Impacts of Preeclampsia on the Brain of the Offspring

Impactos da pré-eclâmpsia no cérebro de nascituros

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Abstract

Preeclampsia (PE) is a significant gestational disorder that causes complications in 3–5% of all human pregnancies. Apart from the immediate risks and complications for mother and fetus, both additionally carry elevated lifelong risks for specific complications. Offspring of PE pregnancies (PE-F1) have higher risks for hypertension, stroke and cognitive impairment compared with well-matched offspring (F1) from uncomplicated pregnancies. Prior to the clinical onset of PE, placental angiokines secreted into the maternal plasma are deviated. In many PE patients this includes deficits in placental growth factor (PGF). Our laboratory found that mice genetically-deleted for PGF (PGF−/−) have altered cerebrovascular and brain neurological development detectable from midgestation to adulthood. We hypothesized that the PGF deficits seen in human PE, deviate fetal cerebrovascular and neurological development in a manner that impairs cognitive functions and elevates stroke risk. Here we summarize the initial analytical outcomes from a pilot study of 8–10 year old male and female PE-F1s and matched controls. Our studies were the first to report magnetic resonance imaging (MRI), magnetic resonance angiography (MRA) and functional brain region assessment by eye movement control and clinical psychometric testing in PE-F1s. Further studies in larger cohorts are essential to define whether there are image-based biomarkers that describe unique anatomical features in PE-F1 brains.

Keywords
► preeclampsia
► PGF
► fetal brain
► MRI
► cognitive development

Resumo

A pré-eclâmpsia (PE) é importante doença gravídica complicando 3–5% de todas as gestações humanas. Além dos riscos imediatos e complicações para a mãe e o feto, a PE associa-se a outros riscos materno-fetais elevados em longo prazo. Nascituros de gestações complicadas por PE (PE-F1) apresentam maiores riscos de desenvolver hipertensão, acidente vascular cerebral e disfunção cognitiva em comparação com prole (F1) de gestações sem complicações. Antes do aparecimento clínico da PE, angiocitocinas placentárias secretadas no plasma materno apresentam-se alteradas. Em muitos pacientes com PE, isso inclui valores plasmáticos reduzidos de Fator de Crescimento Placentário (PGF). Nosso laboratório identificou que camundongos


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Introduction

Preeclampsia (PE) is a significant clinical gestational disorder occurring in 3–5% of all human pregnancies, and is among the leading causes of maternal and fetal morbidity and mortality.\(^1\) PE accounts for up to 12% of all annual global maternal deaths,\(^2\) and up to 25% of all annual global fetal and neonatal deaths.\(^3\) PE by definition is new-onset hypertension (\(>140/90\) mmHg) and at least one of: proteinuria (\(>300\) mg/day), thrombocytopenia (\(<10^5/\mu L\)) renal insufficiency (serum creatinine \(>1.1\) mg/dL), impaired liver function, pulmonary edema, headaches or visual disturbances during the 20th week of gestation.\(^4\) The systemic presentation of PE has been best explained as systemic vascular inflammation.\(^5\) Numerous disturbances in the levels of angiogenic molecules precede and accompany clinical signs, such as low placental growth factor (PGF) and elevation of the soluble form of its receptor (sFLT1).\(^6,7\)

Beyond the immediate gestational complications of PE, numerous long-term maternal complications have been identified.\(^8–10\) Women who have experienced PE have significantly increased risks of developing future cardiovascular risk factors such as dyslipidemia,\(^11\) hypertension\(^12\) and metabolic disease.\(^13\) PE at least doubles the risk of future heart disease,\(^14,15\) elevating lifetime risks for coronary artery disease,\(^16\) cardiovascular disease and stroke.\(^17\)

This review explores the impact of preeclampsia upon the brain of offspring from preeclamptic gestations. While PE’s impact on brain vascular and neurological development occurs during fetal life, postnatal brain assessments are used to study the legacy of the preeclamptic gestation in the offspring. Preliminary outcomes of a novel pilot investigation that we undertook in 8–10-year-old children born from singleton PE gestations (PE-F1) are also discussed.\(^18\) These studies were the first to report Magnetic Resonance Imaging (MRI), Magnetic Resonance Angiography (MRA) and functional brain region assessment using eye movement control testing in PE-F1s.\(^19\) A mechanistic pathway for PE-induced deviations in brain development is proposed from a mouse model of PGF deficiency. Finally, some ideas regarding screening and future approaches for potential therapeutic interventions are introduced, should further studies of larger populations support the pilot study findings.

Methods

The MEDLINE database to March 2016 was searched for articles published in English between 1990–2016 that focused on preeclampsia but additionally mentioned brain function or cognition of offspring. The search terms used were: preeclampsia, cognition, cognitive tests, fetal, brain, offspring, children, newborn, and eye movements (or any of its synonyms). This database screen identified 277 publications. After reading the title and/or abstract, many of these publications were judged to be relevant and were not included in this review. A total of 57 articles were relevant and provided the basis for this review.

Preeclampsia (PE) and Effects on PE-F1s

Offspring born to PE pregnancies (PE-F1s) exhibit elevated lifetime risks for several health disorders and impaired functional capacities across multiple body systems, including the cardiovascular, endocrine and neurological systems.\(^20–22\) In particular, PE-F1s from the Helsinki Birth Cohort\(^23\) are reported to have deficits in cognitive function and elevated stroke risk.\(^24–26\)

During childhood and young adulthood, PE-F1s display a body mass index (BMI) 0.6 kg/m\(^2\) higher than children born to uncomplicated pregnancies.\(^27\) These PE-F1s also display 2.5 mmHg higher systolic and 1.4 mmHg higher diastolic blood pressure during the same timeframe.\(^27\) These increases in pressure translate to an approximate 2-fold increased risk of stroke in adulthood.\(^26\)

PE-F1s are also more likely to experience cerebrovascular and cognitive related disorders than offspring born to non-PE pregnancies matched for gestation length and current age or maternal hypertension during the index pregnancy.\(^28\) As children, PE-F1s exhibit deficits in several cognitive function domains,\(^28\) including verbal reasoning.\(^29\) They also score lower for total intelligence quotient (IQ)\(^30\) and mental development indices (MDI).\(^31,32\) As these children move through adolescence and adulthood, deficits in verbal and arithmetic reasoning persist.\(^28\) During adulthood and up to old age, PE-F1s display more depressive symptoms and higher rates of cognitive decline.\(^24\)

- Table 1 summarizes key findings related to the PE-F1 brain and cognition that have been published to date.
Table 1 Brain and cognition related main findings previously reported in humans. PE-F1s

<table>
<thead>
<tr>
<th>Author (year)</th>
<th>Findings in PE-F1s</th>
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<tbody>
<tr>
<td>Many et al(^{10}) (2003)</td>
<td>Lower IQ scores</td>
</tr>
<tr>
<td>Kajantie et al(^{26}) (2009)</td>
<td>Smaller head circumferences at birth; elevated risk of stroke</td>
</tr>
<tr>
<td>Tuovinen et al(^{44}) (2010)</td>
<td>Higher rates of depression</td>
</tr>
<tr>
<td>Whitehouse et al(^{29}) (2012)</td>
<td>Reduced verbal ability</td>
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<tr>
<td>Tuovinen et al(^{22}) (2012)</td>
<td>Greater cognitive decline at old ages</td>
</tr>
<tr>
<td>Morsing and Maršáľ(^{12}) (2014)</td>
<td>Lower mean verbal IQ (VIQ) and lower full scale IQ (FSIQ)</td>
</tr>
<tr>
<td>Rätsep et al(^{18}) (2015)</td>
<td>Enlarged brain regional volumes in five regions (cerebellum, temporal lobe, brainstem and right and left amygdala)</td>
</tr>
<tr>
<td>Rätsep et al(^{19}) (2016)</td>
<td>Deficits in working memory and visuospatial processing</td>
</tr>
</tbody>
</table>

Abbreviations: PE-F1, offspring of preeclampsia pregnancies; IQ, intelligence quotient; VIQ, verbal intelligence quotient; FSIQ, lower full scale intelligence quotient.

Preeclampsia (PE) and Placenta Growth Factor (PGF) - A Close Relationship

During human pregnancy, several angiogenic factors become expressed at increased levels to support the growth, development and viability of the conceptus through an uncomplicated pregnancy. In particular, vascular endothelial growth factor (VEGF) and its related family member, placental growth factor (PGF), are highly expressed at predictable times over pregnancy. A deficiency in maternal plasma PGF has been linked to an increased likelihood of PE. The primary source for the gestational elevation of angiokines is the placenta.

PGF serves as a biomarker predictive of PE, particularly when combined with clinical factors such as blood pressure or other angiogenic markers such as soluble fms-like tyrosine kinase-1 (sFLT) and soluble endoglin (sENG). In addition, low maternal levels of PGF in early to mid-pregnancy are thought to be a marker for distinguishing between two types of PE.

Mouse studies suggest that the normal gestational roles of PGF are to optimize vascular development within decidua basalis and to sustain normal maternal cardiac function in late gestation. In mice genetically engineered to be PGF-deficient (PGF \(-/\)) , our laboratory identified reduced and aberrant vascular branching in the decidua basalis during early pregnancy (gestation days (GD) 6.5–9.5 of the term 19 day mouse gestation) and enlarged labyrinthine vascular spaces in late placentas (GD 15.5–18.5). The latter finding was interpreted as a negative feature resulting from reduced vascular branching into capillaries in the placental exchange region.

Pilot Study Brain and Cognitive Outcomes for PE-F1s

A recent pilot study used 8–10 year old children (n = 20) whose mothers were part of a research cohort, the Preeclampsia Network (PE-NET). Five boys and five girls whose mothers’ pregnancies were complicated by PE (n = 10) were matched by sex, gestation and current age with children who experienced normal pregnancies (n = 10). All pregnancies were singleton. All of the children were assessed by the same protocol, which included clinical cognitive testing (NEPSY II), eye movement control tests and Magnetic Resonance Imaging and Angiography (MRI/MRA) data collection sequences. The consented and assented children underwent MRI/MRA to evaluate their brain structural and vascular anatomy. The PE-F1s exhibited enlarged brain regional volumes in five regions (cerebellum, temporal lobe, brainstem and right and left amygdalae) when compared with their cohort-matched controls. Diffusion Tensor Image (DTI) analysis suggests further alterations (manuscript in preparation).

Time of flight (TOF) MRA analysis data revealed significant differences in the occipital and parietal lobes. In both the occipital and parietal lobes, the mean vessel radius was significantly shorter in the PE group (control: 0.50 ± 0.01 mm versus PE: 0.45 ± 0.01 mm, p = 0.004; control: 0.55 ± 0.01 mm versus PE 0.52 ± 0.01 mm, p = 0.025 respectively). To understand the significance of the MRA findings, an MRI sequence for Arterial Spin Labeling (ASL) could be conducted, and would determine if brain perfusion is lower in regions with smaller caliber blood vessels.

Psychometric testing outcomes revealed overall deficits in working memory and visuospatial processing amongst PE-F1s. These deficits appear to correlate with the anatomic alterations identified in our MRI analysis within the occipital lobe, parietal lobe, cerebellum and brain stem. Eye movement control impairments also correlate with several of the regions that showed structural deviation in PE-F1s.

Plasma samples collected at term from the mothers of 12 study participants were available for PGF quantification by enzyme linked immunosorbent assay. Samples from the PE mothers had significantly lower levels of PGF than samples from the control mothers (control: 221.0 ± 46.6 pg/mL versus PE: 37.2 ± 21.5 pg/mL, p = 0.024). This suggests that at least some of the PE-F1 study participants had aberrant placental PGF production. The PGF overall data were non-correlative and do not indicate that fetal PGF levels were lower than normal. The data also do not exclude the possibilities that other angiogenic or neurodevelopmental pathways provide a primary etiology to explain the findings of this pilot study. Power calculations based on these data...
estimate 76 pairs of PE-F1 and control children would be needed within a single time window of PE (that is, term or late preterm) to validate the observed anatomic differences.

**Preeclampsia-deviated Brain Development – A Proposed Pathway**

Fetal brain undergoes great constitutional change and growth during development to build the structural, vascular and neurological anatomy that supports future autonomic and cognitive functions. Because the brain requires oxygen and nutrients provided by circulation, development of the fetal cerebral circulation is coincident with brain neural development, and both tissues share many common molecular pathways. Axon induction, guidance and arterial specification all use VEGF and PGF.

Fetal human brain development begins with the formation of the neural tube at gestational age (GA)3–4 weeks. Following neural tube closure, the rostral portion differentiates into three vesicles that eventually form the forebrain, midbrain and hindbrain. Simultaneously, the fetal cerebral vasculature forms. Six pairs of brachial arches, each containing primitive branchial arch arteries, are present at approximately GA18 days. The internal carotid arteries, which supply the anterior portion of the developing fetal brain, begin to form at GA24 days. At GA28 days, each internal carotid artery splits into anterior and posterior divisions, driven by the formation of the brain stem and the occipital lobe. Once the main branches of the cerebral arterial tree have formed, the fetal brain utilizes this blood supply to grow rapidly. Neurons begin to migrate along glial cell scaffolds, and the brain folds into gyri and sulci. In circumstances of maternal stress and uteroplacental nutrient deprivation, human fetal brain development takes priority over other tissues, with evidence of head sparing effects.

PGF-deficient mice (PGF–/–) display altered fetal brain vascular development by mid pregnancy (gestation day 10.5). The earliest deviations are in vessels of the hindbrain, then during development of the Circle of Willis (CW) and, in the early postpartum interval, the growing vascular plexus of the retina. These vascular alterations include narrower lumens, deficient collateral branching and atypical crossovers, deviations that persist into adulthood. Neuronal tissue structure is unlikely to be normal in these mice because adult differences in regional brain volumes have been detected by MRI, and behavior of the mice is atypical. Vascular alterations may underlie these neurological tissue changes, or they may be due to absence of PGF in the neurons themselves. PGF has been studied as a member in the cytokine network of Wallerian Degeneration. After injury, PGF–/– mice have decreased Schwann cell proliferation and present less macrophage invasion than matched inbred controls, resulting in poorer functional recovery.

Overall, these mouse studies suggest PGF plays a crucial role in neurovascular development, and that its deficiency contributes to long-term neuropathology. Since PGF appears to occur at all stages of brain development, it is unclear exactly when the phase of suboptimal human gestational PGF occurs, but reports have indicated it to be around the end of the first trimester. Overlapping of timeframes for suboptimal placental PGF production and CW development occur in humans.

The Circle of Willis is formed between the first and second months of pregnancy (40–55 days), and low PGF levels are present in maternal plasma in the majority of women who develop PE during the first trimester and as early as week 7. If genetic and/or epigenetic mechanisms down-regulate PGF prior to embryonic gastrulation, the derivatives of both inner cell mass and trophoderm cell lineages would be expected to have similar PGF deficient phenotypes. Therefore, vascular alterations in F1s are to be expected from PGF-insufficient PE pregnancies. To date, PGF levels across gestation have been followed in human placenta and maternal plasma but not the fetus.

We hypothesized that the mechanisms deviating placenta-derived angiokines in the circulating women who progress to PE are established prior to blastocyst formation, and are as equally expressed in the tissue derivatives of the inner cell mass, that is, the fetus (Fig. 1), as in the placenta. This hypothesis implicates deviations in fetal synthesis of PGF or other angiokines, rather than maternal hypertension or deviated placental angiokine synthesis as potential mechanisms that could compromise brain vascular development, brain structure and cognitive functions of PE-F1s. This hypothesis additionally suggests that deviations in PE-F1 vascular development are widespread and not restricted to the brain. This may be a component of the elevated cardiovascular disease risk reported in PE-F1s.

**Significance**

The combination of MRI studies with neurocognitive tests is a proven approach to identify and link deviations in brain structure with cognition for several childhood conditions. Data from the detailed pilot studies that combined MRI/MRA, eye movement control and cognitive function tests of 8–10 year old PE-F1s strongly suggest that extension of this approach to PE-F1s will identify consistent vascular and neuroanatomic anomalies, that is, an image phenotype of the PE-F1 brain. This phenotype may explain the cognitive deviations and elevated stroke risk that have been reported in pediatric or adult PE-F1 populations.

The neurocognitive subtests chosen for the pilot study were those used for children participating in NeuroDevNet’s studies. NeuroDevNet is a Canadian Centers of Excellence Consortium that studies neuro- logical brain development and function in children with Fetal Alcohol Spectrum Disorder, Cerebral Palsy and Autism Spectrum Disorder. Selection of these testing paradigms gives the additional exciting possibility for future research outcome comparisons between PE-F1s and children with other major neurodevelopmental disorders.

Further work to define image-based biomarkers describing a unique PE-F1 brain anatomy could lead to personalized interventions and therapies aimed at preventing the development of fetal brain aberrations or their postnatal amelioration. For both the parents and the PE-F1 child, knowledge of the impacts of a PE gestational complication may lead to brain-region specific enhanced educational support to improve the
child’s typical academic and social progress, and to the consideration of stroke prevention prophylaxis at younger ages. The relatively small sample size and racial homogeneity (Caucasian) of the pilot study participants are weaknesses.

Conclusion

Gestations that include fetal exposure to PE appear to elevate risk for altering cerebral vascular and neuroanatomy during development. Such changes during fetal life may explain the postnatal findings of elevated risks for stroke and specific deviations in cognitive functioning, including visual spatial processing and memory. A large, appropriately controlled, cohort study is needed to validate the current findings. Characterization of a PE-F1 “brain imaging signature” could eventually help identify individuals who may need enhanced educational or medical support.

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