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Dual gold/photoredox-catalyzed C(sp)-H arylation of terminal alkynes with diazonium salts†

Adrian Tlahuext-Aca,‡ab Matthew N. Hopkinson,‡a Basudev Sahooab and Frank Glorius*a

The arylation of alkyl and aromatic terminal alkynes by a dual gold/photoredox catalytic system is described. Using aryldiazonium salts as readily available aryl sources, a range of diversely-functionalized arylalkynes could be synthesized under mild, base-free reaction conditions using visible light from simple household sources or even sunlight. This process, which exhibits a broad scope and functional group tolerance, expands the range of transformations amenable to dual gold/photoredox catalysis to those involving C-H bond functionalization and demonstrates the potential of this concept to access Au^I/Au^{III} redox chemistry under mild, redox-neutral conditions.

Introduction

Homogeneous gold catalysis has received significant attention in the last two decades. Due to their carbophilic π -acidity, Au^I and Au^{III} complexes catalyze the addition of a variety of oxygen-, nitrogen-, carbon-, and sulfur-based nucleophiles to unsaturated molecules, such as alkynes, allenes and alkenes, giving rapid and efficient access to complex molecular architectures.1 A common feature regarding the nature of the catalytically active Au species is that it does not easily undergo changes in oxidation state during the course of these reactions. However, there has been significant recent interest in developing Au^I/Au^{III} catalytic processes with the aim of expanding the repertoire of gold-mediated processes, mimicking conventional Mⁿ/Mⁿ⁺² redox cycles of the type commonly invoked in catalysis by other late transition metals.2 Nevertheless, in contrast to its isoelectronic counterpart Pd⁰, the oxidation of Au^I generally requires the use of strong oxidative conditions due to the high redox potential of the Au^{I}/Au^{III} redox couple ($E_0 = 1.41 \text{ V}$).³ A common strategy to access catalytically active Au^{III} species in situ from Au^{I} complexes has been to use strong external oxidants, such as Selectfluor®, tBuOOH, or hypervalent iodine reagents. Applying this concept, a range of oxidative homo- and heterocoupling reactions have been developed over the last few years where the oxidative coupling event follows a conventional gold-mediated

Despite the success of these approaches, the use of strong external oxidants, typically in super-stoichiometric amounts, inevitably reduces their attractiveness with regards to functional group tolerance and cost, while most examples are also conducted at high reaction temperatures. With the goal of developing milder gold-catalyzed coupling reactions that do not require external oxidants, our group recently described the use of a dual gold/photoredox catalysis strategy to access AuI/AuIII redox cycles.5,6 Using this concept, we have developed

Scheme 1 Gold-catalyzed C-C and C-X bond-forming coupling reactions proposed to proceed via Au¹/Au¹¹¹ redox cycles.

nucleophilic addition reaction to a C-C multiple bond, circumventing protodeauration and giving efficient access to difunctionalized products (Scheme 1a). This strategy has also been successfully applied in a selection of coupling reactions of unfunctionalized arenes and alkynes, exploiting the wellestablished ability of gold to activate C(sp²)-H or C(sp)-H bonds

a) Oxidative Heterofunctionalization b) Oxidative C(sp2)-H or C(sp)-H Functionalization of Carbon-Carbon Multiple Bonds [Au] Catalyst Strong Oxidant Strong Oxidant Nucleophile, R1-M/H Nu R1-M/H c) Dual Au/Photoredox-Catalyzed Redox Neutral Heterofunctionalization of Alkenes -XH Photocatalyst Visible Light ArN₂BF₄ Ar₂IBF₄ d) This work: Dual Au/Photoredox C(sp)-H Arylation Process L-Au-C $[Ru(bpy)_3](PF_6)_2$ $\dot{N}_2 \bar{B} F_4$ R = Arvl or Alkvl + Readily Available Arylating Agents

^aOrganisch-Chemisches Institut, Westfälische Wilhelms-Universität Münster, Corrensstraße 40, 48149 Münster, Germany. E-mail: glorius@uni-muenster.de

^bNRW Graduate School of Chemistry, Westfälische Wilhelms-Universität Münster, Wilhelm-Klemm-Strasse 10, 48149 Münster, Germany

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[‡] These authors contributed equally.

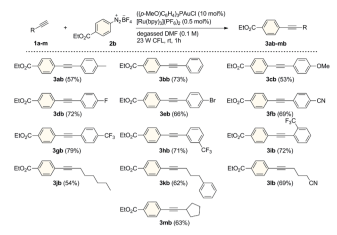
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intramolecular aminoarylation and both intra-5a and intermolecular5b oxyarvlation reactions of alkenes to afford arylated heterocycles and β-aryl ethers employing aryldiazonium or diaryliodonium salts as general arylating agents (Scheme 1c). This dual catalytic system, which has since been further exploited by the Toste group in an impressive arylative ring expansion^{7a} and C-P bond-forming cross-coupling,^{7b} operates under mild and overall redox neutral conditions and uses abundant visible light as an energy source. Building on these studies, we sought to expand the scope of dual gold/photoredox catalysis onto new classes of transformations, exploiting different aspects of the rich chemistry of gold. In this regard, we turned our attention to the development of novel visible lightmediated cross-coupling reactions involving gold-catalyzed C-H bond activation and identified the well-established ability of gold(I) to activate the C(sp)-H bond of terminal alkynes as a promising avenue of investigation. Arylation of the resulting alkynylgold complexes would deliver cross-coupled arylalkyne products in a dual gold and photoredox-catalyzed analogue of the widely-employed Sonogashira-Hagihara reaction.^{8,9} As demonstrated below, this process benefits from mild conditions (room temperature, household light bulb) and broad functional group tolerance while making use of readily-available aryldiazonium salts as the arene coupling partner. Furthermore, by exploiting a non-conventional stepwise oxidation process of Au^I complexes into active Au^{III} species by means of organic radicals generated under photoredox catalysis, we have developed a very fast arylating system that proceeds under

Results and discussion

base-free conditions.10

In a preliminary experiment, p-tolylacetylene (1a) was reacted in the presence of benzenediazonium tetrafluoroborate (2a) under visible light irradiation from a simple household desk lamp fitted with a 23 W fluorescent light bulb (CFL). Upon treatment with 10 and 2.5 mol% of Ph₃PAuCl and [Ru(bpy)₃](PF₆)₂, respectively, in degassed methanol for 2 h at room temperature, we were pleased to observe that the cross-coupled product 3aa was formed in 46% yield (entry 1). Following a short screen, (see the ESI† for details of additional experiments), DMF was identified as the optimum solvent for this transformation (entry 2). A range of different Au^I catalysts bearing various ligands were then evaluated (entries 3-7). Interestingly, the gold(1) chloride complex possessing the comparatively electron-rich phosphine $((p-MeO)C_6H_4)_3P$ led to the highest yield of 3aa (83%), while the trialkylphosphine Cy₃P was also tolerated (entry 4). This trend might be related to the stabilization of the Au^{III} center by these strongly donating ligands during the Au^I/Au^{III} redox cycle. The combination of the Au^I complex ((p-MeO)C₆H₄)₃PAuCl and the photoredox catalyst [Ru(bpy)3](PF6)2 turned out to be both a highly selective and active C(sp)-H arylating system, as 3aa could be isolated in 80% yield using very low loadings of [Ru(bpy)₃](PF₆)₂ (0.5 mol%) after a reaction time of only 1 hour and without the formation of the divne homocoupling product of 1a (entry 8).11 Furthermore, this system does not require the addition of an external base and proceeds under acidic



Scheme 2 Dual Au/photoredox-catalyzed arylation of different alkynes. General conditions: alkyne (1a-m, 0.30 mmol), 2b (1.2 mmol), [Ru(bpy)₃](PF₆)₂ (0.5 mol%), ($(p-MeO)C_6H_4$)₃PAuCl (10 mol%) and degassed DMF (3.0 mL). Isolated yields.

conditions. Interestingly, the reaction also operated efficiently using visible light from a variety of different sources. Under both blue LED (5 W, $\lambda_{\rm max}=465$ nm) and even sunlight irradiation, the formation of the cross coupled product **3aa** was

Table 1 Reaction optimization^a

| Entry | Au(ı) complex | Photocatalyst | Yield ^b (%) |
|-----------------|--|--|------------------------|
| 1 ^c | Ph₃PAuCl | [Ru(bpy) ₃](PF ₆) ₂ | 46 |
| 2 | Ph ₃ PAuCl | $[Ru(bpy)_3](PF_6)_2$ | 78 |
| 3 | ((p-MeO)C ₆ H ₄) ₃ PAuCl | $[Ru(bpy)_3](PF_6)_2$ | 83 |
| 4 | Cy ₃ PAuCl | $[Ru(bpy)_3](PF_6)_2$ | 72 |
| 5 | XPhosAuCl | $[Ru(bpy)_3](PF_6)_2$ | _ |
| 6 | (PhO) ₃ PAuCl | $[Ru(bpy)_3](PF_6)_2$ | _ |
| 7 | IPrAuCl | $[Ru(bpy)_3](PF_6)_2$ | _ |
| 8^d | ((p-MeO)C ₆ H ₄) ₃ PAuCl | $[Ru(bpy)_3](PF_6)_2$ | 83 (80) |
| 9^e | ((p-MeO)C ₆ H ₄) ₃ PAuCl | $[Ru(bpy)_3](PF_6)_2$ | 80 |
| 10 ^f | ((p-MeO)C ₆ H ₄) ₃ PAuCl | $[Ru(bpy)_3](PF_6)_2$ | 71 |
| 11^g | $((p\text{-MeO})C_6H_4)_3$ PAuCl | $[Ru(bpy)_3](PF_6)_2$ | 60 |
| 12^h | $((p\text{-MeO})C_6H_4)_3$ PAuCl | $[Ru(bpy)_3](PF_6)_2$ | 82 |
| 13^{i} | ((p-MeO)C ₆ H ₄) ₃ PAuCl | [Ir(ppy) ₂ (dtbbpy)]PF ₆ | Trace |
| 14 | ((p-MeO)C ₆ H ₄) ₃ PAuCl | Eosin Y | _ |
| 15 | $((p\text{-MeO})C_6H_4)_3$ PAuCl | Fluorescein | _ |

 a General conditions: 1a (0.10 mmol), 2a (0.40 mmol), photocatalyst (2.5 mol%), gold complex (10 mol%) and degassed DMF (1.0 mL). b NMR yields using CH₂Br₂ as internal standard. Isolated yield in parentheses. c Degassed methanol (1.0 mL). d [Ru(bpy)₃](PF₆)₂ (0.5 mol%), 1 h. e [Ru(bpy)₃](PF₆)₂ (0.5 mol%), 5 W blue LEDs, 1 h. f [Ru(bpy)₃](PF₆)₂ (0.5 mol%), sunlight, 8 h. g [Ru(bpy)₃](PF₆)₂ (0.5 mol%), air, 1 h. h [Ru(bpy)₃](PF₆)₂ (2.5 mol%), air, 1 h. h With Ph₂IBF₄ (0.40 mmol), [Ir(ppy)₂(dtbbpy)]PF₆ (5 mol%). See the ESI for further details. ppy = 2-phenylpyridine, dtbbpy = 4,4'-di-tert-butyl-2,2'-bipyridine, XPhos = 2-dicyclohexylphosphino-2',4',6'-triisopropylbiphenyl, IPr = 1,3-bis(2,6-diisopropylphenyl)-imidazol-2-ylidene.

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observed in good yields (entries 9-10). In addition, the dual gold/photoredox C(sp)-H arylation process showed a remarkable robustness towards oxygen, with 3aa being obtained in 60% yield under the same conditions (10 mol% ((p-MeO)C₆- H_4)₃PAuCl and 0.5 mol% [Ru(bpy)₃](PF₆)₂) under an atmosphere of air (entry 11). Upon increasing the loading of [Ru(bpy)₃](PF₆)₂ to 2.5 mol%, the yield of 3aa recovered to 82% yield, indicating that careful exclusion of oxygen is not required to obtain high yields when a slightly increased loading of the photocatalyst is employed (entry 12). In contrast to our previous studies on alkene oxyarylation, attempts to switch the aryl coupling partner to diaryliodonium salts or use less expensive organic dyes were not successful for this transformation and the combination of aryldiazonium salts and [Ru(bpy)3](PF6)2 was employed for further experiments (entries 13-15).

We next evaluated the reaction scope studying a variety of alkyl and aromatic terminal alkynes using 4-(ethoxycarbonyl) benzenediazonium tetrafluoroborate (2b) as the arylating reagent (Scheme 2). With arylalkyne substrates, the reaction was found to tolerate electron-donating and withdrawing substituents on the aryl ring and the corresponding coupled products were isolated in good yields (53-79%) after just 1 h of visible light irradiation. Moreover, as exemplified for the trifluoromethyl-substituted compounds 3gb, 3hb and 3ib, the reaction proceeded with substituents at each of the para-metaand ortho-positions with seemingly comparable efficiency. Similar good results were obtained for a range of alkylsubstituted alkynes. Derivatives 3jb-mb were prepared in 54-69% yields with both primary and secondary alkyl groups being tolerated. In all cases, no diyne compounds resulting from homocoupling of the aryl- or alkylalkynes were observed during the reactions.

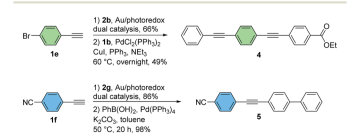
Our attention turned next to an investigation of the scope of the reaction towards a range of substituted aryldiazonium salts

((p-MeO)C₆H₄)₃PAuCl (10 mol%) [Ru(bpy)₃](PF₆)₂ (0.5 mol%) sed DMF (0.1 M) 3fd (73%) 3fa (68%) 3fc (61%) (67%)^[a] 3ff (56%) 3fg (86%) (67%)[a] MeO₂C 3fh (64%) 3fi (74%) 3fj (40%) (65%)[a (49%)[c]

Scheme 3 Dual Au/photoredox-catalyzed arylation of 1f and 1c with substituted benzenediazonium salts. General conditions: alkyne (0.30 mmol), 2a, 2c-l (1.2 mmol), $[Ru(bpy)_3](PF_6)_2$ (0.5 mol%), $((p-MeO)C_6-$ H₄)₃PAuCl (10 mol%) and degassed DMF (3.0 mL). [a] Under air using 2.5 mol% of $[Ru(bpy)_3](PF_6)_2$ in 1 h. [b] $[Ru(bpy)_3](PF_6)_2$ (2.5 mol%), 16 h of reaction. [c] $[Ru(bpy)_3](PF_6)_2$ (2.5 mol%), 5 h under 5 W blue LEDs irradiation. Isolated yields

(2a, 2c-l) using 4-ethynylbenzonitrile 1f as a standard alkyne coupling partner (Scheme 3). We were pleased to find that aryl sources bearing a variety of electronically diverse functional groups reacted successfully, delivering the corresponding crosscoupled products 3fa, 3fc-fj in good yields up to 86%. At this point, the efficiency of the reaction when performed using electron rich groups at both the aromatic alkyne and the diazonium salt was tested by reacting 4-ethynylanisole (1c) with 2,3-dihydrobenzo[b][1,4]dioxine-6-diazonium tetrafluoroborate (2k) under the optimized reaction conditions (Scheme 3). We observed only poor conversions of 1c unless higher loadings of [Ru(bpy)₃](PF₆)₂ (2.5 mol%) and longer irradiation times (16 h) were used. Under these conditions, the coupled product 3ck was isolated in 27% yield. Interestingly, the catalytic activity of this system was increased using blue LEDs (5 W, $\lambda_{max} = 465$ nm) as the light source. After 5 h of irradiation, 3ck was formed in an improved 49% isolated yield. This new set of reaction conditions was also successfully applied for the synthesis of the coupled product 3cl (50% isolated yield). The efficiency of the reaction was evaluated for some substrates under an atmosphere of air using a slightly higher loading of the photocatalyst $[Ru(bpy)_3](PF_6)_2$ (2.5 mol%, conditions from Table 1, entry 12). Interestingly, under these conditions, the arylated products 3fd, 3fg and 3fi were still obtained in good yields (65-67%), demonstrating that efficient catalytic activity can be maintained without rigorous extrusion of oxygen. Sterically-demanding aryldiazonium salts, however, are seemingly less well tolerated under these conditions with the ortho-substituted 2-(methoxyearbonyl)benzenediazonium tetrafluoroborate reacting sluggishly to afford the coupled product 3fj in a moderate yield of 40%.

In addition to its base-free and mild nature, an important feature of the dual gold/photoredox catalytic system is its compatibility with halogenated substrates. In contrast to palladium(0) or many other low-valent late transition metals, gold(I) does not generally undergo conventional oxidative addition to aryl halides under homogeneous conditions. 3,9df,g As such, the brominated diarylalkynes 3eb and 3fg, obtained from a bromoarylalkyne and a bromo-substituted aryldiazonium salt respectively, could be isolated in good yields using this method without competitive cleavage of the C-Br bond. As demonstrated by representative Sonogashira-Hagihara and Suzuki-Miyaura coupling reactions in Scheme 4, these substrates can be readily functionalized further using palladium catalysis.12



Scheme 4 Further manipulation of cross-coupled products (see the ESI† for experimental details).

Scheme 5 Mechanistic proposal

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Overall, this two-step protocol is complementary to stepwise palladium-catalyzed coupling sequences, making use of readilyavailable aryldiazonium salts rather than dihalogenated aryl iodide precursors (Scheme 4).

Alkynes

In accordance with previous studies on dual Au/photoredox catalysis,5,7 we envisage a mechanism involving a photoredoxinduced homogeneous Au^I/Au^{III} redox cycle (Scheme 5).¹³ Upon irradiation with visible light, photoexcitation of the [Ru(bpy)₃]²⁺ photocatalyst takes place to generate the excited form *[Ru(bpy)₃]²⁺ which then undergoes single electron transfer (SET) with one equivalent of the aryldiazonium salt to deliver an aryl radical and the oxidized species [Ru(bpy)₃]³⁺. The photogenerated aryl radical then adds to the Au^I catalyst to give the AuII species A.15 Interestingly, quantum yield measurements at 450 nm by chemical actinometry gave a value of 3.6, reflecting the very important contribution of radical chain processes in this dual catalytic system.16 Therefore, we propose that a second SET process between the open-shell species A with another equivalent of the aryldiazonium salt mainly contributes to the formation of the electrophilic cationic Au^{III} intermediate B. Nevertheless, regeneration of the oxidized [Ru(bpy)₃]³⁺ to the [Ru(bpy)₃]²⁺ photoredox catalyst is still feasible by reaction with **A.** The cationic intermediate **B** is expected to be an excellent π -Lewis acid and coordinates the alkyne substrate, activating it towards the formation of the $\sigma\text{-bonded}$ alkynyl-Au^III complex C upon deprotonation. Intermediate C finally undergoes reductive elimination,¹⁷ regenerating the Au^I catalyst and delivering the cross-coupled product.

Conclusions

In conclusion, we have developed an efficient dual gold/photoredox catalysis methodology for the arylation of terminal alkynes using readily-available aryldiazonium salts. This overall redox neutral cross-coupling process shows broad functional group tolerance, operates at room temperature and is mediated by abundant visible light from a household light bulb or even

sunlight. In addition, the base-free nature of the reaction and tolerance of halogenated substrates may be beneficial in the design of cross-coupling sequences. This reaction represents the first application of dual gold/photoredox catalysis for the activation of C–H bonds and further demonstrates the potential of this methodology to access Au^I/Au^{III} redox cycles under mild conditions without the need for external oxidants.

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- 15 Seminal study on the trapping of carbon-based radicals by gold complexes: (a) C. Aprile, M. Boronat, B. Ferrer, A. Corma and H. García, J. Am. Chem. Soc., 2006, 128, 8388; recent report on the oxidation of gold(1) by the CF₃ radical: (b) M. S. Winston, W. J. Wolf and F. D. Toste, J. Am. Chem. Soc., 2014, 136, 7777.
- 16 For interesting discussions on the prevalence of radical chain processes in photoredox catalysis see: (*a*) M. Majek, F. Filace and A. J. von Wangelin, *Beilstein J. Org. Chem.*, 2014, **10**, 981; (*b*) M. A. Cismesia and T. P. Yoon, *Chem. Sci.*, 2015, **6**, 5426. For quantum yield measurements see the ESI.†
- 17 Report on reductive elimination from these complexes: Y. Fuchita, Y. Utsunomiya and M. Yasutake, *J. Chem. Soc.*, *Dalton Trans.*, 2001, 2330.