

<b>Zeitschrift:</b>	Mitteilungen aus dem Gebiete der Lebensmitteluntersuchung und Hygiene = Travaux de chimie alimentaire et d'hygiène
<b>Herausgeber:</b>	Bundesamt für Gesundheit
<b>Band:</b>	78 (1987)
<b>Heft:</b>	2
<b>Artikel:</b>	GC/MS characterization of pyrrolizidine alkaloids in some species of asteraceae
<b>Autor:</b>	Schmid, P. / Lüthy, J. / Zweifel, U.
<b>DOI:</b>	<a href="https://doi.org/10.5169/seals-982985">https://doi.org/10.5169/seals-982985</a>

### Nutzungsbedingungen

Die ETH-Bibliothek ist die Anbieterin der digitalisierten Zeitschriften auf E-Periodica. Sie besitzt keine Urheberrechte an den Zeitschriften und ist nicht verantwortlich für deren Inhalte. Die Rechte liegen in der Regel bei den Herausgebern beziehungsweise den externen Rechteinhabern. Das Veröffentlichen von Bildern in Print- und Online-Publikationen sowie auf Social Media-Kanälen oder Webseiten ist nur mit vorheriger Genehmigung der Rechteinhaber erlaubt. [Mehr erfahren](#)

### Conditions d'utilisation

L'ETH Library est le fournisseur des revues numérisées. Elle ne détient aucun droit d'auteur sur les revues et n'est pas responsable de leur contenu. En règle générale, les droits sont détenus par les éditeurs ou les détenteurs de droits externes. La reproduction d'images dans des publications imprimées ou en ligne ainsi que sur des canaux de médias sociaux ou des sites web n'est autorisée qu'avec l'accord préalable des détenteurs des droits. [En savoir plus](#)

### Terms of use

The ETH Library is the provider of the digitised journals. It does not own any copyrights to the journals and is not responsible for their content. The rights usually lie with the publishers or the external rights holders. Publishing images in print and online publications, as well as on social media channels or websites, is only permitted with the prior consent of the rights holders. [Find out more](#)

**Download PDF:** 23.08.2025

**ETH-Bibliothek Zürich, E-Periodica, <https://www.e-periodica.ch>**

## GC/MS Characterization of Pyrrolizidine Alkaloids in some Species of Asteraceae

P. Schmid, J. Lüthy\*, U. Zweifel, A. Bettschart\*\* and Ch. Schlatter

Institute of Toxicology, Swiss Federal Institute of Technology and  
University of Zurich, Schwerzenbach

### Introduction

The pyrrolizidine alkaloids (PAs) are a class of compounds occurring in a large variety through a wide range of the flora. Until now more than 200 different PAs have been isolated and identified, mainly in plants of the family *Compositae*. Within the *Compositae* the occurrence of PAs is restricted to just two of the tribes, the *Eupatoriae* and the *Senecioneae* (1). An actual review on the state of research on the PAs is given by Robins (2). In cases of accidental intake of certain PA containing plants by humans the alkaloids have shown high hepatotoxicity (3, 4). Similar observations are reported on poisonings of cattle by hay containing *Senecio alpinus* (5). At comparable doses the PAs cause similar symptoms in laboratory animals (mice and rats) (6). In long-time studies at subacute dose levels carcinogenic effects have been observed with the PAs monocrotaline and lasiocarpine (7). Our own investigations have demonstrated that certain medical herbs and their extracts used in medicaments contain PAs (8, 9).

In the present paper we report on the analytical investigation of PAs from different plant species with gas chromatography/mass spectrometry (GC/MS) using electron impact (EI) as well as positive and negative ion chemical ionization (CI). This study is also part of a project on using the PA content and pattern of alpine plants for their chemotaxonomic characterization.

\* heutige Adresse: Bundesamt für Gesundheitswesen, Bern

\*\* Engel-Apotheke, CH-8840 Einsiedeln

## Experimental

### Plant samples

Aerial parts of *Homogyne alpina* (L.) Cass., *Adenostyles Alliariae* (Gouan) Kerner, *Senecio Doronicum* L. and *Senecio Fuchsii* Gmelin (possibly a bastard of *Senecio Fuchsii* and *Senecio nemorensis* which occur frequently in Switzerland) were collected near Einsiedeln in the Swiss prealpine region (samples of these plants are stored in a herbarium by A. B.). *Eupatorium cannabinum* L. (*Herba Eupatorii cannabini conc.*, Dixa AG, CH-9014 St. Gallen, Switzerland) is a commercial sample of unknown origin and botanically not further characterized. Treatment of the plant material included extraction of the PAs, purification and reduction of the N-oxides as described earlier (8, 10).

### Gas Chromatography/Mass Spectrometry

Gas chromatograph: Carlo Erba 4160

Glass capillary column: 20 m × 0.30 mm, SE-54 film thickness 0.15 µm

Carrier gas: hydrogen, linear flow velocity 50 cm/s

Injection: 1 µl, split ratio 1:10

Temperature of injector: 250 °C

Temperature programme: injection at 140 °C, 4 °C/min up to 260 °C

Mass spectrometer: Finnigan 4510 with data system Incos

Temperature of ion source: 150 °C

Reagent gas: methane, ion source pressure 0.60 torr

Mass range: m/e 41 to 450, cycle time 1.5 s.

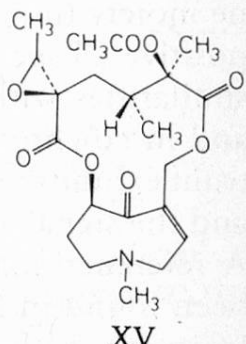
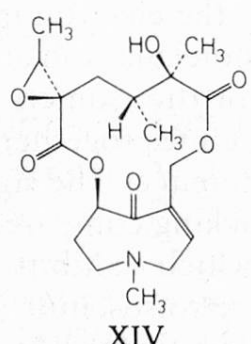
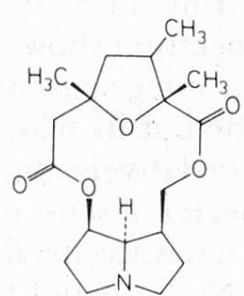
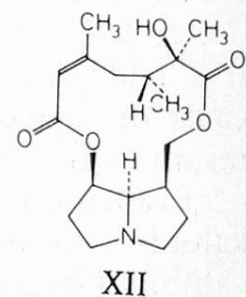
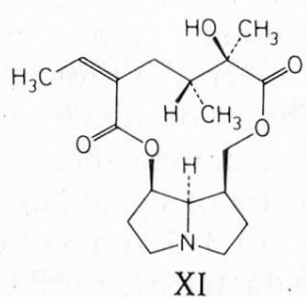
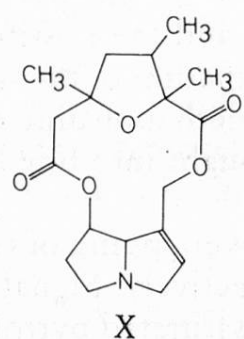
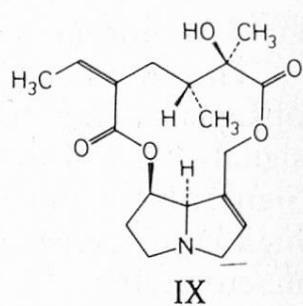
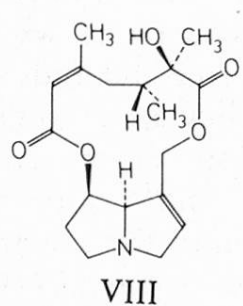
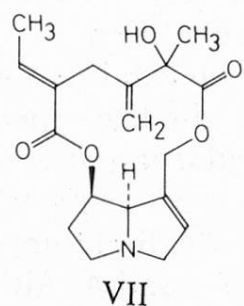
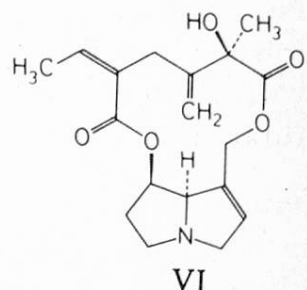
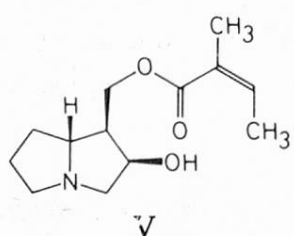
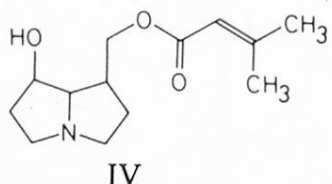
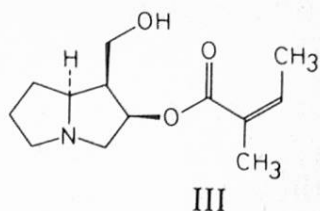
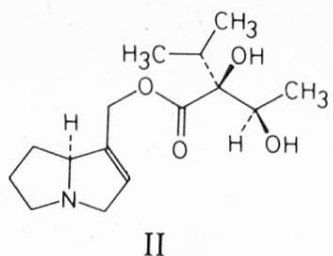
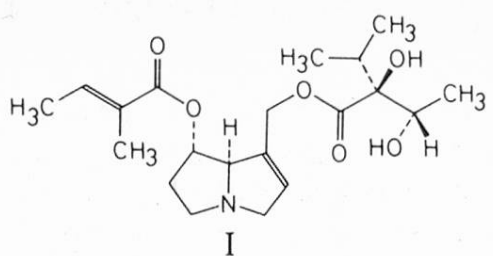
## Results and Discussion

### *Adenostyles Alliariae*

The chromatogram shows 5 signals due to PAs (see table 1). The mass spectra of peaks No 1 and 2 have revealed senencionine (IX) and seneciphylline (VI) respectively (fig. 1); identity was confirmed by co-gas chromatography of the two original compounds isolated from *Senecio alpinus* (11); signal No 3 is due to platiphylline (XI, M<sup>+</sup> 337). The mass spectrum of peak No 4 indicates a molecular ion at m/e 333, but a fragment pattern corresponding to an unidentified seneciphylline isomer, for example spartiodine (VII) (12). Mass spectrum and chromatographic behaviour of peak No 5 agree with the corresponding data of 12-acetyl-seneciphylline, a compound which has already been tentatively identified in *Senecio alpinus* (10). In an earlier investigation seneciphylline and platiphylline have been identified in a sample of *A. Alliariae* of Russian origin (13).

Table 1. Mass spectra of pyrrolizidine alkaloids

Nr.	RT(min)	Amt.	Identification (cc: co-chromatography; ms mass spectrum) or mass spectrum
<i>Adenostyles Alliariae</i>			
1	15.96	++	Senecionine cc, ms
2	16.25	+++	Seneciphylline cc, ms
3	16.73	+	Platiphylline cc, ms
4	16.84	+	M = 333
5	18.60	++	Acetylseneciphylline ms
<i>Homogyne alpina</i>			
1	8.74	++	M = 237
2	22.20	+++	Petasitenine cc, ms
3	25.93	++	Neopetasitenine ms
<i>Senecio Doronicum</i>			
1	4.77	+++	239 (2, M), 140 (21), 139 (24), 109 (9), 108 (100), 84 (9), 83 (30), 82 (4), 81 (4), 80 (12), 70 (15), 68 (4), 55 (23), 53 (5), 43 (6), 42 (7), 41 (9).
2	5.13	++	Macrophylline (9-angelylmacronecine) ms
3	5.53	++	see signal no. 1
4	5.69	++	see signal no. 2
5	10.95	+	see text
6	11.32	++	M = 335
7	11.88	+	M = 337
8	12.07	++	Doroneneine ms
9	12.27	++	Bulgarsenine ms
10	12.50	+	M = 337
11	12.86	++	M = 337
12	13.03	+	M = 353
13	14.76	+	M = 353
<i>Senecio Fuchsii</i>			
1	10.73	+	7-Angelylretronecine ms
2	11.15	+	see signal no. 3
3	12.00	+++	Senecioylplatynecine, fuchsisenecionine ms
4	19.00	+	M = 319
5	22.31	+	M = 337
6	22.88	+	Platiphylline cc, ms
7	24.35	+++	337 (0.1, M), 237 (12), 222 (12), 140 (40), 139 (29), 138 (100), 123 (30), 122 (52), 121 (7), 120 (5), 108 (6), 97 (8), 96 (32), 95 (35), 83 (29), 82 (70), 80 (6), 55 (43), 53 (12), 43 (7), 41 (19).
8	24.62	++	M = 337
9	24.92	+	see signal no. 9
10	25.22	++	M = 337
11	25.50	+	see signal no. 10
<i>Eupatorium cannabinum</i>			
1	23.75	++	Supinine ms
2	25.69	+++	381 (M), 336 (5), 238 (4), 221 (35), 220 (68), 141 (9), 138 (6), 136 (49), 122 (5), 121 (93), 120 (100), 119 (54), 118 (8), 95 (7), 94 (36), 93 (58), 83 (20), 82 (8), 80 (15), 67 (8), 55 (38), 53 (10), 45 (10), 43 (34), 42 (18), 41 (11).
3	26.09	+++	see signal no. 2
4	26.75	++	see signal no. 2
5	27.08	++	see signal no. 2



*Fig. 1.* Molecular formulas of pyrrolizidine alkaloids. I = 7-angelylheliotridine-viridiflorate, II = supinine, III = petasinine, IV = fuchsisenecionine, V = macrophylline, VI = seneciphylline, VII = spartiodine, VIII = doronenine, IX = senecionine, X = retroisosenine, XI = platyphylline, XII = bulgarsenine, XIII = nemorensine, XIV = petasitenine, XV = neopetasitenine

### *Homogyne alpina*

The chromatogram shows two main signals which correspond to petasitenine (XIV, signal No 2) and neopetasitenine (XV, signal No 3). The mass spectra of these two signals are identical with the spectra given by Yamada et al. (14) for petasitenine and neopetasitenine from *Petasites japonicus*. The identity of signal No 2 was ensured by co-chromatography together with native petasitenine. The lower temperature range of the chromatogram contains a minor signal (No 1) with a molecular mass of 237 daltons due to the retronecyl ester of a monounsaturated C<sub>5</sub>-carboxylic acid (angelic, tiglic or senecic acid).

### *Senecio Doronicum*

*S. Doronicum* has already been investigated by Röder et al. (15): the two isologic PAs doronenine and bulgarsenine were isolated and identified.

The chromatogram of the free alkaloids from *S. Doronicum* exhibits a wealth of signals being due to PAs; they are arranged in two groups: the mass spectra of signals No 1–4 in the first group correspond in their molecular ion at m/e 239. Signals No 1 and 3 and No 2 and 4 respectively exhibit each mass spectra with high coincidence. Comparison of the spectra with data from the literature yields macrophylline (V, 9-angelylmacronecine) (16) for signals No 2 and 4. Although no reference data were available for signals No 1 and 3 the fragment pattern in the mass spectra can be understood. The signals at m/e 140 and 139 possibly are due to a loss of an angelyl or isomeric moiety and to an elimination of the respective acid after a hydrogen transfer. A following loss of a methanol unit would cause the signal at m/e 108. These considerations can be brought into line by petasinine (III) (17) or 2-angelylmacronecine.

Evaluation of the mass spectra of signal No 5 shows an overlapping of two alkaloids with molecular masses of 335 and 337 daltons respectively. Signal No 6 exhibits a fragment pattern characteristic for PAs with an unsaturated pyrrolizidine moiety (m/e 93, 119, 136 in the electron impact mass spectrum, m/e 120 in the positive CI spectrum) and a molecular ion at m/e 335. The spectrum shows many similarities with the spectra of the isomers senecionine and integerrimine (10), and in co-chromatography it elutes together with senecionine but its mass spectrum exhibits two striking differences: the signal at m/e 119 is relatively abundant and the signal at m/e 246 is lacking compared to the mass spectra of senecionine. A reference mass spectrum which exhibits these two characteristic details has been found in the literature: retroisosenine (X) (17). Signals No 7, 10 and 12 give fragments with masses characteristic for PAs of the platynecine type (m/e 122, 138 and 140) and molecular ions at m/e 337. The mass spectrum of signal No 8 corresponds with the mass spectrum of doronenine (VIII) (15) and signal No 9 gives a spectrum which is assigned to bulgarsenine (XII) (18). For the spectra of the remaining signals in the chromatogram no references could be found in the literature. The last signal in the chromatogram (No 13) gives a mass spectrum which

points to a PA containing a platynecine moiety, for example a hydroxibulgarsenine isomer.

### *Senecio Fuchsii*

The PA content of *S. Fuchsii*, a species of the group of *S. nemorensis* has already been investigated by several authors (see table 2).

The main signal in the chromatogram (No 7) is due to an alkaloid not reported in the literature until now. Its mass spectrum indicates a molecular mass of 337 daltons, and the fragment pattern points to a saturated PA with a platynecine moiety (*m/e* 96, 122, 138, 140).

Comparison with nemorensine (23) from *S. nemorensis* (co-chromatography, mass spectra) revealed a different behaviour. The reference mass spectra of other isobar PAs as neoplatyphylline (24), hastacine (24) and bulgarsenine (18) exhibit differing fragment patterns. Signals No 8 and 9 give identical spectra indicating isomers of the unknown PA. Signals No 5 and 6 again show the same molecular ions  $M^+$  337). Signal No 6 corresponds to platyphylline, whereas for signal No 5 no reference mass spectrum was available. Signals No 10 and 11 contain very similar spectra of apparently isomeric compounds with a molecular mass of 353 daltons. Evaluation of the fragment pattern in the lower mass region leads to a saturated PA, tentatively an epoxy or hydroxy analogue of the unknown PA in signal No 7. The main PA in the lower temperature range of the chromatogram (No 3) corresponds to 7-senecioylplatynecine (fuchsisenencionine (IV)) (20). Signal No 1 exhibits a molecular ion at *m/e* 237, with a mass spectrum corresponding to the spectrum of 7-angelylretronecine reported by Pedersen et al. (25). Due to its mass spectrum signal No 2 corresponds to an isomer of fuchsisenencionine. Although reference data from the literature are lacking a platynecine esterified with angelic (or an isomeric) acid is supposed.

Table 2. PA content of some species and subspecies of *S. nemorensis* (after Robins (2))

<i>Senecio nemorensis L.</i>	
Jacquiniana (Rchb.) Durand	Nomorensine (XIII) [23]
bulgaricus (Vel.) Stoj. et Stef.	Nomorensine (XIII), bulgarsenine (XII), retroisosenine (X) [18, 23]
subdecurrens Griseb.	Nomorensine (XIII), oxynomorensine (N-oxide of nemorensine), retroisosenine (X), bulgarsenine (XII) [16, 22, 23]
Fuchsii Gmelin	Senencionine (IX), fuchsisenencionine (IV) [20, 21, 23]
Fuchsii Gmelin	Two unknown isomeric PAs, fuchsisenencionine (IV), bulgarsenine (XII), and minor PAs (this paper)
nova (Zlatnik)	Nomorensine (XIII) [23]

A comparison of these results with the data on the investigation of *S. nemorensis* from the literature is presented in table 2. Three different alkaloid patterns have been established for *S. Fuchsii*. Klasek and coworkers (23) have shown the presence of nemorensine, which is also a constituent in other subspecies and varieties of *S. nemorensis*. Röder and coworkers identified fuchsisenecionine as main constituent and senecionine in *S. Fuchsii* (20, 21, 26). In a later publication the same authors reported the g. l. c. behaviour of PAs from some *Senecio* species (27). They found that fuchsisenecionine was eluted after senecionine. This is in strong contrast to the elution sequence observed in the chromatogram of the present paper (see table 1). A possible explanation for this contradiction is that the retention of fuchsisenecionine in (27) resembles much more the retention of a PA with a higher molecular mass like the main PA in the present paper (signal No 7) than the relatively small fuchsisenecionine.

### *Eupatorium cannabinum*

For *E. cannabinum*, another representative of the family *compositae* the PA constituents echinatine and supinine are reported by Pedersen et al. (28). Our own investigation of a Swiss sample of this plant revealed some supplementary results.

Signal No 1 in the chromatogram gives a mass spectrum with the molecular ion and the fragment pattern characteristic for supinine (II), which could be verified by co-chromatography with native supinine. Four apparently isomeric compounds with a molecular mass of 381 daltons and practical identical fragmentation pattern in the mass spectra were detected and established with chemical ionisation (signals No 2-5). They are supposed to correspond to isomers of 7-angelylheliotridine-viridiflorate (I). A mass spectrum corresponding to echinatine as found by Pedersen is lacking in our results.

### *Acknowledgement*

The authors are indebted to Dr. C. C. J. Culvenor, Parkville, Australia, for a gift of authentic platyphylline and supinine, and to Dr. M. Neuenschwander, University of Bern, Switzerland, for a gift of authentic petasitenine, and to Dr. R. Gottlieb, Dr. Gustav Klein, Arzneimittelforschung, Zell-Harmersbach (GFR) for a gift of authentic nemorensine.

### *Summary*

The occurrence of pyrrolizidine alkaloids in the plant species *Adenostyles Alliariae*, *Homogyne alpina*, *Senecio Doronicum*, *Senecio Fuchsii*, and *Eupatorium cannabinum* was studied using GC/MS. EI and CI mass spectra allow a rapid identification of the alkaloids in most cases or can give important indications to the structures at least.

## Zusammenfassung

Das Vorkommen von Pyrrolizidinalkaloiden in den Pflanzenarten *Adenostyles Alliariae*, *Homogyne alpina*, *Senecio Doronicum*, *Senecio Fuchsii* und *Eupatorium cannabinum* wurde mittels GC/MS studiert. EI- und CI-Massenspektren der zahlreich gefundenen Alkaloide erlauben in den meisten Fällen deren rasche Identifizierung oder geben zumindest wichtige Hinweise zur Struktur.

## Résumé

La présence d'alcaloïdes pyrrolizidiniques dans les sortes de plantes suivantes *Adenostyles Alliariae*, *Homogyne alpina*, *Senecio Doronicum*, *Senecio Fuchsii* et *Eupatorium cannabinum* a été étudiée à l'aide de la GC/MS. Les spectres de masse EI et CI des nombreux alcaloïdes trouvés ont permis dans la plupart des cas de les identifier rapidement ou nous donnent pour le moins des indications importantes quant à leur structure.

## Literature

1. Robins, D. J. In: The biology and chemistry of the *Compositae* (ed. V. H. Heywood et al.), p. 831ff. Academic Press, London 1977.
2. Robins, D. J.: The pyrrolizidine alkaloids. Fortschr. Chem. org. Naturstoffe, Vol. 41 (1982).
3. Mohabbat, O., Shafiq Younos, M., Merzad, A. A., Srivastava, R. N., Sedig Gholam Ghaos and Aram, G. N.: An outbreak of hepatic venoocclusive disease in northwestern Afghanistan. Lancet 269–271 (1976).
4. Tandon, B. N., Tandon, R. K., Tandon, H. D., Narndranathan, M. and Joshi, Y. K.: An epidemic of venoocclusive disease of liver in central India. Lancet 271–272 (1976).
5. Pohlenz, J., Lüthy, J., Minder, H. P. and Bivetti, A.: Enzootische Leberzirrhose beim Rind, verursacht durch Pyrrolizidinalkaloiide nach Aufnahme von *Senecio alpinus* (Alpenkreuzkraut). Schweiz. Arch. Tierheilk. 122, 183–193 (1980).
6. Bull, L. B., Culvenor, C. C. J. and Dick, A. T.: The pyrrolizidine alkaloids, p. 54. North-Holland, Amsterdam 1968.
7. IARC Monographs on the evaluation of carcinogenic risk of chemicals to man, Vol. 10, p. 265–342, Lyon 1976.
8. Lüthy, J., Zweifel, U., Schmid, P. and Schlatter, Ch.: Pyrrolizidin-Alkaloiide in *Petasites hybridus* L. und *P. albus* L. Pharm. Acta Helv. 58, 98–100 (1983).
9. Lüthy, J., Brauchli, J., Schmid, P. and Schlatter, Ch.: Pyrrolizidin-Alkaloiide in Arzneipflanzen der Boraginaceen: *Borago officinalis* L. und *Pulmonaria officinalis* L. Pharm. Acta Helv. 59, 242–246 (1984).
10. Lüthy, J., Zweifel, U., Karlhuber, B. and Schlatter, Ch.: Pyrrolizidine alkaloids of *Senecio alpinus* L. and their detection in feedingstuffs. J. Agric. Food Chem. 29, 302–305 (1981).
11. Heim, T.: Zu Metabolismus, Kinetik und Übergang in Milch der Pyrrolizidinalkaloiide von *Senecio alpinus* bei der Ratte. Thesis ETH 7461, Zürich 1984.
12. Warren, F. L.: The pyrrolizidine alkaloids. Fortschr. Chem. org. Naturstoffe 12, 198 (1955).

13. Yakhoutova, L. D., Pimenov, M. G. and Sapunova, L. A.: Khim. Prir. Soedin, 122 (1976).
14. Yamada, K., Tatematsu, H., Suzuki, M., Hirata, Y., Haga, M. and Hirono, I.: Isolation and the structures of two new alkaloids: petasitenine and neopetasitenine from *Petasites Japonicus* Maxim. Chem. Letters 461–464 (1976).
15. Röder, E., Wiedenfeld, H. and Frisse, M.: Pyrrolizidinalkaloide aus *Senecio Doronicum*. Phytochemistry 19, 1275–1277 (1980).
16. Aasen, A. J., Culvenor, C. C. J. and Smith, L. W.: The saturated pyrrolizidine diols. I. Spectral studies and the conversion of an ester of dihydroxyheliotridane into the (+)-enantiomer of hastanecine. J. Org. Chem. 34, 4137–4143 (1969).
17. Yamada, K., Tatematsu, H., Unno, R. and Hirata, Y.: Petasinine and petasinoside, two minor alkaloids from *Petasites Japonicus* Maxim. Tetrahedron Letters 46, 4543–4546 (1978).
18. Nghia, N. T., Sedmera, P., Klásek, A., Boeva, A., Drjanovska, L., Dolejš, L. and Šantavý, F.: Bulgarsenine and Retroisosenine, alkaloids from *Senecion nemorensis* L. var. bulgaricus. Coll. Czech. Chem. Commun. 41, 2952–2963 (1976).
19. Müller, A.: Zur Kenntnis der Senecio-Arten. Heil Gewürzpflanzen 7, 1–24 (1924).
20. Röder, E. und Wiedenfeld, H.: Isolierung und Strukturaufklärung des Alkaloids Fuchsise-necionin aus *Senecio Fuchsii*. Phytochemistry 16, 1462–1463 (1977).
21. Wiedenfeld, H.: Isolierung, Strukturaufklärung und Partialsynthese eines Alkaloids aus *Senecio Fuchsii*. Thesis, Bonn 1976.
22. Klásek, A., Sedmera, P., Vokoun, J., Boeva, A., Dvorackova, S. and Šantavý, F.: Oxynemo-reنسine, an alkaloid from *Senecio nemorensis* L. Coll. Czech. Chem. Commun. 45, 548–558 (1980).
23. Klásek, A., Sedmera, P., Boeva, A. and Šantavý, F.: Pyrrolizidine alkaloids. XX. Nemoren-sine, an alkaloid from *Senecio nemorensis* L. Coll. Czech. Chem. Commun. 38, 2504–2512 (1973).
24. Culvenor, C. C. J., Koretskaga, N. I., Smith, L. W. and Utkin, L. M.: The structures of the pyrrolizidine alkaloids neoplatyphylline and hastacine. Aust. J. Chem. 21, 1671–1673 (1968).
25. Pedersen, E. and Larsen, E.: Mass Spectrometry of some pyrrolizidine alkaloids. Org. Mass Spectrom. 4, 249–256 (1970).
26. Wiedenfeld, H. and Röder, E.: Das Pyrrolizidinalkoid Senecionin aus *Senecio Fuchsii*. Phytochemistry 18, 1083–1084 (1979).
27. Wiedenfeld, H., Pastewka, U., Stengl, P. and Röder, E.: Zur gaschromatographischen Be-stimmung der Pyrrolizidinalkaloide aus einigen Senecioarten. Planta med. 41, 124–128 (1981).
28. Pedersen, E.: Echinatine and Supinine: pyrrolizidine alkaloids from *Eupatorium canabi-num*. Phytochemistry 14, 2086–2087 (1975).

Dr. J. Lüthy  
 Bundesamt für Gesundheitswesen  
 Abteilung Lebensmittelkontrolle  
 Postfach 2644  
 CH-3001 Bern

Dr. P. Schmid  
 U. Zweifel  
 Prof. Dr. Ch. Schlatter  
 Institut für Toxikologie  
 der ETH und Universität Zürich  
 Schorenstrasse 16  
 CH-8603 Schwerzenbach