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Hemilabile MIC^N ligands allow oxidant-free Au(I)/Au(III) arylation-lactonization of γ -alkenoic acids

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Oxidant-free Au-catalyzed reactions are emerging as a new synthetic tool for innovative organic transformations. Still, a deeper mechanistic understanding is needed for a rational design of these processes. Here we describe the synthesis of two Au(I) complexes bearing bidentated hemilabile MIC^N ligands, [Au¹(MIC^N)Cl], and their ability to stabilize square-planar Au(III) species (MIC = mesoionic carbene). The presence of the hemilabile N-ligand contributed to stabilize the ensuing Au(III) species acting as a five-membered ring chelate upon its coordination to the metal center. The Au(III) complexes can be obtained either by using external oxidants or, alternatively, by means of feasible oxidative addition with strained biphenylene C_{sp2}-C_{sp2} bonds as well as with aryl iodides. Based on the fundamental knowledge gained on the redox properties on these Au(I)/Au(III) systems, we successfully develop a novel Au(I)-catalytic procedure for the synthesis of γ -substituted γ -butyrolactones through the arylation-lactonization reaction of the corresponding γ -alkenoic acid. The oxidative addition of the aryl iodide, which in turn is allowed by the hemilabile nature of the MIC^N ligand, is an essential step for this transformation.

Introduction

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Gold catalysis has been dominated by the Lewis acidity of gold towards activation of alkenes and alkynes for nucleophilic attacks.¹⁻⁶ On the other hand, redox Au(I)/Au(III) catalysis has been less explored and mainly achieved in the presence of sacrificial external oxidants,⁷⁻¹⁷ especially due to the reluctance of Au(I) to undergo oxidative addition.¹⁸⁻²¹ In this regard, several strategies have been explored to overcome this limitation,²²⁻⁴⁶ among which the chelation-assisted strategy in pre-designed ligands³⁰⁻³⁴ or the oxidative addition using strained molecules such as biphenylene stand out.³⁵ Bourissou pioneered in 2014 the intramolecular oxidative addition of C_{sp2} -X bonds (X = Br, I) at Au(I) by utilizing rigid 8-halonaphtyl phosphine model substrates, which suitably place the C_{sp2}-X bond close to the Au(I) atom upon coordination to the phosphine moiety. In this manner, facile stabilization of the (P,C)-cyclometalated Au(III) products is obtained via oxidative addition.³² In the same line, Hashmi and Hermange used a similar strategy taking advantage of pendant pyridine groups in monophosphine ligands to

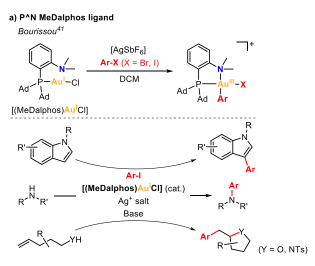
stabilize Au(III) systems upon reaction of Au(I) complexes with aryldiazonium salts.^{47, 48} In these examples, the subtle change from linear to square-planar geometry is crucial for the facile reactivity of Au(I) species towards a key oxidative addition process. Noteworthy, P^N MeDalphos ligand showed great performance at promoting the oxidative addition of aryl halides at Au(I), as well as at stabilizing Au(III) intermediates, which paved the way to Au(I)/Au(III) catalytic transformations using the (MeDalphos)AuCl catalyst (Scheme 1a).⁴¹ For instance, in 2019 Bourissou and coworkers reported the regioselective Au(I)/Au(III) catalyzed C3 arylation of indoles, which is rarely attained by other transition metals (Scheme 1a),⁴⁹ additionally they described the Au(I)/Au(III) catalyzed C3 allylation of indoles using allyl alcohols or allyl acetates.⁵⁰ Significantly, two independent reports by the groups of Patil⁵¹ and Bourissou⁵² disclosed the C_{sp2}-N cross-coupling reactions of aryl iodides with amines mediated by the MeDalphos-enabled Au(I)/Au(III) catalysis (Scheme 1a). In 2020, Bourissou and Patil groups independently showed that the oxidative addition of aryl iodides and π -activation of olefins can be merged by the Au(I)/Au(III) platform using (MeDalphos)AuCl as catalyst, which involves the oxy- and aminoarylation reactions of alkenols and alkenamines (Scheme 1a).^{53, 54} The 1,2-heteroarylation of alkenes was also achieved using external alcohols and amines.⁵⁴⁻⁵⁶ Interestingly, (MeDalphos)AuCl catalyst has proven to be a key factor in the development of other transformations such as 1,2-diarylation of alkenes⁵⁷ and trifluoromethylthiolation, and trifluoromethylselenolation of organohalides.⁵⁸ Very recently, Shi and Patil groups developed new chiral-hemilabile P^N-ligands to access to enantioselective Au(I)/Au(III) catalysis.^{59, 60}

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[†]Electronic Supplementary Information (ESI) available: for materials, instrumentation, experimental procedures and spectroscopic characterization of all compounds. CCDC 2163608 (**4a-OAc**), 2163609 (**2a**), 2163610 (**2b**), 2163611 (**4a-Cl**), 2163612 (**8a-OMe**), 2163613 (**3b-biphenylene**), 2163614 (**5b**), 2163615 (**6b**), 2163616 (**8b-OMe**), 2163617 (**3a**), 2163877 (**16**) and 2176932 (*cis*-**7a-Cl**) contain the supplementary crystallographic data for this paper. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif. See DOI: 10.1039/x0xx00000x

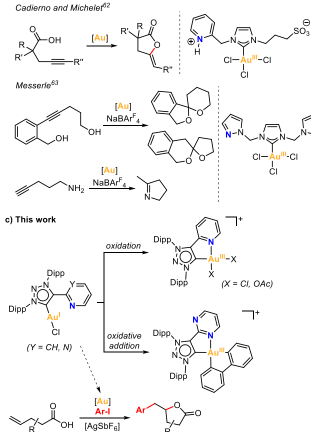
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b) NHC^N ligand

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Scheme 1. (a) Oxidative addition of aryl halides to a Au(I) complex bearing the hemilabile P^N MeDalphos ligand, and selected examples of its catalytic activity. (b) Selected examples of gold complexes bearing hemilabile NHC^N ligands. (c) Ability of hemilabile (MIC^N) ligands to stabilize Au(III) complexes obtained from Au(I) *via* oxidation or *via* oxidative addition, and application of the (MIC^N)Au(I) complexes to the arylation-lactonization reaction of γ -alkenoic acids.

In contrast to the known P^N ligands, the use of bidentate C^N or C^O ligands⁶¹ remains scarcely explored, and the few representative examples are described in Scheme 1b. Cadierno and Michelet studied the catalytic activity of a (NHC^N)Au(III) complex containing a N-heterocyclic carbene ligand (NHC) with a pyridinium side arm in the cyclization of γ -alkynoic acids

(Scheme 1b),⁶² while Messerle and coworkers explored the dihydroalkoxylation and hydroamination Catalyzed by Scheffff) complex bearing a NHC ligand with two pendant pyrazole arms (Scheme 1b).⁶³ Alternatively, Bertrand and coworkers showed that a hemilabile bidentate cyclic (alkyl)(amino)carbene (CAAC) Au(I) complex undergoes oxidative addition of biphenylene.⁴⁰

More recently, Bourissou and coworkers reported the reactivity of a Au(I) complex containing a N3-alkylated mesoionic 1,2,3triazol-5-ylidene bearing a pendant pyridine group.⁶⁴ However, the complex did not undergo oxidative addition towards iodobenzene, thus precluding its exploitation in oxidant-free reactivity. In fact, the formation of a dimeric Au(I) species was observed instead of the formation of the oxidative addition Au(III) product. On the other hand, the Au(I) complex was reactive towards a strong oxidant such as PhICl₂, forming a mononuclear square-planar Au(III) species. The latter experiment demonstrated that mesoionic carbene ligands bearing N-based hemilabile groups (MIC[^]N) can potentially be suitable platforms to stabilize Au(III) species.

Based on the abovementioned, we herein explore the reactivity of two new (MIC^N)Au(I) complexes bearing hemilabile pyridine or pyrimidine pendant groups, their ability to stabilize (MIC^N)Au(III) species and their performance in oxidant-free oxidative addition of aryl halides and strained C-C bonds. After proving the successful ability to sustain oxidant-free Au(I)/Au(III) redox processes, we further extended its application to arylation-lactonization reactions of γ -alkenoic acids (Scheme 1c).

Results and discussion

Synthesis and characterization of (MIC^N)Au(I) complexes

Initially, two 1,2,3-triazolium ligand salt precursors (MIC^N) bearing 2,6-diisopropylphenyl substituents (Dipp) at the outer nitrogens of the azolium ring and either a pyridine (1a) or a pyrimidine (1b) side arm were prepared by following an adapted method reported in the literature (Figure 1a).⁶⁵ We reacted 1,3-bis(2,6-diisopropylphenyl)triaz-1-ene with 2ethynylpyridine or 2-ethynylpyrimidine in the presence of tertbutyl hypochlorite (^tBuOCl) and anhydrous potassium hexafluorophosphate (KPF₆), affording the desired products in high and moderate yield (99 % for 1a and 67% for 1b). In order to obtain the corresponding (MIC^N)Au(I) complexes, we attempted the Au(I) coordination through a transmetalation reaction to Ag(I) species. For this purpose, we reacted the corresponding triazolium salts with silver oxide (Ag₂O) in the presence of cesium carbonate (Cs₂CO₃) and potassium chloride (KCl) as halide source. Then, the addition of dimethylsulfide gold(I) chloride [AuCl(SMe2)] immediately produced the precipitation of the silver halide. After purification, the desired Au(I) complexes 2a and 2b were obtained in 32% and 35% yield, respectively. The ¹H NMR spectra of the complexes showed the absence of the acidic proton of the triazolium salts, indicating that the coordination to gold had occurred. The ¹³C¹H NMR spectrum of 2a showed the characteristic signal of the metalated carbon at 162.1 ppm, while in the case of the

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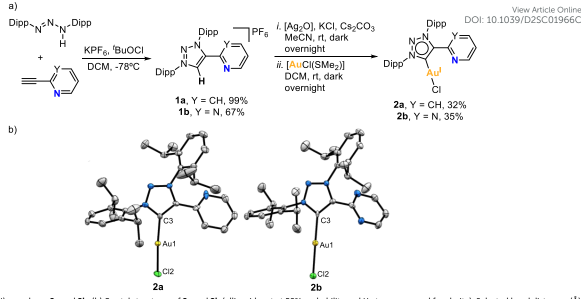


Fig. 1. (a) Synthesis of Au(I) complexes 2a and 2b. (b) Crystal structures of 2a and 2b (ellipsoids set at 50% probability and H atoms removed for clarity). Selected bond distances (Å): for 2a, Au1-C3 1.982(6), Au1-Cl2 2.2854(17); for 2b, Au1-C3 2.006(4), Au1-Cl2 2.2728(16).

pyrimidine derivative **2b**, such signal was slightly shifted downfield, appearing at 165.3 ppm. Furthermore, the molecular structure of both Au(I) complexes **2a** and **2b** was unambiguously determined by X-ray diffraction analyses (Figure 1b). Both complexes are isostructural, showing the mesoionic carbene ligand coordinated to an Au(I) atom, and a chlorine ligand completing the linear coordination sphere around the metal (C-Au-Cl angle is 174.31(18)° for **2a** and 179.30(9)° for **2b**). Noticeably, no coordination of the hemilabile pyridine or pyrimidine moieties at Au(I) was observed.

Then, Au(I) complexes **2a** and **2b** were reacted with silver hexafluoroantimonate (AgSbF₆) in CD_2CI_2 at room temperature (Figure 2a), seeking the engagement of the pendant pyrimidine or pyridine groups in the coordination to Au(I). The ¹H NMR spectra of the corresponding complexes showed some changes in the aromatic and aliphatic region. In particular, the signals of the pyridine or pyrimidine fragments were shifted due to gold coordination. The most significant change was observed in the ¹³C{¹H} NMR spectra. The signal of the carbenic carbon was shifted from 162.1 to 157.2 ppm in the case of the pyridine derivative (**3b**).

X-ray diffraction analysis of complex **3a** showed a head-to-tail dimeric Au(I) species where each ligand is bridging two metals (Figure 2). The Au(I) centers adopted a linear dicoordinate arrangement, showing an aurophilic interaction (Au…Au length of 2.8212(4) Å). Analogously to Bourissou's (MIC^N)Au(I) complex,⁶⁴ the formation of a dimeric Au(I) species was favored rather than a mononuclear Au(I) having both the MIC and py or pym pendant groups coordinated.

Reactivity of (MIC^N)Au(I) complexes with external oxidants and towards oxidative addition

In order to promote the coordination of the side arms of our MIC^N ligands, we attempted the 2 e⁻ chemical oxidation of the Au(I) complexes to mononuclear Au(III). To that end, we studied

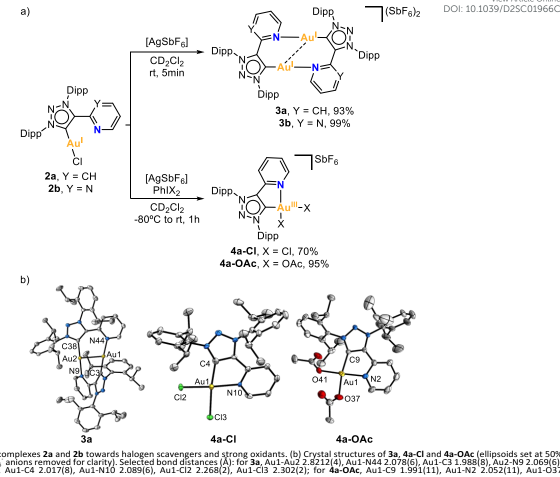
the reactivity of complex 2a against a 2 e⁻ oxidant, namely PhIX₂ (X = Cl or OAc) (Figure 2a). The reaction was performed at low temperature, from -80 °C to rt, in the presence of AgSbF₆ as halogen scavenger. Gratifyingly, the oxidation of Au(I) to Au(III) took place, with the concomitant coordination of the pendant N-moiety to gold. The ¹³C{¹H} NMR spectra of the resulting complexes 4a-Cl and 4a-OAc showed the characteristic signal of the carbenic carbon at higher field when compared to the Au(I) analogue 2a (147.3 ppm for 4a-Cl and 137.1 ppm for 4a-OAc vs 162.1 ppm for 2a). The molecular structures of complexes 4a-Cl and 4a-OAc were elucidated by X-ray diffraction analyses (Figure 2b). Both complexes showed a similar arrangement, i.e. the bidentate MIC^N ligand coordinated to Au(III), and two chlorides or acetates completing its tetracoordinated Au(III) center, which featured a slightly distorted square planar geometry. The C_{carbene}-Au(III) lengths were slightly longer than those found in the Au(I) analogue (2.017(8) Å for 4a-Cl and 1.991(11) Å for 4a-OAc vs 1.982(6) Å for 2a).

At this point, we had demonstrated that these MIC^N ligands are indeed suitable platforms to stabilize Au(III) species. Thus, we were encouraged to explore the possibility of obtaining Au(III) complexes via oxidative addition. With this aim, we first attempted the oxidative addition of complex 2b towards the strained C_{sp2} - C_{sp2} bond of biphenylene, by mixing equimolar amount of 2b and biphenylene in dichloromethane, in the presence of a halide scavenger, from -80 °C to room temperature for 10 minutes (Figure 3a). Unfortunately, we did not observe the desired Au(III) center with a biphenyl moiety but, interestingly, crystallization of the reaction mixture revealed the co-crystallization of the dimeric complex 3b with intact biphenylene (3b·biphenylene) (Figure 3b). Even after 5 hours at 50 °C, the outcome of the reaction remained the same. To our delight, when the reaction was heated at 90 °C for 2 hours in 1,2-dichloroethane, the Au(I) center underwent oxidative addition of the strained biphenylene Csp2-Csp2 bond, forming the expected Au(III) complex 5b (Figure 3a). Notably,

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(a) Reactivity of Au(I) complexes **2a** and **2b** towards halogen scavengers and strong oxidants. (b) Crystal structures of **3a**, **4a-Cl** and **4a-OAc** (ellipsoids set at 50% bility, H atoms and SbF₆ anions removed for clarity). Selected bond distances (Å): for **3a**, Au1-Au2 2.8212(4), Au1-N44 2.078(6), Au1-C3 1.988(8), Au2-N9 2.069(6), 38 1.990(8); for **4a-Cl**, Au1-C4 2.017(8), Au1-N10 2.089(6), Au1-Cl 2.268(2), Au1-Cl 2.302(2); for **4a-OAc**, Au1-C9 1.991(11), Au1-N2 2.052(11), Au1-O37 2.032(10), Au1-041 1.998(9)

the presence of a chloride source (such as KCl or tetrabutylammonium chloride) displaced the hemilabile pyrimidine, affording a neutral Au(III) species, 6b. By adding AgSbF₆ as halide scavenger, we could reversibly form complex 5b, also reinforcing the hemilabile character of the pyrimidine moiety. Indeed, the NMR spectra of 5b showed a dynamic behaviour of the pyrimidine pendant arm, even at 248 K, as indicated by ¹H,¹H-NOESY experiment (Figure S49). Both the cationic 5b and the neutral 6b Au(III) species were characterized by X-ray diffraction analyses (Figure 3b), confirming the weak coordination of the pyrimidine moiety with a long Au(III)-N_{pyrimidine} bond of 2.254 Å for **5b**. The carbene resonance signals were downfield shifted to 182.8 and 180.1 ppm for 5b and 6b, respectively.

We then explored the possibility of Au(I) complexes 2a and 2b to undergo oxidative addition of C_{sp2}-I bonds (Figure 4a). For this purpose, we reacted the Au(I) complexes with para-substituted iodoaryls (R = OMe, Me, F) in the presence of $AgSbF_{6}$, in 1,2dichloroethane and heating at 120 °C overnight. Unexpectedly, no Au-containing compounds were isolated, but on the contrary, the ¹H NMR and mass spectra were consistent with the full conversion to three triazolium salts (8a-R, 9a, and 1a when 2a was used, and 8b-R, 9b and 1b when 2b was used; R = OMe, Me, F) (see yields in Figure 4a). When using 2a with 4iodoanisole, the triazolium salts 8a-OMe and 9a were

unambiguously characterized by X-ray diffraction. Both species co-crystallized, and the single crystal X-ray analysis revealed the presence of 8a-OMe and 9a in a 0.89:0.11 ratio (Figure 4b). Likewise, when 2b was reacted with 4-iodoanisole, the major product was 8b-OMe with a 65% NMR yield, whereas 9b and 1b were both obtained in 16% NMR yield. In this case, 8b-OMe could be isolated and characterized by ¹H NMR, HRMS and Xray diffraction (Figure 4b). The formation of such compounds suggested that gold(I) compounds underwent oxidative addition of the aryl iodide followed by a fast reductive elimination of the mesoionic carbene ligand either with the aryl or iodide moiety coordinated in cis, respectively (Figure 4a). Interestingly, products 8a-R/8b-R were obtained in higher yields than 9a/9b, indicating that after the oxidative addition the major product is the one with the aryl ligand in cis (cis-7a-I or cis-7b-I), probably due to the trans effect of the NHC ligand. By an alternative route, 47 the analogous complex cis-7a-Cl was synthesized through the oxidative addition of an aryldiazonium salt promoted by blue LED irradiation (λ = 447 nm). This methodology has the advantage to occur at 25 °C, allowing the isolation and fully characterization of the Au(III) species cis-7a-CI (see XRD in Figure 4b). When cis-7a-CI was heated at 120 °C overnight in 1,2-DCE, product 8a-OMe was formed in 96% yield. This reaction supports that the formation of 8a-R/8b-R and 9a/9b from Au(I) complexes 2a/2b

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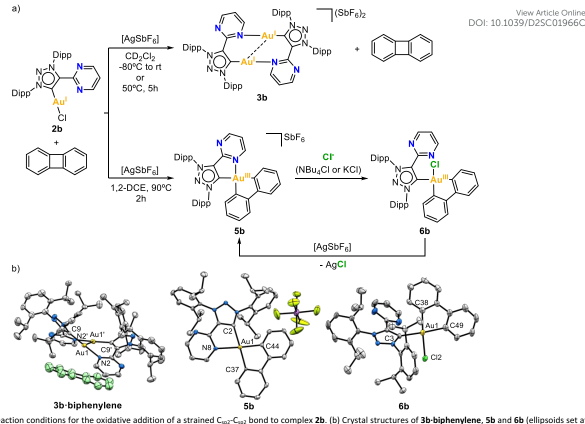


Fig. 3. (a) Study of the reaction conditions for the oxidative addition of a strained Csp2-Csp2 bond to complex 2b. (b) Crystal structures of 3b-biphenylene, 5b and 6b (ellipsoids set at 50% probability. H atoms removed for clarity: in the case of **3b-biphenvlene** the SbF₆ anions have been removed and the biphenvlene molecule is painted in green for clarity). Selected bond distances (Å): for 3b-biphenylene, Au1-Au1' 2.8768(10), Au1-N2 2.105(8), Au1-C9 2.027(10); for 5b, Au1-C2 2.108(5), Au1-N8 2.254(5), Au1-C37 2.066(5), Au1-C44 2.019(5); for 6b, Au1-C3 2.080(6), Au1-Cl2 2.376(3), Au1-C38 2.072(7), Au1-C49 2.057(7).

proceeds via an oxidative addition/reductive elimination pathway. We hypothesize that the hemilabile character of the nitrogen atom and the fast decomposition of the Au(III) at high temperatures promote the reductive elimination process, which has only been observed to our knowledge for a 1,2,3-triazol-5-ylidene based palladium complex.⁶⁶⁻⁶⁸ Nevertheless, our observations sharply contrasted with the lack of reactivity described by Bourissou for a similar (MIC^N)Au(I) system with aryl iodides.⁶⁴

Arylation-lactonization of y-alkenoic acids

Based on the successful catalytic application of the P^N hemilabile ligand strategy, we envisioned that our hemilabile (MIC^N)Au(I) complexes may also mediate a cascade reaction involving the key oxidative addition step. Recently, Bourissou and coworkers described the coupling/cyclization reaction of aryl iodides with alkenol and alkenamine compounds catalyzed by a hemilabile P^N MeDalPhos/Au(I) system.⁵³ Additionally, in 2017, Shi and coworkers reported a system that afforded coupling/cyclization products upon reacting aryldiazonium salts with either alkenamines, alkenols or alkenoic acids, using [(Ph₃P)AuCl] under photo-free conditions.⁶⁹ Thus, we envisioned that our MIC^N/Au(I) system should afford γ-benzyl- γ -butyrolactone products when reacting aryl iodides with γ alkenoic acids, via coupling/lactonization, provided the oxidative addition step takes place (see reaction scheme embedded in Table 1). Our first attempts consisted in a stoichiometric reaction between 4-iodoanisole, 4-pentenoic

acid, complex 2b, K₃PO₄, and AgSbF₆ at 80 °C. When the reactions were carried out in the presence of an excess of base or AgSbF₆, we did not observe any conversion, probably due to the rapid decomposition of the Au(I) complex. We also found that by using chlorinated solvents such as 1,2-dichloroethane, the conversion reached up to 49% (Table S3). We determined by ¹H NMR analysis that the catalyst decomposes under the reaction conditions, forming the triazolium salt 1b. Interestingly, the conversion was significantly higher using 2,2,2-trifluoroethanol (TFE) (up to 92%) instead of 1,2dichloroethane (Table 1 and S3). When the reaction was carried out at 100 °C, the conversion was quantitative, but the yield was very similar to that found at 80 °C (67% yield at 100 °C vs 63% yield at 80 °C, entries 1-2 in Table 1), and we also observed the formation of the triazolium salt 8b-OMe. This was a strong indication that complex 2b underwent an oxidative addition of the C_{sp2} -I bond, but rapid reductive elimination to form the coupling product 8b-OMe occurred.

With these results in hand, we explored the possibility to perform the catalytic version of the arylation-lactonization of γ alkenoic acids. Thus, we carried out the reaction decreasing the loading of 2b, first using substoichiometric 70 mol% of the gold complex. Product 10 was obtained in 65% yield (entry 3), so effectively a quantitative transformation was achieved with respect to Au loading. By using 40 mol% of Au(I) complex the yield was 44%, reaching about 1 TON (entry 4). Interestingly,

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View Article Online DOI: 10.1039/D2SC01966C a) Oxidative addition Œ Ð R [AgSbF₆] Dipp 1,2-DCE, 120°C Dipp overnight ςι (R = OMe, Me, F) Dipp Elusive 2a Y = CH 2b, Y = N species trans-7a-l cis-7a-N₂BF₄ cis-7btrans-7b-l Oxidative addition CD₂Cl₂ 25°C, 16h Reductive elimination (fast) hv (blue LED) 🗖 ŻМе BF₄ Æ Œ Ð Dipp C Din Dip AuX 1.2-DCE, 120°C Dip overnight Reductive elimination Dipp Dipp Dipp ¥ ÓMe Au⁰ 8a-R, Y = CH 9a, Y = CH 1a, Y = CH cis-7a-Cl 9b, Y = N 1b, Y = N 8b-R, Y = N From **2a** / **2b** R = OMe: 8a-OMe 41% 9a 3% 1a 24% 8b-OMe 65% 9b 16% 1b 16% R = Me 8a-Me 34% 9a 3% 1a 31% R = F8a-F 12% 9a traces 1a 66% From cis-7a-Cl 8a-OMe 96% b) CI2 8a-OMe cis-7a-C 8b-OMe 9a

Fig. 4. (a) Reactivity towards aryl iodides and decomposition route of Au(I) species 2a and 2b, and synthesis of cis-7a-Cl by oxidative addition of an aryldiazonium salt. NMR yield of the reactions calculated using 1,3,5-trimethoxybenzene as internal standard. (b) Crystal structures of cis-7a-Cl, 8a-OMe, 9a and 8b-OMe (ellipsoids set at 50% probability, H atoms removed for clarity; for cis-7a-Cl, BF4 anion is removed for clarity; for 8a-OMe and 9a, SbF6 anion is not shown for clarity). Selected bond distances for cis-7a-Cl (Å): Au1-C10 2.009(7), Au1-N3 2.154(7), Au1-C38 2.006(8), Au1-Cl2 2.303(2).

using a lower amount of 2b (10 mol%) the yield was around 28% (~3 TON, entry 5). Similarly, by using hexafluoro-2-propanol (HFIP) the yield was 29% (~3 TON, entry 9); therefore, the best optimized results were obtained with fluorinated alcohols.

To exclude the possibility of oxidizing Au(I) to Au(III) in the presence of Ag(I), complex 2a (1 eq) was reacted with AgSbF₆ (15 eq) in HFIP at 80 °C for 16 hours. Quantitative formation of the Au(I) dimer 3a was observed by NMR, confirming the role of silver as halide scavenger (see Scheme S12).

The scope of the reaction was examined with different aryl halides and γ -alkenoic acids (Figure 5). The arylationlactonization using iodobenzene and complex 2b reached 50% yield (~5 TON) in product 11. As expected, by increasing the

catalyst loading to 20 mol%, the yield of 11 increased up to 87%, thus maintaining the turnover number (~4-5 TON). Gratifyingly, the best yields were reached using 4-iodotoluene, obtaining product 12 in 60% yield using 2a (~6 TON) and 81% yield using 2b (~8 TON). However, the presence of the strong electronwithdrawing group CF_3 in the para-position of the iodoaryl produced product 13 in a lower yield (~13%). Following this trend,^{36,38} 1-iodo-4-nitrobenzene produced a more dramatic drop of the yield of the corresponding product **14** (1%).

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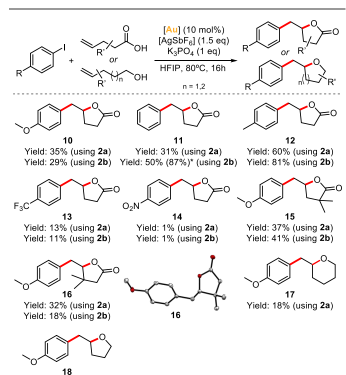
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 Table 1. Synthesis of v-benzyl-v-butyrolactone 10 promoted by complex 2b.

	+			$\frac{2b}{F_{6}, K_{3}PO_{4}}$	
Entry	[Ag] (eq)	[Au] mol%	K₃PO₄ (eq)	Solvent	Conversion % (Yield %)
1 ^ª	2.8	100	1.0	TFE	92(63)
2 ^{a,b}	2.6	100	1.2	TFE	>99(67)
3 [°]	2.2	70	1.2	TFE	81(65)
4 ^c	2.0	40	1.1	TFE	79(44)
5 ^b	1.5	10	1.0	TFE	51(28)
6	1.5	10	0.5	TFE	51(24)
7 ^c	1.5	10	1.1	TFE	54(15)
8 ^c	1.1	1	0.5	TFE	38(1.6)
9	1.5	10	1.0	HFIP	56(29)

Reaction conditions: [Ar-I] = 0.08 M, [4-pentenoic acid] = 0.08 M, 80 °C, 16h. V = 0.55 – 1.40 mL (¹H NMR yield obtained with 1,3,5-trimethoxybenzene as internal standard). ^a[Ar-I] = 0.02 M, [4-pentenoic acid] = 0.1 M, V = 1 mL. ^bT = 100 °C. ^cReaction time = 24h.

Interestingly, the γ -alkenoic acid with α -methyl groups reacted smoothly, yielding 37% of **15** using **2a** and 41% of **15** using **2b**. Also, complexes **2a** and **2b** performed very similar, except in the case of using iodobenzene and 4-pentenoic acid, yet complex **2b** is almost two-fold more active



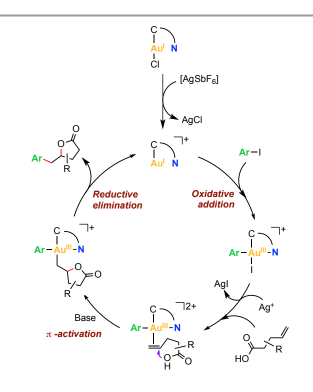
Yield: 25% (using 2a)

Fig. 5. Scope of the arylation-lactonization of alkenoic acids (10 – 16) and oxyarylation of alkenes (17 – 18). *NMR yield using 20% of [Au] is shown in parenthesis. Crystal structure of 16 is shown (ellipsoids set at 50% probability, H atoms removed for clarity).

than **2a** (31% vs 50% yield of compound **11**, for **2a** render **2b**, respectively). The molecular structure of compound **16** was determined by X-ray diffraction analysis and revealed the formation of the (R)-enantiomer by spontaneous resolution.

Finally, in order to broaden the scope of these (MIC^N)Au(I) complexes, we tested complex **2a** in the oxyarylation of alkenols and 1,2-diarylaltion of alkenes under the same catalytic conditions (Scheme S11). The oxyarylation products **17** and **18** were obtained in low but significant yields (18 - 25%, ~2 TON) (Figure 5 and Table S6). On the contrary, the 1,2-diarylation reaction was unsuccessful.

Based on the fundamental understanding of the redox behaviour of complexes 2a and 2b, and the previously reported gold-catalyzed processes combining aryl halide oxidative addition and π -activation of alkenes, ⁵³ a mechanistic proposal is depicted in Figure 6. First, the oxidative addition of the C_{sp2}-I bond into a cationic gold(I) species occurs, affording a cationic arylgold(III) species. Then, the silver salt abstracts the iodide from this Au(III) species to generate a vacant site that can be occupied by the π -coordination of the olefin. Subsequent intramolecular y-lactonization is enabled by the presence of base and the π -activation of the alkene produced by the gold center, to yield an alkylarylgold(III) intermediate. The latter is proposed to undergo a reductive elimination step that affords the y-benzyl-y-butyrolactone product and closes the catalytic cycle by regenerating the initial catalytically active hemilabile [Au^l(MIC^N)]⁺ species.



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Fig. 6. Proposed reaction mechanism for the Au(I)/Au(III)-mediated arylation-lactonization of alkenoic acids.

To gain mechanistic insight, the stoichiometric reaction of *cis*-**7a-Cl** with $AgSbF_6$ (1.2 eq), 4-pentenoic acid (1.3 eq) and K_3PO_4

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(1.0 eq) was carried out in HFIP at 80 °C. After 16 hours, γ -benzyl- γ -butyrolactone 10 was obtained in 73% yield (Scheme S13). Therefore, it is plausible to propose the formation of an arylgold(III) intermediate formed by an oxidative addition reaction of the $C_{\rm sp2}$ -I bond to Au(I).

Conclusions

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We synthesized two Au(I) complexes bearing bidentate hemilabile MIC^N ligands, [Au^I(MIC^N)CI], **2a** and **2b**, and explored their redox reactivity. The formation of the Au(III) complexes **4a-CI** and **4a-OAc** was achieved by using 2 e⁻ oxidants of the type PhIX₂ (X = Cl, OAc), demonstrating a suitable design of the MIC^N ligand to stabilize Au(III) centers. Next, we showed that effective oxidative addition of a strained C_{sp2}-C_{sp2} bond was obtained when reacting **2b** with biphenylene in the presence of a halide scavenger and heating at 90 °C. In this case, complex **5b** was obtained as the desired ensuing Au(III) species.

On the other hand, the oxidative addition of aryl iodides to Au(I) required a temperature of 120 °C, and under these conditions the reductive elimination reaction of Au(III) species is also favored, yielding a series of triazolium salts (**8a-R/8b-R**, **9a/9b** and **1a/1b**). Although the expected Au(III) complexes were not directly detected, we independently synthesized the analogous Au(III) complex *cis*-7a-Cl, which after heating produced the same triazolium product (**8a-OMe**), thus supporting the oxidative addition/reductive elimination pathway for aryl iodides.

Finally, we combined all the redox knowledge gained with this system to develop a novel reaction for the synthesis of γ -benzyl- γ -butyrolactones upon arylation-lactonization of γ -alkenoic acids. The reaction was catalyzed by **2a** and **2b** (10 mol% catalyst loading), obtaining up to 8 TON. The catalytic activity shown by MIC^N/Au system is significant because it shows that readily available hemilabile C^N ligands are a real alternative to hemilabile P^N ligands, where MeDalphos system stands out among catalytic examples of oxidant-free oxidative addition of aryl halides allowed by the presence of pendant N-groups. Current efforts are devoted to extending the application of these hemilabile [Au¹(MIC^N)CI] complexes to novel catalytic transformations.

Data availability

Data for this work, including experimental procedures, NMR spectra and Crystallographic data, are provide in the ESI.

Author Contributions

P.F., H.V. and G.G.-B. performed the experiments. All authors discussed the results and wrote the manuscript.

Conflicts of interest

There are no conflicts to declare

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