Identified mutations in CYP11B1 gene in two Tunisian patients with 11-beta hydroxylase deficiency

H Ennaifer1, I Kammoun1, M Kharrat2, N Cheikhrouhou4, M Khatib1, M Trabelsi2, C Ben Slama1

1-Department of endocrinology and metabolic diseases, National Institute of Nutrition, Tunis, Tunisia
2-Laboratory of Human Genetics, University of medicine of Tunis, Tunisia
3-Department of congenital and hereditary diseases, Charles-Nicole Hospital, Tunis, Tunisia

Introduction:

11-Beta hydroxylase deficiency (11-OHD), a rare autosomal recessive disorder, is caused by CYP11B1 mutations. The incidence of 11-OHD in overall population is approximately 1 in 100,000–200,000 [1]. Based on clinical manifestations, 11-OHD is classified as classic and non-classic forms.

We studied the mutations of CYP11B1 gene in two patients with classic 11β-OHD.

The hypothesis of 11β-OHD deficiency was considered and confirmed by genetic exploration. A non-sense mutation p.G379V of the CYP11B1 gene was found.

Discussion:

In classic 11-OHD, a reduction of adrenal cortisol synthesis leads to elevated plasma ACTH levels, which result in an increased production of cortisol precursors in zona fasciculate. Therefore, the androgen synthesis is increased and hyperandrogenemia appears. Moreover, the increased ACTH secretion also contributes to higher levels of deoxycortisone (DOC) and 11-deoxycortisol [2]. Thus, classic 11-OHD is characterized by severe virilization in newborn females and precocious pseudopuberty in both sexes. The DOC overproduction causes hypertension and hypokalemia.

Non-classic 11-OHD is usually characterized by slight increase of serum androgen, mild hirsutism and irregular menses. However, elevated blood pressure is rarely observed in the mild form. CYP11B1 gene consists of nine exons and encodes a protein of 503 amino acids. CYP11B1 gene is located on chromosome 8q22, approximately 40 kb apart from the aldosterone synthase gene (CYP11B2) [3]. Todate, more than 50 mutations have been reported in patients with 11-OHD, which are clustered in exons 2, 6–8 [4].

The first case is particularly interesting because of the delay of diagnosis. The 11β-hydroxylase deficiency diagnosis is to be considered when hypertension is associated with hypokalemia and hypogonadism, even in adult patients. A particular challenge in the diagnosis of classic 11OHD is the lack of standardized diagnostic work-up; thus, rare and untypical cases are likely to be missed. Early diagnosis and start of disease-specific treatment is important to avoid severe long-term consequences such as hyperandrogenism and potentially hypertension.

The second case shows the relationship between surgical outcome and psychological development. Severely virilized cases may initially be assigned as males, and once such assignment has been made, it may be difficult to reverse, thereby sacrificing fertility. Early medical, psychological and surgical treatment of children with virilizing congenital adrenal hyperplasia should enable them to become normal adults [5].

References: