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Mayuri M. Naik, Vijayendra P. Kamat, Santosh G. Tilve

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## **Graphical Abstract**



## **Copper-mediated synthesis of coursetans** *via* C(sp<sup>2</sup>)-H functionalization: **Protective group free route to coursetrol and 4'**-*O*-methylcoursetrol

Mayuri M. Naik<sup>a</sup>, Vijayendra P. Kamat<sup>a</sup> and Santosh G. Tilve<sup>a, b,</sup> \*

<sup>a</sup> Department of Chemistry, Goa University, Taleigao-Plateau, Goa 403 206, India <sup>b</sup> Organic Chemistry Department, RUDN University, 6 Miklukcho-Maklaya str., Moscow 117198, Russian Federation \*Phone: 0832-6519317; Email: stilve@unigoa.ac.in

Key words: Coumestans, copper acetate, C-H functionalization, coumestrol, 4'-Omethylcoumestrol

**Abstract**: A simple and efficient two step synthesis of coumestans is described. The key reaction in the synthesis is the use of easily available  $Cu(OAc)_2$  for C-H functionalization of 3-(2-hydroxyphenyl)coumarin to give coumestan ring system *via* formal oxidative cyclization. This approach provided a short protective group free route to naturally occurring coumestrol and 4'-*O*-methylcoumestrol.

**1. Introduction**: *6H*-Benzofuro[3,2-*c*]chromen-6-ones, commonly known as coumestans are polycyclic ring systems having a coumarin ring and a benzofuran ring fused together. This ring system is found in many naturally occurring compounds distributed widely in plants.<sup>1</sup> Members belonging to this class of compounds includes wedelolactone, coumestrol, psoralidin, medicagol, lucernol, 4'-O-methylcoumestrol, desmethylwedelolactone, etc. (Figure 1). They exhibit numerous biological activities such as anticancer, estrogenic, phytoalexin activities, anti-venom, antibacterial, antifungal, cytotoxic, and antidepressant.<sup>2</sup> Some coumestans inhibit protein-tyrosine phosphatase 1B<sup>3a</sup> and some are used in the treatment of liver diseases.<sup>3b</sup> These immense biological activities of coumestans have been attracting chemists since decades as an interesting synthetic target.



Natural coumestans	<b>к</b> 1	<b>R</b> <sub>2</sub>	<b>R</b> 3	$\mathbf{K}_4$	$R_5$
Wedelolactone	OH	Н	OMe	OH	OH
Coumestrol	Н	Н	OH	Н	OH
Psoralidin	Н	prenyl	OH	Η	OH
Medicagol	Н	Н	OH	O-Cl	H <sub>2</sub> -O
Lucernol	Н	OH	OH	Н	OH
4'-O-Methylcoumestrol	Н	Н	OH	Η	OMe
Desmethylwedelolactone	OH	Н	OH	OH	OH

Fig 1. Naturally occurring coumestans.

Coumestans have been constructed *via* several synthetic approaches.<sup>4,5</sup> However many of these methods have their own limitations such as multistep syntheses, expensive reagents/catalysts usage, hazardous metal catalysts, difficulty in handling of reagents and/or its excessive requirement, troublesome reaction work up and product isolation. Hence continuous search for new method/reagent/catalyst for coumestan synthesis is pursued.

## 2. Results and discussion:

Our retrosynthetic analysis of coumestan ring system 1 suggested a straight forward two step approach *via* oxidative cyclization of 3-(2-hydroxyphenyl)coumarin 2 (Scheme 1). The required 3-(2-hydroxyphenyl)coumarin 2 can be conveniently obtained by condensation of salicylaldehyde 3 and 2-coumaranone 4 or 2-hydroxyphenylacetic acid 5.



Scheme 1. Retrosynthetic analysis of coumestan 1.

For the oxidative cyclization of **2** four conditions are reported in literature (Scheme 2).<sup>6</sup> First Pb(OAc)<sub>4</sub> in refluxing anhydrous benzene,<sup>6a</sup> later DDQ in refluxing anhydrous benzene,<sup>6b</sup> then PdCl<sub>2</sub> in presence of sodium acetate in DMF at 150  $^{\circ}C^{6c}$  and recently iodine in refluxing pyridine.<sup>6d</sup> These oxidative cyclization methods have some limitations including low product yields, utilization of expensive reagent and limited substrate scope. Hence there is a need to develop a method involving a suitable reagent which can overcome these limitations. To our knowledge Cu(OAc)<sub>2</sub> was not employed for coumestan synthesis *via* oxidative cyclization until the completion of this work.



1. Pb(OAc)<sub>4</sub> (1.5 equiv), anhyd. benzene (30 mL), reflux, 30 min<sup>6a</sup>

4. I<sub>2</sub> (1.0 equiv), anhy. pyridine, reflux, 15  $h^{6d}$ 

<sup>2.</sup> DDQ (1.0 equiv), benzene, reflux, 72  $h^{6b}$ 

<sup>3.</sup> PdCl<sub>2</sub> (1.0 equiv), NaOAc (13.6 equiv.), DMF, 150 °C, 24 h<sup>6c</sup>

## Scheme 2. Reported syntheses of coursetans 1 from 2 using various oxidizing agents.

Copper salts have been widely used in organic reactions owing to its cheap availability and low toxicity. Several reviews have appeared on its role either as reagent and/or catalyst.<sup>7</sup> In particular Cu(OAc)<sub>2</sub> is a mild reagent/catalyst known for the synthesis of several heterocycles.<sup>8</sup> It has gained considerable attention for its role in the intramolecular C-O cyclization *via* C-H functionalization<sup>8c-d</sup> for the construction of heterocyclic compounds. For example a combination of 1.2 equiv Cu(OAc)<sub>2</sub> and 0.2 equiv of Zn(OTf)<sub>2</sub> in toluene:DMSO (20:1) has provided 11-*H*-benzofuro[3,2-*b*]chromen-11-ones and 6*H*-benzofuro[2,3-*c*]chromen-6-ones *via* insertion of oxygen into electron rich aromatic rings.<sup>8c</sup> Interestingly, it has not been explored for the isomeric naturally occurring 6*H*-benzofuro[3,2-*c*]chromen-6-ones (coumestans). Continuing our interest in copper chemistry<sup>9</sup> we envisioned the role of Cu(OAc)<sub>2</sub> in the intramolecular C-O cyclization of 3-(2-hydroxyphenyl)coumarin **2** to form coumestan **1** under forcing conditions.

Table 1. Optimization of the reaction conditions for the synthesis of coumestans<sup>a</sup>

Fntry	Reagent	Solvent	Temperature	Time	Vield
Entry	Keagent	Borvent			
				( <b>h</b> )	(%)
1	Cu(OAc) <sub>2</sub>	Toluene	<mark>110</mark>	24	7
2 <sup>c</sup>	$\frac{\text{Cu(OAc)}_2}{\text{Cu(OTf)}_2}$	Toluene:DMSO	120	24	7
		(20:1)			
3	Cu(OAc) <sub>2</sub>	<i>p</i> -Xylene	<mark>138</mark>	24	20
4	Cu(OAc) <sub>2</sub>	Acetic acid	<mark>118</mark>	24	0
5	Cu(OAc) <sub>2</sub>	DMF	<mark>153</mark>	24	15
6	Cu(OAc) <sub>2</sub>	DMA	<mark>165</mark>	24	6
7	Cu(OAc) <sub>2</sub>	o-DCB	<mark>180</mark>	24	30
8	Cu(OAc) <sub>2</sub>	DMSO	<mark>189</mark>	24	ND
9	Cu(OAc) <sub>2</sub>	Diphenyl ether	<mark>258</mark>	6	76
10	Cu(OAc) <sub>2</sub> .H <sub>2</sub> O	Diphenyl ether	<mark>258</mark>	5	55
11	CuCl <sub>2</sub>	Diphenyl ether	<mark>258</mark>	24	69
12	CuBr <sub>2</sub>	Diphenyl ether	<mark>258</mark>	24	48
13	CuI	Diphenyl ether	<mark>258</mark>	24	19
14	Cu <sub>2</sub> O	Diphenyl ether	<mark>258</mark>	24	75

Y		conditions	
~ 2a	HO		1a -

15	CuO	Diphenyl ether	<mark>258</mark>	16	60
16	Cu(OTf) <sub>2</sub>	<i>p</i> -xylene	<mark>138</mark>	24	40
17	Cu(OTf) <sub>2</sub>	Diphenyl ether	<mark>258</mark>	16	63
18	Cu (nanopowder)	Diphenyl ether	<mark>258</mark>	24	74
19	Cu (metal powder)	Diphenyl ether	<mark>258</mark>	24	72
20	Pd(OAc) <sub>2</sub>	Diphenyl ether	<mark>258</mark>	18	86
21 <sup>d</sup>	$Pd(OAc)_2$	Diphenyl ether	<mark>258</mark>	24	RI
22 <sup>d</sup>	PdCl <sub>2</sub>	Diphenyl ether	<mark>258</mark>	24	RI
23 <sup>e</sup>	10% Pd/C	Diphenyl ether	<mark>258</mark>	12	RI
24 <sup>f</sup>	10% Pd/C	Diphenyl ether	<mark>258</mark>	12	RI
25 <sup>g</sup>	10% Pd/C	Diphenyl ether	<mark>258</mark>	12	RI
26 <sup>h</sup>	10% Pd/C	Diphenyl ether	<mark>258</mark>	12	78

<sup>a</sup> Conditions: 2a (0.4 mmol), reagent (0.4 mmol) and solvent 10 mL, under open air.

<sup>b</sup> Isolated yields.

<sup>c</sup> 0.48 mmol of Cu(OAc)<sub>2</sub>:0.08 mmol of Zn(OTf)<sub>2</sub> were used.

<sup>*d*</sup> 0.04 mmol of reagent was used.

e, f, g, h 10, 20, 30, 50 wt% of reagent was used respectively.

ND: Not determined.

RI: Reaction incomplete.

At the outset 3-(2-hydroxyphenyl)-2*H*-chromen-2-one **2a** was treated with anhydrous  $Cu(OAc)_2$  (1.0 equiv) in toluene under refluxing condition. To our delight the desired product **1a** was formed albeit in poor yield (Table 1, entry 1). Encouraged with this finding, we examined the reported  $Cu(OAc)_2$ :Zn(OTf)<sub>2</sub> in toluene:DMSO (20:1) procedure used for the synthesis of 11-*H*-benzofuro[3,2-*b*]chromen-11-ones.<sup>8c</sup> However, the yield did not increase. The reported C-H activation were achieved on aryl systems which were electron rich systems while in the present case we had to activate an elecron deficient double bond. Hence we envisaged that higher temperature of the reaction may give us the desired prouct. Hence, solvents having higher boiling points (entries 3-9) were examined. Among these solvents, diphenyl ether proved to be the ideal solvent as the product yield increased substantially to 76% (entry 9). With this result, we further screened different metal reagents (entries 9-26). Employing Cu(OAc)<sub>2</sub>.H<sub>2</sub>O diminished the product yield. Alternative Cu reagents such as CuCl<sub>2</sub>, CuBr<sub>2</sub>, CuI, Cu<sub>2</sub>O and CuO also showed the formation of product among which Cu<sub>2</sub>O gave highest yield of 75% (entries 10-15). The more reactive Cu(OTf)<sub>2</sub> gave 40% yield on

refluxing in *p*-xylene (entry 16) and in diphenyl ether the yield was increased to 63% with reduced time (entry 17). Cu metal also proved effective for this cyclization (entries 18-19).

Due to the shorter reaction time and the easy availability, anhydrous  $Cu(OAc)_2$  was chosen for further studies with respect to its concentration and temperature (Table 2). It must be noted that the absence of  $Cu(OAc)_2$  showed no product formation (entry 1). Catalytic amount of  $Cu(OAc)_2$  was effective in producing 66 and 70 % yields with longer reaction time (entries 2-3) and 1.0 equiv proved to be optimum  $Cu(OAc)_2$  concentration giving maximum yield at a faster rate (entry 4). Furthermore on increasing its concentration resulted in decreased product yield (entries 5-6). With 1.0 equiv of  $Cu(OAc)_2$  reactions were performed at different temperatures (entries 7-10) which revealed that the product yield increases with increase in temperature. Hence 258  $^{\circ}C$  was found to be the optimum temperature (entry 4).

Entry	Temperature	Cu(OAc) <sub>2</sub>	Time	Yield
	( <sup>0</sup> C)	(equiv.)	(h)	(%) <sup>b</sup>
1	<mark>258</mark>	0.0	6	00
2	258	0.2	24	66
3	258	0.5	12	70
4	258	<b>1.0</b>	6	76
5	258	1.2	11	70
6	258	2.0	14	56
7	100	1.0	24	25
8	150	1.0	24	50
9	170-180	1.0	12	62
10	200	<mark>1.0</mark>	10	64

Table 2. Optimization of Cu(OAc)<sub>2</sub> concentration and temperature for the synthesis of coumestan 1a from 2a<sup>a</sup>

<sup>a</sup> Conditions: **2a** (0.4 mmol),  $Cu(OAc)_2$  and diphenyl ether (10 mL) at above mentioned temperature under open air.

<sup>b</sup> Isolated yields

The optimum reaction conditions were then studied for the substrate scope (Table 3). Substituents on both the phenyl rings A and B were evaluated. Study on ring A revealed that the electron releasing methoxy and ethoxy substituent's reacted smoothly to give the desired products **1b-1d** in good yields. Naphthol group was very reactive to provide the expected coumestan **1e** in 80% yield. Monomethyl and dimethyl substituents were also successfully converted into the desired products **1f** and **1g**. Coumestans **1h-1j** were formed when

dimethoxy and methylenedioxy substitutions were examined on ring A. Hydroxy substituents also reacted to produce the required coumestans **1k** and **1l** without any need of protection thus exhibiting good efficiency and practicability of this method. Coumestans bearing electron withdrawing bromo and chloro groups **1m** and **1n** were synthesised in 53 and 67% yields respectively. Strong electron withdrawing nitro group was quite reactive enough to offer the desired product **1o**.

Table 3. Synthesis of coursetans via C-H activation using Cu(OAc)<sub>2</sub><sup>a,b</sup>



Sr. No.	Substituted 3-(2- hydroxyphenyl)-2 <i>H</i> - chromen-2-one (2)	Time (h)	Coumestan (1)	Yield <sup>a</sup> (%)
1	2a HO	6		76
2	МеО 0 0 2b но	13	MeO O O O D	66
3	мео но но	6		72
4	2d HO	13	1d of o	70
5	2e HO	8		80
6	2f но	10		65

7	2g HO	17		67
8	OMe MeO O O O	6	OMe MeO C C C C C C C C C C C C C C C C C C C	80
	2h но		1h 0-1-	
9	MeO MeO 2i HO	8	MeO O O MeO O Ii	77
10	2j но	14		61
11	он 0-0 2k но	16		62
12		14		68
13	Вг +0 0 2m но	24 18 °	Br OFO 1m	30 53 °
14	сі но сі но сі	24		67
15	о <sub>2</sub> N 20 но	13	$10 \xrightarrow{O \to O} 0$	54
16	2р но Оме	5		70
	MeO. (0. (0.			
17	2q HO OMe	5		65



<sup>a</sup> Conditions: **2** (0.4 mmol),  $Cu(OAc)_2$  (0.4 mmol) and diphenyl ether (10 mL), reflux under open air for 4-24 h. <sup>b</sup> Isolated yields.

<sup>c</sup>Cu(OAc)<sub>2</sub> (0.6 mmol) was used.

On successfully synthesising above derivatives, we went on to explore the substitution on ring B. When methoxy substituent was employed on ring B with or without substituents on ring A, reaction went on smoothly to afford diverse coursetans **1p-1u** in moderate to good

yields. Among these the isolation of naturally occurring<sup>10</sup> coumestrol dimethyl ether **1q**, dimethyl ether of sativol **1t** and trimethyl ether of lucernol **1u** was quite pleasing. The synthesis of coumestrol **1y** from compound **1q** is well known.<sup>11</sup> The presence of methyl group on ring B also successfully delivered coumestans **1v** and **1w** in 65 and 60% yields respectively without affecting the side chain.

Encouraged by the formation of hydroxyl coumestans **1k** and **1l** we applied this methodology towards the protective group free synthesis of naturally occurring 4'-*O*-methylcoumestrol<sup>1g,12</sup> **1x** and coumestrol<sup>1c,13</sup> **1y**. 4'-*O*-Methylcoumestrol was isolated from *alfalfa* and various other plant species whereas coumestrol was isolated from *alfalfa*, *ladino* clover and many forage crops. The higher binding affinity of coumestrol for ER $\beta$  than other phytoestrogens makes it one of the most potent phytoestrogen.<sup>14</sup> Several total syntheses of coumestrol<sup>4f,4q,11,15</sup> while a few of 4'-*O*-methylcoumestrol<sup>4n,15d,16</sup> have been reported. On subjecting the necessary starting materials to the above reaction conditions it was endearing to get both coumestans **1x** and **1y** in 59 and 55% yields respectively thus eliminating the need of protection-deprotection stratergies adopted in earlier reported synthetic methods. As most of the naturally occurring coumestans contain hydroxyl and/or methoxy group/s, our methodology provides a broad scope for synthesis of such natural members of coumestan family.

The protocol was then successfully tested for the preparation of **1a** from **2a** on a larger scale (2.0 mmol) thus demonstrating its utility. Further, a one pot procedure was attempted by mixing 2- coumaranone **4**, salicylaldehyde **3a**, Cu(OAc)<sub>2</sub> and NEt<sub>3</sub> in diphenyl ether as the solvent system. However, no formation of **1a** was observed (Scheme 3-*A*). Hence a stepwise one pot approach was developed wherein NEt<sub>3</sub> was removed before addition of Cu(OAc)<sub>2</sub> giving product **1a** in good yield (Scheme 3-*B*).



Scheme 3. Stepwise one pot synthesis of coumestan 1a

It is known that the presence of other metal impurity or "homeopathic" metal can also be responsible for such results.<sup>17</sup> Initially we had studied Pd metal for this oxidative cyclization. Pd(OAc)<sub>2</sub> when used in stoichiometric quantities gave coumestan in 86% yield (Table 1entry 20). Catalytic amount of Pd(OAc)<sub>2</sub> or PdCl<sub>2</sub> resulted in incomplete reaction (entries 21-22). Similar findings were also observed for Pd/C (entries 23-26). When 50 wt% of 10% Pd/C was used 78% of coumestan was formed after 12 h (entry 26). However, due to requirement of stoichiometric amount of expensive Pd metal we had not pursued it further. Subsequently it was concluded that Cu(OAc)<sub>2</sub> was the best source for the present C-H activation. But, it was essential to confirm that presence of Pd impurity was not responsible for this Cu(OAc)<sub>2</sub> mediated cylization. Hence ICP-MS analysis of Cu(OAc)<sub>2</sub> sample was carried out. It showed the presence of 1817.97 ppb palladium in Cu(OAc)<sub>2</sub>. Hence to study whether Pd has any role in this oxidative cyclization we added 0.01 equiv of Pd(OAc)<sub>2</sub> in Cu(OAc)<sub>2</sub> and repeated the experiment. However, there was not much change in the yield of the product or duration of reaction. Similar was the case when 0.1 equiv of PdCl<sub>2</sub> was added. Previously, Gong et al.<sup>6c</sup> had employed stoichiometric amount of PdCl<sub>2</sub> for such cyclization. Our attempt to use PdCl<sub>2</sub> under Wacker oxidation conditions [CuCl<sub>2</sub> in DMF:DMA (1:1) solvent] at 150 °C produced only trace amount of product. All these results suggest the trace amount of Pd impurity in Cu(OAc)<sub>2</sub> may not be responsible factor for this oxidative cyclization. However the role of Pd present in ppb level in contact with copper and a synergestic effect of this cannot be ruled out completely. Further study on the synergestic effects of other metal dopants/additives is a subject of future study.

To check whether the mechanism is following a radical pathway the reaction was performed in presence of a radical scavenger TEMPO. It had no effect on the yield of product **1a** suggesting an alternative mechanism. Based on this observation a speculative mechanism is proposed for this intramolecular C-O cyclization (Scheme 4). Cu(II) from Cu(OAc)<sub>2</sub> binds to electronegative hydroxyl oxygen atom of substrate **2** with the liberation of one molecule of acetic acid to form intermediate **6**. Intramolecular oxidative addition of copper to the C-H bond renders **8** via the elimination of another molecule of acetic acid from **7**. Finally the reductive elimination of **8** furnishes coumestan **1** and metallic copper. The formation of metallic copper was confirmed from recorded XRD (see supporting information) of the residue left after the reaction and also from the copper mirror deposits on the walls of the

reaction flask. The presence of  $Cu_2O$  seen in the XRD could be due to the aerobic oxidation of Cu at high reaction temperature.

Recently, after our studies, Zou *et al.*<sup>18</sup> have reported a synthesis of naturally occurring coumestrol and aureol *via* a combination of a Perkin condensation and a Cu(II)-catalyzed hydroxylation/aerobic oxidative microwave mediated coupling reaction.



Scheme 4. Proposed mechanism for the formation of coumestans using Cu(OAc)<sub>2</sub>

#### **3.** Conclusion

In conclusion, we have developed an efficient methodology for the synthesis of coumestans. The method implements economical  $Cu(OAc)_2$  as the reagent in absence of any additional reagent/additive in diphenyl ether solvent *via* C-H activation. Although the reaction temperature is high, simple reaction procedure, large substrate scope, effortless product isolation & good yields make this method attractive over reported methods. Also direct synthesis of hydroxy substituted coumestans without protection strategies makes the method noteworthy. Additional advantages are the one pot synthesis and possible use of catalytic amount of  $Cu(OAc)_2$ . Further scope of  $Cu(OAc)_2$  as an oxidative cyclizing agent employing ligands will be undertaken in near future.

#### 4. Experimental

**4.1. General remarks.** All the compounds were characterized by spectral analysis (IR; <sup>1</sup>H NMR; <sup>13</sup>C NMR) and comparison of their melting points with the literature reports. All the

melting points were uncorrected. Infrared (IR) spectra were recorded in a FTIR instrument using KBr and wavenumbers given in cm<sup>-1</sup>. <sup>1</sup>H (400 MHz) and <sup>13</sup>C (100 MHz) NMR spectra were recorded in DMSO- $d_0$ /CDCl<sub>3</sub> as solvent and TMS as an internal standard. Coupling constants are reported in Hz. High resolution mass spectra (HRMS) were recorded on Q-TOF MS instrument with an electrospray source in ESI mode at IISc, Bangalore. Quantification of palladium impurity in Cu(OAc)<sub>2</sub> was detected by using ICP-MS. All the solvents were distilled prior to use. Column chromatography was performed on 60-120 mesh silica gel. All the chemical reagents and diphenyl ether were purchased from commercial sources and used without further purification unless otherwise stated. Anhydrous Cu(OAc)<sub>2</sub> was purchased from Sigma Aldrich. Substrates **2a** to **2o** were synthesized using literature procedure from 2coumaranone **4** and substituted salicylaldehyde derivatives **3**.<sup>19</sup> Similarly **2p** to **2y** were synthesized from substituted salicylaldehyde derivatives **3** and 2-hydroxyphenylacetic acid derivatives **5** using literature procedure.<sup>6c</sup> All the reactions were carried out under atmospheric conditions without any special cautions unless otherwise stated.

#### **4.2. General procedures**

## (I) General procedure for the synthesis of substrates 2a-20<sup>19</sup>

Substituted salicylaldehyde derivative **3** (455 mg, 3.7 mmol) and 2-coumaranone **4** (500 mg, 3.7 mmol) were mixed together in a round bottom flask. To it triethylamine (15 mL) was added and refluxed for 1 h. After 1 h triethylamine was removed under vacuum and the crude solid was recrystallized from ethanol to afford pure product 3-(2-hydroxyphenyl)-2*H*-chromen-2-one **2a-2o**.

## (II) General procedure for the synthesis of substrates 2p-2y<sup>6c</sup>

Substituted salicylaldehyde derivative **3** (122 mg, 1.0 mmol), substituted 2hydroxyphenylacetic acid **5** (182 mg, 1.0 mmol), sodium acetate (410 mg, 5.0 mmol) and acetic anhydride (245 mg, 2.4 mmol) were mixed together in a round bottom flask. To it acetic acid (4 mL) was added and refluxed for 24 h. After 24 h the solvent was removed under vacuum and water was added to it. The crude solid obtained was filtered and then loaded on column (eluent: petroleum ether/ethyl acetate) to afford pure product 3-(2hydroxyphenyl)-2*H*-chromen-2-one **2p-2y**.

#### (III) General procedure for the synthesis of coumestans 1a-1y

Cu(OAc)<sub>2</sub> (77 mg, 0.4 mmol) was added to substituted 3-(2-hydroxyphenyl)-2*H*-chromen-2one **2a-2y** (100 mg, 0.4 mmol) in a 25 mL round bottom flask. To it 10 mL of diphenyl ether was added. The resulting mixture was then heated to reflux for 4-24 h. After completion of the reaction (monitored by TLC) the reaction mass was cooled to room temperature. Without any further work up it was directly loaded on column (eluent: petroleum ether/ethyl acetate) to afford pure product 6*H*-benzofuro[3,2-*c*]chromen-6-one **1a-1y**.

#### (IV) Procedure for the synthesis of coumestan 1a on 0.5 g scale

Cu(OAc)<sub>2</sub> (382 mg, 2.1 mmol) was added to 3-(2-hydroxyphenyl)-2*H*-chromen-2-one **2a** (500 mg, 2.1 mmol) in a 50 mL round bottom flask. To it 20 mL of diphenyl ether was added. The resulting mixture was then heated to reflux for 21 h. After completion of the reaction (monitored by TLC) the reaction mass was cooled to room temperature. Without any further work up it was directly loaded on column (eluent: petroleum ether/ethyl acetate, v/v = 10/1.5) to afford product 6*H*-benzofuro[3,2-*c*]chromen-6-one **1a** as colorless solid (296 mg, 60%).

#### (V) Procedure for stepwise one pot synthesis of coumestan 1a

2-coumaranone **4** (57 mg, 0.4 mmol) and salicylaldehyde **3a** (52 mg, 0.4 mmol) were mixed together in a 25 mL round bottom flask. To it triethylamine (5 mL) was added and refluxed for 1 h. Triethylamine was removed under vacuum and to the product formed 3-(2-hydroxyphenyl)-2*H*-chromen-2-one **2a**, Cu(OAc)<sub>2</sub> (77 mg, 0.4 mmol) and 10 mL of diphenyl ether were added. The resulting mixture was then heated to reflux for 6 h. After completion of the reaction (monitored by TLC) the reaction mass was cooled to room temperature. Without any further work up it was directly loaded on column (eluent: petroleum ether/ethyl acetate, v/v = 10/1.5) to afford product 6*H*-benzofuro[3,2-*c*]chromen-6-one **1a** as colorless solid (70 mg, 71%).

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#### 6. References and Notes

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