Clinical Approach for Evaluation and Management of Disorders of Sex Development

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ABSTRACT

"It's a boy" and "It's a girl" are words that are heard every second of every day all around the world. However, it is very distressing when the birth attendants are unable to make such a pronouncement because of disorders of sex development (DSD). DSD are congenital conditions associated with an atypical development of chromosomal, gonadal or anatomical sex. Normal sex development progresses in steps from conception to the complete development of the fetal external genitalia; any disturbance in any of these steps can lead to DSD. Ambiguous genitalia are the most common type of DSD and it is a challenging clinical diagnosis for the pediatric endocrinologist. A newborn baby with ambiguous genitalia is often a surprise for both the medical team and the parents, frequently described as an emergency. The condition needs a special approach in terms of counseling the parents appropriately, evaluation and management. The Chicago Conference (2006) recommended new nomenclature and a classification for DSD, as the old nomenclature was confusing for doctors and parents, and sometimes pejorative. The new classification is based on karyotyping and gonadal structure, improving understanding of the underlying pathogenic mechanisms. The rapid progression of genetic diagnosis of DSD using advanced techniques such as next-generation sequencing (NGS) allows more appropriate diagnosis and genetic counseling for families. The focus of the article is a review of normal sex development, DSD classification, clinical approach, genetic assessment, sex assignment, surgical management and risk of germ cell tumor development.

INTRODUCTION

Normal sex development and differentiation

Sex differentiation is a complex phenomenon that follows five steps: 1-Genetic sex, the conjugation of a sperm and the ovum gives rise to the fertilized egg which has either a 46,XYor 46,XX chromosome (1). 2-Formation of undifferentiated structures, during the first six weeks of gestation the male and female fetus develops similar sex elements, which consist of the following potential structures: I) The gonadal ridge which develops in the testis or ovary. II) Germ cells, which eventually develop into spermatocytes or oocytes. III) Two sets of internal sex ducts (Wolffian ducts in males and Mullerian ducts in females). IV) External genitalia including the genital tubercle, urethral folds, labioscrotal folds and urogenital sinus. 3-Gonadal differentiation, the bipotential gonad develops into testis in the presence of a Y chromosome, which carries the SRY gene (Sex-determining Region Y) responsible for testicular formation. In the absence of a Y chromosome or SRY gene, the gonad develops into an ovary (2). 4-Gonadal hormones, at 7 weeks of gestation the testis secrete two hormones, testosterone and anti-Müllerian hormone (AMH). The testosterone produced by the Leydig cells of the fetal testes and plays an essential role in the differentiation of Wolffian ducts to the epididymis, vasa deferentia and seminal vesicle. The AMH, on the other hand, is secreted by the Sertoli cells of the fetal testes and is responsible for the regression of Müllerian ducts. In females, the Müllerian ducts develop spontaneously into fallopian tubes, uterus and upper third of the vagina and the Wolffian ducts disappear (3). 5-Phenotypic sex differentiation, the cells of the external genitalia of the male fetus has the ability to metabolize testosterone to dihydrotestosterone (DHT), which is
more potent than testosterone and induces masculinization of external genitalia at about 9-12 weeks gestation. In a male fetus, the genital tubercle lengthens and thickens to form the penis, the urethral folds grow to become the penile urethra and the genital swellings enlarge to form the scrotal swelling which fuses in the midline to form the scrotum (4). In female fetuses, the absence of circulating testosterone allows the genital tubercle to develop into the clitoris, the urethral folds into the labia minora and the genital swelling into the labia majora.

Nomenclature and classification of DSD
The preexisting nomenclature of DSD (Table 1) such as intersex, sex reversal, pseudohermaphroditism and hermaphroditism was pejorative and confusing to parents, patients and doctors. As a consequence, a re-examination of DSD nomenclature and different classifications took place. In 2006, international experts in pediatric endocrinology met at the Chicago Conference under the auspices of the European Society for Pediatric Endocrinology and the Pediatric Endocrine Society and recommended new nomenclature (Table 1) for DSD, which was defined as "congenital conditions in which the development of chromosomal, gonadal, or anatomic sex is atypical" (5, 6). This definition is wide and includes any disorder that results in abnormal sex differentiation.

The current nomenclature and classifications (Table 1) are clear, understandable for physicians, patients and their families, and provide information regarding the patient's chromosome and gonadal structure, but is not very specific regarding etiology and depend on the physician's approach and facilities for diagnosis.

**Table 1. Revised nomenclature**

<table>
<thead>
<tr>
<th>Previous</th>
<th>Current</th>
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<tbody>
<tr>
<td>Intersex</td>
<td>Disorders of sex development</td>
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<tr>
<td>Female pseudohermaphrodite</td>
<td>46XX DSD</td>
</tr>
<tr>
<td>Male pseudohermaphrodite</td>
<td>46XY DSD</td>
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<tr>
<td>True hermaphrodite</td>
<td>Ovotesticular DSD</td>
</tr>
<tr>
<td>XX male sex reversal</td>
<td>46XX testicular</td>
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<tr>
<td>46XYcompletegonadal</td>
<td>DSD</td>
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<td>dysgenesis</td>
<td>DSD clinical approach</td>
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**Multidisciplinary team**
The best way to approach DSD is through a multidisciplinary team, which includes a neonatologist, endocrinologist, pediatric surgeon, psychologist and other ad hoc members (7). The first step is to counsel the shocked parents, as the birth of a baby with ambiguous genitalia is unexpected for both parents and the medical team. Parents need support and usually have many questions. The team leader should meet with them and provide initial information and support including offering the family the opportunity to have a supportive family member or a close friend to join them. Also, the role of the leader is to advise the parents to delay the registration of the birth, discuss with them the diagnosis, initial treatment plan, findings relevant to sex assignment, the team's recommendations and options, and to provide information and current evidence regarding gender identity outcomes in similar cases.

**History**
A complete detailed history from the parents is essential for an appropriate diagnosis (5, 8) including unexplained sudden infant death, ambiguity, infertility, precocious puberty, amenorrhea, hirsutism, maternal exposure to hormones (particularly progestagens and steroids), (9, 10) or consanguinity. The possibility of aromatase deficiency or maternal androgen secreting tumors with a history of maternal virilization during pregnancy should be considered (5, 11, 12).

**Physical examination**
Parents should be informed in advance about the method of examination and it should be demonstrated to them. In a warm place, with the baby in the frog leg position, a general inspection is performed focusing on the genital anatomy (5, 8). After inspection, palpate for the gonads within the inguinal canal, scrotum, and labioscrotal folds, using the appropriate technique. The gonads are sometimes not palpable. A palpable gonad is most likely testis or rarely an ovotestis because a streak gonad and ovaries do not descend. Measure an abnormal phallic width and stretched length, examine for fused labia or biffed scrotum and determine the position of urethral opening and other orifices in the perineum. The Prader classification (Fig. 1) provides a useful guide to classify the degree of female virilization. The findings range
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from 1 (mild clitoromegaly) to 5 (complete virilization with urethral opening up at the tip of enlarged phallus) (13). The patient is classified according to the palpable or nonpalpable gonads approach (Fig. 2) for differential diagnosis and investigation purposes.

**Fig. 1.** The degree of virilization of external genitalia of females as proposed by Prader. In Prader 1 the only abnormality is mild clitoromegaly with separate openings for urethra and vagina. Praders 2 through 4 demonstrates the progression of virilization from mild to severe. Prader 5 is a markedly enlarged phallus with penile urethra.

**Fig. 2.** Algorithm for basic investigation for patient with DSD based on gonadal palpation. Abbreviations: DSD, disorder of sex development; hCG, human chorionic gonadotropin. Abbreviations: CAH, congenital adrenal hyperplasia; CAH, congenital adrenal hyperplasia; Tumors (adrenal/ovarian); Aromatase deficiency; Other cases of CAH.
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Fig. 3. Algorithm for diagnosing DSD in patients with impalpable gonads by incorporating findings from karyotyping, imaging studies and other laboratory analyses. Abbreviations: CAH, congenital adrenal hyperplasia; DSD, disorder of sex development; MGD, mixed gonadal dysgenesis.

Basic Investigation
The basic diagnostic tests for all patients are karyotyping and pelvic ultrasound to evaluate the presence of Mullerian structures. Other tests that should be done are, for example, the measurement of serum levels of sodium, potassium and 17-hydroxyprogesterone (17-OHP) on the third day of life or later for patients with nonpalpable gonads. This is important because congenital adrenal hyperplasia (CAH), the most common DSD, may result in a life-threatening salt-losing crisis. In addition, a patient with palpable gonads should undergo a human chorionic gonadotropin (hCG) stimulation test to assess the gonadal production of androgens. Possible diagnosis and further investigation depend on the results of the initial investigations (Fig. 3, 4).

46 XX DSD (female pseudohermaphroditism)
Patients in this category have a normal female karyotype and Mullerian structures but variable degrees of virilization of external genitalia with nonpalpable gonads. An elevated serum 17-OHP indicates the presence of CAH, the frequent cause of 46 XX DSD (14, 15, 16, 17, 18) due to 21-alpha hydroxylase-CYP21 deficiency, which leads to the accumulation of 17-OHP and excessive production of androgens (Fig. 5). Most patients have the salt-wasting form, which can present with a life-threatening adrenal crisis in the second or third week of life (19). If the levels of 17-OHP are normal, other types of CAH should be investigated. For example, an enzymatic defect in the adrenal cortisol synthesis pathway 11β-Hydroxysteroid (11β-OH) or 3β-hydroxysteroid dehydrogenase (3β-HSD), can lead to the excessive production of androgens and the accumulation of their precursors (Fig. 5). Other rare causes of 46XX DSD include medication, tumors (adrenal or ovarian) and aromatase deficiency. The identification of such causes can be accomplished through a complete history, physical examination and the demonstration of a high androgen level.

Fig. 4. Algorithm for diagnosing DSD in patients with palpable gonads by incorporating findings from karyotyping, imaging studies and other laboratory analyses. Either targeted gene analysis or targeted next-generation sequencing is preferable for accurate diagnosis of known gene disorder before an untargeted analysis is performed. Abbreviations: DSD, disorder of sex development; MGD, mixed gonadal dysgenesis; FISH, fluorescent in situ hybridization; hCG, human chorionic gonadotropin; SRY, sex-
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determining region Y protein.

**Fig. 5.** The androgen pathway and the mechanism of virilization caused by the most common form of CAH (21-OH deficiency, 3β-OSH deficiency and 11β-OH deficiency). Note that accumulation of precursors above the enzymatic defect lead to virilization by excessive production of androgen.

**Fig. 6.** Mechanism of virilization of the male external genitalia. Triangular shaped boxes demonstrate the main causes of undervirilization. Abbreviations: PAIS, partial androgen insensitivity syndrome; CAIS, complete androgen insensitivity syndrome

**46XY DSD (male pseudohermaphroditism)**

Patients in this category have a normal male karyotype and absence of Mullerian structures but with variable degrees of undervirilization of external genitalia with palpable gonads in most of cases. Fig. 6 illustrates the mechanism of virilization of the male external genitalia and the main causes of undervirilization. An hCG stimulation test is a reliable test to determine the gonadal function and it assists in diagnosing the most common causes of 46XY DSD [partial androgen insensitivity syndrome (PAIS) and 5α-reductase deficiency (5-ARD)]. In PAIS, testosterone (T) and dihydrotestosterone (DHT) levels are high while in 5-ARD, the ratio of T to DHT is high, unresponsive testosterone indicates gonadal failure due to either structural or hormonal synthesis defects and elevated
testosterone precursors may indicate a defect of testicular testosterone synthesis (Fig. 6). There is no consensus about the way how an hCG stimulation test should be conducted, but the protocol we used is daily intramuscular hCG 1000–1500 units for three consecutive days, and measuring basal serum dehydroepiandrosteronesulfate, androstenedione, T and DHT 24 hours after the last injection (20). Another protocol, twice-weekly injections for 3 weeks, can also be used but it takes much longer. In case of normal hCG results, an adrenocorticotropic hormone (ACTH) stimulation test demonstrates a specific defect in adrenal testosterone syntheses such as 17α-hydroxylase and 17,20-lyase. An objective of the rapid advances in genetic studies for DSD in recent years was to make a definitive diagnosis and provide genetic counseling for families. The 46XY DSD mutation, in particular, has been identified in the androgen receptors in about 50% of PAIS and in the 5-alpha reductase gene (SRD5A2). Although genetic testing is an excellent tool that can aid in the diagnosis, it is expensive, not widely available and cannot replace the clinical approach.

**Ovotesticular disorder of sexual development (true hermaphroditism)**

It is a rare disorder of sexual differentiation with geographic variations in the incidence and karyotyping (21). It constitutes about 3-10% of all DSD (6, 22, 23). However, in Southern Africa, this condition occurs in 51% of DSD patients (21). It is characterized by the presence of gonads of ovotestis or a combination of testis on one side and an ovary or ovotestis on the other side. The main clinical presentation is ambiguous genitalia and a unilateral palpable gonad, which can be either be an ovotestis or testis, usually located in the right side (24, 25, 26). Karyotyping is variable. 46XX is the most frequent and constitutes about 60% of cases (27-30), followed by mosaicism 46XX/46XY (25-30%) and 46XY (10-15%) (30,31). Internal sex duct development usually corresponds with the adjacent gonad. Wolffian duct structures typically develop on the gonadal side that contain testicular tissue whereas Müllarian duct structures tend to be observed on the gonadal side that do not contain testicular tissue (32). The patients with ovotesticular DSD could be found in both the palpable and nonpalpable gonads groups (Fig.3, 4) but 46XX DSD with a palpable gonad should raise a strong suspicion of ovotesticular DSD. An uncertain Müllarian duct structure is identified with ultrasound and the androgen concentration could be normal for the male range if testicular tissue is predominant. A diagnosis of ovotesticular DSD is dependent solely on the gonadal biopsy results, which should show the presence of ovarian follicles and testicular seminiferous tubules (27).

**46XX testicular DSD**

It is a rare disorder with an estimated prevalence of 1/20,000 males. The nomenclature indicates that patients with 46XX testicular DSD have a 46XX karyotype and testes. The appearance of external genitalia depends on the presence of the SRY-gene. Approximately 80-90% of patients is SRY-gene positive (33) and has normal male external genitalia. They usually present with infertility, delayed puberty, gynecomastia and a short stature (34). Endocrine tests show hypergonadotrophic hypogonadism due to testicular failure (35, 32). On the other hand, patients who are SRY-gene negative (10-20%), usually have ambiguous genitalia presenting at birth or infancy (36, 37). Clinically patients will be in the palpable gonads group (Fig.3). Patients with 46 XX karyotype and absent normal female internal ducts should raise the suspicion of 46 XX testicular DSD diagnosis, which could be confirmed by identifying the SRY-gene using fluorescence in situ hybridization (FISH) or chromosomal microarray (CMA).

**46XY complete gonadal dysgenesis (CGD)**

Patients with 46XY CGD are phenotypically females; they have normal female internal ducts (fallopian tubes, uterus and vagina) and external female genitalia. The patient cannot be identified at birth or childhood and is only diagnosed in adolescence or adulthood when they present with primary amenorrhea, delayed puberty but with a normal stature and the absence of dysmorphic features. This disorder is also referred to as Swyer syndrome, named after Swyer and colleagues who initially described the female appearance of a patient with negative X-chromatin, normal stature and having streak gonads (38). The majority of cases of XY CGD are of unknown etiology; SRY mutations and deletions have been identified in about 10-20% of the cases (39, 40). Other deletions that were described include NR5A1
Sex assignment
At birth or infancy sex assignment depends on the following factors: 1) Potential for sexual activity. 2) Potential for fertility. 3) The feasibility of constructive surgery. 4) The type of DSD. 5) Prenatal androgen exposure. 6) Family culture (19, 48). For 46 XX CAH, the most frequent condition of DSD, the consensus is to raise individuals as female because 90% of patients have a female gender identity, however, about 5% of patients have gender dysphoria (49). Most patients with 46 XX CAH have the potential for normal sexual function and fertility. However, some publications recommend male gender assignment for markedly virilized patients (Prader 5 and some Prader 4) (13). The topic is controversial, but currently, the recommendation is to raise all patients with 46XX CAH as females whatever the stage of virilization (23). The recommendation for patients with 46XY complete androgen insensitivity (CAIS) or complete gonadal dysgenesis is to be raised as females as they are accepting themselves as female, have not been exposed in utero to an effective androgen that may affect their identity [48, 50], are phenotypically females and have an adjusted sexual life. However, some of these individuals have been reported as having a male gender identity. Patients with 46XY 5α reductase or 17β hydroxysteroid dehydrogenase-3 deficiency are usually recommended to be raised as males even with severe undervirilization (51). The recommendations reflect the potential for virilization at puberty (33) and the possibility for fertility (52, 53) and are supported by literature, reporting that more than 60% of 5α-reductase-deficient patients and 50% of 17β-hydroxysteroid dehydrogenase-3 deficiency patients, who are assigned as female in infancy, ultimately change their gender role (54). However, there are some case reports about patients with 17β hydroxysteroid dehydrogenase-3 deficiency that were assigned females (41). For patients with incomplete gonadal dysgenesis, PAIS and androgen biosynthetic defects, there is still no consensus or recommendations regarding sex assignment, about 25% of individuals are dissatisfied with the sex of rearing whether they are raised as a boy or a girl (55). The decision of sex assignment in patients with ovotesticular DSD depends on the degree of gonadal function and its effect on genital development, the possibility of fertility and normal sexual activity. The same should be considering for patients with mixed gonadal dysgenesis (MGD). Individuals with cloacalexstrophy are usually raised as females (65%) but there is variability in gender identity outcomes (19).

Surgical management
The time for repair with feminizing genitoplasty, which include clitoroplasty, labioplasty and vaginoplasty is not well defined; most surgeons prefer early surgery in infancy and in a specialized center for patients with significant virilization (Prader 3 or higher) (19, 56, 7). Studies reported that early surgery may improve the relationship between children and their parents and relieve parental distress (57-59). It also assists in avoiding complications from the connection between the peritoneum and the urinary tract and benefits from the estrogen effect on tissue in early infancy (56). Several surveys demonstrated that surgery early in life for patients with CAH does not affect their physical and mental quality of life more than other surgical and nonsurgical chronic medical conditions (60, 61). Most women with CAH favor surgery before adolescence (21, 22) although some studies show poor functional and cosmetic results of genitoplasty done in infancy (59, 60) with a high rate of re-intervention in adulthood (62). Other surgeons prefer deferring at least vaginoplasty until puberty, as most of the patients require vaginal dilatation at that time (63-65) if the surgery is done early. The evidence for either approach is limited. Vaginal dilatation should be avoided before puberty. There is no systematic evidence for prophylactic removal of asymptomatic discordant structures, such as a Mullerian remnants or utriculus. The testes should be removed at the time of diagnosis for patients with CAIS (26), PAIS raised female and androgen biosynthetic defects raised female to prevent malignancy in adulthood, associated hernia and psychological problems with the presence of testes. The streak gonads are also at risk of
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malignancy and the current recommendations are that they have to be removed in early childhood for patients with MGD raised male (66) and females with gonadal dysgenesis and Y chromosome (64). A scrotal testis is at increased risk for developing into a malignancy in patients with gonadal dysgenesis and testicular biopsy is recommended at puberty for early diagnosis of premalignant lesion (67). In patients with bilateral ovotestes, separation of ovarian and testicular tissue is performed early in life, as patients are potentially fertile from functional ovarian tissue (66, 68).

Genetics
In the near future, genetic diagnosis of DSD will become the primary and most accurate approach for diagnosis. The rapid progression of genetic techniques enables diagnosis of complex diseases such as DSD through next-generation sequencing (NGS), which includes: 1) chromosomal microarray, which allows the detection of copy number variants (CNVs) microduplications and microdeletions as small as 50 kb (69, 70). In two studies examining a large number of patients with DSD (116 and 23, respectively), CNVs were identified in 13% to 21.5% of the cases (71, 72). 2) Coding regions of the genome (whole-exome sequencing) represent 1%-2% of the entire genome, but contain around 85% of the mutations that cause known genetic disorders (73, 74) and identify significant variants in novel DSD genes (75). 3) Targeted gene sequencing (TGS) is an accurate and fast way to diagnose DSD if the causative gene is known. The coding sequences of 35 known genes involved in sex determination have been developed (76). The best example of advanced progress in genetic diagnosis of DSD is the ability to diagnose the fetal CAH as early as 5 weeks 6 days of gestation by targeted massively parallel sequencing (MPS) of DNA in maternal plasma (77).

Risk of gonadal malignancy
Patients with DSD are at increased risk for gonadal malignancies especially germ cell tumors such as dysgerminomas, seminomas and nonseminomas. The risk in general depends on: 1) the presence of a Y chromosome with undervirilization particularly gonadal dysgenesis and ovotestis (78). 2) The location of the gonad(s). The risk of malignancy is 20%-30% for patients with complete gonadal dysgenesis and 15%-20% for those with mixed gonadal dysgenesis (45, 79). Gonadoblastomas are the most common gonadal tumors and they develop within the first to second decade of life and a gonadectomy is the current management (80). Abdominal or pelvic testis is also at increased risk of tumor development and should be either removed in patients raised as females or descended into the scrotum if raised as males. The scrotal testis is at risk of malignancy in patients with gonadal dysgenesis, and a biopsy is required at the time of puberty to diagnose the tumor early (81). Patients with ovotestis or CAIS have the lowest risk (0.5%) of developing malignancies (40, 48).

Conclusion
Much progress has been made in the knowledge and diagnosis of individuals with DSD. DSD conditions are complex and need a multidisciplinary team with knowledgeable and skillful experts in the management of DSD and specialized centers with the best facilities for diagnosis and management. Parents and patients should be informed in detail about each step in the management and should be informed about the diagnosis and enough time to answer any inquiry. A clinical algorithm facilitates making the final diagnosis and avoids unnecessary investigation and reserve resources. The goal of DSD management in infancy should be to maintain stable gender identity, preserve potential sexual function and fertility and provide cosmetic and functioning external genitalia. Management includes different but important components such as diagnosis, family support, sex assignment and the risk of malignancy. Each component needs accuracy and cautious decision-making. Future studies are essential to be able to determine the cause and the most appropriate management for each individual patient.

Declaration of interest
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