A PATIENT WITH TURNER SYNDROME(45X/46XX) AND CONGENITAL ADRENAL HYPERPLASIA: A CASE REPORT AND A LITERATURE REVIEW

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INTRODUCTION

Turner syndrome is caused by partial or complete loss of the second sexual chromosome which leads to genital system malformation and infertility. 21-hydroxylase deficiency is a well-known cause of disorder of sexual development in genotypic female neonaates. A 8-month-old patient suffering from both 45,X/46,XX Turner’s syndrome and virilization form of CAH was referred to our hospital. She was born spontaneously at full term with normal birth measurement. After birth, she showed normal in growth and development until her mom found ambiguous genitalia when she was 8-month-old. Genital examination revealed a 3cm phallus, normal external urerital orifice and incomplete fusion of labia. An association between Turner syndrome (TS) and Congenital Adrenal Hyperplasia (CAH) due to 21-hydroxylase deficiency is rare. The aim of our study is to report this patient and stress this rare possibility.

AIM

- Summarize a rare case of CAH combined with TS, through the description of clinical manifestations and treatment outcome to strengthen clinicians’ understanding of this disease.
- By comparing the similarities and differences between this patient and the reported case to further summarize the clinical characteristics of the two diseases occurring in one patient at the same time.

METHOD

Through genetic testing and literature review, we confirmed our diagnosis and summarized relevant experience to guide our clinical work.

RESULTS

The patient was 2 months old when she came at the first time for clinical check, for the ambiguous external genitals with a hypertrophic clitoris still after her birth. The external genitalia was ambiguous; a clitoris was 1.5cm in length, absent labia majora and atretic opening of the vagina. Below the clitoris, the external orifice of urethra opening was normal. The result of karyotype was also made by outer hospital showed 45X/46XX, which revealed that this patient should be diagnosed with TS.

In this case, the sanger sequencing of CYP21A2 and karyotyping (including the sex determining region Y gene, SRY) must be verified to identify the etiology. Target sequencing of CYP21A2, followed by bioinformatics analysis, filtering against public databases, and biological analysis showed a homozygote missense mutation located at c.518T>A, p.Ile173Asn. the same time, the the karyotyping result showed a pattern of 45X/46XX like before, while the result of SRY analysis was negative. Therefore, we clarified that this girl was diagnosed with CAH combining with TS.

CONCLUSIONS

This case shows a rare combination of TS and CAH. A female patient with TS, if whose main manifestation is virilization, the SRY gene should be verified. If the SRY gene is negative, the simple virilizing type of CAH needs to be considered. It reminds us that if a known genetic disease cannot explain all the clinical manifestations, it is necessary to consider the possibility that another genetic disease exists at the same time, although we generally use a diagnosed disease to explain all the symptoms of the patients in the clinical diagnosis. Multidisciplinary combination therapy is particularly important in the treatment process, including endocrinology, urology, gynecology and assisted reproductive center.

REFERENCES


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CONTACT INFORMATION

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