DOI: 10.1002/jms.4425

# **RESEARCH ARTICLE**



# Unexpected detection of 3-aroylbenzofuran side products in the preparation of 2-arylbenzofurans: Identification, characterization, and comparison with chalcone's fragmentation patterns using EI/MS<sup>n</sup>

Michela Begala 问 | Giovanna Lucia Delogu

Department of Life and Environmental Sciences, Unit of Drug Sciences, University of Cagliari, Cagliari, Italy

#### Correspondence

Michela Begala, Department of Life and Environmental Sciences, University of Cagliari, via Ospedale 72, 09124, Cagliari, Italy. Email: michelabegala@unica.it

Funding information University of Cagliari

### Abstract

A gas chromatography-mass spectrometry study of the intramolecular Wittig reaction revealed, together with the expected 2-phenylbenzofuran, the formation of an unexpected side product that has not been reported until now. This study reports the identification of the by-product, ie, the 3-benzoyl-2-phenylbenzofuran, on the base of its mass spectrometric behaviour using a combination of electron ionization, exact mass measurement, multiple stage mass spectrometry, and labelled compounds. This study reports the common fragmentation pathways and discusses possible fragment structures of characteristic ions from a series of 3-aroyl-2-arylbenzofuran derivatives obtained as by-product under Wittig conditions. Emphasis is laid on the formation and structure investigation of the [M-H]<sup>+</sup> and [M-OH]<sup>+</sup> ions. Our results showed interesting analogies with the mass spectrometric behaviour of chalcones.

### KEYWORDS

[M-OH]<sup>+</sup>, 3-aroylbenzofurans, chalcone-like fragmentations, intramolecular Wittig reaction, MS<sup>n</sup>

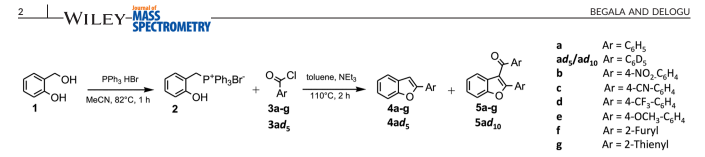
# **1** | INTRODUCTION

2-Phenyl[b]benzofurans serve as core structures of many natural and artificial compounds of biological and medical importance.<sup>1</sup> In the course of a program directed towards the synthesis of novel monoamine oxidase and butyrylcholinesterase inhibitors,<sup>2,3</sup> we planned to synthesize 2-phenylbenzofurans using an intramolecular Wittig procedure due to the accessibility and simplicity of this methodology.<sup>4,5</sup> In particular, compounds **4a-g** were prepared from the appropriate triphenylphosphonium salt **2** and the commercially available aroyl chlorides **3a-g** (Scheme 1).

The reaction mixtures were analysed by gas chromatography (GC) coupled with mass spectrometry (MS) because the reaction products are susceptible to analysis by GC for their volatility and thermal stability. However, while developing this procedure, GC/MS analysis of the reaction mixture, obtained from the salt **2** and the benzoyl chloride **3a**, revealed that, together with the desired product of cyclization **4a**, the unexpected side-product **5a** was present. The same trend was observed in the crude reaction mixtures of the main products **4b-g**, obtained from the reaction of **2** with a variety of aroyl chlorides (**3b-g**). Although the Wittig reaction has been described in several papers regarding the preparation of 2-arylbenzofuran derivatives, the formation of secondary products was not mentioned.<sup>4,5</sup>

Identification of unknown peaks is useful to permit discovery of novel or unexpected reaction products and play an important role in comprehending the reaction mechanism by which this reaction evolves. However, isolation and purification of sufficiently large quantities of by-product compounds for their unambiguous identification and characterization by different instrumental techniques, such as infrared spectroscopy and nuclear magnetic resonance, is a very complex and time-consuming process. Nowadays, GC-MS coupled to multiple stage mass spectrometry (MS<sup>n</sup>) is an invaluable tool for the rapid identification and structural characterization of unknown compounds, especially when low amount of analyte or not very pure samples are

1



SCHEME 1 Synthetic route towards 2-phenylbenzofurans 4a-g and 3-benzoyl-2-phenylbenzofurans 5a-g

available for the analysis. Hence, we have undertaken a detailed investigation on the unknown side products aimed to identify their structure and to elucidate their mass spectrometric behaviour. To this aim, EI-MS<sup>n</sup> experiments were performed using a quadrupole ion trap (IT) analyser as it is very efficient multiple-stage mass spectrometers.<sup>6</sup>

### 2 | EXPERIMENTAL

### 2.1 | MS analysis

All experiments were performed with a Varian Saturn 2000 IT mass spectrometer, operating under electron ionization (EI) conditions (electron energy 70 eV, emission current 20 mA, ion-trap temperature 200°C, manifold temperature 80°C, and automatic gain control target 21.000) with the IT operating in scan mode (scan range from m/z 40–400 at a scan rate of 1 scan/s), coupled with a Varian 3800 gas chromatograph (Varian, Walnut Creek, CA).

Collision-induced dissociation (CID) experiments were carried out by using helium as the collision gas (gas purity [He] was 99.9999%). For  $MS^n$  experiments, the supplementary rf voltage (45-55 V) was varied in such a way that the relative abundance of the surviving precursor ions was 5–15%. An isolation time of 10 milliseconds, excitation time of 40 milliseconds and an isolation width of 3 *m/z* for the precursor ion were used. Each MS/MS spectrum was an average of 5 scans.

Compounds **5a-g** (1- $\mu$ L aliquots of 1.0 × 10<sup>-5</sup> M solutions in dichloromethane) were introduced into the gas chromatographer inlet. An Agilent J&W VF-5 ms low-bleed/MS GC capillary column (30 m, 0.25 mm i.d., 0.25-mm film thickness) (Agilent Technologies Inc., Wilmington, DE, USA) was used. The oven temperature was programmed from 150°C (held for 2 min) to 310°C at 30°C/minute (held for 2 min). The temperature was then ramped to 350 at 20°C/minute. The transfer line was maintained at 180°C and the injector port (30:1 split) at 290°C.

HRMS: MS analyses were performed on a Agilent 6520 LC-ESI(+)-QTOF-MS operated in the positive ion mode. Compounds (1 mg) were dissolved in acetonitrile (CH<sub>3</sub>CN, 0.5 ml), and after injection mass spectral data were acquired in the range m/z of 100–1500 with an acquisition rate of 1.35 spectra/s, averaging 10 000 transients. The source parameters were adjusted as follows: drying gas temperature 250°C, drying gas flow rate 5 L/min, nebulizer pressure 45 psi, and fragmentor voltage 150 V.

### 2.2 | Materials and reagents

All reagents and solvents were purchased from Sigma-Aldrich (Sigma-Aldrich Srl, Milan, Italy) and Alfa Aesar (Thermo Fisher, Kandel, Germany) and used without further purification.

Compounds **4a** and **5a** were obtained by 2-benzyloxy-benzyl triphenyl phosphomium bromide **2** (1.1 mmol) and benzoyl chlorides (1.1 mmol) in the presence of  $Et_3N$  in toluene using the Wittig methodology described by Hercouet and Le Corre.<sup>4,5</sup> Compounds **4b-g** and **5b-g** were prepared using an adaptation of this procedure.<sup>7</sup>

2-(Hydroxyldideuterium-methyl) phenol  $1d_2$  was prepared starting from the methyl 2-hydroxybenzoate in THF at 0°C using lithium aluminum deuteride, as described in literature.<sup>8</sup>

The reference compound 3-benzoyl-2-phenyl-benzofuran was prepared by direct Friedel-Craft acylation of the 2-phenylbenzofuran **4a** with benzoyl chloride using  $AICI_3$  as Lewis acid in anhydrous dichloromethane.<sup>9</sup>

3-Benzoyl-2-phenyl-d<sub>5</sub>-benzofuran  $5ad_5$  was synthesized in the same way as 3-benzoyl-2-phenyl-benzofuran starting from 2-phenylbenzofuran-d<sub>5</sub>  $4ad_5$ .

2-Phenylbenzofuran-d<sub>5</sub> **4ad**<sub>5</sub> and 3-benzoyl-d<sub>5</sub>-2-phenyl-d<sub>5</sub>benzofuran **5ad**<sub>10</sub> were prepared starting from 2-benzyloxy-benzyl triphenyl phosphomium bromide **2** and benzoyl chloride-d<sub>5</sub> in toluene and Et<sub>3</sub>N using the procedure reported elsewhere.<sup>7</sup>

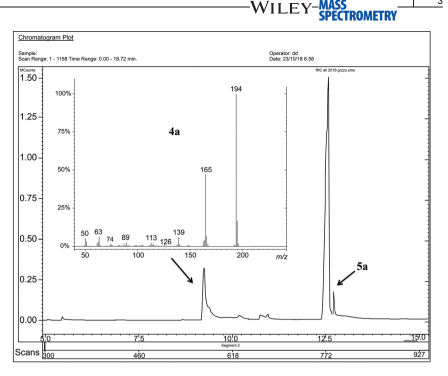
3-Phenyl-4H-chromen-4-one **6** (isoflavone) was prepared by Suzuki-Miyaura cross-coupling of 3-iodochromone with benzenboronic acid catalysed by Pd(0)/C.<sup>10,11</sup>

### 3 | RESULTS AND DISCUSSION

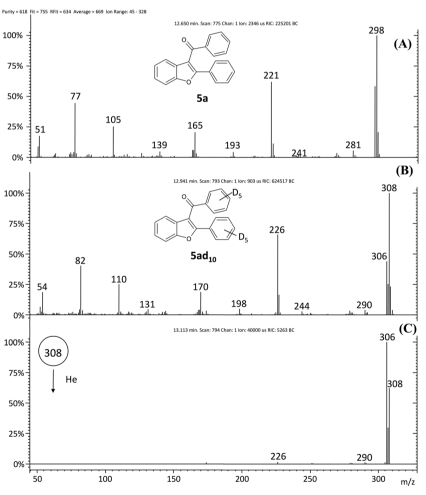
### 3.1 | Identification of the by-product 5a

A typical gas chromatogram of the reaction mixture from the triphenylphosphonium salt of 2-hydroxybenzyl alcohol and the benzoyl chloride is presented in Figure 1. As can be seen, together with the desired 2-phenylbenzofuran **4a** ( $T_R$  9.2 min) and the triphenylposphine oxide ( $T_R$  12.5 min), one further peak is present at higher retention time ( $T_R$  12.6 min), corresponding to the unexpected product **5a**.

By comparing the data obtained from the El mass spectra of compounds **4a** and **5a** (Figures 1 and 2A), it was enabled to define which part of the molecule of 2-phenylbenzofuran **4a** was also present in the structure of the unknown compound **5a**. In particularly, we focused



**FIGURE 1** Gas chromatography-mass spectrometry analysis of the reaction mixture along with the mass spectrum of the 2-phenylbenzofurane **4a** (9.36 min)



**FIGURE 2** Electron ionization mass spectra of compounds (A) 5a (*m*/*z* 298) and (B) the labelled  $5ad_{10}$  (*m*/*z* 308). (C) CID of  $5ad_{10}$ 

our attention on the diagnostic ion at m/z 193 in the mass spectrum of 5a (Figure 2A). This ion most likely corresponds to the intact 2-phenylbenzofuran molecule deprived of a H atom. In fact, under

MS/MS experiments, m/z 193 exhibits the loss of CO typical of the benzofuran nucleus to give the ion at m/z 165, which is also observed in the El and MS/MS mass spectra of the 2-phenylbenzofuran **4a** (M<sup>+.</sup>

# 4 WILEY-MASS SPECTROMETRY

m/z 194).<sup>12-14</sup> As the sum of m/z value of the fragment ions at m/z 193 and m/z 105, absent in the mass spectrum of **4a**, is equal to m/z M<sup>+</sup>. (m/z 298), these two ions could be complementary ions, representing two parts of the molecule; the rest of the structure was determined by comparison of the data achieved from the identification of the ion at m/z 105, with that from accurate mass measurement of the unknown compound **5a**. Under CID experiments, the m/z 105 ion decomposes into ions at m/z 77 and 51 thus indicating the presence of a benzoyl group (C<sub>6</sub>H<sub>5</sub>CO) in the structure of **5a**.<sup>15</sup> Accordingly, accurate mass measurement for the side product **5a** reveals a C<sub>21</sub>H<sub>14</sub>O<sub>2</sub> (calc. 299.1066 exp. 299.1066) composition, showing that the mass difference between **5a** and the 2-phenylbenzofuran **4a** (C<sub>14</sub>H<sub>10</sub>O) was in seven carbon atoms, four hydrogen atoms, and one oxygen (104 Da) suggesting the addition of a benzoyl group to the main product **4a**.

Even more, when the reaction was performed using the labelled benzoyl chloride- $d_5$  **3** $ad_5$  in place of the benzoyl chloride **3**a, we found that the molecular ion shifted from m/z 298, for compound **5**a, to m/z 308, for the labelled compound **5** $ad_{10}$ , thus confirming that two molecules of the labelled acyl chloride contribute to the structure of the side product (Figure 2B). The general structure of acylated 2-phenylbenzofuran was than proposed for the unknown compound **5**a.

The position of the acyl group was then investigated using the 2-hydroxybenzyl alcohol- $d_2$   $1d_2$  as starting reagent. The Wittig reaction so performed leads to the unlabelled acyl benzofuran **5a** at m/z 298. These data clearly show the lack of methine hydrogen in the furan ring of compound **5a**, thus demonstrating that the benzoyl group is attached to the C-3 position of the benzofuran nucleus. Our results indicate, therefore, that the side products **5a** most likely have the 3-aroylbenzofuran structure depicted in Scheme 1. The model 3-benzoyl-2-phenylbenzofuran was prepared by Friedel-Craft acylation of the 2-phenylbenzofuran **4a** and was analysed by MS, as a final confirmation of the proposed structure for **5a**. A detailed study aimed to provide a much deeper insight into the reaction mechanism that lead to the 3-aroyl derivative have been reported elsewhere.<sup>7</sup>

These findings are of particular interest since the 3-arovI-2arylbenzofuran scaffold constitutes the core of many pharmaceutical candidates such as amiodarone,<sup>16</sup> LY 320135.<sup>17</sup> drug benzbromarone,<sup>18</sup> and SKF-64346.<sup>19</sup> Moreover, from synthetic viewpoint, this procedure represents a new regioselective and versatile synthetic approach to 3-acyl derivatives.<sup>7</sup> However, to the best of our knowledge, mass spectrometric investigation of 3-aroylbenzofurans has received only little attention with most of the emphasis devoted to the plasma and urine detection of few biologically active compounds, benziodarone, desethylamiodarone,<sup>20,21</sup> ie. amiodarone, and benzbromarone.22

A detailed study dealing with the fragmentation of aroyl benzofurans has also appeared; however, it only covered the ESI MS/MS behaviour of 2-aroyl derivatives.<sup>23</sup>

Therefore, as a material for further investigations, we present a systematic and detailed study on the El-induced fragmentation of 3-benzoyl-2-phenylbenzofurans. To this aim, we selected several ring substituted 3-acyl derivatives obtained by our procedure and subjected them to El MS<sup>n</sup> experiments using an IT mass spectrometer.

### 3.2 | Fragmentation typical for 3-aroyl-2-arylbenzofurans

The EI mass spectra of compounds 5a-g are depicted in Table 1. We divided the structure into three parts: the benzofuran scaffold, the aryl group (ring A), and the aroyl group (ring B) (see Scheme 2 referred to compound 5a, Ar =  $C_6H_5$ ). Two sets of acylium ions were observed resulting from the two competing  $\alpha$ -cleavages next to the carbonyl group, ie, the [M-Ar]<sup>+</sup> ions *a* and the ArC=O<sup>+</sup> ions *b*. Both types of acylium ions under MS/MS conditions undergo the typical loss of CO to generate, respectively, the 2-phenylbenzofuran fragment ions *c* (complementary to ion *b*) and the phenylium ions *d* (complementary to ion *a*), which, therefore, allow an easy identification of the original intact molecule (vide supra). The only exception to this trend was observed for the nitro derivatives 5b. Under CID experiments, each of the acylium ions from **5b**, ie, the  $[M-Ar]^+$  ion of m/z 266 and the 4-nitrobenzoylcation of m/z 150, does not show the direct losses of CO. Instead, they both form characteristic fragment ions at m/z 236 and 220 and at m/z 120 and 104, respectively, due to the losses of NO and of NO2, which only subsequently lose CO molecule (see the Supporting Information).<sup>24</sup>

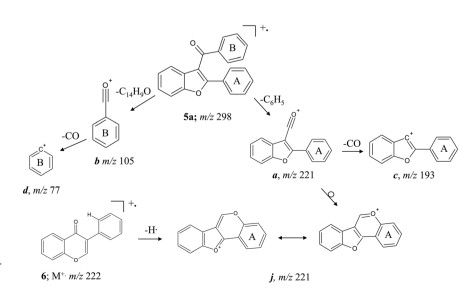
We also examined how the intensity of ions *b* is influenced by substituents on rings B or by heteroaromatic rings, for instance, furyl and thienyl moieties. Concerning compounds **5b-e**, we found that the intensity of the acylium ions *b* is strongly influenced by the nature of the substituent on the benzoyl group (Table 1). In the mass spectrum of compound **5e**, bearing the methoxy group on the ring B, the *p*-methoxybenzoyl cation (*m*/*z* 135) is particularly intense (relative abundance, R.A. 36%). This behaviour can be explained with the ability of the *p*-methoxy group to exert resonance stabilization of the *p*-methoxybenzoyl cation. The mass spectra of the heteroaryl compounds **5f** and **5g** exhibit the ArC=O<sup>+</sup> ions rather intense (R.A. 44 and 37%) reasonably because the 2-furyl and 2-thienyl substituents have electronic effect comparable with that of *p*-anisyl group, well recognized as an overall electron-donating system.

In the case of compounds **5b-d** bearing electron withdrawing groups, NO<sub>2</sub>, CN, and CF<sub>3</sub>, the formation of the ArC=O<sup>+</sup> ions is strongly suppressed (R.A. 4–16%) in favour of the [M-Ar]<sup>+</sup> ions (R.A.34–63%). Moreover, the stronger the electron-withdrawing effect on the ring B, the lower is the relative abundance of the benzoyl cation **b**.

The  $[M-Ar]^+$  acylium ions *a* constitute the most prominent fragment ions in the mass spectra of all compounds (R.A. 34–66%), with the only exception of compounds **5e-g**, for which the formation of ion *b* is the most favoured process (*vide supra*). A possible representation of the  $[M-Ar]^+$  ions is the resonance-stabilized oxonium structure *j* depicted in Scheme 2, which could account for the high intensities observed. We therefore thought of interest to compare the CID mass spectrum of the  $[M-Ar]^+$  ions from **5a** with that of the oxonium ion at *m/z* 221 formed in the El mass spectrum of isoflavone **6** (*m/z* 222),<sup>25</sup> used as reference ion (Figure 3). As can be seen, the CID mass spectra are very similar, proving that the cyclization to structure *j* occurs to some extents. There is, however, a difference in the relative intensity 
 TABLE 1
 EI (70 eV) mass spectra of compounds 5a-g

	5a	5ad <sub>10</sub>	5ad₅	5b	5c	5d	5e	5f	5g
M <sup>+.</sup>	298 (100)	308 (100)	303 (100)	388 (100)	348 (100)	434 (100)	358 (100)	278 (100)	310 (100)
[M-H] <sup>+</sup>	297 (53)	307(20)	302 (29)	387 (18)	347 (40)	433 (44)	357 (2)	277 (3)	309 (6)
[M-D] <sup>+</sup>		306 (44)	301 (34)						
$[M-CH_3]^+$							343 (5)		
[M-H-CH <sub>3</sub> ] <sup>+.</sup>							342 (10)		
$[M-OH]^+$	281 (5)			371 (5)	331 (2)	417 (3)	341 (4)	261 (5)	293 (5)
$[M-OD]^+$		290 (4)	285 (3)						
$[M-F]^+$						415 (7)			
[M-CO] <sup>+.</sup>								250 (55)	282 (9)
[M-NO] <sup>+</sup>				358 (5)					281 (6)
$[M-SH]^+$									277 (18)
$[M-CH_3-CO]^+$							315 (5)		
[M-H-NO <sub>2</sub> ] <sup>+.</sup>				341 (17)					
$[M-CO-CHO]^+$								221 (55)	
$[M-CF_3]^+$						365 (12)			
[M-Ar] <sup>+</sup> a	221 (56)	226 (66)	226 (62)	266 (34)	246 (54)	289 (63)	251 (14)	211 (7)	227 (11)
[a-CO] <sup>+</sup> c	193 (5)	198 (5)	198 (3)	-	218 (2)	261 (2)	223 (3)	183 (2)	199 (2)
[ <b>a</b> -NO] <sup>+.</sup>				236 (20)					
[a-CH <sub>3</sub> CO] <sup>+.</sup>							208 (4)		
[ <b>a</b> -NO <sub>2</sub> ] <sup>+.</sup>				220 (5)					
[ <b>c</b> -CO] <sup>+</sup>	165 (20)	170 (21)	170 (20)	-	190 (15)	233 (8)	195 (9)	155 (12)	171 (6)
ArCO <sup>+</sup> b	105 (24)		105 (25)	150 (4)	130 (12)	173 (16)	135 (36)	95 (44)	111 (37)
C <sub>6</sub> D <sub>5</sub> CO⁺ <b>b</b>		110 (25)							
[ <b>b</b> -CO] <sup>+</sup> <b>d</b>	77 (31)	82 (37)	77 (38)	-	102 (20)	145 (19)	107 (13)	67 (2)	83 (7)
[ <b>b</b> -NO] <sup>+.</sup>				120 (7)					
[ <b>b</b> -NO <sub>2</sub> ] <sup>+.</sup>				104 (5)					

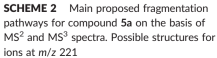
Abbreviation: EI: electron ionization.

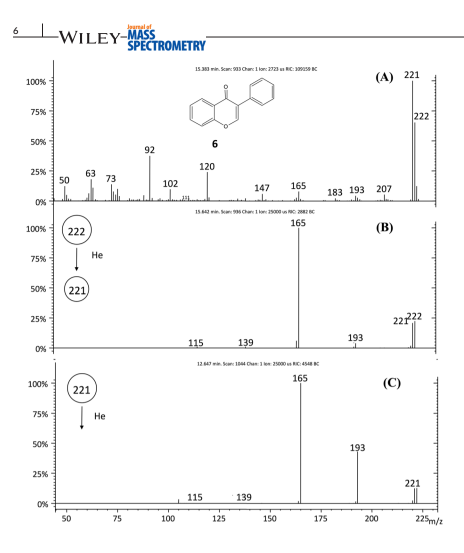


5

ROMETRY

Wiley





of the m/z 193 ion. This ion that we assign to the CO loss is more abundant for the compound **5a** (45% vs 4%). These data can be rationalized by the fact that part of the m/z 221 ion population of **5a** posses the open structure *a*, which undergoes easier this fragmentation.

In addition to the  $\alpha$ -cleavages, characteristic for carbonyl compounds, some further interesting losses were observed under EI-MS and MS/MS experiments, ie, the losses of H<sup>-</sup> and of OH<sup>-</sup> radicals from the molecular ion.

### 3.3 | Elimination of a hydrogen radical

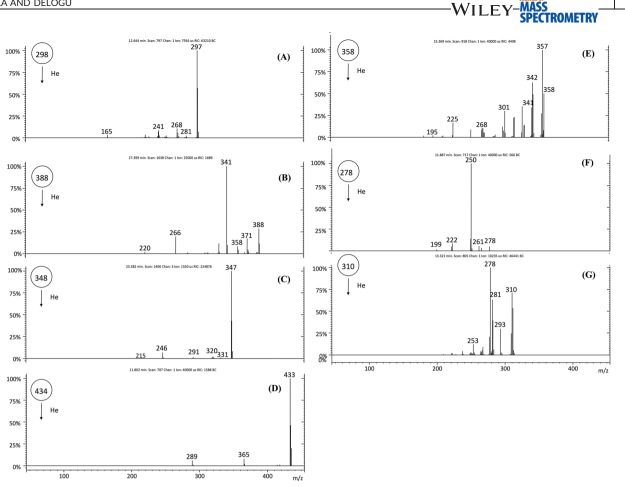
The EI mass spectra of the 3-benzoyl-benzofurans **5a-e** show intense  $[M-H]^+$  ions (Table 1). In the MS/MS spectra, the H<sup>-</sup> loss was even more pronounced (Figure 4). Only for compounds **5b** and **5e**, the H<sup>-</sup> loss occurred together with the loss of the substituent or of part of it as they show the total loss of 47 and 16 amu from the molecular ion. We suggest that the first step is the loss of a hydrogen atom and that NO<sub>2</sub> or CH<sub>3</sub> are lost from the even-electron ion so formed. Thus, the high abundant ions at m/z 341 and m/z 342 were attributed to the [M-H-NO<sub>2</sub>]<sup>+</sup> and the [M-H-CH<sub>3</sub>]<sup>+</sup> for compounds **5b** and **5e**, respectively (Figures 4B and 4E).

**FIGURE 3** (A) Electron ionization mass spectrum of isoflavone **6**. Collision-induced dissociation of the m/z 221 ions of (B) isoflavone **6** and (C) compound **5**a

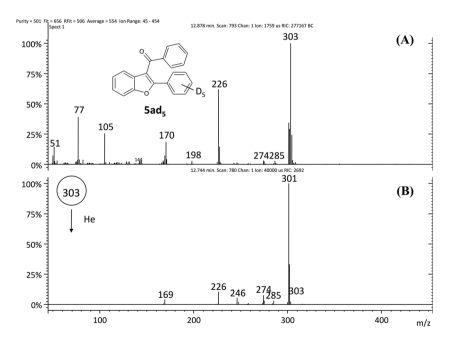
To get insight into the structure of the [M-H]<sup>+</sup> specie, we first investigated the origin of the H<sup>-</sup> lost. To this aim, the MS/MS spectrum of the unsubstituted compound 5a (m/z 298; Figure 4A) was compared with that of its deuterium labelled analogue 5ad<sub>10</sub> (rings A and B =  $C_6D_5$ ; m/z 308. Figure 2C). Under this condition, 5a formed the stable ion at m/z 297, with a R.A. of 100%. The loss of 2 amu (m/z 306) from the molecular ion of the labelled derivatives 5ad<sub>10</sub> (m/z 308) clearly indicates that the source of the H<sup> $\cdot$ </sup> eliminated is the phenyl group in position 2 (ring A) and/or the aroyl group in position 3 (ring B). MS/MS experiments on the 3-benzoyl-2-phenyl- $d_5$ benzofuran **5ad**<sub>5</sub> (Figure 5B) has helped to further elucidate the origin of the H<sup>-</sup> loss. Under MS<sup>2</sup> experiments the molecular ion of the labelled derivative **5ad**<sub>5</sub> (ring A =  $C_6D_5$ ; m/z 303) shows the preferential loss of deuterium radical from the ring A (m/z 301 RA 100%), rather than the loss of H<sup> $\cdot$ </sup> from the benzoyl group (ring B; m/z 302 R. A. 25%). This result indicates that the former process requires less energy than the latter and consequently is more favoured. In fact, MS<sup>n</sup> experiments performed using an IT, privilege processes with the lowest critical energies.<sup>26</sup> Accordingly, a different trend was observed in the full-scan EI mass spectrum of 5ad<sub>5</sub>, which displayed the ions at m/z 302 and 301 with similar intensities (R.A. 29 and 34%; Figure 5A).

We supposed that the driving force for the loss of the aromatic hydrogen radical preferentially from ring A lay in the formation of a highly resonance-stabilized ion, ie, the oxonium ion k reported in





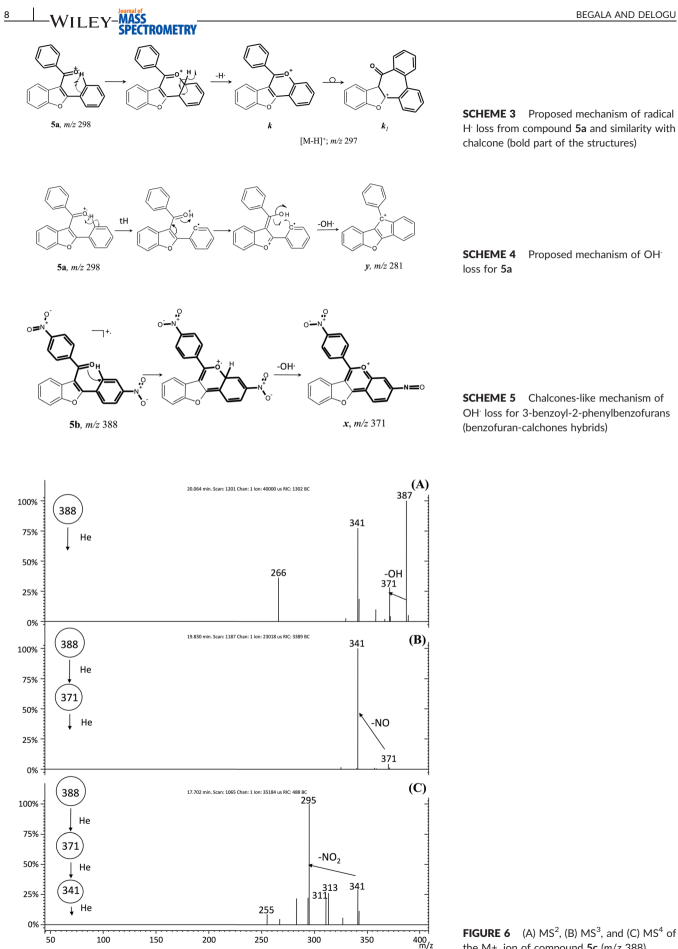
**FIGURE 4** MS/MS of compounds **5a-g**. A, MS<sup>2</sup> of **5a** (*m*/*z* 298); (B) MS<sup>2</sup> of **5b** (*m*/*z* 388); (C) MS<sup>2</sup> of **5c** (*m*/*z* 348); (D) MS<sup>2</sup> of **5d** (*m*/*z* 434); (E) MS<sup>2</sup> of **5e** (*m*/*z* 358); (F) MS<sup>2</sup> of **5f** (*m*/*z* 278); (G) MS<sup>2</sup> of **5g** (*m*/*z* 310)



**FIGURE 5** (A) Electron ionization mass spectrum and (B) MS/MS of compound **5***ad*<sub>5</sub> (*m*/*z* 303)

Scheme 3. A possible mechanism consists in an intramolecular aromatic substitution reaction; this might occur through the interaction of the carbonyl group with the 2' position of the phenyl ring A to form a six-membered ring and consequent hydrogen radical loss that would result in restoration of aromaticity to ring A. Analogous rearrangement process was also postulated to occur by Ronayne<sup>27</sup> and co-workers and by Van de Sande<sup>28</sup> and co-workers for chalcones and led to the formation of an oxonium ion. In fact, 3-benzoyl-2-phenylbenzofurans

7



the M+. ion of compound 5c (m/z 388)

can be considered as benzofuran-chalcones hybrids, where the C2-C3 double bond of the benzofuran nucleus fixes in the *cis* disposition the chalcones-type double bond (bold part of the ions in Scheme 3). It must be point out that the aromatic oxonium ion *k* could further rearrange to the structure  $k_1$ , as demonstrated to occur in chalcones and 3-flavene by Traldi and co-workers.<sup>29</sup> Unfortunately, unlike chalcones, CID experiments on the [M-H]<sup>+</sup> ion lead to the losses of CHO<sup>-</sup> and CO that agree with both structures *k* and *k*<sub>1</sub>.

# 3.4 | Elimination of hydroxyl radical: 3-aroyl-2-aylbenzofurans vs calchones

All compounds under investigation exhibit unusual and fairly small [M-17] peaks (Table 1). With the only exception of compounds 5a, 5c, and 5d, this behaviour is somewhat increased in intensity under MS/MS conditions (R.A. 2-30%; Figure 4). The deuterium labelling results using the simplest 3-benzoyl-benzofuran 5a indicate that a deuterium was involved in this process. As can be seen, in the mass spectrum, the deuterated analogue  $5ad_{10}$  (Figure 2B), ions at m/z281 shifted to m/z 290 reflecting the loss of OD<sup> $\cdot$ </sup> from the molecular ion. The most reasonable inference is that the oxygen involved in the OH<sup>-</sup> loss arises from the carbonyl group. In fact, compounds 5a, as well as compounds 5f and 5g, have no additional substituents, which could themselves produce [M-OH]<sup>+</sup> ion. To the best of our knowledge, only few papers have been reported on the involvement of a carbonyl oxygen atom in the formation of dehydroxylated ions. Bowie and White described the formation of [M-H<sub>2</sub>O]<sup>+</sup> and [M-OH] <sup>+</sup> ions in the mass spectra of aromatic carbonyl compounds, which contain an ortho (or peri) methoxy-substituent.<sup>30</sup> More recently. Zenchevich and Pushkareva reported that compounds bearing -CO-CH=CH-N (CH<sub>3</sub>)<sub>2</sub> structural fragments exhibit unusual peaks of [M-OH]<sup>+</sup> ions formed by a rearrangement with the migration of a hydrogen atom.<sup>31</sup>

To get insight into the mechanism of formation of the  $[M-OH]^+$ ion, further experiments were performed on the labelled compound **5ad**<sub>5</sub>. In the MS/MS mass spectrum of **5ad**<sub>5</sub>, the formation of ion at *m/z* 285 [M-OD]<sup>+</sup> clearly indicates that the hydrogen lost derives from ring A (Figure 5B).

One of the possible mechanisms rationalizing these results is reported in Scheme 4. The carbonyl undergoes loss of a hydroxyl radical after abstraction of the hydrogen from ring A by McLafferty type rearrangement<sup>32</sup>; the ionic species so generated is plausibly represented as the ion with the structure **y**, which contains a new five-membered ring.

In the case of *p*-nitro-derivative **5b**, some considerations have to be done. Compound **5b** has two nitro groups in the *para* positions of ring-A and B, respectively, which may be themselves involved in a M-OH process. The OH loss is commonly observed in nitro-aromatic derivatives, due to an *ortho* effect.<sup>33-37</sup> However, the proximity effect described in these studies occurs between aromatic nitro function and a group in its proximity, ie, in *ortho* or *peri* positions, and, therefore, appears unlikely for *p*-nitro-compound **5b** where the distances

between the NO<sub>2</sub> substituents and the phenyl rings are rather much greater.

WILEY

Interestingly, Baldas and Porter<sup>38</sup> reported that the spectra of 3and 4-nitrocalchones display the loss of hydroxyl radical and that this fragmentation is initiated by the transfer of a hydrogen atom from the 2-position of the phenyl ring to the nitro group via a cyclized intermediate.

Considering the striking analogy between chalcone and **5a** (*vide supra*), we expected, for p-NO<sub>2</sub> derivative **5b**, a similar mechanism to that reported for nitrochalcones,<sup>38</sup> that involves the nitro group rather than the carbonyl. A chalcone-like mechanism was therefore proposed for compound **5b** (Scheme 5).

This behaviour is in agreement with the collisional experiments performed on compound **5b** (Figure 6). Under MS<sup>3</sup> experiments, the [M-OH]<sup>+</sup> ion (*m*/*z* 371) loses NO<sup>-</sup> to form the *m*/*z* 341. MS<sup>4</sup> experiments reveal that the *m*/*z* 341 ion fragments by the loss of a nitro radical (*m*/*z* 295) and of NO<sup>-</sup> (*m*/*z* 311). However, the sequential loses of NO<sup>-</sup> and NO<sub>2</sub><sup>-</sup> from the [M-OH]<sup>+</sup> ion is also consistent with the mechanism that involves the carbonyl group (Scheme 4), which, therefore, cannot be ruled out for the nitro derivative **5b**.

# 4 | CONCLUSIONS

Electron ionization and collision-induced dissociation tandem mass spectrometry have proved to be highly effective for a complete structure assignment of 3-benzoyl-2-phenylbenzofuran, a side product formed in the preparation of 2-phenylbenzofurans under Wittig conditions. Our study showed that beside the alfa cleavages typical of carbonyl compounds, the unusual losses of H<sup>-</sup> and of OH<sup>-</sup> constitute diagnostic fragmentations. These losses were attributed to the interaction between the carbonyl of the 3-acyl group and the 2-phenyl ring, thus reflecting their proximity in the structure of the benzofuran nucleus. We also evidenced that 3-arovl-2-avlbenzofurans can be considered as benzofuran-calchones hybrids and that their El fragmentation routes have strong similarities compared with those of chalcones. However, some differences were also underlined. In particular, we found that the OH<sup>-</sup> loss from the molecular ion of 3-aroyl-2arylbenzofurans involves the carbonyl group in position 3. Only nitro-chalcones were reported to show the [M-OH]<sup>+</sup> ion, and it was suggested that the source of the OH<sup>-</sup> loss was exclusively the nitro group. In the case of the analogous 3-benzoyl-2-phenylbenzofuran derivatives bearing the NO<sub>2</sub> group, MS<sup>n</sup> experiments demonstrated that a nitro-chalcone-like mechanism cannot be excluded.

### ACKNOWLEDGMENTS

The present work was partially supported by Fondo integrativo per la ricerca (Fir)-Annualità 2018-University of Cagliari. The authors thank Professor Pierluigi Caboni from the Department of Life and Environmental Sciences, University of Cagliari, for the accurate mass measurements.

ISS FCTROMETR

### 

### ORCID

Michela Begala b https://orcid.org/0000-0002-8213-2335

### REFERENCES

- 1. Jiang Y, Gao B, Huang W, Liang Y, Huang G, Ma Y. Simple, convenient, and efficient synthesis of 2-arylsubstituted benzo[b]furans. *Synth Commun.* 2009;39:197-204.
- Delogu GL, Pintus F, Mayán L, et al. MAO inhibitory activity of bromo-2phenylbenzofurans: synthesis, in vitro study, and docking calculations. *Med Chem Commun.* 2017;8(9):1788-1796.
- Kumar A, Pintus F, Di Petrillo A, et al. Novel 2-pheynlbenzofuran derivatives as selective butyrylcholinesterase inhibitors for Alzheimer's disease. Sci Rep. 2018;8(1):4424-4436.
- Hercouet A, Le Corre M. Une nouvelle voie d'accès aux benzofurannes. Tetrahedron Lett. 1979;20(23):2145-2148.
- 5. Hercouet A, Le Corre M. Etude des ω acyloxybenzylidènetriphénylphosphoranes. Nouvelle voie d'accès aux benzofurannes. *Tetrahedron*. 1981;37(16):2867-2873.
- March RM, Todd JFJ eds. Practical Aspects of Ion Trap Mass Spectrometry, Volume 1. Boca Raton, FL: CRC Press; 1995.
- Begala M, Caboni PL, Matos MJ, Delogu GL. Unexpected one-step synthesis of 3-benzoyl-2-phenylbenzofurans under Wittig conditions. *Tetrahedron Lett.* 2018;59(18):1711-1714.
- Shen Z, Dornan P, Khan HA, Woo TK, Dong VM. Mechanistic insights into the rhodium catalyzed intramolecular ketone hydroacylation. J Am Chem Soc. 2009;131(3):1077-1091.
- Thévenin M, Thoret S, Grellier P, Dubois J. Synthesis of polysubstituted benzofuran derivatives as novel inhibitors of parasitic growth. *Bioorg Med Chem.* 2013;21(17):4885-4892.
- Mutai P, Pavadai E, Wiid I, Ngwane A, Baker B, Chibale K. Synthesis, antimycobacterial evaluation and pharmacophore modeling of analogues of the natural product formononetin. *Bioorg Med Chem Lett*. 2015;25(12):2510-2513.
- Felpin FX. Practical and efficient Suzuki-Miyaura cross-coupling of 2iodocycloenones with arylboronic acids catalyzed by recyclable Pd(0)/C. J Org Chem. 2005;70(21):8575-8578.
- Porter QN. In: Taylor EC, Weissberger A, eds. Mass Spectrometry of Heterocyclic Compounds. second ed. New York: Wiley-Interscience; 1985:177-180.
- Vebrel J, Roche M, Gore J. Comportement de quelques arylbenzo[b]furannes sous l'impact électronique. Org Mass Spectrom. 1977; 12(12):751-755.
- Begala M, Tocco G, Meli G, Podda G, Urru SAM. 2-Substituted benzofuran fragment ion formation in the electron ionization mass spectra of 6-alkyl- and 6-aryldibenzo(d,f)(1,3)dioxepine derivatives. 1. Spirocyclization of the molecular ions in the gas phase. *Rapid Commun Mass Spectrom*. 2007;21(8):1414-1420.
- Klasinc L, Stefanovic D, Adam S, Güsten H. The mechanism of CO loss in the electron impact-induced fragmentation and its methyl ether. Org Mass Spectrom. 1976;11(2):171-174.
- Singh SN, Fletcher RD, Fisher SG, et al. Amiodarone in patients with congestive heart failure and asymptomatic ventricular arrhythmia. Survival trial of antiarrhythmic therapy in congestive heart failure. N Engl J Med. 1995;333(2):77-82.
- Felder CC, Joyce KE, Briley EM, et al. LY320135, a novel cannabinoid cb1 receptor antagonist, unmasks coupling of the CB1 receptor to stimulation of cAMP accumulation. *Pharmacol Exp Ther.* 1998;284: 291-297.

- Heel RC, Brogden RN, Speight TM, Avery GS. Benzbromarone: a review of its pharmacological properties and therapeutic use in gout and hyperuricaemia. *Drugs.* 1977;14(5):349-366.
- 19. Brenner LM, Brush CK, United States Patent, 1997;4,001,426.
- Niessen MAW, Correa Ceballos RA. Interpretation of MS-MS mass spectra of drugs and pesticides. New York, United States: John Wiley & Sons, Inc; 2017. https://doi.org/10.1002/9781119294269.
- Kuhn J, Götting C, Kleesiek K. Simultaneous measurement of amiodarone and desethylamiodarone in human plasma and serum by stable isotope dilution liquid chromatography-tandem mass spectrometry assay. J Pharm Biomed Anal. 2010;51(1):210-216.
- De Vries JX, Walter-Sack I, Ittensohn A. Analysis of benzbromarone in human plasma and urine by high performance liquid chromatography and gas chromatography-mass spectrometry. J Chromatogr. 1987; 417:420-427.
- Dias HJ, Vieira TM, Crevelin EJ, Donate PM, Vessecchi R, Crotti AEM. Fragmentation of 2-aroylbenzofuran derivatives by electrospray ionization tandem mass spectrometry. J Mass Spectrom. 2017;52(12): 809-816.
- Moraes LAB, Sabino AA, Meurer EC, Eberlin MN. Absolute configuration assignment of ortho, meta, or para isomer by mass spectrometry. *J Mass Spectrom*. 2005;16:431-436.
- Itagaki Y, Kurokawa T, Sasaki S, Chang CT, Chen FC. The mass spectra of chalcones, flavones and isoflavones. *Bull Chem Soc Japan*. 1966; 39(3):538-543.
- Gronowska J, Paradisi C, Traldi P, Vettori U. A study of relevant parameters in collisional-activation of ions in the ion trap mass spectrometer. *Rapid Commun Mass Spectrom.* 1990;4(9):306-313.
- Ronayne J, Williams DH, Bowie JH. Studies in mass spectrometry. XIX. Evidence for the occurrence of aromatic substitution reactions upon Electron impact. J Am Chem Soc. 1966;88(21):4980-4984.
- Van De Sande C, Serum JW, Vandewalle M. Studies in organic mass spectrometry-XII. Mass spectra of chalcones and flavanones. The isomerisation of 2'-Hydroxy-Chalcone and flavanone. Org Mass Spectrom. 1972;6(12):1333-1346.
- Ardanaz CE, Traldi P, Vettori U, Kavka J, Guidugli F. The ion-trap mass spectrometer in ion structure studies. The case of [M-H]<sup>+</sup> ions from chalcone. *Rapid Commun Mass Spectrom*. 1991;5(1):5-10.
- Bowie JH, White PY. Electron impact studies. Part XXXIX. Proximity effects in the mass spectra of aromatic carbonyl compounds containing adjacent Methoxysubstituents. J Chem SOC(B). 1969;0:89-93.
- Zenkevich IG, Pushkareva TI. New possibilities of dimethylformamide dimethylacetal as a derivatization agent for gas chromatography/mass spectrometry analysis. J Anal Chem. 2016;71(14):1341-1351.
- McLafferty FW. Mass spectrometric analysis. Molecular rearrangements. Anal Chem. 1959;3(1):82-87.
- Beynon JH, Saundersa RA, Topham A, William AE. The dissociation of o-nitrotoluene under Electron impact. J Chem Soc. 1965;0:6403-6405.
- Butcher AR, Thomas CB. The loss of a hydroxyl group from the molecular ions of alkylnitrobenzenes. Org Mass Spectrom. 1979;14(8):448-454.
- McLuckey SA, Glish GL. The effect of charge on hydroxyl loss from ortho-substituted nitrobenzene ions. Org Mass Spectrom. 1987;22(4): 224-228.
- Danikiewicz W. Formation of benzimidazole derivatives during electron ionization induced fragmentation and pyrolysis of N-benzyl-onitroaniline. *Rapid Commun Mass Spectrom.* 1998;12(11):689-694.
- Srzic D. Ortho effect of the nitro group in the fragmentation of 1-phenylnitrophenyl-2-pyrazolines. Org Mass Spectrom. 1986;21(7):411-413.

 Baldas J, Porter QN. Mass spectrometric studies. XIV. Nitrochalcones. Aust J Chem. 1979;32(10):2249-2256.

# SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.

**How to cite this article:** Begala M, Delogu GL. Unexpected detection of 3-aroylbenzofuran side products in the preparation of 2-arylbenzofurans: Identification, characterization, and comparison with chalcone's fragmentation patterns using EI/MS<sup>n</sup>. *J Mass Spectrom*. 2019;1–11. <u>https://doi.org/10.1002/jms.4425</u>