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Investigating chemical diversity: o-propargylphenols as key compounds in the divergent synthesis of 2-substituted benzofurans and chromenes†

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In this study, we explored and optimized a MW-enhanced divergent approach for the synthesis of 2-substituted benzofurans and chromenes, starting from seventeen substituted o-propargylphenols characterized by a monoaryl substitution on the propargylic sp³ carbon. Firstly, we developed a robust platform for the preparation of a library of o-propargylphenols. Under basic conditions, o-propargylphenols reacted regioselectively to yield benzofurans in yields ranging from 43% to 100%. Conversely, under cationic gold catalysis, we were able to obtain the corresponding 4*H*-chromenes, albeit in more variable yields (from 25% to 93%) and slightly lower regioselectively. We also proposed plausible mechanisms to explain the divergent outcomes observed. Our findings underscore the potential of diversity-oriented synthesis in the investigation of molecular complexity. Our neglected o-propargylphenols have proven to be versatile and strategic starting materials for accessing oxygen-containing heterocyclic scaffolds through intramolecular cyclization reactions.

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Introduction

Small organic molecules can exert powerful effects on the functions of macromolecules that compose and regulate the living systems. Synthetic organic chemists can access these compounds by exploiting different approaches. Target-Oriented Synthesis¹ (TOS) is the approach of choice when the aim is to prepare small molecules with predetermined macromolecule-perturbing properties. For example, a natural compound can be extracted from natural matrices, purified, and then structurally characterized. Once its activity has been well-defined, it can become a target for a chemical synthesis. Thus, the objective of TOS is to prepare a molecule able to access a *precise* region of the chemical space of the macromolecular target. For example, TOS is the key strategy in total synthesis studies.

A similar approach, aware of medicinal and combinatorial chemistry, aims to explore a *dense* region of the chemical space in proximity to a region known to have useful properties.

The template for the identification of lead compounds may be a natural product, a known drug, or an in-silico-designed structure developed from a mechanistic hypothesis and/or a crystal structure determination of the macromolecular target of interest. This strategy can also be considered a different sort of TOS.

However, as pointed out by Schreiber and Buke in their pivotal review,² despite these approaches have led to great advances in the chemical and life sciences fields, a question remains unanswered: “*Are the regions of chemistry space defined by natural products and known drugs, which have been so intensely scrutinized to date, the best or most fertile regions for discovering small-molecules that modulate macromolecular function in useful ways?*”

The first step to answer this question is to recognize that the problem of exploring broader regions of the chemical space is different than the problem of accessing precise or dense regions, so different chemical tools and distinct solutions are necessary. Diversity-Oriented Synthesis³ (DOS) is a strategy that allows obtaining a broad distribution of different compounds, resulting in a wider exploration of the chemical space, generating, for example, skeletal diversity. Two main approaches allow the generation of skeletal diversity; the first is the Reagent-Based Approach, which involves the use of different reagents to transform a common substrate with the potential for diverse reactivity into structurally diverse molecules (Fig. 1A). The second, called Substrate-Based Approach,

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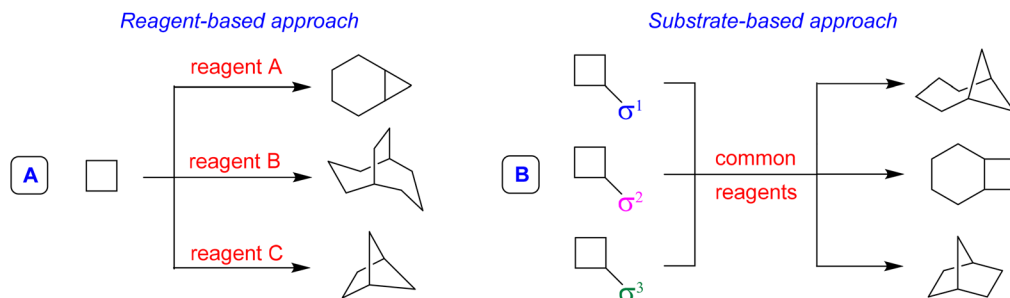


Fig. 1 Two general approaches for generating skeletal diversity.

foresees the transformation of different substrates characterized by the presence of different substituents that pre-encode skeletal information (called σ elements) into a collection of products having distinct molecular skeletons by using common reaction conditions (Fig. 1B).

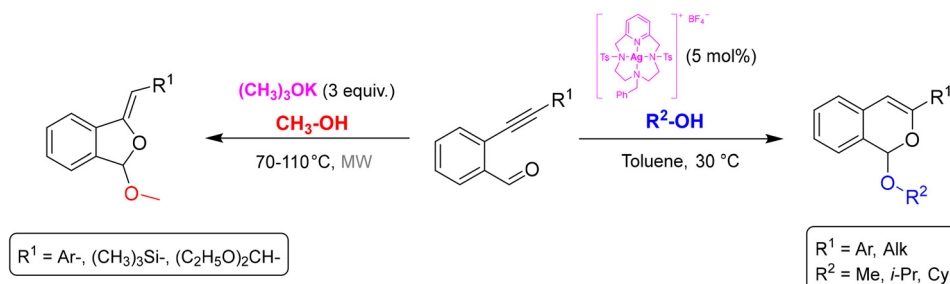
The Reagent-Based Approach (A)⁴ is probably the most used tool in DOS to achieve skeletal diversity.

An example of the application of the reagent-based approach (A) from our research group is the divergent synthesis of isobenzofurans and isochromenes starting from 2-alkynylbenzaldehydes and alcohols. By changing the reaction conditions, it was possible to selectively obtain the two different regioisomeric heterocycles. Isobenzofurans have been obtained in good yields by a base-promoted domino nucleophilic addition/5-*exo-dig* cyclization sequence, under microwave heating at 70–110 °C (Scheme 1, left).⁵ Starting from the same substrates, the reaction catalyzed by an original [Ag(I)(Pc-L)] complex at 30 °C resulted in the selective 6-*endo-dig* cyclization with formation of the regioisomeric isochromene derivatives (Scheme 1, right).⁶

Based on these premises, and in connection with the research interests of our group in diversity-oriented synthesis and discovery of novel strategies for the preparation of heterocycles starting from arylalkynes bearing a proximate nucleophile,⁷ in this work we developed a divergent approach for the regioselective synthesis of 3-unsubstituted-benzofuran and 4-unsubstituted-chromene nuclei starting from different 2-propargylphenols (Scheme 2).

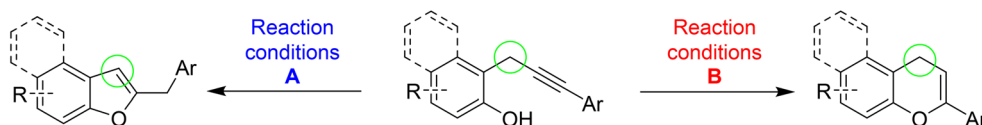
Simple 2-propargylphenols, *i.e.* characterized by the presence of a monoarylsupstitution on the propargylic sp^3 carbon,

have been selected as strategic starting material for the following reasons: (1) their chemistry has not received enough focus,⁸ (2) their structure displays great potential as versatile precursors for different reactions and cyclization paths, (3) the nucleophilic attitude of phenol oxygen is modulable by changing the reaction conditions, (4) cyclization products could represent relevant nuclei in the field of biologically active compounds. In the literature, there are some examples of base-promoted cyclization of propargylphenols to give the corresponding benzofurans through a 5-*exo-dig* cyclization.⁹ On the other hand, only one example of gold-catalyzed¹⁰ 6-*endo-dig* cyclization of propargylphenol to give the corresponding six-member oxygenated heterocycles has been reported, and regarding other metal catalysts, another isolated example describes this cyclization promoted by a ruthenium-based catalyst.¹¹ Interestingly, should be underlined that in all the above-mentioned examples the starting propargylphenols are always characterized by a double substitution on the propargylic sp^3 carbon. These compounds are strongly different from our propargylphenols regarding both the synthesis and the reactivity. Conversely, to the best of our knowledge, starting from simple monoarylsupstituted 2-propargylphenols such as those used in this study, there are only a few examples of synthesis of simple oxygen-containing heterocycles such as dihydrobenzofurans,¹² whereas no example regarding the preparation of benzofurans and chromenes. Thus, in this work, three main essays have been developed: (a) the optimization of a general and robust protocol to generate a library of differently substituted 2-propargylphenols, (b) the development of a selective approach to 3-unsubstituted-benzofurans



Scheme 1 Divergent synthesis of isobenzofurans and isochromenes from 2-alkynylbenzaldehydes.





Scheme 2 This work: divergent synthesis of benzofurans and chromenes from 2-propargylphenols.

and (c) the development of a divergent approach to isomeric 4-unsubstituted-chromenes. In this paper, we describe the results of our efforts.

Results and discussion

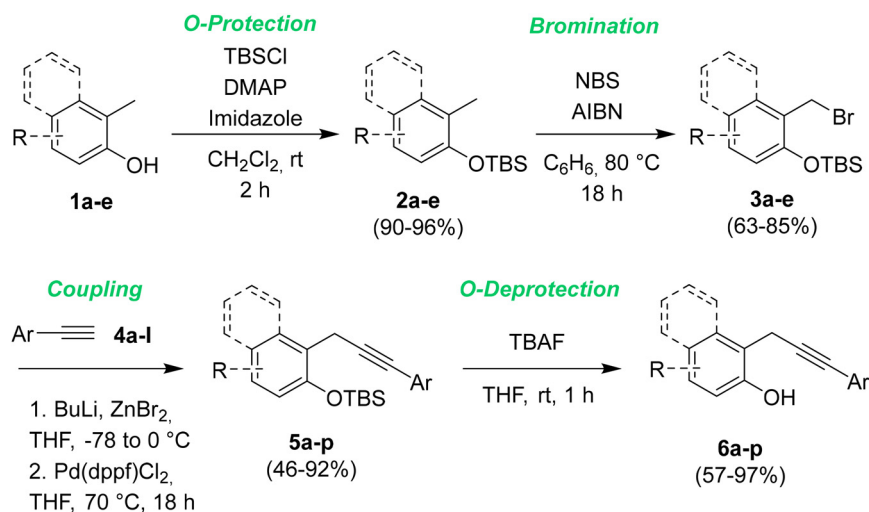
Firstly we prepared a library of sixteen 2-propargylphenols **6a–p**, characterized by the presence of different ED and EW groups in different positions on both the arene moieties. The general synthetic pathway is described in Scheme 3.

The protection of phenol derivatives **1a–e** was obtained by the reaction of *o*-cresols **1a–e** with *tert*-butyldimethylsilylchloride (TBSCl) in the presence of 4-dimethylaminopyridine (DMAP) and imidazole in DCM at rt. By this approach, we were able to synthesize five different substituted *tert*-butyldimethyl(2-tolyloxy)silanes **2a–e** in excellent yields, which were brominated by NBS and AIBN in refluxing benzene for 18 h to give intermediates **3a–e** in very good yields.¹³ The cross-coupling of compounds **3a–e** with different terminal alkynes **4a–l** has been optimized based on a literature procedure reported by Negishi.¹⁴ The alkynylzinc bromide intermediates have been obtained by treatment of **4a–l** with butyllithium and ZnCl₂. Next, a Pd(dppf)Cl₂ catalyzed cross-coupling with (2-(bromomethyl)phenoxy)(*tert*-butyl)dimethylsilanes **3a–e** in refluxing THF overnight gave the sixteen protected *o*-propargyl phenols **5a–p** in yields ranging from 40 to 91%. The following deprotection¹⁵ by treatment with TBAF in THF at rt gave the desired 2-propargylphenols **6a–p** in yields

ranging from good to excellent. 4-Amino-2-(3-phenylprop-2-yn-1-yl)phenol **6q** was obtained by the chemical reduction of **6p** employing iron powder in water (see ESI† for detailed procedures and full characterization data).

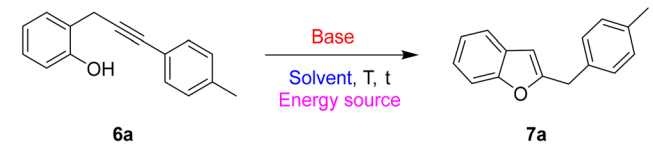
The screening of the most favourable reaction conditions for the base-promoted cyclization of 2-propargylphenols was performed using 2-(3-(*p*-tolyl)prop-2-yn-1-yl)phenol (**6a**) as a model substrate. The reactions were performed on a 0.2 mmol scale testing different organic and inorganic bases, in different solvents and modifying the reaction temperature and the energy source. Table 1 displays a selection of the most representative results of this screening.

We started the study by using already established conditions for related base-promoted cyclizations, *i.e.* DMF as the solvent and cesium carbonate as the base.¹⁶ After overnight stirring at rt the starting material **6a** was quantitatively recovered unreacted (Table 1, entry 1). By changing the solvent (acetonitrile) and raising the reaction temperature to 70 °C, the desired product **7a** was obtained in an encouraging 46% yield (Table 1, entry 2). Then, we tested the activity of some other organic and inorganic bases (Table 1, entries 3–5); potassium carbonate was demonstrated to be the base of choice, yielding the desired product in 75% yield (Table 1, entry 3). Next, we evaluated the effect of the solvent (Table 1, entries 6–9), discovering that the presence of a polar aprotic media is mandatory for the success of the reaction. So, while the reaction in dichloromethane and THF failed (Table 1, entries 6 and 7), and a polar protic solvent such as methanol gave very poor results (Table 1, entry 8), dimethyl sulfoxide gave an



Scheme 3 General synthetic path to 2-propargylphenols **6**.



Table 1 Screening of the optimal reaction conditions for base-promoted cyclization of propargylphenols


Entry	Solvent	Base (10 mol%)	T (°C)	Energy source	t	7a yield ^a (%)	6a rec. (%)
1	DMF	CsCO ₃	RT	Oil bath	16 h	—	Quant. ^b
2	CH ₃ CN	CsCO ₃	70	Oil bath	16 h	46	—
3	CH ₃ CN	K ₂ CO ₃	70	Oil bath	16 h	75	—
4	CH ₃ CN	KOH	70	Oil bath	16 h	35	—
5	CH ₃ CN	TEA	70	Oil bath	16 h	—	Quant. ^b
6	DCE	K ₂ CO ₃	70	Oil bath	16 h	—	Quant. ^b
7	THF	K ₂ CO ₃	70	Oil bath	20 h	—	Quant. ^b
8	MeOH	K ₂ CO ₃	70	Oil bath	24 h	12	76
9	DMSO	K ₂ CO ₃	70	Oil bath	2 h	79	—
10	DMF	K ₂ CO ₃	70	Oil bath	4.5 h	82	—
11	DMF	K ₂ CO ₃	90	Oil bath	1 h	93	—
12	DMF	K ₂ CO ₃	90	MW	30 min	64	—
13	DMF	K ₂ CO ₃	90	MW	10 min	83	—
14	DMF	K ₂ CO ₃	90	MW	5 min	83	—
15	DMF	K₂CO₃	70	MW	5 min	89	—

^a Referred to pure isolated product. ^b TLC analysis displays the presence of a single spot referred to unreacted starting material **6a**; this result is also confirmed by ¹H-NMR spectra of the reaction crude.

interesting 76% yield of the desired product in 2 hours (Table 1, entry 9). The best results in terms of yield, time, and cleanness of the reaction have been obtained with DMF (82% yield) in 4.5 hours (Table 1, entry 10). Yields could be further increased by raising the reaction temperature to 90 °C; under these conditions, the reaction was complete in only 1 hour with a satisfactory 93% yield (Table 1, entry 11). The promising best reaction conditions (polar aprotic solvent and heating) seemed to be ideal for testing the approach under microwave heating¹⁷ (entries 12–15). It is well recognized that MAOS (Microwave Assisted Organic Synthesis) can lead to some important advantages, such as a reduction of reaction times and by-product formation. In the transformation under study, the efficiency of dielectric heating allowed a dramatic drop in reaction times and a slight reduction in reaction temperature. Under dielectric heating, the best results were obtained in a very short time (5 min) at 70 °C (Table 1, entry 15), hence with an overall gain in terms of sustainability of the process.

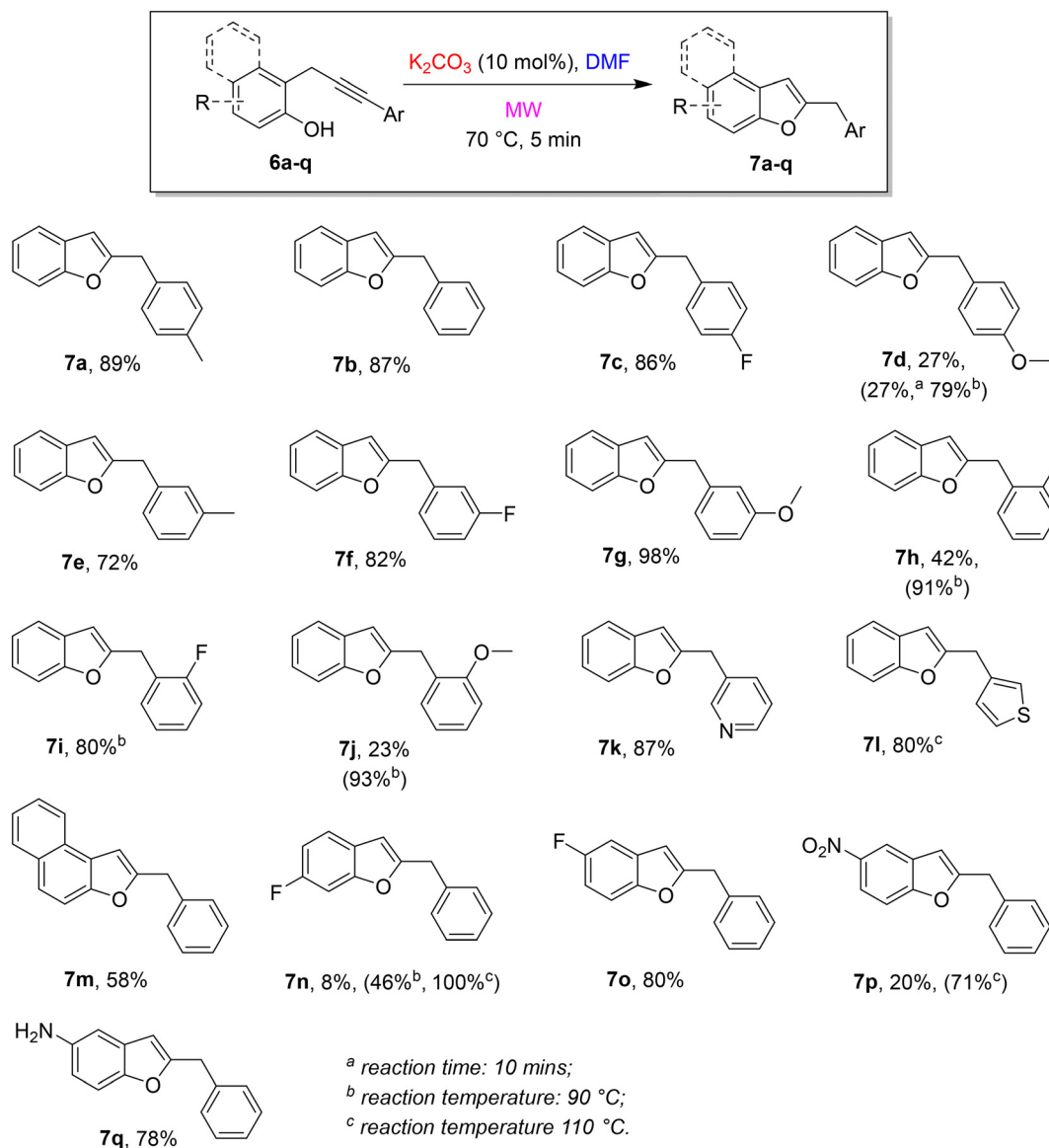
With optimal reaction conditions in hands, the scope and limitations of the approach were explored by changing the substitution on the terminal alkyne and the phenol moiety. The results are summarized in the following Scheme 4.

We made a systematic study with different substitutions on the aryl group at the alkyne terminus. Steric and electronic features have been investigated by changing the electronic nature of the substituent (neutral, electron-donating, and electron-withdrawing) and its position (*para*, *meta*, and *ortho*) on the phenyl ring. The approach was demonstrated to be robust and high-yielding, despite the different substitutions on the aryl group at the alkyne terminus. In the presence of a neutral or an EW group, the cycloisomerization gave the corresponding

benzofurans in very good yields under the standard conditions (**7a**, **7b**, **7c**, **7e**, **7f**, **7g**). Conversely, an ED group on the aryl group in a conjugate position resulted in poor yields (see **7d** and **7j**). This behaviour is probably related to the effect of the electronic nature of the substituent on the electrophilicity of the distal sp carbon involved in the cyclization.¹⁸ In these cases, a rise of the reaction time to 10 minutes at 70 °C did not have any beneficial effect, whereas an increase of the temperature to 90 °C resulted in a dramatic increase in yields (**7d**). A related behaviour was observed also in the presence of a heteroaryl substituent on the alkynyl terminus: an electron-poor heterocycle such as the 3-pyridyl group did not affect the reaction under standard conditions (**7k**, 87%), on the contrary, to obtain good yields in the presence of an electron-rich heterocycle such as 3-thiophenyl, the reaction temperature should be raised to 110 °C (**7l**, 80%). Finally, we tested some modifications on the phenol unit. The shift from *ortho*-propargyl phenol to 1-propargyl-2-naphthol was well-tolerated (**7m**). The results in the presence of a fluorine atom on the phenol moiety depend on the position on the ring (**7n** and **7o**) and an increase of the reaction temperature was required when the fluorine atom is in *meta* to the hydroxy group. A nitro group in *para* to the hydroxy group required a higher temperature to reach a satisfying yield (**7p**) whereas the amino group in the same position was tolerated under standard reaction conditions (**7q**).

Next, to develop a divergent synthesis starting from the same starting materials, we explored the possibility of preparing the regioisomeric chromenes through a metal-catalyzed 6-*endo dig* cyclization. Also in this case, the **6a** was chosen as the model compound and we tested different catalysts, solvents, and conditions to obtain the best reaction results (Table 2). All





Scheme 4 Scope and limitation in the synthesis of benzofurans.

the reactions were conducted with 0.2 mmol of **6a**, 5 mol% loading of the metal catalyst in 2 mL of anhydrous solvent ($c = 0.1$ M), under a nitrogen atmosphere.

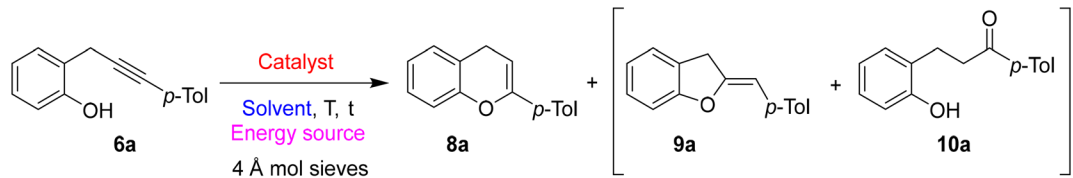
The first reaction was performed in the presence of JohnPhosAuNTf₂ (5 mol%) in DCE at rt (Table 2, entry 1). However, after 24 hours, we obtained only a 21% yield of the desired product **8a** beside a series of by-products and a 13% yield of the 2-hydroxydihydrochalcones **10a** probably arising from the hydrolysis of **8a** triggered by adventitious water in the reaction mixture. To overcome the formation by-product **10a**, 4 Å molecular sieves were added to the reaction mixture, and to increase the yield the temperature was raised to 70 °C (Table 2, entry 2). Under these conditions, the desired product **8a** was obtained in 75% yield beside a small amount of the corresponding 2,3-dihydrobenzofuran with exocyclic double bond **9a**.

Other metal catalysts were tested; however, neither Cu(i), Ag(i), Fe(III), or Au(III) salts gave any positive results, and the starting material was recovered unreacted after 24 hours of reaction at 70 °C (Table 2, entries 3–6).

We next screened other cationic gold(i) catalysts (*i.e.* Ph₃PAuNTf₂, IPrAuNTf₂, and P(OAr)₃AuNTf₂; Ar = 2,4-di-*tert*-butylphenyl) were tested (Table 2, entries 7–9). In all cases, the results were inferior to JohnPhosAuNTf₂, moreover using IPrAuNTf₂ a small amount of regioisomeric benzofuran **7a** was observed (Table 2, entry 8).

Three different solvents were used in the presence of JohnPhosAuNTf₂ as the catalyst at 70 °C. The use of toluene resulted in a slight decrease in the reaction yield with an increase in the formation of isomeric benzofuran compound **7a** (Table 2, entry 10). Conversely, the use of THF increased the conversion rate but strongly reduced the regioselectivity, giving



Table 2 Screening of the optimal reaction conditions for metal-catalyzed cyclization of propargylphenols


Entry	Solvent	Catalyst (5 mol%)	T (°C)	Energy source	t	8a yield ^a (%)	9a yield ^a (%)	10a yield ^a (%)
1	DCE	JohnPhosAuNTf ₂	RT	Oil bath	24 h	21 ^b	—	13
2	DCE	JohnPhosAuNTf ₂	70	Oil bath	2 h	75	5	—
3	DCE	AgSbF ₆	70	Oil bath	24 h	NR ^c	—	—
4	DCE	CuBr	70	Oil bath	24 h	NR ^c	—	—
5	DCE	Fe(OTf) ₃	70	Oil bath	24 h	NR ^c	—	—
6	DCE	NaAuCl ₄	70	Oil bath	24 h	NR ^c	—	—
7	DCE	Ph ₃ PAuNTf ₂	70	Oil bath	24 h	26	—	—
8	DCE	IPrAuNTf ₂	70	Oil bath	3 h	31 ^d	—	—
9	DCE	P(OAr) ₃ AuNTf ₂	70	Oil bath	2 h	47 ^e	7	—
10	Toulene	JohnPhosAuNTf ₂	70	Oil bath	2 h	68 ^f	—	—
11	THF	JohnPhosAuNTf ₂	70	Oil bath	2 h	60	27	—
12	DMF	JohnPhosAuNTf ₂	70	Oil bath	1 h	52 ^e	—	—
13	DCE	JohnPhosAuNTf ₂	70	MW	15 min	61	6	—
14	DCE	JohnPhosAuNTf ₂	85	MW	15 min	68	5	—
15	DCE	JohnPhosAuNTf ₂	100	MW	15 min	79	9	—

^a Referred to pure isolated product. ^b The reaction was performed without 4 Å MS. ^c Starting material was almost quantitatively recovered. ^d 15% of 7a was obtained. ^e Beside unidentified by-products. ^f 10% of 7a was obtained.

rise to a huge amount of the regioisomeric 2,3-dihydrobenzofuran 9a (Table 2, entry 11). The more polar DMF allowed to recover the regioselectivity of the reaction but with modest yields (Table 2, entry 12).

We also tried to shift from traditional to dielectric heating to reduce the reaction time and increase yields. Firstly, the reaction was performed at 70 °C for 15 minutes giving rise to a good 61% yield of 8a and 6% yield of 9a (Table 2, entry 13). Increasing the temperature to 85 °C a slight increase of the yield of 8a to 68% was observed (Table 2, entry 14) beside a reduced amount of by-product 9a. Finally, a reaction temperature of 100 °C allowed to obtain the best result in terms of conversion (88%) and yield of desired 8a (79%), with only a little increase of by-product 9a (Table 2, entry 15).

Thus, with optimal reaction conditions in hands, the scope and limitations of the divergent approach were explored (Scheme 5).

Different substituents showed different effects based on their positions and steric/electronic properties. However, it is difficult to find a rationale to describe accurately the differences in yields and selectivity. The substitution in the *para* position of the phenyl at the alkyne terminus gave rise to the desired chromenes 8a,c,d in very good yields and selectivity (isomeric dihydrobenzofurans 9 absent or <10% yield). Shifting to the substitution in *meta* position, the methyl group allowed the formation of 8e in 82% yield beside a 10% of 9e, while both ED and EW groups caused a decrease in the yields (8f 39%, and 8g 46%, respectively). Moreover, the regioselectivity of the reaction of 6f is one of the poorest, giving rise to the formation of 9f in 35% yield. When the phenyl ring at the alkyne terminus is substituted at the *ortho* position, the outcomes strongly depend on

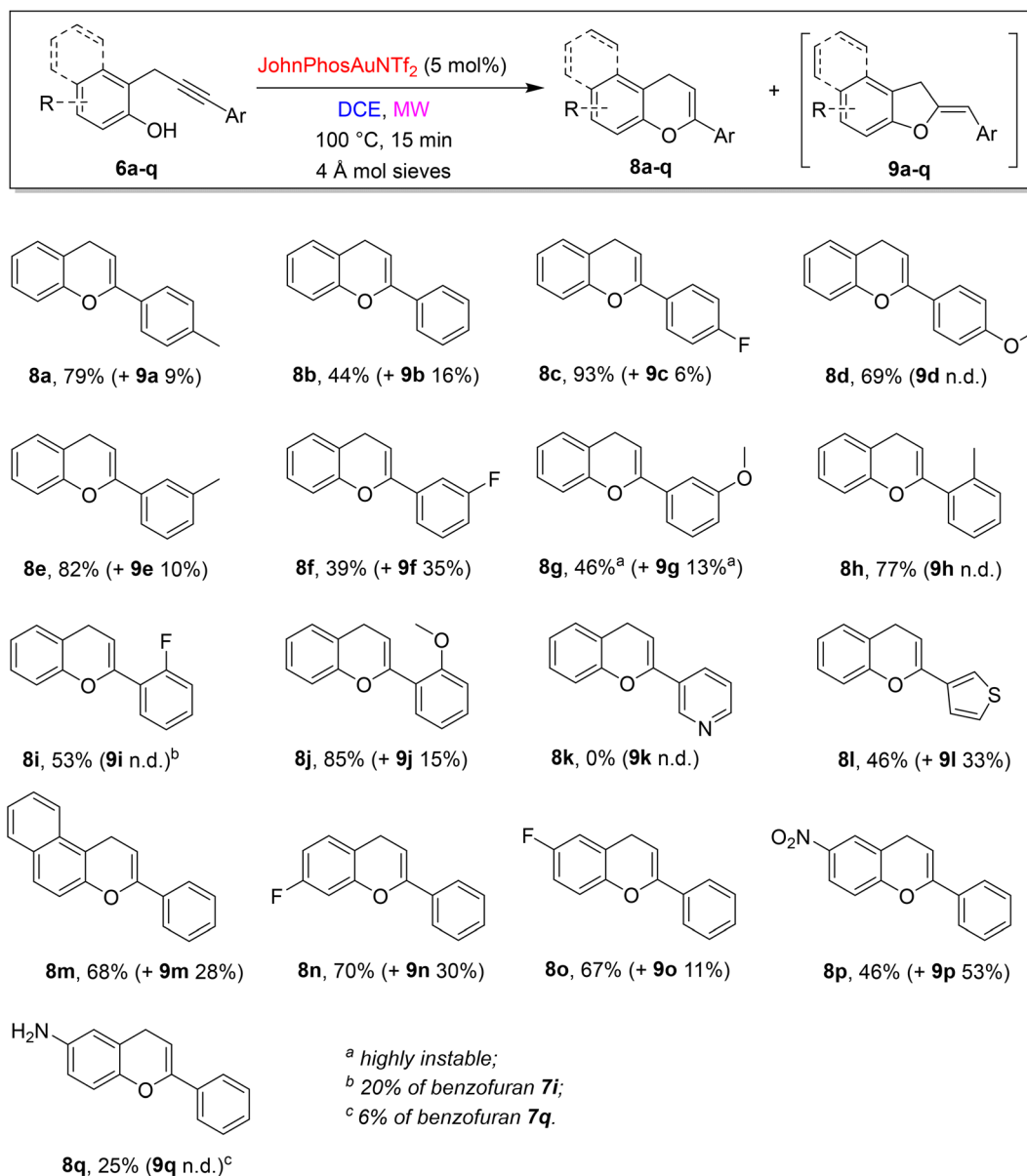
the electronic feature of the substituent: EWG showed worse results than EDG, but in general, the selectivity is high (8h–j). The substitution of the aryl ring with a heteroaryl moiety gave on the whole modest results: propargylphenol 6k, bearing a pyridyl 3-moiety on the alkyne terminus was unreactive and the starting material was fully recovered, while the presence of a 3-thiophenyl substituent induces the formation of the corresponding chromene derivative 8l in a moderate 46% yield, besides a consistent 33% yield of 9l. The failure of the reaction of 6k could be ascribed to a plausible inactivation/complexation of the gold catalyst from the pyridine nitrogen.

The investigation of the scope and limitations proceeded with modifications on phenol moiety. The reaction of 1-propargyl-2-naphthol 6m was high-yielding but poorly selective and led to the formation of a mixture of 8m and 9m in 68% and 28% yield, respectively. A similar behaviour was observed starting from fluorine-containing propargylphenols 6n, whereas a simple change in the position of the fluorine on the phenol ring (6o) resulted in a slight reduction of the amount of dihydrobenzofuran by-product formed (9o).

A reversed regioselectivity was obtained with 6p, bearing a nitro group, with the formation of 9p as the main product (53% yield) and 8p in only 46% yield. Finally, the presence of the amino group showed a quite high selectivity, being 8q the main product obtained, but a low reaction yield (25%), due to the presence of some unidentified by-products in the reaction mixture. The *Z* stereochemistry of dihydrobenzofurans 9 was determined through a NOESY experiment on product 9p and extended to the entire series by analogy (see ESI† for details).

Taking into account the previous literature findings and based on the experimental results, plausible reaction mecha-





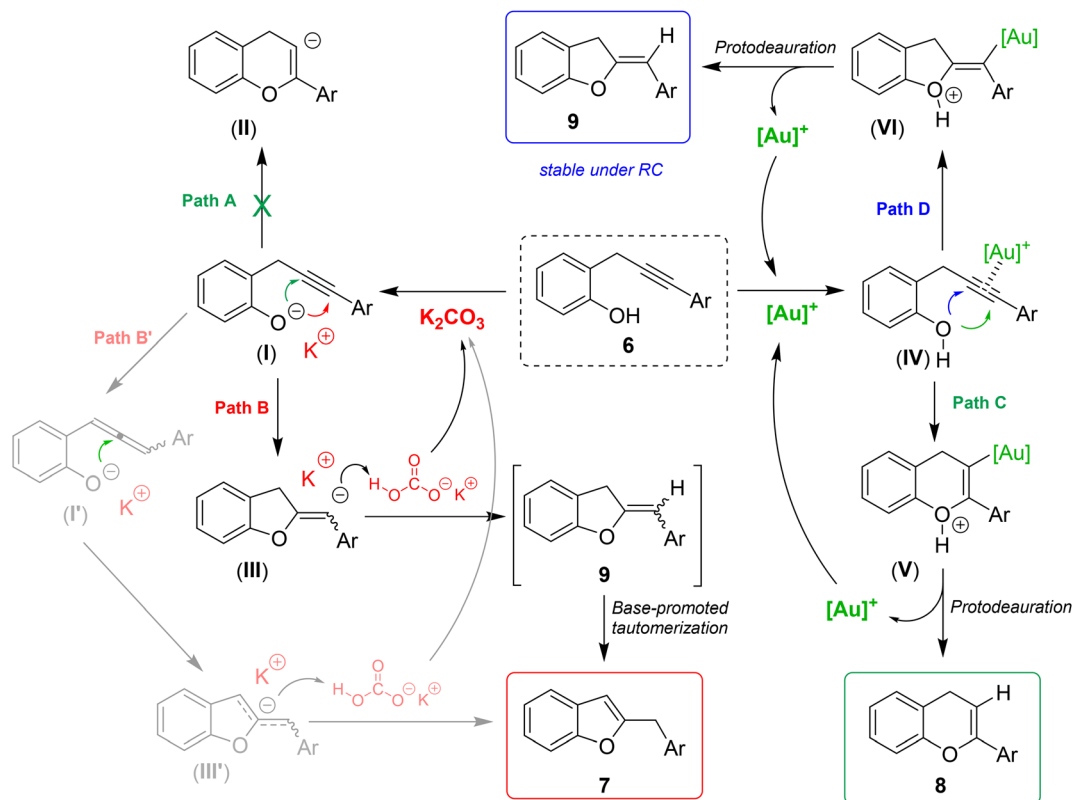
Scheme 5 Scope and limitation in the synthesis of chromenes.

nisms for the divergent formation of isomeric benzofurans 7 chromenes 8 and dihydrobenzofurans 9 are proposed in Scheme 6.

The formation of benzofurans 7 can be justified as follows: potassium carbonate ($pK_b = 3.75$) is a base strong enough to partially deprotonate the 2-propargylphenol 6 ($pK_a \cong 10$) to give the phenoxide intermediate **I**. The latter can theoretically follow two cyclization paths. Path A involves a 6-*endo-dig* cyclization to give a 4*H*-chromene anion intermediate (**II**), whereas path B, through a 5-*exo-dig* cyclization mode, provides a 2-methylene-2,3-dihydrobenzofuran anion (**III**). As already observed in similar base-promoted cyclizations,¹⁹ the 5-*exo-dig* mode is preferred, probably because the alkenyl α -anion (**III**) is stabilized by resonance from the conjugated aryl group. Next, potassium bicarbonate, arising from the phenol deprotona-

tion, can provide the proton to give the neutral 2-methylene-2,3-dihydrobenzofuran derivative 9, which under the basic reaction conditions undergoes a quick base-mediated tautomerization with formation of the aromatic benzofuran nucleus 7 stabilized by resonance. We cannot exclude *a priori* an alternative mechanism that involves the base-promoted formation of an allene intermediate (**I'**),²⁰ which could evolve by intramolecular nucleophilic attack of the oxy anion to the central carbon of the allene to a delocalized benzylic/allylic anionic intermediate (**III'**) and then directly to benzofuran 7 by protonation (Path B'). However, this alternative path seems to be more unlikely because, as already reported by Arai and Shioiri in 2000, moderate bases such as K₂CO₃ are quite ineffective in efficiently promoting such transformation on these systems.²¹





Scheme 6 Proposed divergent reaction mechanisms.

The formation of chromenes involves the activation of the triple bond by the cationic gold to give the π -complex (IV).²² The complexation with the metal enhances the electrophilic properties of C_{sp} carbons that undergo a nucleophilic attack from phenolic oxygen through a 6-*endo-dig* or a 5-*exo-dig* mechanism to give oxo-cationic intermediates (V) (Path C)²³ and/or (VI) (Path D), respectively. The following protodemetalation gives the neutral product chromene 8 and/or the 2,3-dihydrobenzofuran 9 and regenerates the cationic gold catalyst. Normally, the formation of the 6-member heterocycle is preferred, nevertheless, when the electron density of the triple bond is strongly perturbed by the presence of electron-withdrawing substituents on the distal alkyne terminus, the 5-*exo-dig* cyclization mode becomes a competitive path and the isomeric 2,3-dihydrobenzofurans 9, are obtained as by-products in not negligible amounts. It is worth noting that under these neutral reaction conditions, the 2,3-dihydrobenzofurans 9 with exocyclic double bond are stable and the base-promoted tautomerization to give the corresponding aromatic benzofurans 7 was not observed. On the other hand, the treatment of isolated 2,3-dihydrobenzofurans 9 under basic conditions resulted in a quick and quantitative isomerization to the aromatized benzofurans 7 (see ESI† for details).

Conclusions

In this work, a MW-enhanced divergent approach for the synthesis of 2-substituted benzofurans and chromenes starting

from variously substituted *o*-propargylphenols has been studied and optimized. Overall, we developed a robust protocol to synthesize *o*-propargylphenols (seventeen examples) as resourceful precursors for the preparation of seventeen benzofurans and sixteen chromenes. Most of them are new entities with no precedents in the literature. The reactions of *o*-propargylphenols under base conditions, give regioselectively the benzofurans (7) in yields ranging from 43 to 98%. Conversely, under cationic gold catalytic conditions, chromenes (8) were achieved in more variable yields (ranging from 25 to 93%), often beside small amounts of the corresponding 2,3-dihydrobenzofuran isomers (9), thus with lower regioselectivity.

The results of this work demonstrated once again the potential of diversity-oriented synthesis for the exploration of molecular complexity. Neglected *o*-propargylphenols were demonstrated to be versatile and strategic starting materials to access simple nuclei and complex heterocyclic scaffolds by intramolecular cyclization reactions. The potential of these intriguing substrates will be further investigated in our lab.

Author contributions

A. G.: investigation, validation, writing – original draft, data curation. E. B.: conceptualization, investigation, validation, writing – original draft, data curation. I. N. and F. T.: investigation, data curation. V. P.: conceptualization, writing – review



& editing. G. A.: conceptualization, funding acquisition, methodology, supervision, writing – original draft.

Data availability

All data supporting the findings of this study, including experimental details, spectroscopic characterization data, and spectra for all compounds, are available in the ESI.†

Conflicts of interest

There are no conflicts to declare.

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