ChemComm



COMMUNICATION

View Article Online



Cite this: Chem. Commun., 2017, **53**. 5733

Received 24th March 2017, Accepted 4th May 2017

DOI: 10.1039/c7cc02244a

rsc li/chemcomm

Alkynyl sulfoxides as α -sulfinyl carbene equivalents: gold-catalysed oxidative cvclopropanation†

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Alkynyl sulfoxides are shown to act as α-sulfinyl metallocarbene synthons under oxidative gold catalysis, enabling reactions that are not available from diazo-precursors. This strategy is exemplified in the synthesis of fused α -sulfinyl cyclopropanes.

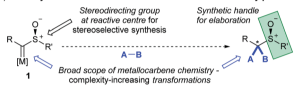
Metallocarbenes underpin a broad range of powerful chemo- and stereoselective transformations in modern organic synthesis.¹ α-Sulfinyl metal carbenes, 1, position a readily-elaborated functional group²⁻⁵ bearing a stereogenic centre at the reactive site (Scheme 1a). However this attractive proposition has yet to be realised using conventional approaches to metal carbene reactivity. Maguire and co-workers established that α -diazo sulfoxides are only isolable when constrained as part of a cyclic system and that they and their resulting \alpha-sulfinyl rhodium carbenes undergo rapid Wolff-like rearrangement (Scheme 1b).⁷ Here we demonstrate how the reactivity patterns of α-sulfinyl carbenes can be accessed from alkynyl sulfoxides under gold catalysis.

The use of a π -acid to chemoselectively activate alkynes in the presence of a nucleophilic oxidant provides an attractive route into α-oxo metal carbene reactivity patterns without the need to install or handle diazo groups.⁸⁻¹¹ One intriguing aspect is that some reactions appear to bypass the actual gold carbene 4 and proceed directly from the vinyl gold carbenoid intermediate 3 (Scheme 1c).¹² We hypothesised that broader applications of α-sulfinyl metal carbene chemistry might therefore be accessible if α -sulfinyl vinyl gold carbenoid 7 could be accessed from alkynyl sulfoxide 6, quenched prior to expulsion of the nucleofuge, and proved less vulnerable to rearrangement than the corresponding metal carbene. This approach presents an interesting challenge as sulfoxides are effective nucleophiles and oxygen-transfer agents in the presence of alkyne-gold complexes¹³ or metal carbenes.¹⁴ For successful application

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† Electronic supplementary information (ESI) available: Experimental procedures and analytical data for new compounds. ¹H and ¹³C NMR spectra. Structural determination and additional catalysis results. CCDC 1528851. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c7cc02244a

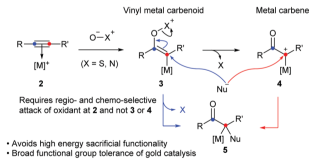
(a) α-Sulfinyl metal carbenes: a desirable but unrealised reactivity profile



(b) Diazo precursors to rhodium carbenes (Maguire)

• Challenging preparation - Unstable except when cyclic (Z = CR2, NR, O)

(c) The alkyne oxidation strategy to α -oxo metal carbenes



(d) Unlocking the potential of α -sulfinyl carbenoids (this work)

regiocontrol?
$$X^+$$
 $O^ X^+$ $A^ A^ A^$

· Readily prepared precursors

· Sulfinyl group must direct and not react

Scheme 1 Approaches to access α -sulfinyl metal carbene type reactivity.

of alkynyl sulfoxide 6 as an α-sulfinyl carbene equivalent, effective π -activation and regionelective oxidation is required, but 6 and 8 must not act as nucleophilic oxidants. 15

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We tested this hypothesis in the oxidative cyclopropanation reaction of readily accessible ene-alkynyl sulfoxides.‡ A reaction survey with 9a identified that the desired cyclopropane-fused thiolane S-oxide was formed as an approximately 6:1 mixture of diastereomers 10a and 10b using 3,5-dichloropyridine-Noxide (11) as stoichiometric oxidant in the presence of various cationic Au(1) catalysts. Phosphite, N-heterocyclic carbene and bulky phosphine ligands all proved effective on the gold, with SPhosAuNTf₂ giving highest yield (Table 1, entries 1–5). Dioxane proved superior to other solvents (entries 5-9) while 11 was more effective than other commonly used pyridine-N-oxide derivatives 12 and 13 (entries 10-13). 16 Changing the temperature had little effect on dr, though conversion stalled at much lower temperatures: at 80 °C the catalyst loading could be halved with little effect, though dropping further was detrimental to conversion of 9a (entry 10). Increasing oxidant loading saw lower yields, likely due to over-oxidation pathways (entry 11).

A range of ene-alkynyl sulfoxides 9a-v were prepared to explore the effect of the alkyne substituent on the reaction (Table 2). Primary, secondary and tertiary alkyl substituents were all accommodated with good conversions at 50 °C (entries 1-6). Notably, cyclopropyl-substituted alkyne 3q gave the same yield and d.r. at room temperature (entry 6). Aryl substituted alkynes were also more reactive, proceeding at room temperature, although higher yields were obtained under the standard conditions (entries 7-20, see ESI† for reactions at room temperature).

Survey of reaction conditions

Entry	Ligand	Solvent	T (°C)	Oxidant	Yield of $\mathbf{10a}^{a}$ (%)
1	L1	Dioxane	65	11	42
2	L2	Dioxane	65	11	45
3	L3	Dioxane	65	11	47
4	L4	Dioxane	65	11	59
5	L5	Dioxane	65	11	66
6	L5	1,2-DCE	60	11	45
7	L5	THF	60	11	37
8	L5	CH_2Cl_2	rt	11	26
9	L5	Toluene	60	11	39
10	L5	Dioxane	80	11	$69^{b,c}$
11	L5	Dioxane	80	11 (2.0 eq.)	54
12	L5	Dioxane	80	12	63
13	L5	Dioxane	80	13	59

^a Reactions performed on a 0.1 mmol scale; yields of the major diastereomer 10a determined by ¹H NMR analysis of the crude reaction mixture using 1,2,4,5-tetramethylbenzene as an internal reference. Overlap prevented accurate determination of dr. b 62% at 2.5 mol% cat. 27% at 1.0 mol% cat. L1 = (tris(2,4-di-tertbutylphenyl)phosphite). L2 = 1,3-bis(2,6-diisopropylphenyl-imidazol-2-ylidene). L3 = 2-dicyclohexylphosphino-2',4',6'-triisopropylbiphenyl. L4 = 2-ditertbutylphosphinobiphenyl. L5 = 2-dicyclohexylphosphino-2',6'-dimethoxybiphenyl (SPhos). ^c Higher concentrations afforded lower yields (42% **10a** at 0.2 M).

In these cases the d.r. was approximately 8:1 as determined by ¹H NMR analysis of the reaction mixture before purification. The aromatic substituent can be either electron-rich or -poor and will accommodate a variety of functionality across all positions. The tolerance of this chemistry is highlighted by the ready inclusion of a 3-bromothiophen-2-yl moiety (entry 20). Furthermore, the reactions of diene-alkynyl sulfoxides 9u/v proceeded smoothly to the desired sulfur heterocycles despite the possibility of competing cycloisomerisation prior to oxidation across one or both of the two 1,6-enyne motifs embedded in the substrates (entries 21 and 22).17

The relative stereochemistry of the major diastereomers 10 and minor diastereomers 10' were assigned using characteristic chemical shifts in the ¹H NMR spectra (see ESI†).

In addition a crystal structure was obtained for major diastereomer 10g (Fig. 1), s confirming the NMR analysis that the sulfoxide oxygen and cyclopropyl methylene are on the same side of the thiolane ring.

The reaction of 9q, bearing an ortho-isopropyl substituent, saw formation of a side-product alongside 10q (Table 2, entry 17) although this was not isolated in sufficient quantity or purity to

Table 2 Substrate scope

Entry	R	9	T (°C)	Time (h)	% Yield of 10 ^{<i>a</i>}
1	ⁿ Bu	9a	50	17	72 (6:1)
2	PhCH ₂ CH ₂	9b	50	3.5	63 (6:1)
3	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	$\mathbf{9c}^{b}$	50	21	70 (10:10:1:1)
4	Cyclohexyl	9d	50	24^c	45 (7:1)
5	^t Bu	9e	50	25	70 (12:1)
6	Cyclopropyl	9f	50	17	$86^{d}(10:1)$
7	Ph	9g	65	0.75	80
8^e	4-MeC_6H_4	9h	23	28	75
9	4-MeOC_6H_4	9i	40	1	78
10^e	4 -AcNH-C $_6$ H $_4$	9j	50	20	79
11	$4-F_3CC_6H_4$	9k	50	17	63
12	$4\text{-MeO}_2\text{CC}_6\text{H}_4$	9 l	50	20	64
13	$4\text{-FC}_6\text{H}_4$	9m	50	17	68
14	3-MeOC_6H_4	9n	50	18	70
15	4-BrC ₆ H ₄	90	50	3	74
16^e	2-BrC ₆ H ₄	9p	23	28	50
17	2^{-i} Pr-C ₆ H ₄	9q	50	28	52 (7:1)
18	2-Naphthyl	9r	50	28	70
19	2-Furyl	9s	50	28	74
20	Br 32,	9t	50	28	73
21	/\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	9u	50	21	63 (10:1)
22	Ph O So	9v	50	21	65 (10:1)

^a Isolated yields after purification by column chromatography. The yields refer to a single diastereomer apart from when diastereomeric ratios are given. ^b **9c** is a 1:1 mixture of diastereomers. ^c Incomplete conversion. ^d The same yield and d.r. were obtained at room temperature. e 2.5 mol% SPhosAuNTf2

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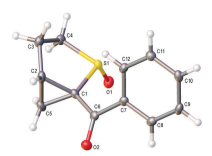


Fig. 1 X-ray crystal structure of major diastereomer 10g

allow full characterisation. We hypothesised that 1,5-hydride transfer from the benzylic position may be competing with cyclopropanation. 18 To test this hypothesis we prepared the methylsulfoxide 12 where cyclopropanation is not possible. The formation of stilbene 13 under the standard reaction conditions is indeed consistent with 1,5-hydride transfer onto a vinyl gold carbenoid (cf. 7) followed by elimination of a proton and protodeauration (Scheme 2). Key resonances in 13 also correlate to those in the side-product from 9q.

The feasibility of using a disubstituted alkene in the cyclopropanation was then explored using styrene 14 (Scheme 3). Under the standard reaction conditions the more heavily substituted cyclopropane 15 was indeed formed, 19 alongside hydroxylated ringopened product 16. Formation of 16 is consistent with the cationic character of a gold carbenoid extending through the alkene and enabling a hydrative cyclisation in the presence of adventitious water.20

A preliminary investigation shows that using alkynyl sulfoxides as α-sulfinyl carbene equivalents is not limited to sulfur heterocycle formation. Under unoptimised conditions, which saw incomplete conversion, 1,5-enyne 17 gave fused carbocyclic ring system 18 as a 1.6:1 mixture of diastereomers (Scheme 4).

In conclusion, the synthetic limitations that have prevented access to desirable aspects of α-sulfinyl metallocarbene reactivity can be bypassed by an oxidative gold catalysis strategy using

Scheme 3 Use of 1,2-disubstituted alkene in oxidative cyclopropanation.

Scheme 2 An alternative reaction pathway consistent with 1,5-hydride transfer

Scheme 4 Synthesis of an α -sulfinyl cyclopropyl-fused cyclopentanone.

readily accessed alkynyl sulfoxides. For the first time α-sulfinyl carbene-like activity is demonstrated through intramolecular cyclopropanation reactions, affording ring-fused cyclopropanes containing α-sulfinylcarbonyl motifs.21 Future work will address the use of this approach in the wider context of carbene reactivity and explore the opportunities arising from the use of enantiopure sulfoxides.22

The authors acknowledge support from the Centre for Chemical and Materials Analysis in the School of Chemistry at University of Birmingham (UoB) and thank Dr Louise Male (UoB) for X-ray crystallography. We thank the UoB for a studentship (MJB).

Notes and references

‡ All sulfoxides were prepared in the racemic series.

§ Crystal structure determination of 10g: crystal data for C12H12O2S $(M = 220.28 \text{ g mol}^{-1})$: triclinic, space group $P\bar{1}$ (no. 2), a = 6.2782(3) Å, $b = 7.1917(3) \text{ Å, } c = 12.4920(6) \text{ Å, } \alpha = 86.275(4)^{\circ}, \ \beta = 75.966(4)^{\circ},$ γ = 66.086(4)°, V = 499.87(4) Å³, Z = 2, T = 100.01(11) K, μ (CuKα) = 2.667 mm⁻¹, D_{calc} = 1.463 g cm⁻³, 7653 reflections measured (7.3° ≤ 2 Θ ≤ 144.236°), 1939 unique ($R_{\rm int}=0.0218$, $R_{\rm sigma}=0.0170$) which were used in all calculations. The final R_1 was 0.0390 ($I>2\sigma(I)$) and w R_2 was 0.0963 (all data). The CIF for the crystal structure of 10g has been deposited with the CCDC and have been given the deposition number CCDC 1528851.

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