HISTOMORPHOLOGICAL CORRELATION OF PIH PLACENTA AND LOW DOSE ASPIRIN TREATED PLACENTA
A.L. Santhi¹, Banumathi²

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ABSTRACT: OBJECTIVES: To correlate the histomorphological changes in PIH Placenta and low dose Aspirin received placenta. METHODS: This prospective study was conducted in 50 patients. They were divided into two groups. 25 placentas were collected from PIH patients and 25 placentas from patients who were treated with prophylactic low dose aspirin [75 mg] from 28th week of gestation. Results; Improved vascular bloodflow with reduced vessel wall changes and infarctswere noticed in the placenta of aspirin group. So there was a good placental weight and baby weight leading to a good neonatal outcome.

KEYWORDS: Placenta, Aspirin, Pregnancy induced Hypertension. [PIH]

INTRODUCTION: Pregnancy induced hypertension [PIH] accounts for 6-8% of all pregnancies. PIH remains a major cause of morbidity and mortality for both mother and baby. Mother may develop DIC, HELLP syndrome, renal or hepatic failure. Baby may suffer from IUGR, IUD and prematurity.¹

Although the cause of PIH remains obscure, it is primarily a placental disorder. During implantation, deficient trophoblastic invasion of maternal spiral arteries leads to underperfusion of uteroplacental circulation and placental ischemia². So in PIH, placenta affected by ischemia shows variety of changes both morphologically and histologically which can be appreciated by light microscopy and special stains.

Women with PIH have deficient intravascular production of prostacyclin, a vasodilator and an excessive production of thromboxane, a vasoconstrictor and platelet agonist. Low dose Aspirin therapy in these patients is beneficial by potent antiplatelet effect. It inhibits platelet production of thromboxane. Thus the loss of maternal and fetal lives to PIH can most often be prevented³.

So detailed study of placenta was undertaken in PIH patients by using special stains. The degree of severity of PIH was correlated clinicopathologically. These histopathological changes in placenta were compared with aspirin treated groups to know the effects of treatment.

AIM AND OBJECTIVES: The principal aim is to study the various histopathological changes that occur in the placenta of PIH patients and compare them with the placenta of PIH patients who received low dose Aspirin. Battery of special stains was used to delineate the hypoxic changes and blood vessel changes in placenta of PIH cases.

MATERIAL AND METHODS: A prospective study of placenta of PIH patients [Mild and Severe] were compared with the placenta of PIH patients on low dose Aspirin 75 mg/day from 28th week of gestation. Placentas were collected in the third trimester after delivery. The age group ranges from 15-32 years, maximum number of cases was between 21- 25 years of age.
Thorough examination of the placenta was done. Weight of the placenta after trimming the membranes, maximum diameter, thickness at the centre, shape, accessory lobe, infarcts [size, location, number], calcification, cyst, tumor and intervillous fibrin. Placentas from normotensive individuals were also collected and used as control. Special stains were also used in control placentas for comparison.

RESULTS:

**Weight of the Placenta:** The mean average weight of the placenta in PIH group was 360 gms and in Aspirin group was 485 gms.

| Weight of the placenta | In PIH Patients group | In Aspirin group |
|------------------------|-----------------------|-----------------|
| 150–250 Gms            | 24%                   | Nil             |
| 250–350 Gms            | 12%                   | 4%              |
| 350–450 Gms            | 44%                   | 28%             |
| 450–550 Gms            | 16%                   | 64%             |
| > 550 Gms              | 4% [Diabetes associated] | 4%              |

**HISTOPATHOLOGICAL FINDINGS OF PLACENTA IN PIH:**

1. **CYTOTROPHOBLASTIC PROLIFERATION:** Minimal proliferation was observed in 72% and severe in 28% of PIH cases. Whenever placenta suffers an ischemic damage as in PIH, germinal zone will be reactivated and cytotrophoblasts will proliferate in an attempt to repair and replace the injured syncytial tissue.

2. **EXCESSIVE NUMBER OF SYNCYTIAL KNOTS:** Minimal increase in syncytial knots were observed in 64% cases and marked increase in 36% of cases.

3. **THICKENING OF BASEMENT MEMBRANE:** Noticed in 50% of cases of PIH which are better appreciated by PAS [Periodic Acid Schiff] staining. Basement membrane protein is secreted by proliferated cytotrophoblasts.

![Fig. 1 & 2: EXCESSIVE SYNCYTIAL KNOTS LOW POWER, HIGH POWER](image-url)
4. ABNORMALITIES OF BLOOD VESSELS:

A. MEDIAL HYPERTROPHY: Concentric onion skin appearance sometimes even totally occluding the lumen. This was best brought out by special stain - VERHOFFE'S elastic stain in 68% of cases.

B. FIBRINOID NECROSIS: Seen in 52% of cases. PTAH [Phosphotungstic acid Haematoxylin] was used to demonstrate this necrosis.
5. **PLACENTAL INFARCTS**: Infarction is most common in PIH [34-60%] and is significant when they are more than 3cm. Old infarcts are firm & white. In PIH, multiple diffuse infarcts were present. Fresh infarcts are red, congested and soft. In our study, fresh infarcts were seen in 12% and old infarcts in 68% of PIH cases. Minimal infarcted areas were observed in aspirin group. Infarcts were associated with significant perinatal morbidity and mortality.

### HISTOPATHOLOGICAL CHANGES

|                          | MILD | SEVERE |
|--------------------------|------|--------|
| CYTOTROPHOBlastic PROLIFERATION | 72%  | 28%    |
| SYNCYTIAL KNOTS          | 64%  | 36%    |
| BASEMENT MEMBRANE THICKENING | 30%  | 26%    |
| MEDIAL HYPERTROPHY       | 38%  | 30%    |
| FIBRINOid NECROSIS       | 28%  | 24%    |
| OLD INFARCTS             | 36%  | 44%    |

### VILLOUS OBSERVATION IN PIH CASES

1. Increased vascularity of villi.
2. Cytotrophoblastic proliferation: Absent in 56% cases and minimal in 32% cases.
3. Syncytial knots: Absent in 68% of cases and minimal in 32% of cases.
4. Fibrinoid Necrosis: 84% of placenta did not show fibrinoid necrosis. 16% showed minimal fibrinoid changes.
5. Medial Hypertrophy: No medial hypertrophy in 76% of cases and minimal in 24% of cases. But there was no occlusion of lumen in these cases. In general, atherosis of blood vessel has reduced.
6. Old Infarcts: Observed only in 8% of Aspirin group.
7. Basement Membrane Thickening: No thickening seen in 72% of cases and minimal in 28% of cases.
**HISTOPATHOLOGICAL CHANGES IN ASPIRIN GROUP**

|                        | ABSENT | MINIMAL |
|------------------------|--------|---------|
| CYTOTROPHOBLASTIC PROLIFERATION | 56%    | 44%     |
| SYNCYTIOTrophic Knots  | 68%    | 32%     |
| BASEMENT MEMBRANE THICKENING | 72%    | 28%     |
| FIBRINOID NECROSIS     | 84%    | 16%     |
| MEDIAL HYPERTROPHY    | 76%    | 24%     |
| OLD INFARCTS          | 92%    | 08%     |

**FETAL OUTCOME:** In 25 cases of PIH, 15 babies were alive, 4 cases of intrauterine Death and 6 cases of deeply asphyxiated babies were noticed. Two asphyxiated were revived and 4 babies were lost. Placenta of intrauterine death fetuses were below average weight. Placenta had retroplacental hematomas and in one case, it occupied almost the entire half of placenta. Histology of these placentas showed massive old infarcts, medial hypertrophy of blood vessels, basement membrane thickening and fibrinoid necrosis.

**BIRTH WEIGHT IN PIH GROUP**

| BIRTH WEIGHT | IN PIH GROUP | PIH PATIENTS ON ASPIRIN |
|--------------|--------------|-------------------------|
| 1.5 – 2 kg   | 24%          | 4%                      |
| 2 – 2.5 kg   | 56%          | 12%                     |
| 2.5 – 3 kg   | 16%          | 68%                     |
| > 3 kg       | 4%           | 16%                     |

**DISCUSSION:** In PIH, major pathological changes are seen in placentas. Villi in the placenta are subjected to reduced maternal uteroplacental blood flow and so they show a characteristic pattern of abnormalities due to hypoxia. There is undue prominence and increase of villous cytотrophобlasts along with thickening of basement membrane. The cytотrophобlastic cells are the stem cells where DNA synthesis and mitotic activity occurs. Syncytотrophoblast being formed by breaking down of limiting membrane of cytотrophобlasts and it is a post mitotically differentiated tissue. Therefore cytотrophобlast is considered to form a germinative zone, though in late stages of pregnancy it is largely quiescent5.
If placenta suffers an ischemic damage as in PIH, the germinative zone will be reactivated and cytotrophoblastic cells proliferate in an attempt to repair and replace the injured syncytial tissue. Basement membrane thickening seen in placenta subjected to ischemia is an incidental byproduct of cytotrophoblastic hyperplasia. Basement protein is almost secreted by cytotrophoblastic cells. The essential response of placenta to ischemia is a reparative one.

PATHOGENESIS OF PIH: Placental ischemia secondary to defective placentation, a prerequisite for the development of preeclampsia has a multifactorial origin. In a normal pregnancy, the trophoblastic invasion is associated with striking changes in the arteries supplying the intervillous space. The endothelium is replaced by trophoblast. By contrast, these changes develop defectively in preeclamptic woman and are also limited to decidual portion of spiral arterioles. Blood vessels are sometime occluded by fibrinoid material and surrounded by foam cells. These features are known as acute atherosis. These changes have been well brought about by special stain, PTAH, which stains fibrinoid material.

One consequence of abnormal invasion of the spiral arteries is deficient uteroplacental circulation. So the villous growth in low oxygen environment as in PIH exhibits the following features:
1. Cytotrophoblastic proliferation
2. Increase in number of syncytial knots
3. Basement membrane Thickening.

ENDOTHELIAL CELL DYSFUNCTION – FINAL COMMON PATHWAY: Many observations point to a central role of endothelial cells in the pathogenesis of PIH. Endothelial injury leads to an increase of potent vasoactive substance such as endothelin, fibronectin and thromboxane. Endothelin is a potent vasoconstrictor and three distinct endothelins exist 1, 2 and 3. Patients with preeclampsia have higher levels of endothelin. A marker of endothelial dysfunction is Fibronectin, a high molecular weight protein. Thromboxane causes vasoconstriction, stimulates platelet aggregation and uterine contractility. Prostacyclin produces opposite physiological effects. Maternal plasma prostacyclin is decreased in both mild and severe cases of preeclampsia whereas thromboxane is increased. Thromboxane is synthesized as a result of endothelial injury.

ROLE OF LOWDOSE ASPIRIN IN PROPHYLAXIS AND EARLY TREATMENT OF PIH: It appears that two vasoconstrictor mechanisms may be operative in woman with PIH i.e. especially in PIH, arachidonic acid is converted via cyclooxygenase pathway into thromboxane A2 with an accompanying reduction in prostacyclin [PGI2] and prostaglandin E2. This pathway is responsive to low dose aspirin therapy. Arachidonic acid may be converted into PG12, PGE2 and TXA2. Low dose aspirin thereby block TXA2 production more than the production of PG12 and PGE2.

Since aspirin has been shown to be beneficial in preventing eclampsia, it is important to be able to predict which patients are at risk for the disease. The most commonly used non-laboratory method is blood pressure measurement. Any rise in blood pressure after 20th week of gestation should raise the concern for preeclampsia. Another means of predicting the risk is based on mean arterial pressure in second trimester. Another routine diagnostic test is roll over test. Pregnant woman whose diastolic pressure increases when they change from left lateral to supine position are
at high risk for preeclampsia. Laboratory methods of prediction include elevated serum uric acid levels, fibronectin levels and decreased urine calcium excretion³.

Therefore low dose aspirin therapy has been proved to be useful in improving the placental blood flow and thereby fetal outcome and baby weight.

CONCLUSION: Low dose Aspirin therapy in PIH patients showed the following findings;

1. Improved vascular blood flow.
2. Reduced vessel wall changes.
3. Reduces the incidence of Placental infarcts.
4. Good Placental weight.
5. Improved Baby weight.
6. Good Neonatal outcome.

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AUTHORS:

1. A.L. Santhi
2. Banumathi

PARTICULARS OF CONTRIBUTORS:

1. Professor and HOD, Department of Pathology, Thanjavur Medical College, Tamilnadu.
2. Professor, Department of Pathology, Thanjavur Medical College, Tamilnadu.

NAME ADDRESS EMAIL ID OF THE CORRESPONDING AUTHOR:

Dr. A.L. Santhi,
10, Philomina Nagar,
Don Bosco School Road,
Thanjavur.
Email-santhiminu@gmail.com

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