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## Access to the Protoilludane Core by Gold-Catalyzed Allene-vinylcyclopropane Cycloisomerization

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Gold(I)-catalyzed allene-vinylcyclopropane cycloisomerization leads to the tricyclic framework of the protoilludanes in a single step by a reaction that involves a cyclopropane ring expansion and a Prins cyclization.

Illudanes and protoilludanes have attracted much attention from the perspective of their biosynthesis,<sup>1</sup> biology,<sup>2</sup> and organic synthesis.<sup>3</sup> Representative members of this numerous family of sesquiterpenes are  $\Delta^6$ -protoilludene (1),<sup>4,5</sup> illudol,<sup>6</sup> repraesentin A (3),<sup>7</sup> and russujaponol D (4)<sup>8</sup> (Figure 1).

Access to this class of compounds still poses important synthetic challenges due to their structural complexity.

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Figure 1. Representative protoilludane sesquiterpenes.

As part of a program on the development of new gold(I)catalyzed cascade reactions for the synthesis of complex sesquiterpenes,<sup>9</sup> we now report a new approach for the construction of the skeleton of this family of natural products in one single step through a gold-catalyzed cycloisomerization.

Cycloisomerization reactions catalyzed by gold(I) complexes have been intensively investigated and represent one of the most powerful methods for the construction of complex molecules in a single step.<sup>10</sup> Among these various transformations, we demonstrated that complex tricyclic compounds such as **6** with an octahydrocyclobuta[*a*]pentalene skeleton can be obtained by a sequence involving a ring expansion and a Prins cyclization from cyclopropylenyne **5** (Scheme 1).<sup>11,12</sup>

Scheme 1. Gold(I)-Catalyzed Cycloisomerization of Cyclopropylenyne 5<sup>11</sup>



Based on this work, we decided to prepare the related cyclization of allenes with vinylcyclopropanes to access the carbon skeleton of the illudanes from a common intermediate. A plausible mechanism for this reaction is depicted in Scheme 2. Coordination of  $AuL^+$  to the allene (I) could trigger a 5-*exo-trig* cyclization to furnish cationic intermediate II, which could undergo a ring expansion to generate III. A Prins cyclization of the vinyl gold with the oxonium cation could give gold(I) carbene intermediate IV. Finally, proton elimination followed by demetalation would provide V or VI. Regarding the relative configuration of II, precedents exist for the formation of related intermediates with both the *cis* or *trans* configuration in gold(I)-catalyzed cycloisomerizations of allenenes.<sup>13</sup>

We first studied the reaction of substrate *E*-7. The cyclization was carried out satisfactorily using [IPrAu(NCPh)]SbF<sub>6</sub> (**B**) (3 mol %) in CH<sub>2</sub>Cl<sub>2</sub> at 0 °C to give **8** in 94% yield as a 6.5:1 mixture of two alkene regioisomers (Scheme 3).<sup>14</sup> On the other hand, cyclization of substrate **9** with two methyl

groups at the allene terminus furnished tricyclic compound **10** in 54% yield as a single isomer.





Scheme 3. Gold(I)-Catalyzed Cyclization of Allene-*E*-vinylcyclopropanes *E*-7 and 9



Although the allene-vinylcyclopropane cyclization provided the desired tricyclic system as originally planned, the relative configuration of **8** and **10** was the opposite to that of the natural protoilludenes. In keeping with the stereospecificity demonstrated in gold(I)-catalyzed cyclization of related enynes,<sup>10</sup> we prepared substrate **Z**-7 from known malonate **11**<sup>15</sup> (Scheme 4). Thus, the anion of malonate **11** was alkylated with mesylate **12** to give allenyl malonate **13** in 86% yield. The acetal was then hydrolyzed to furnish aldehyde **14** (88%), which was alkenylated with phosphonate **15** to yield ketone **16** in almost quantitative yield with the desired Z configuration.<sup>12b,16</sup> Finally, formation of silvlenol ether with TBSOTf and Et<sub>3</sub>N, followed by

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<sup>(14)</sup> The minor isomer **8b** could not be isolated, and its structure was tentatively assigned as a derivative of 3,7b-dimethyl-1,2,4a,5,6,7,7a,7b-octahydro-2aH-cyclobuta[e]inden-2a-ol (stereoisomer of **18a**, Scheme 5), with the trans configuration between the five- and six-membered rings.

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Simmons-Smith cyclopropanation, led to Z-7 (67% over two steps).

Scheme 4. Synthesis of Substrate Z-7 MsC F  $OM_{c}$ OMe F NaH, THF/DMF



Cyclization of Z-7 was best carried out in 1,2-dichloroethane at 23 °C in the presence of catalyst A (3 mol %).<sup>17</sup> The reaction led to the formation of two major products 18a and 18b in a 2:1 ratio (82%, NMR yield), which could not be separated by chromatography (Scheme 5). Their structures were assigned by transformation into crystalline derivatives. First, reduction of the malonate with LiAlH<sub>4</sub> led to the isolation of crystalline diol 19 in 52% yield (2 steps from Z-7), whose structure was determined by X-ray diffraction, which confirmed the trans configuration at the junction between the five- and six-membered rings (Figure 2).<sup>18a</sup> On the other hand, desilylation with tris(dimethylamino)sulfonium difluorotrimethylsilicate (TASF), followed by esterification with *p*-nitrobenzoyl chloride and recrystallization, furnished ester 20 in 20% yield (2 steps from Z-7), whose structure was again assigned by X-ray diffraction (Figure 2).<sup>18b</sup> Hydrogenation of 19 in an autoclave under 50 bar of H<sub>2</sub> using a Pearlman catalyst furnished 21 in 95% yield as a single diastereoisomer. Monotosylation of the diol followed by LiAlH<sub>4</sub> reduction gave alcohol 22 in 60% yield as a 1:1 mixture of diastereoisomers at the newly formed stereocenter, which is structurally related to russujaponol D (4).

We also explored the introduction of the oxygen functionality on the six-membered ring by hydroboration/ oxidation (Scheme 6). The reaction led to a mixture of several diastereomeric alcohols, which could not be separated by chromatography. However, treatment of this mixture with Dess-Martin periodinane gave two ketones 23a and 23b in 59% and 19% yields respectively, whose

(18) (a) X-Ray crystal structure of 19: CCDC 953503. (b) X-Ray

Scheme 5. Gold(I)-Catalyzed Cyclization of Z-7





Figure 2. X-ray structures of 19 (a) and 20 (b).

relative configurations were determined by NOE analysis. The TBS group of the major compound 23a was removed with TASF, and the resulting alcohol was converted into

<sup>(17)</sup> See Supporting Information for a screening of catalysts.

crystal structure of 20: CCDC 953502.

an unstable mesylate intermediate, which was eliminated with DBU to furnish enone 24. No epimerization was observed under these conditions to give the corresponding *cis*-stereoisomer of 24.<sup>19</sup>



Our results demonstrate that intermediates **IIa** and **IIb** do not undergo equilibration and that the cyclopropylcarbenium to cyclobutane ring expansions occurs stereospecifically to form **IVa** or **IVb**, by intramolecular Prins reaction (Scheme 7). Regarding the configuration of cyclopentanes **IIa** and **IIb**, formation of **18a** as the major product in the cyclization of **Z-7** suggests that the gold(I)catalyzed allene-vinylcyclopropane cycloisomerization leads to intermediates **II** with the *trans*-relative configuration. However, compounds **8a** and **18b** could arise from either *trans*- or *cis*-configured intermediates. Scheme 7. Stereospecific Cyclizations of E- and Z-I



In summary, we have shown that the gold-catalyzed allene-vinylcyclopropane cycloisomerization leads directly to complex tricyclic compounds with the skeleton of the protoilludanes. Ongoing work is focused on exploring new routes and catalysts to access the desired *cis*-fusion as well as on developing asymmetric syntheses of these natural compounds.

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**Supporting Information Available.** Experimental procedures and characterization data for all new compounds. This material is available free of charge via the Internet at http://pubs.acs.org.

<sup>(19)</sup> DFT calculations (B3LYP, 6-31G\*) on model *cis*-and *trans*-**25** show that the trans isomer is the most stable ( $\Delta\Delta H^{\circ} = 2.5 \text{ kcal} \cdot \text{mol}^{-1}$ ).

The authors declare no competing financial interest.