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Acyl Migration versus Epoxidation in Gold Catalysis: Facile, Switchable, and Atom-Economic Synthesis of Acylindoles and Quinoline Derivatives

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Abstract: We report a switchable synthesis of acylindoles and quinoline derivatives via gold-catalyzed annulations of anthranils and ynamides. a-Imino gold carbenes, generated in situ from anthranils and an N,O-coordinated gold(III) catalyst, undergo electrophilic attack to the aryl π -bond, followed by unexpected and highly selective 1,4- or 1,3-acyl migrations to form 6-acylindoles or 5-acylindoles. With the (2-biphenyl)di-tert-butylphosphine (JohnPhos) ligand, gold(I) carbenes experienced carbene/carbonyl additions to deliver quinoline oxides. Some of these epoxides are valuable substrates for the preparation of 3-hydroxylquinolines, quinolin-3(4H)-ones, and polycyclic compounds via facile in situ rearrangements. The reaction can be efficiently conducted on a gram scale and the obtained products are valuable substrates for preparing other potentially useful compounds. A computational study explained the unexpected selectivities and the dependency of the reaction pathway on the oxidation state and ligands of gold. With gold(III) the barrier for the formation of the strained oxirane ring is too high; whereas with gold(I) this transition state becomes accessible. Furthermore, energetic barriers to migration of the substituents on the intermediate sigma-complexes support the observed substitution pattern in the final product.

Introduction

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Acyl groups are moieties of fundamental importance in organic chemistry. In contrast to classic Friedel–Crafts acylation reactions,^[1] the migration of an acyl group is an atom-economic technique for preparing aryl aldehydes and ketones. In gold catalysis, propargylic esters undergo 1,2- or 1,3-acyloxy migration by nucleophilic attack of the carbonyl oxygen atom at the gold-activated C=C triple bond, affording a gold carbene or an allene, which are both highly reactive for subsequent functionalization, and enable the generation of

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structural complexity.^[2] A gold-triggered 1,3-acyloxy shift and 1,5-acyl shift cascade for preparing δ -diketones was devised using more complex propargylic esters.^[3] Our group developed a gold-catalyzed tandem 1,3-acyloxy/1,5-acyloxy migration for the synthesis of pyrrolidin-2-ones.^[4] 1,2-Acyl migrations for the synthesis of azepino[3,4-*b*]indol-1-ones and dihydro- γ -carbolines were also explored by our group^[5a] and the group of Liu et al.^[5b] To date, although gold-catalyzed 1,2-and 1,5-acyl, as well 1,3- and 1,5-acyloxy, migrations have been studied well, neither a 1,4- nor a 1,3-acyl migration on a benzene ring has been described to the best of our knowledge.

The development of efficient synthetic methods through α -imino gold carbene intermediates has become an important tool for the construction of heterocycles. Gold-promoted nitrene transfer from nucleophilic nitrene equivalents, such as pyridine-based aza-ylides,^[6] azides,^[7] 2*H*-azirines,^[8] isoxazole derivatives,^[9] and sulfilimines^[10] to alkynes, generates α -imino gold carbenes. These can efficiently undergo nucleophilic attack, C-H insertion, or cyclopropanation. However, no carbene/carbonyl additions of these gold carbenes to form epoxides have been reported in the literature. In 2015, the group of Ye et al.^[9a] reported a formal [3+2] annulation of isoxazoles and ynamides for the synthesis of acylpyrroles (R =H, Scheme 1, entry 1). When $R \neq H$, deacylated products are formed. Subsequently, diverse aza-heterocycles have been prepared from isoxazole derivatives by labile N-O bond cleavage.^[9b-t] Anthranils, readily available and useful nitrenetransfer reagents, have attracted considerable interest over the last three years. Either an attack at the nitrogen atom forms α-imino gold carbenes and subsequent C-H insertions gives 7-acylindoles (Scheme 1, entry 2),^[9b] or this versatile reagent undergoes oxygen attack to give quinoline oxides through seven-membered ring intermediates (Scheme 1, entry 3).^[9i] Inspired by previous reports,^[9-11] we herein report

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Scheme 1. Gold-catalyzed annulations of alkynes and isoxazole derivatives by N-O bond cleavage.

the unprecedented site-selective trapping of such α -imino gold carbenes for a divergent synthesis of acylindoles or epoxides via selective acyl shifts and epoxidations (Scheme 1, entry 4).

Results and Discussion

We initially envisioned the synthesis of 2-aminoindole from 7-methylanthranil 1a and N-phenylynamide 2ac through a [3+2] C-H annulation (Scheme 2, path a). However, a quinoline oxide 5b (confirmed by X-ray crystallog-



Scheme 2. The initial design and the unexpected result.

raphy)^[12] was formed instead via a carbene-complex-mediated epoxidation (Scheme 2, path b). An optimization of the reaction conditions was conducted with 7-methylanthranil 1a and N-methylynamide 2a in the presence of various catalysts LAuCl/AgX and solvents (Table 1). At room temperature and in toluene, IPrAuCl/AgNTf2 produced quinoline oxide 5a in 20% yield, along with the unexpected 6-acylindole 3a (13% yield) via an unprecedented 1,4-acyl migration (Table 1, entry 1). When PPh₃ was replaced by (2-biphenyl)di-tert-butylphosphine (JohnPhos), the conversion rates of 1a were higher than 80% and the ratios of 5a/3a increased dramatically (Table 1, entries 2-5). A change of the silver activators (AgX; X = OTs, OTf, BF₄, SbF₆) delivered **5**a/3a with high selectivity (Table 1, entries 6-9), and John-PhosAuCl/AgSbF₆ performed best (Table 1, entry 9). Solvent

Table 1: Optimization of the reaction conditions^[a,b].

	∖_N ^{_Ts}		Ph	.0
	ili –	gold catalyst (5 mol%)		.∕ I Ph
N ^N	Щ	solvent, RT, 12 h		
1-	Ρ'n			
Ia	2a		3a	5a

Entry	Catalyst	Solvent	Conversion 1 a [%]	Yield 3 a [%]	Yield 5 a [%]
1	IPrAuCl/AgNTf ₂	toluene	34	13	20
2	Ph ₃ PAuCl/AgNTf ₂	toluene	81	8	39
3	L1AuCl/AgNTf ₂	toluene	87	11	50
4	L2AuCl/AgNTf ₂	toluene	90	< 5	29
5	L3AuCl/AgNTf ₂	toluene	80	< 5	51
6	L3AuCl/AgOTs	toluene	52	< 5	26
7	L3AuCl/AgOTf	toluene	83	< 5	64
8	L3AuCl/AgBF₄	toluene	84	< 5	42
9	L3AuCl/AgSbF ₆	toluene	87	< 5	75
10	L3AuCl/AgSbF ₆	THF	75	< 5	56
11	L3AuCl/AgSbF ₆	MeCN	97	9	77
12	L3AuCl/AgSbF ₆	CH_2CI_2	>99	< 5	95 (88) ^[c]
13	$NaAuCl_4 \cdot 2H_2O$	toluene	92	69	< 5
14	PicAuCl ₂	toluene	>99	78	9
15	PicAuCl ₂	THF	>99	42	16
16	PicAuCl ₂	MeCN	>99	61	20
17	PicAuCl ₂	DCE	>99	54	< 5
18	PicAuCl ₂	CH_2Cl_2	>99	91 (87) ^[c]	< 5
19	AuBr ₃	CH ₂ Cl ₂	86	60	16

Acetonitrile (MeCN), 1,2-dichloroethane (DCE), tetrahydrofuran (THF), tosylate (OTs), triflate (OTf), bis(trifluoromethylsulfonyl)imide (NTf₂).



[a] Reaction conditions: 1a (0.10 mmol), 2a (0.12 mmol), catalyst (0.005 mmol), solvent (1.0 mL, 0.1 M), RT. [b] The yield and conversion rate were determined by ¹H NMR spectroscopy with 1,3,5-trimethoxylbenzene as the internal standard. [c] Yields of isolated products (0.2 mmol).

screening revealed CH₂Cl₂ as the optimal choice (Table 1, entries 10–12), affording **5a** in 95% NMR yield (88% isolated yield on a 0.2 mmol scale). Interestingly, gold(III) catalysts precipitously reversed the reaction to give the totally unexpected 6-acylindole **3a** as a major product (Table 1, entries 13–18). Best among the gold(III) catalysts and solvents was PicAuCl₂,^[13] giving **3a** in 91% NMR yield (87% isolated yield on a 0.2 mmol scale) in CH₂Cl₂ (Table 1, entry 18). Alkyl-substituted substrates only afforded products of a 1.2-H shift.

The scope for the synthesis of acylindoles was first investigated under the optimized reaction conditions (Table 1, entry 18). As shown in Scheme 3, anthranil **1a** reacted smoothly with a range of *N*-sulfonyl ynamides **2a–d** at room temperature, affording the corresponding 6-acylindoles **3a–d** in 87–94% yields. While *N*-arylynamides **2j–n** gave the desired products **3j–n** in moderate yields, *N*-alkylynamides **2e–i** showed higher efficiency. Yields of 50–99% were achieved with different ynamides **2o–x** bearing a wide range of functionalities (Me, *t*Bu, OEt, F, Cl, Br, or CF₃) at the *ortho-, meta-*, or *para*-positions on the arene rings. Dimethoxy ynamide **2y** also underwent the transformation efficiently.^[12] Compared to phenyl-substituted substrates, alkyl-substituted substrates only afforded the products of a 1,2-*H* shift; for



Scheme 3. Scope for the synthesis of acylindoles.^[a,b] [a] Reaction conditions: **1** (0.20 mmol, 1.0 equiv), **2** (0.24 mmol, 1.2 equiv), PicAuCl₂ (3.9 mg, 5 mol%), CH₂Cl₂ (2 mL, 0.1 m). [b] Yield of isolated product. [c] 4-nitrophenylsufonyl (Ns). [d] Reaction temperature = 50 °C. Key: protecting group (PG), tosly (Ts), mesityl (Ms), nosyl (Ns).

example, the alkylynamide **2ac** gave the unsaturated amidine **8**. The oxazolidinone-derived ynamide **2ab** was well tolerated to produce **3ab** (66%). 7-Methoxyl- and 7-chloroanthranil are appropriate substrates for this reaction. Disubstituted anthranils **1d** and **1e** were converted into the targets **3ae** and **3af** with 99% yields. Unexpectedly, by blocking the 6-position of anthranils, 1,3-acyl migrations proceeded and yielded 5-acylindoles **4a–c**^[12] in 74–99% yields. Acetyl also migrated successfully and 5-acetylindole **4d** was obtained in 42% yield. Notably the tricyclic product **4e** was delivered in 99% yield.

We then explored the scope for the synthesis of quinoline derivatives by using JohnPhosAuCl/AgSbF₆ as the best catalyst (Scheme 4). In the case of ynamides 2, electronically diverse aromatic and aliphatic substituents at the R¹ position, and different protecting groups, were tolerated well to afford 5a, 5c-e, and 6a-e in 56-97% yields. Some of these quinoline oxides 5 underwent facile rearrangements^[14] to 3-hydroxylquinolines 6a-e.^[12] The polycyclic compound 10 was formed by an intramolecular nucleophilic ring opening reaction of epoxide 5d (Scheme 5). Ynamides containing either an electron-withdrawing group (F, CF₃, or Cl) or an electrondonating group (CH₃ or OCH₂CH₃) on the arene ring $(\mathbf{R}^2 = \mathbf{Ar})$ were easily converted to epoxides **5 f**-**j** and rearranged products 6 f-h. The epoxidation reaction was applied to phenyl ynamides (R^2 = phenyl) and heterocyclic ynamide (R^2 = thiophen-3-yl), and the terminal ynamide $(\mathbf{R}^2 = \mathbf{H})$ also reacted well (89% of 3-hydroxylquinoline **6j**). An alkyl ynamide delivered unsaturated amidine 8 in 47% yield instead of the desired quinoline product. Various anthranils were then examined. Anthranil 1j, without any



Scheme 4. Scope for the synthesis of quinoline derivatives.^[a,b] [a] General procedure: **1** (0.24 mmol, 1.2 equiv) and **2** (0.20 mmol, 1.0 equiv) were added to a well-stirred mixture of JohnPhosAuCl (5.3 mg, 5 mol%) and AgSbF₆ (3.4 mg, 5 mol%) in CH₂Cl₂ (2 mL, 0.1 M). [b] Yield of isolated product. [c] Catalyst (10 mol%), reaction temperature = 50 °C. [d] The yield when using 5 mol% PicAuCl₂ is shown in parentheses.

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Scheme 5. Synthesis of polycyclic compound **10** and quinolin-3(4*H*)ones **7.**^[a,b] [a] Reaction conditions: **1** (0.24 mmol, 1.2 equiv), **2** (0.20 mmol, 1.0 equiv), JohnPhosAuCl (5.3 mg, 5 mol%), AgSbF₆ (3.4 mg, 5 mol%), CH₂Cl₂ (2 mL, 0.1 м). [b] Yield of isolated product. [c] Catalyst (10 mol%), reaction temperature = 50 °C.

substituents, gave epoxide 5k in moderate yield. 7-Chloroand 7-bromoanthranils led to the corresponding products 5l(98%) and 5m (80%). 3-Methyl-substituted anthranils 1i and 1g delivered quinolin-3(4*H*)-ones 7a and 7b through epoxidations and subsequent rearrangements (Scheme 5).

Non-polarized alkynes were also tested (Scheme 6). At higher temperatures, both PicAuCl₂ and JohnPhosAuCl/ AgSbF₆ were able to catalyze the annulation of **1a** and phenylacetylene to 3-hydroxylquinoline **6k** in 30% yield in a slow reaction. 10 mol% Methanesulfonic acid (MsOH) was beneficial for substrate conversion and product formation. Diphenylacetylene also afforded **61** in 60% yield.



 $\textit{Scheme 6.}\xspace$ Tests with non-polarized alkynes. $^{[a]}$ [a] Yield of isolated products.

The gram-scale reaction between **1a** and **2a** (using only 2 mol% gold catalyst) gave **3a** in 83% yield (Scheme 7). Even more useful compounds can be prepared from the obtained acylindoles and quinoline oxides. The NaBH₄-mediated reduction of **3a** afforded 6-hydroxymethylindole **3a'** in excellent yield, LiAlH₄ reduced **3a** to 6,7-dimethylindole **3a'''** in 74% yield at higher temperature. The acyl group of **3a** could be converted to an alkynyl group through a Seyferth–Gilbert homologation. Treatment of quinoline oxide **5b** with 3 equivalents of LiAlH₄ provided facile access to tetrahydroquinolin-3-ol **9** via reduction/deamination processes.

To gain more insight into the reaction mechanism,^[15] we conducted two crossover experiments (Scheme 8). With respect to the 1,4-acyl migration, we conducted a crossover



 $\textit{Scheme 7.}\ \mbox{Gram-scale}$ synthesis and important transformations of $3\,a$ and $5\,b.$



Scheme 8. Crossover experiments.

experiment using [D]-1a and 1d reacting with 2a. For the 1,3-acyl migration process, the experiment employed a mixture containing an equimolar amount of [D]-1k and 1f in the reaction with ynamide 2a. A ¹H NMR spectroscopy and mass spectrometry study of the obtained products [D]-3a, 3ae, 4a, and [D]-4e showed no scrambling of deuterium; both of these two acyl migration processes are intramolecular.

Finally, we conducted a detailed computational study of the reaction mechanism at the M06-D3/def2-TZVP//M06/ LANL2DZ,6-31G(d) level in CH_2Cl_2 . We commenced our DFT-based mechanistic study by investigating which coordinating atom of the ligand on complex **i** is involved in the substitution reaction—a question that has not been studied in the many applications of **i** in catalysis. To this end, four trigonal bipyramidal transition structures TS_i-TS_{iv} were calculated for which the potential energies are reported underneath each structure in Figure 1.

Our calculations indicate that \mathbf{TS}_i is lower in energy than the other three transition states, suggesting that the substitution reaction should mostly occur through \mathbf{TS}_i to give intermediate **iii** (Figure 2) from which the remainder of the catalytic reaction proceeds. Once structure **iii** is formed, it is subjected to a nucleophilic attack by substrate **iv** to afford **v** through transition structure \mathbf{TS}_{ii-v} which lies only 6.8 kcal mol⁻¹ above **iii**. The ensuing intermediate then undergoes N–O bond cleavage via transition structure $\mathbf{TS}_{v\cdot vi}$ with an overall activation free energy of 14.4 kcal mol⁻¹, leading to exergonic formation of intermediate **vi**.



Figure 1. The potential energies (kcalmol^-1) for transition structures $TS_{i\text{-}}TS_{i\text{v}}$



Figure 2. Free energy profile for the substitution reaction between i and ii, nucleophilic attack of iv at iii, and finally cleavage of the O–N bond in v forming vi. Gibbs free energies (potential energies) are in kcal mol⁻¹.

Intermediate **vi** is a branching point for two pathways (Figure 3) as follows: 1) nucleophilic attack of the carbonyl group followed by a deauration reaction via **TS**_{vii} (pathway A) and 2) nucleophilic attack of the aryl ring to afford intermediate **ix** (pathway B). Our calculations show that the carbonyl group is a better nucleophile than the aryl ring but formation of **vii** is a dead end, from which the deauration process is very energy consuming with $\Delta G^+ > 30$ kcal mol⁻¹. This low reactivity toward C–C coupling (deauration with formation of a strained three-membered oxirane ring) is ascribed to the high electrophilicity of the Au^{III} center. Thus, although formation of **vii** is feasible during the reaction, it



Figure 3. Free energy profile for competition between two pathways A and B. Gibbs free energies (potential energies) are in $kcalmol^{-1}$.

undergoes a reverse reaction to give vi from which ix is formed via transition structure TS_{vix} .

Attempts to locate transition structure TS_{vii} were unsuccessful. Thus, the energy of this transition structure was estimated by the scan of the potential energy surface, as illustrated in Figure 4.



Figure 4. Scan of the potential energy surface for formation of the C–C bond starting from structure **vii**.

Intermediate **ix** is a branching point for two competitive pathways (Figure 5). The formyl group can migrate along either the anticlockwise (pathway C) or clockwise (pathway D) direction. Pathway C was calculated to be about 4.4 kcal mol⁻¹ higher in energy than pathway D, implying that the formyl migration preferably takes place via pathway D. A potentially competing methyl shift would have a much higher barrier **TS**_{xi-xi'} to provide **xii**'. The preference for pathway D over C can be easily explained by the natural population analysis (NPA) of intermediate **ix** (Figure 5). The analysis shows that the partial charge of the carbon atom bonded to the sp² nitrogen is much more electron deficient than other centers, resulting in the formyl group preferentially migrating along the clockwise direction.



Figure 5. Free energy profile for competition between two pathways C and D. The NPA charges for intermediate **ix** are given in pink. Gibbs free energies (potential energies) are in kcalmol⁻¹.

Once **xii** is formed, it can be consumed in two ways (Figure 6). It can be involved in another formyl migration (pathway E) or in a deprotonation process assisted by a base-like substrate iv (pathway F). In agreement with the experi-

ment, pathway F was determined to be more favorable. Indeed, this pathway is preceded by protodeauration (proton transfer from **xiii** to the sp² nitrogen atom of **xiv** in a barrierless reaction) and finally produces the experimentally observed product. Based on these findings, we proposed a catalytic cycle for the acyl migration processes (Figure 7).

At this juncture, we are interested in understanding why the gold(I) complex mainly gives the epoxide product, which contrasts with the behavior of the gold(III) complex. This difference stems from the fact that the gold(I) complex contains a less electrophilic center compared with the gold(III) complex. As a consequence, the deauration reaction that starts from intermediate xix, and proceeds via C-C coupling, takes place more easily. Our calculations support this claim by showing that the energies of $TS_{xviii-xx}$ and $TS_{xvii-xviii}$ are com-



Figure 6. Free energy profile (kcal mol^-) for competition between two pathways E and F.



Figure 7. A proposed catalytic cycle derived from the DFT calculations.

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Figure 8. Free energy profile for competition between two pathways A' and B' where gold(I) is used as the catalyst. Gibbs free energies (potential energies) are in kcalmol⁻¹.

parable, resulting in both pathways A' and B' being competitive (Figure 8).

Conclusion

In summary, catalyst-controlled, tunable acyl migrations and epoxidations were discovered that occur via α -imino gold carbene intermediates. This efficient, scalable, atom-economic reaction shows high regioselectivity and broad scope, and it operates under mild conditions with easily available substrates. The method opens novel and concise routes to complicated acylindoles, quinoline oxide derivatives, polycyclic compounds, and related compounds—all of which are potentially useful in medicinal chemistry and drug discovery. A computational study fully supported the observed chemoselectivity for gold(III), as well as the selectivity switch when using gold(I) catalysts. The effects explaining these selectivities and selectivity switches will be inspiring for further research in the field.

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Conflict of interest

The authors declare no conflict of interest.

Keywords: acyl migrations · anthranils · epoxidations · gold carbenes · gold catalysis

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