Integrating clinical and genetic approaches in the diagnosis of 46,XY disorders of sex development

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Abstract

46,XY differences and/or disorders of sex development (DSD) are clinically and genetically heterogeneous conditions. Although complete androgen insensitivity syndrome has a strong genotype–phenotype correlation, the other types of 46,XY DSD are less well defined, and thus, the precise diagnosis is challenging. This study focused on comparing the relationship between clinical assessment and genetic findings in a cohort of well-phenotyped patients with 46,XY DSD. The study was an analysis of clinical investigations followed by genetic testing performed on 35 patients presenting to a single center. The clinical assessment included external masculinization score (EMS), endocrine profiling and radiological evaluation. Array-comparative genomic hybridization (array-CGH) and sequencing of DSD-related genes were performed. Using an integrated approach, reaching the definitive diagnosis was possible in 12 children. The correlation between clinical and genetic findings was higher in patients with a more severe phenotype (median EMS 2.5 vs 6; P=0.04). However, in 13 children, at least one variant of uncertain significance was identified, and most times this variant did not correspond to the original clinical diagnosis. In three patients, the genetic studies guided further clinical assessment which resulted in a reclassification of initial clinical diagnosis. Furthermore, we identified eight patients harboring variants in more than one DSD genes, which was not seen in controls (2.5%; P=0.0003). In summary, taking into account potential challenges in reaching the definitive diagnosis in 46,XY DSD, only integrated approach seems to be the best routine practice.

Introduction

Differences and/or disorders of sex development (DSD) represent rare congenital conditions in which gonadal, chromosomal or anatomical sex is atypical. Due to its heterogeneous clinical presentation and genetic architecture, achieving a definitive diagnosis is challenging, especially when associated with 46,XY karyotype (1). Indeed, with the exception of several conditions with well-defined phenotypes that point to a specific underlying...
genetic cause, i.e. complete androgen insensitivity (CAIS) (2), in the majority of cases neither the clinical presentation nor the phenotype-genotype correlation is indicative enough to predict a precise diagnosis.

The observed phenotypic heterogeneity is partially explained by complex mechanisms that control male sex development, and when disturbed may result in disorder of gonadal development (DGD), disorder of androgen synthesis (DAS) or disorder of androgen action (DAA) (1). Similar to the diverse pathophysiology, the genetic background varies with more than 40 genes having been implicated in its pathogenicity and associated with different modes of inheritance including autosomal dominant, autosomal recessive, X-linked and de novo (1). Moreover, some DSD conditions may result from copy number variations (CNVs), especially in cases with gonadal dysgenesis or in the presence of associated malformations (3, 4, 5). In addition, recent findings (6, 7, 8) indicate oligogenicity, defined as mutation in more than one gene, as a possible mode of inheritance in DSD.

Therefore, in the light of phenotypic variability and the multiple genes associated with the phenotype, the introduction of next-generation sequencing (NGS) methods into the field of DSD significantly improved the diagnostic yield (6, 9). While whole-exome and/or genome sequencing (WES/WGS) are still reserved for the research environment, targeted gene panels are becoming widely used in the diagnostic algorithms of 46,XY DSD on a routine basis (10). However, as their utility was mostly proven in research genetic centers, there are still missing data regarding its application in a clinical setting, where reporting the genetic result to the patient and their family might be complicated by low sensitivity and/or specificity of endocrine tests that are unable to fully define the phenotype (11), potentially inconclusive results with the identification of a variant of uncertain significance or the increasingly stronger argument for oligogenicity as a possible mode of inheritance. This study aimed to compare the relationship between the clinical diagnosis (based on physical examination, biochemical and radiological assessment) to the result of genetic assessment using array-comparative genomic hybridization (array-CGH) and sequencing 37 genes known to underlie 46,XY DSD in a cohort of 35 patients.

Material and methods

Study group

The study group initially consisted of 36 patients (34 probands) diagnosed with 46,XY DSD who were referred to a multidisciplinary clinic at a single pediatric endocrinology department for further investigations. Patients presented with either atypical genitalia in infants and children or atypical sexual development in adolescents. The appearance of external genitalia was described using the external masculinization score (EMS) (12), and consistent with UK guidance (13), only those with an EMS less than 11 were included. During the routine genetic testing prior to this study, one patient with complete gonadal dysgenesis was found to harbor a causative mutation in the SRY gene, and therefore, was excluded from the present study. Among 35 patients, 21 were raised as boys, while the remaining 14 were raised as girls. The cohort was classified according to above mentioned guidelines (13), with an exception of a diagnosis of a non-specific disorder of undermasculinization (NSDUM) assigned to those boys with normal testicular function before the analysis of the AR gene (14), and a diagnosis of syndromic 46,XY DSD assigned to those with multiple (>1) associated conditions. The definition of small for gestational age (SGA) was a birth weight less than the tenth centile.

The study was approved by the Poznan University of Medical Sciences Medical Ethical Committee (Registration number 505/13) and informed consent was obtained from all participants.

Endocrine investigations

Standard hormonal measurements were performed using commercial kits. Luteinizing hormone (LH), follicle-stimulating hormone (FSH) and estradiol (E2) were assessed using chemiluminescent microparticle immunoassay (CMIA) by Architect (Abbott Laboratories) and dihydrotestosterone (DHT) by DHT-optimized ELISA (DRG Instruments GmbH, Germany). Androstenedione (A) and testosterone were measured by radioimmunoassays: Active Androstenedione RIA (Beckman Coulter, Immunotech) and TESTO-RIA-CT (DIAsource ImmunoAssays S.A., Belgium), respectively. Anti-Müllerian hormone (AMH) and inhibin B were evaluated by ELx800 Absorbance Microplate Reader (BioTek Instruments) using Gen II ELISA assay (Beckman Coulter, Diagnostic Systems Laboratories, Inc.). The protocol for the hCG test was based on the administration of 1000 IU of hCG (Pregnyl) intramuscularly for three consecutive days with blood drawn immediately prior to the test on day one and 24 h after the last injection. An adequate increase of testosterone after hCG was considered to be sufficient if testosterone was over 3.5 (nmol/L) (15). Testosterone/DHT ratio was recognized
as elevated if over 25 (16), while testosterone/A ratio was considered as decreased if below 0.8 (17). The GnRH test was conducting using an intravenous administration of 2.5 µg of gonadorelinum (Relefact LH-RH) per kilogram (max. 100 µg) with a blood drawn prior to the test and at 30min and at 60min after injection. The assessment of internal genitalia (presence or absence of uterus, localization of undescended gonads) was performed using transabdominal ultrasound examination.

Genetic investigations

After a thorough literature and database search (OMIM, HGMD and ClinVar), a list of 37 genes implicated in 46,XY DSD and included in the TruSight One sequencing kit (Illumina) were selected for evaluation (Table 1). If the gene was only reported in a single human case or family, then strong supporting in vitro or in vivo data must also be present for the gene to be included (CBX2, FGFR2, WWOX, and TSPY1) as stated in ‘Guidelines for investigating causality of sequence variants in human disease’ (18). Following other studies on the genetic etiology of 46,XY DSD (6, 7), we also evaluated a list of 21 genes implicated in congenital hypogonadotropic hypogonadism (CHH) (Supplementary Table 1, see section on supplementary data given at the end of this article), a condition that shares some clinical presentations with DSD (i.e. cryptorchidism or micropenis) to further study their contribution. Reference transcript numbers for all genes evaluated are noted in Supplementary Table 2.

Blood samples were collected and frozen at −20°C until the analysis, and molecular studies were performed on genomic DNA isolated from peripheral blood leukocytes using FlexiGene DNA Kit (QIAGEN).

Array-CGH was performed using a custom whole genome 180 K oligonucleotide array (Sure Print G3 Human 4x180K) manufactured by Agilent Technologies. The median probe spacing for the arrays was 11 kb. Digestion, labeling and hybridization were completed according to manufacturer’s protocols. Scanning and image analysis were performed as per oligonucleotide array-CGH protocol (Agilent, version 4.0). Microarrays were scanned using the Agilent Scanner. Agilent CytoGenomics Analytics software (v4.0.3.12) was used to visualize, detect and analyze the CNVs from microarray profiles using Tiff images from data. Identified CNVs were cross-checked with data gathered from the following databases: DGV (Database of Genomic Variants) (http://dgv.tcag.ca/dgv/app/home), ISCA (International Standard for Cytogenomic Arrays) (http://dbsearch.clinicalgenome.org/search), NCBI (National Center for Biotechnology Information) (https://ncbi.nlm.nih.gov), as well as a local database of CNV.

For sequencing of the 37 DSD-related genes, patients' DNA samples were normalized to 5 ng/µL in 10 µL of Tris–HCl 10 mM pH 8.5 (total DNA mass 50 ng). Libraries were constructed for NGS using the TruSight One Sequencing Panel Kit (Illumina, Inc.) according to the manufacturer’s protocol, and 150 basepair paired-end reads were generated on the MiSeq instrument (Illumina). Sample runs were validated as successful if the sequencing run yield was >7.5 Gb, had >96% of reads aligning to the human genome and had an overall 30× coverage of >95%. Each run included 3 samples. Only variants with a read depth coverage of >10× and a quality score of >95 passed quality control. Given these stringent parameters for genotype calls, Sanger sequencing was not required for further confirmation (19).

The raw data (fastq files) from the MiSeq sequencing of the TruSight One libraries were uploaded to Saphetor (www.saphetor.com, Lausanne, Switzerland); however, a mask was applied to the data such that only the 37 DSD-related genes were analyzed. This analysis consisted of read alignments, basepair calling and annotation with HGVS (Human Genome Variation Society) nomenclature, function and conservation. Splice-junction (+6 base pairs) and non-synonymous variants with a minor allele frequency (MAF) <1% in Genome Aggregation Database (GnomAD) were evaluated. Of note, a missense variant was considered to be damaging if it was predicted to be damaging by one of the two in silico predictive algorithms, SIFT and/or MutationTaster. The variants were classified into five categories according to the American College of Medical Genetics (ACMG) guidelines (20) using the InterVar program (http://wintervar.wglab.org/): pathogenic, likely pathogenic and variants of uncertain significance (VUS), likely benign, and benign. Variants classified as benign or likely benign were not included in this study. Additionally, as previously described (6), any expansions in the first exon of the AR gene of CAG over 27 repeats (21) or CGG over 24 repeats (22) were classified as VUS, as their contribution into DSD phenotype is still uncertain. A definitive (genetically confirmed) diagnosis was considered to be achieved when pathogenic or likely pathogenic variants were identified.

Oligogenicity of the DSD cohort was assessed against the ‘Cohorte Lausannoise’ (CoLaus) control population, consisting of 405 participants of mixed European origin, phenotyped as described by Firmann et al., in whom whole exome sequencing was performed (23).
Consistent with the DSD cases in the study, only genotype calls with read depths >10× and quality scores >95 were used. Additionally, the 37 DSD-related genes were covered in both the whole exome data of the CoLaus cohort and the TruSight One Sequencing Panel Kit of the DSD cohort to an average read depth of >20× for >99% of the nucleotides in the genes.

Case vs control statistical analysis for an enrichment of individual variants or oligogenicity was performed using a two-tailed Fisher’s exact test. The Mann–Whitney U test was performed to assess the difference in EMS in cases reaching the definitive diagnosis vs cases with no final diagnosis. A statistic of $P<0.05$ was considered to be significant.
Results

Clinical approach based on physical and endocrine evaluation

After structured clinical assessment followed by thorough endocrine evaluation including biochemical and radiological tests, each case was attributed a clinical diagnosis. All data regarding phenotyping are provided in Supplementary Tables 3 and 4. The median EMS was 4.5 (0–9). In all cases with low plasma level of testosterone (basal value if assessed during expected minipuberty/puberty or after hCG test if assessed during childhood) and AMH, a subcategory of DGD was attributed. Among this subgroup, there were (i) five patients with testicular regression syndrome (TRS), as despite lack of testicles they presented with male external genitalia and no Müllerian structures were identified on the ultrasound and (ii) four patients with complete gonadal dysgenesis (CGD), as they presented with female external genitalia, high plasma gonadotropins and an infantile uterus. Additionally, for patients with low plasma level of testosterone or high testosterone/DHT ratio, an AMH serum level within normal ranges, and no Müllerian structures, a subcategory of DAS (disorder of androgen synthesis) was established. Namely, there were three patients suspected for a 5α-reductase deficiency (5αRD) and one patient with undetectable testosterone plasma level and no uterus on the ultrasound indicating proper level of AMH during fetal life. In addition, a subcategory of disorder of androgen action (DAA) was attributed to those with proper gonadal function reflected by normal to high plasma levels of testosterone and AMH and no Müllerian structures. Namely, six patients with atypical genitalia received a phenotypic diagnosis of NSDUM, and eight patients were diagnosed with either a complete or partial form of androgen insensitivity syndrome (CAIS or PAIS) depending on the appearance of the external genitalia. Lastly, there were eight children that received a phenotypic diagnosis of syndromic DSD, as they presented with multiple associated conditions (i.e.: psychomotor delay, dysmorphic features, congenital heart defect). Additionally, although the majority of children presented with normal adrenal function, there were two syndromic patients with a primary (ID 32) and a secondary adrenal insufficiency (ID 30). (Fig. 1, Supplementary Tables 3 and 4).

Array-CGH

Given the incidence of genomic rearrangements in patients with DGD and syndromic forms of DSD, array-CGH was first performed on 10 of the 17 patients with CGD, TRS or sDSD (Fig. 2). No pathogenic genomic structural abnormalities were identified in these patients.

Sequencing results and their relationship to clinical diagnosis

Sequencing of the 37 genes implicated in DSD was performed successfully in all 35 patients (Fig. 2). The results were filtered according to the strategy described in the methods, including ACMG variant classification. This analysis identified 11 pathogenic variants, 2 likely pathogenic variants and 17 VUS (Table 2). Interestingly, each patient harbored at least one variant in 15 out of the 37 DSD-related genes, and most frequently, these were missense mutations. There were recurrent mutations in the AR and HSD17B3 genes in unrelated patients (Table 2).

As a definitive genetic diagnosis was considered when pathogenic or likely pathogenic variants were identified, 12 cases were assigned a definitive genetic diagnosis. In the remaining patients, the final diagnosis remained at the clinical level, as either no variant was identified (n=15) or at least one VUS was found (n=8) (Fig. 3). In the majority of cases, the variant was classified as VUS primarily due to a discordance between the phenotype and genetic mutation or, in cases with expansion of the number of CAG repeats, due to undetermined significance of its contribution to the DSD phenotype. Overall, 13 patients were found to harbor at least one VUS.

Among those with a genetically confirmed diagnosis (n=12), nine patients had a clinical and endocrine evaluation that enabled an accurate initial identification of the underlying genetic cause. Namely, in seven patients suspected for CAIS and one for NSDUM, the pathogenic variant was identified in the AR gene, as well as one case classified with DAS was found to harbor compound heterozygote variants in the HSD17B3 gene. However, in three patients, the result of the genetic assessment led to the re-evaluation of the initial clinical diagnosis which subsequently led to diagnostic reclassification. Specifically, in patients suspected for a disorder of androgen action (ID 14 and 27), variants in the WT1 and NR5A1 genes, respectively, were identified indicating a disorder of gonadal dysgenesis. Additionally, a patient initially classified with 5α-reductase deficiency (ID 11) was found to harbor a likely pathogenic variant in the HOXA13 gene implicated in syndromic forms of DSD.
Factors correlating with achieving a definitive genetic diagnosis

Patients for whom a definitive genetic diagnosis was rendered had a significantly lower mean EMS score (i.e., had a more severe phenotype), than patients with either a VUS or with no variant (2.5 vs 6, \( P = 0.04 \)). Additionally, the highest genetic diagnostic yield was seen in cases initially suspected of DAA (10 patients out of 14), followed by patients classified as DAS (two patients out of four). However, a final genetic diagnosis was not possible in any of the patients suspected of DGD or sDSD.

Oligogenic inheritance in DSD patients

There were eight patients (23%) harboring a mutation in more than one DSD gene (Table 2). To evaluate whether the DSD cohort was statistically enriched for oligogenicity, we compared the results to exome sequencing data from the 247 male participants in the ‘Cohorte Lausannoise’ CoLaus control population. As ACMG criteria are not intended for use in non-monogenic disorders (20), variants in 37 DSD-related genes were filtered for non-synonymous variants with a MAF <1.0%, including nonsense, splice-site (±6 base pairs) and missense variants found to be damaging in at least one of the two protein prediction programs, SIFT or MutationTaster. There was a statistical enrichment in oligogenic variants in our DSD cohort compared to CoLaus controls (23% vs 2.5%; \( P = 0.0003 \)).

Variants in known CHH genes

Despite the fact that DSD patients typically do not present with hypogonadotropic hypogonadism, there is a small overlap in clinical presentation, namely the presence of micropenis and/or cryptorchidism. Therefore, several groups have previously explored a potential genetic overlap in these patients related to the known CHH genes (6, 7). Similarly, we evaluated our DSD cohort, within which four heterozygous and one hemizygous rare variants in the known CHH genes were identified in five DSD patients (14%) who actually presented with high gonadotrophin plasma level (Supplementary Tables 4 and 5). There was no statistically significant difference...
between the DSD cohort and the CoLaus controls (14% vs 20%, $P=0.3$) at the individual variant level, gene level or overall panel of genes.

**Discussion**

Despite the introduction of NGS methods into the clinical setting, the elucidation of genetic background in 46,XY DSD remains challenging. We presented the results on the integrated approach combining clinical and genetic evaluation of a cohort of 46,XY DSD patients systematically assessed in one center. This integrated approach resulted in identification of genetic findings in one-third of patients, while the final diagnosis in rest of them remained clinical only. As expected, a wide spectrum of phenotypes was observed leading to further subcategorization of the DSD cohort into DGD, DAS, DAA and sDSD. Thus, although the sample size was limited, the cohort appeared to be representative.

Importantly, the NGS of 37 DSD-related genes produced valuable results. The diagnostic yield (i.e. percentage of samples with pathogenic or likely pathogenic genetic results) in our cohort was 34% (12/35), which is comparable to other reports (24, 25). In the majority of patients with a definitive diagnosis (9/12), physical and endocrine evaluation was sensitive and/or specific enough to clearly identify the underlying genetic cause. However, with the exception of one case classified as DAS, this was only true for patients suspected of DAA – a condition with the strongest correlation between phenotype and genotype when presenting with a CAIS phenotype (2).

Given the variable specificity of endocrine tests to accurately diagnose a phenotype (i.e. androgens’ ratios in the diagnosis of DAS (11, 26, 27)) and since some conditions may present with a wide spectrum of phenotypes (i.e. phenotypes associated with genes such as NR5A1 (28) or WT1 (29)), an accurate diagnosis may only be possible with parallel genetic evaluation. Interestingly, the genetic results obtained in the present study lead to the re-evaluation of the clinical diagnosis in 3 patients, and the subsequent reclassification of their diagnosis. Specifically, one patient (ID 11) with distal limb anomalies, an abnormal testosterone/DHT ratio and initially suspected of 5αRD was finally accurately diagnosed with a hand-foot-genital syndrome after the identification of a pathogenic variant in the HOXA13 gene (30). Of note, a clinical diagnosis of suspected 5αRD was not genetically confirmed in any of the patients in our cohort, further confirming the limited utility of testosterone/DHT ratios in the diagnosis of 5α-reductase deficiency. Furthermore, two patients (ID 14 and 27) were initially suspected as having DAA, but were found to harbor likely pathogenic mutations in the WT1 and NR5A1 genes, respectively. It has been established that mutations in both genes are implicated in a broad spectrum of phenotypes. In the case of mutations in the WT1 gene, phenotypes can range from hypospadias (29, 31) to much more severe phenotypes such as WAGR (Wilms tumor, aniridia, genitourinary anomalies, and mental retardation), Frasier or Denys-Drash syndromes (1, 29). Phenotypes for patients carrying mutations in the NR5A1 gene cover a spectrum of different degrees of gonadal dysgenesis, with or without associated adrenal insufficiency (28).
Table 2  Variants in 37 DSD-related genes identified in 35 patients with 46,XY DSD.

<table>
<thead>
<tr>
<th>ID</th>
<th>Sex</th>
<th>DSD category</th>
<th>Clinical diagnosis</th>
<th>DSD gene</th>
<th>Inheritance</th>
<th>GnomAD frequency</th>
<th>DNA change</th>
<th>Protein change</th>
<th>SIFT</th>
<th>MT</th>
<th>ACMG classification</th>
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<td>VUS</td>
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<td>XL</td>
<td>(CAG)</td>
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<td>VUS</td>
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<td>HOXA13</td>
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</tr>
<tr>
<td>13</td>
<td>F</td>
<td>DAS</td>
<td>5αRD</td>
<td>αRD</td>
<td>AD          c.539C&gt;T</td>
<td>p.Pro180Leu</td>
<td>D</td>
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<td>D</td>
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<td>Likely pathogenic</td>
</tr>
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<td>p.Phe439Leu</td>
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<td>HSD17B3</td>
<td>AR</td>
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<td>p.Ile438delinsGlyArgPro</td>
<td>Pathogenic</td>
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<td>p.Leu202Arg</td>
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<td>XL</td>
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<td>c.1822C&gt;T</td>
<td>p.Arg608*</td>
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<td>XL</td>
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<td>34</td>
<td>F</td>
<td>DAA</td>
<td>NSDUM</td>
<td>AR</td>
<td>XL</td>
<td>0.03%</td>
<td>c.1822C&gt;T</td>
<td>p.Arg608*</td>
<td>Pathogenic</td>
<td>D</td>
<td>T</td>
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<td>35</td>
<td>M</td>
<td>DAA</td>
<td>NSDUM</td>
<td>AR</td>
<td>XL</td>
<td>0.03%</td>
<td>c.2222C&gt;T</td>
<td>p.Asp696Aan</td>
<td>Pathogenic</td>
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All variants listed are heterozygous, with the exception of changes in AR and ARX, which are hemizygous as these genes are on the X chromosome. The number of CAG repeats leading to the expansion of the polyglutamine tract in the AR gene is specified with the index number.

**Abbreviations:** DSD, disorder of sex development; GS, gonadal development; T, testicular regression syndrome; U, unknown; VUS, variant of uncertain significance; XL, X-linked.

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Our study showed that patients with a more severe phenotype (mean EMS score 2.5) were more likely to have a conclusive genetic diagnosis than those with less severe phenotypes (mean EMS score 6), $P=0.04$. This lower diagnostic yield in children with a less severe phenotype is likely due to several reasons. Since it is difficult to definitively draw a line between the DSD and the mild and common urogenital defect such as hypospadias (14) that can be caused by environmental factors as well as genetic polymorphisms (31, 32), the enrollment of cases with signs of undermasculinization into the study group may have resulted in a bias. However, as it is known that the degree of undermasculinization reflected by lower EMS does not always increase the likelihood of a diagnostic yield (5, 33), it is debatable to exclude mild DSD cases from systematic assessment.

The diagnostic rate differed among DSD subcategories. The highest diagnostic rate (10 cases out of 14) was seen among suspected DAA patients, followed by the DAS subgroup (two cases out of four). Despite the small sample size, these findings are consistent with the results of a targeted gene study on the largest DSD cohort to date (6). However, in contrast with other report, no definitive genetic diagnosis was able to be rendered in patients with syndromic DSD or DGD. One of the reasons might have been the inclusion criteria, as only those patients with CGD who were ‘SRY negative’ prior to the study were enrolled. Since the SRY gene constitutes the primary candidate gene for this disorder (1), the likelihood of identifying the genetic etiology of gonadal dysgenesis among ‘SRY negative’ cases is lower. Furthermore, the subgroup with DGD was represented by cases with TRS, a condition in which environmental factors also play a role (34). Thus, although in some patients the genetic causes were identified in TRS (35, 36), the possible presence of non-genetic effects lowers the odds of a genetic diagnosis.

In addition, although the published incidence of genomic rearrangements in DGD or sDSD patients varies between 20 and 30% (3, 4, 5, 11), our current study did not identify any pathogenic CNV in these patients. These negative array-CGH results are likely due the limited number of cases assessed ($n=10$). Finally, the full spectrum of the genetic determinants of DGD and sDSD is yet to be elucidated; thus, a genetic diagnosis can only be given for a minority of cases (1).

The sequencing of 37 DSD-related genes led to the identification of eight patients (23%) exhibiting oligogenic inheritance of more than one mutation in DSD genes, and this was not observed in controls (2.5%; $P=0.0003$). This finding further reinforces recent studies demonstrating that oligogenicity is present in DSD patients and may at least partly explain the incomplete penetrance and variable expressivity within and across DSD families (6, 7, 8). It is important to note, however, that our current study evaluated oligogenicity in only DSD genes, and did not include variants in known CHH genes. The fact that this strong association is present even when not including CHH genes (as previous studies have done (6, 7)), coupled with a statistically equal number of CHH variants in both our DSD cohort and the CoLauS control population suggests a minimal or at best modifying role of CHH genes in the pathogenesis of DSD. Further, the role of CHH genes in DSD is debatable, given their distinct pathogenesis and endocrine profiles, as well as our results showing a nearly identical frequency of rare CHH variants in both DSD patients and CoLauS controls. However, larger genetic studies coupled with in vivo and/or in vitro functional studies will help to further clarify this.

In summary, the study clearly showed that an integrated approach is the best routine practice in the
diagnosis of 46,XY DSD. In those cases with a wide phenotype variability and reduced utility of conventional endocrine tests to fully determine the phenotype, reaching the definitive diagnosis may only be possible with parallel genetic testing. On the other hand, substantial prevalence of variants of uncertain significance or of possible oligogenic nature with different contribution to the phenotype underlay the importance of thorough clinical assessment. Therefore, in order to overcome potential obstacles in rendering an accurate diagnosis, data from clinical, biochemical and genetic assessment should be analyzed in an integrative manner. This approach should be then followed by reporting the data and comparing with other centers. Furthermore, only one third of patients harbor pathogenic mutations in DSD genes. The remaining patients argue for multi-national studies to identify additional genes involved in the pathogenesis of DSD.

Supplementary data
This is linked to the online version of the paper at https://doi.org/10.1530/EC-18-0472.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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