CASE REPORT

Primary Partial Empty Sella presenting with Prepubertal Hypogonadotropic Hypogonadism: A Case Report

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Abstract

Primary partial empty sella occurs when less than 50% of an enlarged or deformed sella turcica is filled with cerebrospinal fluid. Its diagnosis is confirmed with magnetic resonance (MR) study of sellar and suprasellar regions or computed tomography (CT) for those with absolute contraindication/s to MR.1

Based on etiology, empty sella can be classified as primary or secondary. Compared to secondary empty sella which is caused by pituitary pathological conditions like previous surgical, pharmacological or radiotherapy treatment, the pathogenesis behind primary empty sella (PES) is unclear.2 Some of the identified mechanisms include incomplete formation of sellar diaphragm and the influence of suprasellar or pituitary promoting factors.1 Its incidence is not that high, ranging from 5.5%-12% as accidental finding in autopsy to 8%-35% in clinical practice. Majority of the prevalence of PES is among women – those with history of at least one completed pregnancy in their physiological history. Moreover, its occurrence in children is also less frequent compared to adults, and it is more or less associated with hypothalamic–pituitary dysfunction, genetic disorders or perinatal complication.2

Key words: PES, hypogonadotropic, hypogonadism, micropenis

INTRODUCTION

Empty sella is a radiologic finding pertaining to an enlarged or deformed sella turcica which can be partially (<50%) or completely (>50%) filled with cerebrospinal fluid.1 Its diagnosis is confirmed with magnetic resonance (MR) study of sellar and suprasellar regions or computed tomography (CT) for those with absolute contraindication/s to MR.2

Based on etiology, empty sella can be classified as primary or secondary. Compared to secondary empty sella which is caused by pituitary pathological conditions like previous surgical, pharmacological or radiotherapy treatment, the pathogenesis behind primary empty sella (PES) is unclear.2 Some of the identified mechanisms include incomplete formation of sellar diaphragm and the influence of suprasellar or pituitary promoting factors.1 Its incidence is not that high, ranging from 5.5%-12% as accidental finding in autopsy to 8%-35% in clinical practice. Majority of the prevalence of PES is among women – those with history of at least one completed pregnancy in their physiological history. Moreover, its occurrence in children is also less frequent compared to adults, and it is more or less associated with hypothalamic–pituitary dysfunction, genetic disorders or perinatal complication.2

CASE

RC, a 20-year-old male, sought consult in our institution for small penile size. He was born full-term via normal vaginal delivery to a 35-year-old G3P3 mother, with no perinatal complications. However, there was intake of an abortifacient (4 tablets of misoprostol) on the 5th week of pregnancy. At birth, there was noted “monggo seed” sized penis and “paper-thin” scrotum. His mother was advised by the local health center physician to seek consult with a surgeon however due to financial difficulties, there were no tests nor consult with a specialist done. His childhood and pre-puberty years were unremarkable. There were no symptoms of palpitations, heat or cold intolerance, polyuria, polydipsia, polyphagia. There were no episodes of frequent urinary tract infection, dysuria, or abdominal pain. There were no instances of elevated blood pressure, headache, change in vision. There were no significant health problems during his childhood except for appendicitis for which he underwent an emergency appendectomy in July 28, 2008 at Ospital ng Makati.

However, at around 14 to 16 years of age, there was persistence of high pitched voice, scant pubic and axillary hair and fat deposition on waist and bilateral breasts. There was minimal growth in penile size at approximately 2-3 centimeters (cm) and a bilateral palpable scrotal sac.

On physical examination, he was ambulatory with stable proportion (Figure 1) with a height of 162 cm and arm span of 171 cms. His mother stands 151 cm while his father at 169 centimeters (cm) and a bilateral palpable scrotal sac.

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Micropenis is defined by Schonfeld and Beebe as Stretched Penile Length (SPL) 2.5 SD less than the mean for age without the presence of any other penile anomalies and presence of internal and external genital organs compatible with a 46 XY karyotype. For an average adult patient, mean stretched penile length is 13.3 cm with 9.3 cm as the calculated 2.5 standard deviation less than the mean for age. The patient has a stretched penile length of 3 cm falling more than 2.5 SD below the mean for adult.

To satisfy the criteria for micropenis, pelvic and inguino-scrotal ultrasound was done to confirm the presence of internal genital organs, which revealed small left testicle measuring 0.9x0.5x0.7 cm with chronic parenchymal cm with computed mid parental height for boys of 166.5 cm. Thyroid gland was not palpable. He has gynecomastia without galactorrhea (Figure 2). Genital and pubic hair development was graded as Tanner Stage 1 (Figure 3). He has a flaccid and stretched penile lengths of 2.5 and 3 cm respectively with width of 4 cm. He has palpable small, firm left testis while non palpable on the right. Neurologic examination was normal except for bilateral anosmia. Evaluation was done by Otorhinolaryngology service which showed recurrent rhinosinusitis. CT scan of paranasal sinuses showed pansinusitis with opacified and widened ostiomeatal units. A trial of steroid therapy was recommended which provided slight relief of anosmia.

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hypogonadism, where the growth plates are not yet fused and there is lack of sex steroids, growth plates of the extremities continue to grow past the usual age of cessation. As a result, there is decreased upper to lower ratio and an increased arm span for height leading to eunuchoid proportion like in the patient.

Aside from this, determination of skeletal development by x-ray of left hand and wrist is also a useful way of establishing the stage of physiological development which in some cases may not be parallel with chronological age. The patient has a delayed bone age (compatible to 16 and 5/12 to 14 and 5/12 year old male) by the Greulich-Pyle method (Figure 4). Estradiol is a product of aromatization of testosterone and it mediates additional effects of testosterone on bone resorption, epiphyseal closure, sexual desire, and fat deposition. From the physical examination and the patient’s bone x-ray, this low estradiol might have contributed to the unfused ossification centers and eunuchoid proportion. Related to this is that androstenedione is converted to testosterone by 17-beta hydroxysteroid dehydrogenase before it gets aromatized to estradiol. However, androstenedione itself can be aromatized to estrogen which might have contributed to gynecomastia and fat deposition.

The patient’s cranial MRI showed a shallow sella with apparent flattening of pituitary gland at its floor (Figure 5). The results of baseline endocrine tests are summarized in

changes and a non visualized right testicle. The prostate gland measures 2.2x1.6x1.7 cm (~3 grams). A whole abdominal CT scan was done with noted left inguinal hernia and ovoid soft tissue density in right inguinal region possibly representing the right testis. Other findings include hepatic steatosis and nephrocalcinosis on the left. Lastly, to satisfy the criteria, chromosome analysis was done revealing a karyotype with no numerical and structural aberrations and an XY sex chromosome complement in all 50 cells examined. Hence the patient has a male karyotype of 46,XY.

However, the patient did not just present with isolated micropenis. Alongside, he has gynecomastia, persistently high pitched voice and underdeveloped adult sexual characteristics. Most authorities accept the definition of delayed puberty as the absence of secondary sexual development at an age 2 SD above the mean age of onset of puberty. This is the age at which 95% of normal children have already entered puberty. Based on etiology, pubertal delay can be classified into constitutional growth delay or hypogonadism. The latter can be further classified into hypogonadotropic or hypergonadotropic hypogonadism.

Though stature is the most obvious change in growth, the ratio of the upper and lower segment also changes significantly. Sex steroids are necessary for increase in growth hormone secretion and they directly stimulate epiphyseal plate’s growth and fusion. In prepubertal hypogonadism, where the growth plates are not yet fused and there is lack of sex steroids, growth plates of the extremities continue to grow past the usual age of cessation. As a result, there is decreased upper to lower ratio and an increased arm span for height leading to eunuchoid proportion like in the patient.

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The patient’s cranial MRI showed a shallow sella with apparent flattening of pituitary gland at its floor (Figure 5). The results of baseline endocrine tests are summarized in
A 24-year-old, 46XY patient was admitted for evaluation of infertility and lack of development of secondary sex characteristics. Hypogonadism was only observed in 2 patients. (20.8% n=5, N=24) and isolated hypogonadotropic hypogonadism was observed in minority (18.75%, n=3) with endocrine problem was also seen in 1 patient, which presented as amenorrhea and erectile dysfunction. 6 Hypogonadism as the most common endocrine problem (19%, n=4, N=21) had documented endocrine abnormalities in which only 22% (n=9, N=40) are male. 4 This marginal number of male patients with PES presenting with endocrine problem was also seen in a study done by Maira et al., where in only 12% was documented. 5

Table 1 showing hypogonadotropic hypogonadism (low testosterone, FSH and LH). Other biochemical results are summarized in Table 2 which showed normal fasting blood sugar, serum sodium, potassium, AST and ALT. While, lipid profile showed normal levels of total cholesterol and triglyceride but elevated LDL.

With these findings, the patient was diagnosed with Primary Empty Sella (PES) (Partial) which is probably congenital based on history of abortifacient use on first trimester of pregnancy, findings of prepubertal hypogonadotropic hypogonadism and MRI showing a partially empty sella.

He was referred to Endocrinology and Urology services. Sex hormone replacement was recommended. He was offered testosterone therapy however this was not yet started due to lack of funds. Steroid was not initiated since serum cortisol is just borderline low and patient is also asymptomatic. Orchietomy as prophylaxis for development of malignancy is not warranted but orchidopexy may be offered to monitor tumor development.

Counseling was offered regarding possible psychosocial impact of the physical changes brought about by the endocrine problem. Since he has a partner it was also important to discuss the impact of the physical changes brought about by the hormonal deficiencies on other organ systems, the patient was endorsed to the succeeding endocrinology rotator and outpatient resident. He was last seen on December 2019 and repeat blood chemistries were requested; however, he was lost to follow up thereafter.

**DISCUSSION**

Data on epidemiology of PES varies based on means of diagnosis. It is usually an incidental finding in 5.5%-12% of autopsy cases. However, using neuroimaging, its overall incidence has been estimated at 12%, while approximately 9-35% if based on clinical findings as reported in various case series. Its female-to-male ratio is 5:1 with peak incidence occurring at 30 to 40 years, occasionally earlier in women. It occurs less frequently in children and is associated with hypothalamic–pituitary dysfunction, genetic disorders or perinatal complications.

The etiology of PES is unclear but some of the etiopathogenic hypotheses identified include: 1. incomplete formation of sellar diaphragm. 2. upper sellar factors (persistent or intermittent intracranial idiopathic hypertension, CSF pulsatility, obesity, systemic hypertension) or 3. pituitary factors (conditions associated with variation of pituitary volume like pregnancy, lactation, menopause, hypophysitis, compensatory pituitary hypertrophy to primary hormonal defect).2

Patients with PES have varied symptoms, and endocrine dysfunction is one the least common presenting manifestations. In a study done by De Marinis et al., only 19% (n=40, N=213) had documented endocrine abnormalities in which only 22% (n=9, N=40) are male. 4 This marginal number of male patients with PES presenting with endocrine problem was also seen in a study done by Maira et al., where in only 12% was documented. 5

The prevalent endocrine problem varies in different studies. In a study done by Radha Rani et al., hypocortisolemia was most common (62.5%, n=10, N=16). In the same study, hypogonadism was observed in minority (18.75%, n=3) which presented as amenorrhea and erectile dysfunction. 6

On the other hand, in a study done by Ghatnatti et al., hyperprolactinemia was the most common dysfunction (20.8%, n=5, N=12) and isolated hypogonadotropic hypogonadism was only observed in 2 patients. 7

One study on PES showed hypogonadotropic hypogonadism as the most common endocrine problem (19%, n=5; N=21) which presents as oligomenorrhea in females and decreased sexual function in males. 8 Micropenis and lack of secondary sex characteristics are rare presentations since PES is seldom seen during prepubertal years. Its peak incidence is notable at postpubertal years hence the usual clinical manifestations of decrease in sexual function or erectile dysfunction are observed. One event was documented in 1973 which is almost similar with our case. A 24-year-old, 46XY patient was admitted for evaluation of infertility and lack of development of secondary sex hormone replacement was recommended. He was last seen on December 2019 and repeat blood chemistries were requested; however, he was lost to follow up thereafter.

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characteristics. He is obese with eunuchoid proportion given the height and arm span. He has gynecomastia, bilateral undescended testes, normal prostate and decreased male body hair, however data on micropenis was not mentioned.9

The treatment for these patients entails hormonal replacement. For this case, testosterone therapy may not improve fertility at this point but it may help produce and maintain virilization and prevent future complications of hypogonadotropic hypogonadism like osteoporosis and cardiovascular problems. Lastly, semen analysis and other hormone dynamic tests are recommended to evaluate fertility potential and other possible hormonal problems respectively.10

CONCLUSION

Primary empty sella itself is a rare disease entity. Presenting as its manifestation, hypogonadotropic hypogonadism is also uncommon, mostly seen late at 30 to 40 years of age and majority in females. Prompt recognition of prepubertal hypogonadotropic hypogonadism at an early age can maximize both surgical and medical management in these patients. However, since the patient in this case sought consult late, it is also important to reiterate at this point alternative options if there are future plans for reproduction as well as methods to prevent future complications of these hormonal imbalances.

Ethical Considerations

Patient consent was obtained before submission of the manuscript.

Statement of Authorship

Both authors certified fulfillment of ICMJE authorship criteria.

Author Disclosure

Both authors declared no conflicts of interest.

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