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Gold-catalysed tuning of reactivity in allenes: 9-endo hydroarylation versus formal 5-exo hydroalkylation†

Cite this: *Chem. Commun.,* 2013, **49**, 1282

Received 30th October 2012, Accepted 19th December 2012

DOI: 10.1039/c2cc37872h

www.rsc.org/chemcomm

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The divergent gold-catalysed reactivity (C_{sp^2} –H *versus* C_{sp^3} –H) of aryloxy-tethered allenes has been uncovered.

The preparation of medium-sized ring polycycles through selective C–H bond activation is a big challenge. In this regard, the gold-catalysed intramolecular hydroarylation of allenes may be a possible solution to produce eight- or nine-membered carbocycles, although this achievement has not yet been accomplished. We present here an unprecedented Au-catalyzed 9-endo carbocyclization of aryl allenes as a powerful synthetic tool to obtain novel nine-membered annulated β -lactam derivatives. In addition, it is shown that the outcome of the reaction (9-endo hydroarylation versus formal 5-exo hydroalkylation) can be modulated by the allene tether.

To explore the reactivity of (aryl)allene-tethered 2-azetidinones 1 towards hydroarylation, we selected acetate 1a as a model substrate. As a first try, we were happy to notice that although the reaction of derivative 1a afforded dienol ester 2a as a major component (34% isolated yield), the intramolecular hydroarylation adduct 3a was also isolated as a minor component (13%) (Scheme 1). Thus, in order to prevent the [3,3]-sigmatropic rearrangement involving the acetate group,³ the hydroxy functionality was protected in the form of methyl ether. Fortunately, treatment of allene 1b with [(Ph₃P)AuNTf₂] in 1,2-dichloroethane at room temperature gave full conversion; benzo[b]oxonine 3b being isolated in 51% yield in a totally selective fashion (Scheme 1).⁴ Clearly, applying microwave irradiation and using deactivated silica gel during purification resulted in an increased 72% yield for adduct 3b. Similar figures were observed

Scheme 1 Controlled intramolecular gold-catalysed hydroarylation of allenyl-tethered arenes **1a–h**. (i) 5 mol% [(Ph₃P)AuNTf₂], DCE, μ wave, 110 °C (RT for **1a**). Ac = COMe. Tf = triflyl. DCE = 1,2-dichloroethane.

for tricyclic products 3c-h without harming the sensitive β -lactam ring (Scheme 1). Remarkably, this rare 9-endo carbocyclization reaction was the only operative cyclization mode.

We also decided to undertake a study of the potential use of more diverse substrates in this novel allene hydroarylation mode. Thus, (aryloxy)allenyl-tethered sugars **4a–e** were studied by using the optimum reaction conditions obtained for (aryloxy)allenyl-tethered 2-azetidinones **1b–h**. Remarkably, we found a divergent reactivity compared with the transformation found with allenes **1**; instead of the expected hydroarylation adducts, tricycles **5** arising from a rare 5-exo hydroalkylation were obtained (Scheme 2). Notably, the direct and selective functionalization of an otherwise inactive C_{sp3}-H bond has been achieved. ⁵ Besides, regioselectivity can be completely

(-)-4e R^1 = Br, R^2 = BnOCH₂

Scheme 2 Controlled intramolecular formal 5-exo hydroalkylation reaction of allenyl-tethered arenes **4a-e** under gold-catalysed conditions.

(+)-5e (41%)

⁽⁺⁾⁻¹a R1 = isobutyl, R2 = Me, R3 = Ac (+)-3a (13%) (+)-3b (72%) (+)-1b R^1 = isobutyl, R^2 = Me, R^3 = Me (+)-1c R^1 = isobutyl, R^2 = Ph, R^3 = Me (+)-3c (60%) (+)-1d $R^1 = Bn, R^2 = Me, R^3 = Me$ (+)-3d (58%) (+)-1e R¹ = Bn, R² = Ph, R³ = Me (+)-3e (69%) (+)-1f R^1 = isopropyl, R^2 = Me, R^3 = Me (+)-3f (65%) (+)-1g R^1 = isopropyl, R^2 = Ph, R^3 = Me (+)-3q (59%) (+)-1h R^1 = 2-methylallyl, R^2 = Ph, R^3 = Me (+)-3h (63%)

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[†] Electronic supplementary information (ESI) available: Experimental procedures, characterization data for new compounds, and copies of NMR spectra. See DOI: 10.1039/c2cc37872h

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Scheme 3 Mechanistic explanation for the gold-catalysed hydroarylation of allenyl-tethered oxyarenes 1.

reversed by using the sugar derivative, thus favouring the cyclization of the aryloxy ether group toward the central allene carbon over the cyclization towards the terminal allene carbon. The reaction of allenes 4 did take place with complete stereoselectivity, representing a selective method to afford fused cyclopentenes 5 bearing a quaternary stereocenter.6 Complete conversion was observed by TLC and ¹H NMR analysis of the crude reaction mixtures of allenols 4, and no side-products from isomerisation or polymerisation reactions were detected. Unfortunately, some decomposition was observed on sensitive tricycles 5 during purification by flash chromatography, which may be responsible for the moderate isolated yields.

A possible pathway for the gold-catalysed achievement of tricycles 3 from allenyl-tethered arenes 1 may initially involve the formation of a complex 1-Au(L) through coordination of the

gold salt to the distal allenic double bond. Next, chemo- and regioselective 9-endo carboauration forms species 6. Attack from the activated 2-position of the arene occurs as a result of the stability of the intermediate oxonium cation type 6. Loss of proton generates neutral species 7, which followed by protonolysis of the carbon-gold bond afforded fused ninemembered cycles 3 with concurrent regeneration of the gold catalyst (Scheme 3).

Density functional theory (DFT) calculations have been carried out to gain more insight into the reaction mechanism of the abovediscussed gold-catalysed 9-endo hydroarylation reaction. The corresponding computed reaction profile of the model allenyl-β-lactam reactant 1M with [(Me₃P)AuNTf₂] as catalyst is illustrated in Fig. 1, which shows the corresponding free energies in CH₂Cl₂ solution (PCM-M06/def2-SVP// PCM-B3LYP/def2-SVP level). Our calculations suggest that the reaction starts with the exergonic coordination of the AuPMe₃ catalyst to the distal double bond of the allenic moiety of 1M $(\Delta G_{298} = -9.4 \text{ kcal mol}^{-1})$. Then, the 9-endo carbocyclization reaction to produce the nine-membered ring tricyclic intermediate 2M occurs through the transition state TS1. This saddle point is associated with the nucleophilic addition of the activated ortho-carbon atom to the electrophilic gold complex. Although the activation barrier of this model reaction is relatively high $(\Delta G_{a,298} = 29.9 \text{ kcal mol}^{-1})^{7}$, this process is kinetically and thermodynamically favoured over the corresponding carbocyclization reactions leading to 7- or 8-membered ring intermediates.8 This result is in agreement with the exclusive formation of nine-membered ring tricyclic compounds 3, as experimentally observed. Intermediate 2M is then transformed into the neutral complex 4M via the initially formed complex 3M (where the NTf₂⁻ anion is weakly bonded to 2M) through transition state TS2. The latter saddle point is associated with

Fig. 1 Computed reaction profile for the reaction of allenyl-β-lactam 1M and [(PMe₃)AuNTf₂] catalyst.¹¹

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Scheme 4 Mechanistic explanation for the gold-catalyzed formal 5-exo hydroalkylation of allenyl-tethered oxyarenes 4.

the easy and exergonic proton abstraction in 2M by the NTf₂ anion ($\Delta G_{a,298} = 1.4 \text{ kcal mol}^{-1}$ and $\Delta G_{298} = -16.9 \text{ kcal mol}^{-1}$, from 3M). The addition of the readily formed NHTf2 to 4M forms complex 5M, which evolves into complex 6M via TS3 (associated with the protonolysis reaction of the carbon-gold bond). The latter process is also highly exergonic (ΔG_{298} = -19.4 kcal mol⁻¹) and proceeds with a very low activation barrier ($\Delta G_{a,298} = 0.9 \text{ kcal mol}^{-1}$). ^{9,10} Thus, it can be concluded that the initial 9-endo carbocyclization reaction constitutes the bottle-neck of the process in view of the corresponding endergonicity and relatively high activation barrier. Finally, the reaction ends up with the release of the AuPMe₃⁺ catalyst, which is coordinated to the endocyclic C=C double bond of 6M, to produce the final tricyclic species 7M.

A mechanistic rationale for the gold-catalysed conversion of allenyl-tethered sugars 4 into fused cyclopentenes 5 is more intricate. It is worth noting that the cyclization affords adducts 5 from an allene umpolung hydrofunctionalization instead of that from the usually preferred conventional hydrofunctionalization. The pathway proposed in Scheme 4 looks valid for the formation of tricycles of type 5. It could be presumed that the initially formed gold complex 4-Au(L), through coordination of the gold salt to the distal allenic double bond, undergoes a 1,6hydride shift (rare transfer of hydride versus normal nucleophilic group attack), giving rise to the oxonium species 8. Intramolecular trapping of the oxonium group by the alkenylgold moiety in intermediates 8 generates cationic species 9, through formal 5-exo hydroalkylation. Finally, demetalation yields fused cyclopentenes 5 and regenerates the gold catalyst (Scheme 4).

Preliminary DFT calculations on the model (aryloxy)allenyltethered sugar species 8M (see Fig. S3 in the ESI†) indicate that the direct 1,6-hydride shift from the distal double bond-coordinated complex 8M-Au occurs with a relatively high activation barrier ($\Delta G_{a,298} = 37.7 \text{ kcal mol}^{-1}$, that is 25.2 kcal mol⁻¹ above the separate reactants 8M and AuPMe₃⁺). This step is followed by the highly exergonic C-C bond forming reaction (i.e. oxonium trapping) via TS5 with an activation barrier of 26.1 kcal mol^{-1} . Further DFT calculations involving the NTf2-mediated 1,6-hydride shift and more realistic species leading to a process more compatible with a reaction at room temperature are currently underway.

In conclusion, the divergent gold-catalysed reactivity (C_{sp2}-H versus C_{sp3}-H) of aryloxy-tethered allenes has been studied. We report herein an efficient gold-catalysed 9-endo carbocyclization to fused tricyclic β-lactams from easily accessible aryl allene substrates under mild conditions. In salient contrast to the reaction of (aryloxy)allenyl-tethered 2-azetidinones, the allenyl sugar derivatives provided the 5-exo hydroalkylation adducts as the sole products. The reactions were found to proceed with complete control of product regio- and chemoselectivity.

Support for this work from the MICINN [CTQ2009-09318, CTQ2010-20714-C02-01, and Consolider-Ingenio (CSD2007-00006)], and CAM (Projects S2009/PPQ-1752 and S2009/PPO-1634) is gratefully acknowledged. S.C. thanks Comunidad Autónoma de Madrid and Fondo Social Europeo for a predoctoral contract. I.F. is a Ramón y Cajal fellow.

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- 8 See Fig. S2 in the ESI[†].
- 9 In sharp contrast, the corresponding proton abstraction and protonolysis reactions in the 7-membered ring formation process proceed with much higher activation barriers ($\Delta G_{a,298} = 16.9$ and 37.1 kcal mol⁻¹). This makes the 7-membered ring formation a non-competitive transformation. See Fig. S2 in the ESI[†].
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