Spontaneous Ovarian Hyperstimulation Syndrome Complicating a Normal Singleton Pregnancy: A Case Report

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Abstract

Ovarian hyperstimulation syndrome (OHSS) is the most common and sometimes life threatening complication of ovulation induction protocols and assisted reproductive procedures. It usually occurs iatrogenically by the use of drugs such as exogenous gonadotropins or less frequently colmiphene citrate to induce multiple ovulation. However spontaneous form of this syndrome is extremely rare in the literature. In this report we describe a case of spontaneous OHSS that occurred in a 26 year old pregnant woman with no underlying disease. When abdominal distension and dyspnea progressed, leading to deleterious laboratory results, the pregnancy was terminated by therapeutic abortion.

Keywords: pregnancy, spontaneous ovarian hyperstimulation syndrome

Case report

A 26-year-old gravida 2 para 1 woman was admitted to our clinic in the 18th week of gestation because of abdominal distension, dyspnea and vaginal bleeding. She stated that abdominal distension and dyspnea have occurred one week before and since then she was hospitalized in a public hospital because of the confirmed bilateral ovarian cysts and ascites. As her clinical conditions progressing, it was decided to terminate the pregnancy by the intravaginal misoprostol administration. In the day of misoprostol administration, as her blood pressure decreases and dyspnea and abdominal distension worsening, she was sent to our clinic.

At the admission, blood pressure was 70/40 mm/Hg and pulse rate was 100 beats/min. Obstetric examination revealed 3 cm of cervical dilatation and vaginal bleeding due to misoprostol administration. Ultrasonographic examination revealed a living fetus, bilaterally enlarged multicystic ovaries (150X90 mm right, 135X96 mm left) and a large amounts of ascites (Figure 1). The AP X-ray of the chest showed basal pleural effusion on both sides. Laboratory studies revealed leucocytosis (18000/mm³), low concentra-
tions of hemoglobin (9.5 g/l), hematocrit (28%), albumin (2.7 g/dl) and total protein (5 g/dl). Blood urea nitrogen, creatinine and liver function tests were in normal range. On the hormone essays of the admission day, the $\beta$-hCG value was extremely above the normal range for gestational age (200,000 mIU/l), estradiol was 5500 pg/ml and the thyroid function tests were normal. The CA-125 level was 2110 IU/ml.

In her obstetric history she delivered a healthy normal baby at full term four years ago. In her present pregnancy she stated that a nuchal oedema was seen and therefore an amniocentesis was performed at the 12th weeks of gestation. Analysis of the amniocentesis revealed normal karyotype. She also stated that she had never taken medications for ovulation induction.

Two hours after the admission, the patient aborted a 400 g macroscopically normal fetus and dilatation and curettage was also performed. (Histopathological examination of fetus, placenta and the curettage material showed no abnormality.) According to Gollan’s classification, we made the diagnosis as severe OHSS (5). She was immediately managed with intravenous fluid replacement, human albumin and low-molecular weight heparin (LMWH). Body weight, abdominal circumference, intake and outputs, ultrasonography and laboratory studies were monitored strictly daily. Interestingly, the full-blown picture of the syndrome developed after the day of the abortion. The $\beta$-hCG value increased to 330,000 mIU/l, hemocoencentration (41%) and thrombocytosis (780,000/mm$^3$) occurred. As the dyspnea and abdominal distension remains after the abortion, ultrasound guided therapeutic paracentesis was performed three times with two days intervals and 5500 cc ascitic fluids was taken totally. Cytological examination of the ascites revealed fluid of theca lutein cyst.

At the end of 4th week of admission, her complaints of dyspnea and abdominal distension decreased, laboratory abnormalities like hemocoencentration and thrombocytosis improved and size of the ovarian cysts and ascites regressed. Three weeks after the discharge, she had no complaint, ultrasonographic and laboratory examination was completely normal.

**Discussion**

OHSS occurs iatrogenically in almost all cases by using the drugs to induce multiple ovulation in infertile patients. This condition may also develop spontaneously, but it is an extremely rare event. Although there are no clear predictive risk factors for the development of OHSS, young age, polycystic ovary syndrome, asthenic habitus, luteal supplementation of hCG, protocols with GnRHa, high level of serum estradiol, multiple follicles and ovarian necklace sign were reported as the possible risk factors (4). This syndrome is characterized by massive transudation of protein-rich fluid (mainly albumin) from the vascular space especially into the peritoneal and pleural cavities. It has been reported that the intensity of the syndrome is related to the degree of the follicular response in the ovaries to the ovulation inducing agents (6).

The exact pathogenesis of OHSS is not yet clearly determined. It has been suggested that vasoactive substances such as histamine, serotonin, prostaglandins, interleukins, TNF-$\alpha$, ovarian renin-angiotensin system and vascular endothelial growth factor (VEGF) which are activated by exogenous gonadotropins can lead to increased vascular permeability and extravascular fluid accumulation in OHSS (4,7-9). In recent years particularly VEGF became more popular on the pathogenesis of OHSS. It has been reported that it is responsible for the significant increase in the capillary permeability, extravascular fluid accumulation, hemocoencentration and elevated plasma concentration of von Willebrand factor, all known complications of OHSS (4,10). Elevated levels of this cytokine were found both in the serum and in the ascitic fluid of patients with severe OHSS (4,10,11). However it has been also reported that the serum concentrations of VEGF does not predict the course of the disease (12). The etiopathogenesis of spontaneous OHSS is less clear. Some authors suggested that poly cystic ovary syndrome (PCOS) could also be a risk factor for spontaneous OHSS (13,14). However some cases developed this condition without underlying PCOS (15-18). Spontaneously developed OHSS has been reported in twin and molar pregnancies in which the endogenous hCG levels were higher than normal (19,20). However, OHSS also has been observed in women with normal or lower than normal hCG concentration. Thus, it is postulated that high concentrations of hCG are not responsible for every case of OHSS (21). Hypothyroidism is another postulated risk factor for the development of spontaneous
OHSS (22,23). Our case did not show any signs of PCOS or hypothyroidism as confirmed by blood and ultrasound examinations we performed 3 months after the abortion.

Our case developed the full-blown picture of OHSS after the termination of the pregnancy, and this picture lasted during two weeks. We could exclude the molar pregnancy by histopathological examination of the placenta and fetus, and ovarian malignancy by cytological examination of the ascitic fluid. We postulated that elevated concentration of β-hCG was the main triggering mechanism for the present case.

The principals of management of OHSS are the same in both spontaneously and iatrogenically developed forms. When OHSS complicates pregnancy, many authors recommend the continuation of the pregnancy, as the syndrome is a self-limited process. In almost all cases, the disease regresses spontaneously with time or delivery. However the management and treatment of each patient is critical as deaths from the syndrome have been reported due to hypovolemia, hemorrhage and thromboembolic phenomena (6). Hospitalization is needed in moderate and severe forms. Strictly monitoring of the hemodynamic status, intravenous crystalloid and albumin infusion, application of LMWH for the prophylaxis of thrombosis are the main steps for the management. Paracentesis of ascitic fluids may have positive effects on the respiratory functions and also on the renal functions by increasing urinary output and reducing blood urea nitrogen (24).

However the conservative management could not be sufficient in every case and termination of the pregnancy might be necessary. (13,19). If the clinical conditions worsening despite the extensive therapy as happened in our case, termination of the pregnancy should be encouraged regardless of the gestational age. Another reported subject with this syndrome is that the exploratory laparotomy was performed in some cases in which ovarian malignancy could not be excluded (16,25). In fact, surgery should be reserved only in cases of ovarian rupture, torsion and intraperitoneal hemorrhage.

References