Study of the Non-Immune Hydrops Fetalis (NIHF) and its Management

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Abstract

Hydrops Fetalis is a Greek term that describes pathological fluid accumulation in fetal soft tissues and serous cavities, peritoneal, pleural, pericardial, scalp and body wall. Hydrops Fetalis is considered as a prenatal form of cardiac failure. It is broadly divided into two groups - Immune Hydrops Fetalis (IHF) and Non-Immune Hydrops Fetalis (NIHF). In previous years most of the Immune Hydrops Fetalis is caused by Erythroblastosis Fetalis secondary to Rh Iso-immunization with presence of maternal circulating red cell antibodies. IHF has now decreased with passive maternal immunization markedly up to 20% (1990).

Recently, Non-immune Hydrops Fetalis (NIHF) is relatively more common and accounts for 90% of Hydrops Fetalis. NIHF is also defined as universal edema unassociated with Erythroblastosis Fetalis. It can be caused by wide variety of factors including fetal disorders (Chromosomal abnormalities, Congenital malformation & Twinning), Cardiovascular, metabolic causes, maternal diseases (Diabetes, Hypothyroidism, Anemias), placental and cord abnormalities and infectious pathogens acquired in-utero (Cytomegalovirus). The incidence of NIHF is estimated at 1 in 3000 pregnancies. The aetiology of Hydrops Foetalis is based on the presence or absence of anaemia leading to heart failure and generalized oedema in the foetus and neonate. In approximately 1/4 of all cases, the cause is not determined.

Keywords: Non-immune Hydrops Fetalis (NIHF); Immune Hydrops Fetalis (IHF); Edema; Pericardial effusion; Peritoneal effusion and Erythroblastosis Fetalis

Introduction

Hydrops Fetalis is an accumulation of extracellular fluid in fetal body cavities like pleural, pericardial, scalp and in body wall [1]. Ballantyne described the first case of Hydrops Fetalis over 100 years ago, later NIHF was first distinguished from IHF by Edit Potter in 1943. He described NIHF as universal edema other than Rh incompatibility [2]. NIHF is associated with accumulation of extracellular fluid in fetal body cavities like pleural,
pericardial, scalp, body wall with increased skin thickness of more than 5mm due to generalized subcutaneous edema and placental enlargement. When first described HF constituted 20% of all cases, but with effective anti D prophylaxis for immune Hydrops, NIHF constitutes 90% cases of fetal Hydrops [3]. NIHF incidence reported as 1 in 3000 pregnancies [4]. Approximately 50% of fetuses with non-immune hydrops fetalis die in utero, and about half of the live born infants survive.

Routine ultrasound has been recommended as the initial diagnosis of NIHF. Placental thickness should be measured in all cases to rule out placental edema. 3D scan helps in determination of Facial dysmorphology which may give clue to inborn errors of metabolism and Cystic hygroma may be in cases of Aneuploidy, fetal echocardiography, examination of maternal blood for fetal erythrocytes, amniocentesis and sampling of fetal blood. We report two of NIHF fetuses about 16-18 weeks of gestational age.

Materials and Methods

The present study was conducted over a period of one year among 1000 live births in Kamineni academy of medical sciences and research Centre. We found two cases of Non-immune Hydrops Fetalis (NIHF) of 16-18 weeks. Hydrops Fetuses are discovered incidentally during Prenatal Ultra sonography in the first and second trimester. The incidence, etiology, management options regarding NIHF were discussed in this article.

Observations and Results

Fetus-1: was a Male NIHF about 16-18 weeks (Figure 1&2).

Obstetric history

A 28 years old mother G3P2L2 with 16-18 weeks of gestation came to the hospital and diagnosed IUD. The First pregnancy was full term, normal vaginal delivery, female 6yrs and is healthy. The second pregnancy was full term, normal vaginal delivery, female 3yrs healthy. The present case was Third pregnancy, spontaneous conception with 18 weeks of gestation. The BP was 110/70mmHg, pulse rate 82/min, blood group was A +ve and Hb – 5.3 gms. The mother with severe anemia and gave history of viral infection during early pregnancy.

P/A Examination: Uterus 20 weeks and FHS was absent.

USG Report

a. Single fetus with no cardiac activity and no fetal movements.
b. Cystic hygromatous changes around the neck.
c. Signs of Fetal Hydrops changes: Oedema is seen in the fetal subcutaneous tissue and with Scalp edema.
d. Fetus -16+ weeks, weight 144gms.
e. Over riding of small bones of skull.
f. Liquor adequate. Placentomegaly, Fundal, Posterior, and with Grade 1 maturity.

Impression: Intra uterine fetal demise with gestational age of 16 weeks 2days, with Hydrops changes. After admission consent was taken for MTP. She spontaneously expelled dead fetus weight 180 gms with edematous and hydrops changes. Placenta and membranes expelled and weighing 140gms. The dead fetus was send to the Anatomy department KAMS& RC for academic purpose.

Impression: Male fetus with Non-immune Hydrops fetalis and cystic hygromatous changes around the neck with Facial dysmorphology (Figure 1&2).
Fetus 2: was a Female NIHF about 16-18 weeks, with multiple congenital anomalies and mother underwent MTP (Figures 3 & 4).

Obstetric history: A 22year old mother G3P1L1 with 16-17 weeks of gestation. The First pregnancy was full term, normal vaginal delivery, and male baby of 18 months old and healthy. The Second pregnancy was a twin gestation of 12 weeks induced abortion for fetal anomalies. The present pregnancy was a spontaneous conception with 16-18 weeks of gestation. The BP was 120/80mmHg, pulse rate 82/min, blood group was A +ve and Hb – 7.2 gms.

P/A Examination: Uterus 16-18 weeks and FHS +ve.

Tiffa Scan Report
a) Single viable fetus with fetal movements. Generalized subcutaneous edema of the entire body with presence of bilateral pleural and pericardial effusion. Signs of Fetal Hydrops changes.

b) Brain: Choroid plexus cyst noted in left lateral ventricle. Multi-loculated cystic lesion noted in the neck not communicating with cranium suggestive of cystic hygroma.

c) Thorax: Both lungs appear echogenic with pleural effusion. Heart: Complex congenital anomaly with single ventricle and mild pericardial effusion. Heart rate: 167 BPM.

d) Fetus 17+ weeks, weight 147gms.

e) Liquor adequate. Placenta Fundal, Anterior, Grade 1 maturity.

Diagnosis: Single live Intra uterine fetus with gestational age 16 weeks 5 days, with Hydrops fetalis and multiple congenital anomalies with facial dysmorphology.

After admission consent was taken for MTP. She underwent medical abortion and expelled fetus and placenta. The Fetus was edematous with Hydrops changes. Placenta and membranes expelled and was 160 gms. The dead fetus was sent to Anatomy department KAMS& RC for academic purpose.

Impression: Female fetus with Non- immune Hydrops fetalis and cystic hygromatous changes around the neck.

Discussion

Hydrops fetalis is a Greek term, that describes pathological fluid accumulation in fetal soft tissues and serous cavities. The more common areas include fluid in the abdomen (ascites), fluid around the lungs (pleural effusion), fluid around the heart (pericardial effusion), or extra fluid under the skin edema (anasarca) or scalp. It is detected by foetal ultrasound if the presence of abnormal fluid collection and skin edema (skin thickness)>5mm [5]. Sonographic findings include Placental thickness >4cms in second trimester or >6cms in third trimester [6,7].
Hydrops fetalis are of two types. In previous years most of cases of Hydrops were caused by severe Erythroblastosis fetalis secondary to Rh isoimmunization. With passive maternal immunization, this type of Immune Hydrops fetalis (IHF) has decreased markedly from 100% to less than 20%. According to Wilson, et al. [8] the commonest cause of Hydrops (IHF) was Rh isoimmunization and mortality rates were as high as 100% and fell to less than 20% by 1990. Other studies shows that there are other materno-fetal blood group incompatibilities can still occur. The changing etiology presently most of the cases of Hydrops fetalis are now caused by Non immune Hydrops Foetalis (NIHF). Later Edith Potter in 1943 described NIHF with universal edema unassociated with Erythroblastosis which now accounts for 90% of cases of Hydrops fetalis [2,3]. The diversity of potential etiologies in the investigations of NIHF requires multi disciplinary approach. NIHF has a favourable outcome in 27.5% of cases with idiopathic NIHF carrying the worst prognosis. Negative prognostic factors described are enlarged biventricular dimension, early manifestation of pericardial effusion, and pulmonary hypoplasia secondary to pleural effusions. Pleural fluid drainage in utero may improve outcome in NIHF and timely performed thoracocentesis shunting procedures are also associated with a 60% survival rate.

If Hydrops develops after 30 weeks of gestation, the mortality rate is approximately 25%. If Hydrops develops before 30 weeks gestation, the mortality rate is more than 90%.

Pathogenesis

The common Pathophysiology underlying the etiology of Hydrops fetalis of Nonimmune type is imbalance in regulation of fluid movement between the vascular and interstitial spaces. It may be increase in interstitial fluid production or decrease in lymphatic return [9]. The most common etiology of NIHF includes cardiovascular, chromosomal /genetic and hematologic abnormalities in the fetus. Others may be structural fetal anomalies of thorax, twinning complications, congenital hepatic cirrhosis, and congenital nephritic syndrome. All leading to severe cardiac failure in fetus. Approximately 50% of fetuses with NIHF die in utero with severe viral infections causing fetal anemia (Parvo virus B19 or TORCH). Others include placental and cord abnormalities and inherited metabolic disorders which may also be cause NIHF [10]. New causes of NIHF are being reported every year but approximately in 50% of cases, no cause can be found despite extensive investigations [11]. Ultrasonography before 20 weeks gestation shows the first sign of increased nuchal translucency, followed by generalized skin edema, particularly at back, wherein skin thickness may be 15 to 20 mm. Mostly associated with complex cardiac defects or abnormal karyotype like trisomy 18 or 21 [12]. Before 20 week of gestation, placental oedema is a consistent finding with skin edema and thickness over 25mm is diagnosed as NIHF. Ultrasound measurement of placental thickness is considered for early diagnosis of alpha thalassemia. Maternal serum sample should be considered in cases of consanguineous marriage, stillborn and recurrent Hydrops [13]. An Inborn error of metabolism (IBM) causes Hydrops either by severe anemia or liver failure. This may account for 10% of total cases. Prognosis for IBM is poor leading to recurrent Hydrops being up to 25% in subsequent pregnancy. Facial dysmorphism may give clue to inborn errors of metabolism. Cystic hygroma may be related to an Aneuploidy (Trisomy 13,18 & 21) or Turner syndrome [14-16].

Women with NIHF may develop Mirror syndrome, an uncommon complication in which mother develops edema that “mirrors” that of her Hydrops fetus. Mirror syndrome may represent a form of preeclampsia with edema, hypertension and proteinuria [17].

The current study reports two fetuses with NIHF associated with cystic hygroma and facial dysmorphology. The First fetus was a Male, about 16-18 weeks of gestation, with intrauterine death. A sonographic investigation identified the fetus as NIHF with cystic hygroma. The mother presented with severe anaemia and h/o viral infection during pregnancy. The final impression was male fetus with 16 -18 weeks of gestation with NIHF and without any congenital anomalies. Main cause of NIHF probably may be severe anaemia and viral infection of the mother which may be spread to the fetus or no cause can be found despite extensive investigations. However Foetus had Cystic hygroma which may be suggestive of Aneuploidy (Trisomy 13,18 & 21) or Turner’s syndrome and Facial dysmorphology which is suggestive of an Inborn metabolic errors.

The Second FETUS was a female, about 16-18 weeks of gestation identified NIHF with severe Cardiac abnormalities, CNS defects and cystic hygroma on ultrasound. Medical termination of pregnancy was performed. The final impression was female fetus with 16 -18 weeks of gestation with NIHF. However this Foetus also had Cystic hygroma and Facial dysmorphology which is a clue for any Aneuploidy (Trisomy 13,18&21) or Turner’s syndrome or In born metabolic errors leading to multiple congenital anomalies. In both the cases the parents were advised for chromosomal and genetic analysis.
Conclusion

NIHF requires a multidisciplinary approach. Evaluation of Hydrops is by antibody screening (Indirect Coombs test) followed by detailed sonography. Ultra Sonographic identification of Hydrops fetalis and placenta can be determined in about 65-85% of cases although this includes postnatal evaluation. Fetal chromosomal analysis and genetic molecular testing can be done to rule out any aneuploidy. Fetus with heart failure needs a multi-vessel doppler study using umbilical artery or vein, middle cerebral artery or ductus venosus. Management is guided by presence or absence of additional anomalies. The prognosis depends on etiology and gestational age.

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