Two Cases of VACTERL Association in Pregnancy with Lymphocyte Therapy

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Abstract—VACTERL association is a rare disorder with various congenital malformations. The etiology remains unknown. Combination of at least three congenital anomalies of the following criteria is required for diagnosis: vertebral defects, anal atresia, cardiac anomalies, tracheo-esophageal fistula, renal anomalies, and limb defects. The first case was 1-day old male neonate with multiple congenital anomalies was born from 28 years old mother. The mother had history of pregnancy with lymphocyte therapy. His anomalies included: defects in thoracic and lumbar vertebral, anal atresia, bilateral hydronephrosis, atrial septal defect, and lower limb abnormality. Another case was born with 32 weeks gestational age from mother with history of pregnancy with lymphocyte therapy. He had thoracic vertebral defect, cardiac anomalies and renal defect. Diagnosis based on clinical finding is VACTERL association. Early diagnosis is very important to investigation and treatment of other coexistence anomalies. VACTERL association in mothers with history of pregnancy with lymphocyte therapy has suggested possibly of relationship between VACTERL association and this method of pregnancy.

Keywords—Anal atresia, tracheo-esophageal fistula, atrial septal defect, lymphocyte therapy.

I. INTRODUCTION

VACTERL association, also named as VATER, VACTER, and VACTERL associations was firstly named in 1973 by American physicians David Weyhe Smith and Linda Quan to explain a combination of some specific birth defects [1]-[4]. VACTERL is a descriptive acronym where each letter represents a birth defect. Thus, V denotes the vertebral defects, A indicates for anal atresia, C is for cardiac defects, TE for tracheo-esophageal fistula, R for renal anomalies, L stands for limb abnormalities and S indicates single umbilical artery. In the beginning of knowing and understanding the disease, it was constituted from four congenital defects and termed as VATER. Years later, the other cases, i.e. cardiac defects (C), limb abnormalities (L) and single umbilical artery (S) were added to its abbreviation. VACTREL association is rarely combined with genital anomalies [5]. Among the mentioned defects, the most common were congenital heart defects with 73.3% and vertebral second with 66.6% [4]-[10]. It is documented that the incidence of the mentioned associated anomalies ranges from about 30% to 60% for low lesions and around 70% for high lesions [11]-[13]. Since there is no clear pathological relationship between these different symptoms, VACTERL is labelled an association and not a syndrome. The association is a disorder affecting multiple organ systems or multiple median and paramedian structures [11], [12]. This disease is known as a fairly common pattern of human malformations, happening in nearly 1.6 per 10,000 births. Some studies also estimated that the incidence of the disease is 1 to 10000 to 40000 [14], [15]. It is rarely seen more than once in one family. The congenital malformations usually seen in such cases are one of the main reasons of death and disabilities during prenatal period.

For diagnosis, at least three cases of the six cases should be occurred. It seems that these anomalies occur in the first 4 weeks of life [16] with mesodermal origin [17]. From disease causation point of view, in some cases, the main reason of these malformations is unknown, including several non-accidental congenital anomalies that there is no special etiology for them [1]-[7]. For some fraction of patients, there is evidence for familial clustering suggestive of inherited factors [18], [19]. It is reported that, roughly 90% of VACTERL association cases take place sporadically, while 10% of cases include familial inheritance [18]. The vast majority of genetic causes described in humans have been reported in isolated individuals or families and overall account for only a small percentage of patients with VACTERL association.

Now, there is no accurate genetic mechanism to explain the disease. Some experts believe that deletion of a small area of genetic material on the long arm of chromosome No. 16 is involved in this disease. Here, there are 4 known genes as FOXF1, FOXC2, MTHFD1 and FOXL1. Investigations show that mutation in FOXF1 gene will result in lung diseases, mutation in FOXC2 gene will result in defects of spine and simultaneous mutation in FOXF1 and FOXC2 will result in kidney abnormalities. Defects in chromosomes Nos. 13 and 17 are involved in VACTERL association [16].

Some reports have suggested that this defects may occur more in children of mothers who taken the cholesterol-lowering statin drugs [20]. Some study reported an increased risk for anorectal malformations and VACTERL association among children born after assisted reproduction techniques [21], [22]. Besides, the disease is often sporadic and its outbreak is more in mothers suffering from diabetes [16]. VACTERL association, however, is most likely caused by multiple parameters. As mentioned, multiple genetic and environmental issues play a role in determining the risk of
evolving this disorder and how severe the disorder will be in an individual.

II. DESCRIPTION

The first case is a 1-day neonate, was born from a 28 years old mother by caesarean section. The mother had history of 7 years of infertility, who is pregnant with lymphocyte therapy. The parents were not relatives and the mother had history of hypothyroidism and used levothyroxine during pregnancy. The mother did not mention to history of diabetes, blood pressure, or congenital malformations in the family.

The neonate had good Apgar score at birth and has been transferred to nursery. In examination of lower extremities, both hips were dislocated and the joints were located as hyper flexion. As shown in Fig. 1, the right knee was dislocated and joints of both knees had hyperextension and some degrees of arthrogyrosis. In the left ankle, clubfoot and in the right ankle, calcaneus valgus was seen. Toes were normal. There are no abnormalities in upper limb. In the examination, male genitalia were with normal phallus. Right testis was undescended and touched in inguinal canal. Left testis was into the scrotum. The anus was closed.

During examination of nose about choanal atresia, the nasogastric tube was passed easily from the left nostril, but 8 french catheter did not pass from the right nostril. Finally, 6 french catheter passed it, which indicated stenosis of right nostril.

In the chest radiograph, which is done by nasogastric tube, esophageal atresia was not seen. As shown in Fig. 2, there is severe involvement between the thoracic and lumbar spine as multiple hemi vertebra, hypoplastic vertebral and scoliosis. Considering several involvements and probable cardiac anomalies, echocardiography was done for the patient, in which small PDA (patent ductus arteriosus) was reported. In renal ultrasound, bilateral renal pelvic dilatation was seen. Considering sever al involvements and probable cardiac anomalies, echocardiography was done for the patient, in which small PDA, small ASD2 (atrial septal defect), severe PH (pulmonary hypertension) and TR (tricuspid regurgitation) were reported.

In the third day of hospitalization, the neonate was suffering from pneumothorax and stated under HFO mode. Finally, the neonate was died at the age of six days due to respiratory failure.

III. DISCUSSION

%70 of the neonates, who suffering from VACTERL, have three symptoms; %25 have 4 symptoms and %10 have 5 symptoms. The first case, suffering vertebral involvement, anal atresia, cardiac involvement, renal involvement, and extremity involvement, had 5 main symptoms and the second case, suffering spine anomalies, cardiac involvement, and renal involvement, had 3 main symptoms.

Spine anomalies were seen in %70 of cases, which is often as hemi vertebra, hypoplastic vertebral and extra vertebral. Cardiac involvement is seen in %75 of cases. The most common demonstrations are VSD, TOF, ASD and coarctation of the aorta. Extremity involvement is often as upper limbs, including anomalies of the thumb and radius bone, which could be unilateral or bilateral.

In the said cases, there was no upper limbs involvement; the only severe lower extremity involvement was seen in the first case that is included in non-common cases.

There are some non-common symptoms in VACTERL association; these symptoms in the first case include genitalia anomalies (cryptorchidism) and narrowing of the nasal passage.
Previous studies have shown that Infants born after In Vitro Fertilisation (IVF) have an increased risk for a congenital malformation [23]. An association between neural tube defects and ovarian stimulation with Clomiphene citrate has been discussed for a very long time [24].

Interestingly, in these cases history of mother infertility and pregnancy by IVF and lymphocyte therapy were seen. Considering that the cause of this disease is not completely understood, lymphocyte therapy may have a role in causing the disease. In other studies that examined mother problems in causing the disease, relationship between diabetes and this disease was confirmed.

In VACTERL patients, all aspects of possible anomalies should be considered. Here, additional cases of disease may be found that sometimes there is inevitable need for faster treatment. It seems that in study of the cause of disease, in addition to genetic factors, motherhood issues should be considered. Lymphocyte therapy to induce pregnancy may have a relationship with cause of the disease; further studies are required to prove.

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REFERENCES


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