Prenatal Exposure to the Viral Mimetic Poly I:C Alters Fetal Brain Cytokine Expression and Postnatal Behaviour

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Evidence that, in a rodent model that more closely resembles human brain development, prenatal infection can lead to behavioural abnormalities in postnatal life.

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Introduction

While the aetiology of mental illnesses such as schizophrenia and autism remains unknown, many epidemiological studies have identified a potential neurodevelopmental origin for these disorders [1–6]. These studies have shown an association between infection during pregnancy and increased risk of the development of mental illness disorders in later life. Given that prenatal infection of varying origin and type each appear able to elicit similar postnatal effects [3–9], it is thought that the specific pathogen is not important, but it is the common inflammatory response that is a key mechanism in provoking an alteration in the trajectory of brain development. As infection and inflammation invariably involve the release of inflammatory cytokines, this is known as the cytokine hypothesis [10–13].

Animal studies have sought to investigate the link between exposure to a prenatal infection and the develop-
administration of poly I:C in the brain of fetal rats and mice after maternal administration of many schizophrenia and autistic patients and 2 cytokines that are constitutively altered in the serum of interleukin (IL)-6 and tumour necrosis factor (TNF)-α, −κB) transcription factors, resulting in the subsequent production of nuclear factor kappa-light-chain-enhancer of activated B cells’ (NF-κB) transcription factors, resulting in the subsequent production of inflammatory cytokines [34–36] − in particular of interleukin (IL)-6 and tumour necrosis factor (TNF)-α, two cytokines that are constitutively altered in the serum of many schizophrenia and autistic patients [37–40], and in the brain of fetal rats and mice after maternal administration of poly I:C [18, 41–44].

We sought to determine whether poly I:C administered to the precocial spiny mouse at mid pregnancy, at a stage when myelination of the brain [26] and adrenocortical differentiation is just beginning [32], had an impact on cytokine expression in the fetal brain, and whether prenatal exposure to this viral mimetic had effects on postnatal behaviour. In particular, we examined the effects of prenatal poly I:C administration on prepulse inhibition, which, in conventional rodents, has been used as a signature of a behavioural disorder closely associated with schizophrenia in humans [21–23, 25]. Furthermore, the current study aimed to extend our previous findings [45] by using a higher dose of poly I:C (5 mg/kg), which has been consistently shown to cause deficits in prepulse inhibition and other behavioural abnormalities in rats and mice [17, 19, 25, 46–48].

Animals and Methods

Animals

This study used spiny mice (Acomys cahirinus) obtained from the breeding colony maintained at Monash Medical Centre. The spiny mice were bred and housed as previously described [49]. Experiments were conducted in accordance with the Australian Code of Practice for the Care and Use of Animals for Scientific Purposes. All procedures had received prior approval from the Monash University Animal Ethics Committee.

Prenatal Treatment

Pregnant dams at day 20 of gestation (term is at 39 days) received a single subcutaneous injection of either phosphate-buffered saline (PBS; control) or 5 mg/kg of poly I:C (polyinosinic-polycytidylic acid potassium salt; Sigma Aldrich, Castle Hill, N.S.W., Australia) in a volume of 5 μl/g body weight. Poly I:C was dissolved in PBS solution to yield the required concentration. Developmental events at 20 days of gestation (mid gestation) in the brains of spiny mice compares with that of the human fetus at early-to-mid gestation [26]. All animals were then returned to their home cages immediately after the injection procedures. The animals either were culled for tissue collection or 24 h after injection when maternal, fetal and placental weights were obtained or the pregnancies were allowed to continue until the dams gave birth naturally, and the behaviour of the offspring was assessed from 20 to 35 days of age. For all behavioural tests a minimum of 6 male and 6 female offspring from each treatment group were assessed.

Behaviour Tests

Assessment of behaviour of the offspring occurred at 20 and 35 days of age, which is prior to weaning at about 40 days of age. As the offspring were assessed by all behavioural tests, they were ordered from least stressful to most stressful. Based on the documented observations from our colony [50] and by others [51], spiny mice of this postnatal age can be considered juvenile and prepubertal.

Open Field Test. Offspring at 20 days of age were tested in an open field (50 × 50 × 40 cm; L × W × H) environment to assess (1) exploratory activity by measuring the distance travelled, and (2) anxiety-like behaviours by determining the amounts of time spent in the central zone versus the outer zone of the field. The central zone was defined arbitrarily by the area of the open field excluding a 10-cm outer perimeter. The open field test was conducted between the hours of 11:00 and 13:00 h, and each pup under test, with their siblings and mother, was habituated to the room in which the open field apparatus was kept by placing the home cage in the room for at least 1 h prior to testing. The lighting level was set at 2.8 lx for all trials. The offspring were placed in the centre of the field at the beginning of each trial, which lasted 10 min. Limelight (Neuroscience Inc., Tokyo, Japan) acquisition software was used to track the movement of the animals throughout the open field trial by identifying the nose, body and tail. Images were captured...
the animal spent all the time exploring the novel object, 0 indicates ing both objects. Thus, a discrimination index of 1 indicates that exploring the novel object, divided by the total time spent explor-
the time spent exploring the familiar object from the time spent
to determine the discrimination index, calculated by subtracting exploring each object for the first 5 min in the recall trial was used
total time spent exploring objects in the
ploration of the block. The total time spent exploring objects in the
ject). The remaining plastic block (which had been explored previ-
period), and during this time 1 of the 2 plastic blocks was replaced
following the completion of the open field test, 2 small plastic
blocks of identical shape and size were placed in the open field, positioned 6 cm from the walls of the arena to ensure an unob-
surred view of the animal at all times. The animal was then placed
back in the open field, and its behaviour recorded for a 10-min period, which served as the learning trial in which the animal ac-
quired information regarding the objects in the open field. Follow-
this, the animal was returned to its home cage for 1 h (retention period), and during this time 1 of the 2 plastic blocks was replaced
with a plastic block of a different shape and colour (the novel object). The remaining plastic block (which had been explored previ-
ously in the learning trial) and the novel object were both wiped down with 70% ethanol to remove olfactory cues. The animal was placed back in the open field, and its behaviour and exploration of the 2 objects was recorded for the next 10 min (termed the ‘recall trial’). In the analysis of the behaviour during the learning and re-
call trials, the nose rather than the body of the animal was tracked using the Limelight software. When the animal’s nose was pointed at and within 2 cm of the block, this was deemed to be active expl-
oration of the block. The total time spent exploring objects in the 10 min of the learning trial was measured. The total time spent exploring each object for the first 5 min in the recall trial was used to determine the discrimination index, calculated by subtracting the time spent exploring the familiar object from the time spent exploring the novel object, divided by the total time spent explor-
ing both objects. Thus, a discrimination index of 1 indicates that the animal spent all the time exploring the novel object, 0 indicates no preference for either object, and −1 represents all time spent exploring the familiar object.

**Novel Object Recognition Test.** Immediately after this open field trial, the novel object recognition test was conducted to assess non-
spatial memory, which is based on the innate exploratory behav-
bour of rodents. The above open field trial was used as the habitu-
ation trial for the novel object recognition test to reduce the con-
tribution of the novelty of the open field to the test. Immediately following the completion of the open field test, 2 small plastic
blocks of identical shape and size were placed in the open field, positioned 6 cm from the walls of the arena to ensure an unob-
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**Elevated Plus Maze.** Animals at 30 days of age were placed on the elevated plus maze apparatus to assess anxiety and fear behav-
bour in a novel and challenging environment. The elevated plus maze consisted of an open top rectangular box of 60 × 30 × 45 cm (L × W × H) divided into 3 chambers of 20 × 30 cm (L × W), with an 8 × 8 cm square opening cut into the internal walls to allow the animal to move between chambers. The chambers were numbered 1–3 from left to right for reference. Chambers 1 and 3 contained large cylinders measuring 40 × 8 cm (H × D) drilled with numer-
ous 0.6-cm (D) holes, in 1 of which the ‘stranger’ spiny mouse was
placed during the test. The stranger spiny mouse was age and sex matched to the test spiny mouse. The social interaction tests con-
sisted of the test spiny mouse being habituated to the social inter-
action apparatus for a 5-min period, after which the spiny mouse was taken out and the stranger spiny mouse placed in the cylinder enclosure in chamber 1 or 3. The placement of the stranger spiny mouse in the enclosure was alternated between chambers 1 and 3 for animals in each treatment group to account for preference to a particular chamber of the apparatus. The test spiny mouse was placed back in the middle of chamber 2 and a video recording was taken as the spiny mouse explored the apparatus. The stranger spiny mouse was available to the test spiny mouse for visual, tactile and olfactory contact through the enclosure. Postacquisition anal-
ysis of the time spent in the stranger, centre or empty chambers was performed and the time spent interacting with the stranger or empty enclosure by the subject spiny mouse was measured using the Limelight software. The chamber and enclosure definitions are shown in figure 1. When the animal’s nose was pointed at and within 2 cm of the stranger spiny mouse’s enclosure, this was deemed to be interaction with the stranger spiny mouse.

**Stranger Object Recognition Test.** The social interaction test was used to assess whether the poly I:C treatment had caused changes in the social behaviours of the offspring at 25 days of age. The apparatus consisted of an open top rectangular box of 60 × 30 × 45 cm (L × W × H) divided into 3 chambers of 20 × 30 cm (L × W), with an 8 × 8 cm square opening cut into the internal walls to allow the animal to move between chambers. The chambers were numbered 1–3 from left to right for reference. Chambers 1 and 3 contained large cylinders measuring 40 × 8 cm (H × D) drilled with numer-
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**Elevated Plus Maze.** Animals at 30 days of age were placed on the elevated plus maze apparatus to assess anxiety and fear behav-
bour in a novel and challenging environment. The elevated plus maze test was conducted between 11:00 and 13:00 h. The elevated plus maze consisted of 2 opposing open arms (40 × 10 cm) and 2 opposing closed arms (also 40 × 10 cm) which are surrounded by 15-cm-high walls. The 4 arms were connected in the shape of a
cross to a central platform (10 × 10 cm). The entire apparatus was elevated on a stand, hidden to the animal’s view, 50 cm above the floor. The test began when the spiny mouse was placed on the central platform of the maze facing an open arm and allowed to move freely for 5 min, during which time its movements were recorded by a video camera placed above the apparatus. Limelight acquisition software was used to track the movement of the animal and assess the distance travelled and time spent in each arm through- out the trial.

Prepulse Inhibition Test. Prepulse inhibition (PPI) of acoustic startle was measured using the SR-LAB startle apparatus (San Diego Instruments, San Diego, Calif., USA) at 35 days of age. The sound-proofed startle chamber contained a clear Plexiglas cylinder resting on a piezoelectric transducer that detected movement of the animal. A computer connected to the apparatus recorded the startle responses and controlled the timing and presentation of the acoustic stimuli. Testing took place between 10:30 and 13:00 h. Each test began with 2 min of apparatus acclimatisation, followed by 5 consecutive pulse-alone trials presented in order to habituate the animal and establish a stable startle response. The animals were then presented with a total of 40 pseudorandom trials consisting of 10 pulse-alone trials, 5 no-stimulus trials (i.e., background noise), and 5 prepulse + startle pulse trials where the amplitude of the prepulse was 72, 74, 78, 82 or 86 dB sound pressure level (SPL), corresponding to 2, 4, 8 and 16 dB SPL above the background noise. The session was concluded with 5 consecutive startle pulse-alone trials. The interval between successive trials ranged from 10 to 30 s, with a mean of 20 s. The startle pulse stimuli was a 40-ms pulse of white noise at 115 dB SPL, and the prepulses were 20-ms bursts of white noise delivered 100 ms before the startle pulse. The average PPI was expressed as the average of percent inhibition of the startle response due to each prepulse, i.e. %PPI = amplitude of startle pulse – (amplitude of each prepulse + startle pulse)/(amplitude of startle pulse) × 100%. A higher %PPI score indicates a greater reduction in startle magnitude in prepulse + pulse trials relative to that in pulse-alone trials. The average startle was expressed as the average amplitude of the startle pulse-alone trials.

Quantitative Real-Time PCR

Samples (placenta, fetal) were obtained at 2 h (n = 5 dams) and 24 h (n = 5 dams) after poly I:C administration to assess cytokine gene expression. Total RNA was extracted and DNase treated using the commercially available RNeasy Kits (Qiagen, Malvern East, Vic., Australia). SYBR green PCR master mix, as per the manufacturer’s instructions (Applied Biosystems), was used to determine relative levels of mRNA expression for NF-κB, TNF-α and IL-6. The primer sequences are listed in Table 1. The PCR assay was optimised by determining the specificity of the primer sets, which were found to amplify products of the expected sizes. Quantitative (q)PCR cycling conditions for all genes consisted of an initial denaturation step of 95°C for 10 min, followed by 40 cycles consisting of 95°C for 15 s and 60°C for 1 min. A dissociation curve was also performed at the end of each run. Samples were run in triplicate for each gene of interest. Data obtained from qPCR were analysed as previously described [52]. The relative expression within the sample (ΔCt) was measured from the qPCR data by subtracting the mean cycle threshold (Ct) for the housekeeping gene (18S) from the mean Ct value for the gene of interest. This value was then inserted into the formula 2-ΔCt to give a final arbitrary expression value, which was then divided by the mean 2-ΔCt of the PBS-2-hour treatment group to give a final value against which the relative expression for each gene of interest was determined for the other time (24 h) and the poly I:C group at 2 and 24 h after treatment.

Statistics

All data are presented as means ± SEM. Maternal, fetal and placenta data and pregnancy outcomes were analysed using an independent t test. Open field test results, novel object recognition test results, average startle and average change of PPI (expressed as

Table 1. Primer sequences for genomic analysis

| Housekeeping gene RN18S1 (RNA, 18S ribosomal 1) | ACACGGACAGGATGACAGA | CAAATCGCTCCACCACTAA |
| NF-κB | TGAGGGATCTGCTGGAAGTC | CCAAGTGCAAGGTTCTGA |
| IL-6 | CAGACCCATCAGGCAAGACA | TGCCTGATCTTCCATCCTC |
| TNF-α | CAAATCGGATGACAAAGCCTG | GAGATCCATCCGGTGGC |

Table 2. Effect of prenatal PBS and poly I:C administration on sickness behaviours and pregnancy outcomes

| Maternal, fetal and placental data | PBS | Poly I:C | p value |
| Maternal body weight | change, g | 0.06±0.70 | -1.78±0.26 | 0.04a |
| Fetal body weight, g | 0.49±0.05 | 0.29±0.03 | 0.01a |
| Placental weight, g | 0.23±0.02 | 0.14±0.01 | 0.004a |
| Placental:fetal body weight ratio | 0.50±0.07 | 0.51±0.07 | 0.87 |

| Pregnancy outcomes | Gestational period, days | 39.00±0.21 | 38.88±0.23 | 0.70 |
| Litter size, n | 2.75±0.48 | 3.25±0.48 | 0.49 |

a Significant difference between poly I:C and PBS treatment groups; b significantly different from pretreatment weight.
percent) were analysed using a two-way analysis of variance (treat-
ment × sex). Gene expression results were analysed using a two-
way analysis of variance (treatment × time). A two-way repeated-
measures analysis of variance was used to analyse postnatal growth 
(treatment × sex × age), social interaction test results (treatment × 
sex × chamber/enclosure) and %PPI for each prepulse intensity 
(treatment × sex × intensity). p < 0.05 was accepted as statistically 
significant unless otherwise stated; however, a p value of <0.10 was 
considered noteworthy.

Results

Maternal, Fetal and Placenta Weight, Pregnancy 
Outcomes and Postnatal Growth

Maternal body weight fell significantly after the poly 
I:C treatment and 24 h after the treatment was signifi-
cantly lower than the maternal weight of the PBS-treated 
dams (t 8 = 2.44, p = 0.04; table 2). There was also a sig-
nificant reduction in fetal body weight (t 13 = 3.116, p = 
0.01) and placental weight (t 13 = 3.520, p = 0.004) at 24 h 
after poly I:C administration. When placental weight was 
taken into account, there was no significant difference be-
tween treatment groups in placental-to-fetal weight ratio 
(t 14 = 1.451, p = 0.17; table 2). There was no effect of treat-
ment on litter size or the length of the duration of gesta-
tion (table 2). Body weight increased in all offspring be-
tween 1 day and 40 days of age (main time effect F 2, 48 = 
1,292, p < 0.0001; table 3), with female offspring being 
near-significantly lighter than their male littermates 
(main sex effect F 1, 24 = 3.98, p = 0.06). There was also a 
tendency for poly I:C animals to be lighter throughout 
postnatal life compared with PBS controls, although this 
did not quite reach significance (main treatment effect 
F 1, 24 = 4.01, p = 0.06).

Gene Expression

The time after treatment, or administration of poly I:C, 
had no effect on NF-κB, IL-6 and TNF-α mRNA expression 
in the placenta (fig. 2a, c, e). Similarly, there was no effect 
on IL-6 mRNA expression in the fetal brain. However, 
there was a significant decrease in NF-κB (F 1, 15 = 6.03, p = 
0.03; fig. 2b) and TNF-α (F 1, 16 = 6.33, p = 0.02; fig. 2f) ex-
pression in the fetal brain of poly I:C-treated animals com-
pared with the PBS controls; a similar decrease occurred at 
both 2 and 24 h after administration of poly I:C.

Open Field Test

The total distance travelled by offspring was not differ-
ent between the treatment groups, but there was a strong 
tendency for female offspring to travel less in the open 
field trial than males (main sex effect F 1, 23 = 3.70, p = 0.06; 
fig. 3a). There was no effect of prenatal treatment or sex 
on the distance travelled by offspring in the central zone of 
the open field (fig. 3b, c).

Novel Object Recognition Test

There was no difference between the treatment groups 
in the time spent exploring objects in the initial learning 
trial (fig. 4a). However, in the second trial, offspring from 
poly I:C-treated mothers spent less time exploring the 
new object than pups from PBS-treated mothers, as 
shown by the significantly lower discrimination index 
(F 1, 23 = 9.829, p < 0.01; fig. 4b). The results were similar 
for male and female offspring.

Social Interaction Test

All offspring recognised the presence of the ‘stranger’, 
as shown in figure 5b, by the significantly greater amount 
of time spent exploring the stranger spiny mouse’s cham-
ber compared with the centre or empty chamber (main effect of 
chamber F 2, 46 = 43.795, p < 0.001). However, unlike all other 
treatment groups, male offspring from the 
poly I:C mothers did not interact significantly more with 
the ‘stranger’ enclosure than with the empty enclosure, so 
that a significant ‘stranger × treatment × sex’ interaction term was identified (F 1, 23 = 5.184, p < 0.05; fig. 5a).

<table>
<thead>
<tr>
<th></th>
<th>PBS</th>
<th>Poly I:C</th>
<th>Main effects</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>male</td>
<td>female</td>
<td></td>
</tr>
<tr>
<td>At 1 day</td>
<td>5.82±0.25</td>
<td>5.6±0.1</td>
<td></td>
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<tr>
<td>At 20 days</td>
<td>18.3±1.37</td>
<td>17.16±1.3</td>
<td></td>
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<tr>
<td>At 40 days</td>
<td>32.05±1.37</td>
<td>28.4±0.6</td>
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</tbody>
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Table 3. Effect of prenatal PBS and poly I:C administration on body weight (g) of offspring at 1, 20 and 40 days of age
Elevated Plus Maze

Prenatal treatment had no effect on the distance travelled or time spent in the open or closed arms of the elevated plus maze (data not shown).

Prepulse Inhibition

Average startle was lower in female mice than in their male littermates (main sex effect $F_{1, 21} = 6.275, p < 0.05$), and the difference between the sexes was similar for the PBS- and poly I:C-treated offspring (fig. 6a). Prenatal treatment with poly I:C significantly reduced %PPI for each prepulse intensity, and also for the average relative decrease in PPI in these offspring, compared with PBS. * Main treatment effect of $p < 0.05$. The data are shown as means ± SEM in all graphs.

![Graphs showing relative expression of NF-κB, IL-6, and TNF-α in the placenta and fetal brain](image)

**Fig. 2.** Relative expression of NF-κB (a, b), IL-6 (c, d) and TNF-α (e, f) in the placenta (a, c, e) and fetal brain (b, d, f) 2 h and 24 h after PBS and poly I:C treatment. There was a significant reduction in NF-κB (b) and TNF-α (f) expression in the fetal brains at 2 and 24 h after poly I:C treatment compared with PBS. * Main treatment effect of $p < 0.05$. The data are shown as means ± SEM in all graphs.
Fig. 3. Total distance travelled (a) and the distance travelled (b) and time spent in the central zone (c) of the open field by male (white columns) and female (dashed columns) offspring prenatally exposed to PBS or poly I:C. There was no main effect of treatment on distance travelled overall and the distance travelled in the central zone of the open field. The data are shown as means ± SEM in all graphs.

Fig. 4. Time spent exploring objects in the learning period (a) and discrimination index measured in the recall trial (b) of the novel object recognition test by male (white columns) and female (dashed columns) offspring prenatally exposed to PBS or poly I:C. Poly I:C animals had a significantly lower discrimination index than PBS animals (p < 0.01), irrespective of sex. ** Main treatment effect of p < 0.01 compared with PBS. The data are shown as means ± SEM in all graphs.
Discussion

This study shows that a virus-like infection during pregnancy, mimicked by using the TLR3 agonist poly I:C, is associated with an acute decrease in cytokine expression in the fetal brain. Cytokines are important inflammatory mediators induced by infection [53, 54]. In addition to their important role in the innate immune response, cytokines participate in a wide range of important functions including glial and neuronal development [54, 55], and in pregnancy the placenta is also an additional source of cytokine production [56, 57]. Specifically, in this study using the precocial spiny mouse, poly I:C administration at mid gestation acutely (at 2 h) resulted in a significant decrease in mRNA levels of the proinflammatory cytokine TNF-α in the fetal brain of the spiny mouse, and this change was also present 24 h after treatment. While a shift in the relative production of pro- and anti-inflammatory cytokines might be expected to affect brain development [54], this was an unexpected finding because other studies using altricial rodents have described increased proinflammatory cytokine production following maternal poly I:C treatment. Although it is not known what causes this downregulation, it is known that cytokines produced by the maternal immune system can activate the fetal HPA axis [58], resulting in the release of glucocorticoids which then inhibit the induction of proinflammatory cytokines while stimulating the production of anti-inflammatory cytokines [59, 60]. This negative feedback relationship could explain the downregulation of proinflammatory cytokines in the brain of the fetal spiny mouse shortly after poly I:C treatment.

As the change in cytokine expression was only found in the fetal brain but not the placenta, the question of how this is brought about remains unclear. Cytokines have been found to be present in rodent fetal brains from early gestation [12, 43], and in vitro studies demonstrate that cytokines and chemokines can be produced by human fetal microglia and astrocytes in response to viral infection [58, 61–64]. Furthermore, cytokines are thought to be critically important in glial cell development [54, 65]. This may explain the increase in astrocytes and activated microglia in the neonatal brains of spiny murine offspring exposed to a lower dose of poly I:C at mid gestation, as reported previously [45].

It was found that fetal bodies and placentas were lighter when collected from poly I:C-treated dams, although it is difficult to determine whether either treatment had an effect on the weight of these tissues without knowing the initial weights of the fetuses and placentas. We believe the fetuses were coincidentally smaller in the poly I:C treatment group and that this decrease in placental and fetal weight was not caused by treatment, which is also shown by the fact that the placental-to-fetal weight ratio was not different between the treatment groups.

After birth, prenatal poly I:C treatment resulted in significant changes in the behaviour of the offspring, including deficits in social interaction, sensorimotor gating, and

![Figure 5](image-url)

Fig. 5. Time spent interacting with the stranger mouse in the stranger enclosure compared with the empty enclosure (a) and time spent in each chamber (b) by male and female offspring prenatally exposed to PBS or poly I:C. All offspring spent significantly more time exploring the stranger enclosure than the empty enclosure, except the male poly I:C animals (a). There was no difference in the time spent in each chamber between the treatment groups or sexes. # p < 0.05 compared with stranger enclosure; * p < 0.001 compared with stranger chamber. The data are shown as means ± SEM in all graphs.
of memory and learning, when tested at 3–5 weeks of postnatal age, a stage of development that is equivalent to late childhood and prepubescence in humans [32]. Whereas previous studies have described behavioural abnormalities in adult animals after poly I:C treatment in pregnancy [18, 20, 22], the emergence of symptoms in childhood and early adolescence is an important finding, since some mental illnesses such as autism, ADHD and schizophrenia, thought to be provoked by maternal viral illness during pregnancy, emerge at this stage of life. While the dose of poly I:C used in this study was higher than that used in our previous study [45], it is similar to that used in other studies with conventional mice or rats which produce altricial offspring. This poly I:C dose (5 mg/kg) had no effect on the duration of pregnancy, litter size and neonatal survival; however, unlike that used in our previous study, this higher dose of poly I:C did cause an acute decrease in maternal body weight, an indicator of sickness behaviour used in previous studies [66]. As prenatal poly I:C administration caused a decrease in postnatal growth, although not significantly, it is important to consider whether this higher dose of poly I:C may have caused a global compromise and, if so, what effect this may have had on postnatal behaviour.

Our test of social interaction was based on social novelty and the natural curiosity of the animal to investigate and interact with a stranger spiny mouse. This test showed that male offspring born to mothers given poly I:C at mid gestation spent less time interacting with the stranger, as indicated by the reduction in sniffing at the stranger’s enclosure, a common method of social investigation for rodents [67]. The awareness of the presence of the stranger by the poly I:C-treated offspring appeared to be unaffected, as the control and poly I:C-treated animals spent a similar amount of time in the chamber containing the stranger spiny mouse. In addition, it is unlikely that the reduction in interaction has been affected by an increase in non-specific fear or anxiety, low curiosity or reduced exploratory activity, because the results from the elevated plus maze and the open field tests showed no difference in these behaviours between PBS and poly I:C animals. The social interaction test thus

![Graph 1: Average startle](image1)

![Graph 2: %PPI](image2)

![Graph 3: %PPI by sex](image3)

**Fig. 6.** Effects of prenatal exposure to PBS or poly I:C on average startle (a) as well as %PPI for each prepulse intensity and average %PPI in male (b) and female (c) offspring. There was no difference in average startle between the treatment groups (a). Poly I:C animals showed a significant decrease in %PPI for each prepulse intensity average %PPI (b, c; p < 0.01), irrespective of sex. * p < 0.05 compared with PBS animals. The data are shown as means ± SEM in all graphs.
shows a resistance of the juvenile, male poly I:C offspring to come into close association with an unknown, ‘stranger’ individual. This result is of particular relevance to schizophrenia and autism, with the age at onset of schizophrenia being earlier in males than in females [68], and with autistic males more severely affected than female patients [69–71].

Abnormal social interaction including reduced interest in peers, difficulty in maintaining social interaction, and unusual modes of social interaction have been used as a diagnostic measure for identifying autism in humans [72], although only few animal studies have investigated the impact of poly I:C given during pregnancy on social behaviours of the offspring. This study shows that a single viral mimetic treatment at mid gestation has an impact that results in altered social behaviour in male offspring at an age that corresponds to the approximate time of puberty in this precocial species. Previous studies in mice showing effects on social behaviours differ in that either the dose of poly I:C was very high (20 mg/kg) [73, 74] or repeated poly I:C treatments were given directly to mouse pups at 2–6 days of postnatal age [75]. Further studies on the social behaviour of these animals in the home cage environment, and in groups of other juvenile or adult spiny mice, could be important in establishing whether the results of this simple ‘stranger’ interaction test extend to more complex social interactions and settings.

PPI was used to assess sensorimotor gating, an index of the ability of the brain to filter and process incoming sensory information. A decrease in PPI is a common feature of neuropsychiatric disorders such as autism and schizophrenia [76–79], and changes in PPI have been used to screen potential antipsychotic drugs in human [80, 81] and animal studies [82, 83]. In the current study we found a significant decrease in PPI in both male and female juvenile offspring born to mothers given poly I:C. This complements previous results in rats, where a reduced PPI has been observed in juvenile and adult offspring after prenatal poly I:C exposure [84], although other studies in juvenile and adult mice found a decreased PPI only in the adults [85]. Importantly, the assumption that deficits in PPI do not emerge until adulthood has not been consistently confirmed, and human patients identified as being at risk of schizophrenia clearly show that sensorimotor gating impairments may be evident prior to the full onset of the condition [86, 87]. Variation in developmental trajectories of the multiple brain regions and of the neurochemical substrates involved in regulating auditory PPI may explain the differences in the emergence of PPI deficits between species. Differences in PPI deficits and other behaviours between species may also be the result of the timing of prenatal poly I:C administration, as the stage of fetal development may be important for prompting changes in brain development that persist after birth.

Memory impairment is a well-documented symptom in mental illnesses such as schizophrenia and autism [88–90], and a significant association between impaired cognition and both schizophrenia and autism have been identified in human patients [57, 91]. Our novel object recognition test conducted on the spiny mouse found that the poly I:C treatment during pregnancy impaired the capacity of 20-day-old male and female offspring to detect a novel object, compared with offspring from control pregnancies. A previous study in mice showed that prenatal treatment with poly I:C affected novel object recognition in adult offspring [75], and, again to our knowledge, this has not previously been tested or shown to occur at more juvenile ages.

In addition to non-spatial recognition memory, human studies have found impairments of working memory and reversal learning in schizophrenic [88, 92, 93] and autistic [89, 94] patients. It would be of interest to assess working memory and reversal learning in a study like the present one to provide further validation of our model.

We have shown a significant behavioural impact arising in the juvenile offspring of spiny mice when the mother was given a dose of poly I:C that caused a downregulation of proinflammatory cytokines in the fetal brain. A future direction would be to determine whether behavioural abnormalities in offspring persist, intensify or attenuate in adulthood and old age. Nonetheless, we believe our findings of behavioural abnormalities in young animals are important as groups at risks of psychosis present with behavioural deficits prior to the onset of a disease [86, 87]. Therefore, this period prior to puberty may provide a unique period of time for the development of interventions or treatments.

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References


68 Loranger AW: Sex difference in age at onset of schizophrenia. Arch Gen Psychiatry 1984;41:157–161.


84 Wolff AR, Bilkey DK: The maternal immune activation (MIA) model of schizophrenia produces pre-pulse inhibition (PPI) deficits in both juvenile and adult rats but these effects are not associated with maternal weight loss. Behav Brain Res 2010;213:323–327.


