Combining nitric oxide release with anti-inflammatory activity preserves nigrostriatal dopaminergic innervation and prevents motor impairment in a 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine model of Parkinson’s disease

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Abstract

Background: Current evidence suggests a role of neuroinflammation in the pathogenesis of Parkinson’s disease (PD) and in the 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP) model of basal ganglia injury. Reportedly, nonsteroidal anti-inflammatory drugs (NSAIDs) mitigate DAergic neurotoxicity in rodent models of PD. Consistent with these findings, epidemiological analysis indicated that certain NSAIDs may prevent or delay the progression of PD. However, a serious impediment of chronic NSAID therapy, particularly in the elderly, is gastric, renal and cardiac toxicity. Nitric oxide (NO)-donating NSAIDs, have a safer profile while maintaining anti-inflammatory activity of parent compounds. We have investigated the oral activity of the NO-donating derivative of flurbiprofen, [2-fluoro-α-methyl (1,1′-biphenyl)-4-acetic-4-(nitrooxy)butyl ester], HCT1026 (30 mg kg⁻¹ daily in rodent chow) in mice exposed to the parkinsonian neurotoxin MPTP.

Methods: Ageing mice were fed with a control, flurbiprofen, or HCT1026 diet starting ten days before MPTP administration and continuing for all the experimental period. Striatal high affinity synaptosomial dopamine uptake, motor coordination assessed with the rotarod, tyrosine hydroxylase (TH)- and dopamine transporter (DAT) fiber staining, stereological cell counts, immunoblotting and gene expression analyses were used to assess MPTP-induced nigrostriatal DAergic toxicity and glial activation 1-40 days post-MPTP.

Results: HCT1026 was well tolerated and did not cause any measurable toxic effect, whereas flurbiprofen fed mice showed severe gastrointestinal side-effects. HCT1026 efficiently counteracted motor impairment and reversed MPTP-induced decreased synaptosomal [³H]dopamine uptake, TH- and DAT-stained fibers in striatum and TH⁺ neuron loss in substantia nigra pars compacta (SNpc), as opposed to age-matched mice fed with a control diet. These effects were associated to a significant decrease in reactive macrophage antigen-1 (Mac-1)-positive microglial cells within the striatum and ventral midbrain, decreased expression of iNOS, Mac-1 and NADPH oxidase (PHOX), and downregulation of 3-Nitrotyrosine, a peroxynitrite finger print, in SNpc DAergic neurons.

Conclusions: Oral treatment with HCT1026 has a safe profile and a significant efficacy in counteracting MPTP-induced dopaminergic (DAergic) neurotoxicity, motor impairment and microglia activation in ageing mice. HCT1026 provides a novel promising approach towards the development of effective pharmacological neuroprotective strategies against PD.

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Background
Selective degeneration of dopaminergic (DAergic) neurons in the substantia nigra pars compacta (SN) is a pathological hallmark of both Parkinson’s disease (PD) and 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP) animal model of PD. The decline of dopamine in the striatum is associated clinically with progressive bradykinesia, tremor, rigidity and postural instability [1]. Current DAergic treatments improve the motor symptoms and quality of life for patients during the early stages of PD but do not prevent the progression of the disease associated with disabling side-effects [2]. With the exception of inherited cases linked to specific gene defects that account for <10% of cases, PD is a sporadic condition of unknown causes. Besides host genetics, environment, age, gender and inflammatory processes are factors affecting disease onset and/or progression [3-18].

Activation of microglia, a hallmark of neuroinflammation, has been demonstrated in the SN of PD patients [19], in human patients exposed to MPTP [20], and in experimental models of PD [21-35]. Accumulation of reactive oxygen species (ROS), inflammatory-associated factors including cyclooxygenase-2 (COX-2) and inducible nitric oxide synthase (iNOS)-derived NO, and pro-inflammatory cytokines (including TNF-α, IL-1β and IFN-γ) in the SN of PD patients further support that a state of chronic inflammation characterizes PD brain [5,22-26]. In addition, elevated expression of macrophage-antigen complex 1 (Mac-1), a β2-integrin family member expressed exclusively in microglia, and NADPH oxidase (PHOX), one of the major sources for production of ROS or related reactive nitric species (RNS) in activated microglia, have been reported in PD animal models [8,11,13-16,22-35]. In keeping with these findings, genetic or pharmacological inhibition of most inflammatory factors, including iNOS, PHOX, Mac-1 and COX-2-derived mediators, significantly attenuated DAergic degeneration in experimental models of PD [27-44]. Conversely, blocking the action of endogenous anti-inflammatory molecules, such as glucocorticoid hormones in transgenic mice expressing a glucocorticoid receptor (GR) antisense RNA, sharply increases microglial activation in response to MPTP, resulting in increased DAergic neuron vulnerability [8,10,11].

Consistent with the inflammation hypothesis, epidemiological analysis has indicated that nonsteroidal anti-inflammatory drugs (NSAIDs) may prevent or delay the progression of PD [6,7,45-52]. NSAIDs are among the most widely used therapeutic agents for the treatment of pain, fever and inflammation. Their effects are largely attributed to the inhibition of the enzymatic activity of COXs, of which there are two isoforms, COX-1 and COX-2. Both enzymes are responsible for arachidonic acid conversion in different prostaglandins (PGs) [53,54]. While COX-1 is constitutively expressed in most tissues, COX-2 is induced during pathophysiological responses to inflammatory stimuli [55]. Both mixed and selective COX-2 inhibitors have been reported to mitigate DAergic neurotoxicity in experimental models of PD; or to reduce LPS-induced neuronal damage [recently reviewed in {45,46}]. Besides targeting COXs, NSAIDs can act in a COX-independent way, which includes activation of the nuclear factor peroxisome proliferator-activated receptor-γ (PPAR-γ), the protection against glutamate and 1-methyl-4-phenylpyridinium ion (MPP’) toxicity, scavenging hydroxyl and NO radicals and dopamine-quinone formation [18,45-48].

Nevertheless, the long-term therapy with non-selective NSAIDs is characterized by significant adverse effects on gastrointestinal tract and kidneys, whereas increased risk of cardiovascular events has been reported with COX-2-selective inhibitors [56], which may limit their clinical use in chronic conditions. The nitric oxide (NO)-NSAID HCT1026 [2-fluoro-α-methyl(1,1'-biphenyl)-4-acetic-4-(nitrooxy)butyl ester], NO-donating furibiprofen, belongs to a novel class of anti-inflammatory agents obtained by derivatization of conventional NSAIDs with a NO-donating moiety which strongly reduce their untoward side effects without altering the anti-inflammatory effectiveness [57-66].

We herein report that HCT1026 has a safer profile and a greater efficacy than its parent compound in rescuing nigrostriatal DAergic neurons from MPTP neurotoxicity and that a shift in microglial pro-inflammatory phenotype is involved in this phenomenon. HCT1026 is safe at the gastrointestinal level, and it has been tested in humans; it is effective on oral administration, and it is thus suited for long-term treatment, thereby representing a promising approach towards the development of effective pharmacological neuroprotective strategies against PD.

Methods
Animals
Young adult (2-5 months of age) and ageing (9-11 month-old) male C57BL/6 (Charles River, Calco, Italy) housed (5 mice/cage) in a temperature (21-23°C), humidity (60%), and light (50/50 light:dark cycle, lights on at 06.00 a.m) controlled room, with controlled access to food and water, were allowed to acclimate one week before the start of the experimental protocol. Studies were conducted in strict accord with the Guide for the Care and Use of Laboratory Animals (NIH), and approved by the Review Boards of the OASI Institute (Troina, Italy). The authors further attest that all efforts
Drug administration
The drugs were compounded in the chow (Teklad 2018 diet, Harlan), the schedule of administration defined according to a pilot experiment conducted to monitor daily food intake, and the dose selected as that producing a full anti-inflammatory effect [66]. The following doses were used: HCT1026 190 ppm in the diet or 30 mg kg\(^{-1}\) day\(^{-1}\) per animal; flurbiprofen 120 ppm or 20 mg kg\(^{-1}\) day\(^{-1}\) per animal. Flurbiprofen dose was equimolar to HCT1026 (MW HCT1026:361.4; flurbiprofen:244.3, ratio HCT1026/flurbiprofen = 1.48). Plain teklas 2018 chow was used as control diet. The treatment started 7-10 days prior MPTP administration and throughout the entire experiment. Food consumption was monitored daily, diets were weighed and food intake calculated daily, body weights recorded.

MPTP administration
Both the acute [67] and the subchronic [68] MPTP injection paradigms, and three different dose-levels (5, 15, or 30 mg kg\(^{-1}\) MPTP-HCl measured as a free base), were selected in order to verify the ability of a preventive administration of HCT1026 to exert neuroprotective effects against MPTP-induced DAergic toxicity (Table 1). The same lot of MPTP-HCl (Sigma, Italy) was used for one experimental series. In a first series of experiments, in the acute protocol, MPTP was systemically injected (i.p.) at a dose of 15 mg/kg\(^{-1}\), 4 times a day, at 2 hr intervals [28,36]. In the subchronic regimen, increasing doses of MPTP were administrated i.p. at 24-h interval, for 5 consecutive days and mice sacrificed 7 days post-treatment. The dose of 15 mg/kg\(^{-1}\) day\(^{-1}\) and the subchronic regimen were then selected to assess longterm effects of HCT1026 in all subsequent experiments in ageing mice [32,42]. Groups of mice fed with the different diets and injected with vehicle (0.9% saline, 2 ml kg\(^{-1}\) intraperitoneally), served as controls (see Table 1). MPTP handling and safety measures were in accordance with published guidelines according to Jackson-Lewis and Przedborski [69].

Sacrifice and tissue processing
Controls and MPTP-treated mice were killed at selected times ranging from 0-40 days post- MPTP treatment (dpt). To study early drug effects on microglia activation during the active degeneration phase, groups of mice were studied 1-7 dpt (Table 1). To monitor the severity of nigrostriatal damage and the survival/neurorescue of nigrostriatal neurons, group of mice were studied 7, 21, 30 and 40 dpt. MPTP-induced motor deficit was assessed with the Rotarod, at -7, +1, +3 and +7 dpt. For neurochemical determinations, heads were cooled by rapid immersion in liquid nitrogen. Thereafter, striata of both sides and ventral mesencephalon, were rapidly removed and frozen at -80°C for subsequent determinations [8]. For histopathological determinations, mice were deeply anesthetized and perfused transcervically, as reported in full details [8].

Determination of drug plasma level
Blood samples were taken at the indicated times and plasma samples were frozen and stored at -80° until the analysis was performed. Plasma 0.1 ml was mixed with 10 μl ketoprofen (internal standard, 1 mg ml\(^{-1}\) stock solution in methanol) and 400 μl of cold methanol/acetonitrile (1:1) mixture, vortexed and centrifuged at 13,000 × g for 10 min at 25°C. HPLC analysis was

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**Table 1** Experimental design, animal number and analyses performed per time-point (tp) within each experimental group

| Analyses               | Subacute MPTP | Subchronic MPTP |
|------------------------|---------------|-----------------|
|                        | Days after MPTP discontinuation |               |
|                        | -7 | 0 | +1 | +3 | +7 | 0 | +1 | +3 | +7 | +21 | +30 | +40 |
| Rotarod                | 10 | + | + | + | + |   |   |   |   |   |   |   |
| Neurochem              | 5 | + | + | + | + | + | + | + | + | + | + | + |
| Immunohistochem.       | 5 |   |   |   |   |   |   |   |   |   |   |   |   |
| Microglia markers      | 5 | + | + | + | + | + | + | + | + | + | + | + |
| Gene expression        | 4 | + | + | + | + | + | + | + | + | + | + | + |
| Western blot           | 4 | + | + | + | + | + | + | + | + | + | + | + |

Young (3-5 month-old) and ageing (9-11 month-old) C57Bl/6 mice fed with a control, flurbiprofen or HCT1026 diets (30 mg kg\(^{-1}\)) starting at -10 d, underwent an MPTP treatment according to the subacute (n = 4 intraperitoneal, i.p., injections of MPTP-HCl, 15 mg kg\(^{-1}\) free base, 2 hours apart in one day) or subchronic (n = 5 intraperitoneal, i.p., injections of MPTP-HCl, at 5, 15 or 30 mg kg\(^{-1}\) free base, every 24 h), injection paradigms. Age-matched mice fed with the different diets received physiologic saline (NaCl, 10 ml kg\(^{-1}\)) and served as controls. Mice number and timepoints (tp) for each experimental protocol for each set of analyses/tp, are indicated. Striatum and ventral midbrains tissues were processed for RT-PCR or western blotting, as reported according to the different determinations. For immunohistochemical analyses, on the day of sacrifice mice were anesthetized and transcervically perfused, the brains processed as indicated. See Methods section for details.
performed on fixed volume of organic extraction mixture. Chromatographic analysis was performed on an Agilent 1100 series system equipped with a Diode array detector operating at 246-nm single wavelength. Separations were achieved with a gradient elution on a Synergi MAX-RP 80A column (150 x 2 mm i.d.; 4 μm) equipped with MAX-RP precolumn (4 x 2 mm i.d.). The mobile phase was acetonitrile and phosphoric acid 0.1%. The flow rate was 1 ml min^{-1} and the column temperature was 25°C. Under these conditions, the retention time of HCT1026, flurbiprofen and internal standard were 5.7, 1.3, 0.8 min, respectively. Only flurbiprofen was detectable in plasma of HCT1026 treated mice. HCT1026 and its des-nitro metabolite (i.e. HCT1027) were undetectable in all plasma samples, as previously reported [70].

Motor behavior analysis with the rotarod
An accelerating rotarod (five-lane accelerating rotarod; Ugo Basile, Comerio, Italy) was used to measure motor balance and coordination in mice. Mice have to keep their balance on a horizontal rotating rod (diameter, 3 cm) and rotation speed was increased every 30 sec by 4 rpm. Five mice were tested at the same time, separated by large disks. A trial starts when the mouse is placed on rotating rod, and it stops when the mouse falls down or when 5 min are completed. Falling down activates a switch that automatically stops a timer. The testing day, each mouse is submitted to 5 trials with an intertrial interval of 30 min. Mice housed five per cage were acclimated to a 12 h shift in light/dark cycle so that the exercise occurred during the animals normal wake period. Saline- and MPTP-treated mice fed with a control or HCT1026 diet (10/experimental group) were assessed for their Rotarod performance on day -7, +1, + 3 and +7 dpt.

High-affinity [3H]dopamine uptake assay
Left and right striata were homogenized in ice-cold prelysis buffer (10 mM Tris, pH 7.5, and 0.32 M sucrose) using a Teflon pestle-glass mortar and homogenized tissue centrifuged for 10 min at 1000 x g at 4°C to remove nuclei. The supernatant containing the synaptosomes was collected and aliquots removed for the determination of protein content [71] and dopamine uptake (total high affinity and mazindol non-inhibitable). Fifty micro-liter of supernatant were diluted in Krebs-Ringer phosphate buffer (16 mM NaH2PO4,16 mM Na2HPO4, 119 mM NaCl, 4.7 mM KCl, 1.8 mM CaCl2, 1.2 mM MgSO4, 1.3 mM EDTA, and 5.6 mM glucose; pH 7.4), and incubated at 37°C in the presence or absence of mazindol (10 μM), a high affinity dopamine up-take inhibitor [8]. [3H]Dopamine (25 nM, specific activity, 20-40 Ci/mmol; Amersham, Arlington Heights, IL) was added in Krebs-Ringer buffer and incubation carried on for 6 min at 37°C. Synaptosomes were collected on pre-soaked nitrocellulose filters by filtration and non-specific radioactivity was washed with Krebs-Ringer phosphate buffer followed by filtration. The filters were then transferred into scintillation vials and measured by liquid scintillation (Cytoshunt; ICN, Costa Mesa, CA) counter (Packard). Specific high-affinity neuronal dopamine uptake is expressed as femtomoles of dopamine uptake per microgram of protein minus the femtomoles of mazindol uptake. Values are represented as % changes in dopamine uptake vs. control.

Measurement of MPP+ levels in the striatum
Mice fed with the different diets were killed 90 min after MPTP injection. Both left and right striata were dissected on ice, placed into a vial containing 250 ul of 0.4 N perchloric acid and sonicated. After centrifugation, MPP+ determinations were carried out by HPLC using 5SCX column (Phenomenex). The mobile phase was composed of H2SO4 0.1 M, triethylamine 0.075 M, and acetonitrile 10% at pH 2.3; the flow was 1.5 ml mm^{-1} [8].

Immunohistochemistry
On the day of sacrifice, mice were anesthetized by intra-peritoneal injection of Nembutal (50 mg/kg). Mice were rapidly perfused transcardially with 0.9% saline, followed by 4% paraformaldehyde in phosphate buffer (pH 7.2 at 4°C). Brains were carefully removed and post-fixed for 2-4 hrs, in 4% paraformaldehyde in phosphate buffer saline, pH 7.2 (PBS) and later placed in 15% sucrose in PBS overnight at 4°C. Tissues were frozen at -80° C. Serial cryostat sections (10 μm, from the olfactory bulb to the end of the medulla), were collected, mounted on poly-L-lysine-coated slides and processed for immunohistochemistry. Identification of the level was made by comparison with the sections of the mouse brain [72]. All immunostaining procedures were carried out on sections incubated in blocking buffer (0.3 - 0.5% Triton X-100, 5% BSA and 5% normal serum in PBS) for 30 min at room temp followed by an overnight incubation with the following primary antibodies in blocking buffer at 4° C: (i) tyrosine hydroxilase (TH) (goat anti-TH, Santa Cruz Biotechnology, Inc. USA 1:1000; rabbit anti-TH Pelfreez, Roger, AR, 1: 2000); (ii) rat anti dopamine transporter DAT, Millipore Corp. USA, 1:1000), as markers of dopaminergic neurons; (iii) membranolytic attack complex of complement (rat anti Mac-1/CD11b, Pharmingen International, Becton Dickinson, USA or anti Mac-1, Serotec, Oxford, UK, 1: 1000), as microglial marker; (iv) inducible nitric oxide synthase (rabbit anti-iNOS, Santa Cruz Biotechnology, Inc, 1: 200), 3-nitrotrosine (rabbit anti-3-NT, Upstate, Lake Placid, NY, US, 1: 200), as NO/peroxynitrite finger-print.
sections were counterstained with nuclear counterstain (Dapi or PI, by Vector Laboratories Inc. Burlingame, CA, USA).

All antibodies, whether used for single or dual labeling procedures, were visualized by immunofluorescence, except for TH-Ab that was also visualized using immunoperoxidase. Adjacent tissue was also stained with cresyl violet to validate TH neuron survival [8,73,74]. Sections were incubated with the indicated dilutions of the antibodies, either alone or in combination as described. After 3 (× 5 min) washes in PBS, primary antibodies were revealed with specific FITC and Cy3 conjugated secondary antibodies 1:100-1:200 dilution. (60 min at room temp). After 3 (× 5 min) washes in PBS, sections were mounted with Gel mounting solution (Biomeda corp. Foster City, CA, USA). In all of these protocols, blanks were processed as for experimental samples except that the primary antibodies were replaced with PBS.

**Loss of TH-positive neurons and striatal DAergic innervation**

Loss of TH-positive (TH⁺) SNpc neurons was determined by serial section analysis of the total number of TH⁺ cells counted throughout the entire rostro-caudal (RC) axis of the murine SNpc (Bregma coordinates: -2.92, -3.08, -3.16, -3.20, -3.40 and -3.52) according to Franklin and Paxinos [72] at 7, 21, 30 and 40 days post-MPTP (dpt) or saline injection [8]. Cell counting was done in both side of the brain for each animal, and then right and left values were added to generate a total DA SNpc neuron count, in a total of five animals per experimental group. TH-labeled neurons were scored as positive only if their cell-body image included well defined nuclear counterstaining. Estimates of total TH⁺-stained and cresyl-violet-stained neurons in the SNpc were calculated using the Abercrombie’s correction [74]. The total number of TH⁺ cell bodies was estimated and examined by two independent researchers, in a blind fashion. Loss of striatal DAergic innervation was assessed by quantification of TH- and DAT-immunofluorescent (IF) signal intensity in 10 μm-thick coronal sections located at 0.5, 0.8 and 1.1 mm from bregma, and analyses carried out by confocal laser microscopy as described [8].

**Confocal laser microscopy and image analysis**

Sections labeled by immunofluorescence were visualized and analyzed with a confocal laser scanning microscope LEICA TCS NT (Version 1.0, Leica Lasertechnik GmbH, Heidelberg, Germany), equipped with an argon/krypton laser using 10 ×, 20 ×, and 40 × and 100 × oil-immersion objectives. Pinhole was set at 1-1.3 for optical sections of 0.48-0.5 μm. For TH⁺ and DAT⁺ fibers in striatum, fluorescence intensity per unit of surface area was determined in 10 randomly selected fields (250,000 μm²) using computer-assisted image analysis software (LEICA). Single lower power scans were followed by 16 to 30 serial optical sectionings. Laser attenuation, pinhole diameter, photomultiplier sensitivity, and off-set were kept constants. The average fluorescence intensity (pixel, mean ± SEM per unit surface area) was measured throughout the stack. Within the same stacks, the background pixel intensity in areas devoid of fibers/cells was determined and subtracted. For assessment of reactive microglial cell number, ameboid-shaped Mac-1⁺ cells [75] were counted in striatal and SNpc coronal sections, cell counts averaged for each animal and the mean number of cells per mm² per animal was estimated. A comparable countable area ranging from 1.90 mm² to 2.00 mm² was analyzed in the different MPTP groups. Double-labelled cells with iNOS and Mac-1, were counted and expressed as above. Dual stained TH⁺ 3-NT⁺ cells were counted and values expressed as a percent double-stained TH⁺ NT⁺ /TH⁺ neurons. Each label was analyzed on a total of 12 sections per mice and in at least 4 mice per group. Analyses were performed by two independent researchers blind to the experiment.

**Semi-quantitative RT-PCR**

To analyze transcript levels, total RNA was isolated from striatum and ventral mid brain using the RNeasy isolation Midi kit (Qiagen, #75144). Tissue samples were homogenized in 1 ml of QIAzol Lysis Reagent (Qiagen, #79306) using a rotor-stator homogenizer. Total RNA was isolated from homogenized tissue samples using RNeasy Lipid Tissue Kit (Qiagen, #74804) including DNase digestion. At the end, RNA samples were redissolved in 30 μl of RNase-free water and their concentrations were determined spectrophotometrically by A260 (Nanodrop-ND 1000). The cDNA was synthesized from 2 μg of total RNA using the Retroscript Kit (Ambion, #AM2224) following the manufacturer’s directions. 250 ng of cDNA were used for PCR (96°C for 1 min for 2 cycles; 96°C for 1 min, 58°C for 4 min; 94°C for 1 min, 58°C for 2,5 min for 35 cycles, with a final extension at 70°C for 10 min) by using Super Taq DNA polymerase (Ambion, #AM1710) with specific primer pairs for TH (F: cttggaatccaaacagaggtggtggttatgtag; amplicon: 620 bp); DAT (F: cagagaggtgtgacgctca; R: ggacagttcctcagacaacc; amplicon: 328 bp), iNOS (F: actgctcccggtgtggtggttatgtag; amplicon: 500 bp) and Classic S18 Standard (amplicon: 495 bp; #Ambion AM1720), to normalize the expression. Samples from PCR reactions were separated electrophoretically on 2% agarose gel containing 0.2 μg ml⁻¹ of ethidium bromide. Fluorescent bands of amplified gene products were captured by using Gel Logic 200
Imaging System (Kodak), values normalized against S18, and ratios expressed as percent of control (saline-injected), within each experimental group.

**Western blot analysis**

Protein extracts were prepared for striatum and ventral midbrain (which included the SNpc) (left and right sides) at the indicated time-intervals after saline or MPTP injections (n = 3 per group). The tissue samples were homogenized in lysis buffer (0.33 M sucrose, 8 mM Hepes, pH 7.4 and protease inhibitors) and quantified using the BCA protein determination method (Bio-Rad, Hercules, CA). Protein samples were diluted to equivalent volumes containing 20 µg of protein and boiled in an equal volume of Laemli SDS boiling buffer (Sigma) for 10 min. Samples were loaded into a 9-12% SDS-polyacrilamide gel and separated by electrophoresis for 3 h at 100 V. Proteins were transferred to polyvinylidene difluoride membrane (Amersham Biosciences, Piscataway, NJ) for 1.5 h at 300 mA. After blocking of nonspecific binding with 5% nonfat dry milk in TBST, the membranes were then probed with the following primary antibodies: rabbit anti-TH (Chemicon); rat anti-DAT (Millipore), rabbit anti-Mac1 (AbCam), mouse anti-gp91phox (BD Transduction Laboratories), β-actin (Cell Signaling). After incubation at room temperature for 1 hr, membranes were washed and treated with appropriate secondary antibodies conjugated with horseradish peroxidase (HRP) and blot were exposed onto radiographic film (Hyperfilm; Amersham Bioscience). Membranes were reprobed for β-actin immunolabeling as an internal control. The bands from the Western blots were densitometrically quantified on X-ray films (Image-Quantity One). The data from experimental band were normalized to β-actin, before statistical analysis of variance and values expressed as % of saline-injected controls.

**Statistical analysis**

Data were analyzed by means of two-way analysis of variance (ANOVA), with group and time as independent variables and given as mean±SEM. Striatal neurochemical data (nmol or pmol mg protein⁻¹) are expressed as % of controls. Comparisons a posteriori between different experiments were made by Student-Newman-Keuls t-test.

**Results**

HCT1026 prevents administration counteracts MPTP-induced down-regulation of high affinity synaptosomal DA uptake in the striatum

We first assessed the short-term effect of the control and medicated diets on striatal high affinity synaptosomal [³H]DA uptake (Figure 1A,B), a sensitive quantitative indicator of DAergic axonal terminal density [76]. In 2-3 month-old mice fed with flurbiprofen or HCT1026 diets and treated with saline, DA uptake levels were not different compared to saline-treated mice fed with the control diet. On the other hand, in mice fed with a control diet and exposed to the subacute MPTP (15 mg kg⁻¹, 4 times a day, at 2 h intervals), we observed after 7 d the severe decrease (-75%) of striatal DA uptake (Figure 1A). In mice fed with flurbiprofen and treated with MPTP, a certain degree of protection was observed, as reflected by the less severe decrease (-54%) of DA uptake. On the other hand, HCT1026 afforded a greater protection, as illustrated by the significantly (p < 0.05) smaller (-25%) decrease of [³H] DA uptake levels as compared to mice fed with the control or flurbiprofen diets. With the subchronic (administration of MPTP at 24-h interval, for 5 consecutive days) regimen, increasing the daily doses of MPTP resulted in a dose-dependent loss of DA uptake levels (Figure 1B). By contrast, mice fed with HCT1026 and exposed to 5 mg kg⁻¹ day⁻¹ for five days, were resistant to MPTP-induced DAergic toxicity, as revealed by the counteraction of the almost 38-48% loss of striatal DA uptake measured in mice fed with the control or flurbiprofen diets. In addition, mice exposed to higher (15 and 30 mg kg⁻¹ day⁻¹) MPTP doses and fed with control or flurbiprofen diets exhibited far greater (p < 0.05) decreases of striatal DA uptake compared with mice fed with HCT1026 (Figure 1B).

The preventive oral administration with HCT1026 was well tolerated and did not cause any measurable toxic effect through the treatment, whereas flurbiprofen-fed mice showed severe gastrointestinal side-effects (bleeding). Due to flurbiprofen toxicity, only HCT1026 was studied in the long-term experimental protocol.

To more closely mimick PD condition, we next assessed the longterm efficacy and safety of HCT1026 in ageing (9-11 month-old) mice. Reportedly, the process of ageing increases DAergic vulnerability to MPTP and limits the repair capacity of the nigrostriatal DAergic system [77,78]. We thus selected the subchronic MPTP regimen, at a dose of 15 mg/kg [28,32,42]. Consistent with previous findings [77,78], ageing mice fed with a control diet did not recovered from MPTP insult, as revealed by an almost 70-75% decrease of DA uptake levels measured up to 40 dpt, whereas in mice fed with HCT1026, a significant degree of protection was measured, as reflected by the significant amelioration of striatal DA uptake levels at all time-points studied (Figure 1C). These data indicate that the neuroprotective activity of HCT1026 was maintained up to 40 dpt. In addition, the longterm administration of HCT1026 does not cause any measurable toxic effects.

HCT1026 inhibits MPTP-induced motor impairment

To verify the ability of HCT1026 to affect MPTP-induced impairment of motor coordination [42,79,80], we assessed the ability to maintain the balance on a rotating cylinder
and to adapt to the rate of locomotor activity, by using the Rotarod test, as described (Figure 1D). Saline- and MPTP-treated mice fed with a control, flurbiprofen or HCT1026 diets (30 mg kg⁻¹) starting at -10 d, underwent an MPTP treatment according to the subacute (A) or subchronic (B) injection paradigms, as described. Age-matched mice fed with the different diets received physiologic saline (NaCl, 10 ml kg⁻¹) and served as controls. Seven days after MPTP discontinuation, loss of DAergic functionality was assessed in striatum measuring high affinity synaptosomal striatal [¹³C]DA uptake [8]. HCT1026 proved to be more potent than its parent compound in counteracting MPTP-induced decreases in striatal DA uptake levels in both the subacute (A) and subchronic (B) protocols. Differences were analyzed by ANOVA followed by Newman-Keuls test, and considered significant when p < 0.05. **p < 0.05 vs saline, ° p < 0.05 vs MPTP + control diet.

C: Ageing mice fed with a control or HCT1026 diets, were submitted to the subchronic MPTP regimen, and striatal DA uptake levels measured 21, 30 and 40 d after MPTP (n = 6/time point). Note the long-lasting counteraction of MPTP-induced striatal toxicity in mice fed with HCT1026 as opposed to the control diet. D: Motor performances on Rotarod of saline- and MPTP-treated mice (n = 10/group) fed with a control or HCT1026 diets. Time of permanence on revolving bars (ordinate) are plotted against pre- and post-treatment days (5 trials/day) during which experiments were performed. Mean and SEM values are reported. Establishment of a motor deficit measured 1-7 dpt, is counteracted by HCT1026. Differences were analyzed as above. ** p < 0.05 vs saline; ° p < 0.05 vs MPTP + control diet.

MPTP-treated mice fed with a control diet exhibited a significant decrease in the mean latency to fall at all time-points tested, compared to saline-injected mice (p < 0.05), defining a motor deficit in MPTP-treated animals (Figure 1D). In MPTP mice fed with HCT1026, the mean latency to fall was significantly (p < 0.05) increased compared to MPTP mice fed with a control diet at all time tested. By 7 dpt, HCT1026-fed mice performed as good as the control mice, indicating a significant reduction of the motor impairment by the preventive treatment with HCT1026.
HCT1026 inhibits MPTP-induced loss of striatal TH and DAT at mRNA and protein levels

Tyrosine hydroxylase (TH) is the rate-limiting enzyme in dopamine biosynthesis and a marker for DA neurons. The dopamine transporter, DAT, is a highly specific marker of projecting DAergic nigrostriatal neurons and thus, its expression is proportional to the loss of striatal dopamine content [76]. Accordingly, we examined TH and DAT striatal expression using RT-PCR, immunohistochemistry coupled to confocal microscopy, and western blot (WB) analyses. RT-PCR of TH (Figure 2A and 2B) and DAT (Figure 2E and 2F) mRNAs in striatum...
Figure 3  HCT1026 inhibits MPTP-induced loss of striatal TH- and DAT- proteins by immunohistochemistry and western blotting.

Ageing C57Bl/6 mice fed with a control (ct) or HCT1026 diets starting at -10 d, underwent an MPTP treatment, as described. At different time-intervals, mice were anesthetized and rapidly perfused, the brains were carefully removed and processed for immunohistochemistry, as described. TH- (A) and DAT- (E) IR in striatum (Str) assessed by immunofluorescent staining and image analysis by confocal Laser microscopy in ageing mice fed with ct or HCT1026 diets, treated with saline or MPTP (n = 5/time point). Fluorescence intensity values (Fl, means ± S.E.M.) are expressed as % of saline. **p < 0.05 vs saline, °p < 0.05 vs MPTP fed with control diet.

B-H: Representative confocal images show loss of TH-IF (revealed by FITC, green) in Str of MPTP mice fed with a ct diet at 40 dpt (C) and a substantial rescue of TH- (D) by HCT1026.

F-H: Representative confocal images show loss of DAT-IF (revealed by FITC, green) in Str of MPTP mice fed with a ct diet at 40 dpt (G) and a substantial rescue of DAT-IF (H) by HCT1026.

I-F: For western blot analysis, at 40 d after saline or MPTP injections in mice fed with the ct or HCT1026 diets, mice were sacrificed and striatal tissue samples processed for WB, as described. The data from experimental bands were normalized to β-actin, before statistical analysis of variance and values expressed as % of saline-injected controls, within each respective group. Note the significant decreased TH (I) and DAT (J) protein levels in MPTP mice fed with a ct diet, whereas a recovery was observed in HCT1026 fed mice. *p < 0.05 vs saline, ** p < 0.05 vs MPTP fed with ct.
Figure 4 HCT1026 preventive administration inhibits MPTP-induced dose-dependent loss of TH⁺ cell bodies in SNpc. Ageing mice fed with a ct or HCT1026 diets were submitted to the subchronic MPTP (5 mg, 15 mg or 30 mg kg⁻¹, for 5 consecutive d) regimen, and DAergic cell survival studied after 7 d. A-E: Representative confocal images of dual staining with TH- (green) and DAT- (red) -Abs of coronal midbrain sections at the level of the SNpc 7 d after MPTP. Note the significant protection afforded by HCT1026 preventive treatment in mice treated with the 5 mg kg⁻¹ dose (see panels G-I) as compared to MPTP mice fed with a ct diet (D-F). F: Survival of DAergic cell bodies in SNpc. The total number of TH⁺ and Nissl⁺ neurons was counted throughout the entire rostro-caudal axis of the SNpc. Treatment groups were averaged (means ± S.E.M.) * p < 0.05 vs saline; **p < 0.05, vs MPTP mice fed with a ct diet. HCT1026 significantly reduced the dose-dependent decrease of TH⁺ and Nissl⁺ neurons.
Figure 5 Long-lasting protection of DAergic cell bodies in HCT1026 fed mice. Ageing mice fed with a ct or HCT1026 diet were submitted to MPTP subchronic regimen (15 mg kg⁻¹ for 5 consecutive d) and sacrificed at different time-intervals after MPTP. A-L. Representative confocal images of dual staining with TH- (green) and DAT- (red) -Abs of coronal midbrain sections at the level of the SNpc 7 and 40 d after MPTP. As observed, MPTP mice fed with a ct diet show a marked loss of TH+DAT+ neurons at 7 d (see panels A-C) and fail to recover 40 d following MPTP injury (G-I). By contrast, HCT1026 afforded a significant and long-lasting protection (see panels D-F and J-L). M. The total number of TH+ and Nissl+ neurons was counted throughout the entire rostro-caudal axis of the SNpc. Treatment groups were averaged (means ± S.E.M.) * p < 0.05 vs saline; **p < 0.05, vs MPTP mice fed with a ct diet. HCT1026 significantly reduced the decrease of TH+ and Nissl+ neurons observed up to 40 d in mice fed with a ct diet.
indicated no significant differences in transcript levels in saline-treated mice fed with control or HCT1026 diets. By contrast, exposure to MPTP induced a significant and longlasting decrease of TH mRNA in ageing mice fed with a control diet after either 7, 21 d (Figure 2A and 2C), 30 or 40 d (Figure 2A and 2D). In analogy with these findings, DAT mRNA levels were sharply down-regulated at 7, 21 (Figure 2E and 2G), 30 and 40 d (Figure 2E and 2H) post- MPTP in mice fed with a control diet. By contrast, in mice fed with HCT1026, TH and DAT mRNAs (Figure 2A, B, C and 2D) and DAT mRNA (Figure 2E, F, G and 2H) levels were significantly increased compared with those measured in MPTP mice fed with a control diet, thus supporting...
Figure 7 HCT1026 inhibits MPTP-induced increased Mac-1, PHOX, iNOX and 3-Nitrotyrosine expression in SNpc.

A-B: Western blotting of phagocyte oxidase PHOX (A) and Mac-1 (B) within the VM at different time-intervals after saline or MPTP injection in mice fed with the control or HCT1026 diet. The data from experimental bands were normalized to β-actin, before statistical analysis of variance. Values are expressed as % of saline-injected controls. Differences were analyzed by ANOVA followed by Newman-Keuls test, and considered significant when p < 0.05. * p < 0.05 compared to saline; ° p < 0.05 vs MPTP fed with the control diet.

C: Semi-quantitative RT-PCR for iNOS. The 250 ng of cDNA were used for PCR, by using Super Taq DNA polymerase with specific primer pairs for iNOS (500 bp) and Classic GADPH Standard (270 bp). Samples from PCR reactions were processed as described. Fluorescent bands of amplified gene products were analyzed, the values normalized against GADPH and ratios expressed as % of control, within each experimental group (C), see text for details. Differences were analyzed by ANOVA as above. ** p < 0.05 vs saline; ° p < 0.05 vs MPTP + control diet.

D: Mean numbers of Mac1+iNOS+ cells within the SNpc in saline and MPTP mice fed with the control or HCT1026 diet. Cell counts obtained as described in Material section.

E-F: Representative confocal images showing double staining with Mac-1 (green) and iNOS (red) in MPTP mice fed with the control (E) or HCT1026 (F) diets.

G: Percent (%) of TH+ neurons colocalizing with 3-nitrotyrosine (3-NT), a peroxynitrite footprint. Dual stained TH+ 3-NT+ neurons were counted as described and values expressed as % of total TH+ neurons.

H-I: Representative confocal images showing dual immunostaining with 3-NT (green) and TH (red) in MPTP mice fed with a ct diet (H) showing that a large proportion of DAergic neurons colocalize (orange-to-yellow) as opposed to TH neurons of mice fed with HCT1026 (E) where no colocalization was observed in the large part of SNpc neurons.
HCT1026-induced striatal DAergic neuroprotection was maintained at long time intervals.

As observed (Figure 3A and 3E), average striatal TH- and DAT-immunofluorescent (IF) signal intensity (pixel ± SEM), did not differ in saline-injected mice fed with either control or HCT1026-mediated diets. Subchronic MPTP treatment decreased TH- (Figure 3A and 3C) and DAT-IF (Figure 3E and 3G). In addition, loss of striatal DAergic innervation lasted up to 40 dpt. By contrast, HCT1026 significantly counteracted the loss of TH (Figure 3A and 3D)- and DAT (Figure 3E and 3H)-IF signal intensity, up to 40 dpt. Changes in TH and DAT proteins were next quantified by WB in saline and MPTP mice fed with the control or HCT1026 diet (Figure 3I and 3J). After 40 d from from MPTP administration, a significant reduction of MPTP-induced downregulation of both markers was observed only in mice fed with HCT1026 diet. These and previous findings support the long-term efficacy of HCT1026 in preserving TH and DAT expression and function in striatum of ageing mice exposed to subchronic MPTP regimen.

MPTP metabolism is not affected by HCT1026 preventive treatment

One of the first limiting factors in MPTP toxicity is the conversion of MPTP into MPP+ by means of the monoamine oxidase B (MAO-B) enzymatic activity. MPP+ is known to gain access into neurons via DAT, and subsequently converted into MPP+ by means of monoamine oxidases, including MAO-B. The resulting MPP+ is then taken up by DAT and transported into the cell cytosol where it accumulates. In our study, MPP+ levels measured 90 min after MPTP injection in mice fed with the different diets and exposed to MPTP. There was no significant difference in striatal MPP+ levels measured 90 min after MPTP injection in mice fed with either the control (110 ± 7 ng mg⁻¹ protein) or HCT1026 diet (124 ± 12 ng mg⁻¹ protein) diets, thereby indicating that the greater protection afforded by HCT1026 might not be attributed to poor MPP+ metabolism.

HCT1026 preventive administration decreases MPTP-induced loss of TH+ cell bodies in SNpc

We next assessed the impact of HCT1026 in MPTP-induced toxicity of nigral DAergic cell bodies (Figure 4 and Figure 5). In mice fed with a control diet, MPTP induced a dose-dependent loss of double-stained TH+ (revealed by FITC, in green) DAT+ (revealed by CY3, in red) cells in SNpc (compare Figure 4A, B and 4C with Figure 4D, E and 4F), whereas MPTP-induced DAergic neurotoxicity was significantly (p < 0.05) reduced in HCT1026-fed mice (Figure 4G, H and 4I). Estimation of the total number of TH+ and DAT+ neurons using the Abercrombie correction, confirmed a dose-dependent reduction of TH+ and DAT+ neurons in MPTP-treated mice fed with control diet, suggesting an actual TH+ neuronal loss rather than loss of TH expression (Figure 4I). By contrast, HCT1026-fed mice exhibited a significantly greater number of TH+ and DAT+ neurons at all doses tested (Figure 4I). These results suggest that a certain number of DAergic neurons could survive the MPTP insult in mice receiving a preventive treatment with HCT1026.

In order to verify the ability of the continuous oral treatment with HCT1026 to maintain its neuroprotective effects, saline and MPTP (15 mg kg⁻¹ day⁻¹ per 5 consecutive days), mice fed with the control or HCT1026-medicated diets. Subchronic MPTP treatment decreased TH- (Figure 5A, B, C and 5M), 21, 30 (Figure 5M), or 40 dpt (Figure 5G, H, I and 5M) MPTP mice fed with the control diet did not recover as reflected by the significant, long-lasting loss of TH+ and DAT+ neurons. In sharp contrast, HCT1026 exerted a significant neuroprotective effect that was maintained up to 40 dpt (see Figure 5D, E, F, and 5I, K, L, M), suggesting increased survival/rescue of SNpc neurons in HCT1026-fed mice.

HCT1026 inhibits MPTP-induced microglial activation

Glial inflammatory mechanisms are thought to contribute to MPTP-induced nigrostriatal DAergic degeneration (see Refs in Background). Indeed, when microglia adopt a pro-inflammatory phenotype, the production and release of a plethora of toxic mediators, including pro-inflammatory cytokines and iNOS-derived NO, can enhance neuronal damage in the SNpc and accelerate the appearance of behavioral symptoms [32,81-83]. The ability of HCT1026 to modulate MPTP-induced microglial activation was next assessed during the early phase of active degeneration [8] using immunohistochemistry, WB and RT-PCR analysis.

Microglial cell number/morphology in striatum and SNpc

Changes in activated microglial cell number and morphologic appearance at both striatal and SNpc levels were assessed using Mac-1-Ab, an integrin receptor known to mediate reactive microgliosis and recognized by CD11b/CD18 antibodies. A significant increase in microglial cell number was observed in the SNpc of MPTP-treated mice in comparison to saline-treated mice. The number of microglial cells was significantly lower in HCT1026-treated mice, indicating a protective effect of HCT1026 on microglial cell number.

In summary, our findings provide evidence for the neuroprotective effects of HCT1026, a common anti-inflammatory drug, in the MPTP model of PD. The preventive administration of HCT1026 significantly prevented MPTP-induced nigrostriatal DAergic degeneration, preserved TH- and DAT-immunofluorescent (IF) signal intensity, and reduced the number of TH+ and DAT+ neurons in the SNpc. These results suggest that HCT1026 could be a potential therapeutic target for the treatment of PD.
more elongated cell body with longer and thinner processes in mice fed with HCT1026 diet (see insets in Figure 6G and 6N). Moreover, the number of ameboid-like Mac-1+ cells was sharply reduced both in striatum (Figure 6A, F and 6G) and midbrain (Figure 6H, M and 6N) of HCT1026-fed mice.

**Mac-1 and PHOX expression in the ventral midbrain (VM)**

Mac-1 is linked to the activation of PHOX, a chief component of MPTP-dependent microglia activation [29-33]. Given that Mac-1 and PHOX act in a synergistic way, and are essential to enhance DAergic degeneration, we next evaluated the expression of both markers within the ventral midbrain by WB, during the maximal phase of glia activation [8]. As previously observed, a sharp increase of Mac-1 and PHOX followed MPTP treatment, as early as 1 dpt, in mice fed with the control diet, with levels remaining high at both 3 and 7 dpt (Figure 7A, B). Conversely, HCT1026 significantly reduced MPTP-induced Mac-1 and PHOX expression in the ventral midbrain, supporting reduced microglial activation during the early phase of DAergic degeneration (Figure 7A, B).

**Expression of pro-inflammatory mediators: iNOS and 3-NT**

The ability of a preventive treatment with HCT1026 to inhibit MPTP-induced up-regulation of iNOS was next evaluated by semi-quantitative RT-PCR analysis 1 d after MPTP challenge. Within the ventral midbrain of salinetreated mice fed with a control or HCT1026 diet, no significant difference was observed in iNOS mRNA levels, whereas MPTP significantly up-regulated iNOS transcription (Figure 7C). On the other hand, the MPTP mice fed with HCT1026 exhibited significantly less iNOS mRNA expression compared to controls (Figure 7C).

When glial superoxide generated by PHOX activation and iNOS/NO are concomitantly active, peroxynitrite (ONOO-), and consequently protein tyrosine nitration and hydroxyl radicals over-production occurs [33]. Hence, the generation of high NO concentration followed by the production of peroxynitrite may be involved in DAergic neuronal cell death. We thus used immunohistochemistry to localize iNOS and 3-nitrotyrosine (3-NT) as fingerprint of iNOS-derived NO and peroxynitrite generation. Dual immunostaining with iNOS and Mac-1 within the SNpc of MPTP mice fed with a control diet, indicated that a significant proportion of reactive Mac-1 cells expressed iNOS, 1-7 d post-MPTP (Figure 7D, E). Dual staining with 3-NT- and TH indicated a sharp increase in the percentage of TH+ cells colocalizing with 3-NT at both 1 and 3 d post-MPTP (Figure 7G, H), while a certain decrease was observed 7 dpt. Conversely, in MPTP mice fed with HCT1026, both iNOS- (Figure 7D, F) and 3-NT- (Figure 7G, I) were significantly downregulated 1-7 dpt within the SNpc.

These results indicated an efficient reduction of MPTP-induced microglial activation, pro-inflammatory marker expression, i.e. Mac-1, PHOX, iNOS and 3-NT formation within the SNpc of mice fed with HCT1026, which might account for the observed nigrostriatal protection.

**Discussion**

Grafting a NO-donating moiety into the structure of flurbiprofen, one of the most potent non-selective anti-inflammatory agents, yielded a drug devoided of side toxicity and endowed with a remarkable neuroprotective activity against MPTP-induced DAergic neurotoxicity, motor impairment and microglia activation in ageing mice. HCT1026 was effective in both the acute and sub-chronic models of MPTP, and its neuroprotective activity lasted up to 40 d, as opposed to age-matched controls indicating the longterm safety and efficacy of the HCT1026, as opposed to flurbiprofen. The different outcome of mice treated with HCT1026 was not due to differences in daily food consumption, or to poor MPP+ metabolism. HCT1026-induced neuroprotection was associated with a marked down-regulation of activated microglial cell number and a signifcant decrease of MPTP-induced pro-inflammatory mediators, including iNOS, Mac-1 and PHOX expression, as well reduced 3-NT formation in SNpc DAergic neurons, suggesting that a switch in microglia pro-inflammatory phenotype might contribute to nigrostriatal neuroprotection.

The experimental study was designed to compare the oral activity of HCT1026 with that of flurbiprofen. However, due to the significant gastric toxicity of flurbiprofen observed in the short-term study, only HCT1026-mediated diet was further studied. In long-term experimental protocols, HCT1026 proved to have a safe profile and a significant efficacy in counteracting MPTP-induced striatal DAergic toxicity.

Nitric oxide (•NO) is a biological molecule known to play a major role in a wide variety of physiological and pathological conditions [84]. Some of its functions include vasodilation of blood vessels, GI mucosal healing and defense. Therefore NSAIDs containing NO-donor groups have been developed to obtain effective treatment of inflammation with reduced GI side effects [57-61]. Indeed, grafting an organic nitrate moiety onto the NSAID scaffold has been shown to result in the release of NO through slow kinetics (in comparison with others NO donors, i.e. sodium nitroprusside, S-nitroso-N-acetyl-D,L, penicillamine), possibly mimicking the physiological levels of NO produced by constitutive NO synthases. Thanks to NO release, and the combination of a balanced inhibition of the two main COX...
isoforms, NO-NSAIDs are endowed with little gastrointestinal and renal toxicity compared to their parents compounds [56-61,65,85]. It is believed that the NO, which is released by the metabolism of nitrate as the compounds are broken down, may counteract the consequences of the NSAID-induced decrease in gastric mucosal PGs [65]. It seems important to mention that a clear identification of the metabolic steps by which NO-NSAIDs produce NO has not been established. Experimental findings obtained, in vivo, show that HCT1026 is metabolized into flurbiprofen and NO species, i.e. nitrates and nitrites, [60,61,70], which are detected in plasma and in brain at 2-4 h [70] and 3 h [86], respectively from drug administration. In vitro, HCT1026 is converted into flurbiprofen with different kinetics depending on the cell assay. In rat plasma, 30 min of incubation is required to fully convert HCT1026 to flurbiprofen [see [63]]. By contrast, approximately 35% of HCT1026 is converted into flurbiprofen within 1 h of incubation in human blood, and 24 h of incubation was necessary to reach the almost complete dissociation of HCT1026 [62,63].

Given the rapid action of HCT1026 demonstrated in vitro, Bernardo et al. [63] suggested the possibility that HCT1026 might reach the brain parenchyma and act on brain cells before being cleaved to the nitrate moiety and flurbiprofen [63]. Alternatively, the metabolites flurbiprofen and NO might act concomitantly by activating parallel pathways that ultimately determine the unique effects of HCT1026 [63]. Other studies have suggested a potential action of the HCT1026 metabolite, 4-hydroxybutyl nitrate, since animal studies have shown that the level of inorganic nitrite in the brain increases after oral administration of HCT1026 [86]. Plasma nitrite itself has been shown to provide a source of NO under certain conditions [see [87]].

Neurochemical, morphological and behavioral changes clearly indicate that with the ageing process, increased vulnerability of the nigrostriatal DAergic system and limited recovery from MPTP injury are observed [4,5,17,18,77,78]. Indeed, the nigrostriatal DAergic neurons exhibit compensatory mechanisms in response to MPTP injury, but the degree of plasticity becomes reduced with age [18,77,78]. Accordingly, a diminished compensatory capacity of nigrostriatal DAergic neurons “as a prelude” to PD is recognized to accompany the process of aging [see [88,89]]. Among the mechanisms at play, increased neuronal vulnerability to degenerative conditions, dysfunction of glia-neuron crosstalk, reduced repair capacity of injured DAergic neurons and/or limited neurogenesis may contribute to the poor recovery observed with age [see [18] for review]. Given the role of the ageing process as a critical risk factor for developing PD, we addressed the efficacy of HCT1026 preventive administration schedule in 9 to 11 month-old mice and found that long-term administration of HCT1026 resulted in a significant DAergic neuroprotection following MPTP insult, at both and SNpc levels. These effects lasted up to 40 dpt, supporting HCT1026 as promising approach towards the development of effective pharmacological neuroprotective strategies against PD.

Variable neuroprotective effects have been, so far, reported for both steroidal and non-steroidal, mixed and COX-2-selective inhibitors in different MPTP-mouse models of PD [37-48]. The mixed COX-1/COX-2 inhibitor indomethacin [39] and a COX-2 selective NSAID, rofecoxib [38] treatment rescued DAergic neurons from MPTP injury. Indomethacin, however, appeared toxic at high doses, and rofecoxib, failed to keep its protective properties when used in the prolonged treatment [90]. Studies on mice deficient with COX-2 showed that COX-2 plays a critical role in animal models of DAergic degeneration [38,32,43]. In particular, the role of increased levels of COX-2 in generating a toxic dopamine-quinone species which was responsible for DAergic neuronal degeneration, was demonstrated, whereas the selective COX-2 inhibitor, rofecoxib, exhibited a neuroprotective effect [38]. In the study of Sanchez-Pernaute [43], the COX-2 antagonist, celecoxib, was capable to prevent or slow down DAergic degeneration induced by intrastriatal administration of 6-OHDA. Accordingly, Vijitruth et al. [42] showed that pharmacological or genetic inhibition of COX-2 was capable to reduce motor impairment and to protect DAergic neuronal cell bodies in the SNpc as well as the striatal TH-stained fibres against MPTP-induced neurotoxicity [42].

On the other hand, ibuprofen, a non selective blocker, was shown to diminish the decline of dopamine content in striatum in the MPTP mouse model of PD, in a dose-dependent manner, and was not toxic to the DAergic system [41]. In accord with these experimental results, ibuprofen, but not other non-selective NSAIDs, was shown to diminish the risk/incidence of PD in men [5,6]. The present results showing the long-term DAergic neuroprotection in ageing mice and the safety profile of HCT1026 are of special interest, given that non-selective NSAIDs long-term therapies are hampered by their significant gastrointestinal, renal and cardiovascular side-effects [56,85].

HCT1026-induced neuroprotection observed in the present study was accompanied by a sharp downregulation of all studied markers of microglial activation including ameboid-like microglial cell number, pro-inflammatory mediators as well as two key harmful elements, MAC-1 and PHOX, likely suggesting that a shift from microglial pro-inflammatory (“armfull”) phenotype might be a major contributing factor.

Indeed, under inflammatory conditions, PHOX is the major source of peroxides in the brain. Activation of
microglial PHOX is synergistic with glial iNOS expression in inducing DAergic neuron death [27-36]. Accordingly, PHOX/Mac-1-deficiency mitigates MPTP-induced DAergic neurotoxicity both in vivo and in vitro [30-33]. Inflammatory stimuli together with ROS and RNS activate nuclear factor-κB (NF-κB) in microglial cells, oligodendrocytes and neurons to promote the transcription of inflammatory cytokines, COX-2, iNOS, and apoptosis-promoting factor including p53/Bax [see [45,46]]. Various studies (reported in Background section) have clearly underlined a “dual key mechanism” whereby simultaneous activation glial iNOS and PHOX synergistically act in killing DAergic neurons [33]. This mechanism may mediate inflammatory degeneration in response to cytokines, bacteria, ATP, arachidonate, whereas neuroprotection was observed by NO and peroxynitrite scavengers [33]. Indeed, nitritative stress is among the factors potentially underlying DAergic neurodegeneration [8,27,28,33-36]. Interestingly, the number of DAergic neurons containing 3-NT increased significantly in rhesus monkey midbrain DAergic neurons with age, suggesting a role for aging-related increase of nitritative damage in the selective vulnerability of SN neurons to degeneration in PD [91]. Here, ageing mice fed with a control diet exhibited a dramatic increase of 3-NT colocalization with TH+ neurons by 3 d after MPTP, corresponding to the active DAergic degeneration phase [27,28,32,33,36], while by 7 d a certain decrease in 3-NT accumulation within TH+ neurons was observed, possibly indicating the end of the degeneration phase. In HCT1026-fed mice, the generation of PHOX and iNOS-derived cytotoxic mediators, including 3-NT accumulation within DAergic neurons, were markedly abated, supporting that a significant proportion of SNpc neurons survived MPTP insult.

Although NSAIDs pharmacological actions are related to their ability to inhibit PG biosynthesis, some of their beneficial therapeutic effects are thought to be mediated by a panel of COX-independent mechanisms. NSAIDs are able to inactivate the transcription NF-κB and activator protein-1 (AP-1), critically involved in the induction of multiple inflammatory gene products involved in the inflammatory response (i.e. iNOS, TNF-α). In addition, NSAIDs in neuronal cells might directly and dose-dependently scavenge ROS and RNS, thereby blocking their detrimental effects [45,46]. On the other hand, high concentrations of NSAIDs such as ibuprofen and indomethacin, activate PPARγ. PPARγ is a ligand activated inhibitory transcription factor that antagonizes the activity of NFκB, AP1, signal transducer and activator of transcription-1 (STAT-1) and nuclear factor of activated T cells (NFAT). PPARγ activation is then associated with a reduction in the expression of several inflammatory genes and the production of inflammatory cytokines and iNOS [45,46]. In particular, in vitro studies have reported that selective PPARγ agonists such as pioglitazone, ibuprofen, or indomethacin, can activate PPARγ in microglia, decreasing the number of activated glial cells [see [45,46]].

The mechanisms that differentiate HCT-1026 from flurbiprofen remains a matter of debate. In vitro studies demonstrated that a low concentration of (1 μM) of HCT1026, but not flurbiprofen, activated PPARγ in primary cultures of rat microglia, with kinetics similar to those of the synthetic agonist, ciglitazone [63], supporting additional anti-inflammatory action through PPARγ [63]. In addition PPARγ agonists were reported to mitigate MPTP-induced DAergic neurotoxicity in different PD models [see [13-15,45,46]]. In the recent studies of Abdul-Hay et al. [87], flurbiprofen was 10-fold less potent than HCT-1026 in inhibiting iNOS induction in RAW 264.7 cell cultures. In LPS/IFNγ-induced primary astroglial cultures, HCT-1026 showed anti-inflammatory potency towards inhibition of cytokine and iNOS elevation, providing similar observations to those in microglial cultures [62]. That the anti-inflammatory activity of HCT-1026 could translate into neuroprotection was further demonstrated in a co-culture experiment with LPS-stimulated RAW cells and a neuroblastoma cell culture, where HCT-1026 was highly efficacious neuroprotectant [87].

Grafting a NO-donating moiety to flurbiprofen was reported to confer additional anti-inflammatory properties [51,52,60-66,87,92,93]. It has been suggested that this effect may depend on the negative feedback regulation exerted by low physiologic concentration of NO (nanomolar range) on different inflammatory mediators such as iNOS and COX-2, as well as on their associated functions [94-97]. Indeed, there is evidence that at low concentrations, NO has anti-inflammatory properties as it inhibits the expression of pro-inflammatory proteins (i.e. COX-2 and iNOS), and it counters the release of pro-inflammatory cytokines, such as TNF-α, in activated macrophages [94-97].

It should be recalled, that NO signaling plays an important role in the functioning of the CNS, and activation of soluble guanylate c elusive NO-donating moiety to flurbiprofen was reported to confer additional anti-inflammatory properties [51,52,60-66,87,92,93]. It has been suggested that this effect may depend on the negative feedback regulation exerted by low physiologic concentration of NO (nanomolar range) on different inflammatory mediators such as iNOS and COX-2, as well as on their associated functions [94-97]. Indeed, there is evidence that at low concentrations, NO has anti-inflammatory properties as it inhibits the expression of pro-inflammatory proteins (i.e. COX-2 and iNOS), and it counters the release of pro-inflammatory cytokines, such as TNF-α, in activated macrophages [94-97].

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Besides the NO-mediated effects, most recently proposed are NO-independent and NSAID-independent actions on NFκB and MAPK/ERK signaling pathways [see [64]]. In their study, Idris and co [64] reported the ability of HCT1026 to inhibit receptor activator of NFκB (RANKL), as well as RANKL-induced activation of NFκB and ERK pathway in LPS-stimulated macrophage cultures. In addition, HCT1026 also inhibited TNF-α, IL-1 and LPS-induced signaling. Interestingly enough, the pathways inhibited by HCT1026 all share a similar kinase complex upstream of the NFκB and ERK pathways and this is the most likely target for the action of HCT1026 [64].

It seems important to underline that inflammatory pathways may become hyperactivated with age and/or become more sensitive to immune/neurotoxic challenge, thereby promoting degeneration [99-102]. Given that with age, dysfunctional microglia and altered glia-neuron crosstalk may contribute to the progression of neuronal degeneration [18], HCT1026 preventive and long-term treatment might thus reduce age-dependent and MPTP-induced increase in oxidative and inflammatory attacks to nigrostriatal DAergic neurons. Of special interest, in view of the role of both systemic and central inflammation in modulating the severity of neuronal insult, including DAergic injury [see [13-15,18,24,81-83]], a potential effect of HCT1026 in influencing systemic inflammation cannot be excluded. In addition, the mitigation of the nitrosative/oxidative status of the nigral microenvironment as revealed by downregulation of Mac-1, PHOX and 3-NT in the VM, likely have beneficial consequences for glial expression of critical neuroprotective/neurotrophic factors [18], thereby supporting TH+ neuron survival/neurorescue, possibly through an amelioration/mitigation of SNpc microenvironment.

After brain injury, the inflammatory environment is recognized to have both detrimental and beneficial effects on neuronal outcome, depending on mouse strain, age and sex of the host, the severity of the lesion, the degree and timing glial activation, the hormonal background, the specific cellular context and intrinsic region-specific neuronal characteristics [see [8-18,21-26,40,77,78,100,103,104]]. In degenerative conditions, glia serve neuroprotective functions including the removal of dead cells by phagocytic activity and the production of neurotrophic factors. By contrast, overactivation of microglia or dysfunctional microglial cells as a consequence of ageing and age-related events within the SN microenvironment, [18,25,100-102] likely increase DAergic neuron vulnerability and/or may limit DAergic self-repair abilities. In vivo experiments have recently shown that intranigral administration of prostaglandin J2 (P2) induces microglia activation, selective degeneration of DAergic neurons in the SNpc, formation of ubiquitin- and α-synuclein-immunoreactive aggregates in the spared DAergic neurons, and locomotor deficit [83]. These and other findings have underlined the role of a transient initiation factor, triggering an active self-perpetuating cycle of chronic neuroinflammation, contributing to DAergic neuronal dysfunction [13,83]. By reducing exacerbation of inflammation, HCT1026 may then improve mitochondrial performance, increase glial-mediated neurotrophic support, thus creating a more favorable milieu for nigrostriatal DAergic neuron survival/rescue.

The present results are in line with data obtained in different animal models of brain inflammation, where HCT1026 significantly reduced neuronal loss and decreased the number of reactive microglial cells to a greater extent than the parent compound, flurbiprofen [66,86,87,92,93]. In experimental allergic encephalomyelitis (EAE), oral treatment with HCT1026 which delayed disease onset and decreased the severity of clinical signs in mice immunized with myelin oligodendrocyte peptide (MOG35-55) [66]. In addition, HCT1026 fed mice exhibited significantly reduced mRNA levels of pro-inflammatory cytokines, caspase-1, and iNOS in blood cells, with reduced number of CNS-infiltrating T cells [66]. Recently, HCT1026 was reported to mitigate amyloid-β-induced toxicity, in cell culture, in vitro, while enhancing cognition in response to cholinergic blockade, in vivo [87]. Other studies reported the ability of HCT1026 to reduce microglia activation and to prevent muscular dystrophy pathology in two murine models [93].

Conclusions

We herein report that oral preventive administration of the NO-donating derivative of flurbiprofen, HCT1026, has a safe profile and a significant efficacy in counteracting MPTP-induced DAergic neurotoxicity, motor impairment and microglia activation in ageing mice. In particular, nigrostriatal DAergic neuroprotection afforded by HCT1026 lasted for 40 d after MPTP administration. Hence, DAergic neurons exhibited an increased ability to resist to the cytotoxic environment caused by MPTP injury, leading to a significant neurorescue observed within the striatum and SNpc, at a morphological, neurochemical, and molecular levels. These effects of HCT1026 were associated with reduced microglial proinflammatory phenotype and reduced formation of the peroxynitrite footprint, 3-NT, within TH+ cell bodies. While further studies are required to clarify the mechanism(s) of HCT1026 neuroprotective effects, the combination of a balanced inhibition of the two main COX isoforms with NO release provides a promising approach towards the development of novel and effective therapeutic strategies against PD.
List of abbreviations
(3-NIT): 3-nitrotyrosine; (CDX-2): Cyclooxygenase-2; (DA): Dopamine; (DAT): Dopamine transporter; (EAE): Experimental Allergic Encephalomyelitis; (GR): Glucocorticoid receptor; (HCT1026): [2-fluoro-α-methyl-(1β-biphenyl)-4-acetic-4-(nitroxy)-butyl ester]; (iNOS): Inducible-nitric oxide synthase; (LPS): Lipopolysaccharide; (Mac-1): Macrophage antigen-1; (MAO-B): Monoamine oxidase B; (MOG35-55): Myelin oligodendrocyte peptide; (MPP+): 1-methyl-4-phenylpyridinium ion; (MPTP): 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine; (NO): Nitric oxide; (NSAIDs): Nonsteroidal anti-inflammatory drugs; (OXO): Peroxynitrite; (PD): Parkinson’s disease; (PGs): Prostaglandins, (PHOX): NADPH oxidase; (P2): Prostaglandin J2; (PPAR-γ): Peroxisome proliferator-activated receptor-γ; (RNS): Reactive nitrogen species; (ROS): Reactive oxygen species; (SNpc): Substantia nigra pars compacta; (TH): Tyrosine hydroxylase; (VM): Ventral midbrain; (WB): Western blot.

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Authors’ contribution
FL participated in the design and Ms editing, in all treatment procedures, data elaboration, gene expression and behavioural studies. CT did the immunohistochemical procedures and confocal image analyses, carried out most of the histological and stellogical estimations and graphic representations. SC participated in all treatment procedures and tissues processing for western blot analyses and elaboration of the data. NT participated in all treatment procedures and daily control for animal foodintake, weights and toxicity control at sacrifice, and participate to immunohistochemical procedures and data elaboration. P-A Serra and MC. Morale were responsible for protocol design and for analyses of stutal neurochemistry. F. Impaginatello participated in protocol design and HCT1026 measurements. BM conceived the study and design, analysed the data and prepared the manuscript. All authors read, discussed and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

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