RESEARCH ARTICLE

PREGNANCY COMPLICATED WITH HEREDITARY SPHEROCYTOSIS – LESSONS LEARNT FROM A CASE AND REVIEW OF LITERATURE.

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Abstract

Hereditary spherocytosis (HS) also known as Minkowski-Chauffard syndrome is a genetically determined disorder of the red blood cell membrane cytoskeleton complex causing Hemolytic anemia. There is a wide spectrum of clinical presentation ranging from mild anemia to severe anemia with splenomegaly and jaundice. Increased Mean Corpuscular Hemoglobin Concentration (MCHC), spherocytes in peripheral blood smear and increased osmotic fragility favour the diagnosis of HS. No causal treatment is yet available for this disease. Mild disease warrants no treatment. Folic acid, supportive treatment and regular annual check up usually suffice for moderate disease while frequent hematological supervision and splenectomy is prudent for severe disease. Pregnancy complicates the management of HS and very limited data is available in the literature regarding this.

Case Presentation:- Authors hereby report a case of 24-year-old 36-weeks gestational age primigravida with intrauterine fetal demise. Workup revealed patient to be suffering from Hereditary Spherocytosis that resulted in hemolysis ultimately causing fetal demise. Pregnancy was terminated and patient was subsequently referred to the Department of Surgery for splenectomy.

Conclusion:- Hereditary Spherocytosis, one of the commonest congenital hemolytic anemia may get decompensated during pregnancy. Due to paucity of data in literature regarding management of HS during pregnancy, it is hard to formulate guidelines for indications of splenectomy during pregnancy. However, authors recommend Splenectomy preferably in second trimester in any pregnant women with HS who is experiencing hemolysis or having symptoms, signs and complications thereof. Obstetricians should exercise high index of suspicion to diagnose this not so uncommon entity as early as pregnancy is diagnosed to yield better outcomes.

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Introduction:-
Hereditary spherocytosis (HS) is a genetically determined disorder of the red blood cell membrane cytoskeleton complex causing Hemolytic anemia. The estimated frequency of HS varies from 1:2000 to 1:5000.1-3 Minkowski and Chauffard in the late 19th century reported families whose peripheral blood revealed numerous spherocytes and have been thus duly credited with the discovery of this not so uncommon disorder.4,5 The red cells were abnormally susceptible to lysis in hypotonic media as revealed by in vitro studies, thereby establishing osmotic fragility test virtually diagnostic of HS.

There is a wide spectrum of clinical presentation ranging from mild anemia to severe anemia with splenomegaly and jaundice. Increased Mean Corpuscular Hemoglobin Concentration (MCHC), spherocytes in peripheral blood smear and increased osmotic fragility favour the diagnosis of HS. Sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE) is usually confirmative.1,2

No causal treatment is yet available for this disease. Mild disease warrants no treatment. Folic acid, supportive treatment and regular annual check up usually suffice for moderate disease while frequent hematological supervision and splenectomy is prudent for severe disease.6 It is a well known fact that intercurrent conditions like, pregnancy and infections may cause decompensation in a previously well compensated disease but unfortunately very limited data is available regarding the course of HS during pregnancy with very few cases reported in the literature till date.7 Authors hereby report a case of pregnancy complicated with intrauterine fetal demise who was later diagnosed to be suffering from HS.

Case report:-
A 24-year-old 36-weeks gestational age primigravida presented in the emergency department with the complaints of decreased fetal movement since morning. She was married for 11 months and had normal menstrual cycles. Past history was insignificant except for one unit packed red blood cell transfusion at the age of 12 years. Examination revealed pallor and icterus. Abdominal examination showed gravid uterus with fundal height corresponding to 32-34 weeks gestational age with longitudinal lie and cephalic presentation. Head was found to be floating. No fetal movements were felt during examination. Fetal Heart sounds could not be auscultated. Patient was relaxed and there was no evidence of uterine contractions. Per speculum examination showed Cervix and Vagina to be healthy. There was no evidence of per vaginal discharge or bleeding. Per vagium examination revealed os to be closed, posteriorly placed and was uneffaced.

Obstetric Ultrasonography examination revealed single intrauterine fetus of 38 weeks 0 day in cephalic presentation with absent fetal cardiac activity and fetal movements suggestive of fetal demise. Complete blood count showed anemia with 9.9 g Hemoglobin/dL, leukocytosis with 15600 total leucocyte counts/ cubic mm. Mean Corpuscular Volume (MCV) was 95.9 fl, Mean Corpuscular Hemoglobin (MCH)was 33.9 pg and Mean Corpuscular Hemoglobin Concentration (MCHC)was 35.4 g/dL. Peripheral blood smear showed increased number of spherocytes (+++) with few polychromatophils. Spherocytes were uniform in size and density without any central pallor. Platelets were adequate in number with 2.81 lacs/ cubic mm. Reticulocytosis was present with 8.0 % reticulocytes. Liver function tests revealed indirect hyperbilirubinemia with 5.7 mg total serum bilirubin/dL and 0.9 mg direct serum bilirubin/dL. Liver enzymes were normal. Renal Function Tests were normal. Serum Lactate dehydrogenase was found to be raised with values of 566.9 Units/L. Iron studies, Glucose-6-Phosphate Dehydrogenase (G6PD) assay and Hemoglobin electrophoresis were normal. Patient tested negative for antinuclear antibody (ANA), Indirect and Direct Coomb’s test. The above findings were strongly suggestive of Hemolytic anemia most probably as a result of Hereditary Spherocytosis. Subsequently Osmotic fragility test was done which concluded Initial lysis (minimal resistance) in 0.745 % NaCl, Complete lysis (maximum resistance) in 0.33 % NaCl and Mean corpuscular fragility in 0.515 % NaCl, thereby confirming the diagnosis of Hereditary Spherocytosis.

While enquiring about the old records, patient showed six months old reports. Complete blood counts showed anemia with 10.6 g Hb /dL, leukocytosis with 12130 TLC/ cumm and 2.64 lacs Platelets / cumm. MCV was 89.6 fl, MCH was 30.0 pg and MCHC was 33.5%. Indirect hyperbilirubinemia with 3.2 mg/dL total serum bilirubin and 1.6 mg/dL direct fraction was seen in LFT. USG abdomen done at the periphery showed minimal hepatomegaly with multiple calculi in Gall bladder with moderate splenomegaly and a single live early intrauterine pregnancy of gestational age 12 weeks 5 days.
A diagnosis of 36 weeks pregnancy with Hereditary Spherocytosis with Intrauterine fetal demise was made and was planned to terminate the pregnancy.

**Treatment and outcome:-**

Induction of labour was started with Dinoprostone (Prostaglandin E2) gel followed by insertion of intracervical catheter. Labour was augmented by Oxytocin infusion and dead fetus was delivered. Postpartum period was uneventful. Patient was prescribed Folic acid 5 mg per day and thereafter was referred to the Department of Surgery for Splenectomy.

**Table 1:** Classification of Hereditary Spherocytosis.

<table>
<thead>
<tr>
<th>CLASSIFICATION</th>
<th>Trait</th>
<th>Mild</th>
<th>Moderate</th>
<th>Severe</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemoglobin (g/dL)</td>
<td>Normal</td>
<td>11-15</td>
<td>8-12</td>
<td>6-8</td>
</tr>
<tr>
<td>Reticulocyte count (%)</td>
<td>Normal (&lt;3%)</td>
<td>3-6</td>
<td>&gt;6</td>
<td>&gt;10</td>
</tr>
<tr>
<td>Bilirubin (μmol/L)</td>
<td>&lt;17</td>
<td>17-34</td>
<td>&gt;34</td>
<td>&gt;51</td>
</tr>
<tr>
<td>Spectrin per erythrocyte (%)</td>
<td>100</td>
<td>80-100</td>
<td>50-80</td>
<td>40-60</td>
</tr>
</tbody>
</table>

**Discussion:-**

Hereditary spherocytosis (HS) is one of the commonest congenital hemolytic anemia caused by genetic mutations in genes encoding red blood cell (RBC) membrane proteins. Various membrane proteins can be affected in HS like spectrin, ankyrin, band 3, and protein 4.2 but spectrin deficiency is reported to be the commonest culprit resulting in HS. Autosomal dominant pattern of inheritance is the most frequent mode of transmission of this genetically heterogeneous disease but few cases of autosomal recessive inheritance have also been reported. The estimated prevalence as reported in various systematic reviews range from 1 in 2000 to 1 in 5000 people.\(^1\)\(^-\)\(^3\)

Clinically, there is a wide spectrum of presentation ranging from asymptomatic mild anemia to fulminant hemolytic anemia with pallor, jaundice and splenomegaly requiring aggressive treatment. Gallstones of pigment type caused by increased unconjugated bilirubin may be the first presentation of this disorder. Family history is variably present with patients reporting history of splenectomy and cholecystectomy in family member. Environmental triggers, like, fatigue, infections, emotional distress and pregnancy have been reported to unmask the previously compensated disease.\(^7\)

The HS is characterized by mild to moderate anemia with increased mean corpuscular hemoglobin concentration (MCHC). Increased MCHC due to cellular dehydration is a hallmark of HS. Reticulocytosis is invariably present signifying increased cell production. Peripheral blood smear reveals spherocytes, which in contrast to spherocytes seen in immune hemolytic anemia and thermal injury are quite uniform in size and density (Fig. –1). Unconjugated hyperbilirubinemia is frequently observed due to hemolysis. Incubated osmotic fragility test is the most sensitive test for HS (Fig. –2).\(^8\) Other tests infrequently used are Acidified glycerol lysis time (AGLT), Flow cytometry (Eosin-5-maleimide binding), Osmotic gradient ektacytometry and membrane analysis by SDS-PAGE.\(^9\)
Fig. 1: Peripheral smear shows increased number of spherocytes (black arrow) and few polychromatophils (red arrow). Spherocytes are uniform in size and density without any central pallor.

Fig. 2: Osmotic Fragility Curve of the patient. The whole curve is shifted to the right. Please note the tailing of fragile erythrocytes, which are lysed at higher concentration of saline (0.7-0.8 %). Normally no lysis occurs at these concentrations.
This disorder despite being discovered almost 2 centuries back, unfortunately, there is still no causative treatment to correct the basic disorder. Decision to offer treatment to the patients suffering from HS is based on the severity of clinical symptoms and complications (Table-1). Folate supplements have been routinely prescribed to the patients suffering from chronic hemolysis and folate therapy is recommended in moderate and severe HS but not in mild HS in the dose of 5 mg per day.

Splenectomy has long been the standard treatment for patients with clinically severe HS and is considered in those with moderate HS. It should not be performed in mild HS. Splenectomy is a double edged sword, at one end it eliminates the sign and symptoms of hemolysis and at the other end it makes individual susceptible for life long risk of infections with capsulated organism, especially, Pneumococci with a mortality of 0.1-0.4%. However, this risk has been reduced by vaccination against pneumococcus, meningococcus and Hemophilus influenzae coupled with prophylactic antibiotics. It has been recommended to vaccinate patients undergoing splenectomy two weeks prior to surgery. Recently, a largest single center study done by Thomas Pincez et al reported safety and success of subtotal splenectomy over total splenectomy in children. Concurrent cholecystectomy is advocated if the patient has symptoms of gall stone disease.

Pregnancy complicates the management of hereditary spherocytosis. Traditionally, many authors have held the view of avoiding splenectomy during pregnancy due to fear of perioperative morbidity associated with the surgery to both mother and fetus. There is paucity of data in literature in this regard but the studies although few have been contrary to the long held concept of avoiding splenectomy during pregnancy. Second trimester is usually preferred for splenectomy, although it can be performed at any gestational age. Less invasive laparoscopic splenectomy is routinely preferred due to its obvious advantages over open splenectomy.

Moore et al (1976) have reported two cases of pregnant women in the second trimester with HS who underwent successful splenectomy and reviewed 5 such cases in literature. Ho-Yen DO (1984) described a family with HS in which 3 sisters had hemolytic episodes during their 6 pregnancies but none was treated with splenectomy and each pregnancy resulted in birth of mature and live infant. Maberry et al (1992) reported 23 women with HS during their 50 pregnancies, none requiring splenectomy during pregnancy. Maternal complications were infrequent and perinatal outcomes were excellent. Pajor et al (1993) have reported 8 patients with HS who had 19 pregnancies in total, 10 before splenectomy and 9 after splenectomy. All were managed conservatively and was found that fetal outcome was more favourable after splenectomy than before splenectomy. Brabec V et al (1999) studied the influence of pregnancy on the course of HS in 21 women during their 44 pregnancies and found that pregnancy caused no problems in the majority, even when complications developed they were not serious. Anemia deteriorated in one-third pregnancies in non-splenectomized women due to increased hemolysis, while complaints were minimal in splenectomized patients. Only two instances of successful laparoscopic splenectomy in a pregnant woman suffering from HS have been reported in literature yet.

The case presented by the authors seems to have been undergoing hemolysis since last six months as evident by anaemia and unconjugated hyperbilirubinemia in routine hematological reports and also there was evidence of hepatosplenomegaly in Ultrasonography study of abdomen but unfortunately it could not be recognized at the periphery. It was sad that patient presented to the authors when fetal demise already occurred. Had she been previously worked up and diagnosed as a case of HS and underwent splenectomy, authors feel that it would have been possible to save the life of fetus.

Conclusion:
Hereditary Spherocytosis one of the commonest congenital hemolytic anemia may get decompensated during pregnancy and pregnancy complicates the management of Hereditary Spherocytosis. Pregnancy is usually well tolerated and need for splenectomy doesn’t arise frequently. Although there is a consensus to subject patient to splenectomy and its inherent risks in clinically moderate and severe HS, the management is still controversial during pregnancy. Given the few studies in literature regarding management of HS during pregnancy, it is hard to formulate guidelines for indications of splenectomy during pregnancy at this juncture. The decision to perform splenectomy in pregnancy is guided at best by the indications in non-pregnant patients of HS. However, authors recommend splenectomy in any pregnant women with HS who is experiencing hemolysis or having symptoms, signs and complications thereof. Second trimester should be preferred for splenectomy. Of note, Obstetricians should exercise high index of suspicion to diagnose this not so uncommon entity as early as pregnancy is diagnosed if a present with signs and symptoms suggestive of hemolysis and plan out further management along with discussing the option of
spleenectomy with patient and her attendants for better perinatal outcome. Not surprisingly, earlier diagnosis and management can commensurate into better outcomes.

References: