Abstract: The use of molecular oxygen in palladium-catalyzed oxidation reactions is highly widespread in organic chemistry. However, the direct reoxidation of palladium by \( \text{O}_2 \) is often kinetically unfavored, thus leading the deactivation of the palladium catalyst during the catalytic cycle. In the present work, we report a highly selective palladium-catalyzed carbocyclization of bisallenes to seven-membered heterocycles under atmospheric pressure of \( \text{O}_2 \). The use of a homogenous hybrid catalyst (Co(salophen)-HQ, HQ = hydroquinone) significantly promotes efficient electron transfer between the palladium catalyst and \( \text{O}_2 \) through a low-energy pathway. This aerobic oxidative transformation shows broad substrate scope and functional group compatibility and allowed the preparation of \( \text{O}^- \)-containing seven-membered rings in good yields in most cases.

Over the past decades, palladium-catalyzed oxidations have emerged as powerful and valuable tools in modern organic synthesis.[1] Various protocols have been developed for the assembly of carbon-carbon and carbon-heteroatom bonds that provide useful applications in biology, medicine, and materials science.[2] However, in most cases the use of stoichiometric oxidants such as \( \text{Cu}^2+ \), \( \text{Ag}^+ \), and peroxides often leads to poor selectivity, low atom economy, and considerable amounts of undesired waste. In the perspective of synthetic organic chemistry, molecular oxygen is an inexpensive, abundant, and highly atom-efficient oxidant, which does not generate any toxic by-products, thus fulfilling the requirements of “green chemistry”.[3] Despite significant advances in palladium-catalyzed aerobic oxidations, a severe problem is that fast aggregation of palladium black from the active palladium species (\( \text{Pd}-\text{H} \) or \( \text{Pd}^0 \)) slows down and finally stops the homogenous reaction.[4] Extensive endeavors for solving this problem have focused on the use of air-stable ligands to restrain \( \text{Pd}^0 \) precipitation during the catalytic cycle (Scheme 1a).[5] Moreover, Jiang and co-workers recently developed an elegant protocol that utilizes an efficient metal-organic framework (MOF) for the stabilization of the palladium catalyst in aerobic functionalizations (Scheme 1b).[6] Although a plethora of strategies to circumvent this oxidation problem have been reported, the development of highly active catalytic systems that enable mild \( \text{Pd}^0 \)-catalyzed aerobic oxidations continues to be an important topic in this area.

Our research group has a long-standing interest in palladium-catalyzed oxidations where molecular oxygen is used as a green oxidant.[7] However, as stated above, the direct reoxidation of palladium by molecular oxygen is often kinetically unfavored.[8] To solve this problem, we and others have employed coupled catalytic systems with electron transfer mediators (metal-macrocycle and quinone) to facilitate the relay of electrons between the palladium catalyst and \( \text{O}_2 \) (Scheme 1c).[8] These coupled catalytic systems have been demonstrated to be highly efficient in Wacker oxidations[9b,9c] alcohol oxida-
oxidative olefin functionalizations, and oxidative C–H activations where molecular oxygen is the oxidant. In 1993, a more efficient hybrid catalyst, involving a cobalt-porphyrin with pendant hydroquinone groups in one molecule, was reported by our group as well. Later on, an improved hybrid Schiff base-hydroquinone [Co(salophen)-HQ] as a redox relay catalyst for aerobic oxidation was reported. The latter bifunctional catalyst led to a faster reaction rate compared to the system with the quinone and metal-macrocycle as separate molecules in Pd-catalyzed aerobic carbocyclization of enallenes and dienallenes.

During the past decade, our research group has been involved in the development of palladium-catalyzed oxidative carbocyclizations of allenes bearing an additional unsaturated moiety, such as an olefin, alkyne, or allene. Interestingly, we found that an assisting group (AG), such as an olefin, alkyne or hydroxyl group, is required to make 6-membered rings from enallenes and allenynes and to make 7-membered rings from bisallenes due to the initial allenic C–H cleavage by Pd (Scheme 2a). Afterward, cascade carbocyclization and transmetallation with a nucleophile can occur and give the cyclic product. However, without an activating group, no reaction takes place with the distal π-bond to give the larger rings. This interesting neighboring group effect allows for palladium-catalyzed carbocyclization in a highly selective manner. We recently reported an efficient approach for the synthesis of seven-membered rings via olefin-assisted palladium-catalyzed carboxylation of bisallenenes. However, O-containing seven-membered rings were not explored in the previous study. Also, there is no example on the use of oxidative carbocyclization of bisallenes for the synthesis of useful organoboron molecules. Herein we report a general and efficient catalytic system for the synthesis of O-containing seven-membered heterocycles via cascade borylative carbocyclizations (Scheme 2b). This catalytic system involves the use of a Pd catalyst with a hybrid ETM catalyst for the biomimetic aerobic oxidations.

Initially, a class of bifunctional Schiff base-hydroquinone catalysts were prepared from the ligand precursor salicylaldehyde-hydroquinone, which was covalently synthesized according to our previous report. As shown in Scheme 3, the synthetic route to these oxidation catalysts commences with condensation of two equivalents of salicylaldehyde-hydroquinone with various diamines such as o-phenylenediamine, 4-chloro and 4-methoxy-o-phenylenediamine, ethylenediamine and N-(3-aminopropyl)-N-methylpropane-1,3-diamine. The resulting salen-hydroquinone ligands could be directly used for the next step without further purification. Subsequently, coordination of these ligands with 3d metal salts (Co, Fe, Ni, and Cu) afforded the corresponding hybrid catalysts (Cat. 1–9) in general yields of 78–95%.

At the outset of our investigations, the palladium-catalyzed aerobic oxidation of a readily accessible bisallene 1a with B(pin) 2a was chosen as the benchmark reaction (Table 1). In the absence of electron transfer mediators (ETMs), we did not observe any formation of the desired borylation product under aerobic conditions and the starting material 1a was recovered in 93% yield (entry 1). When catalytic amounts of p-benzoquinone (BQ) was added, the seven-membered carbocycle 3a was selectively formed in low yield (15%, entry 2). Notably, the use of}

![Scheme 2](image)

**Scheme 2.** (a) Palladium-catalyzed oxidative carbocyclization with an assisting group; (b) Current work.

![Scheme 3](image)

**Scheme 3.** Synthesis of Schiff base-hydroquinone hybrid catalysts.

![Diagram](image)

**Diagram.**

![Table 1](image)

**Table 1.** Evaluation of different electron transfer mediators (ETMs) for borylative carbocyclization of bisallene 1a.

| Entry | ETMs | Yield of 3a [%] |
|-------|------|----------------|
| 1     | none | 0              |
| 2     | 20 mol % BQ | 15          |
| 3     | 10 mol % Co(salophen), 20 mol % BQ | 48         |
| 4     | 10 mol % Co(salophen), 20 mol % HQ | 35         |
| 5     | 10 mol % Co(salophen)-HQ (Cat. 1) | 78         |
| 6     | 100 mol % BQ | 78          |

[a] Unless otherwise noted, the following reaction condition were employed with 1a (0.1 mmol, 1.0 equiv), B(pin) 2a (0.13 mmol, 1.3 equiv), Pd(DAC2) (5 mol%), ETMs (10–20 mol%) in 0.1 M acetone, O2 (1 atm) at room temperature (25 °C) for 30 h. (b) Yields were determined by 1H NMR using anisole as an internal standard. (c) 93% of starting material 1a was recovered. (d) <1% of starting material 1a was recovered.
of cobalt-based salophen complex with $p$-benzoquinone (BQ) or hydroquinone (HQ) as separate ETMs led to increased but moderate yields of 3a (48 and 35% yields, entries 3 and 4). To our delight, the bifunctional cobalt catalyst (Co(salophen)-HQ), combining a metal-macrocycle and a quinone moiety, was found to be a highly active catalyst and afforded the desired product 3a in 78% yield (entry 5). Importantly, this aerobic catalytic system appeared to be equally effective in comparison to the application of stoichiometric amounts of BQ as oxidant (78%, entry 6), therefore highlighting the advancement of this efficient and sustainable method.

It is noteworthy that the Schiff base-hydroquinone hybrid catalyst (Co(salophen)-HQ, Cat. 1) leads to significant improvement of the transformation of 1a to 3a (78% yield). This interesting result motivated us to further explore the catalytic performance of various hybrid catalysts synthesized above (vide supra). As shown in Figure 1, compared to Co-based catalyst (Cat. 1), other metal-based catalysts with Fe, Ni and Cu (Cat. 2–4) were less effective in this reaction. This result indicates that the metal center of the salen catalyst has a significant influence in the oxygen activation step. Notably, functional groups such as Cl (Cat. 5) or OMe (Cat. 6) on the diamine backbone resulted in little change of the reaction yields (77% and 74%, respectively). The hybrid catalysts with aliphatic diamine backbones, such as ethylenediamine (Cat. 7), (±)-trans-1,2-diaminocyclohexane (Cat. 8) and N-(3-aminopropyl)-N-methylpropane-1,3-diamine (Cat. 9) exhibited slightly lower reactivity (58–63%). Therefore, the bifunctional catalyst Co(salophen)-HQ was found to be the best performing catalyst in this aerobic carbocyclization to a seven-membered heterocycle.

After optimizing of the reaction conditions, we explored the substrate scope with the optimal oxidation catalyst Co(salophen)-HQ (Scheme 4). The benchmark borylative carbocyclization gave the corresponding product 3a in 75% isolated yield. With $R^2$ being a benzyl or a cyclohexyl group, the corresponding products 3b and 3c were obtained in 78% and 60% yields, respectively. Under the optimal aerobic reaction conditions, cyclohexyldiene bisallene 1d afforded 3d in only 30% yield. We attribute this diminished reactivity to the increased steric bulk of the cyclohexyl group. Not only can a terminal olefin act as an assisting group, but also internal olefins were found to promote the reaction as shown by the formation of products 3e and 3f in 59% and 65% yield, respectively. In addition to an ester group on $R^2$, various functional groups at $R^2$ such as alkyl 1g, hydroxyl 1h, silyl 1i, acetate 1j, sulfonamide 1k, and imide 1l were compatible with the reaction conditions, highlighting the broad substrate scope of this protocol (52–75% yields). As an example of late-stage oxidative functionalization, an estrone-derived substrate 1m efficiently participated in this reaction to afford a functionalized complex molecule 3m in a useful yield (78%). Unfortunately, attempts to synthesize the N-containing seven-membered heterocycle gave the corresponding product 3a in 75% isolated yield. With $R^2$ being a benzyl or a cyclohexyl group, the corresponding products 3b and 3c were obtained in 78% and 60% yields, respectively. Under the optimal aerobic reaction conditions, cyclohexyldiene bisallene 1d afforded 3d in only 30% yield. We attribute this diminished reactivity to the increased steric bulk of the cyclohexyl group. Not only can a terminal olefin act as an assisting group, but also internal olefins were found to promote the reaction as shown by the formation of products 3e and 3f in 59% and 65% yield, respectively. In addition to an ester group on $R^2$, various functional groups at $R^2$ such as alkyl 1g, hydroxyl 1h, silyl 1i, acetate 1j, sulfonamide 1k, and imide 1l were compatible with the reaction conditions, highlighting the broad substrate scope of this protocol (52–75% yields). As an example of late-stage oxidative functionalization, an estrone-derived substrate 1m efficiently participated in this reaction to afford a functionalized complex molecule 3m in a useful yield (78%). Unfortunately, attempts to synthesize the N-containing seven-membered heterocycle gave the corresponding product 3a in 75% isolated yield. With $R^2$ being a benzyl or a cyclohexyl group, the corresponding products 3b and 3c were obtained in 78% and 60% yields, respectively. Under the optimal aerobic reaction conditions, cyclohexyldiene bisallene 1d afforded 3d in only 30% yield. We attribute this diminished reactivity to the increased steric bulk of the cyclohexyl group. Not only can a terminal olefin act as an assisting group, but also internal olefins were found to promote the reaction as shown by the formation of products 3e and 3f in 59% and 65% yield, respectively. In addition to an ester group on $R^2$, various functional groups at $R^2$ such as alkyl 1g, hydroxyl 1h, silyl 1i, acetate 1j, sulfonamide 1k, and imide 1l were compatible with the reaction conditions, highlighting the broad substrate scope of this protocol (52–75% yields). As an example of late-stage oxidative functionalization, an estrone-derived substrate 1m efficiently participated in this reaction to afford a functionalized complex molecule 3m in a useful yield (78%).
using the nitrogen analogue of 3g (where oxygen in the ring had been replaced by NTs) were unsuccessful (See Supporting Information). Bis(neopentyl glycolato)diboron and bis(hexylene glycolato)diboron also worked as the borylating reagent and afforded the corresponding borylation products 3n and 3o in moderate yields (40% and 52%, respectively). The seven-membered borylation product from 1s was not observed due to the steric hindrance of substrate 1s.

To further confirm the effect of the pending olefin group in bisallene 1, we carried out control experiments with 1a' as the substrate, in which the vinyl group in 1a had been replaced by an ethyl group [Eq. (1)]. Attempted reaction of 1a' under standard conditions did not give any product. This result shows that the pending olefin is an indispensable element for this oxidative carbocyclization, which is in accordance with our previous work.\(^{[17]}\)

In analogy with our previous report,\(^{[15]}\) there was a rate enhancement with the hybrid catalyst compared to the use of Co(salophen) and quinone as separate molecules as shown in Figure 2. Co(salophen)-HQ hybrid (Cat. 1) resulted in fast borylative carbocyclization of 1g to 3g (Figure 2a), as well as the direct carbocyclization of 1s to 6 ending with β-H elimination (Figure 2b). These results indicate that the intramolecular electron transfer between the hydroquinone and the oxidized metal-macrocycle of this bifunctional catalyst leads to a more efficient palladium reoxidation process under aerobic conditions.\(^{[23]}\)

On the basis of these experimental findings, a possible mechanism for the oxidative coupling reaction is proposed in Scheme 6. Initially, the coordination of allene and olefin units to the Pd\(^{II}\) center leads to a chelate palladium complex Int-I. This special coordination of the close-by olefin to Pd\(^{II}\) is essential for triggering the allenic C(sp\(^3\))–H cleavage and generating a vinylpalladium intermediate Int-II.\(^{[17]}\) Next, the envisioned ligand exchange of olefin by the distal allene moiety takes place to give Int-III. Subsequent carbocyclization of Int-III by the second allene insertion gives a seven-membered (p-allyl)palladium intermediate Int-IV. Reaction of Int-IV with a trapping reagent such as B\(_2\)pin\(_2\), PhB(OH)\(_2\) or CO/MeOH provides the target product and a Pd\(^{0}\) species, respectively. Finally, with the assistance of the cobalt hybrid catalyst, aerobic oxidation of Pd\(^{0}\) regenerates Pd\(^{II}\) to close the catalytic cycle.

In addition to borylative oxidative carbocyclization, the use of phenylboronic acid as a transmetallating agent under similar reaction conditions also afforded the carbocyclization-arylation product 4 in 48% yield (Scheme 5a). Furthermore, an efficient cascade reaction of bisallene 1a via an oxidative carbocyclization-methoxycarbonylation is demonstrated here (Scheme 5b) and gave the seven-membered carbocycle 5 in 78% yield. In the absence of a trapping reagent, an intramolecular oxidative coupling ending with β-H elimination produced a highly conjugated seven-membered ring 6 in 79% yield (Scheme 5c).

Scheme 5. Synthesis of seven membered rings via arylative, carbonylative and beta-H eliminative carbocyclization. Reaction conditions: (a) bisallene 1a (1.0 equiv), PhB(OH)\(_2\) (1.3 equiv), Pd(OAc)\(_2\) (5 mol%), Cat. 1 (10 mol%), in 0.1 M acetone, O\(_2\) (1 atm) at 25 °C for 30 h; (b) bisallene 1a (1.0 equiv), Pd(OAc)\(_2\) (5 mol%), DMSO (20 mol%), BQ (1.5 equiv), MeOH (5.0 equiv), CO (1 atm) in 0.1 M DCE at 25 °C for 30 h; (c) bisallene 1s (1.0 equiv), Pd(OAc)\(_2\) (5 mol%), Cat. 1 (10 mol%) in 0.1 M THF, O\(_2\) (1 atm) at 25 °C for 30 h.

Figure 2. Reaction profiles with different ETMs: (a) Borylative carbocyclization of 1g to 3g; (b) Direct intramolecular carbocyclization of 1s to 6. For details, please see the supporting information.
In summary, we have developed an efficient and selective palladium-catalyzed carbocyclization of bisallenones under aerobic oxidative conditions. This reaction avoids stoichiometric amounts of non-environmentally friendly oxidants (Cu(I), Ag(I), peroxide etc.) for the activation of allicenic C–H bonds. The use of molecular oxygen as a green oxidant allows for synthesis of important seven-membered heterocycles in moderate to good yields. The key to success of this transformation is the application of a special hybrid electron transfer mediator [Co(salophen)-HQ]—a bifunctional catalyst consisting of a metal-macrocycle and quinone moieties. This hybrid catalyst significantly facilitates the reoxidation of Pd(0) to Pd(II) using molecular oxygen as the terminal oxidant. In view of the high reaction efficiency and selectivity, this protocol is expected to complement the current approach for oxidative functionalization in the synthesis of natural products and pharmacologically active substances.

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Conflict of interest

The authors declare no conflict of interest.

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