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Origins of unique gold-catalysed chemo- and siteselective C-H functionalization of phenols with diazo compounds†

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In past decade, gold revealed more and more unique properties in carbene chemistry. It was disclosed in our recent communication (*J. Am. Chem. Soc.* 2014, **136**, 6904) that gold carbenes have unprecedented chemo- and site-selectivity and ligand effect toward the functionalization of C–H bonds in phenols. In this full article, we report a comprehensively combined theoretical and experimental study on the mechanism of the insertion of gold carbenes into C–H and O–H bonds in phenol. It significantly revealed that the ligands have an important effect on C–H insertion and the reaction proceeds through a pathway involving the formation of an enolate-like intermediate. Moreover, two water molecules serving as a proton shuttle are believed to be the key issue for achieving chemoselective C–H functionalization, which is strongly supported by the DFT calculations and control experiments. It is the first time that a clear explanation is given about the prominent catalysis of gold carbenes toward C–H functionalization based on a theoretical and experimental study.

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Introduction

Phenols are not only frequently found in natural products and pharmaceutics, but are also powerful versatile platforms in organic synthesis.1 Thus, the catalytic functionalization of phenols in a chemo- and site-selective manner is the most encompassing issue in organic chemistry.2 In this context, the transition-metal-promoted decomposition of diazo compounds to generate metal carbenes has broad applications³ in the direct site-selective functionalization of phenols. In most cases, O-H insertion takes place catalysed by a series of transition-metal catalysts such as copper, palladium, iron, and rhodium complexes.4 In contrast, very recently, we and Shi independently found the unprecedented C(sp²)-H bond functionalization of phenols with α-diazoesters,5 furnishing para-substituted phenols Pro-c8 via para C-H functionalization rather than the O-H insertion product Pro-o8. Meanwhile, the chemoselectivity is heavily dependent on the nature of the ligand of the gold

In a previous work on rhodium catalysed intramolecular aromatic C(sp²)-H insertion reactions, Padwa and Doyle assumed that the reactions were triggered by the electrophilic addition of the metallocarbene intermediate to the aromatic ring, then followed by a [1,2]-H migration step.9 Nevertheless, the reaction mechanisms for the prominent activity of gold carbenes toward C-H bonds still remain elusive. On the other hand, despite Yu and coworkers¹o having performed the mechanistic study of the copper- and rhodium-catalysed O-H insertion of water with carbene using DFT calculations, the mechanism of gold-catalyzed O-H insertion has not yet been explored. In principle, there are three pivotal questions

catalyst6 (Table 1). Although Nakamura and Pérez etc.7 have theoretically studied the mechanism of C(sp³)-H insertion with carbenes, and Shi and co-workers5b analysed the electronic structures of gold carbenes by using Density Functional Theory (DFT) calculations, no comprehensive mechanistic insight on the overall pathways of aromatic C(sp2)-H insertion with carbenes has been provided so far. To further understand the origin of this unique behaviour of gold carbenes8 in site selective C-H functionalization of phenol, we decided to carry out DFT calculations and an experimental study to gain insights on the mechanism. Herein, we present a combined computational and experimental study to elucidate the origin of the ligand effect on the chemoselectivity. Furthermore, the trace amount of water in the reaction mixture serving as a proton shuttle is proposed to be the key to give rise to the product **Pro-c8** of C-H insertion, which is supported by our current DFT calculations and control experiments.

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Table 1 $C(sp^2)$ -H bond functionalization of phenols with α -diazoesters under various conditions

Entry	Catalyst (5 mol%)	Time	Pro-c8 ^a	Pro-o8 ^a
1 ^b	Cu(OTf) ₂	5 min	0	11
2^b	Cu(OTf)·toluene	5 min	0	9
3^b	[Pd(CH ₃ CN) ₂ Cl ₂]	12 h	0	33
4	Fe(OTf) ₂	12 h	NR	
5 ^c	$FeCl_2$	12 h	0	0
6^b	$[Rh_2(OAc)_4]$	5 min	0	36
7	Ph₃PAuSbF ₆	5 min	33	45
8	$(PhO)_3PAuSbF_6$	5 min	82	0

^a NMR yield. ^b The conversion of **1** is 100% and the major product is dimer of **1**. ^c Trace amount of dimer from **1** was detected.

unsolved in this transformation: (1) is the mechanism of $C(sp^2)$ –H insertion with gold carbenes the same as that with other metals such as rhodium? (2) What is the origin of the ligand effect on the chemoselectivity? (3) Besides the ligand, any other factors affect the chemoselectivity?

Computational methods

All of the DFT calculations were carried out using the Gaussian 09 software package.¹¹ The geometric structure of the intermediates and transition states were optimized and located by using

the M06 functional, 12 since the M06 functional has been demonstrated to generate accurate results for organometallics especially in the description of noncovalent interactions. The Lanl2dz basis set13 combined with the relativistic effective core potential for inner electrons and double zeta basis for covalent electrons was used to describe the heavy elements Au and P, and the 6-31G* basis set14 was used to describe the nonmetallic elements C, N, O and H. Frequency analyses were also performed based on the structures obtained in the gas phase to confirm that the intermediates are local minima and the transition states have only one imaginary frequency. Intrinsic reaction coordinate (IRC) calculations¹⁵ were performed to make sure that the crucial transition states connect the correct reactants and products. The solvent effect of CH2Cl2 was evaluated with single point calculations using the integral equation formalism model (IEFPCM)¹⁶ with the dielectric constant $\varepsilon =$ 8.93 based on the structures in the gas phase. All discussed energy values of intermediates and transition states throughout this paper are the Gibbs free energies in units of kcal mol⁻¹ that were calculated at the temperature of 298 K, including the corrections of solvation free energies from the IEFPCM model. The 3D images of the calculated structures were plotted using GaussView.17 More structural details of intermediates and transition states can be found in the ESI.†

Results and discussion

On the pathway of C-H functionalization catalysed by (PhO)₃PAuSbF₆

It has been widely accepted that metal catalysts could catalyse diazo compounds to release N_2 and form reactive Au-carbenes,

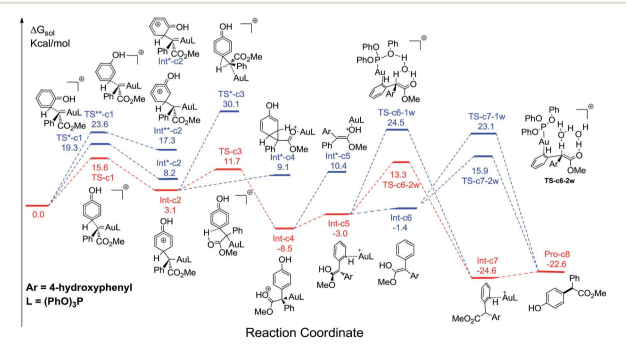


Fig. 1 The calculated free energy profiles ΔG and the corresponding structures of intermediates and transition states along different possible C-H insertion pathways catalyzed by (PhO)₃PAuSbF₆. The reasonable pathways are shown in red and other possible pathways are in blue. All values of free energies are relative to the gold carbene and phenol reactants and in the units of kcal mol⁻¹.

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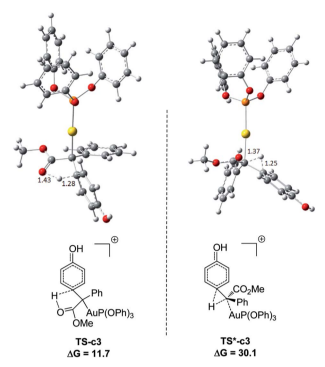


Fig. 2 Optimized structures of transition states TS-c3 and TS*-c3 corresponding to Fig. 1. The distances are in the units of angstroms and the values of free energies ΔG are in kcal mol⁻¹.

which were regarded as the precursors responsible for C-H bond insertion. 5,8a-e To account for the generation of crucial precursors preceding C-H insertion, we calculated the reaction pathways of α -diazoesters and metal catalysts AuL (L = (PhO)₃P and Ph₃P) in solution (Fig. S1 of ESI†). The calculated free energy profiles indicate that the generation of Au-carbene species is quite facile and drastically exothermic, with the low

barriers of 6.5 and 11.4 kcal mol⁻¹, respectively. Therefore, the gold carbenes and phenol are regarded as the reactants for the subsequent C-H bond insertion reactions and the sum of their free energies is set as the benchmark, with the value of 0.0 kcal mol^{-1} .

Fig. 1 displays the calculated free energy profiles for the C(sp²)-H functionalization of phenol with a gold carbene with (PhO)₃P as the ligand, where the structural details of intermediates and transition states are provided in the ESI.† The previous theoretical investigations on the C(sp3)-H insertion of rhodium, copper or silver carbenes, performed by Nakamura and Pérez, indicate that the first step of the reactions involves the formation of σ -bond complexes of carbenes and the C(sp³)-H bond.7 Recently, Pérez and coworkers suggested that the mechanism of aromatic C(sp²)-H bond insertion might be entirely different from that of aliphatic C(sp3)-H bond insertion.8c In our DFT calculations, a transition state TS-c1 with a barrier of 15.6 kcal mol⁻¹ is located for the addition of the gold carbene at the para-position of phenol. TS-c1 represents the transition state of the direct $C(sp^2)-C(sp^2)$ coupling of phenol and the gold carbene, leading to the final para-C-H insertion product Int-c2. Accordingly, the addition of the carbene at the ortho site of phenol gives rise to an intermediate Int*-c2 with a higher energy (8.2 kcal mol⁻¹ vs. Int-c2's 3.1 kcal mol^{-1}) via the transition state **TS*-c1** with 19.3 kcal mol^{-1} . It is not surprising to find that the meta-position of phenol is the most unfavourable site for addition with the gold carbene, which needs an activation energy barrier up to 23.6 kcal mol⁻¹. Such high activation barriers of TS*-c1 and TS**-c1 prevent the carbenes from adding at the ortho- and meta-positions of phenol. The current DFT results are consistent with our experimental observations that the addition at the para-sites of substituted phenols was the predominant even with bulky steric hindrance (i.e. 3,5-dimethyl phenol).5a According to the previous calculations,10 the intermediate Int-c2 may be

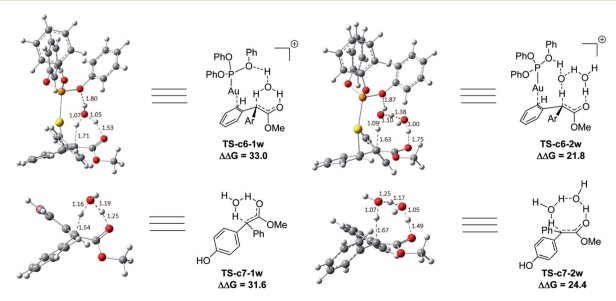


Fig. 3 The structures of TS-c6-1w, TS-c6-2w, TS-c7-1w and TS-c7-2w in the pathways of C-H functionalization of phenol catalysed by $(PhO)_3PAuSbF_6$. The values of the free energy barriers $\Delta\Delta G$ are in kcal mol^{-1} .

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Scheme 1 Control experiments.

isomerized to the enolate via the mechanism of [1,3]-metal migration and yield a stable intermediate. However, it is found that such a stable intermediate hardly exists after the [1,3]-Au migration in Int-c2, since a kind of cyclopropanation product of the Büchner reaction is obtained, as shown by Int*-c4.18 Meanwhile, this process of [1,3]-Au migration seems to be energetically unfavorable, with the endothermicity of 9.1 kcal mol⁻¹ to give **Int*-c4**. Additionally, in order to examine whether the commonly accepted mechanism of direct [1,2]-H shift is feasible, we located the transition state TS*-c3 with the corresponding structure displayed in Fig. 2. Our calculated results imply that TS*-c3 has a considerable barrier of 30.1 kcal mol⁻¹, which could definitely rule out the mechanism of direct [1,2]-H shift in Int-c2. During the course of optimization, we happened to hypothesize that the carbonyl oxygen atom may intend to directly abstract the proton at the original para-site of phenol *via* the five-membered transition state **TS-c3** in Fig. 2, to generate **Int-c4**. This process is quite facile according to the current calculations, merely with a small barrier of 8.6 kcal mol⁻¹. Also, **Int-c4** is more stable than **Int*-c4** by 17.6 kcal mol⁻¹. Subsequently, the intermediate **Int-c4** undergoes an unexpected isomerization to the enol **Int-c5** through [1,3]-migration of the metal complex (PhO)₃PAu⁺ to the phenyl ring rather than to **Int*-c5**.

Two possible pathways might exist between the enol Int-c5 and the final product Pro-c8. One is the gold-associated pathway, where the proton transfers through TS-c6-1w or TSc6-2w to yield the intermediate Int-c7. Then, the metal complex (PhO)₃PAu⁺ dissociates from **Int-c7** to yield **Pro-c8**. The other is the gold-free pathway, where the catalyst (PhO)₃PAu⁺ first dissociates from Int-c5 to yield the gold-free Int-c6 and then proton transfer occurs via TS-c7-1w or TS-c7-2w to yield Pro-c8. As far as the two pathways are concerned, the key question is: how does the proton transfer from the hydroxyl to the final carbon in the enol? In this work, we propose that the water in the solvent plays a crucial role in undertaking the remote proton shuttle. Based on this proposal, we located the one-water assisted transition states TS-c6-1w and TS-c7-1w, as well as the two-water assisted transition states TS-c6-2w and TS-c7-2w along the reaction pathways, with the corresponding activation energy barriers of the elementary steps displayed in Fig. 3. The DFT results indicate that TS-c6-1w and TS-c7-1w involving one water molecule has higher energy barriers than two-water assisted TS-c6-2w and TS-c7-2w. Fig. 3 shows that the carboxyl group in TS-c6-1w and TS-c7-1w has to distort from the planar structures to form hydrogen bonds with H2O, but they still remain planar in TS-c6-2w and TS-c7-2w. The structural distortion in TS-c6-1w and TS-c7-1w might be the reason why they are in high energy states. Interestingly, it is found that the oxygen atoms in the ligand (PhO)₃P form hydrogen bonding

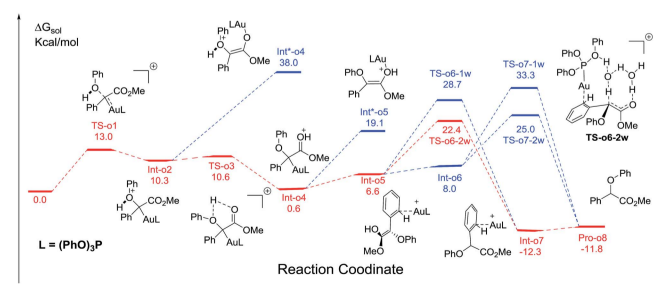


Fig. 4 The calculated free energy profiles ΔG and the corresponding structures of the intermediates and transition states along different possible O–H insertion pathways catalyzed by (PhO)₃PAuSbF₆. The reasonable pathway is shown in red and other possible pathways are in blue. All values of free energies are relative to the gold carbene and phenol reactants and in the units of kcal mol⁻¹.

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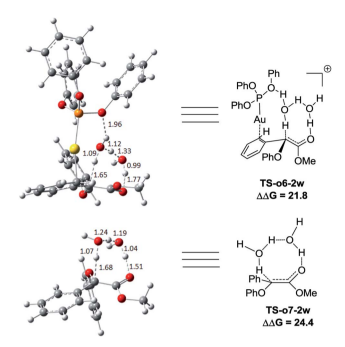


Fig. 5 The structures of TS-o6-2w and TS-o7-2w in the proposed O-H functionalization of phenols catalysed by (PhO)₃PAuSbF₆. The distances are in the units of angstroms and the values of free energy barriers $\Delta \Delta G$ are in kcal mol^{-1}

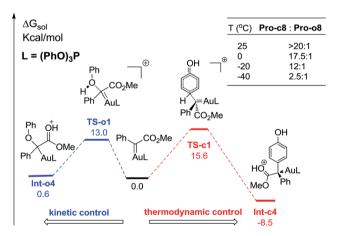


Fig. 6 The reaction temperature on the chemoselectivity and the competitive pathways for the formation of two key intermediates. All values of free energies are relative to the gold carbene and phenol reactants and in the units of kcal mol⁻¹

Table 2 The calculated free energetic spans $\Delta\Delta G$ of hydrogen transfer through transition states (TSs) assisted by one water molecule and two water molecules in Fig. 6. The values of $\Delta\Delta G$ are in the units of $kcal\ mol^{-1}$

TSs	TS-c14-2w	TS-c15-2w	TS-o14-2w	TS-o15-2w
$\Delta\Delta G$	24.1	22.0	20.5	18.0

interactions with the shuttle water molecules in TS-c6-2w, while this effect does not exist in the other located transition state TS*-c6-2w (Fig. S2 in ESI†). The calculated distance between the

oxygen and hydrogen atoms in TS-c6-2w of Fig. 3 is 1.87 Å and the energy of the transition state is 2.2 kcal mol⁻¹ lower than that of TS*-c6-2w, indicating a possible stabilization effect of the metal complex (PhO)₃PAu⁺ for the water chain in **TS-c6-2w**.

Control experiments were carried out in our lab (Scheme 1) to demonstrate the proposed mechanisms of gold carbene insertion into the C-H bond of phenol. When 1 equivalent of D₂O was added into the reaction system, product **Pro-o8** was deuterated with the percentage of 22% (eqn (1)). The relatively low deuterium ratio might be attributed to the existence of trace amount of H2O19 that could also undertake the function of proton transfer. Additionally, it is well known that the hydrogen in the OH group of phenol could readily exchange with the deuterium in D2O, which leads to an increased H2O or DOH amount in solution. If a great number of H2O molecules participate in the process of the proton shuttle, the ratio of deuterated product Pro-o8 will definitely decrease. We also considered the possibility whether the phenols can serve as the proton shuttle. However, the calculated energy barrier between the transition state structure **TS-c6-2p** (ESI†) and **Int-c5** for [1,3]-H shift is highly up to 36.1 kcal mol⁻¹, which implies that it hardly occurs. Eqn (2) manifests that the direct proton exchange of Pro-08 with D₂O is impossible under the reaction conditions. More importantly, eqn (3) demonstrates that the reaction of 1 and deuterated phenol 2-5d gives rise to the product Pro-o8 without deuterium, which supports that direct [1,2]-H shift⁹ (TS*-c3) does indeed not occur in this case. The control experiments demonstrate that this kind of C-H bond insertion reactions governed by a gold carbene follows a novel mechanism involving a water assisting process, rather than previous commonly proposed direct [1,2]-H shift.5,8a,8c,9

Proposed pathway of O-H insertion catalysed by (PhO)₃PAuSbF₆

We next turn to address the question why the O-H insertion of phenol does not occur with (PhO)₃PAuSbF₆ as the catalyst. In Fig. 4, the calculated results show that oxonium ylide20 Int-o2 formed through TS-o1 with a modest barrier. In the case of O-H bond insertion, a possible migration of complex LAu⁺ to the carbonyl group in Int-o2 is also taken into account, but the resulted structure Int*-04 is quite high in energy. The oxonium ylide Int-o2 tends to isomerize to the more stable Int-o4 via TSo3. The migration of the gold catalyst to the phenyl group in Into4 leads to the isomer Int-o5. It should be noted that Int*-o5 is quite energetically unstable, with the gold complex attached to the oxygen atom in the hydroxyl group. Similarly to the mechanism of C-H insertion in Fig. 1, the rearrangements of Int-o5 to Pro-o8 pass through the gold-associated or gold-free pathways via the specific water-assisted transition states. We also locate the two-water assisted transition states TS-o6-2w for the gold-associated pathway and TS-07-2w for the gold-free pathway, respectively (Fig. 5). The activation barrier of TS-o6-2w (15.8 kcal mol^{-1}) is relatively lower that of **TS-o7-2w** (18.4 kcal mol⁻¹), indicating that the transformation of **Int-o5** to **Pro-o8** favours the gold-associated pathway via TS-o6-2w rather than TS-07-2w.

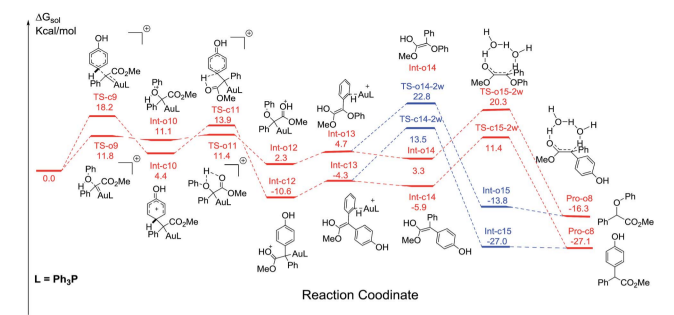


Fig. 7 The competitive pathways of C-H and O-H insertion catalyzed by $Ph_3PAuSbF_6$. The two reasonable pathways are shown in red and the other possible pathways in blue. All values of free energies are relative to the gold carbene and phenol reactants and in the units of kcal mol⁻¹.

Rationale of the chemoselectivity

Comparing these two competitive reaction pathways in Fig. 1 and 4, it is revealed that both rearrangements from Int-c5 or Into5 to the final products are the rate-determining steps in the overall pathways, with almost the same total barriers of 21.8 kcal mol⁻¹. Nevertheless, the DFT calculations reveal that the thermal stability of the intermediate of C-H insertion remarkably differs from that of O-H insertion. For example, the intermediate **Int-c4** is more stable than **Int-o4** by 9.1 kcal mol⁻¹, with an exothermic energy of 8.5 kcal mol⁻¹. As for the final products, the energy of **Pro-c8** is 10.8 kcal mol⁻¹ lower than that of the O-H insertion product Pro-o8. Besides, the endothermicity of TS-c6-2w is 13.3 kcal mol⁻¹, whereas the corresponding energy of **TS-o6-2w** is up to 22.4 kcal mol⁻¹, relative to the sum of the reactants. Such a large endothermicity of TS-06-2w means that it is more difficult to pass through it to reach the product though collision in solution. The calculated reaction pathways in Fig. 1 and 4 reveal that the stability of intermediates plays an important role in chemoselectivity and accounts for the unique chemoselectivity toward C(sp²)-H insertion.

In order to further understand the chemoselectivity, the reactions of phenol with α -phenyldiazoester catalysed by $(PhO)_3PAuSbF_6$ at different temperatures were also carried out. The ratios of **Pro-c8**: **Pro-o8** decreased from >20:1 to 2.5:1 along with lowering the reaction temperature from rt to $-40\,^{\circ}$ C (Fig. 6; for details, see ESI†). These results indicated that the C–H functionalization process was more favored at high temperature than at low temperature. According to the free energy showed in Fig. 1 and 4, **Int-c4** and **Int-o4** should be the key intermediates for C–H functionalization and O–H insertion. As shown in Fig. 6, **Int-c4** was more stable via a higher barrier whilst **Int-o4** was more unstable via a lower barrier. That is, the

formation of **Int-c4** was the process of thermodynamic control, leading to C–H functionalization. In contrast, the generation of **Int-o4** underwent kinetic control, leading to O–H insertion. The control experiments at different temperatures and DFT calculations are in excellent agreement.

Two competitive pathways catalyzed by Ph₃PAuSbF₆

Additionally, the mechanisms of C-H and O-H insertions of phenol catalyzed by Ph₃PAuSbF₆ are also shown in Fig. 7, with the corresponding barriers listed in Table 2. On the one hand, compared to the corresponding pathways in Fig. 1 and 4, we found that the ligand Ph₃P has a remarkable electronic effect on the change of dynamics barriers of electrophilic addition to intermediates Int-o12 and Int-c12. For C-H insertion, the addition barrier via **TS-c9** was enhanced to be 18.2 kcal mol^{-1} , 2.6 kcal mol⁻¹ higher than that of **TS-c1**. In contrast, the barrier of O-H insertion via TS-09 decreases from 13.0 to 11.8 kcal mol⁻¹. It is no doubt that the change of the barriers of addition is disadvantageous to C-H insertion but advantageous to O-H insertion. On the other hand, the calculated barrier of hydrogen transfer via **TS-o15-2w** in O-H insertion is 18.0 kcal mol⁻¹, relatively lower than the barrier of 22.0 kcal mol⁻¹ for C-H insertion. Therefore, the reaction of C-H insertion shown in Fig. 7 is significantly slowed down in comparison to the case when it is catalysed by (PhO)₃PAuSbF₆ in Fig. 1. Product of C-H insertion is still more stable than the counterpart of O-H insertion in terms of thermodynamics according to the current DFT calculations. Considering both the dynamic and thermodynamic effects, the calculated pathways (Fig. 7) could account for the low chemoselectivity for Pro-c8 and Pro-o8 catalysed by Ph₃PAuSbF₆ that has been observed in our experiments (Table 1, entry 7).

Conclusions

In summary, the origins of the unique chemoselectivity of C-H bond functionalization of phenol with diazo compounds catalyzed by (PhO)₃PAuSbF₆ are due to the high thermodynamic stability of intermediates and products, as well as the modest energy barrier of the rate-determining step. It is proposed for the first time that the reactions of gold carbenes with phenol pass through the specific pathways involving the formation of stable enols. The hydrogen transfers in the rearrangements from enol to final products follow a novel reaction pathway with two water molecules serving as a proton shuttle rather than the previous commonly proposed direct [1,2]-H shift. The DFT calculations also reveal that the oxygen atoms in the ligand (PhO)₃P play an important role in stabilizing the transition states of hydrogen transfer. The observed ligand effects of metal complexes on the reaction pathways are well rationalized based on the results of the present DFT calculations. The novel mechanistic insights delineated above will refine the model of gold catalyst in carbene chemistry and contribute to the future development of this field as well as the rational designing in chemo- and site-selective organic transformation.

Acknowledgements

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