Organic & Biomolecular Chemistry



PAPER

View Article Online
View Journal | View Issue



Cite this: *Org. Biomol. Chem.*, 2024, **22**, 2264

Δ -Keto-acid/hydroxy-lactone isomerization in some lichen depsides, depsidones and diphenyl ethers†

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Received 12th December 2023, Accepted 6th February 2024 DOI: 10.1039/d3ob02026f

rsc.li/obc

In some compounds in lichens, the carboxylic acid is *ortho*-substituted by an 2-oxoalkyl chain. This particular structure induces the existence of δ -keto-acid **ka** or hydroxy-lactone **hl** isomers, clearly identified by their NMR data and chemical properties, such as dehydration, methylation and behaviour in thermal conditions. Internal hydrogen bonding between the carboxylic acid and substituent in the *ortho'* position is proposed as an isomerization modulator: an H-bond acceptor (OCH₃) leads to **ka** isomers, whereas **hl** isomers are obtained with an H-bond donor (OH).

Introduction

Depsides, depsidones and diphenyl ethers are naturally occurring polyfunctionalized aromatic compounds produced by lichens^{1,2} or, more rarely, by fungi.³ In some depsides, the C-6 position of the B ring (more commonly referred to C-6') is substituted by a 2-oxoalkyl chain, as shown in Fig. 1 for seven specialized metabolites selected for this study. This unusual chain was introduced as a linear acyl-CoA precursor involved in the polyenzymatic system of polyketide synthase (PKS).^{4,5} The presence of a free carboxylic acid on C-1', in the *ortho* position to the 2-oxoalkyl chain induces the δ -keto-acid (ka)/hydroxy-lactone (hl) isomerization (Scheme 1). The factors influencing the position of this equilibrium are not well understood although this phenomenon, wrongly termed "tautomerism", has been reported.⁶⁻¹¹ Hence, the occurrence of ka and/or hl isomers should be explored further.

In the present study, new results are presented allowing the establishment of the structure and stability of compounds depicted in Fig. 1. Thus, the first observation of the hydroxylactone isomers (hl) of 2'-O-methylmicrophyllinic 2 and of 2'-O-methylsuperphyllinic 3 acids, as well as the obtention of α -collatollic 5 and α -alectoronic 6 acids as δ -keto-acid open form (ka), is reported, and their 1 H NMR data analyzed. Then,

Results

The structures of compounds 1-4 ⁶ and 5-7 ⁷⁻¹¹ (Fig. 1), isolated after acetone extraction and purification by column

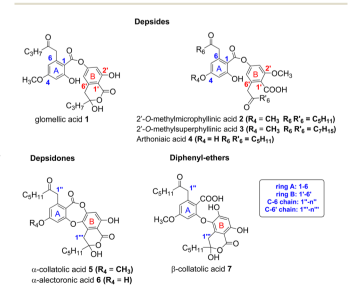


Fig. 1 Structure of depsides, depsidones and diphenyl-ethers bearing a 2-oxoalkyl chain considered in this study. Other metabolites are listed in $S1.\dot{\tau}$

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†Electronic supplementary information (ESI) available. CCDC 2241570. For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.org/10.1039/d3ob02026f

to validate **hl** and **ka** structures, the synthesis of some new derivatives is described. Finally, the hydrogen bonding between the carboxylic acid and vicinal $O-R_{2'}$ ($R_{2'} = H$ or CH_3) is proposed as the main isomerization modulator.

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Scheme 1 Δ -Keto-acid/hydroxy-lactone (ka/hl) isomerism.

chromatography and/or preparative thin-layer chromatography, were confirmed by NMR analyses and matched values stated in the literature.

Hydroxylactones isomers 2hl and 3hl

The isomers **2hl** and **3hl** were observed only in their ¹H NMR spectra in a 15/85% mixture with their corresponding keto-acid isomers **2ka** and **3ka**, whereas **4hl** has never been observed. The occurrence of these **hl** forms could be assessed by the downfield shifting of H-1" and H-3" signals in relation to the transformation of the ketone on C-2" into a hemi-acetal. The C-2" chirality resulted in an AB system for the diastereotopic CH₂-1", whereas these protons were enantiotopic in the **ka** form, leading to a singlet. As an example, the spectrum of **3ka** + **3hl** mixture is shown on Fig. 2.

On the spectra of the 3"'-O-methyl derivatives of 1, 5 and 6 (vide infra), an AB system was observed for CH₂-1"' and CH₂-3"'.

Keto-acid isomers 5ka and 6ka

If pure depsidones **5hl** and **6hl** were dissolved in Culberson's solvent G (toluene, EtOAc, HCOOH: 70/40/4),¹ they were obtained as δ-keto-acid isomers **5ka** and **6ka** after evaporation of the solvent under reduced pressure. Likewise, only keto-acid forms could be observed when **5** and **6** were purified from acetone extracts of lichens using the same solvent system (respectively from *Tephrolema atra* and *Ochrolecchia parella*). For example, an enlarged version of the 2–4 ppm range of the ¹H NMR spectra of **5hl** and **5ka** is depicted in Fig. 3. The recovery of **5ka** and **6ka** after G solvent evaporation was observed when (CD₃)₂CO was used as an NMR solvent. Interestingly, in the case of depside **1**, open isomers **1ka** have never been

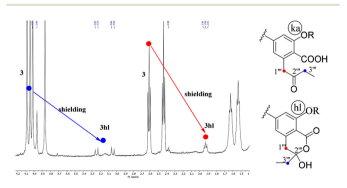


Fig. 2 Expansion of the 1 H NMR spectrum (2–4 ppm, 500 MHz, CDCl₃) of the mixture 3ka + 3hl.

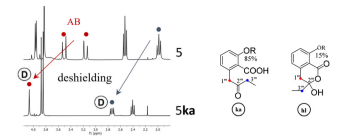


Fig. 3 Expansion of the 1H NMR data (300 MHz) of 5 (5hl) and 5ka (CDCl $_3$). 9: partial H/D exchange.

obtained. The open structures **5ka** and **6ka** were clearly attested in ¹H NMR by the signal deshielding of 2H-1" and 2H-3" induced by the carbonyl in position 2".

Finally, the hybridization change of C-1" resulting in significant differences in ¹H NMR data allowed the unequivocal attribution of hl or ka isomers, as illustrated for CH2-1" and CH₂-3", in Table 1. To explain the coalescence (noted ^a in Table 1) occurring in the spectra of hl isomers, a "tautomerism equilibrium" has been previously proposed. Nevertheless, this phenomenon is the result of inversion of the hydroxy-lactone ring. This explanation was clearly attested with 67, 78 and 1 (see S4†) by the transformation at low temperature of the coalescent CH2-1" signal into an AB system. The spectra of the ka isomers of 5 and 6 were recorded in d6-acetone, and the disappearance and/or intensity reduction of CH2-1" and CH2-3" signals were observed (Fig. 3). A keto-enol tautomerism, which induced a H/D exchange (noted @ in Table 1 and Fig. 3) and discussed below, could be proposed (see S4†). On ¹³C NMR spectra, ring inversion and tautomerism resulted in the disappearance of signals but in some cases C-2" was observed near 206 ppm (ka) or 105 ppm (hl).

Reactivity in relation to the hydroxy-lactone structure (see S6†)

Most reactions were performed with collatolic acids 5 and 7 taken as examples, but new derivatives of α -alectoronic acid 6 and of glomellic acid 1 were also prepared.

Dehydration and O-methylation of the hydroxy-lactone ring

First, dehydration and *O*-methylation of the hydroxy-lactone isomers of **1**, **5** and **6** were performed to confirm the stability of the six-membered ring.

- In a study dedicated to the lichenochemical investigation of *Tephromela atra*, ⁷ the presence of the α -deoxycollatolic acid (= α -collatone) was reported. In the present study, such water removal from acids 5 and 7 was achieved by heating these compounds in a CHCl₃ or CDCl₃/CF₃COOH 80/20 mixture at 55 °C, thereby yielding α -collatone 8 and β -collatone 9. The latter was obtained as crystals suitable for X-ray analysis (Fig. 4) clearly disclosing the hydrogen bonding between OH-2' and C=O-1'.
- Using the *O*-methylation method previously undertaken, 9 in the structural determination of α -alecteronic acid $\mathbf{6}$

Table 1 Selected ¹H NMR data of hl or ka isomers (detailed spectra in S2 and S3†)

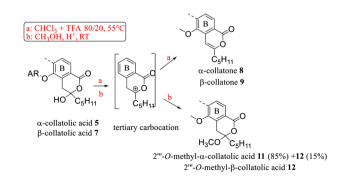
CH_2	1	2hl	3hl	5hl	6hl	7hl
1′′′	$3.07-3.25$ AB^a	3.03-3.22 AB, <i>J</i> = 16	3.04-3.24 AB, <i>J</i> = 16	3.18-3.52 AB, <i>J</i> = 16.2	3.55 (s ^a)	$3.06-3.20$ $AB^a, J = 16$
3‴	1.98 (t ^a)	1.91 $(t, J = 8)$	1.93 $(t, J = 8.2)$	2 (t, J = 8.2)	1.99 (t ^a)	1.96 $(t, J = 8)$
$\overline{\text{CH}_2}$	2ka	3ka		4	5ka ®	6ka
1'''	4.00 (s)	4.02 (s))	3.85 (s)	4.04 (s)	4.28 (s ^a)
3‴	2.42	2.44		2.51	2.36	2.44
	t, J = 7.5	t, J = 7.	5	t, J = 7.5	s^a	$t^a, J = 7.5$

 $\delta_{\rm H}$ in ppm, J in Hz. ^a Coalescent. ©: partial H/D exchange.

Fig. 4 ORTEP diagram of β-collatone 9 (drawn with ellipsoids at 50% probability). H-bonding site is emphasized in the dashed blue circle.

(CH₃OH, H⁺ catalytic at room temperature), 2"'-O-methyl derivatives 10, 11 and 12 from 1, 5 and 7 (Schemes 2 and 3) could be obtained, respectively, in fair yields using CF₃COOH as a catalyst. As previously reported, diastereotopic CH2 appeared as an AB system. Interestingly, the same tertiary carbocation intermediate could be postulated in the dehydration and nucleophilic attack of CH3OH.

In these conditions (CH₃OH, H⁺), chemical evolution occurred with 10 and 11. Regarding the specific example of 10, the rapid cleavage of the ester bond led to glomellin⁶ 13 and to the analogous structure of penidiolactol¹² 13' (Scheme 3). In the case of 11, analogous transesterification partially led to 12 (Schemes 2 and 5).



Scheme 2 Reactivity of the hydroxy-lactone ring of 5 and 7.

Scheme 3 O-Methylation of 1 and degradation of 10.

The methyl ester of 2 is the main metabolite of Porpidia contraponenda,6 but those of 1, 5 and 7 were never observed under our conditions. Moreover, the synthesis of methyl α-alectoronate 14 has been reported if diazomethane is used with 6. The authors suggested that the methylation occurred on the carboxylic acid of 6ka after hydroxy-lactone ring opening. However, another mechanistic sequence in accordance with the hl structure (Scheme 4) can be envisaged in which the driving force would be the OH-lactone deprotonation of 6hl leading to CH3+, which could methylate the alectoronate.

Thermal isomerization of hydroxy-lactones: examples of α-collatolic acid 5 and physodic acid

The behavior in thermal conditions of α -collatolic acid 5 and physodic acid were compared. The previously observed⁷ isomerization of α-collatolic acid 5 into corresponding β-isomer 7 was confirmed herein by heating in refluxing toluene or d8toluene (Scheme 5). With respect to the degradation of 10, involvement of an enol intermediate, assisted by the eventual acidic catalysis of free COOH-1', can be postulated. In contrast,

Scheme 4 Synthesis of methyl α -alectoronate 14 using CH₂N₂. Mechanistic proposition.

Scheme 5 Thermal isomerization of 5 and decarboxylation of physodic acid

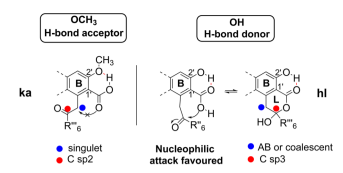
physodic acid was isomerized and decarboxylated to give physodone,^{‡ 15} in same thermal conditions. In this case, acid deprotonation, followed by a electrophilic substitution, has been proposed according to the literature. 13 Then, this decarboxylation absence with 5 and 6 is another strong argument in favor of the cyclic structure (Scheme 5 and S5†). Mixtures of α (depsidones) and β isomers (diphenyl-ethers) of 5 and 6 are also signaled in acetonic extracts of lichens (for example, in *Xanthoparmelia glabrans*¹⁴)

Reactivity in relation of the δ -keto-acid structure

¹H NMR spectra, recorded in d6-acetone, revealed a property in accordance with the open ka structure of 2. Thus, disappearance of CH2-1" and CH2-3" signals was observed (see S4†). This phenomenon, also attested in ²H NMR, can be explained by their deuteriation involving an enol. The NMR solvent would provide deuterium atoms and tautomerism would be catalysed by the close carboxylic acid. Thus, H/D exchange never occurred on the C-1" and C-3" of the A ring bearing an ester on C-1. The less easy enolization of CH₂-3" was slower and incomplete. Compounds 5ka and 6ka presented such an H/D exchange.

Discussion

The results concerning the preparation and characterization of 2hl, 3hl, 5ka and 6ka, as well as those devoted to the stability of the hl and ka isomers, must be explained. The occurrence of a hydroxy group, acting as an H-bond donor (1, 5-7), is in relation to the hydroxy-lactone ring hl, whereas a methoxy substituent, acting as an H-bond acceptor (2-4), is instead associated with δ-keto-acid ka isomers, as clearly attested by NMR data (Scheme 2). Then, the stabilization energy due to the hydrogen bonding (Scheme 6) between O-R2' and COOH-1' could be the main reason accounting for the isomerization control. If $R_{2'}$ = H (1, 5–7), then the hydrogen bonding between OH-2' and COOH-1' promoted the nucleophilic attack of the



Scheme 6 Influence of hydrogen bonding on isomerization.

hydroxy group of the carboxyl on the ketone to give hydroxylactone. In contrast, with $R_{2'} = CH_3$ (2-4), this hydroxy group is bonded with the methoxy group, and δ-keto-acid remains untransformed. Such hydrogen bonding has been widely discussed12,15 and attested through the X-ray structure of 5-methylmellein¹⁶ and β -collatone 9 (Fig. 4).

To explain obtaining 5ka and 6ka during G solvent evaporation, one can propose that, in this solvent, HCOOH would induce breaking of H-bonds, whereas EtOAc would act as an H-bond acceptor. Finally, in depsidones 5 and 6, the ether bridge would induce steric hinderance, disfavouring COOH-1' attack on the ketone and preventing 5hl and 6hl obtention. On the contrary, the ether absence in depside 1hl could explain why 1ka has never been obtained.

Conclusions

In conclusion, it appears that the structure of specialized metabolites bearing a δ-keto-acid moiety on ring B is the result of isomerization controlled by internal hydrogen bonding. This latter involves the acid function and its ortho OH or OCH₃ substituent. If a methoxy is present in C-2', they should be described as keto-acids (ka). If C-2' is bearing an OH, they should be termed hydroxy-lactones hl according to their reactivity. Thus, in the latter case, we suggest to delete the frequent mention of tautomerism. To standardize the nomenclature of these compounds according to their reactivity, we propose to name unusual isomers using the prefix "iso" as follows: 5ka = α -isocollatolic acid, **6ka** = α -isoalectoronic acid, **2hl** = 2'-Omethylisomicrophyllinic acid, and 3hl = 2'-O-methylisosuperphyllinic acid. Finally, this study is another demonstration⁶ of the interest of the extraction of natural products which allows original structural and/or mechanistic chemical studies.

Experimental

Reagents

All reagents were of high quality and purchased from commercial suppliers. They were used as received without further purification or purified/dried according to standard procedures.

[‡] Personal communication. See the NMR spectra in S5.†

Preparation of extracts and samples

Lichen thalli (often with minerals) were ground and extracted thrice with warm acetone. After drying, the solvent was evaporated under reduced pressure. Isolation of 1-4 has been previously reported. Depsidone 5 was obtained from Tephromela atra (Huds.) Hafellner, whereas depsidone 6 from Ocholechia parella (L.) Massal using the same procedure described previously.^{7,8} Both were collected on coastal rocks at Lancieux (France, department 22). To perform the methylation of 2, Porpidia cinereoatra was collected in Néant-sur-Yvel (France, department 56).

TLC analysis

Secondary metabolites were identified by thin-layer chromatography (TLC) using standards. Extracts and standards were loaded onto an aluminium sheet (10 × 20 cm) coated by silica gel 60F254 (Merck). Thereafter, plates were developed with solvent C (toluene/acetic acid: 85/15) or solvent G (toluene/ ethyl acetate/formic acid: 139/83/8). All TLC plates were observed under UV light at 254 and 365 nm (12 W, 6 W at 365 nm, 6 W at 254 nm; Fischer Bioblock Scientific). TLC examples are given in ESI.†

Experimental $R_{\rm f}$ values in solvent C: 1 (0.30), 5 (0.35), 6 (0.16), 7 (0.22), 8 (0.40), 9 (0.51), 10 (0.51), 13 (0.55), 13' (0.32), 11 (0.49), and 12 (0.39) (see TLC on ESI†).

Purifications using liquid chromatography

Silica gel 60 (Merck) and mainly solvent system C¹ were used for the purification of different extracts. Combined fractions with selected metabolites were washed once with water to eliminate acids, dried over MgSO4 and evaporated under reduced pressure. Preparative TLC was also performed using the same

Tephromela atra: 19.4 g; extract: 1.35 g; 5 after chromatography using solvent C or G: 220 mg recrystallization. In toluol: 160 mg.

Ochrolecchia parella: 8.3 g; extract: 390 mg; 6 after chromatography in solvent C or G: 45 mg.

Porpidia cinereoatra: extract: 320 mg; after chromatography in solvent C. 2: 34 mg and confluentic acid: 160 mg.

High resolution mass spectrometry (HRMS) and tandem mass spectrometry (MS-MS)

The mass spectra of standards were recorded CRMPO-ScanMat (Rennes, France) on an Orbitrap Thermo Fisher Scientific Q-Exactive instrument with an ESI (electrospray Ionization) source in negative mode by direct introduction at 5-10 µg mL⁻¹. Samples were prepared in methanol at 10 $\mu g \, mL^{-1}$.

Synthesis procedures

Reactions were performed on a small amount of extraction products; hence, yields are not given. Only the quantities obtained after working up are indicated.

Isomerization of 5 and 6. The open form 5ka and 6ka were first obtained during chromatographic purifications performed using G solvent (twice for 5ka and once for 6ka). After evaporation of the solvent under reduced pressure, the residue was dissolved in d6-acetone and obtained NMR spectra corresponded to 5ka and 6ka. If CDCl₃ was used, the obtained NMR spectra corresponded to 5 and 6. The same result was obtained if pure 5 and 6 were dissolved in G solvent.

Isomerization of 2 and 3. The cyclic form 2hl and 3hl were observed as traces on NMR spectra recorded in CDCl3 of 2 and 3 obtained after chromatographic purification using C solvent.

Dehydration of 5 and 7. The loss of water was performed in a mixture of CHCl₃/CF₃COOH (2 mL/0.5 mL) at 50 °C. The reactions were monitored by TLC: after 5-7 days (no more conversion), the reaction was stopped. After water addition, the organic layer was separated, dried over MgSO4 and evaporated under reduced pressure. The residue was purified by column chromatography using solvent C. NMR spectra of the transformation of 5 and 7 in CDCl3/CF3COOH are shown in ESI.†

5 (20 mg) gave α -collatone = deoxycollatolic acid 8 (11 mg) with similar data to those previously reported.7 Presence of residues of 5 and of 10 was noted on TLC.

7 (20 mg) gave β -collatone 9 (14 mg).

O-Methylations of 1, 5 and 7. The methylation of OH-3" was performed in CH₃OH (2-3 mL) at room temperature with a catalytic amount of CF₃COOH. The reactions were monitored by TLC: for high conversion, 5 days were needed. Every day, one drop of CF₃COOH was added to enhance the reaction. Methanol was evaporated under reduced pressure, and the residue was purified by column chromatography using solvent C.

1 (15 mg) gave 2"'-O-methylglomellic acid 10 (12 mg). After 1 month, along with 10 (4 mg), hydrolysis compounds glomelline 13 (3 mg) and 13' (3 mg) were obtained.

7 (30 mg) gave 2"'-O-methyl-β-collatolic acid 12 (24 mg) and 5 gave 2'''-O-methyl-α-collatolic acid 11 (14 mg) as a white precipitate after filtration. Isomerization of 11 into 12 was also observed by TLC.

Thermal isomerization of 5. The depsidone 5 was allowed to react in refluxing toluene for 1 week. After solvent evaporation under reduced pressure, the residue was purified by column chromatography using solvent C. Yield of 7: 40-50%. Evolution in d8-toluene at 80 °C is shown in ESI.†

X-ray data

β-Collatone 9 CCDC 2241570.†

¹H, ²H and ¹³C NMR spectra

Spectra were recorded at 300/75 MHz or 500/100 MHz (cryoprobe) respectively, using CDCl₃ or d_{δ} -acetone. Shifts (δ values) are given in parts per million (ppm) and coupling constants (J values) are given in Hertz (Hz). The multiplicity of signals was reported as follows: s (singlet), d (doublet), t (triplet), q (quadruplet), quint (quintet), sext (sextet), m (multiplet), * (broad or coalescent signal), dt (doublet of triplet), and td (triplet of doublet). The references were: $CDCl_3/7.26$ ppm and d_{6}

acetone/2.05 ppm. In some compounds, coalescent signals were missing or badly resolved in $^{13}{\rm C}$ NMR.

2hl and 3hl in mixture with 2ka and 3ka (15/85%). Only some signals were visible.

2'-O-Methylisomicrophyllinic acid **2hl** $C_{30}H_{38}O_9$ M_W : 542.62 g mol⁻¹. 3-Hydroxy-8-methoxy-3-pentyl-3*H*-2-benzopyran-1(4*H*)-on-6-yl and 2-hydroxy-4-methoxy-6-(2-oxoheptyl) benzoate. ¹**H NMR** (300 MHz, CDCl₃): 3.22 and 3.03 (AB, J = 16 Hz, 2H), 1.89 (t, 2H). In mixture with **2ka** (15/85%).

2'-O-Methylsuperphyllinic acid **3hl** $C_{34}H_{46}O_9$ M_W : 598.72 g mol⁻¹. 3-Heptyl-3-hydroxy-8-methoxy-3-3H-2-benzopyran-1(4H)-on-6-yl and 2-hydroxy-4-methoxy-6-(2-oxononyl)benzoate. ¹**H NMR** (500 MHz, CDCl₃): 3.24 and 3.04 (AB, J = 16 Hz, 2H), 1.93 (m, 2H). ¹³**C NMR** (125 MHz, CDCl₃): 41.1 and 37.8. In mixture with **3ka** (15/85%).

In ¹³C NMR of **5ka** and **6ka**, some signals of sp² carbons were not visible, particularly C'''-1 and C'''-2.

α-Isocollatolic acid **5ka**: [522-52-1] $C_{28}H_{32}O_9$ $C_{29}H_{34}O_9$ M_W : 526.58 g mol⁻¹. 8-Hydroxy-3-methoxy-11-oxo-1,6-bis(2-oxoheptyl)-11H-dibenzo[b,e][1,4]dioxepin-7-carboxylic acid. ¹H NMR (300 MHz, CD_3COCD_3): 6.87 (d, J = 2.4 Hz, 1H), 6.78 (d, J = 2.4 Hz, 1H), 6.70 (s, 1H), 4.08 (s, 2H), 4.04 (s*, 2H), 3.88 (s, 3H), 2.55 (t, J = 7.5 Hz, 2H), 2.36 (s*, 1.6H), 1.59 (m, 4H), 1.32 (m, 8H), and 0.89 (t, 6H). ¹³C NMR (100 MHz, CD_3COCD_3): 163.5, 162.2, 162.1, 141.4, 141.0, 115.8, 114.2, 106.9, 104.2, 55.5, 54.6, 47.2, 42.0, 41.1, 31.5, 31.2, 23.1, 22.3, 22.2, 13.4, and 13.4.

α-Isoalectoronic acid 6ka: [668-80-4] $C_{28}H_{32}O_9$ 55 g mol⁻¹. M_W : 512.55 g mol⁻¹. 3,8-Dihydroxy-11-oxo-1,6-bis(2-oxoheptyl)-11H-dibenzo[b,e][1,4]dioxepin-7-carboxylic acid. ¹H NMR (300 MHz, CD₃COCD₃): 8.16 (s, 0.4H), 6.75 (d, J = 2.4 Hz, 1H), 6.63 (d, J = 2.4 Hz, 1H), 6.00 (s, 1H), 4.00 (s, 2H), 4.27 (s*, 2H), 3.97 (s, 2H), 2.52 (t, J = 7.5 Hz, 2H), 2.44 (t*, 1.8H), 1.55 (m, 4H), 1.29 (m, 8H), 0.88 (t, J = 7.4 Hz, 3H), and 0.82 (t, J = 7.4 Hz, 3H). ¹³C NMR (100 MHz, CD₃COCD₃): 162.6, 162.5, 162.1, 141.7, 141.1, 117.0, 112.9, 106.7, 105.8, 54.6, 47.3, 41.9, 41.5, 31.4, 31.2, 23.1, 22.3, 22.3, 13.4, and 13.4.

β-Collatone 9 [520-08-1] ¹**H NMR** (300 MHz, CDCl₃): 11.04 (s, 1H), 9.21 (s, 1H), 6.57 (s, 1H), 6.51 (d, J = 2.3 Hz, 1H), 6.37 (d, J = 2.3 Hz, 1H), 6.36 (s, 1H), 6.24 (s, 1H), 3.80 (s, 3H), 2.53 (t, J = 7.5 Hz, 2H), 2.51 (t, J = 7.5 Hz, 2H), 1.70 (m, 4H), 1.37 (m, 4H), 1.33 (m, 4H), 0.93 (t, J = 6.9 Hz, 3H), and 0.91 (t, J = 7.2 Hz, 3H). ¹³C **NMR** (100 MHz, CDCl₃): 166.1, 165.9, 162.8, 161.7, 161.1, 159.8, 159.2, 157.8, 142.2, 132.0, 130.2, 104.6, 104.3, 103.5, 103.3, 102.9, 98.6, 98.4, 56.0, 33.7, 33.4, 31.3, 31.2, 26.6, 22.5, 22.5, 14.1, and 14.0. **HRMS** [M - H] C₂₉H₃₁O₈ calc. 507.20244 found. 507.2024.

2"'-**O-Methylglomellic acid 10** $C_{26}H_{30}O_{9}$ M_{W} : 486.51 g mol⁻¹. (8-Hydroxy-3-methoxy-3-propyl-3H-2-benzopyran-1(4H)-on-6-yl and 2-hydroxy-4-methoxy-6-(2-oxopentyl)benzoate). ¹**H NMR** (300 MHz, CDCl₃): 11.21 (s, 1H), 11.19 (s, 1H), 6.62 (d, J = 2.4 Hz, 1H), 6.46 (d, J = 2.4 Hz, 1H), 6.45 (d, J = 2.4 Hz, 1H), 6.3 (d, J = 2.4 Hz, 1H), 4.03 (s, 2H), 3.84 (s, 3H), 3.18 (m, J_{AB} = 16 Hz, 1H), 3.08 (m, J_{AB} = 16 Hz, 1H), 2.39 (m, 2H), 2.02 and 1.98 (2 m, 2H), 1.54 (m, 6H), 1.43 (m, 2H), 1.39 (m, 4H), 1.02 (t, 3H), and 1.02 (t, 3H). ¹³C **NMR** (100 MHz, CDCl₃): 207.2, 168.8, 168.6, 166.8, 165.2, 163.6, 160.0, 155.4, 139.9, 139.0, 113.6,

112.9, 109.5, 107.6, 106.6, 104.3, 100.2, 51.4, 50.1, 44.5, 37.3, 36.3, 17.2, 17.0, 14.2, and 13.8. **HRMS** $[M - H]^ C_{26}H_{29}O_9$ calc. 485.18171 found. 485.1822.

Compound 13' $C_{13}H_{16}O_5$ M_W : 252.27 g mol⁻¹. 3-Pentyl-3,6,8-trihydroxy-3H-2-benzopyrane-1(4H)-one ¹H NMR (300 MHz, CDCl₃): 11.17 (s, 1H), 6.29 (d, J = 2.4 Hz, 1H), 6.20 (d, J = 2.4 Hz, 1H), 5.55 (s,1H), 3.35 (s, 3H), 3.11 (m, J_{AB} = 16 Hz, 1H), 3.00 (m, J_{AB} = 16 Hz, 1H), 1.44 (m, 6H), 1.44 (t, 3H), and 1.01 (t, 3H). ¹³C NMR (75 MHz, CDCl₃): 168.8, 164.4, 162.5, 140.4, 107.4, 107.1, 101.9, 101.8, 50.1, 37.3, 36.3, 17.0, and 14.2. HRMS [M - H]⁻ $C_{13}H_{15}O_5$ calc. 251.0925 found. 251.0924.

2"'-*O*-Methyl-α-collatolic acid 11 $C_{30}H_{36}O_9$ M_W : 540.62 g mol⁻¹. (2,11-Dimethoxy-5-hydroxy-9-(2-oxoheptyl)-2-pentyl-4H,8H-[1,4]benzodioxepino[2,3-f][2]benzopyran-4(1H),8-dione). ¹H NMR (300 MHz, CDCl₃): 11.01 (s, 1H), 6.79 (s, 1H), 6.58 and 6.59 (m, 2H), 3.99 and 3.97 (AB, J_{AB} = 17 Hz, 2H), 3.84 (s, 2H), 3.53 and 3.00 (AB, J_{AB} = 17 Hz, 2H), 3.34 (s, 3H), 2.54 (t, J = 7 Hz, 2H), 2.07–1.93 (mAB, 2H), 1.7–1.2 (m, 12H), 0.94, (t, J = 7 Hz, 3H), and 0.89 (t, J = 7 Hz, 3H). ¹³C NMR (75 MHz, CDCl₃): 168.1, 163.4, 161.7, 160.3, 150.5, 141.4, 129.7, 115.0 107.9, 107.5, 104.8, 55.8, 50.0, 48.0, 42.8, 35.0, 31.7, 31.3, 30.4, 23.3, 23.2, 22.5, 22.5, 14.0, and 14.0. HRMS [M – H]⁻ $C_{30}H_{35}O_9$ calc. 539.22866 found. 539.2279.

2"'-O-Methyl-β-collatolic acid 12 $C_{30}H_{36}O_9$ M_W : 540.62 g mol⁻¹. (6,8-dihydroxy-3-methoxy-5-(6-methoxy-1-oxo-3-pentyl-1H-2-benzopyran-8-yl)oxy-3,6,8-trihydroxy-3-pentyl-1H-2-benzopyran-1(1H)-one). ¹H NMR (500 MHz, CDCl₃): 11.12 (s, 1H), 6.58 (s*, 1H), 6.51 (d, J = 2.4 Hz), 6.50 (s, 1H), 6.21 (s, 1H), 3.79 (s, 3H), 3.35 (s, 3H), 3.24 (d, J_{AB} = 16 Hz, 1H), 2.99 (d, J_{AB} = 16 Hz, 1H), 2.51 (t, J = 7.4, 2H), 1.96 (m, 2H), 1.70 (m, 2H), 1.34 (m, 12H), 0.91 (t, J = 7.5, 3H), and 0.88 (t, J = 7.5, 3H). ¹³C NMR (75 MHz, CDCl₃): 168.3, 166.0, 162.6, 162.2, 160.6, 159.4, 157.0, 141.7, 134.3, 131.2, 107.0, 103.8, 103.4, 103.2, 102.8, 99.9, 35.0, 33.3, 31.7, 31.5, 31.2, 29.7, 26.5, 23.1, 22.4, 22.4, and 14.0 HRMS [M – H] $C_{30}H_{35}O_9$ calc. 539.22866 found. 539.2287.

Data concerning C-1"2"3" (Fig. 3)

Glomellic acid 1 [52589-14-7] **Proposed DCI**: 3,8-dihydroxy-3-propyl-3*H*-2-benzopyran-1(4*H*)-on-6-yl, and 2-hydroxy-4-methoxy-6-(2-oxopentyl)benzoate. 1 **H NMR** (300 MHz, CDCl₃): 4.04 (s, 2H), 3.22*(s, 2H), 2.39 (t, J = 7.5 Hz, 2H), 1.98* (m, 2H). 13 **C NMR** (80 MHz, CD₃COCD₃): 106.5, 43.5, and 37.2. 6

2'-O-Methylmicrophyllinic acid 2 [79579-62-7] ¹**H NMR** (300 MHz, CDCl₃): 4.07 (s, 2H), 4.03 (s, 2H), 2.59 (t, J = 7.5 Hz, 2H), and 2.42 (t, J = 7.5 Hz, 2H). ¹³C **NMR** (80 MHz, CDCl₃): 51.2 and 42.7. ⁶

2'-O-Methylsuperphyllinic acid 3 [108529-20-0] 1 H NMR (300 MHz, CDCl₃): 4.06 (s, 2H), 4.02 (s, 2H), 2.59 (t, J = 7.5 Hz, 2H), and 2.41 (m, 2H). 13 C NMR (100 MHz, CDCl₃): 207.3, 49.2, and 42.7. 6

Arthoniaic acid 4 [25556-24-5] ¹**H NMR** (300 MHz, CDCl₃): 4.17 (s, 2H), 3.85 (s, 2H), 2.51 (t, J = 7.5 Hz, 2H), and 2.48 (t, J = 7.5 Hz, 2H). ¹³**C NMR** (80 MHz, CDCl₃): 207.4, 47.7, and 42.5. ⁶

Author contributions

Project design by P. Uriac. Synthesis by P. Uriac. Characterization by S. Ferron and P. Jehan. Crystallographic experiments and analyses by T. Roisnel. The manuscript was written by P. Uriac and S. Tomasi with edits from all authors.

Conflicts of interest

There are no conflicts of interest to declare.

Acknowledgements

Dr Pierre Le Pogam (BIoCIS, Paris Saclay), Dr Elsa Catyan for ²H NMR and Dr Arnaud Bondon (CorInt, ISCR) for the recording of 500 MHz NMR spectra and fruitful discussions. Part of this work has been performed using the PRISM core facility (Biogenouest, Univ Rennes, Univ Angers, INRAE, CNRS, France).

References

- S. Huneck and I. Yoshimura, *Identification of Lichen Substances*, Springer Berlin Heidelberg, Berlin, Heidelberg, 1996.
- 2 M. J. Calcott, D. F. Ackerley, A. Knight, R. A. Keyzers and J. G. Owen, *Chem. Soc. Rev.*, 2018, 47, 1730–1760.

- 3 Y. Liang, Y. Zheng, Y. Shen, Q. Li, Y. Lu, S. Ye, X.-N. Li, D. Li, C. Chen, H. Zhu and Y. Zhang, *Tetrahedron Lett.*, 2022, 105, 154046.
- 4 D. Armaleo, X. Sun and C. Culberson, *Mycologia*, 2011, **103**, 741–754
- 5 D. A. Herbst, C. A. Townsend and T. Maier, *Nat. Prod. Rep.*, 2018, 35, 1046–1069.
- 6 S. Ferron, P. Jéhan, X. Guillory and P. Uriac, *Phytochemistry*, 2022, 198, 113139.
- 7 M. Millot, S. Tomasi, S. Sinbandhit and J. Boustie, Phytochem. Lett., 2008, 1, 139-143.
- 8 M. Millot, S. Tomasi, K. Articus, I. Rouaud, A. Bernard and J. Boustie, *J. Nat. Prod.*, 2007, **70**, 316–318.
- 9 J. Elix, B. Ferguson and M. Sargent, Aust. J. Chem., 1974, 27, 2403.
- 10 S. Huneck, A. Porzel and H. T. Lumbsch, *Hergozia*, 1997, 39–43
- 11 O. E. Krivoshchekova, N. P. Mishchenko, L. S. Stepanenko and O. B. Maksimov, *Chem. Nat. Compd.*, 1983, **19**, 11–16.
- 12 A. Saeed, Monatsh. Chem., 2003, 134, 457-463.
- 13 G. E. Dunn, E. G. Janzen and W. Rodewald, Can. J. Chem., 1968, 46, 2905–2909.
- 14 S. Hellou, P. Uriac, F. Le Dévéhat, A. Sauvager, P. Jéhan, A. Zebboudj, J. Boustie and J. Esnault, *Herzogia*, 2019, 32, 485–502.
- 15 K. Mustafa, H. G. Kjaergaard, N. B. Perry and R. T. Weavers, *Tetrahedron*, 2003, **59**, 6113–6120.
- 16 K. Krohn, R. Bahramsari, U. Flörke, K. Ludewig, C. Kliche-Spory, A. Michel, H.-J. Aust, S. Draeger, B. Schulz and S. Antus, *Phytochemistry*, 1997, 45, 313–320.