Effect of Inflammatory Stress on Neural Stem Cells Behaviour: A Rescue Effect of Phosphatidylcholine by Modulating Neuronal Plasticity.

Dario Magaquian  
Instituto de Biologia Molecular y Celular de Rosario (IBR,CONICET) Ocampo y Esmeralda and Facultad de Cs. Bioquímicas y Farmacéuticas, Universidad Nacional de Rosario, Rosario, Argentina.

Susana Delgado Ocaña  
Instituto de Biologia Molecular y Celular de Rosario (IBR,CONICET) Ocampo y Esmeralda and Facultad de Cs. Bioquímicas y Farmacéuticas, Universidad Nacional de Rosario, Rosario, Argentina.

Consuelo Perez  
Instituto de Biologia Molecular y Celular de Rosario (IBR,CONICET) Ocampo y Esmeralda and Facultad de Cs. Bioquímicas y Farmacéuticas, Universidad Nacional de Rosario, Rosario, Argentina.

Claudia Banchio (endars@ibr-conicet.gov.ar)  
Instituto de Biologia Molecular y Celular de Rosario (IBR,CONICET) Ocampo y Esmeralda and Facultad de Cs. Bioquímicas y Farmacéuticas, Universidad Nacional de Rosario, Rosario, Argentina.

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Abstract

The balances between NSCs growth and differentiation, and between glial and neuronal differentiation play a key role for brain regeneration after any pathological conditions. It is well known that the nervous tissue shows a poor recovery after injury due to the factors present in the wounded microenvironment, particularly inflammatory factors, that prevent neuronal differentiation. Thus, it is essential to generate a favourable condition for NSCs and conduct them to differentiate towards functional neurons. Here, we show that neuroinflammation has no effect on NSCs proliferation but induces an aberrant neuronal differentiation that gives rise to dystrophic, non-functional neurons. This is perhaps the initial step of brain failure associate to many neurological disorders. Interestingly, we demonstrate that phosphatidylcholine (PtdCho)-enriched media enhances neuronal differentiation even under inflammatory stress by modifying the commitment of post-mitotic cells. The pro-neurogenic effect of PtdCho increases the population of healthy normal neurons. In addition, we provide evidences that this phospholipid ameliorates the damage of neurons and, in consequence, modulates neuronal plasticity. These results contribute to our understanding of NSCs behaviour under inflammatory conditions, opening up new venues to improve neurogenic capacity in the brain.

Introduction

Despite its diverse presentation, inflammation is a common feature across several neuropa-thological processes and has been implicated as a critical mechanism responsible for the progression of neurodegenerative disorders including Parkinson’s disease, Alzheimer’s disease multiple sclerosis\(^1,2,3\) as well as traumatic brain injury\(^4,5\) and stroke\(^2,6,7\). Neuroinflammation is considered a double-edged sword, with protective as well as detrimental effects on the nervous system, especially during repair and recovery. In response to different types of injuries that cause neurons and oligodendrocytes death, activated astrocytes and the resident immune-like glial cells, the microglia, proliferate and generate proinflammatory cytokines (such as IL-1, IL-6, IFN-\(\gamma\) and TNF-\(\alpha\)), chemokines, prostaglandins, and free oxygen radicals, often leading to the development of cerebral damage, and promoting macrophages infiltration\(^8\). Both kinds of cells act as a host defence mechanism eliminating cellular debris and releasing inflammatory factors. These factors finally activate astrocytes, which proliferate and form the glia scar to define a dense limiting border between the healthy and damaged tissue. Two major niches of neural stem cells (NSCs) that support neurogenesis are in the subventricular zone and in the dentate gyrus of the hippocampus of the adult mammalian brain\(^9,10\). NSCs are multipotent self-renewing cells that have a regenerative potential because they can proliferate, migrate and differentiate into neurons, astrocytes or oligodendrocytes and thus, promote functional and structural repair of the injured tissue. Several studies have evidenced a cross-talk between immune modulators and NSCs fate\(^11,12,13\). In response to inflammatory reactions, it was shown that the glia scar could prevent tissue regeneration by NSCs\(^8\), and that LPS-induced neuroinflammation caused synapse loss by a mechanism dependent of microglia activation and IL-1\(\beta\) secretion\(^14\). In this scenario, understanding the NSCs response to these
conditions and mechanisms involved in the integration into the injured brain will be critical for the
development of effective therapeutic strategies using stem cells.

We have previously demonstrated that phospholipids affect the fate of post-mitotic neural precursors;
specifically, phosphatidylcholine (PtdCho) promotes neuronal differentiation at expenses of astroglial
and unspecified precursors\textsuperscript{15}. As the loss of neurons is the detrimental outcome of brain injuries and
neurodegenerative diseases, we asked whether PtdCho could still favour neuronal differentiation under
inflammatory conditions, and thus prevent or restore tissue damage in this context. By different
approaches we have demonstrated that under pro-inflammatory culturing conditions there is an increase
in neuronal differentiation that could support a renewal mechanism needed for tissue reparation.
Strikingly, under the same conditions, neurons also display an aberrant morphology that could reflect the
deleterious effect of neuroinflammation. Interestingly, addition of liposome of egg-source PtdCho further
induces neuronal differentiation and also rescues the morphological and functional deficit by modulating
neuronal plasticity.

Results

Effect of inflammatory stress on NSCs proliferation

The balance between NSCs proliferation and differentiation is essential for tissue repair\textsuperscript{16}, and up to
know is not clear how it is affected by inflammation. To that end, we incubated NSCs under normal
proliferative condition (in the presence of EGF and FGF, neurosphere culture) supplemented with different
concentrations (% V/V) of macrophages-activated media (AM) or with media without activation as a
control (UM). We confirmed by RT-PCR that macrophages activated with LPS express and, as a
consequence, secrete IL-1\textbeta, IL-6 and TNF-\textalpha to the media as previously demonstrated\textsuperscript{17,18}
(Supplementary figure 1A). Treated cells were incubated for 96 hours, and after this time, cell viability was
analysed by MTT assay; UM plus LPS was also included to evaluate LPS toxicity as control
(Supplementary figure 1B). To induce a moderate stress, cells were treated with AM (20% V/V) and, after
96 hours, the neurosphere's diameter was measured as a growth parameter (Figure 1B). As Figures 1A, B
and D show, there is no significant change in the proliferation rate in the presence of different
concentrations (V/V) of activated media respect to media without activation and control. To evaluate the
role of each IL individually, similar analyses were performed in UM supplemented with IL-1\textbeta and/or IL-6.
The concentrations were determined by assaying cell viability by MTT (Supplementary figure 1C). As
Figures 1C and D show, these ILs, as components of the inflammatory condition, do not affect the
capacity of NSCs to proliferate as no changes were detected in the neurosphere's diameter under the
assayed conditions.

Effect of inflammatory stress on neuronal differentiation of
NSCs
Neuronal differentiation is key in neural tissue regeneration after injuries\textsuperscript{19}. To investigate this process under pro-inflammatory conditions, we analysed neuronal differentiation by immunocytochemistry using βIII-tubulin as neuronal lineage marker. Cells were incubated for 72 hours in media supplemented with macrophages-activated media (AM-20% V/V), media without LPS activation (UM-20% V/V) or control media. The quantification analysis demonstrated that in the presence of AM there is a significant increase in the percentage of neurons in comparison with the UM or the control (Figure 2A). Similar results were observed at shorter time points (Supplementary figure 2). However, a morphological observation revealed that neurons incubated with AM display a different shape than neurons at the control or neurons incubated with UM, with apparent tubulin disorganization, dystrophy with increased soma size (Figure 4) and presence of vacuoles\textsuperscript{20,21}. We therefore hypothesized that neuronal differentiation could be aberrant under pro-inflammatory stress, leading to cell dystrophy. To determine whether the observed effect is a direct action of LPS or the inflammatory components (IL-1β and IL-6), we evaluated neuronal differentiation in the presence or absence of each molecule individually. As Figure 3 shows, treatment with UM supplemented with the indicated concentration of LPS, IL-1β and/or IL-6 did not affect the rate of neuronal differentiation nor the morphology of the neurons.

**Phosphatidylcholine enhances neuronal differentiation and ameliorates neuronal alterations caused by inflammatory conditions**

We have previously demonstrated that PtdCho, as liposomes supplemented in the media, regulates the fate of post-mitotic precursor cells, inducing neurogenesis\textsuperscript{15}. To test the effect of this molecule under pro-inflammatory stress conditions, we incubated NSCs under each condition in the presence of egg-source PtdCho (50 μM) and counted the resulting βIII-tubulin expressing cells. As shown in Figure 2A, the pro-neurogenic effect of PtdCho shown in normal and UM control conditions, is also observed under inflammation, reaching the highest levels of neuronal differentiation. Interestingly, the aberrant phenotype observed in AM was ameliorated when cells were incubated in the presence of PtdCho (Figure 2B, lower panel). A detailed morphometric analysis demonstrated that the soma size was restored in the presence of PtdCho (Figure 4A and B). More interestingly, the percentage of dystrophic neurons, decreases with PtdCho treatment, resulting in a significant increase in the phenotypically normal neurons (Figure 4B) and suggesting that under inflammatory conditions, PtdCho not only regulates the fate of post-mitotic cells increasing neuronal differentiation\textsuperscript{15}, but also rescues the dystrophic neurons, turning its morphology back to the normal.

**PtdCho restores synaptic defect caused by inflammation**

We next examined whether the observed defect in neuronal morphology is associated with alteration in neuronal functionality. The amount and localization of synaptophysin protein plays a critical role in
synapse formation\textsuperscript{22, 23}, exo-endocytosis of synaptic vesicles\textsuperscript{24}, neural plasticity\textsuperscript{25, 26}, memory\textsuperscript{27}, motor development, behavioural features and cognitive impairments\textsuperscript{28}. Neurons were immunostained for the presynaptic protein synaptophysin, and the relative number of signal puncta on neurites was counted and analysed by western blot. As Figure 5 shows, incubation with AM clearly decreases the number of synaptophysin-containing vesicles and its levels of expression relative to the control. Interestingly, incubation with PtdCho restores the level of expression and location of synaptophysin under inflammatory conditions. This result clearly suggests that PtdCho has two effects: induces neurogenesis and improves function of sick/damaged neurons.

We have previously demonstrated that lipid treatment 24 hours after plating the cells did not affect neuronal differentiation, indicating a narrow time-window of response in post-mitotic cells\textsuperscript{15}. To confirm the effect of PtdCho independent of the promotion of neurogenesis, we quantified the percentage of neurons and the morphology adding PtdCho 24 hours post inflammatory condition. As expected, PtdCho does not increase the percentage of neurons (Figure 6A)\textsuperscript{15}, but clearly altered the balance between healthy/normal neurons and dystrophic, pushing it to the normal population (Figure 6B).

**Discussion**

NSCs have a fundamental role after nervous tissue damage as they have the potential for regeneration owing to their capacity of self-renewal and differentiation into neurons\textsuperscript{8, 29}. However, this extraordinary capacity is limited under pathological conditions due to the factors present in the wounded microenvironment that can affect NSCs survival, proliferation and differentiation\textsuperscript{30, 31, 32, 33}.

In this report, we provide details of the NSCs behaviour under inflammatory condition, a common scenario of many acute and chronic brain diseases. Furthermore, we provide evidence that PtdCho treatment could target NSCs conducting them towards functional neurons and also restoring the morphological deficit caused by inflammation.

Neuroinflammation can either affect the niche or the NSCs directly, with the end result of altered NSCs proliferation and/or differentiation\textsuperscript{1, 34}. We demonstrated that after incubation of NSCs with AM or with individual cytokines, cell viability (Supplementary Fig. 1) and the rate of NSCs proliferation are not affected (Fig. 1). These results differ from previous demonstrating that proinflammatory cytokines reduce the number of new born neurons in the dentate gyrus in adult mice due to the restrain of the cell cycle\textsuperscript{35}. As NSCs proliferation depends on the cell progression, we assumed that under our study condition cell cycle progression is not affected. In addition, the observed discrepancy could base on the different origin of the NSCs utilized.

The in deep study of the cellular mechanism leading to neuronal dysfunction under inflammatory condition is essential for the development of novel therapies. These experiments demonstrated that incubation of NSCs with 20\% V/V of AM, but not with ILs individually (Fig. 3), induces aberrant neuronal differentiation, that give rise to dystrophic neurons (Fig. 2). The relatively constant number of neurons in
AM-treated cultures during different periods of times (day 1 to day 3) (Supplementary Fig. 2) suggests that AM does not affect specific step of neuronal differentiation process. Rather, it seems to be a very dynamic sequence of morphological changes with a constant progression to dystrophic morphology (Fig. 2 and Supplementary Fig. 2). It is well known that LPS activates microglia, and the consequent ILs secretion affects neuronal differentiation\textsuperscript{36,37,38}, we discard this effect as ILs and LPS alone did not affect neuronal differentiation of NSCs, nor the morphology of the neurons (Fig. 3).

Currently, lipids are taking a leading role in the nervous system. They have been shown to intervene in cellular functions such as proliferation, differentiation, cell cycle and act as pro-resolution lipid mediators in inflammatory events\textsuperscript{15,39,40,41,42}. As this maladaptive neuronal plasticity that takes place under inflammation could be the reason of many brain failures, we evaluated the effect of PtdCho on NSCs differentiation. We demonstrated that PtdCho induces neuronal differentiation under inflammatory condition increasing the percentage of healthy non-dystrophic neurons (Fig. 4). Therefore, PtdCho changes the fate of post-mitotic cells increasing neurogenesis by turning on the PKA/CREB signalling pathway even under inflammatory conditions (AM). In fact, the percentage of βIII-tubulin positive cells decrease in the presence of PKA inhibitor (KT5720) (Supplementary Fig. 3). This specific effect of PtdCho could favour and increase the replacement of damaged neurons favouring NSCs-dependant neurogenesis. More interestingly, this phospholipid ameliorates the damage of neurons and, in consequence, modulates neuronal plasticity: in fact, treatment with PtdCho even 24 hours post inflammatory condition, restores the soma diameter and increases synaptophysin expression and distribution in neurons (Figs. 4 and 5). Hence, it decreases the amount of dystrophic neurons by a mechanism independent of PKA activity and NSCs differentiation (Fig. 6 and Supplementary Fig. 3). This capacity to modulate neuronal plasticity was also described for choline in the treatment of Rett Syndrome\textsuperscript{43}.

In this stage, a repeated question arises: is choline\textsuperscript{43}, CDP-Choline\textsuperscript{44} or PtdCho\textsuperscript{15} the key molecule for neuronal plasticity? Even though, choline has three main contributory roles in maintaining the cellular physiology in neurons: as precursor for the synthesis of the neurotransmitter acetylcholine\textsuperscript{45}; as a key donor for methylation of DNA and regulation of gene expression\textsuperscript{46}; and for the synthesis of PtdCho\textsuperscript{47}, we propose that choline and CDP-Choline regulate the fate of NSCs and induces neurogenesis by its conversion into PtdCho. Experiments done in the presence of Hemicholineum-3 (a choline kinase inhibitor)\textsuperscript{48} or Edelfosine\textsuperscript{42} demonstrated that choline is unable to promote neuronal differentiation when the Kennedy pathway was blocked (Supplementary Fig. 3); however, PtdCho activates neurogenesis even under inhibition of these enzymes. This results clearly demonstrate that choline need to be converted in PtdCho to affect the fate of NSCs.

In conclusion, more research need to be done to understand the molecular mechanism of PtdCho as modulator of neuronal plasticity, but, considering that loss and damage of neurons are the major consequence of acute and chronic neuroinflammation, these results might open a door to develop new therapeutic approaches.
Methods

Chemicals and antibodies.

Dulbecco's modified medium/Ham's F12 (DMEM/F12 1:1), Dulbecco's Modified Eagle's Medium (DMEM), B27 and anti-rabbit Alexa Fluor® 488-labeled were purchased from Life Technologies Corporation (Carlsbad, CA, USA). Fetal bovine serum (FBS) from Internegocios (Buenos Aires, Argentina). Rabbit anti-β-Tubulin III antibody from Sigma (St. Louis, MO, USA), mouse anti-synaptophysin, mouse anti-β-Actin and KT5720 from Santa Cruz (Dallas, Texas, USA) and anti-mouse Cy3-labeled from Millipore (Massachusetts, USA). Quick-Zol from Kalium (Buenos Aires, Argentina). RNase-free RQ1 DNase enzyme, Reverse Transcriptase enzyme M-MLV from Promega (Wisconsin, USA). TAQ polymerase buffer, dNTPs and TAQ polymerase from TransGen Biotech (Beijing, China). Protease inhibitor cocktail, poly-D-lysine (PDL), epidermal growth factor (EGF), human basic fibroblast growth factor (bFGF), lipopolysaccharide (LPS) and Phosphatidylcholine (P3556) from egg yolk source were from Sigma (St. Louis, MO, USA). As specified in product information, they have a purity over 99% and a fatty acid content of approximately 33% palmitic, 13% stearic, 31% oleic, and 15% linoleic. In addition the detailed fatty acid composition of the mixture of egg yolk phosphatidylcholine and phosphatidylethanolamine has been recently described.

Animal studies and fetal neural stem cells culture.

All animal experiments and related experimental protocols were approved by the Bioethics Commission for the Management and Use of Laboratory Animals of National University of Rosario, Argentina (N 6060/89). All procedures were carried out in accordance with the approved guidelines (Guide for the care and use of Laboratory Animals- 8° edition-e National Academies Press-Washington DC 2011 and in compliance with the ARRIVE guidelines). Time pregnant female C57/BL6 mice (gestation day 13) were sacrificed by cervical dislocation under supervision of the Animal Care and Use Committee. Neurospheres were obtained from E13 cortical cells as previously described. Briefly, the lateral portion of the dorsal telencephalon (cortex) of embryonic day 13 mouse C57/BL6 was isolated. The cortices were chemically disrupted adding trypsin (0.05% w/v) for 5 minutes and then mechanically disrupted into single cells by repeated pipetting in medium DMEM/F12 (1:1) containing 10% FBS, penicillin G (100 units/ml) and streptomycin (100 μg/ml). Cells were centrifuged at 1000 rpm for 5 min and the pellet resuspended in serum-free medium DMEM/F12 (1:1). Dissociated cells were cultured at a density of 5 × 10^4 cells/ml in medium DMEM/F12 (1:1) supplemented with B27, 10 ng/ml bFGF and 10 ng/ml EGF, at 37 °C in a humidified 5% CO₂ incubator. Within 7 days, cells grew as free coating neurospheres that were then collected by centrifugation, and chemically and mechanically dissociated to obtain a new passage. For cell differentiation, neurospheres were chemically and mechanically dissociated. After counting, 2.5 × 10^5 cells were plated on poly-D-lysine (PDL) (10 μg/ml)-coated 24 well plates, or 5 × 10^4 cells were plated on PDL (10 μg/ml)-coated 96 well plates in medium DMEM/F12 (1:1) supplemented with B27.
Macrophages culture and LPS-induced stimulation.

The mouse cell line Raw 264.7 (ATCC® TIB-71™) was cultured in DMEM 10% FBS supplemented with penicillin G (100 units/ml), streptomycin (100 µg/ml) (proliferation conditions) and maintained in a 5% CO₂ humidified incubator at 37 °C. For activation, cells were grown to 80% confluence in petri dishes with DMEM medium supplemented with 10% FBS. At this time, cells were centrifuged at 1500 rpm for 10 minutes. The cell pellet was resuspended in 1 ml of DMEM/F12 medium and cells were transferred to a new plate containing DMEM/F12 stem cell medium (in the absence of FBS, B-27 and growth factors). For stimulation, pure LPS was added in a final concentration of 1 µg/ml. After 18 hours of incubation, cells were centrifugated at 1000 rpm for 5 minutes and the culture medium was filtered through 0.22 µm filters (Sartorius) and stored immediately at -80 °C.

Total RNA isolation, DNase treatment and retrotranscription reaction

Murine macrophage RAW 264.7 total RNA was extracted in Quick-Zol (Kalium) following the supplier's specifications. Briefly, cells were resuspended in 1 ml Quick-Zol and incubated for 5 minutes at room temperature. Then, 0.2 ml of chloroform was added and they were centrifuged at 12,000 rpm for 10 minutes at 4 °C. Then, the aqueous phase was transferred to a new tube, 0.5 ml of isopropanol was added and the samples were incubated for 24 hours at -20 °C. The following day, they were centrifuged at 12,000 rpm for 10 minutes at 4 °C, the pellet was washed with 75% ethanol and centrifuged again at 12,000 rpm for 5 minutes at 4 °C. To evaluate the quality and quantity of the RNA obtained, absorbance measurements were made at 230, 260 and 280 nm (NanoVue Plus, General Electrics). Next, 1 µg of RNA was seeded on a 1.8% agarose gel to evaluate its integrity. To remove DNA from the samples, 10 µg of the RNA / DNA mixture was treated for 2 hours with the RNase-free RQ1 DNase enzyme (Promega) at 37 °C. The reaction was then stopped by incubating the sample at 65 °C for 10 minutes. Once the treatment was completed, the RNA concentration was determined spectrophotometrically (NanoVue Plus, General Electrics). For the reverse transcription reaction, 2 µg of RNA was treated with the Reverse Transcriptase enzyme M-MLV (Promega) strictly following manufacturer's instructions. For a final volume of 25 µl: oligo dT 20 ng/µl, dNTPs 0.5 mM each, buffer M-MLV 5 X, RNase inhibitor 0.8 U/µl and M-MLV Reverse Transcriptase 8 U/µl were added.

Polymerase chain reaction (PCR)

The PCR reactions were performed in the following buffer for a final volume of 50 µl: 1X TAQ polymerase buffer, 3 mM MgCl₂, 25 µM dNTPs each, 0.4 pmol/µl oligonucleotides each, 0.2 U/µl of TAQ polymerase (Easy TAQ, TransGen Biotech). Gene Amp (Parkin-Elmer, Shelton, CT, USA) or My Cycler (BioRad, USA) thermal cyclers were used. The oligonucleotides sequences (5'-3') used are: IL-1β Forward TTCAGGCAGGCAGTATCCTC, IL-1β Reverse GAAGGTCCACGGGAAAGACAC; IL-6 Forward
TAGTCCTCCTACCCCAATTTCC, IL-6 Reverse TTGGTCCTTAGCCACTCCTTC; TNF-α Forward CCTGTAGCCCACGTCTAG, TNF-α Reverse GGGAGTAGACAAGGTACAACCC. The amplification reaction started with an initial denaturation for 10 min at 94 °C followed by 35 cycles of denaturation at 94 °C for 1 min, annealing at 60 °C for 30 s and elongation at 72 °C for 30 s. The last cycle was followed by a 10-min extension step at 72 °C. The amplified products were analyzed by ethidium bromide-stained agarose gel electrophoresis.

Cell viability and proliferation assays

Cell viability was assessed by MTT-reduction assay. After cell treatment, MTT (5 mg/ml) was added to the cell culture medium at a final concentration of 0.5 mg/ml and incubated for 4 hours at 37°C, 5% CO$_2$. The assay was stopped by replacing the MTT-containing medium with DMSO. The extent of MTT reduction was measured spectrophotometrically at 570 nm$^{53}$. Results are expressed as a percentage of the control.

Proliferation of NSCs was assayed by measuring neurosphere’s diameter$^{54}$. Briefly, 5000 living cells were seeded per well in 24-well plates and cultured for up to 96 hours to evaluate the expansion rates. Size of 100 neurospheres (expressed as neurosphere’s diameters) was measured in three independent experiments. Images were taken with a microscope Olympus BH-2 and analysed using the freeware image J (National Institutes of Health, freeware).

Liposome preparation

Concentrated lipid stocks were prepared as previously described$^{55}$. Briefly, pure lipids were diluted in chloroform and dried in acid-washed glass centrifuge tubes under a stream of nitrogen. Phospholipid samples were suspended at 2–6 mM in phosphate-buffered saline at pH 7.2 and sonicated twice for 5 min at power setting 0.2–0.5% amplitude. All samples were sterilized with 0.22 μm-pore filters (Sartorius). The recovery of phospholipids after filtration was typically 90% or more.

Immunocytochemistry

Cells were cultured on PDL (10 μg/ml)-coated glass coverslips in 24-well plates as previously described$^{15}$. After different time of incubation, cells were fixed in 4% (w/v) paraformaldehyde-sucrose for 30 min at room temperature, permeabilized with 0.2% Triton X100 and blocked for 1 hour in 5% BSA. Cells were incubated with the primary antibody overnight at 4 °C followed by incubation with the fluorescently labelled secondary antibody for 1 hour at room temperature. Primary and secondary antibodies were diluted as follows: rabbit anti-βIII-tubulin (1:500) mouse anti-synaptophysin (1:300), anti-rabbit Alexa Fluor® 488-labeled (1:500) and anti-mouse Cy3-labeled (1:300). To visualize nuclei, cells were
counterstained and mounted with ProLong Gold antifade reagent containing DAPI (Molecular probes, Life technologies).

Microscopy and Image Analysis

Micrographs were acquired using a confocal microscope (Zeiss LSM 880) or the Nikon Model Eclipse 800 microscope and quantitative analyses were performed with Image J” (NIH). Cells were counted from twenty randomly selected fields per well for each individual experiment. At least three independent experiments were performed. The percentage of neuronal cell population was calculated against the DAPI-positive total cell number which includes undifferentiated stem cells and differentiated neurons. Cells bearing at least one neurite equal or longer than the soma diameter were considered to be differentiated. Soma size and number of synaptophysin containing vesicles were measured and counted manually. To differentiate dystrophic and normal neuronal populations, 100 neurons from each experiment were manually selected and analysed.

Western blot analysis

Western blot experiment was developed following protocols previously described\textsuperscript{15,42}. Neurosphere-derived cells were plated at a density of $1.5 \times 10^5$ and cultured on PDL-coated 35 mm culture dishes under differentiation conditions. Three days later, cells were collected, resuspended in lysis buffer (50 mM Tris-HCl pH 8.0, 50 mM KCl, 10 mM EDTA, Nonidet P-40 1%, 20 mM NaF, 1 mM Na$_3$VO$_4$, 1 mM PMSF and 1:1000 protease inhibitor cocktail) and sonicated five times at 5% amplitude for 5 s (Sonics and Materials Inc–Vibra CellTM). Protein concentration was determined using bovine serum albumin (BSA) as standard protein and “PierceTM BCA Protein Assay Kit (Thermo Scientific)“. 20 μg of cell lysate were resolved on 12% SDS-polyacrilamide gel electrophoresis (PAGE) and transferred to a nitrocellulose membrane (Amersham, GE Healthcare). After blocking overnight with 5% non-fat milk in 0.1% Tween TBS and washing, blots were incubated with anti-synaptophysin (1:300) overnight at 4 °C. Peroxidase-conjugated anti-mouse IgG (1:8000, Jackson Immuno Research) was used as secondary antibody. For loading protein control anti-β-Actin (1:6000) was used and developed with secondary antibody peroxidase-conjugated anti-mouse IgG (1:8000, Jackson Immuno Research). Labelled proteins were detected with chemiluminescence reagents (AmershamTM ECLTM Prime Western Blotting Detection Reagent, GE Healthcare).

Statistical analysis

Data represent the mean value ± SEM of at least three independent experiments and each individual experiment was performed in technical triplicate. Statistical significance was determined by either Student’s t-test or One-Way ANOVA followed by Tukey’s test using Prism (GraphPad Software Inc.). p-values lower than 0.05 were considered statistically significant.
Declarations

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Author Contributions

D.M., S.D.O. and C.P. performed all the experiments and analysed data; C.B. designed and supervised research, acquired funding and wrote the manuscript. All authors have read and agreed to the published version of the manuscript.

References

1. Covacu, R. & Brundin, L. Effects of Neuroinflammation on Neural Stem Cells. The Neuroscientist: a review journal bringing neurobiology, neurology and psychiatry, 23, 27–39 (2017).
2. Kaur, D., Sharma, V. & Deshmukh, R. Activation of microglia and astrocytes: a roadway to neuroinflammation and Alzheimer's disease., 27, 663–677 (2019).
3. Skaper, S. D., Facci, L., Zusso, M. & Giusti, P. An Inflammation-Centric View of Neurological Disease: Beyond the Neuron. Frontiers in cellular neuroscience, 12, 72 (2018).
4. McKee, A. C. & Daneshvar, D. H. The neuropathology of traumatic brain injury. Handbook of clinical neurology, 127, 45–66 (2015).
5. Dinet, V., Petry, K. G. & Badaut, J. Brain-Immune Interactions and Neuroinflammation After Traumatic Brain Injury. Frontiers in neuroscience, 13, 1178 (2019).
6. Dong, R., Huang, R., Wang, J., Liu, H. & Xu, Z. Effects of Microglial Activation and Polarization on Brain Injury After Stroke. Frontiers in neurology, 12, 620948 (2021).
7. Toman, N. G., Grande, A. W. & Low, W. C. Neural Repair in Stroke. Cell transplantation, 28, 1123–1126 (2019).
8. Xiao, L., Saiki, C. & Ide, R. Stem cell therapy for central nerve system injuries: glial cells hold the key. Neural regeneration research, 9, 1253–1260 (2014).
9. Riquelme, P. A., Drapeau, E. & Doetsch, F. Brain micro-ecologies: neural stem cell niches in the adult mammalian brain. Philosophical transactions of the Royal Society of London Series B, Biological sciences, 363, 123–137 (2008).
10. Fuentelalba, L. C., Obernier, K. & Alvarez-Buylla, A. Adult neural stem cells bridge their niche. Cell stem cell, 10, 698–708 (2012).
11. Carpentier, P. A. & Palmer, T. D. Immune influence on adult neural stem cell regulation and function., 64, 79–92 (2009).
12. Kokaia, Z., Martino, G., Schwartz, M. & Lindvall, O. Cross-talk between neural stem cells and immune cells: the key to better brain repair? Nature neuroscience, 15, 1078–1087 (2012).
13. Bonzano, S. et al. Neuron-Astroglia Cell Fate Decision in the Adult Mouse Hippocampal Neurogenic Niche Is Cell-Intrinsically Controlled by COUP-TFI In Vivo. Cell reports, 24, 329–341 (2018).
14. Sheppard, O., Coleman, M. P. & Durrant, C. S. Lipopolysaccharide-induced neuroinflammation induces presynaptic disruption through a direct action on brain tissue involving microglia-derived interleukin 1 beta. Journal of neuroinflammation, 16, 106 (2019).
15. Montaner, A. et al. Specific Phospholipids Regulate the Acquisition of Neuronal and Astroglial Identities in Post-Mitotic Cells. Scientific reports, 8, 460 (2018).
16. Hardwick, L. J., Ali, F. R., Azzarelli, R. & Philpott, A. Cell cycle regulation of proliferation versus differentiation in the central nervous system. Cell and tissue research, 359, 187–200 (2015).
17. Bailey, L. O., Washburn, N. R., Simon, C. G. Jr., Chan, E. S. & Wang, F. W. Quantification of inflammatory cellular responses using real-time polymerase chain reaction. Journal of biomedical materials research Part A, 69, 305–313 (2004).
18. Han, S. et al. Procyanidin A1 Alleviates Inflammatory Response induced by LPS through NF-kappaB, MAPK, and Nrf2/HO-1 Pathways in RAW264.7 cells. Scientific reports, 9, 15087 (2019).
19. Wei, Z. Z. et al. Regulatory role of the JNK-STAT1/3 signaling in neuronal differentiation of cultured mouse embryonic stem cells. Cellular and molecular neurobiology, 34, 881–893 (2014).
20. Grace, E., Rabiner, C. & Busciglio, J. Characterization of neuronal dystrophy induced by fibrillar amyloid beta: implications for Alzheimer's disease., 114, 265–273 (2002).
21. Wu, H. Y. et al. Amyloid beta induces the morphological neurodegenerative triad of spine loss, dendritic simplification, and neuritic dystrophies through calcineurin activation. The Journal of neuroscience: the official journal of the Society for Neuroscience, 30, 2636–2649 (2010).
22. Tarsa, L. & Goda, Y. Synaptophysin regulates activity-dependent synapse formation in cultured hippocampal neurons. Proceedings of the National Academy of Sciences of the United States of America, 99, 1012–1016 (2002).
23. Kwon, S. E. & Chapman, E. R. Synaptophysin regulates the kinetics of synaptic vesicle endocytosis in central neurons., 70, 847–854 (2011).
24. Valtorta, F., Pennuto, M., Bonanomi, D. & Benfenati, F. Synaptophysin: leading actor or walk-on role in synaptic vesicle exocytosis? BioEssays: news and reviews in molecular, cellular and developmental biology, 26, 445–453 (2004).
25. Li, Q. et al. Impaired Cognitive Function and Altered Hippocampal Synaptic Plasticity in Mice Lacking Dermatan Sulfotransferase Chst14/D4st1. Frontiers in molecular neuroscience, 12, 26 (2019).
26. Wang, A. et al. Early-Life Stress Alters Synaptic Plasticity and mTOR Signaling: Correlation With Anxiety-Like and Cognition-Related Behavior. Frontiers in genetics, 11, 590068 (2020).
27. Farajdokht, F. et al. Sericin protects against acute sleep deprivation-induced memory impairment via enhancement of hippocampal synaptic protein levels and inhibition of oxidative stress and neuroinflammation in mice. *Brain research bulletin, 174*, 203–211 (2021).

28. Granja, M. G. et al. Inflammatory, synaptic, motor, and behavioral alterations induced by gestational sepsis on the offspring at different stages of life. *Journal of neuroinflammation, 18*, 60 (2021).

29. Desai, A. & McConnell, S. Progressive restriction in fate potential by neural progenitors during cerebral cortical development. *Development, 127*, 2863–2872 (2000).

30. Huang, L. & Wang, G. The Effects of Different Factors on the Behavior of Neural Stem Cells. *Stem cells international* 2017, 9497325 (2017).

31. Lee, J-T., Tsai, C-K. & Chou, C-H. Development of Neural Stem Cell-Based Therapies for Parkinson’s Disease. *Transl Neurodegener* 3;4:16(2018).

32. Bernabeu-Zornoza, A. et al. Physiological and pathological effects of amyloid-beta species in neural stem cell biology. *Neural regeneration research, 14*, 2035–2042 (2019).

33. Ottoboni, L., von Wunster, B. & Martino, G. Therapeutic Plasticity of Neural Stem Cells. *Frontiers in neurology, 11*, 148 (2020).

34. Aimone, J. et al. Regulation and function of adult neurogenesis: from genes to cognition. *Physiol Rev, 94*, 991–1026 (2014).

35. Zonis, S. et al. p21Cip restrains hippocampal neurogenesis and protects neuronal progenitors from apoptosis during acute systemic inflammation. *Hippocampus, 23*, 1383–1394 (2013).

36. Perez-Dominguez, M., Avila-Munoz, E., Dominguez-Rivas, E. & Zepeda, A. The detrimental effects of lipopolysaccharide-induced neuroinflammation on adult hippocampal neurogenesis depend on the duration of the pro-inflammatory response. *Neural regeneration research, 14*, 817–825 (2019).

37. Chen, B. et al. Neuroprotective effects of natural compounds on neurotoxin-induced oxidative stress and cell apoptosis. *Nutr Neurosci, 2020*, 1–22 (2020).

38. Hernandez Baltazar, D. et al. Does lipopolysaccharide-based neuroinflammation induce microglia polarization? *Folia neuropathologica, 58*, 113–122 (2020).

39. Knobloch, M. The Role of Lipid Metabolism for Neural Stem Cell Regulation. *Brain plasticity, 3*, 61–71 (2017).

40. Lo Van, A., Hachem, M., Lagarde, M. & Bernoud-Hubac, N. Omega-3 Docosahexaenoic Acid Is a Mediator of Fate-Decision of Adult Neural Stem Cells. *International journal of molecular sciences* 20, (2019).

41. Tiberi, M. & Chiurchiu, V. Specialized Pro-resolving Lipid Mediators and Glial Cells: Emerging Candidates for Brain Homeostasis and Repair. *Frontiers in cellular neuroscience, 15*, 673549 (2021).

42. Paoletti, L. et al. Lysophosphatidylcholine Drives Neuroblast Cell Fate. *Molecular neurobiology, 53*, 6316–6331 (2016).

43. Chin, E. W. M., Lim, W. M., Ma, D., Rosales, F. J. & Goh, E. L. K. Choline Rescues Behavioural Deficits in a Mouse Model of Rett Syndrome by Modulating Neuronal Plasticity. *Molecular neurobiology, 56*,
44. Gutierrez-Fernandez, M. et al. CDP-choline treatment induces brain plasticity markers expression in experimental animal stroke. *Neurochemistry international, 60*, 310–317 (2012).

45. Löffelholz, K., Klein, J. & Köppen, A. Chapter 23: Choline, a precursor of acetylcholine and phospholipids in the brain. *98*, 197–200 (1993).

46. Blusztajn, J. & Mellott, T. Choline nutrition programs brain development via DNA and histone methylation. *Cent Nerv Syst Agents Med Chem, 12*, 82–94 (2012).

47. Marcucci, H., Paoletti, L., Jackowski, S. & Banchio, C. Phosphatidylcholine biosynthesis during neuronal differentiation and its role in cell fate determination. *The Journal of biological chemistry, 285*, 25382–25393 (2010).

48. Domizi, P., Malizia, F., Chazarreta-Cifre, L., Diacovich, L. & Banchio, C. KDM2B regulates choline kinase expression and neuronal differentiation of neuroblastoma cells. *Plos one, 14*, e0210207 (2019).

49. Arana, M. R. et al. Coordinated induction of GST and MRP2 by cAMP in Caco-2 cells: Role of protein kinase A signaling pathway and toxicological relevance. *Toxicology and applied pharmacology, 287*, 178–190 (2015).

50. Chojnacka, A., Gladkowski, W. & Grudniewska, A. Lipase-Catalyzed Transesterification of Egg-Yolk Phosphatidylcholine with Concentrate of n-3 Polyunsaturated Fatty Acids from Cod Liver Oil. *Molecules22*, (2017).

51. Ali, A. H. et al. Identification of phospholipids classes and molecular species in different types of egg yolk by using UPLC-Q-TOF-MS. *Food chemistry, 221*, 58–66 (2017).

52. Costa, M. R., Wen, G., Lepier, A., Schroeder, T. & Gotz, M. Par-complex proteins promote proliferative progenitor divisions in the developing mouse cerebral cortex. *Development, 135*, 11–22 (2008).

53. Mosmann, T. Rapid colorimetric assay for cellular growth and survival: application to proliferation and cytotoxicity assays. *J Immunol Methods, 65*, 55–63 (1983).

54. Xiong, F. et al. Optimal time for passaging neurospheres based on primary neural stem cell cultures., *63*, 621–631 (2011).

55. Esko, J., Nishijima, M. & Raetz, C. Animal cells dependent on exogenous phosphatidylcholine for membrane biogenesis. *Proceedings of the National Academy of Sciences of the United States of America, 79*, 1698–1702 (1982).

**Figures**
NSCs proliferation is not affected by inflammation. a) After incubating the NSCs with different dilutions of medium obtained from LPS-stimulated macrophages (AM) or without stimulation as a control (UM) during 96 hours, proliferation was analysed by measuring Neurosphere's diameter. Graph represents the neurosphere's diameter measured in three independent experiments. b) Diameter of the Neurospheres of NSCs exposed to 20% V/V of UM and AM or control during 96 hours. Graph represents the neurosphere's diameter measured in three independent experiments. c) Diameter of the Neurospheres of NSCs exposed to 50 ng/ml of IL-1β and/or IL-6 during 96 hours. Graph represents the neurosphere's diameter measured in three independent experiments. d) Representative images (10X) of neurospheres incubated in the indicated conditions. Scale bars: 100 μm.
NSCs differentiation is affected by inflammation and restored by PtdCho. a) Percentage of β-III tubulin positive cells analysed by immunocytochemistry coupled to fluorescence microscopy of NSCs exposed to 20% V/V of AM and UM in the presence or in the absence of PtdCho (50 μM) during 72 hours. Graph represents the percentage of neuronal differentiation measured in five independent experiments. Data were presented as mean ± SEM. ***p < 0.001; **p < 0.01; *p < 0.05. b) Representative images (40X) of the immunofluorescence assays with the neuronal marker (β-III Tubulin, green) nuclei (DAPI, blue). Yellow arrow indicates normal neuron and red arrow indicates dystrophic neuron. Scale bars: 20 μm.

Figure 3

NSCs differentiation is not affected by ILs. a) Percentage of β-III tubulin positive cells analysed by immunocytochemistry coupled to fluorescence microscopy of NSCs exposed 72 hours to 20% V/V of UM supplemented with the indicated ILs (50 ng/ml) or LPS (1 µg/ml). Graph represents the percentage of neuronal differentiation measured in three independent experiments. b) Representative images (40X) of the immunofluorescence assays with the neuronal marker (β-III tubulin, green) and nuclei (DAPI, blue). Scale bars: 30 μm.
Figure 4

PtdCho rescues neuronal function induced by AM. a) Quantification of neuron’s soma of cells incubated under the indicated condition. Graph represents the area of neuronal soma measured in four independent experiments. Data were presented as mean ± SEM. **p < 0.01; *p < 0.05. b) Percentage of normal neurons (black bars) and dystrophic neurons (grey bars) after 72 hours in culture under the indicated conditions *p<0.05 (Student’s T-test). c) Representative images (100X) of neurons incubated under the indicated conditions Scale bars: 10 μm.

Figure 5
PtdCho supplementation restores synaptic defects caused by AM. a) Quantification of Synaptophysin vesicles per 10 μm of neurite length. Graph represents the number of synaptophysin-containing vesicles in 10 μm axon length measured in four independent experiments. Data were presented as mean ± SEM. **p < 0.01; *p < 0.05. b) Representative images (63X) of the immunofluorescence assays with the neuronal marker (β-III tubulin, green), synaptic vesicle marker (synaptophysin, red) and nuclei (DAPI, blue). Scale bars: 1 μm. c) Representative image of Western Blot showing synaptophysin level in NSCs exposed to 20% V/V of AM in the presence or in the absence of PtdCho (50 μM) during 72 hours. The gels/blots displayed here are cropped, and without high-contrast (overexposure). The full length gels and blots are included in a Supplementary Information file. Densitometric analysis, *p < 0.05 (Student's T-test).

Figure 6

PtdCho restores morphology of dystrophic neurons. a) Percentage of β-III Tubulin positive cells analysed by immunocytochemistry coupled to fluorescence microscopy of NSCs exposed to 20% V/V of AM when PtdCho was added later on, after 24 h of culture and incubated for 72 hours. Graph represents the percentage of neuronal differentiation measured in three independent experiments. Data were presented
as mean ± SEM. ***p < 0.001. b) Percentage of normal neurons (black bars) and dystrophic neurons (grey bars) after 72 hours in culture under the indicated conditions. ***p < 0.001 (Student’s T-test). c) Representative images (40X) of neurons incubated under the indicated conditions Scale bars: 30 μm.

**Supplementary Files**

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