Case Report

Cryptorchidism is a Useful Clue for Idiopathic Hypogonadotropic Hypogonadism in Pituitary Stalk Thickening

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Abstract

Pituitary stalk lesions can represent a wide range of pathologies. The exact cause is often unknown due to hesitancy to proceed with biopsy. We present a 16-year-old adolescent who presented with delayed puberty, short stature and bilateral cryptorchidism. He was found to have a thickened pituitary stalk of uncertain etiology with partial hypopituitarism (gonadotrophin and growth hormone deficiency) on further assessment. The presence of bilateral cryptorchidism and micropenis represents lack of "mini puberty," a phenomenon of activation of the hypothalamic-pituitary-gonadal (HPG) axis in-utero or within the first few months of life.1 2 These key clinical features have been useful to establish an early temporal relationship and suggest a congenital origin of disease. This enabled a more conservative approach of surveillance to be employed as opposed to invasive pathological examination with pituitary stalk biopsy.

Key words: pituitary disease, hypopituitarism, cryptorchidism, growth hormone, hypogonadotrophic hypogonadism

INTRODUCTION

Pituitary stalk lesions comprise a broad spectrum of diseases that impose a great challenge to the treating physician due to its critical location and pivotal role in the hypothalamic-pituitary (HPA) axis. This proves more of a conundrum in children and adolescents whereby a prompt diagnosis is pertinent to avoid long term repercussions of disease progression and hypopituitarism. The presence of hypopituitarism associated with thickened pituitary stalk implies compromise to the function of the gland which may be permanent or reversible in some cases depending on the etiology and duration of disease. The exact etiology of these lesions often unknown despite rigorous work up, typically warranting a pituitary stalk biopsy as the next step of management. In many cases, there is reluctance to proceed with pathological examination due to the risk of complications. Therefore, the presence of specific clinical features in addition to radiological and laboratory investigations is critical to establish the temporal relationship of insult to the HPA axis and determine possible origin of disease. Bilateral cryptorchidism and micropenis which represents lack of ‘mini puberty’ are key features supporting an early insult to the hypothalamic-pituitary-gonadal (HPG) axis in patients with hypogonadotrophic hypogonadism.1 2

CASE

We present a 16-year-old boy referred from urology for further evaluation of hypogonadotropic hypogonadism. He was initially diagnosed with bilateral cryptorchidism aged 2, and his parents opted for conservative management. There was no spontaneous descent of testes in the next few years and he was eventually lost to follow up. He was referred again to urology aged 12, for recurrent urinary tract infections and was diagnosed with vesico-ureteric reflux (VUR) warranting bilateral ureteric stenting and implantation. There were no other structural or functional abnormalities of the renal system identified. Cryptorchidism was reevaluated at this point and he underwent bilateral orchidopexy, with successful descent only on the left side. Routine clinical and hormonal assessment done at this juncture revealed hypogonadotrophic hypogonadism, warranting an endocrine consult.

Detailed history taking from parents and physical examination elicited the following salient clinical features. He was born full-term with an uneventful antenatal history. His height and weight remained within the 25th and 50th centile till the age of 10. A decline in growth rate was noticed as he approached adolescence, whereby he was one of the shortest amongst his peers in secondary school. He had been a slow learner, with poor academic performance since primary school. His parents had noticed him having low energy levels and fatigue in the later years. No history of anosmia or hearing deficits was elicited. He had no history of polyuria or polydipsia to suggest cranial diabetes insipidus. A full systemic review did not reveal symptoms to suggest intra-cranial mass effect such as chronic headaches, visual disturbances or neurological deficits. He had no past history of intracranial pathology, trauma or radiation. Apart from the history of VUR and cryptorchidism, he did not suffer from any chronic diseases nor receive long term immunosuppression/steroids or chemotherapy. He has no family history of short stature nor delayed puberty. He has one younger sibling, a 12-year-old female who has achieved puberty and is of appropriate height. His anthropometric measurements of weight and height were charted as 58 kg (between 5-10th weight and height were charted as 58 kg (between 5-10th centile till the age of 10. A decline in growth rate was noticed as he approached adolescence, whereby he was one of the shortest amongst his peers in secondary school. He had been a slow learner, with poor academic performance since primary school. His parents had noticed him having low energy levels and fatigue in the later years. No history of anosmia or hearing deficits was elicited. He had no history of polyuria or polydipsia to suggest cranial diabetes insipidus. A full systemic review did not reveal symptoms to suggest intra-cranial mass effect such as chronic headaches, visual disturbances or neurological deficits. He had no past history of intracranial pathology, trauma or radiation. Apart from the history of VUR and cryptorchidism, he did not suffer from any chronic diseases nor receive long term immunosuppression/steroids or chemotherapy. He has no family history of short stature nor delayed puberty. He has one younger sibling, a 12-year-old female who has achieved puberty and is of appropriate height. His anthropometric measurements of weight and height were charted as 58 kg (between 5-10th
representing gonadotroph deficiency early in life coupled with non-suspicious laboratory and imaging workup was highly suggestive of congenital origin of disease. Therefore, a decision was made not to proceed with pituitary stalk biopsy unless there is evidence of disease extension or progressive loss of pituitary function. Growth hormone, norditrophin at a dose of 0.025 mcg/kg/day was initiated in this boy aiming to achieve near adult height. He showed good response to therapy with a 4 cm increment in height after 3 months. Puberty induction will be initiated once acceptable near adult height is achieved. Serial MRI pituitary and pituitary hormone panel will be repeated every 6 months to monitor progression of disease. He is also planned for removal of the right testis due to risk of malignancy and poor function.

As an alternative test, Glucagon Stimulation Test (GST) was chosen. GST showed a peak GH of 6.94 ng/ml (>10 ng/ml) at 90 mins confirming his growth hormone deficiency. Bone age assessment showed his skeletal age to lag at the age of 12-13 years with unfused epiphyseal plates. A MRI pituitary ordered showed an enlarged pituitary stalk extending to the tuber cinereum with homogenous enhancement post gadolinium contrast. The stalk measured 13 mm (AP diameter) at the point of insertion at the infundibulum with uniformed pattern of thickening. The pituitary gland itself was not hypoplastic, with no focal enhancement on post contrast study. The optic nerve, chiasm and tract was not thickened, with no evidence of compression. Therefore, at this point it was concluded that this patient had thickened pituitary stalk with partial hypopituitarism (hypogonadotropic hypogonadism and growth hormone deficiency).

The etiology of thickened pituitary stalk was worked up extensively involving laboratory and radiological investigations. His complete blood count showed no evidence of hematological dyscrasias. His liver and renal biochemistry was normal. Both inflammatory and tumor markers were not raised. There was no evidence of mediastinal or lung mass on chest x-ray. Ultrasound showed very small testes in the right inguinal region and left scrotum measuring 1.2-1.5 cm, both testes had no suspicious malignant features or associated lymphadenopathy. In summary, there were no red flags to suggest an inflammatory or neoplastic process. A multidisciplinary discussion was held to discuss the role of pituitary stalk biopsy for him. The presence of pre-pubertal features representing gonadotroph deficiency early in life coupled with non-suspicious laboratory and imaging workup was highly suggestive of congenital origin of disease. Therefore, a decision was made not to proceed with pituitary stalk biopsy unless there is evidence of disease extension or progressive loss of pituitary function.

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| Table 1. Anterior pituitary hormone panel |
| Hormone | Result | Reference range |
|-----------|--------|-----------------|
| FSH | 2.2 IU/L | 1.5-12.9 |
| LH | 0.58 IU/L | 1.3-9.8 |
| Testosterone | <0.1 nmol/L | <28.8 |
| IGF-1 | 273 ug/L | 287-673 |
| Prolactin | 196.8 mIU/L | 86-324 |
| Am cortisol | 569.9 nmol/L | 145.4-619.4 |
| Thyroid function test | TSH 2.94 mIU/L, Free T4 15.76 pmol/L | TSH 0.52-4.30, T4 12.8-21.0 |

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DISCUSSION

The etiology of thickened pituitary stalks is often broadly divided into 3 categories: neoplastic, inflammatory or infectious and congenital. Two large case series based on retrospective review involving both adults and children with pituitary stalk lesions have showed neoplastic lesions to be the leading cause of pathology. A similar finding was found in a study done in Korea, specifically in the pediatric population below the age of 18 with pituitary stalk lesions. However, Turcu et al., also found 39% of pituitary stalk lesions in a retrospective review with unclear pathology. The role of pituitary stalk biopsy often remains controversial due to the risk involved. Despite using a minimally invasive endoscopic approach, the procedure, coupled with manipulation of the pituitary stalk, imposes a risk of cranial diabetes insipidus, hypopituitarism, cerebrospinal fluid (CSF) leak and meningitis in patients. In addition, there is approximately a 10% risk of a negative biopsy, whereby a diagnosis is not histologically conclusive despite adequate tissue sample.

Radiological features on MRI is often the key in guiding the diagnosis of pituitary stalk lesions. It is also an essential tool to monitor progression of disease or response to treatment in these patients. Specific patterns of enhancement such as uniform, V-shaped, round or diamond, and pyramidal has been associated with various pathologies in literature. The strongest association found was between the congenital lesions and round pattern of enhancement in the Mayo experience by Turcu et al. Extent of lesions was also associated with hypopituitarism, and the lesions with hypothalamic extension have been found to have the highest risk of hypopituitarism.

Congenital cryptorchidism is one of the most common congenital malformation in boys. Its prevalence at birth among boys with a birth weight more than 2,500 g has been reported to range between 1.8% and 8.4%. It is associated with reduced concentration of testosterone and sperm quality in adulthood. There are numerous causes for congenital cryptorchidism including disorders involving sex chromosomes, gonadal development, decrease in androgen synthesis and action, structural defects and hypogonadotropic hypogonadism.

Typically, pituitary gonadotropes start producing FSH and LH around 9 weeks of gestation, reaching their peaks at mid-gestation with a concomitant rise in testosterone. This peak of testosterone is essential for descent of testes in utero. The activation of hypothalamic-pituitary-gonadal (HPG) axis occurs approximately 1 week after birth with a second peak at 1–3 months, a period also synonymous with ‘mini-puberty’. This post-natal surge is essential for testicular descent and penile growth. In those with congenital hypogonadotropic hypogonadism, the lack of gonadotrophins results in the arrest of testicular descent and penile growth in utero and in the neonatal period.

CONCLUSION

In our patient, the lack of ‘mini puberty’ described above suggests the insult to the HPG axis occurred early in life, pointing towards a congenital origin of disease. Despite the radiological features not being typically described in literature for congenital lesions, the extension of the lesion to the hypothalamus accounts for the development of partial hypopituitarism in this patient. In addition, the negative results for both inflammatory and neoplasm workup further supports this diagnosis. With congenital disease as our probable diagnosis, we have chosen a more conservative approach of management with surveillance as described above, alleviating the need for pituitary stalk biopsy at the moment.

Ethical Consideration

Patient consent was obtained before submission of the manuscript.

Statement of Authorship

All authors certified fulfilment of ICMJE authorship criteria.

Author Disclosure

The authors declared no conflict of interest.

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