**22q11.2 Deletion Syndrome: Copy Number Variation Analysis in Two Patients**

**Abstract**

**Background:** 22q11.2 deletion syndrome is the most common microdeletion syndrome. It is caused by an autosomal dominant microdeletion of chromosome 22 at the long arm 11.2 band. It is difficult to diagnose of this syndrome because of its multiple phenotypes.

**Methods and findings:** In this study, we identified two cases of 22q11.2 deletion syndrome by array based comparative genomic hybridization (aCGH). These two cases exhibited atypical anomaly examined by aCGH and they exhibited different presentations although aCGH showed the same deletion.

**Conclusion:** 22q11.2 deletion syndrome is a difficult diagnosed disease which could be efficiently detected by aCGH. The patients could exhibit different phenotypes even when aCGH showed the same deletion. These findings will provide useful information to better understand 22q11.2 deletion syndrome.

**Keywords:** 22q11.2 deletion syndrome; Copy number variation; Array based comparative genomic hybridization (aCGH)

**Received:** January 28, 2016; **Accepted:** May 13, 2016; **Published:** May 16, 2016

**Introduction**

22q11.2 chromosomal region contains several paralogous low copy repeats (LCRs), which predisposes to de novo genomic rearrangement at this locus. Hemizygous deletions of this region, also known as copy number variations (CNVs), results in 22q11.2 deletion syndrome (22q11.2 DS). The 22q11.2 DS affects about 1 in 4000 live births. Patients of 22q11.2 DS have high prevalence of intellectual disability and psychiatric disorders [1] and death related to this syndrome has also been reported. The 22q11.2 DS is much variable with more than 180 features. It showed wide spectrum of clinical presentations ranging from mild to severe, which led to difficult diagnosis of this disease. In most cases, 22q11.2 DS is caused by a hemizygous deletion of 3 million base pairs of DNA that encompasses approximately 40 known genes. Smaller nested deletions of 1.5 million base pairs that span 34 known genes also were reported [2,3].

In this study, we reported two cases of 22q11.2 DS showed atypical anomaly of 2.5 M deletion identified by array based comparative genomic hybridization (aCGH). And these two cases showed different presentations although aCGH revealed the same deletions.

**Case Description**

The patient 1 shows atypical characteristic features for 22q11.2 DS. He is an one-year old boy who exhibits intellectual disabilities, developmental delay, dyskinesia, patent foramen ovale, laryngeal cartilage dysplasia, varus, gastric torsion, cerebral palsy and hearing loss. After peripheral blood was collected from this patient, both Karyotyping analysis and aCGH analysis were performed, while karyotyping analysis showed negative result.

The second sample was from a pregnant woman at 20 weeks of gestation. Fetal ultrasound indicated abnormality of eyes structures and ascites. The pregnant chose definitive prenatal diagnosis through amniocentesis after received genetic counseling. Karyotyping analysis of amniotic fluid did not detect any abnormal. Afterwards, aCGH was carried out with amniotic fluid.
with Feature Extraction software (v10.7). Results were further analyzed by Agilent Genomic Workbench (v6.5) and Agilent Cytogenomics (v2.7.7.0) based on Human Genome build 19, and further interpreted by databases consultation.

aCGH analysis showed that patient one covered about 83.3% deletion of the typically deleted region (TDR) of 22q11.2 deletion syndrome (Figures 1A and 1B). The fetal also covered about 83.3% of the TDR (Figure 1), which is the same as the patient one showed by aCGH.

Comments

aCGH is a powerful tool to detect 22q11.2 deletion syndrome. Different patients could show same deletion examined by aCGH even when they exhibited different presentations.

Ethical Consideration

Ethical approval was obtained from the Ethics Committee of Henan Provincial People’s Hospital, China. The patients involved in this study signed the consent documents.

Discussion

It is known that there are 4 LCRs in proximal region and 4 another smaller LCRs in distal region in 22q11 chromosome based on their chromosome order. Different size and different location of deletions usually exhibited different phenotypes. In this study, the two patients showed different clinical presentations while they shared the same deletion detected by aCGH. Since 22q11.2 DS is a disease that has different phenotypes among different patients, it is difficult in diagnosis. Some mild and atypical phenotypes may not be diagnosed in time and correctly in this disease. Therefore, array based comparative genomic hybridization is a powerful technology that used to identify 22q11.2 DS.

So far, the mechanism of 22q11.2 deletion syndrome remains largely elusive. It was suggested that LCRs mediated rearrangements of 22q11.2 [4]. As how LCRs mediated deletions in 22q11.2 need to be further explored. 4 proximal LCRs show a complex modular structure, and the modular share high level of sequence identity. This may lead to misalignment during meiosis and unequal interchromosomal recombination. So far, LCR mediated non-allelic homologous recombination (NAHR) in genomic rearrangement has already been widely accepted [5]. Afterwards, it is reported that 4 distal LCRs also were involved in this deletion syndrome [6]. Further studies would provide more information that helps to better understand the mechanism of 22q11.2 DS.

Conclusion

In conclusion, 22q11.2 DS is a variable disease exhibited different phenotypes even when aCGH showed the same deletions. aCGH is a pangenomic screening technology with higher resolution and rapid breakpoint localization compared to traditional methods in genomic disorders. It is a good option when the phenotype is atypical and not a clear diagnosis is obtained in clinical in 22q11.2 DS, and even other deletion syndromes.
References:


