Catalyst-dependent selectivity in sulfonium ylide cycloisomerization reactions†

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Divergent catalysis is an emerging field whereby access to structurally diverse compounds from a common precursor is achieved through controlled reaction pathways. Herein we present an unusual example of π-acid catalyst dependent selectivity in the cycloisomerization of alkene sulfonium ylides. Computational mechanistic studies revealed how the ability of palladium to cycle through oxidation states largely controls the selectivity.

Introduction

Divergent catalysis provides a quick access to structurally different compounds from a common precursor through controlled reaction pathways. It is considered a highly attractive tool for the discovery of functional materials and novel drugs. However, the conceptual development of such processes is still rare in the literature.

A distinctive reactivity feature of Au(i), Pt(ii) and Pd(ii) complexes is their propensity towards electrophilic activation of unsaturated C–C bonds. Although these metals are intrinsically different from one another, it is not unusual that their catalysis of C=C or C≡C activation often leads to similar products (a notable exception being ene-yne cyclisations). Nonetheless, Au(i) catalysts only have one free coordination site, whereas Pd(ii) and Pt(ii) salts are square planar and can have two coordination sites. Moreover, Pd(ii) and Pt(ii) salts often behave as bifunctional Lewis acids and even form σ-complexes with hetero-atoms. Last but not least, Pd and Pt complexes can undergo oxidation state changes far more readily than gold.

Sulfonium ylides have long occupied a privileged position in organic synthesis. In recent years, stabilised sulfonium ylides have seen their reactivity greatly expanded through transition-metal-mediated activation. We and others have shown that electrophilic activation of alkene- and alkyne-tethered sulfur ylides by gold catalysts affords cyclopropanes and furans. Nonetheless, cyclopropanation is still dominated by the chemistry of diazo compounds using transition metal catalysis. The resulting metalcarbenes are mild and selective cyclopropanating agents. As a notable exception, the cyclopropanation of olefins containing a sulfur atom does not lead to the desired cyclopropane product. Instead, the sulfonium ylide is formed with complete chemoselectivity (Scheme 1). In face of this limitation of metal carbenoid strategies, we became interested in the possibility of achieving cycloisomerization of the corresponding sulfonium ylides through π-acid catalysis. Herein we report on an intriguing reactivity switch that allows the divergent catalytic synthesis of cyclopropanes or dihydrafurans at will from the same precursor as well as a mechanistic study of these transformations.

Results and discussion

Based on our prior success using gold catalysis for the activation of sulfonium ylides, we started our investigations with sulfonium ylide 1a (Scheme 1) and screened several gold catalysts. Using Echavarren’s catalyst in toluene at 75 °C, the reaction proceeded smoothly to the desired cyclopropane in excellent 95% yield (for other gold catalysts, see the ESI† for details).

Different π-acid catalysts were screened in attempts to optimize the reaction further. While AuCl3 and Hg(OTf)2 only gave low yields of the cyclopropane, Pd(ii) and Pt(ii) salts surprisingly led to formation of a different product 3a (Scheme 1).
1). Using PdCl₂(MeCN)₂ in 1,4-dioxane enabled full conversion and an isolated yield of 82%.²

From the outset, the catalyst-controlled divergence was intriguing and we performed experiments aimed at elucidating the mechanism of these reactions. Critically, conversion of the cyclopropane product 2a to the dihydrofuran 3a is not possible under Pd(II) catalysis under the reaction conditions, nor in the presence of sulfonium ylide 1b. This suggests that neither the palladium catalyst nor an active species formed in the reaction, is responsible for cyclopropane isomerization to the corresponding dihydrofuran and such an isomerization can therefore be excluded as a possible reaction pathway (Scheme 2a).¹³

Suspicious of the involvement of carbene intermediates,¹⁴,¹⁵ we probed the reaction of diazo compound 4 in the presence of the homoallyl sulfide 6 under our reaction conditions (Scheme 2b). Full consumption of the diazo compound was observed, however, no trace of the dihydrofuran 3a, nor sulfonium ylide 1a was obtained. Similarly, no dihydrofuran was obtained from the combination of (phenylmethyl)sulfonium ylide 5 and homoallyl sulfide 6, suggesting that carbene intermediates are not involved.¹⁶

Selected isotope labelling experiments are depicted in Scheme 3. Substrate 1a-d, bearing a deuterium label on the terminal position of the double bond, was synthesized and exposed to the two sets of conditions. In both cases, the deuterium label ended up at the expected position. Interestingly, the high 20 : 1 E/Z ratio of the substrate resulted in cyclopropane and dihydrofuran products with a similar 3 : 1 trans/cis ratio. This suggests that, while the reaction is not fully stereospecific, both pathways might share a common cyclization step.

The mechanism of both reactions was further studied by DFT calculations¹⁷ (see ESI for a full account of the Computational Details and references), employing a ketoester substrate (methyl substituents on both the ketone and the ester group) as model and replacing JohnPhos by PPhMe₂, for computational expediency.¹⁸ The free energy profiles obtained for cyclopropanation with the Au catalyst is represented in Fig. 1.

As shown, following π-coordination of Au to the C=C double bond, an attack of the ylide C-atom to the internal olefinic carbon forms intermediate B, an Au-alkyl complex with a 5-member tetrahydrothiophenium (SC₄) ring. After a conformational rearrangement (B→B'), nucleophilic attack of the coordinated C-atom to the formerly ylidic carbon results in cleavage of the C–S bond and formation of the cyclopropane product. This second step has a barrier of 22 kcal mol⁻¹ and represents the rate-determining event of the process. The overall reaction A→C is exergonic with ΔＧᵣₑₓ = -13 kcal mol⁻¹.¹⁹

A pathway for dihydrofuran formation under Au catalysis was also calculated.¹¹ However, comparison of the two mechanisms shows a clear preference for the cyclopropanation reaction, due to a 21 kcal mol⁻¹ lower overall barrier. Aiming to elucidate the differences between Au and Pd, the corresponding mechanisms with Pd catalysis were also addressed by means of DFT calculations. The free energy profiles obtained are represented in Fig. 2 and 3.

After substitution of two acetonitrile ligands in [PdCl₂(MeCN)₂] by the chelating substrate, the first step of the mechanism parallels that found for Au catalysis – formation of the SC₄ ring after nucleophilic attack of the ylide C-atom to the
internal olefinic carbon. The associated barrier is significant (25 kcal mol$^{-1}$, relative to the separated reactants) but the process is exergonic, with F less stable than E by 5 kcal mol$^{-1}$. In the following step there is C–S bond breaking in the substrate with concomitant coordination of the ylide C-atom, forming a metallacyclobutane G. That formally corresponds to an oxidative addition process, with a change from Pd$^{II}$ (in F) to Pd$^{IV}$ (in G). In the corresponding transition state, TS$_{FG}$, both S–C bond breaking as well as Pd–C coordination are rather incipient. In fact, the S–C bond distance in TS$_{FG}$, is only 0.34 Å longer than in F, and the Pd–C separation is 0.46 Å longer than it will become in intermediate G. The process involves a significant rearrangement of the ligand. In F, there is a square planar geometry around the metal whereas in G there is a square pyramidal geometry with the two chlorides, the O-atom and the former ylide C-atom occupying the four base positions and the olefin terminal C-atom in the apical position. The barrier for this second step is modest, TS$_{FG}$ being 16 kcal mol$^{-1}$ less stable than the initial reactants, and it corresponds to an exergonic process with $\Delta G = -7$ kcal mol$^{-1}$.

The free energy profile for the formation of both the cyclopropane and dihydrofuran products from common Pd$^{IV}$ intermediate G’ (a conformer of G) is represented in Fig. 3. Pd-catalysed cyclopropanation (Fig. 3, left) occurs in a single step, from G’ to H, in a formal reductive elimination process that brings the metal back to Pd$^{II}$. In H, the product is coordinated by both a C–H bond (σ-complex) and the ketone O-atom. In the corresponding transition state, TS$_{GH}$, the formation of the new C–C bond is well advanced, with a distance of 1.94 Å, only 0.41 Å longer than the final one, in H.

Formation of the dihydrofuran (Fig. 3, right) starts with attack of the S-atom onto the coordinated carbon and formation of a stabilized Pd enolate (G’ $\rightarrow$I). This is formally a reductive step, as the metal changes from Pd$^{IV}$ to Pd$^{II}$ – regenerating the characteristic square planar environment. The associated barrier is negligible (1 kcal mol$^{-1}$) and the step is significantly exergonic. Following conformational change at S (I $\rightarrow$I’), attack from the ketone oxygen with concomitant C–S bond breaking, from I to J, yields the dihydrofuran product (as a π-complex). Both TS$_{GI}$ and TS$_{IJ}$ are 3 kcal mol$^{-1}$ more stable than TS$_{GH}$. This indicates that dihydrofuran formation is slightly more favourable than cyclopropanation, for the Pd catalyst. Overall, the free energy balance indicates an exergonic process ($\Delta G_{R} = -16$ kcal mol$^{-1}$).

Taken together, the mechanisms found for the Pd catalyst involve oxidation of the metal from Pd$^{II}$ to Pd$^{IV}$ in an intermediate step. This is accomplished with C–S bond breaking prior to C–C coupling and formation of a metallacyclobutane (in intermediates G/G’). As a result, the sulfur atom is freed for formation of a tetrahydrothiophenium intermediate, effectively providing a natural leaving group for the final attack from the ketone oxygen that produces the dihydrofuran. On the other hand, gold oxidation from Au$^{I}$ to Au$^{III}$ is much less favourable and therefore the favourable route becomes cyclopropanation.

At this juncture, we decided to investigate the scope of both transformations. As shown in Scheme 4, the cyclopropanation reaction is quite general and works well for a wide variety of substrates, tolerating electron-rich or electron-poor arene moieties or a hindered ketone. An electronically diverse range of substituents ($\mathbf{R}$) was tolerated on sulfur. Even with an extra carbon in the tether, cyclization could still be achieved, albeit only in 25% yield. Unfortunately, di- and trisubstituted alkenes did not lead to the desired product in synthetically useful yields.

Interestingly, exquisite selectivity was observed in the formation of cyclopropane 2f (Scheme 4). Noteworthy, the corresponding diphenylsulfonium ylide was an excellent substrate...
for cyclopropanation of tethered allyl esters in our prior work. The observed reactivity could be explained by the difference in calculated activation energy (17.8 vs. 6 kcal mol\(^{-1}\)). This is also qualitatively apparent from a cursory analysis of reaction temperature. Indeed, while the reactions in the present report are performed at 75 °C, intramolecular cyclopropanation of the allyl ester requires 100 °C.

Next, we surveyed the palladium(II)-catalysed cyclization leading to dihydrofurans (Scheme 5). A qualitative difference in reactivity between electron-poor and electron-rich substrates was observed, with electron-donating substituents requiring a decrease in temperature to avoid side product formation. Selectivity analogous to that of the cyclopropanation process was observed for 3f, for which the substrate carries two double bonds susceptible of activation. Malonate derivatives (cf. 3g) gave a complex mixture of products with no dihydrofuran detected. Interestingly, bicyclic product 3o was formed in low yield, while cyclopropanation of the same substrate didn’t take place.

Conclusions

In summary, we have presented an unusual example of \(\pi\)-acid catalyst-dependent selectivity in the cycloisomerization of alkene-tethered sulfonium ylides. While gold catalysis leads to cyclopropane products, palladium(II) leads to a unique dihydrofuran synthesis. Gold effectively enables the formal cyclopropanation of sulfur-containing olefins, a reaction currently beyond the realm of diazo chemistry; palladium unlocks an oxidative addition/reductive elimination catalytic cycle rarely seen in sulfonium ylide chemistry. Computational studies clarified the mechanism and shed light on the factors responsible for the selectivity.

Conflicts of interest

There are no conflicts to declare.

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12 PtCl₂/PtCl₄ were selective for dihydrofuran formation in lower yields. Rh, Ir, Cu or Ag displayed virtually no reactivity. See ESI† for details.

13 Classical Lewis acids (e.g. FeCl₃ or Sc(OTf)₃) could convert 2a into dihydrofuran 3a in moderate yield. See ESI† for details.

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16 The simplicity of the model used in the calculations prevents the discussion of the reaction stereochemistry.

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18 Transition states for gold oxidation, i.e., the equivalents of TS_{SG} for the Au system, have associated barriers above 40 kcal mol⁻¹.