The wide clinical expression spectrum presented in two families with NR5A1 mutation

**Context:** Patients with NR5A1 mutations present a wide spectrum of phenotype. The gene is frequently associated with 46, XY disorders of sex development (DSD).

**Objectives:** To analyze the clinical expression and NR5A1 mutations in two families with probands presenting 46, XY DSD.

**Results:** Two probands with 46, XY gonadal dysgenesis from two unconsanguineous families showed spontaneous puberty, normal levels of ACTH, cortisol, testosterone, increased FSH and decreased LH/FSH, AMH and INHB. In Family 1, a novel pathogenic variant, p.C33X, was identified in the proband. By determining other members in this family, the variant was found in the proband’s mother, two elder sisters and two nephews. However, no obvious anomaly was presented in the eldest sister with the variant while her two sons showed hypospadias. In Family 2, the proband inherited a reported pathogenic variant, p.M11 from his mother who delivered 4 children by natural pregnancy. The self-reported family puberty and fertility history was negative until we calculated the exact menopause age of mothers at 36 and 41 years of age respectively.

**Conclusion:** The 46, XY DSD patients with NR5A1 variants show highly variable phenotypes, from complete female genitalia to ambiguous external genitals/hypospadias. While 46, XX individual may have inconspicuous symptoms. Normal fertility is not a reliable negative family history on this disease. The identified novel pathogenic variant, p.C33X in patient 46, XY DSD and 46, XX primary ovarian insufficiency extends the genotypic and phenotypic spectrum of NR5A1. This study also highlights the critical role of NR5A1 protein in gonadal development and differentiation.

**Keywords:** NR5A1 gene, pathogenic variant, phenotype, gonadal dysgenesis

**Introduction**

Disorders of sex development (DSDs) are a group of heterogeneous diseases characterized by a rare set of conditions in which the chromosomal, gonadal, and phenotypic sex is atypical [1]. Nuclear receptor subfamily 5 group A member 1 (NR5A1, OMIM184757) plays a crucial role in regulating adrenal development, gonad determination and differentiation [2]. NR5A1 gene maps on 9q33 and has one non-translated exon (exon 1) and six other coding exons (exon 2–7) [3]. It encodes the steroidogenic factor 1 (SF-1) consisting of 461 amino acid residuals that regulate several steps of adrenal and gonadal development at early stages of embryonic development. The SF1 has two zinc-finger DNA-binding domains (DBD), a ligand-binding domain (LBD), two functional activation domains (AF-1 and AF-2), an accessory region and a hinge region [4]. The DBD contains a core with two Cys4-zinc-finger motifs and a highly conserved Ftz-F1 box motif potentially involved in interaction with DNA [5]. NR5A1 is highly conserved among different species, and the homology of mice is up to 95% of human NR5A1 gene [6].

In 46, XY individuals, SF-1 activates the expression of anti-Müllerian hormone (AMH) in Sertoli cells, resulting in the regression of Müllerian structures, and inducing production of steroidogenic enzymes in Leydig cells, facilitating the virilization of external genitalia and testes descending [7]. The protein functions in 46, XX individuals mainly as a promotor for follicle development and maturation. Therefore, individuals with NR5A1 mutation are mainly characterized by 46, XY DSD or 46, XX premature ovarian failure [23,25,27,28].

Achermann [8,9] reported initially in 1999 and 2002 that heterozygous mutations of Yanning Song1,2, Lijun Fan1,2, Xiu Zhao2, Xiaoya Ren1,2, Beibei Zhang1,2 & Chunxiu Gong*1,2,3

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NR5A1 gene could cause adrenal insufficiency and 46, XY severe testicular dysplasia. Then the following researches found heterozygous NR5A1 mutations could cause ambiguous genitalia, gonadal dysgenesis without adrenal insufficiency in 46, XY patient [10-14]. Further, Achermann [9] studied a consanguineous pedigree and found that the proband with the homozygous mutation (R92Q) of NR5A1 showed adrenal insufficiency and gonadal dysgenesis while the heterozygous parents and sister expressed no significant clinical phenotype, which is consistent with the results presented by Sadovsky et al. [15] who found that homozygous mutation of Nr5a1 in mice could have profound adrenal insufficiency and gonadal development, while heterozygous mutation only cause gonadal dysgenesis. So, they hypothesized that the SF-1 function in developmental pathways in humans is dosage sensitive. Most recently, case reports indicated that NR5A1 mutations can be associated with a wide spectrum of phenotypes, including 46,XX premature ovarian failure and even sex reversal like testicular DSD or ovotesticular DSD [16-21].

**Results**

**Clinical features**

The proband 1 was a social girl (Subject II-11) presented at the age of 17 years old with primary amenorrhea and virilized for 4 years. When she was born, normal genitals were presented, and the height and weight were in normal as female peers. At age of 13, she showed hoarseness, clitoris hypertrophy but no breast development. As her mother had breast development until 16 years old, so no attention was paid on. Till 17 years old, she got a karyotyping result of 46, XY at a local hospital, and then was referred to our hospital. She was boyish and always played with boys since very young school age according to her mother's description. Physical examination showed normal blood pressure (110/60 mmHg), armpit hair sparse, bilateral breast Tanner 1 and the clitoris enlargement like the penis (length of 5.6 cm). The urethra was open at the perineum (FIGURE 1). The karyotype was 46, XY. The mother of the proband 2 (Subject I-2) had menarche at 17 years old, natural pregnancy at 21 and was menopause at 36 years. Subject II-3 and subject II-7 died a few days after birth because of unknown reasons. The age of subject II-1, II-4, II-5, II-8, II-9, II-10 were 31, 30, 29, 26, 22, 21 years old and the menarche were 16-17 years old, respectively. Both subject III-1 and subject III-2 had hypospadias and had been repaired.

The proband 2 (Subject II-4) came from an unconsanguineous family, presenting with signs of virilization at the age of 10.5 years. She presented at birth with ambiguous external genitalia and was reared as female. Recently, she presented mustache, voice deepen and external genitalia changes. Physical examination showed the height was 148 cm and the weight was 50 kg. The bilateral breast was in Tanner 1 with apenism-like 3-cm long clitoris. The urethra was open at the perineum (FIGURE 1). The karyotype was 46, XY. The mother of the proband 2 (Subject I-2) had regular menstrual and natural pregnancy at the age of 23 years and was menopause at 41 years old.

**Subjects and methods**

**Subjects**

Two probands of 46, XY DSD and their family member were recruited in this study and informed consent was obtained. All the family members declined to take more clinical examinations except genetic testing.

**Methods**

Clinical information of the two probands were collected, including family history, physical examination, serum electrolyte and thyroid function, Adrenocorticotropic hormone (ACTH), Cortisol (Cor), Testosterone (T), Dihydrotestosterone (DHT), Anti Müllerian Hormone (AMH), Inhibin B (INHB), pelvic and testicular ultrasound etc. For genetic testing of the subjects, the AidLab DN01 kit was used to extract the genomic DNA from 2 ml of the peripheral blood in the EDTA anticoagulant tube. According to the primers and PCR reaction conditions as reported [22], all the exons were amplified. The PCR products were detected by 2% agarose gel electrophoresis and were sequenced. The obtained sequences were aligned and mapped to the reference of GeneBank (NM_004959.4). Clustal Omega (http://www.ebi.ac.uk/Tools/msa/clustalo/) was used to analyze the conservation of the novel variant.
The wide clinical expression spectrum presented in two families with NR5A1 mutation

years. SubjectII-1 presented with ambiguous sex external genitalia and died at 3 month with unknown reason. Subject II-2, II-3 were miscarried because of unknown reasons. SubjectII-5 was a normal boy. The electrolyte and thyroid function of the two probands were normal. The B-ultrasound examination of renal and adrenal gland was normal. Clinical features of the two families were shown in TABLE 1.

**Sequence analysis of the NR5A1 gene**

A heterozygous nonsense variant, c.99C>A, p.C33X (FIGURE 2A) was identified in Proband1, subjectI-2, II-1, II-3 and III-1, III-2. This mutation is predicted to alter the protein sequence and create a premature termination, that is c.99C>A coding the p. C33X then leading to deletion of 228 amino acids in SF-1 protein. The multiple species alignment suggested that cysteine on this site and the following 228 amino acid residuals were highly conservative (FIGURE 2B). The variant was not found in HGMD and ExAC database. Proband2 and her mother had a heterozygous variant, c.3G>A, p.M1I (FIGURE 2A), which has been identified as pathogenic variation in HGMD and clinvar database. This variant is located at the initiation codon, which destroys the initiation of translation, leading to protein translation defects [23].

Proband1 had been operated being a boy by his own decision after understanding the situation and had repaired hypospadias in August. Proband2 was 10 years old and...
Table 1. Clinical Data of the 2 family members.

<table>
<thead>
<tr>
<th></th>
<th>Family 1</th>
<th>Family 2</th>
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<tbody>
<tr>
<td></td>
<td>Subject I-2</td>
<td>Subject II-1</td>
</tr>
<tr>
<td>Family number</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age at presentation(y)</td>
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<td>--</td>
</tr>
<tr>
<td>Sex of rearing</td>
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<td>female</td>
</tr>
<tr>
<td>Karyotype</td>
<td>46XX</td>
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<td>Karyotype</td>
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<td>Karyotype</td>
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<tr>
<td>PhenoType</td>
<td>Primary ovarian</td>
<td>normal female</td>
</tr>
<tr>
<td>Karyotype</td>
<td>insufficiency</td>
<td></td>
</tr>
<tr>
<td>External genitalia</td>
<td>female</td>
<td>female</td>
</tr>
<tr>
<td>LH(IU/L) (≤ 11.3)</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>FSH(IU/L) (≤ 13.0)</td>
<td>--</td>
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<tr>
<td>T (ng/dl) (&gt;110)</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>AMH (ng/ml) (&gt;1.66)</td>
<td>--</td>
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<tr>
<td>INHB (pg/ml) (19.67-147.62)</td>
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<tr>
<td>ACTH (pg/ml) (0-46)</td>
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<tr>
<td>COR (ng/dl) (5-25)</td>
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<tr>
<td>DHT (pg/ml)</td>
<td>--</td>
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</tr>
<tr>
<td>NRSA1 mutation</td>
<td>c.99C&gt;T,p,C33X</td>
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The wide clinical expression spectrum presented in two families with NR5A1 mutation

Discussion

Widely variable of NR5A1-related phenotypes were well documented in 46, XY patients [17,24-27]. And the spectrum of disorders associated with NR5A1 variants in 46, XX individuals was further expanded by the demonstration of the specific recurrent heterozygous p.Arg92Trp NR5A1 variant in association with variable degrees of testis development in 46, XX patients from unrelated families [20]. In this research, the proband1 presented with complete female genitalia at birth and appeared masculinization in adolescence. Her two brothers died several days after birth of unknown causes. It is difficult to rule out whether NR5A1 mutation leading to adrenal insufficiency that adrenal crisis causing death. Two nephews of the proband 1 carried the variant showed different degrees of hypospadias. These results indicated that regardless of the same variant in the family, different 46, XY individuals may present different phenotypes.

The two probands' mothers showed premature ovarian failure (POF) and earlier menopause, suggesting that screening NR5A1 gene is necessary in POF patients, as proposed in other research [28]. However, due to the variable phenotypes and incomplete penetrance, some female carriers may be easily misdiagnosed because of normal fertility preserved in 46, XX individual, as shown in the two mothers and the II-1 of family 1. Thus, negative family history, normal fertility, and normal puberty are not always reliable criteria. These features in the two families made us uncertain at the beginning until we got their menopause ages. Moreover, the unreliable negative family history may come from concealment of the mother due to pressure from the family. For the sisters who carried the same mutant allele, we should pay more attention to signs of POF as indicated from their mother. Regular follow-up, evaluation and appropriate treatments should be taken when necessary.

After the gonadal differentiation into testis of a 46, XY individual, SF-1 was expressed in the early testes continuously, maintaining the expression of SOX 9 together with SRY. In 7 weeks of pregnancy, SF-1 activates the expression of AMH, which induces the degeneration of the Müllerian structure and activates the steroid enzyme synthesis to maintain the development of masculine external genitals from the stage of 8 weeks [29]. The damage of Leydig cells at the early stage of embryonic development caused by pathogenic NR5A1 mutations can lead to inadequate secretion of T; therefore, patients with NR5A1 mutations always present deficiency in virilization and are reared as female in most cases. However, at puberty they show hoarseness, prominent Adam’s apple and hypertrophy of the clitoris [31,32], similar situations as indicated in the probands of this study.

Most patients have no remnants of Müllerian structure attributing to sufficient AMH produced at the early stage of embryonic development to degenerate the Müllerian structure. Previous research found the function of Storli cells gradually decreases with age [30], which can be inferred from the patients in this study. Absence of uterus in the two probands indicated AMH secretion by Storli cells were enough during embryonic development. Elevated levels of both LH and FSH with expected low level of suggested a condition of hypergonadotropic hypogonadism. A few studies have confirmed that 46, XY patients with NR5A1 mutation can present spontaneous puberty [31,32] and continuous elevation of FSH level in the follow-up, suggesting damage of Storli cells and the resulting dyszoospermia. This also explains why rare males with NR5A1 mutation have fertility function [33-35]. For the two probands in our study, we observed that LH/FSH ratio, AMH and INHB levels decreased significantly with a normal level of testosterone, indicating the damage of Storli cells may be more severely than that of Leydig cells. Both the two cases showed spontaneous puberty and virilization probably induced by the residual cells, consistent with the former research [31-32].

To date, nearly 120 NR5A1 variants have been documented in Human Gene Mutation Database (http://www.hgmd.cf.ac.uk/ac/gene.php?gene=NR5A1), including nonsense variants (the most common), missense variants, small deletions and duplications, small indels, and splicing variants. The genotype–phenotype relationship is unclear due to deficiency of studies of phenotypic diversity with large content of human cases. In this study, we identified a
novel heterozygous nonsense variant (c.99C>A, p.C33X) in family 1 and a known pathogenic variant (c.3G>A, p.M1I) in the second family.

Generally, mutations in the DBD region of NR5A1 directly interacts with DNA and may alter patterns or recognition of specific DNA sequences, resulting in reduced expression of downstream target genes and disruption of gonadal development regulation. The variant, p.C33X is located in DBD region, directly combined with the first zinc finger protein. It is known that cysteine affects the 3D structure of the whole protein. Its sulfenyl reacts with other groups to form disulfide bond, maintaining protein folding and stability. Cysteine residues also play an important role in cross-linked proteins, which increase the rigidity of protein, and increase the resistance to hydrolysis. Therefore, cysteine changes are not only lead to the early termination of the protein translation, but also lose the ability of DNA to combine with the zinc finger, causing loss of the whole protein function. Although no functional verification was seen, multiple sequence comparisons indicated that the amino acid in this locus was conservative. Combined with family information, clinical symptoms and the nature of this mutation (nonsense mutation), we identified the novel mutation as pathogenic.

The c.3G>A p.M3I was first reported in a family by Lourenco et al. [23] in 2009. Both the cases (46, XX girls), carrying a heterozygous mutation (c.3G>A p.M3I), presented a partial gonadal dysgenesis and primary ovarian insufficiency respectively (irregular menstruation, menstruation ended at 15 years old). Their mother carrying the same mutation without phenotype was interpreted as incomplete penetrance [11]. Translation initiation at this codon is predicted to truncate the N-terminal of the protein, eliminating the DNA-binding domain.

DSD sex rearing has been a hot topic of debate. Many masculine females with NR5A1 variants chose to change gender [25,31]. As showed above, the boys with NR5A1 mutations presented spontaneous puberty, and other similar patients with preserved fertility had been reported [31,32], hinting certain advantages in rearing those 46, XY patients with NR5A1 mutation as boys. In this study, proband 1 has taken repair operation for hypospadias and proband 2 has been waiting until old enough to make further evaluation.

**Conclusion**

In this study, we identified a novel nonsense variant and a known pathogenic variant of NR5A1 gene from two unrelated families. The 46, XY DSD patients with NR5A1 variants show highly variable phenotypes, from complete female genitalia to ambiguous external genitals/hypospadias. The 46, XX individuals with NR5A1 variants could be misdiagnosed due to atypical clinical signs. The identified novel pathogenic variant and detailed clinical characteristics in patients extend the genotypic and phenotypic spectrum of NR5A1. Screening of NR5A1 gene could improve diagnosis of DSD.

**Conflict of interest statement**

All authors declare that no conflicts of interest exist regarding this work.

**Acknowledgments**

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The wide clinical expression spectrum presented in two families with NR5A1 mutation

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