Prenatal Diagnosis of Isolated Hypospadias by Using “Tulip Sign” with Two and Three-Dimensional Ultrasonography

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ABSTRACT

Hypospadias is an anomaly of the male urogenital tract which occurs as a result of deficient closure of the urethral groove. Hypospadias is the most commonly seen urogenital anomaly in male neonates, but it is rarely diagnosed during prenatal period. Herein, a severe hypospadias case which was diagnosed by using 2D (two-dimensional) and 3D (three-dimensional) ultrasonography at 28 weeks of pregnancy with the aid of “tulip sign” finding was presented. 2D ultrasonography of a 29 year old, primigravid woman at 28 weeks of gestation revealed an incurved, short penis with blunt tip located between the two scrotal folds and there was no additional anomaly. Findings were compatible with the “tulip sign” and case was diagnosed as isolated hypospadias. Isolated hypospadias was confirmed after delivery and surgical correction was planned. Prenatal diagnosis of hypospadias and relevant literature has been reviewed.

Keywords: Hypospadias, Ultrasonography, Prenatal diagnosis, Tulip sign


Introduction

Hypospadias is a congenital defect of male genitalia, which is characterized by the opening of the urinary tract on the ventral side of the penis instead of the tip. It is the most commonly seen urogenital anomaly in male neonates with an estimated prevalence of 0.20%-0.30% in live births.1,2 However, it is rarely diagnosed during prenatal period.

Prenatal diagnosis of hypospadias with other anomalies or without has been reported before.3-6 Meizner et al. presented a specific sonographic indication of severe hypospadias and termed this finding as “tulip sign”.7

Although ultrasonographic prenatal diagnosis of hypospadias is known for a long time and hypospadias associated with other anomalies are generally diagnosed prenatally without difficulty, there are a few case reports about prenatal diagnosis of isolated hypospadias cases. We presented a severe hypospadias case which was diagnosed by detecting “tulip sign” at 28 weeks of pregnancy by using two and three dimensional ultrasonography. Despite the fact that fetal gender determination is done almost routinely in Turkey, this is the first prenatal diagnosis of isolated hypospadias case as far as is known in the country. Also, it is one of the few case reports about prenatal diagnosis of isolated hypospadias by using “tulip sign” finding after it was described by Meizner et al.7

Case Report

A 29 year old, primigravid, healthy woman with no significant past medical and obstetric history was admitted for ultrasonographic evaluation at 28 weeks of gestation. Fetal growth was compatible with 28 weeks of gestation. Two-dimensional ultrasonography revealed an incurved, short penis with blunt tip located between the two scrotal folds (Figure 1A and B). There was no additional abnormality in the detailed sonographic examination of the fetus. 3D sonographic evaluation was done to better define the fetal external genitalia and to confirm the diagnosis (Figure 2). Ultrasonographic external genitalia findings resembled the “tulip sign” and the case was diagnosed as isolated hypospadias.

A male newborn weighing 2520 grams was delivered by cesarean section because of intrauterine growth restriction and fetal distress at 38 weeks. Hypospadias was confirmed after delivery and additional anomaly was not detected (Figure 3). Postnatal karyotyping revealed 46, XY. The baby is doing well and waiting for surgical correction.
Hypospadias is a developmental anomaly of the male urethra and characterized by the location of urethral meatus on the ventral side of the penis instead of the tip. Hypospadias develops as a result of a failure of closure of the urethral groove. The urethra may terminate between the proximal glans and perineum and is called as glandular, penile, penoscrotal, or perineal hypospadias according to termination site. Estimated prevalence of hypospadias is 0.20%-0.30% in live births.\(^1,2\) Hypospadias is seen more commonly in whites and in monozygotic twins.\(^8,9\) Hypospadias is usually seen as an isolated abnormality. However, prenatal diagnosis of hypospadias is commonly done in cases associated with other anomalies or dysmorphic syndromes.\(^4,10-12\) Prenatal diagnosis of hypospadias is usually made in late second or third trimester of pregnancy.

Hypospadias is one of the most common urogenital anomalies, but diagnosis is often missed before birth. Ultrasonographic features of hypospadias was described before as; abnormal ventral curvature of the distal penis, small penis with ventral incurring, small lateral folds, abnormal distal morphology of the penis, abnormal fetal micturition.\(^5,6,13\)
Meizner et al. presented a specific sonographic appearance of severe hypospadias and termed this finding as “tulip sign” and they gave sonographic markers for prenatal diagnosis of hypospadias as following:
- Absence of the normal pointed morphology of the penile shaft with blunted tip.
- Abnormal curvature of the penis.
- A small penile shaft.
- Two parallel echogenic lines corresponding to lateral folds of the dermal remains of the prepuce.
- Ventral deflection of urinary stream.”

Three of these findings were observed by using ultrasound in this case. These findings were abnormal curvature of the penis, small penile shaft and penile shaft with blunted tip. Multiplanar and surface-rendered images of the external genitalia were acquired in the axial, coronal and midsagittal planes (Figure 2). Three dimensional ultrasound examinations in the surface rendered mode allowed us to establish more distinct appearance of external genitalia and to confirm the diagnosis of hypospadias. According to Meizner et al. the “tulip sign” represents the most severe form of hypospadias. Therefore; they proposed that the “tulip sign” may help to distinguish between severe hypospadias and other forms of genital abnormalities. When hypospadias repair surgery is performed at an early stage, the cosmetic and functional results are excellent. Therefore, correct diagnosis of severe hypospadias by using the “tulip sign” will reassure the parent especially in ambiguous genitalia cases.

Only one publication could be found about prenatal diagnosis of hypospadias in our country. In this publication, Aslan et al. reported prenatal diagnosis of Wolf-Hirschhorn syndrome (4p-) in association with congenital hypospadias and foot deformity. Ultrasonographic appearance of hypospadias was defined as “irregular distal penis” in this case report. In our country, where families are very curious about the gender of fetus, fetal gender determination is done almost routinely during prenatal ultrasound examination. It is very interesting that despite almost routine fetal gender determination, this is the first prenatal diagnosis of isolated hypospadias case as far as is known in our country. Also, only one case report could be found about “tulip sign” and prenatal diagnosis of hypospadias by ultrasonography in the literature. The most probable reason for this is that ultrasonography of genitalia include sex determination but not anomaly detection. According to Devesa et al.; the purpose of prenatal evaluation of external genitalia should not be only to determine the sex, but also to detect anomalies. Therefore, targeted ultrasound should include fetal sex determination and external genitalia evaluation of anomalies.

Hypospadias may be associated by some syndromes and chromosome abnormalities. Therefore, prenatal diagnosis is important to plan karyotyping, to look for association with any possible dysmorphic syndromes and to reassure the parents. Three dimensional ultrasound may improve the diagnostic accuracy by enabling direct visualization of hypospadias and should be used whenever possible. Prenatal ultrasound should include not only fetal sex determination but also detection of abnormal genitalia.

References