Copy Number Variations in Genetic Diagnosis of Congenital Adrenal Hyperplasia Children

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Background: Congenital adrenal hyperplasia (CAH) is a monogenic disorder caused by genetic diversity in the CYP21A2 gene, with 21-hydroxylase deficiency (21-OHD) as the most common type. Early sex assignment and early diagnosis of different genetic variations with a proper technique are important to reduce mortality and morbidity. Proper early sex identification reduces emotional, social, and psychological stress.

Aim: Detection of a spectrum of aberrations in the CYP21A2 gene, including copy number variations, gene conversion, chimeric genes, and point variations.

Methods: The CYP21A2 gene was screened using MLPA assay in 112 unrelated Egyptian children with 21-OHD CAH (33 males and 79 females).

Results: In the studied group, 79.5% were diagnosed within the first month of life. 46.8% of the genetic females were misdiagnosed as males. Among the copy number variation results, large deletions in 15.4% and three types of chimeric genes in 9% (CH-1, CH-7, and CAH-X CH-1) were detected. Regarding gene dosage, one copy of CYP21A2 was found in 5 cases (4.5%), three copies were detected in 7 cases (6.3%), and one case (0.9%) showed four copies. Eight common genetic variants were identified, I2G, large deletions, large gene conversion (LGC), I172N, F306 + T, -113 SNP, 8bp Del, and exon 6 cluster (V237E and M239K) with an allelic frequency of 32.62%, 15.45%, 7.30%, 3.00%, 2.58%, 2.15%, 0.86%, and 0.86%, respectively.

Conclusion: High prevalence of copy number variations highlights the added value of using MLPA in routine laboratory diagnosis of CAH patients.

Keywords: 21-Hydroxylase deficiency, CYP21A2, multiplex ligation-dependent probe amplification, copy number variations, congenital adrenal hyperplasia

INTRODUCTION

Congenital adrenal hyperplasia (CAH) represents a group of common defects in the adrenal gland steroidogenesis; the main form is 21-hydroxylase enzyme deficiency (21-OHD; OMIM #201910) (Chan et al., 2011). Patients with the classic phenotype can present with life-threatening salt-wasting (SW) and simple virilizing (SV) types depending on the enzymatic activity of the 21-OH enzyme (Rabbani et al., 2011). There is a higher prevalence of CAH in Egypt (1 in 1,209 live births) and Arab
countries due to a higher rate of consanguinity (Temtamy and Aghan, 2012) (Tayel et al., 2011) (Alshabab et al., 2015).

In males, CAH is a medical emergency, with an urgent need for early diagnosis as it can be fatal in cases with SW-CAH. In females, CAH-SV can also be considered a social emergency (Kutney et al., 2016). The sex of a newborn is usually determined at birth on the basis of external genital appearance. Thus, children with atypical genitalia frequently require reassignment of sex because of incorrect original identification. As sex is a fundamental attribute of human life, its reversal after the original assignment is usually associated with marked emotional, social, and psychological stress with accompanying shame (Wisniewski, 2017).

First-line diagnostic tests for CAH include karyotype analysis, hormonal evaluation (from 48 h after birth) for 17-hydroxyprogesterone (17OHP), androgens, basal cortisol, and adrenocorticotropic hormone (ACTH), and ultrasound (for detecting Mullerian structures). The definitive diagnosis of 21OHD CAH is through the assessment of serum 17OHP levels (levels above 10 ng/dl are diagnostic). Additional imaging tests may be recommended, including genogram, retrograde urethrogram, or cystoscopy/vaginoscopy (in females with atypical genitalia). When the gonads are not detected by ultrasound, magnetic resonance imaging is indicated. Molecular studies are performed to check for gene mutations (Guerrero-Fernández et al., 2018).

Management of CAH cases includes lifelong hormonal therapy and reconstructive surgery for female cases with atypical genitalia (feminizing genitoplasty) (Savanelli et al., 2018). This is best done in early infancy to confer an early physical appearance consistent with the gender sex of rearing and to cause less psychological trauma than if delayed (Speiser et al., 2018; Musa et al., 2020).

While CAH-21OH is a monogenic disorder, the human 21-OH gene (cytochrome P450 family 21 subfamily A member 2, CYP21A2) encodes the 21-OH enzyme, and a highly homologous inactive pseudogene, CYP21A1P, is located closely adjacent to the CYP21A2 gene, approximately 30 kb apart in tandem arrangement with the C4A and C4B genes (Blaskó et al., 2009; Choi et al., 2016; Daae et al., 2018). These arrangements comprise the most frequent bimodular RCCX of the RP1-C4-CYP21A1P-TNXA-RP2-C4-CYP21A2-TNXB gene sequence in 69% of alleles in Caucasians (Lee et al., 2010; Tsai et al., 2012). Recombination events commonly occur when a loop forms in the linear DNA of the chromosome. The active CYP21A2 and inactive CYP21A1P genes lie against each other, allowing the transfer of genetic material between them. The advantage of this physiological event is that it increases immunological diversity over time. However, the CYP21A2 gene sits adjacent to the HLA locus and becomes a passive participant in this process (Huynh et al., 2009).

The high degree of sequence similarity between active and pseudogene allows two types of intergenic recombination events that are responsible for about 95% of the mutations associated with 21-OHD (Vrzalová et al., 2011). The first is gene conversion events: approximately 65%–70% of the deleterious mutations are derived from pseudogene CYP21A1P due to small gene conversions, including I2G (28%), I172N (9%), V281L (9%), Q318X (4%), R356W (4%), E6 cluster [I235N, V236E, and M238K] (4%), 8bp Del (3%), P30L (2%), and F306 + T (1%) (Xu et al., 2013). If these transfers comprise a large region, it is called a “large-scale conversion” (Toraman et al., 2013).

When targeted mutation analysis detects multiple mutations, it is possible that the mutations are most likely arising from gene conversion (Nimkarn et al., 1993). The second is unequal crossing over during meiosis: the remaining 20%–25% of the intergenic recombination is represented by CYP21A2 gene deletions or CYP21A1P/CYP21A2 chimeric genes (Concolino et al., 2009). Nine different CYP21A1P/CYP21A2 chimeric genes were described according to their breakpoints (CH-1 to CH-9) (Chen et al., 2012; Veldhuisen et al., 2015). The third common variation in CYP21A2 is spontaneous point mutations (Xu et al., 2013).

Multiplex Ligation-dependent Probe Amplification (MLPA) assay with the addition of common point-mutation (PMS) MLPA probes represents the gold standard in CAH diagnosis and a sensitive tool for identification of chimerical genes (Choi et al., 2016) (Erlandson et al., 2008).

The aim of this study is to perform genetic analysis of CAH 21-OHD Egyptian children to detect a spectrum of aberrations in the CYP21A2 gene, including copy number variations, gene conversion, gene deletion, chimeric genes, and point variations using MLPA analysis.

### MATERIALS AND METHODS

Editorial policies and ethical considerations: children with CAH were recruited from the Diabetes Endocrine Metabolism Pediatric Unit (DEMPU), Children’s Hospital, Cairo University, after obtaining written informed consent from all participants or their legal guardians. The study was approved by the Research Ethics Committee of Cairo University. The diagnosis of CAH due to 21-OHD (salt wasting and simple virilizing) was based on both clinical and biochemical examinations (elevated 17-hydroxyprogesterone).

Three millilitres of blood were collected from each subject in a sterile EDTA vacutainer for the genotyping technique. DNA was extracted using the DNA extraction kit QIAamp (DNA Blood Mini Kit) supplied by QIAGEN (Thermo Fisher Scientific, USA) according to the manufacturer’s instructions. Then Selective Adaptor Ligation, Selective Amplification (SALSA) MLPA Probemix P050-C1 CAH (Catalog no. P050-100R) in combination with the SALSA MLPA EK1 reagent kit (EK1-FAM) supplied by MRC-Holland was used.

Multiplex ligation-dependent probe amplification analysis of the CYP21A2 gene included the following steps according to the manufacturer’s instructions: DNA denaturation, hybridization reaction, ligation reaction, PCR reaction, and fragment separation by capillary electrophoresis, ABI 3500 Genetic Analyzer (Applied Biosystems, Thermo Fisher Scientific, USA).
Bioinformatics and Statistical Analysis
The raw data were analysed using Coffalyser.net software, the MLPA analysis tool developed at MRC-Holland. It was used for data analysis in combination with the appropriate lot-specific MLPA Coffalyser sheet. They were available on www.mlpa.com.

The nomenclature of the variants reported in the current study are according to the Human Genome Variation Society (HGVS) (denDunnen et al., 2016) as follows (NM_000500.8): 113 SNP (rs1246774295, c.-113G > A), I2G (rs6467, c.293–13C > G), 8bp Del (rs387906510, c.332_339delGAGACTAC, c.332_339del), I172N (rs6475, c.518T > A), Exon 6 cluster variants [V237E (rs12530380, c.713T > A), M239K (rs6476), c.719T > A], and F306 + T (rs267606756, c.923dup).

Data were analysed using IBM SPSS Advanced Statistics version 21.0 (IBM Corp., Armonk, NY). Qualitative data were expressed as frequency and percentage.

RESULTS
One hundred and twelve probands with 21-OHD were recruited for the study. The majority (79.5%, 89/122) were diagnosed in the first month of life. 50/112 (44.6%) were diagnosed at birth (day 0) and 39/112 (34.8%) between 1–30 days. Seventy-six (67.9%) patients were born to consanguineous parents. By karyotyping, the genetically confirmed females constituted 79/112 (70.5%) and 33/112 (29.5%) were males. The female-to-male ratio was 2.4:1, and 37/79 (46.8%) of genetic females were misdiagnosed as males.

TABLE 1 | The agreement between MLPA and karyotype among studied cases (n = 112).

| Karyotype | MLPA | Total |
|-----------|------|-------|
| Male, n (%) | Female, n (%) |       |
| 46XY | 32 (97) | 1 (3) | 33 |
| 46XX | 7 (8.9) | 72 (91.1) | 79 |
| Total | 39 | 73 | 112 |

Kappa value* (p value) 0.837 (<0.001)

*Kappa value: < 0: less than chance agreement; 0.01–0.2: slight agreement; 0.21–0.4: fair agreement; 0.41–0.6: moderate agreement; 0.61–0.8: substantial agreement; 0.81–0.99: almost perfect.

The majority of females presented with ambiguous genitalia (78/79, 98.7%). In most cases, 49/79 (62%) presented with Prader stage III, followed by stage IV (16.5%). The remaining cases presented with Prader stage II (10.1%), I (7.6%), and V (2.5%), while 4/33 males (12.1%) presented with precocious puberty.

All cases in the current study had the classical form of 21-OHD. Eighty-eight out of 112 patients (78.6%) were SW, and 24/112 (21.4%) cases had the SV form. The phenotype of all cases with large gene conversions and/or deletions is SW except for one case that was SV (LGC 1-3 exon on one allele and Del 1-7 exon Del on the other allele). Figure 1 shows the ratio chart result of 1–7 exon deletion (Del 1-7).

Regarding the gene dosage, 67% of cases (75/112) had two copies of the active gene. Three copies and one copy of CYP21A2 were found in 7/112 (6.3%) and 5/112 (4.5%) cases, respectively. Zero copies were reported in 24/112 (21.4%), and four copies

FIGURE 1 | Ratio chart showing the homozygous 1–7 exon Del CYP21A2 gene.
were reported in only one patient (0.9%). Genotype results of cases with zero copies of the active gene show homozygous LGC or large deletions.

The majority of single nucleotide variants (SNV) were in a homozygous state, 29/112 (25.9%). Heterozygous SNV was found in 11/112 (9.8%) cases. Compound heterozygous SNV and compound homozygous single nucleotide variants were present in 8/112 (7.1%) and 2/112 (1.8%) cases, respectively. The frequency of genotype distribution of CYP21A2 gene variants among cases is presented in Table 1.

## DISCUSSION

To the best of our knowledge, the study includes the largest sample size among Arab countries and the first one to use MLPA assay for the diagnosis of copy number variations in a group of 21-OHD CAH children in Egypt. One hundred and twelve consecutive unrelated 21-OHD CAH patients were enrolled in the present study. The results revealed that 89/112 (79.5%) were diagnosed within the first month of life and highlight the necessity of a neonatal screening program for CAH in Egypt.

All cases in the current study had the classical form of 21-OHD. Eighty-eight out of 112 patients (78.6%) were salt wasting (SW) and 24 out of 112 (21.4%) cases had the simple virilizing (SV) form. Fortunately, affected males are fewer than females, as males with the SW form are at higher risk of adrenal crises and may pass away unnoticed without proper diagnosis. In our study, the female-to-male ratio was 2:1. This is in accordance with previous studies that found the female-to-male ratio was 3:1 (Al-obaidi et al., 2016).

The wrong sex identification of affected females reared as males in 46.8% of cases can be explained by the high prevalence of female ambiguous genitalia. In the present study, all female cases presented with ambiguous genitalia, except for one case that received prenatal treatment. Reassignment of gender to the female sex in cases with 46XX 21OHD CAH is recommended (Freire et al., 2018) and occurred in 36/37 of the genetic females that were misdiagnosed as males. One genetic female only continued as a male after surgical operations at the request of the parents.

One of the potential roles of MLPA is genetic sex identification in both parents. This is the first to use MLPA first one to use MLPA.
females). Ultrasound can detect the presence of Mullerian structures as a guide for the female internal genitalia. However, it is operator dependent, and sometimes, it can be very difficult to visualize the tiny uterus and ovaries in neonates.

Regarding genotyping, the most frequent variant in the present study was I2G, with an allelic frequency of 31%. The second common variant was large deletions (15.45%), and the third most frequent variant was LGC (7.3%) (Table 2). In 2002, Krone et al.'s study showed similar results (Krone et al., 2002).

Copy number variants presented in 15% of cases as simple (11.6%) or combined with SNVs. These simple cases may be missed in genetic diagnosis when using commercial molecular techniques that only test for SNVs. Thus, copy number variations also need to be checked in the cohort due to the high frequency of this cohort (Gao et al., 2019).

New et al. highlighted the high frequency of homozygous I2G variants in the Middle Eastern population (New et al., 2013). The I2G mutation presented in 31% of cases alone, or combined with other point mutations (9%) or with copy number variations (LGC or deletions) (3.5%). The E6 cluster variants (p.V237E, p.M239K) showed an allele frequency of 0.86%. The homozygous I2G (as the common mutation) and compound heterozygous for I2G and E6 (the least frequent point mutation) mutations cause the same disease phenotype (SW). The genotype frequency among different phenotypes in the studied cases is summarized in Supplementary Table S1.

The lack of identified variants in the present study highlights the need for further genetic studies to detect such a discrepancy between the genotype and phenotype. This may be explained by the presence of common variants of the CYP21A2 gene that were not detected by MLPA probes of probemix P050-C1 CAH as a mutation, P.Q318X, in addition to rare variants that were not tested. Moreover, variants within the promoter region of the gene may be present.

Sequencing is essential to optimize molecular diagnosis in CAH patients. It is considered the only available method that allows for a 100% detection rate for all single nucleotide variants of CYP21A2, either common or rare. Direct sequencing in combination with MLPA offers the highest diagnostic information (Khattab et al., 2016). The gene dosage of the active gene and pseudogene reported in the current study using the MLPA technique showed the CYP21A2 copy number was zero in 24/112 (21.4%). Three copies were found in 7 of the 112 patients (6.3%). To our knowledge, no studies investigate the clinical importance of pseudogene copy number role in the CAH. Pseudogene dosage analysis is important in the differentiation between large deletions and LGC and to avoid misinterpretation of the wild-type sequence of a pseudogene as a duplication of CYP21A2. The present study results showed three copies of CYP21A1P in 21 out of 112 patients (18.8%) and four copies in 4 out of 112 (3.6%).

An added diagnostic potential of using the MLPA technique is that the pseudogene analysis was performed simultaneously with CYP21A2 analysis, so this can help to detect the extent of deletion and different forms of chimeric genes. In the present study, a 30-KB Del-produced chimeric gene (CH-1) was detected in 6/112
cases and a chimeric gene (CH-7) in one case. Coeli-Lacchini et al. conducted a study using the MLPA method. The results revealed a 30-kb Del in 3/90 (3.3%) and a complete CYP21A2 deletion in one of the patients (Coeli-Lacchini et al., 2013). Deletions typically extend approximately 30 kb and have their breakpoint somewhere between exons 3 and 8 of the CYP21A1P pseudogene and ending somewhere at the corresponding point in CYP21A2, thus yielding a single remaining gene in which the 5’ end corresponds to the CYP21A1P pseudogene and the 3’ end corresponds to the CYP21A2 gene. To date, nine types of CYP21A1P/CYP21A2 chimera (CH1–CH9) with different junction sites have been identified (New et al., 2013). Furthermore, loss of MLPA probes specific to the coding regions of the CYP21A2 gene, which constituted loss of exons 1–3, is Del 1–3, loss of exons 1–6 is Del 1–6, and loss of exons 1–8 is Del 1–8 variants in CYP21A2 (Balraj et al., 2013). The classic TNXA/TNXB chimeric gene is a 120 bp deletion at the boundary of exon 35 and intron 35 (CAH-X CH-1). The CYP21A2 gene is completely deleted and replaced by the CYP21A1P pseudogene (Merke et al., 2013). The 148 and 318 nucleotide (nt) probes of TNXB exon 35 of the MLPA probemix (P050-C1 CAH) are located within the 121 nt sequence that is absent in the TNXA pseudogene. Using the MLPA technique, the prevalence of large deletions and large gene conversions in the present study was found in 29/112 (25.9%) patients with the SW guardian/next of kin.

The present study is the first to report large gene Del in Egyptian CAH patients, extending to exon 35 of the TNXB gene that produced a chimeric gene (CAH-X CH-1) in 3/112 patients. All cases with large gene deletions presented with the SW form (Figure 2). This molecular diagnosis coincides with Ehlers–Danlos syndrome (EDS). These results highlight the importance of a complete clinical evaluation for CAH children. Clinical evaluation for connective tissue dysplasia should be routinely performed in CAH patients, especially those harboring a CYP21A2 deletion (Merke et al., 2013) as CAH-X recently considered a subtype of CAH (Lao and Merke, 2021; Miller, 2021). Merke et al. reported CAH-X syndrome in 7% of CAH patients. Patients with CAH-X were more likely to have joint hypermobility, chronic joint pain, multiple joint dislocations, and structural cardiac valves.

**CONCLUSION**

Early genetic sex assignment, genetic diagnosis, and phenotype prediction are important in the early treatment of CAH patients. Due to the sophisticated mechanisms of genetic variations in the CYP21A2 gene, there is an urgent need to use MLPA as a multipotent and cost-effective laboratory routine tool for the molecular diagnosis of 21-OHD CAH in countries with a high frequency of CAH for proper diagnosis of cases. Recommendations for tailoring the panel of MLPA probe mix to include missed important variants will add to its potent diagnostic effect.

**DATA AVAILABILITY STATEMENT**

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author.

**ETHICS STATEMENT**

The studies involving human participants were reviewed and approved by the clinical and chemical pathology ethical committee at Kasr al Ainy Hospital. Written informed consent to participate in this study was provided by the participants’ legal guardian/next of kin.

**AUTHOR CONTRIBUTIONS**

AT: substantial contributions to the conception or design of the work and acquisition and analysis of data. IM: drafting the work and revising it critically for important intellectual content. NM: diagnosing of patients and data collection. FE: substantial contributions to the conception or design of the work and acquisition of data. MH: substantial contributions to the conception or design of the work and acquisition of data. SA: drafting the work or revising it critically for important intellectual content. AI: contribution in lab work. HS: data collection. BL: provided approval for publication of the content. YE: drafting the work. TR: contribution in lab work. HS: data collection. BL: provided approval for publication of the content. YE: drafting the work.

**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fgene.2022.785570/full#supplementary-material

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