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## Ruthenium-catalysed cyclisation reactions of 1,11-dien-6-yne s leading to biinden es<sup>†</sup>

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1,2-Bis(2-allylphenyl)ethynes undergo cycloisomerisation reactions in the presence of  $\text{Cp}^*\text{Ru}(\text{ii})$  catalysts to produce 2,2'-dimethyl-3H,3'H-1,1'-biinden es. On the other hand, tandem ring-closing metathesis of 1,2-bis(2-allylphenyl)ethynes using the Hoveyda–Grubbs 2nd generation catalyst led to the formation of 2,2'-unsubstituted biinden es. Various symmetrical and unsymmetrical bicyclic dienes were prepared by these ruthenium-based cyclisation methods.

## Introduction

3H,3'H-1,1'-Biinden es have previously been prepared by the oxidative homocouplings of (1H-inden-1-yl)lithiums to yield diastereomeric mixtures of 1H,1'H-1,1'-biinden es, followed by base-promoted double bond isomerisation.<sup>1</sup> There are fewer than fifty known biinden es, and some of them have been used as ligands for transition metals,<sup>2</sup> while a biindene-derived diol has been used as a chiral ligand in the titanium(IV)-catalysed enantioselective additions of diethylzinc to aldehydes.<sup>3</sup>

Transition-metal-catalysed cycloisomerisation reactions of enynes are powerful tools for the synthesis of various carbon- and heterocyclic compounds.<sup>4</sup> This method allows for the rapid atom-economical construction of a complex cyclic structure from a linear substrate. The ring-closing metathesis (RCM) of dienes and enynes revolutionised the way in which cycloalkenes are assembled, and has been extremely useful in modern organic synthesis.<sup>5</sup> Herein, we report that 1,11-dien-6-yne s can undergo both cycloisomerisation and tandem RCM reactions catalysed by ruthenium complexes. Notably, these reactions are used to prepare 1,1'-biinden es from 1,2-bis(2-allylphenyl)ethynes.

## Results and discussion

When 1,2-bis(2-allylphenyl)ethyne (**1a**)<sup>6</sup> was heated at 60 °C in EtOH in the presence of 5 mol%  $\text{CpRuCl}(\text{PPh}_3)_2$  for 24 h, it cycloisomerised to afford 2,2'-dimethyl-3H,3'H-1,1'-biindene (**2a**) in 21% yield (Table 1, entry 1). The use of  $\text{Cp}^*\text{RuCl}(\text{PPh}_3)_2$  improved the yield of **2a** to 45% (entry 2), while the reaction in

the presence of  $\text{Cp}^*\text{RuCl}(\text{cod})$  afforded **2a** in 34% yield (entry 3). The use of a cationic ruthenium catalyst generated *in situ* from  $\text{Cp}^*\text{RuCl}(\text{cod})$  and  $\text{NaPF}_6$  gave **2a** in 55% yield (entry 4); indeed, preformed cationic  $[\text{Cp}^*\text{Ru}(\text{MeCN})_3]\text{PF}_6$  exhibited comparable activity (entry 5). The effect of the phosphine ligand was next examined; **2a** was formed in 38% yield when the  $\text{Cp}^*\text{RuCl}(\text{cod})$ -BINAP catalyst system was used (entry 6); however, use of  $\text{P}(\text{C}_6\text{F}_5)_3$  increased the yield of **2a** to 61%, together with a 24% yield of the [2 + 2 + 2] cycloadduct **3a** (entry 7).<sup>7,8</sup> The reaction performed in MeOH in the presence of  $\text{Cp}^*\text{RuCl}(\text{cod})$ - $\text{P}(\text{C}_6\text{F}_5)_3$  furnished **2a** in 85% isolated yield without any noticeable amount of **3a** (entry 8). A similar result was obtained when the reaction was performed at 40 °C (entry 9). Interestingly, the reaction delivered cycloadduct **3a** as the major product when performed in *i*-PrOH (entries 10 and 11).<sup>9</sup>

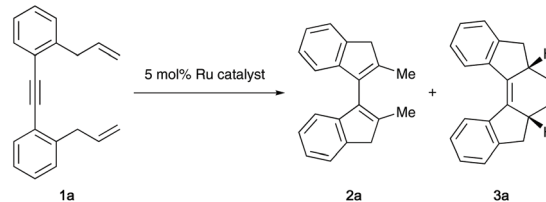
With the optimised reaction conditions in hand, various diallyl diphenylacetylenes **1b-l** bearing substituents on their benzene rings were subjected to the ruthenium-catalysed cycloisomerisation conditions (Table 2). The reaction of 1,2-bis(2-allyl-4-methylphenyl)ethyne (**1b**) afforded tetramethylbiindene **2b** in 67% yield (entry 1), whereas symmetrical dienynes **1c** and **1d** bearing methyl or methoxy groups at the 5 positions of their benzene rings afforded **2c** and **2d**, respectively, in good yields (entries 2 and 3). In contrast, the reactions of chloro- and trifluoromethyl-substituted dienynes **1e** and **1f** formed the [2 + 2 + 2] cycloadducts **3** as major products under the standard conditions (**1e**: **2e** 24% + **3e** 41%; **1f**: **3f** 87%). The cycloisomerisation products from **1e** and **1f** were obtained as the major products in yields of 44% and 29%, respectively, when the reaction was performed with  $[\text{Cp}^*\text{Ru}(\text{MeCN})_3]\text{PF}_6$  (entries 4 and 5). The naphthalene derivative **1g** was also converted into the corresponding product **2g** (entry 6), while unsymmetrically substituted biinden es **2h-l** were similarly prepared by cycloisomerising dienynes **1h-l** (entries 7–11).

The cycloisomerisation conditions were successfully applied to dienye **1m** devoid of *o*-phenylene tethers, which

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Table 1 Ruthenium-catalysed cycloisomerisation of 1,2-bis(2-allylphenyl)ethyne (1a)<sup>a</sup>

Entry	Ru catalyst	Ligand (mol%)	Additive	Solvent	Temp. (°C)	Time (h)	Yield <sup>b</sup> (%) of 2a	Yield <sup>b</sup> (%) of 3a
1	CpRuCl(PPh <sub>3</sub> ) <sub>2</sub>	—	—	EtOH	60	24	21	
2	Cp <sup>*</sup> RuCl(PPh <sub>3</sub> ) <sub>2</sub>	—	—	EtOH	60	4	45	
3	Cp <sup>*</sup> RuCl(cod)	—	—	EtOH	60	24	34	
4	Cp <sup>*</sup> RuCl(cod)	—	NaPF <sub>6</sub>	EtOH	60	24	55	
5	[Cp <sup>*</sup> Ru(MeCN) <sub>3</sub> ]PF <sub>6</sub>	—	—	EtOH	60	12	56	
6	Cp <sup>*</sup> RuCl(cod)	rac-BINAP (5)	—	EtOH	60	24	38	
7	Cp <sup>*</sup> RuCl(cod)	P(C <sub>6</sub> F <sub>5</sub> ) <sub>3</sub> (10)	—	EtOH	60	24	61	24
8	Cp <sup>*</sup> RuCl(cod)	P(C <sub>6</sub> F <sub>5</sub> ) <sub>3</sub> (10)	—	MeOH	60	24	85	
9	Cp <sup>*</sup> RuCl(cod)	P(C <sub>6</sub> F <sub>5</sub> ) <sub>3</sub> (10)	—	MeOH	40	24	87	
10	Cp <sup>*</sup> RuCl(cod)	P(C <sub>6</sub> F <sub>5</sub> ) <sub>3</sub> (10)	—	<i>i</i> -PrOH	60	24	11	55
11	Cp <sup>*</sup> RuCl(cod)	P(C <sub>6</sub> F <sub>5</sub> ) <sub>3</sub> (10)	—	<i>i</i> -PrOH	40	24	14	75

<sup>a</sup> Reaction conditions: 1a (0.050 mmol), ruthenium catalyst (2.5 μmol, 5 mol%), ligand (Ru : P = 1 : 2), solvent (0.5 mL, 0.1 M). <sup>b</sup> Isolated yield.

led to the formation of 1,1'-bicyclopentene **2m** in 82% yield in the presence of [Cp<sup>\*</sup>Ru(MeCN)<sub>3</sub>]PF<sub>6</sub> (Scheme 1, (a)). The alternative cycloisomerisation product **4m** was obtained in 55% yield when **1m** was reacted at 0 °C (Scheme 1, (b)).<sup>10,11</sup> Heating **4m** in the presence of the ruthenium catalyst gave **2m** in 49% yield, but no isomerisation was observed in the absence of the ruthenium catalyst. Based on these results as well as previous studies, we conclude that 2-methylene-1,1'-bi(cyclopentylidene) **4m** is the initial cycloisomerisation product, and that **4m** is also catalytically isomerised to **2m** by the ruthenium catalyst.

Two possible reaction pathways can be proposed for the ruthenium-catalysed cycloisomerisation of 1,11-dien-6-yne **1** (Scheme 2). Path (a) involves the formation of a hydroruthenium species from the catalyst and MeOH,<sup>12</sup> a Markovnikov hydroruthenation to the C=C bond of **1** to form **A**, consecutive carboruthenation (through **B** to **C**), β-hydride elimination that releases **4**, and the final double bond isomerisation of **4** to afford product **2**. On the other hand, in path (b), diyne **1** first undergoes oxidative cyclisation on ruthenium to generate the ruthenacyclopentene species **D**. The unreacted alkene moiety in **D** then inserts into the Ru-C(sp<sup>2</sup>) bond to give the ruthenacycloheptene intermediate **E**. Subsequent β-hydride elimination (to form **F**) followed by reductive elimination yields **4**, which then isomerises to **2** catalysed by a hydroruthenium species. Alternatively, β-hydride elimination from **D** generates alkenylruthenium hydride **G**, which also leads to **4** through intramolecular carboruthenation (to **F**) or hydroruthenation (to **H**). Reductive elimination from intermediate **E** is possible, which gives rise to the [2 + 2 + 2] cycloadduct **3**.

Diene **1n** or **1o**, in which one allyl group is replaced with a crotyl or a methallyl group, was found to be unreact-

tive toward cycloisomerisation, which reveals that the reaction is limited to diynes with unsubstituted C=C double bonds (Chart 1). Moreover, 1,2-bis[2-(vinyloxy)phenyl] ethyne (**1p**) also failed to react, and a complex mixture of products was obtained when unsymmetrical diyne **1q**, bearing malonate and *o*-phenylene tethers, was reacted.<sup>13</sup>

We have been interested in the catalytic syntheses of silole derivatives<sup>14</sup> and the cycloisomerisation of bis-silicon-bridged **1r** was envisaged as a method for the synthesis of a bi(1-siladene).<sup>15</sup> However, the reaction of **1r** under conditions similar to those described above led to a totally different outcome: 1,1',2,2'-tetrahydro-4,4'-bi(1-silanaphthalene) **5** was obtained in 43% yield as the sole product after full conversion of **1r** (Scheme 3). The silanaphthalene **5** may have formed through a stitching reaction mediated by a hydroruthenium species in a manner analogous to the path (a) in Scheme 2, but with initial anti-Markovnikov hydroruthenation.

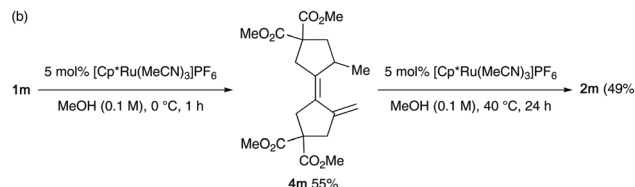
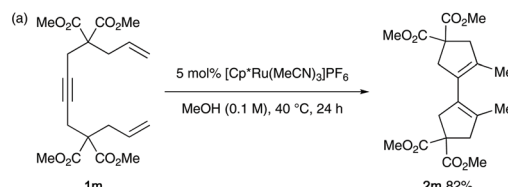
Tandem ring-closing metathesis (RCM) of 1,11-dien-6-yne that form 1,1'-bicyclopentene derivatives has previously been studied,<sup>16</sup> but those of 1,2-bis(2-allylphenyl) ethynes have, to the best of our knowledge, never been examined. If allowed, this reaction provides a route to 3,3'H-1,1'-biindenes that lack substituents at their 2 and 2' positions, which is complementary to the cycloisomerisation of **1**. Tandem RCM of **1a** in the presence of the Hoveyda-Grubbs 2nd generation catalyst at 100 °C in toluene (0.1 M) afforded the desired biindene **6a** in 60% yield (Table 3, entry 1). A lower concentration of **1a** resulted in an improved yield of **6a**, and 0.02 M was found to be optimal for the present reaction (entries 2 and 3). Other Grubbs catalysts were not suitable for this transformation (entries 4 and 5), while the reaction with 3 mol% catalyst gave a



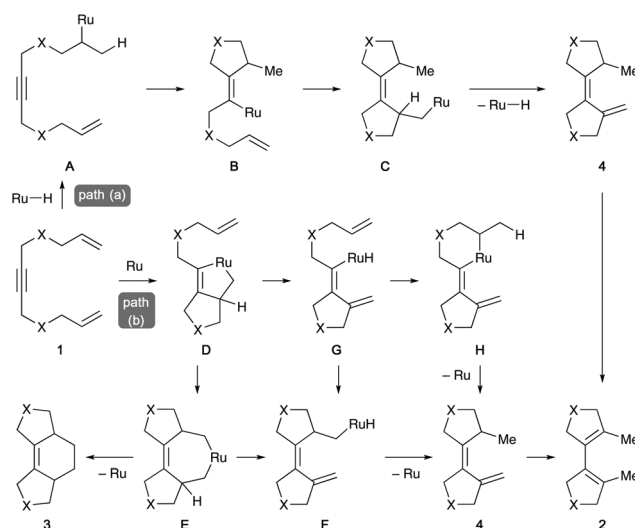
**Table 2** Cycloisomerisation of dienynes **1**

Entry	Dienyne 1	Product 2	Yield <sup>a</sup> (%)
1			67
2			
3			72
4 <sup>b,c,d</sup>			63
5 <sup>b,d</sup>			44
6 <sup>e</sup>			29
7			39
8			90
9			86
10 <sup>d</sup>			84
11 <sup>d</sup>			55
			45

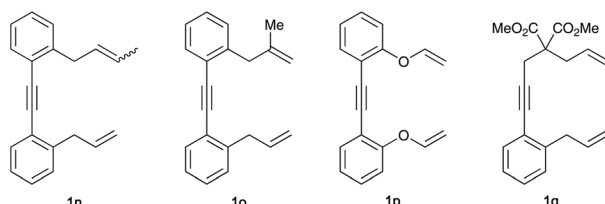
<sup>a</sup> Isolated yield (average of two runs). <sup>b</sup> 5 mol% [Cp\*Ru(MeCN)<sub>3</sub>]PF<sub>6</sub> was used as catalyst. <sup>c</sup> Reaction was performed at 60 °C. <sup>d</sup> The crude reaction mixtures contained byproducts such as 3. <sup>e</sup> Reaction was performed at 80 °C in MeOH (0.05 M).



**Scheme 1** Cycloisomerisation of **1m**.



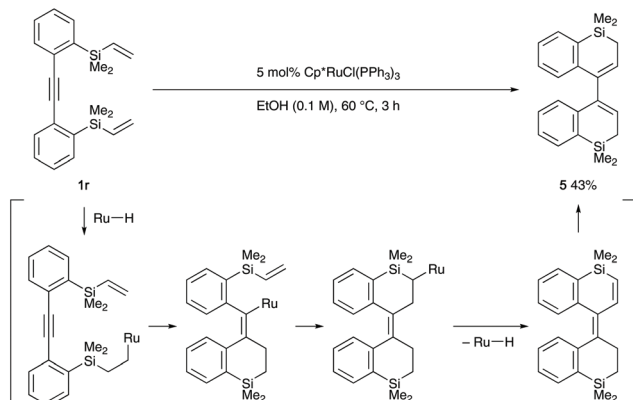
**Scheme 2** Possible reaction pathways for the cycloisomerisation of **1m** ( $X = C_6\text{F}_5\text{CO}_2\text{Me}_2$ )



**Chart 1** Dienynes that failed to undergo ruthenium-catalysed cycloisomerisation

similar result (entry 6). As for the reaction temperature, 100 °C was found to be the best among those examined for the RCM of **1** (entries 6–8).

Various diallyl diphenylacetylenes **1b–l**, which were successfully cycloisomerised (*vide supra*), were examined under the RCM conditions (Table 4). Symmetrical (**1b–g**) and unsymme-

Scheme 3 Cycloisomerisation of **1r**.Table 3 Tandem RCM of **1a**

Entry	Grubbs catalyst (mol%)	Conc. (M)	Temp. (°C)	Time (h)	Yield <sup>a</sup> (%)
1	Hoveyda–Grubbs 2nd cat. (5)	0.1	100	0.5	60
2	Hoveyda–Grubbs 2nd cat. (5)	0.04	100	1	67
3	Hoveyda–Grubbs 2nd cat. (5)	0.02	100	3	77
4	Grubbs 2nd cat. (5)	0.02	100	6	44
5	Stewart–Grubbs cat. (5)	0.02	100	6	26
6	Hoveyda–Grubbs cat. 2nd (3)	0.02	100	6	82
7	Hoveyda–Grubbs cat. 2nd (3)	0.02	90	6	67
8	Hoveyda–Grubbs cat. 2nd (3)	0.02	110	6	74

<sup>a</sup> Isolated yield.

trical (**1h–l**) diynes were converted through tandem RCM into biindenenes **6b–l** in yields ranging from 63% to 96%. Furthermore, diynes **1o–q**, which failed to cycloisomerise, also reacted to afford the corresponding metathesis products **4o–q**, respectively, in good yields. However, the attempted tandem RCM of the bis-silicon-bridged **1r** resulted in no conversion under various metathesis conditions.

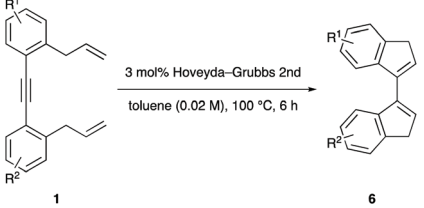
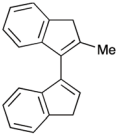
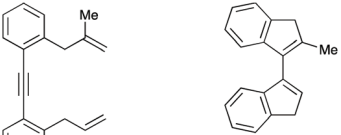
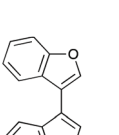
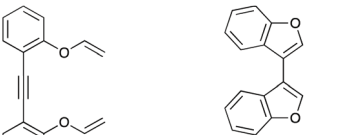
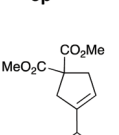
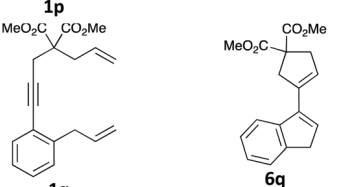
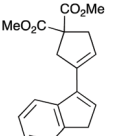
## Conclusions

In conclusion, we developed ruthenium-catalysed cycloisomerisation and tandem-RCM methods for the synthesis of bicyclic conjugated dienes, in which two rings (cycloalkenes) are constructed. Cycloisomerisation of 1,11-dien-6-yne afforded 2,2'-dimethyl-[1,1'-bi(cyclopentene)] derivatives catalysed by  $\text{Cp}^*\text{Ru}$ . On the other hand, 2,2'-unsubstituted bicyclopentenes were prepared through the tandem RCM of 1,11-dien-6-yne with the Hoveyda–Grubbs catalyst.<sup>17</sup>

Table 4 Tandem RCM of diynes **1**

Entry	Dienyne <b>1</b>	Product <b>6</b>	Yield <sup>a</sup> (%)
1	<b>1b</b>	<b>6b</b>	74
2	<b>1c</b> (R = Me)	<b>6c</b>	86
3	<b>1d</b> (R = OMe)	<b>6d</b>	63
4	<b>1e</b> (R = Cl)	<b>6e</b>	96
5	<b>1f</b> (R = CF <sub>3</sub> )	<b>6f</b>	83
6	<b>1g</b>	<b>6g</b>	68
7	<b>1h</b>	<b>6h</b>	90
8	<b>1i</b> (R = Me)	<b>6i</b>	77
9	<b>1j</b> (R = OMe)	<b>6j</b>	74
10	<b>1k</b> (R = Cl)	<b>6k</b>	80
11	<b>1l</b> (R = CF <sub>3</sub> )	<b>6l</b>	85

Table 4 (Contd.)

Entry	Dienyne <b>1</b>	Product <b>6</b>	Yield <sup>a</sup> (%)
12			75
13			93
14			85
			

<sup>a</sup> Isolated yield (average of two runs).

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

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8 Results with other phosphine ligands (**2a/1a + 3a**): PPh<sub>3</sub> (12%/30%); P(c-Hex)<sub>3</sub> (8%/38%); P(OPh)<sub>3</sub> (18%/69%); DPPE (6%/30%); DPPBZ (25%/62%).

9 The formation of a ruthenium hydride might be suppressed in *i*-PrOH in our case; hence the reaction is directed to proceed via the path (b) in Scheme 2. The results in other solvents are as follows: *t*-BuOH (**2a** not detected, **3a** 11%); no reaction in (CF<sub>3</sub>)<sub>2</sub>CHOH.

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