DIPHALLUS: A REPORT OF AN UNUSUAL CASE

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ABSTRACT

We report an unusual case of diphallus in a five year old boy. The two organs appeared separately, one orthotopically in the pubic area and the other in the perineum with no urinary meatus. A perineal hypospadias with meatus in the anterior anal verge was also present. The right testis was undescended and the scrotum was mildly bifid. The patient was continent for both voiding and defecation. Micturition was via the hypospadiac perineal meatus under voluntary control. A review of published cases suggests that this may be the first example of complete diphallia with two separate phalli in different locations.

Keywords: Diphallus, Pseudodiphallia, Duplication of penis, Perineal hypospadias.


INTRODUCTION

Diphallia, also referred to as duplication, is a rare congenital anomaly, as there have been less than 100 reports of diphallus in the literature.16 It has been estimated that in the United States its incidence is one per every 5 million live births. The first documented report of this anomaly dates back to 1602.20 Associated defects are common, including hypospadias, epispadias, bifid scrotum, vesical extrophy, diastasis pubis, absence or duplication of the bladder and urethra, renal agenesis, ectopia and horseshoe kidneys, duplication of the hindgut and spinal dysraphism.

Schneider proposed three different groups: (1) bifid glans, (2) bifid phallus, and (3) complete diphallia or double penis. The term pseudodiphallia proposed by Villanova and Raventos includes cases exhibiting an accessory ectopic penis.

According to Abdel, two main groups exist, true diphallia (partial and complete) and bifid phallus (partial and complete—bifid glans, bifid body).6

Our case does not fit any of these classifications and may be the first example of complete diphallia with two separate phalli in different locations.

Case Report

A five year old boy was referred to our center because of bizarre genitalia. There were two separate phalli, one

Fig. 1. Two distinct phalli are shown.
Diphallus

The left testis was normal in size and consistency, but the right testis was undescended. The patient was continent for both voiding and defecation day and night. Micturition was via the perineal meatus under voluntary control. Physical examination of the heart, lungs, limbs, spine and CNS was normal. Rectal examination revealed normal anal sphincter tonicity and an intact prostate gland.

Family history was negative for any congenital anomalies. Furthermore, the patient’s mother had no infections, nor had she been exposed to any. She also denied any drug consumption during an uneventful pregnancy. Chromosomal study revealed normal X chromosome and appropriate sexual chromatin.

Evaluation of the patient’s upper urinary tract with excretory urogram revealed normal kidneys and ureters with no ectopia or fusion and no osseous abnormalities in vertebrae or pelvic bones (Fig. 3).

Retrograde cystogram showed normal bladder with no evidence of vesicoureteral reflux (Fig. 4). On endoscopy there was a straight narrow tube as the posterior urethra with no evidence of verumontanum.

Ultrasound study of the liver, pancreas and adrenals was normal. Chest x-ray disclosed no abnormalities.

After explaining the situation of the patient to his parents, we asked them for evidence of erection. His mother reported nocturnal tumescence in both phalli.

Surgical exploration was performed, and revealed that there were two corporal tissues within the ectopic phallus which terminated in the perineum. The ectopic phallus was excised completely and a right orchiopexy was done at the
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Fig. 5. Picture showing the patient after ectopic phallic excision and right orchiopexy.

Fig. 6. Low-power view of the specimen demonstrating corporal tissue in the upper left, with skin layers in the lower right side of the picture.

same time (Fig. 5).

The anatomopathologic report was as follows: Gross specimen covered with skin measured 5 cm long and 1 cm wide. A glans penis was located at one end, and corpora cavernosal bodies were seen on cross section. Microscopic examination showed distinct dilated vessels with fibroelastic wall embedded in fibromuscular tissue which was totally surrounded by skin. The diagnosis was diphallus or double penis.

We plan to perform urethroplasty with bladder mucosal graft after six months.

CONCLUSION

Review of literature revealed that complete diphallus is a very rare anomaly. Some of the previous reports of this anomaly were unique in some aspects; i.e., association with a third urethra, duplicated bladder, spinal dysraphism, etc. We believe that our case may be the first example of a complete diphallus locations.

REFERENCES

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