The Role of Fetal Neurology in the Progress of Perinatal Medicine

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Abstract

The prevalence of cerebral palsy has not decreased despite major improvements in clinical care in antenatal/ neonatal period as well as intrapartum period. In about 70% of cases, cerebral palsy results from events occurring before birth that can disrupt normal development of the brain. The antepartum risk factors should include fetal brain maldevelopment and intrauterine brain injuries, which are unclassifiable into congenital brain anomalies and may exist inconspicuously during pregnancy and even after birth. Especially, neuronal migration disorder and acquired brain damage in utero should be responsible for postnatal neurological impairment. Imaging technologies including three dimensional ultrasound have been remarkably improved and contributed to prenatal evaluation of fetal Central Nervous System (CNS) development and assessment of CNS abnormalities in utero. In this article, objective and precise imaging diagnoses of fetal CNS including migration disorders and acquired brain damages. Furthermore, 3D bidirectional power Doppler angiography has depicted fine cerebral vessels of medullary veins which may relate with timing of insult as well as with postnatal neurological prognosis. It is promising to clarify the developmental mechanism of CNS damages with advanced ultrasound diagnostic techniques in the near future. Postnatal unexplained neurological deficits may strongly relate with intrauterine brain development therefore fetal neurology has great responsibility and an important role in perinatal medicine.

Keywords: Fetus; Neurology; Congenital; Brain; Vascularity; Ultrasound

Introduction

In perinatal medicine, one of the most important roles is "to lessen the number of infants with neurological impairment. Cerebral Palsy (CP) is a non-progressive motor disability and its prevalence has remained largely unchanged despite major improvements in clinical care in antenatal/neonatal period as well as intrapartum period. The Surveillance of Cerebral Palsy in Europe (SCPE) reported that the current prevalence of CP is similar in six countries in the registry (2.12 to 2.45 per 1000 live births) as is the increasing trend in time from well below 2 per 1000 (1.7) live births in the 70's to well above 2 per 1000 live births (2.4) in the 90's [1]. The SCPE also reported that the proportion of low-birth weight infants among all children with CP is rising: 32% of all cases in 1966 and 50% in 1989. It can be very difficult to give an exact reason as to why the brain has been injured or failed to develop. In some instances, there may be no obvious single reason of cerebral palsy. It is generally accepted that causes of cerebral palsy can be multiple and complex. These can include infection in the early part of pregnancy, oxygen deprivation to the brain, abnormal brain development, and restricted intrauterine growth. Furthermore, multiple risk factors such as premature birth, multiple pregnancy, advanced maternal age, low birth weight have been considered.

In about 70% of cases, cerebral palsy results from events occurring before birth that can disrupt normal development of the brain. Contrary to common belief, lack of oxygen reaching the fetus during labor and delivery contributes to only a small minority of cases of cerebral palsy, according to a 2003 report by the American College of Obstetricians and Gynecologists (ACOG) and the American Academy of Pediatrics (AAP) [2]. They showed a multivariate analysis of 164 cases of neonatal encephalopathy showed 69% had only antepartum risk factors, 25% had both antepartum risk factors and evidence for intrapartum hypoxia, 4% had evidence of intrapartum hypoxia in the absence of preconceptional or antepartum factors that might also have contributed to neonatal encephalopathy and 2 % had no recognized risk factors for neonatal encephalopathy. The antepartum risk factors should include fetal brain maldevelopment and intrauterine brain injuries, which are unclassifiable into congenital brain anomalies and may exist inconspicuously during pregnancy and even after birth.

Imaging technologies have been remarkably improved and contributed to prenatal evaluation of fetal central nervous system (CNS) development and assessment of CNS abnormalities in utero. Introduction of high-frequency transvaginal transducer has contributed to establishing "sonoembryology" [3] and recent general use of transvaginal sonography in early pregnancy enabled early diagnoses of major fetal anomalies [4]. In the middle and late pregnancy, fetal CNS is generally evaluated through maternal abdominal wall. The brain, however, is three-dimensional structure, and should be assessed in basic three planes of sagittal, coronal and axial sections. Sonographic assessment of the fetal brain in the sagittal and coronal sections requires an approach from fetal parietal direction. Transvaginal sonography of the fetal brain opened a new field in medicine, "neurosonography" [5] Transvaginal approach to the normal fetal brain during the second and third trimester was introduced in the beginning of 1990s [6,7]. Transvaginal observation of the fetal brain offers sagittal and coronal views of the brain from fetal parietal direction through the fontanelles and/or the sagittal suture as ultrasound windows [8,9]. Three dimensional (3D) ultrasound combined with transvaginal approach has greatly contributed to neuroimaging diagnoses [10-14]. Fetal 3D neuroimaging has revealed congenital brain anomalies, such as cranium bifidum, holoprosencephaly, agenesis of the corpus callosum, Chiari malformation, and Dandy-Walker complex in a category of primary/ secondary neurolucation and proencephalic development [15]. Magnetic

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Received June 12, 2013; Accepted June 25, 2013; Published June 30, 2013

Citation: Pooh RK (2013) The Role of Fetal Neurology in the Progress of Perinatal Medicine. J Health Med Informat S11: 015. doi:10.4172/2157-7420.S11-015

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resonance imaging is the other great modality to demonstrate fetal CNS during pregnancy [16,17]. However, beside congenital anomalies, acquired brain damages caused by hypoxic-ischemic episodes and migration disorders should be responsible for postnatal cerebral palsy. It is a difficult task to provide a precise prediction of subsequent development of cerebral palsy after a given antepartum event or complication. Fetal heart rate monitoring cannot reveal the presence of encephalopathy, and neuroimaging by ultrasound and MR imaging is the most reliable modality for disclosure of silent encephalopathy. In many cases with cerebral palsy with acquired brain insults, especially term-delivered infants with reactive fetal heart rate tracing and good Apgar score at delivery, recent imaging study have confirmed the presence of brain insult in utero, suggesting that the majority of cerebral palsy are of antenpartum rather than intrapartum in origin. Furthermore, mal arrangement of nerve cells, so called migration disorder, with unknown silent causes has not been well clarified even by neuroimaging during fetal period. The author believes that acquired brain insults and migration disorders bears very grave responsibility in cerebral palsy and neurological deficit after birth, and investigation and explication of fetal brain must be the key to the mystery of fetal brain. In this article, the up-to-date fetal neurological evaluation and the future perspectives are described.

Method

The representative of fetal neurological evaluation is definitely neuroimaging by ultrasound. 3D ultrasound is one of the most attractive modalities in the field of fetal ultrasound imaging. Automatic scan by dedicated 3D transducer produces motor driven automatic sweeping and is called as a fan scan. With this method, shifting and/ or changing angle of the transducer is not required during scanning and scan duration lasts only several seconds. After acquisition of the target organ, multiplanar imaging analysis and tomographic imaging analysis are possible. Combination of both transvaginal high-resolution sonography and 3D ultrasound may be a great diagnostic tool for evaluation of three-dimensional structure of fetal CNS [18,19]. Recent advanced 3D ultrasound equipments have several useful functions such as surface anatomy imaging of spinal development, cranial surface and brain surface, bony structural imaging of the calvaria and vertebrae, multiplanar imaging of the intracranial structure, tomographic ultrasound imaging of fetal brain in any cutting section, thick slice imaging of the intracranial structures (Volume contrast imaging, VCI), simultaneous volume contrast imaging of the same section or vertical section of fetal brain structure, volume calculation [20,21] of target organs such as intracranial cavity, ventricle, choroid plexus and intracranial lesions, inversion mode (the cystic portions within the volume are displayed in their entirety as an echogenic area, while the grayscale portions of the image are rendered as transparent [22-24]) demonstrating hypoechoic parts such as ventricles and cystic lesions, and 3D power Doppler angiography of the intracranial vascularity [25,26]. By utilizing those 3D functions, objective and precise fetal neurological assessment can be done.

Results

Fetal brain demonstrated by 3D ultrasound

In early pregnancy, 3D volume imaging shows surface anatomy of small fetus and intracranial structure in the mid-sagittal section (Figure 1). Obviously different appearance of the brain surface at 19 and 30 weeks of gestation are demonstrated by 3D surface anatomy imaging (Figure 2). Tomographic ultrasound imaging of the brain structure is quite helpful to understand intracranial detailed brain structure (Figure 3).
fetal brain circulation in the axial and sagittal sections at 31 weeks by 3D bidirectional power Doppler. Furthermore, recent high-frequent transvaginal neuroscan has been able to demonstrate the medullary vessels from the cortex towards subependymal area. Medullary veins are demonstrated between pial mater and longitudinal vein of Schlesinger in the anterior coronal cutting section and between brain surface and periventricular subependymal area in the parasagittal section [27]. Medullary vessels are detectable from early second trimester and they are rapidly developing during second trimester. Before 20 weeks, medullary veins are demonstrated as red color or red/blue colors by bidirectional power Doppler (Figure 7, left) and thereafter remarkable rapid development is seen into shower-like vessels (Figure 7, right).

Migration disorders

Abnormal migration of neurons in the developing brain and nervous system causes various brain maldevelopment. Lissencephaly is one of representatives of migration disorders. Figure 8 shows lissencephaly at 30 weeks of gestation with no gyral pattern. Figures 9 and 10, show non-uniformed migration disorder, which resulted in severe neurological prognosis after birth. It has been believed that phenotypes of migration disorders appear in late pregnancy when gyral pattern is visualized. However, the author has experienced some cases with obvious phenotypes of migration disorder from early second trimester. Figure 11 shows maldeveloped brain with migration disorder at 18 weeks of gestation. Prenatal neuroimaging clearly matches the histological findings.

Acquired brain insults

Acquired brain insults can occur at any gestational age. Figure 12-15 show cases of encephalopathy in various gestations. In Figure 12, early brain hemorrhage with unknown cause may lead to split of brain and intraventricular hemorrhage. Intrauterine encephalopathy can result from various causes, such as vein of Galen aneurysmal malformation.
ventriculomegaly. The postnatal neurological prognosis has been good for 4 years.

**Comments**

Many brain malformations are closely related to neuronal migration disorders [28]. Neuronal migration disorders are caused by the abnormal migration of neurons in the developing brain and nervous system. Neurons must migrate from the areas where they are born to the areas where they will settle into their proper neural circuits. Neuronal migration, which occurs as early as the second month of gestation, is controlled by a complex assortment of chemical guides and signals. When these signals are absent or incorrect, neurons do not end up where they belong. This can result in structurally abnormal or missing areas of the brain in the cerebral hemispheres, cerebellum, brainstem, or hippocampus, including schizencephaly, porencephaly, lissencephaly, agyria, macrogyria, pachygyria, microgyria, micropolygyria, neuronal heterotopias (including band heterotopia), agenesis of the corpus callosum, and agenesis of the cranial nerves.

In terms of encephalopathy or cerebral palsy, ‘timing of brain insult, antepartum, intrapartum or postpartum?’ is one of the serious

(Figure 13), monochorionic twins with one-twin death (Figure 14), high cardiac output due to cardiovascular anomaly (Figure 15) and other causes. Those cases with intrauterine severe encephalopathy resulted in severe cerebral palsy or antenatal/neonatal death.

**Intracranial vascularity with brain abnormality**

Medullary veins shown in Figure 7 develop according to brain tissue development. The case shown in Figure 16 was referred at 34 weeks of gestation due to fetal ventriculomegaly with intrauterine growth retardation. Transvaginal TUI with VCI (upper Figure 16) showed multiple intracerebral calcifications with delayed gyral formation and medullary venous maldevelopment with 19-20 week-appearance. Intrauterine cytomegalo viral infection was strongly suspected by neuroimaging despite maternal negative CMV IgM. Amniocentesis resulted in positive CMV-PCR and fetal CMV infection was confirmed. The estimated timing of fetal CMV infection may have been at 19-20 weeks or before, because of the appearance of medullary veins. The postnatal neurological prognosis has been poor. Figure 17 shows the case with abnormal brain morphology with normal medullary venous development. Medullary veins appear normal despite obvious

**Figure 9:** Shows non-uniformed migration disorder, which resulted in severe neurological prognosis after birth.

**Figure 10:** Figure showed non-uniformed migration disorder, which resulted in severe neurological.

**Figure 11:** Shows maldeveloped brain with migration disorder at 18 weeks of gestation.

**Figure 12:** Show cases of encephalopathy in various gestations.
controversial issues including medico-socio-legal-ethical problems [15]. Although brain insults may relate to antepartum events in a substantial number of term infants with hypoxic-ischemic encephalopathy, the timing of insult cannot always be certain.

The author first reported brain circulation demonstrated by Transvaginal 2D power Doppler in 1996 [29]. Thereafter, transvaginal 3D power Doppler assessment of fetal brain vascularity was successful [30]. Recently, owing to the advanced technology of bidirectional power Doppler, furthermore sophisticated 3D angiostructural image have been able to be demonstrated with vascular direction [27,30]. The author first demonstrated the fine medullary veins in 2008, running from the cortex towards the subependymal area by bidirectional power Doppler with 3D angiostructural imaging [31]. In the fetal brain, the medullary veins within the deeper cerebral white matter are more developed than the subcortical veins located within the subcortical white matter [32]. The mall development of medullary veins may indicate brain developmental abnormality, may predict subsequent hydrocephalus,
and may predict postnatal neurological deficit. The author has been investigating assessment of medullary veins in normal and abnormal brain structure and it is expected that the investigation will lead to one of solutions in the relations between the fetal brain development and postnatal neurological prognosis.

Conclusion

As described in this article, transvaginal high-resolution ultrasound technology has greatly contributed to fetal neuroscience. It is promising to clarify the developmental mechanism of CNS damages with advanced ultrasound diagnostic techniques in the near future. Neurological prognosis should be longitudinally and carefully evaluated according to precise diagnoses [33]. Considering future development in the field of fetal neurology, those precise morphological detection by transvaginal high-resolution neuroimaging should be combined with four-dimensional ultrasound research on fetal behavior [34-36] and molecular genetics which has recently been remarkably contributed to prenatal diagnosis as Sonogenetics [37,38]. Postnatal unexplained neurological deficits may strongly relate with intrauterine brain development therefore fetal neurology has great responsibility and an important role in perinatal medicine.

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