AGRICULTURAL AND FOOD CHEMISTRY

Chemical Composition of the Essential Oils of Serbian Wild-Growing Artemisia absinthium and Artemisia vulgaris

Polina Blagojević, Niko Radulović, Radosav Palić, and Gordana Stojanović*

Department of Chemistry, Faculty of Science and Mathematics, Višegradska 33, 18000 Niš, Serbia and Montenegro

The chemical composition of the aerial and root essential oils, hydrodistilled from *Artemisia absinthium* L. and *Artemisia vulgaris* L. (wild-growing populations from Serbia), were studied by gas chromatography, gas chromatography–mass spectrometry, and ¹³C nuclear magnetic resonance. During the storage of plant material under controlled conditions, a significant decrease of essential oil yields (isolated directly after drying and after 1 year of storage) and significant differences in their chemical compositions were observed. A possible mechanism for the observed oil component interconversion has been discussed. The noticeable differences in the chemical composition of the oils isolated from roots and aerial parts of *A. absinthium* and *A. vulgaris* were also correlated with the diverging biosynthetic pathways of volatiles in the respective plant organs. The antimicrobial activities against the common human pathogens of all of the isolated oils were tested according to National Committee on Clinical Laboratory Standards. The oils showed a broad spectrum of antimicrobial activity against the tested strains. Therefore, these oils can be used as flavor and fragrance ingredients.

KEYWORDS: Artemisia absinthium; Artemisia vulgaris; essential oil; root; storage period; plant organ specification; antimicrobial activity

INTRODUCTION

According to the Council Directive 88/388/EEC (1), on the approximation of the laws of the Member States relating to flavorings for use in foodstuffs and source materials for their production, the addition of thujone-containing plants was reallowed in the European Union. For this reason, an increase in the industrial consumption of Artemisia absinthium L. (wormwood), Asteraceae, limited in the last century as the result of absinth prohibition (2, 3), could be expected. However, the A. absinthium plant material used in industries is still in many instances harvested from wild populations, with variable, nonstandardized thujone (both α - and β -isomers) contents (4). This can be seen from the numerous formerly published data on the variability of A. absinthium essential oil composition considered in detail in the Results and Discussion section. There are no previous reports on the composition of the essential oil isolated from A. absinthium and Artemisia vulgaris L. [locally known as "beli (white) pelin" and "crni (black) pelin", respectively] growing wild in Serbia.

In addition to the renowned wormwood application in preparation of absinth and related beverages, *A. absinthium* has been used since ancient times for medical purposes. From the ethnopharmacological point of view, *A. absinthium* has been used for its antihelmintic, stomachic, antibacterial, antifeedant, antifertility, antipyretic, cytostatic, antitumor, and antimalarial

actions (3, 5). In this sense, it has been reported that *A. vulgaris* (mugwort) possesses similar uses to *A. absinthium*. The rational justification of the widespread use of the mentioned herbs was assessed by testing the antibacterial and antifungal activities of the isolated oils against seven common human pathogens.

Both *A. absinthium* and *A. vulgaris* represent economically important plant species; hence, the investigation of the influence of storage time on the chemical compositions of their volatile oils deserves attention. Because both roots and aerial parts of *A. vulgaris* are used as an herbal remedy (5), we found it interesting to determine the differences in the chemical compositions of the oils isolated from roots and aerial parts. It is surprising that the root essential oil of *A. vulgaris*, a widespread and investigated plant species, has not previously been analyzed. It has been reported that the thujone type monoterpenoids (including thujone itself) were found in only trace amounts in *A. absinthium* root oil (*6*, 7). The possibility of utilizing *A. absinthium* root oil in flavoring or for medical purposes provoked us to reexamine the oil of *A. absinthium* roots as well.

In light of the above-mentioned information, the aim of the present study was (i) to investigate the essential oil chemotypes of wild-growing *A. absinthium* and *A. vulgaris* from Serbia, using gas chromatography (GC), gas chromatography-mass spectrometry (GC-MS), and ¹³C nuclear magnetic resonance (NMR) analysis; (ii) to asses their antimicrobial activity; and (iii) to clarify the influence of plant organ specification and storage on essential oil yield and composition.

^{*} To whom correspondence should be addressed. Tel: +381 63 8949353. Fax: +381 18 533014. E-mail: stgocaus@yahoo.com.

MATERIALS AND METHODS

Plant Materials and Isolation of Essential Oils. The investigation of two populations of *A. absinthium* growing spontaneously at two different locations in the southeastern Serbia was carried out. For this purpose, plant material was collected in the village Mokra (Bela Palanka) (aerial parts; June, 2002; 313 m above sea level) and at the banks of the river Nišava in the urban surroundings of Niš (whole plant; July, 2003; 199 m above sea level). *A. vulgaris* (entire herb) was gathered at Nišava riverside in the city of Niš in July, 2003.

All mentioned species were collected in the full-blooming stage. The voucher specimens were deposited at the Herbarium of the Faculty of Biology, University of Belgrade (BEOU): 16030 (*A. vulgaris*), 16033 (*A. absinthium*, Mokra), and 16034 (*A. absinthium*, Niš).

Immediately after the specimens were air-dried at room temperature (to constant moisture content), the plant material was subjected to hydrodistillation for 2.5 h using a Clevenger type apparatus. In the case where the influence of prolonged storage period was investigated, the dried herbs were additionally kept, in a sealed container, for a total of 1 year prior to oil isolation at ambient temperature (25 ± 2 °C) and without exposure to direct sunlight (*A. absinthium*, Mokra). The oils were dried over anhydrous MgSO₄ and kept at 4 °C until analysis.

Throughout the text, the following abbreviations have been used to designate the obtained essential oils: Aa1 and Aa2 (oils from aerial parts of *A. absinthium*, Mokra, isolated immediately after drying the plant material, and stored for 1 year, respectively); Aa3 and Aa3R (oils from aerial parts and roots of *A. absinthium* oil, Nišava riverside, respectively); and Av and AvR (oils from aerial parts and roots of *A. vulgaris*, Nišava riverside, respectively). Yields of the isolated oils (w/w of dry plant material) and the results of GC, GC-MS, and ¹³C NMR analyses are presented in **Table 1**.

GC and GC-MS. The GC-MS analysis of the oils was performed using a Hewlett-Packard 5890 series II gas chromatograph equipped with SPB-1 (30 m \times 0.25 mm i.d., 0.25 μ m film thickness) and DB-5 $(30 \text{ m} \times 0.2 \text{ mm i.d.}, 0.25 \,\mu\text{m}$ film thickness) fused silica capillary columns directly coupled to a mass selective detector MSD 5971A from the same company, which was operated in EI mode (70 eV). Helium was the carrier gas at a flow rate of 1 mL/min. The injector temperature was 250 °C, and the oven temperature was programmed as follows: held isothermal at 50 °C for 3 min, then gradually increased to 250 °C at 5 °C/min, and finally held isothermal at 250 °C for 15 min. The volume injected was 0.1 μ L of the 10% solution, diluted with diethyl ether. The conditions for the GC-flame ionization detection (FID) were the same as for the GC-MS except H₂ was the carrier gas. The area percentage was obtained electronically from the GC-FID response without the use of an internal standard or correction factors. The compound content in the oils hydrodistilled from A. absinthium (Mokra) before and after 1 year of storage (Aa1 and Aa2 oils, respectively) was additionally expressed as the relative amount of identified components to the dry weight of plant material (mmol/100 g), and the corresponding values are given in Table 1.

NMR Spectroscopy of Carbon-13. ¹³C NMR spectra of the essential oils were recorded on a Varian Mercury 300 spectrometer, operating at 75.462 MHz, equipped with a 5 mm probe in benzene- d_6 (around 50 mg in 0.5 mL of C₆D₆). Chemical shifts are expressed in δ (ppm) downfield from tetramethylsilane as an internal standard. Parameters: pulse width, 6.0 μ s (flip angle, 45°); acquisition time, 1.705 s; relaxation delay D1, 0.294 s (total recycling time 1.999 s); for 64 K data table with a spectral width of 18863.2 Hz; globally optimized alternating-phase rectangular pulses were used; digital resolution, 0.469 Hz/pt; and 5000 scans were accumulated. An exponential multiplication of the free induction decay with the line broadening of 1.0 Hz was applied before Fourier transformation.

Identification Procedure. The linear retention indices (RIs) for all of the compounds were determined by coinjection of the sample with a solution containing the homologous series of C_8-C_{25} *n*-alkanes. Individual identification of components was based on comparison of their mass spectra (MS) with those of Wiley 275 MS and MassFinder 2.3 (8) libraries and with those described by Adams (9), as well as on comparison of their RIs (10) with literature values (9, 11) and, wherever possible, by coinjection with a reference sample (ref; see **Table 1** for

the method of identification of each individual compound). The identity of some compounds was also confirmed by 13 C NMR in the same way as it was described in the ref *12*.

Antimicrobial Activity. The disk diffusion method according to the NCCLS (13) was employed for the determination of in vitro antimicrobial activity of the essential oils and available pure constituents [in 1:10 and 1:30 dilution, w/v (mg/ μ L)], completely following the method and using the same panel of laboratory control strains (see Results and Discussion) as previously reported in ref 12. Standard disks of doxycycline, tiamulin, erythromycin, and nystatin (origin, Institute of Immunology and Virology "Torlak", 30 μ g of the active component, 6 mm diameter) were individually used as positive controls, while the disks imbued with 50 μ L of pure diethyl ether were used as negative controls. Each test was performed in triplicate and repeated three times, and results were analyzed for statistical significance.

RESULTS AND DISCUSSION

Influence of Storage. Because the increase in A. absinthium exploitation in the food industry can be expected, it is of great importance to establish whether significant changes in oil composition and yield during storage of wormwood occur. After 1 year of plant material storage, the oil yield dropped from 0.29 to 0.08 (%, w/w), as well as the number of detected components. With some observed exceptions, all constituents were found in reduced quantities in Aa2 oil (prolonged storage time) (Table 1, values in parentheses). This could be due to, at least partially, the evaporation of the oil constituents during storage. Factors such as light, temperature, and moisture greatly influence the emission of volatiles, and in general, physical properties of one particular compound determine the rate of its release into the atmosphere (14). If the decrease in the amount of components is only the consequence of their volatility properties, it is reasonable to expect a larger decrease in the amount for those constituents that are characterized with higher vapor pressures. Indeed, the fraction ratio of oxygenated monoterpenes that are less volatile than the parent hydrocarbons to hydrocarbon monoterpenes in Aa1 and Aa2 oils was 6.3 and 17.5, respectively, indicating a 2.8 times higher rate of emission of the hydrocarbon fraction. This relationship seems to stand also within the sesquiterpene fraction but is, as expected, less pronounced (0.6 in Aa1 and 0.7 in Aa2 oil). The general ratio of the monoterpene to the sesquiterpene fraction does not, however, follow this rule (the fraction ratio of total monoterpenes to the total sesquiterpenes is 6.1 in Aa1 oil and 21.9 in Aa2 oil). However, this simplified approach could be misleading since some monoterpenoids are less volatile than some other sesquiterpenes.

The relative amounts of certain individual components were higher in the oil isolated from plant material after 1 year of storage: cis- and trans-linalooloxides (furanoid) (1 and 2), 1,8cineole (3), neryl 2-methylpropanoate (4), and geranyl 3methylbutanoate (5) (Figure 1). This increase could be the outcome of chemical transformations of initially existing compounds. All of these transformations could be associated with certain existing metabolic pathways and connected with the action of enzymes or could be temperature, humidity, and light driven alone. Most probably, at the beginning of the drying process, both enzymatic and nonenzymatic reactions took a role in the terpenoid transformation. However, it could be expected that, as a consequence of complete plant dehydration, enzymatic transformations were stopped [enzyme inactivation in dry plant material is very probable (15)]. Figures 2 and 3 show straightforward reaction pathways that could interconnect oxides of linalool (1 and 2) with its potential precursors: $cis-\beta$ epoxyocimene (6), linalool (7), or β -myrcene (8), components

Table 1. Composition of the A. absinthium and A. vulgaris Essential Oils

					A. absinthiu	т	A. v.	Igaris		
		R	а		yield in parentheses					
	compound	SPB-1	DB-5	Aa1 (0.29)	Aa2 (0.08)	Aa3 (0.26)	Aa3R (0.22)	Av (0.06)	AvR (0.04)	identification method ^b
1	hexanal ^c	769	801	d					0.2	RI, MS
2	santolinatriene	910	909					4.0		RI, MS
3	α-thujene	938	928	0.2 (0.33) ^e	0.6 (0.26)	0.4		1.0		RI, MS
4	α-pinene	947	940	0.2 (0.33)	0.4 (0.17)	0.8	0.6	5.9	0.1	RI, MS, ref
с 6	α-renchene"	959 961	950 953	0.1 (0.16)	0.3 (0.13)	2.3	23.3	0.1	tr' tr	RI, MS, "C NIVIR RI MS ref
7	sabinene	992	976	8.1 (13.23)	3.3 (1.42)	10.8	0.2	13.7	u	RI, MS, ¹³ C NMR, ref
8	β -pinene	997	980	011 (10120)	010 (1112)		1.7	tr	tr	RI, MS, ref
9	β -myrcene	1016	994	0.9 (1.47)		0.3	2.3	1.3		RI, MS, ref
10	α-phellandrene ^c	1028	1006	0.7 (1.14)		0.5	3.4			RI, MS
11	<i>p</i> -cymene	1047	1027	1.2 (1.96)	0.5 (0.22)	6.7	4.1	tr	tr	RI, MS, ref
12	1,8-CINEOIE	1050	1034	1.0 (1.63)	16.3 (7.01)			28.9	0.3 tr	RI, MS, ¹³ C NMR, ref
14	v-terninene ^c	1033	1059	0 2 (0 33)				03	u	RI, MIS, TEI RI MS
15	<i>trans</i> -sabinene hvdrate ^c	1087	1069	0.3 (0.49)				0.0	tr	RI, MS
16	cis-linalooloxide (furanoid) ^c	1089	1070		1.5 (0.64)				tr	RI, MS
17	fenchone ^c	1097	1088						tr	RI, MS
18	trans-linalooloxide (furanoid) ^c	1100	1090		1.5 (0.64)					RI, MS
19	a-thujone	1113	1102	0.9 (1.47)	0.6 (0.26)	1.8		1.2		RI, MS, ref
20 21		1123	1104	4.1 (0.70) 19.8 (32.33)	20.2 (0.09)	63.4		13.5		RI, MO, IEI RI MS ¹³ C NMR ref
22	camphor	1129	1143	10.0 (02.00)	20.2 (0.03)	00.4		1.4	0.4	RI, MS, ref
23	$cis-\beta$ -epoxyocimene ^c	1141	1139	10.7 (17.47)					••••	RI, MS, ¹³ C NMR
24	trans-sabinol ^c	1148	1140	2.5 (4.08)						RI, MS
25	albene ^c	1161	1152						0.6	RI, MS
26	lavandulol ^c	1167	1170	1.2 (1.96)	0.0 (0.00)			0.0		RI, MS
27	4-terpineoi	1173	1178	1.7 (2.78)	2.3 (0.99)			2.2		RI, MS, ref
29	trans-sabinenehvdrate acetate ^c	1244	1229	2.0 (4.07)				2.5		RI, MS, ICI
30	bornyl acetate	1258	1280			0.3	4.2	tr	5.0	RI, MS, ref
31	trans-sabinyl acetate ^c	1264	1290	8.8 (14.37)	15.5 (6.66)					RI, MS, ¹³ C NMR
32	isobornyl acetate ^c	1268	1290						tr	RI, MS
33	linalyl acetate	1270	1272			0.1	7.7		0.0	RI, MS, ¹³ C NMR, ref
34	7α-slipniperfol-5-ene ^c	1307	1300						0.3	RI, MS RI MS
36	silphin-1-ene ^c	1329	1320						1.0	RI MS
37	eugenol	1346	1356						0.3	RI, MS, ref
38	linalyl propanoate ^c	1346	1363				8.2			RI, MS, ¹³ C NMR
39	isobornyl propanoate ^c	1349	1366						0.8	RI, MS
40	pethybrene ^c	1362	1371						tr	RI, MS
41	silphiperfol-6-ene	1365	1375					0.6	tr	RI, MS PL MS rof
42 42	modhenhene ^c	1369	1385					0.0	12	RI, MO, IEI RI MS
43	β -bourbonene ^c	1374	1384					tr		RI, MS
44	α -isocomene ^c	1377	1388						4.0	RI, MS, ¹³ C NMR
45	linalyl 2-methylpropanoate ^c	1379	1378				1.0			RI, MS
46	β -elemene ^c	1380	1393					1.4	2.0	RI, MS
47	petasitene	1385	1400						0.7	RI, MS DI MS
40	β -isocomene ^c	1390	1402						4.9	RI, MS
50	α -gurjunene ^c	1400	1409					tr	tr	RI, MS
51	linalyl butanoate ^c	1402	1414	1.2 (1.96)	1.8 (0.77)	0.7	14.4			RI, MS, ¹³ C NMR
52	α-cedrene ^c	1405	1411						0.2	RI, MS
53	β -caryophyllene	1408	1418	3.0 (4.90)		1.2		2.3	1.2	RI, MS, ref
54	α -santalene	1410	1424						tr 0.4	RI, MS
56	eni-B-santalene ^c	1424	1420						0.4	RI, MS
57	α -humulene	1440	1444	0.7 (1.14)				0.5	1.6	RI, MS, ref
58	β -santalene ^c	1445	1462	()					0.8	RI, MS
59	aromadendrenec	1447	1434					0.9		RI, MS
60	β -bisabolene ^c	1462	1496						1.7	RI, MS
61	γ -numulene γ	1464	1481		0 5 (0 22)		0.0	1.1	1.1	RI, MS DI MS
02 63	$(7F)$ - α -famesene ^c	1409	1475 1462	3 3 (5 30)	0.0 (0.22) 2.5 (1.08)		∠.ŏ			RI MS
64	β -selinene ^c	1470	1474	0.8 (1.31)	2.0 (1.00)	2.9		4.7	2.0	RI, MS
65	, allo-aromadendrene ^c	1471	1477	()					tr	RI, MS
66	bicyclogermacrene ^c	1477	1491						0.7	RI, MS
67	γ -gurjunene ^c	1476	1479		10 - ()		<u>.</u>	2.8		RI, MS
68	Inalyl 3-methylbutanoate ^c	1484	1473	7.5 (12.25)	12.5 (5.38)	4.5	21.1		0 4	
09 70	bornyi δ-metnyibutanoate° δ-cadinene ^c	1400 1500	1400 1505					1 2	0.4 1 2	RI, IVIO, "CINIVIK RI MS
10	o dumeno	1000	1000					1.0	ч.Ј	N, WO

Table 1. (Continued)

					A. absinthiu	m	A. vulgaris					
		RI	а		yield in parentheses							
	compound	SPB-1	DB-5	Aa1 (0.29)	Aa2 (0.08)	Aa3 (0.26)	Aa3R (0.22)	Av (0.06)	AvR (0.04)	identification method ^b		
71	<i>cis</i> -α-bisabolene ^c	1521	1504						tr	RI, MS		
72	myrtenyl 3-methylbutanoate	1532	1521	4 4 (7 40)					0.8	RI, MS		
73	neryl 3-methylbutanoate	1553	1535	4.4 (7.18)					12.2	RI, MO		
74 75		1003	1504	3.7 (0.04)				65	13.2	RI, IVIS, "C INIVIR DI MS rof		
75	carpolication of the second se	1507	1500	1 5 (2 15)	12 0 (5 55)	1 1	4.0	0.0	0.6			
70	deranyl 2-methylbutanoate ^c	1503	1570	0.0 (2.45)	3 3 (1 /2)	0.4	4.0		0.0 tr	RI, IVIO, CONIVIR		
78	bumulene ovide II ^c	1506	1505	0.9 (1.47)	3.3 (1.42)	0.4	0.1	1 2	u 25	RI, MS		
79	B-eudesmol	1647	1650					1.2	10.0	RI MS ¹³ C NMR ref		
80	valeranone ^c	1665	1672						5.5	RI MS ¹³ C NMR		
81	α -bisabolol ^c	1686	1681	0.7 (1.14)					6.0	RL MS, ¹³ C NMR		
82	chamazulene ^c	1728	1714	1.0 (1.63)					0.0	RL MS		
83	(E.E)-farnesal ^c	1735	1730	0.8 (1.31)						RI, MS		
84	(E,E)-farnesvl acetate ^c	1828	1844						tr	RI. MS		
85	trans-nerolidyl propanoate ^c	1829	1850						tr	RI, MS		
86	hexahydrofarnesyl acetone ^c	1838	1843						tr	RI, MS		
87	(Z)-nuciferyl propanoate ^c	1875	1893	1.2 (1.96)	0.3 (0.13)					RI, MS		
88	methyl palmitate	1908	1910						tr	RI, MS, ref		
89	ethyl palmitate	1968	1990						tr	RI, MS, ref		
90	(E)-nuciferyl 2-methylpropanoate ^c	1992	1997	0.6 (0.98)	0.4 (0.17)					RI, MS		
91	(E)-nuciferyl butanoate ^c	2004	2012	1.7 (2.78)	1.1 (0.47)	0.7				RI, MS		
92	(E,E)-farnesyl 3-methylbutanoate ^c	2040	2058						tr	RI, MS		
93	methyl linoleate	2071	2097						tr	RI, MS, ref		
94	ethyl linoleate	2139	2155						tr	RI, MS, ref		
	totally identified			98.4	98.5	98.9	99.1	99.3	86.7			
	no. of components			35	22	18	16	29	63			
	monoterpenes			84.6	94.2	94.1	99.1	76.0	30.5			
	hydrocarbons			11.6	5.1	21.8	35.6	26.0	0.1			
	oxygenated (without esters)			45	42.6	65.2	0.0	47.5	0.7			
	esters			28	46.5	7.1	63.5	2.5	29.7			
	sesquiterpenes			13.8	4.3	4.8	0.0	23.3	55.7			
	nyurucarDons			0.0 5.0	2.5	4.1	0.0	1.1	31.1			
	othor			0.0	1.0	0.7	0.0	0.0	24 0 5			
	ouiei			0.0	0.0	0.0	0.0	0.0	0.5			

^a RIs relative to *n*-alkanes on SPB-1 and DB-5 capillary columns. ^b Methods: RI, identification based on RI comparison with literature data on at least one column (SBP-1, DB-5); MS, identification based on mass spectra comparison with those of Wiley 275 MS and MassFinder 2.3 (*8*) library as well as with those described by Adams (*9*); ¹³C NMR, identification based on ¹³C NMR spectra comparison; and ref, coinjection with an authentic sample. ^c Tentatively identified by the combination of MS and RI on two columns and/or ¹³C NMR. ^d Compound not found. ^e The values in parentheses represent the compound content expressed in mmol/100 g of dry plant material.^f Less then 0.1%.

found only in Aa1 oil. After initial epoxidation of linalool, the subsequent in situ conversion of the epoxide to the furanooxides was reported previously (16). A similar chemical connection can be devised to explain the notable increase in the amount of 1,8-cineole (1.63 mmol/100 g in Aa1 and 7.01 mmol/100 g in Aa2 oil). The heat of formation of 1,8-cineole, as compared with those of *trans*-sabinol, *trans*-sabinene hydrate, linalool, nerol, geraniol, and *cis*- β -epoxyocimene, indicates that reactions that would have 1,8-cineole as a product, and some of abovementioned compounds as substrates, would tend toward 1,8-cineole as the thermodynamically more stable compound.

The amounts of α -thujene, α -pinene, and α -fenchene were decreased during storage. However, it was less than one might expect. A possible explanation in the case of α -pinene and α -fenchene could be found in the transformation of myrcene (17) or in the rearrangement reactions of sabinene (corresponding amount dramatically decreased in Aa2 oil) that would lead to the ring strain release due to the opening of the small threemember cycle. An additional explanation in the case of α -thujene could probably be sought in the transformations of *trans*-sabinene hydrate and *trans*-sabinyl acetate.

Even though the relative amounts of neryl 2-methyl propanoate (4) and geranyl 3-methyl butanoate (5) are higher in Aa2 than in Aa1 oil, the total amount of esters of acyclic monoterpenes decreased from 20.33 (Aa1 oil) to 12.33 mmol/ 100 g of dry plant material. The mentioned increase could be the consequence of isomerization processes of linally and neryl esters found in Aa1 oil.

Plant Organ Specification. Although the use of thujonecontaining plants is reallowed in the European Union, there are restrictions on the maximum acceptable levels of thujones in final products for consumption (1). Previously reported data concerning major contributors of A. absinthium (collected in different countries) oils are listed in **Table 2** (18-24), and as it can be seen, high variability, not only in quantity of the major compounds but also in the corresponding class, can be observed. It is, thus, clear why it is necessary to investigate wild wormwood populations for the essential oil composition prior to their application in food and beverage production.

The main constituents found in the essential oil isolated from the aerial parts of *A. absinthium* collected at Mokra were β -thujone (19.8%), *cis*- β -epoxyocimene (10.7%), *trans*-sabinyl acetate (8.8%), sabinene (8.1%), and linalyl 3-methylbutanoate (7.5%). The oil was dominated by monoterpenes (84.6%), most of all belonging to the thujane type compounds (40.6%). The highest amounts of β -thujone (63.4%), sabinene (10.8%), and *p*-cymene (6.8%) were detected in the aerial parts oil obtained from wormwood herbs gathered at the Nišava banks (Aa3). Aa3



Figure 1. Monoterpenes found in the *A. absinthium* essential oil from Mokra (Aa2) with an increased amount after the storage period: 1, *cis*-linalooloxide (furanoid); 2, *trans*-linalooloxide (furanoid); 3, 1,8-cineole; 4, neryl 2-methylpropanoate; and 5, geranyl 3-methylbutanoate.



Figure 2. Possible mechanism of linalool oxides (1 and 2) formation from $cis-\beta$ -epoxyocimene (6).

oil, similar to Aa1, is rich in monoterpenes of the thujane group (77.4%). All of the oils isolated from aerial parts of *A*. *absinthium* contained nuciferyl esters (**Figure 4**). These rather uncommon essential oil constituents, however, seem to be quite characteristic for species belonging to the genus *Artemisia* (24).

The oil isolated from aerial parts of *A. vulgaris* was characterized with a high amount of 1,8-cineole (28.9%), sabinene (13.7%), β -thujone (13.5%), and β -caryophyllene oxide (6.5%). Literature data on previously analyzed mugwort oils are summarized in **Table 3** (21, 23, 25–29).

As far as the food industry is concerned, every factor influencing the organoleptic properties of the oil, or its physiological action, no matter if it is genetic or environmental in nature, is of immense importance in plant selection. Having this in mind and the differences in composition of the oils isolated from aerial parts of *A. absinthium* gathered in Nišava banks and in Mokra, as well as from *A. vulgaris*, from previously reported ones (**Tables 2** and **3**), these plant populations could be considered to represent new chemotypes.

The oils obtained from *A. vulgaris* and *A. absinthium* (**Tables 3** and **2**) gathered in Egypt had α -phellandrene as the main compound (25). β -Myrcene, the major contributor of wormwood oil from the Republic of Bashkortostan, was also found in high amounts in mugworth oil from the same country (23). α -Phellandrene and β -myrcene were not found in such a significant amount in other *A. vulgaris* oils (**Table 3**). Camphor, the major component of *A. vulgaris* oil from Italy, was also one of the major contributors in the Italian *A. absinthium* oil (21). The

resemblance in essential oil compositions of two different plant species, geographically closely related, could indicate a similar high degree of susceptibility of the two *Artemisia* species to external factors. To some extent, this applies to the populations of *A. absinthium* and *A. vulgaris* collected in Niš.

The major component of wormwood root oil was α -fenchene (23.3%), but the most interesting property of this oil was the domination of aliphatic esters. The sum of linalyl (53.4%) (30), bornyl (4.2%), geranyl (4.1%), and neryl (2.8%) esters represents 64.5% of the total oil. In addition to the esters, wormwood root oil contained only hydrocarbon monoterpenes [acyclic (2.3%), p-menthane (7.5%), pinane (0.8%), and thujane (0.2%) type skeletons] and no free (not bound in an ester) oxygenated monoterpenes and no sesquiterpenes at all. This is unusual knowing that monoterpene formation is bound to the presence of active chloroplasts, and the latter cannot be expected in underground plant organs. Contrary to the oil isolated from aerial parts of the same herb, where thujane type monoterpenes were far most prevailing (altogether 76.9% of total oil), a single thujane type compound found in root oil was sabinene but in a very small amount (0.2% of total oil). This difference could be of great importance, since thujone is related with many malicious attributes (2, 31). Toxicological and dermatological studies of linalyl esters, agents that are known and widely used in fragrance and, in some cases, flavor ingredients, utilized in decorative cosmetic, fine fragrances, and other toiletries, as well as in noncosmetic products (32-34), unlike those of thujone, have not revealed that the use of these compounds would have serious effects on human health (32-35).

The composition of Aa3R root essential oil seems to be intermediate between those isolated from normal (main compound was α -fenchene) and transformed (only constituents reported were neryl esters) roots of *A. absinthium* investigated by Kennedy et al. (6) and is in agreement with their proposed hypothesis on the maturity of the root system as the factor governing the oil compositions. However, because they have identified only 62% of the normal and 53% of the transformed root oil components, these hypotheses (maturity of root system or translocation effects) should be taken with some reserve. It is especially difficult to accept the hypothesis based only upon translocation effects knowing that β -thujone, the component that



Figure 3. Possible mechanism of *cis*- and *trans*-linalooloxide (1 and 2) formation from linalool (7) and β -myrcene (8) [via 6,7-epoxymyrcene (9) as a reaction intermediate]. Path a: Markovnikov's addition of water to C3 methylenic double bond and subsequent epoxide opening by the nucleophilic attack of the formed alcohol. Path b: Epoxyde ring opening as a result of the nucleophilic attack of water in position 7 and subsequent furanoid ring closure.

Table 2.	Previously	Reported	Data or	the <i>i</i>	Α.	absinthium	Essential	Oil ^a
Composit	tion from D	ifferent Co	ountries ^b					

	% (w/w) in essential oil									
component	1	2	3	4	5	6	7	8		
sabinene	5.5°	0.1	d	0.3	tre	0.8–9.8	7.4	_		
β -myrcene	0.1			1.2	tr	6.8–20	33.2	0.2		
α-thujone	0.4				tr	2.8–16	20.8	0.2		
linalool	5.9	1.6	10.5	7.4	1.7	2.1-5.1	1.5	0.2		
β -thujone	26.0				1.3	5.5-12.8	13.7	0.5		
cis-6,7-epoxyocimene	24.1	30.0		31.1	25	0-6.7				
trans-6,7-epoxyocimene	1.9	2.3		4.0		0-0.7				
camphor					17	0-0.2		1.4		
terpinene-4-ol	1.0		12		tr	0.2-0.5	0.4	1.8		
cis-chrysanthenyl acetate	1.2	15.5		43.4	22			0.1		
bicyclo[2.2.1]hept- 2-en-7-ol		8.5								
α-phellandrene	0.2		50.5		tr	0.2-0.7	0.3			
	12				+r	0.4-21.3	66			
B converbullence	4.5		1 1 2	06	u 1/	15 15	0.0			
chamazulono	10		4.12	0.0	1.4	02 08	0.0	17 9		
canvonbyllene ovide	1.0	28	0.52		10.0	0.2-0.0		17.0		
	0.5	2.0	0.52		10.0			9.0		
								5 1		
neryl propanoate		3.9				0.52–1.3 ^f		J.1		

^{*a*} Essential oils were obtained from plant material collected at the full bloom from aerial parts of plant. ^{*b*} Key: 1, Croatia (18); 2, France (18); 3, Egypt (19); 4, Spain (20); 5, Italy (21); 6, Siberia (22); 7, Republic of Bashkortostan (23); and 8, Turkey (24). ^{*c*} The four most dominant components in every oil are given in bold. ^{*d*} Component not identified. ^{*e*} Compound found in traces. ^{*f*} Other esters of nerol were present in much higher amounts.

represents up to 63.4% of the oil isolated from the aerial parts of wormwood collected at Nišava banks, was not found in the corresponding root oil at all. It is reasonable to expect that volatiles found in the aerial plant parts would be also present, in some amount, in the root system if translocation channels are present and the transport works both ways.

The main components of mugwort root oil were neryl 2-methylbutanoate (13.2%), β -eudesmol (10.0%), and bornyl



Figure 4. Esters found in the *A. absinthium* essential oils from Mokra: 10-(Z)-nuciferyl propanoate, 11-(E)-nuciferyl 2-methylpropanoate, and 12-(Z)-nuciferyl butanoate.

3-methylbutanoate (8.4%). None of these substances was found in the oil obtained from aerial parts of this plant species. Worthy of mention is the fact that, contrary to the other examined oils, this one was characterized with a high percentage of sesquiterpenes (55.1%) and that esters prevailed in the monoterpenoid fraction (29.7%). Another interesting characteristic of the root oil was also the presence of the rather unusual compounds presented in **Figure 5**: petasitene (**13**), albene (**14**), and triquinane sesquiterpenes, 7 α -silphiperfol-5-ene (**15**), silphiperfol-6-ene (**16**), modhephene (**17**), pethybrene (**18**), presilphiperfol-7-ene (**19**), silphin-1-ene (**20**), α -isocomene (**21**), and β -isocomene (**22**). Triquinane sesquiterpenes were previously also found in some other species from the *Artemisia* genus (*36*, *37*).

In the oil isolated from aerial parts of *A. vulgaris*, the cooccurrence of β -caryophyllene and caryophyllene oxide, and the total lack of triquinane sesquiterpenes, suggests that the only fate of β -caryophyllene is the oxidation to caryophyllene oxide. In the root oil, however, caryophyllene oxide was not detected,

Table 3.	Previous	ly Report	ed Data	on the	e A. vu	lgaris E	essential	Oila
Composi	tion from	Different	Countrie	es ^b				

			% (w	/w) in essential oil					
component	1	2	3	4	5	6	7	8	
α-thujene	2.1	0.8	0.4				0.2	4.1°	
α-pinene	2.0	53.7	1.3	d		15.1	7.6	0.1	
camphene	9.1	1.8	4.2				2.0	3.9	
β -pinene	2.1	7.4	1.8				11.7	1.0	
β -myrcene	0.1	8.8	1.3				1.9		
α-phellandrene	tr ^e	1.0	17.3			6.3			
1,8-cineole	3.9	1.0	3.6			11.7	24.1		
artemisia alcohol	tr							4.5	
α-thujone	tr				0.4	8.5		1.3	
trans-chrysanthenol		13.1							
β -thujone	0.1	0.5			1.2	20.8			
camphor	47.7	0.5	tr		9.2	8.7	13.2	38.7	
lyratol			15.1						
trans-verbenol	7.0								
isoborneol							0.3	8.2	
borneol	0.4	0.6			8.9	2.4	2.1		
α-terpineol	0.3		0.5			0.2	7.5		
verbenone	8.6		tr						
γ-elemene			8.8						
humulene			8.8			0.2			
isobornyl 2-methylbutyrate				5.3					
caryophyllene oxide	2.2		tr	31.1	2.3	0.6	0.3		
2-heptadecanone				5.1					
hexadecanoic acid				6.3					
trans-isoelemicin			15.1						

^a Essential oils were obtained from plant material collected at the full bloom from aerial parts of the plant. ^b Key: 1, Italy (21); 2, Republic of Bashkortostan (23); 3, Egypt (25); 4, Cuba (26); 5, France (27); 6, Croatia (27); 7, Vietnam (28); and 8, India (29) (oil obtained from flowers). ^c The four most dominant components in every oil are given in bold. ^d Component not identified. ^e Compound found in traces.

but instead, considerable amounts of sesquiterpenes 15-22 (Figure 5) were present. Triquinane sesquiterpenes also originate from β -caryophyllene (38) and are thus the products of a biosynthetic branch that could be considered as an adversary to one resulting in caryophyllene oxide formation. This suggests that separate and distinct enzymatic pathways operate in the root system and the aerial parts of the plant.

Antimicrobial Activities. The oils and individual terpenes identified in the oils were tested at two doses (1:10 and 1:30 dilution, 50 μ L), and the results of the antimicrobial assays are given in Table 4. Essential oils isolated from both A. absinthium populations, collected at Mokra and Nišava banks (Aa1 and Aa3 oils, respectively), showed a high inhibitory effect on bacterial and fungal growth, with Aa3 being slightly more active against all tested microorganisms, except in the cases of Escherichia coli and Staphylococcus aureus. This difference could be the consequence of a notably higher β -thujone content in the oil from A. absinthium gathered at Nišava riverside (64.4%) as compared to the oil from Mokra (19.8%). Both thujone isomers showed high antimicrobial activity (Table 4) corroborating this assumption, and in general, it is thought that thujones are recognized as wormwood oil active principles affecting microbial growth (18). However, the observed high activity of Aa1 oil could be correlated as well to the presence of additional two known active constituents, linalool and β -caryophyllene, which also showed good results in the present antimicrobial testing (Table 4). In most cases, A. absinthium (Mokra) oil with prolonged storage time (Aa2) was more active then Aa1 oil. Exceptions include the activity against E. coli, Candida albicans, and S. aureus. Aa2 as compared to Aa1 oil had a considerably higher content of 1,8-cineole, a well-known antimicrobial agent (39), although our findings did not confirm this. A higher



Figure 5. Some unusual compounds found in the root oil from *A. vulgaris*: **13**, petasitene; **14**, albene and triquinane sesquiterpenes; **15**, 7α -silphiperfol-5-ene; **16**, silphiperfol-5-ene; **17**, modhephene; **18**, pethybrene; **19**, presilphiperfol-7-ene; **20**, silphin-1-ene; **21**, α -isocomene; and **22**, β -isocomene.

susceptibility of *E. coli* to Aa1 as compared to Aa2 and Aa3 oils could be connected to the presence of linalool, a monoterpene alcohol apparently showing a high activity against this bacterium (**Table 4**). Although *A. absinthium* root oil (Aa3R) completely lacked thujones, it exhibited a relatively high and nonselective activity. Recent research (*40*) on the antimicrobial activity of geraniol and geranyl derivatives including esters of isobutyric, butyric, propanoic, and acetic acids suggests that the corresponding linalyl esters identified in Aa3R oil could likewise be held responsible for the observed activity of this oil. The antimicrobial action of its main components is yet to be tested, and possible synergistic effects should not be excluded. The high antimicrobial activity of this oil indicates that wormwood roots could have a potential medicinal use.

It seems that the presence of 1,8-cineole and β -thujone ensured that the oil isolated from mugworth aerial parts has a good and nonselective activity against the bacteria and fungi tested. An unexpectedly low activity of *A. vulgaris* root oil was in correlation with the absence of thujones and a rather low concentration of 1,8-cineole. Furthermore, the demonstrated static and noncidal activity of bornyl acetate (5.0%), one of the major constituents of AvR oil, is in agreement with the overall low AvR oil activity. It is reasonable to presume that bornyl 3-methylbutanoate (8.4%) would have comparable influence on microbial growth. Compounds with triquinane carbon skeletons, related to those identified in *A. vulgaris* root oil, represented in **Figure 5** (15, 16, 18, 20, 21, and 22), are reported to be biologically important sesquiterpene antibiotics (*41*). Considering all of these facts, it would be interesting to investigate the

Table 4. Antimicrobial Activity^a of the Examined Oils and Corresponding Pure Constituents

							Sample								
	micro- <i>Escherichia</i> organism ^b coli 95		Salmo enter ATCC	onella itidis 13076	Pseudo aerug ATCC	monas inosa 9027	Klebs pneum ATCC 1	<i>iella</i> oniae 10031	Staphylo aure ATCC	ococcus eus 6538	Car albi ATCC	ndida cans 10231	Aspe ni ATCC	rgillus ger 16404	
	sample	C ^c	S ^d	С	S	С	S	С	S	С	S	С	S	С	S
A. absinthium A. vulgaris	Aa1 Aa2 Aa3 Aa3R Av Av	37 27 29 30 32 16	<i>e</i> 33 36 18	24 30 28 25 25	29 37	1 25 26 28 20 31 13.5	:10 dilution 27 40 29 35 16	n 26 27 36 22 26 13.5	30 34 16	24 24 29 26 26 13.5	27 28 31 29 16	26 25 30 29 40	13.5	25 29 31 24 30	28 31
						1	:30 dilutio	n							
A. absinthium A. vulgaris	Aa1 Aa2 Aa3 Aa3R Av AvR	23 16 18 18 22.5	18 20 24 25	15 22 19 16 17	18	15 16 17 27.5	22 17 14	17 19 21 22	15 26	26 16 19 17 17	28 20 18	15 18 16 14 20		16 18 17 21	15
							Standard								
	micro- organism ^b	Escl co	herichia oli 95	Salr ent ATCO	nonella eritidis C 13076	Pseudomonas aeruginosa ATCC 9027		Klebsiella pneumoniae ATCC 10031		Staphylococcus aureus ATCC 6538		Candida albicans ATCC 10231		Aspergillus niger ATCC 16404	
	sample	C ^c	S ^d	С	S	С	S	С	S	С	S	С	S	С	S
β-caryophyllene caryophyllene oxide α-humulene linalool linalooloxide (furanoid) bornyl acetate camphor α-thujone 1,8-cineole α-pinene <i>p</i> -cymene eugenol		16 18 37 25 38 25 24 34	26 36 18	30 25 34 25 28 25 25 38	35 32 37 20	14 20 33 18 24 23 36	:10 dilutio 18 24 18 22 31 17 Antibiotics	n 21 27 21 24 23 37	16 29 30 31	14 20 24 29 31 29 31 29	17 30 17	40 22 26 27 28	21 21 34 18 39	21 34 19 26 26 26 30	18 39
			Salmon	ella	Pseud	lomonas	KI	ebsiella	S	taphylococc	us	Candid	la	Asper	gillus
micro- organism ^b	Escherichi coli 95	a	enteriti ATCC 13	dis 3076	aeru ATC	iginosa C 9027	pne ATC	eumoniae CC 10031		ATCC 6538	-	albicar ATCC 10	ns 231	nig ATCC	<i>er</i> 16404
sample	C ^c S	S ^d	С	S	С	S	С	S		С	S	С	S	С	S
doxycycline tiamulin erythromycin nystatin	29 10 27 nt	2 1 2	28 10 24 nt		27.5 11 22	nt	27 10 24	nt	28 10 20	nt		nt ^f nt nt 18		n n 17	t t t

^a Antimicrobial activities are represented as the inhibition zones, mm, including the disk diameter, 6 mm. ^b All strains were obtained from the American Type Culture Collection (MD), except for the *E. coli* 95, which was acquired from the Institute of Immunology and Virology "Torlak" (Belgrade). ^c Bacterio- and fungicidal zones. ^d Bacterio- and fungistatic zones. ^e No activity observed. ¹ Not tested.

antimicrobial activities of the pure terpenoids found in AvR oil, since possible antagonism and/or synergism of the oil constituents can be expected. Despite the fact that the root oil of *A. vulgaris* had shown the weakest inhibitory effect on microbial growth as compared to all other examined oils, the existence of other possible specific therapeutic properties of the mugwort root essential oil cannot be excluded.

Although, according to the data presented in **Table 4**, it can be concluded that the antimicrobial activity of some of the analyzed flavor compounds (in given concentrations) is comparable with those of well-known antibiotics such as erythromycin, it is very important to emphasize that flavors cannot be applied in the same way as antibiotics (orally) because of selfdosage limitations influenced by their toxicity, resorption properties, smell, and taste.

ACKNOWLEDGMENT

Tanja Mitrović is acknowledged for collecting *A. absinthium* plant material from Mokra.

LITERATURE CITED

 European Council. Council Directive (EEC) No. 88/388 on the approximation of the laws of the Member States relating to flavorings for use in foodstuff and to source materials for their production. *Off. J. Eur. Commun.* **1988**, *L184*, 61–66.

- (2) Patocka, J.; Plucar, B. Pharmacology and toxicology of absinth. J. Appl. Biomed. 2003, 1, 199–205.
- (3) Tucakov, J. Lecenje Biljem. Fitoterapija; Izdavacko preduzece Rad: Beograd, Jugoslavija, 1973; pp 402, 535–537.
- (4) Nin, S.; Afraioli, P.; Bosetto, M. Quantitative determination of some essential oil components of selected *Artemisia absinthium* plants. J. Essent. Oil Res. 1995, 7, 271–277.
- (5) Kovacevic, N. Osnovi Farmakognozije; Srpska skolska knjiga: Beograd, Srbija i Crna Gora, 2004; pp 274–276.
- (6) Kennedy, A.; Deans, S.; Svoboda, K.; Gray, A.; Waterman, P. Volatile oils from normal and transformed root of *Artemisia absinthium*. *Phytochemistry* **1993**, *32* (6), 1449–1451.
- (7) Nin, S.; Bennici, A.; Roselli, G.; Mariotti, D.; Schiff, S.; Magherini, R. Agrobacterium-mediated transformation of *Arte-misia absinthium* L. (wormwood) and production of secondary metabolites. *Plant Cell Rep.* **1997**, *16*, 725–730.
- (8) Hochmuth, D. H.; Joulain, D.; Konig, W. A. MassFinder Software and Data Bank; University of Hamburg: Hamburg, Germany, 2003.
- (9) Adams, R. P. Identification of Essential Oil Components by Gas Chromatography and Mass Spectroscopy; Allured Pub Corp.: Carol Stream IL, 2001.
- (10) Van Den Dool, H.; Kratz, P. D. A generalization of the retention index system including linear temperature programmed gas liquid partition chromatography. J. Chromatogr. 1963, 11, 463– 471.
- (11) Cavaleiro, C.; Salgueiro, L.; Barroso, J. G.; Figueiredo, A. C.; Pedro, L. G.; Fontihna, S. S.; Bighelli, A.; Casanova, J.; Looman, A.; Scheffer, J. J. C. Composition of the essential oil of *Juniperus cedrus* Webb & Berth. grown on Madeira. *Flavour Fragrance J.* **2002**, *17*, 111–114.
- (12) Radulovic, N.; Stojanovic, G.; Palic, R. Composition and atimicrobial activity of *Equisetum* arvense L. essential oil. *Phytother. Res.* 2006, 20 (1), 85–88.
- (13) NCCLS (National Committee for Clinical Laboratory Standards). Performance standards for antimicrobial disk susceptibility testing. *6th International Supplement*; NCCLS: Wayne, PA, 1997; M2-A6.
- (14) Dudareva, N.; Pichersky, E.; Gershenzon, J. Biochemisty of plant volatiles. *Plant Physiol.* **2004**, *135*, 1893–1902.
- (15) Voet, D.; Voet, J. G. *Biochemistry*; John Wiley & Sons: New York, 1995.
- (16) Schofield, L. J.; Kerton, O. J.; McMorn, P.; Bethell, D.; Ellwood, S.; Hutchings, G. J. Oxidation of α-hydroxy containing monoterpenes using titanium silicate catalysts: Comments on regioselectivity and the role of acidity. *J. Chem. Soc. Perkins Trans.* 2 2002, *8*, 1475–1481.
- (17) Dieckmann, R. H.; Palamand, S. R. Autoxidation of some constituents of hops. I. The monoterpene hydrocarbon, myrcene. *J. Agric. Food Chem.* **1974**, *22* (3), 498–503.
- (18) Juteau, F.; Jerkovic, I.; Masotti, V.; Milos, M.; Mastelic, J.; Bessiere, J. M. Composition and antimicrobial activity of the essential oil of *Artemisia absinthium* from Croatia and France. *Planta Med.* **2003**, *69*, 158–161.
- (19) Aboutabl, E. A.; El Azzouny, A. M.; El Dahmy, S. I. Constituents of the essential oil of *Artemisia absinthium* L. grown in Egypt. *J. Essent. Oil–Bear. Plants* **1998**, *1* (2–3), 82–86.
- (20) Arino, A.; Arberas, I.; Renobales, G.; Dominguez, J. B. Influence of extraction method and storage conditions on the volatile oil of wormwod (*Artemisia absinthium* L.). *Eur. Food Res. Technol.* **1999**, 209, 126–129.
- (21) Mucciarely, M.; Caramiello, R.; Maffei, M. Essential oils from some *Artemisia* species growing spontaneously in northwest Italy. *Flavour Fragrance J.* 1995, 10, 25–32.
- (22) Khanina, M. A.; Serykh, E. A.; Pokrovsky, L. M.; Tkachev, A. V. New data on chemical composition of *Artemisia absinthium* L. from Siberia essential oil. *Khim. Rast. Syr'ya* 2000, *3*, 33–40.

- (23) Khalilov, L. M.; Paramonov, E. A.; Khalilova, A. Z.; Odinokov, V. N.; Muldashev, A. A.; Baltaev, U. A.; Dzehemilev, U. M. Identification and biological activity of volatile organic compounds emitted by plants and insects. IV. Composition of vapor isolated from certain species of *Artemisia* plants. *Chem. Nat. Compd.* **2001**, *37* (4), 339–342.
- (24) Kordali, S.; Cakir, A.; Mavi, A.; Kilic, H.; Yildirim, A. Screening of chemical composition and antifungal and antioxidant activities of the essential oils from three Turkish *Artemisia* species. J. Agric. Food Chem. 2005, 53, 1408–1416.
- (25) Aboutabl, E. A.; El-Azzouny, A.; El Dahmy, S. I. Composition of the essential oil of *Artemisia vulgaris* L. grown in Egypt. J. *Essent. Oil-Bear. Plants* **1998**, 1 (1), 21–27.
- (26) Pino, J. A.; Rosado, A.; Fuentes, V. Composition of the essential oil of *Artemisia vulgaris* L. herb from Cuba. J. Essent. Oil Res. **1999**, 11, 477–478.
- (27) Jerkovic, I.; Mastelic, J.; Milos, M.; Juteau, F.; Masotti, V.; Viano, J. Chemical variability of *Artemisia vulgaris* L. essential oils originated from the Mediterranean area of France and Croatia. *Flavour Fragrance J.* **2003**, *18*, 436–440.
- (28) Thao, N. T. P.; Thuy, N. T.; Hoi, T. M.; Thai, T. H.; Muselli, A.; Bighelli, A.; Castola, V.; Casanova, J. Artemisia vulgaris L. from Vietnam: Chemical variability and composition of the oil along the vegetative life of the plant. J. Essent. Oil Res. 2004, 16 (4), 358–361.
- (29) Haider, F.; Dwivedi, P. D.; Naqvi, A. A.; Bagchi, G. D. Essential oil composition of *Artemisia vulgaris* harvested at different growth periods under Indo-gangetic plain conditions. *J. Essent. Oil Res.* 2003, *15* (6), 376–378.
- (30) Reviewing the available literature brought to our attention a paper of Fraternale et al. (42). In their work, they determined the chemical composition of Peucedanum verticillare. One of the components that they identified was linalyl 3-methyl butanoate, which had a RI of 1395 for this ester on a SPB-1 fused silica column. The RI for the same compound obtained in this study was 1484. Although the experimental conditions were to some extent different, we find them insufficient to cause such a large difference in the value of RIs. As the mentioned component was among the most abundant in A. absinthium essential oil, the confirmation of its identity was also accomplished using ¹³C NMR spectra, in addition to the GC and GC-MS analysis. Moreover, altogether, six different linalyl esters have been found in the analyzed oils, and an excellent correspondence was established between the RIs on a DB-5 column of some of the identified linalyl esters with literature data (43, 44). This strongly suggests that the identification of linalyl 3-methyl butanoate by Fraternale and co-workers (42) may need revision. The RI of their component better corresponds to that for linalyl 2-methyl propanoate or linalyl butanoate.
- (31) Hold, K. M.; Sirisoma, N. S.; Ikeda, T.; Narahashi, T.; Casida, J. E. α-Thujone (the active component of absinthe): γ-Aminobutiric acid type A receptor modulation and metabolic detoxification. *Proc. Natl. Acad. Sci. U.S.A.* 2000, 97 (8), 3826–3831.
- (32) Letizia, C. S.; Cocchiara, J.; Lalko, J.; Api, A. M. Fragrance material review on linalool. *Food Chem. Toxicol.* 2003, *41*, 943– 964.
- (33) Letizia, C. S.; Cocchiara, J.; Lalko, J.; Api, A. M. Fragrance material review on linalyl isovalerate. *Food Chem. Toxicol.* 2003, 41, 1011–1015.
- (34) Letizia, C. S.; Cocchiara, J.; Lalko, J.; Api, A. M. Fragrance material review on linalyl propionate. *Food Chem. Toxicol.* 2003, 41, 1023–1027.
- (35) Bickers, D.; Calow, P.; Greim, H.; Hanifin, J. M.; Rogers, A. E.; Saurat, J. H.; Sipes, I.; Smith, R. L.; Tagami, H. A toxicologic and dermatologic assessment of linalool and related esters when used as fragrance ingredients. *Food Chem. Toxicol.* 2003, *41*, 919–942.
- (36) Weyersahl, P.; Marschall, H.; Schroder, M.; Wahlburg, H.; Kaul, V. K. The sesquiterpene fraction of the essential oil of *Artemisia laciniata* Willd. *Flavour Fragrance J.* **1997**, *12*, 315–325.

- (37) Marco, J. A.; Sanz-Cervera, J. F.; Morante, M. D.; Garcia-Lliso, V.; Valles-Xirau, J.; Jakupovic, J. Tricyclic sesquiterpenes from *Artemisia chamaemelifolia. Phytochemistry* **1996**, *41* (3), 837– 844.
- (38) Coates, R. M.; Ho, Z.; Klobus, M.; Wilom, S. Stereochemistry and reactions of presilphiperfolanol: A branch point marker in triquinane sesquiterpene biogenesis. J. Am. Chem. Soc. 1996, 118, 9249–9254.
- (39) Pattnaik, S.; Subramanyam, V. R.; Bapaji, M.; Kole, C. R. Antibacterial and antifungal activity of aromatic constituents of essential oils. *Microbios* **1997**, *89* (358), 39–46.
- (40) Jirovetz, L.; Buchbauer, G.; Schmidt, E.; Denkova, Z.; Stoyanova, A. S.; Nikolova, R.; Geissler, M. Purity and antimicrobial activities of geraniol and various geranyl derivatives. 36th International Symposium on Essential Oils, Budapest, 2005; p 41, L-5-02.
- (41) Harrington-Frost, N. M.; Pattenden, G. A new synthesis of pentalene using a novel tandem cyclisation involving ketene radical intermediates. *Tetrahedron Lett.* 2000, 41, 403–405.

- (42) Fraternale, D.; Giamperi, L.; Ricci, D.; Manunta, A. Composition of the essential oil of *Peucedanum verticillare*. *Biochem. Syst. Ecol.* 2000, 28, 143–147.
- (43) Baranauskiene, R.; Venskutonis, P. R.; Viskelis, P.; Dambrauskiene, E. Influence of nitrogen fertilizers on the yield and composition of thyme (*Thymus vulgaris*). J. Agric. Food Chem. 2003, 51, 7751–7758.
- (44) Javidnia, K.; Miri, R.; Kamalinejad, M.; Nasiri, A. Composition of the essential oil of *Salvia mirzayanii* Rech. f., & Esfand from Iran. *Flavour Fragrance J.* 2002, *17*, 465–467.

Received for review January 16, 2006. Revised manuscript received April 28, 2006. Accepted May 8, 2006. We thank the Ministry of Science and Environmental Protection of the Republic of Serbia for financial support (Project 142054B).

JF060123O