**OBJECTIVES**

To report a case of hypogonadism reversal and spontaneous puberty in a craniopharyngioma patient who suffered from hypopituitarism as a result of complete surgical resection.

**METHODS**

A 13-year-old boy with hypopituitarism was evaluated for right testicular sensitivity. At the age of 6 the child presented with growth arrest and a craniopharyngioma causing pituitary insufficiency was diagnosed. Treatment with hydrocortisone and thyroxine supplementation was initiated and he was subjected to a transphenoidal surgery that resulted in complete tumor resection (Figure 2). Tumor histology indicated an adamantinous craniopharyngioma, which constitutes the predominant type occurring in children and adolescents. Postsurgical biochemical testing confirmed panhypopituitarism and he was prescribed levothyroxine, hydrocortisone and desmopressin supplementation. Recombinant growth hormone was initiated 2 years following surgery, since post surgical MRI did not depict any sign of pituitary tissue and growth had not resumed (Figure 1). Gonadotropins had been undetectable since diagnosis and the last evaluation had confirmed prepubertal status (Table 1). On clinical examination increased testicular volume was found (10x12mm) with clinical puberty and mild sensitivity was confirmed. The Scrotum ultrasound revealed normal appearing testes with a maximal diameter of 4.5cm left and 4cm right. Testosterone and adrenal androgen levels were below normal limits but a GnRH stimulation test (2.5μg/kg) provoked a fourfold increase of LH (Table 1). Onset of puberty was suspected.

**RESULTS**

Six months later, the onset of puberty was confirmed, by measurement of gonadotropins (FSH = 1.4 U/L, LH = 1.8 U/L, testosterone (98 ng/dl) and inhibin-B (216 pg/ml). The GnRH stimulation test provoked a tenfold increase in LH. The results of biochemical testing combined with the testicular volume, compared to normative values, indicated the onset of puberty in our patient (Table 2). The patient’s treatment regarding his other hormonal deficits was continued. However, the testes enlargement together with the LH increase suggested a possible restoration of the gonadal axis, thus rendering testosterone supplementation unnecessary. Proper follow-up was scheduled to evaluate adequacy of gonadal steroid production.

**CONCLUSIONS**

Recovery of pituitary function in craniopharyngioma patients who had panhypopituitarism both prior and after complete surgical resection of the tumor has only rarely been reported. Nevertheless, this case depicts that one should consider the possibility of a functional gonadal axis prior to steroid supplementation for puberty induction to children suffering from hypopituitarism due to craniopharyngioma diagnosed and treated before the onset of puberty.

**REFERENCES:**