$ showed a similar trend, with a H-L gap of 0.32 eV, indicating that [Au24Pt (PET)₁₈]⁰ and [Au₂₄Pd(PET)₁₈]⁰ have very similar electronic states (Fig. 12A). To investigate the origin of the small H-L gap, DFT calculations were performed to simulate the energy levels of the six- and eight-electron systems (Fig. 12B). $[Au_{25}(SCH_3)_{18}]^-$, $[Au_{24}Pd(SCH_3)_{18}]^{2-}$, and $[Au_{24}Pt(SCH_3)_{18}]^{2-}$ exhibit stable superatomic eight-electron systems (1S21P6), whereas the six-electron systems, including $[Au_{24}Pd(SCH_3)_{18}]^0$ and [Au₂₄Pt(SCH₃)₁₈]⁰, have two electrons less than eight-electron systems. Therefore, they split from a triple degenerate HOMO into a doubly degenerate HOMO and LUMO, functioning as a superatomic 1P orbital (Fig. 12B). The H-L gap cannot be determined from the optical absorption spectrum because the dipole selection rule forbids optical HOMO-LUMO transitions. However, the H-L gaps of $[Au_{24}Pt(PET)_{18}]^0$ and $[Au_{24}Pd$ $(PET)_{18}^{10}$ can be revealed by SWV, because they are not forbidden electrochemically. Therefore, this electrochemical method is particularly useful for $Au_n(SR)_m$, where the HOMO-LUMO

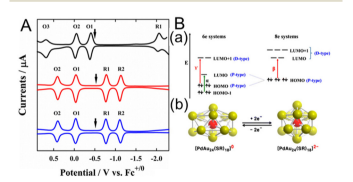


Fig. 12 (A) Square-wave voltammograms of [Au₂₅(SC₆H₁₃)₁₈]⁻ (black), $[PdAu_{24}(SC_6H_{13})_{18}]^0$ (red), and $[PtAu_{24}(SC_6H_{13})_{18}]^0$ (blue). The electrochemical measurements were performed in CH2Cl2 containing 0.1 M TBAPF₆. The arrows indicate the solution open-circuit potentials. (B) (a) Electronic energy levels of the six-electron and eight-electron systems. α , β , and γ denote the optical transitions occurring in the six- and eightelectron systems. (b) Schematic showing Jahn-Teller-like distortion in the core (e.g., PdAu₁₂) predicted for six-electron [PdAu₂₄(SR)₁₈]⁰ (left), which undergoes a structural change to nearly spherical eight-electron [PdAu₂₄(SR)₁₈]²⁻ upon reduction (right). The vertical compression of [PdAu₂₄(SR)₁₈]⁰ (left) is exaggerated. Reproduced with permission from ref. 110. Copyright 2015, American Chemical Society.

transition is optically forbidden. In addition, they reported the doping effects of Pt and Pd atoms for Au₃₈(SR)₂₄. The H-L gap of $[Au_{36}Pt_2(SC_6H_{13})_{24}]^{2-}$ measured by voltammetry was slightly larger than that of undoped $[Au_{38}(SC_6H_{13})_{24}]^0$, whereas the H-L gap of [Au₃₆Pd₂(SC₆H₁₃)₂₄]⁰ was significantly smaller (0.26 V), resulting in the appearance of a near-infrared band at ~0.8 eV.

The change in the H-L gap has also been determined by electrochemical methods for doping with multiple different atoms. In 2020, Negishi and co-workers¹¹⁵ determined the H-L gaps of $[Au_{24}Pt(SR)_{18}]^0$ and $[Au_{24-x}PtCu_x(PET)_{18}]^0$ by DPV measurements (Fig. 13A). The H-L gap of the NC hardly changed by Cu substitution (Fig. 13A-C). The states of the HOMO and LUMO in [Au₂₄Pt(SR)₁₈]⁰ are caused by the dissolution of the degenerate electronic states owing to the symmetry lowering of the metal core (Fig. 13C, Jahn-Teller effect).³⁹ It was presumed that substitution of Cu at the core surface Au did not significantly change the magnitude of the strain in the metal core, resulting in the H-L gap hardly changing. In 2020, Tsukuda and co-workers¹¹⁶ revealed the electronic structure of [Au₂₃PtCd(PET)₁₈] by optical spectroscopy and electrochemical measurements (Fig. 14A and B). The first and second oxidation potentials (O1 and O2) of [Au23PtCd (PET)₁₈ were 0.110 and 0.175 V higher than those of [Au₂₅(PET)₁₈]⁻, respectively (Fig. 14B). This indicates that the 1P superatomic orbitals of Pt@CdAu₁₁ in [Au₂₃PtCd(PET)₁₈] are slightly more stable than those of Au@Au₁₂ in [Au₂₅(PET)₁₈]⁻. The electrochemically determined H-L gap of [Au23PtCd(PET)18] was 1.45 eV, which was in good agreement with the optically determined H–L gap (1.41 eV). This is sig-

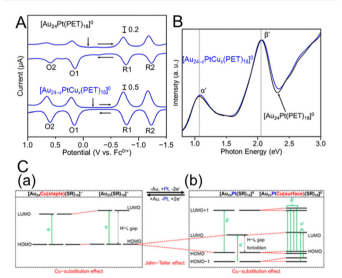


Fig. 13 (A) DPV curves of $[Au_{24}Pt(PET)_{18}]^0$ and $[Au_{24-x}PtCu_x(PET)_{18}]^0$ (x= 0.9). The vertical arrows indicate the solution open-circuit potentials, and the horizontal arrows indicate the scan direction. (B) Optical absorption spectra of $[Au_{24-x}PtCu_x(PET)_{18}]^0$ (x = 0.9) (blue) and $[Au_{24}Pt]^0$ (PET)₁₈]⁰ (black). (C) Schematic of the effect of Cu substitution on the structure of $[Au_{25-x}Cu_{x}(SR)_{18}]^{-}$ and (a) $[Au_{24-x}PtCu_x(SR)_{18}]^0$. Reproduced with permission from ref. 115. Copyright 2020, American Chemical Society.

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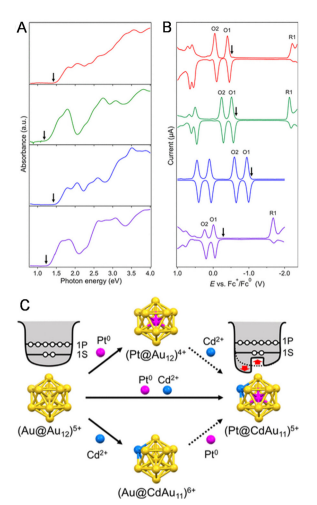


Fig. 14 (A) Ultraviolet–visible–near-infrared spectra and (B) DPV curves of $[Au_{23}PtCd(PET)_{18}]^-$ (red), $[Au_{25}(PET)_{18}]^-$ (green), $[Au_{24}Pt(PET)_{18}]^0$ (blue), and $[Au_{24}Cd(PET)_{18}]^0$ (purple) in benzene and CH_2Cl_2 , respectively. The downward arrows in panels (A) and (B) represent the corresponding optical H–L gaps and open-circuit potentials, respectively. (C) Schematic illustration of the jellium potentials of the (Au@Au₁₂)⁵⁺ and (Pt@CdAu₁₁)⁵⁺ superatoms. Reproduced with permission from ref. 116. Copyright 2020, American Chemical Society.

nificantly larger than the H–L gap (1.30 eV) of $[Au_{25}(PET)_{18}]^-$. However, the H–L gap of $[Au_{24}Cd(PET)_{18}]^0$ (1.29 eV) was almost the same as that of $[Au_{25}(PET)_{18}]^-$. The H–L gap of the Pt@CdAu₁₁ superatomic core is larger than that of Au@Au₁₂, mainly because of Pt doping at the centre (Fig. 14C), which is in good agreement with the electronic structure changes observed by optical absorption spectroscopy and electrochemical measurements.

The Ag replacing effect on the electronic structure of $\operatorname{Au}_n(\operatorname{SR})_m$ was also investigated by Dass and co-workers. ⁹⁰ using $\operatorname{Au}_{144-x}\operatorname{Ag}_x(\operatorname{PET})_{60}$ (x=46). The electrochemical gap of $\operatorname{Au}_{144}(\operatorname{PET})_{60}$ was approximately 0.37 V, while that of $\operatorname{Au}_{144-x}\operatorname{Ag}_x(\operatorname{PET})_{60}$ was approximately 0.38 V. A previous study of the band gap of $\operatorname{Au}_{144-x}\operatorname{Ag}_x(\operatorname{PET})_{60}$ using a combination of optical absorption spectroscopy and DFT calculations showed that the optical band gap is only moderately affected by Ag

doping in this size range, 117 which was consistent with previous reports.

In summary, previous studies indicate that doping the centre of $\mathrm{Au}_{25}(\mathrm{SR})_{18}$ with Pt or Pd as a heteroatom has a significant effect on the electronic structure. It is expected that doping multiple elements can create an even greater variety of electronic states if the doping positions of the different elements can be controlled. The effects of doping with different elements have been found to be more pronounced for $\mathrm{Au}_n(\mathrm{SR})_m$ with smaller size.

3.6 Geometric structure effect on the electronic structure of thiolate-protected gold nanoclusters

Park and Lee¹⁰¹ compared the electronic states of Au₂₅ NCs with two Au₁₃ icosahedra connected via vertex sharing $([Au_{25}(PPh_3)_{10}(SC_6H_{13})_5Cl_2]^{2+})$ and one Au_{13} icosahedral core $([Au_{25}(SC_6H_{13})_{18}]^-)$ (Fig. 15). In this case, a series of $[Au_{25}(PPh_3)_{10}(SR)_5Cl_2]^{2+}$ NCs protected with SR (SR = SC₆H₁₃, PET, 11-hydroxy-1-undecanethiolate (MU), and 10-carboxy-1-decanethiolate (MUA)) was prepared. SWV measurements were performed with $[Au_{25}(SC_6H_{13})_{18}]^-$ and $[Au_{25}(PPh_3)_{10}(SC_6H_{13})_5Cl_2]^{2+}$ in 0.1 M TBAPF₆/CH₂Cl₂ at -78 °C. The results showed that the solution open-circuit potential of the $[Au_{25}(PPh_3)_{10}(SC_6H_{13})_5Cl_2]^{2+}$ was approximately 0.37 V vs. 1,4benzoquinone (BQ)^{-/0}, and a series of oxidation peaks at 0.91 V (O1), 1.13 V (O2), 1.45 V (O3), and 1.62 V (O4) vs. BQ^{-/0} was observed, corresponding to oxidation of +3, +4, +5, and +6 for $[Au_{25}(PPh_3)_{10}(SC_6H_{13})_5Cl_2]^{2+}$, respectively (Fig. 15B). From this voltammogram, the electrochemical energy gap was determined to be 1.54 V from the difference between the initial oxidation (O1) and reduction (R1) potentials (Fig. 15B). Compared with [Au₂₅(SC₆H₁₃)₁₈]⁻, the oxidation and reduction peaks (O1 and R1) of [Au₂₅(PPh₃)₁₀(SC₆H₁₃)₅Cl₂]²⁺ were observed at significantly

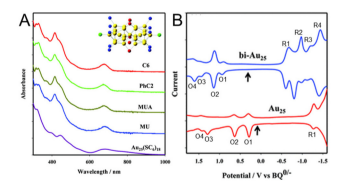


Fig. 15 (A) Absorption spectra of $[Au_{25}(PPh_3)_{10}(SR)_5Cl_2]^{2^+}$ (SR = SC_6H_{13} , PET, MUA, and MU) and $Au_{25}(SC_6H_{13})_{18}$ in CH_2Cl_2 . All of the spectra have been normalized to unity at 300 nm and offset for clarity. The insert shows the core structure of $Au_{25}(PPh_3)_{10}(SR)_5Cl_2$. (B) Square-wave voltammograms of $[Au_{25}(PPh_3)_{10}(SC_6H_{13})_5Cl_2]^{2^+}$ (top) and $[Au_{25}(SC_6H_{13})_{18}]^-$ (bottom) in 0.1 M TBAPF₆ in degassed CH_2Cl_2 at -78 °C. The arrows indicate the solution open-circuit potentials. The open-circuit potential of Au_{25} indicates that the Au_{25} NC is an anionic form, $[Au_{25}(SC_6H_{13})_{18}]^-$. The concentration of $[Au_{25}(PPh_3)_{10}(SC_6H_{13})_5Cl_2]^{2^+}$ was 4 mg mL⁻¹, and the tick interval on the current axis is 0.2 μA. Reproduced with permission from ref. 101. Copyright 2012, American Chemical Society.

more positive potentials. The electrochemical energy gap of $[Au_{25}(PPh_3)_{10}(SC_6H_{13})_5Cl_2]^{2+}$ (1.54 V) was smaller than that of $[Au_{25}(SC_6H_{13})_{18}]^-$ (1.65 V), indicating that these NCs had clearly different electronic energy structures. The electrochemical energy gap of $[Au_{25}(PPh_3)_{10}(SC_6H_{13})_5Cl_2]^{2+}$ was also smaller than that of the icosahedral Au13 NC assigned to $Au_{13}(PPh_3)_4(SC_{12}H_{25})_2Cl_2$ (1.76 V).⁶⁹ Furthermore, the potential gap between O2 and O3 (0.32 V) was significantly smaller than the potential gap of $[Au_{25}(SC_6H_{13})_{18}]^-$ (0.65 V). From the difference between the O2-O3 gap and O1-O2 gap, the charging energy of 0.10 eV was obtained. The electronic energy levels adjacent to the HOMO of $[Au_{25}(PPh_3)_{10}(SC_6H_{13})_5Cl_2]^{2+}$ were revealed to be degenerate or close to each other owing to their high symmetry.118 Although there were differences in their charge states and some ligands, such as SRs, phosphines, and halogens, in the Au₂₅ NCs with different geometries, the difference in their electronic structures was revealed. Furthermore, for such [Au₂₅(PPh₃)₁₀(SR)₅Cl₂]²⁺ NCs, it has been reported that Pd doping decreases the H-L gap owing to splitting of the degenerate HOMO. 119

The effect of the $Au_n(SR)_m$ core structure on the electrochemical gap was investigated by Wu and co-workers. 120 The electronic structures of Au₂₈(S-c-C₆H₁₁)₂₀ (S-c-C₆H₁₁ = cyclohexanethiolate), $Au_{28}(TBBT)_{20}$, $Au_{25}(PET)_{18}$, $Au_{44}(DMBT)_{26}$ (DMBT = 2,4-dimethylbenzenethiolate), Au₄₄(TBBT)₂₈, Au₄₉(DMBT)₂₇, Au₃₈(PET)₂₄, and Au₃₆(TBBT)₂₄ were investigated. The results showed that the $Au_n(SR)_m$ NC with a kernel of the fcc structure has a larger electrochemical gap than the $Au_n(SR)_m$ NC with a kernel of the non-fcc structure.

Conclusions and outlook

Electrochemical techniques can be used to determine the H-L gaps of metal NCs, and the electronic structures of various $Au_n(SR)_m$ have been elucidated by electrochemical measurements, as well as by optical absorption spectroscopy and DFT calculations. Bulk Au does not have a band gap, but a molecule-like H-L gap forms in small Au_n(SR)_m NCs, and the H-L gap tends to increase with decreasing size. Au₂₅(SR)₁₈ is currently the most studied $Au_n(SR)_m$ NC. In $Au_{25}(SR)_{18}$, molecular changes of R in the SR ligand do not significantly change the H-L gap, but the electronic state greatly changes when S is a different atom or group (e.g., Se, Te, and C \equiv CR). Substitution of different elements for Au (i.e., alloying) also changes the H-L gap. In particular, substitution of Pt or Pd for Au in Au₂₅(SR)₁₈ significantly reduces the H-L gap owing to dissolution of the degeneracy of the electronic states. Furthermore, in $Au_n(SR)_m$, where the HOMO-LUMO transition is optically forbidden, the H-L gaps, which cannot be calculated from the optical spectra, can be determined by electrochemical methods.

These electrochemical methods have also been applied to Au NCs protected by phosphine and stibine, and to NCs based on Ag, 130–133 Cu, 134 and group 10 elements. 135–139 In the future, these electrochemical techniques are expected to

reveal the electronic states of metal NCs protected by other ligands and composed of other metals. 140-144

The electrochemical gap also varies depending on the ligand, solvent, electrolyte, measurement mode, temperature, and other factors. This is also the same when determining the H-L gaps from optical absorption spectra, and care should be taken when comparing the H-L gap values obtained in different studies. More accurate H-L gap calculations are required through the development of models that more accurately take the charging energy into account and a unified measurement method. 61,145,146

The electronic structures of $Au_n(SR)_m$ NCs in solution have already been extensively studied. However, it is desirable to clarify the electronic states of $Au_n(SR)_m$ NCs on catalyst supports when considering their use in catalytic reactions in the gas or solid phase. 135,147-151 In this case, it is expected that the electronic/geometric state of $Au_n(SR)_m$ NCs will also significantly change during catalytic reaction. For instance, ligands have also been reported to desorb from $Au_n(SR)_m$ NCs upon applying potential152 and the catalytic properties to be depends on the electronic state of $Au_n(SR)_m$ NCs. 153 It is expected that techniques such as pulse voltammetry can be used to elucidate the mechanism of such electrochemical catalytic reactions. If the electronic states of $Au_n(SR)_m$ NCs during catalytic reaction could be investigated, it would be possible to clarify the details of the catalytic reaction mechanisms predicted by theoretical calculations. 154 It is expected that the electronic states of various $Au_n(SR)_m$ NCs will be clarified by electrochemical methods and these measurement techniques will be further developed in the future.

Conflicts of interest

There are no conflicts to declare.

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