Expression of e-NOS in Human Placentas of Idiopathic Intra Uterine Growth Restriction at Term

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Abstract

Endothelial nitric oxide synthase (e-NOS) is a chemical agent that involves in placental angiogenesis. It is conveyed through embryogenesis. In the contemporary study, we intended to examine the character of placental angiogenesis in the progress of intrauterine growth restriction (IUGR) by means of matching the levels of expression of e-NOS in normal-term pregnancy and IUGR placentas. Angiogenesis is one dominant factor in normal embryogenesis and hence the wellbeing of the newborn. The expression of e-NOS was planned using the streptavidin-biotin-peroxidase technique in placental tissues identified as normal (n = 33) and IUGR (n = 33) cases from mothers looking apparently normal. All were chosen at term pregnancy and obtained between 2nd of February of 2018 and 31th of January 2019 at selected hospitals. Outcomes were appraised semi-quantitatively. The expression of e-NOS marker as an immunohistochemistry total score and staining percentage were significantly greater (p < 0.05) in epithelial surface, smooth muscle cells of fetal vessels and the connective tissue terminal villous core of the IUGR placentas when matched with placentas collected from normal pregnancies at term. Encountered placental histochemical changes regarding the expression of the e-NOS angiogenic factor for idiopathic IUGR newborns raised the suspicion that it was caused by pure placental factors and could represent further requirement for Nitric Oxide to dilate fetal vessels to optimize functional requirement during placental insufficiency. The noticed Increased expression of e-NOS may be the result of inadequate uteroplacental perfusion supporting the proposal that abnormal angiogenesis plays a role in the pathophysiology of IUGR.

Keywords: Idiopathic Iugr, E-Nos, Placenta, Terminal Villi, Immunohistochemistry.

Introduction

Intrauterine growth restriction/retardation (IUGR) is a complex placental vascular disorder resulting in limited weight at birth, preterm delivery, with highly expected perinatal morbidity and mortality ¹-⁴. IUGR could be triggered by different fetal, maternal, and placental influences ¹-³,⁵. Angiogenesis is regarded as a placental factor acting essentially in the development of IUGR ²-⁴,⁶,⁷. Angiogenesis involves the branching of new microvessels from present larger blood vessels. It is an imperative aspect in normal embryogenesis ⁸-¹⁰. Angiogenesis affects greatly the development of the villous vascular tree and the development of chorionic terminal villi. Placental vascular branching early in embryonic life and continues all over gestation ⁹,¹¹. Precise angiogenesis up regulators and down regulators adjust this process ⁴,⁸,⁹. Fetal growth restriction happens because of the failure of elongation, branching, or dilatation of the capillary circulation of terminal villi ⁴. e-NOS has been acknowledged as stimulator for angiogenesis ⁸. It is highly expressed throughout embryogenesis and fetal development ¹,⁶,¹⁰,¹². It synthesizes nitric oxide (NO) solely in the placenta ¹³,¹⁴, signifying that e-NOS is in charge for the increased NO in the placenta. e-NOS is chiefly accountable for the propagation of NO in the vascular endothelial layer of fetal vessels in the placenta ¹⁵,¹⁶,¹⁷. The e-NOS-derived NO is likewise a strong vasodilator compound in the perfused feto-placental vasculature which is empowered by smooth muscles ¹⁸, that is essential for the upkeep of decreased resistance at the fetoplacental interface ¹⁹. Initial revisions have demonstrated that drug conveyed e-NOS suppression results in signs resembling preeclampsia and growth restricted fetuses in rats ²⁰-²⁵.

Patients and Methods: This work was carried out through Department of Anatomy, Medical Faculty,
University of Al-Mustansiriyah, in assistance of Gynecology and Obstetrics department at Fatima Al-Zahraa Administrative Hospital, Al- Khadhraa Private Hospital and Al- Yarmook Teaching Hospital in Baghdad, Iraq. Permitted by the scientific committee of each of these medical institutes. A total of 66 mothers, along with their newborns and placentas were incorporated in this study. All were chosen at term pregnancy (of 38-40 weeks), they were non-smoker, non-diabetic, normotensive apparently healthy and normal women according to their history, clinical assessments, laboratory tests and ultrasound check. Fetal state as IUGR likewise was verified by Doppler ultrasonic exam without apparent fetal anomalies. Any mother suffered from difficult or delayed labor been ruled out from this work. A consent was gained verbally from each mother to be a component of this work. All fetal IUGR was considered to be idiopathic, given that there were no apparent maternal nor fetal basis. Placentas were assigned into two groups. The 1st embraced 33 placentas of newborns having normal average body weight (Control), whereas the 2nd group holds 33 placentas of newborns having idiopathic IUGR based on the basis of a valued fetal weight of less than the gestational tenth percentile, decreased amnion, and proved diminished end diastolic flow velocity of the umbilical artery by Doppler study confirmed by successive obstetric ultrasound examination. Every tissue block measured roughly 1×1×1 centimeter to be fixed to be stained by Immunohistochemistry e-NOS marker. Then, from every tissue block five sequential sections of about 4 μm thicknesses were obtained and were collected on positive charged slides. After de-waxing, the sections were rehydrated gradually. Immunohistochemical staining achieved by the avidin-biotin-peroxidase technique. 3% hydrogen peroxide blocking agent was used to block Endogenous peroxidase. Kept in oven for 15 minutes in sodium citrate buffer solution (pH=6.0) for antigen retrieval. Rabbit polyclonal antibodies reactive with e-NOS (Abnova) was added to the sections and incubated for one hour at room temperature. Sections were washed with phosphate-buffered saline with Tween, then a secondary antibody added for 20 minutes and after that with a biotin-streptavidin complex for half an hour at room temperature. 3,3-diaminobenzidine tetrahydrochloride (DAB) was used to visualize reaction. Finally, the sections were counterstained with hematoxylin and mounted. The intensity and localization of the staining reaction in Syncytiotrophoblasts (epithelial cells), vascular smooth muscle cells (vessels) and chorionic villous stromal connective tissue cells (C.T.) were calculated by two pathologists blinded to the drive of the study. Immunohistochemical reaction of e-NOS antibody scored semiquantitatively for intensity of staining: 0 / negative = no staining, 1+ = weak positive, 2+ = moderately positive; 3+ strongly positive. The total score of immunohistochemistry staining obtained from multiplying the staining intensity with staining percentage for each region in high power field. Statistical analysis achieved by using SPSS v24. Continuous variables were illustrated as mean ± SD and the categorical variables were demonstrated in percentages. Chi-square test was used to compare groups for immunohistochemical outcomes. P - values < 0.05 were considered statistically significant.

**Results**

e-NOS immunohistochemistry cytoplasmic staining was perceived from both control and idiopathic IUGR tissues (Table 1, and Figures 1,2 and 3). A highly significant increase in the expression of e-NOS in IUGR placentas seen in syncytiotrophoblasts (epithelial cells) when compared to control group with p value = 0.003, Also, increase in staining percentage in the IUGR group but no statistical significance with p value = 0.077 difference when compared to the control groups. Regardless, the total score of e-NOS marker in epithelium was significantly higher in IUGR group than that in control groups (p = 0.004). regarding the fetal vessels, the staining intensities of e-NOS marker were significantly higher in IUGR group with p value = 0.02 when compared to the control group. Staining percentage of e-NOS marker in vessels was also significantly higher in IUGR group than that in control groups with p value =0.016. The foregone total score of e-NOS marker in vessels was significantly higher in IUGR group than in control groups (p=0.007).
Figure (1): Difference in staining intensity of e-NOS between IUGR & Control tissues (* p value <0.05)

Figure (2): Difference in staining percentage of e-NOS between IUGR & Control tissues.

Table (1): Difference in staining total score of e-NOS between IUGR & Control tissues.

<table>
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<tr>
<th>localization</th>
<th>Mean</th>
<th>Std. Deviation</th>
<th>p Value</th>
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<tr>
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<td>Control</td>
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<td>1.62</td>
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<tr>
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<tr>
<td>IUGR</td>
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Figure (3): e-NOS staining reaction expressed in terminal villi of placenta, A; showing weak e-NOS staining in control group, B; showing e-NOS stronger expression in idiopathic IUGR group (DAB, A; B; × 400).
Discussion

Many previous studies had documented that a pure placental factor was the solitary cause in some IUGR cases. IUGR incidence was noticed 3% regarding developed countries, though, it scopes 15-20% in developing countries, it partakes in more than 26% stillbirths and congenital anomaly. Not to forget it yields many lasting health issues for grownups. Therefore, the precise diagnosis, and proper management of IUGR-complicated pregnancies are important. Regarding e-NOS expression in the epithelial cells, our results differ from the study of Myatt et al (1997) as they had found no differences in the intensity or type of staining. Regarding the fetal vessels, our results agree with Myatt et al (1997) and Rutherford and colleagues (1995). However, other authors have found decreased levels of e-NOS in vessels in these condition like Giannubilo et al. (2008) and Noris et al. (2004). We hypothesize that the increased e-NOS expression in their vessels is not a disease-specific feature, but perhaps an adaptive response to increased resistance and poor perfusion. It seemed to be no previous works measured the expression of e-NOS in the villus core of terminal chorionic villi of term placenta neither in normal nor in pathological cases like IUGR. The significant elevation in expression of this enzyme, e-NOS, in this region particularly could be explained by its rapid diffusion from endothelial cells to the core, it had been found that it down regulates fibroblast production to decrease villus fibrosis decreasing the fibroblastic activity which in turn reducing the density of the villus core to enhance feto-maternal exchange. It is possible that increased expression of e-NOS seen in our results in idiopathic IUGR placentas at term in all mentioned regions impinge upon regulation of placental vascular reactivity by the smooth muscle of fetal vessels and hence blood flow regulation. In support of this, it had been shown that concentrations of nitrate, a breakdown product of NO, were significantly higher at delivery in umbilical venous blood from idiopathic IUGR pregnancies compared with controls. The observed increased expression of e-NOS that we saw in placentas of Idiopathic IUGR could encourage pathological angiogenesis.

Conclusion

Toward elucidate the pathogenesis of idiopathic IUGR and the associated placental vascular insufficiency, it is needed to clarify the regulatory mechanism of placental vascular development. The noted increased staining expression of e-NOS designates pathological angiogenic action, due to lacking proper uteroplacental perfusion, resulting in the formation of idiopathic IUGR.

Ethical Clearance: was approved and granted from Al-Mustansirya University, Al-Rusafa and Al-Karkh health Directorate application forms as a requirement to obtain PhD degree in Anatomy, Histology and Embryology. Research projects are reviewed and conducted ethically. This research is not published yet.

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Conflict of Interest: Nil

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