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A new gold-catalysed azidation of allenes†

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A new gold-catalysed azidation reaction of allenes is presented as a new highly modular approach for the synthesis of substituted allyl derivatives containing nitrogen from simple precursors.

Organic azides are common building blocks for the synthesis of natural products and nitrogen-containing heterocycles of pharmacological relevance. As classic azides, allyl azides are useful precursors of many functional groups (e.g. amines, nitriles), and can be used in copper-catalysed 1,3-dipolar additions to alkynes for the synthesis of allyl triazoles. The classic synthesis of allyl azides involves the substitution of primary or secondary allyl derivatives with nucleophilic azides, like NaN₃ or TMSN₃ (Scheme 1a). However these methods have limitations, and mixtures of regioisomers are often obtained. Pd- and Mo-catalysed allylic azidations have been also described with better results in the selectivity and scope.

Allenes are versatile starting materials for developing sequential reactions, affording new and efficient methods for the preparation of complex compounds of synthetic or biological importance. In this area, gold-catalysed intermolecular addition of nucleophiles to allenes is an important reaction that has been widely developed in the last few years, especially for oxygen nucleophiles, although the nitrogen and carbon-based nucleophile versions have also been described. We envisioned that nucleophilic azides could be used in a similar gold-catalysed reaction with allenes, as a new method for the synthesis of allyl azides. We report here the first

Scheme 1 Different approaches for the synthesis of allyl azides.

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examples of this concept and our efforts to develop this new methodology as a highly modular approach for the synthesis of allyl azides with extra functionalities that will enable a straightforward increase in molecular complexity from simple allenes (Scheme 1b).

The challenge of this reaction is that, in contrast with alcohols or amines, the available nucleophilic azides, NaN₃ or TMSN₃, do not have the necessary proton to furnish the protonolysis of the vinyl–gold complex intermediate and therefore, the catalytic cycle cannot be completed. In order to close the catalytic cycle, the volatile, toxic and explosive hydrazoic acid, HN₃, has to be cautiously generated *in situ*. Nevertheless, protonolysis of vinyl–gold(1) complexes is only one way to close the catalytic cycle and other electrophiles can be employed to break the C–Au bond without the need for hazardous reagents.

Using commercially available cyclohexylallene (1a) and Ph₃PAuCl/AgOTf, we first investigated the optimal conditions for the hydroazidation reaction. We found that TMSN₃ was the best source of the nucleophilic azide, and trifluoroacetic acid (TFA) in DCM was the best proton source to provide clean complete conversion to the expected allyl azide 2a, with its regioisomer 2a', together with allyl trifluoroacetate, 3a, plus acetamide 4a as an unexpected product (Scheme 2).¹⁵

Scheme 2 Gold-catalysed hydroazidation of cyclohexylallene.

AgOTf seemed to be the most effective chloride abstractor in this reaction, and was used for catalyst screening. ¹⁶ NHC-Au complexes did not favour the azidation reaction, and **3a** was obtained as the major component in low conversions. The best result was obtained with (PhO)₃PAuCl as the catalyst. In a final approach to get the optimum conditions, we found that the best ratio of products and

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Table 1 Scope of the gold-catalysed hydroazidation of allenes

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Entry ^a	Allene	Yield (%), 2:2' (ratio)
1	R = Cy 1a	59, $2a:2a'(3.7:1)^b$
2	R = n-hexyl, 1b	48, 2b : 2b ' $(1.9:1)^c$
3	R = n-octyl, 1c	47, $2\mathbf{c} : 2\mathbf{c}' \ (1.8 : 1)^d$
4	R = Ph, 1d	62 , 2d
5	$R = (MeO_2C)_2CHCH_2$, 1e	43, $2e:2e'(3.8:1)^c$
6	$R = CH_2$ -N-phthalamide, 1f	48, 2f ; 7, 2f ' ^c
7	$R = (Boc)_2 NCH_2, 1g$	38 , 2g ^e
8	R = R' = Me, 1h	$2\mathbf{h} : 2\mathbf{h}' (3.7:1)^f$
9	R = Ph, R'' = n-octyl, 1i	76 , 2i
10	R = Bn, R'' = n-Pr, 1j	67, 2j : 2j ′ (1.3:1)
11	R = Bn, R'' = i-Pr, 1k	55, 2k : 2k ' (1.67:1)
12	$R = Ph(CH_2)_2, R'' = n-Pr, 11$	70, 2l : 2l ' (1:1.13)
13	R = Ph(CH2)2, R'' = i-Pr, 1m	80, $2m : 2m' (1:1)$
14^g	$R = p\text{-}CF_3Ph$, 1n	74, 2n
15^g	$R = p\text{-}CF_3Ph, R'' = n\text{-}Pr, 10$	67 , 20
16	R = p-ClPh, $R'' = n$ -octyl, 1p	72, 2p

 a 3 observed in traces (<5%) in the reaction mixture. b 4a isolated as a by-product (15%). c Methyl ketones isolated as by-products: 4b' (R = n-hexyl, 16%), 4e' (R = (MeO_2C)_2CHCH_2, 17%), 4f' (R = CH_2-N-phthalamide, 32%). d Run without water (0.08 M). e Deprotection of one Boc- was observed under reaction conditions. f 100% conversion; products not isolated due to volatility issues. g Reactions carried out at 30 °C for 60 h.

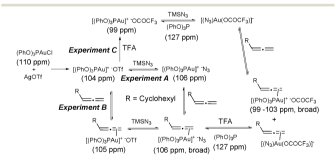
yields was obtained changing the concentration, lowering the temperature, adding up to 5 equivalents of water as additive, ^{17,18} and purifying the mixture over basic alumina. Under the best conditions, we investigated the scope of the reaction with different substituted allenes (Table 1).

The reaction works for mono- and disubstituted allenes with different functionalities, but functional groups sensitive to acid are not well tolerated. Allyl azides 2 from the attack to the less substituted carbon of the allene were obtained as the major or only product in moderate to good yields. The regioselectivity of the reaction seems to be dominated by electronic effects: allenes directly substituted with a phenyl group gave only one regioisomer (entries 4, 9, 14, 15 and 16, Table 1). However, when the phenyl group was moved away from the allenic system (entries 10 to 13, Table 1), both regioisomers were observed, even in the case of bulkier substituents (R" = i-Pr, entries 11 and 13).

It is known that allyl azides equilibrate in solution, even at room temperature, by a [3,3] sigmatropic rearrangement.¹⁹ Monitoring the reaction of allene 1a by 1H NMR we observed that although both allyl azides are formed in the gold-catalysed reaction, the concentration of allyl azide 2a' increases at the beginning and then decreases (2a' = 31% at 1 h, 2a' = 17%, at 3 h), isomerising into 2a during the reaction time.²⁰ We postulate that amide 4a comes from the gold-catalysed Schmidt reaction of the methyl ketone,²¹ formed by protonation of the allene at the terminal position and attack of H_2O , with the *in situ* formed HN_3 .

To investigate the possible formation of LAuN₃ complexes, ²² we carried out a preliminary NMR study: three stoichiometric experiments were performed at room temperature in which the order of addition of the components was varied. ¹H and ³¹P NMR spectra were recorded after each addition (Scheme 3).

Comparison of the reaction profile by analysis of the ³¹P NMR spectra in experiments A and B, where the addition of the allene and the azide were reversed, suggests that the azide coordinates to the gold in both cases, even when the allene is present in the reaction, and therefore an inner-sphere mechanism cannot be ruled out.23 Interestingly, after addition of TFA both reactions exhibited a broad signal in 31P NMR around 100 ppm together with a sharp singlet at 127 ppm from the free (PhO)₂P, which grew over the reaction time.²⁴ This suggests the CF₃CO₂⁻ is capable of displacing either the (PhO)₃P or the N₃ from the gold to reach a complex equilibrium where at least three allene-gold complexes might be involved with different ligands/counterions.²⁵ The exchange of TfO⁻ by CF₃CO₂⁻ and the displacement of (PhO)₃P were also observed in experiment C, where TFA was added first. Further shift of the 31P NMR signal after addition of the TFA suggests the formation of the complex (PhO)₃PAu⁺· OCOCF₃. The signal from free (PhO)₃P was also observed in this reaction when azide was added, suggesting an equilibrium with the complex $[(N_3)Au(OCOCF_3)]^-$. When the allene was added to this mixture a similar reaction profile to the experiments A and B was observed. Analysing the signals of the ¹H NMR spectra we could not observe the proposed vinyl-gold intermediate, but clean conversion to the products, suggesting fast protonolysis. A similar rate of formation of the different products was observed in all experiments after all the components had been added, implying similar catalytic species involved after addition of the acid and suggesting that coordination with the allene and/or the N₃ might be the rate limiting step for the reaction. When we followed the reaction using NMR under catalytic conditions, a broad signal around 100 ppm, and free (PhO)₃P (127 ppm) were observed in ³¹P NMR spectra, which suggests that the complicated equilibrium between the complexes shown in Scheme 3 might be existing under catalytic conditions.26

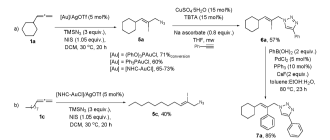


Scheme 3 ³¹P NMR-profile for stoichiometric experiments.

The important utility of this transformation is illustrated by the straightforward synthesis of iodinated allyl azides. By using NIS as the electrophile in the gold-catalysed azidation of allenes, ^{14,27} an extra functionality can be introduced in one step, making this method a very useful tool for the orthogonal functionalization of simple molecules into complex structures. The reaction of allene 1a with TMSN₃, in the presence of NIS using different gold complexes afforded the vinyl iodide 5a as the only regioisomer (Scheme 4a). Vinyl iodide 5c was obtained in moderate yield using the NHC-Au catalyst (Scheme 4b).²⁸

To further demonstrate the potential of this method as a highly modular approach, we carried out the synthesis of derivatives **6a** and

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Scheme 4 Orthogonal functionalization of allenes using the goldcatalysed azidation methodology.

7a by using a click²⁹ and a Suzuki-Miyaura cross-coupling³⁰ reaction with product 5a (Scheme 4a).

In conclusion, we have found that the hydroazidation of allenes to obtain allyl azides is possible by using (PhO)3PAuCl in the presence of TMSN₃ and TFA. The reaction works well for substituted allenes, but functional groups sensitive to acid are not well tolerated. Preliminary mechanistic studies point out to a complex mechanism with equilibrium between several allene-gold-azide complexes, and an unusual displacement of the phosphite ligand from the goldcoordination sphere. Further investigation into this and a more detailed mechanistic study is underway in our laboratories. Further transformations involving the azide and diverse cross-couplings at different stages of the process are envisioned³¹ and will be of general interest to the synthetic chemistry community.

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