First Direct Evidence for the Mechanism of Δ^1 -Tetrahydrocannabinolic Acid Biosynthesis

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Numerous plausible hypotheses have been advanced regarding the biogenesis of Δ^1 -tetrahydrocannabinol (Δ^1 -THC, **1a**), the psychoactive constituent of marihuana (Cannabis sativa L.);1 however, they all lack experimental support. Thus all hypothetical biogenetic schemes assume that Δ^1 -tetrahydrocannabinolic acid (Δ^1 -THCA, **1b**), the precursor of Δ^1 -THC, is formed by cyclization from cannabidiolic acid (2).² We now present experimental evidence that establishes that 1b is actually formed from cannabigerolic acid (3b) through oxidocyclization by a new enzyme, Δ^1 -THCA synthase, and that 2, a plausible intermediate, is not a substrate for this enzymatic reaction.

Since the enzymes that catalyze the formation of 1b have not been purified and studied, we first attempted to identify an enzyme in C. sativa that could cyclize 2 into 1b. Using various extraction and assay conditions we were unable to obtain such an enzyme preparation; however, by contrast, when 3b was used as a substrate,³ the crude enzyme preparation extracted with 1 M CaCl₂ exhibited a potent Δ^1 -THCA-producing activity.⁴ We assume therefore that 1b is derived biosynthetically by the oxidative cyclization of **3b** rather than by the ring closure of **2**.

In order to characterize the oxidocyclase (Δ^1 -THCA synthase), we attempted to purify this new enzyme. Leaf buds of the Mexican Cannabis strain were used for the extraction of the enzyme,⁵ as they showed a much higher enzyme activity than any other parts of this plant. The Δ^1 -THCA synthase was purified to homogeneity by chromatography on (diethylaminoethyl)cellulose, phenyl Sepharose CL-4B, and hydroxylapatite.6 The purified enzyme migrated on sodium dodecyl sulfatepolyacrylamide gel electrophoresis (SDS-PAGE) as a single band with a molecular mass of 75 kD (Figure 1). The native molecular mass determined by gel filtration using Sephadex G-75 was 74 kDa, indicating that Δ^1 -THCA synthase is a monomeric enzyme. The isoelectric point for this enzyme was determined as 6.4 by isoelectric focusing. The purified enzyme was sequenced by Edman degradation, and the following NH2terminal sequence was identified: Asn-Pro-Arg-Glu-Asn-Phe-



Figure 1. SDS-PAGE (12.5% gel) of purified Δ^1 -THCA synthase. The gel was stained with Coomassie Blue. Lane 1: molecular mass standards. Lane 2: purified Δ^1 -THCA synthase (1 μ g). Numbers at left indicate positions of molecular mass standards in kilodaltons.

Leu-Lys-x-Phe-Ser-Lys-His-Ile-Pro-Asn.7 The ∆1-THCA synthase is the first enzyme involved in cannabinoid biogenesis to be identified and purified. The Δ^1 -THCA (1b) produced by the enzymatic reaction⁸ was found to be identical with 1b isolated from the plant by comparison of their CD spectra as well as their FAB-MS. Therefore, the purified Δ^1 -THCA synthase was confirmed to catalyze the stereoselective conversion of 3b to 1b.

In order to evaluate the kinetic properties of Δ^1 -THCA synthase, the enzyme activity was measured under various conditions (Table 1). Since the Δ^1 -THCA synthase activity was highest between pH 5.5 and 6.0, an assay system with 200 μ M 3b at pH 6.0 was chosen as a standard.⁹ Ions such as Mg²⁺ and Mn²⁺, which are known to be cofactors for monoterpene cyclase, 10 had little influence on the Δ^1 -THCA synthase activity, whereas Hg²⁺ strongly inhibited the production of 1b. It is noteworthy that Δ^1 -THCA synthase is not inhibited by the absence of molecular oxygen, contrary to oxygenases and oxidases which absolutely require molecular oxygen for the

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⁽³⁾ Owing to the low CBGA content in C. sativa, 3b was chemically synthesized by coupling of olivetol (Sigma) and geraniol (Aldrich),¹ followed by carboxylation with methylmagnesium carbonate.

⁽⁴⁾ The enzyme activity was determined by HPLC. HPLC was carried out under essentially the same conditions as previously described,¹⁴ but 85% aqueous CH₃CN containing 50 mM phosphoric acid was used for this study as a solvent system.

⁽⁵⁾ Extraction was carried out at 4 °C. Leaf buds (66 g) of C. sativa (Mexican strain) were homogenized with a Waring blender at high speed together with 800 mL of 1 M CaCl₂ containing 10 mM mercaptoethanol. The homogenate was centrifuged at 20000 g for 15 min. The supernatant fluid was concentrated by ultrafiltration and dialyzed against 10 mM aqueous mercaptoethanol. During dialysis, most of the Δ^1 -THCA synthase activity precipitated. The resulting precipitates were collected by centrifugation at 49000g for 60 min and resuspended in 10 mM sodium phosphate buffer (pH 7.0) containing 3 mM mercaptoethanol. The insoluble material was removed by centrifugation at 49000g for 60 min, and the supernatant (crude enzyme extracts) was used for the purification of the Δ^1 -THCA synthase.

⁽⁶⁾ All purification procedures were conducted at 4 °C. The following buffers were used for the purification of the Δ^1 -THCA synthase: buffer A, 10 mM sodium phosphate buffer (pH 7.0) containing 3 mM mercaptoethanol; buffer B, buffer A containing 0.8 M ammonium sulfate; buffer C, 250 mM sodium phosphate buffer (pH 7.0) containing 3 mM mercaptoethanol. The crude enzyme extracts as described in footnote 5 were applied to a Whatman DE-52 chromatographic column (2.5 × 24 cm) using buffer A (flow rate: 1.7 mL/min). The enzyme was eluted soon after the void peak (void volume: 65 mL). This step gave a 3-fold purification, while approximately 45% of loaded activity was recovered by this procedure. The Δ^1 -THCA synthase active fractions were concentrated by ultrafiltration. After ammonium sulfate was added to the concentrated sample to raise the concentration to 0.8 M, the sample was loaded onto a phenyl Sepharose CL-4B column (Pharmacia; 1.5 × 14 cm) previously equilibrated with buffer B. Δ^1 -THCA synthase was eluted by a 500-mL linear gradient of buffer B to buffer A (flow rate: 1.5 mL/min), resulting in an enrichment of ca. 152fold. The Δ^1 -THCA synthase active fractions eluted at about 0.3 M ammonium sulfate were concentrated by ultrafiltration and dialyzed against buffer A. A final purification was achieved by hydroxylapatite chromatography (Nacalai Tesque, Japan; 1.0 × 8 cm) pre-equilibrated with buffer buffer A to buffer C followed at a flow rate of 1.2 mL/min. The enzyme was eluted at about 230 mM phosphate buffer. The overall yield of Δ^1 -THCA synthase from this purification scheme was 3.3%. The purified enzyme (250 µg) was stored at 4 °C in buffer A

⁽⁷⁾ The letter "x" indicates that the identity of the residue is ambiguous. (8) In the presence of the purified enzyme (50 μ g), the substrate buffer (500 mL), which was composed of 100 mM sodium phosphate buffer (pH 6.0), 200 µM 3b, and 0.1% (w/v) Triton X-100, was incubated for 6 h at 30 °C. The reaction mixture was partitioned with AcOEt. The AcOEt layer was subjected to HPLC⁴ to yield **1b** (0.5 mg). (9) Standard assay consisted of the substrate buffer⁸ (500 μ L) and enzyme

solution (100 μ L). After the sample was incubated for 2 h at 30 °C, the reaction was stopped by the addition of 600 μ L of MeOH.

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Table 1. Δ^1 -THCA Synthase Activity under Various Conditions

rel act. ^a	conditns	rel act. ^a
100	1 mM NAD	101
104	1 mM NADP	101
103	1 mM FAD	64
4	1 mM FMN	70
103	$1 \text{ mM H}_2\text{O}_2$	78
91	N_2^b	99
	rel act. ^a 100 104 103 4 103 91	rel act. ^a conditns 100 1 mM NAD 104 1 mM NADP 103 1 mM FAD 4 1 mM FMN 103 1 mM H ₂ O ₂ 91 N ₂ ^b

^a Data were obtained means of three replicated assays. ^b To remove molecular oxygen the standard assay solution was saturated with N₂ gas.

oxidation of substrates. Furthermore, addition of hydrogen peroxide did not stimulate the enzyme, indicating that Δ^1 -THCA synthase is not peroxidase. Therefore, Δ^1 -THCA synthase should be a dehydrogenase, although coenzymes such as NAD, NADP, FAD, and FMN did not stimulate the enzyme activity. Inhibition by both flavins was shown to be due to nonenzymatical degradation of **3b**. Purified Δ^1 -THCA synthese displayed a high activity (V_{max} : 2.68 nkat/(mg of enzyme)) and a high affinity (K_m : 134 μ M) for 3b.¹¹ Cannabinerolic acid (4b),¹² which is a Z isomer of 3b, is also converted to 1b by Δ^1 -THCA synthase, although lower specificity (V_{max} : 0.37 nkat/ (mg of enzyme), $K_{\rm m}$: 254 μ M) for 4b was found as compared to 3b. The C-1/C-2 double bond of 4b has the same configuration as that of 1b, suggesting that 4b could be an intermediate in the oxidocyclization of 3b into 1b. However, the lower activity for 4b shows that the enzymatic reaction does not proceed from 3b through 4b to 1b. We propose that 1b is formed via a common intermediate in the reactions of both 3b and 4b (Figure 2). Recently, kinetic properties similar to those reported above were found for a monoterpene cyclase (limonene

823.



Figure 2. Mechanism of Δ^{1} -THCA (1b) biosynthesis by Δ^{1} -THCA synthase.

synthase), which catalyzes limonene formation with higher V_{max} and lower K_m values for geranyl pyrophosphate as compared to neryl pyrophosphate.^{10b} The limonene synthase, whose full amino acid sequence was revealed,^{10c} did not possess a region similar to the NH₂-terminal sequence of Δ^1 -THCA synthase. Various monoterpene cyclases including limonene synthase have been purified and characterized; however, cyclization by such enzymes was not accompanied by an oxidative reaction.¹⁰ The Δ^1 -THCA synthase described here is apparently a unique cyclase. Δ^1 -THCA synthase did not convert the neutral cannabinoids cannabigerol (3a) and cannabinerol (4a) to 1a, indicating that the presence of the carboxyl group in the substrate is essential for enzymatic cyclization of the terpene moieties.

In conclusion, we have identified C. sativa a new oxidocyclase, which converts 3b and 4b to 1b. This reaction does not proceed through the generally postulated precursor 2. These findings cast serious doubt on the hypothesis that 2 is a biosynthetic precursor of 1b.

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⁽¹¹⁾ V_{max} and K_{m} values were determined by Lineweaver-Burk doublereciprocal plots of the velocity curves of the THCA-producing reaction with concentration of substrates. The assays were carried out with 0.34 μ g of the purified enzyme.

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