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Chlorophyll Biodegradation Products from Hamamelis Virgianana Autumnal Leaves

Abstract: The biodegradation of bioorganic solid waste involves several million tons of senescent plant leaves every autumn. Chemical evidence about bioorganic matter contained in the senescent leaves remains undetermined. The biodegradation of senescent leaves comprises a series of biodegradation transitions that bring about changes in leaf texture, metabolic changes and color. Leaves turn yellow as a result of chlorophyll biodegradation. Chlorophyll biodegradation products, in of Hamamelis virginiana, the autumnal leaves Hamamelidaceae, were investigated. Here is a report on one chlorophyll biodegradation product isolated from yellow Hamamelis virginiana, Hamamelidaceae autumnal leaves. The structure of the isolated chlorophyll biodegradation product was elucidated by spectroscopic and spectrometric data. The isolated chlorophyll biodegradation product was an UNCC (Urobilinogenic Non fluorescent Chlorophyll Catabolite).

Keywords: UNCC, biodegradation, Hamamelis virginiana, Hamamelidaceae

1. INTRODUCTION

Chlorophyll biodegradation has been observed in senescent leaves of several plant families: Altingiaceae[1], Amaranthaceae[2, Brassicaceae[4], 3], Cercidiphyllaceae[5, Gramineae[7], 6], Hamamelidaceae[8] and Solanaceae[9, 10]. From the practical stand point, colorless chlorophyll biodegradation products isolated can be divided into two groups: non fluorescent chlorophyll catabolites (NCCs)[1, 2, 3, 4, 5, 6, 7, 9, 10] and urobilinogenic non – fluorescent chlorophyll catabolites further (UNCCs)[8]. NCCs can subdivided into two groups: glucosylated ones[4, 9, 10] and aglicons[1, 2, 3, 5, 6, 7]. Aglicons can further be subdivided into two group: the ones where modification on the lateral vinyl group has not occurred[1, 2, 5, 6] and others where the lateral vinyl group was oxidized into 1, 2 – diols[3, 7]. The chlorophyll biodegradation product isolated from *Parrotia persica*, Hamamelidaceae was an UNCC[8]. The purpose of this paper is to compare chlorophyll biodegradation product isolated from *Hamamelis virginiana*, Hamamelidaceae autumnal leaves with the chlorophyll biodegradation product isolated from *Parrotia persica*, Hamamelidaceae autumnal leaves and to observe the differences

2. EXPERIMENTAL PART

General

General part is the same as described previously [8].



Plant material

Hamamelis virginiana, Hamamelidaceae, leaves were collected during the autumn (2004) from the Botanical Garden of Fribourg, Switzerland.

Extraction and Isolation

Extraction and isolation were the same as for Parrotia persica autumnal leaves[8], with the following differences: Hamamelis virginiana Hamamelidaceae leaves (120.92 g dry weight, 175.00 g "fresh" weight) were frozen with liquid nitrogen, grinded and extracted with methanol. Crude Hamamelis virginiana extract obtained after evaporation of dichloromethane (t<40°C) yielded 230 mg. Crude extract revealed the presence of a chlorophyll biodegradation product with a spot at R_f= 0.52 on TLC. Prepurification was done on MPLC and 11.8 mg of the prepurified chlorophyll biodegradation product was obtained. Final purification was done by semi - preparative HPLC and the chlorophyll biodegradation product eluted at 73 min. was collected to obtain 1.07 mg of the pure chlorophyll biodegradation product.

3. RESULTS AND DISCUSSION

The isolation procedure for Hamamelis virginiana autumnal leaves was the same as for the *Parrotia persica* autumnal leaves[8]. The LC - MS analysis of the crude Hamamelis virginiana autumnal extract was subjected to RP - C₈ analytical column under the same acquisition parameters and elution solvent mixture as for the Parrotia persica autumnal leaves' extract[8]. The major compound in LC - MS chromatogram of Hamamelis virginiana autumnal leaves' extract gave molecular ion $[M+H]^+$ corresponding to m/z 633 like the UNCC present in Parrotia persica autumnal leaves' extract[8]. The Hamamelis virginiana UNCC eluted at 59.3 min. (Figure 1) and Parrotia persica UNCC eluted at 57.5 min. (Figure

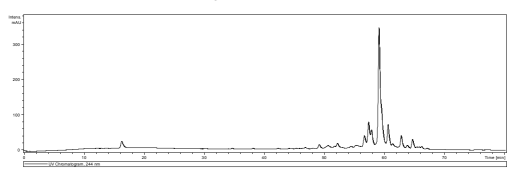


Figure 1. UV chromatogram of Hamamelis virginiana crude leaves' extract, extracted at $\lambda = 244$ nm

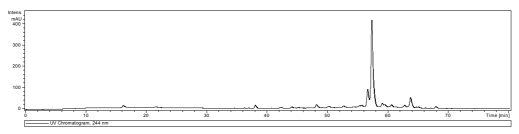


Figure 2. UV chromatogram of Parrotia persica crude leaves' extract, extracted at $\lambda = 244$ nm



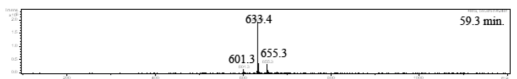


Figure 3. ESIMS of Hamamelis virginiana crude leaves' extract, extracted at 59.3min.

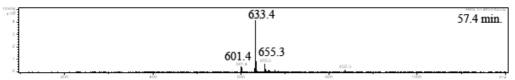


Figure 4. ESIMS of Hamamelis virginiana crude leaves' extract, extracted at 57.4min.

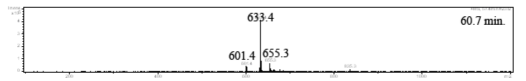


Figure 5. ESIMS of Hamamelis virginiana crude leaves' extract, extracted at 60.7min.

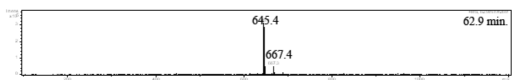


Figure 6. ESIMS of Hamamelis virginiana crude leaves' extract, extracted at 62.9min.

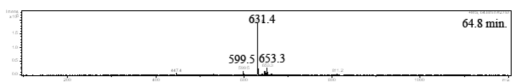


Figure 7. ESIMS of Hamamelis virginiana crude leaves' extract, extracted at 64.8min.

UNCC present in *Hamamelis virginiana* autumnal leaves (Figure 1) eluting at 57.4 min. had the *m/z* 633 (Figure 4) and corresponded to the *Parrotia persica* UNCC (Figure 2), in further text UNCC – *Pp* acronym will be used. The major UNCC in *Hamamelis virginiana* eluted at 59.3 minutes had also the *m/z* 633 (in further text UNCC – *Hvir* acronym will be used) (Figure 3). Another UNCC with the *m/z* 633 was present in *Hamamelis virginiana*

crude leaf extract (Figure 5), along with the NCC with the m/z 645, also called Cj-NCC-1 [5, 6], major chlorophyll biodegradation product from Cercidiphyllum japonicum, Cercidiphillaceae (Figure 6). The compound with the m/z 631 has yet not been characterized (Figure 7).

The major UNCC present in *Hamamelis virginiana* crude leaves' extract was subjected to MPLC (230 mg) to yield 11.8 mg of the prepurified UNCC. The final

purification was performed by the means of the semi – preparative HPLC to give 1.07 mg of the pure UNCC– *Hvir*. The UNCC –

Hvir eluted at 73 min. (Figure 8). The UNCC – *Pp* eluted at 52 min. (Figure 9)

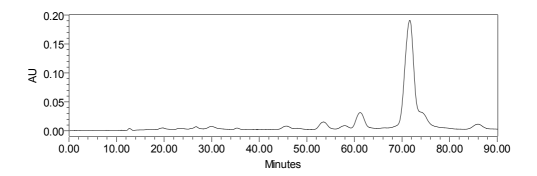


Figure 8. UV chromatogram of the prepurified Hamamelis virginiana leaves' extract, extracted at $\lambda = 244$ nm

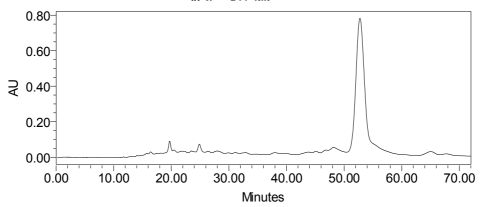


Figure 9. UV chromatogram of the prepurified Parrotia persica leaves' extract, extracted at $\lambda = 244$ nm [8]

Constitution of UNCC - Hvir

The UNCC – Hvir was obtained as orange amorphous solid. The High Resolution ElectroSpray Ionisation Mass Spectrometry (HRESIMS) sprectra showed a molecular ion at m/z 655.2738 for the molecular formula $C_{34}H_{40}N_4O_8Na$ [M+Na]⁺, calculated m/z 655.2738, Δ 0.00 ppm.

Elucidation of the UNCC – *Hvir* structure by NMR data.

In the proton spectrum, there were slight differences in the chemical shifts

between the two UNCCs isolated, one from *Parrotia persica* and the other from *Hamamelis virginiana* autumnal leaves. The NMR spectra of the two UNCCs were measured at the same temperature and in the same solvent CD₃OD-d₄. The compound isolated from *Hamamelis virginiana* autumnal leaves was an isomer of UNCC – *Pp* [8]. The UNCC – *Hvir* proton spectrum is shown in the Figure **10**.



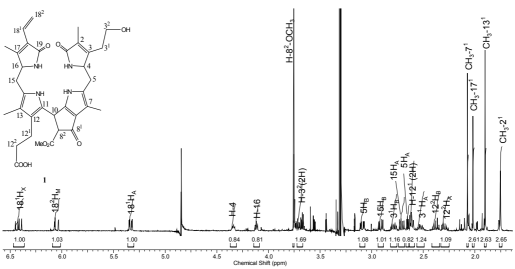


Figure 10. The high resolution proton spectrum of the UNCC - Hvir

The multiplicity of the proton H-16 signal of the UNCC – Hvir was a triplet while in the UNCC – Pp proton spectrum a doublet of doublets was observed (Figure 11). When small interprotonal couplings are

underestimated, the multiplicity of the UNCC – *Hvir* H-4 proton signal is *triplet*, while in case of UNCC – *Pp* it is the *doublet of doublets*.

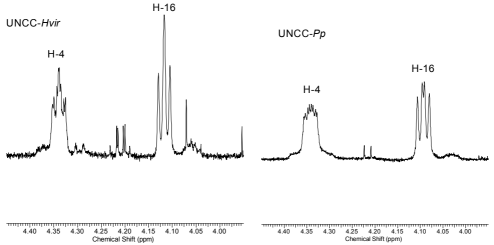


Figure 11. Proton spectra in the region of H-4 and H-16 of the UNCC – Hvir (left) and UNCC – Pp (right).

The difference between two isomers was observed in CD spectra. The chromophore absorbing at λ =244 nm

had $\Delta \varepsilon$ positive in case of UNCC-Pp and in case of UNCC-Hvir the $\Delta \varepsilon$ was negative (Figure 12 and 13).

H/C	S multiplicity	I/Uz)	2	DEPT	COCV (H \H)
	$\delta_{\rm H}$, multiplicity	J(Hz)	δ_{C}	DEFI	COSY (H→H)
1					
21	1.75		0.5		2 11
2 2 ¹ 3 3 ¹	1.75 s		8.5	<u> </u>	$3^{1}H_{A}$
3		6 4 4 2 0	21.1		0 0 0 0 0
31	$2.52 dd H_A$	6.4;13.9	31.1	↓ ↓	$2^{1}, 3^{1}H_{B}, 3^{2};$
	2.50 1.111	6.4;13.3			$3^{1}H_{A}, 3^{2}$
• 2	2.79 <i>dd</i> H _B	60.10.1	<i>-</i>		
3^2	3.67-3.74 m	6.8;13.4	61.4	↓	$3^{1}H_{A,}3^{1}H_{B}$
	H _A and H _B	7.3;13.9			
4	4.34 tt	4.8; 1.4	60.6	1	5H _A , 5H _B
5	$2.65 dd H_A$	8.5;14.9	29.8	↓	4, 5H _A ;
	$3.09 dd H_{\rm B}$	14.9;4.8			4, 5H _B
6					
7					
7 ¹	2.07 s		9.6	1	
8					
8 8 ¹ 8 ² 8 ³ 8 ⁴					
8 ²	3.79 d	3.2			
83					
84	3.75 s		52.9	↑	
9				'	
10	overlapped by		37.4	1	
	the water peak				
11	l and water pour				
12					
12 ¹	2.62 t H _A and	7.7	36.7		12 ¹ H _B ; 12 ¹ H _A
12	H _B	7.7	30.7	*	12 11 _B , 12 11 _A
12 ²	$2.29 dt H_A$	7.5;15.6	20.8		$12^{1}H_{B}, 12^{2}H_{B};$
12	$2.38 dt H_{\rm B}$	8.0;15.9	20.0	*	$12^{11}H_{A}$, $12^{2}H_{A}$
12 ³	2.50 W 11B	5.0,15.7			
13					
131	1.90 s		9.5	1	
14	1.703		7.5	1	
15	2.71 <i>dd</i> H _A	6.9;14.7	29.5	1	15H _B , 16;
13	$2.71 dd H_A$ $2.91 dd H_B$	5.4;14.7	29.3	+	15H _B , 16, 15H _B , 16
16	4.12 <i>t</i>	6.0	61.6	1	15 H _A , 15 H _B , 17 ¹ *
	7.12 1	0.0	01.0	1	13 ПД, 13 ПВ, 17
17 17 ¹	2.02 g		12.8	1	16*
	2.02 s		12.0	<u> </u>	10.
18	(42.11	17.0	107.0		10211 10211
18 ¹	6.42 <i>dd</i>	17.8; 11.7	127.0	1	$18^2 H_A, 18^2 H_B$
18 ²	5.33 <i>dd</i> H _A	11.7,2.4	119.2		18 ² H _{B.} 18 ¹ ; 18 ² H _{A.}
	$6.05 dd H_{\rm B}$	17.8,2.4	117.2	*	18 ¹
19	0.00 WW 11B	17.0,2.1			
	U (500 MH-) and	 3 <i>C (</i> 125 M	77 \ 77.55	1	D. J. of A. a. UNICC Ho

Table 3. ¹H (500 MHz) and ¹³C (125 MHz) NMR data in CD₃OD-d₄ of the UNCC-Hvir DEPT data: CH and CH₃ are phased up, CH₂ is phased down *cross peak of low intensity



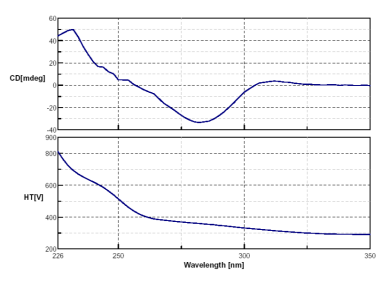


Figure 12. CD and UV spectra of 0.1 µmoldm⁻³ UNCC-Pp in methanol

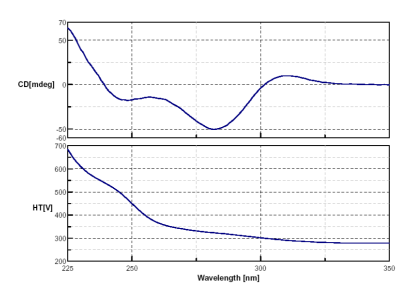


Figure 13. CD and UV spectra of 0.03 µmoldm⁻³ UNCC-Hvir in methanol

Spectroscopic data

UV – Vis in methanol, $C = 3 \cdot 10^{-8}$ mol dm⁻³; $\lambda_{\text{max}}[\text{nm}]$ (log ϵ): 244 (7.24), 283 (7.03).

CD in methanol, $C = 3.10^{-8}$ mol dm⁻³; $\lambda_{\text{max}}[\text{nm}]$ ($\Delta \epsilon$): 244 (-16.7), 283 (-51).

4. CONCLUSIONS

The UNCC was isolated from *Hamamelis virginiana* autumnal leaves and was named UNCC – *Hvir*. Its structure was determined by spectroscopic and spectrometric data. The UNCC – *Hvir* was compared to



the UNCC isolated from *Parrotia persica* (UNCC – Pp) autumnal leaves. The isolated UNCC – Hvir differs from UNCC – Pp in few physico – chemical characteristics. Under the same elution conditions, on the same stationary phase, on the analytical scale, the UNCC – Hvir eluted at 59.0 min. (The capacity factor 2.60) and UNCC – Pp at 57.3 min. (The capacity factor 2.49). Under the same elution conditions, on the same stationary phase, on the semi – preparative scale, UNCC – Hvir eluted at 73 min. And UNCC – Pp at 52 min. The main difference in the proton NMR spectrum was the multiplicity of the proton H-4. When the

interprotonal couplings underestimated the multiplicity of UNCC -Hvir H-4 is a triplet and the multiplicity of UNCC - Pp is a doublet of doubles. The other difference in the proton spectrum was the multiplicity of the proton H-16. In the case of the UNCC - Hvir the multiplicity of the proton H-16 was a triplet and in the case of the UNCC - Pp the multiplicity was doublet of doublets. The UNCC - Hvir isolated from Hamamelis virginiana autumnal leaves is an isomer of the UNCC Pp isolated from Parrotia persica autumnal leaves[8].

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