Etiology of ambiguous genitalia in newborns with XX karyotype: A cross-sectional observational study.

Waqas Imran Khan¹, Abdur Rehman², Erum Afzal³, Asim Khurshid⁴, Sidra Anjum⁵, Fazal ur Rehman⁶

ABSTRACT… Objective: To determine the etiology of ambiguous genitalia in newborn females with XX karyotype. Study Design: Cross Sectional Observational study. Setting: Children Hospital and Institute of Child Health Multan (CH& ICH). Period: July 2018 to December 2019. Material & Methods: Fifty two patients were enrolled after taking informed consent from parents/guardians. Various causes of ambiguous genitalia like CAH, maternal intake of virilizing drugs during pregnancy and isolated local genital defect were noted. Results: Out of 52 patients, mean gestational age was 38.23 ± 2.36 weeks and 10 (20.2 %) had gestational age up to 36 weeks while 42 (80.8%) had gestational age more than 36 weeks. Mean age was 12.37 ± 4.21 days. Mean weight was 2965.43 ± 412.23 grams while 24 (46.1%) had weight up to 2500 grams and 28 (53.9%) had weight more than 2500 grams. Congenital adrenal hyperplasia in our study cases was noted in 45 (86.5%). Congenital local genital defects in our study cases were noted in 6 (11.5%) while maternal ingestion of virilizing drugs during pregnancy was noted in 1 (2%). CAH was not associated with gestational age or weight of newborn with p value 0.680 and 0.595 respectively. Conclusion: Congenital adrenal hyperplasia in our study cases was the commonest reason of ambiguous genitalia in newborn females with XX karyotype followed by congenital local defects and maternal use of virilizing drugs. High index of suspicion is required for the diagnosis of potential life threatening disorder like CAH.

Key words: Ambiguous Genitalia, Causes, Neonates, XX Karyotype.

INTRODUCTION

Disorders of sex development (DSD) are characterized as inherent conditions in which development of sex of an individual is unusual.¹ A few newborns are born into this world with ambiguous genitalia. This implies that the visible genitals—penis and testes or vagina and clitoris—aren’t obviously either male or female. This condition occurs roughly once in every 4,500 births² in countries like Saudi Arabia and Egypt where there is high rate of consanguinity. On the other hand, its prevalence as reported in Germany is low, i.e. once in every 10,000 births.³

At origination, determination of sex depends on chromosomes that are inherited from their parents. An individual is considered as female having XX karyotype, on the other hand an XY karyotype as male. After fertilization of an ovum the sex of an individual is determined long before the gonads are developing. Ambiguous genitalia can develop if hormones controlling this process are disturbed.⁴ ⁵

Neonatal genital ambiguity needs widespread and urgent investigations for critical disorder like congenital adrenal hyperplasia (CAH) which is a frequent cause of DSD.⁶ ⁷ Aromatase insufficiency, intake of androgens or progestins, virilizing adrenal and ovarian tumors, or luteomas are infrequent causes of ambiguous genitalia. Lastly, ovotesticular and 46, XX testicular DSD are seen. Extensive clinical, hormonal, radiological, chromosomal, and genetic evaluations are hence required. However, for precise diagnosis and treatment an urgent evaluation is needed with high index of suspicion about life threatening
condition like CAH. Satisfactory management and prompt gender assignment will have a good impact on psychosocial wellbeing of parents. As they have a key role is decision making process they should be involved in each step of diagnosis and treatment.

The objective of my research is to determine the etiology of ambiguous genitalia in newborn females with XX karyotype.

**MATERIAL & METHODS**

This cross sectional observational study was completed in department of pediatric endocrinology and neonatology at The Children hospital & the Institute of Child health Multan (CH& ICH) from July 2018 to December 2019. Fifty two patients were enrolled after taking informed consent from parents/guardians, according to following criteria: newborn babies that presents within the first 28 days of life and Karyotype proven XX females that are clinically diagnosed of having ambiguous genitalia. All the cases aged more than 28 days, as well as having normal genitalia or those having palpable gonads and not giving consent to participate in this study by patient's/guardians were excluded from this study.

The obstetric history included any endocrine problem in mother (Cushingoid features or virilization), or use of medications for recurrent abortion or contraceptive pills). As some of the causes of DSD are recessively inherited, family histories of genital anomalies, neonatal deaths, atypical pubertal development, or infertility were specifically asked. A complete physical examination of the baby was done to assess for any dysmorphic features. The genital examination included localization of gonads, size of clitoris, urethral opening and vaginal adhesions. Pelvic ultrasonography for internal genital organs, karyotyping, serum electrolytes, hormonal levels both basal and ACTH stimulated (17 hydroxy progesterone, plasma rennin testosterone and dehydroepiandrosterone sulphate-DHEAS) were done. All the data was noted on a predesigned Performa.

Data was entertained in SPSS version 20 for analysis. Quantitative variables are represented as mean and standard deviation. Qualitative variables are represented as frequencies and percentages i.e. causes of ambiguous genitalia in the newborn XX females. These causes include congenital adrenal hyperplasia, maternal intake of virilizing drugs during pregnancy and isolated local genital defect. The effect modifiers like environmental exposure to drugs, congenital deficiency of an enzyme and genetic defect causing gross dysmorphism were controlled by stratification of data. Chi square test was applied to find relationship of causes of ambiguous genitalia with weight and gestational age of study subjects. P value ≤ 0.05 is taken as significant. This study was affirmed by the institutional Ethical committee of CH& ICH, Multan.

**RESULTS**

A total of 52 patients were enrolled in this study. Table-I shows mean gestational age of 38.23 ± 2.36 weeks and 10 (20.2%) had gestational age up to 36 weeks while 42 (80.8%) had gestational age more than 36 weeks. Mean age was 12.37 ± 4.21 days and age range was 3-24 days. Mean weight was 2965.43 ± 412.23 grams while 24 (46.1%) had weight up to 2500 grams and 28 (53.9%) had weight more than 2500 grams.

Table-II shows various causes of ambiguous genitalia. CAH in our study cases was noted in 45 (86.5%). Congenital local genital defects in our study cases were noted in 6 (11.5%) while maternal ingestion of virilizing drugs during pregnancy was noted in 1 (2%) of our study cases. These causes of ambiguous genitalia were stratified with regards to gestational age, weight. (Table-III).

<table>
<thead>
<tr>
<th>Mean Age</th>
<th>12.37 ± 4.21 days</th>
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<tr>
<td>Range</td>
<td>3-24 days</td>
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<tr>
<td>Gestational age</td>
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<tr>
<td>≤ 36 weeks</td>
<td>10 (20.2%)</td>
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<td>&gt; 36 weeks</td>
<td>42 (80.8%)</td>
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<tr>
<td>Weight</td>
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<tr>
<td>≤ 2500 grams</td>
<td>24 (46.1%)</td>
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<td>&gt; 2500 gm</td>
<td>28 (53.9%)</td>
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Table-I. Descriptive statistics of study cases (n=52).
DISCUSSION

At the time of fertilization the genetic sex of a person is determined. Accordingly a 46XX karyotype would form into a phenotypic female, and 46XY to a phenotypic male embryo. DSD develop because of a dissimilarity between the genetic, gonadal and phenotypic sex and are the outcome of early disorder in the programming of sex determination.\(^\text{10,11}\) One of clinical emergencies in infants with genital ambiguity, for example CAH\(^\text{12}\) and certain dysmorphic syndromes which may be life threatening to child’s life, and uncertain sex may cause devastating psychosocial impacts on patients and parents. A skilled multidisciplinary team\(^\text{13,14}\) is necessary for the comprehensive care of kids with ambiguous genitalia which is typically found in teaching hospitals.

In our study the most frequent reason of ambiguous genitalia is CAH 45 (86.5%). These results are similar to a study done in Syria which also showed high rate of CAH in ambiguous genitalia.\(^\text{15}\) Congenital local genital defects in our study cases were noted in 6 (11.5%) while maternal ingestion of virilizing drugs during pregnancy was noted in 1 (2%) of our study cases. Literature review also suggests that CAH is the most frequent reason of ambiguous genitalia with XX karyotype.\(^\text{16,17}\) Similar findings are seen in present study that 86.5% of cases were CAH.

CAH was not associated with gestational age or weight of newborn with p value 0.680 and 0.595 respectively. Similar results were also observed in congenital local defects and maternal intake of virilizing drugs. Poyrazoglu S. et al\(^\text{18}\) and Miles HL et al.\(^\text{19}\) also reported no association of birth weight with various causes of ambiguous genitalia.

In countries like Pakistan with meager resources prompt diagnosis and management of ambiguous genitalia is warranted to avoid life threatening complication of CAH and psychological support for the family. High risk families should be offered prenatal testing and antenatal treatment to avoid genital ambiguity. Development of specialized centers and trained staff with expertise in dealing with disorders of sexual differentiation is essential for caring such patients. The limitations of my study are small number of patients as this is a rare condition. Due to rapidly evolving genetic testing, new technology should be incorporated in the diagnostic procedure. Due to diversity of causes of ambiguous genitalia the subgroups are small but it shows the challenge of diagnosing such children.

CONCLUSION

Congenital adrenal hyperplasia in our study cases was the most commonest reason of ambiguous genitalia in newborn females with XX karyotype followed by congenital local defects and maternal use of virilizing drugs. High index of suspicion is required for the diagnosis of potential life
threatening disorder like CAH.

References


## AUTHORSHIP AND CONTRIBUTION DECLARATION

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<th>Sr. #</th>
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<th>Author(s) Signature</th>
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<tbody>
<tr>
<td>1</td>
<td>Waqas Imran Khan</td>
<td>Literature search, Study design, Data acquisition, review, final approval.</td>
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<tr>
<td>2</td>
<td>Abdur Rehman</td>
<td>Data acquisition, Critical review, Approval.</td>
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<td>3</td>
<td>Erum Afzal</td>
<td>Data acquisition, Critical review, Approval.</td>
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<td>4</td>
<td>Asim Khurshid</td>
<td>Data acquisition, Critical review, Approval.</td>
<td></td>
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<tr>
<td>5</td>
<td>Sidra Anjum</td>
<td>Data acquisition, Critical review, Approval.</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>Fazal ur Rehman</td>
<td>Data acquisition, Critical review, Approval.</td>
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