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Generation of Gold Carbenes in Water: Efficient Intermolecular Trapping of the α-Oxo Gold Carbenoids by Indoles and Anilines

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The efficient intermolecular reaction of gold carbene intermediates, generated via gold-catalyzed alkyne oxidation, with indoles and anilines has been realized in aqueous media. Importantly, it was revealed for the first time that water could dramatically suppress the undesired overoxidation, providing a general and practical solution to the problem of overoxidation in the gold-catalyzed intermolecular alkyne oxidation

10 with external nucleophiles. This strategy was successfully applied to the formal synthesis of the Pfizer's chiral endothelin antagonist UK-350,926.

Introduction

The generation of α-oxo gold carbenes via gold-catalyzed intermolecular alkyne oxidation represents a significant advance

- 15 in gold catalysis,¹ and various efficient synthetic methods have been developed based on this approach by $Zhang² Liu³$ and others.4-5 Despite these significant achievements, intermolecular alkyne oxidation with external nucleophiles remains a particularly challenging task. There are two major reasons for this:
- 20 (1) most external nucleophiles can also attack alkynes directly, catalyzed by gold species. (2) This type of α -oxo gold carbenes is highly reactive and often suffers the competing overoxidation of the carbene center by the very oxidant^{5,3g} and also many other side reactions. To date, successful intermolecular examples have
- 25 been quite scarce, $4a$ except for some reactions which have to rely on P,N- or P,S-bidentate ligands and slow addition of oxidants via syringe pump^{2b,c} or using the external nucleophiles as the reaction solvent to minimize the second oxidation.²¹

Recently, the use of an environmentally friendly reaction $_{30}$ medium, especially water, has attracted much attention.⁶ Although many types of transition metal-catalyzed reactions, including olefin metathesis, Mizoroki-Heck reaction, Suzuki reaction, and Sonogashira reaction in water have been achieved, the investigation of gold-catalyzed reactions conducted in

- 35 aqueous medium is still limited⁷ and generally focused on reactions catalyzed by the water-soluble gold complex.⁸ Competitive hydration of the substrates is probably the major reason for this paucity when gold-catalyzed transformations of alkynes are performed in water.⁹ Herein we present the first
- 40 example of generation of gold carbenes through intermolecular alkyne oxidation in aqueous media, and efficient intermolecular trapping of the α-oxo gold carbenes by indoles and anilines. Importantly, it was revealed for the first time that water could dramatically suppress the undesired overoxidation of gold 45 carbene and make this approach become a viable synthetic
- methodology, which is complementary to the traditional

generation of metal carbenoids in organic solvents from diazo compounds. The synthetic utility is also demonstrated by the formal synthesis of the Pfizer's chiral endothelin antagonist UK-50 350,926.

Results and discussion

During the last decade, a lot of attention has been paid to the development of transition-metal (M = Rh, Ru, Cu, Pd, Fe, In) catalyzed C-H functionalization of indoles by carbenoids derived 55 from diazo compounds, allowing an efficient access to indole derivatives.¹⁰⁻¹¹ However, the use of hazardous, not easily accessible and potentially explosive α-diazo ketone precursors here severely limits its further synthetic applications and the molecular flexibility. Moreover, it remains challenging to control 60 the regioselectivity and chemoselectivity of the C-H insertion reaction, N-H insertion reaction and the cyclopropanation reaction. Therefore, the development of alternative approach is still highly desirable. We envisioned that using the abovementioned oxidative approach α-oxo gold carbenoids of type **I** 65 would be readily accessed by using ynamide **2**, instead of the corresponding hazardous and potentially explosive α-diazoamide, and its subsequent C-H functionalization of indoles **1** would deliver **3** in a non-diazo approach (Scheme 1).

At the outset, we used *N*-methylindole **1a** and ynamide substrate **2a** as the reacting partners and 2-bromopyridine *N*-

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oxide as the oxidant (see SI for the screening of different oxidants and solvents), and some of the results are listed in Table 1. Initially, diketone compound **3aa** was detected as a major product with various typical gold catalysts with a range of electronic and

- 5 steric characteristics (Table 1, entries 1-8). To suppress the overoxidation of ynamide into the diketone, we then performed the reaction by slow addition of the oxidant into the reaction mixture using a syringe pump, which was utilized in previous intermolecular trapping of α -oxo gold carbenes.^{2b,c} However, in
- 10 this case, only enamide **3ab** was produced through a goldcatalyzed direct addition of indole **1a** to alkyne **2a** (Table 1, entries $9-10$.¹² To our delight, it was found that the reaction yields could be substantially improved when the reaction was run

Table 1 Optimization of reaction conditions*^a*

Reaction run in vials with DCE and/or deionized water; $[2a] = 0.1$ M. Measured by ¹H NMR using diethyl phthalate as the internal standard. ^c The oxidant was introduced into the reaction mixture over a 1 h period using a syringe pump. *^d*20 The mixture of oxidant and **1a** were introduced into the reaction mixture over a 1 h period using a syringe pump.

in 10:1 DCE-H₂O mixture solvent (Table 1, entries 11-13). Further studies revealed that the increase of the ratio of $H₂O/DCE$

offered a further improvement (Table 1, entries 14-19) and 25 excellent yield could be achieved using 10:1 H₂O-DCE as the solvent (Table 1, entries 17-19). Strikingly, the reaction could proceed equally well even in neat H_2O when 5 mol % IPrAuNTf₂ was employed as the catalyst and the desired indole derivative **3a** could be furnished in 88% yield (Table 1, entry 21). Notably, 30 without a gold catalyst, the reaction failed to give even a trace of **3a**, and PtCl₂ and AgNTf₂ were not effective in promoting this

reaction (8 and 25%, respectively). With the optimized reaction conditions in hand, the scope of the transformation was explored with a range of indoles and 35 ynamides. As shown in Table 2, the reaction proceeded smoothly with various substrates, and the yields ranged from 70% to 97%. Importantly, only C3-alkylated products were formed in all cases. We first examined various substituted indoles with ynamide substrate **2a** and were pleased to find that the reaction provided 40 the desired indole derivatives **3a**-**3j** in good to excellent yields (Table 2, entries 1-10). In the case of the indoles bearing strong electron-withdrawing groups, improved yields could be obtained by using slight excess of ynamide **2a** (Table 2, entries 6-7). Then, various ynamides **2** were investigated and the reaction also 45 proceeded well, affording the corresponding products **3k**-**3s** in good to excellent yields (Table 2, entries 11-19). In case of 3-en-1-ynamide **2d**, the reaction gave α, β-unsaturated imide **3m** as a single isomer in 88 % yield (Table 2, entry 13). Notably, when $R³$ is an alkyl group, the desired **3p** and **3q** could be formed in 93% 50 and 87% yield, respectively and no carbene insertion into a CH bond was observed (Table 2, entries 16-17). In addition, ynamides containing a Ms or a Bs group also worked well to give the desired **3r** and **3s** in 70% and 90% yields, respectively (Table 2, entries 18-19). Other *N*-substituted indoles were also suitable 55 substrates for this reaction to furnish the corresponding functionalized indoles **3t**-**3u** in good yields (Table 2, entries 20- 21). Even for unprotected indole substrate, this reaction still led to a respectable 70% yield and no *N*-alkylated product was formed (Table 2, entry 22).^{11m,n} It should be mentioned that in 60 some cases, slightly improved yields could be achieved by using 1.05 equiv of indole in order to prohibit the background enamide formation (Table 2, entry 9, entries 11-12 and entry 22). To further test the practicality of the current catalytic system, the reaction was carried out in a gram scale in the presence of 2.5 65 mol % gold catalyst and the desired product **3a** was afforded in 92% yield, highlighting the synthetic utility of this chemistry (Table 2, entry 1). Finally, X-ray diffraction studies of product **3b** further confirmed our structural assignments.¹³

This chemistry can also be extended to 2-substituted indole 70 substrate **1m** under the optimized reaction conditions, leading to the formation of the desired product **3w** in 76% yield (eqn (1)).

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^{*a*} Reactions run in vials; $[2] = 0.1$ M; isolated yields are reported. *b* 5.0 5 mmol scale, 2.5 mol % gold catalyst was used, 4 h. \degree Indole 1 (1.0 equiv), ynamide **2a** (1.5 equiv, added in two portions), 2 h. *^d* 1.05 equiv of indole **1** was used. e Using H₂O/DCE = 10/1 as the solvent.

The significance of this chemistry is additionally demonstrated by its application to the formal synthesis of the Pfizer's chiral ¹⁰ endothelin antagonist UK-350,926 $(3z)$,¹⁴ as depicted in Scheme

^{2. 2-}Ynamide substrate **2k** was prepared from 2-methoxy-4 methylbenzenesulfonyl chloride^{14e} in a two-step process involving a copper-catalyzed amidation of alkynyl bromide compound. Then, the reaction of the ynamide **2k**, indole **1n**, and 15 oxidant in the presence of 5 mol % IPrAuNTf₂ in water at 80 °C for 4 h gave the compound **3x** in 65% yield. Indole **3x** was subjected to hydrogenation to afford **3y**, which has been transformed to **3z** in one step using a dynamic resolution process.14a This total synthesis of racemic UK-350,926 20 constitutes a short, four-step sequence from the above sulfonyl chloride in a 42% overall yield.

Scheme 2 Formal synthesis of Pfizer's chiral endothelin antagonist UK-350,926. Reagents and conditions: (a) $BnNH₂$ (1.1 equiv), Et₃N, rt, 4 h; 5-25 (bromoethynyl)benzo[d][1,3]dioxole (1.0 equiv), CuSO4•5H2O (0.1 equiv), 1,10-phenanthroline (0.1 equiv), K_2CO_3 (3.0 equiv), toluene, 80 °C, 10 h; (b) IPrAuNTf₂ (5 mol %), benzyl 1-methyl-1H-indole-6carboxylate **1n** (1.0 equiv), **2k** (1.5 equiv), 2-bromopyridine *N*-oxide (2.0 equiv), H_2O , 80 °C, 4 h.

Two plausible mechanisms to rationalize the formation of indole **3a** are proposed (Scheme 3). Path a is generally believed for the gold-catalyzed oxidative processes involving the α–oxo gold carbene intermediate **B**, which could be further trapped by indole **1a** to deliver the final **3a**. Another pathway, path b, may 35 involve the generation of intermediate **C** by a direct intermolecular $S_N 2$ ' reaction. The intermediate C then undergoes protondeauration to furnish product **3a** along with the regeneration of the gold catalyst.

⁴⁰**Scheme 3** Possible reaction mechanisms.

To probe the feasibility of the two paths (a and b), we first prepared the diazo compound **2l**. As shown in eqn (2), it was

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found that the reaction of **2l** with indole **1a** indeed produced the desired **3a** in 48% yield. In particular, 62% yield could be achieved in the presence of 0.5 equiv of oxidant, which was supposed to facilitate the generation of gold carbene **B** through a

- s stabilization effect as previously reported by Liu.^{3b} Moreover, a similar water effect was observed as in its absence the reaction efficiency decreased dramatically and the ketoimide **3aa** was formed in a much increased yield. In addition, the reaction of **1a** with **2m** only resulted in the formation of α, β-unsaturated imide
- 10 product (eqn (3)).⁴ⁱ These results suggest that the reaction presumably goes through the path a and the α –oxo gold carbene intermediate **B** is most likely generated. Notably, the reaction pathway involving trapping of α-oxo gold carbenes by water, followed by nucleophilic attack by indole is much less possible
- 15 because the reaction of α-hydroxyimide **3aa'** with **1a** failed to produce **3a** and also no **3aa'** formation was observed in the reaction system (eqn (4)). More importantly, the inactivity of insertion to the O-H bond of water guaranteed further transformations of the derived gold carbene in aqueous solution.¹
- 20 In addition, we also prepared **3aa** and subjected it to the optimal reaction conditions, and no **3a** was obtained, which rules out diketone **3aa** as an intermediate on the way to product **3a** (eqn (4)).

- 25 While the role of H_2O in this reaction remains to be clearly established, we suspected that the water-soluble organic oxide (2 bromopyridine *N*-oxide) in this system exists mainly in water phase whereas the catalyst and internal alkynes and indole nucleophiles form insoluble organic mesophases, in which the 30 reactions take place. As a result, this mesophase system maintains
- proper concentration of the external oxidants to avoid a double oxidation of alkynes and favors the nucleophilic addition of indole to gold carbenes, which can be equivalent to reaction conditions using slow addition of an oxidant.¹⁶ Indeed, it was
- 35 found that when the reaction was run in water by using waterinsoluble N -oxides (even at 80 \degree C), significant formation of diketone **3aa** was observed (eqn (5)). In addition, kinetic experiments showed that the reaction proceeded much slower in water than in DCE (5 min *vs* 1 h, see SI for the detailed kinetic 40 study of the reaction), further supporting the above speculation
- that the concentration of the water-soluble *N*-oxides in the reaction phase was kept at a low level.

We next considered the possibility of extending the reaction to 45 other electron-rich aromatics. To our delight, the reaction of the ynamide **2a** with *N*,*N*-dimethylaniline **4a** in the presence of 5 mol % IPrAuNTf₂ and 2 equiv of 2-bromopyridine N-oxide could proceed smoothly in water, delivering the desired **5a** in 78% NMR yield, as depicted in eqn (6). In contrast, when the reaction 50 was carried out in DCE, **5a** was obtained only in 10% yield and significant formation of diketone **3aa** could be observed, thus exhibiting again that the role of water here serves to dramatically promote this oxidative gold catalysis. The use of 3 equiv of **4a** minimized the formation of hydration byproduct **3ac**, 18,5a leading 55 to a further improved yield.

 b Yield of isolated 5a was 89%

There are several well-documented examples of intramolecular trapping of α-oxo gold carbenoids by aromatic substitution reaction, $17,24,56$ however, to our best knowledge, examples of the 60 intermolecular version have not been reported. We then examined the scope of this intermolecular aromatic substitution of *N*, *N*disubstituted anilines **4** with α-oxo gold carbenoids generated via oxidation of ynamides **2**. As summarized in Table 3, all of the reactions took place smoothly and afforded the corresponding **5** 65 in largely good to excellent yields. In entries 1-5, this reaction works satisfactorily with various types of *N*,*N*-dimethyl aniline derivatives **4b**-**4f**; their electrophilic aromatic substitution products **5b**-**5f** were obtained in 63-90% yields. The reactions were also extensible to different ynamides **2**, which delivered the 70 desired **5g**-**5l** in 78-96% yields (Table 3, entries 6-11). Notably, the reaction with 3-en-1-ynamide **2d** afforded α, β-unsaturated imide **5i** as a single isomer in 86 % yield (Table 3, entry 8 vs Table 2, entry 13). Entries 12 and 13 show our reactions with Ms and Bs protected ynamides **2i**-**2j**, resulting in excellent yields of 75 the desired products **5m**-**5n** (97% and 90% isolated yields). This reaction also worked well for other types of *N*,*N*-disubstituted anilines **4g**-**4h**, giving the desired products **5o**-**5p** in 94% and 70% yields, respectively (Table 3, entries 14-15). The molecular

catalysis on the secondary aniline **6a**, no aromatic substitution product was observed. Instead, the intermolecular N–H insertion product **7a** was isolated albeit in low yield. As shown in eqn (7), 10 water indeed significantly inhibited the formation of diketone byproduct **3aa**, however, the formation of background enamide **3ad** became dominant, which could be rationalized by a goldcatalyzed direct addition of *N*-methylaniline **6a** to ynamide **2a**. Gratifyingly, the yield could be substantially improved just by a

 15 simple tuning of the ratio of $H₂O/DCE$.

Dh

Ts

 DCF

 $H₂O$ $DCE/H₂O = 1/1$

 $2a$

Conditions

 b Yield of isolated **7a** was 81%.</sup>

 $6a(2 \text{equiv})$

5 mol % IPrAuNTf solvent, 80 °C, 2 h

(2 equiv)

 $7a$

75%

 $30%$

83%^b

^a Measured by ¹H NMR using diethyl phthalate as the internal standard.

structure of **5f** was confirmed by X-ray diffraction.¹³

Table 3 Reaction scope with different tertiary anilines **4** and ynamides **2***^a*

Table 4 Reaction scope with different secondary or primary anilines **6** and ynamides **2***^a*

Conclusions

In summary, the challenging intermolecular trapping of α-oxo gold carbenes by external nucleophiles such as indoles and 45 anilines is achieved. Different from the well known Cu, Rh, and Ru carbenoids, which were formed from metal-catalyzed

Although intramolecular trapping of this type of gold carbenes via facile $O-H^{2i,j}$ and $N-H^{19,2g}$ insertions has been well

Ρh

'N'

 $\frac{1}{P}h$

Ts

3ac

 $1%$

48%

 $< 2%$

Ph

 326

ฅ่ ∧'n
Ph

7a

 $Yield^a$

3aa

18%

 $1%$

 $1%$

 (7)

decomposition of diazo compounds and normally in organic solvents, this carbenoid species is generated through goldcatalyzed intermolecular alkyne oxidation and its relevant reactions can be performed in aqueous media, thus making this

- 5 non-diazo protocol very practical and environmentally friendly. Significantly, it was revealed that water could dramatically inhibit the formation of diketone byproducts, which might serve as a general solution to the overoxidation in oxidative gold catalysis. This new method enables a succinct and practical
- 10 formal synthesis of the Pfizer's chiral endothelin antagonist UK-350,926. Studies to elucidate the detailed mechanism and further application of this water-assisted protocol for the oxidative gold catalysis will be pursued further in our laboratory.

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