Recent Advances in the Reactions of Cyclic Carbynes

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Review

The acyclic organic alkynes and carbyne bonds exhibit linear shapes. Metallabenzynes and metallapentalynes are six- or five-membered metallacycles containing carbynes, whose carbene-carbon bond angles are less than 180°. Such distortion results in considerable ring strain, resulting in the unprecedented reactivity compared with acyclic carbynes. Meanwhile, the aromaticity of these metallacycles would stabilize the ring system. The fascinating combination of ring strain and aromaticity would lead to interesting reactivities. This mini review summarized recent findings on the reactivity of the metal–carbon triple bonds and the aromatic ring system. In the case of metallabenzynes, aromaticity would prevail over ring strain. The reactions are similar to those of organic aromatics, especially in electrophilic reactions. Meanwhile, fragmentation of metallacarbynes might be observed via migratory insertion if the aromaticity of metallacarbynes is strongly affected.

In the case of metallapentalynes, the extremely small bond angle would result in high reactivity of the carbyne moiety, which would undergo typical reactions for organic alkynes, including interaction with coinage metal complexes, electrophilic reactions, nucleophilic reactions and cycloaddition reactions, whereas the strong aromaticity ensured the integrity of the bicyclic framework of metallapentalynes throughout all reported reaction conditions.

Keywords: cyclic carbynes; reactivities; metallabenzynes; metallapentalynes

1. Introduction

The alkyne (carbon-carbon triple bond), which is a fundamental functional group in organic chemistry, is involved in a large number of reactions in organic chemistry [1–5]. Carbyne complexes, i.e., transition metal complexes with metal-carbon triple bonds, can be described as analogs of alkynes, whereas a transition metal replaces one of the sp carbons. They have attracted considerable attention because of their remarkable features and their significance as catalysts or reagents for various types of organic transformations [5–11]. Part of the reactivities of carbynes paralleled that of their organic parents, such as nucleophilic reactions, electrophilic reactions, photochemistry, oxidation and reduction, reactions with chalcogenides, reactions with unsaturated organic substrates (cycloadditions) and substitution on the α carbon (Figure 1). However, the incorporation of metal into the triple would lead to fascinating properties, which were quite different from organic alkynes. For example, late-transition-metal carbyne species were applied as catalysts in alkyne metathesis or alkyne polymerization reactions. Carbynes might be coupled or reactive to other metal complexes to afford bi- or multi-metallic complexes. Ligand substitution is one of the fundamental reactions in organometallic chemistry, which is certainly involved in the reactions of carbynes. In addition, carbyne ligands might be cleaved from metal centers to give various organic products (Figure 1).
The alkyne moiety is normally linear due to the sp hybridization of acetylenic carbon. Ring strain would be raised from the introduction of alkyne moiety into an organic cycle. As a result, limited insights could be gained into the reactivity of organic cyclic alkynes due to the instability caused by the high ring tension [12,13]. The incorporation of a metal center into the acetylenic carbon would stabilize the strained cyclic carbon cycles containing alkynes. The cyclic carbynes obtained until now could be classified as six-membered metallabenzynes [14–21] A and five-membered metallapentalynes [22–24] B (Figure 2). In addition, the first metallapyridyne [25] was documented by Xia and coworkers, providing one precious example of cyclic carbynes.

![Figure 2. Strained metallacycles with carbyne bond.](image)

The difference between acyclic carbynes and cyclic carbynes might be due to the ring strain. The chemical properties of cyclic carbynes are summarized in Figure 3. Among the limited examples, the reaction types of cyclic carbynes are similar to those of acyclic carbynes except that in most cases the “migratory insertion” [26] would not stop at the carbene complexes and lead to the transformation of aromatic metallacycles into cyclopentadienes (Cp) complexes [27].

![Figure 3. Reactions of cyclic carbynes.](image)

Carbynes are usually recognized as important intermediates in many transition-metal-catalyzed organic reactions [28,29]. The thorough investigation of the reactivities and properties might provide a...
new perspective for the application of late-transition-metal carbyne species as prosperous catalysts in various organic reactions. The chemistry of metallabenzynes has been fully illustrated and nicely reviewed [14–21]. However, there is no review paper covering the entirety of cyclic carbynes (metallabenzynes and metallapentalynes). This account deals with them together for the first time and the scope has been limited to the reactivities of metallabenzynes and metallapentalynes, seeking to provide a general picture of reactivity of the metal–carbon triple bonds or the entire aromatic system.

2. Reactivity of Six-Membered Metallabenzynes

Metallabenzynes have been recognized as transition metal analogs of benzyne in which a C atom is replaced by an isolobal transition metal fragment. The chemical structure of metallabenzynes exhibits aromaticity, which would lead to the similar reactions occurring in their organic counterparts. On the other hand, the carbyne moiety (M≡C, metal-carbon triple bond) is reactive to several reagents and might involve regular reactions that normal acyclic carbynes would experience. The first metallabenzene was secured by Jia [30] and co-workers in 2001. Then, many interesting metallabenzynes were produced and well-characterized. (Figure 4) [31–40] Most of the metallabenzynes are osmabenzynes (1–6) [30–38]. Rhenabenzynes (7), which are present in very limited amounts, were reported by Jia’s group [39,40]. Moreover, only osmabenzynes and rhenabenzynes have been discovered to date.

![Figure 4. Metallabenzynes reported to date.](image)

The cationic osmabenzyne 8 was formed by treatment of metallabenzyne 1 [30] with two equivalent HBF₄ in wet dichloromethane. The anionic chloride ligand was replaced by the neutral water with the aid of Bronsted acid [41]. Similarly, the dicationic osmabenzyne 9 was prepared by the replacement of chloride ligands in 1 with 2,2′-bipyridine (bipy) in the presence of thallium triflate (TlOTf) [42]. The exchange of phosphate ligands was much easier, and no reagents and catalysts were needed. Excess PCy₃ in refluxing benzene would simply kick out the triphenylphosphine and result in neutral osmabenzyne 10 [43]. The transformations are depicted in Scheme 1.
2.1. Electrophilic Substitution Reaction

Trimethylsilyl group (TMS) was usually recognized as the “equivalent” of cationic hydrogen and labile to electrophiles in organic reactions. In the presence of excess tetrafluoroboric acid and water, metallabenzyne 1 [30] was converted to cationic metallabenzyne 11 via electrophilic desilylation and subsequent ligand exchange. Sodium chloride would provide anionic chloride ion to avoid ligand exchange to facilitate the formation of neutral desilylated osmabenzyne 12. Treatment of isolated 11 with NaCl also gave 12. Bromination of metallabenzyne 1 with excess elemental bromine readily provides brominated osmabenzyne 13, in which both the Me₃Si and Cl in 1 have been replaced by Br [41]. Nitrosation of metallabenzyne 1 was carried out in the presence of excess nitrosonium salt (NOBF₄) and sodium chloride at low temperature to furnish osmabenzyne 14 in which the SiMe₃ substituent at C4 was replaced by Br [41]. Nitration of metallabenzyne 1 took place at C4 to afford osmabenzyne 15 under similar reaction conditions except that nitronium salt (NO₂BF₄) was applied. Surprisingly, the biclyclic species 18 was isolated together with osmabenzyne 15. A plausible mechanism was rationalized. Initially, electrophilic substitution of NO₂⁺ produced osmabenzyne 15, which might undergo migratory insertion reaction and coordination of anionic oxygen atom to give intermediate 16. Subsequent single electron transfer generated the radical 17, which would abstract a hydrogen atom from solvent to give the osmium complex 18 [44]. The reactions of osmabenzyne 1 is summarized in Scheme 2.

Scheme 2. Electrophilic reactions of osmabenzyne 1.
It is well known that C-SiMe₃ bonds on an aromatic ring are more reactive than C-H bonds towards electrophiles. In order to further investigate the reactivity of metallabenzynes without SiMe₃ group, osmabenzyne 3f was subjected to electrophilic conditions (Scheme 3). Bromination and nitration were carried out under similar conditions and provided the identical brominated product 13 and mononitrated osmabenzyne 19, respectively [44]. Treatment of osmabenzyne 3f with CF₃SO₂D in the presence of sodium chloride produced 3,5-dideuterated osmabenzyne 20 [41], which indicates that the carbons of OsC or OsCH were not attacked by the acid under the protonation condition.

![Scheme 3. Electrophilic reactions of osmabenzyne 3f.](image)

Chlorination of osmabenzyne 1 and 3f with excess HCl/H₂O₂ furnished the same 3,5-dichlorinated product 21, while C₂ monochlorinated product 22 could be obtained when the ratio of AlCl₃/H₂O₂ was carefully controlled [44]. Noteworthy is that the carbyne moiety was attacked by oxygen atoms in the presence of hydrogen peroxide to afford metalla-oxirene fragments in the chlorination reactions (Scheme 4).

![Scheme 4. Chlorination reactions of osmabenzyynes 1 and 3f.](image)

The electrophilic substitution reactions of metallabenzynes resemble the organic arenes and the metal-carbon triple bonds stayed intact under most electrophilic conditions except when highly oxidative H₂O₂ was applied. These phenomena demonstrate the aromatic properties of metallabenzynes.

### 2.2. Nucleophilic Reaction

The carbyne moiety in metallabenzyne could be attacked by nucleophiles. The first nucleophilic reactions of metallabenzynes were explored by Jia (Scheme 5) [42]. In order to increase the electrophilicity of metallacyclic, the neutral osmabenzyne 1 was transformed to dicationic osmabenzyne 9 by ligand exchange, which was quite electrophilic and readily reacted with nucleophiles (Scheme 5). Water, a weak nucleophile, participated in the nucleophilic attack of the carbyne atom to generate metallaphenol 23, which precipitated from the solvent system (H₂O/THF = 3:4). Desilylation was also observed on the SiMe₃ adjacent to carbyne carbon, which might be attributed to the higher nucleophilicity of the...
Si-bound carbon. The metallaphenol 23 could be transformed to acyclic complex 24 in the presence of catalytic HBF$_4$. Protonation and enol-ketone transformation might give intermediate 25, which was deprotonated to afford the ketone form 26. A six-electron retro-electrocyclization reaction of 26 would furnish acyclic complex 24. Moreover, osmabenzyne could be transformed to acyclic complex 24 directly if better solubility (H$_2$O/THF = 1:15) and longer reaction time (two days) were provided. The combination of methanol and potassium carbonate not only provided methoxide to conduct the nucleophile addition to the carbyne carbon, but also removed the trimethylsilyl group adjacent to the carbyne carbon, resulting in the similar product 27. The hydride provided by NaBH$_4$ attacked the carbyne carbon of osmabenzyne to give the metallabenzene intermediate 28, which would undergo migratory insertion reaction and subsequent rearrangement to afford cyclopentadienyl complex 29. The regioselectivity of nucleophilic attack is C1 (carbyne carbon) in Scheme 5.

Xia and coworkers reported nucleophilic addition to metallabenzynes with different regioselectivity (C3) [36]. They prepared osmabenzyne 4 and subjected it to nucleophilic addition with stronger nucleophiles, such as ethyllithium (EtLi) or sodium methanethiolate (NaSMe). Not surprisingly, the carbyne carbon (C3) was attacked to generate isoosmabenzene 30 or 31. If primary amine was applied, it would kick out the chloride ligand to give the dicationic osmabenzyne 32, which is more electrophilic than 4. Hence, the nucleophilic attack of primary amine on C3 and subsequent rearrangement of isoosmabenzene 33 furnished the ring-opened product 34 or 35 (Scheme 6).
2.3. Migratory Insertion Reactions

Metallabenzenes and metallabenzynes usually exhibit rearrangement to the corresponding cyclopentadienyl complexes, which is the major decomposition pathway. The migratory insertion step involves reductive elimination, in which the two carbon atoms adjacent to metal undergo the coupling reaction to form cyclopentadienyl moiety. In 2007, Jia and Lin [37] discovered that reduction of complex 36 and subsequent cyclometallation gave hydrido osmabenzyne intermediate 37, which underwent hydride shift to attack the carbyne carbon to afford metallabenzene intermediate 38. Intermediate 38 would undergo migratory insertion reaction and subsequent rearrangement to afford cyclopentadienyl (Cp) complex 39 (Scheme 7).

Metallabenzenes could also undergo migratory insertion reactions to give carbene complexes. The substituent effect was illustrated by Jia and Lin. As depicted in Scheme 8, osmabenzyne bearing bulky substituents (t-butyl or 1-adamantyl) para to the metal center would slowly convert to the corresponding osmium carbene complexes 40a and 40b at room temperature in solution [38]. The steric hindrance of the bulky substituents might prevent the osmium carbene complexes from further transformation [45,46].
The ligand effect played an important role in the rearrangement of osmabenzyne 1 to the corresponding cyclopentadienyl complex 41, which was demonstrated by Jia [43] as shown in Scheme 9. Ligand exchange reaction of stable osmabenzyne 1 with Mo(CO)₆ would replace one phosphine ligand with carbon monoxide and deliver reactive osmabenzyne intermediate 42, which underwent migratory insertion involving the two metal-bonded carbons to give the carbene intermediate 43. Migratory insertion of the carbene into the Os-Cl bond in 43 with a subsequent rearrangement to η⁵ coordination would furnish Cp complex 41.

Recent, Xia and coworkers documented a series of five-membered cyclic metal carbyne complexes, i.e., osmapentalynes [47–51] and ruthenapentalynes [52]. The carbine-carbon bond angles in the metallapentalynes are around 130°, which are much smaller than those of the acyclic metal carbynes (180°). The large ring strain associated with extreme distortion of the metal carbyne unit lead to the high reactivity of the metal–carbon triple bond, which exhibits a unique performance.

3. Reactivity of Five-Membered Metallapentalynes

The metal-carbon triple bond possesses an “alkyne-like” character and can react with an external metal precursor to generate bimetallic adducts, in which osmium-carbon triple bond coordinated with the coinage metal center. Xia reported that osmapentalyne 44 [48] was treated with cuprous chloride to afford hetero bimetallic adduct 45, in which the chloride bonded to osmium also coordinated with copper atoms. As different silver or gold precursors were subjected to osmapentalyne 44, the corresponding hetero bimetallic adduct 46 or 47 was formed in a similar way. The interaction between cyclic osmacarbyne with coinage metal is weak, which is supported by the fact that in the presence of ligands (PPh₃ for 45 or (n-Bu)₄NCl for 46–47) the bimetallic complex readily dissociates and osmapentalyne 44 would be regenerated (Scheme 10) [53]. Ruthenapentalynes 48a–b readily reacted with CuCl to afford similar bimetallic complexes 49a and 49b, except that the chloride bonded to ruthenium did not coordinate with copper [52].
3.2. Electrophilic Reaction

Alkynes, which exhibit electrophilic properties, would generate alkenes upon treatment of electrophiles. Carbynes and metallapentalynes are both sensitive to acids. The acyclic carbynes might generally form carbenes, while the shift of metal–carbon triple bond would be observed in metallapentalynes. In most cases, the fused rings of metallapentalynes would be reserved, which might be attributed to the higher stability derived from the aromaticity.

Upon treatment of osmapentalynes $50a$–$c$ \[47\] with $\text{HBF}_4\cdot\text{H}_2\text{O}$ at room temperature, the metal–carbon triple bond could shift from one ring to another to generate osmapentalynes $51a$–$c$ \[54\]. The plausible mechanism was that the osmapentalyne was protonated to give the osmapentalene intermediate $52$ bearing a 16-electron osmium center, which was not stable enough according to the 18-electron transition metal rule. Subsequent elimination of H$_2$ would furnish osmapentalynes $51a$–$c$ with an 18-electron osmium center. Fortunately, the 16-electron osmapentalenes, complex $52a$ and $52d$, were captured and characterized by the reaction of osmapentalyne $50$ or $51$ with $\text{HBF}_4\cdot\text{Et}_2\text{O}$ or $\text{AlCl}_3$ (Scheme 11). The metal carbyne bond shift reaction of the ruthenapentalyne $48b$ is similar to that of osmapentalynes, as is the mechanism \[52\].

The halogenation of metallapentalynes \[55\] would further demonstrate the electrophilicity of metallapentalynes. Treatment of osmapentalyne $51a$ \[47\] with ICl or elemental Br$_2$ led to the formation of the corresponding halogenated osmapentalene $55$ or $56$, respectively, which could be viewed as the first examples of metallaiodirenium and metallabromirenium ions (Scheme 12). The well-characterized bromocarbene complex $56$ or iodocarbene complex $55$ are similar to the generally proposed intermediates in the halogenation of alkynes, which would further demonstrate the “alkyne-like” properties of metal–carbon triple bond in metallapentalynes. The nucleophilic substitution reaction of $55$ with Bu$_4$NBr would generate brominated complex $56$. However, the application of Bu$_4$NCl to the halogenated osmapentalynes $55$–$56$ would result in the elimination of the halogen cation and the regeneration of osmapentalyne $51a$. Moreover, $51a$ also reacted with elemental selenium at $60^\circ\text{C}$, leading to the formation of Se-containing osmapentalene $57$ \[56\], which was regarded as the first example of $\sigma$-aromaticity dominating in an unsaturated Se-containing ring.
The reactions of metal carbynes and carbon–carbon triple bonds tended to provide cycloaddition intermediates or products. In contrast, Xia and coworkers documented a fantastic reaction between osmapentalynes and terminal alkynes in 2020, giving rise to acyclic addition products 59 and 61 (Scheme 13) [57]. High efficiency, regio- and stereoselectivity were observed, furnishing exclusively trans and anti-Markovnikov products in excellent yields. These reactions possess good functional tolerance for both alkynes and osmapentalynes and are easy to handle (air atmosphere and ambient conditions). The plausible mechanism is the electrophilic addition of cyclic carbyne by the terminal alkynes under the synergistic effect of protons on the basis of experimental observations as well as theoretical calculations. The application of the reactions provides an easy access to functionalized \( \pi - \pi \) conjugated systems, which exhibit great significance in functional materials.
3.3. Nucleophilic Reaction

The nonlinear distortion of the carbyne carbon angle of the osmapentalyne also facilitates the nucleophilic reaction. In contrast to the 16-electron osmapentalene 52 secured by protonation [54], 18-electron osmapentalenes 62 and 63 were achieved by the nucleophilic attack of the methanethioxide (MeO⁻) and the methanoxide (MeO⁻) anions towards the carbyne carbon of osmapentalyne 50a under the atmosphere of carbon monoxide. [54] Xia and coworkers reported the transformation of osmapentalyne to osmafulvenallene 64 (Scheme 14), which constituted the pioneering example of well-characterized metallafulvenallenes [58].

$$\text{[Os]} = \text{OsCl(PPh}_3\text{)}_2$$

Scheme 13. Reactions of alkynes with osmapentalynes.

Scheme 14. Nucleophilic reactions of osmapentalyne to form 62–64.

The rationale of the mechanism is depicted in Scheme 15. The combination of cesium carbonate and methanol would generate methanoxide, which conducted the first nucleophilic addition to a carbyne carbon, affording the metallapentalene intermediate 63. Subsequent coordination with a terminal alkyne led to the π-alkyne complex 65, which would form the vinylidene η¹-complex 66 via isomerization. The ω-carbon within the five-membered metallacycle was attacked by the second
methanol molecule. Sequential ring opening, protonation and coordination of the ester group furnished the final fused metallafulvenallene complex 64.

![Scheme 15. Proposed mechanism for the formation of 64.](image)

Very recently, oxygenation of osmapentalynes was reported by Xia’s group (Scheme 16). The first example was the oxygenation of the osmium carbolong complex ring [59]. Applying pyridine N-oxide as the oxidant, oxygen atoms were transferred to the metal–carbon triple bond in a nucleophilic reaction. Meanwhile, the labile aldehyde was oxidized to carboxylic acid, affording unprecedented OCOCO-type pentadentate chelates 69. The second example was the direct oxygenation of the metal center in the carbolong complex 70 [60]. The mixture of osmapentalyne 70 and excess sodium methoxide under oxygen atmosphere resulted in almost quantitative formation of osmapentene 71, in which the dioxygen was bound to the metal center. The plausible mechanism was depicted as the nucleophilic attack at carbyne carbon atom by sodium methoxide, which led to the reduction of metal center and dissociation of chloride, which facilitated the coordination and the following reduction of dioxygen.

![Scheme 16. Oxygenation of osmapentalynes.](image)

The direct attack of a free isocyanide on the carbyne carbon atom of the osmapentalyne 50a would lead to the formation of η2-iminoketenyl metallapentalene intermediates as well as the unprecedented metallaindene derivatives (Scheme 17). The steric hindrance of the bulky N-substituents tert-butyl or 1-adamantyl inhibited the bending at the isocyanide nitrogen, which prevented the formation of a η2-iminoketenyl structure and led to the osmapentene 72a–b. In comparison to bulky isocyanides, other less bulky isocyanides could afford the desired metallacyclopropenimine complex 73a–e, which were often regarded as the intermediates in nucleophile-induced carbyne-isocyanide coupling.
The same research group extended the metallapentalene and metallaindene system to ruthenium except that the η2-iminoketenyl species were not observed [62]. It is noteworthy that the addition of excess isocyanides at elevated temperature led to metallaindene derivatives, which represented the first metal-bridged polycyclic metallaaromatics [61]. Treatment of metallapentalyne 50a with the first isocyanide provided η2-iminoketenyl metallapentalene 73. The second isocyanide would coordinate with the osmium center to afford intermediate 74, which underwent sequential isocyanide insertion and coordination of a third isocyanide to generate intermediate 75. The exocyclic imine group could kick out one triphenylphosphine and coordinate with the osmium center, followed by subsequent aromatization to furnish metallaindene 77 (Scheme 18). The same research group extended the metallapentalene and metallaindene system to ruthenium except that the η2-iminoketenyl species were not observed [62].

The metal-carbon triple bond moiety in carbyne shifted complex 51a is also sensitive to nucleophilic attack (Scheme 19). Thus, products 78 and 79 were formed by treating osmapentalyne 51a with sodium hydrosulfide and benzylamine, respectively [47]. Noteworthy is that in the presence of aniline and
cesium carbonate, osmapentalyne 51a was transformed to the novel metal-bridged polycyclic aromatic system 80, in which the metal center is shared by three aromatic five-membered rings [63]. The reaction involved the nucleophilic addition of aniline to the carbyne moiety and subsequent C-H bond activation (oxidative addition), providing complex 80a. Treatment of osmapentalyne 51a and phenol produced the similar product 80b via a paralleled mechanism. A number of metal-bridged polycyclic aromatics, including different substitutions on the aromatic ring or larger fused-ring systems, were achieved with the same synthetic strategy [63–65].

![Scheme 19. Nucleophilic reactions of osmapentalyne 51a to form 78–80.](image_url)

In the presence of arene nucleophile X, the lactone-fused osmapentalyne 81 [66] was converted to metal bridgehead polycyclic π conjugate systems 82. Further transformations with t-BuOK or cyclohexyl isocyanide would lead to the neutral complex 83 or coordinated complex 84, respectively (Scheme 20). The delocalization of the above mentioned three metalla-aromatics could be used to switch the charge transport pathway of single-molecule junctions and thus tune the charge transport abilities significantly [67], which shed light on the potential applications of metal-bridged polycyclic aromatics in materials science.

![Scheme 20. Nucleophilic reactions of osmapentalyne 81 and further transformation.](image_url)

The ruthenium-carbon triple bond moiety in ruthenapentalynes showed nucleophilic reactivity. The reaction of ruthenapentalyne 48a with sodium thiophenoxide under carbon monoxide atmosphere resulted in the generation of the nucleophilic addition product ruthenapentalene 85. On the basis of the rich and unique reactivity of ruthenapentalynes, Xia and coworkers designed and carried out fantastic cascade cyclization reactions with bidentate nucleophiles (sodium cyanate or sodium dicyanamide) to afford annulation complexes 86 or 87, respectively (Scheme 21) [52].
Scheme 21. Nucleophilic reactions of ruthenapentalyne 43.

A plausible cascade cyclization mechanism is depicted in Scheme 22 [52]. The first cyanate ion initiated the nucleophilic attack at the carbyne carbon of 48a, leading to the formation of ruthenapentalene intermediate 88, which could be further attacked by the second cyanate ion to give intermediate 89. The Cl ligand dissociates from intermediate 89 to form the intermediate 90, which contains a vacant site at the ruthenium center. The subsequent coordination of the OCN group with the metal center could result in the final polycyclic complex 86. The mechanism to access the annulation product 87 might parallel that of the above cascade cyclization reaction, additionally, the nucleophilic attack was initiated by the N(CN)$_2$ ion.

Scheme 22. Proposed mechanism for the formation of 86.

3.4. Cycloaddition Reactions

The metallapentalynes exhibit rich metal carbyne reactivities due to the extreme strain in the fused five-membered ring containing a metal carbyne bond. The driving force for the cycloaddition reaction of metallapentalynes with the alkynes could be attributed to the release of the large ring strain in the five-membered ring of metallapentalynes. Xia and coworkers documented a number of cycloaddition reactions involving the metal-carbon triple bond in metallapentalynes [52,66,68–73].

Treatment of the cationic osmapentalyne 51a with propionic acid or ethoxyacetylene provided the planar products 91a and 91b, respectively, through unprecedented [2+2] cycloaddition reactions of alkynes with a late-transition-metal carbyne complex [68]. It is noteworthy that two classical anti-aromatic frameworks, cyclobutadiene and pentalene, were stabilized by incorporating one transition metal fragment into the anti-aromatic systems, thus providing novel aromatic metallacycles. Complexes 91a and 91b constituted the first example of [2+2] cycloaddition products formed between a late-transition-metal carbyne with alkynes. Further insertion was not observed in the presence of excess alkynes, thus preventing the formation of larger metallacycles. The reaction could be extended to early-transition-metal carbyne complexes. Treatment of ruthenapentalyne 93 with ethoxyacetylene furnished complex 94, which constitute the first [2+2] cycloaddition reaction of ruthenium carbyne complex with an alkyne [52]. The first [2+2] cycloaddition reaction of metallapentalyne 51a with nitrosoarenes (Ar–N=O) was documented by Xia et al. [69], affording well-characterized...
metallapentalenoxazetes 92a–c (Scheme 23). Faster reaction rate was observed provided that the
electron-donating substituents were introduced to the para-position of the nitrosophenyl ring (92b–c),
which could be attributed to the enhanced nucleophilicity of the nitrogen atom.

The [2+2] cycloadditions of osmapentalynes with alkynes were further investigated with different reaction conditions. In the presence of water and oxygen, osmapentalyne and arylacetylene were transformed to the first α-metallapentalenofuran 95. Furthermore, 18O labeling experiments suggest that the oxygen atom in the furan ring comes from water [66]. With the synthetic methodology in hand, modifications were carried out to install olefinic groups in the metallapentalenofurans, which were then subjected to polymerization to provide metallapolymers exhibiting excellent stimuli-responsive properties [74,75]. The lactone-fused metallapentalynes 81 and 96 were formed if tetrafluoroboric acid was also present. Interestingly, the metal–carbon triple bond in osmapentalynes 51a was shifted to another five-membered ring in complex 81 and 96 accompanied by the formation of new lactone ring (Scheme 24).

Scheme 23. The [2+2] cycloaddition reactions of metallapentalynes with alkynes and nitrosoarenes.

Scheme 24. Synthesis of complexes 81, 95–96.
In 2018, Xia and coworkers advanced the above [2+2] cycloaddition to establish the first example of a [2+2+2] cycloaddition reaction of an alkyne with a late-transition-metal carbyne complex [48]. Upon treatment of \( \text{NH}_4\text{PF}_6 \), neutral osmapentalyne \( 44 \) underwent [2+2] cycloaddition with the first alkyne. Meanwhile, the chloride ligand of the osmium center was kicked out to afford a cationic intermediate \( 97 \), which exhibited potent reactivity to be inserted by the second alkyne, affording non-planar eleven-carbon framework \( 98 \). Subsequent migratory insertion furnished the thermally stable \( \eta^3\)-Cp complex \( 99 \) (Scheme 25).

![Scheme 25. The [2+2+2] cycloaddition of metallasapentalyne 44 with alkynes.](image)

In 2019, the same research group reported the formal [2+2+2] cycloaddition reaction of a metal-carbyne triple bond with nitriles, demonstrating the “alkyne-like” properties of metal–carbon triple bonds. Treatment of osmapentalyne \( 51a \) with 2,2-diphenylacetonitrile in the presence of sodium hydroxide resulted in the formation of tricyclic metallapyrazine \( 100 \) (Scheme 26) [70].

![Scheme 26. The [2+2+2] cycloaddition of metallasapentalyne 51a with nitriles.](image)

The reaction mechanism was rationalized as follows (Scheme 27). The deprotonation of 2,2-diphenylacetonitrile with sodium hydroxide provided the ketenimine intermediate as a nucleophile, which could attack the carbyne carbon to give intermediate \( 101 \). The chloride ligand was kicked out and replaced with second molecule of 2,2-diphenylacetonitrile to afford intermediate \( 102 \), which would facilitate the 6π electrocyclization reaction to furnish metallapyrazine complex \( 100 \).

Very recently, Xia’s group enriched the “alkyne like” reactivity of metal–carbon triple bond. Osmapentalyne \( 103 \) was prepared from the deprotonation of metallasapentalene, furnishing the first cyclic metal carbyne in tricyclic metalla-aromatics with a bridgehead metal, which was subjected to a range of diversely functionalized azides. Excellent yields, regioselectivity and compatibility were observed to provide an unprecedented series of polycyclic aromatics (\( 104a–e \)), in which a bridgehead metal center was shared by four five-membered aromatic rings, thus representing a typical “metalla-click” reaction (Scheme 28) [71]. The construction of polycyclic aromatics that share a bridgehead atom with more than three rings has never been accomplished before this work, which therefore broke the record.
The substrate scope was examined with electron-withdrawing substituents to give the expected product, substantiated by theoretical calculations. Based on the results achieved, the Simmons–Smith reactions were also extended to ruthenapentalynes [72]. These results demonstrate that the reactions were sensitive to electronic effects, which was substantiated by theoretical calculations. Based on the results achieved, the Simmons–Smith reactions were also extended to ruthenapentalynes [72].

During the preparation of this manuscript, Xia and coworkers secured the Simmons-Smith reactions of substrates with metal–carbon triple bonds for the first time [72] (Scheme 29). Osmapentalyne 51a was transformed into cyclopropenation product 105a in the presence of classical Fukuyama reagent. The substrate scope was examined with electron-withdrawing substituents to give the expected cyclopropene products 105b-d in good yields. When the substituents were switched to carbinols, the Fukuyama reagent failed, and the more reactive Shi’s reagent was used to furnish the desired product 107. These results demonstrate that the reactions were sensitive to electronic effect, which was substantiated by theoretical calculations. Based on the results achieved, the Simmons–Smith reactions were also extended to ruthenapentalynes [72].

**Scheme 27.** Mechanism of [2+2+2] cycloaddition of metallapentalyne 51a with nitriles.

**Scheme 28.** Metalla-click reactions.

**Scheme 29.** Simmons-Smith reactions.
Unprecedented [3+1] cycloadditions of metalla-azirines with terminal alkynes were documented by Xia’s [73] group (Scheme 30). Osmapentalyne 50a was transformed into the metalla-azirine 108 in the presence of sodium azide. The reaction was the first synthesis of a metalla-azirine by [2+1] cycloaddition reaction of a metal carbyne with an azide and paralleled the first [3+2] cycloadditions of late-transition-metal carbyne with organic azides [71]. In the presence of AgBF4, a series of substituted terminal alkynes as well as metalla-azirine 108 were transformed to various tetracyclic complexes 109a–d. Moreover, the five coordinating atoms lie in the equatorial plane in the CCCCX-type (X = N, O, S) carbolong complexes obtained and the reaction provides a valuable supplement to the construction of planar pentadentate chelates.

![Scheme 30. Formal [3+1] cycloadditions.](image)

4. Conclusions

Fantastic molecular architectures characterized with aromaticity have drawn considerable attention from the synthetic community, thus producing a huge number of aromatic structures with elegance as well as variety. As commonly sensed, the organic alkyne exhibits a linear shape and the bond angles around the acetylenic carbons are 180°. Many efforts have been devoted to the chemistry of cyclic complexes containing metal-carbon triple bonds, in which the carbine-carbon bond angles were less than 180° since the first example of metallabenzyne in 2001. Cyclic complexes containing carbynes could be classified into six-membered metallacarbynes and five-membered metallapentalynes. The carbine-carbon bond angles of metallabenzenes are around 148°. Such a record was held for 12 years until the discovery of metallapentalynes, whose carbine-carbon bond angles were bent to around 130°, far away from the ideal value of 180°. Such extreme distortion results in considerable large ring strain, resulting in the unprecedented high reactivity.

The reaction types of metallabenzenes could be classified into four types, such as ligand exchange reactions, electrophilic reactions, nucleophilic reactions and migratory insertion reactions. Ligand exchange reactions are the fundamental reactions in organometallic chemistry and were unsurprisingly involved in the reactions of metallabenzenes and metallapentalynes. Due to the ambiphilic properties of carbynes, both electrophilic and nucleophilic reactions were observed in the reactions of metallabenzenes and metallapentalynes, which resemble the classical metal carbyne complex. Metallabenzenes exhibited excellent aromatic properties and the whole cyclic system acted as an entire entity under common electrophilic conditions, thus resembling the electrophilic substitution
reactions displayed upon organic aromatics. The carbyne moiety stayed intact, whereas the aromatic rings were attacked by electrophiles. One exception that was observed was that the carbyne moiety was epoxidized upon treatment of hydrogen peroxide. In most cases, the nucleophiles would attack the carbyne carbon. However, “softer” nucleophiles might attack the para position of the six-membered ring system, thus demonstrating the aromaticity of metallabenzenes. Meanwhile, the aromaticity of osmabenzenes was strongly affected by the corresponding substituents of metallocycles and ligands of metal center, leading to the decomposition of metallacarbyne ring systems to form cyclopentadiene complexes via migratory insertion reactions. In sharp contrast, the ring systems of metallapentalynes were never broken throughout all the reported reactions, which might be attributed to the higher stability of metallapentalynes derived from their strong aromaticity despite the extremely strained bond angles of metallapentalynes. The metal–carbon triple bond in metallapentalynes possessed “alkyne-like” character, which was demonstrated by the formation of osmapentalyne-coinage metal complexes and cycloaddition reactions.

Recent advances in the reactivity of a series of cyclic complexes containing carbenes described herein deepened our understanding of the nature of metallabenzenes and metallapentalynes and resulted in the production of a number of novel metallocycles. However, the research on the reactivity of metallabenzenes and metallapentalynes is still limited compared with that of organic alkynes and metal-carbon triple bonds. Additionally, the reactivity studies of these unique species need further exploration, and the isolation of other metalla-aromatics is also anticipated via the corresponding research. Considering that metallocycles are regarded as intermediates in many transition-metal-catalyzed reactions, this study provides a new perspective for the application of late-transition-metal carbyne species as catalysts.

The unique structure leads to novel properties. Novel metallocycles derived from the reactivity studies of metallapentalynes exhibited great application prospects in various areas, such as near-infrared dyes [47], photothermal therapy [74–77], phototherapy [78], and self-healing materials [79], enabling their potential applications in materials science and biomedicine. Future developments of metallabenzenes and metallapentalynes and exploration of their potential applications in catalysis [60], materials science and biomedicine would not only extend our perception of aromaticity in metallocycles, but also further enrich fundamental organometallic chemistry.

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