

Received 1st June 2015  
Accepted 17th July 2015DOI: 10.1039/c5sc01950h  
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## Gold-catalyzed formal $[4\pi + 2\pi]$ -cycloadditions of propiolate derivatives with unactivated nitriles<sup>†</sup>

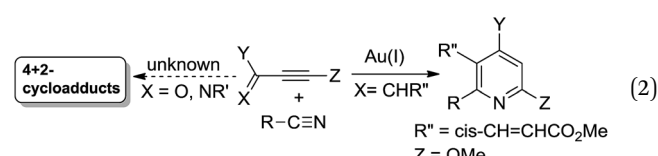
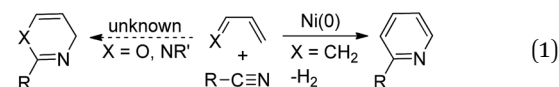
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Gold-catalyzed hetero- $[4\pi + 2\pi]$ -cycloadditions of *tert*-butyl propiolates with unactivated nitriles are described; the resulting 6*H*-1,3-oxazin-6-ones are not easily accessible *via* conventional methods. This new finding enables a one-pot gold-catalyzed synthesis of highly substituted pyridines through sequential gold-catalyzed reactions of *tert*-butyl propiolates with nitriles, and then with electron-deficient alkynes in the same solvent. The utility of these  $[4 + 2]$ -cycloadditions is further expanded with various aldehydes, ketones and 2-phenyloxetane, yielding satisfactory yields of cycloadducts.

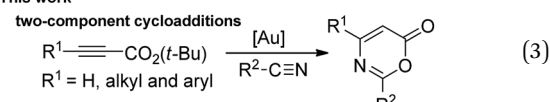
### Introduction

Metal-catalyzed  $[4\pi + 2\pi]$ -cycloadditions are powerful tools for the construction of carbo- or heterocyclic frameworks.<sup>1,2</sup> Although common nitriles and alkynes represent common triple bond motifs, nitriles are generally less reactive than alkynes in catalytic  $[4\pi + 2\pi]$ -cycloadditions; the chemical stability of nitriles is reflected by their bond energy (854 kJ mol<sup>-1</sup>), being larger than that of alkynes (835 kJ mol<sup>-1</sup>).<sup>3</sup> For instance, thermal  $[4\pi + 2\pi]$ -cycloadditions of dienes with unactivated nitriles required 600 °C (2 min) to give pyridine derivatives in 0.1–0.5% yields.<sup>4a</sup> In the context of catalytic  $[4\pi + 2\pi]$ -cycloadditions, not surprisingly, only one literature report documents both nitrile/1,3-diene and nitrile/1,3-ynye systems (eqn (1) and (2)).<sup>4b,c</sup> Ogoshi reported the first formal  $[4 + 2]$ -cycloadditions of common nitriles with dienes using Ni(0) catalysts (eqn (1)).<sup>4b</sup> Although Barluenga and Aguilar reported formal  $[4\pi + 2\pi]$ -cycloadditions of some 3-en-1-ynes with unactivated nitriles,<sup>4c</sup> such highly functionalized 3-en-1-ynes (X = *cis*-unsaturated ester, Z = alkoxy) are too specialized to reflect the reaction generality (eqn (2)). The  $[4\pi + 2\pi]$ -nitrile cycloadditions still remain an unsolved task for O- and N-substituted analogues of 1,3-dienes and 1,3-ynyes (X = O, NR', eqn (1) and (2)).<sup>5</sup> In a significant advance, we here report the gold-catalyzed formal hetero- $[4\pi + 2\pi]$ -cycloadditions<sup>6,7</sup> of various propiolates with nitriles to afford 6*H*-1,3-oxazin-6-ones efficiently (eqn (3)).<sup>8</sup> These findings enable the development of new cascade cycloadditions using three  $\pi$ -motifs including propiolates, nitriles and alkynes, yielding highly substituted pyridine derivatives. Notably, 6*H*-1,3-oxazin-6-ones are useful intermediates in various organic reactions whereas highly substituted pyridines

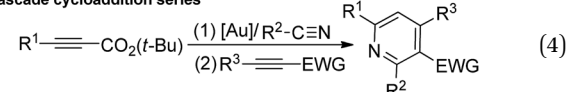
are important structural cores commonly found in many bioactive molecules (see ESI Fig. S1<sup>†</sup>);<sup>9,10</sup> their availability from convenient *t*-butyl propiolates increases the synthetic utility of this gold catalysis.



This work



cascade cycloaddition series



### Results and discussion

We envisage that direct  $[4\pi + 2\pi]$ -cycloadditions of propiolate derivatives with nitriles provide the most convenient synthesis of 6*H*-1,3-oxazin-6-ones such as 3; the current procedures rely mainly on thermal rearrangement of *N*-acyl  $\beta$ -

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† Electronic supplementary information (ESI) available. CCDC 1062512, 1062513 and 1062772. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c5sc01950h



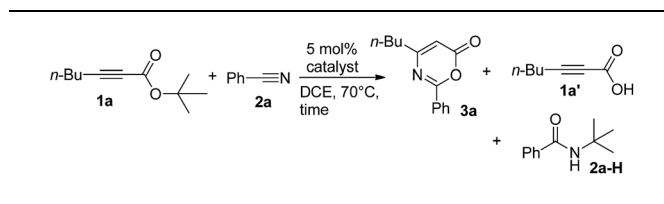
lactams.<sup>8a–d</sup> To test the feasibility, as shown in Table 1, *tert*-butyl hept-2-ynoate (**1a**, 1 equiv.) was treated with benzonitrile **2a** (3 equiv.) and  $\text{AuCl}_3$  (5 mol%) in hot DCE ( $70^\circ\text{C}$ , 16 h), affording the desired product **3a** in only a small yield (5%) together with the initial **1a** in 45% recovery (entry 1). The use of  $\text{PPh}_3\text{AuCl}/\text{AgSbF}_6$  significantly increased the yield of the desired **3a** to 51% (entry 2). We also examined other cationic gold catalysts (5 mol%) including  $\text{IPrAuCl}/\text{AgSbF}_6$  and  $\text{P}(t\text{-Bu})_2(o\text{-biphenyl})\text{AuCl}/\text{AgSbF}_6$ , yielding compound **3a** in 64% and 85% yields, respectively (see entries 3 and 4). With the alteration of the silver salts as in  $\text{P}(t\text{-Bu})_2(o\text{-biphenyl})\text{AuCl}/\text{AgX}$  ( $\text{X} = \text{NTf}_2$  and  $\text{OTf}$ ), the product yields slightly decreased to 77% and 72%, respectively (entries 5 and 6).  $\text{AgSbF}_6$  ( $70^\circ\text{C}$ , 24 h) and  $\text{Zn}(\text{OTf})_2$  (19 h) were found to be inactive in DCE, leading to a recovery of the starting compound **1a** in 72–75% yield (entries 7 and 8). The use of  $\text{In}(\text{OTf})_3$ ,  $\text{Sc}(\text{OTf})_3$  and  $\text{TfOH}$  in DCE gave hept-2-ynoic acid **1a'** in 65–72% yield and amide species **2a-H** (25–35% yield) along with unreacted starting compound **1a** (5–15% yield, entries 9–11). The yields of compound **3a** varied with the solvents ( $70^\circ\text{C}$ ), with 65% in toluene (22 h), 82% in  $\text{C}_6\text{H}_5\text{Cl}$  (18 h) and 56% in 1,4-dioxane (19 h, entries 12–14).

Table 2 assesses the reaction generality using various propiolate derivatives with varied nitriles. We first examined the reactions with unsubstituted propiolate species **1b**; its cycloaddition with benzonitrile **2a** proceeded smoothly to form the formal cycloadduct **3b** in 65% yield (entry 1). The reaction scope is extensible to aliphatically substituted

propiolate species **1c–1e** ( $\text{R} = \text{isopropyl, cyclopropyl and cyclohexyl}$ ), yielding the desired products **3c–3e** in satisfactory yields (77–85%, entries 2–4). This formal cycloaddition is also applicable to alkenyl-substituted propiolate **1f** to afford the corresponding product **3f** in 68% yield (entry 5). We tested the reactions on various phenyl-substituted propiolate species **1g–1j** bearing various *para*-substituents ( $\text{X} = \text{H, OMe, F and Cl}$ ); their resulting cycloadducts **3g–3j** were obtained in satisfactory yields (65–72%, entries 6–9). We performed an X-ray diffraction study of product **3g** to confirm its molecular structure.<sup>11</sup> We also prepared 2- and 3-thienyl-substituted propiolate derivatives **1k** and **1l**; their reactions with benzonitrile afforded cycloadducts **3k** and **3l** in reasonable yields (entries 10 and 11, 55–58%). Entries 12–15 show the tests of *tert*-butyl hept-2-ynoate **1a** with benzonitriles **2b–2e** bearing various *para*-substituents ( $\text{X} = \text{OMe, Me, CO}_2\text{Me, Cl}$ ) that afforded the desired cycloadducts **3m–3p** in satisfactory yields (62–76%). These catalytic cycloadditions were compatible with disparate nitriles including cyclohexyl nitrile (**2f**), cinnamonnitrile (**2g**) and 3-thienyl nitrile (**2h**), affording the expected products **3q–3s** in satisfactory yields (66–78%, entries 16–18).

As inferred from the chemistry of *2H*-pyran-2-ones,<sup>12,13</sup> one representative compound **3a** (1 equiv.) was treated with diethyl but-2-ynedioate (4 equiv.) in hot *p*-xylene ( $150^\circ\text{C}$ , 10 h) to afford tetrasubstituted pyridine **5a** in 96% yield; this reaction sequence presumably proceeds with intermediate **I** that is prone to a loss of  $\text{CO}_2$  (eqn (5)). As chlorobenzene is also an effective solvent for such a nitrile/propiolate cycloaddition (Table 1, entry 9), we developed a

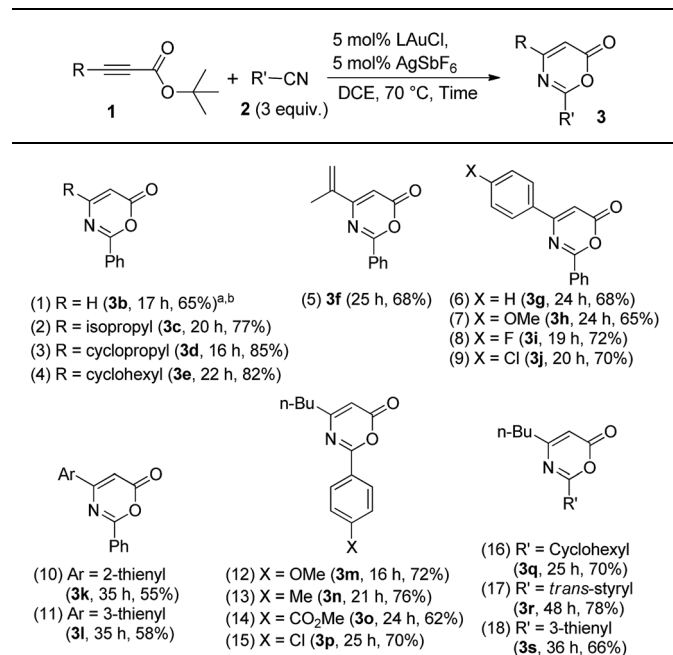
Table 1 Tests of propiolate derivatives with gold catalysts



Entries	Catalyst	Solvent	Time (h)	Yields <sup>a,b</sup> (%)			
				<b>1a</b>	<b>3a</b>	<b>1a'</b>	<b>2a-H</b>
1	$\text{AuCl}_3$	DCE	16	45	5	—	—
2	$\text{Ph}_3\text{PAuCl}/\text{AgSbF}_6$	DCE	12	—	51	—	—
3	$\text{IPrAuCl}/\text{AgSbF}_6$	DCE	19	—	64	—	—
4	$\text{LAuCl}/\text{AgSbF}_6$	DCE	18	—	85	—	—
5	$\text{LAuCl}/\text{AgNTf}_2$	DCE	20	—	77	—	—
6	$\text{LAuCl}/\text{AgOTf}$	DCE	22	—	72	—	—
7	$\text{AgSbF}_6$	DCE	24	75	—	—	—
8	$\text{Zn}(\text{OTf})_2^c$	DCE	19	72	—	—	—
9	$\text{In}(\text{OTf})_3^c$	DCE	18	15	—	72	35
10	$\text{Sc}(\text{OTf})_3^c$	DCE	22	10	—	65	32
11	$\text{HOTf}^c$	DCE	15	5	—	67	25
12	$\text{LAuCl}/\text{AgSbF}_6$	Toluene	22	—	65	—	—
13	$\text{LAuCl}/\text{AgSbF}_6$	$\text{C}_6\text{H}_5\text{Cl}$	18	—	82	—	—
14	$\text{LAuCl}/\text{AgSbF}_6$	1,4-Dioxane	19	—	56	—	—

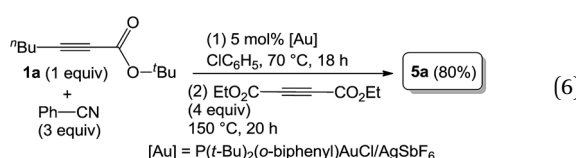
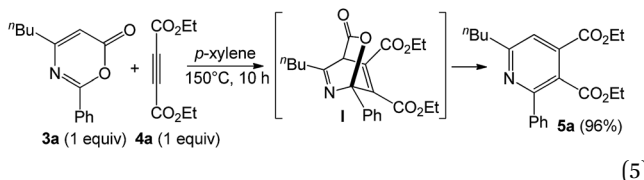
<sup>a</sup>  $[\mathbf{1a}] = 0.18 \text{ M}$ . <sup>b</sup> Product yields are reported after purification using a silica column.  $\text{IPr} = 1,3\text{-bis}(\text{diisopropyl phenyl})\text{-imidazol-2-ylidene}$ ,  $\text{L} = \text{P}(t\text{-Bu})_2(o\text{-biphenyl})$ ,  $\text{Tf} = \text{trifluoromethanesulfonyl}$ . <sup>c</sup> Reactions carried out at room temperature.

Table 2 Formal cycloadditions of various propiolates with nitriles



<sup>a</sup>  $[\mathbf{2}] = 0.18 \text{ M}$ . <sup>b</sup> Product yields are reported after purification using a silica column.  $\text{L} = \text{P}(t\text{-Bu})_2(o\text{-biphenyl})$ .

one-pot reaction involving the prior heating of a chlorobenzene solution of propiolate derivative **1a**, benzonitrile (3 equiv.) and  $P(t\text{-Bu})_2(o\text{-biphenyl})\text{AuCl/AgSbF}_6$  (5 mol%) at 70 °C (18 h) in a sealed tube to ensure a complete consumption of starting compound **1a**; to this solution was added diethyl but-2-yne dioate (4 equiv.) with further heating at 150 °C for 20 h. This one-pot process delivered the desired pyridine **5a** in 80% yield (eqn (6)). If the three reactants in the same proportions were heated together with a gold catalyst in hot chlorobenzene (150 °C, 20 h), the yield of **5a** was decreased to 38% yield.

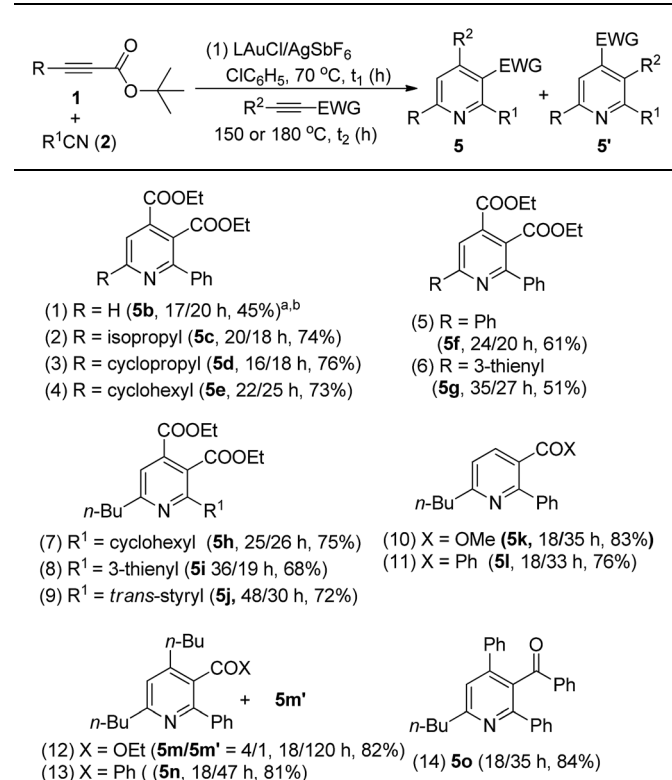


The easy operation of this one-pot reaction inspires us to examine the scope of the reaction using various propiolates, nitriles and alkynes; the results are summarized in Table 3. The procedures follow exactly that described in eqn (6). In the second stage of heating, the temperature is 150 °C for entries 1–7 and 180 °C for entries 8–12. Entry 1 shows the compatibility of these cycloadditions with unsubstituted propiolate derivative **1b** ( $R = H$ ) that reacted sequentially with benzonitrile (**2a**) and diethyl but-2-yne dioate (**4a**) to yield the desired pyridine **5b** in 45% yield. We also tested the reactions on various alkyl-substituted propiolates **1c–1e** ( $R = \text{isopropyl, cyclopropyl and cyclohexyl}$ ) that reacted with the same alkyne and benzonitrile to afford the desired pyridine species **5c–5e** in 73–76% yields (entries 2–4). The reaction is further applicable to aryl-substituted propiolates **1g** and **1l** ( $R = \text{Ph, 3-thienyl}$ ) to deliver the desired pyridines **5f** and **5g** in 61% and 51% yield, respectively (entries 5 and 6). We tested the reactions of model propiolate (**1a**) and diethyl but-2-yne dioate (**4a**) with various nitriles ( $R^1 = \text{cyclohexyl, 3-thienyl and } trans\text{-styryl}$ ), affording the expected pyridine products **5h–5j** in satisfactory yields (68–75%, entries 7–9). The reactions were extensible to various unsymmetric alkynes **4b–4f** that reacted with propiolate (**1a**) and benzonitrile (**2a**) with excellent or high regioselectivity (entries 11–15). The reactions worked well for terminal alkynes **4b** ( $\text{EWG} = \text{COOMe}$ ) and **4c** ( $\text{EWG} = \text{COPh}$ ) to afford the desired pyridines **5k** and **5l** as single regioisomers, with respective yields of 83% and 76% (entries 10 and 11). For *n*-butyl propiolate **4d**, this one-pot sequence gave two inseparable isomeric products **5m/5m'** = 4/1, in a combined 82% yield (entry 12). For the other *n*-butyl and phenyl-substituted

ynones **4e** and **4f** ( $\text{EWG} = \text{COPh}$ ), their reactions afforded **5n** and **5o** with excellent regioselectivity and satisfactory yields (81–84%) (entries 13–14). The structures of representative compounds **5m** and **5n** were confirmed by proton NOE effects whereas the structure of cycloadduct **5o** was elucidated with an HMBC experiment (see ESI†).

As nitriles are weakly nucleophilic, we envisage that aldehydes and ketones might be applicable substrates. To our pleasure, gold-catalyzed reactions of 3-phenylpropiolate **1g** with benzaldehyde, phenyl methyl ketone and acetone in hot dichloroethane (DCE) proceeded smoothly to afford formal cycloadducts **6a–6c** in high yields (86–89%, eqn (7)). The structure of compound **6a** was determined by X-ray diffraction.<sup>11</sup> These carbonyl cycloadditions were also applicable to alkyl-substituted propiolates (**1a**) and (**1e**), yielding the desired compounds **6d** and **6e** in 87% and 77% yield, respectively (eqn (8)). Such a reaction was, notably, accessible to an eight-membered oxacyclic compound **6f** with 2.5 mol% 1,3-bis(diisopropyl phenyl)-imidazol-2-ylidene  $\text{AuSbF}_6$ ; it was isolated as a single regioisomer with 67% yield with 2-phenyloxetane (3 equiv.) and its molecular structure has been confirmed by X-ray diffraction.<sup>11</sup> The compatibility of this gold catalysis with aldehydes, ketones and oxetanes truly reflects a broad applicability of these cycloadditions.

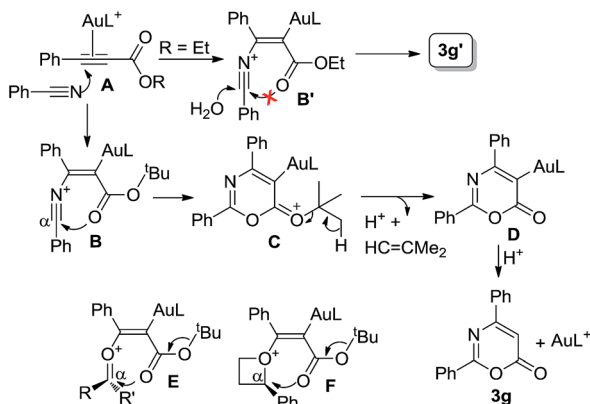
Table 3 One-pot operations with nitriles, propiolates and alkynes



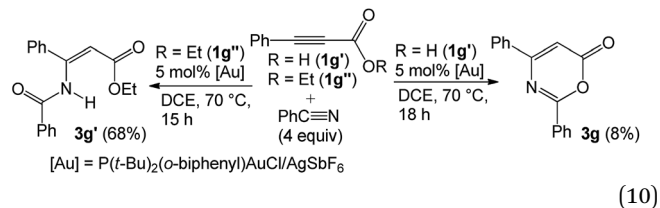
<sup>a</sup> 5 mol% gold catalyst,  $\text{L} = P(t\text{-Bu})_2(o\text{-biphenyl})$ ,  $\text{R}^1\text{CN}$  (3 equiv.),  $\text{R}^2\text{---EWG}$  (4 equiv.), 150 °C for entries 1–9 and 180 °C for entries 10–14.

<sup>b</sup> These data correspond to the reaction time  $t_1/t_2$ .





Scheme 1 A postulated reaction mechanism.



(10)

The control experiments in eqn (10) indicate a mechanism involving a prior formation of nitrilium species **B** *via* a  $\pi$ -alkyne activation, proceeding with an attack of nitrile at the gold- $\pi$ -alkyne species **A**. As shown in Scheme 1, we postulate that the *tert*-butoxy group of species **B** increases the nucleophilicity of a carbonyl group to attack this nitrilium moiety efficiently. This process releases a *tert*-butyl cation to induce a demetalation to form the observed cycloadduct **3g**. Beside nitriles, various aldehydes, ketones and oxetanes are more reactive than alkenes upon comparison of their applicable propiolates. We postulate that these nucleophiles generate intermediates **B**, **E** and **F** bearing a large positive charge on the reacting  $C_{\alpha}$ -carbons because of their adjacent oxonium and ammonium centers. We envisage that the propiolate cycloadditions match well with those nucleophiles that can develop highly polarized carbocations through  $\pi$ -alkyne activations.

## Conclusions

Unactivated nitriles are known to be stable triple-bond species, and their [4 + 2]-cycloadditions with 4 $\pi$ -bond motifs and other small molecules have few successful examples.<sup>15</sup> This work reports the hetero-[4 $\pi$  + 2 $\pi$ ]-cycloadditions of *tert*-butyl propiolates and nitriles, yielding useful 6*H*-1,3-oxazin-6-ones, which are not readily prepared with current methods.<sup>8</sup> This new finding enables a one-pot gold-catalyzed synthesis of highly substituted pyridines through sequential reactions of *tert*-butyl propiolates with nitriles, and then with electron-deficient alkynes in the same solvent. The utility of these [4 + 2]-cycloadditions is further expanded with various aldehydes, ketones and 2-phenyloxetane, yielding satisfactory yields of cycloadducts. This work provides a new version of *tert*-butyl propiolates that feature useful four-atom building blocks with polar  $\pi$ -bond motifs such as nitriles, aldehydes and ketones, although their reactions with alkenes were reported to be restrictive.<sup>8</sup>

## Acknowledgements

We thank the National Science Council, Taiwan, for financial support of this work.

## Notes and references

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Prior to this work, Shin reported gold-catalyzed [4 + 2]-cycloadditions of alkenes with propionic acid, which was, however, the only applicable substrate.<sup>6a</sup> Here, we employ diverse propiolate substrates to comply with not only nitriles but also aldehydes, ketones and oxetanes. To understand this discrepancy, we performed the reaction of 3-phenylpropionic acid (**1g'**) with benzonitrile with the same gold catalyst in DCE, but the yield of the desired compound **1g** was only 8%, much smaller than that (68%) of its *tert*-butoxy derivative **1g''** (Table 2, entry 5). Clearly, prior transformations of *tert*-butoxy propiolates to the propiolate acids do not occur in the course of the reactions. For ethyl propiolate **1g''**, its corresponding reaction with benzonitrile gave the amide-addition product **3g'** in 68% yield (eqn (10)); under this condition, benzonitrile was not effectively transformed into benzamide with this gold catalyst.<sup>14</sup>





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