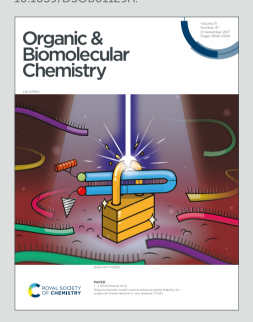


# Organic & Biomolecular Chemistry



**Accepted Manuscript** 

This article can be cited before page numbers have been issued, to do this please use: J. C. Morales-Solís, M. Ordoñez, E. Marqués López and R. P. Herrera, *Org. Biomol. Chem.*, 2025, DOI: 10.1039/D5OB01129A.



This is an Accepted Manuscript, which has been through the Royal Society of Chemistry peer review process and has been accepted for publication.

Accepted Manuscripts are published online shortly after acceptance, before technical editing, formatting and proof reading. Using this free service, authors can make their results available to the community, in citable form, before we publish the edited article. We will replace this Accepted Manuscript with the edited and formatted Advance Article as soon as it is available.

You can find more information about Accepted Manuscripts in the <u>Information for Authors</u>.

Please note that technical editing may introduce minor changes to the text and/or graphics, which may alter content. The journal's standard <u>Terms & Conditions</u> and the <u>Ethical guidelines</u> still apply. In no event shall the Royal Society of Chemistry be held responsible for any errors or omissions in this Accepted Manuscript or any consequences arising from the use of any information it contains.



View Article Online DOI: 10.1039/D5OB01129A

## **REVIEW**

Received 00th January 20xx.

# Organocatalysed C-2 and C-3 Functionalisation of Chromones

Juan Carlos Morales-Solís, a,b Mario Ordoñez, Eugenia Marqués-López\*a and Raquel P. Herrera\*a

Accepted 00th January 20xx

DOI: 10.1039/x0xx00000x

This review highlights the recent advancements in organocatalysed Michael addition reactions involving chromones, an important class of heterocyclic compounds with significant biological and pharmacological properties. Chromones, with their versatile conjugated structure, act as both electrophilic and nucleophilic partners, providing an ideal platform to synthesise diverse stereochemically enriched molecular frameworks. The work highlights various organocatalysed examples using chiral carbenes, phosphoric acids, thioureas and squaramides, that have demonstrated high efficiency and stereocontrol in a range of reactions. Mechanistic insights into how these catalytic systems activate the chromone scaffold and promote enantioselective transformations are also discussed in detail. Finally, we underline emerging trends and future directions in this research area, addressing current limitations, such as the need for more efficient catalytic systems and broader substrate compatibility.

#### Introduction

The field of organocatalysis has gained significant attention since the early 2000s.¹ Organocatalytic methods represent an exciting and sustainable approach to chemical synthesis of complex and chiral molecules, which are particularly important in medicine and the pharmaceutical industry,² where the stereochemistry of a molecule can drastically affect its biological activity.³ Additionally, this field is recognised for its advantages in terms of environmental impact, reaction versatility, and access to enantiomerically enriched products, making it a valuable tool for chemists in academic, industrial, and pharmaceutical research.

Among the plethora of organic reaction developed in this field, the Michael addition reaction is a cornerstone transformation in organic chemistry, particularly valued for its ability to create carbon-carbon and carbon-heteroatom bonds.<sup>4</sup> This kind of reaction provides a powerful alternative for building complex molecular frameworks, especially in the synthesis of natural products, pharmaceuticals, and advanced materials. One interesting application of the Michael addition involves the use of chromones as electrophilic (nucleophilic) partners.<sup>5</sup>

Chromones (1-benzopyran-4-ones) are heterocyclic compounds containing a benzopyran core with a ketone group. The chromone skeleton is found in a wide variety of naturally occurring compounds, including flavonoids, coumarins, and certain alkaloids, many of which exhibit valuable

**Scheme 1** Activated electrophilic/nucleophilic **C-2** and **C-3** positions for conjugated addition reactions in chromones.

One key feature of chromones is the conjugation between the carbonyl group at C-4 and the adjacent double bond. This conjugation not only stabilises the molecule but also broadens the reactivity potential of the chromone nucleus.

The resulting chromanone adducts also constitute an important structural motif found in many bioactive and natural compounds, making this reaction valuable for the synthesis of pharmacologically active molecules (Figure 1). This reaction opens new pathways for constructing complex molecules, including those with potential anticancer, antiviral, or anti-inflammatory properties, further highlighting the importance

pharmacological properties.<sup>6</sup> These properties make chromones an attractive target for chemical modification through reactions like the Michael addition, enabling the construction of new molecules with potential biological activity. In the context of the Michael reaction, chromones can act as both nucleophiles and electrophiles, although they are more commonly employed as electrophilic partners. The most reactive site in chromones for nucleophilic additions is the C-2 position, which is activated due to electron withdrawal from the carbonyl group at C-4 (Scheme 1). Therefore, in this review, we aim to highlight the chameleonic behaviour exhibited by these fascinating structures.

a. Department of Organic Chemistry. Laboratorio de Organocatálisis Asimétrica. Instituto de Síntesis Química y Catálisis Homogénea (CSIC-University of Zaragoza). C/ Pedro Cerbuna 12, 50009 Zaragoza, Spain. E-mail: <a href="mmaamarq@unizar.es">mmaamarq@unizar.es</a> & raquelph@unizar.es

b. Centro de Investigaciones Químicas-IICBA, Universidad Autónoma del Estado de Morelos, Av. Universidad 1001, 62209 Cuernavaca, Morelos, México.

<sup>†</sup> Footnotes relating to the title and/or authors should appear here. Supplementary Information available: [details of any supplementary information available should be included here]. See DOI: 10.1039/x0xx00000x

This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence

Dpen Access Article. Published on 23 September 2025. Downloaded on 9/24/2025 6:03:17 PM

of the Michael addition reaction in the context of chromone chemistry.<sup>8</sup>

View Article Online
DOI: 10.1039/D50B01129A

Journal Name

# 

Aposphaerin A: R = R' = OH, R" = Me Aposphaerin B: R = OEt, R' = R" = H

Figure 1 Structure of bioactive chromanone natural products.

Recent advances in asymmetric Michael additions using chromones have focused on the development of enantioselective variants, allowing for the creation of chiral centres in a controlled manner. In this context, asymmetric transition-metal complexes have also been employed to achieve high enantioselectivity, leading to significant advancements in this area of research. The racemic approach using silyl compounds to activate the carbonyl group of this structure also deserves special mention, as it represents a pioneering approach for activating this conjugated skeleton and has served as a source of inspiration for further research in this area. The server is a server of the server is a server in the server is a server of the server is a server in the server in the server is a server in the server in the server is a server in the server is a server in the server in the server is a server in th

Flindersiachromanone

In addition, organocatalysts have been successfully developed to enhance the efficiency and selectivity of Michael additions with chromones. These catalytic strategies allow for mild reaction conditions, improved yields, and stereocontrol over the product, making them highly attractive for large-scale and industrial applications.

Although other valuable reviews have been published covering various aspects of chromone chemistry,<sup>5</sup> such as annulation reactions and C–H activation processes, there remains a lack of a comprehensive overview specifically focused on organocatalytic approaches. This gap has motivated us to undertake the present review, with a particular emphasis on the use of chiral organocatalysts. Our aim is to highlight recent advances and provide a detailed analysis of the progress made in organocatalysed C-2 and C-3 functionalisation of chromones.

Michael addition reactions using chromones as electrophilic C-2 synthons

In the context of chromones acting as electrophilic C-2 synthons, the Michael addition reaction is an interesting synthetic approach that usually requires organometallic reagents, as in the Zimmerman's strategy, which uses Zn(OTf)<sub>2</sub> as a catalyst,<sup>11</sup> or in Feringa's method,<sup>9c</sup> which employs a copper catalyst to produce 2-alkylchromanones. However, recent advances in the C-H functionalisation of chromones under metal-free conditions, which deserve special mention, have been developed by Han's group in 2014.<sup>12</sup> They demonstrated that the reaction of chromone 1a and cyclohexane (2a) in the presence of 4 eq. of di-tert-butyl peroxide (DTBP) at 120 °C for 24 h, yielded 2-cyclohexylchroman-4-one (3aa) in 72% yield (Scheme 2).

(+)-Pinostrobin

Scheme 2 Synthesis of 2-cyclohexylchroman-4-one (3aa)  $\emph{via}$  sp³ C-H functionalisation.

Attempts to increase the yield by using additional DTBP or alternative oxidants such as 2,3-dichloro-5,6-dicyano-1,4-benzoquinone (DDQ), NaClO, K<sub>2</sub>S<sub>2</sub>O<sub>8</sub>, 1,4-benzoquinone (BQ), or O<sub>2</sub> were unsuccessful. The method exhibited broad applicability, tolerating a variety of chromones **1a-h** and alkanes **2a-d**. Chromones bearing a methyl or bromine substituent at the C-6 position, as well as methoxy groups at the C-5 and C-7 positions, afforded the corresponding chromanones **3** with

This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence

Open Access Article. Published on 23 September 2025. Downloaded on 9/24/2025 6:03:17 PM

**ARTICLE** 

yields ranging from 58% to 83% (see the original article for a complete scope of the process).12 Notably, the electronwithdrawing nature of the bromine at C-6 enhanced the yield,

whereas steric hindrance and electron-donating groups, such as a methoxy substituent at C-5, seemed 🕬 🗚 🗗 📆 🛣 impact on the yield (Figure 2).

The proposed mechanism proceeds through a radical pathway (Scheme 3). Initial homolysis of DTBP 4 would generate a tertbutoxy radical 5·, which would abstract a hydrogen atom from cyclohexane 2a to form a cyclohexane radical 2a. This radical would add to chromone 1a, yielding an intermediate 6aa that would react further with the tert-butanol to form an enol intermediate 7aa. Subsequent keto-enol tautomerisation would produce the final 2-alkylchromanone product 3aa.

t-BuOH 7aa 6aa

Scheme 3 Mechanism via radical course.

In other hand, the gem-difluoroalkyl moiety have shown great importance in drug design to enhance the metabolic stability and biological properties of molecules. 13 Therefore, Hao and coworkers have selectively introduced a gem-difluoroalkyl moiety by developing an efficient metal-free Michael addition of difluoroenoxysilanes chromone 8a to using trifluoromethanesulfonic acid (HOTf) as a catalyst (Scheme 4).14 the reaction of chromone 1a with difluoroenoxysilane 8a in dichloromethane was tested with 10 mol% of different Lewis acids as Fe(OTf)<sub>3</sub>, Al(OTf)<sub>3</sub>, Bi(OTf)<sub>3</sub>, Cu(OTf)<sub>3</sub>, and trimethylsilyl trifluoromethansulfonate (TMSOTf)

over 72 h, obtaining the 2-difluoroalkylated chroman-4-one 9aa in moderate yields. Notably, among all of them, the best yield was obtained with TMSOTf (72%), prompting further exploration to enhance the yields by exploring the Brønsted acid catalyst HOTf in dichloromethane (78%) and hexane (85%) over 27 h. However, the optimal reaction conditions were established using 5 mol% of HOTf in THF as solvent, after 9 h of reaction, resulting in the synthesis of 2-difluoroalkylated chroman-4-one 9aa with 76% yield (Scheme 4).

Scheme 4 Highly efficient Michael addition of difluoroenoxysilane 8a to chromone

After establishing the optimal reaction conditions, the scope of the reaction was evaluated (Figure 3). It was found that chromones substituted at the C-7 position (1a,h,i) with electron-donating groups (H, Me, OMe) led the desired products 9aa, 9ha and 9ia at higher yields compared to those bearing electron-withdrawing groups (F, Cl, Br) (9ja, 9ka, 9la). Furthermore, electron-withdrawing groups (F, Cl, Br, NO<sub>2</sub>) at the C-6 position (9da, 9ea, 9fa, 9ga) resulted a higher yield than those with electron-donating groups (Me, OMe) (9ba, 9ca). phenyl-substituted the scope of Subsequently. difluoroenoxysilanes 8b-e was evaluated, revealing that substituents in para position improved the yield, achieving values between 89 and 93% (9ab, 9ac and 9ad) (Figure 3).

ARTICLE Journal Name

**Figure 3** Highly efficient Michael addition of difluoroenoxysilane **8a-e** to chromones **1a-l**.

**9ad** 11 h, 89%

9ae

13 h. 79%

However, the need to prepare enantiomerically cle pure compounds requires new stereoselective 10 strategies 01 that consider both environmental impact and reaction versatility. Consequently, as illustrated in the following methods, asymmetric organocatalysis has played a crucial role in addressing this challenge.

# Enantioselective Michael addition reactions using chromones as electrophilic C-2 synthons

This approach enables the synthesis of highly functionalised, stereochemically enriched compounds that are crucial intermediates in pharmaceuticals, natural products, and advanced materials.<sup>2,3</sup>

Interestingly, based on the anion-binding ability of silanediols as catalysts, <sup>15</sup> Mattson and co-workers pioneered the first enantioselective C-2 functionalisation of chromones. <sup>16</sup> Their approach uses enolsilyl acetals as nucleophiles within a catalytic silanediol ion-pair strategy. In this methodology, the reaction of chromone **1a** with a trialkylsilyl triflate **10** in toluene, at 60 °C, afforded its 4-siloxybenzopyrylium triflate **11a**, which was captured with the silanediol catalyst **12** to form a chiral ion pair **13**. <sup>16</sup> Subsequently, the addition of the silyl acetal **14** at -78 °C provided the corresponding 2-alkylchroman-4-one **15a** (Scheme 5). Notable, the nature of silyl group present on both the silyl acetal **14** and the silyl triflate **10** significantly influenced the enantioselectivity of the reaction, where the tri-*iso*-propyl group was better than *tert*-butyldimethylsilyl group.

Scheme 5 Enantioselective C-2 functionalisation of chromones 1a via chiral ion-pair 13.

Further studies exploring the scope of this reaction, using 20 mol% of catalyst **12**, provided evidence about the effect of the substituents on the chromone backbone (Figure 4). In fact, electron-withdrawing substituents such as Br, F, and Cl at the C-6 position (**15c**, **15e**, **15f**) led to higher yields compared to electron-donating groups like H and Me (**15a**, **15b**). Notably, the

highest yield was achieved when a CF<sub>3</sub> group was introduced at the C-7 position (15h). Additionally, the enantioselectivity improved in the presence of electron-withdrawing groups (Br, F, Cl, NO<sub>2</sub>) at the C-6 position (15c, 15e, 15f, 15g). The best enantiomeric excess (ee) was observed for the 3,5-( $F_3C$ )Ph-substituted derivative (15d) (Figure 4).

Journal Name ARTICLE

View Article Online DOI: 10 1039/D50801129A 
$$F_3$$
C  $F_3$ C

The absolute configuration of the newly formed stereogenic centre at the C-2 position of chromanones **15** was determined by X-ray crystallography of its iminochromanone derivative **16a** (Scheme 6). This derivative was obtained by reacting a racemic (*rac*) mixture of 2-substituted chromanone **15c** with (*R*)-2-methylpropane-2-sulfinamide using Ti(OMe)<sub>4</sub> as catalyst in THF at reflux, affording the iminochromanone **16a** in 80% yield and a diastereomeric ratio (dr) of 55:45. The diastereoisomers were then separated by crystallisation from dichloromethane and hexanes. Finally, the X-ray analysis and the deprotection of the iminochromanone **16a** revealed the absolute configuration (*S*) of its chromanone **15c** (Scheme 6).

Scheme 6 Chiral resolution of chroman-4-one  ${\bf 15c}$  by formation of the iminochromanone  ${\bf 16a}.$ 

In the context of chromone as C-2 electrophilic synthon, the presence of a carboxylic acid at the C-3 position has demonstrated significant potential in enhancing the activation of the Michael acceptor, facilitating the synthesis of novel 2-substituted chroman-4-ones. Albrecht's group has been a pioneer in developing a decarboxylative Michael addition using malonic acid half-thioesters in their first enantioselective approach.<sup>17</sup> Subsequently, Albrecht and co-workers explored

the Michael addition of azlactones 18 to chromone-3-carboxylic acids 17 to synthesise novel  $\alpha$ -amino acid moieties. 18 Initially, the reaction between chromone 1a with azlactone 18a was carried out in the presence of 20 mol% of quinine in THF at room temperature. However, no reaction was observed under these conditions. Introducing a carboxylic acid moiety at the C-3 position of chromone 17a successfully facilitated the formation of the desired product 20aa in over 95% yield via a decarboxylative Michael addition, albeit initially with low diastereoselectivity. In this context, to improve the enantioselectivity of the process, they conducted a screening of organocatalysts. Notably, cinchonidine-derived squaramides 19 significantly improved the stereoselectivity, yielding the 2-substituted chromone 20aa with >95% yield, a diastereomeric ratio of >20:1, and 82% enantiomeric excess (Scheme 7).18

Scheme 7 Decarboxylative Michael addition of azlactone  ${\bf 18a}$  to chromone-3-carboxylic acid  ${\bf 17a}$ .

The method showed a broad scope in the preparation of 2-substituted chromanones **20** (Figures 5 and 6). In all cases, the final products were obtained with moderate-to-high yields.

ARTICLE Journal Name

Notably, good diastereoselectivities were achieved with the model azlactone **18a**, although lower enantioselectivity was observed in all cases compared to product **20aa** (Figure 5). Additionally, electron-withdrawing groups (Br, F, Cl) in the chromone-3-carboxylic acid (**17c,17e,17f**) gave the products **20** in higher yield than those with an electron-donating group (Me)

in **17b** (Figure 6). Despite these variations, the best outcome in this series was still achieved with the reaction in both product **20aa** (see the original article for a complete scope of the process).

Figure 5 Selected examples from the general scope of the decarboxylative Michael addition of azlactones 18a-f to chromone-3-carboxylic acid 17a

Figure 6 Selected examples from the general scope of the decarboxylative Michael addition of azlactone 18a to chromone-3-carboxylic acids 17a-e.

Furthermore, the absolute configuration of the newly formed stereogenic centre was confirmed by X-ray crystallographic analysis of the chromanone **20ah**. Finally, the hydrolysis of the azlactone **20aa** using TsOH in a MeOH:CHCl<sub>3</sub> (2:1) at 40 °C produced the  $\alpha,\alpha$ -disubstituted amino acid **21aa** (Scheme 8).

**Scheme 8** Azlactone-ring opening to afford protected  $\alpha, \alpha$ -disubstituted amino acid **21aa**.

Additionally, a transition state model **22** was proposed by the authors to explain the observed stereoselectivity (Figure 7). It is suggested that the chromone-3-carboxylic acid **17a** would interact with the cinchona catalyst **19** via hydrogen bonding with the NHs of the squaramide. At the same time, the tertiary amine moiety, on the quinuclidine ring of **19**, would promote the deprotonation of the azlactone **18a**. The simultaneous interaction between the protonated catalyst and the resulting enolate, forming an ion pair, would facilitate the enantioselective Michael addition.

Figure 7 Proposed transition state 22 to explain the observed stereochemistry in 20.

Contemporaneously, García Mancheño and co-workers reported a highly enantioselective nucleophilic dearomatisation reaction of pyrylium salts using a chiral multicoordination triazole anion-binding catalyst **23** to afford 2-alkyl chromanones **26**. Initially, the reaction was performed as previously described by Akiba, Wherein the active benzopyrylium ion was generated through the reaction of chromone **1a** with a silyl derivative [tert-butyldimethylsilyl trifluoromethanesulfonate (TBSOTf)] in the presence of catalytic amounts of chiral H-donor catalyst **23** and 2,4,6-collidine at 60 °C in toluene. The optimal conditions involved the use of TBSOTf with a remarkably low catalyst loadings (1 mol%) of the multicoordination triazole **23**, followed by the reaction with a silyl ketene acetal **25a** at -78 °C,

Journal Name ARTICLE

affording the 2-alkyl chromanone **26aa** in 90% yield and 94% ee (Scheme 9).

Scheme 9 General reaction by chiral ion-pair 24 to obtain chromanone 26aa.

The scope of the reaction was investigated by testing various silyl ketene acetals **25a-e**. The substitution pattern on the nucleophile was found to significantly influence on the enantioselectivity of the process, with more sterically hindered nucleophiles proceeding smoothly and exhibiting lower enantioselectivity (**25c-e**) than smaller nucleophiles (**25a-b**). Chromones substituted at C-6 position with either electrondonating and electron-withdrawing groups delivered final products **26** in high yields and with moderate to excellent enantioselectivities (Figure 8). Interestingly, the method also proved effective for challenging thiochromones and 4-pyrones, producing final products with moderate yields but good to excellent enantioselectivity (see the original article for a complete scope of the process). <sup>19</sup>

**26ae**, R' = *i*-Pr 95% yield, 40% ee

Figure 8 Selected examples from the general scope using chiral ion-pair to obtain chromanones 24-26.

# Enantioselective conjugated addition reactions using chromones as nucleophilic C-2 synthons

Additionally, enantioselective Michael addition reactions using chromones as nucleophilic synthons are also of significant importance in synthetic organic chemistry, as they provide a powerful and efficient approach for constructing structurally complex and biologically relevant molecules with high stereocontrol.<sup>20</sup>

Recently, in the field of C-H functionalisation to obtain new 2functionalised chromones, 21 Zlotin and co-workers, 22 reported a metal-free C-2 functionalisation of 3-hydroxychromones 27 via electrophilic olefination with electron-deficient olefins, such as nitroolefins **28** (Scheme 10) or  $\beta$ , $\gamma$ -unsaturated- $\alpha$ -keto esters 32 (Scheme 11), using a recyclable C2-symmetric squaramide 29 as organocatalyst. The reaction was optimised between the 3hydroxychromone 27a and nitroolefin 28a in the presence of 2 mol% of the chiral C2-symmetric squaramide 29 at room temperature in EtOH (95%). This process afforded 2-alkyl-3hydroxychromone 30aa in 97% yield and 94% ee (Scheme 10). Transition state 31 was proposed by the authors to explain the absolute configuration observed in the final products 30. In this mechanism, the bifunctional squaramide 29 would deprotonate the chromone 27a through its tertiary amino group. Then, the catalyst would interact with the enolate to influence the geometry of the active complex. Simultaneously, strong hydrogen bonds formed between the electron-deficient olefins 28a and the NH groups of the squaramide, would drive the nucleophilic attack determining the absolute configuration of final products 30.

ARTICLE Journal Name

31

**Scheme 10** Model reaction for the formation of chromenone **30aa** through C2-symmetric squaramide catalyst.

DOI: 10.1039/D5OB01129A

With the best reaction conditions established, the scope of this methodology was evaluated with various 3-hydroxychromones **27a-c** and several nitroolefins **28a-f**, leading to the corresponding 2-alkyl-chromones **30** with yields of up to 99% and excellent enantioselectivity (90-99% ee, see the original article for a complete scope of the process). In general, this method demonstrated an excellent tolerance to electronic effects of the substituents in the 3-hydroxychromones **27a-c** and nitroolefins **28a-f** (Figure 9).<sup>22</sup>

Figure 9 Selected examples from the general scope using chiral squaramide 31 to obtain chromenone derivatives 30.

Scheme 11 General scope using  $\beta$ , $\gamma$ -unsaturated- $\alpha$ -keto esters 32a,b to obtain chromenone derivatives 33 and 34.

Additionally,  $\beta$ , $\gamma$ -unsaturated- $\alpha$ -keto esters **32a,b** showed almost quantitative yield and high enantioselectivity, leading to

the formation of the corresponding Michael adducts **33** and **34** in a lactonisation/ring opening equilibrium (Scheme 11).

Journal Name ARTICLE

View Article Online

Following the same approach of chromones as nucleophilic C-2 synthon, Rafiński and co-workers reported a very interesting asymmetric organocatalysed conjugated addition of 3-hydroxychromones **24** via chiral  $\alpha,\beta$ -unsaturated acyl azoliums. <sup>23</sup> In this methodology, initially, 3-hydroxychromone **27a** reacted with ynal **35a** in the presence of a carbene, generated from chiral aminoindanol-derived triazolium salt **36**, using a proton sponge (PS) as sterically hindered and non-nucleophilic base, in dichloromethane. Subsequent treatment with MeOH, afforded the C-2 alkylated 3-hydroxychromone **37aa** in 99% yield (as determined by <sup>1</sup>H NMR analysis), and 95% ee (Scheme **12**).

Scheme 12 General reaction by chiral carbene to obtain chromenone 37aa.

Under the best reaction conditions, this strategy showed a great scope as shown in Figure 10 (see the original article for a complete scope of the process). Notably, various 3-hydroxychromones 27 were tested with different substituents

at C-6 and C-7 position (27a-i), affording the desired products 37 in 66-99% yield and excellent enantioselectivities (56-99% ee). Notably, a methoxy group at C-5 position (27h) gave its respective chromone 37ha in 97% yield and 99% ee. Additionally, a series of ynals with electron-withdrawing (35bd) and electron-donating groups (35e-g) on the para position of the phenyl ring worked well. However, the CF3 moiety (35h) exhibited only moderate selectivity. A similar trend was observed for the phenyl ortho-substituted ynals (35i-I) giving high stereoselectivities. However, phenyl meta-substituted ynals (35m and 35n) showed lower enantioselectivity without affecting the yield (37am and 37an). The reaction also showed good tolerance for challenging ynals bearing alkyl substituents, yielding the desired C2-alkylated chromenones in moderate to high yields with consistent enantioselectivities. To further extend this approach, the ring opening was performed using different oxygen or nitrogen nucleophiles, successfully generating the corresponding chromenones without compromising the enantioselectivity (see the original article for complete scope of the process).<sup>23</sup> Notably, trimethylsilylpropynal, used as precursor for  $\alpha,\beta$ -unsaturated acyl azolium, efficiently produced the C2-alkylated product 37, despite undergoing complete desilylation under the reaction conditions. In this regard, the silyl derivative serves as a safer alternative to propynal, which is known for its explosive nature. The absolute stereochemistry of the chiral centre was determined through single-crystal X-ray analysis.

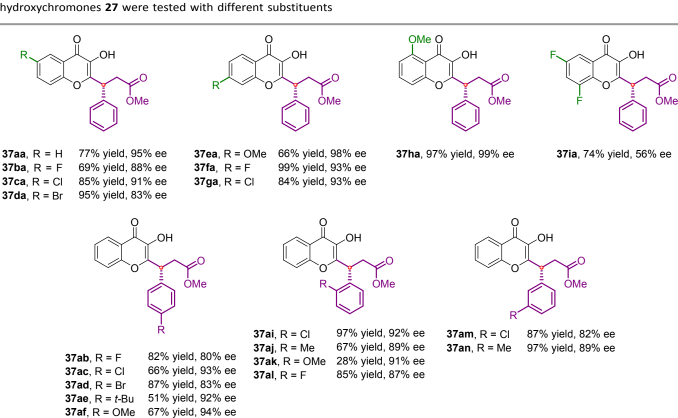


Figure 10 Selected examples from the general scope using a chiral carbene to obtain chromenone derivatives 37.

95% yield, 93% ee

66% yield, 63% ee

**37ag**, R = Me

37ah, R = CF<sub>3</sub>

View Article Online DOI: 10.1039/D5OB01129A

## **REVIEW**

The authors propose a reaction mechanism that begins with the formation of the  $\alpha,\beta$ -unsaturated acyl azolium 38 via redox isomerisation of ynal 35a. This intermediate undergoes a 1,2addition with 3-hydroxychromone 27a, yielding intermediate 39. A subsequent Coates-Claisen rearrangement generates the 2-alkyl-chroman-3,4-dione 40, which, through tautomerisation and lactonisation of amide 41, forms the tricyclic chromone 42. Finally, reaction with a nucleophile (MeOH) produces the C2functionalised 3-hydroxychromone 37aa with enantioselectivity (Scheme 13). The authors proposed this mechanism based on common experimental observations, particularly the formation of esterification byproducts, which is typically associated with 1,2-addition pathways. This behaviour is consistent with the involvement of an NHC catalyst. In their study, esterification side reactions were also observed, supporting the idea that the reaction does not proceed via a nucleophilic 1,4-addition of the enol to the catalytically generated  $\alpha,\beta$ -unsaturated acyl azolium. Instead, it likely follows a 1,2-addition route, leading to O-acylation products. Additionally, based on the [3,3]-sigmatropic rearrangement mechanism previously described by Bode for kojic acid derivatives,<sup>24</sup> the authors suggest that a similar rearrangement could be occurring in the reaction between hydroxychromenones 27 and  $\alpha,\beta$ -unsaturated acyl azoliums 38.

Scheme 13. Proposed mechanism for obtaining C2-functionalised chromenone derivative 37aa

37aa, 77% yield, 95% ee

Simultaneously, **Jiang** and co-workers design enantioselective reductive cross coupling of chromones 1 and electron deficient olefins 38, applying a visible-light-driven cooperative photoredox system in combination with a chiral Brønsted acid catalyst.<sup>25</sup> The reaction was performed using chromone 1a and 2-vinylpyridine 38a as model substrates. A dicyanopyrazine-derived chromophore (DPZ) 39 (2 mol%) was employed as photoredox catalyst in the presence of 20 mol% of the chiral phosphoric acid (CPA) 40 and the Hantzsch ester 41 in dichloromethane at -25 °C. The reaction was conducted under 3 W blue LED light in an oxygen-free environment for 60 hours, affording the 2-alkylchromanone 42aa with a 63% yield and 97% of enantiomeric excess (Scheme 14).

**Scheme 14** Model enantioselective reductive cross coupling of chromone **1a** with 2-vinylpyridine **38a**.

This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence.

Article. Published on 23 September 2025. Downloaded on 9/24/2025 6:03:17 PM

Journal Name ARTICLE

The scope of this asymmetric cross coupling protocol was evaluated using several chromones **1a-i** containing electron-withdrawing and electron-donating substituents, affording the 2-alkylchromanones **42** in moderate to high yields with excellent enantioselectivities. Furthermore, to improve the yield of the reaction, **2**-vinylpyridines substituted with a methyl group at the C-3 and C-4 position were tested **(38b** and **38c)**. The best result **(93%** yield, 96% ee) was obtained with the

pyridine substituted at C-3 (42ab). On the other hand, when the reaction was performed with a thiochromone, the best result was using the 2-vinylpyridine 38c bearing a methyl group at C-4 position affording thiochromanone 43lc. Moreover, different substituted thiochromones exhibited excellent enantioselectivities, albeit with lower yield (43aa-ga, 43hb and 43ic) (Figure 11).

Figure 11 Scope of the enantioselective reductive cross coupling of chromones 1 to obtain 2-alkylchromanone derivatives 42 and 2-alkylthiochromanones 43.

Based on the success of this process, the authors proposed a reaction mechanism involving radical intermediates generated under photoredox conditions (Scheme 15). Simultaneously, the chiral phosphoric acid 40 would form hydrogen-bonding interactions with the substrates, stabilising the intermediate and organising the reactive components in a well-defined transition state (TS 46). This precise arrangement ensures enantioselective C-2 functionalisation of the chromone 1a. This dual catalytic system, combining photoredox and CPA catalysis, enables effective control over both reactivity and selectivity, facilitating the formation of C-2-functionalised chromones 42 with excellent stereochemical outcomes.

**Scheme 15** Proposed mechanism through radical intermediates.

# Chromones as nucleophilic C-3 synthons and ulterior Michael addition reactions

Addition reactions using chromones as nucleophilic C-3 synthons are highly valuable in organic synthesis, offering a

versatile strategy to access diverse, stereochemically enriched scaffolds that are crucial for the development of bioactive compounds and natural product derivatives.

In this context, homoisoflavonoids<sup>26</sup> are an important subclass of flavonoids with significant potential in drug design due to their demonstrated biological activities, including antioxidant,<sup>27</sup> cytotoxic<sup>28</sup> or antimicrobial effects (Figure 12).<sup>29</sup>

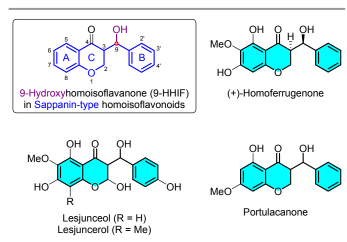


Figure 12 Potential homoisoflavonoids with biological properties.

In this framework, Seo and co-workers<sup>30</sup> have established a diastereodivergent synthesis of *anti*- and *syn*-9-hydroxyflavanones **49** and **53**, respectively, starting from chromones **1** and carbaldehydes **47** *via* a reductive aldol reaction or the Morita-Baylis-Hillman (MBH) reaction, followed by a *syn*-selective Michael reduction. Initially, they demonstrated the feasibility of the 3,9-*anti* selective synthetic route. Under reductive conditions, the enolate of chromone **1** was generated using Na, K or Li tri-*sec*-butylborohydride

**ARTICLE Journal Name** 

(Selectride) in THF at -78 °C, which provided the (E)-enolate. This enolate subsequently reacted with arylaldehydes 47, affording the anti-3,9-hydroxyflavanones 49 through a Zimmerman-Traxler transition state 48, giving the products from 59 to 97% yield and up to 1:14 diastereomeric ratio (Scheme 16).

Scheme 16 Transition state 48 in the reductive aldol reaction conducting to the anti-product 49.

This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence.

Open Access Article. Published on 23 September 2025. Downloaded on 9/24/2025 6:03:17 PM

On the other hand, it is very important to consider that  $\alpha$ substituent-β-hydroxy ketones, such as the aldol chromone derivatives 50, are highly prone to H-3 deprotonation (pK<sub>a</sub> 14.8), which lead to  $\beta$ -elimination or retro-oxa-Michael reaction. To optimise the reaction conditions to obtain the desired synproducts 53, the MBH reaction was carried out using 1,8diazabicyclo[5.4.0]undec-7-ene (DBU) in MeOH at room temperature to produce the enone intermediates 50. A subsequent Michael reduction with 3 eq. of L-Selectride

facilitated the formation of a highly organised achelated transition state **51**, positioning H-3 on the ଡpposite/face ଖ୍ୟାଧାର phenyl group. Various proton sources were then tested, with phenol giving rise to the highest product formation. Additionally, 2-methylbenzimidazole (52a) provided comparable yield and with an improved dr compared to MeOH, NH<sub>4</sub>Cl or AcOH, which instead led to retro-aldol by-products or degradation (Scheme 17).

Scheme 17 Synthesis of the syn-product 53 by MBH/Michael reduction strategy.

Finally, higher anti- and syn-selectivity was observed in both methods using electron-rich aromatic rings than electrondeficient aromatic rings. However, the MeO group located on the C-5 position of chromones and/or C-2' position of the carbaldehyde got worse diastereoselectivity due to possible involvement in the cation chelation during the stereochemistrydetermining step (Scheme 18, see the original article for a complete scope of the process).30

**Scheme 18** Selected examples from the general scope of the highly efficient MBH reaction for obtaining 3-alkylchromanones **53**. a) For details in the Supporting Information of ref.30. b) yield of the MBH; c) Combined yield of both diastereoisomers of *syn*-selective **1**,4-reduction; d) Result obtained by using **52b** as a PS.

Furthermore, Chen and co-workers<sup>31</sup> initially reported the catalysed reactions of chromone-derived MBH adducts with synthesis of chiral 3-aminomethylene-flavanones by In(III)-

amines via a tandem allylic amination/ring opening/oxa-

This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence.

Open Access Article. Published on 23 September 2025. Downloaded on 9/24/2025 6:03:17 PM

**Journal Name ARTICLE** 

Michael addition reactions in a one pot process.<sup>32</sup> In the next step of synthesis of this type of compounds, an organocatalytic oxa-Michael addition was employed.<sup>33</sup> This reaction began with the treatment of chromone-derived MBH carbonate 54a with naphthylmethylenamine (55a) in the presence of 20 mol% of cinchonidine-derived thiourea catalyst in toluene at 40 °C for 60 h, affording the product 57a in 74% yield and 36% ee. To improve the enantioselectivity, as reported by Zhu and Lu,34

trifunctional catalysts containing primary amino acid units were found to be more effective for asymmetric conjugate addition than bifunctional thioureas. Following this method, different trifunctional catalysts were tested, resulting in a significant improvement in enantioselectivity when trifunctional cinchonidine-derived catalyst 56 was used, producing the 3aminomethylene-flavone 57a in 75% yield and 82% ee (Scheme

Scheme 19 Enantioselective organocatalytic synthesis of 3-aminomethylene-flavanones 57a

With the optimal reaction conditions, several carbonates 54 and amines 55 were examined (Figure 13), yielding moderate to high yields and enantioselectivities for all products 57 (see the original article for a complete scope of the process). The absolute configuration of the new stereogenic centre at C-2 position was then confirmed as R by X-ray crystallographic analysis of product 57c. Drawing on these findings, the authors suggested a plausible transition state 58 for the tandem reaction (Scheme 19). Firstly, amines (ArCH2NH2) would react with the MBH carbonate, which is activated by the thioureabased catalyst 56, through hydrogen bond formation, leading to the generation of intermediate 58 (Scheme 19). This step is followed by an intramolecular oxa-Michael addition, stablishing multiple hydrogen bonds between the trifunctional catalyst and the intermediate 58. These interactions enhance the reaction rate and ensure precise stereochemical control, facilitating the oxa-Michael addition to the  $\alpha,\beta\text{-unsaturated}$  imine produced in situ, attacking from the Si face of the double bond.

**57h**, R = Me 72% yield, 89% ee

Figure 13 Selected examples from the general scope for the enantioselective organocatalytic synthesis of 3-aminomethylene-flavanones

View Article Online DOI: 10.1039/D5OB01129A

## **Conclusions**

This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence

Article. Published on 23 September 2025. Downloaded on 9/24/2025 6:03:17 PM

The Michael addition reactions involving chromones represent a cornerstone in modern organic synthesis, offering an interesting alternative for the construction of complex, stereochemically enriched molecules. The remarkable versatility of chromones, acting as electrophilic or nucleophilic partners, has enabled the development of methodologies applicable in pharmaceuticals, natural products, and materials science.

Recent advancements have been driven by innovative organocatalytic systems that leverage non-metal-based strategies for asymmetric transformations, providing a sustainable and highly efficient approach to chemical synthesis. Despite these successes, several challenges remain, including improving the scalability of reactions, expanding the substrate scope to obtain more functionalised chromones, and achieving higher enantioselectivity even under milder or greener conditions. Future research in enantioselective Michael addition reactions involving chromones should focus on several key areas to address current challenges and unlock new opportunities. One priority is the development of novel organocatalysts with enhanced activity, selectivity, and environmental compatibility. These catalysts should not only improve reaction efficiency but also align with principles of green chemistry by reducing waste and utilising sustainable resources, such as greener solvents.

Additionally, a deeper understanding of the reaction mechanisms is essential. Employing advanced computational techniques could shed light on the underlying processes, helping to refine transition state models and guide the rational design of more effective catalytic systems.

Moreover, exploring tandem or cascade reactions to achieve multi-functionalised products in a single synthetic step could pave the way for even more complex and diverse molecular architectures.

By addressing these priorities, future research can further establish enantioselective Michael addition reactions as a cornerstone methodology in modern synthetic organic chemistry, with broad applications in both academia and industry.

## **Author contributions**

All authors wrote and approved the paper.

#### Conflicts of interest

There are no conflicts to declare.

#### Acknowledgements

The authors thank Ministerio de Ciencia, Innovación y Universidades and Agencia Estatal de Investigación for the project PID2023-147471NB-I00 funded by MICIU/AEI10.13039/501100011033, Gobierno de Aragón-Fondo Social Europeo (Research Group E07\_23R) and Secretaría de Ciencia, Humanidades Tecnología e Innovación (SECIHTI) for financial support through projects 286614 and 140607. J.C.M.S. also thanks SECIHTI for predoctoral scholarship 1004402. We gratefully acknowledge funkderulz for the free use of their image shared on Pixabay, which was used to illustrate this work.

## **Notes and references**

- For recent reviews in this area of research, see: (a) S.-H. Xiang and B. Tan, Advances in asymmetric organocatalysis over the last 10 years, Nat. Commun., 2020, 11, 3786; (b) A. Antenucci, S. Dughera and P. Renzi, Green chemistry meets asymmetric organocatalysis: A critical overview on catalysts Synthesis, ChemSusChem, 2021, 14, 2785–2853; (c) Z. Chen, Q.-Q. Yang, W. Du and Y.-C. Chen, Asymmetric organocatalysis involving double activation, Tetrahedron Chem, 2022, 2, 100017; (d) S. Ardevines, E. Marqués-López and R. P. Herrera, Horizons in asymmetric organocatalysis: En route to the sustainability and new applications, Catalysts, 2022, 12, 101; (e) O. García Mancheño and M. Waser, Recent developments and trends in asymmetric organocatalysis, Eur. J. Org. Chem., 2023, 26, e202200950.
- (a) A. Acarlone and L. Bernardi, Enantioselective organocatalytic approaches to active pharmaceutical ingredients selected industrial examples, *Phys. Sci. Rev.*, 2019, 4, 20180097; (b) B. Han, X.-H. He, Y.-Q. Liu, G. He, C. Peng and J.-L. Li, Asymmetric organocatalysis: an enabling technology for medicinal chemistry, *Chem. Soc. Rev.*, 2021, 50, 1522–1586; (c) A. J. Burke, Asymmetric organocatalysis in drug discovery and development for active pharmaceutical ingredients, *Expert Opin. Drug Discov.*, 2022, 18, 37–46; (d) E. Reyes, L. Prieto and A. Milelli, Asymmetric organocatalysis: A survival guide to medicinal chemists, *Molecules*, 2023, 28, 271.
- 3 K. A. Scott, N. Ropek, B. Melillo, S. L. Schreiber, B. F. Cravatt and E. V. Vinogradova, Stereochemical diversity as a source of discovery in chemical biology, *Curr. Res. Chem. Biol.*, 2022, 2, 100028.
- For selected reviews, see: (a) D. R. López, D. Sadaba, I. Delso, R. P. Herrera, T. Tejero and P. Merino, Asymmetric organocatalytic synthesis of y-nitrocarbonyl compounds through Michael and Domino reactions, Tetrahedron: Asymmetry, 2010, 21, 2561-2610; (b) Y. Zhang and W. Wang, Recent advances in organocatalytic asymmetric Michael reactions, Catal. Sci. Technol., 2012, 2, 42-53; (c) Y. Wang and D.-M. Du, Recent advances in organocatalytic asymmetric oxa-Michael addition triggered cascade reactions, Org. Chem. Front., 2020, 7, 3266-3283; (d) T. Das, S. Mohapatra, N. P. Mishra, S. Nayak and B. P. Raiguru, Recent advances in organocatalytic asymmetric Michael addition reactions to  $\alpha$ , $\beta$ -unsaturated nitroolefins, ChemistrySelect, 2021, 6, 3745-3781; (e) P. Sharma, R. Gupta and R. K. Bansal, Recent advances in organocatalytic asymmetric aza-Michael reactions of amines and amides, Beilstein J. Org. Chem., 2021, 17, 2585-2610; (f) S. D. Pasuparthy and B. Maiti, Enantioselective organocatalytic Michael addition reactions catalyzed by proline/prolinol/supported proline based organocatalysts:

This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence

Article. Published on 23 September 2025. Downloaded on 9/24/2025 6:03:17 PM

Journal Name ARTICLE

- An overview, *ChemistrySelect*, 2022, **7**, e202104261; (g) R. Ballini, A. Palmieri and M. Petrini, Catalysts' evolution in the asymmetric conjugate addition of nitroalkanes to electron-poor alkenes, *Org. Chem. Front.*, 2022, **9**, 6077–6113.
- For very elegant and interesting reviews about annulation reactions using chromones, see: (a) M. Zhang, Y. Gong, W. Zhou, Y. Zhou and X.-L. Liu, Recent advances of chromone-based reactants in the catalytic asymmetric domino annulation reaction, *Org. Chem. Front.*, 2021, **8**, 3968–3989; (b) P. Langer, Domino reactions of chromones with activated carbonyl compounds, *Beilstein J. Org. Chem.*, 2024, **20**, 1256–1269. For a very interesting review about C-H activation using metal catalysts, see: (c) A. T. Benny and E. K. Radhakrishnan, Advances in the site-selective C-5, C-3 and C-2 functionalization of chromones via sp<sup>2</sup> C-H activation, *RSC Adv.*, 2022, **12**, 3343–3358.
- 6 (a) M. Costa, T. A. Dias, A. Brito and F. Proença, Biological importance of structurally diversified chromenes, *Eur. J. Med. Chem.*, 2016, 123, 487–507; (b) J. Yang, J. Lai, W. Kong and S. Li, Asymmetric synthesis of Sakuranetin-relevant flavanones for the identification of new chiral antifungal leads, *J. Agric. Food Chem.*, 2022, 70, 3409–3419.
- 7 (a) S. Emami and Z. Ghanbarimasir, Recent advances of chroman-4-one derivatives: Synthetic approaches and bioactivities, *Eur. J. Med. Chem.*, 2015, **93**, 539–563; (b) S. Kamboj and R. Singh, Chromanone-a prerogative therapeutic scaffold: An overview, *Arab. J. Sci. Eng.*, 2022, **47**, 75–111.
- For the synthesis of 4-chromanone-derived compounds following other strategies, see: E. J. Diana, U. S. Kanchana and T. V. Mathew, Current developments in the synthesis of 4-chromanone-derived compounds, *Org. Biomol. Chem.*, 2021, **19**, 7995–8008.
  - For selected metal-catalysed examples, see: (a) J. Chen, J. Chen, F. Lang, X. Zhang, L. Cun, J. Zhu, J. Deng and J. Liao, A C2-symmetric chiral bis-sulfoxide ligand in a rhodiumcatalyzed reaction: Asymmetric 1,4-addition of sodium tetraarylborates to chromenones, J. Am. Chem. Soc., 2010, 132, 4552-4553; (b) T. Korenaga, K. Hayashi, Y. Akaki, R. Maenishi and T. Sakai, Highly enantioselective and efficient synthesis of flavanones including Pinostrobin through the rhodium-catalyzed asymmetric 1,4-addition, Org. Lett., 2011, 13, 2022–2025; (c) C. Vila, V. Hornillos, M. Fañanás-Mastral and B. L. Feringa, Catalytic asymmetric conjugate addition of Grignard reagents to chromones, Chem. Commun., 2013, 49, 5933-5935; (d) J. C. Holder, A. N. Marziale, M. Gatti, B. Mao and B. M. Stoltz, Palladium-catalyzed asymmetric conjugate addition of arylboronic acids to heterocyclic acceptors, Chem. Eur. J., 2013, 19, 74-77; (e) Q. He, C. M. So, Z. Bian, T. Hayashi and J. Wang, Rhodium/chiral diene-catalyzed asymmetric 1,4-addition of arylboronic acids to chromones: A highly enantioselective pathway for accessing chiral flavanones, Chem. Asian J., 2015, 10, 540-543; (f) L. G. DeRatt, M. Pappoppula and A. Aponick, A facile enantioselective alkynylation of chromones, Angew. Chem. Int. Ed., 2019, 58, 8416-8420; (g) B. M. Trost, E. Gnanamani, C. A. Kalnmals, C.-I. J. Hung and J. S. Tracy, Direct enantioand diastereoselective vinylogous addition of butenolides to chromones catalyzed by Zn-prophenol, J. Am. Chem. Soc., 2019, 141, 1489-1493; (h) Y. Guan, J. W. Attard and A. E. Mattson, Copper bis(oxazoline)-catalyzed enantioselective alkynylation of benzopyrylium ions, Chem. Eur. J., 2020, 26, 1742-1747; (i) Y. Guan, T. A. Buivydas, R. F. Lalisse, J. W. Attard, R. Ali, C. Stern, C. M. Hadad and A. Mattson, Robust, enantioselective construction of challenging, biologically relevant tertiary ether stereocenters, ACS Catal., 2021, 11, 6325-6333; (j) J. Cui, N. Kumagai, T. Watanabe and M. Shibasaki, Direct catalytic asymmetric and anti-selective

vinylogous addition of butenolides to chromones, Chem. Sci.,

- 2020, **11**, 7170–7176; (k) Y. Mo, Q. Chen, J. Li, D. Ye Y. Zhoures. Dong, X. Liu and X. Feng, Asymmetric catalytic confired addition of cyanide to chromones and β-substituted cyclohexenones, *ACS Catal.*, 2023, **13**, 877–886; (I) S. Samanta, J. Cui, H. Noda, T. Watanabe and M. Shibasaki, Asymmetric *syn*-selective vinylogous addition of butenolides to chromones via Al-Li-BINOL catalysis, *J. Org. Chem.*, 2023, **88**, 1177–1184.
- 10 (a) H. Iwasaki, T. Kume, Y. Yamamoto and K.-y. Akiba, Reaction of 4-t-butyldimethylsiloxy-1-benzopyrylium salt with enol silyl ethers and active methylenes, Tetrahedron Lett., 1987, 28, 6355-6358; (b) Y.-G. Lee, K. Ishimaru, H. Iwasaki, K. Ohkata and K.-y. Akiba, Tandem reactions in 4siloxy-1-benzopyrylium salts: Introduction of substituents and cyclohexene and cyclopentane annulation in chromones, J. Org. Chem., 1991, **56**, 2058–2066; (c) I. Mori, Y. Zimura, T. Nakano, S.-i. Matsunaga, G. Iwasalfi, A. Ogawa and K. Hayakawa, Trimethylsilyl triflate promoted 1,4-addition of silyl phosphites to cyclic enones, Tetrahedron Lett., 1997, 38, 3543-3546; (d) T. Qin, R. P. Johnson and J. A. Porco, Jr., Vinylogous addition of siloxyfurans to benzopyryliums: A concise approach to the tetrahydroxanthone natural products, J. Am. Chem. Soc., 2011, 133, 1714-1717; (e) J. Liu, Z. Li, P. Tong, Z. Xie, Y. Zhang and Y. Li, TMSI-promoted vinylogous Michael addition of siloxyfuran to 2-substituted chromones: A general approach for the total synthesis of chromanone lactone natural products, J. Org. Chem., 2015, 80, 1632-1643; (f) E. Kerste, M. P. Beller and U. Koert, Second generation total synthesis of (–)-Preussochromone D, Eur. J. Org. Chem., 2020, 2020, 3699-3711; (g) R. O. Argüello-Velasco, J. C. Morales-Solís, M. Muñoz-Vidales, J. L. Viveros-Ceballos, I. Romero-Estudillo and M. Ordoñez, An efficient synthesis of cis-4-hydroxyphosphonic and cis-4hydroxyphosphinic analogs of pipecolic acid from cyclic enaminones, Amino Acids, 2022, 54, 299-310; (h) M. Ordoñez, R. O. Argüello-Velasco, T. Miranda-Blancas, I. Romero-Estudillo and V. Labastida-Galván, First stereoselective synthesis of diethyl cis- and trans-(4-hydroxy-1,2,3,4-tetrahydroquinolin-2-yl)phosphonates and ethyl phenylphosphinates from Quinolin-4(1H)-one, Synthesis, 2023, 55, 4181-4190.
- 11 J. R. Zimmerman, M. Manpadi and R. Spatney, Tin-free radical reactions under minimal solvent conditions for the synthesis of substituted chromones and coumarins, *Green Chem.*, 2011, **13**, 3103–3106.
- 12 J. Zhao, H. Fang, P. Qian, J. Han and Y. Pan, Metal-free oxidative C(sp³)—H bond functionalization of alkanes and conjugate addition to chromones, *Org. Lett.*, 2014, 16, 5342— 5345.
- 13 (a) K. Ando, F. Koike, F. Kondo and H. Takayama, An improved synthesis of 24, 24-fifluoro-1α,25-dihydroxyvitamin D<sub>3</sub> from readily available vitamin D<sub>2</sub>, Chem. Pharm. Bull., 1995, 43, 189–192; (b) Y. Zhou, J. Wang, Z. Gu, S. Wang, W. Zhu, J. L. Aceña, V. A. Soloshonok, K. Izawa and H. Liu, Next generation of fluorine-containing pharmaceuticals, compounds currently in phase II–III clinical trials of major pharmaceutical companies: New structural trends and therapeutic areas, Chem. Rev., 2016, 116, 422–518; (c) Y. Zafrani, D. Yeffet, G. Sod-Moriah, A. Berliner, D. Amir, D. Marciano, E. Gershonov and S. Saphier, Difluoromethyl bioisostere: Examining the "lipophilic hydrogen bond donor" concept, J. Med. Chem., 2017, 60, 797–804.
- 14 X.-Y. Wang, M. Yang, Y. Zhou, J. Zhou and Y.-J. Hao, A highly efficient metal-free selective 1,4-addition of difluoroenoxysilanes to chromones, *Org. Biomol. Chem.*, 2023, 21, 1033–1037.

**ARTICLE Journal Name** 

15 (a) S.-i. Kondo, T. Harada, R. Tanaka and M. Unno, Anion recognition by a silanediol-based receptor, Org. Lett., 2006, 8, 4621-4624; (b) A. G. Schafer, J. M. Wieting and A. E. Mattson, Silanediols: A new class of hydrogen bond donor catalysts, Org. Lett., 2011, 13, 5228-5231; (c) K. Brak and E.

- 25 K. Cao, C. Li, D. Tian, X. Zhao, Y. Yin and Z. Jiang, Catalytic enantioselective reductive cross coupling of electrondeficient olefins, Org. Lett., 2022, 24, 4788-4792.
- 26 (a) B. M. Abegaz, J. Mutanyatta-Comar and M. Nindi, Naturally occurring homoisoflavonoids: Phytochemistry, biological activities and synthesis, Nat. Prod. Commun., 2007, 2, 475-498; (b) K. Du Toit, S. E. Drewes and J. Bodenstein, The chemical structures, plant origins, ethnobotany and biological activities of homoisoflavanones, Nat. Prod. Res., 2010, 24, 457-490; (c) B. M. Abegaz and H. H. Kinfe, Naturally occurring homoisoflavonoids: Phytochemistry, biological activities, and synthesis (Part II), Nat. Prod. Commun., 2019, 14, 1-20; (d) F. C. Demidoff, P. R. R. Costa and G. S. Caleffi, Advances in the

- synthesis of rearranged homoisoflavonoids, Org. Biomol. Chem., 2024, 22, 4839-4863. DOI: 10.1039/D5OB01129A
- 27 M. Machala, R. Kubínová, P. Hořavová and V. Suchý, Chemoprotective potentials of homoisoflavonoids and chalcones of Dracaena cinnabari: modulations of drugmetabolizing enzymes and antioxidant activity, Phytoter.
- 28 J. Mutanyatta, B. G. Matapa, D. D. Shushu and B. M. Abegaz, Homoisoflavonoids and xanthones from the tubers of wild and in vitro regenerated Ledebouria graminifolia and cytotoxic activities of some of the homoisoflavonoids. Phytochemistry 2003, 62, 797-804.
- 29 (a) G. O'Donell, F. Bucar and S. Gibbons, Phytochemistry and antimycobacterial activity of Chlorophytum inornatum, Phytochemistry, 2006, 67, 178-182; (b) V. L. N. Reddy, V. Ravikanth, V. V. N. S. J. Lakshmi, M. Suryanarayan and Y. Venkateswarlu, Inhibitory activity of homoisoflavonoids from Caesalpinia sappan against Beauveria bassiana, Fitoterapia,
- 30 S. Lee, S. Kwon, J. Hur, and S.-Y. Seo, Diastereodivergent synthesis of syn- and anti-9-hydroxyhomoisoflavanone and its application to the total syntheses of (±)-Homoferrugenone and (±)-Portulacanone F, Adv. Synth.
- 31 (a) C. Wu, Y. Liu, H. Zeng, L. Liu, D. Wang and Y. Chen, In(III)-Catalyzed tandem reaction of chromone-derived Morita-Baylis-Hillman alcohols with amines, Org. Biomol. Chem., 2011, 9, 253-256; (b) C. Wu, H. Zeng, L. Liu, D. Wang and Y. Chen, Tandem allylic amination/ring-opening/oxa-Michael addition reactions of chromone-derived Morita-Baylis-Hillman acetates with amines, Tetrahedron, 2011, 67, 1231-
- 32 For a non-racemic version of a similar process, see also: J. Lei, Y. Li, L.-J. He, Y.-F. Luo, D.-Y. Tang, W. Yan, H.-K. Lin, H.-y. Li, Z.-Z. Chen and Z.-G. Xu, Expeditious access of chromone analogues via a Michael addition-driven multicomponent reaction, Org. Chem. Front., 2020, 7, 987-992.
- 33 N.-j. Zhong, L. Liu, D. Wang and Y.-J. Chen, Enantioselective tandem reaction of chromone-derived Morita-Baylis-Hillman carbonates with benzylamines catalyzed by a trifunctional organocatalyst: the synthesis of chiral 3aminomethylene-flavanones, Chem. Commun., 2013, 49,
- 34 Q. Zhu and Y. X. Lu, Stereocontrolled creation of all-carbon quaternary stereocenters by organocatalytic conjugate addition of oxindoles to vinyl sulfone, Angew. Chem. Int. Ed.,

Open Access Article. Published on 23 September 2025. Downloaded on 9/24/2025 6:03:17 PM.

PM-NC

PM-NC

This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence.

No primary research results, software or code have been included and no new data  $_{\text{DOI: }10.1039/\text{D5OB01129A}}^{\text{View Article Online}}$  were generated or analysed as part of this review.