N-heterocyclic carbene-stabilized metal nanoparticles within porous organic cages for catalytic application

Tong Liu¹,†, Sha Bai¹,†, Le Zhang¹, F. Ekkehardt Hahn¹,2 and Ying-Feng Han¹,*

ABSTRACT
Tuning the surface-embellishing ligands of metal nanoparticles (NPs) is a powerful strategy to modulate their morphology and surface electronic and functional features, impacting their catalytic activity and selectivity. In this work, we report the design and synthesis of a polytriazolium organic cage PIC-T, capable of stabilizing PdNPs within its discrete cavity. The obtained material (denoted Pd@PCC-T) is highly durable and monodispersed with narrow particle-size distribution of 2.06 ± 0.02 nm, exhibiting excellent catalytic performance and recyclability in the Sonogashira coupling and tandem reaction to synthesize benzofuran derivatives. Further investigation indicates that the modulation of N-heterocyclic carbene sites embedded in the organic cage has an impact on NPs’ catalytic efficiency, thus providing a novel methodology to design superior NP catalysts.

Keywords: metal nanoparticle, porous organic cage, catalytic reaction, N-heterocyclic carbene

INTRODUCTION
In recent years, porous organic cages with intrinsic inner cavities and tunable surfaces have emerged as a new class of microporous material with promising applications in recognition [1], molecular separation [2] and catalysis [3]. Differently to other porous materials such as metal-organic frameworks and zeolites [4,5], porous organic cages exhibit intriguing features, including structural tunability, thermal and chemical stability, and unique solution processability, making them prominent candidates for encapsulating diverse metal nanoparticles (MNPs) within their nanoporous cavities [6–13]. MNPs possess high surface area/volume ratios with many active catalytic sites, giving them exceptional catalytic activity compared with their metal counterparts [14]. Although they have been demonstrated to be critical heterogeneous catalysts for a wide array of reactions, MNPs still suffer from difficulties related to agglomeration and consequent loss of catalytic activity during catalytic processes [15]. In this regard, porous organic cages offer protecting shells for the encapsulated MNPs with minimum coverage and greater surface accessibility within their uniform but flexible cavities.

Over the past few decades, N-heterocyclic carbenes (NHCs) have established themselves as one of the most potent ligands in many catalytic systems [16–19], as well as metal-NHC-stabilized MNPs [20–27]. However, the development of discrete 3D NHC-based architectures to encapsulate MNPs with superior catalytic performance is still in its infancy [28]. Recently, our group developed an efficient supramolecular strategy named the metal-carbene template approach (MCTA) to synthesize polyanilidazolium organic cages (PICA) [29–32]. Using this strategy, a new type of PIC material with different shapes and sizes can be readily tailored by the rational design of NHC precursors. PICs have good solubility and stability in most common solvents and have the potential to support MNPs [29]. More importantly, their in-situ-generated poly-NHC sites are essential in controlling the nucleation and growth of MNPs. The strong σ-donating ability of the NHCs further enhances the interactions between the shells (cages) and the MNPs [33], consequently leading to encapsulated MNPs within polycarbene cages (PCCs) with unique properties.

Compared to the imidazolium counterparts, replacing a carbon atom with an electronegative
Figure 1. Modulation of the NHC sites embedded in Pd@PCC-T (X = N) for tunable catalytic performance.

The binding energy values at 340.6 and 335.3 eV correspond well with the typical 1H NMR analysis proved the anchoring between Pd centers and 1,2,4-triazolin-5-ylidene and the formation of NHC species by the disappearance the characteristic proton of 1,2,4-triazolium as well as the broadening and shifting of the relevant protons of PIC-T (Fig. S3), as previously reported [8,30]. In addition, two-dimensional diffusion-ordered spectroscopy revealed that the as-synthesized Pd@PCC-T has a similar diffusion coefficient to the reported Au@PCC-T [32], which confirms the similar size and shape of Pd@PCC-T and Au@PCC-T (Fig. S3). It provides important evidence to support the idea that the formed PdNPs are entrapped within the cage. Finally, the Pd@PCC-T was isolated, after centrifugation, as a black powder. Remarkably, this material was well dispersed in methanol or ethanol and stable for six months without agglomeration or any significant color change under ambient conditions (Fig. S4).

The particle diameter and size distribution of Pd@PCC-T were analyzed by transmission electron microscopy (TEM) micrographs. The image presented well-dispersed PdNPs with a narrow size distribution of 2.06 ± 0.02 nm (Fig. 2b), matching well with the optimized cage structure (Fig. S5). This indicates that the cage template can offer a protecting shell for the size-controlled synthesis of PdNPs and the majority of PdNPs are probably anchored within the cage cavities [9]. The scanning electron microscopy (SEM) image also depicted a rough morphology consisting of granular particles of as-prepared Pd@PCC-T (Fig. S6); the palladium loading is estimated to be ~22 wt%, determined by inductive coupled plasma mass spectrometry (ICP-MS) [6]. Additional energy dispersive X-ray elemental mapping analysis showed that the formed PdNPs were highly distributed inside the cage cavities of PCC-T (Fig. S7). The survey X-ray photoelectron spectroscopy (XPS) of Pd@PCC-T then confirmed the composition of Pd, C, N and O elements in the sample (Fig. S8).

Furthermore, a Pd 3d XPS spectrum was performed to identify the chemical states of Pd in Pd@PCC-T. The binding energy values at 340.6 and 335.3 eV correspond well with the typical
spin-orbit pairs of 3d_{5/2} and 3d_{3/2}, indicative of its zero-oxidation state (Fig. 2c). To gain further insight into the electronic coupling between PCC-T and PdNPs, N1s XPS analysis was performed (Fig. 2d). The binding energy assigned to the carbene N shifted from 401.6 eV to 400.7 eV after forming Pd@PCC-T, definitely demonstrating the anchoring of PdNPs by the organic cage [39,40]. Moreover, high-resolution TEM (HRTEM) images revealed PdNPs with a d-spacing of 0.22 nm, corresponding to the (111) lattice planes of zero-valent Pd (Fig. S9) [9,41]. The powder X-ray diffraction (PXRD) pattern further disclosed that the crystal form of Pd@PCC-T is a face-centered cubic (fcc) structure (Fig. S10).

To evaluate the role of the cage scaffold in the synthesis of PdNPs, controlled experiments were performed. Under standard conditions, immediate precipitation caused by the aggregation of the PdNPs was observed in the absence of the PIC-T template (Fig. S11). In the second case, only aggregated amorphous Pd⁰ species were detected when the monodentate 1,2,4-triazol-5-ylidene-derived NHC ligands were used as templates (Fig. S12). These results further demonstrate the crucial role of the cavity confining effect in the nucleation, controlled synthesis and stabilization of PdNPs. In addition, Pd@PCC-T was stable even after exposure to 1-dodecanethiol (DDT, C₅ = 5 mM in methanol) for 24 h, according to the TEM, XPS and PXRD analyses (Fig. S14). This synthetic strategy was further applied in preparing poly-imidazolium cage (PIC-I) stabilized NPs, Pd@PCC-I. This material, characterized in a similar manner by XPS, XRD, TEM and ICP-MS (Fig. S13), exhibited comparative particle sizes (2.06 ± 0.01 nm) and Pd loading (∼21 wt%) to Pd@PCC-T.

**Catalytic performance investigation into the Sonogashira reaction**

To date, PdNPs have been demonstrated as superior catalysts in a variety of organic reactions [42]. For the purpose of evaluating the catalytic activity of...
Table 1. The substrate scope of Pd@PCC-T catalyzed Sonogashira reaction.\(^a\)

| Entry | X   | R\(_1\) | R\(_2\) | Time (h) | Product | Yield (%)\(^b\) |
|-------|-----|---------|---------|----------|---------|------------------|
| 1     | Br  | H       | H       | 6        | 3a      | 99               |
| 2     | Br  | H       | 4-CH\(_3\) | 6        | 3b      | 99               |
| 3     | Br  | H       | 4-OCH\(_3\) | 6        | 3c      | >99              |
| 4     | Br  | H       | 4-F     | 6        | 3d      | 90               |
| 5     | Br  | H       | 4-Cl    | 6        | 3e      | 92               |
| 6     | Br  | H       | 4-CF\(_3\) | 6        | 3f      | 89               |
| 7     | Br  | H       | 4-C(CH\(_3\))\(_3\) | 6        | 3g      | 97               |
| 8     | Br  | H       | 4-pyridyl | 6        | 3h      | 92               |
| 9     | Br  | H       | 3-F     | 6        | 3i      | 93               |
| 10    | Br  | H       | 3-OCH\(_3\) | 6        | 3j      | 96               |
| 11    | Br  | H       | 4-OCH\(_3\) | 6        | 3k      | 95               |
| 12    | Br  | 4-Cl    | 4-OCH\(_3\) | 6        | 3l      | 88               |
| 13    | I   | H       | H       | 1        | 3a      | >99              |
| 14    | I   | H       | 4-OCH\(_3\) | 1        | 3c      | >99              |
| 15    | I   | H       | 4-CF\(_3\) | 1        | 3f      | 96               |
| 16    | I   | H       | 4-C(CH\(_3\))\(_3\) | 1        | 3g      | >99              |
| 17    | I   | H       | 4-pyridyl | 1        | 3h      | 97               |
| 18    | I   | 4-Cl    | 4-OCH\(_3\) | 1        | 3l      | 98               |

\(^a\)Reaction conditions: aryl halide (2.5 mmol), phenylacetylene (3.0 mmol), NEt\(_3\) (7.5 mmol), DMF (5.0 mL) and Pd@PCC-T (0.5 mol% Pd, 6.0 mg).

\(^b\)Yield determined by gas chromatography (GC) analysis.

Catalytic performance investigation into the tandem reaction

The benzofuran motif is a key structural unit in various biologically active natural products and pharmaceutical compounds, such as amurenensin H, anigopreissin A, amiodarone and so on [45]. The synthesis of these benzofuran derivatives, therefore, has attracted a lot of attention. Among the different methodologies, domino Sonogashira alkynylation coupling has been established as a green approach with high atom economy. Encouraged by the above results, the Pd@PCC-T was further used to synthesize 2-arylbenzofuran derivatives, ubiquitous in regular prescription and potential new drugs [46]. Preliminary studies focused on the reaction between 2-iodophenol and phenylacetylene as benchmark substrates. After screening a couple of reaction parameters (Supplementary Data, Table S3), the reaction carried out in the presence of Pd@PCC-T (0.5 mol% Pd) with Cs\(_2\)CO\(_3\) as a base provided the best result (6a, 98% yield) via a one-pot method (Table S3, entry 5).

Although the size-dependent catalytic activity of NPs has been intensively investigated, the functional sites modulation of organic cage supports to tailor the catalytic performance of encapsulated metal NPs has been seldom explored [47]. To investigate this issue, Pd@PCC-I, having a comparable particle size
**Table 2.** The tandem reaction for the synthesis of benzofuran derivatives catalyzed by different Pd catalysts.3

| Entry | Catalyst                  | Co-catalyst | Yield (%) |
|-------|---------------------------|-------------|-----------|
| 1     | Pd@PCC-T                  | /           | 98        |
| 2     | Pd@PCC-I                  | /           | 61        |
| 3     | PdCl2                     | /           | 22        |
| 4     | Pd/C                      | /           | 11        |
| 5     | [Ph3P]2PdCl2              | /           | 0         |
| 6     | [Ph3P]2PdCl2              | CuI         | 28        |
| 7     | Pd(OAc)2                 | /           | 32        |

*Reaction conditions: 2-iodophenol (2.5 mmol), phenylacetylene (3.0 mmol), Cs2CO3 (7.5 mmol), DMSO (6.0 mL) and Pd catalyst (0.5 mol% Pd). Yield determined by GC analysis.*

**Figure 3.** (a) Time course of the formation of 2-phenylbenzofuran by using Pd@PCC-T and Pd@PCC-I respectively. (b) Recyclability of the Pd@PCC-T catalyst in the tandem reaction.

and Pd loading to Pd@PCC-T but different NHC sites, was employed as a catalyst (Table 2, entry 2). Surprisingly, the yield of the model tandem reaction significantly decreased from 98% to 61% under identical conditions. The time-dependent reaction profiles further reveal that Pd@PCC-T exhibits much better catalytic activity in the overall conversion and reaction rate (4 h, 98%), while Pd@PCC-I only affords 73% yield even after a lengthened reaction time of 10 h (Fig. 3a). Thus, the cage-stabilized PdNPs do greatly enhance the catalytic efficiency in this reaction, benefitting from subtle modulation of NHC sites. For comparison, several commonly used Pd catalysts were also tested under the established conditions. Strikingly, the yields decreased from 98% for Pd@PCC-T to 22%, 11% and 0% for PdCl2, Pd/C and [Ph3P]2PdCl2, respectively (Table 2, entries 3–5). Generally, adding a Cu(I) co-catalyst could accelerate the Sonogashira reaction as previously reported [48]. Indeed, combing a co-catalyst CuI with [Ph3P]2PdCl2 leads to a 28% yield in this system (Table 2, entry 6); however, this is still inferior. Despite a slight increase with Pd(OAc)2 (32%, Table 2, entry 7), the yield is much lower than the Pd@PCC-I catalyst (61%). In contrast, the better catalytic performance of Pd@PCCs highlights the critical role of cage confinement in enhancing catalytic efficiency in the tandem reaction. After the reaction completed, the residue solution exhibited no catalytic activity after separation of Pd@PCC-T from the reaction mixture, excluding the leaching of the homogeneous metal species. These findings indicate that modulation of the chemical environment (electronic effect, steric effect etc.) around PdNPs [49,50] via ligand modification at the atomic level can regulate the interaction between metal NPs and substrates for sequential catalysis, more or less.

Pd@PCC-T also exhibited unique catalytic reactivity in synthesizing 2-arylbenzofuran derivatives bearing diverse functional groups (Fig. 4). The electron-withdrawing groups, that is, F (6b and 6c), Cl (6d) and CF3 (6f), and electron-donating groups, such as Me (6e), OMe (6i), gave the corresponding 2-arylbenzofuran with excellent yields (92%–98%). Furthermore, 2-iodophenol with different substituents (Cl and Me) were evaluated. It is interesting to note that 2-iodophenol containing an electron-donating group (Me; 6m, 6n, 6o) gave yields (90%–97%) that were slightly higher than those containing an electron-withdrawing group (Cl; 6j, 6k, 6l; 85%–92%), indicating that the electronic effect of the starting materials also slightly affects the yields.

**Catalyst recyclability**

The reusability of Pd@PCC-T was further evaluated. In each case, Pd@PCC-T was recovered by centrifugation at the end of the reaction and reused directly, without any activation. Indeed, no significant loss of catalytic activity was observed even after six successive cycles, affording high conversions for both coupling and tandem reactions (Figs 3b and S15). At the end of the six catalytic cycles, the catalysts were analyzed by TEM, XPS and PXRD, and no apparent changes in the size, morphology and structure were detected (Figs S16 and S17). In addition, the Pd content was determined to be <300 ppb by ICP-MS analysis in the residual reaction solution after the sixth cycling experiments (see Supplementary Data for details). The small amount of Pd leaching further suggests the adequate stabilization of ultrafine PdNPs within the confined cage cavity and the strong interactions between NPs and 1,2,4-triazol-5-ylidene-based NHC sites to suppress the Pd leaching and enhance the catalyst stability.

**CONCLUSION**

In summary, we presented our method to construct a soluble polytriazolium cage (PIC-T) with six 1,2,4-triazolium groups and a fascinating cavity capable...
Figure 4. The substrate scope of the Pd@PCC-T catalyzed tandem reaction for the synthesis of benzofuran derivatives. Reaction conditions: 2-iodophenol (2.5 mmol), phenylacetylene (3.0 mmol), Cs$_2$CO$_3$ (7.5 mmol), DMSO (6.0 mL) and Pd@PCC-T (0.5 mol% Pd, 6.0 mg). Yield determined by GC analysis.

METHODS

Synthesis of Pd@PCC-T and Pd@PCC-I

In a typical synthesis, PIC-T (20.7 mg, 0.008 mmol) was first dissolved in 10.0 mL CH$_3$CN (colorless), and a solution of palladium acetate in CH$_3$CN (18.0 mg, 0.08 mmol, red-orange, 3.0 mL) was then added with continuous stirring for 2 h. For the resulting mixture, a freshly prepared solution of NaBH$_4$ (30.3 mg in 3.0 mL of methanol, 0.8 mmol) was then added dropwise and the solution color immediately changed from red-orange to deep brown. After vigorous stirring for 3 h, the mixture was evaporated to dryness to afford a dark solid after washing with water and methanol. Finally, the solid was dried under vacuum overnight to obtain Pd@PCC-T as a black solid powder. This method was extended to synthesize Pd@PCC-I by using PIC-I as a precursor.

The general procedure of the catalytic Sonogashira reaction

Aryl halides (2.5 mmol), phenylacetylene derivatives (3.0 mmol), NEt$_3$ (7.5 mmol, 758.9 mg), Pd@PCC-T (0.5 mol% Pd, 6.0 mg) and DMF (5.0 mL) were added into a 50 mL flask under air atmosphere and heated to 40°C. After completion of the reaction, distilled water was poured into the reaction mixture, which was then extracted by dichloromethane. The organic layer was then collected, washed with brine and dried over anhydrous MgSO$_4$. The reaction yield was monitored by analyzing the sample with GC analysis. For examination of the reusability of Pd@PCC-T, the heterogeneous mixture was centrifugated after the reaction and the recovered catalyst was then reused for the subsequent reaction with fresh substrates and solvent.

The general procedure of the catalytic tandem reaction for synthesizing 2-arylbenzofuran derivatives

A 50 mL flask was charged with Pd@PCC-T (0.5 mol%, 6.0 mg), 2-iodophenol derivatives (2.5 mmol), phenylacetylene derivatives (3.0 mmol), Cs$_2$CO$_3$ (7.5 mmol, 2.4 g) and DMSO (6.0 mL). After stirring the mixture under air atmosphere at 60°C for 4 h, the product was obtained through a similar process to the Sonogashira reaction and the reaction yield was then monitored by analyzing the sample with GC analysis. The reusability of Pd@PCC-T was continuously evaluated.

Catalytic performance investigation into different Pd catalysts

A 50 mL Schlenk tube was charged with the Pd catalyst (0.5 mol%), 2-iodophenol (2.5 mmol), phenylacetylene (3.0 mmol), Cs$_2$CO$_3$ (7.5 mmol, 2.4 g) and DMSO (6.0 mL). After stirring the mixture under standard conditions, the reaction yield was monitored by analyzing the product with GC analysis.

SUPPLEMENTARY DATA

Supplementary data are available at NSR online.
FUNDING

This work was supported by the National Natural Science Fund for Distinguished Young Scholars of China (22052107), the National Youth Top-Notch Talent Support Program of China, the Key Science and Technology Innovation Team of Shaanxi Province (2019TD-007 and 2019JLZ-02) and the FM&EM International Joint Laboratory of Northwest University.

AUTHOR CONTRIBUTIONS

Y.-F.H. conceived this study and designed the experiments. T.L. performed material synthesis and characterization with S.B. and L.Z.’s assistance. Y.-F.H., S.B., F.E.H. and T.L. analyzed the data and co-wrote the manuscript. All authors discussed the results and reviewed the manuscript.

Conflict of interest statement. None declared.

REFERENCES

1. Wang H, Jin Y and Sun N et al. Post-synthetic modification of porous organic cages. Chem Soc Rev 2021; 50: 8874–86.
2. Tozawa T, Jones JTA and Swamy SI et al. Porous organic cages. Nat Mater 2009; 8: 973–8.
3. Bai S, Wang YY and Han YF. Template synthesis of metal nanoparticles within organic cage. Chin J Chem 2019; 37: 1289–90.
4. Hasell T and Cooper AI. Porous organic cages: soluble, modular and molecular pores. Nat Rev Mater 2016; 1: 16053.
5. Hua M, Wang S and Gong Y et al. Hierarchically porous organic cages. Angew Chem Int Ed 2021; 60: 12490–7.
6. Mondal B and Mukherjee PS. Cage encapsulated gold nanoparticles as heterogeneous photocatalyst for facile and selective reduction of nitroarenes to azo compounds. J Am Chem Soc 2018; 140: 12592–601.
7. McCaffrey R, Long H and Jin Y et al. Template synthesis of gold nanoparticles with an organic molecular cage. J Am Chem Soc 2014; 136: 1782–5.
8. Yang X, Sun JK and Kitta M et al. Encapsulating highly catalytically active metal nano-clusters inside porous organic cages. Nat Catal 2018; 1: 214–20.
9. Mondal B, Acharya K and Howlader P et al. Molecular cage impregnated palladium nanoparticles: efficient, additive-free heterogeneous catalysts for cyanation of aryl halides. J Am Chem Soc 2016; 138: 1709–16.
10. Qiu L, McCaffrey R and Jin Y et al. Cage-templated synthesis of highly stable palladium nanoparticles and their catalytic activities in Suzuki–Miyaura coupling. Chem Sci 2018; 9: 676–80.
11. Zhang Y, Xiong Y and Ge J et al. Porous organic cage stabilised palladium nanoparticles: efficient heterogeneous catalysts for carbonylation reaction of aryl halides. Chem Commun 2018; 54: 2786–9.
12. Chen GJ, Xin WL and Wang JS et al. Visible-light triggered selective reduction of nitroarenes to azo compounds catalysed by Ag@organic molecular cages. Chem Commun 2019; 55: 3586–9.
13. Ichijo T, Sato S and Fujita M et al. Size-, mass-, and density-controlled preparation of TiO2 nanoparticles in a spherical coordination template. J Am Chem Soc 2013; 135: 6786–9.
14. Yue Q, Li J and Zhang Y et al. Plasmon-inspired nanoengineering of functional yolk-shell microspheres with magnetic core and mesoporous silica shell. J Am Chem Soc 2017; 139: 15486–93.
15. Sun Z, Sun B and Qiao M et al. A general chelate-assisted co-assembly to metallic nanoparticles-incorporated ordered mesoporous carbon catalysts for Fischer-Tropsch synthesis. J Am Chem Soc 2012; 134: 17553–60.
16. Peris E. Smart N-heterocyclic carbene ligands in catalysis. Chem Rev 2018; 118: 9888–10031.
17. Chernyshov VM, Denisova EA and Eremin DB et al. The key role of R-NHC coupling (R = C, H, heteroatom) and M-NHC bond cleavage in the evolution of M/NHC complexes and formation of catalytically active species. Chem Sci 2020; 11: 6957–77.
18. Kashin AS and Ananikov VP. Monitoring chemical reactions in liquid media using electron microscopy. Nat Rev Chem 2019; 3: 624–37.
19. Prima DO, Madiyeva M and Burykina JV et al. Evidence for ‘cocktail'-type catalysis in Buchwald–Hartwig reaction. A mechanistic study. Catal Sci Technol 2021; 11: 7171–88.
20. An YY, Yu JG and Han YF. Recent advances in the chemistry of N-heterocyclic-carbene-functionalized metal-nanoparticles and their applications. Chin J Chem 2019; 37: 76–87.
21. Sun N, Zhang ST and Simon F et al. Poly(3-hexylthiophene) functionalized with N-heterocyclic carbenes as robust and conductive ligands for the stabilization of gold nanoparticles. Angew Chem Int Ed 2021; 60: 3912–7.
22. Man RWY, Li CH and MacLean MWA et al. Ultrastable gold nanoparticles modified by bidentate N-heterocyclic carbene ligands. J Am Chem Soc 2018; 140: 1576–9.
23. He C, Wu QJ and Mao MJ et al. Multifunctional gold nanoparticles@imidazolium-based cationic covalent triazine frameworks for efficient tandem reactions. CCS Chem 2020; 2: 2368–80.
24. He C, Liang J and Zou YH et al. Metal-organic frameworks bonded with metal N-heterocyclic carbene for efficient catalysis. Natl Sci Rev 2022; 9: nwab157.
25. Asensio JM, Tricard S and Coppol Y et al. Knight shift in 13C NMR resonances confirms the coordination of N-heterocyclic carbene ligands to water-soluble palladium nanoparticles. Angew Chem Int Ed 2017; 56: 865–9.
26. Lara P, Rivada-Wheelaghan D and Conejero S et al. Ruthenium nanoparticles stabilized by N-heterocyclic carbene: ligand location and influence on reactivity. Angew Chem Int Ed 2011; 50: 12080–4.
27. Möller N, Rühl A and Lamping S et al. Stabilization of high oxidation state upconversion nanoparticles by N-heterocyclic carbenes. Angew Chem Int Ed 2017; 56: 4356–60.
28. Koy M, Bellotti P and Das M et al. N-heterocyclic carbenes as tunable ligands for catalytic metal surfaces. Natl Catalysis 2019; 4: 352–63.
29. Zhang L, Das R and Li CT et al. C3-symmetric assemblies from trigonal polycarbene ligands and M(III) ions for the synthesis of three-dimensional polyimidazolium cations. Angew Chem Int Ed 2019; 58: 13360–4.
30. Gan MM, Liu JQ and Zhang L et al. Preparation and post-assembly modification of metallosupramolecular assemblies from poly(N-heterocyclic carbene) ligands. Chem Rev 2018; 118: 9587–641.

31. Sun LY, Sinha N and Yan T et al. Template synthesis of three-dimensional hexakisimidazolium cages. Angew Chem Int Ed 2018; 57: 5161–5.

32. Gou XX, Liu T and Wang YY et al. Ultrastable and highly catalytically active N-heterocyclic-carbene-stabilized gold nanoparticles in confined spaces. Angew Chem Int Ed 2020; 59: 16683–9.

33. Hopkinson MN, Richter C and Schedler M et al. An overview of N-heterocyclic carbenes. Nature 2014; 510: 485–96.

34. Li Y, Yang T and Li N et al. Multistimuli-responsive fluorescent organometallic assemblies based on mesoionic carbene-decorated tetraphenylethene ligands and their applications in cell imaging. CCS Chem 2021; 3: 957–68.

35. Tenne M, Metz S and Münster I et al. Phosphorescent platinum(II) complexes based on C/A+/Lambda1 Aryltriazol-5-ylidenes. Organometallics 2013; 32: 6257–64.

36. Hollóczki O, Kelemen Z and Nyulászi L. On the organocatalytic activity of N-heterocyclic carbenes: role of sulfur in thiamine. J Org Chem 2012; 77: 6014–22.

37. Dröge T and Glorus F. Measure of all rings–N-heterocyclic carbenes. Angew Chem Int Ed 2010; 49: 6940–52.

38. Maji B, Breugst M and Mayr H. N-heterocyclic carbenes: organocatalysts with moderate nucleophilicity but extraordinarily high Lewis basicity. Angew Chem Int Ed 2011; 50: 6915–9.

39. Crudden CM, Horton JH and Ebralidze II et al. Ultra stable self-assembled monolayers of N-heterocyclic carbenes on gold. Nat Chem 2014; 6: 409–14.

40. Rühling A, Schaepe K and Rakers L et al. Modular bidentate hybrid NHC-thioether ligands for the stabilization of palladium nanoparticles in various solvents. Angew Chem Int Ed 2016; 55: 5856–60.

41. Wang X, Ruditskiy A and Xia Y. Rational design and synthesis of noble-metal nanoframes for catalytic and photonic applications. Natl Sci Rev 2016; 3: 520–33.

42. Feng Y, Shao Q and Huang B et al. Surface engineering at the interface of core/shell nanoparticles promotes hydrogen peroxide generation. Natl Sci Rev 2018; 5: 895–906.

43. Hamasaka G, Roy D and Tazawa A et al. Arylation of terminal alkynes by aryl iodides catalyzed by parts-per-million loading of palladium acetate. ACS Catal 2019; 9: 11640–6.

44. Fernández E, Rivero-Crespo MA and Dominguez I et al. Base-controlled Heck, Suzuki, and Sonogashira reactions catalyzed by ligand-free platinum or palladium single atom and sub-nanometer clusters. J Am Chem Soc 2019; 141: 1928–40.

45. Bhargava S and Rathore D. Synthetic routes and biological activities of benzo-furan and its derivatives: a review. Lett Org Chem 2017; 14: 381–402.

46. Ji G, Duan Y and Zhang S et al. Synthesis of benzo-furans from terminal alkynes and iodophenols catalyzed by recyclable palladium nanoparticles supported on N, O-dual doped hierarchical porous carbon under copper- and ligand-free conditions. Catal Today 2019; 330: 101–8.

47. Li L, Li Z and Yang W et al. Integration of Pd nanoparticles with engineered pore walls in MOFs for enhanced catalysis. Chem 2021; 7: 686–98.

48. Chinchilla R and Nájera C. The Sonogashira reaction: a booming methodology in synthetic organic chemistry. Chem Rev 2007; 107: 874–922.

49. Cai G, Ding M and Wu Q et al. Encapsulating soluble active species into hollow crystalline porous capsules beyond integration of homogeneous and heterogeneous catalysis. Natl Sci Rev 2020; 7: 37–45.

50. Unger Y, Meyer D and Strassner T. Blue phosphorescent platinum (II) tetracarbene complexes with bis(triazoline-5-ylidene) ligands. Dalton Trans 2010; 39: 4295–301.