INTRODUCTION

Birth is among the most complex biological mechanisms in mammals, with major and abrupt hormonal, physiological, immunological, microbiotal, and vascular changes. It is associated with a massive release of stress molecules and hormones, including oxytocin required to trigger labor and mother-infant interactions and to compensate for the potentially deleterious actions of stress (1). Astonishingly, we have little information on how the brain is prepared for and endures this critical step. Several indirect lines of evidence obtained in humans suggest that brain growth velocity slows during the third trimester, possibly in an effort to avoid an exacerbated brain volume during vaginal delivery (2). This is followed by a spurt of growth after birth that correlates with synapse density (3). Despite their importance, human epidemiological investigations fall short in determining the nature of the events occurring during labor and birth. To understand whether and how the brain is prepared to meet that challenge, experimental studies comparing brain and neuronal parameters shortly before and after birth are essential. Gaining a better understanding of the cellular events occurring during labor and birth is also important from a clinical and public health perspective. Thus, epidemiological studies suggest higher incidence of autism spectrum disorders (ASD) and other developmental disorders following birth-related alterations, including C-sections (4), preterm delivery (5), or delivery complications (6). Experimental observations also suggest that labor and birth are critical periods, playing an important role in the pathogenesis of brain disorders (7).

Here, we compared neuronal morphology and brain volume shortly before and after birth in control (CTL) animals and observed that they are similar, suggesting a “growth stop signal” in preparation for labor and birth. We then tested the hypothesis that this stop signal is alleviated by intrauterine injections of valproate (VPA) in a timed pregnant rat. In this model, we observed a growth during the same short period. We showed that hippocampal neuronal morphology and cerebral volumes, including in the hippocampus and neocortex, were increased by labor and birth in ASD conditions. The NKCC1 chloride importer antagonist bumetanide, which attenuates ASD severity in children (8) and animal models of ASD (9, 10), reduced some of these changes, suggesting that high levels of chloride during labor and birth are pathogenic.

RESULTS

CA3 pyramidal neuronal growth is altered in the VPA condition

Using a rat model of ASD by injection in utero of VPA (9), we first reconstructed CA3 hippocampal pyramidal cells. These neurons have been extensively investigated during the perinatal period to determine electrical and behavioral features of ASD (9, 11). Referring to birth as the period between embryonic day 21 (E21) and the first postnatal day P0 (-24 hours) that includes labor and birth, we compared E21 CTL versus E21 VPA, E21 CTL versus P0 CTL, E21 VPA versus P0 VPA, and P0 CTL versus P0 VPA conditions (table S1). Despite a high degree of heterogeneity (Fig. 1, A to D, and figs. S1 and S2), we found no significant differences between E21 and P0 CTL neurons in all parameters investigated (Fig. 1, E to I, and figs. S3, A to C). There was a trend for E21 VPA neurons to be smaller than E21 CTL neurons but without reaching statistical significance, except for the Sholl profile (Fig. 1E). On the other hand, the Sholl profile of P0 VPA neurons was shifted to higher values [E21 CTL versus E21 VPA (P < 0.0001), E21 CTL versus P0 CTL (P = 0.0684), E21 VPA versus P0 VPA (P < 0.0001), and P0 CTL versus P0 VPA (P = 0.0163), Kolmogorov-Smirnov test; Fig. 1E]. In P0 VPA neurons, the total apical dendrite length (Fig. 1F), enclosing radius (Fig. 1G), and critical radius (Fig. 1H) were significantly higher compared with those in P0 CTL [P = 0.0156, 0.0110, and 0.0079, respectively; Kruskal-Wallis tests followed by Dunn’s posttests (KWD)] and E21 VPA (P < 0.0001, P < 0.0001, and P = 0.0004, respectively, KWD). However, the number of apical nodes (Fig. 1I) was not different from P0 CTL or E21 VPA, indicating that the apical arbor complexity remained unchanged. Soma volumes were also significantly increased in P0 VPA compared with E21 VPA (P = 0.0027, KWD;
**Fig. 1. CA3 pyramidal neuron morphology is affected during birth in the VPA condition.** (A to D) Representative neuron reconstructions from E21 CTL (A), E21 VPA-treated (B), P0 CTL (C), and P0 VPA-treated (D) rats; axons are presented in blue. Scale bar, 50 µm. (E) Sholl profiles of apical dendrites illustrating shifts in the number of crossing dendrites along the radial axe occurring in the VPA conditions. (F to I) Mean values for the total apical dendrite length (F), the apical dendrite enclosing radius length (G), the apical dendrite critical radius (H), and the number of apical dendrite nodes (I). Error bars denote SEM. Kruskal-Wallis test followed by Dunn’s posttest, *P < 0.05, **P < 0.01, ***P < 0.001, and ****P < 0.0001. ns, not significant (P > 0.05). (J and K) Long versus short cell clustering based on apical morphology. (J) Percentage of neurons distributed in the long cell cluster (the short cell cluster proportion is shown in white bars on top) for each condition. (K) Histogram showing the distribution of the long cell cluster proportion in 1000 randomizations of cell affiliation among all conditions where E21 and P0 VPA are found out of the range of possibilities for CTL conditions. Dashed lines indicate the 99% confidence interval.
Microglia do not undergo major alterations during birth

Converging evidence from many studies suggests an involvement of microglia in neurodevelopmental disorders including autism (14). Since microglia can affect progenitor cell proliferation and neuronal growth during brain development (15), we assessed if these cells could be affected during birth in CTL and VPA animals. Using the iDISCO clearing method (16), we counted, for each condition, the total number and computed the density of IBA1 immunolabeled cells before and after birth for each condition in four different structures, including the hippocampus, neocortex, dorsolateral striatum, and whole brain (movie S1 and table S3). In CTL and VPA pups, the densities of microglia in these structures were similar before and after birth (Fig. 2, A and C, and fig. S4, A and B). Microglia activation state was then evaluated on the basis of the cellular area and volume (table S3) (17). Despite a significant reduction of hippocampal microglia activation in P0 VPA animals compared with P0 CTL [area, P0 CTL versus P0 VPA (P = 0.0182); volume, P0 CTL versus P0 VPA (P = 0.0193); KWD], we observed no differences in other brain structures for each condition (Fig. 2, B and D). Together, these results demonstrate that only hippocampal microglia activation state is affected in VPA neonates, suggesting a potential effect of VPA linked to birth. However, to which extent these cells might contribute to the CA3 neuronal growth remains to be clarified.

Volumetric alterations around birth in the VPA condition

Toddlers, children, and adolescents with ASD have bigger brains than normotypic ones (18–20), but whether these changes occur around birth is not known. To evaluate this, we used the same clearing method to compare, before and after birth, the volume of the hippocampus, neocortex, dorsolateral striatum, lateral ventricles, and whole brain in CTL and VPA-treated rats (movie S2 and table S4). In CTL rats, these volumes were not significantly different before and after birth [hippocampus, E21 CTL versus P0 CTL (P = 0.7944); whole brain and neocortical volumes of P0 VPA pups, abolishing the volume increase observed during birth in the VPA condition [hippocampus: P0 VPA versus P0 VPA + BUM (P = 0.0116); neocortex: P0 VPA versus P0 VPA + BUM (P = 0.0242); KWD] (Fig. 3B). Maternal bumetanide treatment also shifted the distribution of cerebral volumes from lognormal back to normal in the population of VPA brains, restoring smaller cerebral structure volumes (Fig. 3C). It also decreased the CA3 volume to CTL level after birth, suggesting that the increased growth observed in this region could be mediated by the excitatory actions of GABA (Fig. 3D). Therefore, maternal bumetanide administration prevents the enhanced growth observed in VPA animals during birth.
Fig. 2. Microglia density is not affected but activation state is reduced at birth in the VPA condition. (A) Representative three-dimensional (3D) pictures of hippocampal (top), neocortical (middle), and whole-brain (bottom) Iba1 immunoreactive cells (in green) and of automatically detected objects (in red) for E21 CTL, E21 VPA, P0 CTL, and P0 VPA conditions. Scale bars, 600 μm. (B) Representative pictures of morphometric-based analysis of hippocampal and neocortical microglia. An alteration of global cellular morphology (including soma and part of the branching processes) is visible in the hippocampus in the P0 VPA group compared with the CTL group, compatible with a reduced activation state. Iba1, green; detected surfaces superimposed, red. Scale bars, 100 μm. (C) Hippocampal, neocortical, and the whole-brain quantification showing no alterations of microglia density (number of cells per mm³) in VPA animals compared with the CTL ones. (D) In the hippocampus, VPA treatment induces a significant decrease of microglial area and volume parameters at birth, correlated with a reduced activated state of these cells. Error bars denote SEM. Kruskal-Wallis test followed by Dunn's posttest, *P < 0.05.
Fig. 3. VPA treatment induces volumetric alterations of the hippocampus, whole brain, and neocortex around birth. (A) Average shape of the hippocampus (top), neocortex (middle), and whole brain (bottom) for E21 CTL, E21 VPA, P0 CTL, P0 VPA, and P0 VPA + BUM conditions. Scale bar, 1 mm. (B) Volume measurements of the hippocampus, neocortex, and whole brain. Error bars denote SEM. Kruskal-Wallis test followed by Dunn’s posttest, *P < 0.05, **P < 0.01, and ****P < 0.0001. ns, not significant (P > 0.05). (C) Distribution of normalized volumes of all structures in each experimental group. After birth in the VPA group, the distribution is changed to lognormal, and normal law is restored by bumetanide. (D) Representation of significant local size changes between groups of interest. Kruskal-Wallis test followed by Dunn’s posttest (P < 0.05, corrected with the Benjamini-Hochberg procedure for multiple comparisons). Red-yellow colors show significant size increase, blue-green colors indicate significant size decrease, and gray color is assigned to regions with nonsignificant size changes. Representative segmented CA3 region in yellow (right).
DISCUSSION
Neuronal and brain “growth stop signal” in preparation to birth

To the best of our knowledge, this is the first description of global brain changes around birth in CTL and in an ASD rodent model. Birth being one of the most complex biological mechanisms in mammals, it is intuitively expected that some alterations in brain functions occur in preparation for it. Yet, despite its importance, our knowledge of the events occurring shortly before and after birth is very limited. Volumetric fluorescence microimaging of whole organs provides a new approach to study how tissues and cells develop and respond to disease. In CTL rats, there is no difference between shortly before and after birth, indicating that brain growth is halted or at least does not occur during that period. We also show that CA3 pyramidal neuron apical dendrites and somata are similar at E21 and P0. The volume of structures, including the hippocampus, neocortex, and striatum, is also unaltered, suggesting a common sequence. Astonishingly, the ventricle volumes are increased. More investigations are much needed to determine whether a given structure possibly indispensable for the functions that must be executed at birth (crawling, suction, and smell) might not show this lack of growth during this period.

Intrauterine VPA abolishes the growth stop signal during birth

In the VPA condition, CA3 pyramidal neurons in offspring at birth had longer apical dendrites and bigger somata but showed no changes in basal dendrites. The volumes of the hippocampus and neocortex were also significantly increased, but not the lateral ventricles, in contrast to CTLs, suggesting major changes of the neuronal during birth. Major morphological alterations have been reported in animal models and postmortem investigations in patients with ASD. Thus, in the VPA rat model, dendritic arborization of neurons in layer 2 of the motor cortex is more complex than in CTL condition (21). Following intrauterine VPA, the number of spines is reduced in the hippocampus, the prefrontal cortex, and the basolateral amygdala but increased in the nucleus accumbens and layer 3 of the prefrontal cortex (22). Maternal immune activation that is associated with increased incidence in ASD also leads to profound cortical malformations that are the direct cause of the deleterious social impairments (23, 24). Postmortem studies in children with ASD (2 to 15 years old) revealed focal patches of abnormal laminar cytoarchitecture and cortical disorganization of neurons (25). Individuals with ASD show at birth an initial excess of mature neurons (26). Similarly, neuron number in the prefrontal cortex was increased by 67% in children with ASD (2 to 16 years old), and brain weight was bigger than in age-matched CTLs (27). Collectively, these studies suggest that the variety of events leading to ASD also lead to morphological cortical abnormalities. In addition, children with ASD have a bigger brain (28, 29). Thus, 15% of boys with ASD have megalencephaly, with a brain size that is disproportionate to the body (18). Also, brain imaging functional connectivity differs already at 6 months of age in babies with ASD (30). The brains of children with ASD undergo an abnormal brain trajectory that includes a period of overgrowth (28). The authors suggest that the enlargement starts at the end of the first postnatal year, but direct evidence on this issue is lacking. Neither the earlier occurrence of these changes nor whether birth constitutes a flag point in this sequence is understood at this time. Present results suggest that bigger brain structures and enhanced neuronal growth are conspicuous early on and that at least in the VPA model, neurons grow and brain structures increase their volumes during birth. The intrauterine insults by a not yet understood mechanism inactivate a putative neuroprotective mechanism occurring during this vulnerable period.

Bumetanide-sensitive excitatory GABA and growth during birth in the VPA condition

The reduction of the volume changes produced in offspring by maternal bumetanide suggests that GABAergic signals and, notably, neuronal [Cl−] levels might contribute to these volume changes. The volumes of the hippocampus, neocortex, and lateral striatum were not increased after treatment, in contrast to CTLs. GABA exerts a trophic action on development, including cell proliferation (31), neuronal migration (32), differentiation (33), network formation (34, 35), and dendritic growth (36–39). This phenomenon is mediated by the excitatory actions of GABA that activate voltage-gated calcium currents and increase intracellular calcium levels, releasing growth factors, notably BDNF (brain-derived neurotrophic factor), that modulate neuronal growth and dendritic extension (40, 41). These effects are mediated by the high [Cl−] levels present in immature neurons and require active NKCC1 chloride import, as they are blocked by the highly specific antagonist bumetanide or by NKCC1 shRNA (short hairpin RNA) manipulations to invalidate the importer (37, 39, 42). Furthermore, in human and mice embryos, there is a depolarizing GABAergic projection from the zona incerta to cortical layer 1 that stimulates dendritic growth from deeper cortical layers, and silencing this activity decreases dendritic growth (38). Therefore, high [Cl−] levels and excitatory actions of GABA are indispensable for embryonic neuronal growth.

However, during birth, this excitatory action might exert potentially deleterious actions. Birth is associated with a massive release of stress molecules (43) that augment [Cl−] levels, leading to neuronal hyperexcitability and potential seizures during a highly vulnerable period (44, 45). Oxytocin induces during birth an abrupt depolarizing-to-hyperpolarizing shift of GABA actions associated with a reduction of [Cl−] levels exerting an antistress action (11). This phenomenon was abolished in the VPA model of ASD and in fragile X mice (9) but was restored by maternal administration of bumetanide during birth, which also attenuated the severity of ASD (9, 10). Here, we show that bumetanide possibly restores the brain growth stop signal, suggesting that the high [Cl−] levels during this critical period are instrumental in the long-term deleterious sequelae of ASD. Bumetanide attenuates ASD severity in children and adolescents (8).

Therefore, the neuroprotective oxytocin-mediated mechanism is alleviated in the VPA model of ASD; GABA depolarizes and excites neurons leading to growth factor release, neuronal growth, and enhanced hippocampal and neocortical volumes. Growth during labor and birth might underlie the defects in neuronal and network wiring and affect an essential period of sensory acquisition and social interactions. Birth itself acts as a triggering signal to accelerate cortical sensory maps and major early functions including suction (46, 47). It is suggested that the reduction of the inhibitory actions of GABA and the aberrant synchronized activity generated consequently affect this operation, leading to long-term sequelae.
CONCLUSION
Despite the intrinsic limitations of species differences, our results stress the need to consider birth as a critical period in the pathogenesis of ASD (7). The present approach provides means to compare genetic or environmental intrauterine insults and determine common rules between different disorders.

MATERIALS AND METHODS
VPA rat model of autism
All experiments were performed in accordance with the European Community Council directives regarding the protection of animals used for experimental and scientific purposes (2010/63/EU).

Time-mated female Wistar rats, housed under standard laboratory conditions, were treated by in utero VPA injection to generate ASD in offspring (9). Briefly, the sodium salt of VPA (Sigma-Aldrich Inc., USA) was dissolved in physiological saline solution to a concentration of 300 mg/ml. Pregnant rats received a single intraperitoneal dose of 600 mg/kg on gestational day 12 (E12). CTL animals were not manipulated. In our laboratory conditions, the day of delivery was on E22, without differences between CTL and VPA-treated females. Bumetanide treatment (2 mg/kg; Sigma-Aldrich) was given to the dams in drinking water at least 12 hours before birth. The offspring were used for experiments on E21 and postnatal day 0 (P0 = E22). Experiments were performed on animals of both sexes. E21 embryos were extracted from the dams by hysterectomy after isoflurane anesthesia (Aerrane, Baxter, France), followed by cervical dislocation. Newborn P0 pups were decapitated shortly after delivery. For morphology, brains were immediately placed in an ice-cold oxygenated solution for slicing. For iDISCO experiments, brains were postfixed in Antigenfix (Diapath, Italy) for overnight fixation and stored at 4°C in phosphate buffered saline (PBS). Thermo Fisher Scientific Inc., USA) with 0.05% sodium azide (Sigma-Aldrich) until use. For each embryo and pup, a biopsy of the tail was kept for sex genotyping (chromosome Y).

Slice preparation and revelation process of biocytin-filled pyramidal neurons
We prepared horizontal brain slices (350 μm thick) from E21 and P0 male and female rats using a Leica VT1200 S (Leica Microsystems, Germany) vibratome in ice-cold oxygenated solution containing 132.5 mM choline chloride, 2.5 mM KCl, 1.23 mM NaH2PO4, 3 mM MgCl2, 0.7 mM CaCl2, 25 mM NaHCO3, and 8 mM glucose. Slices were then left to recover for 1 hour at room temperature (22° to 25°C) in artificial cerebrospinal fluid containing 126 mM NaCl, 3.5 mM KCl, 1.2 mM NaH2PO4, 2 mM CaCl2, 1 mM MgCl2, 11 mM glucose, and 25 mM NaHCO3 (pH 7.4). All solutions were bubbled with 95% O2/5% CO2. Borosilicate glass pipettes (4 to 7 megohm) were filled with a solution containing 130 mM K-glucosone, 10 mM Na-glucosone, 7 mM NaCl, 4 mM Mg-ATP (adenosine 5′-triphosphate), 10 mM Heps, 4 mM phosphocreatine, and 0.3 mM Na-GTP (guanosine 5′-triphosphate) (pH 7.3) with KOH, 285 to 290 mOsm. Biocytin (0.5 to 1%; Sigma-Aldrich) was added to the solution just before patching the cells.

After biocytin diffusion into the cells (10 min), slices were transferred into Antigenfix for overnight fixation. Aldehyde fixation causes nonuniform brain tissue shrinkage (48), making it very difficult to apply any correction factor. Thus, special care was taken to respect the exact equal time of fixation and processing for each slice to decrease, as much as possible, the shrinkage variability (49).

After fixation, slices were sequentially washed in PBS, 0.3M glycine in PBS, and PBS. They were incubated for 1 hour at room temperature in PBS containing 10% of normal goat serum (NGS; Jackson ImmunoResearch Inc., USA), 1% bovine serum albumin (BSA; Sigma-Aldrich), and 0.3% of Triton X-100 (Sigma-Aldrich). They were then incubated overnight at 4°C, with streptavidin coupled with Alexa 647 (1/1000; Thermo Fisher Scientific Inc., USA) diluted in PBS containing 1% NGS and 1% BSA and 0.3% of Triton X-100. CTL slices without any biocytin were systematically added in each experiment to check the streptavidin specificity. Last, slices were washed in PBS and incubated for 10 min at room temperature with Hoechst staining solution (1/1000; Sigma-Aldrich) diluted in PBS. They were washed in PBS, mounted on slides, and coveredslipped using Fluoromount-G (Electron Microscopy Sciences, USA) as mounting medium.

Image acquisition and morphology analyses
Confocal images were acquired on an SP5X Leica microscope (Leica Microsystems) using the 647-nm band of a white laser for excitation of Alexa 647 (spectral detection, 657 to 750 nm) and a diode 405 nm for excitation of Hoechst staining (spectral detection, 411 to 479 nm). High-magnification images (1024 × 1024 pixel) of Alexa 647 signal were acquired at 200 Hz using a 63× oil immersion objective, pinhole set to “Airy 1.” Stacks for dendrite analysis were obtained by scanning with a line average of 2 and a z step of 0.50 μm.

Morphology analyses were blindly performed to avoid any bias. Neurons were selected according to their localization in the stratum pyramidale of CA3 and the quality of their staining: dying cells, damaged cells, and interneurons were systematically excluded. Patches were done in the entire CA3 area to guaranty homogeneity in cell origins between ages and conditions. Thus, similar proportions of cells were found localized in the CA3a, b versus CA3c subregions (less than 4% of difference; fig. S3D). The position of each cell was noted, as it was shown that CA3a, b, and c exhibit a gradient in pyramidal cell morphology (50). Somata depth positions inside the stratum pyramidale were also carefully noted. In juvenile and adult rodents, deep pyramidal cells (near stratum oriens) are known to be older and longer than superficial neurons (near stratum lucidum/ radiatum; (51, 52)).

For dendrite and somata analysis, stacks were imported in the open-source platform Fiji (1.50e, Java 1.8.0_60, 64 bit; https://fiji.sc/) and stitched together (https://imagej.net/Image_Stitching). Without any additional preprocessing, somata reconstruction was performed using the ROI manager tool, and dendrite reconstruction and measurements were performed using the semiautomatic “Simple Neurite Tracer” plugin (http://imagej.net/Simple_Neurite_Tracer). On the basis of the reconstructions, a Sholl analysis of apical dendrites was then performed using the “Sholl Analysis” plugin (http://imagej.net/Sholl_Analysis). Apical morphological parameters (table S1) were also used to perform a cluster analysis.

K-means clustering was performed in MATLAB (Mathworks Inc., USA) using Euclidean distance, 100 times iterations and 100 times replicates. Clustering was aimed at creating two clusters of cells based on the six following morphological properties (variables): total apical dendrite length, apical dendrite nodes, apical primary dendrite length, ending radius, critical radius, and maximum crossing dendrites. Each variable was normalized by its own maximum, with
resulting values between 0 and 1. In each cluster, the average total apical dendrite length was used as reference to identify the cluster with longer and the one with shorter average total apical dendrite length. To create a surrogate statistic on the clustering analysis, the group labeling of the cells (i.e., E21 CTL, E21 VPA, P0 CTL, and P0 VPA) was reassigned randomly between the list of cells while keeping the same number of cells per group. The same clustering procedure was then performed over 10,000 surrogate cell lists, and the distribution of the cells in each cluster (long versus short neurons) per group was calculated so that a 99% confidence interval could be estimated.

### iDISCO protocol

Whole-brain clearing was performed according to the previously reported 3DISCO and iDISCO+ clearing procedures (16, 53, 54) with slight modifications. Samples were dehydrated in a graded series (20, 40, 60, 80, and 100%) of methanol (Sigma-Aldrich) diluted in PBS during 1.5 hours each, at room temperature. They were then incubated overnight at room temperature on a platform shaker in a solution of PBSG-T [PBS containing 0.2% gelatin (Sigma-Aldrich), 0.5% Triton X-100, 0.02% sodium azide (Sigma-Aldrich)] for 2 days. Next, samples were transferred to PBSG-T containing Iba1 primary antibody (1:500; rabbit anti-Iba1, Wako Chemicals Inc., USA) to label microglia and placed at 37°C, with rotation at 100 rpm, for 10 days. This was followed by six washes of 1 hour in PBSG-T at room temperature. Samples were then incubated in secondary antibody (donkey anti-rabbit Alexa 647) diluted in PBSG-T for 2 days at 37°C. After six washes of 1 hour in PBSG-T at room temperature, samples were stored at 4°C in PBS until use. For tissue clearing, all incubation steps were performed at room temperature using a 15-ml polypropylene tube (Dutscher SAS, France) covered with aluminum foil to avoid exposure to light. Samples were dehydrated in a graded series (20, 40, 60, 80, and 100%) of methanol diluted in PBS during 1 hour. This was followed by a delipidation step of 20 min in dichloromethane (DCM; Sigma-Aldrich). Samples were then transferred to 100% DCM until they have sunk. Last, samples were cleared overnight in dibenzylether (DBE; Sigma-Aldrich) and stored in DBE at room temperature in the dark.

### 3DISCO imaging and image processing

Three-dimensional (3D) imaging was performed with an ultramicroscope (LaVision BioTec GmbH, Germany) using InspectorPro software (LaVision BioTec). The light sheet was generated by a laser (wavelengths, 488, 555, and 647 nm; Coherent Sapphire Laser, LaVision BioTec) and two cylindrical lenses. A binocular stereomicroscope (MXV10, Olympus, Japan), with a 2× objective (MVPLAPO, Olympus), was used at magnification ×2.5. Samples were placed in an imaging tank made of 100% quartz (LaVision BioTec) filled with DBE and illuminated from the side by the laser light. Images were acquired with a pcO.edge sCMOS charge-coupled device camera (2560 × 2160 pixel size, LaVision BioTec). The step size between each image was fixed at 2 μm. 3D image quantifications were generated using Imaris x64 software (version 9.2.1, Bitplane, Switzerland). Each resulting Imaris file was 16-bit images and was subsequently converted to 8 bits. A 3D reconstruction of each sample was performed using the volume rendering tool (Imaris). Segmentation-based tracing of the whole brain, neocortex, hippocampus, dorsolateral striatum, and lateral ventricles was performed using the Imaris “Surface” module in manual creation mode. Border delineation of each structure was achieved on the autofluorescent channel generated by a 488-nm light exposure. Anatomical frontiers of each brain structure were followed according to the Allen Brain Atlas [Allen Institute (55)] and adapted to the developing rat brain. 3D pictures were generated using the “snapshot” tool (Imaris).

Image registration to compute average shapes of the whole brain, hippocampus, and neocortex was performed using a python script, MeshLab software (www.meshlab.net/), and the SPHARM module of 3DSlicer software (www.slicer.org/). First, the triangulation data of reconstructed surfaces were exported from Imaris. Then, the quality of mesh was checked by PyMesh library of python. When needed, we used MeshLab to modify the mesh quality and for visualization. Then, the surfaces in each group were registered using the SPHARM module. SPHARM establishes an optimized bijective mapping of each surface to the unit sphere. Consequently, for each point on a surface, corresponding points on other surfaces were specified. Last, surfaces were aligned on the basis of the first-order ellipsoid, and the average of each coordinate component was computed in python.

### Morphometric analysis of microglia

Automated microglia cell detection was performed with the Imaris “spots” feature using an estimated XY diameter of 20 μm, and a threshold was applied on the basis of the intensity at the center of the spot, Gaussian filtered by 3/4 of spot radius, allowing an accurate detection of visible cells without including the background signal. Measurement of microglia reactivity was performed according to previously reported work (18), which describes “resting” microglia as small with elaborate ramifications and the “activated” as bigger, more amoeboid cells with retracted processes. To quantitatively describe this morphological switch, we analyzed cell area and volume that are morphological parameters expected to be modified when cells become activated. This morphometric measurement of microglia activation was achieved with the Imaris “Surface” tool using a diameter of the largest sphere of 20 μm. Upper and lower thresholds of surface area and volume were determined as 3/2 of interquartile range below and above the first and third quartiles, respectively.

### Local shape analysis

To investigate the local size changes of the hippocampus, the shortest distance of each point of every hippocampal surface to an individual medial axis, created by the SPHARM module through the hippocampus, was computed (fig. S7) (56). The intergroup comparisons of local distance maps in corresponding points were performed by using the Kruskal-Wallis test followed by post hoc Dunn’s test (P < 0.05). The P values were corrected for multiple comparisons by using the Benjamini-Hochberg procedure. The significant local signed difference maps are shown in Fig. 3D.

### Movie processing

Movies were generated using Imaris x64 software (version 8.0.1, Bitplane). Stack images were first converted to an Imaris file (.ims) using ImarisFileConverter, and the 3D reconstruction was performed using the “volume rendering” function. Optical slices were obtained using the “orthoslicer” tool. To isolate a specific region of the tissue, the surface tool was used and the mask option was selected. Movies were then generated using the “animation” tools. Movie reconstruction with .tiff series was done with Fiji, and addition of titles and transitions have been done with iMovie (version 10.1.9, Apple Inc., USA).
Statistical analysis

Statistics were performed in Prism 6 (GraphPad Software Inc., USA). After checking normality using the Shapiro-Wilk test, multiple comparisons of parameters between each group were carried out using Kruskal-Wallis tests, followed by post hoc Dunn’s tests. Sholl profiles were compared using the Kolmogorov-Smirnov test, male versus female parameter comparisons were made using the Mann-Whitney test, and correlation analyses were performed using the Spearman test since data were nonparametric. Exact P values were specified when possible and considered significant when <0.05. Results are shown as means ± SEM in tables S1 to S4. Images were edited using Photoshop CC and Illustrator CC (Adobe Systems Software Ireland Ltd., Ireland).

**Supplementary Materials**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/5/1/eaa0394/DC1

Table S1. Morphological parameters based on dendrite reconstruction.

Table S2. Comparison of males (M) versus females (F) pyramidal cell morphology.

Table S3. Microglia number and density (per mm²) and area (μm²).

Table S4. Volume quantifications (μm³) for all brain structures.

Fig. S1. Reconstructions of CA3 pyramidal neurons of E21 and P0 CTL rats.

Fig. S2. Reconstructions of CA3 pyramidal neurons of E21 and P0 VPA-treated rats.

Fig. S3. Nonaopical pyramidal cell morphology alterations and cell position within the CA3 zone.

Fig. S4. Effect of birth on microglia density and volume in the dorsolateral striatum of CTL and VPA-treated rats.

Fig. S5. The IDISCO clearing method enables concurrent 3D volume analyses of brains of rat pups.

Fig. S6. Lateral ventricle changes among all conditions (E21 CTL, E21 VPA, P0 CTL, P0 VPA, and P0 VPA + BUM).

Fig. S7. Medial axis and distance map of hippocampus.

Movie S1. Spot detection of hippocampal microgliosis and morphometric-based microgliosis analysis.

Movie S2. Imaris segmentation of all brain structures.

**References and Notes**


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Data and materials availability: All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. Additional data related to this paper may be requested from the authors.

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