CHEMISTRY

Constraint of a ruthenium-carbon triple bond to a five-membered ring

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The incorporation of a metal-carbon triple bond into a ring system is challenging because of the linear nature of triple bonds. To date, the synthesis of these complexes has been limited to those containing third-row transition metal centers, namely, osmium and rhenium. We report the synthesis and full characterization of the first cyclic metal carbyne complex with a second-row transition metal center, ruthenapentalyne. It shows a bond angle of 130.2(3)° around the sp-hybridized carbyne carbon, which represents the recorded smallest angle of second-row transition metal carbyne complexes, as it deviates nearly 50° from the original angle (180°). Density functional theory calculations suggest that the inherent aromatic nature of these metallacycles with bent $Ru \equiv C-C$ moieties enhances their stability. Reactivity studies showed striking observations, such as ambiphilic reactivity, a metal-carbon triple bond shift, and a [2 + 2] cycloaddition reaction with alkyne and cascade cyclization reactions with ambident nucleophiles.

INTRODUCTION

Transition metal complexes containing metal-carbon triple bonds (namely, metal carbyne or metal alkylidyne complexes) hold an important place in chemistry because they stimulate fundamental investigations of metal-carbon interactions and are valuable in synthetic chemistry as homogeneous catalysts or precursors for interesting organometallic compounds (1-4). Accordingly, the chemistry of the metal-carbon triple bond has been an area of intense interest and has attracted tremendous attention in both industry and academia since the seminal work by Fischer et al. (5). Although a wide variety of metal carbyne complexes are known, most documented examples feature acyclic structures. The linear geometry of the sp-hybridized carbyne carbon makes the synthesis of cyclic metal carbyne complexes particularly difficult. Thus, cyclic metal carbyne complexes remained elusive until the pioneering works by Wen et al. (6) (Fig. 1). Until now, well-defined cyclic metal carbynes were only known for third-row transition metal centers osmium and rhenium, namely, osmabenzynes (7–9), rhenabenzynes (10), osmapyridynes (11), osmapentalynes (12, 13), and dirhenadehydro [12] annulenes (14), whereas these complexes with non-third-row transition metals have not yet been accomplished.

The lack of non-third-row transition metal carbynes is not surprising. Traditionally, the strength of transition metal-carbon bonds decreases with the ascending of a column in the periodic table (15), which results in relative lability of the first- and second-row organometallic compounds in comparison with their third-row analogs (16, 17). In particular, cyclic metal carbyne complexes with first- and second-row transition metals are considerably less stable than their third-row counterparts both thermodynamically and kinetically, as demonstrated by computational analysis (18, 19). Thus, intrinsic instability of the non-third-row transition metal carbynes is responsible for their absence, which greatly restrained their synthesis and further exploration.

During our recent studies on the chemistry of metallacyclic complexes, we found that typically unstable species could show enhanced

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stability owing to metalla-aromaticity, which yielded a number of exceptional organometallic complexes with osmium and ruthenium centers (12, 20, 21). These results prompted us to explore the construction of cyclic metal carbyne complexes with second-row transition metal centers using our synthetic strategies for metalla-aromatics. Herein, we report the synthesis and characterization of unprecedented ruthenium-based cyclic carbyne complexes, that is, ruthenapentalynes. These ruthenapentalynes exhibit good stability in the solid state, although the bent Ru \equiv C–C moiety shows a bond angle of only 130.2(3)°, which deviates considerably from the linear nature of the sp-hybridized carbyne carbon. Experimental and computational studies revealed the inherent aromaticity of these cyclic ruthenium carbyne complexes. Synergistic interplay of aromaticity and ring strain results in the unique reactivities of ruthenapentalyne, providing access to a series of novel polycyclic aromatic molecules with a second-row transition metal.

RESULTS

Synthesis and characterization of ruthenapentalynes

We recently established an efficient aromaticity-driven method for the construction of cyclic osmium carbyne complexes based on multiyne chains (termed carbolongs) (13). We envisioned that the powerful ligating ability of carbolongs would lead to the realization of cyclic metal carbyne complexes with second-row transition metal centers. As shown in Fig. 2A, we formed the expected cyclic ruthenium carbyne complex **2a** and isolated it as a red solid in 65% yield when a solution of carbolong **1a** was treated with RuCl₂(PPh₃)₃ and excess PPh₃ at room temperature (RT) for 3 hours. We also obtained a similar cyclic ruthenium carbyne complex, **2b**, upon reacting **1b** with RuCl₂(PPh₃)₃.

High-resolution mass spectrometry (HRMS), elemental analysis (EA), and nuclear magnetic resonance (NMR) spectroscopy supported the identification of **2a** and **2b**. In the ¹H NMR spectrum, we observed the two resonances from the metallacyclic skeletons at 13.06 and 7.32 parts per million (ppm) (**2a**) and 13.17 and 7.35 ppm (**2b**), which are located in the metalla-aromatic region and are comparable to those of osmapentalynes (*12, 13*). The ¹³C NMR spectrum exhibits characteristic low-field signals at 355.69 (**2a**) and 355.57 (**2b**) ppm for the carbyne carbons, which are comparable to the signals of previously reported ruthenium carbyne complexes (*22, 23*). However, these

chemical shifts are significantly low field-shifted in comparison to those of sp-hybridized carbon atoms in closely related complexes, such as cyclic metal vinylidene complexes (24) and cyclic allenes and their metal complexes reported by Bertrand and colleagues (25, 26).

Single-crystal x-ray diffraction analysis determined the solid-state structure of **2a** (Fig. 2B). The structure of **2a** contains two fused fivemembered metallacycles with good planarity, as reflected by the small mean deviation (0.015 Å) from the least-squares plane through Ru and C1–C7. The carbon-carbon bond distances within the metallacycles [1.360(7) to 1.412(6) Å] lie between typical single- and double-bond distances. The corresponding carbon-carbon bond lengths, together with the planarity of the metallabicycle, indicate extensive electronic delocalization within the ruthenapentalyne ring in **2a**. Consistent with

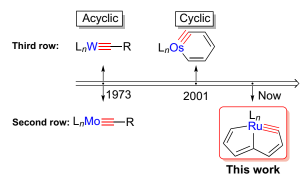


Fig. 1. Development of transition metal carbyne complexes. L, ligand.

other cyclic metal carbyne complexes (6-13), ruthenapentalyne 2a could be represented by two resonance structures, namely, 2A with a Ru \equiv C unit and **2B** with a Ru=C=C unit (Fig. 2A). Because of the extensive electronic delocalization within the metallacycles, as well as the ring strain of small-sized cyclic metal carbyne, the Ru-C1 bond length [1.833(4) Å] in 2a is longer than those of terminal ruthenium carbynes (1.660 to 1.766 Å) (bond length ranges in this article are all based on a search of the Cambridge Structural Database, version 5.39 in November 2017), whereas it is comparable to the Os≡C triple bond length of the first osmabenzyne [1.815(4) Å] (6) and osmapentalyne [1.845(3) Å] (12) with aromaticity. The comparability of Os=C and Ru≡C bond lengths within similar overall structures is explained by the similar atom radii of ruthenium (1.34 Å) and osmium (1.35 Å) atoms (27) and exemplified by the identical metal-carbon triple bond lengths of cationic carbyne complex [Cp*(PEt₃)₂Ru≡CCH=CPh₂]²⁺ $(1.765 \text{ Å}) (28) \text{ and } [Cp(P^{i}Pr_{3})_{2}Os \equiv CCH = CPh_{2}]^{2+} (1.764 \text{ Å}) (29). The$ Ru-C4 [2.104(4) Å] and Ru-C7 [2.036(4) Å] bond distances in 2a are within the range of the distances of typical Ru-C(vinyl) bonds (1.973 to 2.202 Å), with the Ru–C7 bond length slightly shorter than that of Ru-C4. Notably, the bond angle at the sp-hybridized carbon atom in 2a is only 130.2(3)°, which, to the best of our knowledge, is far smaller than the previously reported smallest bond angle for a carbyne carbon bonded with a second-row transition metal [152.3(3)°] (30). Thus, ruthenapentalyne 2a can be regarded as the first example of a cyclic metal carbyne complex with a second-row transition metal center, and its exceptional bond angle is the smallest on record for second-row transition metal carbyne complexes.

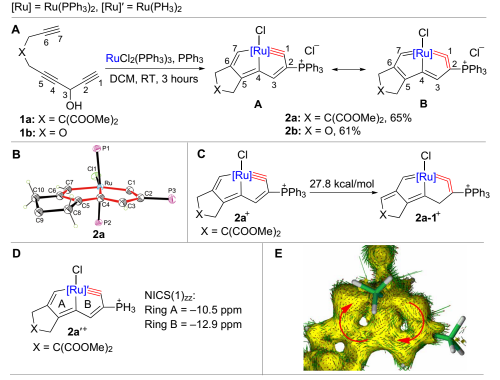


Fig. 2. The synthesis, structure, and aromaticity of ruthenapentalynes 2. (A) Synthesis of ruthenapentalynes 2 from carbolongs 1. DCM, dichloromethane. (B) X-ray molecular structure for the cation of ruthenapentalyne 2a (the ellipsoids are drawn at the 50% probability level; phenyl groups and ester groups are omitted for clarity; the detailed structure is presented in fig. S1). (C) ASE evaluation of the aromaticity of ruthenapentalyne 2a. (D) Nucleus-independent chemical shift (NICS)(1)_{zz} evaluations of aromaticity of model complex 2a'. (E) AICD plot of model complex 2a' with an isosurface value of 0.03. The magnetic field vector is orthogonal to the ring plane and points upward (aromatic species exhibit clockwise diatropic circulations).

Evaluation of the aromaticity of ruthenapentalyne 2a by density functional theory calculations

Formation of small rings containing sp-hybridized carbon atoms is often disfavored because of strain. However, in this case, ruthenapentalyne 2a exhibits good thermal stability (the solid sample is persistent in air at 60°C for at least 3 hours). We surmised that the inherent aromaticity of 2a significantly enhances the stabilization of the fivemembered ring with the Ru≡C unit. We performed density functional theory (DFT) computations to evaluate the aromaticity of ruthenapentalyne 2a. As shown in Fig. 2C, the aromatic stabilization energy (ASE) of 2a (27.8 kcal/mol) is comparable to those of other reported metalla-aromatics (31). We also calculated the nucleusindependent chemical shift along the z axis at 1 Å above the ring critical point $[NICS(1)_{zz}]$ (32) (we used the average value when the environments above and below the ring centers were not equivalent) of model compound 2a', which was simplified by replacing the PPh₃ groups with PH₃ groups (Fig. 2D). Both the considerable ASE value and negative nucleus-independent chemical shift (NICS) values suggest that the metallacycles in ruthenapentalyne 2a are aromatic. In addition, the anisotropy of the induced current density (AICD) analysis confirmed the aromaticity of 2a (33). The obvious diatropic ring current (clockwise vectors) demonstrated the aromatic character, within the metallabicyclic moiety of model ruthenapentalyne 2a' (Fig. 2E).

Considering the tremendous progress in metalla-aromatic chemistry, it is quite peculiar that a great number of metalla-aromatic species contain third-row transition metals (*34*), whereas only a few of them contain first- and second-row transition metal centers (*35–39*). Ruthenapentalyne **2a** represents an important complement to the metalla-aromatic family and distinguishes itself by furnishing a ruthenium carbyne moiety within a five-membered ring. We also conducted DFT calculations to elucidate the formation of ruthenapentalyne **2a** (fig. S7). The computed free-energy profile of the reaction shows that the stability of aromatic ruthenapentalyne is crucial for the overall reaction, which can be classified as an aromaticity-driven process.

Ambiphilic reactivity and [2 + 2] cycloaddition reactions with alkynes

Although ruthenapentalynes 2 are stable in the solid state, the metal carbyne moieties of 2 are highly reactive toward both nucleophiles and electrophiles in solution. As shown in Fig. 3A, the reaction of 2 with sodium thiophenoxide under an atmosphere of CO led to the formation of the nucleophilic addition product ruthenapentalenes 3, which were characterized by NMR spectroscopy. The ¹H NMR spectra showed signals for the metallacycles at 10.87 and 7.83 ppm for 3a and 10.92 and 7.67 ppm for 3b. All the NMR shifts are in the typical region for metalla-aromatics, illustrating the aromaticity of 3. Singlecrystal x-ray analysis confirmed the structure of 3b and disclosed the electronic delocalization within the metallacycles (Fig. 3B). The crystal structure revealed the planarity of the metallabicycles (the mean deviation from the least-squares plane of Ru and C1–C7 is 0.033 Å). The bond lengths of Ru-C1 [2.089(3) Å] and Ru-C7 [2.087(3) Å] are identical, both of them falling within the same range as those of previous reported ruthenabenzenes (1.910 to 2.110 Å) (20, 36). Complexes 3 also show good stability toward air, water, and heat. We evaluated the aromaticity of 3 using DFT calculations (figs. S9 to S11). Despite the great recent interest in such aromatic species, the metallapentalene examples reported are limited to only osmapentalenes (21). The synthesis of 3 demonstrates that the incorporation of metals other than osmium into pentalene rings is feasible.

Ruthenapentalynes are also capable of reacting with copper(I) halides. As shown in Fig. 3A, ruthenapentalynes 2 readily reacted with CuCl to afford bimetallic complexes 4, which were isolated in moderate yield (4a, 87%; 4b, 81%). The NMR spectra and solid-state structure established 4 as bimetallacyclopropene derivatives. Taking 4a as an example, the C^7H and C^3H signals observed at 13.01 and 7.54 ppm compare well with those in the ¹H NMR spectroscopy of the parent complex 2a (13.06 and 7.32 ppm), whereas the resonance of C^1 (325.76 ppm) is significantly upshifted in comparison with that of 2a (355.69 ppm), as determined by ¹³C NMR spectroscopy. We also characterized complex 4a using single-crystal x-ray diffraction (Fig. 3C). The Ru-C1 bond length [1.907(2) Å] is 0.074 Å longer than that in the parent complex 2a [1.833(4) Å], and the Ru–Cu1 bond length of 2.5333(4) Å is shorter than the sum of the covalent radii of Ru and Cu atoms (2.62 Å), revealing the contribution from two resonance forms, A (with a bimetallacyclopropene unit) and **B** (with the ruthenacarbyne coordinated to the copper center), as shown in fig. S13. The deviation from the least-squares plane through Cu1, Ru, and C1-C7 is only 0.023 Å, indicating that the Cu atom exists in the same plane as the five-membered metallabicycle. The Cu1-C1 bond length of 1.919(2) Å is within the range of typical Cu-C(aryl) bond lengths (1.849 to 2.020 Å). These results suggest that there is a strong interaction between the CuCl and Ru-C moieties and that resonance form A should be more important. To the best of our knowledge, complexes 4 represent the first ruthenium carbyne Cu(I) adducts, even though their osmium counterparts were obtained nearly 40 years ago (40, 41).

Although a series of typical electrophiles, including MeI, MeOTf, EtOTf, Me₃OBF₄, and Et₃OBF₄, cannot react with ruthenium carbyne moieties of ruthenapentalynes 2, we attempted to seek other electrophilic reactions. The metal-carbon triple bond of 2 could shift from the original ring to the other fused ring in the presence of acid. Such an extraordinary metal carbyne bond shift reaction has been observed only in osmium species (21). As shown in Fig. 3A, treatment of ruthenapentalyne 2b with dichloroacetic acid at 0°C for 20 min formed new ruthenapentalyne 5, which was characterized in situ. Ruthenapentalyne 5 is not stable in solution and persists only for 4 hours in the reaction mixture. Solution NMR spectra showed the expected chemical shift (13.03 ppm) for $C^{1}H$ in ¹H NMR and the slightly up-field signal at 345.72 ppm in ¹³C NMR for the carbyne carbon in comparison with that of the parent ruthenapentalyne 2b (355.49 ppm for C^1). In addition, the HRMS shows an ion peak at mass/charge ratio (m/z) 1053.1892 corresponding to $[C_{63}H_{51}ClOP_3Ru]^+$. To understand the mechanistic aspects of the conversion of 2b to 5, we performed the experiments starting from deuterium-labeled acid. As shown in Fig. 3A, the isotopic-labeling experiment suggests that the hydrogen atom at the original carbyne carbon position in the product 5 comes from the acid proton. Thus, we think that the electrophilic attack of acid, on the carbyne carbon of 2a, initiates the conversion reaction and produces a metal center 16e ruthenapentalene intermediate (Fig. 3B), owing to the ambiphilic reactivity of 2a. The following elimination of proton from the C7 position of ruthenapentalene intermediate could yield the new ruthenapentalyne 5. The proposed addition/elimination mechanism is similar to the metal carbyne bond shift reaction of osmapentalyne (21).

As suggested by the reactions of **2** with sodium thiophenoxide, CuCl, and acids (Cl₂CHCOOH and CF₃COOD), ruthenapentalynes are ambiphilic. In general, metal carbyne complexes are either nucleophilic or electrophilic: Only a few of them can react with both nucleophiles and electrophiles (12, 42, 43). We assumed that the ambiphilic character $[Ru] = Ru(PPh_3)_2$

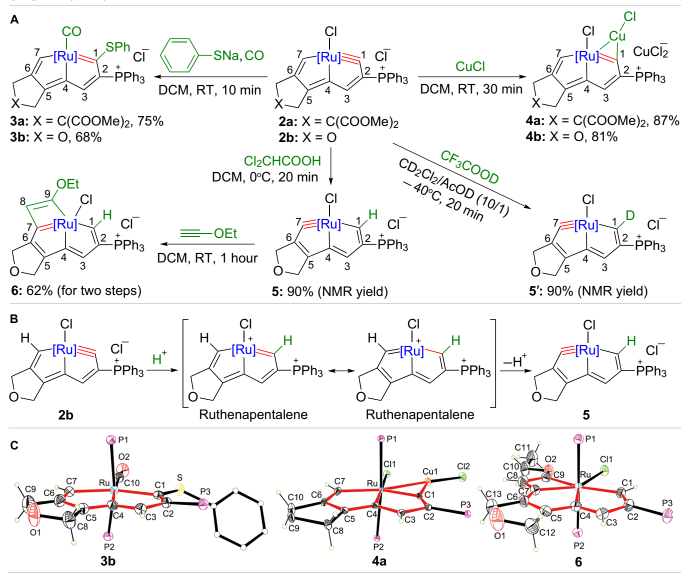


Fig. 3. Ambiphilic reactivity and [2 + 2] cycloaddition reaction of ruthenapentalynes. (A) Reactions of 2 with sodium thiophenoxide and CuCl, Cl₂CHCOOH, and CF₃COOD. (B) The proposed mechanism for the metal-carbon triple bond shift reaction of ruthenapentalyne 2b in the presence of acid. (C) X-ray crystal structures of the cations of 3b, 4a, and 6 (the ellipsoids are drawn at the 50% probability level; phenyl groups of 3b, 4a, and 6 and ester groups of 4a are omitted for clarity; the detailed structures are presented in figs. S2 to S4 for 3b, 4a, and 6, respectively).

of **2** should be attributed to their extremely distorted $Ru \equiv C-C$ moieties, which may result in a strong tendency to relieve the ring strain by transformation of the sp-hybridized carbyne carbons.

Since the thermal lability of **5** precluded its crystallographic characterization, we carried out the reactivity studies to further confirm its identity. Treatment of in situ prepared **5** with HC≡COEt in dichloromethane at RT for 1 hour afforded complex **6** (Fig. 3A). Complex **6** is stable enough to undergo NMR spectroscopy and single-crystal x-ray diffraction analysis. The core of **6** contains a four-membered ruthenacyclobutadiene ring (Fig. 3C), which is achieved via the [2 + 2] cycloaddition of the alkyne with the metal carbyne unit of **5**. Four Ru–C bonds are located in the equatorial plane in **4**, among which Ru–C1 [2.101(4) Å] and Ru–C7 [2.099(4) Å] are comparable and within the range of those of previous reported ruthenabenzenes (1.910 to 2.110 Å) (20, 36), whereas the other two [Ru–C4, 2.143(4) Å; Ru–C9, 2.191(4) Å] are much longer. The carbon–carbon lengths of the metallatricycle fall within the range of 1.332(6) to 1.433(6) Å with slight alternation between C–C single and C=C double bonds.

The ruthenacyclobutadiene moiety of **6** supports the assignment of the location of the metal-carbon triple bond in ruthenapentalyne **5**. Although ruthenacyclobutadiene complexes have fascinated theoretical chemists (44, 45), only one example has been achieved experimentally, which was prepared by tetramerization of phenylacetylide residues at ruthenium centers (46). For [2 + 2] cycloadditions of alkynes with metal carbyne complexes, there is only one previously reported example for the late-transition metal carbynes (47), although examples of early-transition metal carbyne complexes have been well documented (48–50). The formation of complex **6** represents the first [2 + 2] cycloaddition reaction of a ruthenium carbyne complex with an alkyne.

Cascade cyclization reactions with ambident nucleophiles

Since the above investigation underlined the peculiarities of the metal carbyne moieties in ruthenapentalynes **2**, we next examined the reactions with ambident substrates to obtain more insight into their reactivities and properties. Accordingly, we first reacted **2** with cyanates. As shown in Fig. 4A, treatment of **2a** with excess NaOCN led to annulation complex 7, which contained two molecules of the cyanate ion.

X-ray crystallographic analysis of 7 showed that the ruthenium center adopts a distorted pentagonal bipyramidal geometry with the equatorial plane vertexes occupied by four carbon atoms and one oxygen atom (Fig. 4B). The carbon-carbon bond lengths within the two five-membered metallacycles range from 1.353(6) to 1.452(6) Å. In the six-membered azametallacycle, the 1.285(6) Å C8–N1 and 1.313(6) Å C1–N2 distances are in the double-bond range and are significantly shorter than C9–N1 [1.391(6) Å] and C9–N2 [1.385(6) Å]. The exocyclic C9–O2 bond [1.239(6) Å] is a typical C=O double bond and shorter than C8–O1 [1.249(5) Å] within the three-membered ring. The Ru–C8 bond [2.040(4) Å] is the shortest Ru–C bond in 7, which suggests a strong interaction between the metal centers and the OCN ligand. In comparison, the other three Ru–C bond lengths in 7 are 2.122(4), 2.087(4), and 2.141(4) Å for Ru–C1, Ru–C4, and Ru–C7, respectively. The structural parameters demonstrate an electron-localized structure of 7, which can be represented by two resonance structures of 7A and 7B (Fig. 4A).

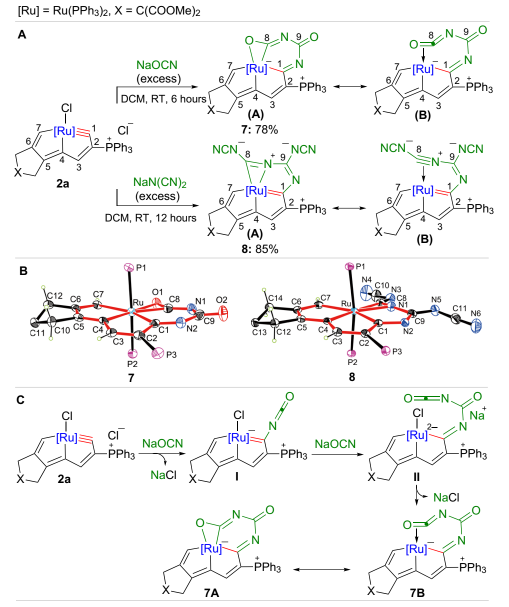


Fig. 4. Cascade cyclization reactions of ruthenapentalyne 2a. (A) Cascade cyclization reactions of 2a with ambident nucleophiles. (B) X-ray molecular structures for complexes 7 and 8 (the ellipsoids are drawn at the 50% probability level; phenyl groups and ester groups are omitted for clarity; the detailed structures are presented in figs. S5 and S6 for 7 and 8, respectively). (C) Proposed mechanism for the formation of 7.

We outlined a proposed cascade cyclization mechanism in Fig. 4C. Initially, a nucleophilic attack of a cyanate ion at the carbyne carbon of **2a** would presumably lead to the formation of ruthenapentalene intermediate **I**, which can be further attacked by a second molecule of cyanate ion to yield intermediate **II**. The following coordination of the OCN group to the metal center, accompanied by the dissociation of the Cl group, could result in the final polycyclic complex 7.

Sodium dicyanamide was shown to have a similar ambident ability to react with ruthenapentalynes 2 (Fig. 4A). We hypothesized that the access to annulation product 8 may proceed via a similar cascade cyclization reaction initiated by the nucleophilic attack of the N(CN)2 ion (see fig. S8). X-ray diffraction analysis of 8 locates a five-membered and a three-membered azametallacycle fused with the ruthenapentalene skeleton (Fig. 4B). The Ru-N1 bond length is 2.021(3) Å, in accordance with an azaruthenacycle featuring Ru-N single bonds (51), whereas the Ru–C8 bond [2.213(3) Å] is slightly longer than the typical Ru-C bond length of a Ru-C=N moiety (1.962 to 2.177 Å). The carbonnitrogen bond lengths of the five-membered azametallacycle [C1-N2, 1.338(4) Å; C9–N1, 1.386(4) Å; C9–N2, 1.362(4) Å] are similar to each other and lie between the typical lengths of carbon-nitrogen single and double bonds, but the carbon-nitrogen distance in the threemembered azametallacycle is 1.283(4) Å. Thus, complex 8 has a delocalized structure with contributions from the two resonance structures 8A and 8B, as shown in Fig. 4A.

The rich reactivities of ruthenapentalyne **2** allow access to a series of unusual polycyclic ruthenacycles, with aromatic stability and structural complexity. We summarized the ultraviolet-visible (UV-vis) absorption spectra of polycyclic ruthenacycles **2** to **4** and **6** to **8** in Fig. 5. All of them exhibit efficient absorption ranging from the UV-vis region. When the π -conjugated framework is extended, the characteristic energy absorption bands are relatively red-shifted. The broad absorption, unique structures, and remarkable stability could facilitate further application of these polycyclic ruthenacycles.

CONCLUSION

In summary, we have synthesized the first cyclic carbyne complexes with a second-row transition metal center, namely, ruthenapentalynes, by one-pot reactions of carbolongs with commercially available $RuCl_2(PPh_3)_3$. The carbyne carbon atom of the ruthenapentalyne

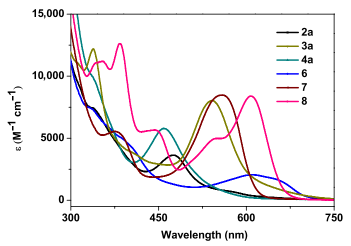


Fig. 5. UV-vis absorption spectra of 2a, 3a, 4a, 6, 7, and 8. Measured in CH_2CI_2 at RT (1.0 × 10⁻⁴ M).

shows a bond angle of only 130.2(3)°, which is greatly reduced compared to the previous record of the smallest angle within a secondrow transition metal carbyne complex. Our experimental observations, together with theoretical calculations, revealed that the inherent aromaticity plays a key role in the stabilization of ruthenapentalynes. We demonstrated that because of the rich and unique reactivity, the ruthenapentalynes could serve as precursors to a family of ruthenacyclic complexes with diverse structural features and distinct properties. This work provides a valuable supplement to the previous paucity of transition metal carbyne complexes and may promote better understanding of the chemistry of cyclic metal carbyne complexes.

MATERIALS AND METHODS

All syntheses were performed under an inert atmosphere (N_2) using standard Schlenk techniques, unless otherwise stated. Solvents were distilled from sodium/benzophenone (hexane and diethyl ether) or calcium hydride (dichloromethane) under N₂ before use. The starting materials 1a and 1b were synthesized according to previously published procedures (13). Other reagents were used as received from commercial sources without further purification. Column chromatography was performed on silica gel (200 to 300 mesh) in air. NMR spectroscopy was performed using a Bruker Advance II 300 spectrometer, a Bruker Advance II 400 spectrometer, a Bruker Advance III 500 spectrometer, or a Bruker Ascend III 600 spectrometer at RT or 0° C. The ¹H and ¹³C NMR chemical shifts (δ) are relative to tetramethylsilane, and the ³¹P NMR chemical shifts are relative to 85% H₃PO₄. The absolute values of the coupling constants are given in hertz. The theoretical molecular ion peak was calculated by Compass Isotope Pattern software supplied by Bruker Company. HRMS was conducted using a Bruker Daltonics Apex Ultra 7.0 T FTMS instrument. EAs were performed on a vario EL III elemental analyzer. Absorption spectra were recorded on a Shimadzu UV2550 UV-vis spectrophotometer.

Representative syntheses are given below. Further details for the syntheses of complexes **2b**, **3b**, **4b**, and **5'** are given in the Supplementary Materials.

Complex 2a

A dichloromethane solution (5 ml) of carbolong 1a (1.65 g, 6.29 mmol) was added slowly to a red solution of RuCl₂(PPh₃)₃ (3.54 g, 3.69 mmol) and PPh₃ (5.02 g, 19.1 mmol) in dichloromethane (150 ml). The reaction mixture was stirred at RT for 3 hours to yield a red solution. The solution was evaporated under vacuum to a volume of approximately 15 ml and then washed with hexane $(3 \times 200 \text{ ml})$ to afford a red solid. The solid was purified by flash chromatography on silica gel (eluent, 20:1 dichloromethane/methanol) to yield complex 2a as a red solid. Yield: 2.89 g, 65%. ¹H NMR (600.1 MHz, CD_2Cl_2): δ 13.06 ppm (s, 1H, C^7H), 7.32 ppm [1H, C^3H , determined by heteronuclear single-quantum coherence (HSQC)], 7.00 to 7.78 ppm (45H of Ph and 1H of $C^{3}H$ mentioned above), 3.63 ppm (s, 6H, COOCH₃), 2.94 ppm [t, ${}^{5}J$ (HP) = 3.82 Hz, 2H, C¹⁰H], and 2.74 ppm (s, 2H, $C^{8}H$). ³¹P NMR (242.9 MHz, $CD_{2}Cl_{2}$): δ 29.62 ppm [d, ⁴J(PP) = 5.89 Hz, RuPPh₃] and 6.35 ppm [t, ${}^{4}J(PP) = 5.89$ Hz, CPPh₃]. ${}^{13}C$ NMR [150.9 MHz, CD₂Cl₂, plus ¹³C DEPT-135, ¹H-¹³C HSQC, ¹H-¹³C heteronuclear multiple-bond correlation (HMBC)]: δ 355.69 ppm [dt, apparent q, ${}^{2}J(CP) = 14.92$ Hz, ${}^{2}J(CP) = 14.92$ Hz, C^{1}], 250.64 ppm (m, C^{7}), 189.01 ppm [dt, ${}^{3}J(CP) = 25.58$ Hz, ${}^{2}J(CP) = 5.09$ Hz, C⁴], 179.74 ppm (s, C⁵), 170.53 ppm (s, COOCH₃), 163.33 ppm

[t, ${}^{3}J(CP) = 3.62$ Hz, C⁶], 153.74 ppm [d, ${}^{2}J(CP) = 14.34$ Hz, C³], 126.98 to 134.52 ppm (Ph), 121.60 ppm [dt, ${}^{1}J(CP) = 92.66$ Hz, ${}^{3}J(CP) =$ 3.51 Hz, C²], 118.32 ppm [d, ${}^{1}J(CP) = 91.08$ Hz, Ph], 62.90 ppm (s, C⁹), 53.13 ppm (s, COOCH₃, obtained by ${}^{13}C$ DEPT-135), 37.80 ppm (s, C⁸), and 36.10 ppm (s, C¹⁰). Analysis calculated (Anal. calculated; %) for C₆₈H₅₇Cl₂O₄P₃Ru: C, 67.89; H, 4.78. Found: C, 67.60; H, 5.15. HRMS [electrospray ionization (ESI)]: m/z calculated for [C₆₈H₅₇ClO₄P₃Ru]⁺, 1167.2212 [M]⁺; found, 1167.2226.

Complex 3a

A mixture of 2a (300 mg, 0.249 mmol) and NaSPh (99 mg, 0.75 mmol) in dichloromethane (10 ml) was stirred at RT for 10 min under an atmosphere of CO to yield a red solution. The solid was removed by filtration. The filtrate was evaporated under vacuum to approximately 2 ml and then purified by column chromatography on silica gel (eluent, 15:1 dichloromethane/methanol) to afford 3a as a red solid. Yield: 244 mg, 75%. ¹H NMR (500.2 MHz, CD₂Cl₂): δ 10.87 ppm (s, 1H, C^7H), 7.83 ppm [d, ${}^{3}J$ (HP) = 5.45 Hz, 1H, $C^{3}H$], 7.04 to 7.68 ppm (45H, Ph), 6.88 ppm $[t, {}^{3}J(HH) = 7.40 \text{ Hz}, 1H, Ph], 6.52 \text{ ppm}$ [t, ³*J*(HH) = 7.40 Hz, 2H, Ph], 5.99 ppm [d, ³*J*(HH) = 7.40 Hz, 2H, Ph], 3.52 ppm (s, 6H, COOCH₃), 2.55 ppm (s, 2H, C¹⁰H), and 2.09 ppm (s, 2H, C⁸H). ³¹P NMR (202.5 MHz, CD₂Cl₂): δ 36.79 ppm (s, RuPPh₃) and 7.28 ppm (s, CPPh₃). ¹³C NMR (125.8 MHz, CD₂Cl₂, plus ¹³C-dept 135, ¹H-¹³C HSQC, and ¹H-¹³C HMBC): δ 249.56 ppm (br, C^1) , 238.51 ppm $[t, {}^2J(CP) = 17.74 \text{ Hz}, C^7]$, 203.96 ppm $[t, {}^2J(CP) =$ 15.06 Hz, RuCO], 197.80 ppm [dt, ${}^{3}J(CP) = 27.45$ Hz, ${}^{2}J(CP) = 7.71$ Hz, C⁴], 180.96 ppm (s, C⁵), 171.71 ppm (s, COOCH₃), 169.85 ppm (s, C⁶), 161.86 ppm [d, ${}^{2}J(CP) = 24.53$ Hz, C³], 139.00 ppm (s, Ph), 128.29 to 135.33 ppm (Ph), 122.35 ppm [d, ¹*J*(CP) = 86.85 Hz, Ph], 121.97 ppm [d, ${}^{1}J(CP) = 59.36$ Hz, C^{2}], 64.08 ppm (s, C^{9}), 53.32 ppm (s, COOCH₃), 38.38 ppm (s, C¹⁰), and 37.62 ppm (s, C⁸). Anal. calculated (%) for C₇₅H₆₂ClO₅SP₃Ru: C, 69.04; H, 4.79. Found: C, 68.92; H, 4.85. HRMS (ESI): m/z calculated for $[C_{75}H_{62}O_5P_3RuS]^+$, 1269.2589 [M]⁺; found, 1269.2588.

Complex 4a

A mixture of complex 2a (300 mg, 0.249 mmol) and CuCl (51 mg, 0.52 mmol) in dichloromethane (10 ml) was stirred for 30 min at RT to yield an orange solution. Excess CuCl was removed by filtration. The filtrate was evaporated under vacuum to approximately 3 ml and washed with diethyl ether $(3 \times 20 \text{ ml})$ to afford 4a as an orange solid. Yield: 303 mg, 87%. ¹H NMR (400.1 MHz, CD₂Cl₂): δ 13.01 ppm [t, ${}^{3}J$ (HP) = 4.55 Hz, 1H, C⁷H], 7.54 ppm (1H, C³H, determined by HSQC), 7.04 to 7.81 ppm (45H of Ph and 1H of $C^{3}H$ mentioned above), 3.64 ppm (s, 6H, COOCH₃), 2.86 ppm [t, ${}^{5}J$ (HP) = 3.76 Hz, 2H, C¹⁰H], and 2.83 ppm (s, 2H, C⁸H). ³¹P NMR (161.9 MHz, CD_2Cl_2): δ 22.84 ppm [d, 4J (PP) = 4.15 Hz, RuPPh₃] and 6.65 ppm $[t, {}^{4}J(PP) = 4.15 \text{ Hz}, CPPh_{3}]. {}^{13}C \text{ NMR} (100.6 \text{ MHz}, CD_{2}Cl_{2}, plus)$ ¹³C-dept 135, ¹H-¹³C HSQC, and ¹H-¹³C HMBC): δ 325.76 ppm (br, C^{1}), 250.58 ppm [t, ${}^{2}J(CP) = 13.02$ Hz, C^{7}], 194.40 ppm $[dt, {}^{3}J(CP) = 22.93 \text{ Hz}, {}^{2}J(CP) = 4.55 \text{ Hz}, C^{4}], 184.57 \text{ ppm } (s, C^{5}),$ 171.01 ppm (s, COOCH₃), 166.52 ppm (s, C⁶), 158.41 ppm $[d, {}^{2}J(CP) = 15.32 \text{ Hz}, C^{3}], 128.58 \text{ to } 135.45 \text{ ppm}$ (Ph), 127.93 ppm $[dt, {}^{1}J(CP) = 87.05 \text{ Hz}, {}^{3}J(CP) = 3.81 \text{ Hz}, C^{2}], 118.10 \text{ ppm} [d, {}^{1}J(CP) =$ 90.36 Hz, Ph], 63.98 ppm (s, C⁹), 53.34 ppm (s, COOCH₃), 39.34 ppm (s, C^{10}) , and 36.79 ppm (s, C^8) . Anal. calculated (%) for $C_{68}H_{57}Cl_4C$ u₂O₄P₃Ru: C, 58.29; H, 4.10. Found: C, 58.22; H, 4.47. HRMS (ESI): *m/z* calculated for [C₆₈H₅₇Cl₂CuO₄P₃Ru]⁺, 1267.1185 [M]⁺; found, 1267.1187.

Complex 5

Cl₂CHCOOH (58 µl, 0.703 mmol) was added to a solution of 2b (300 mg, 0.275 mmol) in dichloromethane (15 ml). The reaction mixture was stirred at 0°C for 20 min to yield a red solution of 5 (in ca. 90% yield based on ¹H and ³¹P NMR), which was characterized by in situ NMR and HRMS (ESI). ¹H NMR (600.1 MHz, CD₂Cl₂): δ 13.03 ppm [ddd, ${}^{2}J(\text{HP}) = 16.85 \text{ Hz}, {}^{2}J(\text{HP}) = 5.72 \text{ Hz}, {}^{2}J(\text{HP}) = 3.00 \text{ Hz}, 1\text{H}, \text{C}^{1}H$ 7.64 ppm (s, 1H, $C^{3}H$), 6.84 to 7.82 ppm (45H of Ph and 1H of $C^{3}H$ mentioned above), 4.05 ppm (s, 2H, C⁸H), and 3.55 ppm (s, 2H, $C^{9}H$). ³¹P NMR (242.9 MHz, CD₂Cl₂): δ 32.53 ppm [d, ⁴J(PP) = 4.93 Hz, RuPPh₃] and 12.16 ppm $[t, {}^{4}J(PP) = 4.93$ Hz, CPPh₃]. {}^{13}C NMR (150.9 MHz, CD₂Cl₂, plus ¹³C-dept 135, ¹H-¹³C HSQC, and ¹H-¹³C HMBC): δ 345.72 ppm (br, C⁷), 249.98 ppm (br, C¹), 193.77 ppm (s, C^5) , 187.73 ppm $[dt, {}^3J(CP) = 22.29 \text{ Hz}, {}^2J(CP) = 5.83 \text{ Hz}, C^4]$, 157.59 ppm (s, C⁶), 143.63 ppm [d, ${}^{2}J(CP) = 22.33$ Hz, C³], 127.07 to 134.34 ppm (Ph), 126.21 ppm [d, ${}^{1}J(CP) = 68.70$ Hz, C^{2}], 117.73 ppm [d, ${}^{1}J(CP) =$ 88.44 Hz, Ph], 69.72 ppm (s, C⁹), and 65.22 ppm (s, C⁸). HRMS (ESI): m/z calculated for $[C_{63}H_{51}ClOP_{3}Ru]^{+}$, 1053.1893 $[M]^{+}$; found, 1053.1892.

Complex 6

Cl₂CHCOOH (39 µl, 0.473 mmol) was added to a solution of 2b (200 mg, 0.184 mmol) in dichloromethane (10 ml). The reaction mixture was stirred at 0°C for 20 min, and then HC≡COEt [71 µl $(\omega/\omega = 50\% n$ -hexane solution), 0.364 mmol] was added. The reaction mixture was further stirred for 1 hour at RT to yield a green solution. The solution was evaporated under vacuum to a volume of approximately 2 ml and washed with hexane $(3 \times 15 \text{ ml})$ to afford a green solid. The solid was purified by chromatography on silica gel (silicone; eluent, 20:1 dichloromethane/methanol) to vield complex 6 as a green solid. Yield: 132 mg, 62%. ¹H NMR (300.1 MHz, CD₂Cl₂): δ 12.40 ppm [dd, ${}^{2}J(HP) = 18.71$ Hz, ${}^{2}J(HP) = 4.23$ Hz, 1H, C¹H], 6.88 to 7.77 ppm (45H, Ph), 6.16 ppm (br, 1H, $C^{3}H$), 5.85 ppm [t, ${}^{4}J$ (HP) = 2.11 Hz, 1H, $C^{8}H$], 3.68 ppm [q, 2H, ³J(HH) = 7.11 Hz, OCH₂CH₃], 3.52 ppm (s, 2H, C¹²H), 3.43 ppm (s, 2H, C¹³H), and 1.23 ppm $[t, {}^{3}J(HH) = 7.11 \text{ Hz}, \text{ OCH}_2\text{CH}_3]. {}^{31}\text{P} \text{ NMR} (121.5 \text{ MHz}, \text{CD}_2\text{Cl}_2):$ δ 19.84 ppm [d, ⁴*J*(PP) = 2.83 Hz, Ru*P*Ph₃] and 6.45 ppm [t, ⁴*J*(PP) = 2.83 Hz, CPPh₃]. ¹³C NMR (75.5 MHz, CD₂Cl₂, plus ¹³C-dept 135, ¹H-¹³C HSQC, and ¹H-¹³C HMBC): δ 253.15 ppm (br, C¹), 212.11 ppm (br, C^9) , 204.92 ppm $[t, {}^2J(CP) = 8.92 \text{ Hz}, C^7]$, 191.02 ppm $[dt, {}^3J(CP) =$ 28.00 Hz, ${}^{2}J(CP) = 9.16$ Hz, C^{4}], 188.04 ppm (s, \overline{C}^{5}), 151.34 ppm $(s, C^{6}), 147.14 \text{ ppm } [d, {}^{2}J(CP) = 24.94 \text{ Hz}, C^{3}], 126.77 \text{ to } 133.90 \text{ ppm}$ (Ph), 125.87 ppm [t, ${}^{3}J(CP) = 3.73$ Hz, C^{8}], 121.36 ppm [d, ${}^{1}J(CP) =$ 71.91 Hz, C^2], 118.69 ppm [d, ${}^1J(CP) = 88.44$ Hz, Ph], 68.74 ppm (s, C¹³), 68.23 ppm (s, OCH₂CH₃), 65.11 ppm (s, C¹²), and 12.96 ppm (s, OCH₂CH₃). Anal. calculated (%) for C₆₇H₅₇Cl₂O₂P₃Ru: C, 69.43; H, 4.96. Found: C, 69.07; H, 5.10. HRMS (ESI): m/z calculated for $[C_{67}H_{57}ClO_2P_3Ru]^+$, 1123.2313; found, 1123.2327.

Complex 7

A mixture of **2a** (300 mg, 0.249 mmol) and sodium cyanate (97 mg, 1.49 mmol) in dichloromethane (10 ml) was stirred for 6 hours at RT to yield a red solution. The solid was removed by filtration. The filtrate was evaporated under vacuum to approximately 2 ml and washed with diethyl ether (3 × 20 ml) to afford 7 as a red solid. Yield: 236 mg, 78%. ¹H NMR (600.1 MHz, CD₂Cl₂): δ 10.47 ppm (s, 1H, C⁷H), 7.57 ppm (s, 1H, C³H), 6.77 to 7.55 ppm (45H, Ph), 3.51 ppm (s, 6H, COOCH₃), 2.66 ppm (s, 2H, C¹⁰H), and 2.42 ppm (s, 2H, C¹²H). ³¹P NMR (242.9 MHz, CD₂Cl₂): δ 29.89 ppm (s, RuPPh₃) and 12.24 ppm (s, CPPh₃). ¹³C

NMR (150.9 MHz, CD₂Cl₂, plus ¹³C-dept 135, ¹H-¹³C HSQC, and ¹H-¹³C HMBC): δ 239.75 ppm (br, C¹), 207.71 ppm [t, ²*J*(CP) = 14.89 Hz, C⁷], 193.32 ppm [t, ²*J*(CP) = 4.59 Hz, C⁸], 181.46 ppm [dt, ³*J*(CP) = 29.91 Hz, ²*J*(CP) = 7.44 Hz, C⁴], 173.36 ppm (s, C⁹), 172.31 ppm (s, COOCH₃), 167.49 ppm (s, C⁵), 162.04 ppm [d, ²*J*(CP) = 19.22 Hz, C³], 156.03 ppm (s, C⁶), 127.19 to 134.67 ppm (Ph), 123.18 ppm [d, ¹*J*(CP) = 90.56 Hz, Ph], 122.91 ppm [d, ¹*J*(CP) = 75.83 Hz, C²], 63.80 ppm (s, C¹¹), 52.61 ppm (s, COOCH₃), 38.10 ppm (s, C¹²), and 37.71 ppm (s, C¹⁰). Anal. calculated (%) for C₇₀H₅₇N₂O₆P₃Ru: N, 2.30; C, 69.13; H, 4.72. Found: N, 2.26; C, 68.81; H, 5.09. HRMS (ESI): *m/z* calculated for [C₇₀H₅₈N₂O₆P₃Ru]⁺, 1217.2565 [M + H]⁺; found, 1217.2568.

Complex 8

A mixture of complex **2a** (300 mg, 0.249 mmol) and sodium dicyanamide (133 mg, 1.49 mmol) in dichloromethane (10 ml) was stirred for 12 hours at RT to yield a navy blue solution. Excess sodium salt was removed by filtration. The filtrate was evaporated under vacuum to approximately 3 ml. Then, the residue was washed with diethyl ether (3×20 ml) to afford **8** as a green solid. Yield: 268 mg, 85%. ¹H NMR (400.1 MHz, CD₂Cl₂): δ 11.26 ppm (s, 1H, C⁷H), 7.62 ppm (s, 1H, C³H), 6.89 to 7.70 ppm (45H of Ph and 1H of C³H mentioned above), 3.46 ppm (s, 6H, COOCH₃), 2.74 ppm (s, 2H, C¹⁴H), and 2.62 ppm (s, 2H, C¹²H). ³¹P NMR (161.9 MHz MHz, CD₂Cl₂): δ 31.74 ppm (s, RuPPh₃) and 10.47 ppm (s, CPPh₃). Unfortunately, the poor solubility of **8** prevented its ¹³C NMR characterization. Anal. calculated (%) for C₇₂H₅₇N₆O₄P₃Ru: N, 6.65; C, 68.40; H, 4.54. Found: N, 7.01; C, 68.60; H, 4.42. HRMS (ESI): *m/z* calculated for [C₇₂H₅₈N₆O₄P₃Ru]⁺, 1265.2790 [M + H]⁺; found, 1265.2783.

SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/ content/full/4/6/eaat0336/DC1 fig. S1. X-ray molecular structure for the cation of complex **2a**.

- fig. S2. X-ray molecular structure for the cation of complex **2b**.
- fig. S3. X-ray molecular structure for the cation of complex **4a**.
- fig. S4. X-ray molecular structure for the cation of complex 6.
- fig. S5. X-ray molecular structure for complex 7.
- fig. S6. X-ray molecular structure for complex 8.
- fig. S7. Gibbs free-energy pathway for the DFT-calculated formation mechanism of complex
- 2a at 298 K.
- fig. S8. A plausible mechanism for the formation of complex 8.
- fig. S9. ASE evaluation of complex 3b.
- fig. S10. NICS evaluations of model complex ${\bf 3b}'.$
- fig. S11. AICD plot of the model complex ${\bf 3b}'$ with an isosurface value of 0.03.
- fig. S12. AICD plot of the model complex **2a**' with an isosurface value of 0.03.
- fig. S13. Resonance structures of complexes **4**.
- fig. S14. Positive-ion ESI-MS spectrum of [**2a**]⁺ [C₆₈H₅₇ClO₄P₃Ru]⁺ measured in methanol.
- fig. S15. Positive-ion ESI-MS spectrum of $[\mathbf{2b}]^+$ $[C_{63}H_{51}ClOP_3Ru]^+$ measured in methanol.
- fig. S16. Positive-ion ESI-MS spectrum of $[3a]^+$ [C₇₅H₆₂O₅P₃RuS]⁺ measured in methanol. fig. S17. Positive-ion ESI-MS spectrum of $[3b]^+$ [C₇₀H₅₆O₂P₃RuS]⁺ measured in methanol.
- fig. S18. Positive-ion ESI-MS spectrum of $[4a]^+$ [C₆₈H₅₇Cl₂CuO₄P₃Ru]⁺ measured in methanol.
- fig. S19. Positive-ion ESI-MS spectrum of [**4b**]⁺ [C₆₃H₅₁Cl₂CuO4²3Ru]⁺ measured in methanol.
- fig. S20. Positive-ion ESI-MS spectrum of $[5]^+$ [C₆₃H₅₁OCIP₃Ru]⁺ measured in dichloromethane.
- fig. S21. Positive-ion ESI-MS spectrum of $[\mathbf{5}']^+$ [C₆₃H₅₀DOClP₃Ru]⁺ measured in dichloromethane.
- fig. S22. Positive-ion ESI-MS spectrum of $[6]^+$ [$C_{67}H_{57}$ ClO₂P₃Ru]⁺ measured in methanol.
- fig. S23. Positive-ion ESI-MS spectrum of $[7 + H]^+$ $[C_{70}H_{58}N_2O_6P_3Ru]^+$ measured in methanol.
- fig. S24. Positive-ion ESI-MS spectrum of $[\mathbf{8} + H]^+ [C_{72}H_{56}N_6O_4P_3Ru]^+$ measured in methanol. fig. S25. ¹H NMR spectrum (600.1 MHz) of complex **2a** in CD₂Cl₂ at RT.
- fig. S26. ³¹P NMR spectrum (242.9 MHz) of complex **2a** in CD_2CI_2 at RT.
- fig. S27. ¹³C NMR spectrum (150.9 MHz) of complex **2a** in CD₂Cl₂ at R1.
- fig. S28. ¹H NMR spectrum (300.1 MHz) of complex **2b** in CD₂Cl₂/CD₃OD (v/v = 3/1) at RT.
- fig. S29. ³¹P NMR spectrum (121.5 MHz) of complex **2b** in CD₂Cl₂/CD₃OD (v/v = 3/1) at RT.
- fig. S30. ¹³C NMR spectrum (75.5 MHz) of complex **2b** in CD₂Cl₂/CD₃OD (v/v = 3/1) at RT.
- fig. S31. ¹H NMR spectrum (500.2 MHz, CD₂Cl₂) of complex **3a** at RT.
- fig. S32. ³¹P NMR spectrum (202.5 MHz, CD_2Cl_2) of complex **3a** at RT.

- fig. S34. ¹H NMR spectrum (300.1 MHz, CD₂Cl₂) of complex **3b** at RT. fig. S35. ³¹P NMR spectrum (121.5 MHz, CD₂Cl₂) of complex **3b** at RT. fig. S36. ¹³C NMR spectrum (75.5 MHz, CD₂Cl₂) of complex **3b** at RT. fig. S37. ¹H NMR spectrum (400.1 MHz, CD₂Cl₂) of complex **4a** at RT. fig. S38. ³¹P NMR spectrum (161.9 MHz, CD₂Cl₂) of complex **4a** at RT. fig. S39. ¹³C NMR spectrum (100.6 MHz, CD₂Cl₂) of complex **4a** at RT. fig. S40. ¹¹H NMR spectrum (600.1 MHz, CD₂Cl₂) of complex **4b** at RT. fig. S41. ³¹P NMR spectrum (242.9 MHz, CD₂Cl₂) of complex **4b** at RT.
- fig. S42. ¹³C NMR spectrum (150.5 MHz, CD_2Cl_2) of complex **4b** at RT.

fig. S33. ¹³C NMR spectrum (125.8 MHz, CD₂Cl₂) of complex **3a** at RT.

- fig. S43. In situ ^1H NMR spectrum (600.1 MHz, CD_2Cl_2) of complex **5** at 0°C.
- fig. S44. In situ ³¹P NMR spectrum (242.9 MHz, CD_2Cl_2) of complex **5** at 0°C.
- fig. S45. In situ ¹³C NMR spectrum (150.9 MHz, CD_2Cl_2) of complex **5** at 0°C.
- fig. S46. In situ ¹H NMR spectrum (500.2 MHz, CD₂Cl₂/CH₃COOD = 10/1) of complex **5**' at 0°C.
- fig. S47. In situ ³¹P NMR spectrum (202.5 MHz, CD₂Cl₂/CH₃COOD = 10/1) of complex **5**' at 0°C.
- fig. S48. ¹H NMR spectrum (300.1 MHz, CD₂Cl₂) of complex **6** at RT. fig. S49. ³¹P NMR spectrum (121.5 MHz, CD₂Cl₂) of complex **6** at RT.
- fig. S50. ¹³C NMR spectrum (75.5 MHz, CD_2Cl_2) of complex **6** at RT.
- fig. S51. ¹H NMR spectrum (600.1 MHz, CD_2Cl_2) of complex **7** at RT.
- fig. S52. ³¹P NMR spectrum (242.9 MHz, CD_2Cl_2) of complex **7** at RT.
- fig. S53. ¹³C NMR spectrum (150.9 MHz, CD₂Cl₂) of complex **7** at RT.
- fig. S54. ¹H NMR spectrum (400.1 MHz, CDCl₃) of complex ${f 8}$ at RT.
- fig. S55. ^{31}P NMR spectrum (161.9 MHz, CDCl_3) of complex ${\bf 8}$ at RT.
- table S1. Crystal data and structure refinement for 2a, 3b, and 4a.
- table S2. Crystal data and structure refinement for 6, 7, and 8.

table S3. Response to the questions raised in the checkCIF reports of complexes **4a**, **7**, and **8**. table S4. Thermal decomposition data of complexes **2** to **4** and **6** to **8** in the solid state. Supplementary Materials and Methods

data file S1. CIF files for complexes 2a, 3b, 4a, 6, 7, and 8.

data file S2. Cartesian coordinate–optimized structures for ASE, NICS, and AICD calculations. data file S3. Cartesian coordinate–optimized structures for mechanism studies. References (52–60)

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