High dose folic acid supplementation of rats alters synaptic transmission and seizure susceptibility in offspring

Fernando Girotto*, Lucas Scott*, Yosef Avchalumov, Jacqueline Harris, Stephanie Iannattone, Chris Drummond-Main, Rose Tabias, Luis Bello-Espinosa, Jong M. Rho, Jörn Davidsen, G. Campbell Teskey & Michael A. Colicos

1Department of Physiology & Pharmacology, Faculty of Medicine, and the Hotchkiss Brain Institute, University of Calgary, Calgary, AB, Canada T2N 4N1, 2Pediatric Neurology and the Alberta Children’s Hospital Research Institute, University of Calgary, Calgary, AB, Canada T2N 4N1, 3Complexity Science Group, Department of Physics and Astronomy, Faculty of Science, University of Calgary, Calgary, AB, Canada T2N 4N1, 4Department of Cell Biology & Anatomy, Faculty of Medicine, and the Hotchkiss Brain Institute, University of Calgary, Calgary, AB, Canada T2N 4N1.

Maternal folic acid supplementation is essential to reduce the risk of neural tube defects. We hypothesize that high levels of folic acid throughout gestation may produce neural networks more susceptible to seizure in offspring. We hence administered large doses of folic acid to rats before and during gestation and found their offspring had a 42% decrease in their seizure threshold. In vitro, acute application of folic acid or its metabolite 4Hfolate to neurons induced hyper-excitability and bursting. Cultured neuronal networks which develop in the presence of a low concentration (50 nM) of 4Hfolate had reduced capacity to stabilize their network dynamics after a burst of high-frequency activity, and an increase in the frequency of mEPSCs. Networks reared in the presence of the folic acid metabolite 5M4Hfolate developed a spontaneous, distinctive bursting pattern, and both metabolites produced an increase in synaptic density.

The discovery that dietary folic acid supplementation before and during pregnancy can dramatically reduce the incidence of neural tube defects has been a major advance for child health 1,2. However since the neural tube closes at day 27 during human gestation3, high levels of folic acid after this point may serve no benefit, and could even be detrimental4. Folic acid and its related compounds are based on pteroic acid conjugated to L-glutamate, the latter being the major excitatory neurotransmitter in the brain, and folic acid itself has been previously demonstrated to have excitatory effects5–7. As there are glutamatergic synapses present in the hippocampus as early as the 3rd trimester8, and given the tight link between neuronal activity and synaptic development, we hypothesized that the continued presence of folic acid during neuronal development might alter the delicate balance of normal network formation and function.

Our experimental approach was as follows. First we demonstrated that folic acid can directly increase neuronal excitability by acutely applying it to cultured neurons while recording their activity. With this established, we pursued the hypothesis that folic acid exposure increases neuronal activity during development, possibly resulting in abnormal brain development, since many aspects of the establishment of neuronal connectivity are dependent on activity. We tested if administration of folic acid to pregnant rats leads to lowered seizure threshold in offspring, using a kainic acid model of seizure induction. We then used an in vitro primary neuronal culture model to investigate the mechanism in detail, in which we allow neuronal cultures to develop in the presence of low concentrations of 4Hfolate or 5M4Hfolate, replicating the in utero conditions. We chose these two metabolites of folic acid for the following reasons. The liver metabolizes folic acid to 4Hfolate, and so for the in vitro study this is the highest-level biologically active form we can use. When added to the media, 4Hfolate will in part be catabolized to pterin and p-aminobenzoylglutamate scission products, the latter of which will go on to produce L-glutamate. Enzymatically, 4Hfolate is also metabolized to 5M4Hfolate, a form commonly studied in terms of its effect on cellular physiology. Therefore in experiments using 4Hfolate we could determine the full spectrum of effects caused by folic acid, and by adding just 5M4Hfolate, we determine the effect of this specific metabolite alone. Similar to the in vivo experiments, we challenge the 4Hfolate and 5M4Hfolate reared cultures with a burst of high frequency activity and determine how well they re-stabilized their activity level. Previous reports have
suggested that folic acid can alter synaptic density\textsuperscript{6,7}, and so we look for structural changes in connectivity, using the distribution patterns of the synaptic maker synaptophysin. Finally, we analyze miniature EPSCs from neurons grown in the presence of 4Hfolate and demonstrate an increase in the frequency of the mEPSC events. We hypothesize that folic acid supplementation enhances neuronal activity and consequently the rate of development, potentially resulting in hyper-excitability and ultimately may lead to a network that is more vulnerable to seizures.

**Results**

**Folic acid and neuronal excitability.** Based on studies from the mid 1980’s\textsuperscript{3,4}, we hypothesized that exposing neurons to folic acid would alter their firing rate. Current clamp recordings were made from \( \sim 3 \) weeks in \textit{vitro} neurons in dissociated hippocampal culture networks, to which either folic acid, 4Hfolate or vehicle alone was acutely added. As previously stated, 4Hfolate was investigated since folic acid is first metabolized in the liver to this active form\textsuperscript{10}. When 10 mM concentrations of folic acid or 4Hfolate were applied to the neuron, a distinct change in the firing pattern of the cell was observed. In neurons which did not show a bursting pattern of activity during the initial recording period, both folic acid and 4Hfolate caused them rapidly (within minutes) to fire in bursts, whereas this was not observed with the controls (Figure 1a,b,c). This effect was protracted, being observable at least 10 minutes following this initial application of the drug (shown for folic acid in Figure 1D). These results suggest that folic acid and its derivatives have the potential acutely to alter the firing rate of individual neurons, and consequently alter the firing dynamics of a neuronal system.

**Maternal exposure to dietary folic acid during gestation.** The observation that folic acid and its derivatives could alter neuronal firing raised the possibility that its presence during gestation may affect development. To examine the potential \textit{in vivo} relevance of this we tested the hypothesis that maternal exposure to dietary folic acid during gestation may make the offspring more vulnerable to a challenge by a convulsant drug. Pups from dams that received folic acid is first metabolized in the liver to this active form\textsuperscript{10}. When added. As previously stated, 4Hfolate was investigated since folic acid is metabolized in the liver to this active form\textsuperscript{10}. When 10 mM concentrations of folic acid or 4Hfolate were applied to the neuron, a distinct change in the firing pattern of the cell was observed. In neurons which did not show a bursting pattern of activity during the initial recording period, both folic acid and 4Hfolate caused them rapidly (within minutes) to fire in bursts, whereas this was not observed with the controls (Figure 1a,b,c). This effect was protracted, being observable at least 10 minutes following this initial application of the drug (shown for folic acid in Figure 1D). These results suggest that folic acid and its derivatives have the potential acutely to alter the firing rate of individual neurons, and consequently alter the firing dynamics of a neuronal system.

![Figure 1](https://example.com/figure1.png)

**Analysis of network activity in 4Hfolate reared cultures.** We sought to determine if the presence of folic acid during the developmental establishment of neuronal connectivity affects the nature of the resulting network formed. To do this we grew neuronal networks from dissociated hippocampi from P0 rat pups (see methods) in media supplemented with low levels of 4Hfolate and characterized the activity patterns of the mature circuits. While several timepoints were tested, we chose 3 week old cultures as this amount of time allows the activity patterns to mature and become more consistent. Analysis of the firing events produces “fingerprints” of neuronal activity as illustrated in Figure 2a–d, where each individual neuron’s activity is represented on one horizontal line over time. A tick mark is added to the plot when a neuron fires. To demonstrate the effectiveness of this method, we monitored culture-wide activity during direct application of bath solution alone (Figure 2a) or 4Hfolate (Figure 2b). These fingerprints show how increased activity (dark vertical band after addition of 4Hfolate) and bursting (vertical bands of synchronous firing of the population of neurons several minutes after the addition of 4Hfolate) can be easily visualized in the entire network. To assay the cultures that had been reared in 4Hfolate, baseline activity was first recorded for two minutes, then all of the neurons in the field of view were stimulated for 30 seconds using photoconductive stimulation (indicated by the red strip on the fingerprints). We continued to record activity for three minutes following the stimulus. Figure 2c shows that control cultures had a healthy level of stochastic spontaneous activity prior to the stimulation, and afterward there was an overall reversion to previous activity levels. In contrast, cultures grown in the presence of 4Hfolate (Figure 2d) had an increase (\( \sim 2.5 \) fold, Figure 2e) in the overall level of activity (\( n = 17, p < 0.05 \)) following stimulation. Since network activity patterns in dissociated cultures can be variable, samples were taken from multiple regions of each chip, from five separate culture rounds, and experiments were performed over the course of eight months to control for the potential of seasonal variation, none of which was observed.

**Analysis of network activity in 5M4Hfolate reared cultures.** As stated above, one of the metabolites of 4Hfolate is 5M4Hfolate, a
form commonly associated with regulation of cellular functions, and it is linked to a variety of disorders, including epilepsy. While the breakdown of 4Hfolate to L-glutamate could be a major component of the observed effects, we also tested 5M4Hfolate in the in vitro assay system. The presence of 5M4Hfolate during culture development had an intriguing effect. Figure 3a,b shows a comparison between a control and a 5M4Hfolate reared culture. While a less dramatic overall enhancement of network activity persisted after stimulation in comparison to 4Hfolate, a distinctive pattern of cell firing was observed. A subpopulation of neurons was seen to fire rapidly for short periods of time (Figure 3b). To quantify this effect we calculated the time intervals between firing events and this data is displayed as a histogram in Figure 3c,d. While the control cultures generated a fairly stochastic distribution of inter-event timings, the 5M4Hfolate reared cultures showed a radically different pattern, with the repeated burst frequencies giving a distinctive population of inter-event times. This pattern was seen in many cultures even in the absence of stimulation (n = 6/8, Figure 3e), and was quantified in terms of their order (entropy) in comparison to controls (Figure 3f). This result suggests that there is a decrease in entropy, or an increase in order, in the firing events in the 5M4Hfolate reared cultures.

Together, the 4Hfolate and the 5M4Hfolate data demonstrate that there are quantitative differences in response to a high-frequency burst of activity between control and folate reared cultures. Control cultures appear to have the ability to maintain their level and pattern of firing rates following a stimulation event, whereas neuronal networks formed in the presence of folate tend to lose this firing rate pattern or the ability to regulate their firing level. This finding can be compared to the in vivo data, which shows the ~2 fold increase in susceptibility to seizure, presented graphically in Figure 2f. Furthermore, there is a qualitative difference between

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**Figure 2** | In vitro and in vivo effect of folic acid on neuronal activity (A–D). Dissociated co-cultures of neurons and glia were grown on silicon wafers, stained with Fluo-4 and the activity recorded. Firing events in each of up to 1000 neurons (cell index number on vertical axis) are recorded, detected with software, and represented by a tick mark. (A) Direct application of extracellular buffer had no substantial effect on the network activity of 3 week old cultures. (B) Similar to the electrophysiological recordings in Figure 1, optical recording from 1000 neurons show the entire network responding to the acute addition of 4Hfolate. (C) and (D) show the effect of rearing cultures in the presence of low levels (50 nM) of 4Hfolate in comparison to controls. Stochastic activity during a 2 minute control period is recorded, followed by a 30 sec stimulation of all neurons at 60 Hz (red stripe), and then 3 minutes post-stimulation. (C) Control cultures reverted rapidly to pre-stimulation levels of activity. (D) Culture grown in the presence of 4Hfolate for 3 weeks prior to testing, showed a continued increase in activity following stimulation. (E) Statistics of the ratio of overall firing rate of the culture after/ before stimulation. Statistics are from n = 14 recordings for each condition, taken from 5 separate culture rounds over the course of 4 months. (F) Results from the in vivo study shown for comparison, showing the increased sensitivity (a decrease in the latency to seizure) of pups from dams supplemented with 4.0 mg of folic acid in 1 ml of sucrose daily (control: 813 sec, folic acid: 469 sec). All values are significantly different at p < 0.05 (t-test).

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neuronal networks reared with each folic acid metabolite, suggesting that each may have a different effect on the connectivity and consequently the nature of the excitability.

Folate reared cultures have increased synaptic density. After identifying a difference in the network activity between control, 4Hfolate and 5M4Hfolate reared cultures, we subsequently did a structural comparison between the conditions to see if there were any overt changes in synaptic distribution. Cultures used for the activity movies were fixed and immunostained with an antibody against synaptophysin (Figure 4a–f). We found that there was a significant increase in synaptic density in both the 4Hfolate and the 5M4Hfolate cultures after 1 week of growth, however this quantitative difference diminished after 2 weeks (Figure 4g,h). Interestingly there was a qualitative change in the folate reared cultures at two weeks. While the control cultures appeared to have a relatively homogeneous distribution (Figure 4d), the folate reared cultures showed a distinct patterning in their synaptic distribution along neuronal processes (Figure 4e,f). While it is tempting to interpret this as an effect of the higher activity levels inducing synaptic pruning and organization, further studies are being pursued to more accurately quantify this phenomenon.

Rearing in the presence of 4Hfolate alters neuronal electrophysiology. The differences in basic structural connectivity at a gross level prompted us to investigate whether synaptic transmission was altered in 4Hfolate reared cultures. We recorded mEPSCs from both cultures (Figure 5a,b), and observed differences in frequency of the events (Figure 5b,c), however there was no significant change in amplitude (Figure 5d,e). This result correlates with the enhanced synaptic density seen in the folate reared cultures during the initial stages of network development. However there is also an increase in mEPSC frequency at 4 weeks, yet only qualitative differences in synaptic distribution at the later timepoint. This suggests other mechanisms might be at work modifying activity in more mature cultures.

Discussion
Folic acid supplementation during the first month of fetal development is critical to ensure the closure of the neural tube, however the necessity to maintain high folate levels after this event has not been substantiated. As an initial step toward investigating the underlying physiology of folic acid’s effect on brain development, we used both in vivo and in vitro rodent paradigms. In an in vivo latency-to-seizure...
Figure 4 | Analysis of synaptic distribution in control and folate reared cultures. 1 and 2 week old (A,D) control, (B,E) 4Hfolate reared and (C,F) 5M4Hfolate reared cultures were immunostained for synaptophysin. Synaptic counts were extracted from 8 regions from each culture, and 3 separate culture rounds were analyzed. Statistics for (G) 1 week and (H) 2 week counts, showing controls significantly lower at one week from both metabolites (n = 24, p < 0.05 (t-test)).

Figure 5 | Miniature EPSC analysis. (A) mEPSC recorded (2 uM TTX, −60 mV voltage clamp) from either 2 week or 4 week old hippocampal cultures from control and folate reared conditions were analyzed to determine (B, C) average frequency and (D, E) average amplitude. Statistical differences (*) were only seen for frequency (n = 6 recordings for each condition, between 50–500 mEPSCs per recording, p < 0.05).
We found that intake of folic acid by pregnant rats resulted in offspring which had seizures twice as fast as controls when challenged by a condition which destabilized their neural activity. While this assay provides a good first indication of the predisposition of the animals to seizures, more in depth analysis of direct brain activity recordings, and their performance in more sensitive assays such as kindling will help define the differences in the offspring exposed to folic acid in utero. It should also be noted that the dose given to the pregnant rats (4 mg/day) is difficult to compare to and substantially higher than that typically taken by to humans (0.4–1 mg/day, up to 5 mg/day in at-risk populations). We do not hypothesize that high levels of folic acid alone are sufficient to result in offspring with a seizure disorder, but rather that it could contribute to this possibility when combined with other destabilizing factors or mutations. Consequently, higher doses were used in the in vitro assay in order to observe any resulting hyperactivity or instability in a normal rat brain background. Future studies could utilize seizure-prone rats, which could more closely duplicate our hypothesized paradigm.

To further investigate this phenomenon we established an in vitro neuronal culture assay system and showed that cultures grown in the presence of 4Hfolate have a reduced capacity to re-regulate network activity patterns following a perturbation, a result similar in principle to the in vivo study. A reduced capacity to stabilize following a high-frequency burst could be problematic, as for the brain to perform normally it must be able to function in multiple states, including bursts of high frequency activity. It must also be able to reestablish a stable level of activity following such bursts.

Our choice of using 4Hfolate was primarily based on the fact that in an in vitro system there is no liver to convert folic acid to its biologically active forms and 4Hfolate is the highest-level metabolite past this point. Cultures were frequently replenished with the compound since its stability in solution is short, and, importantly, it is possible that the catabolites of 4Hfolate are in fact causal to the system instability. For example it is quite plausible that the presence of higher levels of L-glutamate could be stimulating a more rapid rate of development of neuronal connectivity. However since we were uncertain of how much of the 4Hfolate gets converted to 5M4Hfolate (a very biologically relevant form) by the cells, we were uncertain of how much of the 4Hfolate gets converted to 5M4Hfolate (a very biologically relevant form) by the cells, we repeated the in vitro activity experiments with direct exposure to low levels of 5M4Hfolate. This produced an interesting result: a subset of cells became very hyperexcitable, firing at high frequency in bursts. This result demonstrates the diversity of effects that could be produced by folic acid supplementation, as it could occur through the developmental of neuronal connectivity, which then leads to a hyper-excitability network. However the full spectrum of the effects of folic acid and its specific metabolites continues to be investigated. As a DNA methylating agent, folic acid is a candidate for inducing epigenetic changes, an interesting factor to consider given the novel presentation of epilepsy in families with no genetic pre-disposition. However a full understanding of these mechanisms is crucial, as previously unrealized advantages of full term folic acid supplementation might exist. For instance, recent studies have suggested a link between continued folic acid supplementation and reduced incidences of autism16. To reiterate, the presence of folic acid during the first trimester is absolutely essential, and the goal of this study is not to make a clinical recommendation for maternal health. Rather we hope this study encourages further investigation of the necessity and consequences of continued folic acid supplementation in late gestation.

Methods

In vitro assay. Young sexually mature female Long-Evans rats were paired with age-matched males and given daily supplements of 4.0 mg/kg of folic acid and 5% sucrose solution, while controls were given sucrose alone. After parturition the oral folic acid was discontinued and the litters culled to 6 male pups. At postnatal day 14 their latency to first behavioral seizure (clonus) following an i.p. injection of 15 mg/kg kainic acid (KA) was determined. This latency to the first behavioral seizure after KA administration, referred to as the seizure onset time (SOT), was defined by the occurrence of forelimb clonus, rearing, and loss of balance. Seizure onset time is a commonly used measure to describe susceptibility to convulsant compounds in rats17 and is related to clear paroxysmal epileptiform discharges18. Pups were sacrificed humanely with a pentobarbital overdose at the end of a one hour observation period. One pup in the control group did not display a seizure and was removed from the subsequent analysis.

In vitro assay. To investigate the cellular- and network-level effects of long term folic acid exposure, we used an in vitro model system based on dissociated cultures of rat hippocampal neurons. Briefly, high density co-cultures of neurons and glia were prepared from P0 pups obtained from Sprague Dawley dams. Neuronal activity experiments with direct exposure to 4Hfolate were repeated in this assay system. To quantify this, we first determined spontaneous neuronal network activity during a two-minute baseline period. Briefly, neuronal activity was recorded from P0 pups from normal Sprague Dawley rats as previously described19. Cultures were grown on silicon wafers to facilitate the firing of sub-populations of neurons at a precise frequency using photoconductive stimulation. Briefly, the technique works by selectively illuminating the region of the wafer to be stimulated, which increases the conductivity of the underlining silicon due to the photoconductive effect. Then a short (2 ms) pulse is applied across the surface of the wafer which initiates action potentials in the neurons within the illuminated region20.

Experimental cultures received 4Hfolate or 5M4Hfolate supplements daily in their growth media at a concentration of 50 nM. Supplements were prepared by dissolving the appropriate amount of the compound in media (BME) to make an 1 μM stock solution. This stock solution became diluted to 50 nM upon addition to the active culture media. The concentration chosen was estimated based on measures of actual folate blood levels21. Using this paradigm, we tested if the presence of folic acid metabolites during development causes a change in the activity patterns of the neuronal networks that develop. To quantify this, we first determined spontaneous network activity during a two-minute baseline period. Briefly, neuronal activity was recorded by Fluo-4 calcium imaging under low magnification over a large (~100 μm2) region of the neuronal culture with a high sensitivity camera. Optical data was recorded at 33 ms intervals, individual cells identified, their intensity traced over time and action potentials identified using custom software written in Matlab.

Immunocytochemistry and synaptic density analysis. Neuronal cultures were fixed in 4% paraformaldehyde and 15% picric acid prepared in PBS, and then subjected to immunofluorescent labeling with an mouse monoclonal anti-synaptophysin antibody (Sigma-Aldrich, St. Louis). Images were taken with a 20× water immersion objective and processed with a Richardson-Lucy algorithm using the DeconvolutionLab plugin (Biomedical Imaging Group, EPFL) for Image (NIH). The deconvoluted images were analyzed with the Graylevel Watershed (Biomedical Imaging Group, EPFL) plugin for ImageJ, which compartmentalized and counted the individual synapses in the image.

Electrophysiology. Whole-cell patch clamp recordings were performed using an Axon Axon Multiclamp 700 b amplifier, 1440A Digidata digitizer and pClamp software (Axon Instruments, Foster City, CA). Recordings were made at 50 kHz, and subsequently filtered at 5 kHz. Extracellular bath solution (EBS) for the recordings contained 135 mM NaCl, 10 mM glucose, 3 mM CaCl2, 5 mM KCl, 2 mM MgCl2, and 5 mM Hepes, pH adjusted to 7.3 with NaOH and adjusted to 310 mosmol with Sorbitol. Patch pipettes were pulled from borosilicate GD-1 glass capillaries (Narishige Scientific, Japan) and filled with intracellular solution containing 100 mM K-glutamate, 1.7 mM KCl, 0.6 mM EGTA, 5 mM MgCl2, 10 mM Hepes, 4 mM ATP and 0.1 mM MgATP.
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Author contributions

M.C. authored the main manuscript text and prepared the figures; F.G. and J.D. designed and performed the in vitro activity pattern and complexity analysis, L.S. performed the long term in vitro experiments, Y.A. performed the mEPSC experiments, J.H. initiated and performed a preliminary study, S.I. performed the synaptic density experiments, C.D.M. investigated the acute folate electrophysiology, R.T. assisted with protein chemistry, L.B.E. performed the long term synaptic density new measurements. The authors declare no competing financial interests.

Additional information

Competing financial interests: The authors declare no competing financial interests.

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