Khan, Javeria Shanawer, Sidra, Siddique, Memoona, Altaf, Aisha, Bacha, Raham, Farooq, Syed Muhammad Yousaf. (2020), Sonographic Comparison of Peak Systolic Velocities of Fetal Middle Cerebral Artery Among Normotensive and Hypertensive Mothers. In: Journal of Health and Medical Sciences, Vol.3, No.1, 26-32.

ISSN 2622-7258

DOI: 10.31014/aior.1994.03.01.93

The online version of this article can be found at: https://www.asianinstituteofresearch.org/
Sonographic Comparison of Peak Systolic Velocities of Fetal Middle Cerebral Artery Among Normotensive and Hypertensive Mothers

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Abstract
Background: Pregnancy-induced hypertension is most common hypertension in expectant women after 20 weeks growing the risk of subsequent hypertension, ischemic heart disease and cerebrovascular disease. Reduced renovation of the spiral artery is particularly well-thought-out as a cause failing. Maternal hypertension effects placental insufficiency, which can disturb the nutrition and the oxygen supply to the fetus. In hypertensive mothers, the potential of placental insufficiency is raised leading to intrauterine growth retardation but peak systolic velocity of middle cerebral artery has no reference value for our population. No matter, Ultrasonography can examine dynamically in real-time. However, the sonographic criteria in practice have no clear-cut numerical value to label IUGR. Objective: To compare peak systolic velocity of fetal middle cerebral artery among normotensive and hypertensive mothers. Setting: The data was collected from the following setups of Pakistan: Mushtaq Medical Imaging Sargodha, PAF Hospital Lahore. Materials and methods: The study was conducted after the approval of ethical committee of the University of Lahore. All patients were enrolled in this study after signing the informed consent form. Toshiba Xario with convex transducer of frequency 3-5 MHz was used. Fetal Middle cerebral artery indices were measured by Pulsed and Power Doppler through trans-abdominal scan. A cross-sectional analytical sampling technique was used and data were further analyzed with the help of the Statistical Package for the Social Sciences version twenty-four (SPSS 24, International Business Machines, Armonk, NY, the United States of America). Results: Among 137 patients, 60(43.8%) patients were hypertensive with mean PSV of the MCA, 35.63 cm/s and 77 (56.2%) normotensive patients with mean PSV of 36.64 cm/s. Conclusions: Peak systolic velocity of fetal middle cerebral artery in hypertensive mothers is less than that of Normotensive mothers.

Keywords: Ultrasound, Pregnancy, Hypertension, Placenta, Peak Systolic Velocity
**Introduction**

When the systolic blood pressure is higher than 140 mm of Hg and diastolic blood pressure gives measurements greater than 90 mm of Hg, this is known as hypertension (Kintiraki et al., 2015). Placenta is important for upholding of fetus during pregnancy and for helping usual growth and progress of fetus. Pregnancy-induced hypertension (PIH) or hypertension during pregnancy is the foremost reason of motherly transient and is a significant aspect in depletion of fetus (Salmani et al., 2014). Hypertension due to pregnancy enhances the threat of subsequent hypertension, ischemic heart disease and cerebrovascular disease (Nelson-Piercy, 2015). The risk of pregnancy-induced hypertension is highest in women younger than 20 years; first conception; those with twin or molar pregnancies; those with hypercholesterolemia, pre gestational diabetes, or obesity; and those with a previous family history of hypertension due to pregnancy (Houry and Salhi, 2014). Placental ischemia/hypoxia is supposed to stimulate the initiation of the motherly vascular endothelium, which causes increase in vascular sensitivity to hormone angiotensin II and decrease in the production of vasodilators, for example, nitric oxide and these endothelial oddities cause generalized vasoconstriction in whole of the body including the kidneys, which have a dire participation in the longstanding maintenance of arterial pressure (Palei et al., 2013). A placenta, but not the fetus, is requisite for the progress of hypertension during pregnancy. Certainly, the only active cure for hypertension during pregnancy is placental delivery. The old concept of the pathogenic processes convoluted in hypertension during pregnancy is that a placenta that is effected by ischemia yields soluble reasons that, when circulating through the mother’s blood circulation, are accountable for the medical indices of the disease (Chaiworapongsa et al., 2014). The source of hypertension through pregnancy is still not known. Reduced remodeling of the spiral artery has particularly been measured as a flaw producing hypertension during pregnancy. Remodeling is a multistep course in which the first decidua associated step should be begun implantation may be occurring (Steegers et al., 2010). Usually, instantly after embedding, ectodermal cells from the outermost epithelial layer of the blastocyst increase to miscellaneous types of trophoblastic cells. Union of cells produces the syncytium below the embryo that has been implanted, which shows that the initial intrusive trophoblastic cell type is migrating into the endometrium of the mother. When the space between the villi is formed, the inner layer of the trophoblasts originating from the trophectodermal layer forms primary villi by production and penetration through the primitive syncytium. All the way through pregnancy, these chief villi make subordinate and tertiary villi (Knöfler and Pollheimer, 2013).

It is testified that the possibilities for disruption of placenta comprise of late age of mother at pregnancy and increased number of children, smoking, multiple gestations, chronic hypertension, pregnancy induced hypertension and oligohydramnios (Hasegawa et al., 2014). Because fetal growth anomalies are often supposed as expressions of dysfunction of the placenta, it is remarkable to notice that placental abruption following pain, bleeding, and, possibly, interruption of fetal gas and nutrient exchange is more corporate in women with chronic hypertension (Seely and Ecker, 2014). Intrauterine growth restriction (IUGR) is frequently triggered by placental insufficiency, in reaction, fetus adjusts its own blood circulation to retain its nutrition and supply of oxygen to the brain (‘brain-sparing’) (Cohen et al., 2015). The Doppler detectable modifications in the fetal circulation related with IUGR and fetal hypoxia include raised resistance in the umbilical artery and peripheral vessel in connotation with reduced resistance in fetal cerebral vessels. This pattern of haemo dynamic adjustment is thought to mirror the “brain sparing” phenomenon in which hypoxic fetuses perfuse the brain, heart and adrenals (Kasegaonkar et al., 2016).

As mentioned before, maternal hypertension can effect placental insufficiency, which in turn can disturb the nutrition and supply of oxygen to the fetus. It is pondered that to stun this insufficiency, fetal vascular remodeling takes place. During this vascular remodeling, most of the blood flow circulates to the vital organs. As middle cerebral artery supplies brain (vital organ), the middle cerebral artery will be targeted in the research to observe the effect of maternal blood flow on fetal blood flow to brain. The supply of blood to the fetal brain can figure the growth of fetal brain. Middle cerebral artery peak systolic velocity may be a better analyst of perinatal impermanence in preterm IUGR than the PI. When the fetus is oxygen deficient, central reorganization of blood circulation outcomes in higher flow of blood to the fetal brain, heart, and adrenal glands, and a reduced flow of blood to the circulations to other tissues and organs of the fetus (Berkley et al., 2012).
Methodology

In this cross-sectional study, data was collected from expectant women of age 20-35 years visiting mentioned diagnostic setups of Lahore and Sargodha during 4 months, from 10th July to 10th October 2019. Sample size of 137 patients was calculated by using standardized formula. The study was started after the approval of ethical committee of the University of Lahore. All the patients were enrolled in this study after signing the informed consent form. Data was collected according to the variable of the age, hypertension, parity, trimesters and presence or absence of IUGR. Ultrasound machine Toshiba Xario with convex transducer of frequency range 3 to 5 MHz was used at Mushtaq Medical Imaging Sargodha and PAF Hospital Lahore, to calculate peak systolic velocities of fetal middle cerebral arteries trans abdominally. All the patients were examined according the American Institute of Ultrasound in Medicine (AIUM) gynecological ultrasound guidelines. Privacy of the patient was observed all the time and remained of top priority throughout the study.

Inclusion Criteria: Hypertensive and non-hypertensive mothers, pregnancy of 2nd and 3rd trimester, age group 20-35 years, nulliparous and multiparous mothers were included in the study.

Exclusion Criteria: Fetus with vital congenital abnormalities, intrauterine death of the babies and anomalies of the chromosomes were not included in the research study.

Data analysis: The collected data was processed thorough Statistical Package for the Social Sciences version 24 (SPSS 24, IBM, Armonk, NY, United States of America) for statistical calculations and analysis. Frequency distribution and descriptive statistics of all involved variables and the incidence were calculated.

Results

In this study total frequency of the patients were 137 pregnant females with Pregnancy of 2nd and 3rd trimester. Individuals were of age group 20-35 years. Mean age of the patients was 29.81±2.81 (25-35years). Table number 1 shows descriptive statistics of normal and hypertensive patients. In total of 137 pregnant patients. Figure 3 and Figure 4 show peak systolic velocities of Middle cerebral artery.

We diagnosed 60(43.8%) patients were hypertensive and 77(56.2%) were normotensive. Figure-1 and table no 1 show the frequency of hypertensive and normotensive patients. Out of 137 females 60(43.8%) patients were diagnosed with hypertension and 77(56.2%) patients were normal (normotensive). Demographic details of Peak systolic velocity of middle cerebral artery are shown in Figure-1 and Figure-2.

Table -1

| Blood pressure | Number of participant s | Mean   | Std. Deviation | Std. Error Mean |
|----------------|-------------------------|--------|----------------|-----------------|
| Normal         | 77                      | 36.6429| 12.27306       | 1.39865         |
| HTN            | 60                      | 35.6300| 17.53961       | 2.26435         |
Figure 1: Comparison of the PSV of the middle cerebral artery and number of participants in the Hypertensive and normal groups of pregnant female

Figure 2: Frequency and Peak Systolic Velocity of the fetal MC
Figure 3: Peak systolic velocity of the fetal Middle cerebral Artery is 57.3 cm/s at 30 weeks of pregnancy in normotensive mothers.

Figure 4: Peak systolic velocity of the fetal Middle cerebral Artery 28.9 cm/s at 27 weeks of pregnancy in hypertensive mothers.
Discussion

This study was performed to know the relationship between increase and decrease of the peak systolic velocities of fetal middle cerebral arteries among normotensive and hypertensive mothers. Among the patients of 137, we concluded that peak systolic velocity of middle cerebral artery is reduced in hypertensive mothers.

Acute severe hypertension is caused by generalized vasoconstriction in arteries and decreased blood flow in utero-placental and resulting in insufficient supply of oxygen at places of mother’s fetal exchange, forwarding the baby to lack of oxygen supply, hypoxia. The research studies submit that the assessment of blood flow in the middle cerebral artery of the fetus is one of finest signs of hazard of contrary perinatal consequence and analyzing the related diseases afterwards.

Michael Fischer, Helen and Sonia presented a research in 2015 for relating hypertension during pregnancy and abnormalities of the neonates. This cohort study involved more than eight hundred thousand pregnancies and was followed up after the birth of the babies. Generally, more than 19 thousand were diagnosed with chronic hypertension; of which, 42.0% were given the treatment of hypertension by drugs. When theses treated mothers and pregnancies were compared to the normal ones, pregnancies with hypertension treatments were at higher risks of abnormalities by birth. In their study of abnormalities of the babies' organs and body parts, the treated and untreated hypertension during pregnancy had a notable rise in the danger of heart abnormalities (OR 1.6, 95% and OR 1.5, 95% respectively). Abnormalities by birth in the neonates were seen in more than 29 thousand patients that were not hypertensive or given drugs for hypertension, four hundred and ninety-one patients that were exposed to antihypertensive treatment, and five hundred and eighty-one pregnancies without the treatment of pregnancy-induced hypertension. Major abnormalities in the babies were of heart, followed by the malformations of muscles, genitals, urinary and the digestive systems (Bateman et al., 2015). Line Sehested and Pedersen studied on the predictions and reasons for intrauterine growth retardation. 73 patients were kept under the study. 78% of the neonates were delivered by C-section. Risk factors of mothers included hypertension during pregnancy, cigarette smoking and abruption of the placenta. 22 babies were diagnosed by hypoglycemia. After follow up of 12 months, 90% of the babies gained good health and 7% had a neurologically poor result. None of the infants were demised (Sehested and Pedersen, 2014).

Another study was conducted by Allanson and Mari Muller in South Africa. They studied intrauterine death and related them with maternal diseases. A total of twenty-three thousand, five hundred and three births were there. There were six hundred and eighty-seven deaths right after delivery. The early neonatal demise was 50.7% more than deaths due to complications of mothers and then resulted in stillbirths (50.4%) and deaths right after birth was 25.8%. Mothers of babies who were died late after births were healthy. Stillbirths were noticed in the mothers who were hypertensive. Early neonatal deaths had healthy mothers. Most of the neonatal deaths occurred due to immaturity (48.7%) plus lack of oxygen supply (40.6%). One hundred and seventy three late deaths were due to intrauterine growth restriction and happened to have a birth weight less than the 10th centile for gestational age (Allanson et al., 2015).

Conclusions

Peak systolic velocity of fetal middle cerebral artery in hypertensive mothers is less than that of Normotensive mothers.

References

ALLANSON, E. R., MULLER, M. & PATTINSON, R. C. 2015. Causes of perinatal mortality and associated maternal complications in a South African province: challenges in predicting poor outcomes. *BMC pregnancy and childbirth*, 15, 37.

BATEMAN, B. T., HUYBRECHTS, K. F., FISCHER, M. A., SEELY, E. W., ECKER, J. L., OBERG, A. S., FRANKLIN, J. M., MOGUN, H. & HERNANDEZ-DIAZ, S. 2015. Chronic hypertension in pregnancy...
and the risk of congenital malformations: a cohort study. *American Journal of Obstetrics & Gynecology*, 212, 337-337.e14.

BERKLEY, E., CHAUHAN, S. P., ABUHAMAD, A. & COMMITTEE, S. F. M.-F. M. P. 2012. Doppler assessment of the fetus with intrauterine growth restriction. *American journal of obstetrics and gynecology*, 206, 300-308.

CHAIWORAPONGSA, T., CHAEMSAITHONG, P., YEO, L. & ROMERO, R. 2014. Pre-eclampsia part 1: current understanding of its pathophysiology. *Nature Reviews Nephrology*, 10, 466.

COHEN, E., BAERTS, W. & VAN BEL, F. 2015. Brain-sparing in intrauterine growth restriction: considerations for the neonatologist. *Neonatology*, 108, 269-276.

HASEGAWA, J., NAKAMURA, M., HAMADA, S., ICHIZUKA, K., MATSUOKA, R., SEKIZAWA, A. & OKAI, T. 2014. Capable of identifying risk factors for placental abruption. *The journal of maternal-fetal & neonatal medicine*, 27, 52-56.

HOURY, D. E. & SALHI, B. A. 2014. Acute complications of pregnancy. *Rosen's Emergency Medicine: Concepts and Clinical Practice*. 8th ed. Philadelphia, PA: Elsevier Saunders.

KASEGAONKAR, M., PATIL, A. & GOSAVI, A. 2016. COMPARATIVE STUDY OF FETOPLACENTAL BLOOD FLOW OF NORMAL AND PREGNANCY INDUCED HYPERTENSIVE MOTHERS BY DOPPLER METHOD, TO PREDICT NEONATAL OUTCOME AND MODE OF DELIVERY. *Int J Anat Res*, 4, 2868-72.

KINTIRAKI, E., PAPAKATSIKA, S., KOTRONIS, G., GOULIS, D. G. & KOTSIS, V. 2015. Pregnancy-induced hypertension. *Hormones (Athens)*, 14, 211-223.

KNÖFLER, M. & POLLHEIMER, J. 2013. Human placental trophoblast invasion and differentiation: a particular focus on Wnt signaling. *Frontiers in genetics*, 4, 190.

NELSON-PIERCY, C. 2015. *Handbook of obstetric medicine*, CRC press.

PALEI, A. C., SPRADLEY, F. T., WARRINGTON, J. P., GEORGE, E. M. & GRANGER, J. P. 2013. Pathophysiology of hypertension in pre-eclampsia: a lesson in integrative physiology. *Acta physiologica*, 208, 224-233.

SALMANI, D., PURUSHOTHAMAN, S., SOMASHEKARA, S. C., GNANAGURUDASAN, E., SUMANGALADEVI, K., HARIKISHAN, R. & VENKATESHWARAREDDY, M. 2014. Study of structural changes in placenta in pregnancy-induced hypertension. *Journal of natural science, biology, and medicine*, 5, 352.

SEELY, E. W. & ECKER, J. 2014. Chronic hypertension in pregnancy. *Circulation*, 129, 1254-1261.

SEHESTED, L. T. & PEDERSEN, P. 2014. Prognosis and risk factors for intrauterine growth retardation. *Dan Med J*, 61, A4826.

STEEGERS, E. A., VON DADELSZEN, P., DUVEKOT, J. J. & PIJNENBORG, R. 2010. Pre-eclampsia. *The Lancet*, 376, 631-644.