Group 2-catalysis for the Atom-Efficient Synthesis of Imidazolidine and Thiazolidine Derivatives

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Abstract
A wide variety of functionalised imidazolidine-2-ones and –thiones, 2-imino-imidazolidines and thiazolidine-2-thiones have been synthesised under very mild reaction conditions using simple and cost-effective alkaline earth bis(amide) precatalysts, [Ae{N(SiMe3)2}2(THF)2] (Ae = Mg, Ca, Sr). The reactions ensue with 100% atom-efficiency as one-pot cascades from simple, commercially available terminal alkyne and heterocumulene reagents. The reactions take place through the initial assembly of propargylamidines, which are utilized in subsequent cyclisation reactions through addition of the isocyanate, isothiocyanate and, in one case, carbon disulphide reagents. This reactivity is deduced to take place through a well-defined sequence of heterocumulene hydroacetylenation and alkyne hydroamination steps which are all mediated at the alkaline earth centre. The rate and regioselectivity of the cyclisation reactions are, thus, found to be heavily dependent upon the identity of the catalytic alkaline earth centre employed. Similarly, the selectivity of the reactions is observed to be profoundly affected by stereoelectronic variations in the individual substrates, albeit via a similar group 2-centred reaction mechanism in all cases studied.

Introduction
Heterocyclic moieties are constituents of two-thirds of the top-selling small molecule pharmaceuticals in the USA.1 The imidazolidine-2,4-dione (hydantoin) moiety is a particularly notable constituent of many biologically active compounds and numerous hydantoin derivatives have been identified as anticonvulsant, antiulcer, antiarrhythmic, antimuscarinic, antiviral and antidiabetic agents.2 Phenytoin, for example, is commonly used in the treatment of epilepsy,3 and aplysinopsin, isolated from the sponge Aplysinopsis reticulate has been shown to exhibit cytotoxicity against cancer cells.4 The agonist of the human androgen receptor, GLPG04924,5 and the hydantoin muscle relaxant, dantrolene,6 have been widely employed in a medical context. Many herbicides also contain hydantoin...
skeleta as an integral part of their structure and they are also applied as intermediates for the synthesis of enantiomerically pure amino acids by dynamic kinetic resolution.\textsuperscript{7,8}

\begin{center}
\begin{tabular}{cc}
Phenytoin & Aplysinopsin \\
\includegraphics[width=0.2\textwidth]{Phenytoin.png} & \includegraphics[width=0.2\textwidth]{Aplysinopsin.png} \\
GLPG04924 & Dantrolene \\
\includegraphics[width=0.2\textwidth]{GLPG04924.png} & \includegraphics[width=0.2\textwidth]{Dantrolene.png}
\end{tabular}
\end{center}

The development of efficient and preferentially catalytic methods for the rapid construction of molecularly diverse hydantoin molecules from simple and inexpensive starting materials is clearly very desirable. In this regard, several transition-metal catalysed methods have been reported. A copper-based amination reaction of esters with di-\textit{tert}-butyldiaziridinone has been shown to afford 1,3,5-trisubstituted hydantoins\textsuperscript{,9} while a palladium-catalysed carbonylation reaction of aldehydes with ureas and carbon monoxide furnishes 5-, 3,5-, and 1,3,5-substituted hydantoins\textsuperscript{10} and a nickel-catalysed protocol starting from acrylates and isocyanates has recently been described\textsuperscript{11}. Of most relevance to the current work several processes have been described in which the hydantoin core is assembled from one molecule of phenylacetylene and two molecules of isocyanate through iron\textsuperscript{,12} ruthenium\textsuperscript{,13} or manganese catalysis\textsuperscript{,14}. Although described in some cases as proceeding as a [2+2+1] cycloaddition, these processes are otherwise mechanistically uncertain.
Scheme 1: Alkaline earth-catalysed intramolecular hydroamination of aminoalkenes (A), hydroacetylation of carbodiimides (B) and catalytic synthesis of imidazolidin-2-ones (C).

The alkaline earth elements (Mg, Ca, Sr and Ba) comprise an alternative suite of underexploited but inexpensive and potentially more sustainable and environmentally benign catalytic elements in their effectively invariant 2+ oxidation state. For some time we have been engaged in a programme of research to develop a defined catalytic and reaction chemistry, derived largely from sequences of metal-centred $\sigma$-bond metathesis and polarized insertion reactions. Since our initial report of the calcium-catalysed intramolecular hydroamination of aminoalkenes (Scheme 1A), we and others have applied alkaline earth pre-catalysts to an ever-growing array molecular catalytic reactions. Of most relevance to the current work, we have previously reported that complex bis(hydantoins) (II, inset Scheme 1) may be synthesised, albeit in a stoichiometric sense, through a magnesium-based cascade reaction between phenylacetylene and an organic isocyanate. Subsequent to this discovery, which was rationalised to take place through a sequence of intramolecular isocyanate hydroacetylenation and intermolecular C≡C and C=C hydroamination steps, we have described in preliminary form an extension of this chemistry to a one pot catalytic regime. In this latter case, reactions catalysed by the readily available bis-hexamethyldisilazides [Ae{N(SiMe$_3$)$_2$}(THF)$_2$] (Ia, Ae = Mg; Ib, Ae = Ca; Ic, Ae = Sr) allowed the facile
synthesis of a variety of highly functionalised imidazolidin-2-ones with 100% atom efficiency (Scheme 1B/C). These reactions take place via the initial catalytic formation of a propargylamidine (III in Scheme 1B),36 whereupon addition of an organic isocyanate initiates a further cascade of reactions involving isocyanate insertion into the catalytically active group 2 amidinate (IV in Schemes 1B/C), intramolecular Ae-N insertion of the alkynyl residue and protonolysis of the cyclised alkaline earth vinyl intermediate by the remaining propargylamidine (III) to regenerate IV and provide the imidazolidin-2-one derivative (V, Scheme 1C). In this contribution we provide a full description of the scope of this reactivity and reveal that this approach is easily extended to alternative sulphur-based heterocumulenes for the synthesis of imidazolidin-2-thiones and thiazole derivatives. Although no

Results and Discussion

We have recently reported the use of the alkaline earth bis(amido) complexes, [Ae[N(SiMe$_3$)$_2$]]$_2$(THF)$_2$ (Ae = Mg Ia, Sr Ib, Ca Ic) for the synthesis of propargylamidines by hydroacetylenation of carbodiimides.36 Following this procedure (N,N'-di-iso-propyl)-phenylpropargylamidine, compound 1, was synthesised in situ using 5 mol% of Ic. Subsequent addition of one molar equivalent of tert-butylisocyanate resulted in quantitative formation of the corresponding imidazolidin-2-one, compound 3, within the first point of analysis at room temperature (Scheme 2; Table 1, entry 3). The resultant $^1$H NMR spectrum displayed two new characteristic benzylidene singlets present in a 22:78 ratio at $\delta$ 6.50 and 5.95 ppm, respectively, coupling in the $^{13}$C{$^1$H} NMR spectrum to signals at $\delta$ 112.4 and 120.9 ppm, respectively. These were assigned by a NOESY experiment to the Z and E isomers of 3, respectively (Figure 1).
The catalyst loading could be lowered to 0.5 mol%, affording (Z)-3 and (E)-3 in the same ratio with an 87% yield from an 18 hour reaction at room temperature (Table 1, entry 4). The reaction of 1 with tert-butylisocyanate using 5 mol% of the analogous magnesium and calcium precatalysts, 1a and 1b, revealed the intermediacy of a linear urea derivative, compound 2, resulting from isocyanate insertion into the amidine within the first point of analysis and presenting a distinctive downfield NH singlet at δ_{1H} 10.48 ppm (Scheme 2). Although the magnesium-catalysed cyclisation to compound 2 took several days to reach completion at room temperature (Table 1, entry 1), compound 3 could not be isolated by methanol quenching of the catalyst as this also reversed the formation of 3. The rate of cyclisation dramatically increased with increasing metal cation size, in the order of Sr > Ca >>> Mg (Table 1, entries 1–3). A minor dependence on metal cation identity was also observed for the Z:E isomer ratio of the product, varying from ca. 1:2 for magnesium to 1:4 for strontium (Table 1, entries 1–3). Once full conversion to 3 was reached the isomer ratio did not change upon heating to 60 °C, suggesting no interconversion between the two isomers (vide infra for a rationale for the independent formation of each isomer). Heating to 80 °C, however, induced tautomerization of the benzylidene and iso-propylimino moieties to the corresponding 5-isopropylidenediamo-4-benzyl-imidazole-2-one, compound 4. Compound 4 displayed a distinctive ¹H NMR benzyl singlet (2H) resonance around 3.3 ppm, which correlated with a ¹³C{¹H} resonance at δ 103.2 ppm, and two distinct iso-propylidene methyl singlets (3H each) at 1.6 and 1.8 ppm, which were shown to be mutually coupled by a COSY experiment.

**Figure 1.** Identification of (Z)-3 and (E)-3 by NOESY NMR spectroscopy.
Table 1. Isocyanate scope for the catalytic synthesis of (5-benzylidene-4-imino)imidazolidin-2-ones from 1 using precatalysts Ia–c in C<sub>6</sub>D<sub>6</sub> at room temperature.

<table>
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<tr>
<th>Entry</th>
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<th>Product</th>
<th>cat (mol%)</th>
<th>Time (h)</th>
<th>NMR yield (%)&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Z:E (%)</th>
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<tr>
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<td>0.1</td>
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<td>22:78</td>
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<tr>
<td>4&lt;sup&gt;b&lt;/sup&gt;</td>
<td>''</td>
<td>Ic</td>
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<td>18</td>
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<tr>
<td>5&lt;sup&gt;b&lt;/sup&gt;</td>
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<td>Ic</td>
<td>0.5</td>
<td>18</td>
<td>92</td>
<td>32:68</td>
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<tr>
<td>6&lt;sup&gt;b&lt;/sup&gt;</td>
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<td>&gt;99</td>
<td>40:60</td>
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<tr>
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<td>iPr&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Ic</td>
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<td>3</td>
<td>98</td>
<td>41:59</td>
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<tr>
<td>8&lt;sup&gt;b&lt;/sup&gt;</td>
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<td>90:10</td>
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<tr>
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<td>Ic</td>
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<td>Ic</td>
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<td>&gt;99</td>
<td>85:15</td>
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<tr>
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<td>Ic</td>
<td>0.5</td>
<td>6</td>
<td>98</td>
<td>98:2</td>
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<tr>
<td>12&lt;sup&gt;b&lt;/sup&gt;</td>
<td>2,6-Pr&lt;sub&gt;2&lt;/sub&gt;C&lt;sub&gt;6&lt;/sub&gt;Ph</td>
<td>Ic</td>
<td>0.5</td>
<td>18</td>
<td>78</td>
<td>90:10</td>
</tr>
</tbody>
</table>

<sup>a</sup> Determined by integration of benzylidene protons against an internal HN(SiMe<sub>3</sub>)<sub>2</sub> standard.<sup>b</sup> Using isolated compound 1.

Encouraged by these results the scope of the isocyanate substrates that could be employed in this reaction was investigated using propargylamidine 1, with 0.5 mol% of Ic at room temperature (Scheme 3). While isocyanate insertion to form the linear analogue of 2 proved virtually instantaneous and independent of isocyanate identity, the rate of subsequent cyclisation to the corresponding imidazolidin-2-one was strongly influenced by the nature of the isocyanate substituent. Whereas the least sterically demanding substrate, ethylisocyanate, afforded a 91% conversion as determined by <sup>1</sup>H NMR spectroscopy within 20 minutes at room temperature (Table 1, entry 9), the bulkiest substrates, tert-butyl-, adamantanyl- and 2,6-di-iso-propylphenylisocyanate, required extended reaction times to reach good conversions (Table 1, entries 4–5 and 12). When the catalyst loading was increased to 5 mol%, however,
all reactions proceeded to completion within less than 5 minutes at room temperature. The Z:E isomer ratio of the products was found to be governed by both steric and electronic factors. Whereas the various arylisocyanates primarily yielded the Z isomer, (Table 1, entries 10–12) the selectivity of isomer formation for reactions with alkylisocyanates was found to be sensitive to the steric demands of the N-bound organic substituent. While isocyanates bearing primary ethyl and n-propyl substituents provided predominant formation of the Z isomer (Table 1, entries 8–9), this shifted to a preference for E isomer formation for secondary and tertiary alkyl substituents (Table 1, entries 1–7). The mesityl derivative, compound 11, however, was formed essentially as a single Z-isomer (Table 1, entries 11) while a maximum E-selectivity of 78% was achieved for the tert-butyl derivative, compound 3 (Table 1, entries 3). It thus appears that, for the alkyl derivatives at least, the orientation of the benzylidene phenyl group depends on the relative steric pressure imposed by the imino-isopropyl and isocyanate alkyl substituents. We tentatively suggest that this notable preference toward Z isomer formation for all aryl derivatives studied possibly reflects a stabilizing π-interaction between the benzylidene phenyl substituent and the N-aryl moieties during heterocycle assembly at the metal centre.

The reactions were easily translated to a preparative scale. The amidine, 1, was prepared on a 10 mmol scale using 2.5 mol% of Ic overnight in hexanes at 80 °C, and crystallized in 88% yield. The subsequent reaction of 1 with phenylisocyanate on a 0.88 mmol scale, using 2.5 mol% of Ic in 0.5 mL of hexane, was highly exothermic and crystallisation of the heterocyclic product, 10, commenced almost instantaneously. After one hour at room temperature the reaction mixture was filtered and the colourless solid dried in vacuo to yield 10 in near quantitative yield (285 mg, 93%). Single crystal X-ray diffraction experiments were performed on samples of compounds 3 and 12 which were obtained from the NMR scale reactions after quenching with methanol, filtration and slow evaporation of the solvent at 4 °C. The results of these experiments, which have been reported previously in preliminary form,19 are displayed in Figure 2 and show (E)-3 (left) and (Z)-12 (right), both of which correspond to the major isomer observed by 1H NMR spectroscopy (Table 1, entries 3 and 12). In both cases bond lengths and angles are within the range expected for these (5-benzylidene-4-imino)-imidazolidin-2-ones.34
Figure 2. ORTEP representations of compounds (Z)-3 (left) and (E)-12 (right). Ellipsoids at 30% probability. Hydrogen atoms omitted for clarity except for the benzylidene proton H4.

The influence of amidine N-substitution on the reaction was evaluated through the synthesis of a number of phenylpropargylamidines from phenylacetylene and commercially available carbodiimides using 5 mol% of Ic in C₆D₆ at 60 °C. Once full conversion to the amidine had been achieved one molar equivalent of iso-propylisocyanate was added and the reaction monitored by ¹H NMR spectroscopy. The rate of insertion and subsequent cyclisation was found to be highly dependent on the steric demands of the amidine N-substituents (Scheme 4). Whereas di(iso-propyl)- and dicyclohexylcarbodiimide-derived phenylpropargylamidines provided instantaneous and quantitative conversion to the respective heterocyclic products 7 and 13 at room temperature (Table 2, entries 1–2), the much more sterically hindered di(tert-butyl) derivative required 40 hours at room temperature to reach only 75% conversion to heterocycle 14 (Table 2, entry 3). Although all three symmetrical (N,N'-dialkyl)phenylpropargylamidines displayed a bias towards the E-isomer, this was significantly more pronounced for the dicyclohexyl and di(tert-butyl) substituted imidazolidin-2-ones, compounds 13 and 14 (Table 2, entries 1–3). Notably the unsymmetrical [1-ethyl-3-(tert-butyl)]phenylpropargylamidine precursor, 15, led to exclusive isocyanate insertion at the less hindered N-ethyl nitrogen atom, yielding a single heterocyclic product, compound 16. The Z:E isomer ratio of this latter species was intermediate between that of the di(isopropyl) and di(tert-butyl) derivatives (Table 2, entry 4). Carbodiimides with less sterically demanding N-substituents, such as di(p-tolyl)carbodiimide and [1-(N,N'-dimethylaminopropyl)-3-tert-butyl]carbodiimide, proved unsuitable for simple
hydroacetylenation in the first step of the reaction. Rather, they underwent twofold carbodiimide insertion followed by intramolecular hydroamination/cyclisation to yield the $N,N'$-[(5-benzylidene-imidazolidin-2,4-ylidene)diamine products 17 and 18 (Scheme 4, Table 2, entries 5–6). Sufficiently selective access to the desired propargylamidines could also not be achieved through application of the less reactive magnesium precatalyst, Ia, although in this case double carbodiimide insertion and the rate of subsequent intramolecular hydroamination were significantly perturbed. 75% $Z$-selectivity was observed in the $N,N',N''$-tetra-phenyl product, 18, in line with the $Z$-selectivity observed for arylisocyanate insertion/cyclisation (Table 2, entry 6). Unfortunately, unlike the unsymmetrical carbodiimide-derived product, 16, no regioselectivity was observed when employing [1-$(N,N'$-dimethylaminopropyl)-3-ethyl] carbodiimide presumably due to the lack of steric discrimination between the two carbodiimide substituents. In this case in the formation of all four possible insertion regioisomers, as well as $E/Z$-isomerism for each of them, was observed (Scheme 4; Table 2, entry 5).
Table 2. Carbodiimide scope for the synthesis of (5-benzylidene-4-imino)imidazolidin-2-ones from phenylacetylene and iso-propylisocyanate using 5 mol% of Ic in C₆D₆ at room temperature.

<table>
<thead>
<tr>
<th>Entry</th>
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<th>R'</th>
<th>Product</th>
<th>Time (h)</th>
<th>NMR yield (%)</th>
<th>Z:E (%)</th>
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<td>1</td>
<td>iPr</td>
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<tr>
<td>5</td>
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<td>(CH₂)₃NMe₂</td>
<td>17ᵇ</td>
<td>4</td>
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<tr>
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<td>18ᵇ</td>
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<td>97</td>
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ᵃ Determined by integration of benzylidene protons against an internal HN(SiMe₃)₂ standard.
ᵇ Two eq. RN=C=NR', 80 °C. Use of Ia instead of Ic did not afford better selectivity for the formation of the desired propargylamidine over the double carbodiimide insertion/cyclisation products 17 and 18.
ᶜ Mixture of all four possible regioisomers.

The influence of acetylenic substitution was also investigated. The strontium-catalysed reaction of iso-propyl- or phenylisocyanate with (N,N-di:iso-propyl)-n-butylpropargylamidine did not provide the desired heterocycles. Rather the reactions stalled after formation of the linear urea molecules, compounds 19 and 20, resulting from isocyanate insertion (Scheme 5). The latter species did not undergo intramolecular hydroamination/cyclisation even after prolonged heating at 100 °C. Similar observations have been made in the intramolecular hydroamination of aminoalkenes, in which cyclisation is hindered by the presence of terminal alkyl substituents on the alkene moiety, whereas terminal aryl substitution promotes cyclisation due to an activating electronic effect.¹⁶,¹⁷

In contrast, aryl or heteroaryl substitution on the acetylenic fragment afforded clean and near-quantitative formation of the desired imidazolidin-2-ones in stepwise reactions with di(iso-propyl)carbodiimide and iso-propylisocyanate using 5 mol% of Ic (Scheme 6). While under these conditions intermolecular hydroamination of the isocyanate with the propargylamidine remained instantaneous and quantitative in all cases, the subsequent intramolecular
cyclisation step proved highly dependent on the substitution pattern of the acetylenic aryl moiety. Ortho-substitution, with substrates such as o-tolylacetylene and 1-naphthylacetylene, resulted in significantly reduced cyclisation rates compared to the parent phenylacetylene (Table 3, entries 2–3). Para-methyl-substitution, however, had no effect on the efficacy of the intramolecular hydroamination step (Table 3, entry 4). It is also notable that the Z:E isomer ratio was unaffected by alkyl or aryl substitution on the phenyl ring (Table 3, entries 1–4). For para-substituents a clear trend of decreasing rate of cyclisation with increasing substituent electron-withdrawing effect was observed; from 1 hour to achieve 96% conversion to the para-chloro derivative, 25, to 24 hours to reach 90% conversion for the para-trifluoromethyl and para-fluoro derivatives, 27 and 28 (Table 3, entries 6–10). This trend contrasts with our previous observations in which the rate of cyclisation during the intramolecular hydroalkoxylation of hydroxyalkynes bearing para-substituted terminal aryl groups was seen to increase with increasingly electron-withdrawing substitution in para position.37 We tentatively ascribe the apparent reversal in reactivity in the present instance to a more remote effect of stronger electron-withdrawing para-substitution on the polarisation of the alkaline earth amidinate nitrogen centres and consequent less facile isocyanate insertion. In mitigation of this hypothesis, monitoring of these reactions by 1H NMR spectroscopy revealed that isocyanate insertion into the more electron-poor amidines is less rapid and provides lower conversion than with the more electron-rich amidines. In contrast meta-fluoro-substitution resulted in only a small decrease in reactivity, yielding 98% of the heterocyclic product, 24, in 1.5 hours (Table 3, entry 5). The nature of the electron-withdrawing substituent also influences the Z:E selectivity, although no discernible pattern could be discriminated.

It is notable that these reactions were tolerant of chloro-substitution on the phenyl ring, providing the potential for further catalytic functionalisation of the aryl moiety. After quenching and extraction with methanol, compound 25 crystallized in good yield (85%) upon solvent evaporation at room temperature. Figure 3 shows the results of a single crystal X-ray diffraction experiment on the major (E)-isomer. Heteroaryl-substituted heterocycles could also be obtained in this manner. N,N'-di-iso-propyl-3-pyridinylpropargylamidine was synthesised using the magnesium precatalyst, Ia, in preference to the strontium precursor, Ic, which was found to provide twofold carbodiimide insertion and cyclisation. Under these conditions, insertion of iso-propylisocyanate and cyclisation were very rapid and effectively quantitative, to yield the pyridine-substituted heterocycle, 29, in excellent yield (Table 3, entry 11). Attempts to synthesise the 2-pyridinyl analogue, however, were less discriminating.
as even the use of Ia resulted in competitive carbodiimide insertion, with ESI-MS analysis detecting molecular weights indicative of 1:1, 1:2, 2:1, 2:2 and even 3:2 acetylene:carbodiimide ratios. The strontium-catalysed reaction of 3-thiophenylacetylene with di(iso-propyl)carbodiimide did provide the desired amidine, allowing for subsequent isocyanate insertion and cyclisation. In this case, however, three different products were obtained: the expected Z and E isomers of heterocycle 30, present in a 63:37 ratio, as well as a new compound which was identified as the Diels-Alder adduct of two molecules of this heterocyclic product.

**Scheme 6**

\[
\begin{align*}
Ar-H + {}^t\text{PrN}=C=N^t\text{Pr} & \rightarrow \text{Ar}^N\text{Pr} \rightarrow \text{Ar}^N\text{Pr} + {}^t\text{PrN}=C=O \rightarrow \text{Ar}^N\text{Pr} \rightarrow \text{Ar}^N\text{Pr} \rightarrow \text{Ar}^N\text{Pr} \\
\text{Ar} = & \text{o-tol } 21, \text{ 1-naphthyl } 22, \text{ p-tol } 23, \text{ 3-FC}_6\text{H}_4 24, \text{ 4-ClC}_6\text{H}_4 25, \text{ 4-OMeC}_6\text{H}_4 26, \text{ 4-CF}_3\text{C}_6\text{H}_4 27, \text{ 4-FC}_6\text{H}_6 28, \text{ 3-pyridinyl } 29, \text{ 3-thiophenyl } 30
\end{align*}
\]

**Table 3.** Arylacetylene scope for the synthesis of (5-benzylidene-4-imino)imidazolidin-2-ones from di(iso-propyl)carbodiimide and iso-propylisocyanate using 5 mol% of Ic in C₆D₆ at room temperature.

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<td>4-CF₃-C₆H₄</td>
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<tr>
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<td>&quot;</td>
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<td>2</td>
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<td>11</td>
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<td>29</td>
<td>25</td>
<td>0.1</td>
<td>&gt;99</td>
<td>38:62</td>
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<tr>
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<td>3-thiophenyl</td>
<td>30</td>
<td>25</td>
<td>0.1</td>
<td>&gt;99</td>
<td>63:37</td>
</tr>
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* Determined by integration of benzylidene protons against an internal HN(SiMe₃)₂ standard.

b 3-pyridinylpropargylamidine was synthesised using 5 mol% of Ia at 40 °C for 1 day, as Ic resulted in double insertion of the carbodiimide and formation of the imidazolidin-2,4-ylidene product analogous to 29.
A stoichiometric reaction between Ia, 2 equivalents of (N,N’-di-isopropyl)phenylpropargylamidine and 2 equivalents of 2,6-di-isopropyl-phenylisocyanate did not yield the expected magnesium insertion complex. Analysis by NMR spectroscopy instead indicated complete consumption of the substrates to form the corresponding N-heterocyclic product, 3, with reformation of Ia. This result suggests that cyclisation is assisted by the protic [HN(SiMe₃)₂] liberated upon protonolysis of Ia with the amidine. A similar concerted insertion/protonolysis cyclisation mechanism has been proposed for the intramolecular hydroamination of aminoalkenes, based on the observation of high KIE values and stoichiometric reactions.¹⁷,²³

In contrast, a further stoichiometric reaction utilising [Mg(CH₂Ph)₂(THF)₂] in place of Ia, and with consequent liberation of the aprotic toluene conjugate acid, yielded the desired homoleptic insertion complex, compound II, in quantitative yield (Scheme 7). An X-ray diffraction experiment, preliminary details of which were included in our previous communication,¹⁹ revealed II to be a distorted square pyramidal magnesium 2-(propargylamidino)imidate complex, with a THF molecule coordinated in the apical position.
Coordination in the basal plane is provided by the oxygen atoms arising from the inserted isocyanate substrates and the imino-nitrogen of the amidinate moieties to form a quasi-planar 6-membered [MgNCNCO] metallacycle (Figure 4). The rather short C-O bond lengths [1.275(4), 1.274(4) Å], lengthened C1-N1 [1.451(4) Å] and C29-N4 bonds [1.452(4) Å] and planarity of the N1 and N4 nitrogen atoms within II suggest some degree of delocalization over the chelate ring. The short C1-N3 [1.289(4) Å] and C29-N6 [1.291(4) Å] bond lengths are clearly indicative of pendant imine functionalities. Isolated samples of complex II also provided similar catalytic activity to Ia for the formation of 3, suggesting it is a catalytic intermediate.

Figure 4. ORTEP representation of compound II. Ellipsoids at 30% probability. Hydrogen atoms omitted for clarity.

A variable temperature \(^1\)H NMR experiment performed on II in \(d_8\)-toluene provided evidence for isocyanate de-insertion at higher temperatures (Scheme 8). A van’t Hoff analysis of this equilibrium provided \(\Delta H^\ddagger = +88\) kJ mol\(^{-1}\) and \(\Delta S^\ddagger = 208\) J K\(^{-1}\) mol\(^{-1}\), yielding a \(\Delta G^\ddagger(298\) K)
value of +26 kJ mol\(^{-1}\) for the de-insertion process. Although a definitive interpretation of this latter value would require deconvolution of both the de-insertion and potential dimerization of the resultant bis(propargylamidinate) species, III, this positive but low free energy of activation at 298 K indicates that the potential for this reversibility is likely to be significant during the course of the catalysis at ambient or slightly elevated temperatures.

**Figure 5.** Proposed mechanism for the independent formation of E- and Z-heterocycles.

Based on the structure of II and the selectivity of the reactions toward the formation of the kinetic imidazolidin-2-one products rather than the thermodynamic 2-iminooxazolidines, we propose that the N,O-chelate, A in Figure 5, first isomerizes to the corresponding N,N-chelate (B) by decooordination of the oxygen atom and rotation of the imidate imino residue to coordinate to the metal centre. The metal-bound nitrogen atom of the amidino fragment must then necessarily decoordinate and rotate to allow the alkyne to interact with the metal centre. This may result in either a C\(_{\text{chair}}\) or a C\(_{\text{boat}}\) conformation, giving rise, upon proton-assisted insertion/cyclisation to the heterocyclic products (E)-D and (Z)-D, respectively. Taking into account all these observations the catalytic cycle presented in Figure 6 may be envisaged.
Figure 6. Mechanism for the alkaline earth-catalysed formation of imidazolidin-2-ones from arylpropargylamidines and isocyanates.

The insertion/cyclisation reactivity was further extended to a variety of isothiocyanates, yielding the corresponding imidazolidin-2-thiones, compounds 31 – 35 (Scheme 9). All such compounds presented a characteristic $^{13}$C NMR thione resonance around 180 ppm, as well as two distinct $^1$H NMR benzylidene singlets at ca. 6.5 ppm and 5.9 ppm corresponding to the Z and E isomers, respectively. Reactions using 0.5 mol% of Ia were significantly slower than for the corresponding isocyanates. Quantitative conversion to the imidazolidin-2-thiones products was, however, achieved within 30 minutes at room temperature 5 mol% of Ic.

Scheme 9

$\text{Ph} = \text{Ph}$ 31, $^4\text{BuC}_6\text{H}_4$ 32, Et 33, $^1\text{Pr}$ 34, Cy 35.
Table 4. Isothiocyanate scope for the catalytic synthesis of (5-benzylidene-4-imino)imidazolidin-2-thiones from 1 using 5 mol% of Ic in C₆D₆.

<table>
<thead>
<tr>
<th>Entry</th>
<th>R</th>
<th>Product</th>
<th>T (°C)</th>
<th>Z/E (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
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<td>25</td>
<td>97:3</td>
</tr>
<tr>
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<td>4-tBu-C₆H₄</td>
<td>32</td>
<td>25</td>
<td>95:5</td>
</tr>
<tr>
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<td>iPr</td>
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<tr>
<td>6</td>
<td>tBu</td>
<td>–</td>
<td>80</td>
<td>–</td>
</tr>
<tr>
<td>7</td>
<td>Ad</td>
<td>–</td>
<td>80</td>
<td>–</td>
</tr>
</tbody>
</table>

Unlike the isocyanate-based reactions, the linear isothiocyanate insertion intermediates analogous to 2 were not distinguishable by ¹H NMR spectroscopy due to broadening of the resonances. The Z:E isomer distribution of the products, quantified by integration of the ¹H NMR singlets of the benzylidene protons, however, followed similar patterns to that of the imidazolidin-2-ones. Aryl substituents and the less sterically hindering ethyl functionality predominantly yielded the Z isomer (Table 4, entries 1-3) while the more sterically demanding iso-propyl and cyclohexyl derivatives yielded the E isomer as the major product (Table 4, entries 4-5). The steric influence of the isothiocyanate alkyl substituent appeared, however, more pronounced than for the corresponding isocyanates, presumably due to the greater steric repulsion of the exocyclic sulfur versus oxygen atom. In contrast to the isocyanate-based reactions no observable isothiocyanate insertion was evident even at high temperature for the larger tert-butyl and adamantyl derivatives (Table 4, entries 6-7). The isolated imidazolidin-2-thiones proved very moisture-sensitive and had to be isolated in a glovebox. An X-ray diffraction analysis experiment performed on single crystals of the 4-tert-butylphenyl derivative yielded the structure of the predominant Z-isomer, (Z)-32, (Figure 7) clearly revealing the maintenance of the thione functionality [C=S 1.654(2) Å].¹⁹
Insight into the possible course of reaction during the synthesis of compounds 31 – 35 was provided by a reaction between the homoleptic calcium (\(N,N'\)-di-iso-propyl)phenylpropargylamidinate dimer, IV, and two molar equivalents of (para-tert-butyl)phenylisothiocyanate in toluene. This process provided clean access to the heteroleptic calcium compound, V (Scheme 10), which a single crystal X-ray crystallographic analysis revealed to be a dimeric calcium species bearing terminal amidinate ligands and bridging (2-propargylglylamidino)imidothioate ligands (Figure 8). The latter units form a six-membered \(N,S\)-chelate, with the sulfur atom bridging between both calcium centres, and coordination to the second alkaline earth centre through the pendant \(N-(4\text{-}\text{tert}-\text{butylphenyl})\text{imino nitrogen}\). The sulphur-containing ligand is broadly analogous to the isocyanate-derived magnesium chelate within the homoleptic species, compound II. We, thus, suggest the formation of the catalytic production of the imidazolidin-2-thione derivatives also follows a similar mechanistic pathway, wherein the imidazolidine products bearing an exocyclic thione moiety are formed in kinetic preference to the more thermodynamically stable thiazole heterocycles.
**Figure 8:** ORTEP representation of compound V. Ellipsoids at 30% probability. Hydrogen atoms and iso-propyl methyl groups omitted for clarity.

Although no evidence for the formation of the alternative thiazolidine isomers was observed during the synthesis of compounds 31 – 35, a further extension of this protocol demonstrated that the thiazolidine ring system was accessible through the strontium-catalysed reaction of carbon disulphide with \((N,N'\text{-di-iso-propyl})\)phenylpropargyl-amidine. At 60 °C, CS$_2$ insertion followed by intramolecular hydrothiolation, proceeded smoothly to yield the (4-benzylidene-5-imino)thiazolidin-2-thione product, 36, which, over a period of 48 hours, fully tautomered to the corresponding 5-benzyl-1,3-thiazole-2-thione, 37 (Scheme 11).

![Scheme 11](image)

**Conclusion**

In conclusion we have demonstrated the applicability of readily available and sustainable alkaline earth bis(amide) precatalysts to the facile one-pot, 100% atom-efficient, stepwise
synthesis of a wide variety of highly functionalized imidazolidin-2-ones, imidazolidin-2-thiones, \(N,N'\)-[(5-benzylidene-imidazolidin-2,4-ylidene)diamines and 1,3-thiazolidin-2-thiones from the simplest commercially available building blocks. Although the current investigation is relatively broad in scope, we suggest that the formation of compound 37 indicates that even broader manifestations of this protocol may be achieved through the incorporation of further alternative heterocumulene substrates. Catalytic access to a variety of valuable heterocyclic derivatives will, thus, be easily accessible in a cost-efficient manner and with complete atom-efficiency. We are continuing to explore these possibilities and will describe our investigations in future publications.

**Associated Content**

**Supporting Information.** Experimental procedures and full characterisation data, details of the X-ray analyses of compounds \(Z\)-3, \(E\)-12, \(E\)-25, \(Z\)-32, II and V (CCDC 1008314–1008316 and CCDC 1051077 - 1051079).

**Acknowledgements**

We thank the EPSRC (UK) for generous funding.

**Notes and References**

Structurally diverse imidazolidine-2-ones and -thiones, 2-imino-imidazolidines and thiazolidine-2-thiones may be synthesised using simple alkaline earth bis(amide) precatalysts with 100% atom-efficiency in one-pot cascades from commercially available terminal alkyne and heterocumulene reagents.