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Boosting Gold(I) Catalysis via Weak Interactions: New Fine-Tunable Impy Ligands

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omogenous gold catalysis has rapidly emerged as an established tool in synthetic organic methodologies.¹⁻⁴ Despite its relatively young age (pioneering studies date back to early 2000), the use of gold(I) species in the electrophilic activation of unsaturated hydrocarbons and cross-coupling reactions has already begun to parallel the use of longer investigated and more consolidated noble metal catalysis. The robustness and adaptability of gold(I) catalysis in multifaceted applications is witnessed daily by its exploitation to enable synthetic methodologies such as photocatalysis, the total synthesis of structurally elaborated bioactive and photoactive molecular motifs, asymmetric catalysis, dual catalysis, electrosynthesis, and flow-chemistry.⁵⁻⁸ This scenario is the consequence of the ongoing request for dedicated gold complexes featuring a predesigned flexible and fine-tunable catalytic performance.⁹ Among others, the use of readily available N-heterocyclic carbenes (NHCs)¹⁰⁻¹³ as ligands in gold chemistry¹⁴⁻¹⁸ deserves a particular mention due to the strong σ -donating properties of the soft singlet carbon atom of NHCs, which results in Au complexes (soft metal) commonly featuring high stabilities and yet unique catalytic activities.^{19,20} Additionally, NHCs can rapidly access impressive organometallic diversity and complexity by incorporating a theoretically unlimited range of functionalities, a key requisite for the development of "on-demand" catalysts. In this wide and dynamic picture, the development of unsymmetrically substituted monodentate NHC ligands is receiving growing credit due to its prompt access to unlimited fine-tunable gold

species and modular stereochemical environments for asymmetric transformations.^{21–23}

The robust and readily accessible imidazo[1,5-a]pyridin-3ylidene platform (ImPy) A (introduced almost simultaneously by Lassaletta²⁴ and Glorius²⁵) covers a prominent role in metal-carbene chemistry,²⁶⁻³³ allowing the installation of steric as well as electronic tools in close proximity to the metal center (i.e., C5-position of the ImPy core). In addition, ImPy shows a higher oxidation resistance and better synthetic accessibility with respect to P-based analogous ligands (Figure 1, top). As a consequence, over the past years fully characterized [Au(I)]- and [Au(III)]-ImPy complexes³⁴⁻³⁶ have been documented with ultimate applications in metal drug chemistry³⁷ and catalysis.³⁸⁻⁴¹ Modulating effects through steric constraints at the C5-position have been extensively investigated; however, the overall impact of electronic parameters have been far less systematically accounted so far and has been limited to the introduction of electron-rich aromatic pendants.42

In the present work, a systematic investigation on a new family of C5-oligoaryl-ImPy gold complexes was carried out,

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Figure 1. Chemical engineerization of fine-tunable C5-oligoaryl–ImPy gold complexes (DIPP, 2,5-diisopropyl-phenyl).

disclosing an unprecedented role of the electron-deficient 3,5- $(CF_3)_2$ -phenyl ring in modulating and enhancing the catalytic activity of cationic gold(I) complexes in several transformations involving carbocationic- or carbene-like organogold intermediates (Figure 1, bottom).¹⁹

In line with our ongoing interest in homogeneous gold catalysis^{43–50} and fine-tunable organometallic adducts via "secondary interactions", ^{51–56} we describe here two series of C5-oligoaryl imidazolium salts **6a–g**, featuring phenyl- (P_n-ImPy) and 2-thienyl- (T_n-ImPy) based pendants (Scheme 1). The role of the electronic properties imprinted by the diversely functionalized π -clouds on the P_n-ImPy and T_n-ImPy carbene series was assessed in three different gold-catalyzed method-

Scheme 1. Synthetic Sequence for the Preparation of P_n and T_n -ImPy·HX 6a-g



ologies by a combined SC-XRD and DFT investigation of the corresponding [Au(I)] complexes.^{57,58}

Imidazopyridinium salts were obtained via initial Suzuki– Miyaura cross-coupling reactions among the pyridyl aldehyde 1 and the corresponding aryl-boronic acids 2a-f. The 6-aryl pyridylcarbaldehydes 3 were isolated in excellent yields (65– 98%) under optimal conditions (Scheme 1a). Then, the corresponding imines 5 were obtained quantitatively under dehydrative conditions (MgSO₄, Et₂O, rt) in the presence of 2,6-(iPr)₂-aniline 4 (Scheme 1b). Finally, the desired imidazolium platforms 6a-f were achieved via a ring-closing strategy in the presence of (CH₂O)_n and TMSX (X = Br, Cl; 74–94%, Scheme 1c). Differently, for the synthesis of the T3-ImPy·HBr 6g, a Pd(0)-catalyzed Suzuki–Miyaura coupling between the Br derivative 7⁴⁰ and the commercially available terthienyl-boronate 8 (69%) was adopted.

Subsequently, the C(5)-substituted imidazo[1,5-a]pyridinium halides 6a-g were subjected to the [Au(I)] complexation (10a-g) via metal metathesis through the corresponding Ag-complexes (9). The protocol (DCM, rt, dark, 16 h) proved highly competent in all cases, yielding the desired air-stable LAuCl complexes in excellent overall yields (81-99%, Scheme 2).

Scheme 2. Two-Step Procedure for the Synthesis of T_n - or P_n ImPy-AuCl complexes 10a-g via a Metal Exchange Protocol of the Silver Analogues 9



Some of us has recently documented the combined role of the cationic [Au(I)] center and basic anions^{59,60} in the chemo-, regio-, and stereoselective dearomatization of 2-naphthols^{61–67} with allenamides,^{68–70} discovering the syner-gistic role of multiple secondary interactions in controlling the mechanistic profile.^{71,72} In this line, we elected the dearomative process of dimethyl-naphthol 12 and allenamide 13 (Scheme 3) as a model control experiment for testing new [ImPyAuCl] complexes 10a–g.⁷³ For these experiments, the corresponding trifluoroacetate complexes 11a–g were generated *in situ* via a chloride scavenging reaction of complexes

Scheme 3. P_n -ImPy-AuCl/AgTFA- and T_n -ImPy-AuCl/AgTFA-Catalyzed Dearomatization of Naphthol 12^{*a*}



^aThe yields are reported as an average of two runs.

From the data collected in the Scheme 3, several interesting conclusions can be drawn. The P_n -ImPyAuCl series provided product 14 from very low to almost quantitative yields (26–95%) with an unexpected correlation to the electronic properties of the pendant group at C5. As a matter of fact, while electron-rich pendants are generally accommodated in the organic ligand to stabilize cationic Au(I) intermediates via stacking interactions,⁷⁴ in our case the incorporation of the 3,5-(CF₃)₂-C₆H₃ unit (11c) resulted as the best catalyst, providing 14 in a 95% yield.

Furthermore, complex **11c** showed the highest initial reaction rate among the complexes tested (**11a–11d**); more precisely, the rate was ca. $2.8 \times$ higher than that of the 3,5-(OMe)₂Ph-containing NHC–[Au(I)] complex **11d** (see the SI for details).

To shed light on this intriguing and unexpected chemical outcome, a dedicated crystallographic investigation was carried out on both series of P_{n^-} and T_n -ImPy-AuCl complexes. Suitable single crystals for the SC-XRD analysis of complexes **10a**-**g** were obtained via vapor diffusion of *n*-hexane in DCM/ toluene solutions of gold adducts. Some representative X-ray structures are depicted in Figure 2 (see the SI for further details).



Figure 2. Representative molecular structures for complexes 10a, 10e, 10b', and 10f. Hydrogen atoms were removed for clarity. The $Ar_{centroid}$ ···Au interaction is shown as black dashed lines. The S_1 ···S₂ interaction (10f) is shown as a red line.

All synthesized [Au(I)] complexes showed a linear bicoordination geometry with the Cl atom *trans* to the NHC ligand and an almost perpendicular orientation of the bulky DIPP unit with respect to the imidazolyl ring (dihedral angles range from 83.84° to 87.79°, Table S11). This common arrangement is a consequence of the rotational restriction faced by the 2,6-diisopropyl substituents. Additionally, intramolecular Au $\cdots\pi$ contacts with the aryl pendants at the C5-position were observed.

In line also with recent findings by Toste and Sigman,⁷⁵ who elegantly disclosed strong structure–catalytic activity correlations in regio-divergent [Au(I)]-catalyzed cycloaddition reactions, we targeted Au–Cl, Au–Cl, and Au—Ar crystallographic distances as potential probe parameters to indirectly

predict or rationalize the catalytic performances of the new ImPy–AuCl complexes. In this study, the cationic $[10b-Au(CH_3CN)](SbF_6)$ complex 10b' (Figure 2) was also taken into account, comparing the influence of the cationic metal center on the crystallographic data.

In Table 1, several structural parameters have been collected for a simplified comparative analysis.

Table 1. Structural Parameters Obtained by SC-XRD for ImPy-AuCl Complexes 10

10	Au–Cl (Å)	Au-C1 (Å)	Au···Ar $(Å)^a$	Ar–ImPy $(^{\circ})^{b}$
10a	2.304(3)	1.983(7)	3.511	56.95
10b	2.2828(9)	1.985(3)	3.412	67.65
10b'	2.019(3)b	1.974(3)	3.384	61.62
10c	2.287(1)	1.982(5)	3.352	64.63
10d	2.2765(9)	1.980(2)	3.353	64.94
10e	2.277(2)	1.981(7)	3.526	57.52
10f	2.267(2)	1.983(6)	3.477	82.58
10g	2.2932(6)	1.970(3)	3.456	73.20

"The distance between [Au(I)] and the center of mass of the aromatic ring at C5, as calculated by Mercury. "The Au–N(ACN) distance.

Interestingly, the Au–C1 distances were scarcely affected by the electronic properties of the ligand (Au–C1 range of 1.970–1.983 Å). The truncated conjugation at C5 between the ImPy core and the aryl ring could be responsible for this trend. The large dihedral angles recorded in the solid state for the two aromatic systems (Ar and ImPy, 56.95–82.58°) are in good agreement with this hypothesis. Similarly, the Au–Cl distances (2.267–2.304 Å) showed only slight variations. In particular, while almost constant Au–Cl distances were observed in the P_n -ImPyAuCl series (**10a–d**), the T_n -ImPyAuCl family shows shorter distances for mono- and bithienyl substituents (**10e** and **10f**) but a longer distance in the case of the *tert*-thienyl pendant (**10g**).

Therefore, Au…Ar contacts were examined across the two series P_n and T_n . First, by increasing the number of aryl units in both series, closer intramolecular contacts were identified. This trend nicely correlates with the electron-richness of oligoaryl cores that increases along with the number of repeating units. Second, a comparison between the 10b' and 10b (entries 2 and 3, Table 1) revealed only a slight contraction in both the C1-Au bond and the Au-Ar contact for the cationic complex, supporting the suitability of the Au-Cl precursors 10a-g for the structure-activity relationship investigation. Interestingly, the introduction of arenes featuring opposite electronic properties such as $3,5-(OMe)_2$ -Ph (10d) and $3,5-(CF_3)_2$ -Ph (10c) caused a shortening of the Au…Ar distances (3.353-3.352 Å) in both cases with respect to unsubstituted 10a (3.511 Å). However, a drastic change in the catalytic outcome was highlighted in the 10a-c series (95% yield with 10c and 26% yield with 10d).

Concerning the T_n -ImPyAuCl series, no direct S(thienyl)... Au contacts have emerged regardless the length of the oligothienyl.^{49,76} Surprisingly, **10f** showed a quite uncommon *cis*-arrangement of the two thienyl sulfur atoms in the solid state (Figure 2, S1...S2 = 3.301 Å).⁷⁷ This feature can be attributed to the fact that the S1...S2 forces play a stabilizing role in the packing efficiency.⁷⁸ On the contrary, complex **10g** bearing a terthienyl chain at C5 shows the conventional *transtrans*-arrangement of the thienyl units (see the SI). The uncommon cis-arrangement observed for the T2 side arm could be tentatively targeted to rationalize the unsatisfactory catalytic performance of 10f-AgTFA (Y = 22%).⁷⁹ Differently, T1 and T3 bearing gold complexes worked smoothly in the dearomative process, delivering 14 in 87% and 74% yields, respectively. The slight drop in the chemical outcome for 10g might be ascribed to the more electron-rich T3 and consequent "shielding" of the electrophilic properties of [Au(I)] center (Au···Ar = 3.536 and 3.456 for 10e and 10g, respectively). Additionally, the photophysical properties of the T3-ImPy series (6g and 10g) were investigated both in solution (DCM) and in solid state (see Figures S1 and S2 and Table S4). Here, a significant drop in the emission quantum yield of the T3 salt 6g (6.2%) was recorded when the metal center was introduced (0.2%, 10g). This phenomenon could be tentatively assigned to a higher efficiency of the nonradiative intersystem crossing deactivation of the fluorescent excited state promoted by the heavy atom effect.

To further analyze the electronic nature of such Au-Ar interactions and to provide a visual representation of them, we resorted to electronic structure calculations using density functional theory on complexes 10a, 10c, and 10d.^{80,81} Geometry optimizations and a topological analysis of the obtained electron density was performed using the atoms in molecules (AIM) and non-covalent interactions (NCIs) models.^{82,83} Calculations were performed at the B3LYP/6-31+G(d) level for main-group atoms, and the SDD basis set and electron core potential were used for the Au atom.^{84,85} We resorted to this compact basis set due to the large size of the complexes and the previous systematic benchmark showing that this particular combination of density functional and basis set, although compact, was able to reproduce gold chemistry with a comparable accuracy to triple- ζ and even larger basis sets.^{86,87} Actually, our calculations faithfully reproduced the experimentally recorded X-ray structures. Not only close steric contacts were observed from the NCIs analysis (Figure 3,



Figure 3. Ring and bond critical points of the electron density of **10c** (left) and the close noncovalent Au $\cdot\cdot\pi$ -aryl steric contact (right).

right) but a full covalent bond-like topology was also detected under the AIM analysis. A bond critical point can be found between the gold atom and the aryl fragment, along with the expected ring critical point to fulfill the Poincaré–Hopf rule. Therefore, both analyses point toward intimate Au…aryl interactions in these complexes (Figure 3, left).

Further support for this interpretation was gathered via a population analysis of the electron density calculated for intermediate A when the aryl C5-pendant was undecorated and also when it was *meta*-disubstituted with CF_3 and OMe groups (Scheme 4). The partial charges found at the electrophilic end of the allenamide framework in these intermediates are fully compatible with the view of increased reactivity due to the electron-withdrawing effects of the C5-

Scheme 4. Proposed Reaction Intermediate A and Its Natural Bond Orbital (NBO) Partial Charge at the Electrophilic CH_2 Terminus with Respect to the C5-Pendant Substitution



pendant when substituted with CF₃. Gratifyingly, charge values at this reactive terminus (0.037, 0.035, and 0.041 au for the -OMe-substituted, unsubstituted, and $-CF_3$ -substituted systems, respectively) correlate well with the observed experimental yields.

In addition, the marked deviation from planarity recorded for the NHC and aryl skeletal unit (see Table 1) precludes a major impact of the arene electronic features on the whole ImPy carbene fragment and elects the Ar…[Au] secondary contacts as responsible for the fine-tuning of the catalytic performance.

Finally, we decided to assess the extensibility of this unprecedent accelerating effect of the CF₃-containg ring to other [Au(I)]-catalyzed manipulations of unsaturated hydrocarbons. In this regards, two chemical transformations based on carbocationic- or carbene-like organogold intermediates,¹⁹ namely olefin cyclopropanation⁸⁸ and the cyclization of alkynylbenzaldehyde,⁸⁹ were analyzed (Scheme 5).

Scheme 5. Olefin Cyclopropanations and Alkynylbenzaldehyde Cyclizations: Examples of the Accelerating Effect of CF₃-Containing Carbene Ligands in [Au(I)] Catalysis



In this regard, the intermolecular olefin cyclopropanation introduced by Toste was first performed in the presence of complexes $10a,c-d/AgSbF_6$ (Scheme 5a). Interestingly, a marked increase of the chemical outcome toward the formation of the cyclopropane *cis*-17 (yield up to 88%) was observed when fluorinated complex 10c was adopted. The same trend was recorded also in the cyclization of the alkynylbenzaldehyde 18 in the presence of isopropanol. Here, $10c/AgNTf2_6$ (5 mol %) provided the acetal 19 in a 68% yield, more than fivefold that obtained with the electron-rich complex 10d (Scheme 5b). Finally, it is worth mentioning that this reactivity trend was not affected by the loading of the gold complex. As a matter of fact, when catalysts 10 were employed in a 2.5 mol % loading, a similar chemical outcome relationship was recorded. 90

In conclusion, we have documented the preparation, characterization (spectroscopic and crystallographic), and use in catalysis of a new class of ImPyAuX complexes featuring diversely substituted aromatic pendants at the C5-position. The impact of electronically different side-arms on the overall catalytic properties of the gold complexes (*i.e.*, dearomatization of naphthols with allenamides) was examined through dedicated kinetic experiments, revealing an unusual accelerating action exerted by the $3,5-(CF_3)_2$ -Ph unit. Activating secondary interactions exerted by the ligand C5-pendant on the gold–alkylidene intermediate were accounted for the experimental outcome that resulted effective in several Aumediated electrophilic activations of unsaturated hydrocarbons.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsorginorgau.1c00052.

General synthetic procedures; photophysical, crystallographic, and computational data; and NMR spectra (PDF)

Accession Codes

CCDC 2091513–2091520 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/cif, or by emailing data_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.

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Notes

The authors declare no competing financial interest.

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