

A CHEMOTAXONOMIC ANALYSIS OF CANNABINOID VARIATION IN *CANNABIS* (CANNABACEAE)¹

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Cannabinoids are important chemotaxonomic markers unique to *Cannabis*. Previous studies show that a plant's dry-weight ratio of Δ^9 -tetrahydrocannabinol (THC) to cannabidiol (CBD) can be assigned to one of three chemotypes and that alleles B_D and B_T encode alloenzymes that catalyze the conversion of cannabigerol to CBD and THC, respectively. In the present study, the frequencies of B_D and B_T in sample populations of 157 *Cannabis* accessions were determined from CBD and THC banding patterns, visualized by starch gel electrophoresis. Gas chromatography was used to quantify cannabinoid levels in 96 of the same accessions. The data were interpreted with respect to previous analyses of genetic and morphological variation in the same germplasm collection. Two biotypes (infraspecific taxa of unassigned rank) of *C. sativa* and four biotypes of *C. indica* were recognized. Mean THC levels and the frequency of B_T were significantly higher in *C. indica* than *C. sativa*. The proportion of high THC/CBD chemotype plants in most accessions assigned to *C. sativa* was <25% and in most accessions assigned to *C. indica* was >25%. Plants with relatively high levels of tetrahydrocannabinol (THCV) and/or cannabidivarin (CBDV) were common only in *C. indica*. This study supports a two-species concept of *Cannabis*.

Key words: cannabinoid; *Cannabis*; chemotaxonomy; evolution; genetics; taxonomy; tetrahydrocannabinol.

Cannabis (Cannabaceae) has been a source of fiber, food, oil, medicine, and inebriant since prehistoric times (Chopra, 1969; Schultes, 1973; Abel, 1980). Whether the genus consists of one or more species is a matter of divided opinion (Schultes et al., 1974; Small and Cronquist, 1976; Emboden, 1981). *Cannabis* strains cultivated for fiber and/or seed production (here referred to as hemp) are commonly differentiated from strains cultivated for medicinal or recreational use, but the evolutionary relationships between these two groups and between cultivated and wild or naturalized (feral) populations are not well understood. To further resolve these issues, a systematic investigation of genetic, morphological, and biochemical variation was conducted on a diverse collection of 157 *Cannabis* accessions grown in a common environment (Hillig, 2004, in press). Chemotaxonomic aspects of that investigation are reported herein. Qualitative and quantitative analyses of cannabinoid variation and a method of characterizing cannabinoid differences among populations based on a simple genetic model provide new evidence regarding the evolution and domestication of this socioeconomically important genus.

Cannabinoids are terpenophenolic compounds unique to *Cannabis*. They are produced by glandular trichomes that occur on most aerial surfaces of the plant (Dayanandan and Kaufman, 1976; Turner et al., 1978). Approximately 61 cannabinoids are known to exist, although some of these are breakdown products or artifacts (Schultes and Hofmann, 1980; Turner et al., 1980). The cannabinoids discussed in this paper are biosynthesized in an acidic (carboxylated) form and are decarboxylated upon heating and drying of harvested plant material (Doorenbos et al., 1971). They are here referred to in their decarboxylated form. Cannabigerol (CBG) is the direct precursor of cannabichromene (CBC), cannabidiol (CBD), and

Δ^9 -tetrahydrocannabinol (THC) (Taura et al., 1995, 1996; Morimoto et al., 1997). A homologous series of compounds with propyl side-chains is biosynthesized from cannabigerovarin (CBGV), including cannabivarichromene (CBCV), cannabidivarin (CBDV), and Δ^9 -tetrahydrocannabivarin (THCV), respectively (Fig. 1A–D) (Schultes and Hofmann, 1980). THC and/or CBD are generally produced in greatest abundance. However, THCV and less commonly CBDV may exceed the levels of THC and/or CBD in some plants (Baker et al., 1980). THC and THCV are primarily responsible for the euphoric effects of marijuana and hashish (McPartland and Russo, 2001).

Chemotaxonomy has a long history of use in the delimitation of *Cannabis* taxa. Lamarck (1785) emphasized the greater inebriant potential of *C. indica* Lam. when he differentiated it from *C. sativa* L. Names and descriptions of other putative species of *Cannabis* have been published (reviewed in Schultes et al., 1974; Small and Cronquist, 1976). Of these, only *C. ruderalis* Janisch. is commonly accepted. Small (1979a) considered the amount of THC produced by *Cannabis* to be an “extremely important” taxonomic character and used gas chromatography (GC) to differentiate *indica* strains from *sativa* strains on the basis of their THC content (Small and Beckstead, 1973a, b; Small et al., 1975; Small and Cronquist, 1976). Small and Cronquist (1976) favored a monospecific concept and assigned these two taxa to subspecies of *C. sativa*.

Numerous biochemical studies of *Cannabis* plants grown from achenes (“seeds”) of known geographic origin have been reported (Fetterman et al., 1971; Fetterman and Turner, 1972; Nordal and Braenden, 1973; Small and Beckstead, 1973a, b; Turner et al., 1973; Turner and Hadley, 1973a, 1974; Boucher et al., 1974; Holley et al., 1975; Small et al., 1975; Rowan and Fairbairn, 1977; Beutler and Der Marderosian, 1978; Clark and Bohm, 1979; Turner et al., 1979; Fournier and Paris, 1980; Hemphill et al., 1980; Veszki et al., 1980; de Meijer et al., 1992). Forensic studies of *Cannabis* examined marijuana and hashish samples of known origin, grown in a range of environments (Jenkins and Patterson, 1973; Poddar et al., 1973; Baker et al., 1980, 1982; Barni-Comparini et al., 1984;

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TABLE 1. Taxonomic circumscription of the *Cannabis* germplasm collection based on a previous analysis of allozyme allele frequencies. The assignment of accessions to putative taxa was previously published (Hillig, 2004).

Putative Taxon	Description
<i>C. indica</i> hemp biotype ^a	Hemp landraces from southern and eastern Asia
<i>C. indica</i> feral biotype ^b	Feral populations from India and Nepal
<i>C. indica</i> NLD biotype ^c	Narrow-leaflet drug (NLD) strains from the Indian subcontinent and other drug producing regions
<i>C. indica</i> WLD biotype ^d	Wide-leaflet drug (WLD) strains from Afghanistan and Pakistan
<i>C. sativa</i> hemp biotype ^e	Hemp landraces from Europe, Asia Minor and central Asia
<i>C. sativa</i> feral biotype ^f	Feral populations from eastern Europe
<i>C. ruderalis</i> ^g	Ruderal populations from central Asia

^a Includes accessions previously assigned to *C. chinensis* Delile.

^b Includes accessions previously assigned to *C. sativa* L. subsp. *indica* (Lam.) Small & Cronq. var. *kafiristanica* (Vav.) Small & Cronq.

^c Includes accessions previously assigned to *C. sativa* subsp. *indica* var. *indica* (Lam.) Wehmer, excluding accessions assigned to *C. indica* sensu Schultes et al. (1974) and Anderson (1980).

^d Includes accessions previously assigned to *C. indica* Lam. sensu Schultes et al. (1974) and Anderson (1980).

^e Includes accessions previously assigned to *C. sativa* subsp. *sativa* var. *sativa* sensu Small and Cronquist (1976), excluding accessions assigned to *C. chinensis*.

^f Includes accessions previously assigned to *C. sativa* subsp. *sativa* var. *spontanea* (Vav.) Small & Cronq., excluding accessions assigned to *C. ruderalis* Janisch.

^g Includes accessions previously assigned to *C. ruderalis*.

netic data. Based on these results, a working hypothesis for a taxonomic circumscription of the *Cannabis* germplasm collection is given in Table 1. This hypothesis represents a synthesis of polytypic treatments of *Cannabis* by Lamarck (1785), Delile (1849), Janischewsky (1924), Vavilov and Bukinich (1929), Schultes et al. (1974), and Anderson (1980). Recognition of *C. sativa* and *C. indica* as separate species is primarily based on allozyme allele frequencies, morphological differences, different geographic ranges, and the fact that putative wild populations of both species have been found within the indigenous range of *Cannabis*, presumed to be in central Asia, the northwest Himalayas, and western China (de Candolle, 1885; Vavilov, 1926; Vavilov and Bukinich, 1929; Zuckovskij, 1962; Hillig, 2004, in press). Putative infraspecific taxa of unassigned rank are referred to as "biotypes" pending a taxonomic revision of the genus, in progress. Chemotaxonomic support for a two-species hypothesis is provided by an analysis of flavonoid variation that detected luteolin C-glycuronide in 30 of 31 plants assignable to *C. sativa*, but not in 21 of 22 plants assignable to *C. indica* (Clark and Bohm, 1979).

Allozyme analysis revealed that the hemp accessions in the germplasm collection under study derive from both the *C. sativa* and *C. indica* gene pools (Hillig, 2004). Hemp landraces from Europe, Asia Minor, and central Asia are assigned to the hemp biotype of *C. sativa* and hemp landraces from southern and eastern Asia are assigned to the hemp biotype of *C. indica*. Accessions of wild or naturalized populations from eastern Europe and the northwest Himalayas are assigned to the feral biotypes of *C. sativa* and *C. indica*, respectively. Ruderal accessions from central Asia are tentatively assigned to *C. ruderalis*, although few morphological differences were found between these accessions and those assigned to the feral biotype of *C. sativa* (Hillig, in press). Plants of accessions cultivated for drug production were characterized as having either narrow lanceolate or linear-lanceolate leaflets or wide oblanceolate leaflets. Both biotypes derive from the *C. indica* gene pool and are morphologically distinct (Anderson, 1980; Hillig, 2004, in press).

The objective of the present study was to characterize qualitative and quantitative patterns of cannabinoid variation and differences in B_D and B_T allele frequencies among putative

taxa, in the same *Cannabis* collection that Hillig (2004, in press) examined in previous genetic and morphological studies. Comparison of genetic, morphological, and biochemical patterns of variation within a single set of accessions was anticipated to advance our understanding of taxonomic and evolutionary relationships within the genus.

MATERIALS AND METHODS

Germplasm collection—A diverse collection of 157 *Cannabis* accessions of known geographic origin was obtained from breeders, researchers, gene banks, and law enforcement agencies. Each accession consisted of a small packet of viable achenes. Passport data, accession codes, and the assignment of accessions to putative taxa are published elsewhere (Hillig, 2004). Industrial hemp strains selected for low THC production were excluded from this investigation. Voucher specimens are deposited in the Indiana University herbarium (IND). The a priori assignment of accessions to species is based on a multivariate analysis of allozyme allele frequencies and on geographic origins, without regard to cannabinoid content (Hillig, 2004). Assignment of accessions to infraspecific taxa (biotypes) is primarily based on geographic origins, morphological traits, and presumed purpose of cultivation.

Sample preparation for GC analysis—*Cannabis* plants were grown in two secure greenhouses at Indiana University, Bloomington, Indiana, USA. The plants were individually grown in 13-cm clay pots in a soil mixture of three parts (by volume) black peat, two parts vermiculite, and one part each of sand and top soil. Each plant received 100 mL of nutrient solution at weekly intervals during the vegetative period of growth, consisting of 25 mL of Dyna-Gro 7-9-5 (Dyna-Gro Corporation, San Pablo, California, USA) and 18 mL of 1 M potassium hydroxide (to neutralize the acidity) per 3.8 L of water. Pistillate plants were isolated from staminate plants before anther dehiscence. The inflorescences of pistillate plants were sampled when resin production of each plant was visually assessed to have reached its peak. The samples were air dried at room temperature and oven dried overnight at 30°C prior to extraction. The primary and larger secondary leaves were removed, and only visibly resinous floral bracts and small subtending leaves were analyzed.

Sample extraction—Sample material (50 mg) was placed in a test tube with 1 mL of chloroform. The plant material was crushed with a glass rod and briefly sonicated to dislodge and/or rupture the resin heads of the glandular trichomes. The sample remained in the solvent at room temperature for at least one hour, and was sonicated again for a few seconds. A 20- μ L volume of extract was transferred to a small test tube, and the solvent was evaporated

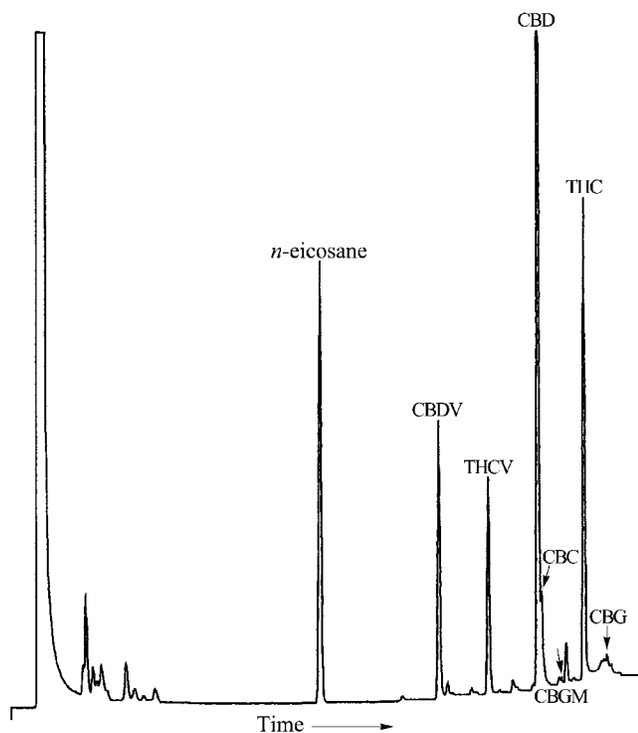


Fig. 2. Gas chromatogram of a chemotype II plant of an Indian accession assigned to the feral biotype of *Cannabis indica*, with relatively high levels of cannabidiol (CBD), Δ^9 -tetrahydrocannabinol (THC), cannabidiol (CBDV), and Δ^9 -tetrahydrocannabinol (THCV). Also labeled are peaks for *n*-eicosane (the internal standard), cannabichromene (CBC), cannabigerol monomethylether (CBGM), and cannabigerol (CBG).

with a gentle stream of nitrogen. The residue was redissolved in 50 μ L of acetone containing 0.25 mg/mL of *n*-eicosane, the internal standard.

GC conditions—Chromatograms were generated with a Hewlett-Packard 5710A gas-liquid chromatograph fitted with a 30 m \times 0.53 mm DB-5ms column (J & W Scientific, Rancho Cordova, California, USA) having a film thickness of 1.5 μ m. Injector and detector port temperatures were 250°C and 300°C, respectively. Carrier and make-up gas flow rates were 5.8 mL/min (He) and 24.2 mL/min (N₂), respectively. The temperature was held at 200°C for 8 min, then increased to 300°C at 4°C/min and held for 4 min. The integrator was calibrated with single concentrations of CBC, CBD, CBG and THC at 0.5, 0.4, 0.5, and 0.4 mg/mL, respectively.

Accessions analyzed by GC—Ninety-six of the 157 *Cannabis* accessions were analyzed by GC. The countries of origin and number of accessions from each country were: Afghanistan-10, Bulgaria-1, China-6, Colombia-2, Gambia-1, Germany-1, Hungary-8, India-5, Italy-4, Jamaica-1, Japan-2, Lesotho-1, Mexico-4, Nepal-2, Nigeria-1, Romania-2, Russia-13, South Africa-3, South Korea-7, Spain-3, Swaziland-1, Thailand-6, Turkey-5, Uganda-2, Ukraine-3, and former Yugoslavia-2. One to seven plants of each accession were analyzed.

Starch gel electrophoresis—A fortuitous result of starch gel electrophoresis for a previous survey of allozyme variation was that THC and CBD migrated into the starch gels and were stained by the dye Fast Black K used to visualize the allozyme banding patterns of leucine aminopeptidase (LAP). Materials and methods for starch gel electrophoresis and a photograph showing cannabinoid banding patterns are published elsewhere (Hillig, 2004). About 10 plants of each of the 157 accessions were assayed, although fewer plants were assayed for accessions obtained late in the investigation. By this method, a

TABLE 2. Retention times (RT) and peak identities for the gas chromatogram in Fig. 2 of a *Cannabis indica* extract.

RT (min)	Peak Identity
13.707	<i>n</i> -Eicosane ^a
19.017	Cannabidiol (CBDV)
21.225	Δ^9 -Tetrahydrocannabinol (THCV)
23.436	Cannabidiol (CBD)
23.594	Cannabichromene (CBC)
[24.543] ^b	Cannabigerol monomethylether (CBGM) ^c
25.457	Δ^9 -Tetrahydrocannabinol (THC)
26.500	Cannabigerol (CBG)

^a Internal standard.

^b Not detected on the chromatogram in Fig. 2. The RT was interpolated from other chromatograms.

^c Presumed to be the peak for CBGM.

larger number of plants and accessions were qualitatively characterized than by GC analysis alone.

Statistical analysis—Statistical analyses were performed with JMP version 5.0 (SAS Institute, 2002). THC vs. CBD levels of individual plants were plotted and separate regression lines were calculated for the three chemotypes. Means and standard deviations of the THC/CBD ratios and the quantitative levels of CBD and THC were determined, as well as the linear correlations between THC and CBD, for each of the three chemotypes. Means and standard deviations of the quantitative levels of CBC, CBD, CBG, THC, and CBD + THC were determined for each of the seven putative taxa in Table 1, as well the sum of the areas under the chromatogram peaks (relative to the internal standard) corresponding to CBDV and THCv, and for the peak corresponding to CBGM. Student's *t* test ($P \leq 0.05$) was used to determine which means were significantly different for pairwise comparisons between taxa. Means and standard deviations of the *B₇* allele frequency were determined for each putative taxon. The estimated proportions of chemotype I individuals in the accessions and putative taxa were determined by squaring the corresponding *B₇* allele frequencies.

RESULTS

A sample chromatogram of a chemotype II plant of an Indian accession assigned to the feral biotype of *C. indica* is shown in Fig. 2. This plant had relatively high levels of CBD, CBDV, THC, and THCv. Table 2 shows the GC retention times of the labeled peaks in Fig. 2.

Qualitative differences—A histogram of the THC/CBD ratios (\log_{10}) of individual plants shows three discrete groups (Fig. 3). Chemotype I plants had THC/CBD ratios greater than 50, chemotype II plants had THC/CBD ratios between 0.25 and 1.50, and chemotype III plants had THC/CBD ratios less than 0.20 (Table 3). THC was detected in all 253 plants analyzed, but CBD was not detected in some chemotype I plants. Figure 4 shows a scatter plot of THC vs. CBD levels of individual plants, with separate regression lines (forced through the origin) for the three chemotypes. A highly significant ($P \leq 0.0001$) linear correlation existed between THC and CBD content for all three chemotypes, although the correlation for chemotype III plants ($r = 0.88$) was substantially greater than for chemotypes I and II ($r = 0.52$ in both cases).

Quantitative differences—Mean THC levels were significantly higher in the narrow-leaflet drug (NLD) and wide-leaflet drug (WLD) biotypes of *C. indica* than in all other taxa (Table 4). The WLD biotype had a significantly higher mean level of CBD + THC than the NLD biotype, while the latter

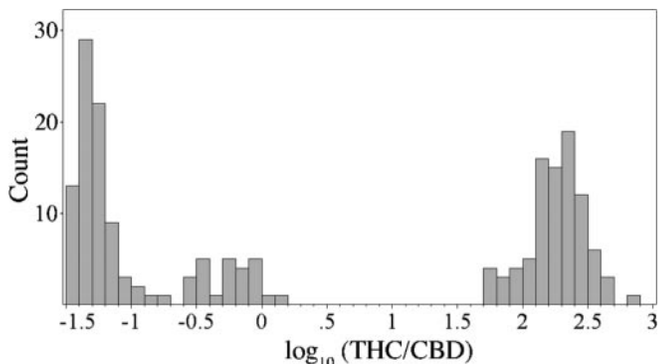


Fig. 3. Histogram of \log_{10} values of the dry-weight ratios of Δ^9 -tetrahydrocannabinol/cannabidiol (THC/CBD) for 194 *Cannabis* plants. Plants in which CBD was not detected were excluded. Plants with \log_{10} values >1.0 were assigned to chemotype I, plants with \log_{10} values <-0.7 were assigned to chemotype III, and plants with intermediate values were assigned to chemotype II.

had a significantly lower mean level of CBD. The hemp and feral biotypes of *C. indica* averaged higher levels of THC than CBD. Accessions assigned to *C. ruderalis* and the feral biotype of *C. sativa* had the lowest mean levels of CBD + THC, but not significantly lower than the hemp biotype of *C. sativa* and the hemp and feral biotypes of *C. indica*.

Minor cannabinoids—The chromatographic peaks for CBC and CBD sometimes merged because of their close retention times. A highly significant correlation ($r = 0.79, P \leq 0.0001$) existed between CBC and CBD content for the pooled values of the chemotype II and III plants in which both cannabinoids were detected, with about 6% as much CBC produced as CBD. The correlation between CBC and CBD content for chemotype I plants was much weaker but still significant ($r = 0.25, P = 0.02$), whereas the correlation between CBC and THC content in this group was not significant ($r = 0.19, P > 0.05$). The hemp biotype of *C. indica* had a significantly higher mean level of CBC than the other taxa in Table 4, except the *C. indica* feral biotype. The mean level of CBG was significantly higher in the NLD biotype of *C. indica* than in the hemp and feral biotypes of *C. sativa*.

CBDV, THCV, and CBGM were not quantified because calibration standards were not available. The apparent levels (i.e., areas under the peaks relative to the internal standard [i.s.]) of these cannabinoids were evaluated and compared (Table 4).

TABLE 3. Arithmetic means, standard deviations, and ranges of the THC/CBD ratios and of the dry-weight percentages of cannabidiol (CBD) and Δ^9 -tetrahydrocannabinol (THC) for chemotype I, II, and III plants. Mature unpollinated pistillate inflorescences from 96 *Cannabis* accessions were sampled. Plants in which CBD was not detected were excluded. Linear regression lines (forced through the origin) and Pearson product-moment correlations (r) of CBD vs. THC are shown for the three chemotypes. N = number of plants analyzed.

	Chemotype I $N = 88$	Chemotype II $N = 25$	Chemotype III $N = 80$
THC/CBD (SD)	205 (106)	0.63 (0.28)	0.05 (0.02)
Range	52–727	0.27–1.32	0.03–0.17
CBD% (SD)	0.04 (0.03)	3.8 (1.7)	4.6 (2.5)
Range%	0.01–0.16	0.9–7.3	1.0–13.6
THC% (SD)	6.3 (3.1)	2.3 (1.2)	0.24 (0.14)
Range%	0.6–14.7	0.6–5.3	0.05–0.69
Regression line	THC% = 137 CBD%	THC% = 0.56 CBD%	THC% = 0.05 CBD%
Correlation (r) ^a	0.52***	0.52***	0.88***

^a *** = Highly significant ($P \leq 0.0001$).

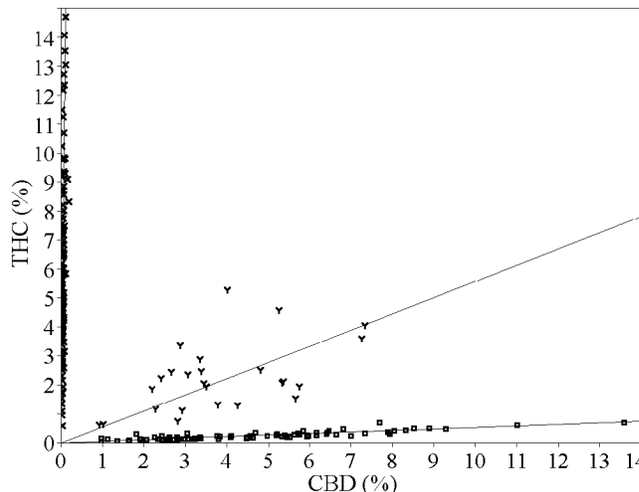


Fig. 4. Plot of Δ^9 -tetrahydrocannabinol (THC) % vs. cannabidiol (CBD) % for 253 *Cannabis* plants. Chemotype I, II, and III plants are marked with an X, Y, and square, respectively. Linear regression lines (forced through the origin) are drawn for each chemotype.

CBDV and/or THCV were detected in all plants that were analyzed. However, all 36 plants with peak areas of (CBDV + THCV)/i.s. >0.30 were of *C. indica* accessions from countries in Asia (Afghanistan, China, India, Nepal, Thailand) or Africa (Gambia, Lesotho, Nigeria, South Africa, Swaziland). The mean peak area of (CBDV + THCV)/i.s. was significantly higher for the feral biotype of *C. indica* than for all other taxa.

Twenty-five of the 29 plants with peak areas of CBGM/i.s. >0.05 were of accessions from Afghanistan, China, Japan, or South Korea. Plants of two accessions from Russia and two accessions from Hungary also had enhanced levels of CBGM.

Genetic analysis—CBD and THC appeared as distinct bands toward the bottom of starch gels stained with Fast Black K, with CBD migrating ahead of THC. CBD stained dark blue, and THC stained dark violet. A diffuse blue band that ran ahead of CBD and THC was visible for plants with high apparent levels of THCV. It is likely that these cannabinoids migrated into the gels in their carboxylated forms, which are water-soluble. The cannabinoid banding patterns were interpreted in accord with the hypothesis that a single gene with two codominant alleles controls the conversion of CBG to THC and CBD. Plants with a strong CBD band were assigned

TABLE 4. Arithmetic means^a, standard deviations, and ranges of the dry-weight percentages of cannabichromene (CBC), cannabidiol (CBD), cannabigerol (CBG), Δ⁹-tetrahydrocannabinol (THC), and (CBD + THC) for 253 *Cannabis* plants assigned to seven putative taxa. Statistics are also given for the peak areas (determined by gas chromatography) relative to the internal standard (i.s.) of cannabidivarin plus Δ⁹-tetrahydrocannabivarin (CBDV + THCV), and of cannabigerol monomethylether (CBGM). *N* = number of plants analyzed.

Compound	<i>C. indica</i> Hemp Biotype <i>N</i> = 45	<i>C. indica</i> Feral Biotype <i>N</i> = 14	<i>C. indica</i> NLD Biotype <i>N</i> = 68	<i>C. indica</i> WLD Biotype <i>N</i> = 40	<i>C. sativa</i> Hemp Biotype <i>N</i> = 62	<i>C. sativa</i> Feral Biotype <i>N</i> = 16	<i>C. ruderalis</i> <i>N</i> = 7
CBC%	0.34 A	0.18 AB	0.19 B	0.17 B	0.18 B	0.13 B	0.07 B
(SD)	(0.47)	(0.27)	(0.21)	(0.25)	(0.24)	(0.20)	(0.10)
Range%	0.0–1.9	0.0–0.9	0.0–0.9	0.0–1.4	0.0–1.2	0.0–0.8	0.0–0.2
CBD%	1.43 BC	1.95 BC	0.02 D	1.21 C	4.01 A	3.62 A	3.02 AB
(SD)	(2.45)	(2.82)	(0.02)	(2.78)	(2.66)	(1.80)	(1.29)
Range%	0.0–8.5	0.0–7.9	0.0–0.1	0.0–11.0	0.0–13.6	1.7–8.3	1.0–4.6
CBG%	0.18 AB	0.22 AB	0.24 A	0.19 AB	0.14 B	0.08 B	0.11 AB
(SD)	(0.20)	(0.23)	(0.27)	(0.32)	(0.16)	(0.11)	(0.16)
Range%	0.0–1.0	0.0–0.7	0.0–1.1	0.0–1.8	0.0–0.7	0.0–0.3	0.0–0.5
THC%	3.54 B	3.04 B	5.48 A	6.49 A	1.16 C	0.39 C	0.17 C
(SD)	(2.58)	(2.12)	(2.41)	(4.09)	(2.05)	(0.61)	(0.08)
Range%	0.1–9.3	0.3–6.0	1.4–12.4	0.1–14.7	0.1–11.5	0.1–2.5	0.1–0.3
(CBD + THC)%	4.97 BC	4.99 BC	5.50 B	7.70 A	5.17 BC	4.01 C	3.19 C
(SD)	(2.61)	(1.91)	(2.42)	(3.45)	(2.59)	(1.83)	(1.37)
Range%	0.6–11.4	1.7–8.2	1.4–12.4	1.7–14.8	1.2–14.3	1.7–8.8	1.0–4.8
(CBDV + THCV)/i.s.	0.19 B	0.90 A	0.25 B	0.14 BC	0.05 C	0.09 BC	0.05 BC
(SD)	(0.35)	(0.80)	(0.40)	(0.30)	(0.06)	(0.10)	(0.05)
Range	0.0–1.6	0.0–2.7	0.0–2.1	0.0–1.4	0.0–0.3	0.0–0.3	0.0–0.1
CBGM/i.s.	0.05 A	0.00 C	0.01 C	0.02 B	0.01 BC	0.00 BC	0.01 BC
(SD)	(0.05)	(0.01)	(0.01)	(0.03)	(0.03)	(0.01)	(0.01)
Range	0.0–0.18	0.0–0.02	0.0–0.05	0.0–0.14	0.0–0.15	0.0–0.03	0.0–0.03

^a Means (in rows) not connected by the same letter are significantly different using Student's *t* test (*P* ≤ 0.05).

the genotype *B_D/B_D*, plants with a strong THC band were assigned the genotype *B_T/B_T*, and plants with both bands at about equal intensity were assigned the genotype *B_D/B_T*.

Twenty-six percent of the 157 sample populations consisted entirely of genotype *B_D/B_D* individuals, and 29% consisted entirely of genotype *B_T/B_T* individuals. The remaining 45% consisted of a combination of genotype *B_D/B_D*, *B_D/B_T*, and/or *B_T/B_T* individuals in various proportions. Only data for *B_T* are reported because the frequency of *B_D* is complementary. The mean frequency of *B_T* ranged from 0.04 in putative *C. ruderalis* to 1.00 in the NLD biotype of *C. indica* (Table 5). It was significantly higher for the NLD biotype than for all other taxa. The mean frequency of *B_T* was also significantly higher for the other three biotypes of *C. indica* than for putative *C. ruderalis* and the two biotypes of *C. sativa*, which did not differ significantly. Estimates of the mean proportion of chemotype I individuals in the *C. indica* biotypes ranged from 56% in the feral biotype to 100% in the NLD biotype, while in *C. sativa* and putative *C. ruderalis* the estimated mean proportion of chemotype I individuals did not exceed 1%. However, es-

timates for individual accessions ranged from 0 to 30% in *C. sativa* and from 18 to 100% in *C. indica* (excluding a Chinese and a Japanese hemp accession that may have been purposely selected for chemotype III fixation.)

DISCUSSION

Chemotaxonomic differentiation—The a priori assignment of accessions to *C. sativa* and *C. indica* based on geographic origins and differences in allozyme allele frequencies is consistent with significant differences in cannabinoid levels and *B_T* allele frequencies between the two species. Recognition of *C. ruderalis* is not supported by the chemotaxonomic evidence because no significant differences were found between ruderal accessions from central Asia and eastern Europe. These results are consistent with Vavilov's two-species concept and his interpretation that *C. ruderalis* is synonymous with *C. sativa* var. *spontanea* Vav. (Vavilov, 1926; Vavilov and Bukinich, 1929).

This study confirms that the THC/CBD ratio of individual *Cannabis* plants can be assigned to one of three discrete chem-

TABLE 5. Arithmetic means^a, standard deviations, and ranges of the *B_T* allele frequencies for sample populations of 157 *Cannabis* accessions assigned to seven putative taxa. Allele *B_T* encodes Δ⁹-tetrahydrocannabinolic-acid synthase. Also shown are statistics estimating the proportion of chemotype I individuals within each taxon. *N* = number of accessions analyzed.

	<i>C. indica</i> Hemp Biotype <i>N</i> = 19	<i>C. indica</i> Feral Biotype <i>N</i> = 5	<i>C. indica</i> NLD Biotype <i>N</i> = 27	<i>C. indica</i> WLD Biotype <i>N</i> = 11	<i>C. sativa</i> Hemp Biotype <i>N</i> = 81	<i>C. sativa</i> Feral Biotype <i>N</i> = 8	<i>C. ruderalis</i> <i>N</i> = 6
<i>B_T</i> frequency	0.79 B	0.75 B	1.00 A	0.85 B	0.11 C	0.08 C	0.04 C
(SD)	(0.32)	(0.21)	(0.00)	(0.28)	(0.14)	(0.15)	(0.08)
Range	0.0–1.0	0.42–1.0	1.0–1.0	0.24–1.0	0.0–0.55	0.0–0.42	0.0–0.20
Chemotype I%	62 BC	56 C	100 A	72 B	1 D	1 D	<1 D
(SD)	37	30	0	37	6	6	2
Range %	27 ^b –100	18–100	100	17–100	0–30	0–18	0–4

^a Means not connected by the same letter are significantly different using Student's *t* test (*P* ≤ 0.05).

^b Excluding accessions Ch-1 and Jp-2 that both had an estimated proportion of chemotype I plants <1%.

otypes. The limits between chemotypes coincide with those reported by Vollner et al. (1986). As expected, plants with high levels of THC were common within the two drug biotypes of *C. indica*. However, plants with relatively high levels of THC were also common within the hemp and feral biotypes of this species. In contrast, most plants assigned to *C. sativa* had relatively low levels of THC. Because chemotype I, II, and III plants were found in both species, the chemotype of an individual plant is of limited use for chemotaxonomic determination of species membership.

It was estimated that chemotype I individuals comprised <25% of each population for all but two of the 89 accessions assigned to *C. sativa* and >25% of each population for all but four of the 62 accessions assigned to *C. indica*. This appears to be a reasonable guideline for differentiation of the two species, in conjunction with other taxonomic traits. The advantage of this approach over the method of Small and Cronquist (1976) for differentiating *sativa* from *indica* (regardless of taxonomic rank) is that the frequency of chemotype I plants in a given population is stable from one generation to the next (assuming Hardy–Weinberg equilibrium) and not influenced by biotic and abiotic factors that affect the quantitative production of cannabinoids. For populations that have been purposely selected for a fixed chemotype, geographic origin and morphological traits are better indicators of species membership (Hillig, 2004, in press).

Small and Beckstead (1973a) reported that chemotype I strains usually originate from countries south of latitude 30°N and that chemotype II and III strains usually originate from countries north of this latitude. In the present study, nearly all accessions having a relatively low B_T frequency (<0.4) originated from latitudes north of 35°N. However, several accessions with a relatively high B_T frequency (>0.4) also originated from above this latitude, including drug accessions from Afghanistan and Pakistan and hemp accessions from China, Japan, and South Korea. It appears that the range of *C. sativa* in Europe and Asia does not extend below about 35°N latitude, whereas the range of *C. indica* extends both above and below this latitude (Hillig, 2004).

The propyl side-chain homologs of CBD and THC are also of chemotaxonomic significance. Elevated levels of CBDV and/or THCV were much more common in plants of *C. indica* than in plants of *C. sativa*. Plants with elevated levels of THCV, sometimes exceeding THC, were detected in all four biotypes of *C. indica*, but not in all accessions. Segregation ratios in F_2 populations from controlled crosses between low and high THCV individuals indicate that at least two loci control this trait (de Meijer et al., 2003; K. Hillig, unpublished data). The gene(s) controlling the enhanced biosynthesis of propyl cannabinoids appear to have originated in *C. indica* and not to have spread appreciably into *C. sativa*.

Small and Beckstead (1973b) reported that plants of several accessions from China, Japan, and South Korea had enhanced levels of CBGM. They assigned these accessions to a fourth chemotype, but Small and Cronquist (1976) did not treat this group as a separate taxon. In the present study, a relatively high apparent level of CBGM was most common in plants assigned to the hemp biotype of *C. indica*. This trait appears to be a useful chemotaxonomic marker for differentiating this taxon.

For the purpose of hypothesis testing, Hillig (2004) assigned hemp landraces from southern and eastern Asia to *C. chinensis* Delile. Although Small and Cronquist (1976) reduced *C. chi-*

nensis to synonymy with *C. sativa* subsp. *sativa* var. *sativa* (together with European hemp), an analysis of allozyme allele frequencies shows that the southern and eastern Asian hemp accessions in the germplasm collection under study derive from the *C. indica* gene pool (Hillig, 2004). In general, the southern and eastern Asian hemp accessions consisted mostly of chemotype I plants, although they tended to produce less THC than accessions assigned to the drug biotypes. This is consistent with the results of Rowan and Fairbairn (1977) who reported relatively high THC/CBD ratios but low THC content in plants grown from birdseed of Chinese origin. Hong and Clarke (1996) reported that hemp strains cultivated in China are generally lower in THC than drug strains, but THC/CBD ratios were not reported. European hemp breeders have struggled for over 50 years to reduce THC levels in Chinese hemp, partially because they did not recognize that eastern Asian hemp is a biotype of *C. indica* (Bredemann et al., 1956).

For systematic studies, it is best to obtain achenes of southern and eastern Asian strains directly from sources in these regions because of the possibility that such strains in European germplasm collections have hybridized with *C. sativa*. The *Cannabis* germplasm collection studied by Small and Beckstead (1973b) included accessions labeled “gigantea,” “sinensis,” “indica,” and “himalayana” that were obtained from European sources, although these labels suggest that these strains were of Asian origin. Small and Beckstead categorized these accessions as chemotype II or III populations, in contrast with accessions obtained directly from China, India, Japan, and South Korea, several of which were categorized as chemotype I populations. A few strains from China and Japan with relatively high THC/CBD ratios were categorized with the moderate to low (<0.5% d.w.) THC strains, apparently due to their late maturity and low resin production. It seems likely that several of the aforementioned strains that Small and Beckstead obtained from European sources were hybridized with *C. sativa* and that several strains from southern and eastern Asian sources that they included in the high THC category were cultivated for fiber and/or seed production, and not for drug production.

Quantitative differences—The cannabinoid levels reported in this investigation are approximately four to five times greater than those reported by Small and Beckstead (1973b). About 20% of the chemotype III plants had THC levels >0.30% d.w. of the sampled plant material, which exceeds the arbitrary divide established by Small and Beckstead between “non-psychotomimetic” and “psychotomimetic” plants. This 0.3% value is used in several countries as an upper limit for allowable THC levels in industrial hemp strains (Small, 1999). The present study shows that an upper limit of 0.8% d.w. of THC would be more consistent with the natural range of variation among chemotype III plants, based on a hypothetical maximum d.w. of 16% CBD in pistillate inflorescences and a 1 : 20 ratio of THC : CBD.

Although CBD was not detected in some chemotype I plants, separate GC analyses in which more highly concentrated extracts were analyzed show that CBD was indeed present in these plants, but below the threshold of detection in the present study (K. Hillig, unpublished data, Indiana University). This may also be the case in other studies that reported the absence of CBD in certain drug strains (summarized in Baker et al., 1980). About 5% as much THC relative to CBD was produced in chemotype III plants, which is close to

the 4% value reported by Fournier and Paris (1980). Reasons for the low level production of CBD in chemotype I plants and of THC in chemotype III plants are unknown. De Meijer et al. (2003) hypothesized that more than one variant of the CBD-acid synthase and/or THC-acid synthase alleles are present in the *Cannabis* gene pool and that these variant alleles encode allozymes with different catalytic efficiencies. They proposed that different combinations of these alleles result in somewhat different THC/CBD ratios among heterozygous (B_D/B_T) individuals. The histogram in Fig. 3 appears to show a bimodal distribution among chemotype II plants, consistent with this hypothesis. However, only two alleles (B_D and B_T) were inferred from the CBD and THC banding patterns visualized by starch gel electrophoresis.

Barni-Comparini et al. (1984) reported the complete absence of CBC in chemotype II and III plants. In the present study, CBC was detected in all three chemotypes but not in all plants. The high correlation ($r = 0.79$) between CBC and CBD levels in chemotype II and III plants has not previously been reported. Curiously, the highest levels of CBC were detected in staminate flowers of chemotype I plants assigned to the hemp biotype of *C. indica* (K. Hillig, unpublished data, Indiana University).

Domestication and evolution—Small and Cronquist (1976) assumed that the *sativa* and *indica* taxa diverged primarily as a result of human selection for fiber and/or seed production on the one hand and for high THC production on the other. However, the present investigation does not support this hypothesis. The high frequency of B_T in the feral biotype of *C. indica* suggests that this allele may have been present at high frequency in the *C. indica* gene pool prior to domestication. Human selection of plants carrying two copies of the B_T allele appears only to be of appreciable significance in the domestication of the NLD biotype. Human selection may have resulted in an increase in the quantitative levels of cannabinoids produced by the WLD biotype, but the average amount of CBD + THC produced by the NLD biotype did not significantly differ from the hemp and feral biotypes of *C. indica*. In fact, the average amount of THC + CBD produced by the NLD accessions was not significantly greater than the average amount of these two cannabinoids produced by the hemp accessions of *C. sativa*. Small and Beckstead (1973b) also reported comparable levels of total cannabinoids in their “psychotomimetic” and “non-psychotomimetic” strains. Plants with enhanced levels of THC were uncommon in most drug accessions, except those from southern Africa and an accession from Afghanistan. This suggests that humans may have selected against this trait in cultivated drug strains.

In contrast with the NLD biotype, the WLD biotype did not have a significantly higher frequency of B_T than the hemp and feral biotypes of *C. indica*. This may be explained by the different products obtained from the two drug biotypes of *C. indica*. NLD strains are usually cultivated for the production of marijuana (pistillate inflorescences), and it is the product of a single plant that is utilized. WLD strains are traditionally cultivated for the production of hashish (detached glandular trichomes), which is the combined product of many plants. A marijuana plant with two C_D alleles (chemotype III) would be ineffectual for its intended use, whereas hashish made from a population of chemotype I, II, and III individuals typically contains more THC than CBD and depending on the proportions of the different chemotypes is more or less psychoactive

(Ek et al., 1972; Clarke, 1998). Thus, human selection is expected to favor chemotype I plants as seed sources for marijuana cultivation, whereas the chemotypes of individual plants cultivated for hashish production are usually unknown to the cultivator (Clarke and Watson, 2002).

The presence of allele B_T in the *C. sativa* gene pool suggests that introgression from *C. indica* might have played a role in the evolution of *C. sativa*. Wind-blown pollen may have contributed to allele migration between the two gene pools (Cabezudo et al., 1997). Relatively high B_T frequencies (range 0.38–0.55) were detected in seven hemp accessions from Turkey, Spain, Italy, former Yugoslavia, and southern Russia, which are assignable to the southern eco-geographical group of *C. sativa* (Davidyan, 1972). Additional allozyme markers and morphological traits typical of *C. indica* were also observed in the southern group of *C. sativa* (Hillig, 2004, in press). *Cannabis indica* may have been introduced into Asia Minor for the purpose of hashish production and hybridized with *C. sativa*. Davidian (1972) cited evidence that *Cannabis* was introduced into Europe by both a northern and a southern route. Introgressed stock may have spread into new areas through trade or human migration (Heiser, 1973).

The patterns of cannabinoid variation provide evidence of progenitor-derivative relationships. The low frequency of B_T and the low levels of propyl cannabinoids in accessions assigned to *C. ruderalis* suggest that this putative taxon could be the progenitor of *C. sativa*, but not of *C. indica*. The feral biotype of *C. sativa* may be comprised of “escaped” populations of cultivated *C. sativa* that have merged with naturalized populations of *C. ruderalis* (Vavilov, 1926). The wide range of cannabinoid variation within feral accessions of *C. indica* suggests that this biotype could be the progenitor of the cultivated biotypes of *C. indica*. The high incidence of plants in this taxon with enhanced levels of propyl cannabinoids suggests that it is not the progenitor of *C. sativa*, in which plants with enhanced levels of propyl cannabinoids are much less common. It is unlikely that the feral biotype of *C. indica* represents an escape of NLD strains from cultivation because the NLD biotype is lacking in cannabinoid variation. The high frequency of B_T in the hemp biotype of *C. indica* suggests the possibility that one or both drug biotypes could have been secondarily derived from this taxon’s gene pool. More feral and cultivated populations of *C. indica* and *C. sativa* will have to be studied to further resolve these issues.

This study of cannabinoid variation supports a two-species concept for *Cannabis*. A taxonomic revision that applies valid scientific names to the biotypes of *C. indica* recognized herein is supported by the chemotaxonomic data.

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