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A gold-catalysed fully intermolecular oxidation

and sulfur-ylide formation sequence on

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An efficient C–O, C–S and C–C bond-forming sequence leads to functionalised compounds bearing sulfur-substituted quaternary carbons. Ynamides are employed as diazo-equivalents to access the [2,3]-sigmatropic rearrangements of allyl sulfonium ylides by a three-component chemoselective oxidation and intermolecular ylide formation.

ynamides\*

The [2,3]-sigmatropic rearrangement of allyl sulfonium ylides is a potent method for the formation of Csp<sup>3</sup>-Csp<sup>3</sup> bonds.<sup>1</sup> A significant hydrocarbon functionalisation process is achieved when the rearrangement is coupled with in situ ylide formation through reaction of a sulfide with a metal carbene, formed in situ from a sacrificial functionality such as the diazo-group (Doyle-Kirmse reaction, Scheme 1).<sup>2,3</sup> We<sup>4</sup> and others<sup>5</sup> have been engaged in efforts to access the synthetic potential of these ylides using methods to generate carbenoids that avoid the pre-installation and use of potentially hazardous high-energy diazo groups. Our initial studies established that sulfur ylides can be prepared from the intermolecular reaction of sulfides with gold-carbenes (Scheme 2).4a,d,6 However, the use of propargylic carboxylates as carbenoid precursors impacted on the subsequent ylidic rearrangements to generally afford products isomeric to those from [2,3]-sigmatropic pathways. More-congested centres and synthetically valuable sulfur-substituted quaternary carbons were inaccessible as terminal alkynes and unsubstituted allyl groups on the sulfide<sup>7</sup> were required.<sup>4a,d</sup> We subsequently established that sigmatropic-rearrangements of sulfur ylides were accessible through an intramolecular gold or platinum catalysed cycloisomerisation of alkynyl allyl sulfoxides.<sup>4b,8</sup> Here we report a diazo-free oxidation-ylide formation sequence to access sulfur-ylide rearrangements by a selective and efficient fully intermolecular transformation of ynamides.

In light of our,<sup>9</sup> and others<sup>10</sup> studies into gold-catalysed intermolecular atom/group-transfer onto ynamides **A**,<sup>11</sup> we questioned







Scheme 2 Intermolecular trapping of gold carbenes with allyl sulfides: ref. 4a,d

whether ynamides could be used to replace diazo compounds in intermolecular sulfonium ylide formation to form quaternary carbons (Schemes 1 and 3).

Significant transformations have resulted from  $\alpha$ -oxo-gold carbene formation by intermolecular oxidation of a C–C triple bond.<sup>9,10,12</sup> While these predominantly feature subsequent intramolecular cyclisations, there are striking exceptions in the terminal alkyne series where the mono-substituted gold carbene is trapped with intermolecular oxygen-, nitrogen- and halide nucleo-philes other than the oxidant.<sup>13</sup> Similar three-component couplings have not however been reported in the ynamide series and it is notable that the resulting 1,1-disubstituted organogold species C/D appears prone to oxidation even when intramolecular pathways are available.<sup>9,10,14</sup> A successful outcome therefore requires a high level of reagent-compatibility and selectivity: the sulfide must not affect

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Scheme 3 A gold-catalysed intermolecular oxidation – intermolecular trapping manifold for ynamides.

activation or oxidation of the ynamide  $(A \rightarrow B \rightarrow C)$ .<sup>15</sup> Ylide formation must then compete successfully with cyclopropanation (G) and oxidation (H) of organogold species C and/or D.

After exploring a variety of parameters including catalyst, solvent and oxidant with various ynamides (see ESI<sup>+</sup>), reaction conditions were found to effect the complexity-increasing cascade from ynamides 1 into the functionalised tertiary thioethers 4 (Scheme 4). Competing double-oxidation to form imido-ketones (H) was observed throughout this study alongside incomplete consumption of 1, and this pathway was exclusively observed using Au(III) or NHC-Au(I) catalysts. However, good conversion and selectivity for the formation of 4 was achieved using a combination of cationic gold phosphite complex Au-I alongside methylpicolinate N-oxide 3. The reactions were run in dichloromethane at room temperature and reagents 1:2:3 were added at the start of the reaction in a near-stoichiometric 1:1.2:1.3 ratio respectively (Scheme 4). Recourse to continual or portion-wise slow-addition strategies that maintain a low concentration of oxidant to favour reaction with the desired nucleophile was avoided. This preference was maintained throughout the study in order to explore the extent and influences on the observed chemoselective preference for sulfide attack at the organogold intermediate.13,16



Scheme 4 The ynamide-based Doyle–Kirmse type reaction.

Ynamides bearing *N*-phenyl and *N*-methyl groups reacted similarly (**4ba** *vs.* **4aa**) while an *N*-benzyl equivalent was less suited affording a lower yield of **4ca** with competing overoxidation observed.<sup>17</sup> The trend between *N*-phenyl and *N*-methyl was maintained in the *p*-tolylsulfonamide series, albeit with slightly higher yields (Scheme 4, **4da** and **4ea**). In both cases, products from cyclisation of **C/D** onto the *N*-substituent were not observed. An ynamide prepared from imidazole was unreactive. Oxazolidinone-derived ynamide **1f** could be used to form **4fa** but greater conversion and selectivity for ylide formation was achieved using a bulky phosphine-derived catalyst **Au-II** in place of the phosphite–gold complex, identifying potentially significant influencing factors between different types of ynamides (see ESI<sup>†</sup>).

The sulfonamide-derived ynamides were used throughout the remainder of this study into structural effects on the relative efficacy of the standard conditions. The reaction remained effective in the presence of *S*-aryl, *S*-benzyl- and *S*-alkyl allyl sulfides **2b–e** as well as bisallylsulfide **2f** (Scheme 5). Unsurprisingly, a more electron-deficient aryl bromide substituent renders the ylide pathway less competitive against oxidation with ~30% yield of the over-oxidation product determined from the crude reaction mixture.

The role of the ynamide *C*-substituent was next tested under the standard conditions (Scheme 6). None of the desired product from ylide formation was observed when anisole-derived ynamide **1g** was reacted with sulfides **2a** or **2b** with the over-oxidation product **5** instead predominating. A preference for over-oxidation against an intramolecular reaction has recently been reported when using *p*-anisole derived ynamide.<sup>10f</sup> In contrast, both inductively and mesomerically electron-withdrawing groups were well tolerated (**4hb** and **4ib**). Alkyl-substituted gold carbenoids are prone to **1**,2-insertion<sup>9a</sup> and this intramolecular pathway was indeed preferred, affording **6** with low conversion of the hexynederived ynamide **1j**. A conjugated ene-ynamide **1k** did not follow a recently reported 4- $\pi$ -electrocyclisation pathway<sup>10c</sup> instead reacting productively to afford functionalised tertiary allylic thioether **4ka**.

By analogy to the effects encountered in diazo-derived metal carbene chemistry, changing the ynamide *C*-substituent was expected to influence the reactivity of the resulting carbenoid.<sup>2</sup> However, the impact of the electronic influence on the chemo-selectivity of these ynamide-based reactions is informative: the formation of the desired compound **4** requires the active organogold species to react preferentially with the polarisable sulfur nucleophile **2** in preference to the dipolar oxygen-nucleophile **3**.



Scheme 5 Effect of the non-migrating group on the allyl sulfide.



Nucleophilic attack can occur either alongside- or after elimination of methylpicolinate (Scheme 3,  $\mathbf{C} \rightarrow \mathbf{E} \nu s$ .  $\mathbf{C} \rightarrow \mathbf{D} \rightarrow \mathbf{E}$ ).<sup>8–10,12,13</sup> Considering the contrasting natures of the nucleophiles, a favourable attack of the sulfide on the vinyl gold carbenoid **C**, bearing a relatively-uncharged carbon centre, appears more likely than on the cationic carbon of gold carbene **D**. The strongly  $\pi$ -acidic phosphite ligand affords significant cationic character to **D** by decreasing the ability of gold to donate electron density.<sup>18</sup> However, these same electronic characteristics should aid the desired chemoselectivity by disfavouring the elimination of methylpicolinate required to form **D**. On this basis, the reactivity of electron-rich ynamide  $\mathbf{1g} (\mathbf{R}^1 = p\text{-MeOC}_6\text{H}_4)$  can be explained by the mesomeric contribution from the anisole MeO-group aiding elimination of the nucleofuge to form **D** and disfavouring ylide formation.

Sulfides with substituted allyl units were then explored to ascertain if a [2,3]-sigmatropic rearrangement was operative. Pleasingly, and in contrast with other systems,<sup>7</sup> both 2-vinyl-tetrahydrothiophene **2g** and cinnamyl derivative **2h** underwent clean reactions. The functionalised 8-membered sulfur heterocycle **4eg** and the tertiary thioether bearing a vicinal tertiary stereogenic centre **4eh** are the products expected from sulfur-ylide formation and [2,3]-sigmatropic rearrangement with allylic inversion (Scheme 7).‡<sup>2,6,19</sup> Interestingly, formation of **4eh** proceeded with higher diastereoselectivity than is normally observed from the



Scheme 7 Reactions of ynamide-derived sulfur ylides displaying allylic inversion. Crystal structure of **4eh** with ellipsoids drawn at the 50% probability level.

reaction of substituted allyl sulfides 2h with metal carbeness derived from donor-acceptor diazocompounds.<sup>6,19</sup> The relative stereochemistry of the major diastereoisomer was confirmed by X-ray analysis (Scheme 7) and is consistent with a preference for the cinnamyl phenyl group to be positioned anti to the amide group across an envelope transition state. Rapid access to these *N*,*N*-disubstituted amide derivatives by the ynamide strategy may therefore provide additional synthetic advantages.

In conclusion, a gold-catalysed cascade reaction is reported for the practical complexity-increasing synthesis of functionalised tertiary thioethers directly from ynamides. Selectivity for sulfur ylide formation over competing processes is rationalised and can be achieved when all reagents are added at the start of the process with a near-stoichiometric reagent loading. The products are consistent with those expected from [2,3]-sigmatropic rearrangement of allyl sulfonium ylides. This report demonstrates that ynamides can be used as synthetically attractive replacements to 1,1-disubstituted diazocompounds in intermolecular processes.

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## Notes and references

‡ Crystal data for **4eh**: C<sub>36</sub>H<sub>31</sub>NO<sub>3</sub>S<sub>2</sub>, M = 589.74, monoclinic, a = 16.0175(3) Å, b = 9.6290(2) Å, c = 19.5279(5) Å,  $\beta = 98.570(1)^{\circ}$ , U = 2978.2(1) Å<sup>3</sup>, T = 120(2) K, space group  $P2_1/c$ , Z = 4, 34 108 reflections measured, 6813 unique ( $R_{int} = 0.0607$ ) which were used in all calculations. The final  $R_1$  was 0.0458 ( $I > 2\sigma(I)$ ) and w $R(F_2)$  was 0.1150 (all data).

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