Thermodynamic and reactivity studies of a tin corrole–cobalt porphyrin heterobimetallic complex†

Zikuan Wang, Zhengmin Yao, Zeyu Lyu, Qinsi Xiong, Bingwu Wang and Xuefeng Fu†

A heterobimetallic complex, (TPFC)Sn–Co(TAP) (TPFC = 5,10,15,20-tetrakis(p-methoxyphenyl)porphyrin), was synthesized. The complex featured a Sn–Co bond with a bond dissociation enthalpy (BDE) of 30.2 ± 0.9 kcal mol\(^{-1}\) and a bond dissociation Gibbs free energy (BDFE) of 21.0 ± 0.2 kcal mol\(^{-1}\), which underwent homolysis to produce the (TPFC)Sn radical and (TAP)Co\(^{n+}\) under either heat or visible light irradiation. The novel tin radical (TPFC)Sn, being the first four-coordinate tin radical observed at room temperature, was studied spectroscopically and computationally. (TPFC)Sn–Co(TAP) promoted the oligomerization of aryl alkynes to give the insertion products (TPFC)Sn–(CH=CH\(\text{Ar}\))\(_n\)-Co(TAP) \((n = 1, 2,\) or 3) as well as 1,3,5-triarylbenzenes. Mechanistic studies revealed a radical chain mechanism involving the (TPFC)Sn radical as the key intermediate.

Introduction

Heterobimetallic complexes featuring metal–metal bonds are interesting compounds possessing unconventional bonding patterns, which enable them to act as catalysts,\(^1\) magnets,\(^2\) supramolecular building blocks\(^3\) and molecular precursors for chemical vapor deposition (CVD).\(^4\) Transition metal (TM)–tin complexes are perhaps one of the most representative classes of such compounds, given the vast scope of their catalytic applications.\(^5,6\) Although the introduction of TM–tin bonds often leads to a marked enhancement in the rate of catalysis compared to the parent TM compounds,\(^5,6\) in most cases the tin-containing group just acts as a spectator ligand, which modifies the reactivity of the TM center through electronic effects rather than directly participating in the reactions. The strong dative and back-donating bonds between the TM and tin atoms decrease the availability of orbitals of the tin atom towards exogenous substrates, and also diminish the possibility of TM–tin bond dissociation. Rational design of TM–tin complexes featuring weak metal–metal bonds might thus help in revealing the “non-innocent” character of stannyl ligands and possibly lead to novel reaction modes which are not attainable by mononuclear TM systems.

Previously, we reported the synthesis of corrole Ge\(\text{m}\) radicals,\(^9\) Ge\(\text{n}\) cation complexes,\(^10\) and a corrole Sn\(\text{n}\) complex.\(^11\) However, the preparation of high valent corrole tin complexes with well-exposed tin centers, i.e. corrole Sn\(\text{m}\) radicals and Sn\(\text{n}\) cations, is impeded by the inert pair effect, i.e. the relativistic contraction of the Sn 5s shell and a corresponding lowering of the 5s energy level, which raises the redox potentials of tin complexes relative to their germanium analogs. Cleavage of weak TM–tin bonds bypasses this difficulty and represents a promising route towards corrole tin radicals. Porphyrin cobalt emerges as an ideal candidate for the construction of weak TM–tin bonds due to (1) the similar electronegativities of Sn (1.96) and Co (1.88),\(^12\) favoring homolytic Sn–Co bond cleavage, (2) the availability of a Co 3d\(_z^2\) orbital as the frontier orbital for σ bonding,\(^13\) and (3) the known ability of the Co–C bonds of organocobalt(m) porphyrin complexes (bond dissociation enthalpies (BDE) ~ 20 kcal mol\(^{-1}\))\(^14\)–\(^16\) to undergo homolysis under both thermal\(^14\) and photochemical\(^17\) conditions. Hence, we envisioned that a complex (cor)Sn–Co(por) (cor = corrole, por = porphyrin) might possess a sufficiently weak Sn–Co bond to be a viable precursor to the hitherto elusive (cor)Sn radical species, featuring a formally trivalent tin center. Moreover, the coproduct (por)Co\(^{n+}\) might assist in substrate activation promoted by (cor)Sn, resulting in novel reaction pathways.\(^18\)

Complexes that feature Sn–Co bonds have been documented previously. While formally double bonded Sn=Co complexes are known, e.g. CpCo(C\(_5\)H\(_4\))=Sn(CH(TMS)\(_3\))\(_2\),\(^19\) most Sn–Co complexes feature covalent, single Sn–Co bonds where back-bonding is believed to be relatively unimportant. Although porphyrin,\(^20\) dimethylglyoxime derivatives\(^21\) and CO\(^2\) have been used as the ligands of the cobalt center, practically all
heterobimetallic Sn–Co complexes feature alkyl or halogen-substituted tin centers. The proposed [cor]Sn–Co[por] complex, with a nitrogen-ligated pentacoordinate tin atom, might thus reveal novel aspects of the chemistry of such bimetallic complexes.

Herein we report the synthesis of the heterobimetallic complex (TPFC)Sn–Co(TAP) (1), TPFC = 5,10,15-tris[pentafluorophenyl]corrole, TAP = 5,10,15,20-tetrakis[p-methoxyphenyl]porphyrin; Scheme 1). Upon heating or visible light irradiation of 1, the active (TPFC)Sn (2) radical as well as (TAP)CoII (3) were generated. The BDE and bond dissociation free energy (BDFE) of the Sn–Co bond were estimated to be 30.2 ± 0.9 kcal mol⁻¹ and 21.0 ± 0.2 kcal mol⁻¹ at 298 K, respectively. The reactivity was exploited in the reaction with alkynes, revealing 1 as a potential alkyne cyclotrimerization catalyst.

Results and discussion

Synthesis, characterization and DFT studies of 1

1 was synthesized by the reaction of the nucleophilic Na[(TPFC)Sn³] (4) with (TAP)CoCl (5) in 77% yield (Scheme 1). The complex showed characteristic¹H NMR signals of the corrole and the porphyrin rings (Fig. S2†), which were shifted away from the typical chemical shift ranges of monomeric (TPFC)Sn–X and (TAP)Co–X complexes, respectively, due to the combined ring current effects of the two aromatic macrocyclic ligands. Interestingly, 1 was ¹¹⁹Sn NMR silent, due to the extensive quadrupolar relaxation by the ⁵⁹Co (I = 7/2) nucleus.

Single crystal XRD results unambiguously confirmed the direct Sn–Co bonding (Fig. 1; the Sn–Co bond length was 2.4559(10) Å, which was within the range of typical Sn–Co single bonds⁴–⁷). The Sn–Co bond was approximately perpendicular to both the corrole and the porphyrin planes, and the two ligand planes were aligned face-to-face. The Sn atom protruded noticeably (0.872 Å) from the N₆-plane of the corrole ligand, comparable with known (TPFC)Sn–X complexes (0.776–0.798 Å),¹¹ while the Co atom was essentially within the porphyrin plane (the distance to the N₁–N₂–N₃–N₄ mean plane was 0.102 Å).

DFT calculations [B3LYP/def2-TZVPP//BP86/def2-TZVP(-f)] were performed to shed light on the bonding between the Sn and Co atoms. The DFT optimized geometry agreed very well with the XRD structure (see Fig. 2 and Table S8†). The Sn–Co bond was a covalent single bond according to natural bonding orbital (NBO) calculations (a Wiberg bond order of 0.60). The Sn–Co bonding NBO comprised 41.5% Sn character and 58.5% Co character, and the sp³–d and sp²–d hybridizations were adopted by the Sn and Co atoms, respectively (Fig. 2). Thus, the polarity of the bond was best described as Sn⁺–Co⁶⁻ with a natural ionicity of ᵢₛₙᵢₛₙₒ = –0.17, a value comparable to those of aliphatic C–H bonds which typically have ᵢᵢ₋ᵢ₋ = 0.16–0.19.²¹ Nevertheless, non-Lewis (i.e. ionic) resonance forms contributed significantly to Sn–Co bonding: the second-order perturbative estimate of interaction energy between the bonding and antibonding NBOs of the Sn–Co bond amounted

![Fig. 1](image1.png)

**Fig. 1** Solid state structure of 1. Hydrogen atoms are omitted for clarity. Selected bond lengths (Å): Sn1–Co1 2.4559(10), Sn1–N5 2.120(5), Sn1–N6 2.107(5), Sn1–N7 2.122(5), Sn1–N8 2.108(5), Co1–N1 1.957(4), Co1–N2 1.960(5), Co1–N3 1.967(4), and Co1–N4 1.955(5).

![Fig. 2](image2.png)

**Fig. 2** Isosurface plot of the Sn–Co σ-bonding NBO of 1 (isovalue = 0.02).
to 31.0 kcal mol\(^{-1}\), leading to a strong occupation (\(n = 0.463\)) of the antibonding NBO. Upon formation of the Sn–Co bond from the open-shell (TPFC)Sn and (TAP)Co fragments, a significant electron density depletion was observed at the tin center, accompanied by an electron density accumulation at the cobalt center (Fig. S45†), which confirmed the charge shift as suggested by NBO analysis. Slight density accumulation and depletion were also observed on the corrole and porphyrin ligands, respectively, which were likely due to changes in metal-to-ligand delocalization effects. The Sn 5d\(_x\) orbitals were nearly unoccupied (\(n < 0.003\)), indicating that Co 3d → Sn 5d back-bonding is negligible. A second-order perturbative estimate yielded a total Co 3d → Sn 5d \(\pi\) back-bonding energy of 1.6 kcal mol\(^{-1}\).

The energetics of Sn–Co bond cleavage pathways for 1, namely heterolytic formation of Sn\(^{II}\) + Co\(^{III}\) and Sn\(^{IV}\) + Co\(^{I}\), and homolytic formation of Sn\(^{III}\) + Co\(^{II}\), were subsequently calculated (Table 1). Apparently, the homolysis pathway was the only thermally accessible mode at practical temperatures, and might also be viable under photochemical conditions. In contrast, the heterolysis pathways were so hindered that even visible light (\(\lambda > 400\) nm, or a photon energy of less than 71 kcal mol\(^{-1}\)) was unlikely to overcome the unfavorable thermodynamics.

**Photolysis of the Sn–Co bond**

Experimentally, the Sn–Co bond of 1 was found to be remarkably photo-labile. An attempt to record the UV-Vis spectrum of 1 in aerated toluene revealed a rapid spectral change within 10 seconds, with isosbestic points at 414, 457, 493 and 610 nm (Fig. 3a). Since 1 was reasonably stable at room temperature under air within a time scale of several hours, the transformation was apparently triggered by light. To elucidate the composition of the photoproducts, we repeated the experiment in C\(_6\)D\(_6\) using a Xe lamp (equipped with a 420–780 nm band pass filter) as the light source, and monitored the reaction course by \(^1\)H NMR. When 1 atm of dry O\(_2\) was used, the signals of 3 and a new (TPFC)Sn species appeared (Fig. S5a†). The chemical shifts of the latter species (8.88, 8.34, 8.14, and 7.82 ppm) resembled those of (TPFC)Sn–CH\(_2\)–Sn(TPFC),\(^{11}\) suggesting that the species was the \(\mu\)-oxo dimer [(TPFC)Sn–O–Sn(TPFC)] (6). This assignment was confirmed by the independent synthesis of 6 from aerobic oxidation of 4 (see the ESI†).

<table>
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<tr>
<th>Product(^a)</th>
<th>(\Delta G) (kcal mol(^{-1}))</th>
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<tr>
<td>[Sn(^{II})](^+) + [Co(^{III})](^+)</td>
<td>75.1</td>
</tr>
<tr>
<td>[Sn(^{IV})](^+) + [Co(^{I})]</td>
<td>19.9</td>
</tr>
<tr>
<td>[Sn(^{III})](^+) + [Co(^{II})]</td>
<td>113.2</td>
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\(^a\) Formal oxidation states are given, which might differ from the actual oxidation states.

**Fig. 3** (a) Photo-oxidation of 1 in aerated toluene over a period of 11 s monitored by stopped-flow UV-Vis absorption spectroscopy. (b) Comparison of the UV-Vis spectra of 3 and 6 with that of the product mixture.
structured Q band absorptions around 550–600 nm, but is in accordance with TDDFT calculated absorptions of 2 (Fig. 5a). Thus, we tentatively attribute the extra signal to complex 2. Such reactivity was also proposed for the photolysis of (py)(dmgH)$_2$Co–SnPh$_3$ (dmg = dimethylglyoximato)$^{18}$ and (OEP)Co–SnPh$_3$ (OEP = 2,3,7,8,12,13,17,18-octaethylporphyrin),$^{20}$ but only indirect evidence for the involvement of tin radicals (by halogen abstraction from haloalkanes, for example) was provided.

Notably, the transient absorption signal uniformly decayed towards zero within a couple of milliseconds (Fig. 5b), suggesting regeneration of 1 through recombination of 2 and 3. The decay kinetics was followed spectrosopically both in the absence and in the presence of various excess amounts of 3. The decay rate was enhanced upon the addition of 3 (Fig. 5c), and exhibited first order dependence on the concentration of the latter (Fig. 5d). The recombination rate constant of 2 and 3 was thus estimated to be $(9.2 \pm 0.2) \times 10^7$ L (mol$^{-1}$ s$^{-1}$) at 298 K, which implied a diffusion-controlled process. Under the assumption of strictly diffusion-controlled behavior, the activation enthalpy of recombination could be taken as the activation enthalpy of the viscous flow of toluene,$^{26}$ which was known accurately to be 1.93 kcal mol$^{-1}$. Thus, the activation entropy of recombination, $-15.60 \pm 0.04$ cal (mol$^{-1}$ K$^{-1}$), could be obtained. In case the diffusion-controlled conditions did not hold.

Fig. 4 Solid state structure of 6. Hydrogen atoms are omitted for clarity. Selected bond lengths (Å) and angles (°): Sn1–O1 1.924(3), Sn2–O1 1.934(3), and Sn1–O1–Sn2 134.46(14).

Fig. 5 Laser flash photolysis of 1. (a) Transient spectrum of a 5.2 × 10$^{-6}$ mol L$^{-1}$ toluene solution of 1, recorded at 10 ns after the laser pulse, compared against the experimental spectra of 1 and 3, and calculated absorptions of 2; (b) time-resolved decay of the transient signal, with [1] = 2.6 × 10$^{-6}$ mol L$^{-1}$; (c) transient kinetic decay plots of 2.6 × 10$^{-6}$ mol L$^{-1}$ of 1 in the presence of various concentrations of 3, measured at 404 nm; and (d) first-order plot of apparent recombination rate constants against [3] (the non-zero intercept was due to the fact that the amount of 3 formed through photolysis was comparable to that of excess 3; see the ESI†).

\[(py)(dmgH)_2Co–SnPh_3\]
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Fig. 6 Side-view of (TPFC)Ge (a) and (TPFC)Sn (b), and spin density plots of (TPFC)Ge (c) and (TPFC)Sn (d).

exactly, slight underestimations of both the activation enthalpy and the activation entropy were expected.

Reported tin radicals mostly feature tricoordinate tin centers with pyramidal or planar conformations, depending on the substituents. Tetracoordinate tin radicals, to the best of our knowledge, have been elusive, except for radiogenic radical cations based on organostannanes, which are only stable in solid matrices at low temperatures. Calculations revealed that adopted a domed structure in close analogy with (TPFC)Ge reported by us, although the doming effect was more severe for the tin complex due to the larger atomic radius of tin (Fig. 6a and b). The two complexes also shared qualitatively similar electronic structures, where significant spin density was delocalized into the \( \psi(\text{a}_{\text{Sn}}) \) corrole \( \pi \) orbital (Fig. 6c and d). Nevertheless, this effect was much more pronounced for Sn (NPA spin population on the Sn atom = 0.11) than for Ge (spin population on the Ge atom = 0.45). Consequently, in contrast to (TPFC)Ge, the dominant resonance form of 2 was \( \text{(TPFC)}^{+}\text{Sn}^{+} \) as indicated by NBO calculations, suggesting significant non-innocent ligand character. The anti-bonding interaction of the Sn sp orbital with the corrole \( \psi(\text{a}_{\text{Sn}}) \) orbital shifted the latter upwards, explaining the red-shifted Soret band appearing at 450–500 nm (Fig. S46†).

The above results may suggest that 2 should behave as a ligand-centered radical. Experiments, in contrast, revealed a tin radical-like behavior. Photolyzing a toluene solution of 1 in the presence of \( N\)-tert-butyl-\( \alpha \)-phenylnitrone (PBN) at \(-50^\circ \text{C}\) led to rapid formation of a signal typical of PBN-trapped radicals (Fig. 7a). At \(-20^\circ \text{C}\), the signals were sharpened due to an increase of the molecular tumbling rate and concomitantly better isotropic averaging, revealing pairs of \( ^{117,119}\text{Sn} \) satellites (Fig. 7b). A spectral fit assuming the isotropic fast-motion regime yielded the parameters \( g = 2.0048 \), \( A^{(119}\text{Sn}) = 5.5 \text{ G} \), \( A^{(113}\text{Sn}) = 14.1 \text{ G} \), and \( A^{(114}\text{Sn}) = 18.0 \text{ G} \). The \( A^{(113}\text{H}) \) was considerably larger than most PBN adducts, including adducts of oxygen (2–3 G) and carbon radicals (3–4 G), while some heavier group 14 radicals were reported to give \( A^{(1}\text{H}) \) values of comparable magnitude (e.g. \( \text{Ph}_2\text{C}^{\text{13}}\text{Ge}, 10.9 \text{ G} \)). Thus, 2 must have been trapped by PBN through the tin atom, rather than one of the ligand atoms, which implied that the tin atom was more reactive than the ligand. Apparently, the favorable steric and limited structural reorganization energy of the domed tin atom overruled the larger spin density of the ligand. This was also in accord with the observed formation of 6 from photo-oxidation of 1, as opposed to ligand oxidation products. The EPR signal due to 3 was observed upon cooling to \(-150^\circ \text{C}\) (Fig. S13†).

Thermolysis of the Sn–Co bond

With evidence for photolysis in hand, we then investigated the thermal lability of 1. Unfortunately, the PBN trapping experiment was not feasible under thermal conditions because the (TPFC)Sn–PBN adduct was unstable above \(-20^\circ \text{C}\). Thus, the radical trap \( 2,2,6,6\)-tetramethylpiperidin-1-yl)oxyl (TEMPO) was used instead. Heating a \( \text{C}_6\text{D}_6 \) solution of 1 and excess TEMPO at \( 70^\circ \text{C} \) gave (TPFC)Sn–TEMPO (8) and 3 as the sole products as evidenced by \( ^{1}\text{H} \) NMR (Fig. S14†). The reaction rate was first order on 1 and zeroth order on TEMPO (Fig. 8), suggesting homolytic Sn–Co bond dissociation followed by fast trapping of the (TPFC)Sn (2) radical by TEMPO (for further evidence of the homolytic nature of the dissociation, see the ESI†). An activation enthalpy of 32.1 \( \pm \) 0.9 kcal mol\(^{-1}\) and an activation entropy of 15 \( \pm \) 3 cal (mol\(^{-1}\) K\(^{-1}\)) were determined by kinetic measurements in toluene-\( d_8 \) at 70–100 °C (Fig. 9). The large
activation entropy confirmed the involvement of a dissociative rate-limiting step, and the value was close to that reported for homolytic dissociation of another metal–metal bond complex stabilized by a conjugated planar ligand, (tmtaa)Rh(tmtaa) (10 ± 1 cal (mol\(^{-1}\) K\(^{-1}\))), tmtaa = dibenzotetramethylaza[14]annulene).\(^{33}\)

Although we are unaware of any existing quantitative kinetic measurements of the homolysis barriers of Sn–Co bonds, the complex (OEP)Co–SnPh\(_3\) was reported to have a half-life of ca. 48 h at 120 °C with (TTP)Co\(^{11}\) (TTP = 5,10,15,20-tetra(p-tolyl) porphyrin) as a radical trap,\(^{28}\) which indicates a stronger Sn–Co bond than that in 1. This might partly be attributed to the spin delocalization in 2 (Fig. 6), which favored the Sn–Co bond homolysis of 1 compared to (OEP)Co–SnPh\(_3\).

**Estimation of the Sn–Co bond energy**

With the knowledge of the kinetics of both forward and backward directions of the reaction, reliable results on the thermodynamics of Sn–Co bond dissociation could be obtained (Table 2). The bond dissociation enthalpy (BDE) of the Sn–Co bond was estimated to be \(\Delta H = 30.2 \pm 0.9\) kcal mol\(^{-1}\) by subtracting the backward activation enthalpy, 1.93 kcal mol\(^{-1}\), from the forward activation enthalpy. A bond dissociation entropy of \(\Delta S = 31 \pm 3\) cal (mol\(^{-1}\) K\(^{-1}\)) was obtained similarly. It might be interesting to note that exactly half of the dissociation entropy was released in the transition state. The bond dissociation Gibbs free energy (BDGE) was calculated to be \(\Delta G = 21.0 \pm 0.2\) kcal mol\(^{-1}\), in excellent agreement with the DFT value (Table 2).\(^{14}\)

**Reaction of 1 with alkynes**

It is well known that stannyl radicals readily undergo addition to alkynes, a fact that was exploited in reactions such as radical hydrostannylations as well as cascade ring closing reactions.\(^{35}\) Indeed, 1 was reactive towards p-fluorophenylacetylene under Xe lamp irradiation to form mono-, bis- and tris-insertion products, delivering complexes 9–11 (Scheme 2). The reaction proceeded under either thermal or photochemical conditions as shown by \(^1\)H NMR and HR-ESI-MS (Fig. S19 and S20†). Column chromatography of the reaction mixture gave, apart from 9–11, three additional fractions: a mixture of two organocobalt porphyrin complexes (12 and 13), whose structures were confirmed by independent syntheses (see the ESI†); a complex mixture of various organotin corrole complexes in which 14 was identified by HR-ESI-MS; and a cyclotrimerization product, 1,3,5-tris[p-fluorophenyl]benzene (15) (Scheme 2).

**Table 2** Kinetic and thermodynamic parameters for Sn–Co bond dissociation of 1

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<thead>
<tr>
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<th>(\Delta H^b)</th>
<th>(\Delta S^b)</th>
<th>(\Delta G_{298}^b)</th>
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<tr>
<td>Dissociation barrier</td>
<td>32.1 ± 0.9</td>
<td>15 ± 3</td>
<td>27.6 ± 0.2</td>
</tr>
<tr>
<td>Recombination barrier</td>
<td>1.93</td>
<td>-15.60 ± 0.04</td>
<td>6.58 ± 0.01</td>
</tr>
<tr>
<td>Thermodynamics of dissociation (exp.)</td>
<td>30.2 ± 0.9</td>
<td>31 ± 3</td>
<td>21.0 ± 0.2</td>
</tr>
<tr>
<td>Thermodynamics of dissociation (DFT)</td>
<td>38.9</td>
<td>63.8</td>
<td>19.9</td>
</tr>
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</table>

\(^{a}\) In kcal mol\(^{-1}\); \(^{b}\) In cal (mol\(^{-1}\) K\(^{-1}\)). \(^{c}\) Obtained by extrapolation of Fig. 9 to 298 K. \(^{d}\) Taken as the activation enthalpy of viscous flow (see text) assuming negligible error. \(^{e}\) Obtained by subtraction of recombination parameters from those of dissociation. \(^{f}\) See ref. 34 for explanations of the discrepancy with experiment.
To our delight, the turnover number (TON) of the cyclo-trimerization product reached 4.3 when \( p \)-methoxyphenylacetylene was used as a substrate. The presence of MeO groups also introduced sufficient polarity differences between the insertion products \( 9^\circ - 11^\circ \) which allowed us to isolate these complexes. Stoichiometric photolysis of the insertion products revealed that \( 9^\circ \) was cleanly photolyzed to yield \( 1 \) and \( p \)-methoxyphenylacetylene, while \( 10^\circ \) reacted sluggishly to give \( 1 \) and an undefined mixture, where neither \( p \)-methoxyphenylacetylene nor the cyclotrimerization product \( 15^\circ \) was found. Unexpectedly, photolysis of \( 11^\circ \), which already possessed an alkyne trimer motif between the metal centers, only gave a small amount (maximum yield 8%) of \( 15^\circ \) and regenerated \( 2 \), along with \( 1 \) and \( 3 \) in 41% and 55% yields, respectively.

Based on the evidence above, we propose that the alkyne insertion initially occurred through the sequential addition of alkyne molecules to the stannyl radical \( 2 \), followed by reversible capturing of the resulting carbon radicals by \( 3 \) (Scheme 2). The latter process regulated the radical concentration in a way similar to that in cobalt-mediated living radical polymerization (CMRP), where the chain radicals generated from successive addition of olefin monomers to an initiator radical are reversibly captured by a cobalt(II) complex, reducing the concentration of the former and thereby suppressing biradical termination. The addition of the third alkyne molecule led to two stereoisomers, \( 11^\circ R - c i s \) and \( 11^\circ R - t r a n s \). Back-biting of \( 11^\circ R - c i s \) yielded the cyclotrimerization product \( 15 \) and regenerated \( 2 \), the cyclization of \( 11^\circ R - t r a n s \) necessarily required a trans-cis isomerization. This explains the low yield of \( 15^\circ \) in the stoichiometric photolysis of \( 11^\circ \), which involved \( 11^\circ R - t r a n s \) rather than \( 11^\circ R - c i s \) (Fig. S38†). The formation of \( 14 \) was due to hydrogen atom transfer (HAT) from \( 11^\circ R - t r a n s \) to \( 3 \), which deactivated the catalyst irreversibly. The by-product, \( \text{TAP-Co-H} (16) \), underwent fast reaction with excess alkyne to yield \( 12 \) and \( 13 \), resembling the catalytic chain transfer (CCT) process in CMRP where insertion of olefins to the Co–H bond is observed instead.

**DFT studies of alkyne cyclotrimerization**

To confirm the proposed mechanism as well as to gain more insight into the back-biting process of \( 11^\circ R - c i s \), we performed a computational study of the reaction using \( p \)-fluorophenylacetylene as a model substrate (Fig. 10). The addition
of the alkyne to 2 was moderately endergonic ($\Delta G = 12.5$ kcal mol$^{-1}$), consistent with the observed reversibility of the formation of 9 from 2. Subsequent alkyne additions, however, were exergonic ($\Delta G = -9.1$ and $-16.1$ kcal mol$^{-1}$) due to the formation of C–C bonds, which drove the aforementioned monoaddition equilibrium to the product side. It is worth noting that while the second alkyne addition gave exclusively the trans isomer 10 due to favorable steric effects of the transition state, the barriers for the formation of 11-rad-trans and 11-rad-cis were almost identical. Accepting 11-rad-trans as a non-productive species, the computed energetics suggested a maximum TON of 2.9 for the alkyne trimerization at 298 K, which was very close to the experimental value (1.6).

The back-biting of 11-rad-cis could occur through two possible pathways: (a) a 5-exo-trig cyclization, which is Baldwin allowed but leads to a five-membered ring as opposed to the observed six-membered ring; or (b) a 6-endo-trig cyclization, which is Baldwin disallowed but in which the product 17-rad (Fig. 10) is in accord with the observed structure of the final product 15. In practice, however, the Baldwin disallowed route (b) showed a much lower reaction barrier than that of route (a), thanks to thermodynamic control. Further calculations showed that if route (a) were operative, it would lead to the 1,2,4-cycloaddition product through ring expansion (Fig. S47†). Since the 1,2,4-product was not observed experimentally, this further confirmed the predominance of route (b) over route (a). Sn–C bond scission of 17-rad readily afforded the aromatized cyclotrimerization product 15 (Fig. 10).

Since only ESI-MS evidence for 14 was available, the experimentally observed HAT process might as well occur from 11-rad-cis or the cyclized product 17-rad instead of 11-rad-trans, because all three reactions would give products with the same molecular formula. HAT from 11-rad-cis to 3 can be safely ruled out since the barrier of the 6-endo-trig cyclization (1.1 kcal mol$^{-1}$) was too low for HAT or any bimolecular reaction to be competitive (typical barriers of HAT processes from carbon radicals to porphyrin cobalt are 10–15 kcal mol$^{-1}$).16 17-rad, on the other hand, possessed an HAT-active C–H bond that was so well-shielded as to preclude any possibility for the HAT process (Fig. S48†). This conclusively demonstrated that the observed CCT product originated exclusively from 11-rad-trans, rather than 11-rad-cis or 17-rad.

Conclusions

In conclusion, we have synthesized an unusual heterobimetallic complex, 1, that possesses a covalent, weakly polar Sn–Co bond. The Sn–Co bond undergoes homolysis under thermal and photochemical conditions, releasing an active, four-coordinate stannyl radical 2 in a reversible manner. The thermodynamics and kinetics of the homolysis were studied extensively using a combination of various spectroscopic methods, yielding a Sn–Co BDE of 30.2 ± 0.9 kcal mol$^{-1}$ and a BDE of 21.0 ± 0.2 kcal mol$^{-1}$ at 298 K.

Four-coordinate tin radicals such as 2 are rare and unstable due to the reluctance of the less Lewis acidic Sn(II) center to accept a fourth ligand. However, the use of the rigid four-coordinate TPFC ligand forced a four-coordinate situation around the tin atom, leading to the first observation of a four-coordinate tin radical at room temperature, with a lifetime of at least several milliseconds. Moreover, 2 exhibits significant noninnocent ligand character contributing further to its stability, yet with uncompromised and essentially metal-centered reactivity. Taking advantage of the reactivity of 2, we show experimentally and computationally that 1 is catalytically active towards the cyclotrimerization of aryl alkynes.

Conflicts of interest

There are no conflicts of interest to declare.

Acknowledgements

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gratefully acknowledge Prof. Ming-Tian Zhang and Ms Han Li for laser flash photolysis experiments, and Mr Jin Xiong for assistance with single crystal structure refinement.

Notes and references

30 In corrole chemistry it is customary to denote the ligand π orbitals with the irreducible representations of the respective orbitals of porphyrin (Ds): A. Ghosh, Chem. Rev., 2017, 117, 3798. In the present paper we give the symmetry representations assuming a symmetric domed corrole (C₄), followed by the approximate D₄h representations in parentheses.
34 Note that the SMD solvation model was used in the calculation of ΔH, which already included the effect of solvation entropy. Unfortunately, this contribution could not be separated out, leading to a positive error in ΔH because the solvation entropy discourages dissociation due to an increase of the solvent accessible surface area. On the other hand, the ideal gas model was used in the calculation of ΔS, as is usually done, resulting in a neglect of solvation contribution (which is negative) and thus a positive error. The two errors exactly canceled out each other, and so ΔG was accurately reproduced.