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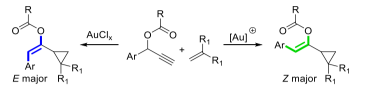


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Gold-catalyzed addition of propargyl acetates to olefins via *O*-acyl migration/cyclopropanation sequence: insight into the diastereoselective formation of the alkene

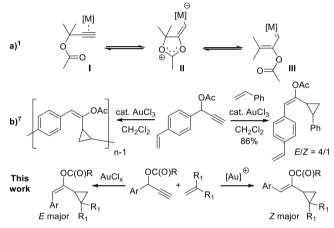
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Abstract. This article discloses a study on the well-known addition of propargyl acetates to olefins via an O-acyl migration/cyclopropanation sequence. Herein, we show that the stereochemical outcome of the olefin is strongly dependent on the gold-catalyst and reaction parameters (concentration, temperature, alkene partner equivalents); *E* and the *Z* isomers can be selectively formed by judicious choice of reaction conditions.

The electrophilic activation of propargyl acetates I by metals to generate metal carbenes III via 1.2-O-acvl migration constitutes a powerful and versatile tool for various valuable synthetic transformations generating molecular diversity (Scheme 1a).¹ The intramolecular transformation was first discovered in 1976 by Ohloff (ZnCl₂).² followed by Rautenstrauch in 1984 (palladium and platinum),³ and re-evidenced by our group in 2002 with PtCl₂.^{1j} The intermolecular trapping of carbene III by an alkene was first reported by Ohe and Uemura with a ruthenium(II) catalyst.⁴ Subsequently, this reaction has witnessed intense developments due to the reactivity of π -acidic metals, mainly gold catalysts,¹ with applications in total syntheses of natural products⁵ and in asymmetric catalysis.⁶ In 2014, we reported the polymerization of bifunctional monomers via gold-catalyzed polycyclopropanation.7 Using AuCl₃ as a catalyst, we obtained conjugated polymers integrating a cyclopropyl/vinyl/phenyl repeating unit. Notably, this transformation provided the E stereoisomer as the major product (Scheme 1b), which marked a sharp contrast with previously reported intermolecular reactions featuring only Z-isomer adducts, generally obtained by using cationic gold(I) catalysts.^{1,6a,8} This piqued our interest since, contrary to cyclopropanation diastereoselectivity that has been well investigated,⁸ alkene stereochemical outcome has rarely been discussed. Unsurprisingly, the only reported E-selective synthesis arose from intramolecular transformations yielding cyclic olefins.¹ To the best of our knowledge, only a few other intermolecular examples provide the E-isomer, yet always as the minor product.9,1g Moreover, Z-selectivity has also been observed when using PtCl₂,¹⁰ cationic rhodium(I)¹¹ or [RuCl₂(CO)₃]₂¹² catalysts.¹³ This intriguing difference between our experimental observations and the results published in the literature led us to study the Z/Ediastereoselectivity of this reaction, and its dependence on reaction conditions.14,8a



Scheme 1: Gold-Catalyzed Migration of Propargyl Acetate

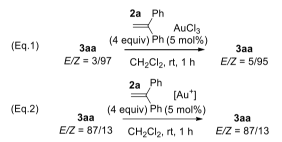
In our study, we selected propargyl acetate 1a and 1,1-disubstituted alkene 2a, as the carbene acceptor, devoid of a prochiral face to suppress complex diastereomeric mixtures. We first screened the gold catalyst. Chlorinated gold complexes (Table 1, entries 1-5) demonstrated a preference for formation of the E-isomer (E/Z ratios ranging from 73/27 to 83/17). Using fully cationic and halide-free gold complexes (entries 6-9) induced an inversion of the stereoselectivity, preferentially furnishing the Zisomer as the major product. When di-cationic [Au(pic)][SbF₆]₂ was used, only the Z-isomer was observed, albeit with very low conversion (entry 6). Using [PPh₃Au][SbF₆] improved conversion, accompanied by the formation of numerous side-products, giving an E/Z ratio of 14/86. When JohnPhos was used as the ligand (entries 8-9), yields increased while the selectivity slightly decreased. Notably, the appearance of the reaction mixture was very different in both cases. When chlorinated gold complexes were used, the color of the reaction mixture turned to purple-toblack aspect only a few minutes after adding the alkyne partner (entries 1 to 5), whereas it remains light-yellow with fully cationic gold (entries 6 to 9). Thus, the in-situ formation of gold clusters or nanoparticles, that are known to catalyze such cycloisomerizations,15 cannot be excluded in the case of chlorinated gold complexes that are known to be unstable in solution.16

Table 1: Catalyst screening

Ph	OAc + Ph Ph Ph equiv) 2a (4 equiv)	$\frac{\text{cat. (5 mol\%)}}{\text{CH}_2\text{Cl}_2 (0.1\text{M}), 1\text{h, rt}} \overset{\text{Ph}^2}{\rightarrow}$		Ph Ph Ph
	cat.	Yield (conversion)	Е	Ζ
1 ^[a]	AuCl ₃	80%	82	18
2 ^[a]	AuCl	74% (82%)	80	20
3 ^[a]	NaAuCl₄	84%	73	27
4 ^[a]	Au(pic)Cl ₂	94%	81	19
5 ^[a]	AupicCl ₂ +AgSbF ₆	65% (80%)	83	17
6	AupicCl ₂ +2AgSbF ₆	(3%)	0	100
7	PPh ₃ AuCl + AgSbF ₆	34% (73%)	14	86
8	[Au]+	89%	32	68
9	[Au]+AgSbF ₆	70% (90%)	31	69

[a]: purple-to-black aspect of the reaction mixture. pic = 2-pyridinecarboxylato; [Au]⁺: [o-Ph-C₆H₄-P(*t*Bu)₂Au(MeCN)][SbF₆]; [Au]: o-Ph-C₆H₄-P(*t*Bu)₂AuCl; cat.: catalyst; Conditions: **1a** (75 mg, 0.43 mmol, 1 equiv), **2a** (0.3 mL, 1.72 mmol, 4 equiv), cat. (22 µmol, 5 mol%), CH₂Cl₂ (4.3 mL), 1h, rt.

When the reaction is complete, the dr is constant and nearly no E-to-Z or Z-to-E isomerization was observed, even by resubjecting products to reaction conditions (Scheme 2, and SI, section 2.2.3). In both cases, **3aa** was fully recovered.



Scheme 2: Configurational stability of cycloisomerization adducts; [Au]⁺: [*o*-Ph-C₆H₄-P(*t*Bu)₂Au(MeCN)][SbF₆]; Conditions: (Eq.1): **3aa** (80 mg, 0.23 mmol, 1 equiv, E/Z = 3/97), **2a** (0.16 mL, 0.90 mmol, 4 equiv), AuCl₃ (3.4 mg, 11 µmol, 5 mol%), CH₂Cl₂ (4.5 mL), 1h, rt. (Eq.2): **3aa** (130 mg, 0.37 mmol, 1 equiv, E/Z = 87/13), **2a** (0.26 mL, 1.47 mmol, 4 equiv), [Au]⁺ (14 mg, 18 µmol, 5 mol%), CH₂Cl₂ (4.5 mL), 1h, rt.

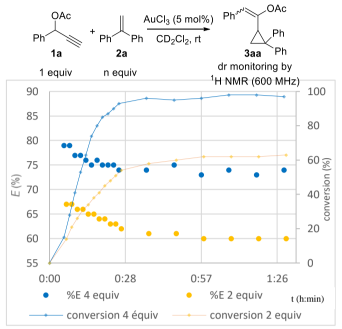
The influence of alkene partner stoichiometry was then evaluated. Interestingly, increasing equivalents of alkene resulted in greater the proportions of *E* diastereoisomer, independent of the catalyst used (Table 2). Changing alkene stoichiometry from 1 to 4 equivalents, increased E/Z dr from 50/50 to 82/18 when AuCl₃ was used (entries 1-3), and from 9/91 to 32/68 using a cationic gold catalyst (entries 4-6).

Table 2: Alkene stoichiometry

	OAc Ph 1a (1 equiv)	+ Pl 2a	$h \qquad Ph \qquad \frac{\text{cat. (5 mol%)}}{\text{CH}_2\text{Cl}_2 (0.1\text{M}), 1\text{h},}$ (n equiv)	►	Ph ³ OAc Ph 3aa Ph
	n	cat.	Yield (conversion)	Е	Ζ
1	1	AuCl₃	(55%)	50	50
2	2	AuCl₃	51% (59%)	71	29
3	4	AuCl ₃	94%	82	18
4	1	[Au]+	68%	9	91
5	2	[Au]+	75%	19	81
6	4	[Au]⁺	89%	32	68

[Au]⁺: [*o*-Ph-C₆H₄-P(*t*Bu)₂Au(MeCN)][SbF₆]; Conditions: **1a** (75 mg, 0.43 mmol, 1 equiv), **2a** (n equiv), cat. (22 μmol, 5 mol%), CH₂Cl₂ (4.3 mL), 1h, rt.

Monitoring the dr during the reaction confirmed these findings as the percentage of E diastereoisomer slowly decreases with increasing conversion, which is consistent with the decrease of alkene equivalents (Figure 1).



Conditions: **1a** (15 mg, 86 µmol, 1 equiv), **2a** (30μ L, 0.17 mmol, 2 equiv (yellow) or 61µL, 0.34 mmol, 4 equiv (blue)), AuCl₃ (1.3 mg, 4.3 µmol, 5 mol%), CD₂Cl₂ (1mL), 300 K).

Figure 1: ¹H NMR monitoring of the diastereoisomeric ratio during the reaction (600 MHz, CD_2Cl_2 , 293 K)

The effect of reaction concentration was also examined (Table 3). Higher concentrations favor formation of the *E* diastereoisomer, irrespective of the catalyst used. Using AuCl₃, the best drs were obtained in concentrated reaction mixtures (entries 1-2, *c* 0.2M), while with cationic gold dilute reaction mixtures resulted in higher diastereoselectivities (entries 9-10, *c* 0.05M). Toluene was found to be the solvent of choice for highest dr, whichever catalyst was used (entries 2 and 10, and SI, section 2.2.4).¹⁷

Table 3: Concentration

OAc		cat. (5 mol%)	Ph
Ph ·	Ph Ph	solvent (c), 1h, rt	∠Ph
1a (1 equiv)	2a (4 equiv)		3aa [`] Ph

	cat.	solvent	С	Yield (conv)	Е	Ζ
1	AuCl₃	CH ₂ Cl ₂	0.2M	88%	88	12
2	AuCl₃	toluene	0.2M	60% (80%)	91	9
3	AuCl₃	CH_2CI_2	0.1M	94%	82	18
4	AuCl₃	toluene	0.1M	95%	87	13
5	AuCl ₃	CH_2CI_2	0.05M	(67%)	76	24
6	[Au]+	CH ₂ Cl ₂	0.2M	74%	42	58
7	[Au] ⁺	CH ₂ Cl ₂	0.1M	89%	32	68
8	[Au] ⁺	toluene	0.1M	51%	11	89
9	[Au]+	CH_2CI_2	0.05M	73%	21	79
10	[Au] ⁺	toluene	0.05M	77%	6	94

$$\label{eq:approx} \begin{split} & [Au]^{\star} : [\textit{o-Ph-C}_6H_4-P(\textit{tBu})_2Au(MeCN)] [SbF_6] ; Conditions: \textbf{1a} (75 mg, 0.43 mmol, 1 equiv), \textbf{2a} (0.3 mL, 1.72 mmol, 4 equiv), cat. (22 µmol, 5 mol%), solvent (\textit{c}), 1h, rt. \end{split}$$

Finally, the effect of temperature was also evaluated (Table 4). Irrespective of the catalyst, low temperature (0°C, entries 3-4 and

8) favors *E*-isomer formation while higher temperature (40°C, entries 1, 5-6) favors the *Z*-isomer. Once again toluene provided the highest dr (entries 3-4; 5-6).

Table 4	Tamanaratura
Table 4:	Temperature

Table 4: Temperature								
	OAc			cat. (5 mol%)	Phun	2.10		
Ph		+ Ph H	Ph solv	ent (0.1M), 1h, T °C	\triangle	Ph		
1a (1 equiv)	2a (4 eq	uiv)		3aa	Ph		
	cat.	solvent	т	Yield (conversion)	Е	Ζ		
1	AuCl ₃	CH_2CI_2	40°C	(50%)	70	30		
2	AuCl₃	CH_2CI_2	rt	94%	82	18		
3	AuCl₃	CH_2CI_2	0°C	(77%)	90	10		
4	AuCl₃	toluene	0°C	(38%)	100	0		
5	[Au]+	toluene	40°C	52% (75%)	4	96		
6	[Au]⁺	CH_2CI_2	40°C	72%	21	79		
7	[Au]+	CH_2CI_2	rt	89%	32	68		
8	[Au]⁺	CH_2Cl_2	0°C	75%	47	53		

$$\label{eq:addition} \begin{split} & [Au]^+: [\textit{o-Ph-C}_6H_4\text{-P}(\textit{t}Bu)_2Au(MeCN)] [SbF_6]; \mbox{ Conditions: } 1a \ (75 \ \mbox{mg}, \ 0.43 \ \mbox{mmol}, \ 1 \ \mbox{equiv}), \ 2a \ (0.3 \ \mbox{mL}, \ 1.72 \ \mbox{mmol}, \ 4 \ \mbox{equiv}), \ cat. \ (22 \ \mbox{µmol}, \ 5 \ \mbox{mol}), \ \mbox{solvent} \ (4.3 \ \mbox{mL}), \ 1h. \end{split}$$

We then screened the scope and limitations (Figure 2). No change in reactivity was observed by changing the acetate function to a benzoate or a pivaloate: the transformation remains stereoselective and predominantly leads to the *E*-isomer when AuCl₃ is used as catalyst, whereas the *Z* product is generated with [Au]⁺ (**3ba** and **3ca**). It was also possible to change the alkene partner to methylenecyclohexane (**3ab** and **3bb**) which displayed similar reaction outcomes, albeit requiring longer reaction times. On the other hand, when 2-methylene-1,3-diphenylpropane was used as the alkene partner, the *Z*-isomer was formed in all cases, an increase in the proportion of *E*-isomer being observed when AuCl₃ was used (**3ac**).

Introduction of a thiophene moiety led to numerous by-products and lower yields ranging from 28% and 48%, however the vinylcyclopropane **3da** was formed in the expected diastereoselective tendency respective of the gold catalyst used.

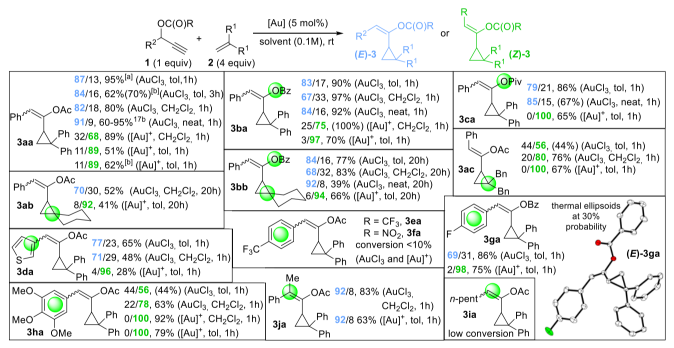
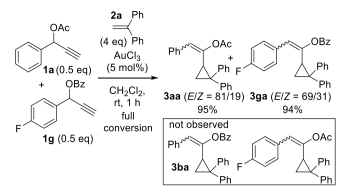


Figure 2. Scope and limitations; [a] 1 0.43 mmol scale, EZ measured on the crude material by ¹H NMR spectroscopy, yield (conversion) (catalyst, solvent, duration); [b] 1 2 mmol scale; tol: toluene, [Au]⁺: [o-Ph-C₆H₄-P(*t*Bu)₂Au(MeCN)][SbF₆].

Variation of substituents on the aryl group was next examined. Electron-withdrawing substituents such as p-nitro or p-trifluoromethyl groups drastically decreased reactivity (3ea and 3fa). However, we succeeded in introducing a p-fluorine moiety that afforded full conversion, albeit with a small increase of the Z-isomer (3ga vs 3ba). The introduction of electron-rich methoxysubstituents led to full conversion, and a selectivity always in favor of the Z-isomer, irrespective of the catalyst used (3ha). Nevertheless, a catalyst effect was observed, [Au]⁺ only led to the Z-isomer, whereas AuCl₃ led to partial formation of the E-product (E/Z = 44/56). Introduction of an alkyl chain at the propargylic position led to very low conversion (3ia).¹⁸ Finally, introduction of a guaternary center led to the same E selectivity in both cases (3ja). This last result could be explained by post-isomerization as suggested by Liu,¹⁹ due to the tetrasubstituted nature of the alkene.

Given that ester scrambling could be responsible for the observed *E*-selectivity during the reaction with $AuCl_{3}$,^{14c} we sought to rule out this possibility. When a 1:1 mixture of propargylic acetate **1a** and benzoate **1g** was subjected to reaction conditions, we only observed the expected products **3aa** and **3ga**, and no structure arising from ester elimination-re-addition (Scheme 3 and SI, section 2.2.6).



 $\label{eq:scheme 3: Ester scrambling evaluation; Conditions: 1a (37.5 mg, 0.215 mmol, 0.5 equiv), 1g (55 mg, 0.215 mmol, 0.5 equiv), 2a (0.3 mL, 1.72 mmol, 4 equiv), AuCl_3 (5.6 mg, 22 \mu mol, 5 mol%), CH_2Cl_2 (4 mL), rt, 1h.$

From these experimental results, we can conclude that the use of a fully cationic Au-catalyst preferentially furnishes the *Z*-product, also favored under thermodynamic conditions (dilute reaction mixture, high temperature) while the use of a chlorinated catalyst such as $AuCl_3$ preferentially leads to the *E* product, generally formed under kinetic conditions (concentrated reaction mixture, low temperature, excess of alkene).

In this work, we focused on the gold-catalyzed 1,2-O-acyl migration/trapping of the intermediate carbene by a double bond leading to vinylcyclopropanes. Interestingly, only a scarce number reports have indicated a relationship between the stereochemistry of the double bond and the nature of the catalyst used: additionally, the formation of the E stereoisomer has rarely been reported in the literature for such an intermolecular transformation. Thus, we studied the factors influencing this and developed conditions allowing to obtain a stereoselective access to both isomers. Notably, we showed that the diastereomeric ratio is a direct function of the nature of the catalyst and substrates, alkene and alkyne equivalents, solvent, concentration and temperature. The stereochemistry of all obtained species is supported by 2D NOESY NMR spectroscopy and X-ray crystallography. The observation of the reaction mixture color, which turned from yellow to purple-to-black in a few minutes when a chlorinated gold catalyst was used suggests the in-situ formation of gold clusters or nanoparticles, which would be responsible of this unusual E-stereochemical outcome. Through a scrambling experiment, we proved that this selectivity did not arise from an ester elimination-re-addition process. DFT calculations are currently underway to rationalize our experimental observations.

Data availability Statement

The data underlying this study are available in the published article and its Supporting Information.

Supporting Information Statement

The Supporting Information is available free of charge at. It includes synthetic procedures, ¹H and ¹³C NMR spectra for all new compounds as well as non-commercially available cycloisomerization substrates and products, X-ray crystallographic analysis of **3ga** (CCDC 2095223), 2D NOESY analysis for diastereoisomers attribution, solvent evaluation, configurational stability study and NMR monitoring of the reaction.

Keywords

cycloisomerization • diastereoselectivity • gold • cyclopropane

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