Maternal Vitamin C Deficiency during Pregnancy Persistently Impairs Hippocampal Neurogenesis in Offspring of Guinea Pigs

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Abstract

While having the highest vitamin C (VitC) concentrations in the body, specific functions of VitC in the brain have only recently been acknowledged. We have shown that postnatal VitC deficiency in guinea pigs causes impairment of hippocampal memory function and leads to 30% less neurons. This study investigates how prenatal VitC deficiency affects postnatal hippocampal development and if any such effect can be reversed by postnatal VitC repletion. Eighty pregnant Dunkin Hartley guinea pig dams were randomized into weight stratified groups receiving High (900 mg) or Low (100 mg) VitC per kg diet. Newborn pups (n = 157) were randomized into a total of four postnatal feeding regimens: High/High (Control); High/Low (Depleted), Low/Low (Deficient); and Low/High (Repleted). Proliferation and migration of newborn cells in the dentate gyrus was assessed by BrdU labeling and hippocampal volumes were determined by stereology. Prenatal VitC deficiency resulted in a significant reduction in postnatal hippocampal volume (P < 0.001) which was not reversed by postnatal repletion. There was no difference in postnatal cellular proliferation and survival rates in the hippocampus between dietary groups, however, migration of newborn cells into the granular layer of the hippocampus dentate gyrus was significantly reduced in prenatally deficient animals (P < 0.01). We conclude that a prenatal VitC deficiency in guinea pigs leads to persistent impairment of postnatal hippocampal development which is not alleviated by postnatal repletion. Our findings place attention on a yet unrecognized consequence of marginal VitC deficiency during pregnancy.

Introduction

Vitamin C (VitC) deficiency can have detrimental effects in the brain of juvenile guinea pigs. We have previously shown that a postnatal VitC deficiency in guinea pigs results in an impaired spatial memory function and a significant reduction in the number of neurons in the cornu ammonis (CA)1, the CA2–3 and the dentate gyrus (DG) of the hippocampus, unveiling a histomorphometric link between VitC deficiency and a negative impact on brain function [1]. During development, the brain is particularly vulnerable to low levels of antioxidants because of a high growth rate resulting in increased cellular metabolism and an immature antioxidant network leading to redox imbalance [2]. Thus, levels of VitC are increased in several tissues, including the brain during early life [3], and a reduced level has been shown to increase markers of oxidative stress [2,4].

The brain undergoes multiple steps of cellular proliferation, differentiation and maturation during development. In vitro data have shown that VitC plays a key role in the regulation of neuronal differentiation, maturation and neurite formation [5–7], implying that a negative impact of VitC deficiency on brain development could be expected. This has also been supported by in vivo findings establishing that VitC transport to the brain is essential for perinatal survival [8,9]. Within the brain, the hippocampus has one of the highest VitC concentrations [10–12] and a corresponding high level of the VitC transporter—sodium dependent vitamin C co-transporter 2 (SVCT2) [13,14].

Like humans, guinea pigs are unable to synthesize VitC due to a mutation in the gene encoding for L-gulonolactone oxidase, making guinea pigs a highly valuable in vivo model for VitC deficiency. The guinea pig is a precocial species with the peak of overall cellular growth in the brain (‘the brain growth spurt’) occurring before birth [15,16]. Based on our previously shown VitC deficiency induced reduction in hippocampal neurons and impairment of Morris Water Maze (MWM) performance in young guinea pigs, we therefore hypothesized that these effects may also occur when deficiency is imposed during gestation, a condition that can be observed in large subpopulations of pregnant women.

The aim of the current study was to investigate the effect of prenatal VitC deficiency on postnatal hippocampal development.
and function in guinea pigs and whether any such effect could be reversed by VitC repletion immediately after birth. Hippocampal development was assessed by determining the rate of proliferation of newborn cells in the DG at postnatal day (P)10, their survival at P27, and calculating hippocampal volumes on P10, P27 and P70. We found that prenatally deficiency negatively affects the postnatal hippocampal volume and cellular migration within the hippocampus. These findings were persistent despite re-introduction of high VitC levels immediately after birth.

Materials and Methods

Ethics
The study was approved by the Danish Animal Experimentation Inspectorate under the Ministry of Justice ( Permit number: 2007/561-1298). Surgery prior to euthanasia was performed under Isoflurane anesthesia and all efforts were made to minimize suffering.

In Vivo Study
Eighty Dunkin Hartley guinea pigs (Charles Rivers Lab, Kieslegg, Germany) at gestational day 18, e.g. within the first trimester (gestation length is 58–78 days [17]), were equipped with a subcutaneous (s.c.) microchip for identification, and randomized into weight stratified dietary groups receiving High (900 mg, n = 30) or Low (100 mg, n = 50) levels of VitC per kg diet (quality controlled diets; Special Diets Services, SDS, Witham, England). Upon arrival, maternal VitC levels in plasma were obtained by blood sampling (39.8±11.3 μM). The dose of 100 mg VitC/kg feed has been shown to result in a non-scorbutic VitC deficiency in guinea pigs even during long-term studies [18,19]. After birth, newborn pups were randomized again to either High (750 mg) or Low (100 mg) VitC per kg diet, giving a total of four different postnatal dietary feeding regimens: High/High (control, CTRL); High/Low (postnatal depletion, DEPL; only male pups), Low/Low (pre/postnatal deficiency, DEF); and Low/High (postnatal repletion, REPL). See Figure 1 for graphical overview of study design.

Pups were weighed and micro-chipped (s.c. in the neck) within 24 hrs after birth. Of the total of 285 live born pups, 168 were included in the current study. There was no statistically significant difference in the number of live born pups between dietary groups or effects of dietary VitC or gender on birth weight of pups. Four dams proved not to have conceived and were subsequently euthanized; one dam was euthanized during gestation.

Male pups (n = 91) were randomly allocated to experimental groups stratified by birth weights. Within the first five days of life, four pups born from High and four pups from Low mothers died, leaving n = 83. Pups were weaned five days after birth, post-natal day (P5, except from DEPL, which were weaned at P2) and group-housed in approved guinea pig housing racks. Pups received BrDU intraperitoneally (100 mg/kg Sigma-Aldrich, St Louis, MO, USA) once daily from P5–P9, as described previously [20]. Pups were euthanized either on P10 or P27. During the study, six animals died or were euthanized. Autopsies revealed gingivitis and dental malformations in two pups, whereas no specific macroscopic findings were recorded for the remaining four pups.

Female pups (n = 77) were randomly allocated to the three dietary groups CTRL, DEF and REPL (n = 22 for each group) and weaned around P7. Twelve animals were allocated from each dietary group (stratified according to birth weights and age) to be tested in a cued MWM. Additional age-matched pups were included in the CTRL and DEF groups (n = 5 and n = 6 respectively) to be tested in a cued MWM. Pups were euthanized between P56–P79 referred to in the text as P70.

Unless stated otherwise, animals were housed in floor pens, allowed feed, dried hay and water ad libitum, and tended several times daily by trained staff. Body weights were recorded at least twice a week. At arrival and two times during gestation, 300 μL blood samples were taken from the v. saphena at its superficial course on tibia in eight random dams of each group for verification of VitC status (data not shown). Likewise, blood samples were taken from female pups around P35 and P63 (data not shown).

Euthanasia
Animals were anesthetized by inhalation of Isoflurane (Isoba Vet 100%, Intervet International, Boxmeer, The Netherlands). Upon the disappearance of voluntary reflexes, thoracotomy was performed and an intracardial blood sample was collected by

Figure 1. Overview of In Vivo Study. Eighty pregnant guinea pigs were weight-stratified into two groups at gestational day 18. Group High (n = 30) was assigned to a diet containing 900 mg of vitamin C per kg diet and group Low (n = 50) received a diet containing 100 mg of vitamin C per kg. Otherwise, diets were identical. Newborn pups were assigned to either of four dietary groups (CTRL, DEF, DEPL or REPL) by weight stratification. Weaned male pups (n = 83) received a daily BrdU injection (100 mg/kg body weight) between post-natal day (P)5 and P9 and were sacrificed either at P10 or P27. Female animals (n = 77) did not receive BrdU. A subset (n = 36) were submitted to two consecutive tests in the non-cued Morris water maze (MWM) initiated at P24 and P52. A cued test (c-MWM) was performed on age-matched pups from both CTRL and DEF groups (n = 5 and n = 6, respectively) at P35. All female pups were sacrificed at approximately two months of age (P70). Biochemical and histochemical analyses were performed as indicated on the scheme.

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5 mL syringe and 18G hypodermic needle previously flushed with 15% tripotassium-EDTA [21]. Euthanasia was achieved by exsanguination and subsequent decapitation. Blood samples were immediately centrifuged and stabilized after sampling. Tissue samples were removed and placed in ice-cold phosphate buffered saline (PBS) prior to freezing on dry ice and subsequent ~80°C. The brain was excised, weighed and divided into left and right hemispheres (by section through the cerebral longitudinal fissure). The hemispheres were randomly assigned to either fixation (4% paraformaldehyde (PFA) in PBS, 0.15 M, pH 7.4 to be transferred to 1% PFA in PBS within 48 hrs) or storage at −80°C. The hemispheres were scanned with a Polaroid print scan 35plus and the hippocampal area was determined in a blinded fashion with ImageJ [Wayne Rasband, NIH). The hippocampal volume for each animal was then calculated by the Cavalieri principle, according to the formula [20]: \(V = \sum \text{areas} \times \text{distance between sections.}\)

To assess proliferation and migration of newborn cells in the hippocampus, immunofluorescence staining was performed as described previously [31] on four out of eight coronal sections covering the depth of the DG, making each region 800 μm apart. Deparaffinized and rehydrated sections were boiled for 15 min in 10 mM sodium citrate buffer, pH 6.0 for optimized antigen retrieval of BrdU, nuclear DNA and cell markers [32]. Sections were co-stained with polyclonal sheep anti-BrdU (Novus MB 500-235) and mouse monoclonal anti-NeuN (Chemicon MAB377) or mouse monoclonal anti-GFAP (Chemicon MAB360), and counterstained with 4′,6-diamidino-2-phenylindole (DAPI) to visualize cell nuclei. The BrdU-positive cells in each part of the DG (hilus, subgranular zone [SGZ] and granular layer [GL]) were counted in a blinded fashion on stitched images taken at 20X magnification. Images were acquired on a wide-field epifluorescence microscope (AxioImager M1) using AxioVision Digital Imaging software (Carl Zeiss MicroImaging GmbH, Jena, Germany).

**Data Analysis**

Statistical analysis of variance was performed on the collected data for animal body weight, brain weight, VitC concentration in brain and plasma, MWM performance, hippocampal volume, and number of BrdU positive cells in the different regions of the DG (Statistica v 7, Statsoft, Tulsa, OH, USA). Two-way ANOVA was applied to the stereological, immunohistochemical and biochemical data. In case of significance, Tukey’s post hoc test was applied. For analysis of MWM data, repeated measures ANOVA was applied to the cued test and acquisition phase of the non-cued MWM. Retention trial of the non-cued MWM was analyzed by one-way ANOVA. Comparison of swim-patterns was done with Fishers exact test. A P-value <0.05 was considered to be statistically significant.

**Results**

**Effect on Body Weight and Brain Weight**

During the study, no animals developed clinical signs of scurvy in accordance with our expectations and previous validation of a diet of 100 mg VitC/kg feed to be non-scorbutic [1,18,19,33]. Postnatally depleted (DEPL) pups exhibited a significant lower body weight than controls (CTRL) (P<0.001), whereas the remaining groups were not significantly different from CTRL (Table 1). VitC deficiency did not have an effect on brain weight, except specifically for postnatally repleted (REPL) animals at P10. As expected, brain and body weight increased with age (p<0.001; Table 1).

**Biochemical Analysis**

Plasma VitC concentration reflected the applied dietary regimes (Table 2), the deficient guinea pigs (DEF and DEPL) being consistently significantly lower in plasma VitC than CTRL counterparts (p<0.001). Plasma Vitamin C concentrations in REPL were also significantly lower than CTRLs (P<0.01 by ANOVA), particularly manifested at P27 (P<0.001). A similar pattern was observed for total ASC concentrations in the brain (DEF and DEPL vs CTRL, P<0.001), except that brain ASC was not lower in REPL, demonstrating the preferential repletion of the brain. Plasma ASC concentrations increased with age (p<0.05), while brain ASC concentration (p<0.001) and DHA declined significantly with age (p<0.01). At P70, MWM animals displayed significantly lower levels of ASC in plasma, but not brain,
Table 1. Brain weight and histological data from the different time-points.

|                  | P10 (male) | P27 (male) | P70 (female) | Two-way ANOVA |
|------------------|------------|------------|--------------|---------------|
|                  | CTRL (n=10)| DEF (n=10) | DEPL (n=11)  | REPL (n=10)   |
| Body weight (g)  | 115.3±14.6 | 124.2±14.4 | 105.4±23.5   | 103.4±19.5    |
|                 | 236.9±38.3 | 244.9±30.2 | 159.3±48.7** | 223.4±49.0    |
|                 | 438.6±68.9 | 406.8±40.7 | 482.1±96.1   |               |
|                | CTRL (n=10) | DEF (n=10) | DEPL (n=7)   | REPL (n=10)   |
| Brain weight (g) | 55.7±3.7   | 51.6±2.3   | 56.5±5.1     | 47.2±5.8**    |
|                 | 59.2±6.2   | 54.1±5.2   | 62.3±7.1     | 53.5±3.7      |
| Hippocampal      | 55.7±40    | 54.4±6.7   | 55.7±40      |               |
| volume (mm³)     | (n=9)      | (n=9)      | (n=9)        |               |

Data is presented as mean±SD and were analyzed by two-way ANOVA using postnatal age and dietary regimen as factors followed by Tukey’s test for posthoc comparisons in case of statistical significance. *p<0.05; **p<0.01; ***p<0.001 compared to CTRL animals. ‘ns’: not significant. Within each time point, one-way ANOVA was used.

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Effect on the Hippocampus

Animals subjected to prenatal VitC deficiency (DEF and REPL) showed an increased ASC oxidation ratio compared to non-mazed animals (P<0.001). ASC oxidation ratio was reduced in DEPL animals at the two time points investigated. Hippocampal volumes were about 10 to 15% lower in the prenatally-deficient animals (DEF and REPL) compared to the other groups (P<0.001; Table 1). Hippocampal volumes were adjusted for total brain weight to identify that the observed effect of prenatal deficiency was specific to the hippocampus and not due to a general retardation of brain growth. Despite this adjustment, hippocampal volume for DEPL animals stayed significantly below the corresponding counterpart (P<0.001) by two-way ANOVA. Hippocampal volume increased from P27 to P70, although no significant increase was observed with age (P>0.05), suggesting that the decrease observed with age was not due to a general retardation of brain growth, but rather due to a prenatal deficiency specific to the hippocampus. To investigate alterations in proliferation and/or survival of newborn cells in the DG, animals were labeled with 5-bromodeoxyuridine (BrdU) between P5 and P9, and sacrificed at either P10 or P27. The DG perimeter was defined as described previously [30]. To investigate altered proliferation and/or survival of newborn cells in the DG, animals were labeled with 5-bromodeoxyuridine (BrdU) between P5 and P9, and sacrificed at either P10 or P27. Sections were stained for BrdU and NeuN (marker for mature neurons) or GAP (marker for astrocytes) and the number of BrdU-positive cells in each sub-region of the DG, i.e., hilus, subgranular zone (SGZ) and granular layer (GL), was calculated. In agreement with previous findings [30], no significant differences in the mean number of BrdU-positive cells in the different regions were observed between the control and the DG of DEF animals, suggesting that the observed effect of prenatal deficiency within the hippocampus was a specific effect on newborn cell proliferation and/or survival. To investigate whether there was a difference in the number of BrdU-positive cells in the different regions within the hippocampus of DEF animals, two-way ANOVA was performed to compare prenatally deficient (DEF and REPL) animals with their corresponding counterparts (P<0.001). Although no significant difference in the number of BrdU-positive cells in the different regions within the hippocampus of DEF animals was observed with age (P>0.05), suggesting that the decrease observed with age was not due to a general retardation of brain growth, but rather due to a prenatal deficiency specific to the hippocampus.
Performance in the Morris Water Maze Test

**Cued morris water maze test.** No statistically significant differences were found between the two groups in any of the analyzed parameters (time to platform localization, total swim distance and cumulative distance to platform).

**Non-cued morris water maze test.** Analysis on quantitative data obtained from the non-cued MWM revealed no significant difference between the three dietary groups at either P24 or P52 tests during the five-day acquisition phase; animals were at the same initial level and equally improved their performance during this learning phase. On the P24 and P52 retention trials (time in platform quadrant, time to first hit on platform area, average distance to platform position and total number of crossings of platform area) we did not find any statistically significant differences between groups, besides that at the P52 test REPL animals displayed a decrease in average swim velocity (cm/sec) significantly different to DEF animals ($P < 0.05$, one-way ANOVA) in the retention trial. There was a significant effect of time between P24 and P52 tests, seen as an overall increase in total swim distance ($P < 0.001$), swim velocity ($P < 0.001$) and frequency of platform crossings ($P < 0.05$), and a reduction in the mean distance to platform area ($P < 0.05$). This corresponds well with the general observation that the animals became more able swimmers with age. However, subsequent reviews of swim patterns [1,22] did not disclose a difference between groups at either tests ($P > 0.05$), the majority in all groups displaying a random swimming pattern.

**Discussion**

The major findings of the present study were that prenatal VitC deficiency affects hippocampal development and leads to a significant and persistent reduction in postnatal hippocampal volume, irrespective of a VitC repletion right after birth. The hippocampal atrophy observed in prenatally VitC-deficient animals may be the consequence of the reduced migration of newborn progenitor cells from the SGZ into the DG.

Our data demonstrates that in spite of selective placental transport of VitC from mother to fetus [36–38], the prenatal brain development of the offspring is indeed sensitive to maternal VitC deficiency. Moreover, our findings indicate that VitC deficiency during second and third trimester may trigger adaptive mechanisms, e.g. by up-regulating expression of the vitamin C transporter SVCT2 protein or other protective mechanisms, to increase postnatal VitC uptake and retention in the brain ensuring a rapid repletion once an adequate VitC supply is provided.

The differences in hippocampal volume between prenatally deficient and non-deficient counterparts were maintained even when corrected for brain weight, indicating that the observed effects are not related to retarded brain growth. Moreover, prenatal VitC deficiency leads to a specific reduction in postnatal hippocampal volume which could be found throughout the investigated dietary regimen. The reduction of hippocampal volume was not due to a change in the rate of proliferation or survival of newly generated cells. In humans, 10–15% reductions in hippocampal volume as those found here are observed during early signs of dementia or long-term ecstasy or cannabis abuse [39–41].

While the postnatal production and survival of neurons was not affected by VitC deficiency, the migration and therefore maturation of newborn cells was negatively affected by prenatal deficiency. We found that the number of BrdU positive cells present in the GL of the DG was significantly lower in prenatally deficient animals (DEF and REPL). Although, ASC is known to affect proliferation and maturation of neuronal precursor cells in vitro [42], this is the first report that describes a negative impact of VitC deficiency on progenitor cell development in vivo. Although not conclusively demonstrated by our data, it is likely that the impaired migration of newborn cells (despite similar production and survival) is responsible for the decreased hippocampal volume in prenatally deficient compared to control animals.

The brain maintains one of the highest concentrations of VitC in the body and is able to favor this organ even during prolonged states of deficiency [2,18,34]. One of the earliest clinical signs of pathological VitC deficiency in guinea pig pups is weight loss [2]. The body weight of DEPL animals was significantly decreased compared to that of CTRL (Table 1). This is surprising since one could imagine that being chronically deficient would be more deleterious than acute postnatal deficiency. One reason could be that these animals were weaned earlier than counterparts (P2 versus P5) resulting in an overall decreased weight gain. However, it may also be that DEPL animals are more sensitive to a sudden decrease in VitC compared to animals who have been accustomed to low levels prenatally. Reviewing brain VitC concentrations at P10, it is evident that repletion occurs highly efficiently, REPL animals being no different than the CTRL group throughout the study. Thus, an alternative explanation could be that that animals subjected to chronically low fetal supply of vitC (DEF animals) have adapted to this environment e.g. by up-regulating SVCT1 & 2 and perhaps ASC recycling to spare as much vitC as possible [14]. The SVCT2 protein is responsible for the majority of the VitC uptake into the brain [43] and its expression, at least in some organs including the cerebellum, has been shown to be increased by VitC depletion [44,45]. Such a protective mechanism may allow REPL and DEF animals to have a very efficient uptake, compared to DEPL counterparts. In contrast, pups exposed to acute deficiency (DEPL animals) do not have these preventive measures readily available and are therefore more susceptible to a sudden decrease in vitC supply, e.g. requiring time to adjust transport and retention mechanisms. An increased sensitivity to deficiency in guinea pigs going from a high in VitC diet to a very low has previously been shown [46,47]. Albeit this did not translate into a reduction of the hippocampal volume in the present study, we believe that an effect of depletion would be visible at two months of age. This is supported by our previous study where a postnatal depletion from P7 resulted in 30% reduction of hippocampal neuronal number compared to control counterparts [1].

Several studies have reported that a reduced hippocampal volume may be linked to impaired cognitive capacity [48,49] and other neurologic disorders [50,51]. We have reported that postnatal VitC depletion for two months is associated with a diminished spatial memory ability in the non-cued MWM and coherent reductions of neurons in the hippocampus DG, CA1 and CA2-3 fields [1]. To investigate a functional effect of prenatal VitC deficiency, the present animals were also submitted to the MWM test. Cued MWM in CTRL and DEF groups displayed equal performances, demonstrating that VitC deficiency did not induce alterations in basic motor or sensory skills [25,27]. This was supported by the non-cued MWM acquisition trial, confirming that animals were equally competent prior to the retention tests. However, in contrast to previous findings, we found no difference in memory performances between dietary groups in the non-cued MWM on either P24 or P52; the majority of the animals being random swimmers. Regardless of the reason, limited information can be retrieved from non-cued tests when almost none of the animals display a spatially persistent swim pattern, as this suggests that the potential window of effect was bypassed in the present.
Table 2. Vitamin C status in guinea pigs at the different postnatal time points.

| Effect of MWM | P10 (male) | P27 (male) | P70 (female) | Two-way ANOVA | P70 (female, MWM) | Effect of MWM |
|---------------|------------|------------|--------------|---------------|-------------------|---------------|
|               | CTRL (n = 10) | DEF (n = 10) | DEPL (n = 11) | REPL (n = 10) | CTRL (n = 10) | DEF (n = 9) | DEPL (n = 7) | REPL (n = 10) | CTRL (n = 10) | DEF (n = 10) | REPL (n = 10) |
| Plasma total ascorbate (µM) | 31.3 ± 9.7 | 3.3 ± 2.9*** | 5.7 ± 6.3*** | 24.6 ± 13.0 | 54.8 ± 20.1 | 3.5 ± 1.4*** | 26.1 ± 1.5*** | 24.1 ± 11.8*** | 51.8 ± 22.4 | 4.0 ± 3.2*** | 57.3 ± 13.6 |
| Brain total ascorbate (µmol/g tissue) | 1.67 ± 0.10 | 0.76 ± 0.20*** | 1.42 ± 0.13** | 1.80 ± 0.14 | 1.58 ± 0.13 | 0.63 ± 0.20*** | 0.83 ± 0.38*** | 1.54 ± 0.09 | 1.34 ± 0.15 | 0.45 ± 0.25*** | 1.45 ± 0.04 |
| Brain ascorbate oxidation ratio (% DHA of total VitC) | 3.7 ± 2.5 | 20.0 ± 10.1*** | 8.4 ± 6.7 | 5.8 ± 2.9 | 4.3 ± 9.4 | 6.7 ± 3.3 | 4.5 ± 6.7 | 3.8 ± 3.5 | 4.5 ± 2.8 | 9.5 ± 46** | 3.1 ± 2.2 |

Data is presented as mean ± SD and were analyzed by two-way ANOVA using postnatal age and dietary regimen as factors followed by Tukey’s test for posthoc comparisons in case of statistical significance.

*p < 0.05; **p < 0.01; ***p < 0.001 compared to CTRL animals. ns: not significant. Within each time point, one-way ANOVA was used. Effect of the MWM trial on biochemical markers were tested by two-way ANOVA using MWM and dietary regimen as factors.

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study. Moreover, we speculate that the difference in hippocampal volume observed here may just not have been large enough to detect an effect by MWM, had we been able to measure it.

Interestingly, we found an overall effect of MWM on ASC homeostasis. Plasma concentrations of ASC in animals subjected to water maze test were significantly lower than those of their non-mazed counterparts \( P < 0.05 \). This is likely to be a transient effect due to a reduced feed intake during testing. However, while having comparable ASC levels, the brain displayed a several-fold higher ASC oxidation ratio in mazed compared to non-mazed animals \( P < 0.001 \), suggesting that the MWM procedure may impose a degree of stress on the animals leading to enhanced oxidative stress in the brain. This finding is potentially interesting and should be investigated further as increased oxidative stress—related to the MWM—may impact the interpretation of cognitive performance tests in general.

In conclusion, we show that prenatal VitC deficiency has a persistent negative effect on postnatal hippocampal development as measured by volume reduction over the two month postnatal period monitored in this study. This is most likely due to impaired
migration of newborn cells into the GL, which may be the consequence of a block in maturation. A marginal deficiency in maternal VitC status leads to a persistent deviation of the hippocampus in offspring supports a pivotal role of VitC during brain development, and emphasizes the requirement for an adequate supply during pregnancy. Marginal vitamin C deficiency is far from uncommon in humans, and may affect large subpopulations including mothers-to-be and young children [52]. This study provides substantial in vivo experimental data on a yet unrecognized consequence of marginal, non-scorbutic VitC deficiency that may prove to have implications for nutritional recommendations to pregnant women.

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Author Contributions
Conceived and designed the experiments: PTN JGS SC JL. Performed the experiments: PTN LV JGS NJ SH MDP SC JL. Analyzed the data: JL SC PTN LV SH. Wrote the paper: PTN JL SC.

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