HU-444, A Novel, Potent Anti-Inflammatory, Non-Psychotropic Cannabinoid

Christeene G. Haj, Percy F. Sumariwalla, Lumír Hanuš, Natalya M. Kogan, Zhana Yektin, Raphael Mechoulam, Mark Feldmann and Ruth Gallily

Institute for Drug Research, Hebrew University Medical Faculty, Jerusalem 91120, Israel (C.G.H., L.H., N.M.K., R.M.), Kennedy Institute of Rheumatology, Hammersmith, London W67BA, United Kingdom (P.F.S., M.F.) and Lautenberg Center for Immunology, Hebrew University Medical Faculty, Jerusalem 91120, Israel (Z.Y., R.G.)
Running title page

A) Anti-inflammatory effects of the HU-444 Cannabinoid

B) Corresponding authors: Prof. Raphael Mechoulam, Institute for Drug Research, Hebrew University Medical Faculty, Jerusalem 91120, Israel. Tel: 00972-2-6758634, Fax: 00972-6758073, Email: mechou@cc.huji.ac.il; Prof. Ruth Gallily, Lautenberg Center for Immunology, Hebrew University Medical Faculty, Jerusalem 91120, Israel. Tel: 00972-2-6758712, Fax: 00972-2-6757296, E-mail: ruthg@ekmd.huji.ac.il.

C) The number of text pages: 28

The number of tables: 1
The number of figures: 7
The number of references: 54
The number of words in the Abstract: 178
The number of words in the Introduction: 520
The number of words in the Discussion: 952

D) Abbreviations: CII, type II collagen; CBD, cannabidiol; CIA, collagen-induced arthritis; Con A, Concanavalin A; HU-444, (-)-8,9-dihydro-CBD-7-oic acid-diacetate; DMEM, Dulbecco’s modified Eagle medium; FCS, fetal calf serum; LPS, lipopolysaccharide; NO, nitric oxide; RA, rheumatoid arthritis; RPMI-1640, Roswell Park Memorial Institute-1640 medium; THC, Δ⁹-tetrahydrocannabinol; TG, thioglycollate; TGMΦ, thioglycollate-elicited peritoneal macrophages; TNF-α, tumor necrosis factor alpha.

E) A recommended section assignment: Inflammation.
A B S T R A C T

Cannabidiol (CBD) is a component of cannabis, which does not cause the typical marijuana-type effects, but has a high potential for use in several therapeutic areas. In contrast to Δ⁹-tetrahydrocannabinol (Δ⁹-THC) it binds very weakly to the CB1 and CB2 cannabinoid receptors. It has potent activity in both in vitro and in vivo anti-inflammatory assays. Thus, it lowers the formation of TNF-α, a proinflammatory cytokine, and was found to be an oral anti-arthritic therapeutic in murine collagen-induced arthritis in vivo. However in acidic media it can cyclize to the psychoactive Δ⁹-THC. We report the synthesis of a novel CBD derivative, HU-444, which cannot be converted by acid cyclization into a Δ⁹-THC-like compound. In vitro HU-444 had anti-inflammatory activity (decrease of reactive oxygen intermediates and inhibition of TNF-α production by macrophages); in vivo it led to suppression of production of TNF-α and amelioration of liver damage as well as lowering of mouse collagen-induced arthritis. HU-444 did not cause Δ⁹-THC-like effects in mice. We believe that HU-444 represents a potential novel drug for rheumatoid arthritis and other inflammatory diseases.
Introduction

Cannabidiol (CBD), a nonpsychotropic constituent present in most *Cannabis sativa* varieties, causes a large number of both central and peripheral pharmacological effects (Mechoulam et al., 2002; Mechoulam et al., 2007; Pertwee, 2005; Zhornitsky and Potvin, 2012; Fernández-Ruiz et al., 2013). CBD is a potent anti-oxidant (Hampson et al., 2000) which may explain – at least in part – its neuroprotective effects in neurodegenerative disorders (Fernández-Ruiz et al., 2013), in amelioration of the progressive degeneration of nigrostriatal dopaminergic neurons occurring in a model of Parkinson's disease (Lastres-Becker et al., 2005), in cerebral ischemia (Braida et al., 2003), in cerebral infarction in mice (Hayakawa et al., 2007), and in hypoxia-ischemia in newborn rats (Pazos et al., 2012). Additional effects of therapeutic relevance, through various mechanisms, are its action in animals on type-1 diabetes (Weiss et al., 2008), on some types of cancer (Massi et al., 2013), on myocardial ischemic reperfusion injury (Durst et al., 2007), on reduction of microglial activation – thus possibly on the progression of Alzheimer's disease (Martín-Moreno et al., 2011), on brain and liver functions in a fulminant hepatic failure-induced model of hepatic encephalopathy (Avraham et al., 2011), on nausea and emesis (Rock et al., 2012) and others.

The anti-epileptic activity of CBD in human patients has been known for nearly 35 years (Cunha et al., 1980), but only recently have marijuana extracts with high levels of CBD been used to suppress pediatric epilepsies (Porter and Jacobson, 2013). In clinical trials CBD has been shown to have anti-schizophrenic (Leweke et al., 2012) and anti-anxiety (Bergamaschi et al., 2011) properties.
Of particular relevance to the results presented here is the potent anti-inflammatory action of CBD (Mechoulam et al., 2002; Mechoulam et al., 2005; Pertwee, 2005; Mechoulam et al., 2005). We have previously shown that, *in vitro*, CBD suppresses lymphocyte proliferation and blocks zymosan-triggered reactive oxygen burst by peritoneal granulocytes (Malfait et al., 2000). *In vivo* results reported include blocking lipopolysaccharide (LPS)-induced rise in serum tumor necrosis factor (TNF-α) in mice as well as the progression of arthritis caused by immunization of mice with type II collagen (CII) (Malfait et al., 2000).

It is somewhat surprising that in spite of the very promising pharmacological effects of CBD and its lack of toxicity it has not been developed as a single drug. CBD is marketed together with Δ⁹-THC (in a 1:1 ratio) by GW Pharmaceuticals as Sativex, sold in Canada and several European countries for spasticity, due to multiple sclerosis (Syed et al., 2014).

We report now the synthesis of a novel CBD derivative, HU-444, using CBD as the starting material, and the evaluation of its anti-inflammatory properties *in vitro* and *in vivo*. This novel compound was chosen as the conversion of the C-1 methyl group into a carboxyl group in the cannabinoid series has been shown to enhance anti-inflammatory activity (Sumariwalla et al., 2004; Burstein et al., 1992) and the reduction of the 8,9-double bond would preclude possible ring cyclization under acidic conditions with one of the phenolic groups leading to psychoactive THC-like compounds.

The results reported now show that HU-444 is a potent anti-inflammatory compound, both *in vitro* and *in vivo*. 

Materials and methods

Reagents. All solvents were purchased from Biolab LTD. (Jerusalem, Israel) and J.T.Baker (Deventer, Holland). Chemicals were purchased from Sigma-Aldrich (Rehovot, Israel), Acros, Holland Moran LTD (Yehud, Israel), Alfa Aesar (Lancashire, UK), Merck (Darmstadt, Germany), J.T. Baker (Center Valley, PA, USA) and Penta (Prague, Czech Republic). They were used without further purification in the reactions except of dry diethyl ether and CH₂Cl₂, which were refluxed over sodium and phosphorous pentoxide (P₂O₅) respectively, and freshly distilled prior to its use.

¹H NMR spectra were obtained on a Bruker AMX 300 MHz apparatus using CDCl₃ (δ=7.25 ppm) and TMS as internal standard for ¹H NMR. Thin-layer chromatography (TLC) was run on silica gel 60F₂₅₄ plates (Merck). Column chromatography was performed on silica gel 60 Å, Merck. The compounds were localized at 254 nm using a UV lamp.

Syntheses. The synthesis of HU-444 is presented in Scheme 1. The synthetic steps a, b, and c are published in Kozela et al. (2015) (In press). Below we describe in detail the subsequent reaction steps d-i.

Syntheses of (-)-8,9-dihydro-CBD-7-oic acid-diacetate (HU-444) (5) and (-)-8,9-dihydro-CBD-7-oic acid (HU-445) (6)

Cannabidiol (CBD). Crystalline CBD was isolated from hashish following the procedure described by Gaoni and Mechoulam (Gaoni and Mechoulam, 1971). Yield 15%; melting point: 62°C; [α]D²⁰ = -56° (CHCl₃).

Synthesis of (-)-8,9-dihydro-7-oxo-CBD-diacetate and (-)-8,9-dihydro-6-oxo-CBD-diacetate (mixture 4). Dry chromium oxide (353 mg, 3.53 mmol) was added to a stirred
solution of dry pyridine (0.951 ml, 11.83 mmol) in dry CH₂Cl₂:DMF (4:1) (4 ml) and stirred at room temperature for 15 min under nitrogen. A solution of mixture 3 (378 mg, 0.91 mmol) in dry CH₂Cl₂:DMF (4:1) (1.5 ml) was added and the reaction was stirred at room temperature for 1 hour. Ethanol (2 ml) was added and the reaction was stirred at room temperature for 10 min. The mixture was filtrated on silica gel covered with thin layer of Na₂SO₄, washed several times with ethyl acetate and evaporated to dryness. The residue was chromatographed on silica gel with (10% ether – petroleum ether) to give mixture 4 without separation with a yield of 63%.

**Synthesis of (-)-8,9-dihydro-CBD-7-oic acid- diacetate – (HU-444) (5)**

NaClO₂ (221 mg, 2.44 mmol) was added in small quantities to a stirred mixture of mixture 4 (235 mg, 0.568 mmol), 2-methyl-2-butene (1.5 ml, 14.22 mmol) and a saturated aqueous solution of KH₂PO₄ (0.67 ml) in tert-butanol (13.37 ml). The reaction was stirred at room temperature for 5 hours, and monitored by TLC. Water was added (60 ml) and the mixture was extracted several times with ethyl acetate. The organic phase was washed with brine, dried over MgSO₄ and filtered. Removal of the solvent under reduced pressure afforded a residue that was chromatographed on silica gel with (30% ether – petroleum ether) to give compound 5 as a white solid. Yield 13%, Mp: 152 °C; [α]D<sup>20</sup> = -57° (CHCl₃); HPLC (60% acetonitrile, 15% water and 25% methanol): t<sub>R</sub> = 7.69 min, 96%. ¹H NMR (CDCl₃): δ 6.94 (1H,s,olefin), 6.88 (2H,s,Ar), 3.58 (1H,m,benzyl), 2.51 (3H, m, allyl + benzyl), 2.22 (6H, s, OAc), 1.97 (1H, m), 1.86 (1H, m), 1.54 (5H, br s), 1.32 (5H, m), 0.98 (9H, t, terminal CH₃). MS m/z: 502 (silylation), 472, 430, 415, 400. Exact mass calculated for C<sub>25</sub>H₃₄O₆ 430.2199, Found 430.2198.
Synthesis of (-)-8,9-dihydro-CBD-7-oic acid – (HU-445) (6). Compound 5 (30 mg, 0.069 mmol) was dissolved in ethanol (8 ml). NaBH₄ (3 mg, 0.092 mmol) was added and the reaction was refluxed for 1 hour. The ethanol was removed under pressure, the residue was diluted with water (10 ml) and the solution was extracted with ether. The combined organic extracts were washed with brine, dried over MgSO₄ and filtered. Removal of the solvents under reduced pressure afforded a residue that was chromatographed on silica gel with (30% ether – petroleum ether) to give compound 6 as a white solid with a yield of 90%, Mp: 88-90°C; 

[α]D²⁰ = -56.2° (CHCl₃); HPLC (60% acetonitrile, 15% water and 25% methanol): tₚ = 8.86 min, 95%. ¹H NMR (CDCl₃): δ 6.94 (1H, s, olefin), 6.88 (2H, s, Ar), 4.12 (2H, br s), 3.59 (1H, m, benzyl), 2.51 (3H, m, allyl + benzyl), 2.22 (3H, s, OAc), 1.97 (1H, m), 1.86 (1H, m), 1.55 (5H, br s), 1.32 (5H, m), 0.98 (9H, t, terminal CH₃). MS m/z: 562 (silylation), 489, 416, 346, 343. Exact mass calculated for C₂₁H₂₈O₄ 346.1987, Found 346.1948.

Binding to the cannabinoid receptors. The binding of HU-444 and HU-445 to the cannabinoid receptors CB1 and CB2 was performed as previously described (Devane et al., 1992; Devane et al., 1992; Bayewitch et al., 1996).

Mice. In Jerusalem C57BL/6 and Sabra female mice, 8-10 weeks old were obtained from Harlan Laboratories, Israel; in London DBA/1 male mice, 12 weeks old, were obtained from Harlan laboratories, UK. Mice were housed in a room with controlled temperature (22 ± 1°C), humidity (60% ± 1%) and light 12h per day. Mice were fed standard animal chow and tap water ad libitum and were kept in a specific pathogen free (SPF) facility. The animal research in Jerusalem was conducted in compliance with
international laws and approved by the Ethical Committee of the Hebrew University Medical School, while in London all animal studies conducted had received prior approval of the local ethical review process committee, and were performed under the guidance of the Home Office Animals (Scientific Procedures) Act 1986 (PPL: 70/5446).

**Macrophages.** Peritoneal cells were harvested from C57BL/6 female mice 4 days after intraperitoneal injection of 1.5 ml of 3% thioglycollate (TG) medium (Difco). The cells (TG macrophages) were washed with phosphate-buffered saline, resuspended in Dulbecco’s modified Eagle medium (DMEM) supplemented with 10% fetal calf serum (FCS), sodium pyruvate, glutamine and antibiotics and plated \((1.2 \times 10^5)\) in 96-microwell flat-bottom plates (Nunc, Roskilde, Denmark). Following 2-3 h incubation at 37°C, the non-adherent cells were removed by intensive rinsing with PBS. About 95% of the adherent cells were macrophages.

**THC-like activity in mice.** THC and other agonists of the CB1 cannabinoid receptor cause a typical tetrad of pharmacological effects in Sabra mice – namely ring immobility (catalepsy), which measures the percentage of time, over 4 min, mice remain immobile on a ring (5.5 cm diameter), the open field test which measures locomotor activity, hypothermia and hot plate latency (antinociception) (Martin et al., 1991; Fride and Mechoulam, 1993). Sabra mice were administered HU-444 i.p. and assayed in the above tetrad.

**Macrophage cell lines.** RAW 264.7 cells, a monocytic-macrophage cell line, derived from BALB/c mice, were obtained from American Type Culture collection (Rockville, MD). The cells were cultured in Dulbecco's modified Eagle medium (DMEM)
supplemented with 10% fetal calf serum (FCS) and sodium pyruvate, glutamine, and antibiotics. For activation, the cells (10^5 cells/microwell) were incubated with 1 μg/ml lipopolysaccharide (LPS, E.coli Sigma, Israel) for 24 h.

**Treatment of macrophages with HU-444.** HU-444 was first dissolved in absolute ethanol (1 mg/50–100 μl ethanol), and the solutions were further diluted with DMEM medium. Various nontoxic concentrations were added to the macrophages, followed by addition of 1 μg/ml LPS dissolved in saline for activation. The macrophages were then cultivated in a humid atmosphere with 5% CO₂ for 24 h. The supernatant fluids were harvested and kept at −20°C until assayed for TNF-α.

**Reactive oxygen species (ROS) production by RAW 264.7 macrophages.** RAW 264.7 cells were removed using a scrapper, washed, and resuspended in Hanks’ balanced salt solution (without phenol red). For measurement of chemiluminescence, 0.5 ml of cell suspension (5 × 10^5 cells) was added to each luminometer tube, together with various doses of HU-444 (dissolved in ethanol and diluted with Hanks’ salt solution). Then, 10 μL of luminol (Sigma) and 30 μL containing 40 μg of zymosan (Sigma) were added to the tubes; the chemiluminescence was measured immediately in a Luminometer (Biolumate LB 95, Berhold, Wilbad, Germany).

**Evaluation of TNF-α levels produced in vitro and in vivo.** Peritoneal thioglycollate macrophages were cultured (1.2x10^5/microwell) for 2-3 h, rinsed in PBS and further incubated in DMEM with 5% FCS, in the presence of varying concentrations of HU-444. LPS was used as a stimulus at 1 μg/ml. The viability of the cells at the end of the experiment was >90%. The supernatants were collected and kept at -20°C until assayed.
To determine TNF-α levels in mice sera, C57BL/6 female mice (8-10 weeks old; 24 mice/experiment), were injected with 5 mg/kg LPS (i.p.) and then with various doses of HU-444. Ninety minutes later the mice were bled; the sera were kept at -20°C until assayed (Malfait et al., 2000). TNF-α levels both in sera and the cultures supernatants were determined by ELISA (R&D Systems, San Diego, California) according to the manufacturer's procedure.

**Nitric oxide (NO) detection assay.** NO generation was determined by measuring the nitrite accumulated in the supernatants (100 μL) of LPS (1 μg/ml) and HU-444-treated RAW macrophages as follows. An equal volume (100 μL) of Griess reagent (1% sulfanilamide, 0.1% naphthalene diamine HCl, 2% H₃PO₄) was added to each supernatant. Following 10 min of incubation at room temperature, the color production was measured at 550 nm with an ELISA reader. The concentration of nitrate was calculated according to a standard curve.

**Liver injury.** C57BL/6 male mice, 8-10 weeks old, total of 48 mice (in three experiments) were injected i.v. with 20 mg/kg Con A, dissolved in pathogen free saline (Tiegs et al., 1992). Immediately thereafter the mice were injected i.p. with various doses of HU-444. Twenty hours later, the mice were bled through the orbital plexus (for collection of serum for liver enzymes and TNFα assays) and sacrificed. The livers were removed, fixed with buffer formalin and stained with haematoxylin & eosin, for microscope evaluation. TNF-α levels in the sera of Con A-treated mice were determined by ELISA (R&D Systems).

**Determination of ALT and AST levels.** The levels of two aminotransferases, alanine aminotransferase (ALT) and aspartate aminotransferase (AST), were assayed in the sera
of Con A-treated mice, with or without HU-444 treatment, by ALT and AST strips respectively (Reforam-Mannheim GmbH, Mannheim, Germany) and quantitated by an automated analyzer (Reflotran Plus, Roche).

**Histological analysis.** The livers of Con A-injected mice, with or without HU-444 treatment, were fixed in 10% buffer-formalin. Paraffin sections of 4-5μm thickness, were stained with haematoxylin and eosin.

For microscopic evaluation, sections were examined (X40) blindly and scored 0-5 as follows: 0-normal liver morphology, 1- minimal changes, 2- mild, few lesions, 3- moderate number of lesions, 4- marked changes in the liver, 5- severe liver damage.

**Collagen-induced arthritis study: induction monitoring and treatment with HU-444.** Collagen-induced arthritis (CIA) was induced in genetically susceptible DBA/1 mice (H-2^a^, Harlan Laboratories, U.K.) by subcutaneous immunization, at sixteen weeks with purified type II chicken collagen emulsified with Freund’s complete adjuvant (100 μl of 2 mg/ml, divided over two sites in the lower dorsal region). Mice were purchased at 12 weeks of age and acclimatized in the animal house over a 20h light/dark cycle. Mice were housed in individual ventilated cages and the ambient temperature was maintained at 21°C. Mice were fed with standard laboratory chow and tap water *ad libitum*. Onset was characterized by visible clinical signs of arthritis, which were a varying degree of redness and swelling of any of the paws. Mice were treated with HU-444 either i.p. (graded doses of 2.5, 5.0 & 10 mg/kg) or orally (15 mg/kg), daily for ten days. For the purpose of administering various doses of HU-444 to mice the average weight of a mouse was considered as 25 g and nine arthritic mice were recruited in all study groups. All treatments were carried out from day 1, which is considered as the day when the first
visible clinical signs of arthritis developed in a mouse. Vehicle (ethanol: cremophor-EL: saline; 1:1:18 v/v/v for i.p. or 1:1:8 v/v/v for p.o. administrations respectively) was administered to a group of arthritic mice used as controls. Vehicle on its own and HU-444 in vehicle were administered in experiments at either 0.1 (oral gavage) or 0.2 (intra-peritoneal) milliliters. All paws were graded daily using previously well-validated clinical score read-outs (Sumariwalla et al., 2004). Hind paw thickness was measured (mm) using a calliper device. All treatment studies had the prior approval from a local ethical review process committee and were conducted in accordance to the Home office approved project license PPL: 70/5446.

**Histology of arthritic feet.** Mice were euthanized at the end of treatments and hind paws fixed in buffered formalin, decalcified in 10% EDTA for 3 to 4 weeks, embedded in paraffin wax, sectioned to 5 µm thicknesses and stained with haematoxylin and eosin. All joints were scored blindly for joint architectural changes by a well validated scoring system (Sumariwalla et al., 2003). In brief, score 0 represented normal joints, 1 mild, change involving synovial hyperplasia, chondrocyte de-nucleation and a few focal bone and cartilage erosions, 2 moderate, change involving extensive synovial hyperplasia and pannus front accompanying large areas of bone and cartilage erosions, 3 severe, total loss of joint architecture. For the purpose of scoring, all joints (distal, proximal phalangeal, first metatarsal, medial cuneiform and tibia-tarsus) present within the stained section were scored. Histological evaluation was done on all mice paw sections from all groups. Percentage of joints involved as normal, mild, moderate and severe were then calculated for each group. Data were presented as percentage joints scored and classified as either protected (normal to mild
arthritic changes) or damaged (moderate to severe arthritic changes). Images were captured using Olympus BX51 optical microscope and DP Controller and Manager Software (Olympus Corporation, version 3.3.1.292/222, USA), at a magnification of 100 X. For publication purposes, only the proximal phalangeal joint images were depicted for all treatment groups.

**Experimental Design and Statistics.** Experiments, both *in vitro* and *in vivo* (4-5 mice/group), were repeated three times and the results were expressed as the mean ± SEM of triplicate values. Evaluation of statistical significance was done by using analysis of variance (ANOVA). Joint histology data were evaluated by Fischer’s exact test. $P \leq 0.05$ values were considered significant.

**Results**

**Chemistry.** CBD, which was extracted from hashish (Gaoni and Mechoulam, 1971) was hydrogenated with Pt (IV) oxide in ethyl acetate to yield 8,9-dihydro-CBD 1 which was then converted into the diacetate 2 by pyridine and acetic anhydride. Compound 2 was oxidized with selenium IV oxide in ethanol to give a mixture of the allylic alcohols 3. The allylic hydroxylation reaction by selenium IV oxide has been widely investigated (Sharpless and Lauer, 1972; Lander et al., 1976; Stephenson and Speth, 1979). Apparently it involves a (2,3)sigmatropic migration reaction of intermediate allylselenilic acids leading mostly to allylic hydroxylation products. It has been widely used for the oxidation of cyclohexenyl systems. As mixture 3 was difficult to separate it was directly oxidized with chromium VI oxide in the presence of pyridine, to give a mixture 4 of the expected aldehyde and ketone. The chromium (VI) oxide-pyridine complex is used as an oxidant for the conversion of primary and secondary alcohols to
aldehydes and ketones (Holum, 1961; Ratcliffe and Rodehorst, 1970) at room temperature. On further oxidation of the mixture 4 with sodium chlorite (Burstein et al., 1992; Pellegata et al., 1986) we obtained the acid diacetate HU-444 (compound 5) from the aldehyde which was easy to separate from the unchanged ketone. The overall yield from CBD was 13% (96% purity by HPLC). The conversion of the diacetate to the free diphenol was carried out by sodium borohydride in ethanol to give HU-445 (compound 6) in 90% yield (95% purity by HPLC). All compounds were purified on silica gel chromatography (Scheme 1).

While the overall yield of HU-444 is relatively low, in view of the ready availability of CBD, its synthesis represents a practical route.

**Binding to the cannabinoid receptors.** Neither 5 (HU-444) nor 6 (HU-445) were found to bind to either the CB1 or the CB2 receptor (Ki above 10 µM).

**Pharmacology.** Compound 5 (HU-444) was first investigated for its possible THC-like psychoactivity in the tetrad assay, which measures cannabinoid-induced hypokinesia, hypothermia and antinociception in a tail flick or hot plate test in mice (Mechoulam et al., 2014; Martin et al., 1991; Fride and Mechoulam, 1993). The mouse tetrad serves as a useful *in vivo* screen for psychotropic cannabinoids, which, in contrast to many other types of drugs, displays potency in all four of these bioassays. As no psychoactivity was noted (data not presented), we evaluated HU-444 in a wide range of assays relevant to inflammation.

The *in vitro* anti-inflammatory assays were mostly done with macrophages, while for the *in vivo* studies we used an animal model for both autoimmune hepatitis and rheumatoid
arthritus. The results showed that HU-444 is a potent anti-inflammatory compound, both
in vitro and in vivo.

**Suppression of reactive oxygen species (ROS) production by HU-444.** To study the
effect of HU-444 on the ability of macrophages to produce ROS, RAW 264.7 cells were
stimulated with zymosan together with various doses of HU-444. A 39% and 62%
decrease of ROS generation was observed in the presence of 20 and 40 µg/ml HU-444,
respectively (Fig. 1A).

**Suppression of nitric oxide (NO) production by HU-444.** To study the effect of HU-
444 on macrophage production of NO, RAW 264.7 cells were incubated for 24 h with
LPS (1 µg/ml) and various doses (5-60 µg/ml) of HU-444. NO production was
suppressed by 51% and 64% following incubation of the cells with 40 and 60 µg/ml HU-
444, respectively (Fig. 1B).

**Inhibition of TNF-α production in macrophages and mice.** TNF-α production by TG
macrophages was determined following their incubation with LPS (1 µg/ml) and various
doses of HU-444. Inhibition of up to 69% was noted upon cell incubation with 40 µg/ml
HU-444 for 24 h (Fig. 2A). HU-444 also reduced the TNF-α serum levels by 34-45% in
C57BL/6 mice after i.p. injection of LPS (100 µg/mouse). Already at 2.5 mg/kg HU-444 a
strong inhibition in TNFα serum levels of 34% was observed, with only a slightly higher
inhibition (45%) when increasing the dose to 10 mg/kg, suggesting that a plateau effect
has been reached (Fig. 2B).

**Effect of HU-444 on Con A-induced liver damage.** Intravenous administration of Con
A to mice causes a CD4+ T cell driven, TNFR1-dependent acute hepatitis, leading to
pathological damage of the liver. This is accompanied by elevation of liver enzymes,
interleukin-2 and inflammatory cytokines (Kusters et al., 1997; Ohta and Sitkovsky, 2001). When Con A treated mice were injection i.p. with 5 mg/kg HU-444, a marked reduction in the ALT and AST aminotransferase serum levels was observed (Fig. 3). Alanine aminotransferase (ALT) was reduced by 87% (Fig. 3A) and aspartate aminotransferase (AST) by 85% (Fig. 3B). We observed a bell shaped dose response with both enzymes, with an optimal dose of 5 mg/kg. A bell-shaped dose response is often observed with CBD drugs (Jamontt et al., 2010). The TNF-\(\alpha\) serum levels in Con A-treated C57BL/6 mice were also reduced after treatment with HU-444. Inhibition of 53% in TNF-\(\alpha\) titers was scored following injection of 5 mg/kg HU-444 (Fig. 4).

**Prevention of Con A-induced liver damage by HU-444.** To study the effect of HU-444 on Con A-induced liver damage, histopathological evaluations of the livers were performed (Table 1 and Fig. 5). Con A treatment caused marked liver damage with necrosis and mononuclear cell infiltration (Fig. 5A). Administration of HU-444 attenuated significantly liver damage and reduced mononuclear infiltration, leading to almost normal histology (Fig. 5C-5D). An optimally preserved normal liver histology was observed after treatment with 5 mg/kg HU-444 (Fig. 5C).

**Treatment with HU-444 provides beneficial outcome in an established mouse model of arthritis.** We explored the potential anti-inflammatory and disease modulating properties of HU-444 using our mouse collagen-induced arthritis (CIA) model (Malfait et al., 2000; Sumariwalla et al., 2004). Administration of HU-444 systemically by intraperitoneal or oral route ameliorated clinical signs of arthritis (Fig. 6). Treatments commenced from day 1 of arthritis, which is considered as the first day of the appearance of visible clinical signs of arthritis (redness/swelling) in any of the paws.
The average day of onset of disease is around day 14-28 post immunization. This period can, however, be variable between mice and experiments. As seen from Figure 6, an inverse proportion dose curve was established with the highest dose of 10 mg/kg, daily being clinically non-beneficial, while the two lower doses of 2.5 and 5.0 mg/kg showed beneficial clinical efficacy. Doses below 2.5 mg/kg were inefficient, indicating for the existence of an inverse bell-shaped curve, as is seen with CBD (Malfait et al., 2000) and a synthetic CBD derivative (Sumariwalla et al., 2004).

Clinical signs of arthritis were recorded daily as a composite clinical score of the four limbs which was significantly reduced at 2.5 and 5.0 mg/kg over the entire 10 day period of HU-444 administration (Fig. 6A). In addition, hind paw thickness (mm) readings recorded as paw swelling changes from day 1 of arthritis (Δ\text{mm}) showed significant reduction with 5 mg/kg but not with 2.5 or 10 mg/kg (Fig. 6B). Only, a modest reduction in paw swelling was recorded upon i.p. injection of vehicle from days 5-10 of CIA. More importantly, daily oral administration of HU-444 at 15 mg/kg to arthritic mice for ten days significantly ameliorated the clinical score and paw swelling profiles (Figs. 6C-6D).

HU-444 also provided clinical benefits in terms of conferring protection to joints and prevention of damage caused by pathological arthritic alterations (Fig. 7). Haematoxylin and eosin staining of joint sections of mice treated i.p. with HU-444 (2.5 mg/kg) showed a significant reduction (36.7%) in damaged joints, with a concomitant increase (18.8%) in protected joints, when compared to a vehicle-treated group (Fig. 7A). Similarly, a 45.5% reduction in damaged joints, and 48.9% increase in protected joints was seen upon oral treatment of arthritic mice with HU-444 (15 mg/kg), when compared
to the vehicle treated group (Fig. 7B). Image analysis of representative proximal phalanx joints from both intraperitoneal and oral treated mice showed preserved joint architecture when compared to vehicle treated mice (Figs. 7C-7D). A non-arthritic joint from an immunized mouse, as well as an arthritic joint from an untreated CIA mouse are shown alongside for comparison (Fig. 7E). However, despite the significant joint protection seen upon treatment with HU-444, some residual synovial hypertrophy remained.

**Discussion**

In the present study, we report the synthesis of a novel resorcinol derivative HU-444 (compound 5) that exhibits anti-inflammatory effects both *in vitro* and *in vivo* using mouse models of rheumatoid arthritis (RA) and inflammation-induced liver damage..

Rheumatoid arthritis is an autoimmune, chronic, debilitating musculoskeletal disease (Feldmann et al., 1995). Increasingly, RA is known to be associated with cardiovascular inflammatory diseases leading to enhanced mortality in patients (Full et al., 2009). Key cellular and molecular components associated with the onset and ongoing inflammatory processes in RA have been identified, though yet the specific antigen/environmental agent triggering the onset of the autoimmune pathways remains uncharacterized (Feldmann and Williams, 2010).

We have previously observed that CBD and its synthetic derivate HU-320 have potent anti-inflammatory effects and ameliorate the clinical signs of rheumatoid arthritis in a mouse model (Malfait et al., 2000; Sumariwalla et al., 2004). The anti-inflammatory disease remitting activity of CBD and HU-320, led us to explore the potential of HU-444, a new, chemically related cannabinoid compound.
We decided to synthesize HU-444 to get a compound with enhanced anti-inflammatory activity that can not be converted under acidic conditions to a psycho-active THC-like molecule. The synthesis uses CBD as the starting material. CBD is formed in the plant (or by heating) from its precursor cannabidiolic acid (CBD acid). CBD acid is present in most Cannabis sativa varieties, including hemp, which is used for industrial purposes. In view of the growing interest in CBD (Porter and Jacobson, 2013), several Cannabis sativa varieties have been developed which contain up to 20% CBD (or CBD acid). Hence CBD is potentially an inexpensive natural product. The synthetic pathway from CBD to HU-444 is short, but the yield is relatively low. In view of the simplicity of the reactions used, the yields can presumably be increased, if needed.

While in the past marijuana and hashish, the most widely used preparations of cannabis containing CBD and the psychoactive ∆⁹-tetrahydrocannabinol (THC) in almost equal amounts (about 2-5%), many marijuana varieties today contain high levels of THC (15-20%), presumably due to mostly illegal commercial interests. These varieties contain almost no CBD, a constituent which does not cause the typical marihuana "high" and is hence apparently of minor interest to the illicit growers.

The in vitro assays used in the present study are well established and widely used in anti-inflammatory research. They parallel to a large extent the assays previously used by us for the examination of the anti-inflammatory properties of CBD (Malfait et al., 2000) and HU-320 (Sumariwalla et al., 2004). We observed that HU-444 is a potent anti-inflammatory drug both in vitro and in vivo. As expected, HU-444 did not cause any of the typical THC-like pharmacological effects in mice.
We show that HU-444 suppresses *in vitro* the generation of ROS and nitric oxide (NO) by RAW mouse monocytes induced by zymosan and inhibits TNF-α production by TG macrophages induced by LPS. *In vivo* production of TNF-α elicited by LPS is also prevented by HU-444. These findings clearly show an immunosuppressive function of HU-444 on macrophage function. The most striking effect following administration of HU-444 was the amelioration of the Con A-mediated liver damage (Fig. 5). Administration of 5 mg/kg HU-444 restored to a considerable extent the normal histology of the liver. This is reflected in the reduced serum levels of the liver enzymes ALT and AST following treatment with HU-444. As Con A-induced hepatitis is considered to mimic autoimmune hepatitis in human (Krawitt, 2006), the marked amelioration of hepatitis by HU-444 seen in our study, suggests a promising curative therapeutic role for this cannabinoid compound in human autoimmune hepatitis.

In light of the encouraging anti-inflammatory properties of HU-444 we decided to explore the therapeutic potential of HU-444 in an animal model of arthritis. Mouse collagen-induced arthritis (CIA) remains a well-characterized and widely used model of RA (Williams, 1998; Williams, 2004). The CIA model is routinely used to evaluate the possible therapeutic potential of novel anti-inflammatory/disease ameliorating compounds for use in RA (Williams, 2007). Indeed, anti-TNF-α, IL-Ra and other antibody-based biologics targeting key cellular and molecular targets had proven their merits originally in this model and then in clinical settings of RA (Taylor et al., 2001). However, most of these treatments are delivered by *i.v.* administration and are beyond the reach of many patients due to their cost factor. Thus, the search for easily prepared, small molecular weight compounds, which can be delivered *per os* continues. A hitherto mostly
untapped wealth of novel compounds is natural products from plants, known to possess anti-inflammatory properties. Indeed, *Cannabis sativa* preparations and pure cannabinoids have been used to relieve symptoms of pain associated with several clinical disease conditions and ailments (Marmor, 1998; Zurier, 2003). We show in this paper that our semi-synthetic compound HU-444 based on the naturally occurring CBD has profound anti-inflammatory effects that relieves paw swelling and arthritis symptoms in collagen-induced arthritis, and may thus be a potent oral drug in the therapy of human rheumatoid arthritis.

**Conclusion**

Our results on the anti-inflammatory potential of HU-444 indicate that it lowers the levels of a variety of inflammatory mediators, such as TNF-α, ROS, NO *in vitro*, and ameliorates arthritis in a mouse model at both the macroscopic and pathological levels, when administered either *i.p.* or orally.

We believe that HU-444, a low molecular weight compound, has the potential to be developed as a novel drug for use in inflammatory conditions, particularly in RA.

**Acknowledgments**

We thank the US National Institute on Drug Abuse for a grant to R.M. (DA 9789), the Kessler Foundation of Boston for a donation to R.M. supporting this project and the Israel Ministry of Science for a stipend to C.H. We thank all staff members of the Biological Services Unit at the Kennedy Institute of Rheumatology (KIR) in London for their help provided in the care and maintenance of laboratory mice. Special thanks to Mr. David Essex for his assistance with histology of arthritic paws. Arthritis Research, U.K. provided a core grant facility to KIR which helped facilitate part of this work.
Authorship Contributions

Participated in research design: Haj, Sumariwalla, Mechoulam, Feldmann, Gallily

Conducted experiments: Haj, Sumariwalla, Hanuš, Kogan, Yektin, Gallily

Performed data analysis: Haj, Sumariwalla, Mechoulam, Feldmann, Gallily

Wrote or contributed to the writing of manuscript: Haj, Sumariwalla, Mechoulam, Gallily.

Address Correspondence to: Prof. Raphael Mechoulam, Institute for Drug Research, Hebrew University Medical Faculty, Jerusalem 91120, Israel. E-mail: mechou@cc.huji.ac.il; Prof. Ruth Gallily, Lautenberg Center for Immunology, Hebrew University Medical Faculty, Jerusalem 91120, Israel. E-mail: ruthg@ekmd.huji.ac.il
References


Footnotes

This work was supported by the US National Institute on Drug Abuse for a grant to R.M. [DA 978]; the Kessler Foundation of Boston for a donation to R.M. supporting this project; and the Israel Ministry of Science for a stipend to C.H.
Legends

Fig. 1. Effect of HU-444 on in vitro macrophage function. (A), Generation of ROS (assayed by chemiluminescence) (Ben-Shabat et al., 2006) by RAW 264.7 mouse macrophage cell line, following stimulation with zymosan, in the absence or presence of HU-444. Control Chemiluminescence was 1370 cpm (counts per minute). *Statistically significant difference with a p value ≤ 0.05. (B), Generation of NO (assayed by Griess reagent) (Ben-Shabat et al., 2006) by RAW 264.7 cells, following stimulation by 1 μg/ml LPS for 24 hrs in the presence of HU-444. The level of NO2/NO3 in the control supernatants was 42.5 nmol/ml. *Statistically significant difference with a p value ≤ 0.05.

Fig. 2. Effect of HU-444 on TNF-α production in vitro and in vivo. (A), TNF-α (determined by ELISA) production by TG macrophages following incubation with 1 μg/ml LPS in presence of HU-444 for 24hrs. Control macrophages (10^5), produced 59±3 pg/ml TNF-α, during the 24 hrs. *Statistically significant difference with a p value ≤ 0.05. (B), TNF-α in mice sera 90 min after injections of LPS and HU-444. TNF titer in the control LPS sera was 732 ± 60pg/ml. *Statistically significant difference with a p value ≤ 0.05.

Fig. 3. Levels of liver aminotransferases (ALT, AST) in sera of Con A-treated mice following injection of HU-444. A), ALT, alanine aminotransferase. *Statistically significant difference with a p value ≤ 0.05. (B), AST, aspartate aminotransferase. *Statistically significant difference with a p value ≤ 0.05.
Fig. 4. TNF-α level in serum of mice 20 hrs after intravenous injection of Con A and intraperitoneal injection of HU-444. *Statistically significant difference with a p value $\leq$ 0.05.

Fig. 5. Histopathology of liver, 20 hrs after Con A injections and HU-444 injections. Staining with haematoxylin & eosin. (A), Con A injected mice. Severe damage can be seen; areas of necrosis and infiltration of inflammatory cells. (B), Injection of 2.5 mg/kg HU-444, concomitant with Con A. Normal liver histology was largely preserved, although some infiltration of inflammatory cells was seen. (C), Injection of 5 mg/kg HU-444, concomitant with Con A. Near normal liver structure seen microscopically. (D), Injection of 10 mg/kg HU-444, concomitant with Con A. Although without area of necrosis, changes of normal liver cell morphology can be seen microscopically.

Fig. 6. Amelioration of clinical signs of arthritis in a mouse CIA model following treatment with HU-444. (A), clinical arthritic score (B), paw swelling ($\Delta$ mm) as a change from day 1 of arthritis over a ten day period following daily intraperitoneal treatment with vehicle (●), and HU-444 at 2.5 (■), 5.0 (▲) and 10 (▼) mg/kg doses. (C), clinical arthritic score (D), paw swelling ($\Delta$ mm) as a change from day 1 of arthritis over a ten day period following daily oral treatment with vehicle (●) and HU-444 at 15 mg/kg (■). All treatments were initiated at day 1 of visible clinical signs of arthritis and lasted ten days. Each symbol represents the mean ± s.e.m. of 9 mice. Statistically significant difference **$p$$<0.05$ (1-way ANOVA with Tukey’s post test), ***$p$$<0.0001$ (2-way ANOVA with Bonferroni’s post test).

Fig. 7. Treatment of established CIA with HU-444 protects against pathological joint damage. Percentage of joints scored as either protected (□) or damaged (▨), following
(A), intraperitoneal treatment at different doses of HU-444 (as indicated in figure) or (B), oral treatment with HU-444 at 15 mg/kg. Statistically significant difference \( *p=0.0332, \quad ***p=0.0006 \) by Fisher’s exact test. Representative H&E stained images (captured at 100X) of proximal phalanx joints from (C), intraperitoneal (D), oral treatment with vehicle or HU-444, as indicated (E), non-arthritic and arthritic joint, as indicated from immunized CIA mice from an unrelated experiment are shown for comparison. Figure annotations refer to C (cartilage), B (bone), E (erosions), P (pannus), SY (synovium) and JS (joint space). Scale bars represent 200 µm optical distances.

**TABLE 1**

Concomitant treatment of mice with HU-444 confers protection against Con A induced liver damage.

<table>
<thead>
<tr>
<th>Treatment (HU-444 mg/kg, i.p.)</th>
<th>Pathological Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Con A</td>
<td>3-4</td>
</tr>
<tr>
<td>Con A + 2.5</td>
<td>3</td>
</tr>
<tr>
<td>Con A + 5.0</td>
<td>1.5</td>
</tr>
<tr>
<td>Con A + 10.0</td>
<td>2.5</td>
</tr>
</tbody>
</table>
Scheme 1. Syntheses of HU-444 and HU-445. Reagents and Conditions: (a) Pt(IV) oxide/H₂, EtOAc, 10psi, rt, 2min; (b) pyridine, acetic anhydride, rt, 12h; (c) SeO₂, EtOH, rt, 3h; (d) pyridine; (e) CrO₃, CH₂Cl₂/DMF, rt, 1h; (f) NaClO₂; (g) 2-methyl-2-butene; (h) KH₂PO₄, i-butanol, rt, 1h; (i) NaBH₄, EtOH, reflux, 1h.
Figure 1 Haj et al.

(A) % Inhibition of ROS

(B) % Inhibition of NO
Figure 2 Haj et al.

A

% Inhibition of TNFα by TGMΦ

\[ \text{µg/ml} \]

B

% Inhibition of TNFα in serum

\[ \text{mg/kg} \]
A-Con A control B-HU444 2.5 mg/kg C-5 mg/kg D-10 mg/kg x40

Arrow in A and B shows infiltration of inflammatory cells.

Negligible infiltration in C and D.

Fig. 5.
Fig. 6.
Fig. 7. (A, B)
Fig. 7. (C-E)