Clinical heterogeneity and intrafamilial variability of Joubert syndrome in two siblings with CPLANE1 variants

Xiujuan Zhang1,2 | Yue Shen3 | Ping Li4 | Ruikun Cai3 | Chao Lu3 | Qian Li3 | Cuixia Chen3 | Yufei Yu3 | Tingting Cheng3,5 | Xian Wang1 | Minna Luo3 | Cuixia Chen3 | Yufei Yu3 | Tingting Cheng3,5 | Xian Wang1 | Minna Luo3

1Department of Physiology and Pathophysiology, School of Basic Medicine Sciences, Key Laboratory of Molecular Cardiovascular Science, Ministry of Education, Peking University, Beijing, China
2Key Laboratory of Cell Differentiation and Apoptosis of Chinese Ministry of Education, Department of Pathophysiology, Shanghai Jiao Tong University School of Medicine, Shanghai, China
3National Human Genetic Resources Center, National Research Institute for Family Planning, Beijing, China
4Department of Developmental Pediatrics, The Second Hospital of Jilin University, Changchun, Jilin, China
5Graduate School of Peking Union Medical College, Beijing, China

Abstract

Background: Joubert syndrome (JBTS) is a rare genetic disorder that is characterized by midbrain-hindbrain malformations. Multiple variants in genes that affect ciliary function contribute to the genetic and clinical heterogeneity of JBTS and its subtypes. However, the correlation between genotype and phenotype has not been elucidated due to the limited number of patients available.

Methods: In this study, we observed different clinical features in two siblings from the same family. The older sibling was classified as a pure JBTS patient, whereas her younger sibling displayed oral-facial-digital defects and was therefore classified as an oral-facial-digital syndrome type VI (OFD VI) patient. Next, we performed human genetic tests to identify the potential pathogenic variants in the two siblings.

Results: Genetic sequencing indicated that both siblings harbored compound heterozygous variants of a missense variant (c.1067C>T, p.S356F) and a frameshift variant (c.8377_8378del, p.E2793Lfs*24) in CPLANE1 (NM_023073.3).

Conclusion: This study reports that two novel CPLANE1 variants are associated with the occurrence of JBTS and OFD VI. These results help elucidate the intrafamilial phenotypic variability associated with CPLANE1 variants.

KEYWORDS

CPLANE1, intrafamilial heterogeneity, Joubert syndrome, OFD VI
Joubert syndrome (JBTS, OMIM# 213300) is a classic ciliopathy that was first identified by Dr. Joubert et al., (1969). JBTS is a rare recessive genetic disorder with an estimated prevalence of 1/80000-1/100000 (Valente et al., 2013). The clinical diagnosis of JBTS by magnetic resonance imaging (MRI) is characterized by distinct cerebellar vermis and brainstem malformation, known as the “molar tooth signs (MTS)” (Romani et al., 2013). Most patients with JBTS exhibit developmental delay, hypotonia, ataxia, abnormal breathing patterns, and/or abnormal eye movements, as well as other variable features, including retinal dystrophy, polydactyly, coloboma, and cystic disease. Variations in more than 35 genes, most of which encode ciliary proteins, have been demonstrated to be associated with the development of JBTS (Wheway et al., 2015). More variations associated with JBTS need to be elucidated.

CPLANE1 (previously known as C5orf42, NM_023073.3) is one of the causative genes for JBTS and oral-facial-digital syndrome VI (OFD VI, OMIM# 277170). OFD VI represents a rare phenotypic subtype of Joubert syndrome and related disorders (Poretti et al., 2012; Romani et al., 2015). The CPLANE1 protein has a putative coiled-coil region and exhibits features of a transmembrane protein. This protein is located at the ciliary transition zone and is involved in the recruitment of peripheral IFT-A proteins to the basal body for cilium assembly (Toriyama et al., 2016). Patients with CPLANE1 variants have been observed to have multiple birth defects typical of ciliopathies (Bonnard et al., 2018). To date, more than 130 different CPLANE1 variants have been identified, and individuals carrying CPLANE1 variants exhibit various phenotypes (Ben-Salem et al., 2014). In this study, we identified a missense variant (c.1067C>T, p.S356F) and a frameshift variant (c.8377_8378del, p.E2793Lfs*24) of CPLANE1 in two siblings. Although the siblings carried the same CPLANE1 variants, the clinical features showed a high degree of intrafamilial variability.

Genomic DNA was isolated by using the QIAamp DNA Blood Mini Kit (Qiagen). Whole-exome libraries were prepared using the Agilent SureSelect Human All Exon V6 kit (Agilent Technologies Inc.), and sequenced on the Illumina NovaSeq 6000 platform. The variants were called and annotated as previously described (Luo et al., 2019). Sanger sequencing was used to further validate the identified variants of CPLANE1 (NM_023073.3). The PCR primers used were as follows: CPLANE1-Exon 9 Forward (5′-CTGCTTGGTGACAGCCCATT-3′) and Reverse (5′-ATGAGATGACAGACTCAAAAGAAA-3′); CPLANE1-Exon 43 Forward (5′-GGTTTGCCAGTGTGGAAATAG-3′) and Reverse (5′-TTCCTCAGTAAATCATGTTCAGTAA-3′).

The proband was a 14-year-old female (18C1). She was born full-term by cesarean section to a healthy couple. She was unable to sit until 1 year of age and could not walk independently until 5 years of age. At the age of 7 years, she underwent a brain MRI, which showed typical MTS with thickening and elongation of the superior cerebellar peduncles (SCP), deepening of the interpeduncular fossa (IF), hypoplasia of the cerebellar vermis, and enlargement of the fourth ventricle (Figure 1a). No other facial deformities or skeletal defects were observed on physical examination, except for oculomotor anomalies. Nor renal/hepatic involvement or retinal degeneration was observed. The Marshalla oral sensorimotor test (MOST) (Marshalla, 2007) indicated reduced strength and poor coordination of her lips, tongue, and jaw, suggesting that she experienced delayed language development, reduced intelligibility of speech, and apraxia of speech (Figure 1c and Table 1). Based on these findings, 18C1 was diagnosed as a pure JBTS patient.

Her brother (18C2) was a 6-year-old boy who exhibited similar neurological symptoms, but his clinical features differed from 18C1. Prenatal ultrasonography at 32 weeks gestation demonstrated ventriculomegaly, and prenatal MRI at 35 weeks gestation showed typical MTS features of cerebellar hypoplasia. The patient was born full-term by cesarean section with a birth weight of 3.8 kg. After birth, polydactyly on both hands and bilateral great toes were observed (Figure 1c). 18C2 exhibited abnormal breathing patterns and...
hypotonia. Physical examination demonstrated the patient's facial deformities, including a prominent forehead, low-set ears, wide nasal bridge, and hypertelorism. Tongue hamartomas was also noticed on examination. In addition, the patient exhibited skeletal abnormalities, such as scoliosis. The patient also had severe deficits in both gross motor skills and language skills. All the clinical features together with the MTS on brain MRI (Figure 1b) confirmed the diagnosis of OFD VI.

3.2 Genetic studies revealed novel heterozygous CPLANE1 variants in two affected individuals

To determine the genetic basis of JBTS and OFD VI in this family, we performed WES on both siblings and validated the variants in all family members by using Sanger sequencing. We observed that both the proband (18C1) and her brother (18C2) carried novel heterozygous variants of CPLANE1, a paternally inherited frameshift variant c.8377_8378del (p. E2793Lfs*24) and a maternally inherited missense variant c.1067C>T (p.S356F), respectively (Figure 2a,b). These two variants have not been reported in public databases, such as gnomAD, ExAC, or 1000 Genomes. Both variants are located outside of the predicted transmembrane and coiled-coil domains. The c.8377_8378del variant causes the loss of the C-terminal domain (pfam15392) of CPLANE1 (Figure 2c) and is classified as a “pathogenic variant” according to the American College of Medical Genetics and Genomics (ACMG) guidelines (Richards et al., 2015). The missense variant c.1067C>T results in the substitution of the highly conserved serine into phenylalanine at residue 356 (Figure 2d), which is predicted to be deleterious by several tools (SIFT = 0.002; PolyPhen = 0.997; CADD PHRED = 23.9; ClinPred = 0.985). Thus, the c.1067C>T variation is annotated to be “likely pathogenic” according to the ACMG guidelines. Further, additional pathogenic variants in other ciliopathy-related genes were not identified. Taken together, the results of this study suggest that these two novel heterozygous variants may be causative for the clinical features of JBTS and OFD VI in these two siblings.
Since 2012, multiple variants of \textit{CPLANE1} have been identified in JBTS families by using WES \citep{Alazami2012, Asadollahi2018, Kroes2016, Liu2020, Ohba2013, Srour2012}. Meanwhile, several novel variants of \textit{CPLANE1} have been reported to be responsible for the OFD VI and milder type of JBTS phenotypes \citep{Bayram2015, Bonnard2018, Lopez2014, Romani2015, Wentzensen2015}. According to the Human Gene Mutation Database (HGMD), more than 125 variants in the \textit{CPLANE1} gene have been confirmed to be associated with JBTS traits, including 74 missense variants, 15 splice-change variants, and 36 insertion-deletion variants. In our study, it was found that two patients carrying the same compound heterozygous variants of \textit{CPLANE1}.

**TABLE 1** Clinical features of the patients with \textit{CPLANE1} variants

| Patient ID | 18C1 | Frequency in JBTS Patients | Frequency in OFD VI Patients |
|-----------|------|-----------------------------|-----------------------------|
| c.1067C>T (p.Ser356Phe) + + | 93.1% | 100% |
| c.8377_8381delGAAA (p.Glu2793IlefsTer31) + + | 100% | 100% |

| Age | 14 years | 6 years |
|-----|----------|--------|
| Sex | Female | Male |
| Brain MRI | MTS | MTS |
| Developmental delay | + | + |
| Intellectual disability | + | + |
| Oculomotor apraxia | + | + |
| Breathing abnormality | – | + |
| Limb abnormality | – | + |
| Scoliosis | – | + |
| Tongue hamartomas | – | + |
| Retinal involvement | – | + |
| Renal involvement | – | + |
| Hepatic involvement | – | + |

**FIGURE 2** Genetic findings and functional analysis of the variants. (a) The pedigree of the family shows two novel variants. (b) Sequence chromatograms show heterozygous variations in \textit{CPLANE1} in the two siblings. (c) The locations of variants in \textit{CPLANE1} and the predicted transmembrane domain (TM), coiled-coil domain (CCD), and Joubert syndrome-associated domain. (d) The amino acid alignments generated by homologene (NCBI) show that the residue affected by the c.1067C>T variant is highly conserved.

4 | DISCUSSION

Since 2012, multiple variants of \textit{CPLANE1} have been identified in JBTS families by using WES \citep{Alazami2012, Asadollahi2018, Kroes2016, Liu2020, Ohba2013, Srour2012}. Meanwhile, several novel variants of \textit{CPLANE1} have been reported to be responsible for the OFD VI and milder type of JBTS phenotypes.
(c.8377_8378del and c.1067C>T) exhibited different clinical features, with 18C1 being diagnosed as a JBTS patient and 18C2 being diagnosed as an OFD VI patient. Due to abnormalities in the localization and function of ciliogenesis and planar polarity effector proteins in the CPLANE1 variants (Toriyama et al., 2016), we surmise that both variants may contribute to the development of JBTS and OFD VI diseases by affecting ciliary function. Considering that rare cases of JBTS or OFD VI have been reported in the Chinese population, these two novel CPLANE1 variants will provide valuable information for future clinical diagnosis.

JBTS is a heterogeneous disorder, and the clinical features may vary between patients. Surprisingly, a high degree of intrafamilial heterogeneity was observed in this family. Although the two siblings exhibited similar neurological symptoms, 18C2 showed more severe deficits in gross motor skills, skeletal development, and respiratory patterns (Figure 1c and Table 1). Romani also reported that patients with CPLANE1 variants exhibited diverse JBTS phenotypes, with the OFD VI phenotype being the strongest (Romani et al., 2015). We surmise that the phenotypic heterogeneity may be due to the presence of genetic modifiers and that the genetic background of patients can alter the expressiveness of CPLANE1 variants (Zaki et al., 2011). Recently, the first modifier of Joubert syndrome, a single locus of Barttin, was revealed for the first time in a mouse model (Ramsbottom et al., 2020). In-depth biostatistical and functional analyses are needed to reveal additional genes that alter CPLANE1 variant-associated disease phenotypes.

In summary, we identified two new CPLANE1 variants from JBTS and OFD VI patients. The two siblings carrying these two variants showed intrafamilial clinical heterogeneity.

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CONFLICT OF INTEREST
The authors declare no conflict of interest.

AUTHOR CONTRIBUTIONS
Xu Ma, Zongfu Cao, Muqing Cao, Minna Luo, and Xian Wang conceived and supervised the project; Ping Li and Minna Luo collected the biological samples and clinical information of the patients; Xiujuan Zhang and Yue Shen analyzed and interpreted the medical data; Xiujuan Zhang, Yue Shen, Chao Lu, Qian Li, and Tingting Cheng prepared the samples and performed human genetic testing; Zongfu Cao, Minna Luo, Ruikun Cai, Cuixia Chen, and Yufei Yu analyzed and interpreted the WES data. Xu Ma, Zongfu Cao, Muqing Cao, Xiujuan Zhang, and Minna Luo wrote the manuscript with the help of all other authors.

DATA AVAILABILITY STATEMENT
The data in this study are available from the corresponding author (X.M.), upon reasonable request.

ORCID
Yue Shen https://orcid.org/0000-0003-2797-9178
Minna Luo https://orcid.org/0000-0003-1855-985X
Muqing Cao https://orcid.org/0000-0002-0352-6548
Xu Ma https://orcid.org/0000-0001-7674-3589

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