

Kratom is the leaves of *M. speciosa* endemic to tropical Southeast Asia. The leaves have been traditionally used for their opium-like effect and coca-like stimulant ability in Thailand and Malaysia, where the leaves are called 'Kratom' and 'Biak-Biak,' respectively [1, 2]. The leaves contain mitragynine (1) and its related indole alkaloids [1, 2], and 7-hydroxymitragynine (2) is reported as the active principle for their opium-like effect [2, 3].

During the course of our study of psychoactive plants [4–6], we investigated the botanical origin of the kratom obtained from the Japanese markets in preparation for its future regulation based on the internal transcribed spacer (ITS) sequence of the nuclear rDNA. In addition, the polymerase chain reaction-restriction fragment length polymorphism (PCR-RFLP) method reported by Sukrong et al. [7] for the rapid identification of *M. speciosa* was applied to the commercial kratom products in order to estimate its accuracy and utility.

Materials and methods

Materials

Details of the referential plants and drugs which are recognised as kratom or related species in Thailand and Malaysia are shown in Table 1, together with their ITS genotypes. Among them, Mt-1 to Mt-3 were collected at Chulalongkorn University, Thailand, and were identified as *M. speciosa* (Mt-1 and Mt-2) and *M. hirsuta* (Mt-3) by Dr. Sumphan Wongseripipatana. Although Mt-1 and Mt-2 are not discriminated taxonomically, they have quite different leaves. So, we separately treated them as a big leaf for Mt-1 and a small leaf for Mt-2 based on their leaf size. Mt-4 to Mt-7 were kindly supplied by Ms. Latifah BT. HJ. Idris at the Pharmaceutical Services Division, Ministry of Health, Malaysia, through the Kanto-Shin'etsu Regional Narcotic Control Office, Japan. They were obtained as dried raw materials. Voucher specimens were deposited at the Division of Pharmacognosy, Phytochemistry and Narcotics,

National Institute of Health Sciences, Japan. A map of the collection area is shown in Fig. S1, Supplementary Material. The phytochemical study of Mt-3 (*M. hirsuta*) has already been reported by Kitajima et al. [8].

Details of the commercial kratoms available in the Japanese markets are shown in Table 2, together with their ITS genotypes. They were purchased from shops on the Internet. They were in powder, dried leaves and gum product forms.

The leaves of magic mint (*S. divinorum*) and chacruna (putative *Psychotria viridis*; Rubiaceae) used for the negative controls in the PCR-RFLP assay were also purchased from shops on the Internet. Magic mint used in this study has already been identified as *S. divinorum* by genetic and phytochemical techniques [6].

rDNA, ITS sequence analysis

Ten mg of each sample was crushed using a mixer mill, MM-300 (Qiagen). The genomic DNA was extracted and purified from the powdered sample using a DNeasy Plant Mini Kit (Qiagen). The ITS region (small subunit rDNA-ITS1-5.8S rDNA-ITS2-large subunit rDNA) of the nuclear rDNA was two-steps (nested) PCR-amplified using the obtained genomic DNA as a template. PCR was separately performed with Gene TaqNT DNA polymerase (Nippon Gene) and KOD DNA Polymerase (Toyobo) using a DNA engine PTC-200 (Bio-Rad). The primer sequences are: ITS-S1 5'-GGA AGT AAA AGT CGT AAC AAG G-3' and ITS-AS1 5'-TTT TCC TCC GCT TAT TGA TAT GC-3' for the first-round PCR; and ITS-S2 5'-TCC GTA GGT GAA CCT GCG G-3' and ITS-AS2 5'-GTA GTC CCG CCT GAC CTG-3' for the second-round PCR. The amplicons were purified by Montage-PCR (Millipore) and were directly sequenced. The cycle sequencing reaction was performed using a BigDye Terminator v3.1 cycle sequencing kit (Applied Biosystems). Subcloning of the amplicon by KOD DNA polymerase into the plasmid vectors was performed using a Zero Blunt PCR Cloning Kit (Invitrogen). The DNA sequences were aligned with a Clustal W program [9].

Table 1 Details of the referential plants and drugs used in this study

Sample no.	Scientific name or crude drug name	Locality	Form	Genotype
Mt-1 ^a	<i>M. speciosa</i> (big leaf)	Bangkok, Thailand	Dried leaf	Type 1
Mt-2 ^a	<i>M. speciosa</i> (small leaf)	Bangkok, Thailand	Dried leaf	Hybrid ^b
Mt-3	<i>M. hirsuta</i>	Bangkok, Thailand	Dried leaf (cutting)	Type 5
Mt-4	Kratom	Perlis, Malaysia	Dried leaf	Type 1
Mt-5	Kratom	Pahang, Malaysia	Dried leaf	Type 2
Mt-6	Kratom	Negeri Semblian, Malaysia	Dried leaf	Type 1
Mt-7	Kratom	Perlis, Malaysia	Dried leaf	Type 1

^a Although they are not discriminated in taxonomically, they have quite different leaves

^b Hybrid between *M. speciosa* and *M. diversifolia*/*M. hirsuta*

Table 2 Details of the commercial kratom products used in this study

Sample no.	Name	Form	Genotype
Mtk-1	Concd. kratom (50–60% pure)	Gum	No amplicon
Mtk-2	Kratom Crush Leaf KR-L	Dried leaves (cutting)	Type 3
Mtk-3	Kratom Gold KR-G	Powder	Type 4
Mtk-4	Kratom KR-1	Powder	Type 3
Mtk-5	Kratom	Dried leaves (cutting)	Type 4
Mtk-6	Kratom	Dried leaves	Type 3
Mtk-7	Mitragaina Extract Powder	Powder	Type 4
Mtk-8	Mitragaina	Dried leaves	Other ^a
Mtk-9	Mitragyna 40× resin	Gum	No amplicon
Mtk-10	Plant Sample Kratom	Dried leaves (cutting)	Type 3
Mtk-11	Kratom EX10	Dried leaves (cutting)	Other ^a
Mtk-12	Concd. Kratom KR-XV	Gum	No amplicon
Mtk-13	Kratom KR-1	Powder	Type 3

^a BLAST search suggests a closely related plant within the same tribe, Naucleaeae s.l. as *Mitragyna* (Table S1)

Table 3 Internal transcribed spacer (ITS) genotypes with nucleotide variable sites found in this study and the international nucleotide sequence database

Genotype	Aligned position															Accession no.
	ITS1						5.8S	ITS2								
	26	42	101	105	118	171	362	452	510	514	516	539	564	565	608	
Type 1	T	C	C	C	A	-	A	G	T	T	A	C	C	A	C	AB249645
Type 2	.	.	.	M	None
Type 3	.	.	.	A	None
Type 4	.	.	Y	M	M	.	.	.	None
Type 5	Y	G	T	.	-	G	C	A	-	C	C	.	T	G	A	None
<i>M. hirsuta</i>	.	G	T	.	-	G	C	A	-	C	C	.	T	G	A	AB249647
<i>M. diversifolia</i>	.	G	T	.	-	G	C	A	-	C	C	.	T	G	A	AB249646

A dot '.' and a hyphen '-' indicate the same nucleotide as *M. speciosa* type 1 sequence and a gap, respectively. M, A/C; Y, C/T

PCR-RFLP assay

The ITS1 DNA fragment was PCR-amplified with a new pair of primers, ITS-S1 and ITS1-AS1 (5'-TAT CCG TTG CCG AGA GTC-3') using Nova Taq DNA polymerase Hot Start (Merck) and Ampdirect Plus (Shimadzu). The resulting PCR products were digested with a restriction enzyme, *Xma*I (New England BioLabs), at 37°C for 4 h. The digested fragments were analysed by 2.0% agarose gel electrophoresis and visualised by ethidium bromide staining.

Results and discussion

ITS sequence analysis

The genus *Mitragyna* consists of ten species around the world. A phylogenetic analysis of the *Mitragyna* plants has already been performed using nuclear rDNA, ITS and

chloroplast DNA, *rbc*L and *trn*T-F sequences [7, 10], and it was shown that *M. speciosa* is well discriminated from other related plants based on the ITS sequence [7].

The ITS sequence of each genotype found in this study is summarised in Table 3, together with those on the international nucleotide sequence database (DDBJ, EMBL and GenBank; INSD), and the genotypes of all samples are shown in Tables 1 and 2. Among the referential plants and drugs (Mt-1 to Mt-7), the ITS sequence of Mt-1, 4, 6 and 7 were identical and the same as that of *M. speciosa* on the INSD (accession no.: AB249645; type 1 in Table 3). The ITS sequence of Mt-3 was also identical to those of *M. hirsuta* and *M. diversifolia* on the INSD (AB249646 and AB249647, respectively), except for one site with nucleotide additivity [11]. The sequence of Mt-2 seemed to be a mixture of the type 1 and *M. hirsuta/diversifolia* sequences on the INSD on direct sequencing. Therefore, the amplicon from Mt-2 was introduced into the plasmid vector and 12 clones were sequenced. As a result, six sequences each of

type 1 and *M. hirsuta/versifolia* were detected, suggesting that Mt-2 was a hybrid between *M. speciosa* and *M. hirsuta/versifolia*. This result was also supported by the contents of 1 and 2 in Mt-1 to Mt-3: 23.4 and 0.124 mg/g of 1 and 2 were present respectively in Mt-1, while 1.6 and 0.031 mg/g were present in Mt-2 and neither 1 nor 2 was detected in Mt-3 (Details will be reported in a future paper). Mt-5 has the same sequence as type 1 except for one site with nucleotide additivity [11]. These data indicate that most of the kratoms from Thailand and Malay peninsula have the type 1 sequence.

Among 13 commercial kratom products (Mtk-1 to -13), no amplicons were obtained from the gum samples (Mtk-1, -9 and -12), which appear to contain no organic tissue. The ITS sequences of Mtk-8 and -11 were identical, and the similarity survey for the sequence using the BLAST search program suggested that these products were made from the same tribe plant (*Naucleaeae sensu lato*) [10] as *Mitragyna* (Table S1, Supplementary Material). The presence of these plants in the commercial kratoms can be dangerous for human health because of the various indole alkaloids with high pharmacological activities are reported from the *Naucleaeae* s.l. plants [12]. Although the ITS sequences of the other samples (Mtk-2 to -7, -10 and -13) showed a great similarity to the type 1 sequence, all of them showed a one or three nucleotide(s) difference from the type 1 sequence (types 3 and 4 in Tables 2 and 3). Probably, a small number of nucleotide substitution(s) in types 3 and 4 sequences is (are) attributed to either the intraspecific nucleotide variation in *M. speciosa* or the closely related plants to *M. speciosa*. Unfortunately, the real truth remains inconclusive from the current study. A more detailed investigation including the wide sampling in other areas such as Sumatra, Borneo and the New Guinea islands might solve this question.

All of the commercial kratoms having types 3 or 4 sequences were found to contain the hallucinogens in the range from 12.1 to 21.4 mg/g of 1 and 0.114 to 0.393 mg/g

of 2 in the LC/MS analysis (details will be reported in a future paper). Therefore, we subsequently examined the accuracy and utility of the reported authentication method of *M. speciosa* [7] in preparation for the future regulation of kratom, although the original species of the commercial kratoms remain unclear.

PCR-RFLP assay of the commercial kratom products

The PCR-RFLP method for the identification of *M. speciosa* established by Sukrong et al., is based on the nucleotide substitution at position 42 among the *Mitragyna* species [7]. Namely, *M. speciosa* has an *Xma*I site (CCCGGG) in the aligned position 42–47, while other species have the sequence GCCGGG in the same position. Although some different ITS genotypes from that in Sukrong et al.'s study [7] were found in the commercial kratoms, there are no nucleotide substitutions at position 42 among all of the ITS genotypes observed in this study. Therefore, it is expected that Sukrong et al.'s method can be utilised for the rapid identification of commercial kratom products. Hence, we applied the commercial kratoms to the method. For the negative controls in the assay, the leaves of chacruna and magic mint were selected because they are hallucinogenic plants normally used in the form of leaves, like the kratom products. The electropherogram of the assay is shown in Fig. 1. The image is also shown in colour in Fig. S2, Supplementary Material. Only the PCR products from the samples which have kratom genotypes were cleaved into two fragments by the *Xma*I digestion, while the original PCR products remained as a single band in the other samples (Fig. 1). The method could clearly distinguish kratoms from the other psychoactive plants (NC1 and NC2) and also from the same family (Mtk-8 and -11) and genus (Mt-3; *M. hirsuta*) plants as *M. speciosa* (Fig. 1). Simultaneously, our data indicate that the method is practical enough for application to processed products, except for gum samples.

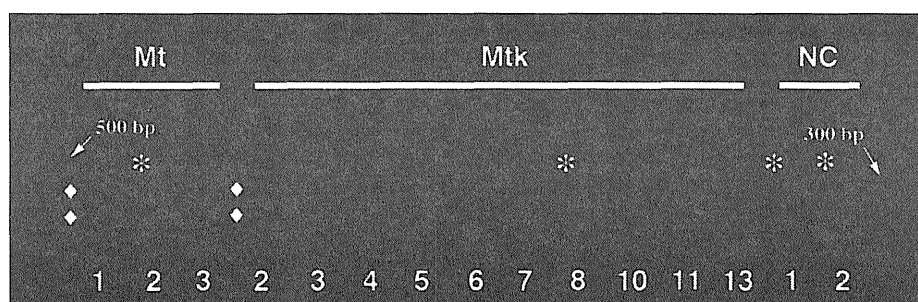


Fig. 1 Electropherogram in the polymerase chain reaction-restriction fragment length polymorphism (PCR-RFLP) assay. NC: negative control. NC1, chacruna (putative *Psychotria viridis*); NC2, magic mint (*Salvia divinorum*). Details of Mt-1 to -3 and Mtk-2 to 13 are

shown in Tables 1 and 2. An asterisk and a diamond indicate the original PCR product and the restriction fragments, respectively. The samples in which types 1 to 4 sequences were detected are shown as underlined numbers

In conclusion, the present study revealed that most of the commercial kratoms available in the Japanese markets are derived from *M. speciosa* or closely related plants and that the reported authentication method for *M. speciosa* utilizing PCR-RFLP has high practicality due to its wide range of application, high accuracy and simplicity.

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References

1. Takayama H, Aimi N, Sakai S (2000) Chemical studies on the analgesic indole alkaloids from the traditional medicine (*Mitragyna speciosa*) used for opium substitute. *Yakugaku Zasshi* 120:959–967
2. Takayama H (2004) Chemistry and pharmacology of analgesic indole alkaloids from the rubiaceous plant, *Mitragyna speciosa*. *Chem Pharm Bull* 52(8):916–928
3. Matsumoto K, Horie S, Ishikawa H, Takayama H, Aimi N, Ponglux D, Watanabe K (2004) Antinociceptive effect of 7-hydroxymitragynine in mice: discovery of an orally active opioid analgesic from the Thai medicinal herb *Mitragyna speciosa*. *Life Sci* 74:2143–2155
4. Maruyama T, Kawahara N, Fukiharu T, Yokoyama K, Makino Y, Goda Y (2005) DNA and chemical analyses of commercial fly agaric-related products. *Shokuhin Eiseigaku Zasshi* 46:49–54
5. Maruyama T, Kawahara N, Yokoyama K, Makino Y, Fukiharu T, Goda Y (2006) Phylogenetic relationship of psychoactive fungi based on rRNA gene for a large subunit and their identification using the TaqMan assay (II). *Forensic Sci Int* 163:51–58
6. Maruyama T, Kamakura H, Kikura-Hanajiri R, Goda Y (2008) Authentication and ultra performance liquid chromatography (UPLC)/MS analysis of magic mint, *Salvia divinorum* and its related plants. *Yakugaku Zasshi* 128:179–183
7. Sukrong S, Zhu S, Ruangrunsi N, Phadungcharoen T, Palanuvej C, Komatsu K (2007) Molecular analysis of the genus *Mitragyna* existing in Thailand based on rDNA ITS sequences and its application to identify a narcotic species: *Mitragyna speciosa*. *Biol Pharm Bull* 30:1284–1288
8. Kitajima M, Nakayama T, Kogure N, Wongseripipatana S, Takayama H (2007) New heteroyohimbine-type oxindole alkaloid from the leaves of Thai *Mitragyna hirsuta*. *J Nat Med* 61:192–195
9. Thompson JD, Higgins DG, Gibson TJ (1994) CLUSTAL W: improving the sensitivity of progressive multiple sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice. *Nucleic Acids Res* 22:4673–4680
10. Razafimandimbison SG, Bremer B (2002) Phylogeny and classification of Naucleaeae s.l. (Rubiaceae) inferred from molecular (ITS, *rbcl*, and *trnT-F*) and morphological data. *Am J Bot* 89:1027–1041
11. Sang T, Crawford DJ, Stuessy TF (1995) Documentation of reticulate evolution in peonies (*Paeonia*) using internal transcribed spacer sequences of nuclear ribosomal DNA: implications for biogeography and concerted evolution. *Proc Natl Acad Sci USA* 92:6813–6817
12. Phillipson JD, Hemingway SR, Ridsdale CE (1982) The chemotaxonomic significance of alkaloids in the Naucleaeae s.l. (Rubiaceae). *J Nat Prod* 45:145–162

Identification of a Cannabinoid Analog as a New Type of Designer Drug in a Herbal Product

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A new type of designer drug, a cannabinoid analog (**1**), was found in a herbal product distributed on the illegal drug market in Japan in expectation of its narcotic effect. The structure of **1** was identified by LC-MS, GC-MS, high-resolution MS, and NMR analyses. Compound **1** showed a molecular weight of 332, and accurate mass measurement exhibited its elemental composition to be $C_{22}H_{36}O_2$. Together, the mass and NMR spectrometric data revealed that **1** was (1*RS*,3*SR*)-3-[4-(1,1-dimethyloctyl)-2-hydroxyphenyl]cyclohexan-1-ol, which was first synthesized in 1979 by a group at Pfizer Inc. and reported as a potent cannabinoid analog possessing cannabinoid receptor binding activity and analgesic activity in the 1990s. This is the first report to identify a cannabinoid analog in an illegal drug.

Key words cannabinoid analog; designer drug; herbal product; (1*RS*,3*SR*)-3-[4-(1,1-dimethyloctyl)-2-hydroxyphenyl]cyclohexan-1-ol; drug abuse

Many types of chemicals are widely distributed and abused as psychotropic substances. In Japan every year this decade, a market survey of illegal drugs is performed by the Ministry of Health, Labour and Welfare.^{1–9} Following the results of the survey, the compound identified and recognized as a designer drug^{10–12} came to be strictly controlled by the Narcotics and Psychotropic Control Law or by the Pharmaceutical Affairs Law as designated substances (Shitei-Yakubutsu).^{13–16} In 2008, seven new designer drugs were classified as narcotics or designated substances, and all of them are analogs of phenylethylamine or tryptamine.

Cannabis sativa L. (cannabis, hemp, marijuana, marihuana) is widely abused around the world because it contains psychoactive cannabinoids, such as Δ^9 -tetrahydrocannabinol (Δ^9 -THC), which contains no amine groups (Fig. 1). In the past few decades, a number of analogs of Δ^9 -THC were synthesized, and their structure–activity relationships were studied.^{17,18} In the 1980s, a group at Pfizer Inc. explored the development of analgesics using potent synthetic cannabinoids.^{19–22} After the discovery of cannabinoid receptors, type 1 (CB₁, central type) and type 2 (CB₂, peripheral

type), as well as the discovery of an endogenous cannabinoid, their physiological roles were elucidated; a number of cannabinoid analogs were then newly synthesized, and their pharmacological activity for the treatment of various diseases was studied.^{23,24}

Recently, cannabis abuse seems to have spread in Japan. In this study, we identified a novel designer drug (**1**) possessing cannabinoid activity as an adulterant in a herbal product (Fig. 1). Compound **1** was first synthesized by Pfizer Inc. in 1979,²⁵ and reported as a cannabinoid analog in the 1990s.^{26–30} Although many designer drugs having phenylethylamine, tryptamine, and piperazine structures have been found,^{10–12} this is the first report to identify a non-nitrogenated compound, a phenylcyclohexane derivative having cannabinoid activity.

Experimental

Chemicals and Reagents HPLC-grade acetonitrile and all other chemicals (analytical grade) were obtained from Wako Pure Chemical Industries, Ltd. (Osaka, Japan). Centrifugal filter devices (Ultrafree-MC, 0.45 μ m filter unit) were from Millipore (Bedford, MA, U.S.A.).

Samples A product was purchased via the Internet (June 2008). The product was described as a herbal mixture and had the appearance of dried plants. The ingredients were listed as “Baybean,” “Blue lotus,” “Dwarf skullcap,” “Indian warrior,” “Lion’s tail,” “Maconha brava,” “Marshmallow,” “Pink lotus,” “Red clover,” “Rose,” “Siberian motherwort,” “Vanilla,” and “Honey.”

Preparation of Sample Solution A product (20 mg) was crushed into powder and extracted with 2 ml of methanol under ultrasonication for 10 min. After centrifugation (5 min at 3000 rpm), the solution was filtered through a centrifugal filter device.

Instrumentation Gas chromatography-mass spectrometry (GC-MS) in the electron impact (EI) mode at 70 eV of electron energy was used. Analysis was performed on a Hewlett-Packard 6890N GC with a 5975 mass selective detector using a capillary column (HP1-MS capillary, 30 m \times 0.25 mm i.d., 0.25 μ m film thickness) and helium gas as a carrier. An initial column temperature of 80 $^{\circ}$ C was employed and the temperature was increased at a rate of 5 $^{\circ}$ C/min to 190 $^{\circ}$ C and at a second rate of 10 $^{\circ}$ C/min to 310 $^{\circ}$ C. Data were obtained in a full scan mode with a scan range of m/z 40–550. An ultra-performance liquid chromatography-electrospray ionization-mass spectrometer (UPLC-ESI-MS), consisting of a Waters ACQUITY UPLC system equipped with a Single Quadrupole Detector (SQD) mass detector and a photo diode array (PDA) (Waters, Milford, MA, U.S.A.), was also used. The sample solutions were separated using a Waters ACQUITY UPLC HSS T3 column (2.1 \times 100 mm i.d., 1.8 μ m; Waters) at 40 $^{\circ}$ C. The following gradient system was used with a mobile phase A (10 mM ammonium formate buffer, pH 3.5) and a mobile phase B (acetonitrile) delivered at 0.3 ml/min; A : B 50 : 50 (0 min)–20 : 80 (20–40 min). The injection volume was 5 μ l. The wavelength of the PDA detector for screening was set from UV 190 to 400 nm, and chromatographic peaks were monitored at UV 254 and 280 nm. Mass analysis by the ESI was used in both a positive and a negative mode. Nitrogen gas was used for desolvation at a flow rate of 600 l/h at 350 $^{\circ}$ C. The capillary voltage was 3000 V, and the cone voltage was 30 V. MS data were recorded in the full scan mode (m/z 150–700). Preparative TLC was carried out using a silica gel plate (silica gel 60, 20 \times 20 cm, 0.5 mm, Merck, Darmstadt, Germany).

Isolation of Compound 1 A product (3 g) was extracted with 100 ml of methanol by ultrasonication for 1 h. After the extraction was performed three times, the supernatant was evaporated to dryness. The extract was subjected to preparative silica gel TLC using $CHCl_3$ –acetone (4/1) as developing solvent. A portion of the silica gel in the TLC plate was taken and eluted with $CHCl_3$ –MeOH (1/1) to give a fraction 1. Repeated fractionation of fr. 1 by preparative silica gel TLC with $CHCl_3$ –MeOH (20/1) gave compound **1** (15 mg) as an off-white solid.

Measurement of Accurate Mass The accurate mass of the target compound was measured by the LTQ Orbitrap XL instrument (Thermo Fisher Scientific Inc., Waltham, MA, U.S.A.) with the direct-infusion ESI positive and negative ion modes under the following conditions: solvent flow rate 5 μ l/min, sheath gas flow rate 20 arb, auxiliary gas flow rate 10 arb, spray voltage 5 kV, capillary temperature 275 $^{\circ}$ C, capillary voltage 4 V, and tube

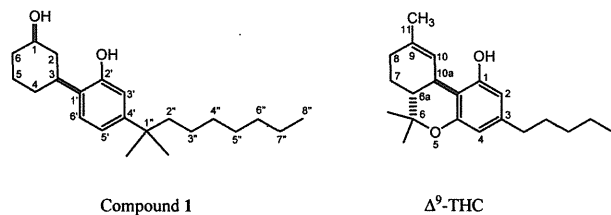


Fig. 1. Structures of Compound **1** and Δ^9 -Tetrahydrocannabinol (Δ^9 -THC)

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lens 60 V. Tyrosine 1,3,6 standard was used as a mass calibrant of FT mass analyzer (resolution=100000), and tyrosine 3 standard was used as a lock mass ion (m/z 508.20783) during the measurement. Theoretical mass and delta value (mmu) were calculated by using the elemental composition tool of Xcalibur/Qual Browser software (Thermo Fisher Scientific Inc.). MS data were recorded in the full scan mode (m/z 100–1000).

NMR Analysis CDCl_3 (99.96%) and $\text{DMSO-}d_6$ (99.96%) were purchased from ISOTEC Inc., which is part of Sigma-Aldrich Inc. (St. Louis, MO, U.S.A.). The NMR spectra were obtained on ECA-600 and ECA-800 spectrometers (JEOL Ltd., Tokyo, Japan). Assignments were made via $^1\text{H-NMR}$, $^{13}\text{C-NMR}$, heteronuclear multiple quantum coherence (HMQC), heteronuclear multiple-bond correlation (HMBC), double quantum filtered correlation spectroscopy (DQF-COSY), and rotating frame nuclear Overhauser effect (ROE) spectra.

Results and Discussion

In the sample solution of the product, an unknown main peak was detected by GC-EI-MS and by LC-ESI-MS analyses. The former found a peak at 47.9 min and showed four ion peaks at m/z (% relative intensity) 332 (16), 314 (14), 233 (80), and 215 (100). On the other hand, the latter detected a peak at 14.5 min and exhibited major ion peaks at m/z 333 $[\text{M}+\text{H}]^+$, 315 $[\text{M}+\text{H}-18]^+$ in the positive scan mode and at m/z 331 $[\text{M}-\text{H}]^-$ in the negative scan mode. The PDA-sliced UV spectrum of the peak exhibited maxima at 220 and 275 nm and minima at 212 and 249 nm. The accurate mass of **1** revealed $[\text{M}+\text{H}]^+$ at m/z 333.27918 in the positive scan mode and $[\text{M}-\text{H}]^-$ at 331.26442 in the negative scan mode, suggesting molecular formulae of $\text{C}_{22}\text{H}_{37}\text{O}_2$ and $\text{C}_{22}\text{H}_{35}\text{O}_2$, respectively. The errors between the observed mass and theoretical mass of $[\text{M}+\text{H}]^+$ and $[\text{M}-\text{H}]^-$ are +0.71 and -0.18 mmu, respectively. The ^1H - and ^{13}C -NMR spectra of **1** exhibited 36 protons and 22 carbons. These results suggested that **1** contained oxygen atoms but no nitrogen atoms.

The ^1H -NMR spectrum of **1** exhibited 36 non-exchangeable protons, including three methyl signals at δ 1.22 (6H, s) and 0.83 (3H, t, $J=7.2$ Hz), as well as ABX-type aromatic proton signals at δ 7.06 (1H, d, $J=8.2$ Hz), 6.84 (1H, dd, $J=8.2, 2.0$ Hz), and 6.67 (1H, d, $J=2.0$ Hz), as shown in Table 1. In addition, the ^1H -NMR spectrum also showed two methine proton signals at δ 2.86 (1H, tt, $J=12.4, 3.1$ Hz) and 3.76 (1H, tt, $J=11.0, 4.1$ Hz), and a characteristic signal assignable to hydroxy proton at δ 4.51 (1H, br d, $J=4.6$ Hz) and 9.01 (1H, br s). The ^{13}C -NMR spectrum of **1** showed 22 carbon signals, including three methyls, ten methylenes, two methines with one oxygenated carbon (δ 71.2) and one quaternary carbon, three aromatic carbons (δ 113.1, 118.5, 126.3), and three quaternary carbons (δ 128.7, 149.1, 152.3). The presence of three partial structures (1,3-substituted cyclohexyl group, 1,1-dimethyloctyl group, and 1,2,4-substituted phenyl) was suggested from its DQF-COSY, HMQC, and HMBC spectra (Table 1, Fig. 2). The connectivity of these groups was deduced from the HMBC spectrum (Table 1, Fig. 2). A methine proton at δ 2.86 (H-3) of the cyclohexyl group correlated to the phenyl carbons at δ 152.3 and 126.3 (C-2', C-6'), and two aromatic protons, at δ 6.67 and 6.84 (H-3', H-5') of the phenyl group, showed correlations to the quaternary carbon at δ 37.3 (C-1'). In addition, the irradiation of the hydroxyl proton at δ 9.01 (2'-OH) resulted in ROE on the aromatic proton (H-3'), as shown in Fig. 3. The relative configuration between two methine protons at C-1 and C-3 established a *cis* configuration by the ROE

Table 1. NMR Data of Compound **1** in CDCl_3 ^{a)}

No.	^{13}C	^1H	HMBC ^{b)}
1	71.2	3.76, 1H, tt, $J=11.0, 4.1$ Hz	2, 6 ^{c)}
2	41.9	ax, 1.44, 1H, m, overlapped eq, 2.16, 1H, br d, $J=11.7$ Hz	3, 4, 1' 1, 3, 4
3	35.3	2.86, 1H, tt, $J=12.4, 3.1$ Hz	2, 4, 1', 2', 6'
4	31.7	ax, 1.30, 1H, m, overlapped eq, 1.82, 1H, d, $J=13.1$ Hz	2, 3, 5, 1' ^{c)} 2
5	24.5	ax, 1.44, 1H, m, overlapped eq, 1.87, 1H, dq, $J=13.4, 3.4$ Hz	1, 4, 6 1, 4, 6
6	35.5	ax, 1.27, 1H, m, overlapped eq, 2.05, 1H, br d, $J=12.0$ Hz	1, 2, 5 1, 2, 4
1'	128.7	—	—
2'	152.3	—	—
3'	113.1	6.67, 1H, d, $J=2.0$ Hz	1', 2', 4', 5', 1''
4'	149.1	—	—
5'	118.5	6.84, 1H, dd, $J=8.2, 2.0$ Hz	1', 2', 3', 6', 1''
6'	126.3	7.06, 1H, d, $J=8.2$ Hz	3, 2', 3', 4', 5'
1''	37.3	—	—
2''	44.6	1.51, 2H, m	4', 1'', 3'', 4'', 1''-(CH_3) ₂
3''	24.7	1.04, 2H, m	2'', 4'', 5''
4''	30.3	1.17, 2H, m, overlapped	2'', 3'', 5''
5''	29.2	1.17, 2H, m, overlapped	3'', 7''
6''	31.9	1.17, 2H, m, overlapped	5'', 7''
7''	22.6	1.24, 2H, m	6'', 8''
8''	14.1	0.83, 3H, t, $J=7.2$ Hz	6'', 7''
1''-(CH_3) ₂	28.9	1.22, 6H, s	4', 1'', 2''
1-OH	—	4.51, 1H, br d, $J=4.6$ Hz ^{d)}	1, 2, 6 ^{d)}
2'-OH	—	9.01, 1H, br s ^{d)}	1', 2', 3' ^{d)}

a) Recorded in CDCl_3 at 600 and 800 MHz (^1H) and 150 and 200 MHz (^{13}C), respectively; data in δ ppm (J in Hz). b) $J=8$ Hz, the proton signal correlated with the indicated carbons. c) $J=4$ Hz. d) Recorded in $\text{DMSO-}d_6$.

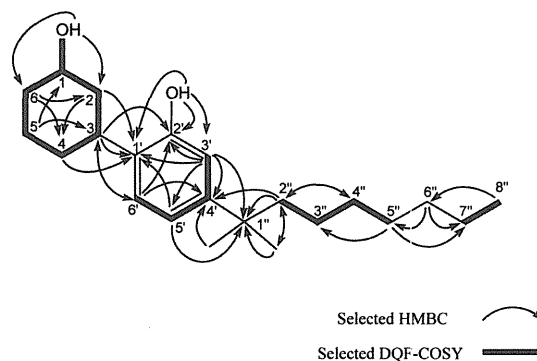


Fig. 2. Selected DQF-COSY and HMBC Correlations of Compound **1**

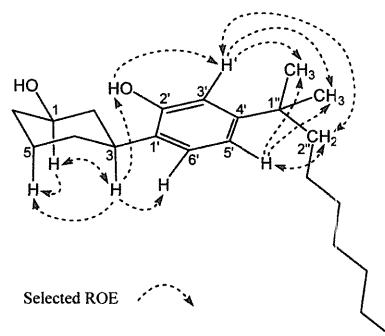


Fig. 3. Selected ROE Correlations of Compound **1**

correlations (Fig. 3). Therefore, the structure of **1** is finally elucidated as (1*RS*,3*SR*)-3-[4-(1,1-dimethyloctyl)-2-hydroxyphenyl]cyclohexan-1-ol.

The deduced structure has already been synthesized by Pfizer Inc. and reported as a cannabinoid analog.^{25,26} Pharmaceutical studies showed that **1** has potent cannabinoid receptor binding activity *in vitro* and analgesic activity *in vivo* in mice.^{27–30} Compton *et al.* reported that compound **1** was approximately 5-fold more potent than Δ^9 -THC at the viewpoint of pharmacological activity.²⁸

This is the first case in which **1** has been detected as a designer drug and an ingredient in a herbal product. Pfizer Inc. has also reported many analogs of **1** and has described their synthesis with pharmacological data.^{19,22,31,32} Additionally, various cannabinoid analogs are synthesized one after another and their pharmacological activity studied for the development of new useful drugs for the treatment of a number of diseases.^{23,24} This situation alerts us that these described cannabinoid analogs other than **1** may be found as designer drugs or adulterants in illegal products in the near future. To avoid health problems and abuse caused by new designer drugs, we have to continuously monitor such compounds during our surveillance.

Acknowledgments Part of this work was supported by a Health and Labor Sciences Research Grant from the Ministry of Health, Labour and Welfare, Japan.

References

- 1) Satake M., *Kokuritsu Iyakuhin Shokuhin Eisei Kenkyusho Hokoku (Bulletin of National Institute of Health Sciences)*, **118**, 204–205 (2000).
- 2) Goda Y., *Kokuritsu Iyakuhin Shokuhin Eisei Kenkyusho Hokoku*, **119**, 121–123 (2001).
- 3) Goda Y., *Kokuritsu Iyakuhin Shokuhin Eisei Kenkyusho Hokoku*, **120**, 143–145 (2002).
- 4) Goda Y., *Kokuritsu Iyakuhin Shokuhin Eisei Kenkyusho Hokoku*, **121**, 152–154 (2003).
- 5) Goda Y., *Kokuritsu Iyakuhin Shokuhin Eisei Kenkyusho Hokoku*, **122**, 70–73 (2004).
- 6) Goda Y., *Kokuritsu Iyakuhin Shokuhin Eisei Kenkyusho Hokoku*, **123**, 80–83 (2005).
- 7) Goda Y., *Kokuritsu Iyakuhin Shokuhin Eisei Kenkyusho Hokoku*, **124**, 94–98 (2006).
- 8) Goda Y., *Kokuritsu Iyakuhin Shokuhin Eisei Kenkyusho Hokoku*, **125**, 129–132 (2007).
- 9) Goda Y., *Kokuritsu Iyakuhin Shokuhin Eisei Kenkyusho Hokoku*, **126**, 137–141 (2008).
- 10) Kikura-Hanajiri R., Kawamura M., Saisho K., Kodama Y., Goda Y., *J. Chromatogr. B*, **855**, 121–126 (2007).
- 11) Uchiyama N., Kikura-Hanajiri R., Kawahara N., Goda Y., *Yakugaku Zasshi*, **128**, 1499–1505 (2008).
- 12) Kikura-Hanajiri R., Hayashi M., Saisho K., Goda Y., *J. Chromatogr. B*, **825**, 29–37 (2005).
- 13) Ministry of Health, Labour and Welfare, Japan, websites: listed in (<http://www.mhlw.go.jp/shingi/2006/11/s1109-4.html>) [in Japanese].
- 14) Kikura-Hanajiri R., Kawamura M., Uchiyama N., Ogata J., Kamakura H., Saisho K., Goda Y., *Yakugaku Zasshi*, **128**, 971–979 (2008).
- 15) Uchiyama N., Kawamura M., Kamakura H., Kikura-Hanajiri R., Goda Y., *Yakugaku Zasshi*, **128**, 981–987 (2008).
- 16) Ministry of Health, Labour and Welfare, Japan, websites: listed in (<http://www.mhlw.go.jp/shingi/2007/09/s0920-4.html>) [in Japanese].
- 17) Razdan R. K., *Pharmacol. Rev.*, **38**, 75–149 (1986).
- 18) Rapaka R. S., Makriyannis A., “NIDA Research Monograph,” Vol. 79, National Institute on Drug Abuse, Rockville, MD, 1987.
- 19) Howlett A. C., Johnson M. R., Melvin L. S., “NIDA Research Monograph,” Vol. 96, National Institute of Drug Abuse, Rockville, MD, 1990, pp. 100–111.
- 20) Johnson M. R., Melvin L. S., Milne G. M., *Life Sci.*, **31**, 1703–1706 (1982).
- 21) Weissman A., Milne G. M., Melvin L. S., *J. Pharmacol. Exp. Ther.*, **223**, 516–523 (1982).
- 22) Melvin L. S., Johnson M. R., Herbert C. A., Milne G. M., Weissman A. A., *J. Med. Chem.*, **27**, 67–71 (1984).
- 23) Pacher P., Batkai S., Kunos G., *Pharmacol. Rev.*, **58**, 389–462 (2006).
- 24) Kulkarni S. K., Ninan I., *Indian J. Pharmacol.*, **33**, 170–184 (2001).
- 25) Herbert C. A., Johnson M. R., Melvin L. S. Jr., DE Patent 2839836 (1979).
- 26) Thomas B. F., Compton D. R., Martin B. R., *J. Pharmacol. Exp. Ther.*, **255**, 624–630 (1990).
- 27) Melvin L. S., Milne G. M., Johnson M. R., Subramaniam B., Wilken G. H., Howlett A. C., *Mol. Pharmacol.*, **44**, 1008–1015 (1993).
- 28) Compton D. R., Johnson M. R., Melvin L. S., Martin B. R., *J. Pharmacol. Exp. Ther.*, **260**, 201–209 (1992).
- 29) Compton D. R., Rice K. C., De Costa B. R., Razdan R. K., Melvin L. S., Johnson M. R., Martin B. R., *J. Pharmacol. Exp. Ther.*, **265**, 218–226 (1993).
- 30) Martin B. R., Compton D. R., Thomas B. F., Prescott W. R., Little P. J., Razdan R. K., Johnson M. R., Melvin L. S., Mechoulam R., Ward S. J., *Pharmacol. Biochem. Behav.*, **40**, 471–478 (1991).
- 31) Howlett A. C., Johnson M. R., Melvin L. S., Milne G. M., *Mol. Pharmacol.*, **33**, 297–302 (1988).
- 32) Devane W. A., Dysarz F. A. 3rd., Johnson M. R., Melvin L. S., Howlett A. C., *Mol. Pharmacol.*, **34**, 605–613 (1988).

Identification of a cannabimimetic indole as a designer drug in a herbal product

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Abstract A cannabimimetic indole has been identified as a new adulterant in a herbal product being sold illegally in Japan for its expected narcotic effect. Liquid chromatography-mass spectrometry and gas chromatography-mass spectrometry analyses indicated that the product contained two major compounds. One was identified as a cannabinoid analog (1*RS*,3*SR*)-3-[4-(1,1-dimethyloctyl)-2-hydroxyphenyl]cyclohexan-1-ol (**1**) by direct comparison with the authentic compound, which we reported previously. The other compound (**2**) showed a molecular weight of 341 daltons, and accurate mass spectral measurements showed its elemental composition to be C₂₄H₂₃NO. Both mass and nuclear magnetic resonance spectrometric data revealed that **2** was 1-pentyl-3-(1-naphthoyl)indole [or naphthalen-1-yl-(1-pentylindol-3-yl)methanone] being identical to JWH-018, which was synthesized by Wiley and coworkers in 1998. This compound was reported as a potent cannabinoid receptor agonist possessing a pharmacological cannabimimetic activity.

Keywords 1-Pentyl-3-(1-naphthoyl)indole · Naphthalen-1-yl-(1-pentylindol-3-yl)methanone · JWH-018 · Cannabimimetic indole · Designer drug · Herbal product

Introduction

Various psychotropic substances are being sold and distributed around the world via the Internet. Most recently, we found a synthetic cannabinoid analog (1*RS*,3*SR*)-3-[4-(1,1-dimethyloctyl)-2-hydroxyphenyl]cyclohexan-1-ol (**1**) [1], which contains no amino groups (Fig. 1), as an adulterant in a herbal product being commercially sold as an incense. This was the first report to identify a synthetic cannabinoid analog in a herbal product distributed on the illegal drug market for its expected narcotic effect. At almost the same time, we found another compound (**2**) that was also thought to be an adulterant in the same type of herbal products. This compound was finally found to be identical to JWH-018, a cannabimimetic aminoalkyl naphthoyl indole derivative; it had been first synthesized by Huffman and coworkers in 1998, and was reported as a potent cannabinoid receptor agonist possessing a cannabimimetic pharmacological activity in vivo [2–5]. Also, in January 2009, the Health Minister of Germany announced that **2** is an active component in a mislabeled mixture of herbs; **2** has been a controlled substance in Germany since 22 January 2009 [6]. However, no scientific report describing the isolation and identification of this compound from herbal products has been published. The present report deals with the details of its identification in a herbal product by various instrumental analyses.

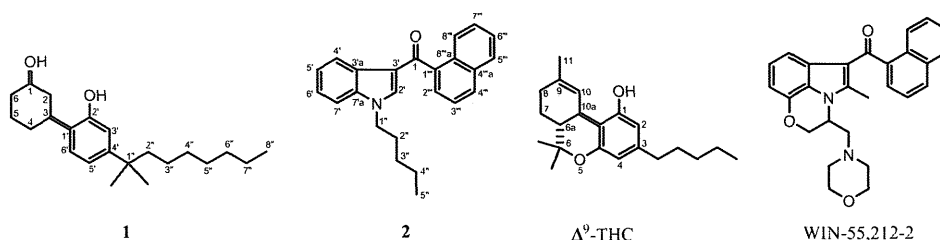
Materials and methods

Materials and preparation

Acetonitrile (high-performance liquid chromatography grade) and all other chemicals (analytical grade) were

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Fig. 1 Structures of detected compounds **1**, **2** and related compounds [Δ^9 -tetrahydrocannabinol (Δ^9 -THC) and WIN-55,212-2]



obtained from Wako (Osaka, Japan). A product, described as a herbal mixture and having the appearance of dried plants, was purchased via the Internet (December 2008). A 10-mg portion of the product was crushed into powder and extracted with 1 ml of methanol under ultrasonication for 10 min. After centrifugation for 5 min at 3000 rpm, the supernatant solution was passed through a centrifugal filter (Ultrafree-MC, 0.45 μ m filter unit, Millipore, Bedford, MA, USA).

Instrumental analyses

Gas chromatography-mass spectrometry (GC-MS) was used in the electron impact (EI) mode at 70 eV of electron energy. The analysis was performed on a Hewlett-Packard 6890N GC with a 5975 mass-selective detector (Agilent, Palo Alto, CA, USA) using a capillary column (HP1-MS capillary, 30 m \times 0.25 mm i.d., 0.25 μ m film thickness, Agilent) and helium as carrier gas. An initial column temperature of 80°C was employed, and it was increased at a rate of 5°C/min to 190°C and then at a second rate of 10°C/min up to 310°C. The data were obtained in the full scan mode with a scan range of m/z 40–550. The analysis was performed under the same conditions as used in the analysis of designated drugs (Shitei-Yakubutsu) controlled by the Pharmaceutical Affairs Law of Japan [7].

The MS analysis was also made by liquid chromatography-electrospray ionization-mass spectrometry (LC-ESI-MS). The instrument consisted of an ACQUITY ultra-performance LC system connected with a single quadrupole mass detector and a photodiode array (PDA) detector (Waters, Milford, MA, USA). The sample solutions were separated using an ACQUITY UPLC HSS T3 column (2.1 mm i.d. \times 100 mm, 1.8 μ m; Waters) protected by a Van Guard column (2.1 mm i.d. \times 5 mm, 1.8 μ m; Waters) at 40°C. The following gradient system was used with mobile phase A (0.1% formic acid in water) and mobile phase B (0.1% formic acid in acetonitrile) delivered at 0.3 ml/min; 50% A/50% B for 3 min, changing to 20% A/80% B over 2 min and held with the final composition over 5 min. The injection volume was 1 μ l. The wavelength of the PDA detector for screening

was set from 190 to 500 nm, and chromatographic peaks were monitored at 275 nm. Mass analysis by ESI was used in both positive and negative modes. Nitrogen gas was used for desolvation at a flow rate of 650 l/h at 350°C. The capillary and cone voltages were 3000 V and 30 V, respectively. MS data were recorded in the full scan mode (m/z 150–700).

The accurate mass spectrum of the target compound was measured using a direct analysis in real time (DART) ion source coupled to a time-of-flight (TOF) mass spectrometer (AccuTOF JMS-100LC, JEOL, Tokyo, Japan) operated in the positive ion mode. The measurements were made with the ion guide peak voltage set at 500 V, the reflectron voltage at 950 V, orifice 1 voltage at 15 V, orifice 2 voltage at 5 V, ring lens voltage at 5 V, and the orifice 1 temperature at 80°C. The mass range was 100–500 daltons. The DART ion source was used at a helium gas flow rate of 2.0 l/min, the gas heater temperature at 250°C, the discharge electrode needle setting at 3200 V, electrode 1 at 100 V, and electrode 2 at 250 V. Internal mass number calibration was achieved using PEG600, and diphenhydramine was used as an internal standard for each analysis.

For nuclear magnetic resonance (NMR) analysis, CDCl_3 (99.96%) was purchased from ISOTEC, a part of Sigma-Aldrich (St. Louis, MO, USA). The NMR spectra were obtained on ECA-600 and ECA-800 spectrometers (JEOL). Assignments were made via ^1H NMR, ^{13}C NMR, heteronuclear multiple quantum coherence (HMQC), heteronuclear multiple-bond correlation (HMBC), double quantum filtered correlation spectroscopy (DQF-COSY), and rotating frame nuclear overhauser effect (ROE) spectra.

Isolation of compound **2**

A 3-g portion of the herbal product was extracted with 100 ml of methanol by ultrasonication for 1 h. After the extraction was repeated three times, the combined supernatant was evaporated to dryness. The extract was loaded on a preparative silica gel thin layer chromatography (TLC) plate (Silica Gel 60, 20 \times 20 cm, 2 mm, Merck, Darmstadt, Germany) using hexane/acetone

(4:1) as developing solvent. A portion of the silica gel in the TLC plate was taken and eluted with CH_2Cl_2 /methanol (2:1) to give fraction 1. Repeated fractionation of fraction 1 by preparative silica gel TLC with hexane/ CH_2Cl_2 (1:20) gave compound **2** (15 mg) as an off-white solid.

Results and discussion

In the sample solution of the product, two major peaks were detected by LC-ESI-MS analysis (Fig. 2a–d). One peak, detected at 7.8 min, exhibited two ion peaks at m/z 333 $[\text{M}+\text{H}]^+$ and at 315 $[\text{M}+\text{H}-18]^+$ in the positive scan mode (Fig. 2e). A comparison with the mass spectrum of the authentic compound revealed that this peak was (1*R*S,3*S*R)-3-[4-(1,1-dimethyloctyl)-2-hydroxyphenyl]cyclohexan-1-ol (**1**) (Fig. 1), which was reported as an adulterant in a herbal product in our previous study [1]. Another unknown peak (**2**) detected

at 7.5 min showed a major peak at m/z 342 $[\text{M}+\text{H}]^+$ (Fig. 2f). The PDA-sliced ultraviolet (UV) spectrum of the peak (**2**) exhibited maxima at 218, 247, and 314 nm and minima at 239 and 285 nm (Fig. 2h). These characteristics were completely different from those of **1** (UV λ_{max} 220, 275 nm; λ_{min} 212, 249 nm, Fig. 2g).

GC-EI-MS analysis showed two major peaks with a peak of α -tocopherol, which had been added as an antioxidant (Fig. 3a). One peak, detected at 47.9 min, showed a mass spectrum with four ion peaks at m/z (relative intensity) 332 (16), 314 (14), 233 (80), and 215 (100) as shown in Fig. 3b, which was identical to the mass spectrum of the authentic compound (**1**). An unknown peak (**2**), detected at 51.7 min, showed a mass spectrum with five ion peaks at m/z 341 (100), 324 (43), 284 (58), 214 (52), and 127 (32), as shown in Fig. 3c.

The accurate mass spectrum measured by TOF-MS showed a protonated molecular ion peak ($[\text{M}+\text{H}]^+$) at m/z 342.18579 in the positive mode, suggesting that the molecular formula of **2** was $\text{C}_{24}\text{H}_{24}\text{NO}$. The error between

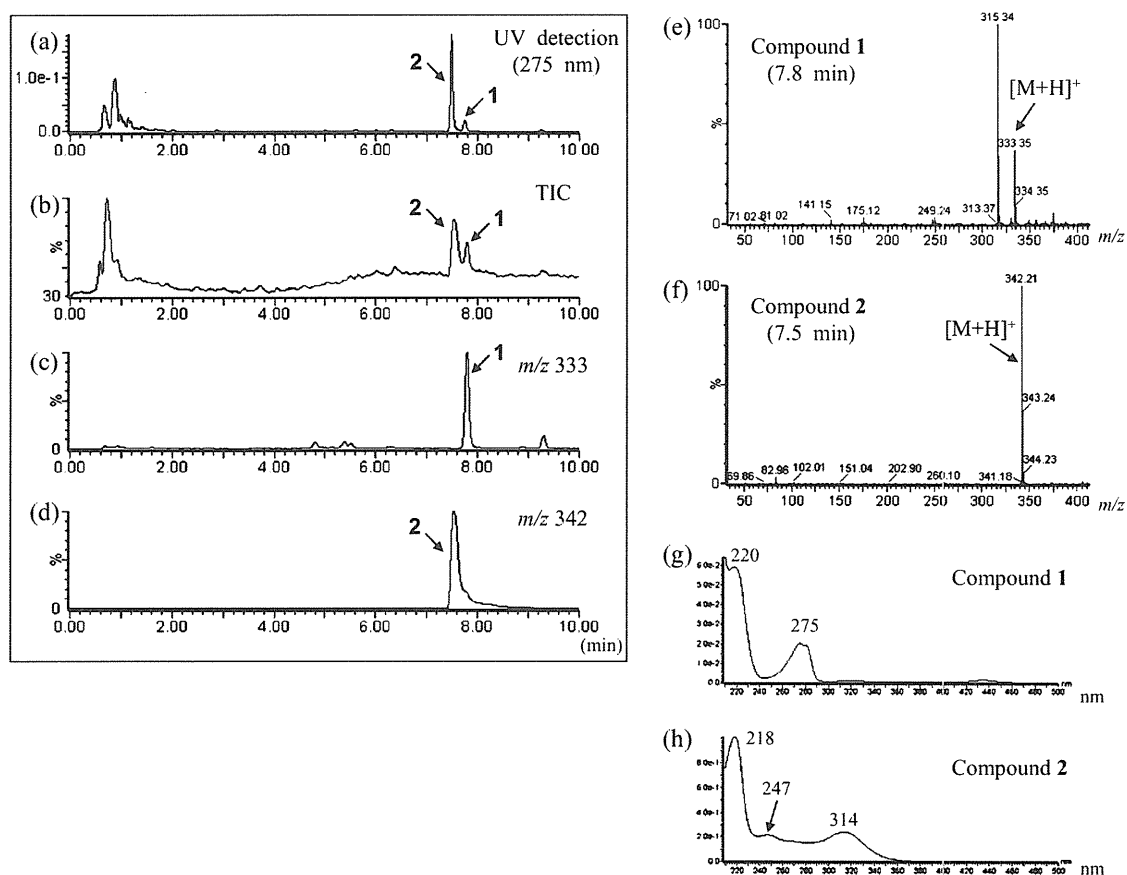


Fig. 2a–h Data from high-performance liquid chromatography with ultraviolet detection (**a**, **g**, **h**) and liquid chromatography-electrospray ionization-mass spectrometry (**b–f**) for the extract of the sample. Total ion chromatogram (**b**), mass chromatograms at

m/z 333 (**1**) (**c**) and m/z 342 (**2**) (**d**), electrospray ionization mass spectra (**e**, **f**) and ultraviolet spectra (**g**, **h**) of each peak are shown

Fig. 3 Total ion chromatogram (a) and electron impact mass spectra of the peaks detected at 47.9 min (1) (b) and 51.7 min (2) (c) measured by gas chromatography-mass spectrometry

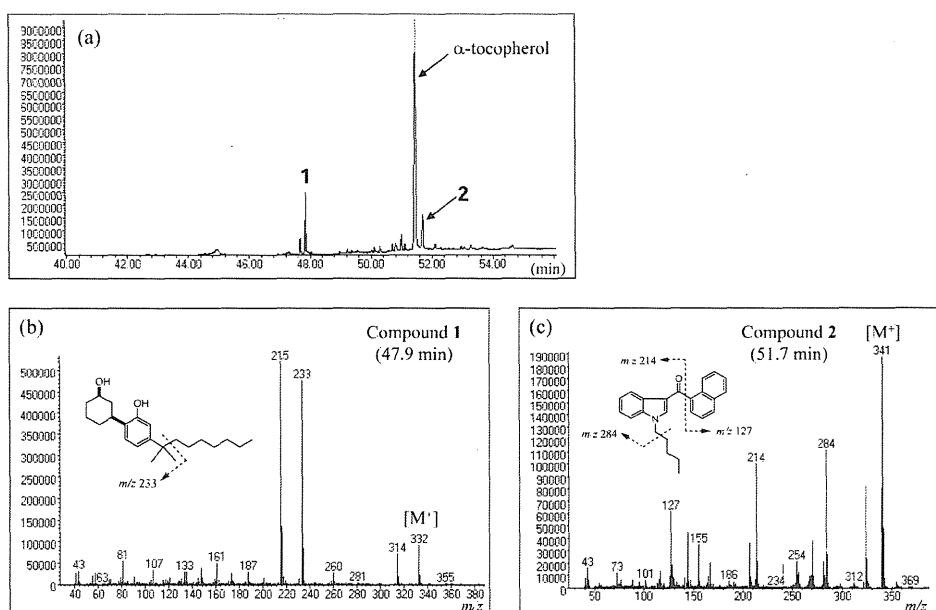


Table 1 Nuclear magnetic resonance data of compound 2

No.	¹³ C	¹ H	HMBC ^a
1	192.0	–	–
2'	137.9	7.33, 1H, s, overlapped	1, 3', 3'a, 7'a, 1''
3'	117.5	–	–
3'a	127.0	–	–
4'	122.9	8.47, 1H, m	3', 3'a, 6', 7'a
5'	122.8	7.35, 1H, m, overlapped	7'
6'	123.6	7.35, 1H, m, overlapped	7'a
7'	110.0	7.38, 1H, m, overlapped	3'a, 5', 7'a
7'a	137.0	–	–
1''	47.2	4.05, 2H, t, <i>J</i> = 7.4 Hz	2', 7'a, 2'', 3''
2''	29.5	1.79, 2H, quint, <i>J</i> = 7.4 Hz	1'', 3'', 4''
3''	28.9	1.24, 2H, m, overlapped	1'', 4'', 5''
4''	22.2	1.28, 2H, m, overlapped	2'', 3'', 5''
5''	13.8	0.83, 3H, t, <i>J</i> = 7.0 Hz	3'', 4''
1'''	139.1	–	–
2'''	125.8	7.64, 1H, dd, <i>J</i> = 7.1, 1.3 Hz	1, 3''', 4''', 8'''a
3'''	124.5	7.51, 1H, dd, <i>J</i> = 8.3, 7.1 Hz, overlapped	1''', 2''', 4'''a
4'''	129.9	7.95, 1H, brd, <i>J</i> = 8.3 Hz	2''', 4'''a, 5''', 8'''a
4'''a	133.7	–	–
5'''	128.1	7.90, 1H, brd, <i>J</i> = 8.3 Hz	4''', 7''', 8'''a
6'''	126.3	7.50, 1H, td, <i>J</i> = 6.9, 1.4 Hz, overlapped	4'''a, 7''', 8'''
7'''	126.7	7.45, 1H, ddd, <i>J</i> = 8.3, 6.9, 1.4 Hz	5''', 8'''a
8'''	126.0	8.17, 1H, brd, <i>J</i> = 8.3 Hz	1''', 4'''a, 6''', 8'''a
8'''a	130.8	–	–

Recorded in CDCl₃ at 600 and 800 MHz (¹H) and 150 and 200 MHz (¹³C), respectively; data in δ ppm

^aFor heteronuclear multiple-bond correlation (HMBC), *J* = 8 Hz, the proton signal correlated with the indicated carbons

the mass number observed and theoretical mass number of [M+H]⁺ was –0.10 amu.

The ¹H NMR spectrum of 2 showed 23 nonexchangeable protons, including a methyl signal at δ 0.83 (3H, t, *J* = 7.0 Hz), AB₂-type aromatic proton signals at δ 7.51 (1H, dd, *J* = 8.3, 7.1 Hz), 7.64 (1H, dd, *J* = 7.1, 1.3 Hz), and 7.95 (1H, brd, *J* = 8.3 Hz), and AA'BB'-type aromatic proton signals at δ 7.45 (1H, ddd, *J* = 8.3, 6.9,

1.4 Hz), 7.50 (1H, td, *J* = 6.9, 1.4 Hz), 7.90 (1H, brd, *J* = 8.3 Hz), and 8.17 (1H, brd, *J* = 8.3 Hz) as shown in Table 1. In addition, the ¹H NMR spectrum also showed three methylene proton signals, at δ 1.24 and 1.28 (each 2H, m) and at 1.79 (2H, quint, *J* = 7.4 Hz), as well as a characteristic methylene signal connected to a nitrogen atom at δ 4.05 (2H, t, *J* = 7.4 Hz). The ¹³C NMR spectrum of 2 showed 24 carbon signals, suggesting the

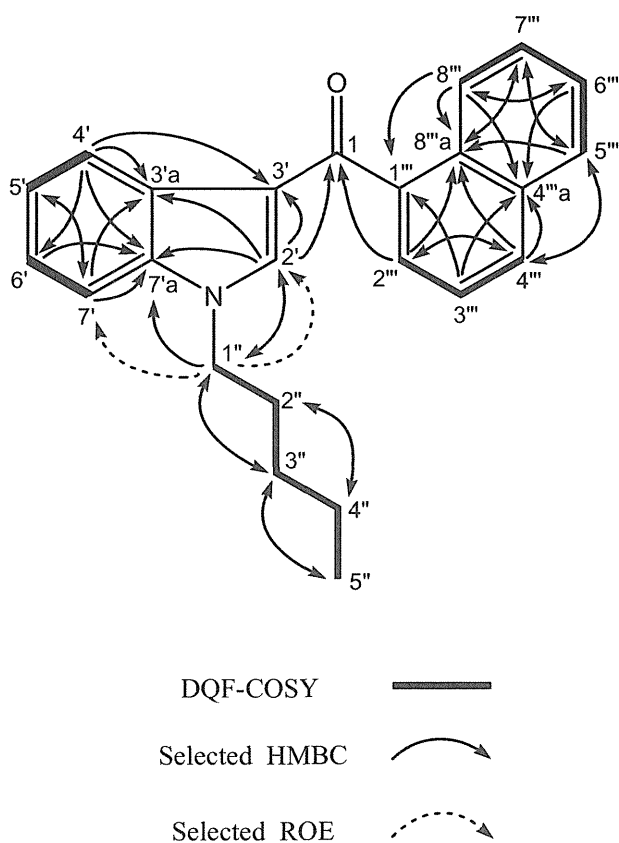


Fig. 4 Selected correlations for compound **2** by two-dimensional nuclear magnetic resonance spectroscopy techniques. *DQF-COSY*, Double quantum filtered correlation spectroscopy; *HMBC*, heteronuclear multiple-bond correlation spectroscopy; *ROE*, rotating frame nuclear overhauser effect spectroscopy

presence of a methyl, 4 methylenes with a nitrogenated carbon (δ 47.2), 12 aromatic carbons (δ 110.0, 122.8, 122.9, 123.6, 124.5, 125.8, 126.0, 126.3, 126.7, 128.1, 129.9, and 137.9), 6 aromatic quaternary carbons (δ 117.5, 127.0, 130.8, 133.7, 137.0, and 139.1), and a carbonyl carbon (δ 192.0). The presence of three partial structures (a 1,3-substituted indole group, a 1-substituted naphthalene group, and an *n*-pentyl group) was suggested from the DQF-COSY, HMQC, and HMBC spectra (Table 1, Fig. 4). The connectivity of these groups and the carbonyl group was deduced from the HMBC spectrum. An aromatic proton at δ 7.33 (H-2') of the indole group correlated to the carbonyl carbon at δ 192.0 (C-1), and the methylene carbon at δ 47.2 (C-1'') of the *n*-pentyl group and an aromatic proton at δ 7.64 (H-2''') of the naphthalene group showed correlations to the carbonyl carbons at δ 192.0 (C-1). In addition, irradiation of the methylene protons at δ 4.05 (H-1'') of the *n*-pentyl group resulted in ROE responses by the aromatic protons (H-2' and H-7') as shown in Fig. 4.

On the basis of the mass spectra (Figs. 2, 3) and NMR data (Table 1, Fig. 4), the structure of **2** was finally elucidated as 1-pentyl-3-(1-naphthoyl)indole [or naphthalen-1-yl-(1-pentylindol-3-yl)methanone]. The deduced compound had been already synthesized and named JWH-018 by Wiley et al. [2] in 1998. This compound is a potent cannabinoid receptor agonist possessing a pharmacological activity of a cannabinoid in vivo [2–5]. Wiley et al. [2] described that **2** showed a 4.5-fold more potent affinity for the CB₁ receptor ($K_i = 9 \pm 5$ nM) than did Δ^9 -tetrahydrocannabinol (Δ^9 -THC, Fig. 1), which is psychoactive and a major constituent of *Cannabis sativa* L. (cannabis, hemp, marijuana, marihuana) ($K_i = 41 \pm 2$ nM). Compound **2** produced potent cannabinoid effects of antinociception, hypomobility, hypothermia, and ring immobility in in vivo assays [2,3]. In the present study, we have identified compound **2** as a designer drug and an adulterant together with **1** in a herbal product.

The synthesis of many analogs of **1** and **2** together with pharmacological data has been already described [2–5,8–11]. In the past few decades, a number of analogs of Δ^9 -THC have been synthesized based on the partially reduced dibenzopyran structure of THC, and their structure–activity relationships were studied [12,13]. In the 1980s, a group at Pfizer explored the development of analgesics using potent synthetic nontraditional cannabinoids, which lack the dibenzopyran structure present in the traditional cannabinoids but exhibit typical cannabinoid pharmacological effects [14–22]. On the other hand, D'Ambra et al. [23] reported in 1992 that aminoalkylindoles, such as WIN-55212-2, were bound to a cannabinoid brain receptor with high affinity (Fig. 1). A subsequent study by Huffman et al. [24] established that an aminoalkyl portion of the molecule, such as WIN-55212-2, could be replaced by an alkyl group to provide indole derivatives that have higher affinity for the brain receptor and exhibit typical cannabinoid pharmacological effects in vivo. These authors also described the structure–activity relationships of indole-derived, pyrrole-derived, and indene-derived cannabinoids [2,3,11]. After the discovery of cannabinoid receptors, CB₁ (central type) and CB₂ (peripheral type), as well as the discovery of endogenous cannabinoids, their physiological roles were elucidated to some extent [25]. A number of cannabinoid analogs, such as derivatives based on THC, indole, pyrrole, indene, and pyrazole, were then newly synthesized and their pharmacological activities applicable to the treatments of various diseases were studied [26,27]. This situation alerts us that these cannabinoid analogs other than **1** and **2** will be found as designer drugs or adulterants in illegal products as cannabis replacements in the near future. To avoid health problems and abuse caused by new designer

drugs, we must continuously monitor such compounds through surveillance.

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References

- Uchiyama N, Kikura-Hanajiri R, Kawahara N, Haishima Y, Goda Y (2009) Identification of a cannabinoid analog as a new type of designer drug in a herbal product. *Chem Pharm Bull* 57(4), (in press)
- Wiley JL, Compton DR, Dai D, Lainton JA, Phillips M, Huffman JW, Martin BR (1998) Structure–activity relationships of indole- and pyrrole-derived cannabinoids. *J Pharmacol Exp Ther* 285:995–1004
- Huffman JW (1999) Cannabimimetic indoles, pyrroles and indenes. *Curr Med Chem* 6:705–720
- Aung MM, Griffin G, Huffman JW, Wu M, Keel C, Yang B, Showalter VM, Abood ME, Martin BR (2000) Influence of the *N*-1 alkyl chain length of cannabimimetic indoles upon CB₁ and CB₂ receptor binding. *Drug Alcohol Depend* 60:133–140
- Huffman JW, Mabon R, Wu MJ, Lu J, Hart R, Hurst DP, Reggio PH, Wiley JL, Martin BR (2003) 3-Indolyl-1-naphthylmethanes: new cannabimimetic indoles provide evidence for aromatic stacking interactions with the CB₁ cannabinoid receptor. *Bioorg Med Chem* 11:539–549
- Zweiundzwanzigste Verordnung, zur Änderung betäubungsmittelrechtlicher Vorschriften (2009), Germany. BGBl I Nr. 3 vom 21.01.2009, 22. BtMÄndV vom 19. Januar 2009, S. 49–50, <http://www.bgblportal.de/BGBl/bgbl1f/bgbl109s0049.pdf>. Accessed 19 Jan 2009
- Kikura-Hanajiri R, Kawamura M, Uchiyama N, Ogata J, Kamakura H, Saisho K, Goda Y (2008) Analytical data of designated substances (Shitei-Yakubutsu) controlled by the Pharmaceutical Affairs Law in Japan, part I: GC-MS and LC-MS. *Yakugaku Zasshi* 128:971–979
- Martin BR, Wiley JL, Beletskaya I, Sim-Selley LJ, Smith FL, Dewey WL, Cottney J, Adams J, Baker J, Hill D, Saha B, Zerkowski J, Mahadevan A, Razdan RK (2006) Pharmacological characterization of novel water-soluble cannabinoids. *J Pharmacol Exp Ther* 318:1230–1239
- Howlett AC, Johnson MR, Melvin LS, Milne GM (1988) Nonclassical cannabinoid analgesics inhibit adenylate cyclase: development of a cannabinoid receptor model. *Mol Pharmacol* 33:297–302
- Devane WA, Dysarz FA, Johnson MR, Melvin LS, Howlett AC (1988) Determination and characterization of a cannabinoid receptor in rat brain. *Mol Pharmacol* 34:605–613
- Huffman JW, Padgett LW (2005) Recent developments in the medicinal chemistry of cannabimimetic indoles, pyrroles and indenes. *Curr Med Chem* 12:1395–1411
- Razdan RK (1986) Structure–activity relationships in cannabinoids. *Pharmacol Rev* 38:75–149
- Rapaka RS, Makriyannis A (1987) Structure–activity relationships of the cannabinoids. *NIDA Res Monogr* 79:1–216
- Thomas BF, Compton DR, Martin BR (1990) Characterization of the lipophilicity of natural and synthetic analogs of Δ^9 -tetrahydrocannabinol and its relationship to pharmacological potency. *J Pharmacol Exp Ther* 255:624–630
- Melvin LS, Milne GM, Johnson MR, Subramaniam B, Wilken GH, Howlett AC (1993) Structure–activity relationships for cannabinoid receptor-binding and analgesic activity: studies of bicyclic cannabinoid analogs. *Mol Pharmacol* 44:1008–1015
- Compton DR, Johnson MR, Melvin LS, Martin BR (1992) Pharmacological profile of a series of bicyclic cannabinoid analogs: classification as cannabimimetic agents. *J Pharmacol Exp Ther* 260:201–209
- Compton DR, Rice KC, De Costa BR, Razdan RK, Melvin LS, Johnson MR, Martin BR (1993) Cannabinoid structure–activity relationships: correlation of receptor binding and in vivo activities. *J Pharmacol Exp Ther* 265:218–226
- Martin BR, Compton DR, Thomas BF, Prescott WR, Little PJ, Razdan RK, Johnson MR, Melvin LS, Mechoulam R, Ward SJ (1991) Behavioral, biochemical, and molecular modeling evaluations of cannabinoid analogs. *Pharmacol Biochem Behav* 40:471–478
- Howlett AC, Johnson MR, Melvin LS (1990) Classical and nonclassical cannabinoids: mechanism of action–brain binding. *NIDA Res Monogr* 96:100–111
- Johnson MR, Melvin LS, Milne GM (1982) Prototype cannabinoid analgesics, prostaglandins and opiates—a search for points of mechanistic interaction. *Life Sci* 31:1703–1706
- Weissman A, Milne GM, Melvin LS (1982) Cannabimimetic activity from CP-47497, a derivative of 3-phenylcyclohexanol. *J Pharmacol Exp Ther* 223:516–523
- Melvin LS, Johnson MR, Herbert CA, Milne GM, Weissman A (1984) A cannabinoid derived prototypical analgesic. *J Med Chem* 27:67–71
- D’Ambra TE, Estep KG, Bell MR, Eissenstat MA, Josef KA, Ward SJ, Haycock DA, Baizman ER, Casiano FM, Beglin NC, Chippari SM, Grego JD, Kullnig RK, Daley GT (1992) Conformationally restrained analogues of pravadolone: nanomolar potent, enantioselective, (aminoalkyl)indole agonists of the cannabinoid receptor. *J Med Chem* 35:124–135
- Huffman JW, Dai D, Martin BR, Compton DR (1994) Design, synthesis and pharmacology of cannabimimetic indoles. *Bioorg Med Chem Lett* 4:563–566
- Maccarrone M (2008) Good news for CB₁ receptors: endogenous agonists are in the right place. *Br J Pharmacol* 153:179–181
- Pacher P, Batkai S, Kunos G (2006) The endocannabinoid system as an emerging target of pharmacotherapy. *Pharmacol Rev* 58:389–462
- Kulkarni SK, Ninan I (2001) Current concepts in cannabinoid pharmacology. *Indian J Pharmacol* 33:170–184

植物系違法ドラッグ製品及び法規制植物試料の Direct Analysis in Real Time (DART)-TOFMS を用いた迅速スクリーニング法の検討

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Simple and Rapid Screening for Psychotropic Natural Products Using Direct Analysis in Real Time (DART)-TOFMS

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Direct Analysis in Real Time (DART) is a novel ionization technique that provides for the rapid ionization of small molecules under ambient conditions. To investigate the trend of non-controlled psychotropic plants of abuse in Japan, a rapid screening method, without sample preparation, was developed using DART-time of flight mass spectrometer (TOFMS) for plant products. The major psychotropic constituents of these products were determined using liquid chromatography-mass spectrometry (LC/MS). As a result of the DART-TOFMS analyses of 36 products, the protonated molecular ions $[M+H]^+$, corresponding to 6 kinds of major hallucinogenic constituents (mescaline, salvinorin A, *N,N*-dimethyltryptamine, harmine, harmaline and lysergamide), were detected in 21 products. It was possible to estimate their accurate elemental compositions through exact mass measurements. These results were consistent with those of the LC/MS analyses and the contents of the 6 psychotropic constituents were in the range from 0.05 to 45 $\mu\text{g}/\text{mg}$. Typical controlled narcotic drugs, tetrahydrocannabinol, opioid alkaloids and psilocin were also directly detected in marijuana cigarette, opium gum and magic mushroom respectively. Although it is difficult to estimate the matrix effects caused by other plant ingredients, the DART-TOFMS could be useful as a simple and rapid screening method for the targeted psychotropic natural products, because it provides the molecular information of the target compounds without time-consuming extraction and pre-treatment steps.

Key words—Direct Analysis in Real Time (DART); TOFMS; LC/MS; psychotropic plants

緒 言

平成 18 年度の薬事法改正により, 平成 19 年 4 月より指定薬物制度が施行され, 違法ドラッグに対する規制が強化されてきている. しかし一方で, 規制が厳しくなった化学合成化合物の代替品として, 植物系違法ドラッグ製品の流通が深刻な問題となっている. 植物系違法ドラッグ製品に関しては, 製品が同一名称であっても異なった植物が流通していたり, 特定の活性成分を含有する植物が多数存在するために同定が困難であったり, 観賞用として流通している植物の近縁種であったりと, 一律に規制することが困難である. このような製品の流通実態を調査するためには, まずは, 個々の植物成分等についての研究を行うとともに, 実際の製品を分析するた

めの, 効率のよいスクリーニング手法の検討が重要な課題である.

近年開発されたイオン化法 Direct Analysis in Real Time (DART™) の動作原理は, 導入された He ガスをニードル電極の放電によりプラズマ化し, 励起状態の中性気体分子として大気ガス中に放出すると, 測定対象物に直接作用しイオン化することによる. 大気圧下で非接触に試料をイオン化でき, さらに質量検出器に time of flight mass spectrometer (TOFMS) を用いることで, 精密質量測定に基づく元素組成推定が可能となる.¹⁾ また, DART は試料表面からイオン化が可能な他手法 fast atom bombardment (FAB) 法及び matrix-assisted laser desorption ionization (MALDI) 法と比較して, マトリックスが不要であり, イオン源と質量検出器間が開放されているため, 液体, 固体等の試料形態を問わず, 間にかざすだけで試料表面がイオン化される利

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点がある。既に DART を使用した薬品,²⁾ 香料,³⁾ 食品添加物,⁴⁾ 薬物,⁵⁾ 培養植物^{6,7)}等への分析適用例が報告されている。

植物系違法ドラッグ製品は、乾燥植物の葉、樹皮、種子、樹脂、粉末など多様な形態で流通している。これらの製品について、抽出操作等の前処理を省いて直接分析が可能であれば、成分スクリーニングの簡便化、迅速化が期待できる。本研究では様々な形態の植物系違法ドラッグ 36 製品を対象として、DART-TOFMS を用いて、代表的な植物由来幻覚成分である *N,N*-dimethyltryptamine (DMT), mescaline, harmine, harmaline, salvinorin A 及び lysergamide (LSA) の検出を試みた (Fig. 1)。また、同製品について、過去に行った LC/MS 分析の結果⁸⁾と比較し、植物系違法ドラッグ製品のスクリーニング分析における DART-TOFMS の有用性について論じた。さらに、法律で規制されている植物試料として、大麻, psilocin, psilocybin 含有キノコ (いわゆるマジックマッシュルーム) 及びあへんに着目し、これら試料中の主活性成分分析に適用して、本法の有用性を検討した。

実験方法

1. 試料 2004–2007 年に市場で流通していた、向精神様活性を標榜する様々な形態の植物系違法ドラッグ 36 製品 [乾燥植物 (葉・花) 細片 11, 粉末 8, 樹脂状 5, 種子 4, 樹皮 3, タバコ状 2, そ

の他 3 試料] を測定試料とした。これらの 36 製品は、LC/MS 分析により、植物由来活性成分及び故意に添加された合成化合物について、既報⁸⁾により同定済みのものである。また、これら製品中に含まれる代表的な植物由来幻覚成分 DMT, mescaline, harmine, harmaline, salvinorin A, LSA の含有量は、LC/MS 分析による既報⁸⁾のものを参照した。大麻取締法、麻薬及び向精神薬取締法及びあへん法等の法律で規制されている植物試料は、大麻 2 種、マジックマッシュルーム 1 種及びあへん 2 種を使用した。

2. 試薬 DMT, mescaline HCl, psilocin, psilocybin は過去に報告した方法⁹⁾で合成したものを使用した。Salvinorin A は徳島文理大学香川薬学部代田 修博士より、LSA は星薬科大学の細江智夫博士より供された。Harmine 及び harmaline は Aldrich 社 (MO, USA) より、 Δ^9 -tetrahydrocannabinol (THC) は Cerilliant 社 (TX, USA) より購入した。Morphine HCl, codeine H₃PO₄, thebaine, papaverine HCl 及び noscapine HCl は塩野義製薬から購入又は供されたものを使用した。各化合物は過去に HPLC, TLC を用いて純度確認を行い、標品として使用しているものを用いた。それぞれの化合物は 1 mg/ml のメタノール溶液を作成し、定性用の標準化合物溶液とした。その他の試薬は HPLC 用又は試薬特級に準じたものを使用した。また、LC/MS 測定時の抽出溶液の膜ろ過フィルターとして UltraFree-MC (孔径 0.45 μ m) Millipore 社製

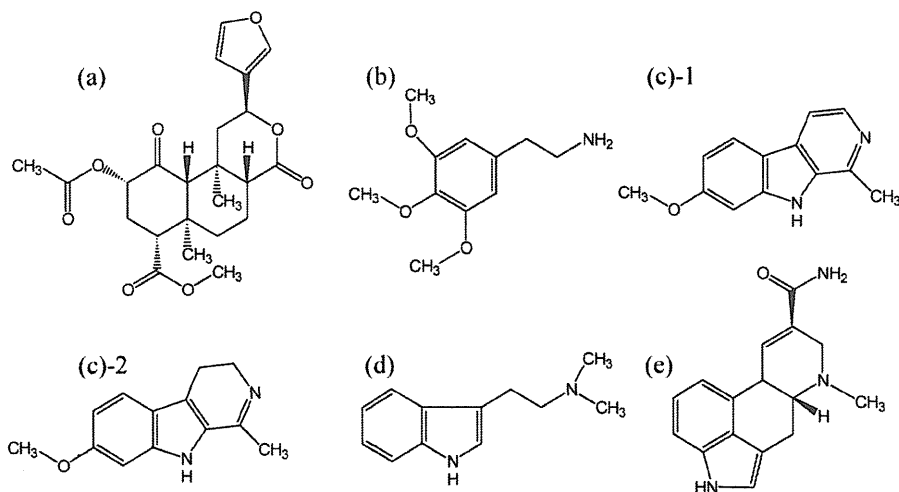


Fig. 1. Chemical Structures of Six Kinds of Major Hallucinogenic Plant Constituents Studied in This Study (a): salvinorin A, (b): mescaline, (c)-1: harmine, (c)-2: harmaline, (d): *N,N*-dimethyltryptamine (DMT), (e): lysergamide (LSA).

(MA, USA) を使用した。

3. DART-TOFMS 測定条件 DART-TOFMS 測定装置として、イオン源 Direct Analysis in Real Time (DART) (日本電子社製) に質量分析計 AccuTOF JMS-T100 (日本電子社製) を連結したものを使用した。測定条件は以下の通りである。なお、質量校正には PEG600 を使用し、各測定の内標準物質として diphenhydramine ($C_{17}H_{21}NO$) 溶液を用いた。

DART 条件

Positive mode; gas flow: He, 2.0 l/min, gas temp.: 200–250°C, needle: 3200 kV, electrode 1: 100 V, electrode 2: 250 V

TOFMS 条件

Positive mode; orifice 1: 15 V, 80°C, orifice 2: 5 V, ring lens: 5 V, ion guide: 500 V, reflectron: 950 V, mass range: 10–1000 (Da)

4. DART-TOFMS 測定方法 測定試料を DART イオン源と TOFMS の間にかざすことで測定した。乾燥した植物試料 (葉, 樹皮, 根) 等は試料片をピンセットでつまみ, そのまま先端をイオン源にかざした。また粉末, 樹脂は薬包紙に包むか, 微量をガラス棒の先端に付着させ測定を行った。種子は乳鉢を用いて粗く砕き, 内容物をガラス棒の先端に微量付着させた。また, 溶液はガラス棒に液体を付着させたのちに先端をかざして測定した。同一試料を 1 回の分析中で数回繰り返し測定し, 得られたスペクトルの精密質量値から化合物の組成推定を行った。試料から分析対象化合物の $[M+H]^+$ に相当するイオンピークが検出された場合には, さらに定性用標準化合物メタノール溶液を使用して分析を行い, 両者の測定値を比較した。

5. LC/MS 測定方法 粉末, 樹脂状の製品は 10 mg, その他の乾燥植物等は乳鉢で粉碎し 20 mg を量り取り, メタノール 2 ml を加えて 10 分間超音波抽出を行った。2000 rpm で 5 分間遠心後, 上清を膜ろ過して LC/MS 測定用試料とした。測定機器に LC/MSD 1100 (Agilent, CA, USA), カラムに Atlantis dC18 (2.1×150 mm, $5 \mu\text{m}$) (Waters, MA, USA) を使用し positive mode で測定を行った。その他の詳細条件は過去の報告に記載した通りである。⁸⁾

結果及び考察

1. 植物系違法ドラッグ製品の分析 DART-TOFMS で試料を測定するに当たり, 2004–2007 年に市場で流通していた様々な形態の植物系違法ドラッグ製品 36 試料を測定試料とした。乾燥した葉, 樹皮, サボテンの皮などからは, 試料の一部をイオン源にかざすと, そのままの状態でも容易に成分ピークが検出できた。種子は表面からでは検出できず, 殻を砕いて中身をガラス棒に微量付着させることで検出可能となった。

DART イオン源を用いた質量分析では, カラムによる分離を行わないので各成分はイオン化されると同時に検出され, スペクトル強度はイオン化の容易さに大きく左右される。¹⁾ また, DART-TOFMS を用いた分析では前処理が不要で, 測定時間も 1 分程度と非常に迅速であった。さらに DART 測定に必要な試料は微量で, 葉 1 枚, 種子 1 粒から目的成分を検出できるため, 同一試料を LC/MS 等で測定可能であり, スクリーニングの信頼性を高めることができる。したがって, 1 次スクリーニングを簡便な DART-TOFMS で行ったのち, LC/MS 等を用いて確認及び定量を行う手法が有効である。

DART-TOFMS の positive mode で植物系違法ドラッグ製品の測定を行った結果, 測定した 36 製品のうち, 分析対象とした植物由来幻覚成分 6 化合物 (DMT, mescaline, harmine, harmaline, salvinorin A, LSA) に相当する $[M+H]^+$ イオンが主なピークとして 21 製品から検出された (Table 1)。本測定結果と過去に報告した LC/MS 分析による成分分析結果⁸⁾を比較すると, 検出化合物は両測定方法で一致した。さらに, 表中にこれら化合物について LC/MS 定量分析の結果を参考値として記載した。DART-TOFMS を用いた定量分析は, TLC を装着した分析についての報告があるものの, その再現性, 定量性は LC/MS に比べて低い。¹⁰⁾ また, DART-TOFMS 分析の検出限界について LC/MS 分析と単純に比較することはできないが, 今回使用した植物試料の対象成分量の範囲ではすべて検出可能であった。

分析対象化合物を検出した代表的な製品について, Fig. 2 に DART-TOFMS スペクトル及び同製品メタノール抽出液の LC/MS スペクトル及びクロマト

Table 1. DART-TOFMS and LC/MS Analyses of the 36 Plant Products Advertized Psychotropic Effects

Compounds	Plant products (Indicated name)	Form	DART-TOFMS			LC/MS ^{b)}	
			Detected ^{a)} (samples)	Accurate mass measurements	Estimated elemental compositions	Detected (samples)	Amount ($\mu\text{g}/\text{mg}$)
Salvinorin A	Salvia	Dried leaves, Cigarette	9	433.1887/ 373.1666	$\text{C}_{23}\text{H}_{29}\text{O}_8/$ $\text{C}_{21}\text{H}_{25}\text{O}_6[\text{M}+\text{H}-60]^+$	9	3.0-23.0
Mescaline	San pedro etc.	Cactus, Powder	3	212.1268	$\text{C}_{11}\text{H}_{18}\text{NO}_3$	3	2.0-17.0
Harmine	Harmala etc.	Bark, Seeds, Powder, Resin	5	213.1037	$\text{C}_{13}\text{H}_{13}\text{N}_2\text{O}$	5	0.8-35.0
Harmaline				215.1276	$\text{C}_{13}\text{H}_{15}\text{N}_2\text{O}$		0.1-45.0
DMT	Ayahuasca etc.	Bark	2	189.1397	$\text{C}_{12}\text{H}_{17}\text{N}_2$	2	11.0, 12.0
LSA	Woodrose etc.	Seeds	2	268.1448	$\text{C}_{16}\text{H}_{18}\text{N}_3\text{O}$	2	0.05, 2.0

a) Detected the ion corresponding to the protonated molecular ion of the targeted compound. b) Data are from ref. 8).

グラムを示した。

「アヤワスカ」を標榜する乾燥植物片 2 製品から、DMT の $[\text{M}+\text{H}]^+$ に相当するイオンをメインピークとして検出し、精密質量の測定値 189.1397 から DMT のプロトン付加体の組成式 $\text{C}_{12}\text{H}_{17}\text{N}_2$ (理論値 189.1392) が推測された [Fig. 2(a)].

Salvinorin A に関しては、DART-TOFMS 測定において $[\text{M}+\text{H}]^+$ に相当するイオンと主ピークとして $[\text{M}+\text{H}-60]^+$ に相当するイオンが 9 製品から検出された。香料が添加されているものや、他の植物と混合している製品 (タバコ状) 等についても分析したが、いずれの試料からも m/z 433 及び 373 の精密質量値が測定可能であり、プロトン付加体の組成式が推定可能であった。LC/MS による定量分析では 9 製品中の salvinorin A の含有量は 3-23 $\mu\text{g}/\text{mg}$ であった。Figure 2(b) に salvinorin A を含有する代表的な製品 (乾燥葉) から得られた DART-TOFMS スペクトル及び LC/MS の測定結果を示した。

LSA の $[\text{M}+\text{H}]^+$ に相当するイオンが検出された 2 製品は外見の異なる種子で woodrose と Rivea corymbosa の表記があり、LC/MS 測定による LSA 含有量はそれぞれ 2 $\mu\text{g}/\text{mg}$, 0.05 $\mu\text{g}/\text{mg}$ と大きく異なった。LSA は分解し易く、LC/MS 測定においては、迅速な抽出操作、測定が求められるが、DART-TOFMS を用いた分析は前処理が不要であり、含有量 0.05 $\mu\text{g}/\text{mg}$ の試料からも検出可能であった [Fig. 2(c)]. また、LSA 含有植物には iso LSA が存在することが報告されている。¹¹⁾ LC/MS 分析では分離可能であっても、DART-TOFMS では異性を区別できないため、同一ピークとして検

出されたと考えられた。

Mescaline では、DART-TOFMS 測定において 3 製品から $[\text{M}+\text{H}]^+$ に相当するピークが検出された。また、精密質量の測定値 212.1285 から組成式 $\text{C}_{11}\text{H}_{18}\text{NO}_3$ (理論値 212.1287) が推測された [Fig. 2(d)]. LC/MS 分析においては乾燥したサボテン試料を粉砕する必要があるが、DART-TOFMS による分析では試料片から容易に測定が可能であった。

Harmine, harmaline は含有が確認された 5 製品のすべてで同時に検出され、LC/MS 測定における含有量に相関なく harmine 由来の $[\text{M}+\text{H}]^+$ ピークに相当する m/z 213 の強度が、harmaline 由来の $[\text{M}+\text{H}]^+$ に相当する m/z 215 より強く観察された [Fig. 2(e)]. Harmine, harmaline 含有植物製品は、種子や樹皮等の植物体及び樹脂や粉末状の製品が存在したが、加工された製品より植物体において、両化合物のピーク強度が高い傾向にあった。

以上の結果から、DART-TOFMS を用いた分析は、今回分析対象とした植物由来幻覚成分を製品中から検出するのに優れた方法と考えられた。特に、Salvinorin A 含有植物 (*Salvia divinorum*) は、Salvinorin A とともに、平成 19 年 4 月より指定薬物として規制されているが、呈色反応やイムノアッセイなどの簡易スクリーニング手法は現在のところ報告されていない。また、Salvinorin A は、植物製品のメタノール抽出物を用いた GC/MS や HPLC 等の分析において、configuration の異なる化合物や分解物が一部検出されることが報告されており、¹²⁾ 様々な形態の製品から前処理を行わず迅速に測定することが可能な DART-TOFMS は、有用な分析手法であると考えられた。しかし、高濃度に添加される

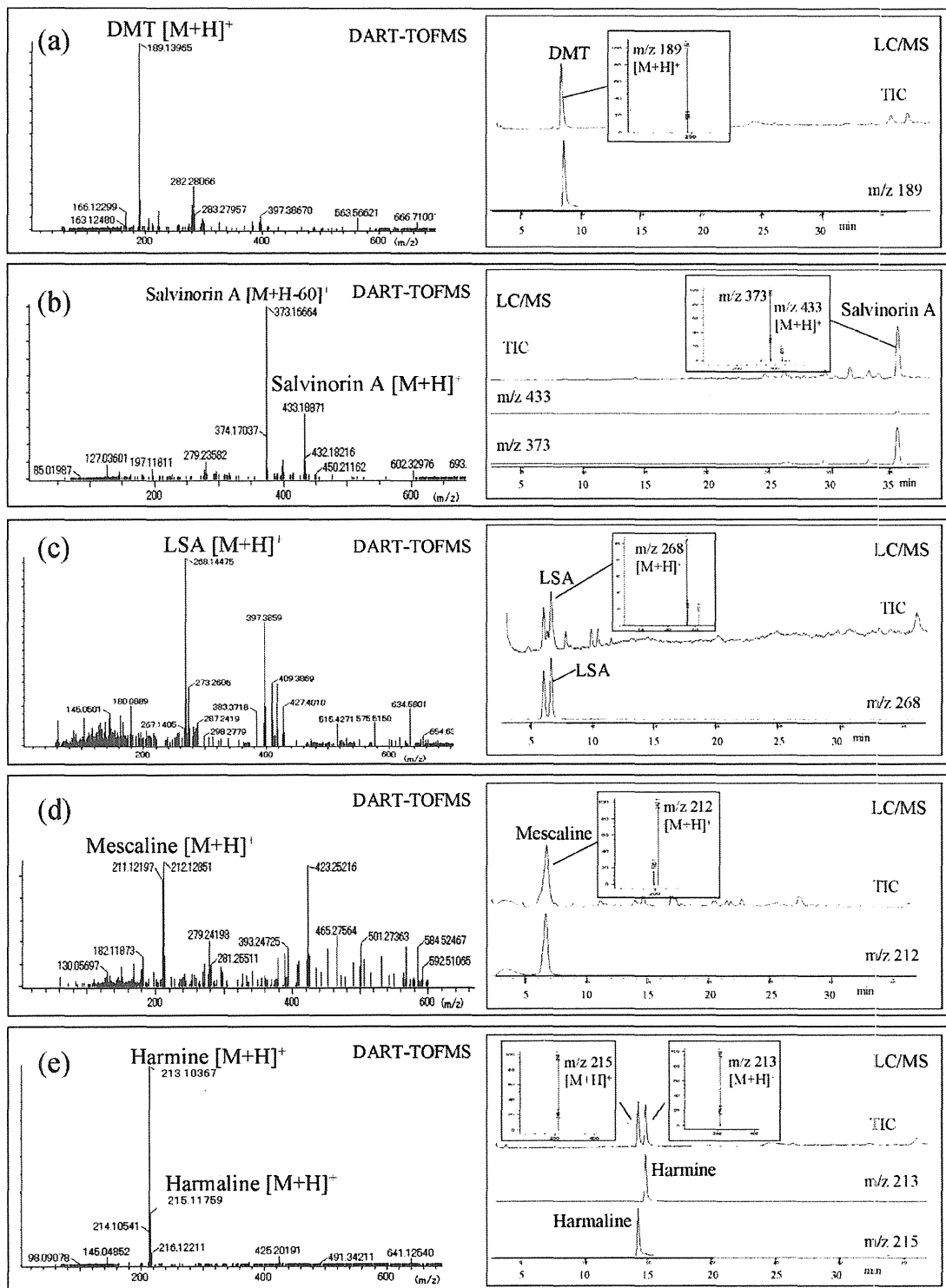


Fig. 2. DART-TOFMS Mass Spectra and LC/MS Ion Chromatograms of the Products

(a): “Ayawaska” (bark sample contained DMT), (b): “Salvia” (dried leaf sample contained salvinorin A), (c): “Rivea corymbosa” (seeds sample contained LSA), (d): “San Pedro” (dried cactus sample contained mescaline), (e): “Harmala” (seeds sample contained harmine and harmaline).

可能性のある合成化合物や常在成分によるイオン化の妨害も否定できないため、強度の大きい他ピークが検出された場合には注意が必要である。また、構

造類似体や異性体の多い合成違法ドラッグ成分においては、組成式が同一な化合物が多く DART-TOFMS のみでは判別できないことに留意しなけれ

ばならない。

2. 法律で規制されている植物試料の分析 法律で規制されている植物試料として、大麻 (*Cannabis sativa* L.), psilocin, psilocybin 含有キノコ (いわゆるマジックマッシュルーム) 及び、アヘン試料の DART-TOFMS による測定を行い Fig. 3 に結果を示した。

大麻試料では、乾燥葉及びタバコ状の試料から、代表的な大麻成分である THC, cannabidiol (CBD) 及び cannabinol (CBN) の $[M+H]^+$ に相当するイオンを主なピークとして検出することが可能であった。また、精密質量の測定値 (315.2330 及び 311.2018) から、THC/CBD 及び CBN のプロトン付加体の組成式 $C_{21}H_{31}O_2$ 及び $C_{21}H_{27}O_2$ (理論値 315.2324 及び 311.2011) が推定可能であった [Fig. 3 (a)].

乾燥キノコ試料は試料片を直接イオン源にかざしたところ、psilocin の $[M+H]^+$ に相当するイオンを主ピークとして検出することが可能であった。また、精密質量の測定値 205.1324 から psilocin のプロトン付加体の組成式 $C_{12}H_{17}N_2O$ (理論値 205.1341) が推定可能であった [Fig. 3 (b)]. しかし、主活性

成分の1つである psilocybin に相当する質量のイオンピークは検出されなかった。Psilocybin は熱等により容易に脱リン酸化され psilocin に分解されるため、GC/MS 分析では psilocin として検出されることが報告されている。¹³⁾ DART-TOFMS による測定でも同様に、標準品の psilocybin 溶液を測定したところ、イオン化時の熱により psilocin として検出されることを確認しており、そのために今回乾燥キノコ試料から psilocybin が検出されなかったと考えられる。

アヘン試料については樹脂状と粉末状の試料を、葉包紙に少量取りイオン源にかざして測定したところ、主要なアヘンアルカロイドの $[M+H]^+$ に相当するイオンが検出された。Morphine ($C_{17}H_{19}NO_3$, 理論値 286.1443, 測定値 286.1431), codeine ($C_{18}H_{21}NO_3$, 理論値 300.1559, 測定値 300.1579), thebaine ($C_{19}H_{21}NO_3$, 理論値 312.1599, 測定値 312.1595), papaverine ($C_{20}H_{21}NO_4$, 理論値 340.1549, 測定値 340.1535) 及び noscapine ($C_{22}H_{23}NO_7$, 理論値 414.1553, 測定値 414.1542) が、測定値よりプロトン付加体の組成式が推定可能であった 5 化合物であ

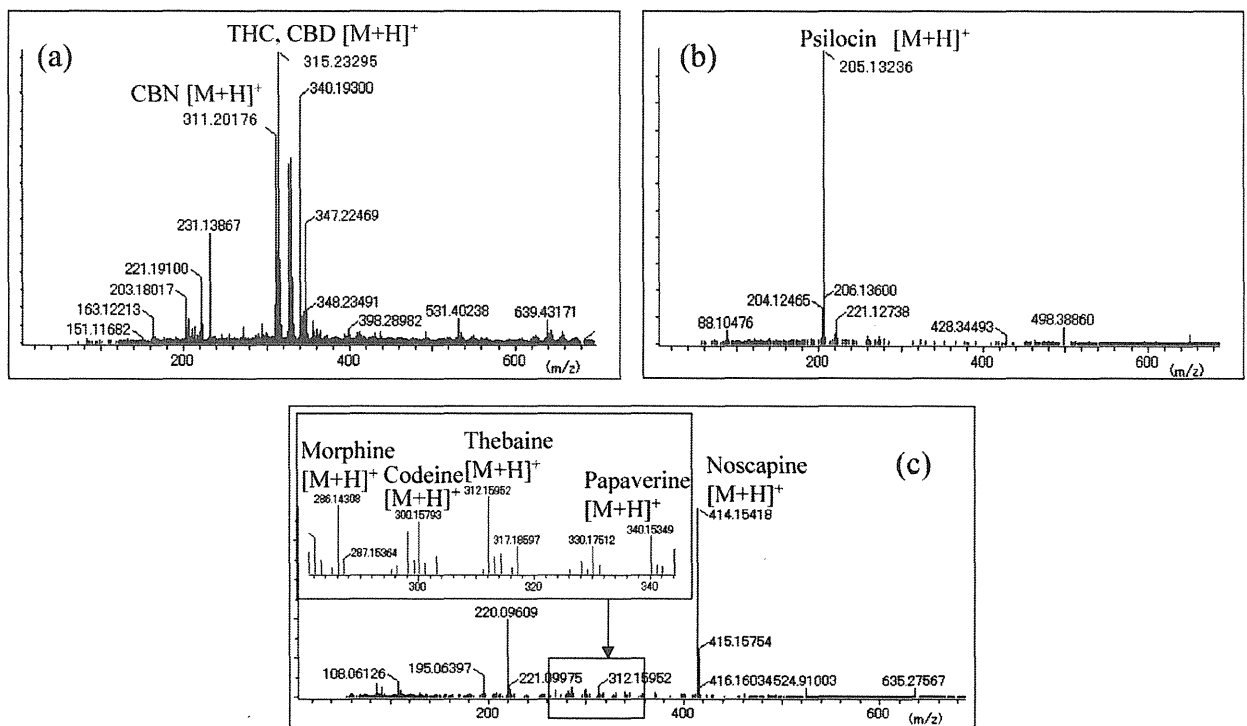


Fig. 3. DART-TOFMS Mass Spectra

(a): marijuana cigarette sample contained CBN, THC and CBD, (b): dried mushroom sample contained psilocin, (c): opium resin contained morphine, codeine, thebaine, papaverin and noscapin.

る [Fig. 3(c)]. 樹脂試料からは noscapine が、粉末試料からは papaverine が主ピークとして検出されたが、アルカロイド中最も含量が高い morphine のイオン強度は低かった。そこで、同濃度に調整したあへんアルカロイド 5 化合物のメタノール溶液を、ガラス棒に付着させて DART-TOFMS で測定したところ、morphine のイオン強度が最も低く、他化合物が検出された 10 $\mu\text{g}/\text{ml}$ において検出ができなかった。したがって、あへん試料を DART で確認する際は、morphine より noscapine 等の成分を指標とするのが望ましいと考えられた。

結 論

DART-TOFMS を用いて、乾燥植物、樹脂、粉末等の形態を有する植物系違法ドラッグ製品及び法規制植物について、抽出操作等の前処理を行わずに測定を行った。その結果、今回測定を行った植物に含有される代表的な幻覚成分や活性成分に相当するピークが検出でき、さらに精密質量値より化合物の組成推定を行うことで、含有成分の推定が可能であった。LC/MS や GC/MS 等他の分析手法と比較すると、DART-TOFMS では前処理が不要で測定時間が短く簡便であり、キャリアオーバーが起き難い等の利点がある。したがって、一次スクリーニングを DART-TOFMS で行ったのち、LC/MS や GC/MS 等の分離分析手段を用いて成分の同定、定量を行うという一連の手法が植物違法ドラッグの分析法として有用であると考えられる。

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REFERENCES

- 1) Cody R. B., Laramée J. A., Durst H. D., *Anal. Chem.*, **77**, 2297–2302 (2005).
- 2) Petucci C., Diffendal J., Kaufman D., Mekonnen B., Terefenko G., Musselman B., *Anal. Chem.*, **79**, 5064–5070 (2007).
- 3) Haefliger O. P., Jeckelmann N., *Rapid Commun. Mass Spectrom.*, **21**, 1361–1366 (2007).
- 4) Vail T., Jones P. R., Sparkman O. D., *J. Anal. Toxicol.*, **31**, 304–312 (2007).
- 5) Bennett M. J., Steiner R. R., *J. Forensic Sci.*, **54**, 370–375 (2009).
- 6) Madhusudanan K. P., Banerjee S., Khanuja S. P., Chattopadhyay S. K., *Biomed. Chromatogr.*, **22**, 596–600 (2008).
- 7) Banerjee S., Madhusudanan K. P., Khanuja S. P., Chattopadhyay S. K., *Biomed. Chromatogr.*, **22**, 250–253 (2008).
- 8) Kawamura M., Kikura-Hanajiri R., Goda Y., *Jpn. J. Food Chem.*, **15**, 73–78 (2008).
- 9) Ono M., Shimamine M., Takahashi K., *Eisei Shikenjo Hokoku*, **91**, 33–41 (1973).
- 10) Morlock G., Ueda Y., *J. Chromatogr. A.*, **1143**, 243–251 (2007).
- 11) Chao J. M., Der Marderosian A. H., *J. Pharm. Sci.*, **62**, 588–591 (1973).
- 12) Kikura-Hanajiri R., Kawamura M., Uchiyama N., Ogata J., Kamakura H., Saisho K., Goda Y., *Yakugaku Zasshi*, **128**, 971–979 (2008).
- 13) Kikura-Hanajiri R., Hayashi M., Saisho K., Goda Y., *J. Chromatogr. B.*, **825**, 29–37 (2005).