3-Hydroxymorphinan, a metabolite of dextromethorphan, protects nigrostriatal pathway against MPTP-elicited damage both in vivo and in vitro

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ABSTRACT We investigated the neuroprotective property of analogs of dextromethorphan (DM) in lipopolysaccharide (LPS) and 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP) models to identify neuroprotective drugs for Parkinson’s disease (PD). In vivo studies showed that daily injections with DM analogs protected dopamine (DA) neurons in substantia nigra pars compacta and restored DA levels in striatum using two different models for PD. Of the five analogs studied, 3-hydroxymorphinan (3-HM), a metabolite of DM, was the most potent, and restored DA neuronal loss and DA depletion up to 90% of the controls. Behavioral studies showed an excellent correlation between potency for preventing toxin-induced decrease in motor activities and neuroprotective effects among the DM analogs studied, of which 3-HM was the most potent in attenuating behavioral damage. In vitro studies revealed two glia-dependent mechanisms for the neuroprotection by 3-HM. First, astroglia mediated the 3-HM-induced neurotrophic effect by increasing the gene expression of neurotrophic factors, which was associated with the increased acetylation of histone H3. Second, microglia participated in 3-HM-mediated neuroprotection by reducing MPTP-elicited reactive microgliosis as evidenced by the decreased production of reactive oxygen species. In summary, we show the potent neuroprotection by 3-HM in LPS and MPTP PD models investigated. With its high efficacy and low toxicity, 3-HM may be a novel therapy for PD.—Zhang, W., Shin, E-J., Wang, T., Lee, P. H., Pang, H., Wie, M-B., Kim, W-K., Kim, S-J., Huang, W-H., Wang, Y., Zhang, W., Hong, J-S., Kim, H-C. 3-Hydroxy morphinan, a metabolite of dextromethorphan, protects nigrostriatal pathway against MPTP-elicited damage both in vivo and in vitro. FASEB J. 20, 2496–2511 (2006)

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Parkinson’s disease (PD) is a neurodegenerative disorder characterized by a chronic and progressive loss of dopamine (DA) neurons in substantia nigra pars compacta (SNpc), leading to movement disorders including dyskinesia, resting tremor, rigidity, and gait disturbance (1). Despite significant advances in understanding the pathological changes in PD, the etiology and the underlying mechanism responsible for the progressive nature of neurodegeneration remain poorly understood. Furthermore, current therapies are limited to relieving PD symptoms and are largely ineffective in halting the process of neurodegeneration.

We have reported that dextromethorphan (DM), a widely used anticough agent, protected DA neurons against inflammagen LPS in midbrain neuron-glia cultures (2) and neurotoxin 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP)-elicited neurotoxicity in vivo (3) via an anti-inflammatory effect by preventing the over-activation of microglia. Furthermore, we recently reported that 3-hydroxymorphinan (3-HM), a metabolite of DM, exerted a more potent neuroprotection than DM against LPS-induced DA neurotoxicity in primary midbrain mixed neuron-glia cultures (4). We attribute the higher potency of 3-HM to its additional neurotrophic effect provided by astroglia.
in addition to the anti-inflammatory activity shared by 3-HM and DM (4).

To continue this research, we investigated the neuroprotective property of a variety of analogs of DM. In this paper, we report that of the five morphinan analogs studied, 3-HM provided the best neuroprotection against LPS- and MPTP-elicited DA neurotoxicity both in vivo and in vitro. Mechanistic studies showed that the neurotrophic effect was due to an increase in the production of several neurotrophic factors from astroglia, which may be associated with the increase in histone acetylation induced by 3-HM as revealed by the increased level of histone 3 protein. Additionally, 3-HM decreased MPTP-induced production of both the extracellular superoxide and intracellular reactive oxygen species (iROS), which may be the basis for the anti-inflammatory mechanism of 3-HM.

MATERIALS AND METHODS

Reagent

3-HM was synthesized as described by Kim et al. (5). All materials related to cell cultures were obtained from Invitrogen (Carlsbad, CA, USA). Reagent from Sigma-Aldrich (St. Louis, MO, USA) included MPTP, MPP+ LPS, cytosine β-d-arabinofuranoside (ara-c), leu-leu methyl ester (LME), superoxide dismutase (SOD), 3,3'-diaminobenzidine, and pentobarbital. Sources for other compounds included: chloral hydrate (Fluka, Buchs, SG, Switzerland), [3H]DA (30 Ci/mmol) (Perkin Elmer Life Sciences, Boston, MA, USA), polyclonal anti-TH antibody (Ab) (a gift from Dr. John Reinhard of GlaxoSmithKline, Research Triangle Park, NC, USA), Vectastain avidin-biotin complex (ABC) kit and biotinylated secondary Ab (Vector Laboratories, Burlingame, CA, USA), fluorescence probe 2',7'-dichloro-hydrofluoreseein (DCFH-DA) (Molecular Probes, Eugene, OR, USA; Calbiochem, San Diego, CA, USA), WST-1 (Dojindo Laboratories, USA), ECL kit (Amersham Pharmacia Biotech., Piscataway, NJ, USA).

Animal treatment

All animals were treated in strict accordance with the NIH Guide for the Care and Use of Laboratory Animals (NIH Guide for the Care and Use of Laboratory Animals). Eight-wk-old male C57BL/6J mice (Bio Genomics Inc., Charles River Technology, Gapyung-Gun, Gyeonggi-Do, South Korea) weighing ~25 ± 3 g were maintained on a 12:12 h light:dark cycle and fed ad libitum. They were adapted for 2 wk to the conditions described above before experimentation. To examine the effect of morphinans on MPTP-induced neurotoxicity, mice received daily MPTP injection [20 mg/kg of MPTP.HCl (16.52 mg/kg as a base), s.c.] for 7 consecutive days. Each morphinan (12 or 24 mg/kg, i.p.) was administered 30 min before every MPTP injection for the last 3 days. Animals were sacrificed 24 h after the last MPTP injection.

For intrastriatal LPS injection, mice were deeply anesthetized using chloral hydrate (300 mg/kg, s.c.), then fixed on the stereotaxic frame (David Kopf Instruments, Tujunga, CA, USA). LPS [(L-7011, Sigma-Aldrich Co, St. Louis, MO, USA) 2 μg in a volume of 2 μl of PBS] was injected bilaterally to 0.7 mm anterior from bregma, 1.0 mm lateral from midline, and 3.4 mm from dura. Each morphinan analog (see structures in Fig. 1) was administered (12 or 24 mg/kg, i.p.) twice, 4 h and 30 min before LPS treatment. Mice were sacrificed 3 wk after LPS injection. Striatal tissues after treatment with MPTP or LPS were rapidly dissected, immediately frozen in liquid nitrogen, then stored at ~80°C until use. The rest of mice were perfused transcardially with 50 ml of PBS, followed by 200 ml of 4% paraformaldehyde in PBS under anaesthesia with pentobarbital (50 mg/kg, i.p.). Perfused brains were post-fixed in 4% paraformaldehyde for 4 h, then immersed in 30% sucrose in PBS until sink.

Locomotor activity

To determine the psychotropic effect of morphinans, mice received a daily injection of each morphinan (20 or 40

**Figure 1.** Structure of morphinan compound DM and its analogs. Structure of morphinan compound DM and its analogs, including 3-hydroxymorphinan (3-HM), 3-allyloxy-17-methylmorphinan (3-AM), 3-cyclopropylmethoxy-17-methylmorphinan (3-CM), 3-methyl-N-methyl-morphinan (DF), and dextrophan (DX), are shown in Fig. 1.
mg/kg, i.p.) for 7 days. Ten minutes after the last injection, locomotor activity was measured for 30 min. After measuring locomotor activity (i.e., 40 min after the last injection), the “absolute turn angular” was analyzed in a 3 min monitoring period using an automated video tracking system (Noldus Information Technology, Wageningen, the Netherlands) to examine locomotor patterns (6). Eight test boxes (40×40×30 cm high) were operated simultaneously by an IBM computer. Animals were studied individually during locomotion in each test box, where they were adapted for 10 min before starting the experiment. The distance traveled in centimeters by the animals in horizontal locomotor activity was analyzed. Data were collected and analyzed between 0900 and 1700 h.

**Conditioned place preference test**

Mice received an i.p. injection of saline just before entering the white or black compartment. Each compound (20 or 40 mg/kg, i.p.) dissolved in saline (0.1 ml/10 g) was administered immediately before the mice were placed in the white compartment. On day 1, the mice were preexposed to the test apparatus for 15 min. The guillotine style doors were raised and mice were allowed to move freely between the two compartments. On day 2, the time each mouse spent in each compartment was recorded for 15 min. On days 3, 5, 7, 9, 11, and 13, the mice were injected with each drug before being confined to the white compartment, the nonpreferred side, for 40 min. On days 4, 6, 8, 10, and 12, the mice were injected with saline before being confined to the black compartment, the preferred side, for 40 min. On day 14, the guillotine doors were raised. The mice were initially placed in the tunnel and the time spent by the mice in the two compartments was recorded for 15 min. The scores were calculated from the differences in the time spent in the white compartment in the testing and pretesting phases (6, 7). Data were analyzed between 0900 and 1700 h.

**Measurement of MPP+ in striatal extract after the last treatment with MPTP**

Striatal tissue samples were homogenized in buffer [0.4 M perchloric containing sodium metabisulphite (0.1%, w/v), EDTA (0.01%, w/v), and cysteine (0.1%, w/v)]. An aliquot (100 μl) of homogenizing buffer was used per milligram of tissue to produce a final tissue concentration of 0.01 g/ml. Samples were centrifuged for 10 min (10,000 rpm; 4°C) and the supernatant was retained for subsequent analysis of MPP+ levels using LC-MS/MS. A stock solution of MPP+ was prepared in homogenization buffer at a concentration of 1.0 mg/ml. This solution was stored at 4°C, all other solutions were prepared by diluting this stock solution with homogenization buffer. Buffer A was composed of 0.1% formic acid in H₂O (v/v). Buffer B consisted of 0.1% formic acid in acetonitrile (v/v). A linear gradient elution profile from 100% A to 100% B over 5 min at a flow rate of 0.5 ml/min was used for the separation, with a 3 min re-equilibration between injections.

The HPLC system was composed of two Jasco model PU-1585 HPLC pumps (Great Dunmow, UK), an HTS PAL autosampler fitted with a twin PAL stackcooler and twin six port injection valves with a 20 μl injection loop (CTC Analytics, Zwingen, Switzerland). The chromatography of MPP+ was carried out using a 100 mm × 2.1 mm i.d., 3 μm, Atlantis HILIC silica column (Waters, Milford, MA, USA), maintained at 40°C. Eluates were detected using a Sciex API 4000 triquadrupole mass spectrometer equipped a Turboion-spray source (TSQ Quantum Ultra, Thermo, USA). The ion spray voltage was set at 5 kV and source temperature at 500°C. The mass spectrometer was operated in the positive ion electrospray mode with the following parameters: declustering potential 60 V, focusing potential 10 V, entrance potential ~10 V, collision cell exit potential 11 V. Nitrogen was used for both the curtain and collision gas with an ion energy of 37 eV. MPP+ was monitored using a molecular reaction monitoring mode with the transition m/z 170.4 → 127.9 (8).

**Immunohistochemistry and cell counting in the SNpc of C57 mice**

The brains were cut on a horizontal sliding microtome into 35 μm transverse free-floating sections. Four to 5 brain sections of SNpc were collected at intervals of 140 μm. Immunohistochemistry was performed as described previously (9). Briefly, prior to incubation with the primary antibodies, sections were preincubated with 0.3% hydrogen peroxide in PBS for 30 min, then in PBS containing 0.4% Triton X-100 for 20 min and 1% normal serum for 20 min. After 48 h incubation with the anti-TH (1:5000) at 4°C, sections were incubated with the biotinylated anti-mouse IgG (1:1000) for 1 h and immersed in Vectastain ABC reagents for 1 h. 3,3’-Diaminobenzidine was used as a chromogen. The bound complex was visualized using 3,3’-diaminobenzidine. Digital images of TH immunoreactive (TH-ir) neurons in SNpc were acquired at 40× magnifications on an Olympus microscope (Olympus®, Tokyo, Japan) using an attached Polaroid digital microscope camera (Polaroid®, Cambridge, MA, USA) and IBM PC. A region of interest (ROI) was created by outlining SNpc using Optimas® version 6.51 (Media Cybernetics, Inc., Silver Spring, MD, USA). Cell counting was performed blindly by two investigators and results were obtained from the average. A mean value for the number of SNpc TH-ir neurons was then deduced by averaging the counts of 4–5 sections for each animal. Results were indicated as % of saline (+saline)-treated group.

**Neurochemical analysis of the contents of striatal neurotransmitters**

The levels of DA and its metabolites 3,4-dihydroxyphenylacetic acid (DOPAC) and homovanillic acid (HVA) were determined by HPLC coupled with electrochemical detection as described (9, 10). Briefly, striatal tissues were sonicated in 0.2 M perchloric acid (PCA) (20% W/V) containing the internal standard 3,4-dihydroxybenzylamine (10 mg wet tissue/ml). The homogenate was centrifuged and a 20 μl aliquot of the supernatant was injected into the HPLC equipped with a 3 μm C₁₈ column. The mobile phase was comprised of 26 ml of acetonitrile, 21 ml of tetrahydrofuran, and 960 ml of 0.15 M monochloroacetic acid (pH 3.0) containing 50 mg/L of EDTA and 200 mg/L of sodium octyl sulfate. The amount of DA, DOPAC, and HVA were determined by comparison of peak height ratio of tissue sample with standards and expressed in nanograms per gram of wet weight of tissue.

**Preparation of cell cultures**

Fisher 344 rat and C57BL/6J mice were purchased from Jackson Laboratories (Bar Harbor, ME, USA). Breeding schedules for the rat and mice were designed to achieve accurate timed-pregnancy of 14 ± 0.5 and 13 ± 0.5 days, respectively. Animals were maintained and bred in a strict pathogen free environment. Housing, breeding, and experimental use of the animals were performed in strict accordance with the National Institutes of Heath guidelines. Various primary cultures used in this study were described below.
Primary midbrain mixed neuron-glia cultures

Primary midbrain neuron-glia cultures were prepared from the brains of embryonic day 14 ± 0.5 days of Fisher 344 rats and 15 ± 0.5 days of C57BL/6J mice as described previously (4). Briefly, the ventral midbrain tissues were removed and dissociated by a mild mechanical trituration. Rat (mouse) cells were seeded at 5 × 10^5/well (1 × 10^5/well) in 24-well (96-well) culture plates precoated with poly-l-lysine (20 μg/ml) and maintained at 37°C in a humidified atmosphere of 5% CO_2 and 95% air in 0.5 ml (200 μl) maintenance medium that consisted of minimum essential medium (MEM) containing 10% heat-inactivated FBS, 10% heat-inactivated horse serum (HS), 1 g/L glucose (Glc), 2 mM l-glutamine, 1 mM sodium pyruvate, 100 μM nonessential amino acids, 50 U/ml penicillin, and 50 μg/ml streptomycin. Three days after the initial seeding, 0.5 ml of fresh maintenance medium was added to each well of 24-well culture plates. Seven (2)-day-old 24 (96)-well cultures were used for treatment. The composition of the cultures at the time of treatment was ~48% astroglia, 11% microglia, and 40% neurons in which ~1% were TH-ir neurons.

Primary midbrain neuron-enriched cultures

To obtain rat primary neuron-enriched cultures, 24 h after seeding the cells, Ara-c was added to a final concentration of 7.5 μM to suppress glial proliferation (11). Three days later cultures were changed back to maintenance medium and were used for treatment 7 days after initial seeding. This method typically can enrich for neurons to >95% purity.

Primary microglia-enriched cultures

Primary microglia-enriched cultures were prepared from the whole brains of 1-day-old Fisher 344 rat pups following our described protocol (12). Briefly, brain tissues were triturated after removing the meninges and blood vessels. Cells (5 × 10^6) were seeded in a 150 cm^2 cultures flask. After a confluent monolayer of glia cells had been obtained, microglia were shaken off, collected (a purity of >98%), and seeded onto 96-well plates for extracellular superoxide and iROS assay.

Primary midbrain astroglia-enriched cultures

After removing loosely attached microglia by shaking three times, the remaining astroglia were detached with trypsin-EDTA and seeded in the same culture medium as that used for microglia cultures. After three consecutive passages, a purity of >98% of astroglia preparation can be achieved and used to measure 3-HM-induced gene expression of neurotrophic factors using reverse transcription (RT) and real-time quantitative PCR.

Primary midbrain reconstituted neuron-microglia cultures

To explore the role of microglia on 3-HM-exerted DA neuroprotection in the primary midbrain mixed neuron-glia culture system, the reconstituted cultures were established by adding 10% (5 × 10^5/well) of microglia back to neuron-enriched cultures as described previously (13). In normal neuron-glia cultures, the composition of microglia is ~10% of the total cells.

Primary midbrain neuron-astroglia cocultures

Rat primary midbrain neuron-astroglia cocultures were obtained by suppressing microglial proliferation with 1.5 mM LME 24 h after seeding the cells, as described previously (13). Three days later cultures were changed back to maintenance medium and used for treatment 7 days after initial seeding. The composition of the cells at the time of treatment was ~54% astrocytes, 1% microglia, and 45% neurons.

DA uptake assay

[^H]DA uptake assay was performed as described previously (12). Cultures were incubated for 20 min at 37°C with 1 μM[^H]DA in Krebs-Ringer buffer (16 mM sodium phosphate, 119 mM NaCl, 4.7 mM KCl, 1.8 mM CaCl_2, 1.2 mM MgSO_4, 1.3 mM EDTA, and 5.6 mM Glc; pH 7.4). After washing three times with ice-cold Krebs-Ringer buffer, cells were collected in NaOH (1N). Radioactivity was determined by liquid scintillation counting. Nonspecific DA uptake observed in the presence of mazindol (10 μM) was subtracted.

Immunocytochemical staining

DA neurons were immunostained with anti-TH Ab. Briefly, 3.7% formaldehyde-fixed cultures were treated with 1% hydrogen peroxide (20 min), followed by sequential incubation with blocking solution (30 min), primary Ab (overnight, 4°C), biotinylated secondary Ab (2 h), and ABC reagents (40 min). Color was developed with 3, 3′-diaminobenzidine. For morphological analysis, images were recorded with an inverted microscope (Nikon, Tokyo, Japan) connected to a charged-coupled device camera (DAGE-MTI of MC, Inc., Michigan City, IN, USA) operated with the MetaMorph software (Universal Imaging Corporation, Downingtown, PA, USA). For visual counting DA neurons, nine representative areas per well of the 24-well plate were counted under the microscope at 100× magnification. To measure the average DA neuronal dendrite, 50 representative DA neurons in each well and three wells for each treatment condition were selected.

Expressions of neurotrophic factors from astroglia using RT and real-time PCR

Astroglia-enriched cultures treated with 3-HM (5 μM) for 3, 6, 12, and 24 h were detached with trypsin-EDTA. Total RNA was isolated with Trizol reagent, followed by purification. Forward and reverse primers for selected neurotrophic factors were designed using ABI Primer Express software (Applied Biosystems, Foster City, CA, USA). The sequences of the primers were β-actin: TCCTCTGAGCCGCAAGTACTCT (F), GCTCAGTAAACGTCGGCTGAA (R); ADNF: TGCA-GTCTGCTAATGCTCCTTCT (F), ACTCCTGATGCTGTTGA-CGTA (R); GDNF: GAGAGAGGAACCGGCAAGCT (F), GTTAGACGGCAACCCGCAACCT (R); NT3: CACCAACGCC-GAAAAGCTAT (F), CACACGGTCTCAGTGTGAC (R); h-bFGF: GAACCGTCGAGTCTGAAT (R), GTCTGGT-TACGTGCCCATACC (R); epidermal growth factor (EGF): GCCACCTGGTCCGAAACA (F), TACATGCACGGCACCAT (R); TGF-β 1: GCCTGAGTGCGCT GCTTCTTGA (F), GAAGGGAAAAGCCCCTGTATTCC (R). Insulin-like growth factor (IGF)-1: ACAGGCTATGCTGCAAGATT (F), GATCA-CAGCTCGAGGAAAGCAA (R); TGF-α: CCAAGGGTCGAGGAAT (R), TGGTGGCGGCTGTAGAC (R). RT and real-time PCR were utilized for analysis of the gene expression of neurotrophic factors using forward and reverse primers for the selected genes (14). Total RNA was reversely transcription
with MuLV reverse transcriptase and oligo-dT primers, and subjected to real-time PCR analysis using SYBR green PCR master mix (Applied Biosystems, Cheshire, UK). The relative differences in expression between groups were determined using cycle time (Ct) values as follows: the Ct values for the genes of interest were first normalized with β-actin of the same sample, then the relative differences between control and treatment groups were calculated and expressed as relative increases, setting control as 100%. Assuming that the Ct value is reflective of the initial starting copy and that there is 100% efficiency, a difference of one cycle is equivalent to a 2-fold difference in starting copy. Standard curve analysis was performed and used for the calculation.

Western blot analysis of acetylated histone H3 by 3-HM

Astroglia-enriched cultures treated with 3-HM (5 μM) for 1, 3, 6, 12, and 24 h were detached by scraping in sampling buffer containing 10 mM HEPES (pH 7.9), 1.5 mM MgCl₂, 10 mM KCl, 1.5 mM phenylmethylsulfonyl fluoride, and 0.5 mM dithiothreitol. Histones were isolated by means of acid extraction. Isolated histones were adjusted to equal protein concentrations and separated by 4–12% polyacrylamide-0.1% SDS gels. The samples were electrotransfered onto PVDF membranes. Blots were incubated with primary Ab against acetylated histone H3, then probed with a horseradish peroxidase-conjugated secondary Ab. Detection was performed by using ECL via ECL kit.

Assay of extracellular superoxide production in mouse primary mixed midbrain neuron-glia cultures

The production of extracellular superoxide was determined by measuring the SOD-inhibited reduction of the tetrazolium salt WST-1, originally reported by Tan and Berridge (15) and adapted for neural cultures by Liu et al. (12, 17). To measure the reactive microgliosis-related release of superoxide, neuron-glia cultures in 96-well culture plates were pretreated for 30 min with vehicle alone or 0.5–1 μM 3-HM prior to treatment with MPTP (0.5 μM) or MPP⁺ (0.25 μM) in the 100 μl treatment medium/well (phenol red-free MEM, containing 2% FBS and 2% HS) following the described protocol (3). Two and 5 days after MPTP or MPP⁺ treatment, the desired concentrations of 3-HM were added back again to the corresponding 3-HM-treated cultures. On days 4 and 7, 75 μl of WST-1 (final concentration: 1 mM) in treatment medium without and with SOD (final concentration: 50 U/ml) was added to each well. Absorbance at 450 nm was read with a SpectraMax Plus microplate spectrophotometer (Molecular Devices, Palo Alto, CA, USA). The difference in absorbance observed in the absence and presence of SOD was considered to be the amount of superoxide produced. Results are expressed as a percentage of vehicle-treated control cultures.

Determination of the levels of iROS in C57BL/6J mouse primary midbrain mixed neuron-glia cultures

The level of iROS in C57BL/6J mouse-derived primary midbrain mixed neuron-glia cultures was measured using a fluorescent probe, DCFHDA, as described (18), with modifications (19). The method of treatment was the same as that for superoxide assay described above. Briefly, 4 and 7 days after treatment, cultures were treated with 10 μM DCFHDA diluted in phenol red-free HBSS containing 2% FBS and 2% HS. After an additional 2 h of incubation at 37°C, fluorescence intensity was measured at 485 nm for excitation and 530 nm for emission using a SpectraMax Gemini XS fluorescence microplate reader (Molecular Devices).

Statistical analysis

Statistical significance of the differences was determined by ANOVA, followed by Bonferroni’s t test using the StatView program (Abacus Concepts, Inc., Berkeley, CA, USA) except for Figs. 2, 3, 9 and Table 1 and Table 2, in which 1-way ANOVA with Fischer’s PLSD test using Statcel® software (OMS, Saitama, Japan) was used. A value of P < 0.05 was considered statistically significant.

RESULTS

3-HM protected against DA neurodegeneration induced by MPTP and LPS in vivo

Two in vivo rodent PD models were used to determine the neuroprotective effect of 3-HM and related morphi-

| Compound                  | DA       | DOPAC    | HVA      |
|---------------------------|----------|----------|----------|
| Saline + saline           | 1041 ± 106 | 145 ± 13 | 118 ± 11 |
| Saline + LPS              | 445 ± 59*  | 54 ± 9*  | 61 ± 8*  |
| DM (12 mg/kg) + LPS       | 626 ± 72  | 65 ± 8   | 79 ± 8   |
| DM (24 mg/kg) + LPS       | 889 ± 74*** | 90 ± 14** | 93 ± 13** |
| 3-HM (12 mg/kg) + LPS     | 689 ± 73  | 80 ± 10  | 80 ± 8   |
| 3-HM (24 mg/kg) + LPS     | 935 ± 85*** | 106 ± 11*** | 99 ± 12** |
| 3-AM (12 mg/kg) + LPS     | 466 ± 58  | 64 ± 8   | 73 ± 8   |
| 3-AM (24 mg/kg) + LPS     | 485 ± 67  | 64 ± 8   | 73 ± 10  |
| 3-CM (12 mg/kg) + LPS     | 498 ± 82  | 69 ± 8   | 79 ± 8   |
| 3-CM (24 mg/kg) + LPS     | 687 ± 53*  | 82 ± 6*  | 87 ± 5*  |
| DF (12 mg/kg) + LPS       | 555 ± 86  | 56 ± 9   | 60 ± 7   |
| DF (24 mg/kg) + LPS       | 552 ± 77  | 55 ± 8   | 63 ± 8   |

*LPS (2 μg in a volume of 2 μl of PBS) was injected into both sides of the striatum of C57BL/6J mice. Each morphinan was administered twice (4 h and 30 min) before intrastriatal injection with LPS. Mice were sacrificed 3 wk after last LPS injection. Each value is the mean ± s.e. from 12 animals. *P < 0.01 vs. saline + saline; **P < 0.05, ***P < 0.02, and ****P < 0.01 vs. saline + LPS, respectively.
nan analogs on DA neurons. To extend our previous in vitro finding that 3-HM protected DA neurons against LPS-induced neurotoxicity (4), a bilateral intrastriatal LPS infusion of the LPS (2 µg in a volume of 2 µl of PBS) in vivo model was first used. LPS infusion caused a significant decrease in the number of TH-ir neurons (26% of control) in SNpc of the injected group compared with the vehicle-injected controls (Fig. 2A).

There was a corresponding reduction in the content of DA (43% of control) and its metabolites, DOPAC (37% of control) and HVA (52% of control), in LPS-injected striatum (Table 1). Daily injections with morphinans resulted in various degrees of protection on DA neurons in SNpc and recovery of DA levels in striatum (Fig. 2A, B and Table 1). Of the five analogs tested, 3-HM was the most potent; it protected DA neurons by up to 75% that of the control (Fig. 2A). LPS-induced reduction in striatal DA levels was dose-dependently protected by 3-HM to 66% in a low dose and 78% in a high dose (Table 2). To determine whether morphinans affect the entry of MPTP to the brain, we had measured the brain MPP⁺ concentrations at different time points after the last injection of MPTP in both control and 3-HM treated mice. The result showed no significant difference of MPP⁺ levels in either group 1, 2, 3, 4, or 6 h after the last injection of MPTP (Fig. 3C).

Locomotor activity and its tracing pattern in LPS- and MPTP-treated animals were measured. Injections of both toxins caused significant hypolocomotor activities (Fig. 2B, Fig. 3B), which were partially recovered with the injections of morphinans. Corresponding to the neuroprotective studies shown above, of the different analogs tested, 3-HM showed the most potent effect in recovering these hypolocomotor activities.

### Table 2. 3-HM is the most potent morphinan in attenuation of the depletion of DA and its metabolites DOPAC and HVA in striatum induced by MPTP in vivo

| Compound | DA (mg/100 mg wet tissue) | DOPAC (mg/100 mg wet tissue) | HVA (mg/100 mg wet tissue) |
|----------|--------------------------|-----------------------------|---------------------------|
| Saline + saline | 1056 ± 96* | 146 ± 15 | 118 ± 13 |
| Saline + LPS | 403 ± 60* | 49 ± 8* | 59 ± 11* |
| DM (12 mg/kg) + LPS | 498 ± 69 | 66 ± 9 | 80 ± 8 |
| DM (24 mg/kg) + LPS | 658 ± 10* (*** | 82 ± 12*** | 92 ± 11*** |
| 3-HM (12 mg/kg) + LPS | 571 ± 82 | 67 ± 9 | 81 ± 8 |
| 3-HM (24 mg/kg) + LPS | 823 ± 57*** | 95 ± 10*** | 101 ± 10*** |
| 3-AM (12 mg/kg) + LPS | 466 ± 70 | 52 ± 7 | 68 ± 8 |
| 3-AM (24 mg/kg) + LPS | 550 ± 80 | 67 ± 8 | 75 ± 9 |
| 3-CM (12 mg/kg) + LPS | 492 ± 65 | 57 ± 6 | 67 ± 9 |
| 3-CM (24 mg/kg) + LPS | 619 ± 47* | 76 ± 7* | 89 ± 7* |
| DF (12 mg/kg) + LPS | 408 ± 46 | 49 ± 7 | 61 ± 8 |
| DF (24 mg/kg) + LPS | 385 ± 65 | 53 ± 6 | 64 ± 7 |

*C57BL/6J mice received daily MPTP injections (20 mg/kg, s.c.) for 7 consecutive days. Each morphinans was administered 30 min before every injection of MPTP for the last 3 days. Animals were sacrificed 24 h after the last MPTP injection. Each value is the mean ± SE from 12 animals. *P < 0.01 vs. saline + saline; **P < 0.05, ***P < 0.02, and ****P < 0.01 vs. saline + MPTP, respectively.

The second in vivo model used involved daily injections of MPTP (20 mg/kg, s.c.), which induced a dramatic decrease of TH-ir neurons (27% of control) in SNpc compared with vehicle-injected controls (Fig. 3A). There was a corresponding decrease in the content of DA (38% of control) and its metabolites, DOPAC (34% of control) and HVA (50% of control), in MPTP-injected striatum (Table 2). Similar to the results in the LPS model, daily injections with morphinans for 3 days resulted in different degrees of protection on DA neurons in SNpc and on DA levels in striatum (Fig. 3A, B and Table 2). Among the different analogs investigated, 3-HM was also the most potent compound protecting against MPTP-induced loss of DA neurons of up to 76% of control (Fig 3A). MPTP-induced decrease in striatal DA levels was dose-dependently protected by 3-HM to 54% in a low dose and 78% in a high dose (Table 2). To determine whether morphinans affect the entry of MPTP to the brain, we had measured the brain MPP⁺ concentrations at different time points after the last injection of MPTP in both control and 3-HM treated mice. The result showed no significant difference of MPP⁺ levels in either group 1, 2, 3, 4, or 6 h after the last injection of MPTP (Fig. 3C).

Locomotor activity and its tracing pattern in LPS- and MPTP-treated animals were measured. Injections of both toxins caused significant hypolocomotor activities (Fig. 2B, Fig. 3B), which were partially recovered with the injections of morphinans. Corresponding to the neuroprotective studies shown above, of the different analogs tested, 3-HM showed the most potent effect in recovering these hypolocomotor activities.

### 3-HM protected against MPTP/MPP⁺-elicited DA neurodegeneration in vitro

Because of its high potency, 3-HM was used as a prototype to further delineate the mechanism of neuroprotection by using rat primary midbrain mixed neuron-gliala cultures. 3-HM (1–5 µM) or vehicle was added to neuron-gliala cultures 30 min prior to the application of MPTP (1 µM)/MPP⁺ (0.5 µM). Seven days later the damage of DA neurons was determined by [3H]DA uptake assay and morphological evaluation, including TH-ir neuron counts and dendrite measurements. As shown in Fig. 4A, MPTP/MPP⁺ reduced DA uptake capacity by 25 to 40%, respectively, which was completely prevented by 1–5 µM (94–129% of control in MPTP treatment and 87–118% of control in MPP⁺ treatment, respectively). 3-HM alone increased DA uptake capacity of up to 185–226% that of control.
Cell count analysis of DA neuronal numbers in the cultures revealed that MPTP/MPP\(^+\) induced loss of DA neurons by nearly 37% and 43%, respectively, compared with vehicle-treated control cultures (Fig. 4B). MPTP/MPP\(^+\)-induced shortening of neurite length was parallel to that of neuronal number (Fig. 4C). Furthermore, the degree of loss of neuronal number and neurite length was comparable to that of DA uptake (Fig. 4A). Pretreatment with 3-HM (5 μM) significantly attenuated MPTP/MPP\(^+\)-induced loss of DA neurons (127% and 138% of control) and neurites (135% and 145% of control) (Fig. 4B). The number of DA neuron and the length of neurites in the cultures treated with 3-HM (5 μM) alone were 216% and 233% vs. control, respectively.

Morphologically, in addition to the reduction in the abundance of DA neurons, the neurites of the remaining DA neurons in MPTP/MPP\(^+\)-treated cultures were significantly damaged compared with vehicle-treated control cultures, which were characterized by shorter, thinner, and broken neurites and, in the worst circumstance, even disappeared. In the MPTP/MPP\(^+\)-treated cultures pretreated with 3-HM, DA neurons were significantly more numerous, and neurites were less damaged than in cultures treated with MPTP/MPP\(^+\) alone (Fig 4C). Cultures treated with 3-HM alone had much more abundant neurites and healthier cell bodies compared with vehicle-treated control cultures (Fig. 4C).

**Neuroprotective effect of 3-HM against MPP\(^+\)-induced DA neurotoxicity was glia dependent**

We previously reported the neurotrophic activity of 3-HM in rat primary midbrain mixed neuron-glia cultures (4), but the mechanism of how 3-HM exerted a neurotrophic effect was not understood. To determine whether the neurotrophic effect of 3-HM contributed to the neuroprotection on DA neurons from MPP\(^+\)-induced damage, the following experiments were performed. In neuron-enriched cultures, MPP\(^+\)-(0.5 μM)-induced direct DA neurotoxicity was revealed by a decrease in DA uptake (59% of control), which was...
similar to that in neuron-glia cultures (50% of control). However, 3-HM protected DA neurons in neuron-glia cultures (101% of control) but not in neuron-enriched cultures (71% of control), indicating that neuroprotection of 3-HM was mediated through glia cells. Based on the finding that 3-HM itself enhanced the capacity of DA uptake in neuron-glia cultures (210% of control) but not in neuron-enriched cultures (113% of control), we conclude there was a close correlation between the neurotrophic action and neuroprotection, and that glial cells were the major players in the neuroprotection afforded by 3-HM (Fig. 5).

Neurotrophic factors released from astroglia contributed to the neuroprotective effect of 3-HM against MPP⁺-induced DA neurotoxicity

In co-cultures of neuron-astroglia (adding 50% of astroglia back to neuron-enriched cultures) and neuron-microglia (adding 10% of microglia back to neuron-enriched cultures), 3-HM (5 μM) alone increased [³H]DA uptake in the cultures with astroglia (199% of control) but not microglia (105% of control), indicating that astroglia, not microglia, were the mediators of the neurotrophic action by 3-HM (Fig. 5). These results
were consistent with the findings that pronounced protection afforded by 3-HM against MPP\(^+\)-induced DA neurotoxicity was observed in neuron-astroglia cocultures (Fig. 5).

Based on this information, we hypothesized that release of neurotrophic factors from astroglia by 3-HM was responsible for the robust neurotrophic effects. Further study using RT and real-time PCR revealed increases in the gene expressions of an array of neurotrophic factors, including endothelial growth factor (epidermal growth factor), neurotrophin 3 (NT3), glia cell line-derived neurotrophic factor (GDNF), astroglia-derived neurotrophic factor (ADNF), transforming growth factor \(\beta_1\) (TGF-\(\beta_1\)) and transforming growth factor \(\alpha\) (TGF-\(\alpha\)), from astroglia treated with 3-HM (5 \(\mu\)M) (Table 3). The peak time of gene expression at the mRNA level for most of the factors was between 3 and 6 h after 3-HM administration. Among the factors examined, the expression of EGF appeared first, which peaked at 3 h (196% of control), followed by GDNF, NT3, TGF-\(\beta_1\), and TGF-\(\alpha\) (167%, 164%, 151%, and 144% of control, respectively); at 6 h after treatment, neurons and measuring the length of neurites. Representative pictures from 3 experiments were showed (C). Results are expressed as a percentage of vehicle-treated control cultures and are the mean ± SE from 3 independent experiments in triplicate. *\(P < 0.05\) vs. vehicle-treated control cultures; **\(P < 0.05\) vs. MPP\(^+\)-treated cultures.

Figure 5. Astroglia is the source of neurotrophic and related neuroprotective effect of 3-HM against MPP\(^+\)-induced DA neurodegeneration. Rat primary midbrain mixed neuron-glia cultures were seeded in a 24-well culture plate at 5 \(\times\) 10\(^5\)/well; primary neuron-enriched cultures were obtained by adding Ara-c to neuron-glia cultures at a final concentration of 7.5 \(\mu\)M to suppress glial proliferation 24 h after seeding the cells; Primary neuron-astroglia cocultures were obtained by suppressing microglial proliferation with 1.5 mM LME 24 h after seeding the cells. Reconstituted neuron-microglia cultures were established by adding 10% (5\(\times\)10\(^4\)/well) of microglia back to neuron-enriched cultures. All above-mentioned cultures were pretreated with 3-HM (5 \(\mu\)M) for 30 min before the addition of MPP\(^+\) (0.5 \(\mu\)M); DA neurotoxicity was quantified by \[^3H\]DA uptake assay 7–9 days after treatment with MPP\(^+\). Results are expressed as a percentage of vehicle-treated control cultures and are the mean ± SE from 4–5 independent experiments in triplicate. *\(P < 0.05\) vs. vehicle-treated control cultures; **\(P < 0.05\) vs. MPP\(^+\)-treated cultures.
the factors with increased gene expression were ADNF, TGF-β1, GDNF, and NT3 (159%, 149%, 146%, and 133% of control, respectively); TGF-β1 was the one that still maintained the level to some degree (131% of control) 12 h after the treatment. The protein levels of most of these neurotrophic factors were below the detecting limit of the assay.

In another study, Western blot analysis (Fig. 6A, B) indicated that 3-HM (5 μg/ml) enhanced the expression of acetylated histone H3 in astroglia by up to 2.7-fold as soon as 1 h after treatment and maintained the level at 2.6-, 2.5-, 2-, and 2.5-fold of control after 3, 6, 12, and 24 h treatment, respectively, whereas valproic acid (VPA) served as a positive control (Chen, P. et al., unpublished results). The observation that the level of total histone H3 showed no change (Fig. 6A, B) indicated that the increased acetylated histone H3 was not due to the alteration of total histone H3.

Prevention of MPTP/MPP⁺-induced microgliosis by 3-HM was through inhibiting ROS production

In neuron-glia cultures, the neuroprotection afforded by 3-HM was increased by 51% increase in DA uptake (Fig. 5), which may due mainly to the neurotrophic action from astroglia indicated by the 35% of increase in DA uptake in neuron-astroglia co-cultures (Fig. 5). Although we did not see the neurotrophic effect in neuron-microglia cocultures because 3-HM alone did not increase the capacity of DA uptake, significant enhancement of DA uptake (>20%) was still found in the cultures treated with MPP⁺ and 3-HM, indicating that microglia was another target in 3-HM-provided neuroprotection against MPP⁺-elicited damage.

MPTP/MPP⁺ have been shown to induce a rapid reactive microgliosis after causing the direct DA neurodegeneration in mice (20, 21). Previous work from our laboratory indicated that MPTP/MPP⁺-induced reactive microgliosis was closely associated with an increased production of oxygen free radicals (20, 22), including extracellular superoxide and iROS from microglia. Therefore, we determined whether 3-HM was able to reduce MPTP/MPP⁺-induced ROS generation.

Production of extracellular superoxide was determined in mouse primary midbrain mixed neuron-glia cultures pretreated for 30 min with 3-HM (0.25–1 μM) prior to the addition of MPTP (0.5 μM)/MPP⁺ (0.25 μM). Fresh 3-HM was added back to the cultures at days 2 and 5 after MPTP/MPP⁺ treatment for replenishment. On days 4 and 7, release of extracellular superoxide from activated microglia was measured using WST-1 as described in Materials and Methods. As shown in Fig. 7A, B, treatment of cultures with 0.5 and 1 μM 3-HM dose-dependently inhibited MPTP (0.5 μM)/MPP⁺ (0.25 μM)-induced superoxide production at the desired time points. 3-HM alone showed no effect on superoxide production.

Next, the effect of 3-HM on MPTP/MPP⁺-induced iROS production was studied. Primary midbrain mixed neuron-glia cultures were treated in the same manner as mentioned above and iROS were determined at days 4 and 7 after MPTP (0.5 μM)/MPP⁺ (0.25 μM) treatment. As shown in Fig. 8A, B, on day 4, 3-HM (0.25–0.5 μM) dose-dependently inhibited the levels of iROS induced by MPTP (0.5 μM). MPP⁺ (0.25 μM)-induced iROS production was also decreased by 0.5–1 μM 3-HM on day 7 in a dose-dependent manner. Hence, these results demonstrated that 3-HM was capable of reducing MPTP/MPP⁺-elicited reactive microgliosis-induced iROS production.

Taken together, these results demonstrated that 3-HM effectively inhibited MPTP/MPP⁺-induced extracellular superoxide and iROS production, and suggested that the inhibition of reactive microgliosis through decreasing ROS productions was another important mechanism associated with the neuroprotective effect of 3-HM in addition to the neurotrophic action.

3-HM did not produce psychotropic side effects

Saline-treated animals exhibited basal locomotor activity and circling activity (Fig. 9A, B) as we demonstrated previously (6, 7). The repeated treatment of DM or its major metabolite, phencyclidine (PCP)-like dextrophan (DX), significantly increased locomotor activity and circling behavior, although these behavioral re-

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**TABLE 3.** 3-HM increases gene expression of several neurotrophic factors from astroglia using RT and real-time PCR analysis

| Neurotrophic factors | 3 h     | 6 h     | 12 h    | 24 h    |
|----------------------|---------|---------|---------|---------|
| EGF                  | 196 ± 34** | 127 ± 13 | 123 ± 15 | 133 ± 3 |
| GDNF                 | 167 ± 13*** | 146 ± 9** | 123 ± 25 | 139 ± 10 |
| NT3                  | 164 ± 10*** | 133 ± 8** | 103 ± 14 | 101 ± 4  |
| TGF-β1               | 151 ± 13** | 149 ± 9** | 131 ± 11* | 125 ± 8  |
| TGF-α                | 144 ± 12** | 88 ± 6   | 115 ± 14 | 76 ± 10  |
| ADNF                 | 124 ± 13  | 159 ± 17** | 109 ± 10 | 125 ± 9  |

*Primary mixed glial cells were prepared from the whole brains of 1-day-old Fisher 344 rat pups. Cells (5 × 10⁷) were seeded in a 150 cm² cultures flask. After a confluent monolayer of glia cells had been obtained, microglia were shaken off. After three consecutive passages, a purity of >98% of astroglia was achieved and treated with 3-HM (5 μM) for 1, 3, 6, 12, and 24 h. Astroglia were then detached with trypsin-EDTA and total RNA was isolated with Trizol reagent, followed by purification. RT and real-time PCR were utilized to detect gene expression of a series of neurotrophic factors using forward and reverse primers for the selected genes. Results are expressed as a percentage of vehicle-treated control cultures and are the mean ± sr from 3 independent experiments in triplicate. *P < 0.05, **P < 0.01, and ***P < 0.005 vs. vehicle-treated control cultures, respectively.
responses were less than PCP (Fig. 9A, B). 3-HM was the one that had the least behavioral responses mentioned above compared with DM and DX (Fig. 9A, B). For conditioned place preference (CPP), DM, DX, and PCP produced significant CPP whereas no CPP was observed in saline-treated animals. 3-HM did not show selective CPP effects that seen in DM- or DX-treated animals, which was comparable to other morphinans, such as 3-AM, 3-CM, and DF. Taken together, behavioral study indicated that 3-HM exerted negligible psychotropic effect compared with parent compound DM or its major metabolite DX.

DISCUSSION

In this study we report that of the five analogs of DM investigated, 3-HM provided the greatest degree of neuroprotection on DA neurons in the nigro-striatal pathway against LPS- and MPTP-elicited damage both in vivo and in vitro. The potent neuroprotective effect of 3-HM was attributed to its neurotrophic factor-releasing activity from astroglia and the inhibition of microgliosis. Note that 3-HM, a natural endogenous metabolite of DM, exerted a more potent neuroprotection than its parent compound.

In the in vivo study, 3-HM almost completely (up to 90% of the control in a high concentration of 24 mg/kg, i.p.) prevented DA neuronal loss in SNpc after MPTP or LPS treatment (Fig. 2A, B, Fig. 3A, B) and protected the DA depletion in striatum (Tables 1 and 2). The degree of neuroprotection by 3-HM shown in this study represents one of the most potent neuroprotective agents that have been reported (23–26). Most of the neuroprotective compounds are capable of protecting DA neurons in SNpc, but have much less efficacy in protecting DA content in striatum (23, 27–31). In this study, 3-HM showed equal potency in protecting DA neurons in SNpc and the DA content in striatum. The efficacy of neuroprotection was well correlated with the amelioration of behavioral damage, which revealed that 3-HM was the most potent one in attenuating both LPS-
Astroglia mediated the 3-HM-induced neurotrophic effect

In the in vitro culture systems containing astroglia, such as neuron-glia and neuron-astroglia cultures, 3-HM alone-treated groups showed a significant neurotrophic effect revealed by the dramatically enhanced DA uptake after 3-HM-alone treatment compared with vehicle-treated control cultures (Fig. 5), which was closely correlated with the increased number of DA neurons and the prolonged length of DA neurites in the 3-HM alone-treated group (Fig. 4B). However, parent compound DM itself did not show neurotrophic effect as indicated by increased DA uptake, DA neuron numbers, and neurites (2, 3). Thus, the neurotrophic effect of 3-HM is critical for explaining the more potent neuroprotective action of 3-HM on DA neurons in the SNpc and DA content in the striatum.

Further study indicated that the astroglia-mediated neurotrophic effect after 3-HM treatment was via increasing the gene expression of an array of neurotrophic factors in astroglia such as EGF, NT3, GDNF, ADNF, TGF-α, and TGF-β1. The magnitudes of the increase in the expression of these neurotrophic factors at the mRNA level were modest; probably the neurotrophic effect of 3-HM could not be attributed to a single factor, since the concentration of any individual factor released might not have been great enough to be effective. Thus, it was likely that the synergy among the released factors may account for the neurotrophic effect of 3-HM. The synergistic neurotrophic effect from the combined trophic factors has been documented (32–35).

To search for the possible mechanism for the neurotrophic action of 3-HM, we studied the acetylation of histone based on the recent findings in our lab about VPA, a drug widely used to treat seizures (36) and manic/depressive phases of bipolar mood disorder (36, 37). Histone deacetylase (HDAC) is an enzyme that catalyzes the removal of acetyl group from lysine residues of histones (38, 39). VPA was reported to directly inhibit HDAC at therapeutic levels, causing histone hyperacetylation (38, 39). HDAC has been strongly implicated in the modulation of gene expression as well as life span in a variety of organisms (40). In addition, VPA has been shown to activate the cell survival factor presumably through inhibition of HDAC (36, 41). Our lab recently found that VPA exerted neuroprotection on DA neurons and released neurotrophic factors from astroglia, which was associated with the increase in acetylation of chromatin histone H3 through the inhibition of HDAC (Chen, P. et al., unpublished results). The ability of VPA to covalently modify histone structures through enhancing acetylation can trigger changes in the gene expression of distinct downstream factors, such as neurotrophic factors (Chen, P. et al., unpublished results).

Astroglia mediated the 3-HM-induced neurotrophic effect

and MPTP-induced hypolocomotor activity. In addition, we have demonstrated that relatively high doses of 3-HM exerted negligible psychotropic side effects compared with DM and its major metabolite, DX, which is an important consideration for the potential for long-term clinical usage in PD patients. Furthermore, both MPTP and intrastral LPS injection models first target dopaminergic terminal field, which in turn cause retrograde neurodegeneration of dopaminergic neurons in the substantia nigra. Our data showing that 3-HM prevents both the degeneration of nigral dopaminergic neurons and the depletion of dopamine level in the striatum suggest a possibility that targeting dopaminergic terminal field can be considered as a strategy for the preventive therapy of PD.

Mechanistic studies using primary midbrain neuron-glial cocultures revealed two glia-dependent mechanisms for the potent neuroprotection afforded by 3-HM in an MPTP model.
increased level. This finding suggested that 3-HM increased gene expression of neurotrophic factors through the increase in acetylation of chromatin histone. Studies are under way to further determine whether 3-HM-induced the increase in the acetylation of histone H3 is due to the inhibition of HDAC or the increase in the activity of histone acetylase.

Microglia participated in 3-HM-mediated neuroprotection against MPTP-induced DA neuronal damage

We previously reported that the neuroprotective effect of DM and 3-HM in primary midbrain neuron-glia cultures treated with LPS was associated with the inhibition of LPS-induced microglia activation (2, 4), which in turn slowed down the inflammatory process and protected DA neurons from LPS-induced damage. In this study, the anti-inflammatory effect of 3-HM in the MPTP model was mediated through a mechanism different from the above-mentioned LPS model. The anti-inflammatory effect of 3-HM in the MPTP model was due to the reduced reactive microgliaosis elicited after MPTP treatment as evidenced by the decrease in the production of ROS from microglia, including extracellular superoxide and iROS. Reactive microgliaosis after MPTP treatment has been documented by our laboratory and others (20, 22, 42, 43). It is believed that the initial damage or death of DA neurons caused by MPTP could signal the activa-
Inhibition of microglial over-activation is a potential target for the neuroprotection of 3-HM

It is generally accepted that neuroinflammation generated by the activation of microglia plays a critical role in neurodegenerative disorders, such as PD. We and others have documented that over-activation of microglia produces an array of proinflammatory and neurotoxic factors such as free radicals, cytokines, chemokines, and prostaglandins, which may damage the surrounding neurons and cause neurodegeneration. Among the various factors released from microglia, superoxide generated by NADPH oxidase plays the most critical role in the subsequent neuroinflammatory process and neurodegeneration by exerting direct damage on DA neurons and promoting the production of other proinflammatory factors such as NO, TNFα, IL-1, and PGE2. Thus, NADPH oxidase is an ideal target for developing anti-inflammatory drugs due to 1) its dual roles played in inflammation-related neuronal damage, and 2) a high degree of cell type specificity because NADPH oxidase is highly expressed in immune cells such as microglia, macrophages, or neutrophils, but is in low abundance in T or B cells and is not found in most of the other cell types. Our research also shows that this anti-inflammatory approach is equally effective in preventing the over-activation of microglia caused by either a direct activation induced by LPS or an indirect reactive microgliosis elicited by MPTP/MPP+.

In summary, we show the potent neuroprotective effect of 3-HM against LPS- and MPTP-induced DA neuronal damage in in vivo and in vitro PD models through dual functions of increasing the gene expression of a series of neurotrophic factors from astroglia and decreasing superoxide and iROS production from microglia caused by the reactive microgliosis. In view of significant attenuation of behavioral damage induced by both LPS and MPTP and the negligible psychotropic side effect of 3-HM, plus the fact that 3-HM, being a small molecule, can be taken orally, 3-HM may serve as a prototype drug for the development of neuroprotective compounds that comprise both neurotrophic and anti-inflammatory properties to slow down or even halt the progression of PD.

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