 EDGE ARTICLE

Cite this: Chem. Sci., 2018, 9, 676

Cage-templated synthesis of highly stable palladium nanoparticles and their catalytic activities in Suzuki–Miyaura coupling†

Li Qiu, Ryan McCaffrey, Yinghua Jin, Yu Gong, Yiming Hu, Hongliang Sun, Wounjhang Park and Wei Zhang †

We report the controlled synthesis of small palladium nanoparticles (PdNPs) with narrow particle size distribution (1.8 ± 0.2 nm) using an organic molecular cage as a template. The well-defined cage structure and thioether anchoring groups inside the cavity are critical for the formation of narrowly distributed PdNPs, offering a confined organic molecular environment and guiding PdNP nucleation and growth. The resulting encapsulated PdNPs are resistant to agglomeration and stable in solution exposed to air at room temperature. When provided with a protective cage shell with minimum surface coverage, such PdNPs are capable of catalyzing organic reactions, showing high catalytic activity in Suzuki–Miyaura coupling reactions.

Results and discussion

Our group has been interested in exploring shape-persistent 3-D organic molecular cages as templates to control encapsulation and growth of nanoparticles. The recent advent of dynamic covalent chemistry (DCvC)22–28 has enabled facile large-scale synthesis of shape-persistent 3-D organic molecular cages, which have attracted tremendous attention as viable candidates for applications such as carbon capture and fullerene separation.29–34 With well-defined and permanently rigid pore structures, such a cage template35–37 can offer a protecting shell with minimum surface coverage, which would be advantageous compared to conventional small organic ligands or macromolecular ligands that form thick, insulating layers on the nanoparticle surface.38,39 Herein, we report robust organic cage-templated synthesis of stable, highly soluble, and narrowly distributed palladium nanoparticles (PdNPs). Their catalytic application in Suzuki–Miyaura cross-coupling reactions was also explored.

Introduction

Metal nanoparticles (NPs) have been widely applied in various disciplines of modern sciences, including catalysis,1–4 diagnostic imaging,5–8 sensing,5–8 magnetic recording,12 electronics13 and optics.14 These materials often exhibit particular physical and chemical characteristics arising from their small size and high surface-to-volume ratio, which are distinct from those of bulk materials.15 The properties of metal NPs highly depend on their size, shape and composition.16–17 thus the synthesis of narrowly distributed particles of a specific structure and composition has become an important research area in nanoscience. Various solution phase methods have been developed for the synthesis of nanoparticles, many of which are based on surface capping ligands and dendritic architectures.1,18–20 Recently, the synthesis of monodisperse nanoparticles has advanced through the use of closed-shell, hollow “ship-in-bottle” structures, such as protein cages, supramolecular DNA assemblies,22,23 and metal-coordination complexes,24 which confine the particles to a specific size and shape. However, the size-controllable preparation of small and narrowly dispersed nanoparticles remains challenging.

†Electronic supplementary information (ESI) available: Detailed experimental materials, general synthetic procedures, TEM images, and spectral characterization data. See DOI: 10.1039/c7sc03148c
††These authors contributed equally.
PdCl$_2^{2-}$ was transferred to the organic phase. A solution of cage 3a (1 equiv.) in CH$_2$Cl$_2$ was added to the above biphasic mixture and stirred for 45 minutes. The orange-red color deepened and the mixture was subsequently reduced with an aqueous solution of sodium borohydride (190 equiv., rt) resulting in a dark brown organic phase with no precipitation, which indicates the efficient reduction of Pd$^{2+}$ and further stabilization of PdNPs by cage molecule 3a. After extraction using CH$_2$Cl$_2$, the resulting PdNP@3a solution was dried over sodium sulfate, precipitated out from ethanol, and characterized by UV-vis, $^1$H NMR (DOSY), HR-TEM and energy-dispersive X-ray spectroscopy (EDS).

The UV-vis absorption spectra of the solution before and after reduction are shown in Fig. 1b. In the absence of cage 3a, the absorption spectrum of tetraethylammonium tetra-chloropalladate(II) in CH$_2$Cl$_2$ shows absorption peaks at 250 nm and 320 nm arising from ligand-to-metal charge-transfer transitions (Pd$^{II}$, red line, Fig. 1b). A transfer agent TOAB to maintain good solubility in benzene). Complete reduction of PdCl$_4^{2-}$ to Pd$^0$ and the formation of PdNPs was supported by the absence of bands from 300–500 nm (PdNP@3a, green line, Fig. 1b), which correlates well with results previously reported in the literature.‡‡

The diameter and size distribution of the resulting PdNP@3a were then analyzed by TEM micrographs (Fig. 1c, S3‡). A solution of PdNP@3a in CH$_2$Cl$_2$ was drop cast onto carbon-coated 300 mesh copper grids (CF300-Cu) and allowed to air dry before the measurements. The TEM image (Fig. 1c) shows well-dispersed PdNPs with an average size of 1.8 nm (over 500 particles counted), which matches well with our computational models showing an internal cavity size of 1.8–2.1 nm (Fig. 1a and d). The formation of such small PdNPs agrees well with the absence of a plasmon peak in the UV-vis spectrum.

Energy-dispersive X-ray spectroscopy (EDS) (Fig. S5†) also confirmed unambiguously the presence of metallic palladium. MALDI-TOF and ESI-MS spectra of PdNP@3a show the presence of various multiply-charged Pd$_{m}@$3a, Pd$_{12}@$3a, Pd$_{18}@$3a, Pd$_{23}@$3a, and Pd$_{16}@$3a, supporting the complexation of PdNPs with the cage. These PdNP@3a complexes are stable and highly soluble in common organic solvents.

Consistent with our previous findings on AuNP@3a,‡† substantial broadening and shifting of not only the protons of the thioether group but also all of the aromatic protons of the cage skeleton were observed in the $^1$H NMR spectra (Fig. S2†) of the above PdNP@3a complex (with a small amount of phase-transfer agent TOAB to maintain good solubility in benzene). This indicates that the cage shell is likely to be tightly wrapped around the PdNP and experiences restricted mobility. $^1$H diffusion-ordered spectroscopy (DOSY) NMR performed on both the free cage 3a and PdNP@3a under the same temperature and concentration (Fig. S6 and S7†) gave very similar diffusion coefficients of $(2.5 \pm 0.5) \times 10^{-10}$ m$^2$ s$^{-1}$ and $(2.4 \pm 0.5) \times 10^{-10}$ m$^2$ s$^{-1}$ respectively, which indicates the similar size and shape of the free cage and PdNP@3a. It therefore also supports the notion that PdNPs rest inside the cage cavity instead of aggregating on the cage surface.

It is expected that thioether groups inside the cage cavity provide a preferential nucleation site for Pd, and further deposition in the spatially confined cavity would provide PdNPs of a similar size to the cage. Additionally, encapsulation of PdNPs in isolated cavities would also prevent their agglomeration and improve the chemical and thermal stability. Control experiments in the absence of cage 3a led to the formation of a black precipitate (Fig. S4†), supporting the notion that the
presence of cage 3a, which contains six amino groups and three thioether groups, is necessary for the growth, size control, and stabilization of PdNPs. In order to understand exactly what structural requirements are necessary for PdNP growth, we performed additional control experiments in the presence of cage 3b, where the only structural difference is that the three thioethers are replaced with methyl groups. We again observed the complete precipitation of PdNPs. A further control experiment, in which only the thioether side piece 2a was used as the stabilizing ligand, also led to similar aggregation and precipitation, further suggesting that the closed cage and multidentate interaction involving thioether groups is necessary for PdNP-cage stabilization.

With the successful formation of stable PdNP@3a, we next explored the use of PdNP@3a in homogenous catalysis. The advances in novel catalytic technologies for more efficient chemical transformations have arguably been some of the most significant developments in the history of modern science. As compared to their bulk counterparts, nanoparticle-based catalysts often exhibit unique and enhanced catalytic properties due to their high surface area, size reduction, and shape variation. Since palladium-mediated catalysis has been widely applied and shown great synthetic power in current organic chemistry,\textsuperscript{25}–\textsuperscript{29} as a proof-of-principle, we chose to use one of the most popular carbon–carbon (C–C) bond formation reactions, the Suzuki–Miyaura cross-coupling reaction, to evaluate the catalytic activity of PdNP@3a. The coupling reactions were carried out using phenylboronic acid (1.5 equiv.) and PdNP@3a (0.01 equiv.) along with various aryl iodides and aryl bromides bearing a variety of functional groups. All reactions were performed in a mixture of toluene/H\textsubscript{2}O (10 : 1, v/v) using Na\textsubscript{2}CO\textsubscript{3} (3 equiv.) as a base. The catalytic activity of PdNP@3a was first investigated using phenylboronic acid and aryl iodides at 100 °C under microwave heating (Table 1, entries 1 and 2). The reaction progress was monitored via TLC and the crude reaction mixtures were analyzed by \textsuperscript{1}H NMR spectroscopy. Although conversion was good after 10–15 minutes of reaction, we found that increasing the reaction time to 30 minutes gave almost quantitative yields. To broaden the scope of the PdNP@3a catalyst, the same conditions were applied to a series of aryl bromide substrates using a temperature of 140 °C. We obtained the desired coupling products in almost quantitative yields in most entries. The analysis of the crude reaction mixture by \textsuperscript{1}H NMR shows little evidence of impurity formation during the coupling except in entry 5, which was seen to contain 1–2% biphenyl by-product.

As a comparison, the commercially available palladium (0) tetrakis catalyst Pd(PPh\textsubscript{4}) was used under the same conditions. We found the conversion to the desired products could only be achieved in yields of 73–85% (Table 1) after 30 min. Although we observed only a modest increase in overall yields using PdNP@3a, we found that a major advantage of this cage-based catalyst is its ability to be stored in solution under atmosphere over a period of several hours with no decrease in activity, thus highlighting its superb stability under ambient conditions. Such stability is remarkably superior to palladium tetrakis catalyst, which must be stored under inert atmosphere with very limited exposure to air. When both catalysts are stored in solution for 2.5 hours (open to air), the activity of PdNP@3a remained the same while the activity of Pd(PPh\textsubscript{4})\textsubscript{4} was decreased to 40% (Table 1, entry 4), further corroborating the stability of the nanocatalyst.

In addition to Pd(PPh\textsubscript{4})\textsubscript{4}, three other commercially available palladium catalysts, Pd(PPh\textsubscript{3})\textsubscript{2}Cl\textsubscript{2}, Pd\textsubscript{2}(dba)\textsubscript{3} and Pd/C were employed for comparison. As shown in entries 1 and 6, while both Pd(PPh\textsubscript{3})\textsubscript{2}Cl\textsubscript{2} and Pd\textsubscript{2}(dba)\textsubscript{3} gave comparably high yields to PdNP@3a for iodotoluene, PdNP@3a showed higher activity for brominated substrates. When compared to the commercial nanoparticle Pd/C (5%), PdNP@3a exhibited a much better performance, not only with higher yields (99% vs. ~20%, entries 1 and 6) but also with a higher reaction rate: the reaction between 4-iodotoluene and phenylboronic acid catalyzed by PdNP@3a produced 4-methyl-1,1'-biphenyl with an 80% conversion after 6 min of reaction with a turnover frequency (TOF) of 800 h\textsuperscript{-1}, which is about 6 times greater than that of Pd/C catalysts (TOF: 130 h\textsuperscript{-1}) under the same conditions. Meanwhile PdNP@3a showed very good reproducibility for the reaction between phenylboronic acid and 4'-bromacetophenone, with a standard deviation (in reaction yield) of only 0.4% in five repeats (entry 6). The catalytic efficiency of PdNP@3a can be attributed to both its high solubility and largely unpassivated surface, which presumably allows for easy substrate diffusion to the nanoparticle surface.\textsuperscript{27} Although precipitation of palladium black was observed occasionally after removal of the reaction vessel from the microwave reactor (which is probably due to the high temperature breaking the bonding interaction between PdNP and the cage), interestingly, the recycled palladium catalyst could be reused at least 3 more times in the coupling reaction without loss of yield (Table S1), which supports the high reactivity of PdNP@3a.

As previously discussed, traditional synthetic methods for PdNPs rely heavily on classic organic supports (e.g. polymers,\textsuperscript{30} dendrimers,\textsuperscript{37–39} micelles,\textsuperscript{40} etc.) to prevent NP agglomeration at the colloidal stage. It has been reported that some dendrimer-stabilized PdNPs show good stability and excellent catalytic activity. For example, Astruc and co-workers demonstrated that PdNPs stabilized by dendritic nanoreactors have high catalytic activity (TOF = 4.5 × 10\textsuperscript{4} h\textsuperscript{-1}) toward the Suzuki–Miyaura reaction of 1,4-bromonitrobenzene and phenylboronic acid.\textsuperscript{30} However, in general, there is a compromise between the stability of the resulting NPs and the available surface accessibility for substrate activation and transformation.\textsuperscript{41,42} For instance, the El-Sayed group found that although G4 dendrimer-encapsulated PdNPs were more stable than polymer-stabilized PdNPs, the catalytic activity was in fact lower.\textsuperscript{44} In our cage-based supports, not only can the particle size be predetermined by the cage cavity size, but further stability and resistance to agglomeration can also be achieved via the well-defined closed architecture of the cage. This is in great contrast to dendritic architectures where increases in generation lead to steric crowding on the dendrimer periphery and subsequent lowering of the activity of the PdNP.\textsuperscript{44} Our study successfully demonstrates the use of organic cage molecules as suitable colloidal chemistry platforms for the controllable synthesis and stabilization of nanometer-sized catalytically active metallic nanoparticles. Due to their highly accessible,
well-defined surface morphology and long shelf-life in solution, cage-encapsulated nanoparticles show great promise as convenient and active catalysts.

**Conclusions**

In summary, we report the synthesis and stabilization of ~1.8 nm-sized PdNPs within the cavity of an organic cage molecule, which show excellent catalytic activity in the Suzuki–Miyaura reaction. The incorporation of metallic nanoparticles within the cavities of well-defined, discrete molecular cage molecules as novel catalytic supports is attractive given their stability and size programmability within the interior cage dimensions. Our cage-template approach will pave new avenues in the development of significantly improved or even novel catalysts for various chemical transformations through the use of different metallic nanoparticle centers. In addition, we envision that such an approach could provide a powerful platform for controlled growth of novel nanostructured materials, which can be used in a range of nanotechnologies, including nanocatalytic applications.

**Conflicts of interest**

There are no conflicts to declare.

**Acknowledgements**

We thank Dr Lily A. Robertson for her assistance with EDS and Garry Morgan for his assistance with TEM. The authors thank Army Research Office (W911NF-12-1-0581) for financial support.

---

**Notes and references**

1. **Metal nanoparticles for catalysis: advances and applications**, ed. F. Tao, In: RSC Catal. Ser., 2014; 17, RSC, 2014.
2. M. Zahnakiran and S. Ozbek, *Nanoscale*, 2011, 3, 3462–3481.
3. R. M. Crooks, M. Q. Zhao, L. Sun, V. Chechik and L. K. Yeung, *Acc. Chem. Res.*, 2001, 34, 181–190.
4. N. Toshima and T. Yonezawa, *New J. Chem.*, 1998, 22, 1179–1201.
5. D. Astruc, F. Lu and J. R. Aranaes, *Angew. Chem., Int. Ed.*, 2005, 44, 7852–7872.
6. N. T. Phan, M. Van Der Sluys and C. W. Jones, *Adv. Synth. Catal.*, 2006, 348, 609–679.
7. *From Bioimaging to Biosensors: Noble Metal Nanoparticles in Biodetection*, ed. L.-K. Chau and H.-T. Chang, Pan Stanford Publishing Pte. Ltd., 2012.
8. Y. I. Park, K. T. Lee, Y. D. Suh and T. Hyeon, *Chem. Soc. Rev.*, 2015, 44, 1302–1317.
9. E. C. Dreaden and M. A. El-Sayed, *Acc. Chem. Res.*, 2012, 45, 1854–1865.
10. A. S. Thakor, J. Jokerst, C. Zavaleta, T. F. Massoud and S. S. Gambhir, *Nano Lett.*, 2011, 11, 4029–4036.
11. P. Alivisatos, *Nat. Biotechnol.*, 2004, 22, 47–52.
12. J. P. Wang, *Proc. IEEE*, 2008, 96, 1847–1863.
13. A. Kamysnyh and S. Magdassi, *Small*, 2014, 10, 3515–3535.
14. K. L. Kelly, E. Coronado, L. L. Zhao and G. C. Schatz, *J. Phys. Chem. B*, 2003, 107, 668–677.
15. **Metal Nanoparticles: Synthesis, Characterization, and Applications**, ed. D. L. Feldheim and C. A. Foss Jr, Marcel Dekker, Inc., 2002.
16. A. R. Tao, S. Habas and P. Yang, *Small*, 2008, 4, 310–325.
