Differential effects of early or late exposure to prenatal maternal immune activation on mouse embryonic neurodevelopment

Elisa Gumaa,b,1, Maude Bordeleau,c, Fernando González Ibáñez,c,d,e, Katherine Picardc,d,e, Emily Snookf, Gabriel Desrosiers-Grégoiref,g, Shoshana Springh, Jason P. Lerchf,i, Brian J. Niemant,j, Gabriel A. Devenylm,n, Marie-Eve Tremblayf,j,k,l, and M. Mallar Chakravartyb,m,p,q

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Exposure to maternal immune activation (MIA) in utero is a risk factor for neurodevelopmental and psychiatric disorders. MIA-induced deficits in adolescent and adult offspring have been well characterized; however, less is known about the effects of MIA exposure on embryo development. To address this gap, we performed high-resolution ex vivo MRI to investigate the effects of early (gestational day [GD]9) and late (GD17) MIA exposure on embryo (GD18) brain structure. We identify striking neuroanatomical changes in the embryo brain, particularly in the late-exposed offspring. We further examined the putative neuroanatomical underpinnings of MIA timing in the hippocampus using electron microscopy and identified differential effects due to MIA timing. An increase in apoptotic cell density was observed in the GD9-exposed offspring, while an increase in the density of neurons and glia with ultrastructural features reflective of increased neuroinflammation and oxidative stress was observed in GD17-exposed offspring, particularly in females. Overall, our findings integrate imaging techniques across different scales to identify differential impact of MIA timing on the earliest stages of neurodevelopment.

maternal immune activation | embryo brain development | structural MRI | electron microscopy | brain development

1. Introduction

Brain development is a remarkable and complex set of processes under the organizational control of genetic, environmental, and immune regulation. The tightly regulated nature and interdependence of these processes make them vulnerable to a variety of risk factors. Converging lines of evidence suggest an association between prenatal exposure to maternal infection and increased risk for a host of neurodevelopmental disorders (5). MIA leads to an increase in maternal proinflammatory cytokines and chemokines, which are thought to interfere with fetal brain development by disturbing its delicate ecosystem, potentially as a consequence of the microglial response (5–9). Exposure to MIA during the sensitive window of in utero brain development may alter neurodevelopmental trajectories, thereby increasing risk for neurodevelopmental disorders later in life (10, 11). Identifying these sensitive windows and their impact on later development is critical to our understanding of the effects of MIA exposure. Previous work from our group has demonstrated that the gestational timing of MIA exposure has a differential impact on offspring brain and behavioral development, with a greater variation observed following early MIA exposure (gestational day [GD]9) relative to exposure later in gestation (GD17) or to saline exposure (11). These differences may be attributable to variation in maternal immune responsiveness and fetal brain development at different gestational timings and to MIA-induced neurodevelopmental variation throughout development (12).

Although there is significant evidence that MIA exposure in utero alters brain development trajectories in both human (13) and animal models (14, 15), it is unclear how soon after MIA exposure these changes can be detected. MIA-induced outcomes have been better characterized in adolescent and adult rodent offspring (12). However, to better understand the initiation and progression of MIA-induced variation in brain development, it is critical to study its impact at the earliest stages of life. Human neuroimaging studies have identified alterations in functional and structural connectivity in the infant brain development.

Significance

Prenatal exposure to maternal infection increases the risk of developing mental health disorders, such as schizophrenia and autism spectrum disorder. Exposure to maternal immune activation has been associated with a number of neuroanatomical deficits in adolescent and adult offspring, with differing effects based on the gestational timing of infection. However, little is known about how the embryo brain is affected. We show, using whole-brain MRI, that maternal immune activation significantly affects brain anatomy. When the exposure occurs early in pregnancy, volume reductions are mainly observed, while the opposite is true for exposure later in pregnancy. Furthermore, we identify alterations to the density of certain classes of neurons and glia, which have been associated with stress and inflammation in the brain.
brain following exposure to chronic, low-grade inflammation (as measured by interleukin-6 levels and/or C-reactive protein in the maternal plasma) (16–18). Even though some observations in early phases of life have been made, there is less information available regarding how MIA exposure impacts the morphogenesis of the fetus. Recent work on this topic suggests that MIA exposure induces acute up-regulation of genes involved in immune signaling, hypoxia, and angiogenesis in the fetal mouse brain (19). Further, alterations in neuronal proliferation, neuronal and glial specification, cortical lamination (19, 20), global messenger RNA translation, and altered nascent proteome synthesis have been reported (21).

These findings suggest that effects of MIA exposure may be detectable in the fetal and neonatal period across mouse and human studies. However, it is unclear whether the transcriptional and histological variation observed in rodents translates to the neuroanatomical changes detected by whole-brain imaging observed in human studies. Furthermore, although gestational timing has been shown to have differential effects in adolescent and adult offspring (11, 22), it is unclear how it affects neurodevelopment in its early phase. A better understanding of the neurodevelopmental sequelae of MIA exposure on very early brain development is of importance in the context of the current COVID-19 pandemic, as mothers who contracted the virus during pregnancy were more likely to have obstetric complications leading to poor fetal health outcomes that may impact downstream brain development, such as low birth weight, intrauterine growth restriction, and preterm birth (23, 24).

To build upon our previous investigations, in which we characterized brain development from adolescence to adulthood (11), we aimed to develop a chronology of how the timing of MIA exposure may impact brain development in utero, using the same gestational exposures as in our previous work. We leveraged structural MRI, an inherently three-dimensional (3D) imaging technique applicable for mouse phenotyping (25). This technique allows for a comparable assay across species, providing a potential avenue for establishing cross-species homology (26). We examine the effects of in utero exposure to early (GD9) or late (GD17) MIA with a viral mimetic, polyinosinic:polycytidylic acid (poly I:C), on embryo brain morphology at GD18 using high-resolution ex vivo whole-brain MRI. To better understand the cellular underpinnings of the volumetric changes identified by MRI, we utilized high-resolution electron microscopy (EM) to examine the density of certain cells, including apoptotic cells, dark neurons, and dark glial cells. These dark cells are unique from other neurons and glia, as they have distinct ultrastructural characteristics reflective of oxidative stress and are only identifiable with EM (27). We focused on dark and apoptotic cells, as they have been identified as a putative marker for neuroinflammation, cellular stress, apoptosis, and disease in the brain parenchyma (27–29). The dorsal hippocampus was selected as the region of interest, as it was differentially affected by GD9 and GD17 MIA timing. Further, alterations in this region have been consistently associated with neuropsychiatric disorders (30, 31), as well as in response to MIA exposure in our previous work (11) and that of others (32). Our results demonstrate neuroanatomical alterations in the GD18 embryo brain following MIA, with differential effects due to timing in many regions, including the dentate gyrus of the hippocampus. Here, we observed a significant increase in the density of apoptotic cells in the GD9-exposed embryos (but not GD17) relative to the control group, while an increase in the density of dark cells (both neurons and glia) was observed in the GD17-exposed offspring. These findings suggest that morphological changes due to MIA exposure are already detectable in the fetal brain and that the timing of MIA exposure may differentially impact the brain both at anatomical and cellular levels.

2. Results

2.1. Differential Effects due to MIA Timing on Embryo Brain Volume. To investigate the effects of early or late prenatal MIA exposure on embryo brain development, pregnant dams were exposed to either poly I:C (P1530-25MG poly I:C sodium salt TLR ligand tested; Sigma-Aldrich) (5 mg/kg, intraperitoneally [i.p.]) or saline solution (NaCl) at GD9 or GD17 (Section 4.1). Embryos from each of these four groups were harvested at GD18 and fixed in order to acquire ex vivo high-resolution structural MRIs at the same timepoint (Section 4.2). Once we confirmed that there were no statistically significant differences between our two control groups, they were combined, leaving us with three treatment groups: saline (SAL), GD9-poly I:C (POL E), and GD17-poly I:C (POL L). We used deformation morphometry to create a consensus average of the group, and the Jacobian determinants were analyzed by using a mixed-effects model to examine relative volumetric differences between groups.

Overall, we observed differing patterns of whole-brain morphology dependent on MIA timing; decreases in brain volume were more prevalent following GD9 exposure, while striking brain-volume increases were observed following GD17 exposure. We observed significant neuroanatomical differences due to MIA timing. The POL E embryos exhibited smaller volumes relative to SAL in the globus pallidus, hippocampus (including the dentate gyrus, as well as more posterior regions), fornix, centromedian thalamic nucleus, and cerebellum. Larger volume was observed in the prelimbic area, lateral septum, subventricular zone, caudate-putamen, sexually dimorphic nucleus of the hypothalamus, basolateral amygdala, and CA1 region of the hippocampus in the POL E group relative to SAL (t = 4.242, <1% false discovery rate [FDR]) (Fig. 1).

GD17 MIA exposure induced very striking volumetric alterations, particularly volumetric increases in the brain of POL L offspring at GD18 (t = 3.234, <1% FDR) relative to SAL offspring. Regions of volume increase included the ventral pallidum, septal plate and lateral septal nucleus, medial and lateral preoptic nuclei, caudate-putamen, globus pallidus, hippocampus (both dentate gyrus and CA1 regions), cingulum, anterior commissure, cortical plate, corpus callosum, external capsule, centromedian thalamic nucleus, and cerebellum. Decreases in volume were observed in the ventral hippocampus, more anterior subregions of the cortical plate, bilateral amygdala, fornix, and ventromedial thalamus (Fig. 1).

Interestingly, POL E and POL L MIA exposures were observed to have opposite effects on brain volume in some regions implicated in neurodevelopmental disorders and identified in previous MIA studies (11, 14), such as the dorsal hippocampus, wherein GD9 exposure decreased volume and GD17 exposure increased volume. Similar observations were made for the centromedian thalamic nucleus. The septal nucleus and caudate-putamen were increased in both MIA-exposed groups. A significant difference between POL E and POL L embryo brain anatomy was also observed (t = 3.590, <1% FDR). These results are fully described in SI Appendix, section 2.2.1 and Fig. 4. Post hoc investigation of sex differences revealed no significant sex-by-group interactions.
chose to use a nonparametric Kruskall–Wallis test (33), rather than using parametric statistics, we were interested in the density (measure per group, there were no group differences in total cell density). For aggregate density average density of dark glial, dark neuronal, and apoptotic cells across slices (i.e., average per embryo). For aggregate density measure per group, there were no group differences in total cell density ($\chi^2 = 0.680, df = 2, P = 0.712$; SI Appendix, Fig. 6). Although the density of dark glial and dark neuron cells appeared to be higher in the POL L offspring, there were no differences in dark glial cell density ($\chi^2 = 3.322, df = 2, P = 0.190$) or dark neuron cell density ($\chi^2 = 0.758, df = 2, P = 0.685$). A significant group effect was observed for apoptotic cell density ($\chi^2 = 6.349, df = 2, P = 0.042$), wherein the POL E offspring had significantly higher density than POL L offspring ($P = 0.053$), as well as the SAL offspring, although not significantly so ($P = 0.117$) (Fig. 2; see SI Appendix, Fig. 7 for more representative images of dark glia, dark neurons, and apoptotic cells).

### 2.2. EM of Embryo Dorsal Hippocampus

Given the differential effects of MIA timing on the dorsal hippocampus, and that this region has been shown to be affected by MIA exposure in our own work (11) and in various neuropsychiatric disorders (31), it was selected for EM investigation of cell density. Specifically, we focused on apoptotic cells, dark neurons, and dark glia, defined by ultrastructural properties only identifiable with EM and indicative of potential oxidative stress and neuroinflammation in the brain parenchyma (27–29); this may suggest a specific vulnerability of these cells during this early neurodevelopmental time frame.

In keeping with norms in the field (due to sample size and distribution) (33), rather than using parametric statistics, we chose to use a nonparametric Kruskall–Wallis test, followed by a pairwise Wilcoxon-rank test to assess group differences in the average density of dark glial, dark neuronal, and apoptotic cells across slices (i.e., average per embryo). For aggregate density measure per group, there were no group differences in total cell density ($\chi^2 = 0.680, df = 2, P = 0.712$; SI Appendix, Fig. 6). Although the density of dark glial and dark neuron cells appeared to be higher in the POL L offspring, there were no differences in dark glial cell density ($\chi^2 = 3.322, df = 2, P = 0.190$) or dark neuron cell density ($\chi^2 = 0.758, df = 2, P = 0.685$). A significant group effect was observed for apoptotic cell density ($\chi^2 = 6.349, df = 2, P = 0.042$), wherein the POL E offspring had significantly higher density than POL L offspring ($P = 0.053$), as well as the SAL offspring, although not significantly so ($P = 0.117$) (Fig. 2; see SI Appendix, Fig. 7 for more representative images of dark glia, dark neurons, and apoptotic cells).

Given that the variance in cell density differed quite drastically between groups, we sought to examine if the distributions in cell density differed between groups using the most granular data we had. Rather than using the pooled data per mouse (as above), we applied the shift function to the density measures acquired from each slice per mouse (3) to maximize variance. The shift function (34) quantifies how two distributions differ based on deciles of the distributions, i.e., it describes how one distribution should be transformed to match the other and estimates how and by how much one distribution must be shifted.

Comparison of cell-density distributions across groups confirmed that there were no overall differences in total cell density between groups (apart from a significant difference between POL E and SAL distributions only at the seventh decile of distribution [$P = 0.045$]; SI Appendix, Tables 2–4).

Interestingly, dark glial density was significantly lower for POL E offspring relative to SAL at higher deciles of the distribution (5th through 9th decile, $P < 0.020$; SI Appendix, Table 5; Fig. 3). Distribution for POL L offspring was not different from SAL; however, they also had significantly more dark glia than POL E at higher deciles of the distribution. Comparison of distributions for dark neurons revealed only subtle differences, with significantly higher density at higher deciles of the distribution for POL E relative to SAL (decile 9, $P = 0.016$; SI Appendix, Table 8) and POL L (deciles 8 and 9, $P < 0.010$; SI Appendix, Table 10), with no differences between POL L and SAL (SI Appendix, Table 9; Fig. 3). Finally, for apoptotic cell density, POL E offspring had significantly
higher density across lower deciles of distribution relative to SAL (deciles 1 to 4, \(P = 0.039\); SI Appendix, Table 11) and across all deciles relative to POL L (\(P < 0.020\); SI Appendix, Table 12). No differences between POL L and SAL were observed (SI Appendix, Table 13; Fig. 3).

Sex differences in distribution were also observed for all cell types (total, dark glia, dark neurons, and apoptotic cells). Of interest, increased density of dark glial cells was observed in the POL L females relative to SAL, while decreased density was observed for the POL E females, further described in SI Appendix, Table 13; Fig. 3.)
Appendix, section 2.3.2 and Fig. 8, with the summary of results per decile provided in SI Appendix, Tables 14–37.

Finally, using immunocytochemical transmission EM, we confirmed that the cells we identified to be apoptotic [those with a pyknotic nucleus (28, 35–37)] also had positive active Caspase 3 staining (Fig. 2), given the role of active Caspase 3 in morphological changes and DNA fragmentation leading to apoptosis (38). Furthermore, we sought to determine whether the dark glial cells identified were colocalized with markers of active microglia using CD11b immunostaining, previously shown to be present in adult contexts of pathology (27). However, despite multiple efforts, no positive staining was identified with transmission EM, potentially due to the difficulty in establishing an EM-compatible immunostaining protocol.

3. Discussion

There is a well-established link between MIA exposure in utero and latent neuroanatomical and behavioral abnormalities that emerge in adolescence or adulthood, with relevance to schizophrenia and ASD pathology (39–41). However, limited work has been conducted in the early neurodevelopmental period (12). We leveraged high-resolution ex vivo MRI and EM to characterize the effects of MIA exposure at two gestational timepoints on the embryo brain at GD18. Our results suggest that the embryo mouse brain undergoes significant remodeling in response to MIA, particularly due to late gestational exposure, coupled with changes in the presence of dark and apoptotic cells in the hippocampus. Elucidating the neurodevelopmental changes across the embryonic periods following MIA exposure is an important step toward our understanding of MIA exposure as a primer of downstream psychopathology and as a risk factor for an array of neuropsychiatric disorders.

Interestingly, we see volume reductions due to early MIA exposure in several brain regions, whereas we see striking volume expansions following late exposure. Since the late-exposed embryo brains were harvested 24 h after immune exposure, we are likely capturing an acute neuroinflammatory or stress response, or an acceleration of brain development in response to the immune stimulus. However, given that there was no difference based on SAL timing, one can assume that any acute effects are attributable to the MIA itself. Interestingly, there is homology between regions affected in the embryo brains and those in which we observed altered neurodevelopmental trajectories from childhood to adulthood in our published work (11). Some of these regions include the striatum/caudate-putamen, hippocampus, lateral septum, cingulate cortex, and cerebellum, many of which have been implicated in neuroimaging studies of humans with schizophrenia.
or ASD (30, 42, 43). Previous animal studies also report that MIA in late gestation increases neuroinflammation in the embryo rat brain and decreases placental function, as measured by T2-signal intensity (44). These findings suggest that increased neuroinflammation and decreased placental function could, in part, be driving some of the volumetric increases in the late exposed embryo brain, which may provide some mechanisms underlying disease pathology.

To gain more insight into the putative cellular underpinnings of the volumetric changes, we performed EM experiments in the dorsal hippocampus, a region highly implicated in neurodevelopmental pathology (31). In addition to the current findings, we previously observed interesting transcriptional changes to genes involved in early neurodevelopmental processes in the dorsal hippocampus of adolescent MIA-exposed mice—although following a different gestational exposure (11). A similar study conducted in nonhuman primates also found that the largest transcriptional changes due to MIA exposure were found in the offspring hippocampus, relative to the prefrontal and anterior cingulate cortex (45), in line with our observations in the mouse (11). Alterations to hippocampal morphology, both at the gross-anatomical and molecular level, and function have been implicated in numerous MIA studies (46, 47). Importantly, dysfunction of this region has also been proposed as a central player in the pathophysiology of schizophrenia, as well as other neuropsychiatric illnesses, wherein dysfunction begins during subclinical stages and worsens as symptoms progress (31). Of course, other brain regions, which may or may not be connected to the hippocampus, play a role as well (30, 42, 43).

Identifying the cellular processes triggered by MIA exposure is critical to our understanding of how this risk factor may alter offspring neurodevelopmental trajectories. By leveraging high-resolution EM techniques, we have an unprecedented opportunity to investigate the brain parenchyma at nanoscale resolution (28). This allows for identification of changes in different cell types and their unique features. We identified differential effects due to gestational MIA timing. In GD9-exposed offspring, where decreased dorsal hippocampal volume was detected, we observed an increase in apoptotic cell density (confirmed via Caspase 3 staining) and a decrease in dark glial cell density compared to SAL offspring. Conversely, in GD17-exposed offspring, which had enlarged dorsal hippocampal volumes, there was a tendency for greater dark-cell density (although not significantly), particularly for dark glial cells in the late-exposed females. These cellular results align well with the volume-based MRI results, wherein increased apoptosis could be linked to the decreased volume in GD9-exposed offspring, while increased dark-cell density could be linked to an acute inflammatory response and increased volume in the GD17-exposed offspring. Importantly, this may point to differences in neuropathological mechanisms in the fetal brain associated with MIA exposure and to some putative sex differences that require further investigation.

We focused our analyses on dark cells, both neurons and glia, as well as apoptotic cells, as these have been frequently detected in response to stress (29), aging (48), and neurodegeneration (49). They are thought to play a role in both normal and pathological synapse and neuronal network formation (49). Dark neurons are typically defined by a darker appearance, with an accumulation of mitochondria and nuclear indentations, associated with structural plasticity (49–51), as well as markers of cellular stress (dilated endoplasmic reticulum and Golgi apparatus) (28). Dark glial cells, particularly microglia, also display cellular markers of stress and have been shown to have hyperramified processes, which often lead to increased physiologically relevant contents, such as synaptic contacts and increased phagocytic capacity (27, 52). Further, reports of microglial reactivity and density in the brains of prenatally immune-challenged animals early in life are mixed, with observations of increased density and motility, as well as no differences (53). Although we were not able to identify colocalization between dark glial and CD11b staining, this does not rule out the possibility that these cells are, in fact, dark microglia, but it necessitates future, cell-specific research. By focusing specifically on dark microglial cells in the future, we may gain better insight into the phenotypic variability of these reactive cells and parse some of the heterogeneity in the literature.

At GD9, corresponding with our first MIA exposure, microglia colonize the brain, initiating their development toward maturity. Interestingly, sex differences in the number and morphology of microglia have been observed, with males showing greater numbers of these cells earlier in development (postnatal day [PND]14) than females, who have more microglia later in development (PND30 to PND60) (54). GD9 also occurs at a time at which the fetal brain is undergoing extensive neural proliferation and migration, which transitions more toward circuit refinement and cortical organization by GD17 (our late-MIA timepoint) (55). Importantly, microglia may play important roles in the regulation of apoptosis (56), as well as the permeability and formation of the blood–brain barrier, which typically takes shape between GD13.5 and GD15.5 in the mouse. Apoptosis is a critical cellular process in early brain and placental development in utero (57). The process of apoptosis has been detected as early as GD5, but increases significantly toward the end of gestation, peaking in early postnatal life before dropping off (58). In contrast with our findings, previous rodent studies using immunohistochemical techniques have actually observed an increase in apoptosis following lipopolysaccharide exposure in late pregnancy in rats (59) and poly I:C exposure at GD17, but not GD9, in mice (22). The discrepancy between these findings and ours may be due to differences in techniques or features used to identify these cells; immunohistochemistry may be less sensitive than EM to detection of ongoing apoptosis.

Exposure to poly I:C at the dose selected (5 mg/kg) typically elicits an acute immune response peaking 3 to 4 h postinjection, resolving, typically, within 24 to 48 h (60). We chose this dose and administration to model a more acute inflammatory exposure and to match our previous work (11, 61). Some of the differences in MIA-exposed offspring may be due to differences in developmental stage of the fetus (discussed in greater detail below) or to the maternal milieu. The maternal immune and endocrine systems undergo numerous changes during gestation. During early pregnancy (corresponding with GD9 exposure), the mother is in a more proinflammatory state due to implantation and placentation, the need to repair uterine epithelium and cycle between cell death and repair. In contrast, the later phase of pregnancy (other than immediately preceding labor and delivery) is characterized by a more anti-inflammatory state, wherein the mother, placenta, and fetus have achieved a symbiotic relationship (62, 63). These differences in maternal environment may influence the effects observed by MIA challenge and may, in part, explain some of the differences in offspring effects observed in our study.

The results presented here should be considered in light of their limitations. The design of our embryo study would be more complete with an assessment of neuroanatomy acutely following the GD9 exposure at GD10; this would allow us to detect whether volume increases, as those detected in the
GD17-exposed offspring, are a response to acute inflammation or specific to that gestational time point. Additionally, examining the brain of GD17-exposed offspring 9 d following MIA exposure, at PND8, could provide a similar delay after MIA before collection for imaging; however, the comparison of embryonic vs. postnatal brain development may have its own confounds. Unfortunately, the embryo is too small for MRI acquisition at GD9 or GD10; however, imaging could be performed by using other techniques, such as optical projection tomography (64). Regarding the EM analysis, we did not look at the extracellular space, which could contribute to the changes in volume detected in the MRI. In future work, cryofixation methods for preserving the EM samples would allow for more in-depth analyses of the extracellular space volume and composition (65).

The cytokine response elicited by our first and second batches of poly I:C differed, in that the second batch (to which a small subset of samples was exposed) did not elicit as strong of an immune response, particularly for the GD17 offspring. Although both batches of poly I:C were used within the 1-y mark for sample collection, the immunostimulatory potential of our second batch was tested when the product was older (2 y) than the first batch (1 y), leading to potential degradation of the material when immunostimulatory potential was tested in a satellite group of animals. An alternative approach to using satellite animals would be to measure cytokine response to poly

Fig. 4. Differences in distribution of dark glial, dark neuron, and apoptotic cell density per group. Distribution of dark glial cell density (A), dark neuron density (B), and apoptotic cell density (C) for all hippocampal slices per animal. The red line identifies the median of the data, while each black bar denotes a decile of distribution. A percentile bootstrapping technique applied to identify the difference in decile between the POL E and SAL for dark glial cell density (D), dark neuron density (E), and apoptotic cell density (F) showing decreased density of dark glia and increased density of apoptotic cells for POL E offspring. Next, POL L and SAL comparisons are shown for dark glia (G), dark neurons (H), and apoptotic cells (I), showing that POL L had higher density than POL E for dark glia and neurons, while POL L has lower apoptotic cell density. *P < 0.05.

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I:C challenge in the experimental dams; this would allow for a more nuanced investigation of the relationship between the strength of the immune response in the mother and the degree of alteration observed in the offspring. Recent work by Mueller et al. (66) has shown that there may be resilience of susceptibility to MIA, depending on the magnitude of maternal inflammatory response. We opted for the use of satellite animals, as we were concerned about introducing an additional intervention and stressor to the experimental dams. Important work from the human neuroimaging literature has shown that effects of maternal stress can be detected at the structural level in the human neonatal brain, both across structure and function. Thus, we wanted to ensure that we were isolating the effects of MIA without an additional confound of stress (67–71). Here, we provide evidence for the presence of neuroanatomical alterations in the embryo brain, differentially affected by timing; however, in future work, it would be of high interest to associate the relationship between maternal cytokine responsiveness and offspring outcomes as a means to parse some of the heterogeneity observed. In addition to investigating the maternal immune response as a variance moderator, investigating the variance within and between litters is of high interest. While we do include litter as a random intercept in our mixed-effects modeling, investigating this more closely may be of high interest for future work. We comprehensively examined the effects of prenatal MIA exposure, a known risk factor for neuropsychiatric disorders, at two gestational timepoints on embryo brain anatomy at the gross morphological and cellular levels. We identified striking neuroanatomical remodeling in the embryo brain, particularly following exposure in late gestation; we also observed sex-dependent alterations in the density of dark neuronal and glial cells in the dorsal hippocampus, with greater cell density in female offspring. This may reflect the initiation of pathological circuit remodeling. These findings show that MIA exposure induces striking neurodevelopmental changes in embryonic development, which may further our understanding of how this risk factor increases the likelihood of developing neuropsychiatric illnesses later in life.

4. Materials and Methods

4.1. Animals, Prenatal Immune Activation, and Sample Preparation. C57BL/6J female and male mice of breeding age (8 to 12 wk old) were subject to timed mating procedures (described in SI Appendix, section 1.1) to generate pregnant dams. Pregnant dams were randomly assigned to one of four treatment groups (see Fig. 4 for experimental design): 1) poly I:C (P1530-25MG poly I:C sodium salt TLR ligand tested; Sigma-Aldrich) (5 mg/kg, i.p.) at GD9 (POL E; seven embryo dams); 2) 0.9% sterile NaCl solution (saline) at GD9 (SAL E; six embryo dams); 3) poly I:C at GD17 (5 mg/kg, i.p.) (POL L; seven embryo dams); or 4) saline at GD17 (SAL L; four embryo dams). Additionally, immunostimulatory potential of poly I:C was confirmed in a separate group of dams (see SI Appendix, section 1.1 for methods and SI Appendix, section 2.1, Table 1, and Figs. 1–3 for results).

4.1.1. Sample preparation for MRI. On GD18, pregnant dams were euthanized; embryos were extracted and postfixed in 4% paraformaldehyde with 2% gadoxinum (MRI contrast agent; Bracco Imaging S.p.A) in phosphate-buffered saline (PBS) for 1 wk. A piece of the yolk sac was collected for genotyping of each embryo to identify the sex of the mouse via presence of the SRY gene (performed by Transnetyx). Collections were performed in two separate cohorts (with two different poly I:C batches from the same supplier outlined in SI Appendix, section 1.2 and Table 1).

4.2. Magnetic Resonance Image Acquisition and Processing. All samples were shipped to the Mouse Imaging Centre (Toronto) for scanning. A multichannel 7.0-T MRI scanner with a 40-cm-diameter bore (Varian Inc.) was used to acquire anatomical images of the entire embryo (whole body). A custom-built 16-coil solenoid array was used to acquire 40-μm3-resolution images from 16 samples concurrently (72) (see SI Appendix, section 1.3.1 for details).

Preprocessed embryo brain images (https://github.com/CoBrALab/documentation/wiki/Embryo-scan-preprocessing) of all subjects in the study were aligned by unbiased deformation-based morphometry using the antsMultivariateTemplateConstruction2.sh tool (https://github.com/CoBrALab/twolevel_ants_dbm) (73). The output of this iterative group-wise registration procedure generates a group average from all the scans in the study, as well as the minimum deformation field that is required to accurately map each individual subject to the average at the voxel level (see 3D representation of group average in SI Appendix, Fig. 5). Relative Jacobian determinants (74), which explicitly model only the nonlinear deformations and remove global linear transformation (attributable to differences in total brain size), were blurred by using a Gaussian kernel at 160-μm full-width-at-half-maximum to better conform to Gaussian assumptions for downstream statistical testing (see SI Appendix, section 1.3.2 for details).

4.3. EM. After MRI scanning, embryo brains (SAL E, n = 4 [two males/two females]; SAL L, n = 7 [three males/four females]; POL E, n = 8 [four males/four females]; and POL L, n = 8 [four males/four females]) were extracted from the fixed samples and further postfixed with 3.5% acrolein in phosphate buffer (100 mM) (pH 7.4) overnight at 4 °C. Postfixed brains were sectioned to 50-μm sagittal slices by using a VT1200S vibratome (Leica Biosystem) and stored at −20 °C in cryoprotectant (30% glycerol and 30% ethylene glycol in PBS [50 mM] [pH 7.4]). Three brain sections in which the dorsal hippocampus was present [coronal section 12-15 (75), roughly equivalent to lateral 2.0 to 2.8 mm (76)], were processed for EM by using osmium-thiocarbohydrazide-osmium postfixation (77) (SI Appendix, section 1.4.1). Samples were sectioned into 70- to 75-nm ultrathin sections by using an Ultracut UC7 ultramicrotome (Leica Biosystems). Three levels of section-rubens were collected at an interval of 10 μm, glued on a specimen mount with conductive carbon adhesive tabs (Electron Microscopy Sciences), and imaged by array tomography at 25-nm resolution with an acceleration voltage of 1.4 kV and current of 1.2 nA using a Zeiss Crossbeam 540 Gemini scanning EM (Zeiss) (three images per embryo).

Images from all the four treatment groups were analyzed blind to the experimental conditions by using QuPath (v0.2.0-m3) software (78). For each picture, region areas were traced and measured to calculate cell density. Cell type and apoptotic state were determined by ultrastructural features. Total cell numbers, dark cells (neuronal and glial cells), and apoptotic cells were then counted within the dorsal hippocampus (CA1, CA3, and dentate gyrus). The percentage of dark cells was acquired anatomical images of the entire embryo (whole body). A custom-built 16-coil solenoid array was used to acquire 40-μm3-resolution images from 16 samples concurrently (72) (see SI Appendix, section 1.3.1 for details).

Preprocessed embryo brain images (https://github.com/CoBrALab/documentation/wiki/Embryo-scan-preprocessing) of all subjects in the study were aligned by unbiased deformation-based morphometry using the antsMultivariateTemplateConstruction2.sh tool (https://github.com/CoBrALab/twolevel_ants_dbm) (73). The output of this iterative group-wise registration procedure generates a group average from all the scans in the study, as well as the minimum deformation field that is required to accurately map each individual subject to the average at the voxel level (see 3D representation of group average in SI Appendix, Fig. 5). Relative Jacobian determinants (74), which explicitly model only the nonlinear deformations and remove global linear transformation (attributable to differences in total brain size), were blurred by using a Gaussian kernel at 160-μm full-width-at-half-maximum to better conform to Gaussian assumptions for downstream statistical testing (see SI Appendix, section 1.3.2 for details).

Table 1. Final sample size for embryo MRI data following quality control

| Group | Males MRI (cohort 1/cohort 2) | Females MRI (cohort 1/cohort 2) | Litters (cohort 1/cohort 2) |
|-------|-------------------------------|--------------------------------|---------------------------|
| SAL E | 14 (13/1)                     | 15 (13/2)                      | 7 (6/1)                   |
| SAL L | 12 (10/2)                     | 12 (6/6)                       | 4 (2/2)                   |
| POL E | 11 (6/5)                      | 17 (9/8)                       | 7 (4/3)                   |
| POL L | 14 (11/3)                     | 17 (15/2)                      | 7 (6/1)                   |

Collection was performed in two rounds with two different batches of poly I:C (same supplier and product). Number indicates total sample size following quality control. The number of samples coming from each collection cohort is noted in parentheses.

4.3. EM. After MRI scanning, embryo brains (SAL E, n = 4 [two males/two females]; SAL L, n = 7 [three males/four females]; POL E, n = 8 [four males/four females]; and POL L, n = 8 [four males/four females]) were extracted from the fixed samples and further postfixed with 3.5% acrolein in phosphate buffer (100 mM) (pH 7.4) overnight at 4 °C. Postfixed brains were sectioned to 50-μm sagittal slices by using a VT1200S vibratome (Leica Biosystem) and stored at −20 °C in cryoprotectant (30% glycerol and 30% ethylene glycol in PBS [50 mM] [pH 7.4]). Three brain sections in which the dorsal hippocampus was present [coronal section 12-15 (75), roughly equivalent to lateral 2.0 to 2.8 mm (76)], were processed for EM by using osmium-thiocarbohydrazide-osmium postfixation (77) (SI Appendix, section 1.4.1). Samples were sectioned into 70- to 75-nm ultrathin sections by using an Ultracut UC7 ultramicrotome (Leica Biosystems). Three levels of section-rubens were collected at an interval of 10 μm, glued on a specimen mount with conductive carbon adhesive tabs (Electron Microscopy Sciences), and imaged by array tomography at 25-nm resolution with an acceleration voltage of 1.4 kV and current of 1.2 nA using a Zeiss Crossbeam 540 Gemini scanning EM (Zeiss) (three images per embryo).

Images from all the four treatment groups were analyzed blind to the experimental conditions by using QuPath (v0.2.0-m3) software (78). For each picture, region areas were traced and measured to calculate cell density. Cell type and apoptotic state were determined by ultrastructural features. Total cell numbers, dark cells (neuronal and glial cells), and apoptotic cells were then counted within the dorsal hippocampus (CA1, CA3, and dentate gyrus). The percentage of dark cells was...
cell population or apoptotic cell number was calculated as a ratio over the total cell population (details in SI Appendix, section 1.4.1). Finally, immunocytochemical transmission EM was used to determine whether cells identified as apoptotic, those with a pyknotic nucleus, were also positive for active Caspase 3 staining. Additionally, to investigate whether dark cells identified were specifically microglia, immunocytochemical staining against CD11b was performed (see SI Appendix, section 1.4.2 for details).

### 4.4. Statistical Analyses.

#### 4.4.1. Neuroimaging data analysis.

Statistical analyses were performed by using the R software package (R version 3.5.1, RMINC version 1.5.2.2; https://www.r-project.org/). To assess the effects of poly i:C exposure at different gestational timepoints on embryo neuroanatomy, we ran a whole-brain voxel-wise linear mixed-effects model ("minLmer"; lme4_1.1.21 package (79)) on the relative Jacobian determinant files using group and sex as fixed effects and number of pups per litter and cohort collection batch as random intercepts. A Satterthwaite approximation was used to compute degrees of freedom for every voxel (using the "minLmerEstimateDF" function). The FDR correction (using "minFDR") was applied to correct for multiple testing (80, 81) (see SI Appendix, section 1.5.1 for details). This analysis was run again with the POL L group as the reference in order to directly compare POL E to POL L differences (SI Appendix, section 2.2.1). Sex differences were explored as a follow-up analysis (SI Appendix, section 1.5.1). Putative differences in organ volume (i.e., lungs, heart, liver, etc.) for the embryos were also investigated by applying deformation-based morphometry (as described above) to the body cavity (SI Appendix, section 1.5.1); however, no differences were observed (SI Appendix, section 2.2.2).

#### 4.4.2. EM. A nonparametric Kruskal–Wallis test followed by a painwise Wilcoxon test was used to assess group differences in the density of dark glial, dark neuroglia, and apoptotic cells in the combined SAL, POL E, and POL L groups averaged per subject across slices (the SAL group was combined to maintain consistency with the MRI findings).

To maximize variance, the shift function was used to compare differences in the distribution of density measures acquired from each slice per mouse (3), rather than using the pooled data per mouse (34). This allows us to quantify how two distributions differ based on deciles of the distributions; i.e., it describes how one distribution should be transformed to match the other and estimates how and by how much one distribution must be shifted. When a significant difference is observed between deciles, it suggests that there is a specific difference in the density of the cell type investigated (i.e., dark glia) between groups; this allows us to determine whether differences are consistent across the entire distribution or more localized to one or both tails or the center. In the context of the cell-density data acquired, this technique allows us to compare groups beyond means or medians by accounting for the variance across the entire distribution of the data; this may provide us with a more nuanced understanding of differences between groups. Three pairwise comparisons were made (SAL–POL E, SAL–POL L, and POL L–POL E) on the distributions for dark glia, dark neuron, and apoptotic cell density. A percentile bootstrap technique was used to derive CIs based on differences in distribution at each decile of the distribution with a corresponding P value. This was then repeated to assess sex differences, so the same comparisons were made in only males and only females, followed by the same percentile bootstrap procedure.

**Data Availability.** Structural MRI and EM cell counts and cell-density data have been deposited in Zenodo, as is the code used for statistical analyses (http://doi.org/10.5281/zenodo.5164621). Instructions for running the ANTs DBN analysis can be found on GitHub (https://github.com/CoBrALab/twolevel-ants-dbms), as can information on MRI preprocessing steps (https://github.com/CoBrALab/documentation/wiki/Embryo-scan-preprocessing).

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