DFT study on the gold(I)-catalyzed cycloaddition and rearrangement reactions of allene-containing allylic silyl ether

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Research Article

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Abstract

The DFT calculation of the B3LYP level was carried out to explore the reaction mechanism of the synthesis of spirocyclo[4, 5]decane skeleton by gold-catalyzed allenyl compounds. The more accurate energy under the CH3CN solvent in the experiment is calculated by the single point energy of the SMD model. Computational studies have shown that the reaction consists of three main steps: intramolecular cycloaddition of the end group carbon atoms of allenyl and vinyl groups, the semipinacol rearrangement process in which the four-membered ring is reconstructed into the five-membered ring, the elimination reaction releases the catalyst and obtains the product. The calculation results show that Zheng et al. reported that the gold-catalyzed synthesis reaction can easily occur under the experimental conditions due to its low activation free energy (12.42–16.79 kcal/mol). Furthermore, it was found that the MOMO(CH₂)₂ substituent has higher reactivity than the corresponding reactant of the phenyl substituent.

1. Introduction

Molecules with a spirocyclo[4, 5]decane skeleton have good biological properties due to the important moiety of the quaternary carbon center, which is widely present in various pharmaceutical molecules and natural bioactive products, such as spirojatamol, erythrodiene and cedrene, etc [1–6]. However, the key part of the structure of the scaffold containing quaternary carbon is difficult to construct directly in the synthesis due to the large steric repulsion [7–11]. The development of an effective synthesis method is also one of the synthesis problems that need to be overcome, and it has also attracted a large number of chemists to carry out related research work [12–15]. Over the years, some synthetic methods have been developed and molecular functionalization studies have been carried out. Especially intramolecular cycloaddition or intermolecular cycloaddition reaction is mostly, such as intramolecular dearomatization cyclization of phenols [16], photo-induced intramolecular [2 + 2] cyclization [17], intermolecular [4 + 2] or [3 + 2] cycloaddition [18–19]. In addition, in order to further improve the reaction efficiency and enhance the diversity of product structures, transition metal catalysis and rearrangement reactions are also widely used in the synthesis of such products. The semipinacol rearrangement is also considered to be a very effective method to build a quaternary carbon skeleton through migration or recombination. Studies have reported that both vinylogous α-ketol and α-hydroxy epoxide alcohol have been used to construct spirocyclo[4.5] decane skeletons through semipinacol rearrangement [20–22].

In order to further explore new rearrangement modes for introducing the required functional groups more effectively, recent studies by Zheng et al. found that Au(I)-catalyzed allyl alcohol compounds (a or b) can easily undergo intramolecular cyclization / semipinacol rearrangement reaction, and high yields (95% and 65%) of spirocyclic [4.5]-decane skeleton (Scheme 1) can be generated by Au(PPh₃)Cl catalyst in CH₃CN solution at room temperature [23]. Based on the experimental research results of Zheng's research team, a key intermediate was given to predict the possible reaction process. In order to further explore the detailed catalytic reaction mechanism, a feasible channel was given to synthesize spirocyclo[4.5]decane skeleton of quaternary carbon center from allene-containing allylic silyl ether (Scheme 2). First, the allylic alcohols reactant (a) is activated by Au(I) catalyst to obtain intermediate I. Then, the allene group and
the terminal carbon of the vinyl group undergo intramolecular cyclization to form intermediate \( \text{II} \). Next, a rearrangement reaction occurred on the four-membered ring in intermediate \( \text{II} \) and the corresponding five-membered ring intermediate \( \text{III} \) was formed, which is also the possible intermediate structure mentioned in the experimental report. Finally, water molecules are added to the intermediate \( \text{III} \) to obtain the final product \( \text{a-p} \) and release the gold (I) catalyst and TESOH molecules.

As far as we know, there is no relevant theoretical research report to explain the new gold-catalyzed synthesis of quaternary carbon-containing framework from allenes reported by Zheng and co-workers. In this article, based on the experimental phenomenon of Zheng et al., we carried out a calculation study of the density functional theory B3LYP method on the mechanism of the cyclization and rearrangement of allene-containing allylic silyl ether catalyzed by gold (I) catalyst. We hope to clarify the specific reaction mechanism and learn more about the factors that control the activity of this reaction through theoretical research.

2. Computational Details

The DFT calculations [24] of all research systems are performed using Gaussian 09 software [25]. Both geometric optimization and frequency calculation are done using the B3LYP calculation method [26] of density functional theory. The SDD basis set containing the effective core relativistic pseudo potential [27] is selected for the Au, while other atoms applied 6-31G \((d, p)\) basis set [28]. We also determined and verified the TS structure to correctly connect the two corresponding stable structures through IRC calculations [29]. In order to obtain the precise energy corresponding to the experimental reaction solvent system, we used a larger basis set to recalculate the single-point energy of all optimized geometries in the corresponding experimental solvents. This is also a widely used method to obtain accurate energy in catalytic reactions [30–34]. The larger basis set of 6-311++G \((d, p)\) (SDD with ECP for Au) in the SMD implicit solvent environment [35] with acetonitrile \((\text{solvent} = \text{CH}_3\text{CN}, \varepsilon = 35.688)\) is used. All energy values discussed in the context are relative Gibbs energy \((\text{kcal/mol})\) corrected by the single-point energy of solvation. The three-dimensional (3D) structures were obtained directly from the calculated output using CYLview software.

3. Results And Discussion

For this catalytic synthesis reaction, experimental studies have proved that \( \text{Au(PPh}_3\text{)Cl} \) is the most effective gold catalyst. According to the experimental conditions, theoretical calculations are based on \( \text{Au(PPh}_3\text{)}^+ \) as the catalyst model, which interacts with the allene reactant \( \text{a} \) in Scheme 1 and the terminal unsaturated bond is activated to obtain the initial reaction complex \( 1\text{a-int} \). The frontier molecular orbital and the electrostatic potential diagram of the reaction complex \( 1\text{a-int} \) are shown in Fig. 1. It can be seen from the LUMO molecular orbital that the \( \pi \) electrons on the terminal \( \text{C=C} \) double bond of the allene group in the molecule \( 1\text{a-int} \) are filled on the \( d \) orbital of \( \text{Au}^+ \), the \( \pi \) bond is activated and the electrons on the orbital have a large overlap degree. In addition, on the two molecular orbitals of HOMO and HOMO-1,
the non-terminal carbon-carbon double bond in the allene group and the vinyl group in the molecule have strong \(\pi\) electrons, especially the vinyl group is easy to interact with activated carbon-carbon bond of the allene group undergoes an intramolecular cycloaddition reaction to complete the ring closure. The energy gap between the HOMO and LUMO orbitals of the molecule is 0.16 hartree, and the smaller energy difference helps the electron transition to complete the cyclization process. The electrostatic potential diagram clearly shows that most of the atoms in the molecule are electropositive, only three oxygen atoms and the \(\pi\) system show weak electronegativity. In the calculation study of this reaction process, \textbf{1a-int} was used as the initial reaction complex to explore the specific reaction mechanism.

3.1 Reaction Mechanism

The free energy profiles of gold(I)-catalyzed intramolecular cycloaddition and rearrangement reactions starting from reactant \textbf{1a-int} is given in Fig. 2. The key geometric geometries giving the main structural parameters are shown in Fig. 3. In \textbf{1a-int}, the two new gold-carbon bonds Au-C1 and Au-C2 formed by the gold(I) ion and allene group are 2.403 and 2.276 Å respectively. The C1-C2 bond is also further activated by Au(PPh\(_3\))^+, elongating to 1.360 Å. From Fig. 2 along the initial complex \textit{1a-int}, the allene group and the terminal carbon atom of the vinyl group undergo an intramolecular cycloaddition reaction through the transition state \textbf{12a-ts} to form the intermediate \textbf{2a-int}. At the same time, the gold atom also transferred to the middle carbon atom of the allene group. The only imaginary frequency of \textbf{12a-ts} is 255.04 i cm\(^{-1}\) and the energy barrier of this step is 12.42 kcal/mol relative to \textbf{1a-int}, and the energy of \textbf{2a-int} is 2.42 kcal/mol. In \textbf{12a-ts}, the distance of the interacting C1-C3 bond is 2.075 Å, the Au atom is also completely transferred to the C2 atom and the Au-C2 bond is shortened to 2.128 Å, while the C1-C2 bond is stretched to 1.449 Å. From \textbf{1a-int} to \textbf{2a-int}, the unsaturated group completes the ring closure to form a cyclohexene six-membered ring structure \textbf{2a-int}, and the C1 and C3 atoms are also transformed from \(sp^2\) to \(sp^3\) hybridization. The C1-C2 and C1-C3 bonds both exhibit the characteristics of a single bond, and the bond lengths in \textbf{2a-int} are 1.517 and 1.563 Å respectively. However, the four-membered ring skeleton in the \textbf{2a-int} structure still has certain instability. He can easily construct the four-membered ring into a more stable five-membered ring through the semipinacol rearrangement process of the transition state \textbf{23a-ts} (imaginary frequency is 194.93 i cm\(^{-1}\)) to obtain a more stable intermediate \textbf{3a-int}. The activation free energy of this step is 9.59 kcal/mol relative to the intermediate \textbf{2a-int}. For \textbf{23a-ts}, the C5-C6 bond tends to dissociate and stretch to 1.758 Å, and C4-C6 has a tendency to form bonds, with a bond length of 2.239 Å. The formation of \textbf{3a-int} structure indicates that the rearrangement process has been completed, and its energy is reduced by 26.23 kcal/mol compared to \textbf{2a-int}, which is a step of strong exothermic process. The spirocyclo[4.5]decane skeleton has been initially formed in the \textbf{3a-int} structure. The C4 atom transitions to the \(sp^3\) hybrid mode and the newly formed C4-C6 bond is 1.554 Å. The six-membered ring and the five-membered ring form two planes that are approximately perpendicular to each other. Then, the H\(^+\) in the water in the reaction system attacks the intermediate \textbf{3a-int} through the transition state \textbf{34a-ts} to obtain the intermediate \textbf{4a-int} and release the catalyst Au(PPh\(_3\))^+. The activation energy of this step is relatively low with 4.44 kcal/mol, The energy of the step of generating \textbf{4a-int} is almost unchanged and only reduced by 1.62 kcal/mol. The only virtual frequency in \textbf{34a-ts} is 768.64 i cm\(^{-1}\), and its vibration
mode is mainly shown in the H between Au and C2 atoms, where the Au-H and H-C2 bond lengths are 1.834 and 1.826 Å, respectively. Finally, the excess OH\(^-\) in the water interacts with the intermediate 4a-int to remove the TESOH molecule to obtain the final product a-p, whose energy continues to decrease by 3.44 kcal/mol.

Throughout the entire catalytic reaction process, the first step of the intramolecular cycloaddition process is the rate-determining step of the entire reaction, and its energy is relatively low at 12.42 kcal/mol. This is consistent with the experimental results that the synthesis reaction can be carried out under the reaction conditions at room temperature, and the product with a high yield of 95% can be obtained. The activation free energy of the second key step of semipinacol rearrangement is lower than that of the first step, and the reaction process of the second step is easier to complete. Moreover, the entire reaction is also an exothermic process with an exothermic amount of 28.87 kcal/mol. This provides a good explanation for the detailed reaction mechanism reported by Zheng et al.

3.2 The influence of the substituents on the reactant allenyl group

According to experimental reports, in order to further investigate the influence of substituents on the reactivity, the MOMO(CH\(_2\))\(_2\) group in the reaction substrate a was replaced by a phenyl group as a new reactant b (Scheme 1), and the reaction mechanism was studied by the same theoretical calculation method. The specific reaction process and mechanism similar to the reaction substrate a is shown in Fig. 4. The geometric structure containing the main parameters is shown in Fig. 5. In the reaction complex 1b-int, the newly formed Au-C1 bond is 0.018 Å longer than that of 1a-int, while the Au-C2 bond is shorter than the corresponding one by 0.025 Å. It can be clearly seen from Fig. 4 that starting from the complex 1b-int and going through the transition states 12b-ts, 23b-ts and 34b-ts along the reaction path, it also goes through three steps, namely intramolecular cycloaddition, semipinacol rearrangement and the elimination process of releasing the catalyst, etc. The activation free energy of the three steps is 16.79, 12.49 and 5.13 kcal/mol respectively, which are 4.37, 2.90 and 0.69 kcal/mol higher than the activation free energy of the 1a-int reaction channel respectively. The reaction system shows that the cycloaddition process (the first step) is still the rate-determining step of the entire catalytic reaction process, and its reaction barrier is also slightly higher than 1a-int. This reaction channel is also a strong exothermic process, and its exothermic heat is almost the same as that of the 1a-int reaction channel, its value is -29.59 kcal/mol, which is only 0.72 kcal/mol higher. From the structural point of view, the basic skeleton of the intermediate and transition state structure is basically similar to that of the 1a-int reaction channel, except that the phenyl group replaces the MOMO(CH\(_2\))\(_2\)- group to cause systematic changes in some parameters. The IRC calculation results confirm that 1b-int, 2b-int, 3b-int and 4b-int are connected to the corresponding intermediates 1b-int, 2b-int, 3b-int and 4b-int, respectively.

The Au(I)-catalyzed reaction of allene-containing allylic silyl ether to synthesize spirocyclo[4.5]decane skeleton has lower activation free energy (channels a and b are 12.42 and 16.79 kcal/mol, respectively). Moreover, the reaction barrier of intramolecular cycloaddition is higher than that of semipinacol rearrangement. However, comparing the reactivity of gold-catalyzed substrates with two different substituents, it is obvious that the activation barrier of MOMO(CH\(_2\))\(_2\)- is 4.37 kcal/mol lower than that of...
phenyl, and it has higher reactivity. The calculated results can well support the experimental report results. The experiment shows that the yields of the two products \textbf{a-p} and \textbf{b-p} are 95% and 65%, respectively. This experimental phenomenon is also consistent with the calculation result that the product \textbf{a-p} has a lower activation energy, and the product \textbf{b-p} has a higher energy barrier.

4. Conclusion

In this work, we provide detailed calculation studies and theoretical analysis of DFT (B3LYP) level for the synthesis of spirocyclo[4.5]decane skeleton catalyzed by gold complexes. Detailed calculations and precise parameters of the geometry and energy of the intermediates and transition states on the reaction channel are given. Based on the experimental report of Zheng et al., the optimal reaction path of the catalytic reaction was given, including three main steps: intramolecular cycloaddition, semipinacol rearrangement, and elimination of the release catalyst. The first step of the reaction, the cycloaddition process is the rate determining step of the entire catalytic cycle. Studies have found that this type of reaction has relatively low activation free energy, and the reaction barrier for the product \textbf{a-p} is 12.42 kcal/mol. For gold catalyst substrates with two different substituents, MOMO(CH2)2- is more reactive than phenyl. The detailed theoretical calculation results are completely consistent with the experimental phenomenon, which provides a good explanation and supplement for the experimental research.

Declarations

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Conflict of interest

The authors declare that they have no competing interests.

Ethics approval

Consent to participate

Consent for publication

Written informed consent for publication was obtained from all participants.

Availability of data and material:

Not applicable.

Code availability
Contributions:

YaMei Zhao: Problem selection, writing and data analysis.

MengDan Huo: Methods, project management, result analysis, manuscript editing.

HongJi Zhou: Data analysis, writing.

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**Figures**

![Figure 1](https://example.com/figure1.png)

**Figure 1**
Frontier molecular orbital and electrostatic potential diagram of the complex 1a-int

Figure 2

Gibbs free energy profiles for the Au(I)-catalyzed intramolecular cyclization and rearrangement of reactant a (R = MOMOCH2CH2- in Scheme 1). The energies are given in kcal/mol.
Figure 3

The optimized structures with the key bond lengths in Å for pathway a are shown in Fig. 2.
Figure 4

Gibbs free energy profiles for the Au(I)-catalyzed intramolecular cyclization and rearrangement of reactant a (R = Ph in Scheme 1). The energies are given in kcal/mol.
Figure 5
The optimized structures with the key bond lengths in Å for pathway b are shown in Fig. 4.

Supplementary Files
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- Schemes1and2.pdf