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Monolayer-Protected Cluster Molecules

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ABSTRACT

In this report, we evaluate the present state of the rapidly emerging field of monolayer-protected cluster (MPC) molecules with regard to their synthesis and monolayer functionalization, their core and monolayer structure, their composition, and their properties. Finally, we canvass some of the important remaining research opportunities involving MPCs.

Introduction

Ligand-capped gold clusters have been known for a number of years; however, the practical formation of stable, isolable monolayer-protected clusters (MPCs) has only recently been demonstrated. In a seminal contribution, Schiffrin and co-workers showed that the classical two-phase colloid preparation of Faraday$^{1b}$ could be combined with contemporary phase transfer and alkanethiol/Au chemistry to prepare, in a simple procedure, very small clusters of gold atoms ($\leq 5$ nm average core diameter) coated with alkanethiolate monolayers.

Alkanethiolate MPCs differ from conventional colloids$^{1b}$ and nanoparticles prepared by other routes (micelle$^{13a}$ and polymer stabilization,$^{10b}$ gas-phase cluster formation,$^{5c}$ stabilization with small organic ligands$^{13a-13b}$) in that they can be repeatedly isolated from and redissolved in common organic solvents without irreversible aggregation or decomposition. This attribute and the air stability of MPCs are difficult to overemphasize, since they allow for handling and derivatization of MPCs in ways familiar to molecular chemists, plus considerable flexibility in characterization relative to previous nanoparticle embodiments.$^{5}$

Metallic and semiconducting nanoparticles represent a scantily explored materials dimension that bridges bulk material and molecular behavior$^{6}$ and offers prospects of novel, and even size-dependent, chemical, electronic, and physical properties. Such properties may, additionally, depend on whether nanoparticles comprise organized monolayer or multilayer films or are solutions of organized or randomly dispersed entities. Following a long-standing interest in nanometer-scale materials,$^{7}$ we adopted Schiffrin’s procedure$^{2}$ in an exploration of the synthesis, characterization, and functionalization of MPCs as a new molecular class. Also of interest are structural analogies of MPCs to self-assembled monolayers on flat surfaces$^{8}$ (2D-SAMs) and to large, spatially defined macromolecules, such as dendrimers.$^{9}$

Progress in these directions by our laboratory and other laboratories forms the core of this Account.$^{10}$

MPC Synthesis

In the original Schiffrin reaction,$^{2}$ addition of dodecanethiol to organic-phase AuCl$_4^-$ (1:1 mole:mole), followed by reduction with BH$_4^-$, leads to dodecanethiolate-protected Au clusters having a 1–3 nm range of core diameters:

$$\text{AuCl}_4^-(\text{toluene}) + \text{RSH} \rightarrow (-\text{Au}_n\text{SR}_m)(\text{polymer}) \quad (1)$$

$$(-\text{Au}_n\text{SR}_m)_0 + \text{BH}_4^- \rightarrow \text{Au}_n(\text{SR})_y \quad (2)$$

Subsequent reports have shown that a wide range of alkanethiolate chainlengths (C3–C24)$^{11a}$ can-functionalized alkanethiols,$^{12}$ and dialkyl disulfides$^{13}$ can be employed in this same protocol (Scheme 1). An example MPC formula is Au$_{14}(\text{S(CH}_2)_3\text{CH}_3)_30$.

The details of reactions 1 and 2 have not been completely dissected; however, the reaction’s behavior is consistent with a nucleation–growth–passivation process: (i) Larger thiol:gold mole ratios give smaller average MPC core sizes,$^{5,14}$ and (ii) fast reductant addition and cooled solutions produce smaller, more monodisperse MPCs,$^{5,15a}$ (iii) Quenching the reaction immediately following reduction produces higher abundances of MPCs having very small core sizes ($\leq 2$ nm).$^{15b,c}$

The Schiffrin reaction tolerates considerable modification in regard to the protecting ligand structures (Scheme 1).$^{16}$ While alkanethiolate MPCs (a) are nonpolar, highly polar ligands (b–d) can yield$^{16c-d}$ water-soluble MPCs in modified syntheses. MPCs with arenethiolate$^{16d-e}$ (e–g) and ($\gamma$-mercaptopropyl)trimethoxysilane ligands (h) have also been prepared.$^{16h}$ Sterically bulky ligands (larger footprint) tend to produce smaller Au core sizes (relative to alkanethiolate MPCs prepared using equal thiol/AuCl$_4^-$ ratios), suggesting$^{16b,c}$ a steric connection (as yet unproven) to the dynamics of core passivation.

The core metal of MPCs can also be varied, and reports of MPCs with Ag$^{17a,b}$ and alloy (Au / Ag, Au / Cu, Au / Ag / Cu, Au / Pt, Au / Pd, and Au / Ag / Cu / Pd) cores (monolayer-protected alloy clusters, MPACs)$^{17b}$ have appeared. Ag MPCs and MPACs, like their MPC counterparts, are

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Characterization of MPCs

The excellent handling properties of MPCs facilitate their structural and compositional analysis, which can be divided into analytical information about the central core and the surrounding monolayer.

Core Size and Shape. The core dimensions of MPCs have been measured—with generally good agreement—by scanning tunneling microscopy (STM), transmission electron microscopy (TEM), small-angle X-ray scattering (SAXS), laser desorption–ionization mass spectrometry (LDI-MS), and X-ray diffraction (XRD). TEM allows concomitant observation of the self-ordering and regular core–core spacing found in thin MPC films.

Knowing and controlling the dispersity of core size is also important. Figure 1 illustrates the use of LDI-MS to track the solubility–fractionation of dodecanethiolate MPC core sizes, conducted in concert with theoretical modeling to predict the core size and shape of isolated fractions. The number of atoms per core tends toward closed shell structures (“magic numbers”) with an equilibrium truncated octahedral shape. As a cautionary note, however, reactions 1 and 2 are based on competing core growth/passivation kinetics, so the occurrence of non-equilibrium MPC shapes cannot be discounted. Analysis of water-soluble MPCs with highly polar ligands is less advanced, but dispersities have been resolved by gel chromatography and capillary electrophoresis and fractions in the former examined with MALDI-MS.

MPC Monolayers. MPC monolayer shells have also been examined with multiple techniques. Most studies have been conducted on polydisperse (core size) MPCs, so the data represent averages. Elemental analysis is consistent with intact RS–ligands (reactions 1 and 2). The average number of ligands per core is derived from a combination of XPS or thermogravimetric and core size analyses. The MPC ligand coverages—50% or more—are larger than the 33% (ligand/surface Au atom) coverage characteristic of 2D-SAMs on Au(111) terraces. Theoretical calculations indicate that this high coverage results from larger ligand/Au binding ratios on core edges and vertexes (confirming our own modeling). Experiments addressing

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the nature of MPC ligands provides strong evidence that, like 2D-SAMS, they should be regarded as covalently-attached thiolate-like units, not adsorbed disulfides. Our reference to the ligand as a thiolate should not be taken as signifying anionic RS⁻Au⁺ moiety.

NMR spectroscopy is particularly informative about the structure and content of MPC monolayers.⁵,¹¹,¹⁹a–d ¹H and ¹³C NMR resonances of MPC monolayers are characteristically broadened relative to those of free alkanethiols; the factors include spin–spin relaxation broadening, a distribution in chemical shifts due to differences in Au–SR binding sites, and a gradient in monolayer packing density from near-core to chain terminus with associated dipolar broadening. Figure 2 illustrates the broadening of ¹H NMR resonances of dodecanethiolate MPCs with increased core size, which for the methyl resonance (δ ~ 0.9) is thought to reflect primarily T₂ broadening. The products of MPC functionalization reactions can be assessed with NMR data, generally more accurately following T₂-induced MPC decomposition to a disulfide solution.²¹

Vibrational spectroscopy has shown that alkanethiolate chains of 2D-SAMS are typically in an all-trans, zigzag configuration. Chain conformations in solid-state MPC films while mostly all-trans, also contain significant (5–25%) gauche defect concentrations at both inner and terminal chain ends, especially for longer chainlengths and above the chain melting temperature. Hydrogen bonding can be an important structural component in MPCs with polar monolayers.¹⁶a, b Monolayer ordering in solid-state MPC samples is also associated with interdigitation of chain domains (or “bundles”) on neighboring cluster molecules.¹⁹c In contrast, monolayers of dissolved alkanethiolate MPCs exhibit disorder comparable to that of liquid alkanes. That MPC monolayers prove to be only rough analogies of 2D-SAMS is likely due to the high proportion of classically defined MPC core surface defect sites (edges, vertexes).²¹

Order–disorder transitions in solid-state alkanethiolate MPCs are detectable with differential scanning calorimetry (DSC) and variable-temperature IR and NMR spectroscopies (Figure 3).¹⁹a–c The transition temperatures increase with methylene chain length (like melting of crystalline alkane hydrocarbons) and, by FTIR, are accompanied by increased gauche defect concentrations.¹⁹a Transition enthalpies (scaled to the organic fraction of the MPC) also increase with methylene chain length, and are smaller than those of corresponding alkanes.¹⁹c

In an elegant order–disorder study, Lennox and coworkers applied variable-temperature solid-state ²H NMR to MPCs with deuterated octadecanethiolate monolayers. The results showed that, as temperature is increased, disorder (increase in gauche bond population) propagates from the chain terminus toward the middle of the chain, leading to eventual chain melting, but does not extend to the tethered sulfur headgroup.

Reactivity of MPCs

Understanding reactivities of MPC monolayers and developing efficient strategies to functionalize them is key to their application in areas such as catalysis and chemical sensing. Toward this end, we have investigated ligand place-exchanges (to form poly-homo- and -hetero-functionalized MPCs),²¹ nucleophilic substitutions,²¹ and ester and amide couplings (Scheme 2).²¹ The resulting MPCs are large molecules with multiple functionalities, spherically organized around a central, metal core, a spatial arrangement suggesting analogy to dendrimers. However, whereas dendrimers are typically more dense at their periphery than core, MPCs are “soft objects with hard cores”.

Place-Exchange Reactions. Ligand place-exchange was a key step in opening up MPC functionalization. For example, MPCs with alkanethiolate monolayers (RS) can be functionalized with R’ groups by the reaction (Scheme 2a)

\[
x(R'SH) + (RS)_mMPC \rightarrow x(RSH) + (R'S)_n(RS)_{m-n}MPC
\]

where \(x\) and \(m\) are the numbers of new and original ligands, respectively.²²a The rate and equilibrium stoichiometry (\(x\)) of reaction 3 are controlled by factors that include the (reaction feed) mole ratio of R’SH to RS units, their relative steric bulk, and R versus R’ chainlengths.²²b, c

Study of ligand place-exchange dynamics and mechanism shows that exchange (i) has a 1:1 stoichiometry, (ii) is an associative (as opposed to dissociative) reaction, (iii) yields the displaced ligand in solution as a thiol, and (iv) does not involve disulfides or oxidized sulfur species. The time-dependent rate of place-exchange was interpreted as reflecting a hierarchy of different core surface binding sites with associated susceptibility (vertexes, edges and terraces) to place-exchange.

Reactivity of ω-Functional Groups. Functional groups on the outermost portion of the MPC ligand sphere can straightforwardly undergo many synthetic transformations. In an early example Schiffrin reported the esterification of p-mercaptophenol-protected MPCs using propionic anhydride (Scheme 2b)

Our laboratory examined steric effects in MPC reactivity with Sₓ₂ reactions of mixed monolayer MPCs containing o-bromoalkanethiols (Scheme 2c). The reactivities are similar to those of primary alkyl halide monomers (RBr),¹ result a sharply divergent from re-
ports\textsuperscript{25} of analogous reactions on 2D-SAMs. The S\textsubscript{0}2 reactivity did slow with increasing steric bulk of primary amine nucleophiles, but even \(\text{o}-\text{bromo sit e s}^\text{buried}^\text{"exposed" ones (ii) These results reflect a monolayer disorder consistent with the results of vibrational spectroscopy of dissolved MPCs.}\textsuperscript{19}

Amide and ester coupling reactions give access\textsuperscript{23} to alkanethiolate/Au-based MPCs bearing multiple copies of a very diverse selection of structural groups. The reactions start from a small and readily prepared (by place-exchange) subset of \(\omega\)-functionalized MPC materials (Scheme 2d). Attractive features of this synthetic strategy are the following: (i) the available amide and ester activating agents are diverse, (ii) a plethora of target molecules with amine, carboxylic acid, or alcohol substituents is available, and (iii) the synthesis (often not facile) of \(\omega\)-functionalized alkanethiol derivatives, required in the place-exchange pathway, is avoided. The coupling strategy has been extended to functionalize water-soluble MPCs\textsuperscript{26} and to CdS nanoparticles.\textsuperscript{27}

Other MPC functionalization strategies include coupling of MPCs with functionalized silane monolayers with mercaptosilane monomers (Scheme 2e),\textsuperscript{166} forming cage-like structures surrounding the gold core. Mirkin’s group has used \(\omega\)-norbornenyl-functionalized MPCs to grow polymeric shells from a MPC surface using a functional-group-tolerant ruthenium carbene ROMP catalyst (Scheme 2f);\textsuperscript{28} these shells can be functionalized with virtually any norbornenyl-containing monomer.

**Chemical Properties of Functionalized MPCs**

MPCs bearing spectroscopic (fluorophores and chromophores) and redox groups (Figure 4) have been prepared using the above reactions.

**Electrochemistry.** Electrode reactions of poly-redox-functionalized clusters are unusual in that many equivalents of redox charge per MPC can be delivered, under diffusion control, to the electrode/solution interface. The few known precedents of poly-electron transfer reactions include soluble redox polymers\textsuperscript{29} and redox-labeled dendrimers.\textsuperscript{9} MPCs have been poly-functionalized with ferrocene (\(\leq 25\) sites per MPC),\textsuperscript{22a,30} anthraquinone (\(\leq 25\)),\textsuperscript{31} phenothiazine (\(\leq 7\)),\textsuperscript{23} and viologen (\(\leq 36\))\textsuperscript{26} groupings. Figure 4a illustrates voltammetry\textsuperscript{30} of a \(\text{Fc}_\text{2}\)-MPC solution; analogous ferrocenes have been examined as Langmuir monolayers at the air/water interface.\textsuperscript{30c} MPCs that are poly-hetero-functionalized\textsuperscript{22b} with mixtures of ferrocene and anthraquinone sites exhibit voltammetry of the two groups, occurring independently of each other. Diffusion coefficients from electrochemical measurements have been compared with those from Taylor dispersion experiments.\textsuperscript{32}

The poly-electron transfer reactions studied so far\textsuperscript{22,30,32} are mass transport-controlled one-electron events that rapidly occur either serially or independently parallel. All attached groups are electroactive.\textsuperscript{23} The electrochemical observations also include currents for charging of the electrical double layers of the MPC Au cores, a characteristic reflecting equivalence of MPCs to “diffusing nanoelectrodes”.

Multiple MPC electron donor and acceptor groups invite applications to mediated electrocatalysis. The basic prospects include concerted poly-electron exchanges with substrates, intramolecular electron transfer reactions during binding of MPCs to substrates (perhaps transiently), and (“cofactor”) MPCs that are “multipurpose” by allowing both electron and proton transfers with a substrate in electron–proton coupled reactions. When the poly-redox groups are weakly electronically coupled to the Au core, mediated electrocatalysis is likely to occur only by sequential one-electron transfers. A significant step toward truly concerted poly-electron transfers will be shortening (or altogether removing) the connecting spacers between the redox groups and the metal-like Au, so as to promote strong electron donor–acceptor/core electronic coupling.

Our first\textsuperscript{11} probe of mediated electrocatalysis with MPCs was based on the reduction of gem-dinitrocyclo-
hexane (DNC) by clusters bearing ca. 25 electrogenerated anthraquinone radical anion (AQ\textsuperscript{-}) sites. This study showed that (i) MPC and monomeric AQ\textsuperscript{-} site reactivities with DNC were nearly the same, and (ii) the observed electrocatalytic currents are enhanced, relative to freely diffusing AQ\textsuperscript{-}, due to the smaller diffusion coefficient of MPC AQ\textsuperscript{-} and ensuing compression of the reaction layer.

**Other Attached Groups.** MPCs can, of course, be endowed with other forms of behavior by choice of the poly-functionality. The carboxylic acids of water-soluble tiopronin MPCs (Scheme 1) can be titrated as weak acids (Figure 4b), revealing electrolyte concentration-dependent charge repulsion affects analogous to those of polyprotic acids.\textsuperscript{16b} Labeling MPCs with fluorophores may lead to reagents useful in chemical analysis. Tiopronin MPCs functionalized with fluorescein\textsuperscript{26} (~3.7/MPC) showed ~50% lower luminescence compared to fluorescein monomer, suggestive of energy transfer to the gold core (Figure 5).

\textsuperscript{a}(a) Place-exchange,\textsuperscript{22} (b) reaction of $p$-mercaptophenol with propionic anhydride,\textsuperscript{16e} (c) $S$\textsubscript{N}2 reaction of $\omega$-bromoalkanethiolated MPCs with primary alylamines,\textsuperscript{21} (d) amide and ester coupling reactions,\textsuperscript{27} (e) siloxane formation reactions,\textsuperscript{16b} and (f) transition-metal-catalyzed ring-opening metathesis polymerization (ROMP).\textsuperscript{28}
No polymer electrolytes can be fashioned by using polyether MPC ligands, such as thiolated poly(ethylene glycol) (Figure 4d). This material dissolves Li⁺ salts and, for reasons not yet understood, is more ionic conductive than the corresponding polyether ligand.

We anticipate that further studies will reveal that chemical properties such as the above are, to some extent, tunable as a function (for example) of coverage on the nanoparticle surface, of core properties (size, charge, metal), and of relative chain lengths of core-to-functional group connections.

**Properties of MPC Cores**

**Core Spectroscopy.** X-ray photoelectron spectroscopy (XPS) of 2D-SAMs shows Au 4f binding energies very near those of Au(0) and S 2p binding energies, consistent with a RS⁻ ligand. Like 2D-SAMs, MPCs give energies distinctive of Au(0) and a surface thiolate, indicating that the bond is somewhere between RS⁻M and RS'M⁺. These results are striking inasmuch as as many as about one-third of the Au atoms in the MPC bear alkanethiolate ligands. Only for the smallest cores is a suggestion of an increased binding energy seen.

UV-visible spectra of nanometer-sized MPCs contain several features. The strong UV absorption of the

**FIGURE 5.** UV/vis spectrum of dodecanethiolate MPC with 3.0 nm average Au core diameter. Inset: Spectra of dodecanethiolate MPCs with alloy cores: (a) AuAg₂₅ (λ_max = 468 nm), (b) AuAg₅₀₄ (504 nm), (c) AuAg₅₅₅ (518 nm), (d) AuAg₉₃₇₄₄ (488 nm). All spectra were acquired at ~10⁻⁶ M concentration in hexane.
Au core decays roughly exponentially into the visible, with a broad surface plasmon (SP) band superimposed at ~500 nm that decreases in intensity and energy (slightly) with decreasing core size (Figure 5). For monolayer-protected clusters with alloy cores, the SP band positions and intensities are, interestingly, intermediate between those of corresponding mono-metal MPCs (Figure 5 inset).

Spectra of very small, monodisperse Au MPCs (1.1–1.9 nm core diameter) develop distinct step-like structures, indicative of more molecule-like properties and transitions to discrete LUMOs of the conduction band. Such spectra have been used to estimate HOMO–LUMO “gap” energies as a function of core size for Au MPCs.

**Core Charging Properties.** The quantization of double layer capacitance charging is one of the more interesting solution electrochemical properties of alkanethiolate MPCs. The capacitance is associated with the ionic space charge formed around an MPC dissolved in an electrolyte solution, upon electronic charging of the core. Because of the small, sub-attofarad (aF) capacitance ($C_{CLU}$) of an MPC, charging of the tiny capacitor by single electrons occurs in potential intervals, $\Delta V = \frac{e}{C_{CLU}}$, that exceed $k_B T$. These potential intervals are readily observable in voltammetry (Figure 6b) of MPC solutions. The discretized (one electron) double layer charging steps are smeared out when the MPC core sizes are not mono-disperse or are too large. Quantized MPC double layer charging is formally analogous to classical STM-based “Coulomb staircase” experiments on single MPCs, but the equivalent circuits differ. Our initial descriptor, “ensemble Coulomb staircase charging”, has been replaced with the simpler and more accurate one, “quantized double layer charging”.

The spacing of the current peaks on the potential axis in Figure 6b reflects the underlying nature of the MPC core charging process. An even peak spacing signifies a
metal-like core whose charging is controlled by electrostatic (i.e., double layer) principles. The evolution of molecule-like properties when the MPC core is very small is seen as a wide spacing between the two current peaks adjacent to the MPC potential-of-zero charge ($E_{PZC}$, ca. –0.2 V vs Ag/AgCl). This electrochemical band gap is consistent with the optical band gap of such MPCs.

It is important to understand that one-electron double layer charging peaks in MPC solutions are formally analogous to those produced by traditional redox reactions (i.e., oxidation of ferrocene). The double layer charging of MPCs is kinetically fast (demonstrably controlled by mass transport), so that the profile of working electrode current versus average potential of MPC cores in the adjacent solution is determined by traditional combinations of the Nernst equation with mass transport relationships. Each core charging step has a “formal potential”, which, referenced to the MPC potential of zero charge ($E_{PZC}$), is:

$$E_{PZC}^{\text{z}} = E_{PZC} + (z - 1/2)e/C_{CLU}$$

where $E_{PZC}^{\text{z}}$, the formal potential of the $z$ ($z - 1$) charge state “couple” corresponds to DPV peak currents (Figure 6b), $z$ is signed such that $z > 0$ and $z < 0$ correspond to core “oxidation” and “reduction”, respectively. Insofar as $C_{CLU}$ is potential independent, which seems to be so for potentials not far removed from $E_{PZC}$, this relation predicts a linear plot of $E_{PZC}^{\text{z}}$ vs charge state, with $C_{CLU}$ determined from its slope. Recent studies of the monolayer chain length dependence of $C_{CLU}$ show that $C_{CLU}$ varies with the dielectric thickness in remarkable agreement with a simple concentric sphere capacitor model (Figure 6c,d). The simple model predicts no dependence in $C_{CLU}$ on solvent or electrolyte changes.

The charge stored on MPC cores can be used for chemical reactions. We have electrolytically stored charge on MPCs in solutions; these solutions can be dried and reconstituted with minimal loss of the MPC core charge upon redissolution. Charges can also be stored on MPCs using chemical reactions. Charged MPCs can be used as quantitative redox reagents; Figure 7 shows an example (without a drying step) of titration of an ethyl ferrocene solution with oxidatively charged MPCs. Charged MPCs also undergo electron transfer reactions with themselves, as shown by observing equilibrium potentials in solution mixtures of differently charged MPCs. All results to date are consistent with the analogy of MPC core electrochemistry with regular redox reactions. Other analogies are of interest; optical and electron transfer dynamics properties of electronically charged MPCs are under active study.

**Solid-State Electronic Conductivity.** Dry (solid-state) films of alkanethiolate MPCs are electronically conductive, to degrees governed by core-to-core electron tunneling through intervening monolayers. Under high potential gradients, current–potential curves have been obtained for dry MPC films that are characteristic of electron hopping conductivity. The electron hopping rates decreased and the activation barrier increased with increasing alkanethiolate chainlength, consistent with a combination of Cermet theory (hopping controlled by thermally activated electron transfer that creates oppositely charged Au cores) and chainlength-dependent tunneling. The MPCs gave a tunneling constant of $\beta = 1.2/\text{Å}$, similar to electron transfer rates studied in 2D-SAMs of $\sigma$-redox group functionalized alkanethiolytes ($\beta \approx 1/\text{Å}$) and confirmed by more recent work. MPCs with mixed core charge states exhibit enhanced electronic conductivity as expected for a “mixed-valent” material.

### Conclusions and Outlook

This report has described recent advances in the MPC field, but there remain numerous unexplored territories. Further understanding of MPC behavior requires expansion of synthetic routes to core-monodisperse MPCs and to those with rationally designed ligand shells and core compositions. Better understanding and control of these and other essential properties may lead to applications to areas such as outlined in Scheme 3. The substantial progress made in the five years since the Schiffrin route.
was published is impressive. Overall, however, we feel that this field is still in its infancy, and new fundamental insights and applications are almost certain to be revealed in the future.

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References


(12) Some reports describe monolayer-stabilized clusters prepared by capping an already-prepared colloid particle with an alkanethiol monolayer; such materials typically flocculate over time or with loss of solvent.


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