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Distribution of Volatile Sulfur Compounds in an Interspecific Hybrid between Onion (*Allium cepa* L.) and Leek (*Allium porrum* L.)

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Volatile sulfur compounds of an interspecific hybrid between *Allium cepa* and *Allium porrum* were analyzed by gas chromatography (GC) and GC/mass spectrometry prior to isolation of the individual oils by a simultaneous distillation/extraxtion (SDE) method. Furthermore, the aroma profiles of various onion and leek cultivars were investigated. Major volatile components detected in onion were 2-methyl-2-pentenal, (*E*)-methyl 1-propenyl disulfide, methyl propyl trisulfide, and propanethiol, whereas dipropyl trisulfide, dipropyl disulfide, and (*E*)-propenyl propyl disulfide predominated in leek oils. According to the higher amount of leek chromosomes in the cell nucleus, the percentages of the measured sulfur volatiles in the hybrid material correspond more to the leek than to the onion flavor profile. Discrimination analysis was successfully applied to classify the three predefined *Allium* groups. It has been found that at minimum five of the most important aroma volatiles were necessary to receive sufficient differentiated clusters without any overlapping areas. This graphical display demonstrates very clearly the influence of the genetic background of the parent varieties on the sulfur volatile composition in the *Allium* hybrid.

Keywords: Interspecific allium hybrid; onion; leek; GC analysis; volatile aroma compounds; discrimination analysis

INTRODUCTION

Allium vegetables, such as onion, garlic, leek, and chives, are traditionally known for their fresh flavors but also for their antibacterial and fungicidal properties. The characteristic flavor of Allium species is due to the volatile oil, which consists mainly of sulfur compounds. When the cells of the Allium plants are ruptured, a reaction between the enzymes (lyase, alliinase) and the individual flavor precursors takes place, resulting in the formation of an individual flavor profile (Tressl et al., 1975; Whitaker, 1976; Boelens et al., 1980; Kuo et al., 1990; Lancaster and Boland, 1990; Stephani and Baltes, 1992) as presented generally in Figure 1. It has been found that these sulfur volatiles play also a defensive role against animal pests and microorganisms (Block, 1985; Carson, 1987).

On the basis of the results from case-control studies performed in different parts of the world, *Allium* vegetable consumption has been found to be associated with a reduction of stomach cancer risk in humans (You et al., 1989; Hansson et al., 1993; Graham et al., 1990). It has been proved that some of the organosulfur compounds present in raw and cooked *Allium* vegetable act as antimutagens in in-vitro models and even as anticarcinogens in in-vivo test systems (Dorant et al., 1993).

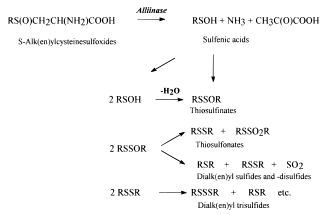


Figure 1. General scheme for the formation of alk(en)yl sulfur components in *Allium* plants.

In the past the composition and formation of volatiles in onion, garlic, leek, and also in ornamental Allium species have been extensively studied (Brodnitz et al., 1969; Freeman, 1975; Yu et al., 1989). It has been found that the characteristic aroma impressions recognized in the individual *Allium* species or varieties are predominantly attributed to a different pattern of nonvolatile alk(en)ylcysteinsulfoxides in the plant. Depending on the species the alk(en)yl groups are mainly a combination of propyl, 1-propenyl, allyl, and methyl substituents. So far as *Allium* species have been analyzed, they can be divided in the following three groups (Freeman and Whenham, 1975): high proportions of propyl/ propenyl cysteinsulfoxide (e.g., onions), high proportions of allyl cysteinsulfoxide (e.g., garlic), and high methyl cysteinsulfoxide portions (e.g., ornamental Allium spe-

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cies). According to these results it must be assumed that the formation of the flavor precursors is genetically controlled. Up to now no single *Allium* species has been described in the literature to produce different sulfur amino acids as the major flavor precursors.

Recently, the aroma composition of an interspecific hybrid between Allium fistolosum and Allium tuberosum, formed by embryo culture, has been investigated (Kobayashi et al., 1997). According to the published results there is only little difference between the compositions of *A. tuberosum* and the hybrid; however, much more of the allyl-substituted sulfur components occur in the hybrid plant, which explains its stronger garlic-like aroma.

The aim of the study reported here is to characterize an interspecific hybrid of Allium cepa and Allium porrum with special respect to its very distinctive flavor profile, influenced by the genetic background of the parent plants.

MATERIALS AND METHODS

Cultivation and Sample Preparation of *Allium* **Plants.** Onion varieties (Hystar, Summit, Stuttgarter Riesen, Vitesso, Romeo, Bristol, Macho, Boston, Trefford) and leek cultivars (Upton, Erik, Lanzelot, Parton, Porbella, Gloria, Nepal, Profina) were cultivated in the experimental garden of the Landesversuchsanstalt Gartenbau at Ditfurt (Germany). The leek variety Pollux was grown under comparable growing conditions in the experimental garden of the Federal Centre for Breeding Research on Cultivated Plants in Quedlinburg (Germany). The interspecific hybrid 99/1 was obtained from the cross of onion Summit with leek Pollux (Peterka et al., 1997).

Normal farming practice was employed. Onion bulbs and leek plants were harvested from July to August. Approximately five to eight randomly taken ripe onions formed one sample. In the case of leek cultivars and hybrid material, the edible parts of approximately four to five plants, freshly homogenized prior to analysis, were taken for one investigation. Two identical replica samples of each variety were analyzed under the same conditions.

The washed Allium samples were cut into pieces and macerated carefully. A sample amount of ~120-160 g was divided in two equivalent portions, each of them transferred into a 250 mL round-bottom flask and blended with 100 mL of water. The volatile constituents were extracted for 3 h at boiling temperature in a Likens-Nickerson water distillation/ solvent extraction (SDE) apparatus. n-Pentane (15 mL, purity = 99.0%) was used as extracting solvent. One milliliter of a 2.5×10^{-3} M internal standard solution (6-methyl-5-hepten-2-one in *n*-pentane) was added to the received aroma extract.

The following authentic reference samples were obtained from commercial suppliers (Oxford Chemicals, Harlepool, U.K., and Dragoco Gerberding & Co. AG, Holzminden, Germany): allyl propyl disulfide, allyl propyl trisulfide, 2,5-dimethylthiophene, dimethyl trisulfide, dipropyl disulfide, dipropyl trisulfide, methyl propyl disulfide, and methyl propyl trisulfide.

Gas Chromatographic and Mass Spectrometric (GC/ MS) Analysis. A Hewlett-Packard chromatograph model HP 5890 Series II equipped with an FID and a fused silica capillary column [ĤP ÎNNOWAX with a 0.5 μ m bonded PEG phase, $60 \text{ m} \times 0.25 \text{ mm}$ (i.d.)] was used to analyze the isolated volatiles. The operating conditions were as follows: injector temperature, 250 °C; detector temperature, 280 °C; H₂ flow rate, 1 mL/min; oven temperature, 80-220 °C raised at 4 °C/ min; injection volume, 1 μ L; split ratio, 1:40. Linear retention indices were calculated against C₇-C₂₀ n-paraffins as references according to the method of van den Dool and Kratz (1963).

Mass spectral analyses were carried out in a Hewlett-Packard gas chromatograph (HP 5890 Series II) directly coupled to a Hewlett-Packard MSD, model HP 5972. Opera-

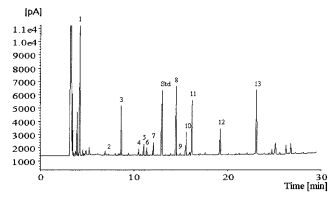


Figure 2. Gas chromatogram of the oil obtained from an interspecific Allium hybrid (A. cepa × A. porrum) by SDE procedure. Peak identification is according to Table 1. Internal standard (Std): 6-methyl-5-hepten-2-one.

tional parameters were as follows: carrier gas, He, 1 mL/min; ionization voltage, 70 eV; source temperature, 195 °C; scan region, 27-300 m/z.

The GC separation conditions applied were the same as described above.

Statistics. Discrimation analysis of the collected analytical data was carried out using the statistical software program SYSTAT 7.0 of SPSS Inc., Chicago, IL.

RESULTS AND DISCUSSION

The oils received from the *Allium* hybrid and from different onion and leek varieties have been analyzed to identify the most important aroma components as well as to describe the hybrid species on the basis of the characteristic profile of volatile sulfur substances. The individual oil constituents are identified by comparing their mass spectra and retention indices with those of reference substances.

Representing the high resolution of the separated components, Figure 2 shows a typical gas chromatogram of an *Allium* hybrid oil received by the SDE procedure. It has to be mentioned here that the sample preparation process was performed very accurately and reproducibly due to the fact that the rates of formation and half-lives of the volatile di-, tri-, and tetrasulfides vary widely (Kallio et al., 1994). The concentrations of the individual sulfur volatiles identified in the studied Allium oils are presented in Table 1. As can be seen, the leek and onion varieties, cultivated under the same growing conditions, do not show any strong differences in their flavor profiles within each species.

The composition of the investigated leek oils corresponds very well with the data already indicated in earlier studies (Schreyen et al., 1976; Stephani and Baltes, 1992). Main components in the leek oil are dipropyl trisulfide, dipropyl disulfide, propanethiol, and (*E*)-1-propenyl propyl disulfide. In a similar way the composition of onion oils received from different varieties shows no significant differences concerning the flavor profile, so the variability of the volatile sulfur components is also here comparatively small within the different cultivars.

Only the variety Stuttgarter Riesen indicates relatively high contents of 2-methyl-2-pentenal, whereas the variety Hystar contains clearly lower contents of this flavor material. As main constituents in the onion oil, 2-methyl-2-pentenal, (*E*)-methyl 1-propenyl disulfide, methyl propyl trisulfide, and propanethiol are detected.

Concerning the GC profile the aroma extract of the Allium hybrid presents more similarity with leek than

Table 1. Composition a of Volatile Compounds Detected in Different Onion and Leek Varieties as well as an Interspecific Hybrid of A. cepa \times A. porrum

13, dipropy trisulfide, RI = 1713	62	72	62	40	46	∞	20	98	96		555		1248	1230	2044	1460	1672	1463	1609	1389	1843
12, methyl propyl trisulfide, RI = 1576	263	378	193	209	189	92	117	214	210		301		428	851	993	182	813	1303	805	006	519
11, (E)-1- propenyl propyl disulfide, $RI = 1473$	93	107	84	99	75	37	83	104	107		383		682	558	957	722	870	884	1056	845	1027
10, (Z)-1- propenyl propyl disulfide, $RI = 1450$	58	93	59	54	54	31	56	74	72		187		377	289	443	365	360	330	261	360	308
9, dimethyl trisulfide, RI = 1430	245	375	124	217	158	82	47	112	66		32		24	112	88	0	72	252	102	107	27
8, dipropyl disulfide, RI = 1413	36	26	35	5	21	0	29	42	46		348		545	896	954	823	918	1016	1298	991	1283
7, (E)-methyl 1-propenyl disulfide, RI = 1322	245	419	178	262	206	135	118	172	153		118		154	230	297	49	223	453	234	339	143
6, (Z)-methyl 1 -propenyl disulfide, $RI = 1298$	179	298	133	196	156	107	88	127	214		80		06	131	160	30	130	229	06	178	09
5, dimethylthiophene, RI = 1285	114	173	136	122	135	118	167	136	131		93		148	106	114	126	84	75	41	97	47
4, methyl propyl disulfide, RI = 1263	42	09	27	29	24	10	18	36	31		52		06	256	237	45	170	569	566	241	161
3, 2-methyl- 2-pentenal, RI = 1190	424	642	510	420	520	401	589	490	509		326		336	378	412	410	301	392	135	358	178
2, dimethyl disulfide, RI = 11110	63	121	31	52	33	16	16	35	25		6		24	92	52	0	30	80	36	22	14
$1,^{b}$ 1-proparethiol, RI c = 855	174	243	168	161	159	104	217	258	246		502		861	932	1099	1083	851	663	602	1028	885
	onion cv. Hvstar	Stuttgarter Riesen	Vitesso	Romeo	Bristol	Macho	Boston	Trefford	Summit	hybrid	$Summit \times Pollux$	leek cv.	Pollux	Upton	Erik	Lanzelot	Parton	Porbella	Gloria	Nepal	Profina

^a Amounts calculated as micrograms per 100 g of fresh material by an internal standard method, average of two samples. ^b Peak number in Figure 2. ^c Retention index.

Figure 3. DA scores of leek (\bigcirc) and onion (\times) varieties as well as samples of the interspecific *Allium* hybrid (+) based on the following volatile aroma components: dipropyl disulfide, (Z)-1-propenyl propyl disulfide, (E)-1-propenyl propyl disulfide, dipropyl trisulfide, propanethiol, and 2-methyl-2-pentenal.

with onion varieties. Major sulfur components of the oil are dipropyl trisulfide, propanethiol, (E)-1-propenyl propyl disulfide, and dipropyl disulfide. In accordance with the composition of leek the percentage of 2-methyl-2-pentenal is comparatively low. This flavor substance, occurring in onion in relatively large amounts, is formed from the amino acid 1-propenyl-L-cysteine sulfoxide by the enzymatic reaction with alliinase (Freeman and Whenham, 1975). Thus, it can be assumed that the different genetic backgrounds of leek and onion are responsible for the different biochemical production yields of the individual alk(en)yl cysteine sulfoxides in the investigated *Allium* plants.

Discrimination analysis (DA) involving all 13 detected volatile compounds analyzed in this study gives some visual evidence of the ability to differ between onion and leek cultivars and the hybrid material as clearly separated factor groups. A differentiation of these three groups can be also achieved when only six of the most important aroma components are involved in the statistical calculations (Figures 3 and 4). The graphical display of the resulting scores using factors 1 and 2

shows that all *Allium* hybrid samples build up a single cluster without any overlapping areas with the clusters of the parents varieties.

It should be noted here that all points in the DA plot correspond to physically different samples, and therefore these samples cover a wide range in terms of analytical reproducibility, variety, time of harvesting, and cultivation influences. However, these factors seem to introduce a variability into the data set, which is not as important as the variability from the different Allium species and the hybrid material, respectively. Regarding the percentages of the individual sulfur volatiles, it can be seen very easily that the hybrid material corresponds more to the leek than to the onion profile. As already mentioned above, this is attributed to the fact that the genetic information coming from the leek supplies a larger contribution for the biogenesis of the analyzed volatile flavor materials or precursors than the onion genome does. The hybrid plants are allotriploid, which means they possess 3x = 24 chromosomes in the cell nucleus. From the allotetraploid leek here come 16 chromosomes, whereas the diploid onion plant contributes to the hybrid only 8 chromosomes.

The results presented in this study clearly demonstrate the potential for DA to facilitate the differentiation between similar *Allium* species or genotypes on the basis of the most characteristic volatile sulfur components. It can be assumed that interspecific *Allium* hybrids are a useful approach to increase the genetic variability of *Allium* flavor characteristics for further improvement in breeding.

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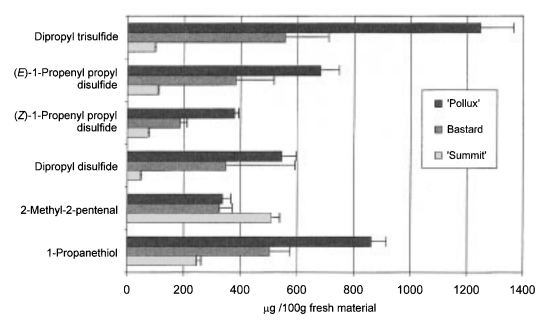


Figure 4. Contents [micrograms per 100 g of fresh material] with confidence limits of most important aroma compounds in the *Allium* hybrid and the relating parent varieties Summit and Pollux.

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Registry No. Supplied by the Author: 1-propanethiol, 75-33-2; dimethyl disulfide, 624-92-0; 2-methyl-2-pentenal, 623-36-9; methyl propyl disulfide, 2179-60-4; 2,5-dimethyl-thiophene, 638-02-8; (*Z*)-methyl 1-propenyl disulfide, 23838-19-9; (*E*)-methyl 1-propenyl disulfide, 23838-19-9; dipropyl disulfide, 629-19-6; dimethyl disulfide, 624-92-0; dimethyl trisulfide, 3658-80-8; (*Z*)-1-propenyl propyl disulfide, 23838-20-2; (*E*)-1-propenyl propyl disulfide, 23838-21-3; methyl propyl trisulfide, 17619-36-2; dipropyl trisulfide, 6028-61-1; 6-methyl-5-hepten-2-one, 110-93-0.

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