Molecular and Cytogenetic Analysis of Romanian Patients with Differences in Sex Development

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Abstract: Differences in sex development (DSD) are often correlated with a genetic etiology. This study aimed to assess the etiology of DSD patients following a protocol of genetic testing. Materials and methods. This study prospectively investigated a total of 267 patients with DSD who presented to Clinical Emergency Hospital for Children Cluj-Napoca between January 2012 and December 2019. Each patient was clinically, biochemically, and morphologically evaluated. As a first intervention, the genetic test included karyotype + SRY testing. A high value of 17-hydroxyprogesterone was found in 39 patients, in whom strip assay analysis of the CYP21A2 gene was subsequently performed. A total of 35 patients were evaluated by chromosomal microarray technique, and 22 patients were evaluated by the NGS of a gene panel. Results. The karyotype analysis established the diagnosis in 15% of the patients, most of whom presented with sex chromosome abnormalities. Genetic testing of CYP21A2 established a confirmation of the diagnosis in 44% of patients tested. SNP array analysis was particularly useful in patients with syndromic DSD; 20% of patients tested presented with pathogenic CNVs or uniparental disomy. Gene panel sequencing established the diagnosis in 11 of the 22 tested patients (50%), and the androgen receptor gene was most often involved in these patients. The genes that presented as pathogenic or likely pathogenic variants or variants of uncertain significance were RSP01, FGFR1, WT1, CHD7, AR, NIPBL, AMHR2, AR, EMX2, CYP17A1, NR0B1, GNRHR, GATA4, and ATG genes. Conclusion. An evaluation following a genetic testing protocol that included karyotype and SRY gene testing, CYP21A2 analysis, chromosomal analysis by microarray, and high-throughput sequencing were useful in establishing the diagnosis, with a spectrum of diagnostic yield depending on the technique (between 15 and 50%). Additionally, new genetic variants not previously described in DSD were observed.

Keywords: differences in sex development; karyotype; SRY; CYP21A2; chromosomal microarray; next-generation sequencing

1. Background

Differences in sex development (DSD), defined by atypical developments of chromosomal, gonadal, or phenotypic sex, are observed in 1:4000 newborns, although this incidence may differ according to the disorders included. Isolated hypospadias or cryptorchidism are more frequently observed, in as many as 1:200 newborns [1–4]. These disorders are at

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the end of the DSD spectrum, and in some cases are not included as DSD, even if in some cases, they represent a milder effect of the same etiology seen in severe DSD.

Clinical features that indicate DSD usually include isolated clitoral hypertrophy, isolated posterior hypospadias, bilateral cryptorchidism or ectopia, and unilateral cryptorchidism/testicular ectopia associated with hypospadias or micro-penis; these signs are often seen in the newborn period [5,6]. Signs that suggest DSD at puberty are virilization of initially female genitalia, pubertal delay, and primary amenorrhea [5,6]. Usually, the clinical investigation aims to examine the following: genital tubercle (size and aspect), labioscrotal folds (their posterior fusion), the urinary meatus (localization), and palpable gonads. Based on these data, the Prader stages—a masculinization scale of the external genitalia and anogenital distance—could be useful indices for the degree of genital virilization [5–7]. Clinical evaluation of the external genitalia is always followed by an imaging evaluation to describe the characteristics of the urogenital sinus, Mullerian structures, gonads, adrenal glands, and sometimes, the renourinary system (to determine the common embryonic origin). This evaluation usually is performed using ultrasounds and MRI, although genitography/genitoscopy or even exploratory laparoscopy and histopathological examination are utilized in some situations [8].

DSD are classified depending on karyotype (i.e., 46,XY DSD; 46,XX DSD; and sex chromosome DSD), and in some cases, DSD is observed as part of other developmental disorders (e.g., syndromic DSD) [9]. 46,XY and 46,XX DSD could be the results of abnormal gonadal development or disorders of steroid synthesis and action. 46,XX disorders of ovarian development include 46,XX testicular and ovotesticular DSD, which are caused by SRY translocation in 85% of testicular 46,XX and 10% of ovotesticular 46,XX, as well as SOX9, RSPO1, or SOX3 disorders. They also include 46,XX gonadal dysgenesis, which, in cases of premature ovarian failure, is due to genes involved in gonadal development, such as FSHR, NR5A1, BMP15, STAG3, and WT1 [9,10]. 46,XY disorders of testicular development include complete and partial gonadal dysgenesis, which are due more often to SRY gene abnormalities and less often to DHH, NR5A1, MAP3K1, CBX2, NROB1, WNT4, or DMRT1 gene abnormalities [9,10]. Sex chromosome DSD includes Turner syndrome (45,X and variants); Klinefelter syndrome (47,XXY and variants); mosaic 45,X/46,XY (mixed gonadal dysgenesis); 46,XX/46,XY (ovotesticular DSD); and other aneuploidies involving sex chromosomes [9].

DSD may not be an endocrine disorder but rather a developmental one, such as cloacal exstrophy (omphalocele-exstrophy-imperforate anus-spinal defects–OEIS spectrum (in 46,XX and 46,XY DSD) and developmental abnormalities of the Mullerian structures (e.g., hypospadias in 46,XY DSD or Mayer-Rokitansky-Küster-Hauser (MRKH) syndrome, vaginal atresia, and Mullerian agenesis in 46,XX DSD). In cases of 46,XX and 46,XY disorders of steroid hormone biosynthesis, CYP21A2 gene mutations are the most common cause (>90% of these cases). Genetic mutations found among the remainder include: STAR (46,XY DSD), 3βHSD (46,XY DSD), CYP11A1 (46,XY DSD), CYP11B1 (46,XX DSD), POR (46,XX and 46,XY DSD), CYP17A1 (46,XY DSD), CYP19A1 (46,XX DSD), 5ARD (46,XY DSD), or 17βHSD3 (46,XY DSD) [9]. Disorders of steroid hormone action in 46,XY patients are most often due to gene mutations in AR, AMHR, or LHR abnormalities.

Morphologic assessment is always based on biochemical and genetic analysis. Biochemical analysis refers primarily to steroid analysis, such as plasma 17-hydroxyprogesterone, testosterone, dihydrotestosterone, androstenedione, DHEA, DHEAs, deoxycorticosterone, plasma renin activity, ACTH, FSH, LH, AMH, and urinary/24-h evaluation of urinary steroid profile (performed by gas chromatography associated with mass spectrometry [GC/MS]) [11]. Plasma sexual steroids are examined with immunoassay or liquid chromatography associated with tandem mass spectrometry (LC-MS/MS) [11]. Genetic testing is based on first-tier testing, as well as karyotype analysis and SRY gene evaluation, which aims to establish the first step in understanding the pathogenetic mechanism [9,12]. Then, depending on the clinical and hormonal picture, the CYP21A2 gene is assessed by Sanger sequencing if 17-hydroxyprogesterone indicates an adrenal
enzymatic block. If not, a gene panel or exome, or whole genome will be performed using next-generation sequencing (NGS) techniques [13]. If DSD is associated with other symptoms (e.g., syndromic DSD), chromosomal analysis by microarray is performed to observe the copy number variants (CNVs) that may have caused the disorder. Knowing the precise diagnosis will help in understanding the prognosis and designing better treatments that consider a precise etiopathogenetic mechanism. Therefore, the aim of this study was to assess the genetic etiology of patients with DSD presented in our service, who were clinically, biochemically, and anatomically evaluated.

2. Materials and Methods

Patients with DSD who presented to Clinical Emergency Hospital for Children Cluj-Napoca between January 2012 and December 2019 were investigated prospectively. The inclusion criteria for the study group were clitoral hypertrophy, posterior hypospadia, bilateral cryptorchidism or ectopia, unilateral cryptorchidism/testicular ectopia associated with hypospadias or micro-penis, puberty delay, and primary amenorrhea. A total of 267 patients were evaluated (Figure 1) with karyotype and SRY testing (using either fluorescent in situ hybridization (FISH) or polymerase chain reaction (PCR) techniques).

Figure 1. The investigation protocol for genetic testing in the study group. (* The patients were not further tested due to the absence of clinical monitoring after their first medical visit).

Patients were evaluated by imaging studies (ultrasound, pelvic MRI), with additional studies (such as exploratory laparotomy and gonadal biopsy) performed depending on the clinical context. Hormonal testing (for 17-hydroxyprogesterone, DHEAS, delta4 androstenedione, testosterone, DHT, and AMH) was also performed for some patients depending on the clinical context.

In 46,XX patients with high 17-hydroxyprogesterone values (greater than 2 ng/mL), testing for 21-hydroxylase deficiency was performed. For some patients, genetic testing
was performed if 17-hydroxyprogesterone was over 10 ng/mL after a stimulation test with synthetic ACTH. Forty-one 46,XX patients were genetically tested using strip assay analysis for the 11 most common mutations of CYP21A2 (Figure 1). For patients for whom karyotype, SRY, and 21-hydroxylase deficiency testing did not establish a diagnosis, the SNP array technique was used to evaluate CNVs (copy number variants) if the patients continued to be evaluated. Some patients declined follow-up investigations precluding obtaining further clinical data or biological samples. These patients had no clinical, hormonal, or morphological differences from the group further tested. Therefore, given limited financial funds for genomic testing (SNP array or high throughput sequencing) provided by our healthcare system, it was decided to perform these tests only for the patients who continued the follow-up for their disorders.

SNP array testing was completed for 35 patients (Figure 1), of whom 22 patients had negative results and were consequently evaluated with gene panel sequencing (TruSight One panel, Illumina) (Figure 1). This panel included known genes associated with human pathology, particularly those associated with DSD.

The research was approved by the ethics committee of “Iuliu Hatieganu” University of Medicine and Pharmacy of Cluj-Napoca. Written informed consent was obtained from the parents of all patients in the study.

2.1. Strip Assay Technique for the CYP21A2 Gene

The DNA from 3 mL of peripheral blood from each patient was purified using a Wizard® Genomic DNA Purification Kit (Promega, Madison, WI, USA). This was performed using a PCR amplification followed by hybridization on a strip containing specific oligonucleotide probes for the 11 most common mutations of the CYP21A2 gene: P30L, I2 splice (I2 G), Del 8 bp E3 (G110del8nt), I172N, Cluster E6 (I236N, V237E, M239K), V281L, L307 frameshift (F306+T), Q318X, R356W, P453S, and R483P (® ViennaLab CAH Strip Assay, Vienna, Austria) [14].

2.2. Chromosomal Microarray Technique

The chromosomal microarray comprised an SNP array, which was performed using an Infinium OmniExpress-24 BeadChip array kit (Illumina, San Diego, CA, USA), and the platform iScan System (Illumina, San Diego, CA, USA). The bioinformatic instrument was Genome Studio software version 2.0 (Illumina, San Diego, CA, USA). This analysis was based on 700,000 markers, and the interpretation of the results was based on the American College of Medical Genetics (ACMG) recommendation [15].

2.3. Genes Panel Sequencing

The sequencing was performed using the TruSight One Kit (Illumina, San Diego, CA, USA), which targets 4800 genes associated with human pathology (12 Mb). This included around 150 genes or candidate genes associated with the clinical phenotype for DSD. The sequencing was performed with the MiSeq platform (Illumina, San Diego, CA, USA) [16] using the manufacturer’s instructions. Bioinformatic analysis was performed using Galaxy bioinformatic platform, and variant interpretation was based on ACMG recommendations [17].

3. Results

A total of 267 patients were evaluated with karyotype and SRY testing. Of these patients, 39 (14.6%) were diagnosed with different chromosomal abnormalities, most of them involving the sex chromosomes (36 of 39 patients; Table 1). The 46,XX karyotype was observed in 104 patients (39%), while the 46,XY karyotype was observed in 122 patients (46%). Two patients with the 46,XX karyotype were SRY positive.
Table 1. Karyotypes observed in DSD patients.

| Karyotype                        | Number of Patients, n |
|----------------------------------|----------------------|
| 45,X                             | 18                   |
| 45,X/46,XY                       | 2                    |
| 45,X/46,XX                       | 4                    |
| 45,X/46,XX/47,XXX                | 3                    |
| 46,Xi(Xp)                        | 1                    |
| 46,XX.dup(XX)(q22q24)            | 1                    |
| 47,XXY                           | 3                    |
| 46,XY/47,XXX                     | 1                    |
| 48,XXXXY                         | 1                    |
| 49,XXXXX                         | 1                    |
| 46,XY.dup(8)(q24.3)              | 1                    |
| 46,XY.del(12)(q13)               | 1                    |
| 46,XY/47,XY,+mar                 | 1                    |
| 46,XX                            | 106                  |
| 46,XY                            | 122                  |

Of patients with the 46,XX karyotype, 39 were tested for the main CYP21A2 mutations (using strip assay). This testing was performed if their basal 17-hydroxyprogesterone was greater than 2 ng/mL or if stimulated 17-hydroxyprogesterone (after synthetic ACTH administration) was greater than 10 ng/mL. In 17 (44%) of the 39 patients, the diagnosis was confirmed.

Of 85 patients with the 46,XX karyotype (without an established etiologic diagnosis, after CYP21A2 testing) and of 122 patients with the 46,XY karyotype, 8 and 27 patients, respectively, were further analyzed using chromosomal microarray (Table 2). The other undiagnosed patients were not tested further for different reasons (i.e., absent biological sample, absent clinical follow-up, limited possibility of genomic testing). Three of the eight 46,XX patients (38%) presented pathogenic CNVs (two patients) or VUS (variant of unknown significance) (one patient); two of these patients presented syndromic 46,XX DSD, while one presented an isolated form (Table 2).

Of the 27 46,XY patients, 10 patients (37%) were diagnosed with pathogenic CNVs (three patients), uniparental disomy (three patients), or VUS (four patients); each of these 10 patients presented syndromic DSD.

Patient p10, 46,XY, presented a recurrent 16p11.2 deletion of 597kb, which has classically been associated with intellectual disability, developmental delay, autism spectrum disorders, and obesity. Cryptorchidism has also been mentioned in patients with this CNV. However, the precise gene responsible for cryptorchidism is not known. Patient p4 presented with disomy of chromosome 7, and his phenotype was concordant with the Russell-Silver phenotype. This syndrome has been associated with cryptorchidism, micropenis, and hypospadias in other patients, and thus the genital phenotype in this patient was considered a result of this syndrome. Two patients, p14 and p21, presented 15 chromosome isodisomy, and further MLPA identified a maternal origin for these chromosomes. The clinical phenotype superposed that of Prader-Willi syndrome, and in this clinical context, the cryptorchidism was due to hypogonadotropic hypogonadism. Patient p18, who had an exon 4 deletion in the OTC gene, presented cryptorchidism. OTC mutations have not been previously described in association with this phenotype; however, the metabolic defect associated with hypotonia at a younger age may have had some association with
cryptorchidism. Patients p3 and p6 presented uncertain CNVs in the 15q11.2 region, but previous data have not supported the involvement of these CNVs in DSD. Patients p24, p26, and p35 presented variants of uncertain significance, with no clear argument for their consideration in a final diagnosis.

P30, a 46,XX patient, presented 1q21.1 deletion, or BP2-BP3 (thrombocytopenia absent radius [TAR] syndrome), and the clinical phenotype of the patient superposed with this syndrome. Genitourinary abnormalities and Mayer-Rokitansky-Küster-Hauser syndrome have been described in TAR syndrome, and this diagnosis was etiologically appropriate for the uterus agenesis described in patient p30. The gene RBM8A is suggested to be responsible for the Mayer-Rokitansky-Küster-Hauser phenotype in TAR syndrome.

Patient p31 was diagnosed with 18p11.32-18p11.31 deletion, an etiological diagnosis for genitourinary abnormality, cryptorchidism, and micro-penis that have been described in other patients with this deletion. The CYP21A2 homozygous deletion in 46,XX patient p32 identified by SNP array technique was also an etiologic diagnosis for this patient. Chromosomal microarray led to a clear etiologic diagnosis for two of eight 46,XX patients (25%) and for five of 27 46,XY patients (19%). Thus, of the 35 patients tested (46,XX and 46,XY), 7 (20%) had a final etiologic diagnosis.

Patients with negative results after SNP array testing were analyzed by gene panel sequencing that included the known and candidate genes involved in DSD (Table 3). Of 22 patients who were analyzed, five patients had the 46,XX karyotype, and 17 patients had the 46,XY karyotype. Three of the five tested 46,XX patients (60%) presented pathogenic variants (two patients) and VUS (one patient). Patient p9 (a 46,XX patient) presented compound heterozygous variants in the RSPO1 gene, which were interpreted as class III variants. These variants included one c.286+1G>A (modifies the splice site between exon 4—intron 4, not noted in gnomAD, described in Clinvar [patient 18085567], predicted as pathogenic by BayesDel_addAF, DANN, EIGEN, FATHMM-MKL, MutationTaster, and scSNV-Splicing) and one c.484A>G (missense variant in exon 6, described in gnomAD as having a very low frequency [f = 0.00000401]). Another 46,XX patient, p12, presented compound heterozygous variants in the FGFR1 gene, one in c.914A>G, which was interpreted as pathogenic (a missense mutation in exon 8, a hot-spot region of the gene, classified as pathogenic by Uniprot, already described in several articles, not found in gnomAD, predicted as pathogenic by BayesDel_addAF, DANN, DEOGEN2, EIGEN, FATHMM-MKL, LIST-S2, M-CAP, MVP, MutationTaster, and SIFT) and one in c.2440A>C, which was interpreted as VUS (missense variant in exon 19, not found in gnomAD, predicted as pathogenic by the prediction platforms BayesDel_addAF, DANN, DEOGEN2, EIGEN, FATHMM-MKL, LIST-S2, M-CAP, MutationAssessor, MutationTaster, and SIFT). Patient p34 presented a heterozygous c.1075G>T variant in the ATM gene, which was interpreted as pathogenic (a null variant in exon 9, not found in gnomAD with pathogenic predictions in BayesDel_addAF, DANN, EIGEN, FATHMM-MKL, and MutationTaster); this gene has been associated with premature ovarian failure, as observed in patient p34.

Of the 17 46,XY patients tested by gene panel sequencing, 13 (76%) presented pathogenic variants (two patients), likely pathogenic variants (seven patients), and VUS (four patients). Patient p1 presented a heterozygous variant, c.437C>A, in the WT1 gene, which was interpreted as VUS (a missense variant in exon 1, not described in gnomAD; 93% of missense variants of the WT1 gene are pathogenic, predicted pathogenic by BayesDel_addAF, DANN, FATHMM-MKL, M-CAP, PrimateAI, and SIFT). Patient p2 presented a heterozygous variant, c.5405-7G>A, in the CHD7 gene, which was classified as likely pathogenic (an intronic variant, classified as pathogenic in Clinvar by several submissions and articles, not found in gnomAD).
Table 2. Pathogenic CNVs and VUS observed in patients analyzed by SNP array analysis.

| Patient | Isolated/Syndromic | Clinical Phenotype | Karyotype | CNV/DUP | Size | Main Genes Included | Interpretation |
|---------|--------------------|--------------------|-----------|---------|------|---------------------|----------------|
| p3      | Syndromic          | Primary amenorrhea, clitoridian hypertrophy, short stature, intellectual disability | 46,XX | arr[GRCh37]15q11.2 (22737333_23226254)x3 | 472 kb | NIPA1, NIPA2, CYFIP1, TUBGCP5 | VUS |
| p4      | Syndromic          | Bilateral cryptorchidism, intellectual disability, craniofacial dysmorphism | 46,XY | UPD7 | - | - | Pathogenic |
| p6      | Syndromic          | Enlarged phallus, urogenital sinus, partial posterior fusion of genital folds (Prader 3), global developmental delay, craniofacial dysmorphism | 46,XY | arr[GRCh37]15q11.2 (22652330_23226254)x3 | 573 kb | NIPA1, NIPA2, CYFIP1, TUBGCP5 | VUS |
| p10     | Syndromic          | Cryptorchidism, micropania, puberty delay, intellectual disability | 46,XY | arr[GRCh37]16p11.2 (29595483_30192561)x1 | 597 kb | PRTT2, KIF22, ALDOA, TRX6, SEZ6L2, TMEM219, MAPK3, SPN, QPRT, MAZ, MVP, and 11 other OMIM genes (proximal 16p11.2 microdeletion syndrome) | Pathogenic |
| p14     | Syndromic          | Bilateral cryptorchidism, short stature, obesity, deafness | 46,XY | UPD15 | - | - | Pathogenic |
| p18     | Syndromic          | Cryptorchidism, obesity, macrocrania, micromelia | 46,XY | arr[GRCh37]Xp11.4 (38230704_38246882)x1 | 16 kb | OTC (exon 4) | Pathogenic |
| p21     | Syndromic          | Micro-penis, bilateral, cryptorchidism, hypotonia, craniofacial dysmorphism | 46,XY | UPD15 | - | - | Pathogenic |
| p24     | Syndromic          | Scrotal hypospadia, cryptorchidism, intellectual disability, language delay, obesity | 46,XY | arr[GRCh37]Xp25.1 (5256116_5391419)x1 | 135 kb | FARS2, LYRM4 | VUS |
| p26     | Syndromic          | Micro-penis, cryptorchidism, craniofacial dysmorphism, intellectual disability | 46,XY | arr[GRCh37]14q11.2 (19401291_20420338)x3 | 1.01 Mb | POTEG | VUS |
| p30     | Syndromic          | Primary amenorrhea, uterus agenesis, forearm agenesis, ectromelia, epilepsy, intellectual disability | 46,XX | arr[GRCh37]1q21.1 (145394955_145755813)x1 | 360 kb | RBM8A, PEX11B, POLR3G, HJV, HEF2A, and other OMIM genes (BP2-BP3 microdeletion syndrome) | Pathogenic |
| p31     | Syndromic          | Micro-penis intellectual disability, craniofacial dysmorphism | 46,XY | arr[GRCh37]18p11.32p11.31 (13034_430081)x1 | 4.3 Mb | SMCHD1, LPIN2, TGIF1, and other OMIM genes included in 18p11.3 region (18p11.3 deletion syndrome) | Pathogenic |
| p32     | Isolated           | Enlarged phallus, partial fusion of genital folds, Prader 2 | 46,XX | arr[GRCh37]Xp21.33 (32005904_32006896)x0 | 0.99 kb | CYP21A2 (exons 1-3) | Pathogenic |
| p35     | Syndromic          | Scrotal hypospadia, cryptorchidism, palatine cleft, skeletal dysplasia | 46,XY | arr[GRCh37]Xp11.4 (41665315_41684603)x1 | 19 kb | CASK (intron 2) | VUS |

CNV—copy number variants, UPD—uniparental disomy, VUS—variant of uncertain significance.
| Patient | Age | Gender | Gonads                        | Muller Derivatives          | External genitalia      | Puberty | FSH LH | Hormonal                              | Other                                                      | Karyotype | SRY | Variant (HGVS)                                         | Zigosity                      | Variant Interpretation (ACMG Criteria) |
|---------|-----|--------|-------------------------------|-----------------------------|-------------------------|---------|--------|-----------|--------------------------------------|------------------------------------------------------------|----------|-----|-----------------------------------------------------|---------------------------------|-------------------------------------|
| p1      | 4   | F      | Bilateral testicular ectopia  | Yes, uterus, Vagina         | Female                  | Tanner 1 Prepuberty level N |        |        | craniofacial dysmorphism, diaphragmatic hernia, calos body hypoplasia | 46,XY + | WT1(NM_024426.5): c.437C>A (p.Trp151Cys) MAF = 0 | Heterozygous               | VUS (PM2, PP2, PP3) |
| p2      | 11  | M      | Bilateral cryptorchidism      | No                          | Micro penis, cryptorchidism | Tanner 1 Prepuberty level N |        |        | Craniofacial dysmorphism             | 46,XY + | CHD7(NM_017780.3): c.5405-7G>A MAF = 0 rs398124321 MAF = 0 | Heterozygous | Likely pathogenic (PP5, PM2) |
| p5      | 1   | F      | Bilateral cryptorchidism      | No                          | Female                  | Tanner 1 Prepuberty level N |        |        | craniofacial dysmorphism, GDD/ID     | 46,XY + | AR(NM_000044.6): c.2071_2073del (p.Asp691del) MAF = 0 | Heterozygous | Likely pathogenic (PM1, PM2, PM4, PP3) |
| p7      | 1.6 | M      | Anorhidia                    | No                          | Micro penis             | Tanner 1 ↑↑↑ Prepuberty level |        |        | craniofacial dysmorphism, GDD/ID     | 46,XY + | NIPBL(NM_133433.4): c.1808del (p.Lys603SerfsTer11) rs727503767 MAF = 0 | Heterozygous | Pathogenic (PVS1, PM2, PP3, PP5) |
| p8      | 0.3 | F      | Bilateral testicular ectopia  | No                          | Clitoridian hypertrophy, 2 orifices, right inguinal hernia | Tanner 1 Prepuberty level N |        |        | Hypercalcemia                         | 46,XY + | AR(NM_000044): c.167T>A (p.Leu56Gln) rs868709351 MAF = 0 | Heterozygous | VUS (PM2, PP2) |
|         |     |        |                               |                             |                         |                      |        |        |                                      |                                      | AR(NM_000044): c.170_171insGCAGCAACCAC (p.L57insGlnGlnGlnGln) rs3032358 MAF = 0 | Heterozygous | VUS (PM2, PP3) |
| Patient | Age | Gender | Gonads | Muller Derivatives | External genitalia | Puberty | FSH LH | Hormonal | Other | Karyotype | SRY | Variant (HGVS) | Zygosity | Variant Interpretation (ACMG Criteria) |
|---------|-----|--------|--------|-------------------|-------------------|---------|--------|----------|-------|-----------|-----|----------------|---------|---------------------------------------|
| p9      | 3.1 | M      | Scrotal testes | No | Penoscrotal hypospadias, micro-penis, bifidus scrotum | Tanner 1 | Prepuberty level | T, DHT ↓ | no | 46,XX | - | RSP01(NM_001242908.2): | Heterozygous | VUS (PM2, PP3, PP5) |
|         |     |        |        |       |       |         |         |          |     |           |    | c.286+1G>A | rs157099690, MAF = 0 |                  |                                     |
|         |     |        |        |       |       |         |         |          |     |           |    | RSP01(NM_001242908.2): | Heterozygous | VUS |                           |
|         |     |        |        |       |       |         |         |          |     |           |    | c.484A>G (p.Lys162Glu) | rs36043533, MAF = 0.0000464 |                  |                                     |
| p11     | 0.3 | F      | Bilateral testicular ectopia | Yes, uterus, vagina | Penoclitoridian organ, one orifice at the base of the gland, labioscrotal folds posteriorly fusioned, not palpable gonads, Prader 4 | Tanner 1 | Prepuberty level | N | no | 46,XY + | AMHR2(NM_020547.3): | Heterozygous | Pathogenic (PVS1, PM2, PP3) |
|         |     |        |        |       |       |         |         |          |     |           |    | c.233-2A>G, MAF = 0 |                  |                                     |
|         |     |        |        |       |       |         |         |          |     |           |    | AMHR2(NM_020547.3): | Heterozygous | Likely pathogenic (PVS1, PM2) |
|         |     |        |        |       |       |         |         |          |     |           |    | c.133delA (p.Thr45GlnfsTer23), MAF = 0 |                  |                                     |
|         |     |        |        |       |       |         |         |          |     |           |    | AMHR2(NM_020547.3): | Heterozygous | VUS (PM2, PP2, PP3) |
|         |     |        |        |       |       |         |         |          |     |           |    | c.137G>T (p.Gly46Val) |                  |                                     |
| p12     | 17  | F      | Not evidenced | Yes, hypoplastic uterus | Female N | Tanner 1 | ↑ | N | no | 46,XX + | FGFR1(NM_001174067.1): | Heterozygous | pathogenic (PS3, PM1, PM2, PP2, PP3, PP5) |
|         |     |        |        |       |       |         |         |          |     |           |    | c.914A>G (p.Glu305Gly) | rs727505369, MAF = 0 |                  |                                     |
|         |     |        |        |       |       |         |         |          |     |           |    | FGFR1(NM_001174067.1): | Heterozygous | VUS (PM2, PP2, PP3) |
|         |     |        |        |       |       |         |         |          |     |           |    | c.2440A>C (p.Thr814Pro) |                  |                                     |
Table 3. Cont.

| Patient | Age | Gender | Gonads | Muller Derivatives | External genitalia | Puberty | FSH LH | Hormonal | Other | Karyotype | SRY | Variant (HGVS) | Ziposity | Variant Interpretation (ACMG Criteria) |
|---------|-----|--------|--------|-------------------|-------------------|---------|--------|----------|-------|-----------|-----|----------------|----------|---------------------------------------|
| p13     | 22  | M      | Bilateral cryptorchidism | No | Masculine Prader 5 | Tanner 5 | N      | N        | no    | 46,XY     | +   | AR(NM_000044.6): | Heterozygous | VUS (PM2, PP2, PP3) |
|         |     |        |        |                   |                  |         |        |          |       |           |     | c.1097A>C (p.Asn366Thr) MAF = 0 |              |                                      |
| p15     | 0.1 | M      | Bilateral cryptorchidism | No | Micro-penis, hypospadias, one orifice at the base, scrotal folds, inguinal hernia | Tanner 1 | Prepuberty level | N      | no    | 46,XY     | +   | NR0B1(NM_000475.5): | Heterozygous | Likely pathogenic (PVS1, PM2) |
|         |     |        |        |                   |                  |         |        |          |       |           |     | c.512G>A (p.Trp171Ter); MAF = 0 |              |                                      |
| p16     | 1.2 | M      | Scrotal testes | No | Micro-penis, penoscrotal hypospadias | Tanner 1 | Prepuberty level | N      | no    | 46,XY     | +   | AR(NM_000044.6): | Heterozygous | Likely pathogenic (PM1, PM2, PP2, PP3) |
|         |     |        |        |                   |                  |         |        |          |       |           |     | c.2415C>G (p.Phe805Leu) MAF = 0 |              |                                      |
| p17     | 4   | M      | Scrotal testes | No | Hypospadias | Tanner 1 | Prepuberty level | N      | Pierre Robin sequence, calos body hypoplasia, hydrocephaly, GDD/ID | 46,XY     | +   | EMX2(NM_004098): | Heterozygous | VUS (PM2, PP3) |
|         |     |        |        |                   |                  |         |        |          |       |           |     | c.614G>A (p.Arg205Gln) MAF = 0 |              |                                      |
Table 3. Cont.

| Patient | Age | Gender | Gonads | Muller Derivatives | External genitalia | Puberty | FSH LH | Hormonal | Other | Karyotype | SRY | Variant (HGVS) | Zygosity | Variant Interpretation (ACMG Criteria) |
|---------|-----|--------|--------|-------------------|-------------------|---------|-------|----------|-------|-----------|-----|----------------|----------|--------------------------------------|
| p19     | 48  | F      | Bilateral cryptorchidism | no | Female, upper region of vagina without external orifice | Tanner 1 | ↑↑↑ | Adrenal insufficiency | arterial hypertension | 46,XY | + | CYP17A1(NM_000102.4): c.1318C>T (p.Arg440Cys) | Heterozygous | Likely pathogenic (PM2, PM5, PP2, PP3) |
|         |     |        |        |                   |                   |         |       |          |       |           |     | CYP17A1(NM_000102.4): c.1214A>G(p.Glu405Gly) | Heterozygous | VUS (PM2, PP2, PP3) |
| p20     | 4   | M      | Bilateral cryptorchidism | No | Male, Prader 5 | Tanner 1 | Prepuberty level | Adrenal insufficiency | 46,XY | + | - | - | - | - | |
| p22     | 15  | M      | Scrotal testes | No | Male N | Tanner 3 | Ginecomastia | N | T↑ estradiol↑ | no | 46,XY | + | - | - | - | - |
| p23     | 40  | F      | Not evidenced | Yes, uterus, vagina | Female N | Tanner 3 | N | N | no | 46,XX | - | - | - | - | - |
| p25     | 1   | M      | Bilateral cryptorchidism | No | micro-penis | Tanner 1 | ↓ | N | no | 46,XY | + | CYP17A1(NM_000102.4): c.236T>C (p.Leu79Pro) | Heterozygous | Likely pathogenic (PM1, PM2, PP2, PP3) |

MAF = 0
| Patient | Age | Gender | Gonads | Muller Derivatives | External genitalia | Puberty | FSH LH | Hormonal | Other | Karyotype | SRY | Variant (HGVS) | Zigosity | Variant Interpretation (ACMG Criteria) |
|---------|-----|--------|--------|-------------------|------------------|---------|-------|---------|-------|-----------|-----|----------------|---------|--------------------------------------|
| p27     | 14  | M      | Bilateral cryptorchidism | No | Micro-penis, penoscrotal hypospadias | Tanner 4, Ginecomastia | N | N | Obesity, short stature, astigmatism, hypermetropia | 46,XY | + | - | - |
| p28     | 6   | M      | Bilateral cryptorchidism | No | penoscrotal hypospadias | Tanner 1 Prepuberty level | N | | Dysmorphism, aortic bicuspidia, ventricular septal defect, GDD/ID | 46,XY | + | GATA4(NM_002052.5): c.698C>A (p.Thr233Lys) Heterozygous | Likely pathogenic (PM1, PM2, PP2, PP3) |
| p29     | 0.8 | M      | Unilateral cryptorchidism | No | Penoclitoridian glans, one orifice at the base, labioscrotal folds, one gonad at the left fold, Prader 3 | Tanner 1 Prepuberty level | N | no | 46,XY | + | - | - |
| p33     | 20  | F      | Ovarian agenesis | Yes, uterus, vagina | Female | Tanner 3 | ↑↑↑ | N | no | 46,XX | - | - | - |
| p34     | 32  | F      | Ovarian agenesis | Yes, uterus, vagina | Female | Tanner 3 | ↑↑↑ | N | no | 46,XX | - | ATM(NM_000051.4): c.1079G>T (p.Glu359Ter) Heterozygous | Pathogenic (PVS1, PM2, PP3) |

VUS—variant of unknown significance, MAF—minor allele frequency, GDD—global developmental delay, ID—intellectual disability, N—normal values, M—male, F—female.
Four 46,XY patients (p5, p8, p13, and p16) presented variants in the AR gene. In patient p5, the variant c.2071_2073del was classified as likely pathogenic (an in-frame deletion, not described in gnomAD, found in the hot-spot region of the AR gene, predicted as pathogenic by phyloP). Patient p8 presented two VUS variants, c.167T>A and c.170_171insGGACGACGCA (not found in gnomAD, predicted as pathogenic by phyloP). In patient p13, the variant c.1097A>C was classified as VUS (missense variant in exon 1, not found in gnomAD, 96% of missense variants in the AR gene are pathogenic, prediction as pathogenic by the prediction platforms BayesDel_addAF, DANN, FATHMM-MKL, LIST-S2, M-CAP, MVP and SIFT). For patient p16, the variant in AR gene was interpreted as likely pathogenic (a missense variant in exon 6, in a known hot-spot mutation, not found in gnomAD and predictions of pathogenicity in: BayesDel_addAF, DANN, FATHMM-MKL, LIST-S2, M-CAP, MVP, MutationTaster, PrimateAI, and SIFT).

Patient p7 presented a pathogenic variant on the NIPBL gene (a null frameshift variant, not found in gnomAD, Clinvar classification as pathogenic, pathogenic prediction on phyloP). Patient p11 presented two variants in the AMHR2 gene, one pathogenic (c.233-2A>G, null variant within splice site, not found in gnomAD, and predictions sustained the pathogenicity: BayesDel_addAF, EIGEN, FATHMM-MKL, MutationTaster, and scSNV-Splicing) and the other likely pathogenic (c.133delA, null frameshift variant, not found in gnomAD). Patient p15 had a likely pathogenic variant, c.512G>A, in the NR0B1 gene (null nonsense variant, not found in gnomAD). Patient p17 presented a VUS variant, c.614G>A, in the EMX2 gene (not found in gnomAD, pathogenicity predictions from: BayesDel_addAF, DANN, DEOGEN2, EIGEN, FATHMM-MKL, LIST-S2, M-CAP, MVP, MutationTaster, and PrimateAI). Patient p19 presented two variants in CYP17A1, one classified as likely pathogenic (c.1318C>T, a missense variant in exon 8, very small frequency on gnomAD, 97% of missense variants are pathogenic, the alternative variant was classified pathogenic by Uniprot, predictions for pathogenicity in: BayesDel_addAF, DANN, DEOGEN2, EIGEN, FATHMM-MKL, LIST-S2, M-CAP, MVP, MutationAssessor, MutationTaster, and SIFT) and the other VUS (c.1214A>G, a missense variant in exon 7, not found in gnomAD with predictions of pathogenicity in DANN, DEOGEN2, EIGEN, FATHMM-MKL, LIST-S2, M-CAP, MutationAssessor, MutationTaster, and SIFT).

Patient p25 presented the variant c.1214A>G in the GNRHR gene, which was noted as likely pathogenic (missense mutation in a functional domain of the gene, not found in gnomAD with predictions of pathogenicity in: BayesDel_addAF, DANN, DEOGEN2, EIGEN, FATHMM-MKL, M-CAP, MVP, MutationAssessor, MutationTaster, and SIFT). However, this gene is autosomal recessive, and the variant, heterozygous, was not sufficient to develop the pathology.

Patient p28 presented the variant c.698C>A in the GATA4 gene, which was classified as likely pathogenic (a missense variant in a functional domain of the gene, not found in gnomAD with predictions of pathogenicity in BayesDel_addAF, DANN, DEOGEN2, EIGEN, FATHMM-MKL, M-CAP, MVP, MutationTaster, PrimateAI and SIFT).

4. Discussion

In this study of 267 patients with DSD, of the 93 patients that completed all applicable tests, a diagnosis was obtained for 87 (94%) of the patients. A total of 174 patients did not pursue genetic testing and remained etiology unknown following karyotyping, SRY testing, and CYP21A2 strip testing. Karyotype testing established the diagnosis in 15% of patients, most of whom presented abnormalities of the sex chromosomes. A value of basal 17-hydroxyprogesterone above the threshold of 2 ng/mL or of stimulated 17-hydroxyprogesterone above 10 ng/mL indicated a deficiency of 21-hydroxylase, and genetic testing of CYP21A2 confirmed the diagnosis in 17 out of 39 patients (44% of CYP21A2 tested patients).

SNP array analysis was particularly useful in DSD patients who presented associated signs (syndromic DSD). Of patients tested by SNP array, 20% received a final etiologic diagnosis, and almost all of them, with one exception, presented syndromic DSD (this...
percentage was similar between 46,XX and 46,XY DSD). Gene panel sequencing, which included DSD-associated genes (known and candidate) established or strongly suggested the etiologic diagnosis in 11 of 22 patients tested (50%). The androgen receptor gene mutations were most observed in the study group, but variants in some genes less commonly associated with DSD were also observed, such as those in the GATA4 gene.

Regarding the percentage of chromosomal abnormalities observed in patients with DSD, a similar result (around 15%) has been found by other studies, and most of these abnormalities involve the sex chromosomes [18,19]. Thus, karyotype has a role not only in establishing the first step in a pathogenetic algorithm (in 46,XX or 46,XY DSD) but also in observing an etiology in a number of cases (in sex chromosome DSD). The karyotype also provides advantages for identifying mosaicism, translocations, and X chromosome structural variants, which in some situations are not easily suggested using chromosomal microarray [13].

Concerning the CNVs observed in the study group, the percentage of pathogenic CNVs and uniparental disomy was 20%, similar to that found in other studies, which ranged from 15–20% [18,20–23]. One study found that chromosomal microarray is similar to classical cytogenetics when identifying Turner syndrome [24]. However, chromosomal microarray analysis has advantages over karyotype regarding the identification of cryptic Y chromosome material in patients with Turner syndrome [13]. For example, 1q21 deletion (involving RBM8A gene), identified in a 46,XX patient, is a described etiology in Mayer-Rokitansky-Küster-Hauser syndrome, which is present in some patients with this BP2-BF3 deletion [25].

Another patient was diagnosed with 21-hydroxylase deficiency due to homozygous deletion identified by SNP array testing. The presence of maternal uniparental disomy of 15 and uniparental disomy of chromosome 7 in three of the 35 patients studied by chromosomal microarray provided useful information, as the phenotype of these disorders (Prader-Willi or Russell-Silver syndromes) includes signs of DSD usually associated with other findings that are less obvious. Concerning the other pathogenic CNVs or VUS described in the study, a clear association was not found between genes included in these regions and DSD; these CNVs or genes have been more frequently associated with neurologic development than that of other systems or organs.

High-throughput sequencing was found to be the most effective in cases of DSD patients due to its high percentages of positive results. The findings of high-throughput sequencing indicated etiologic and pathogenetic mechanisms, which could be very important for designing optimal therapy and achieving a better prognosis. Of the study’s patients, 50% had a pathogenic or likely pathogenic variant responsible for their clinical development, and a similar percentage was observed in a large international cohort of DSD patients [26]. However, this percentage was higher than that observed in other studies on DSD patients (around 30%) [27–32]. This higher percentage may have been due to the fact that the patients tested by sequencing comprised those with a severe DSD phenotype [28]. A similar percentage of diagnosis was found for both 46,XY and 46,XX patients. The genes that presented pathogenic or likely pathogenic variants or VUS were RSPO1, FGFR1, WT1, CHD7, AR, NIPBL, AMHR2, AR, EMX2, CYP17A1, NR0B1, GNRHR, GATA4, and ATM genes. As in other studies, the frequency of AR gene mutations was higher than that of other genes [28,33–35].

The variant described in the GATA4 gene, c.698C>A (p.Thr233Lys), fell within the N-terminal zinc finger region, one that was recently proved by van der Bergen to have a pathogenic effect in 46,XY DSD, unlike variants in other regions of this gene, which usually have a benign contribution [36]. Another pathogenic variant identified in the present study was in the NIPBL gene; this variant is associated with Cornelia de Lange syndrome. The genetic diagnosis for this patient was due to genital abnormalities seen and less to the dysmorphic and neuropsychiatric signs classically associated with this syndrome.

A limitation of the present study was the impossibility of evaluating the parents to establish the de novo or inherited characteristics of the unknown variants, as well as
the inability to perform functional studies for these variants. However, an important contribution of this study was that it was the first in the authors' country to perform a well-defined algorithm for genetic testing in DSD.

In conclusion, an evaluation following a genetic testing algorithm including karyotype and SRY gene testing, CYP21A2 analysis, chromosomal analysis by microarray, and next-generation sequencing provided a diagnosis for 87 patients with a spectrum of diagnostic yield between 15 and 50%, depending on the technique. Additionally, new genetic variants not previously described in DSD patients were observed.

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