Control of Mitochondrial pH by Uncoupling Protein 4 in Astrocytes Promotes Neuronal Survival*

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Background: The role of uncoupling proteins (UCP) in the brain is unclear.

Results: UCPs, present in astrocytes, mediate the intramitochondrial acidification leading to a decrease in mitochondrial ATP production.

Conclusion: Astrocyte pH regulation promotes ATP synthesis by glycolysis whose final product, lactate, increases neuronal survival.

Significance: We describe a new role for a brain uncoupling protein.

Brain activity is energetically costly and requires a steady and highly regulated flow of energy equivalents between neural cells. It is believed that a substantial share of cerebral glucose, the major source of energy of the brain, will preferentially be metabolized in astrocytes via aerobic glycolysis. The aim of this study was to evaluate whether uncoupling proteins (UCPs), located in the inner membrane of mitochondria, play a role in setting up the metabolic response pattern of astrocytes. UCPs are believed to mediate the transmembrane transfer of protons, resulting in the uncoupling of oxidative phosphorylation from ATP production. UCPs are therefore potentially important regulators of energy fluxes. The main UCP isoforms expressed in the brain are UCP2, UCP4, and UCP5. We examined in particular the role of UCP4 in neuron-astrocyte metabolic coupling and measured a range of functional metabolic parameters including mitochondrial electrical potential and pH, reactive oxygen species production, NAD/NADH ratio, ATP/ADP ratio, CO₂ and lactate production, and oxygen consumption rate. In brief, we found that UCP4 regulates the intramitochondrial pH of astrocytes, which acidifies as a consequence of glutamate uptake, with the main consequence of reducing efficiency of mitochondrial ATP production. The diminished ATP production is effectively compensated by enhancement of glycolysis. This nonoxidative production of energy is not associated with deleterious H₂O₂ production. We show that astrocytes expressing more UCP4 produced more lactate, which is used as an energy source by neurons, and had the ability to enhance neuronal survival.

The uncoupling phenomenon of respiration from ATP production in favor of the production of heat (1) has been well described for the uncoupling protein 1 (UCP1, thermogenin)² (2, 3). This phenomenon is controlled by the sympathetic nervous system and is mainly present in specific tissues dedicated to the production of heat, such as brown adipose tissue.

The common term “uncoupling protein” is a legacy of the analysis of sequence homology with UCP1. It is used for other UCP isoforms whose role is less established and not necessarily related to heat production. As far as UCP2 is concerned, its role is related to temperature control (4) and regulation of reactive oxygen species (ROS) (5) in the tissues studied. UCP3 expression correlates with free fatty acid production (6) with a decoupling role primarily in muscle (7); its expression is marginal in brain (8). The role of UCP4 and UCP5 is still largely unknown. In brain, UCP4 and 5 mRNA levels are modulated by diet or temperature changes (9). A role in the regulation of ROS has been described in neural cells (10, 11).

Neurons and astrocytes present different metabolic profiles (12). Neurons are highly oxidative, producing ROS, whereas astrocytes are predominantly glycolytic. In the brain, a role in ROS and temperature control have been described for UCP2 (4). UCP4 and UCP5 are well expressed in the brain (13, 14), and roles in ROS and calcium level regulation (10, 15) have been described. In terms of brain cellular localization, immunocytochemistry has revealed that UCP4 is mainly expressed in neurons and less so in astrocytes (16). UCP5 is present in neuronal cell lines (17, 18); in the rodent brain, UCP5 expression is observed in the dorsomedial hypothalamic nucleus, hippocampus, paraventricular thalamic nucleus, mediodorsal thalamic nucleus, and ventromedial hypothalamus (13, 19).

In summary, only limited information is currently available regarding the role of UCPs in the brain, with the main observations with a cellular resolution being limited to neurons. In this

‡ The abbreviations used are: UCP, uncoupling protein; ROS, reactive oxygen species; OCR, oxygen consumption rate; Tricine, N-[2-hydroxy-1,1-bis(hydroxymethyl)ethyl]glycine; MTT, 3-(4,5-dimethylthiazol-2-ylo)-2,5-diphenyltetrazolium bromide; Bicine, N,N-bis(2-hydroxyethyl)glycine.

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THE JOURNAL OF BIOLOGICAL CHEMISTRY VOL. 289, NO. 45, pp. 31014 –31028, November 7, 2014 © 2014 by The American Society for Biochemistry and Molecular Biology, Inc. Published in the U.S.A.
study, we have investigated the roles of UCP4 and UCP5 in astrocytes, a cell type central to brain energy metabolism.

We have previously shown (20) that uptake into astrocytes of neuronally released glutamate results in an intramitochondrial pH decrease in astrocytes rapidly reducing the effectiveness of the oxidative phosphorylation. In the present report, we show that this effect of glutamate involves UCP4. Indeed, we observed that the decrease in oxidative efficiency following glutamate uptake into astrocytes is controlled by UCP4, which reduces mitochondrial respiration by lowering the intramitochondrial pH. Reducing mitochondrial efficiency promotes aerobic glycolysis in astrocytes. Lactate produced through glycolysis can be used by neurons as an energy substrate consistent with the astrocyte-neuron lactate shuttle (21). These results indicate that UCP4 plays a key role in astrocyte-neuron metabolic coupling (12).

**EXPERIMENTAL PROCEDURES**

**Primary Culture of Astrocytes**—Primary cultures of cortical astrocytes were prepared from newborns (1–2 days old) Swiss Albino mice (OF1; Charles River Laboratories) as previously described (22). The cells were seeded at a density of \(10^5\) cells/cm\(^2\). Depending on the experiments, the cells were seeded on 35-mm Petri dishes or multiwell plates of 6, 12, 24, or 48 wells. The plates were incubated at 37 °C in an atmosphere containing 5% CO\(_2\) and 95% air. The cells were rinsed 3–5 days after seeding, and the medium was changed twice a week until cells reached confluency at 21 days in vitro. Experiments were performed on confluent cells.

**Primary Cultures of Neurons and Co-culture**—Primary cultures of cortical neurons were prepared from embryonic day 17 OF1 mice embryos (Charles River Laboratories). Pregnant mice were sacrificed by cervical dislocation. The abdomen was opened, and the placenta containing the embryos was removed and put in a Petri dish. The embryos were rapidly recovered, and the cut heads were placed in Petri dish containing 25 ml of ice-cold Hanks’ buffered salt solution supplemented with penicillin and streptomycin. The embryo brains were separated from the skulls, and olfactory bulbs, striata, hippocampi, and meninges were removed. The isolated cortices were minced in ice-cold Hanks’ buffered salt solution supplemented with penicillin and streptomycin. The embryo brains were isolated from the skulls, and olfactory bulbs, striata, hippocampi, and meninges were removed. The isolated cortices were minced in ice-cold Hanks’ buffered salt solution supplemented with penicillin and streptomycin. The embryo brains were separated from the skulls, and olfactory bulbs, striata, hippocampi, and meninges were removed. The isolated cortices were minced in ice-cold Hanks’ buffered salt solution supplemented with penicillin and streptomycin. The embryo brains were separated from the skulls, and olfactory bulbs, striata, hippocampi, and meninges were removed. The isolated cortices were minced in ice-cold Hanks’ buffered salt solution supplemented with penicillin and streptomycin. The embryo brains were separated from the skulls, and olfactory bulbs, striata, hippocampi, and meninges were removed. The isolated cortices were minced in ice-cold Hanks’ buffered salt solution supplemented with penicillin and streptomycin. The embryo brains were separated from the skulls, and olfactory bulbs, striata, hippocampi, and meninges were removed. The isolated cortices were minced in ice-cold Hanks’ buffered salt solution supplemented with penicillin and streptomycin. The embryo brains were separated from the skulls, and olfactory bulbs, striata, hippocampi, and meninges were removed. The isolated cortices were minced in ice-cold Hanks’ buffered salt solution supplemented with penicillin and streptomycin.
**Lentiviral DNA Constructs—** mRNA for UCP4 and UCP5 were isolated from brain of OF1 mice. Reverse transcription was prepared by using 200 ng of extracted mRNA in 50 µl of total RT. Specific cDNA was obtained by PCR with the following primers: mUCP4FoEcoRI, 5'-GAA TTC TGC TGA ATG CCT ATC GCC; mUCP4ReSphI, 5'-GCA TGC ATG GCC TGA TCT CAC TC; mUCP5FoEcoRI, 5'-GAA TTC TGC ATG CAA GAC GTA GAT CGG ATA CGC GTA AGC GTA ATC CGG AAC GTC GTA CGG ATA CGC GTA TGG AAC GTC ATA TGG GTA. The next step in the lentiviral construction is the use of the Gateway System. The gene of interest is thus fused to three contiguous copies of the HA epitope (AGC GTA GTC AGG TAC ATC GTA GAG GTA AGC GTA ATC CGG AAC GTA AGC GTA ATC CGG ATA CGC GTA GTC TGG AAC GTA ATA TGG GTA). The mature antisense coded by the shUCP4 virus was ATT TCT TTG AAA CGA ACA TCG. As a control of gene silencing, we infected astrocytes with a lentivirus made with a vector containing no sequence.

**Western Blots—** When astrocytes reached confluence at day 21 in vitro, 35-mm Petri dishes of infected cells were placed on ice and rinsed with ice-cold PBS. Cells were homogenized and proteins were extracted with 200 µl of lysis buffer (30 mM HEPES, 210 mM sucrose, 40 mM NaCl, 2 mM EDTA, 1% SDS, and a mixture of protease inhibitors (Complete 11257000; Roche). Each condition was made in duplicate, and two dishes were mixed. Protein samples were sonicated and frozen at -80 °C. 10 µl of each sample were mixed to 10 µl of 2× loading buffer (100 mM Tris-Cl, pH 6.8, 200 mM DTT, 4% SDS, 0.1% bromphenol blue, 10% glycerol) and heated at 100 °C for 4 min before loading onto an Invitrogen NuPage 10% gel (NP0301BOX). When proteins were loaded after mitochondrial isolation following the mitochondria isolation kit (MITOISO1; Sigma-Aldrich) protocol, 2.5 µg of protein were loaded. Gel was run at 100 volts at 4 °C. After running, proteins were transferred on nitrocellulose membranes (84261514; Whatman) in a Tris-glycine transfer buffer (0.4M glycine, 20 mM Tris-Cl, pH 8.9, 300 mM NaCl) and then incubated overnight at 4 °C with primary antibodies polyclonal anti-HA (1:4000) (AB9110; Abcam), monoclonal anti-β actin (1:5000) (A5441; Sigma-Aldrich), or monoclonal anti-cytochrome c oxidase (1:1000) (MS407; Mitosciences) diluted in the same blocking buffer. After several washes in PBS, secondary antibodies anti-mouse IgG IRDye 800 (1:5000) (610-132-121; Bioconcept) and anti-rabbit IgG Alexafluor 680 (1:5000) (Juro, Lucerne, Switzerland) were applied for 2 h at room temperature and protected from light. After washing, membranes were scanned with the Odyssey Infrared Imaging System (LI-COR Biosciences).

**Immunocytochemistry—** Astrocytes grown on 20-mm coverslips and infected with UCP lentiviral constructs were carefully rinsed with PBS before being fixed in cold parformaldehdyde 4% for 15 min at room temperature. Treatment with PBS, 0.1% Triton for 15 min was performed to permeabilize the cell membranes. After three PBS washes, fixed cells were then treated with casein (0.5% in PBS) for 1 h at room temperature to block nonspecific sites. Fixed cultures were then incubated overnight at 4 °C in the presence of the primary antibodies polyclonal anti-HA (1:500) (AB9110; Abcam), polyclonal GFAP (1:500) (Z0334; Dako), or monoclonal anti-cytochrome c oxidase (1:500) (MS407; Mitosciences) diluted in the same blocking buffer. After several washes in PBS, secondary antibodies anti-mouse IgG IRDye 800 (1:5000) (610-132-121; Bioconcept) and anti-rabbit IgG Alexafluor 680 (1:5000) (Juro, Lucerne, Switzerland) were applied for 2 h at room temperature and protected from light. After washing, membranes were scanned with the Odyssey Infrared Imaging System (LI-COR Biosciences).
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and mitochondria was calculated with the Manders’ coefficient which is the part of the intensity in each channel that coincides with a number of intensity in the other channel (with 0 indicating no co-localization and 1 indicating perfect co-localization).

Glucose Uptake Assay—Glucose uptake was measured as previously described (23). The effect of glutamate on astrocytic glucose uptake was measured in parallel in other Petri dishes by adding glutamate 200 μM in the medium containing [3H]2-deoxyglucose for 20 min of incubation. Other Petri dishes were used to measure the portion of glucose uptake that is not linked to glucose transporter by addition of the glucose transporter inhibitor cytochalasin B (C6762; Sigma-Aldrich) 25 μM during 20 min of incubation. The fraction of glucose transported is calculated by subtracting the fraction of glucose uptake that is not inhibited by the cytochalasin B. Glucose uptake was normalized to the protein content.

Glycogen Content—Glycogen content was measured enzymatically as previously described (22). In brief, astrocytes grown on 35-mm Petri dishes and previously infected with UCP lentiviral constructs were washed three times with ice-cold PBS and lysed by sonication in HCl 30 mM. Two aliquots of 100 μl were sampled. 300 μl of 0.1 M acetate buffer, pH 4.65, was added to the first aliquot, whereas the same buffer containing 1% amyloglucosidase (10 mg/ml) was added to the other one. After incubation at room temperature for 30 min, 2 ml of 0.1 x Tris-HCl buffer, pH 8.1, containing 3.3 mM MgCl2, 0.33 mM ATP, 38 μM NADP, 4 μg/ml hexokinase, and 2 μg/ml glucose-6-phosphate dehydrogenase were added, and the mixture was incubated at room temperature for 30 min. Fluorescence associated with the formation of NADPH was read on a fluorometer using excitation wavelength at 340 nm and emission wavelength at 450 nm. Glycogen was normalized to protein content and was expressed as glycosyl units originating from glycogen. The results are expressed as percentages of control values.

Mitochondrial Potential Measurement—Astrocytes grown on a multpile of 48 wells and previously infected with UCP lentiviral constructs were rinsed carefully and incubated 1 h in phenol red-free DMEM (D5030; Sigma) supplemented with 2 mM glucose and 44 mM NaHCO3. After rinsing, cells were incubated in the same medium containing 5 μg/ml JC-1 dye (T-3168; Invitrogen) for 30 min at 37 °C in an atmosphere containing 5% CO2 and 95% air and protected from light. The JC-1 dye exhibits potential dependent accumulation in mitochondria, indicated by a fluorescence emission shift from green to red (24, 25). At low potential, monomers are formed that display green fluorescence, whereas at higher potential, JC-1 aggregates and exhibits a red fluorescence. Cells were carefully rinsed with PBS to remove any extracellular dye, and fluorescence was directly detected on the multiplate using a fluorometer (Safir 2; Tecan). The wavelengths used are emission 497/ excitation 594 for the red fluorescence and emission 497/ excitation 524 for the green fluorescence. The mitochondrial potential was estimated by calculating the ratio of green to red fluorescence. The results are expressed as percentages of control values.

ATP/ADP Ratio—ATP content was measured enzymatically with the luciferase. In the presence of ATP, Mg2+, and oxygen, luciferin is oxygenated by luciferase into oxy Luciferin. This reaction emits light that is proportional to the amount of ATP. ATP was determined with the CellTiter-Glo Luminescent cell viability assay (Promega) with some modifications. Astrocytes grown on multplate of 48 wells and previously infected were rinsed and incubated 1 h at 37 °C in an atmosphere containing 5% CO2 and 95% air in DMEM (D5030; Sigma-Aldrich) containing 44 mM NaHCO3 and 2 mM glucose. At the end of the incubation, medium was removed, and 200 μl of Tricine buffer solution (40 mM Tricine, 3 mM EDTA, 85 mM NaCl, 3.6 mM KCl, 100 mM NaF, and 0.1% saponin (84510; Sigma-Aldrich), pH 7.4) was put in each well. Cells were lysed by saponin effect and by pipetting. The saponin is a “soft” soap that breaks cell membranes but preserves the enzymatic reaction. Each sample was separated in two: one part for ATP measure and the other part for ATP + ADP measure. 90-μl aliquots were distributed in a black-walled 96-well type microplates (PerkinElmer Life Sciences). For the ATP + ADP measure, 10 μl of converting solution (100 mM Tricine, 100 mM MgSO4, 25 mM KCl, 1 mM phosphoenolpyruvate, and 100 units/ml pyruvate kinase), pH 7.75, was added in each well, whereas the same solution without phosphoenolpyruvate and pyruvate kinase was added to the samples for ATP measure. An incubation of 5 min at room temperature was performed before adding 10 μl of MgCl2 solution (4 mM Tricine and 100 mM MgCl2). Finally, 100 μl of CellTiter-Glo reagent (G7571; Promega) was added, and luminescence was immediately detected with a luminometer (Safire 2; Tecan). Luminescence was measured in a kinetic way determined by 20 readings at intervals of 1 min. Luminescence read at the plateau were taken to calculate the ATP/ADP ratio. ADP values were calculated by subtracting ATP values from ATP + ADP values. The results are expressed as percentages of control values.

Hydrogen Peroxide Quantification—H2O2 accumulated in the medium and was detected enzymatically with Amplex red (26, 27). Oxidation of Amplex red is catalyzed by the horseradish peroxidase in the presence of H2O2 into highly fluorescent resorufin. When reaching confluency, astrocytes grown on multplates of 24 wells and previously infected with UCP lentiviral constructs were carefully rinsed before being incubated 4 h in phenol red free Neurobasal medium supplemented with GlutaMAX and B27 without antioxidant (B27 minus AO) and containing 10 μM Amplex red (A12222; Invitrogen) and 1 unit/ml horseradish peroxidase (type II, P8250; Sigma-Aldrich). At the end of the incubation, 100 μl were loaded on a black-walled 96-well plate (PerkinElmer Life Sciences), and fluorescence was detected using wavelengths at 545 nm for excitation and 590 nm for emission. The results are expressed as percentages of control values.

NAD+/NADH Cycling Assay—NADH and NAD+ were measured enzymatically as previously described (28) with some modifications. NAD+ is reduced by alcohol dehydrogenase in the presence of ethanol. NAD+ is renewed by the reduction of phenazine ethosulfate, whose oxidized form is restored by the reduction of MTT. Reduced MTT is visualized by absorbance at 570 nm. Infected astrocytes grown on 35-mm Petri dish and reaching confluency were incubated 24 h in phenol red-free Neurobasal medium without glucose (0050128DJ; Invitrogen) supplemented with 5 mM glucose, B27, and GlutaMAX. The
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cells were then put on ice and washed three times with ice-cold PBS. Cells were recovered in 600 μl of carbonate-bicarbonate (20 mM to 100 mM) buffer, pH 10, containing 10 mM nicotinamide (72340; Sigma-Aldrich) and were directly frozen to disrupt cell membrane. Once thawed, the samples were kept on ice and separated in two parts: one for the dosage of NADH + NAD⁺ and the other for NADH only. Samples for NADH detection were heated at 60 °C for 30 min to destroy NAD⁺ and were kept on ice. 50 μl of each samples were loaded on a 96-well microplate, and the reaction was initiated by addition of 150 μl of reaction mix composed of 133 mM Bicine (14871; Sigma-Aldrich), 5.33 mM EDTA, 0.56 mM MTT (M5655; Sigma-Aldrich), 2.21 mM phenazine ethosulfate (0210095601; MP Biomedicals), 667 mM ethanol, and 40 units/ml alcohol dehydrogenase (A3263; Sigma-Aldrich). Reduction of MTT was followed by measuring absorbance at 570-mm for intervals of 30 s. NAD⁺ values were calculated by subtracting NADH from NADH + NAD⁺ values. Concentrations were determined from a standard curve from 0.1 to 2 μM NADH (10107735001; Roche) and were normalized to the protein content. The results are expressed as percentages of control conditions.

CO₂ Production Assay—CO₂ production was measured by incorporation of radioactive labeled glucose as previously described (29). This technique enabled us to distinguish the quantity of CO₂ produced by glucose metabolism via the TCA cycle between those produced by the pentose phosphate pathway. D-[1-14C]Glucose produces 14CO₂ via the two pathways, whereas 14CO₂ produced by D-[6-14C]glucose is exclusively provided by the TCA cycle. CO₂ produced specifically by the TCA cycle is determined by metabolism of D-[6-14C]glucose, whereas use of D-[1-14C]glucose gives the total amount of CO₂ formed. When reaching confluency, medium of astrocytes growing on 35-mm Petri dishes was changed for DMEM (D5030; Sigma-Aldrich) containing 2.5 mM glucose, 7.5 mM NaHCO₃, 5 mM HEPES, and 10 ml/liter antibiotic/antimycotic solution, and cells were incubated 2 h at 37 °C in an atmosphere containing 5% CO₂ and 95% air. The medium was then replaced by 1 ml of the same medium supplemented with 2.5 μCi/ml D-[1-14C] (CFA349; Amersham Biosciences) glucose or D-[6-14C]glucose (CFA351; Amersham Biosciences), and culture dishes were placed in sealed glass containers and incubated for 2 h at 37 °C. 0.2 M HCl was injected in dishes containing the respiring cells to stop the reaction, and radioactive 14CO₂ was captured by addition of 1 ml of Carbo-Sorb (PerkinElmer Life Sciences) against the walls of the container. These were kept aside 1 h for equilibration, and two aliquots of 400 μl of Carbo-Sorb were taken for radioactivity assay by liquid scintillation counting (Permafluor E⁺; PerkinElmer Life Sciences). The values were normalized to the protein content, and the results are expressed as percentages of control values.

Oxygen Consumption Rate—The oxygen consumption rate (OCR) was assessed using a noninvasive extracellular flux analyzer (XF24; Seahorse Biosciences, North Billerica, MA). It is composed of two calibrated optical sensors that directly measure the oxygen consumption rate in a 7-μl microchamber with limited diffusion. The depletion of oxygen in the medium is then measured several times during a defined period. Injector ports surrounding the sensor of each well can be used to automatically deliver drugs during the experiment. The cells were plated in XF24 wells (XF24 FluxPaks, 100850-001; Bucher, Basel, Switzerland) of 0.38 cm² area per well and incubated for 1 h in a non-CO₂ incubator with unbuffered medium before loaded in the XF24 Analyzer. For astrocytes, the medium was composed of DMEM (D5030; Sigma-Aldrich) supplemented with 32 mm NaCl, 15 mg/liter Phenol red, 10 mM D-2-deoxyglucose, 5 mM glucose, and 5 mM pyruvate. For neuronal cultures, medium was composed of DMEM supplemented with 32 mm NaCl, 15 mg/liter Phenol red, and 0.5 mM glucose. During the measure, 200 μM glutamate was first added in astrocytes culture. In neurons, glucose and/or lactate were injected at energetically equivalent concentrations as indicated. In all cases, later injection of 5 μM oligomycin (75352; Sigma-Aldrich) and 2 μM carbonyl cyanide-4-(trifluoromethoxy)phenylhydrazone (C2920; Sigma-Aldrich) provided estimation of the success of the experiment. Effect of glutamate was normalized to the basal OCR, as well as the effect of glucose and/or lactate in neurons. The results are expressed as percentages of control values.

Mitochondrial Matrix pH—Two-week-old astrocytes were placed in 2 ml of antibiotic-free and serum-free DMEM with FuGENE (Roche) and DNA encoding for genetically encoded pH indicator MitoSypHer (30). Quantities of DNA/FuGENE were 2 μg of DNA/12 μl of FuGENE for MitoSypHer and 4 μg of DNA/8 μl of FuGENE for MIMS-EYFP. After 4 h, the medium was changed with DMEM plus 10% serum, and the cells were used 2–3 days after transfection for pure astrocyte culture and 14 days for mixed neuron-astrocyte cultures. Mitochondrial matrix pH was measured by fluorescence microscopy as previously described (20). Briefly, cells were placed in a thermostatted chamber designed for rapid exchange of perfusion solutions (31) and superfused at 35 °C. Experimental solution contained 160 mM NaCl, 5.4 mM KCl, 20 mM HEPES, 1.3 mM CaCl₂, 0.8 mM MgSO₄, 0.78 mM NaH₂PO₄, 5 mM glucose (pH 7.4) and was bubbled with air. MitoSypHer fluorescence was sequentially excited at 490 and 420 nm and detected at >515 nm. At the end of each experiment, in situ calibration was performed by permeabilization of cells with 10 μM monensin and 3 μM gliclazide and superfused with a series of calibration solutions of given pH values (pH 5.9, 7.0 7.5, and 8.0). Calibration solutions contained 20 mM NaCl, 125 mM KCl, 0.5 mM MgCl₂, 0.2 mM EGTA, 20 mM HEPES and bubbled with air.

Lactate Release Assay—Lactate released by astrocytes previously infected with UCP lentiviral constructs was determined with an enzymatic method, first described by Rosenberg and Rush (32) with some modifications. Lactate dehydrogenase oxidizes lactate into pyruvate in the presence of NAD⁺. The resulting NADH is measured by fluorometry. 24 h after co-culture initiation, 10 μl of medium was taken and mixed with 190 μl of medium. This was mixed with 1 ml of 0.2 M glycine-semicarbazide, pH 10 buffer containing 2 mg/ml NAD and 13.75 units/ml lactate dehydrogenase and was incubated at 40 °C for 1 h. 200 μl of this reaction was transferred into a black-walled 96-well plate (PerkinElmer Life Sciences). The formation of NADH was measured by fluorescence (Safire 2; Tecan) using excitation wavelength at 340 nm and emission wavelength at 450 nm. Absolute values were determined from a standard curve, and
lactate release in the medium was normalized to the DNA quantity in each Petri dish using the PicoGreen method. For this normalization, cells were incubated for 5 min in PBS containing PicoGreen solution (P7589; Invitrogen) diluted 1:200. After washing, fluorescence was measured using excitation wavelength at 480 nm and emission wavelength at 520 nm. The results are expressed as percentages of control values.

Viability Test: Calcein Assay—48 h after co-culture initiation, coverslips with neurons were transferred to new wells containing Neurobasal medium supplemented with GlutaMAX, B27 without antioxidant (B27 minus AO), and 1 μg/ml calcein-AM. Neurons were carefully rinsed, and calcein fluorescence was measured directly in the multwell plate with a fluorescence reader (Safire 2; Tecan) with excitation wavelength of 490 nm and emission of 515 nm. The calcein fluorescence was normalized to the cell quantity in each well by adding the nucleus marker DAPI (D9542; Sigma-Aldrich). Neurons were incubated for 10 min in PBS containing DAPI 7 μg/ml, and after rinsing, DAPI fluorescence was measured using excitation wavelength at 360 nm and emission at 461 nm. The results are expressed as percentages of control values.

Viability Test: MTT Reduction Assay—Viability of neurons in co-culture were measured with the MTT (M5655; Sigma-Aldrich) reduction assay (33). 48 h after co-culture initiation, co-culture were measured with the MTT (M5655; Sigma-Aldrich) supplemented with 2 mM glucose, 44 mM NaHCO3, and 20 nM Mitotracker Red 580 (M-22425; Invitrogen). Cells in culture were rinsed, and calcein fluorescence was measured from cytosolic fraction and run on a polyacrylamide gel. Mitochondrial localization of the overexpressed uncoupling proteins was also confirmed by immunohistochemistry in astrocytes infected with the corresponding UCP construct, detected by a band close to 32 kDa.

Mitochondrial Localization of Overexpressed UCPs in Astrocytes—Expression of various UCPs in the brain has been previously reported (8, 34). The results presented in Table 1 summarize the expression levels for each UCP isoform in astrocyte and in neuron mouse primary cultures. UCP2, UCP4, and UCP5 are the main UCP isoforms and are expressed both in astrocytes and neurons.

Overexpression of the various isoforms of uncoupling protein in primary cultures of astrocytes through viral transfection results in an increased expression of the corresponding UCP isoform as determined by RT-PCR. The increase ranged from a 250-fold increase in mRNA for UCP2, UCP4, and UCP5 to 30,000-fold for UCP1 and UCP3 (one should note that the expression levels of these two latter isoforms were extremely low before transfection). Using silencing constructs for UCP4 or UCP5 (see “Experimental Procedures”), we observed a reduction in the corresponding UCPs mRNAs (data not shown).

We also analyzed the expression and the subcellular localization of overexpressed uncoupling proteins by Western blotting and immunohistochemistry. For increased specificity, we used an hemagglutinin epitope (HA tag) (see “Experimental Procedures”) fused to UCP sequences that allows us to determine the expression of each overexpressed isoform. As seen in Fig. 1A, all isoforms are present in astrocytes infected with the corresponding UCP construct, detected by a band close to 32 kDa. For the UCP5 protein, two bands were observed in some blots, most likely resulting from alternative splicing (9).

UCP4 and UCP5 overexpression was also determined in mitochondrial preparations. Mitochondrial fraction was separated from cytosolic fraction and run on a polyacrylamide gel. Antibody against cytochrome oxidase was used as a marker of the mitochondrial fraction. Uncoupling proteins 4 and 5 were highly enriched in the purified mitochondrial fraction (Fig. 1B). Mitochondrial localization of the overexpressed uncoupling proteins was also confirmed by immunohistochemistry in astrocytes. Primary antibodies directed against the HA tag and the mitochondrial cytochrome oxidase were used for co-localization analysis. The UCP4-HA (Fig. 1C) is revealed in green, whereas cytochrome oxidase is revealed in red, and co-localization appears in white. The results in Fig. 1C showed that UCP4-HA co-localized with the cytochrome oxidase, suggesting that the overexpressed uncoupling protein was targeted to the mitochondria.

We tested the viability of the lentivirus-infected astrocytes using the calcein test and found that it did not affect cell survival.

### RESULTS

**Mitochondrial Localization of Overexpressed UCPs in Astrocytes**—Expression of various UCPs in the brain has been previously reported (8, 34). The results presented in Table 1 summarize the expression levels for each UCP isoform in astrocyte and in neuron mouse primary cultures. UCP2, UCP4, and UCP5 are the main UCP isoforms and are expressed both in astrocytes and neurons.

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We tested the viability of the lentivirus-infected astrocytes using the calcein test and found that it did not affect cell survival.

### Table 1

| UCP   | Astrocytes | Neurons |
|-------|------------|---------|
| UCP1  | 0.33 ± 0.02| 1.72 ± 0.27 |
| UCP2  | 1.83 ± 0.3 | 119 ± 45  |
| UCP3  | 0.08 ± 0.02| 0.06 ± 0.02 |
| UCP4  | 100 ± 2    | 321 ± 35  |
| UCP5  | 291 ± 12   | 478 ± 32  |

**UCP4 Regulates Intramitochondrial pH**

To assess real differences level of mRNA for the various UCP isoforms, we convert quantitative PCR data in copy number. To evaluate the differences between the different UCP isoforms, we have arbitrary set the UCP4 copy number to 100. The other UCPs isoforms are shown as percentages compared with astrocyte UCP4 expression level. This representation highlights the differences of each UCP isoform between astrocytes and neurons expression levels.
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**FIGURE 1. Mitochondrial localization of overexpressed UCPs in astrocytes.** A, Western blot showing the expression of each HA-tagged isoform of UCPs. B, localization of overexpressed UCP. HA-tagged overexpressed UCP4 and UCP5 were present in the total fraction and were purified in the mitochondrial fraction. C, immunohistochemistry showing the co-localization of UCP4-HA (green) and cytochrome oxidase (red). Thresholded Manders’s coefficient (green versus red channel) was 0.93, indicating a very high degree of correlation. Ctrl, control; Cyt., cytosolic fraction; Mito., mitochondrial fraction; Tot., total fraction.

It has been shown that during neuronal activation, glutamate exposure increased lactate release from astrocytes (21). Lactate release was measured at the end of the 20-min glucose uptake experiments in control and UCPs overexpressing astrocytes stimulated or not with glutamate. Overexpression of uncoupling proteins did not modify the basal or glutamate-stimulated astrocytic lactate release. In contrast, overexpression of UCP4 or UCP5 silencing was without effect on basal glucose uptake. However, UCP4 silencing significantly reduced glutamate-stimulated glucose uptake by 34 ± 5.4% compared with control astrocytes (Fig. 2A, arrow).

It has been shown that astrocytes can metabolically adapt drawing on their glycogen stocks. Glucose metabolism. Glutamate promotes glucose uptake into astrocytes (21). Overexpression of different UCP isoforms did not affect glucose uptake either under basal conditions or following glutamate stimulation (Fig. 2A) as revealed by the [3H]2-deoxyglucose experiments. Similarly, UCP4 or UCP5 silencing was without effect on basal glucose uptake. However, UCP4 silencing significantly reduced glutamate-stimulated glucose uptake by 34 ± 5.4% compared with control astrocytes (Fig. 2A, arrow).

Glucose uptake by astrocytes and neurons. Glucose uptake, lactate release, and glycogen content were measured as an index of astrocytic glucose metabolism. Glutamate promotes glucose uptake into astrocytes on the metabolic coupling between astrocytes and neurons. Glucose uptake, lactate release, and glycogen content were measured as an index of astrocytic glucose metabolism. Glutamate promotes glucose uptake into astrocytes on the metabolic coupling between astrocytes and neurons. Glucose uptake, lactate release, and glycogen content were measured as an index of astrocytic glucose metabolism.

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overexpression had no effect on the mitochondrial potential of astrocytes (not shown). This suggests that UCP activity is tissue-specific. Silencing of UCP4 and UCP5 had the opposite effect to overexpression, resulting in an increase in the mitochondrial potential by 10% (Fig. 3A, gray columns). These results confirm the implication of UCP4 and UCP5 in mitochondrial potential in astrocytes.

In mitochondria, ATP production is closely linked to the proton gradient across the inner mitochondrial membrane. Because ATP synthase requires the proton motive force to produce ATP, it is expected that a reduction in mitochondrial potential leads to a reduction in ATP formation. Accordingly, overexpression of UCP4 or UCP5 significantly decreases the ATP/ADP ratio in astrocytes by 12–14% (Fig. 3B, black columns). These results suggested a leak of protons from the intramitochondrial matrix via uncoupling proteins, which led to a decreased activity of the FoF1-ATP synthase. Silencing yielded the reverse effect for both UCP4 and 5, i.e. increased the ATP/ADP ratio by 20%, supporting this conclusion (Fig. 3B, gray columns).

A possible role for the uncoupling protein described in the literature (for review, see Ref. 37) is the reduction of ROS production. Indeed, a mild uncoupling has been shown to have a moderating effect on oxidative stress. We tested this protective effect for the two brain isoforms of uncoupling proteins in astrocytes. We chose to measure superoxide anion production as an index of ROS production. However, because of its very short half-life, we quantified H2O2 release as an alternative. Overexpression of UCP4 or UCP5 significantly reduced H2O2 release into the medium of astrocyte cultures by 9–11% compared with control cells (Fig. 3C, black columns). In contrast, silencing of UCP4 and UCP5 resulted in enhanced H2O2 release by 127 and 133%, respectively, of control cells (Fig. 3C, gray columns).

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**A** Effects of UCP2, UCP4, and UCP5 on astrocytic glucose metabolism and the effect of silencing with shUCP4,5 on respective mRNA levels. A, glucose uptake before (black bars) and after (gray bars) glutamate exposure. The presence of glutamate increased glucose uptake by 45%. UCPs overexpression had no effect on glutamate-stimulated glucose uptake. The results are means of triplicates of eight independent experiments. Silencing of UCP4 reduced glutamate-stimulated glucose uptake compared with control conditions. We have subtracted from each value represented by an histogram the value of passive glucose uptake obtained when glucose transporter are blocked with cytochalasin B. B, overexpression of UCP4 and UCP5 reduced glycogen content compared with control conditions. The results are means of triplicates of six independent experiments. C, mRNA level of UCP4 and UCP5 genes in astrocytes expressing either the shUCP4 or shUCP5. Ctrl, control; 2-DG, 2-deoxyglucose; glut, glutamate.
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**Figure 3.** UCP4 and UCP5 reduced the mitochondrial function in astrocytes. A, mitochondrial electrical potential was reduced in the presence of overexpressed UCP4 and UCP5 (black bars). Overexpression of UCP2 did not modify the electrical potential of astrocytes. Control astrocytes are infected with a virus containing the GFP sequence. The silencing of these two isoforms increased the mitochondrial potential (gray bars), and control astrocytes are infected with a virus targeting the GFP sequence. The results are means of quadruplicates of at least 12 independent experiments. B, the ATP/ADP ratio was reduced in the presence of overexpressed UCP4 and UCP5 (black bars); opposite effects were obtained with the silencing (gray bars). Overexpression of UCP2 did not modify the ATP/ADP ratio of astrocytes. The results are means of quadruplicate of at least 12 independent experiments. C, UCP4 and UCP5 overexpression (black bars) reduced H$_2$O$_2$ level found in medium, whereas silencing of UCP4 or UCP5 (gray bars) increased the release of H$_2$O$_2$. The results are means of quadruplicate of at least 14 independent experiments. Ctrl, control.

**A**

**B**

**C**

hydrogenase and pyruvate dehydrogenase. The TCA cycle is an important source of NADH providing reducing equivalents to the respiratory chain. Astrocytes overexpressing UCP4 showed a 33% reduction of the NAD$^+$/NADH ratio compared with control astrocytes (Fig. 4A, black columns). In contrast, UCP4 silencing increased the ratio by 148% of control levels (Fig. 4A, gray columns), suggesting that the cell is in a more reduced state in these conditions. Modulation of UCP5 expression was without effect.

The decrease in the NAD$^+$/NADH ratio observed following UCP4 overexpression could result both from an increase in the TCA cycle activity and/or a reduced activity of the respiratory chain. Because we found that oxidative phosphorylation was decreased (mitochondrial potential and ATP/ADP ratio) following UCP4 overexpression, we wanted to investigate whether the TCA cycle activity was reduced as well. To test this hypothesis, we measured the CO$_2$ production from glucose radioactively labeled in C1 or C6 to differentiate CO$_2$ production caused by the TCA cycle from total CO$_2$ produced. We observed that CO$_2$ production by the TCA cycle was increased by 31% in the presence of overexpressed UCP4 (Fig. 4B). Overexpression of UCP5 had no significant effect.

**Effects of UCP4 and UCP5 on Mitochondrial Respiration in Astrocytes and Relation to Mitochondrial Acidification**—Having determined that UCP4 and UCP5 had an uncoupling activity by modifying the mitochondrial potential, the ATP/ADP ratio, the H$_2$O$_2$ level, and NAD/NADH ratio and that UCP4 also affected the TCA cycle, we set out to determine the role of UCPs in the final element of the electron transport chain, namely the oxygen consumption. We analyzed the effect of overexpressed or silenced UCP4 and UCP5 in astrocytes on the OCR. We measured the respiration profile of astrocytes for 2 h including application of various compounds. We compared the OCR of control astrocytes and astrocytes overexpressing or silenced for UCP4 after 200 μM glutamate stimulation (Fig. 5A). We had previously shown that glutamate reduced the OCR in astrocytes, a phenomenon dependent on the mitochondrial matrix acidification (20). We could thus assume that if the mitochondrial matrix acidification and the mitochondrial potential reduction were linked and somehow controlled via an UCP, we could expect that the astrocytic OCR decrease in the presence of glutamate would be enhanced when overexpressing uncoupling proteins. We therefore measured the basal and glutamate-stimulated OCR in astrocytes overexpressing or silenced for UCP4 and UCP5. Glutamate application on astrocyte reduced OCR by 11% compared with basal level (Fig. 5A). The OCR represented in Fig. 5B are showing that in normal medium astrocyte overexpressing UCP4 do have and higher OCR, because the ATP/ADP ratio is reduced and mitochondrial potential is reduced. The silencing of UCP4 in these conditions also shows a higher OCR than in control condition, because we made these respiration measurements in the presence of deoxy-glucose, which blocks glycolysis. Abundance or reduction of UCP4 in astrocyte modified the OCR response to glutamate (Fig. 5C). Indeed, in the presence of glutamate, overexpression of UCP4 decreased OCR by 16%; conversely, silencing of UCP4 decreased the OCR by only 2%.

As previously mentioned, the effects of glutamate on oxygen consumption rate were shown to depend on the mitochondrial matrix acidification (20). We tested the implication of uncoupling proteins on this acidification by measuring the mitochondrial pH in astrocytes overexpressing UCP4 or UCP5, before
and after stimulation by 200 μM glutamate. Overexpression of UCP4 reduced significantly the basal mitochondrial matrix pH from pH 7.31 ± 0.06 in control cells to pH 7.05 ± 0.08 in infected cells (Fig. 6). UCP4 silencing did not significantly affect basal mitochondrial matrix pH, nor did UCP5 overexpression or silencing. In control cells, glutamate application decreased mitochondrial matrix to pH 6.64 ± 0.1, whereas UCP4 silencing decreased pH to 6.88 ± 0.05, therefore causing a milder acidification. UCP5 did not significantly modify the basal or the glutamate-stimulated matrix pH neither in overexpression nor in silencing. These results indicate that UCP4 is involved in the glutamate-induced mitochondrial matrix acidification, which is the likely cause of its effect on mitochondrial respiration.

UCP4-mediated Lactate Release from Astrocytes Improve Neuronal Survival—It has already been demonstrated that astrocyte-derived lactate is an important energy substrate for neurons (12, 38–41). As an index of metabolic activity in neurons, we measured the OCR of neurons exposed to the two most relevant energy substrates, namely glucose and lactate. As reported in Fig. 7A, the OCR is markedly increased in neurons exposed to lactate (50 mM) alone compared with a mix of glucose (12.5 mM) and lactate (25 mM) or to glucose alone (25 mM). All substrates or mixtures thereof were tested at equicaloric final concentrations. Given the role of astrocytes in providing lactate to neurons, we set out to study the possible involvement of astrocytic uncoupling protein level of expression on lactate production and in neuronal survival.

The release of lactate, accumulated for 24 h, was increased by 16% in astrocytes overexpressing UCP4 compared with control astrocytes (Fig. 7B, black columns). UCP5 overexpression had no significant effect on lactate release. Silencing of UCP4 or UCP5 reduced lactate release to 88 ± 4.2 and 80 ± 2.6% of control values, respectively (Fig. 7B, gray columns).

Because neurons exposed to lactate increased their respiratory rate (Fig. 7A), we measured the viability of neurons cocultured with astrocytes overexpressing or silenced for UCP4/UCP5. Overexpression of UCP4 or UCP5 in astrocytes increased the neuronal survival by nearly 20–35% compared with control astrocytes (Fig. 7C, black columns). In contrast, UCP4 or UCP5 silencing in astrocytes reduced significantly the viability of co-cultured neurons by 33 and 24%, respectively (Fig. 7C, gray columns).

DISCUSSION

The decoupling of proton gradient dedicated to the production of ATP in favor of a decrease in entropy and hence an increase in temperature is a well described (1, 42, 43) phenomenon in brown adipose tissue and is caused by the presence of UCP1 in this tissue. In brain cells, UCP1 is marginally expressed, and its possible role in the CNS remains unclear.

The UCP2 isoform role in the brain is still debated, but it appears to be linked either to controlling the local temperature to increase conduction velocity (4) or to controlling ROS production (5). Regarding this control of ROS by UCP2, this role
seems to be measured not only in the CNS, but in most studied tissues. UCP3 isoform appears to be expressed exclusively in muscle and is found certainly in a nonspecific manner in the brain. Our overexpression of the five isoforms with an HA tag in astrocytes in culture was used to validate the method of expression of the protein and also allowed confirmation that none of the commercial antibodies directed against one or the other UCPs isoforms was really specific.

The primary aim of this study was to elucidate the role of UCP4 and UCP5 isoforms in the CNS. Localization of UCP4 and UCP5 in the mouse brain (16, 19) demonstrates an important RNA expression that is likely to be translated into a functional role. Brain cell metabolism is tightly regulated, and its perturbation can lead to dysfunction and maybe to neurodegenerative diseases as discussed by Andrews et al. (44). It therefore becomes important to understand UCP4 and UCP5 functions, mechanisms of action, and, in particular, their involvement in metabolism in astrocytes, given the intimate metabolic relationship between astrocytes and neurons.

We first confirmed the earlier observation (21) that glutamate leads to a stimulation of glucose uptake by 20% and found that the same happens in astrocytes overexpressing UCP4 or UCP5. However, when endogenous UCP4 was reduced using sh-UCP4 (Fig. 2A), this enhancement of glucose uptake caused by glutamate did not occur. This observation was parallel to lactate release, which was similarly influenced by UCP4. Thus, we hypothesized that UCP4 operates a control on respiration (Fig. 5A) by tuning intramitochondrial pH level as we observed (20) in the presence of glutamate.

The reduced level of glycogen in astrocytes overexpressing UCP4 or UCP5 may be a consequence of the mitochondrial uncoupling observed and hence to the reduction of the available ATP in cell. One ATP is necessary to transform glucose to glucose 1-phosphate, which is the first step in the glycogen synthesis.
At the astrocyte mitochondrial level, the effects observed during the overexpression of UCP4 or 5 are decreased mitochondrial electrical potential, decreased ATP production, and decreased H₂O₂ production (Fig. 8). One can deduce from these results that overexpression of UCP4 or 5 decreases the efficiency of oxidative phosphorylation. Under these conditions, the Krebs cycle is enhanced, which results in a decrease of the NAD/NADH ratio, and also by an increase in the CO₂ production. This increase in NADH and CO₂ from the Krebs cycle is possible because there is no exhaustion of the substrates (ADP, NAD, and acetyl-CoA) and no negative regulation by ATP. It also indicates that the production of reducing equivalents (NADH) has not reached the threshold of inhibition as evidenced by the accumulation of CO₂ coming from the Krebs cycle that keeps functioning.

All mitochondrial indices tested (ATP production, H₂O₂ production, mitochondrial potential, and NAD/NADH ratio) were confirmed by the opposite effects elicited by UCP4 or UCP5.
silencing, demonstrating the essential role of UCPs in astrocytes in the mitochondrial respiration control. According to the results presented in this study, the mode of action of UCP4 on mitochondria in astrocytes can be summarized in an uncoupling of oxidative phosphorylation. It is likely that the same applies to UCP5, also expressed by astrocyte mitochondria.

The next issue is to identify a functional role for these UCPs. We have previously shown (20) that astrocytes OCR decreased by 20% following glutamate application and that this decrease in respiration was due to mitochondrial acidification. We now show that the decrease in pH is controlled by UCP4. Overexpression of UCP4 actually decreased basal intramitochondrial pH, and glutamate application caused an additional acidification to reach the same pH 6.6 as in the nontransfected astrocytes. This pH value likely represents the lower limit of matrix acidification. This decrease in pH in all cases had the same effect of reducing the respiration by 20%. The demonstration of the role played by UCP4 in controlling respiration via the intramitochondrial pH is even more striking when we reduced the UCP4 expression. In this case, the intramitochondrial pH drop was less pronounced in the presence of glutamate, and respiration is consequently not decreased to the same extent.

The results of the UCP5 experiments do not allow clear-cut conclusions regarding its mode of action. It is noteworthy that UCP5 can be found in two different variants (9). For this study, we cloned the full-length sequence in astrocyte; it is conceivable that the UCP5 short variant is the form present in astrocytes, whereas the long form is present in neurons, or the other way round. Further investigations need to be undertaken to clarify by which mechanism astrocyte UCP5 increases neuronal viability.

UCP4 function would then be that of reducing astrocyte respiration in the presence of glutamate, through a decrease in intramitochondrial pH. To compensate for the impaired oxidative energy production, astrocytes enhance glycolysis, and therefore their production of lactate. Lactate produced by astrocytes is used by neurons, as demonstrated by the OCR increase caused by lactate in neuronal cultures and has the beneficial effect of enhancing their survival rate, as shown in Fig. 7C.

**CONCLUSION**

In summary, the release of glutamate in synapses and reuptake by astrocytes induces a mechanism of decreased respiration in astrocyte, an effect mediated by UCP4, and possibly UCP5, whose consequence is to promote energy production by glycolysis for producing preferentially lactate that can be in turn used by neurons (Fig. 8). Overall, control of pH by the intramitochondrial UCP4 in astrocytes may be considered as a mechanism that helps sustain neuronal survival. Therefore, it is not surprising that a modification in UCP4 expression was found at the onset of neurodegenerative diseases (11, 45, 46). This work explains for the first time a role for the presence of UCP4 in the brain.
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