Combined glucocorticoid and mineralocorticoid deficiency related to a new NNT mutation: a case report

E. Doye 1, CL. Gay 1, S. Castets 1, F. Roucher-Boulez 2, Y. Morel 2, I. Plotton 2, M. Nicolino 1

1 CHU Hospices Civils de Lyon, Service d’Endocrinologie Pédiatrique, Lyon, France; 2 Hospices Civils de Lyon, Laboratoire d’Endocrinologie Moléculaire et Maladies Rares, Lyon, France

BACKGROUND

Familial glucocorticoid deficiency is an autosomal recessive disorder characterized by specific failure of adrenocortical glucocorticoid production in response to adrenocorticotropic hormone (ACTH). Mutations of the NNT (nicotinamide nucleotide transhydrogenase) gene have recently been implicated in familial glucocorticoid deficiency.

OBJECTIVE

To describe a new case of familial glucocorticoid deficiency with combined mineralocorticoid insufficiency and extra adrenal manifestations.

RESULTS

Suffering from a febrile viral respiratory disease, an eight-month-old boy presented with status epilepticus caused by hypoglycemia. Multiple medical complications occurred, and invasive ventilation was required for 18 days. The results of blood tests performed during hypoglycemia revealed adrenal insufficiency. Renin and aldosterone levels were high but considered consistent with the mild hyponatremia and severe hypotension. Subsequent measurements revealed persistent high renin levels with low aldosterone concentration and hyponatremia, confirming the diagnosis of partial mineralocorticoid deficiency. ACTH levels remained high after six months of suitable treatment.

His parents are consanguineous and his father has a glucocorticoid deficiency since he was 18 months old, without mineralocorticoid deficiency during childhood. Genetic analysis revealed a new homozygous NNT mutation (p.Arg129*) for the child and his father.

Associated disorders:
- Thyroid stimulating hormone deficiency.
- Bradycardic sinus rhythm without cardiopathy.
- Neurological sequelae including severe hypotonia.

CONCLUSION

This case illustrates combined glucocorticoid and mineralocorticoid deficiency related to a new NNT mutation and underlines intra-familial phenotype heterogeneity. NNT gene should be considered when the most common etiologies of adrenal deficiency have been eliminated even if there is mineralocorticoid deficiency, in order to limit the serious consequences by a delayed diagnosis especially in offspring and to investigate any associated disorders.

References