Solution-phase synthesis of $\text{Al}_{13}^-$ using a dendrimer template

Tetsuya Kambe$^{1,2}$, Naoki Haruta$^2$, Takane Imaoka$^{1,2,3}$ & Kimihisa Yamamoto$^{1,2}$

Superatoms, clusters that mimic the properties of elements different to those of which they are composed, have the potential to serve as building blocks for unprecedented materials with tunable properties. The development of a method for the solution-phase synthesis of superatoms would be an indispensable achievement for the future progress of this research field. Here we report the fabrication of aluminum clusters in solution using a dendrimer template, producing $\text{Al}_{13}^-$, which is the most well-known superatom. The $\text{Al}_{13}^-$ cluster is identified using mass spectrometry and scanning transmission electron microscopy, and X-ray photoelectron spectroscopy is used to measure the binding energies. The superatomic stability of $\text{Al}_{13}^-$ is demonstrated by evaluating its tendency toward oxidation. In addition, the synthesis of $\text{Al}_{13}^-$ in solution enables electrochemical measurements, the results of which suggest oxidation of $\text{Al}_{13}^-$. This solution-phase synthesis of $\text{Al}_{13}^-$ superatoms has a significant role for the experimental development of cluster science.
Superatoms, clusters that exhibit properties similar to those of elements different to those of which they are composed, have the potential to serve as building blocks for unprecedented materials with tunable properties. Studies on superatoms have been mainly theoretical and have been supported by mass detection in the gas phase under high-vacuum conditions\(^{1,2}\). With regard to further progress in this research field, achieving a solution-phase synthetic method for superatoms is important. The approach using noble metals Au, Ag, and Cu produces ligand-protected clusters, and although they have produced cluster crystals\(^{3,4}\), these are thermodynamically stable magic clusters. Therefore, a more advanced synthetic strategy is required.

Experimental studies on superatoms started with the synthesis of vaporized sodium and aluminum clusters\(^{5,6}\), and have attracted wide attention since the discovery of the Al\(_{13}\) halogen-like superatom by Khanna et al\(^{7}\). This was followed by the synthesis of Al\(_{7}\), A\(_{3}\), and KO\(_{2}\)–\(^{8,9}\). Superatom properties can be estimated using a simple theory referred to as the Jellium model\(^{10}\) that defines superatomic orbitals (e.g., 1S, 1P, 1D, 2S, 1F, 2P) in a highly symmetric cluster. For example, the Al\(_{13}\) cluster has 39 valence electrons and exhibits a halogenic nature based on its superatomic orbitals (1S\(^2\), 1P\(^6\), 1D\(^{10}\), 2S\(^2\), 1F\(^{14}\), 2P\(^3\)), resulting in a reactivity with \(I^–\) and the high stability of the mono anionic species Al\(_{13}^–\)\(^{11,12}\).

The aluminum clusters with anionic or cationic states have been extensively studied and confirmed by gas-phase synthesis and quantum chemical calculations\(^{13–15}\). Such a study also revealed photoelectron spectroscopy about the energy levels of the valence electrons using laser light, and the superatom character of Al\(_{13}\)—containing an extraordinary stability against oxidation\(^{16–18}\). On the other hand, the solution-phase synthesis has advantages because the synthesis is scalable without any special apparatus. Although poly(vinylpyrrolidone) or tetrahydrofuran (THF) units containing an extraordinary stability against oxidation\(^{16–18}\) are superatom exists in solution.

**Results**

**Fabrication of aluminum clusters using DPAG4.** The preparation method is illustrated in Fig. 1a. The number of Al atoms in the clusters was controlled using the stepwise assembly feature of the DPAs. The fourth-generation DPA dendrimer (DPAG4) and that with a pyridine core (pyDPAG4) were used (Supplementary Fig. 1). The assembly of AlCl\(_3\) was confirmed by UV–Vis titration; the absorption change in a THF solution of the DPAG4 or pyDPAG4 during the addition of AlCl\(_3\) indicated complexation behavior between the imine site and AlCl\(_3\) in a radial fashion, and a multi-step shift of the isosbestic point provided evidence that the targeted amounts of the AlCl\(_3\) were coordinated in the DPAGs\(^{14}\). The DPAG4 enabled the controlled assembly of 4, 12, 28, and 60 units of AlCl\(_3\) (Supplementary Fig. 2). In contrast, the pyDPAG4, which has a pyridine core that can coordinate one additional metal salt, enabled the assembly of 13 units of AlCl\(_3\) in the dendritic capsule. Supplementary Figure 3 shows assembly of AlCl\(_3\) from 0 to 61 equivalents. The obvious change in the isosbestic point at 13 equivalents demonstrates preparation of 13AlCl\(_3\)–pyDPAG4 in solution. The AlCl\(_3\) units positioned in the DPAs were reduced into clusters by a benzophenone ketyl radical prepared using sodium metal in THF. The fabricated aluminum clusters were then subjected to various experiments. The aluminum clusters with 14 atoms were prepared using the DPAG4. Even though the branched structure of the DPAG4 was not the best for 14 atom clusters, the enough ability to fabricate the cluster as the major product with small deviation was already reported in our previous work\(^{19}\).

The fabricated cluster was protected by the dendrimer shell; however, the superatomic Al\(_{13}^–\) could be detected using negative-mode matrix-assisted laser desorption/ionization (MALDI) mass spectrometry at \(m/z=350.9\) via dissociation from the DPA template during the laser desorption process. This peak was overlapped by a fragment of the pyDPAG4, but the formation of Al\(_{13}^–\) in solution was still confirmed (Fig. 1b). Al\(_{13}^–\) was specifically observed when the 13AlCl\(_3\)–pyDPAG4 complex was reduced, whereas it was not detected for 12AlCl\(_3\)–pyDPAG4 and 14AlCl\(_3\)–pyDPAG4 (Supplementary Fig. 4). In addition, the Al\(_{13}^–\) cluster was detected using high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM). The observed cluster sizes were measured to be 1.0 ± 0.1 nm (Fig. 1c, d). The cluster sizes were reasonable considering the calculated diameter of Al\(_{13}^–\) (0.95 nm, see Methods). These results confirm that the Al\(_{13}^–\) superatom exists in solution.

**Electronic state.** The oxidation states of the clusters were estimated from XPS Al 2p spectra. The XPS samples were prepared on glassy carbon substrates by repetitive drop-casting. The spectra and binding energies (BEs) of the aluminum clusters which were synthesized using 4, 12, 13, 28, and 60 AlCl\(_3\) at the DPAs are summarized in Fig. 2a, b, respectively. All the aluminum clusters possessed BEs equal or lower than that of bulk aluminum, thereby indicating the presence of reduced clusters. In addition, the BE for \(n=13\) cluster seems to be different from the BEs for \(n=4, 12\), and 28 atomic clusters (Fig. 2b). Although the XPS spectra generally reflect the surface state of materials, it is not applied to these aluminum clusters because they have lower than or equal to 60 atoms with the diameters of about 1 nm. Considering that the photoelectrons can escape from about a 1–4 nm thick material\(^{35}\), almost all atoms in the clusters were analyzed.

To interpret these XPS results in a theoretical manner, population analyses for aluminum clusters including Al\(_{13}^–\) were performed (Supplementary Fig. 5). The employed structures were based on the literatures\(^{34,35}\). To estimate the charge distribution effect on the 2p binding energies, we have calculated the 2p orbital levels. Although core-hole relaxation in the final state has not been taken into account, simple correlation between core levels and binding energies has been reported\(^{36}\). In the case of Al\(_{13}^–\), the 2p levels of the inner atom are apart from those of surface atoms (Supplementary Fig. 6). This difference offers possibility of two peaks in the XPS spectrum. However, only one broad peak was detected in our experiments. The observed peak could be ascribed to 12 surface atoms or the mixture of 12 surface atoms and an inner atom. Also, in the cases of anionic clusters in other sizes, the non-uniform charge distributions yield different 2p levels, as shown in Supplementary Fig. 7a. However, all the 2p levels are...
close and overlap to be one broad peak. The size dependence of the 2p levels of anionic clusters is consistent with our experimental results. This is in contrast to the neutral clusters (Supplementary Fig. 7b).

These shifts of binding energy in the XPS measurements and calculated energy levels could be considered by the influence of the negative charges. The charges per atom were simply calculated based on the assumption that the anionic charges are fully delocalized in the aluminum clusters (Supplementary Fig. 8). The anionic charges per atom increased as the number of atoms decreased. This tendency was in agreement with the observed BEs. Therefore, these BEs are assignable to monovalent or more than divalent anionic states.

**Stability.** The reactivity of the aluminum clusters with oxygen was evaluated by monitoring the change in the XPS Al 2p peaks. The samples deposited on the substrates with the DPAs were exposed to air and subjected to Ar sputtering (2 kV, 2 min) to remove any surface impurities before the measurement. The results are shown in Fig. 3. The peaks corresponding to the oxidized species from Al_{12} and Al_{14} clusters increased within several tens of hours, whereas the peak for Al_{13}^{-} maintained its position. This result demonstrates the specific stability of the Al_{13}^{-} clusters toward oxygen and is consistent with the superatomic nature observed in the gas phase^{18}. The clusters observed within the DPAs were deposited with ketyl radical salts; however, they did not affect the stability because the radical anions were quickly oxidized in air within several minutes (Supplementary Fig. 9). The extraordinary stability of Al_{13}^{-} was well theoretically investigated in previous reports^{37–39}. In addition, inclusive studies of aluminum clusters were also reported with their local and global minima^{44,40,41}. They fully supported our result.

**Electrochemical oxidation.** The oxidation processes of the prepared aluminum clusters in the DPAs were investigated by potentiodynamic electrochemical measurements in THF

---

**Fig. 1** Fabrication of Al_{13}^{-} using the dendrimer template. **a** Illustration of dendritic poly-phenylazomethines including an aluminum cluster in the solution state. **b** Negative-mode matrix-assisted laser desorption/ionization mass spectra of Al_{13}^{-} and blank. 1,8,9-Trihydroxyanthracene was used as the matrix. **c** A high-angle annular dark-field scanning transmission electron microscopy image of Al_{13}^{-} on a TEM grid covered with a thin carbon film (scale bar: 5 nm). **d** Histogram of the observed particle sizes in the STEM sample.
results, the voltammogram of AlCl₃ measured in THF showed a reduction and Al₂O₃. The fabricated clusters were protected by the DPA template. It is useful for synthesis of clusters in solution. The oxidation states of the aluminum clusters were determined from BEs measured in XPS Al 2p spectra. The stability and reactivity of Al₁₃ − were compared with those of other aluminum clusters. The particularly high stability of Al₁₃ − was demonstrated by air exposure and through theoretical calculations. In addition, the aluminum clusters in solution exhibited redox behavior derived from Al₁₃−/Al₁₃ in DPAG₄. The solution-phase synthesis of Al₁₃− superatoms has a significant role for the experimental development of cluster science.

Discussion
The fabricated clusters were protected by the DPA template. It is useful for synthesis of clusters in solution, however, the features have possibility to be different from the inherent character. Considering the isolation of clusters from the DPAs, stability of the clusters is a crucial factor.

In conclusion, aluminum clusters, including the superatomic Al₁₃− species, were fabricated in solution by utilizing the controlled metal assembly of DPA templates. To the best of our knowledge, this is the first example of the synthesis of the benchmark Al₁₃− superatom in solution. The Al₁₃− cluster was identified by mass spectrometry and STEM measurements. In addition, specific features of the Al₁₃− in the XPS, oxidation reaction and electrochemical measurement demonstrated its successful synthesis in solution. The oxidation states of the aluminum clusters were determined from BEs measured in XPS Al 2p spectra. The stability and reactivity of Al₁₃− were compared with those of other aluminum clusters. The particularly high stability of Al₁₃− was demonstrated by air exposure and through theoretical calculations. In addition, the aluminum clusters in solution exhibited redox behavior derived from Al₁₃−/Al₁₃ in DPAG₄. The solution-phase synthesis of Al₁₃− superatoms has a significant role for the experimental development of cluster science.

Methods
Materials. pyDPAG₄ and DPAG₄ were synthesized by a convergent method. The fourth-generation phenylazomethine dendrons were connected to the core molecules. The detailed procedures are described in previous reports. AlCl₃ (ultra dry) was purchased from Alfa Aesar, a Johnson Matthey Company. Dehydrated THF (stabilizer-free) was obtained from Kanto Chemicals. Sodium metal, benzophenone and tetrabutylammonium hexafluorophosphate were obtained from Wako Pure Chemical Industries, Ltd. and TCI Co., Ltd.

Assembly of AlCl₃ in pyDPAG₄. The experiment was conducted in an N₂-filled glove box. A THF solution of AlCl₃ (2.60 mM) and pyDPAG₄ (4.01 μM) was prepared. An appropriate equivalent of AlCl₃ solution was then added to the DPAG₄ solution. Complexation between AlCl₃ and DPAG₄ was monitored using UV–Vis spectroscopy.

Assembly of AlCl₃ in DPAG₄. The experimental procedure was conducted in the same way as that with pyDPAG₄. The concentrations of AlCl₃/THF and DPAG₄/THF were 5.41 mM and 5.06 μM, respectively.

Fabrication of aluminum clusters. pyDPAG₄ was used for the Al₁₃− cluster, and DPAG₄ was used for the Al₆, Al₁₂, Al₁₄, Al₂₈, and Al₆₀ clusters. The inferior number (n) of Alₙ reflects the added equivalent of the AlCl₃ in the DPAs.
Dendrimer solutions in THF (3–8 μM) were prepared in an Ar-filled glove box. The required equivalents of AlCl₃ in THF were added to each dendrimer solution to prepare the AlCl₃-complex dendrimer. The benzophenone ketyl radical was prepared by the addition of an excess amount of sodium metal into the solution of benzophenone in THF (0.3 M). The THF solution of benzophenone ketyl radicals (1 mL) was added to the prepared AlCl₃-complex solution (1 mL) to fabricate aluminum clusters at the DPAs. Samples for STEM measurements were prepared by the use of 60 equivalents of benzophenone to AlCl₃ for reduction.

**Characterization.** UV–Vis spectra were recorded at 20 °C using Shimadzu UV-3600 and UV-3100PC spectrometers with a quartz cell having an optical length of 1 cm. XPS spectra were measured with a PHI 5000 VersaProbe (Ulvac-Phi, Inc.). Al Kα (15 kV, 25 W) radiation was used as the X-ray source. The beam was focused on a 100 μm² area. Samples were sputtered with an Ar ion gun to remove the oxidized surface prior to the measurements. The spectra were analyzed with the MultiPak software (Physical Electronics), and were standardized according to the Au 4f₇/₂ peak at 84.0 eV. Background subtract, peak smoothing and fitting were performed.

**Fig. 3** Change in the XPS Al 2p peaks upon air exposure. The graphs show the spectral change for the Alₙ clusters. a n = 13, b n = 12, and c n = 14. The exposure times were 1–2 min, 5 h, 24–25 h, and 46–48 h. The vertical axes are normalized intensity. The spectra for Al₁₂ and Al₁₄ samples could be fitted by two peaks derived from the initial reduction peak (blue dashed line) and oxidation peak (red dashed line). Ratio of the oxidized (red squares) to the as-prepared (blue diamonds) peaks for d Al₁₂ and e Al₁₄ samples.

**Fig. 4** Oxidation waves of the fabricated anionic aluminum cluster samples. They correspond to Alₙ clusters. a n = 12 (6 μM), b n = 13 (4 μM), and c n = 14 (4 μM) in THF. The potential sweep was started from −1.7 or −1.8 V (open circuit potentials) at the scan speed of 0.025 V/s. Black bars indicate 10 μA in a and c, and 5 μA in b. NaPF₆ was used as an electrolyte (0.05 M).
used to estimate peak areas. STEM images were obtained using a transmission electron microscope (JEOL, ARM-200F) and the HAADF method. STEM samples were deposited on a super high-resolution carbon film with a Cu mesh (Okenoshi Co.). Cyclic voltammetry was performed using a BAS ALST7050 analyzer. A glassy carbon disc electrode and platinum wire were used as the working and counter electrodes, respectively. An Ag+/Ag (0.01 M AgNO3 in 0.1 M Bu4NCIO4/acetonitrile) electrode was used as the reference electrode. NaPF6 (0.05 M) was used as the electrolyte.

Calculations. Geometry optimizations for Aln, A12n, Al13n, and Aln in anionic and neutral states were carried out with the B3LYP/P6–31 G(d, p) level of theory. The employed structures were based on the past literatures. Vibrational analyses were also conducted for all the optimized structures and no imaginary frequency appeared. For the theoretical interpretation of the observed XPS data, Mulliken and natural population analyses were performed. In addition, the 2p orbital levels were calculated by single-point energy calculations with the GB3LYP/DZP-DKH level of theory within fourth-order Douglas–Kroll–Hess relativistic approximation, which is aimed at the incorporation of spin-orbit coupling. The diameter of Al13n+ was estimated using the Monte-Carlo method, in which the molecular volume was defined as the volume inside a contour with a density of 10–6 electrons/Bohr3. All the calculations were conducted with Gaussian 16.

Data availability. The data that support the findings of this study are available from the corresponding author upon reasonable request.

Received: 16 February 2017 Accepted: 15 November 2017
Published online: 11 December 2017

References

1. Khanna, S. N. & Jena, P. Atomic clusters: building blocks for a class of solids. Phys. Rev. B 51, 13705 (1995).
2. Reveles, J. U. et al. Designer magnetic superatom. Nat. Chem. 1, 310–315 (2009).
3. Yang, H. et al. Ligand-stabilized Au12Cl6− and bimetallic nanoclusters: ligand engineering to control the exposure of metal sites. J. Am. Chem. Soc. 135, 9568–9571 (2013).
4. Yang, H. et al. All-thiol-stabilized Au46 and Au12Al8 nanoparticles with single-crystal structures. Nat. Commun. 4, 2422 (2013).
5. de Heer, W. A. The physics of simple metal clusters: experimental aspects and simple models. Rev. Mod. Phys. 65, 611–676 (1993).
6. Li, X., Wu, H., Wang, X.-B. & Wang, L.-S. s-p Hybridization and electron shell structures in aluminum clusters: a photoelectron spectroscopy study. Phys. Rev. Lett. 81, 1909–1912 (1998).
7. Bergeron, D. E., Castleman Jr, A. W., Morisato, T. & Khanna, S. N. Formation of Al13n+: evidence for the superhalogen character of Al13n. Science 304, 84–87 (2004).
8. Reveles, J. U., Khanna, S. N., Roach, P. J. & Castleman, A. W. Jr. Multiple valence superatoms. PNAS 103, 18405–18410 (2006).
9. Castleman, A. W. Jr. et al. From designer clusters to synthetic crystalline nanoassemblies. Nano Lett. 7, 2734–2741 (2007).
10. Reber, A. C., Khanna, S. N. & Castleman, A. W. Jr. Superatom compounds, clusters, and assemblies: ultra alkali motifs and architectures. J. Am. Chem. Soc. 129, 10189–10194 (2007).
11. Knight, W. D. et al. Electronic shell structure and abundances of sodium clusters. Phys. Rev. Lett. 52, 2141–2143 (1984).
12. Castleman, A. W. Jr. & Khanna, S. N. Clusters, superatoms, and building blocks of new materials. J. Phys. Chem. C 113, 2666–2675 (2009).
13. Dietz, T. G., Duncan, M. A., Powers, D. E. & Smalley, R. E. Laser production of supersonic metal cluster beams. J. Chem. Phys. 74, 6511–6512 (1981).
14. Jarrold, M. F., Bower, J. E. & Kraus, J. S. Collision induced dissociation of metal cluster ions: Bare aluminum clusters, Aln+, (n = 3–26). J. Chem. Phys. 86, 3876–3885 (1987).
15. Iturrioz, M. P., Lopez, M. J., Alonso, J. A. & Soler, J. M. Electronic and atomic structure of Na, Mg, Al and Pb clusters. Z. Phys. D 11, 163–174 (1989).
16. Gantefer, G., Gauza, M., Mieines-Broer, K. H. & Lutz, H. O. Photoelectron spectroscopy of jet-cooled aluminum cluster anions. Z. Phys. D 9, 253–261 (1988).
17. Leuchttner, R. E., Harms, A. C. & Castleman, A. W. Jr. Aluminum cluster reactions. J. Chem. Phys. 94, 1093–1101 (1991).
18. Leuchttner, R. E., Harms, A. C. & Castleman, A. W. Jr. Thermal metal cluster anion reations: behavior of aluminum clusters with oxygen. J. Chem. Phys. 91, 2753–2754 (1989).
Division, Technical Department, Tokyo Institute of Technology, for the STEM analysis. The XPS measurement was conducted at the Advanced Characterization Nanotechnology Platform of the University of Tokyo, supported by the "Nanotechnology Platform" of the Ministry of Education, Culture, Sports, Science and Technology (MEXT), Japan.

Author contributions
T.K. carried out the experiments except for theoretical calculations. N.H. conducted the calculations. T.K., N.H., T.I., and K.Y. wrote the manuscript. K.Y. supervised this work and project through discussing the concept with T.K.

Additional information
Supplementary Information accompanies this paper at https://doi.org/10.1038/s41467-017-02250-4.

Competing interests: The authors declare no competing financial interests.

Reprints and permission information is available online at http://npg.nature.com/reprintsandpermissions/

Publisher's note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2017