Cyclization of 1,6-Enynes Catalyzed by Gold Nanoparticles Supported on TiO₂: Significant Changes in Selectivity and Mechanism, as Compared to Homogeneous Au-Catalysis

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Gold nanoparticles supported on TiO₂ (1.2 mol %) catalyze, for the first time under heterogeneous conditions, the cycloisomerization of a series of 1,6-enynes in high yields. In several cases, the product selectivity differs significantly as compared to homogeneous Au(I)-catalysis. Based on product analysis and stereoisotopic studies it is proposed that the major or exclusive pathway involves a 5-exo cyclization mode to form stereoselectively gold cyclopropyl carbenes that undergo a single cleavage pathway, in contrast to homogeneous Au-catalysis where the double cleavage pathway operates substantially.

The homogeneous Au(I)-catalyzed cyclization of 1,6 enynes has been extensively studied during the past decade, revealing a variety of unprecedented modes of skeletal rearrangements.¹ The puzzling mechanistic peculiarity of

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cycloisomerizations and the discussions over the nature of the intermediates (carbenes or carbocations) continue to fascinate.2 Under heterogenized conditions the reaction has been marginally studied. A specific example was recently presented using a gold(I) complex covalently bound on a polystyrene resin that provides the typical product selectivity of homogeneous $Au(I)$ catalysts.³ The identification of stabilized ionic gold species on metal oxide supported gold nanoparticles $(Au NPs)$,⁴ and the fact that epoxides⁵ and silanes⁶ are readily activated by Au NPs on titania $(Au/TiO₂)$,⁷ let us recently examine its catalytic

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⁽⁷⁾ Au/TiO₂ (\sim 1 wt % in Au) is commercially available.

efficiency in heterogeneous alkyne activation. Thus, we found that $Au/TiO₂$ catalyzes the cycloisomerization of aryl propargyl ethers into $(2H)$ -chromenes.⁸ Byproducts from oxidative dimerization promoted by molecular oxygen were also formed. It was postulated that formation of (2H)-chromenes is catalyzed by positively charged Au species, whereas dimeric $2H, 2'H-3, 3'$ -bichromenes derive via a redox $\text{Au}^{\text{III}} - \text{Au}^{\text{I}}$ catalytic cycle. Although Au/CeO_2 have reported to be inactive against 1,6-enyne cyclization, we were prompted to test $Au/TiO₂$ as a catalyst for such a purpose and develop the first heterogeneous goldcatalyzed approach in this well-studied under homogeneous conditions reaction.

To our delight, gold nanoparticles (1.2% mol in overall Au content) catalyze in high yields the cyclization of a series of 1,6-enynes in refluxing DCE, which provides once more proof for the existence of ionic gold species on $Au/TiO₂$ as the potent catalytic sites. However, on many occasions the observed product selectivity differs significantly from that under homogeneous Au(I) conditions. The initial examples using 1,6-enynes bearing a simple propargylmoiety are presented in Scheme 1. Parent enyne 1 provides exclusively the five-membered ring carbocycle 1a in 94% isolated yield. This selectivity resembles its Pd(II)-, Pt(II)-, Ir(I)-, In(III)-, or Ru(II)-catalyzed cyclization¹⁰ in contradiction to homogeneous $Au(I)$ -catalysis,¹¹ which provides primarily a six-membered ring cyclization product, with 1a being a minor one. For the case of 2, diene 2a is formed as occurs under homogeneous conditions, 12 accompanied by \sim 5% of 2b. Crotyl-substituted 3 (E/Z = 3/1) not only led primarily to product $3a^{11}$ ($E/Z \approx 3/1$) but also formed a minor amount (12%) of the sixmembered 3b ($Z/E \approx 3/1$). Product 3b had been reported as a minor one in the In-catalyzed cyclization of 3^{10d} and as major in the Ru-catalyzed¹³ cyclization. Cinnamylsubstituted 4 decomposes under the reaction conditions suffering oxidative cleavage of the double bond. Enyne 5 mainly affords the nonconjugated diene 5a (65% relative yield), conjugated diene $5b$ (5%), a minor fraction of two six-membered ring isomers 5c and 5d (total 23%), oxidative dimerization product $5e^{14}$ (5%) as an equimolar mixture of diastereomers (meso and dl), and a small fraction (\sim 2%) of two other dimers (isomeric to 5e) as

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revealed by GC-MS analysis. Diene 5a is produced in the $PtCl₂$ -catalyzed cyclization of 5^{15} Under homogeneous Au(I)-conditions 5a was seen only if the reaction solvent is polar $DMSO₁₁¹¹$ while in DCM 5b (a minor product under our reaction conditions) is exclusively formed.

Scheme 1. Cyclization of Enynes $1-5^a$ Catalyzed by Au/TiO₂^{b,c}

 a Z = C(COOMe)₂. ^b1.2 mol % of catalyst. ^cDCE, 70 °C.

For further exploration we examined the cyclization of internal enynes 6-11 (Scheme 2). Generally, their reaction rates are lower; however with one exception high yields were obtained. The cyclization of 6 provides a striking difference among homogeneous and heterogeneous gold catalysis. In the presence of $Au/TiO₂$, diene 6a (single cleavage product) is exclusively formed, in contrast to homogeneous Au(I)-conditions, where the isomeric (E) -3a is formed via a double cleavage rearrangement mechanism.16 An identical cyclization mode was found in the case of 7 which selectively provides the acid-sensitive carbocycle 7a in 75% yield. Enyne 8 affords primarily a mixture of five-membered ring allene 8a (major), the sixmembered nonconjugated diene 8b (minor), and a mixture of three dimeric products (GC-MS) in an approximately 6% relative ratio, which could not be separated and characterized properly. The formation of allenes in the gold-catalyzed cyclization of some benzyl-substituted

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Scheme 2. Cyclization of Enynes $6-11^a$ Catalyzed by Au/TiO₂^{*b,c*}

 a Z = C(COOMe)₂. ^b1.2% mol of catalyst. ^cDCE, 70 °C. ^dA mixture of three products from an oxidative dimerization pathway was also formed in ∼6% relative yield (GC-MS analysis).

internal enynes has been recently demonstrated.¹⁷ Concerning phenyl substituted internal enynes, 9 lead to the five-membered ring carbocycle 9a in 81% isolated yield, in contrast to homogeneous Au(I)-catalysis where a fused bicyclic cyclobutene derivative¹⁸ is primarily formed. It is also known that 9 mainly cyclizes into 9a yet under Rh(II) catalysis conditions.19 Substrate 10 was unreactive even under prolonged refluxing conditions, while 11 afforded the fused tricycle 11a in 83% yield, in accordance to the homogeneous gold catalysis conditions. $2f,18$

Useful mechanistic information was acquired upon studying the $Au/TiO₂-catalyzed$ cycloisomerization of some deuterium labeled 1,6-enynes, presented in Scheme 3. Generally, sp-C deuterium labeled terminal alkynes suffer gradual isotopic depletion under our reaction conditions; thus for substrates requiring a prolonged time to cyclize, mechanistic conclusions could not be drawn.

Scheme 3. Cyclization of Some Deuterium Labelled 1,6-Enynes

Partial deuterium exchange in terminal alkynes is also known under homogeneous $Au(I)$ -conditions²⁰ and most likely involves the reversible formation of Au-acetylides.²¹ Despite the partial deuterium depletion²² on the starting materials for the case of enynes 2-d and 5-d, tentative conclusions regarding the disposition of the D-atom were acquired. Thus, in product 2a-d, deuterium resides on the exocyclic secondary sp^2 -carbon of the pentacycle. In the cyclization of 5-d, deuterium is stereoselectively attached on the exocyclic double bond of the major product 5a-d (tentative assignment due to the 70% isotopic depletion in cyclization product; see Supporting Information). An identical stereochemical conclusion has been reported in the cyclization of $5-d$ catalyzed by a homogeneous $Au(I)$ complex in $DMSO¹¹$. This stereoselective deuterium disposition excluded the participation of an Alder ene- type process. Additionally, the stereoselective introduction of D in the side allyl chain of enynes 1 and 6, either by $LiAlD₄$ addition to propargyl alcohol²³ (synthesis of $1-d$) or by $LiAlH₄$ addition to propargyl alcohol followed by quenching with D_2O (synthesis of 1'-d and 6-d), was achieved.²⁴ Regarding the cyclization of 1-d, the D-atom appeared exclusively on the endocyclic secondary sp^2 -carbon, whereas, for $1'-d$ and $6-d$, stereospecific disposition of the stereochemical integrity of the terminal olefinic carbon atom was found in cyclized products $1/a-d$ and $6a-d$, respectively.

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Scheme 4. A General Mechanistic Scenario for the Cyclization of 1,6-Enynes Catalyzed by Au/TiO2

Prior to the mechanistic discussion, we propose that the stereoselective disposition of H and D in product 5a-d indicates catalysis by monodentate to the enynes, gold(I) bearing, oxidized nanoclusters.²⁵ Based on the product analysis of Schemes $1-2$, and the stereoisotopic results in Scheme 3, we present in Scheme 4 a general mechanistic scenario for the $Au/TiO₂$ -catalyzed cycloisomerization of 1,6-enynes, considering catalysis by Au(I) species. It is clear that in all cases the major pathway is a 5-exo cyclization to form stereoselectively cyclopropyl carbene I, as proposed earlier.^{1f} Depending on the substituents, I may undergo ring expansion to cyclobutyl tertiary carbocation II or isomerize to monocyclic III (typical example, the cyclization of 5). Carbocation II in turn eliminates Au^+ in a conrotatory fashion²⁶ (typical examples, the cyclization of 1'-*d* and 6-*d*). If $R_1 = H$ and R_2 , R_3 , and $R_4 = Me$, carbocation III undergoes an intramolecular hydride

shift¹⁷ to form allene **8a**. A single cleavage pathway (especially in the cyclization of 1 and 6) forming an exocyclic carbocation, as suggested 2a,b under homogeneous Au(I)-catalysis conditions, is less likely to participate under our heterogeneous conditions due to the requirement for the formation of a pseudoprimary carbocation (see Supporting Information, p S38). Analogously, the most likely fate of the minor 6-endo cyclization mode is to form stereoselectively cyclopropyl carbene IV, which depending on the substituents may lead either to fused bicyclic carbocation V or to monocyclic VI. The minor products of oxidative dimerization (cyclization of enynes 5 and 8) more likely arise via a redox $Au^{III} - Au^{I}$ catalytic cycle primarily from deprotonated intermediate III, as postulated earlier in the case of aryl propargyl ethers.⁸ Therefore, $Au/TiO₂$ catalyzes the cyclization of nonprenyl-substituted 1,6-enynes exclusively via a single skeletal rearrangement of gold cyclopropyl carbenes into cyclobutyl carbocations, in contrast to homogeneous catalysis by Au(I) and Au(III) complexes or salts that proceed via mixed single and double cleavage pathways. 27 It is possible that the polar support plays a crucial role in our case, through stabilizing the transformation of cyclopropyl carbenes into rearranged cyclobutyl carbocations or, alternatively, their opening into carbocation III for the case of prenyl-substituted enynes.

In conclusion, we have presented for the first time a heterogeneous gold-based catalytic approach regarding the cyclization of 1,6-enynes. Product analysis and stereoisotopic studies reveal that the main pathway involves an initial 5-exo cyclization by Au(I) species to form stereoselectively gold cyclopropyl carbenes. Isomerization of carbenes into bicyclic cyclobutyl carbocations via a single cleavage pathway, followed by conrotatory ring opening, results in the stereoselective disposition of the terminal olefinic bond of reacting enynes. Depending on the substituents, ring opening of cyclopropyl carbenes into open carbocations may also take place. By contrast, based on the literature data, in the homogeneous Au-catalyzed cyclization of 1,6-enynes, the double cleavage pathway operates substantially. The current work exemplifies the unique nature of Au nanoparticles in catalysis. 28

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Supporting Information Available. Copies of ${}^{1}H, {}^{13}C$ NMR of all products and key compounds. This material is available free of charge via the Internet at http://pubs. acs.org.

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