Mirror Syndrome - A Rare Case

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Received: January 19, 2018; Published: January 24, 2018
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Abstract
Mirror syndrome is a maternal condition which develops as a result of fetal condition, the main feature being edema; both mother and fetus appear as mirror image of each other. Initially it was described as maternal edema in response to fetal edema due to Rhesus isoimmunisation [1]. This clinical condition was discovered as early as 1892 by John W Ballantyne who found and explained the association of maternal edema in pregnancy with fetal and placental hydrops due to rhesus immunization [2]. Subsequently this rare condition has been found in cases of fetal hydrops due to various causes. One such uncommon case of ‘Mirror Syndrome’ due to non immune hydrops is being presented.

Keywords: Ballantyne syndrome; Fetal Hydrops, Preeclampsia

Introduction
Mirror syndrome is a maternal condition which develops as a result of fetal condition, the main feature being maternal and fetal edema. Both appear as mirror image of each other hence named so. The term ‘Mirror Syndrome’ was first introduced by O’Driscoll who found this condition in cases of fetal hydrops due to Rh Isoimmunisation [1]. This clinical condition was discovered as early as 1892 by John W Ballantyne who found and explained the association of maternal edema in pregnancy with fetal and placental hydrops due to rhesus immunization [2]. Subsequently this rare condition has been found in cases of fetal hydrops due to various causes. One such uncommon case of ‘Mirror Syndrome’ due to non immune hydrops is being presented.

Case Report
27 years old gravida 2 para 1 was a booked case at a peripheral outpatient clinic. She was on regular follow up there when the medical attendant detected that patient had gained excessive weight and was hypertensive hence she was referred to this hospital. At that time she was 29 weeks pregnant and did not have any complaint except swelling feet of insidious onset of two weeks duration. Her past medical and surgical history was unremarkable. She had one normal delivery three years back. Premonitory symptoms or breathing difficulty were not present. Her weight was 87 kg (pre pregnancy her weight was 63 Kg), BP 170/110 mm of Hg, her BP was recorded as normal on the previous visit which was four weeks prior, and there was mild pallor. She had excessive pedal edema up to mid thighs and abdominal wall. There was mild pallor otherwise general and systemic examination was essentially normal. Obstetrical examination showed fundal height of 34 weeks (at gestation of 29 weeks) with abdominal wall edema. Fetal parts were felt but with difficulty due to excessive abdominal wall swelling. Fetal heart rate was normal. Her blood group was A Rh positive. She was admitted to the obstetrical ward for detailed investigations, rest and close monitoring. She was started on Labetalol. Her Hb was 8.3 gm%, with normal leukocyte count but platelets were low (77,000/ml), prothrombin time and INR were normal. Blood sugar, kidney function tests and liver function tests, HIV and other viral markers were normal.

Ultrasound done that day revealed fetal ascites, bilateral pleural effusion and skin edema. Head circumference and femur length were corresponding to 25 weeks gestation but abdominal circumference was equal to 32 weeks size. Heart, spine and kidneys were normal and amniotic fluid was normal. Placenta was located fundo posteriorly. The cause of fetal hydrops was not clearly understood, Rh isoimmunisation and ABO incompatibility were ruled out. TORCH test and mother’s Hb electrophoresis were normal. Facilities for amniocentesis, karyotype, and fetal echocardiography were not available. Parents were counselled about poor fetal prognosis and risk to mother; they agreed for termination. Since Bishop’s score was poor cervical ripening was initiated. With priming doses only she went into labour and developed manifestations of revealed type abruptio placenta during late first stage of labour, lost about 300 ml of fresh blood. Augmentation after ARM was done and she progressed normally and delivered a male still born baby weighing 1.5 kg, baby had massive distention of abdomen due to ascites (Figure 1). Placenta was markedly enlarged and weighed...
1.2 Kg (Figure 2). She improved after delivery; her blood pressure became normal without antihypertensive after eight days. She was reviewed after two weeks when her swelling had disappeared completely. Postmortem examination of baby was not agreed to and HPE of placenta was unremarkable.

**Figure 1:** Fetal hydrops (Abdominal distension due to Ascites).

**Figure 2:** Hydropic and enlarged placenta weighing 1.2 Kg (Baby’s weight 1.5 Kg).

**Discussion**

Mirror syndrome, an exceptionally rare condition [3]. Is described as the fetal hydrops and maternal edema along with excessive weight gain as found in preeclampsia [4]. This condition is different from preeclampsia as it is associated with hemodilution [5]. Where as preeclampsia is linked with hem concentration. Edamptic fits are unknown in mirror syndrome. Clinical entity of Mirror syndrome is known for more than 100 years yet the causation is not clearly understood [6]. Only after development of ultrasound, the diagnosis of this condition has been possible. Basically it is the fetal hydrops which is the initiating event in causation and manifestation of mirror syndrome. The condition is also called triple edema [7], as it affects fetus, placenta and mother; all having fluid accumulation. The name “Mirror Syndrome” refers to the similarity between maternal edema and fetal hydrops. Fetal hydrops may be due to immunological or non immunological causes [3], though when it was first described it was due to Rh isoimmunisation. Many non immunological causes have been described such as fetal cardiac defects, arrhythmias, aneurysm of vein of Galen, sacrococcygeal teratoma, placental chorisoangioma, infection with human parvovirus, Coxsackie B and cytomegalovirus. Mirror syndrome has been reported in multiple pregnancies due to supraventricular tachycardia in one fetus, TTTS (Twin to Twin Transfusion Syndrome) in a dichorionic triplet pregnancy, and unknown etiology in one fetus in a dichorionic pregnancy [8]. Congenital mesoblastic nephroma (CMN), an extremely rare cause of mirror syndrome has been reported by Takahashi [9]. It is the fetal hydrops irrespective of its etiology which is responsible for maternal condition [7].

Mirror syndrome is associated with a marked increase in fetal mortality and maternal morbidity. Treatment is usually in the interest of the mother unless the fetus has some correctable condition while in utero. Maternal symptoms improve dramatically after delivery or fetal death [10,11]. Braun T [8] mentioned that maternal symptoms associated with mirror syndrome disappeared shortly after treatment of fetal condition or delivery (mean 8.9 days). Sometimes delivery of the baby is difficult due to massive fetal ascites causing obstruction. Delivery is possible in these cases only after aspiration of ascetic fluid [12]. When the specific cause of fetal hydrops can be identified and corrected by in utero treatment, reversion of Ballantyne syndrome can occur [13]. Fetal treatment by intrauterine transfusion for hydrops and anemia caused by parovirus infection and Diamond-Blackfan anemia led to the disappearance of the Mirror syndrome [14]. Placement of a peritoneal- amniotic shunt resolved the fetal hydrops and maternal mirror syndrome [15]. In utero treatment of hydropic fetus can result in disappearance of mirror syndrome. When the specific cause of fetal hydrops cannot be identified and corrected, immediate delivery is necessary in order to avoid fetal death and maternal complications. The genesis of this syndrome remains unknown and elusive though many postulations have been made like elevated serum hCG levels [16]. Placental hypoxia [17]. High levels of anti-angiogenic factor [18]. Many times it is not possible to find the cause of fetal hydrops even after detailed investigations.

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