Four Years of Foetal Autopsies in A Peripheral Teaching Hospital: A Retrospective Analysis

Priya Subashchandrabose, Jayaganesh Parthasarathy* and Chitra Srinivasan
Department of Pathology, Saveetha Medical College, Chennai, India

ABSTRACT

Background: The aim of our study was to assess the prevalent causes of foetal demise in our region, so as to facilitate necessary prenatal measures to reduce the Intra uterine deaths.

Methods: A retrospective study on 28 foetuses, received in the Department of Pathology, either because of intra uterine death or those which were medically terminated for various abnormal prenatal findings, for a period of 4 years from November 2011 to November 2015. Autopsy report findings including full external examination, anthropometric measurements, gross, microscopic details of different organs including placenta and ultrasound reports were analysed.

Results: Total number of foetal autopsies was twenty eight. Most prevalent cause of death was congenital anomalies (39.3%) followed by placental insufficiency (17.8%) and umbilical cord anomaly (10.7%). The most frequent multiple system anomalies were central nervous system defect and musculoskeletal system malformations. Foetal autopsy provided additional findings in 8 cases (28.57%).

Conclusion: Foetal autopsy is a vital procedure in confirming the cause of foetal death. It also provides additional findings not identified through prenatal ultrasound. It provides the treating clinician and parents with very crucial information on implications for future pregnancies.

Keywords: Congenital Anomalies, Foetal Autopsy, Intra Uterine Death, Placental Abnormalities, Prenatal Diagnosis.

Introduction

Foetal autopsy is very valuable because it gives the parents, the cause for the death of their baby. Foetal death is classified as early (less than 22 weeks of gestational age), intermediate (22 – 27 weeks of gestational age) and late (greater than 28 weeks of gestational age). The early deaths are referred to as abortions and the late and intermediate deaths are termed together as Intra uterine deaths (IUD) or still births. Congenital anomalies are an important cause of mortality in developing countries. Improvement in prenatal care can reduce the incidence to substantial amount. Genetic disorders or obstetric conditions that may cause recurrent threat to future pregnancies can be recognized. IUD contributes substantially to perinatal mortality thus proving to be a good indicator of quality of antenatal care provided and also to assess the amount of pregnancy loss. The main objectives of the foetal autopsy include identification the gestational age and congenital abnormalities, document growth and development, assess and establish the cause of death and probable risk of recurrence. Our study was hence performed to identify the prevalent causes of IUD on foetal autopsies in our region, categorise them based on causes and thus help in taking essential steps to reduce the IUD by improvising the prenatal care.

Materials and Methods

After obtaining the institutional ethics committee, this retrospective study on 28 foetal autopsies performed in the department of Pathology, Saveetha Medical College, over a period of three years from November 2012 to November 2015 was performed. The cases, for which all the autopsy data were available, were included in the study. Macerated and autolyzed foetuses were excluded from the study. All the details in autopsy reports including gestational age, sex of the foetus, external examination findings, anthropometric measurements, internal examination findings, findings of the examination of head & neck, brain and spinal cord, results of examination of head & neck, brain and spinal cord, results of examination of placenta and umbilical cord were retrieved from the departmental achieves and analysed.

Cunningham & Hollier classification was used to divide cases into foetal, placental or maternal categories as: Foetal (25-40%) - chromosomal anomalies, non-chromosomal birth defects, non-immune hydrops, infections; Placental (25-35%) - abruption, foetal-maternal haemorrhage, cord accident, placental insufficiency, intra partum asphyxia, placenta previa, twin to twin transfusion, chorioamnionitis; Maternal (5-10%) - diabetes, hypertensive disorders, trauma, abnormal labour, sepsis, uterine rupture, post-term pregnancy, drugs, antiphospholipid antibodies, unexplained.
Result

Over a period of four years from 2011 to 2015, twenty eight cases of foetal autopsies were performed in the department of pathology. Mean maternal age was 25.3 years, youngest was 20 years and oldest was 36 years. Mode of delivery in almost all cases (27 cases) was vaginal and only one case was lower segment caesarean section (LSCS). Among 28 cases, eleven were therapeutic terminations, sixteen were intra-uterine deaths and one was perinatal death. We received twenty male foetuses and eight female foetuses and hence the male: female ratio was 2.5:1 in our series.

Most of the foetal deaths in our study, fourteen cases (50%) were early foetal deaths, twelve cases (42.9%) were intermediate and two were late foetal deaths (7.1%). The number of cases in various categories and causes of foetal death have been shown in table 1. The most prevalent cause of death was congenital anomalies (11 cases, 39.3%) followed by placental insufficiency (5 cases, 17.8%), Umbilical cord anomaly (3 cases, 10.7%). Foetal causes were implicated in highest number of cases (46.5%).

Congenital anomalies were common between gestational ages of 18-26 weeks. Birth weights ranging between 250-1000 grams (67.8%) were showing most congenital anomalies. The most common congenital anomaly detected was central nervous system and musculoskeletal system. The various anomalies detected according to the different systems are shown in table 2.

We observed seven interesting cases including: Meckel Gruber syndrome, Arnold chiari malformation type 2, Double outlet right ventricle with ventricular septal defect, Twin – twin transfusion syndrome, Cystic hygroma, Gastrochisis and a case of floppy infant.

Placental insufficiency was the most common placental abnormality observed and umbilical cord anomalies with single artery were seen in three cases.

Anemia was the commonest maternal cause associated with congenital anomalies, followed by preeclampsia, polyhydramnios, oligohydromnios and one case was positive for CMV. In one case, the mother was a known case of Hemophilia with history of losing the first baby and this foetus was the second conception. However, chorionic villus biopsy done at 16 weeks showed the foetus is not a case of haemophilia. The two cases with chorioamnionitis gave history of premature rupture of membranes.

Routine antenatal checkup included ultrasonography, and the reports were retrieved from the archives and analysed. There was a good correlation between the ultrasound findings and autopsy findings.

The different morphologic findings including the various gross abnormalities and the microscopic findings in the placenta, cord and foetuses that were seen at autopsy are shown in the table 3.

### Table 1: Distribution of cases in various categories and causes of foetal death.

| Categories | Causes                  | No. of cases | Percentage (%) |
|------------|-------------------------|--------------|----------------|
| Foetal (46.5%) | Congenital Anomalies  | 11           | 39.3           |
|            | Infection               | 1            | 3.6            |
|            | Non – immune hydrops    | 1            | 3.6            |
| Placental (39.2%) | Twin to twin transfusion | 1            | 3.6            |
|            | Placental Insufficiency | 5            | 17.8           |
|            | Chorioamnionitis        | 2            | 7.1            |
|            | Umbilical cord anomaly  | 3            | 10.7           |
| Maternal (14.3%) | Diabetes               | 1            | 3.6            |
|            | Pre-eclampsia           | 1            | 7.1            |
|            | Pregnancy induced       | 1            | 1.9            |
|            | Hypertension            | 1            |                |
|            | Haemophilia             | 1            | 3.6            |
| Total      |                         | 28           | 100            |

### Table 2: System wise distribution of congenital anomalies.

| Defects                          | No of cases (total 11 cases) |
|----------------------------------|------------------------------|
| Central nervous defects          |                              |
| Spina bifida with Meningomyelocele | 1                            |
| Spina bifida with Rachischisis   | 1                            |
| Arnold chiari malformation       | 1                            |
## Foetal Autopsies

### Defects and No of cases

| Defects                        | No of cases (total 11 cases) |
|--------------------------------|-------------------------------|
| **Musculoskeletal defects**    |                               |
| Gastrochisis                   | 1                             |
| Unspecified                    | 1                             |
| **Urogenital defects**         |                               |
| Bilateral dysplastic kidneys   | 1                             |
| **Cardiovascular system defect** |                             |
| Double outlet right ventricle with ventricular septal defect | 1 |
| **Lymphatic system defects**   |                               |
| Cystic hygroma                 | 1                             |
| Multiple system defects        |                               |

### Table 3: Morphological features in foetus, placenta and umbilical cord on autopsy.

| S.No | Cause of foetal death                        | Morphologic findings in foetuses in autopsy | Morphologic findings in cord and placenta in autopsy |
|------|----------------------------------------------|--------------------------------------------|-----------------------------------------------------|
| 1.   | Congenital anomalies (11 cases)              | Bilateral dysplastic kidneys               | Gross: Cord, placenta and membranes were normal      |
|      |                                              | Gross: External and cut surface showed multiple thin walled cysts involving the whole parenchyma (Figure 1A) | Micros: The Placenta, cord and membranes showed no significant pathology. |
|      |                                              | Micros: Immature tubules lined by a single layer of cuboidal epithelium and surrounded by immature mesenchyme. Many cystic spaces lined by cuboidal cells and few scattered glomeruli were seen. (Figure 1B) |                                                     |
|      | Diaphragmatic hernia with spina bifida       | Gross: Spina bifida with exposure of spinal cord starting from mid thoracic to lumbar region. Left lobe liver, stomach, proximal intestine and spleen were seen to herniate into the left thorax with displacement of thoracic structures in to right side. Left lung is hypoplastic (Figure 2) | Gross: Cord, placenta and membranes were normal. Micros: Placenta showed features of mild insufficiency with focal areas of calcification |
|      |                                              | Micros: All organs were within normal limits. |                                                     |
|      |                                              | Gross: Cord, placenta and membranes were normal. Micros: Placenta showed features of mild insufficiency with focal areas of chorangiosis |                                                     |
|      | Meckel Gruber syndrome                       | Gross: Occipital meningocele (Figure 3A), Bilateral dysplastic kidneys, Marked oligohydrominios, Polydactyly in all four limbs (Figure 3B, 3C) and Hypoplastic lungs. | Gross: Placenta showed marginal insertion of cord. Micros: Areas of infarction and focal areas of chorangiosis |
|      |                                              | Micros: Sections from lungs were normal, sections from kidney showed primitive glomeruli and primitive tubules with dilated tubules lined by flattened cells and surrounded by immature mesenchyme. Many cystic spaces lined by cuboidal cells were noted. |                                                     |
|      | Anencephaly with facial abnormalities and polydactyly | Gross: Absence of cranium with exposure of brain substance. Occiput was bifid with a central defect. Low set ears, cleft lip and cleft palate were noted. All four limbs showed polydactyly. | Gross: Cord, placenta and membranes were normal. Micros: Placenta showed features of mild insufficiency with focal areas of calcification |
|      |                                              | Micros: All organs were within normal limits. |                                                     |
|      | Spina bifida with Rachischisis               | Gross: Spine showed spina bifida and rachischisis from D10 level to S1 level with absence of posterior spinal elements (Figure 4A). | Gross: Cord, placenta and membranes were normal. Micros: Placenta showed focal areas of necrosis and calcification (Figure 4B). |
|      |                                              | Micros: All organs were within normal limits. |                                                     |
| S.No | Cause of foetal death | Morphologic findings in foetuses in autopsy | Morphologic findings in cord and placenta in autopsy |
|------|----------------------|-------------------------------------------|-----------------------------------------------------|
|      |                      |                                            |                                                     |
|      | **Gastrochisis**     | Gross: Anterior abdominal wall showed a midline defect with intestines protruding through the opening (Figure 5). **Micros:** All organs were within normal limits. | Gross: Placenta was bigger in size and weight for the gestational age (Figure 5). **Micros:** Placenta showed features of mild insufficiency. |
|      |                      |                                            |                                                     |
|      | **Arnold chiari malformation - Type 2** | Gross: Cerebellar tonsils were seen herniating through the foramen magnum (Figure 6A). Spina bifida was seen in the Lumbosacral region with meningo myelocoele. **Micros:** All organs were within normal limits. | Gross: Cord, placenta and membranes were normal **Micros:** The Placenta, cord and membranes showed no significant pathology. |
|      |                      |                                            |                                                     |
|      | **Double outlet right ventricle with ventricular septal defect** | Gross: Heart – right ventricle showed both opening of aorta, pulmonary artery and a ventricular septal defect was noted.(Figure 7) **Micros:** All organs were within normal limits. | Gross: Cord showed Single umbilical artery **Micro:** Cord showed Single umbilical artery |
|      |                      |                                            |                                                     |
|      | **Cystic hygroma with Hydrops feta** | Gross: Nuchal odema was seen. Whole foetus was oedematous (Hydrops feta). A large sac measuring 8X6cms was seen attached to the posterior aspect of neck and filled with clear fluid. The lining of the cyst was smooth and was filled with clear colourless fluid. (Figure 8) **Micros:** All organs were within normal limits. | Gross: Cord, placenta and membranes were normal **Micros:** The Placenta, cord and membranes showed no significant pathology. |
|      |                      |                                            |                                                     |
|      | **Spina bifida with Meningomyelocele** | Gross: Spina bifida was seen in the Lumbosacral region with meningo myelocoele (Figure 9A). **Micros:** All organs were within normal limits. | Gross: Cord, placenta and membranes were normal **Micros:** Placenta showing features of vascular insufficiency mainly chorangiosis (Figure 9B, 9C). |
|      |                      |                                            |                                                     |
|      | **Unexplained –**    | Gross: The foetus was floppy. No other abnormalities were made out. **Micros:** Sections from the skeletal muscle showed features of pan fascicular atrophy. However tissue sent for electron microscopy to a referral centre failed to any specific findings. | Gross: Cord, placenta and membranes were normal **Micros:** The Placenta, cord and membranes showed no significant pathology. |
|      |                      |                                            |                                                     |
|      | **Infection**        | Gross: All the organs were unremarkable **Micros:** Parenchymal organs showed heavy interstitial mononuclear lymphoplasmacytic infiltrates. | Gross: Cord, placenta and membranes were normal **Micros:** Cord showed mild focal mononuclear inflammation. Placenta showed intervillious lymphoplasmacytic infiltrates. |
|      |                      |                                            |                                                     |
|      | **Non – immune hydrops** | Gross: Nuchal odema was seen. Whole foetus was oedematous (Hydrops feta). **Micros:** All organs were within normal limits. | Gross: Cord, placenta and membranes were normal **Micros:** The Placenta, cord and membranes showed no significant pathology. |
|      |                      |                                            |                                                     |
|      | **Twin to twin transfusion syndrome** | Gross: One foetus was congested and plethoric and the other was pale and anaemic (Figure 10 ) **Micros:** All organs were within normal limits. | Gross: Cord, placenta and membranes were normal **Micros:** Placenta was monoamnionic and monochorionic with features of mild insufficiency. |
| S.No | Cause of foetal death                  | Morphologic findings in foetuses in autopsy | Morphologic findings in cord and placenta in autopsy |
|------|--------------------------------------|-------------------------------------------|-----------------------------------------------------|
| 5.   | Placental Insufficiency (5 cases)     | Gross: Features of Intra Uterine Growth Retardation<br>Micros: All organs were within normal limits. | Gross: Cord, placenta and membranes were normal<br>Micros: Increased syncytial knots, increase in terminal mature villi, decreased vasculature, areas of infarction, necrosis and fibrosis, foci of dystrophic calcification. (Figure 11) |
| 6.   | Chorioamnionitis (2 cases)           | Gross: All organs were normal<br>Micros: All parenchymal organs were normal except for a mild mononuclear inflammatory infiltrates in interstitial spaces of the lung. | Gross: Placenta showed focal areas of infarction and fibrinous deposits.<br>Micros: Presence of areas of fibrinoid necrosis and neutrophilic infiltrates in the amnion, chorionic plate and decidua (Figure 12). Some of the villi also showed neutrophilic infiltrates. |
| 7.   | Umbilical cord anomaly (3 cases)     | Gross: No abnormality detected<br>Micros: All the organs were within normal limits. | Gross: Two cases had single umbilical artery and more case had true knots in the cord. Micros: Single umbilical artery was noted. |
| 8.   | Maternal Diabetes mellitus           | Gross: No abnormality detected<br>Micros: All the organs were within normal limits. | Gross: Cord, placenta and membranes were normal<br>Micros: Placenta showing features of vascular insufficiency mainly chorangiosis. |
| 9.   | Maternal Pre-eclampsia (1 case)<br>Pregnancy induced hypertension (1 case) | Gross: Foetus showed features if IUGR.<br>Micros: All organs were unremarkable. | Gross: Cord, placenta and membranes were normal<br>Micros: Placenta showing features of vascular insufficiency mainly infarct, calcification and chorangiosis |
| 10.  | Maternal Haemophilia                 | Gross: No abnormality detected<br>Micros: All the organs were within normal limits. | Gross: Cord, placenta and membranes were normal<br>Micros: Placenta showing features of insufficiency including, increased syncytial knots, increase in terminal mature villi, decreased vasculature, areas of infarction, necrosis and fibrosis, foci of dystrophic calcification. |

Fig. 1: Bilateral dysplastic kidneys. A – Gross image showing cut surface with multiple thin walled cysts, involving the whole parenchyma. B – microscopic picture showing renal tissue with immature tubules surrounded by immature mesenchyme and cystic spaces with few scattered glomeruli (H&E 40X). C – microscopic picture showing dilated tubules lined by single layer of cuboidal epithelium with surrounding by immature mesenchyme (H&E 100X).
Fig. 2: Meckel Gruber syndrome. A- Gross image showing meningocele. B- Gross image showing polydactyly (Six digits) in upper limbs. C- Gross image showing polydactyly (Six digits) in lower limbs.

Fig. 3: Arnold chiari malformation - Type 2. A – Gross image showing the herniation of cerebellar tonsils through the foramen magnum unto upper cervical vertebra (arrows). B – Ultrasound image demonstrating banana sign. C - Ultrasound image demonstrating lemon sign.

Fig. 4: Double outlet right ventricle with ventricular septal defect. A – Gross image showing communication between the two ventricles. B – Gross image showing opening of both aorta and pulmonary artery into right ventricle as pointed by the two sticks.
Discussion

Foetal autopsy contributes a lot to the diagnosis of IUD and congenital anomalies are a most important cause of perinatal deaths. If performed meticulously along with detailed history from clinician, the foetal autopsies can identify the cause of foetal death in most of the cases. In a study by Faye Petersen et al. the cause of death could be detected by autopsy in 94% of the cases. 

In developing countries like India, congenital malformations remain an important cause to contribute to 25 to 30% cases of perinatal death. Major congenital anomalies were seen in 3% of neonates. Multiple congenital anomalies with risk of recurrence in the subsequent pregnancies were noted in 0.7% of the neonates. In our study also congenital anomalies was the leading cause and accounted for about 39.3% of the cases and multiple anomalies were found in 10.7%, which is much higher than in literature, probably because of regional variations and increased prevalence of consanguinity in our region.

In their study, Faye Petersen et al. also detected that one third of the foetal deaths were caused by congenital structural anomalies of which the most common ones were the neural tube defects, hydrops foetalis and congenital heart diseases. Neural tube defects had a five percent risk of recurrence in subsequent pregnancies. In one of our case where the foetus was diagnosed to have spina bifida and meningomyelocoele, also had a history of previous sibling with spina bifida. The most common congenital anomalies were Central nervous system (CNS) disorders accounting for 20 – 45%. Nielson et al. reported 34.8% in 2006, Ceylaner et al. reported 26.4% in 2007, Tomatir et al reported 31.1% in 2009 and Andola et al. reported 34.09% in 2013. Our current study shows 21.4%. We had a case of Arnold chiari malformation type II.

In the series of Tuncbilek E et al, the most frequent congenital malformations were those in the musculoskeletal system. On the other hand, malformations of the musculoskeletal system were the second most common in our series. One case was diagnosed by ultrasonography and confirmed by autopsy. But the other case was diagnosed only by autopsy.

About 25 – 35% of the foetal deaths were due to the abnormalities in the cord, Placenta and membranes. Anomalous foetuses also had abnormal findings in the placenta. The main abnormalities in placenta included placential infection and placental insufficiency, which was associated with intra uterine growth retardation. Soma et al. Identified that the most common lesion found in pregnancy induced hypertension (PIH) and pre eclampsia was placental infarction (54.7%), which was also supported by the study of Wentworth, who identified placental infarction in placenta of 67% of women with severe PIH and in 11.7% of women with mild PIH. In our study, placental causes per se, accounted for 39.2% of foetal deaths and there were placental abnormalities in addition to congenital malformations in 21.4% of foetuses. The most common abnormality in placenta in our study was placental insufficiency. The features of placental insufficiency include increase in syncytial knots also referred to as the Tenny-Parker change. It is termed as increase in syncytial knots, when more than thirty percent of the tertiary villi possess syncytial buds. Another finding is in insufficiency are calcifications and literature states that this placental finding is associated with foetal distress.

In their study, Bengston et al. documented chorioamnionitis in 45.8% of cases with premature rupture of membranes. And out of this 49.1% foetuses died thus had perinatal mortality indicating that the chorioamnionitis is an important cause of intrauterine death. Premature rupture of membranes and pre term labour is the most common cause of amnionitis. In our study chorioamnionitis was seen in two cases, both had history of premature rupture of membranes. Nonspecific mild inflammation was also seen in one case. It’s well documented that infection is one of the important causes of neonatal deaths and still births. Cytomegalovirus...
recurrence, and thus plays a crucial role in counselling the examination and added importantly about the chance of findings that were not diagnosed by prenatal ultrasound. Data shows that the autopsy examination gave additional and can be helpful to counsel the parents. Literature subsequent pregnancies can be detected with foetal autopsy

which may indicate the possibility of recurrence in subsequent pregnancies can be detected with foetal autopsy and can be helpful to counsel the parents. Literature data shows that the autopsy examination gave additional findings that were not diagnosed by prenatal ultrasound examination and added importantly about the chance of recurrence, and thus plays a crucial role in counselling the parents.

**Conclusion**

To conclude, Foetal autopsy is a vital procedure in confirming the cause of foetal death and thus plays a crucial role in prenatal workup. It also provides additional findings not identified through prenatal ultrasound, which can modify the cause of death and thus the risk of recurrence in subsequent conceptions. It is an important tool in providing the treating clinician and parents with very crucial information on implications for future pregnancies.

**Acknowledgements**

I deeply acknowledge my Professor and HOD with all technical staff of department of pathology for helping out in the processing and writing up of this article.

I also acknowledge the department of obstetrics for providing the fetuses for conducting of fetal autopsies to help in the diagnosis of causes of fetal demise.

**Reference**

1. Fatima U, Sherwani R, Khan T, Zaheer S: Foetal Autopsy—Categories and Causes of Death. J Clin Diagn Res 2014; 8:FC05-08.
2. Ceylaner G, Ceylaner S, Gunyeli I, Ekici E, Celasun B, Danisman N: Evaluation of 2407 fetuses in a Turkish population. Prenat Diagn 2007; 27: 800-7.
3. Brodie M, Laing IA, Keeling JW, Meckenzie KJ: Ten years of neonatal autopsies in tertiary referral centre: retrospective study. BMJ 2002; 324:761–3.
4. Siebert JR: Perinatal, fetal and embryonic autopsy: in Gilbert-Barness E, Kapur RP, Oligny LL, Siebert JR, eds. Potter’s pathology of the fetus, infant, and child. Philadelphia: Mosby Elsevier; 2007, 685–729.
5. Cunninham FG, Hollier LM: Fetal Death: in Cunninham FG, Williams, Whitridge J, eds. Williams Obstetrics. Norwalk Ct: Appleton & Lange; 1997.
6. Faye-Petersen OM, Guinn DA, Wenstrom KD: The value of perinatal Autopsy. Obstet Gynecol 1999; 94:915–20.
7. Shankar VH, Phadke SR: Clinical utility of fetal autopsy and comparison with prenatal ultrasound findings. J Perinatol 2006; 26:224–29.
8. Nielsen LA, Maroun LL, Broholm H, Laursen H, Graem N: Neural tube defect and associated anomalies in a fetal perinatal autopsy series. AMPIIS 2006; 114:239-46.
9. Ceylaner S, Ceylaner G, Gunyeli I,Ekici E,Tug M, Taner D, Ekerbicer H, Mollamahmutoglu L, Danisman N: Postmortem evaluation of 220 prenataly diagnoses fetuses with neural tube defect detection of associated anomalies in a Turkish population. Prenat Diagn 2006; 26:147–53.
10. Nayak SR, Garg N: Determination of antepartum fetal death. J Obstet Gynecol India 2010; 60:494-7.
11. Tuncbilek E, Boduroglu K, Alkiasifoglu M: Results of the Turkish congenital malformation survey. Turk J Pediatr 1999; 41:287-97.
12. Allessandri LM, Stanley FJ, Garner JB, Newham J, Walters BNJ: A case-control study of unexplained antepartum stillbirths. Br J Obstet Gynecol 1992; 99:711–8.
13. Bonetti LR, Ferrari P, Trani N, Maccio L, Laura S, Giuliana S, Facchinetti F, Rivasi F: The role of fetal autopsy and
placental examination in the causes of fetal death: a retrospective study of 132 cases of stillbirths. Arch Gynecol Obstet 2011; 283:231–41.

14. Soma H, Yoshida K, Mukaida T, Tabuchi Y: Morphologic changes in the hypertensive placenta. Contrib. Gynecol. Obstet 1982; 9:58–75.

15. Wentworth P: Placental infarction and Toxemia of pregnancy. Am J Obstet Gynecol 1967; 99:318–26.

16. Tenney B, Parker F: The placental in Toxemia of pregnancy. Am J Obstet Gynecol 1940; 39:1000-5.

17. Bengtson J.M, Van Marter LJ, Barss VA, Greene MF, Tuomala RE, Epstein MF: Pregnancy outcome after premature rupture of membranes at or before 26 weeks gestation. Obstet Gynaecol 1989; 73:921–7.

18. Grandjean H, Larroque D, Levi S: The performance of routine ultrasonographic screening of pregnancies in the Eurofetus study. Am J Obstet Gynaecol 1999; 81:446-54.

*Corresponding author:*  
Dr P Jayaganesh, Postal Address: No 47, Jani Jhan Khan Road, Royapettah, Chennai 600014 INDIA  
Phone: +91 9884115987  
Email: jayaganeshp@gmail.com

Financial or other Competing Interests: None.