Introduction

Zellweger spectrum disorder (ZSD) is a phenotypic continuum including Zellweger syndrome (ZS; OMIM #214100), neonatal adrenoleukodystrophy (NALD; OMIM #202370) and infantile Refsum disease (IRD, OMIM #266510). ZS is the most severe form of ZSD and is characterized by craniofacial dysmorphism and severe neurological abnormalities (1,2). Life expectancy in infants with ZS is severely limited, with rarely a one-year survival (3). Patients with ZS have severe hypotonia, difficulty feeding, hepatomegaly, ocular disorders, and seizures. In addition, this phenotype is characterized by craniofacial abnormalities, including widely fontanelle, cranial sutures, prominent forehead, and low-set ears (4).

ZSD is an autosomal recessive inherited disorder caused by a genetic defect (called PEX) involved in peroxisomal biosynthesis. Up to now, there have been 14 ZSD-PEX gene genotype-phenotype correlation in neonates with Zellweger syndrome

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Background: Zellweger syndrome (ZS) is commonly manifested as facial deformities, hypotonia, and liver dysfunction. However, ZS caused by PEX26 gene mutation shows a broad and dispersed clinical pattern. In this study, the PEX26 gene in ZS was analyzed to enrich its clinical characteristics. Meanwhile, phenotypic and genotypic characteristics of Zellweger spectrum disorder (ZSD) induced by PEX26 mutation were evaluated.

Methods: The clinical data of newborn with ZS in our hospital were analyzed retrospectively. We performed WES and found that the infant carried the PEX26 gene variant. We searched the biomedical literature databases (PubMed, Web of Science, and EMBASE) to compare clinical features and genotypes.

Results: The neonate developed facial deformities, hypotonia, feeding difficulties, and seizures. Her homozygous variant was found in the PEX26 gene (NM_017929: exon2: c.34del) inherited from both parents. Electronic databases, including our case, reported 32 pathogenic variants in PEX26. We found that variation c.292C> T accounted for the largest proportion of PEX26 mutations (16/66, 24.24%). The proportion of deleterious mutations in ZS patients was significantly higher than that in NALD and IRD patients.

Conclusions: We identified pathogenic variations in the PEX26 gene and expanded the known mutant spectrum. By comparing patients with PEX26 mutations, the study determined that a significantly higher percentage of deleterious mutations in ZS was associated with severe clinical phenotypic characteristics.

Keywords: PEX26; Zellweger syndrome (ZS); Zellweger spectrum disorder (ZSD); newborn

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genes associated with ZSD (5). These PEX genes encode proteins required for peroxisomal biogenesis, including membrane biogenesis, division, and include the mechanism of peroxisomal matrix introduction (6,7). Among all ZSD patients, PEX1 mutation was the most common cause, accounting for 58%. Nevertheless, PEX26 mutations account for only 3% of total ZSD (8). Therefore, the phenotype of ZS caused by PEX26 gene mutation is wide and dispersed.

In this study, we reported a neonatal ZS case with a novel homozygous mutation in PEX26 gene. We also searched and retrospectively analyzed PEX26 mutations in neonatal cases diagnosed with ZS from the database. This research enriches the clinical features and evaluates the phenotypic and genotypic associations induced by ZSD with PEX26 mutations.

We present the following article in accordance with the MDAR reporting checklist (available at https://dx.doi.org/10.21037/tp-21-103).

Methods

Case presentation

We investigated the case of a near-term infant, female, born at 36+6 weeks gestation, weighing 2,230 g. The infant was transferred to the Neonatal Intensive Care Unit (NICU) of the Children’s Hospital of Fudan University at 3 hours after birth, due to polypnea and birth asphyxia. Mom had normal prenatal care. Fetal ultrasonography at 28 weeks revealed bilateral cerebral ventriculomegaly, and fetal magnetic resonance imaging (MRI) manifested enlargement of posterior horn of right ventricle. Her parents were healthy and non-consanguineous. The infant was born with hypotonia and had no spontaneous respiration, required intubation at birth. Her Apgar scores were 4 at 1 minute, 4 at 5 minutes and 8 at 10 minutes respectively. Physical examination on admission revealed large anterior and posterior fontanel with prominent forehead and widely separated sutures, low-set ears, systemic hypotonia, minimal spontaneous movement, and transverse palmar creases on the left hand.

Initial laboratory workup demonstrated hyperbilirubinemia (327.6 μmol/L) and elevated liver enzymes (aspartate aminotransferase, 245.3 IU/L; alanine aminotransferase, 143.3 IU/L), suggesting hepatic dysfunction. Urine organic acids revealed obvious elevated 4-hydroxyphenyllactic acid. Very long chain fatty acid of plasma analysis showed normal C22 and C24, but dominant elevated C26 (>11.69 nmol/L, normal <1.3 nmol/L), C24/C22 (2.32, normal ≤1.39), C26/C22 (>0.649, normal ≤0.023). The chest X-ray showed right pneumothorax. Abdominal ultrasound showed indistinct structure of bilateral kidneys. Head ultrasound indicated bilateral cerebral ventriculomegaly, which confirmed by fetal cerebral MRI (Figure 1). A bleeding spot was found on the right fundus by ophthalmic examination. During hospitalization, she was initially treated with mechanical ventilated and ursodeoxycholic acid. She was discharged home with low flow nasal prone oxygen and fed by nasogastric tube due to hypotonia and weak uncoordinated suck reflex.

The infant developed an epileptic seizure when she was eight months old and was brought to our hospital for treatment. Video-electroencephalogram (VEEG) showed hypersrrhythmia during both wakefulness and sleep, especially in the posterior area. Moreover, focal seizures were detected during sleep (Figure 2). With palliative care, she died at 9 months of age from dyspnea and...
cardiopulmonary arrest.

**Whole exome sequencing (WES)**

A trio WES was performed by WuXi NextCODE Genomics, Shanghai, China (CLIA Lab ID: 99D2064856) using a previous described protocol. Briefly, exome capture was performed using the Agilent SureSelect Human All Exon V5, Illumina TruSeq Rapid PE Cluster, and SBS kits (Agilent Technologies, Santa Clara, CA, USA). WES was performed on the Illumina HiSeq 2000/2500 platform. Reads were aligned to the human genome reference sequence (GRCh37/hg19 build of UCSC Genome Browser; http://genome.ucsc.edu) with the Burrows-Wheeler Aligner v.0.6.2. Duplicate paired-end reads were marked with Picard v.1.55 (https://broadinstitute.github.io/picard/). The Genome Analysis Toolkit v.2.3–9 was used for base quality score recalibration, indel realignment, and variant discovery. Variants were annotated using a pipeline developed in-house and filtered in the Exome Variant Server, gnomAD, Exome Aggregation Consortium, or the dbSNP databases. The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013). The study was approved by the ethics committee of Children's Hospital of Fudan University (IRB No.2021-237). The parents of the patient provided written informed consent.

**Literature review and data analysis**

A comprehensive literature search was performed to identify the relevant researches and case reports in PubMed, the Web of Science and EMBASE from inception to August 1, 2020. We used the following search strategy “((PEX26 OR PEX26 gene) AND ((Zellweger syndrome) OR (ZS) OR (ZSD) OR (Peroxisome biogenesis disorder) OR (Zellweger spectrum disorder)))”. In the Chinese database, literature researches were conducted by using the “PEX26”, “Zellweger syndrome”, and “Peroxisome biogenesis disorder” as keywords from the inception to August 1, 2020, and no relevant literatures were found.

**Statistical analysis**

The genotypes of ZSD with the mutation of PEX26 and

![Figure 2 VEEG. VEEG findings at the 8 months of age. Wakefulness (A) and asleep periods (B), with hypsarrhythmia characterized by asynchronous, multifocal high amplitude slow waves, intermingled with multifocal spikes, which was prominent in the posterior. (C) Epileptic spasms appear quite typical in morphology and a focal seizure was detected during sleep. VEEG, video-electroencephalogram.](image)
their clinical subtypes were compared. Differences in the occurrence of deleterious variants in ZS groups and NALD/IRD groups were analyzed by using Fisher exact test. A two-sided P value was used to test for statistical significance (P<0.05).

Results

Whole exome sequencing

Our case depended on clinical manifestations and laboratory studies for diagnosis, and confirmation is based on the homozygosity variant in the PEX26 gene (NM_017929: exon2: c.34del) that we identified. The variant was validated by Sanger sequencing on a 3500XL instrument, and both parents are heterozygous carriers. We identified this variant as pathogenic variant because the type of this variant is nonsense, which loss of function is a known mechanism of this gene. Moreover, this variant is absent from ExAC, the 1000 Genomes database, and HGMD, ClinVar database.

Clinical characteristic of newborns with ZS caused by PEX26 mutation

A total of 9 articles referring to 31 cases of ZSD with PEX26 pathogenic variants were identified form the electronic database (9-17). However, after careful filtering, only four ZS infants were described in detail in postnatal clinical manifestations. Other babies were mentioned only briefly about the mutation site or were not found to have neonatal presentation. These four infants are from Saudi Arabia (9), Tunisian (10) and America (11,12). Finally, combined with our case, the clinical phenotypes of 5 ZS infants were summarized and shown in Table 1 (male =4, female =1). All (100%, 5/5) infants were characterized by hypotonia, with 80% (4/5) having difficulty feeding, but only 40% (2/5) presenting with seizures. All (100%, 5/5) infants were born with facial deformities and liver dysfunction. 80% (4/5) of the patients were characterized by enlarged fontanelles and prominent forehead. 60% (3/5) of patients had wide cranial sutures and low-set ears. Kidney abnormalities occurred in only 40% of cases (2/5), only one of the patients existed renal cysts. Other systems involved are ventricular dilation, fundus lesions, and skeletal abnormalities (such as bilateral clubfeet and chondrodyplasia). All plasma analyses of very long chain fatty acids showed elevated C26 and/or C24. The prognosis is poor, with three out of five infants failing to survive the first year of life.
Among the 32 pathogenic variants (our case and the 31 previously reported ones were reviewed), 9 cases were diagnosed as ZS, 4 as NALD, 8 as IRD, and 11 cases were not clinically diagnosed, as shown in Table 2. All showed recessive models, 21 (65.6%) homozygous variants and 11 (34.4%) complex heterozygous variants were identified. It is worth noting that all the mutations in ZS are homozygous, although it contains several types of variation (frameshift, nonsense, missense, and splicing mutations). Therefore, complex heterozygous mutations were found only in NALD and IRD cases, mainly in IRD cases (7/8, 87.5%).

Currently, 22 disease-causing mutations of ZSD have been reported in the PEX26 gene identified in these 32 patients. Among them, missense variant accounted for 45.5% (10/22), nonsense variant accounted for 22.7% (5/22), frameshift variant accounted for 22.7% (5/22), and splicing variant accounted for 9.1% (2/22). Notably, the missense variant was the most common form of all mutations (42/66, 63.64%) and was identified in 24 patients (3 with ZS, 4 with NALD, 8 with IRD, and 9 with unknown diagnosis). C. 292C>T variant was detected in both NALD and IRD, accounting for the largest proportion (16/66, 24.24%), but it has not been reported in ZS.

Of these 32 patients, 9 were diagnosed with ZS, of which 3 had missense mutations (33.3%, 6/18), and the remaining 6 had harmful mutations (66.7%, 12/18), including nonsense, splices, or frameshift mutations. Another 12 patients were diagnosed with NALD or IRD, with missense mutations and deleterious mutations accounting for 84.6% (22/26) and 15.4% (4/26) of the total mutations, respectively (Table 2). The proportion of harmful mutations in ZS patients was significantly higher than that in NALD and IRD patients (P<0.001) (Table 3).

### Table 2 Statistics on the mutation sites of PEX26 gene mutation leading to ZSD

| Case | ZS | NALD | IRD | Undescribed |
|------|----|------|-----|-------------|
|      | Mutation | Zygosity | Mutation | Zygosity | Mutation | Zygosity | Mutation | Zygosity |
| 1    | c.265G>A. p. (Gly89Arg) | Hom | c.292C>T. p. (Arg98Trp) | Hom | c.2T>C. p. (Met1Thr), c.134T>C. p. (Leu45Pro) | Het | c.292C>T. p. (Arg98Trp) | Hom |
| 2    | c.265G>A. p. (Gly89Arg) | Hom | c.292C>T. p. (Arg98Trp) | Hom | c.457C>G. p. (Leu153Val) | Het | c.296G>A. p. (Trp99*) | Hom |
| 3    | c.230+1G>T | Hom | c.292C>T. p. (Arg98Trp) | Hom | c.457C>G. p. (Leu153Val) | Het | c.192-216del, c.353C>G | Het |
| 4    | c.256T>C. p. (Cys86Arg) | Het | c.292C>T. p. (Arg98Trp), c.134T>C. p. (Leu45Pro) | Het | c.292C>T. p. (Arg98Trp), c.255insT | Het | c.37_38del. p. (Arg13Glyfs*101), c.667+2T>C | Het |
| 5    | c.296G>A. p. (Trp99*) | Hom |                  |                  | c.292C>T. p. (Arg98Trp), c.131T>C. p. (Leu44Pro) | Het | c.292C>T. p. (Arg98Trp), c.574C>T. p. (Arg192*) | Het |
| 6    | c.T35insC | Hom |                  |                  | c.292C>T. p. (Arg98Trp), c.131T>C. p. (Leu44Pro) | Het | c.153C>A. p. (Phe51Leu) | Hom |
| 7    | c.37_38del. p. (Arg13Glyfs*101) | Hom |                  |                  | c.292C>T. p. (Arg98Trp), c.426_548dup122bpinsT | Het | c.153C>A. p. (Phe51Leu) | Hom |
| 8    | c.315 G>A. p. (Trp105*) | Hom |                  |                  | c.292C>T. p. (Arg98Trp) | Het | c.153C>A. p. (Phe51Leu) | Hom |
| 9    | c.34del. p. (Leu12Serfs*70) | Hom |                  |                  |                  |                  | c.153C>A. p. (Phe51Leu) | Hom |
| 10   |                  |                  |                  |                  |                  |                  | c.153C>A. p. (Phe51Leu) | Hom |
| 11   |                  |                  |                  |                  |                  |                  | c.153C>A. p. (Phe51Leu) | Hom |
| Sum  | 9 | 4 | 8 | 11 |

Case reported this time. Mutation sites are collected from references (5-13). Het, heterozygous; Hom, homozygous.
Discussion

Considering its pathophysiology, ZSD has been classified into three categories. The first group: ZS, critically sick newborns, most of whom do not survive beyond the first year. Group 2: NALD, infantile, usually has adrenal cortex atrophy, and most children die before the age of three to five years. Group 3: IRD, mild. Most of those affected suffer from ataxia, and many of them can go into adolescence (3,18,19). The differences between the three diseases were due to the severity of the clinical phenotype and the mutation of the genotype. Recently, Heimler syndrome, a very mild and unique phenotype associated with defects in the PEX gene, has been reported. Heimler syndrome is considered to be part of the ZSD continuum (20,21), but is not generally recognized.

ZS is a multisystem disease that is apparent at birth. Findings include severe characteristic craniofacial deformities, hypotonia, and seizures. Hepatomegaly and renal cortical cyst accounted for 80% and 70% respectively (3). Eye abnormalities such as cataracts and sensorineural hearing loss are typical but not always recognizable when first presented (22). Our literature review found that in all ZS infants with PEX26 mutations, hypotonia, facial deformities, abnormal liver function and elevated C26 / C24 were prominent signs. However, the incidence of seizures in the neonatal period (40%) is lower than that of hypotonia and liver disease, which may be related to atypical at the neonatal period or too early death to present. Meanwhile, PEX1 was the most involved gene, accounting for nearly 70% of patients with ZSD. Preuss et al. found that renal cysts are typical of PEX1 mutated infants (23), which is different from our results (only one of five renal cysts exists). With respect to bone signs, two-fifths of the newborns in our study showed bilateral clubfeet, which has not been reported in ZS infants with PEX1 mutation. Brain MRI or ultrasound revealed polymicrogyria and ventriculomegaly in 80% of ZS infants with PEX26 mutations. These brain disorders may contribute to the diagnosis of the disease.

PEX26 encodes a 34-kDa peroxisomal membrane protein of 305 amino acids, that assists in the localization of PEX1-PEX6 complexes to the peroxisomal membrane (6,15,24). The C-terminal part of PEX26 is exposed to the matrix side with one transmembrane domain (aa252-269) located in the peroxisomal membrane, while the N-terminal part is cytosol-exposed and vital in its biological activity. The PEX6 binding domain located in the N-terminal region of the protein corresponds to the amino acid 29-174 (14,17,25), which is essential for binding to PEX6. The PEX1-PEX6 AAA ATPase complex was recruited and anchored to the peroxisomal membrane (26,27). However, Tamura et al. believed that PEX26 residues amino acids13–48 was sufficient to bind to PEX6 (28).

The strongly represented mutation in ZSD with PEX26 mutations is the missense mutation c.292C>T, which changes the arginine located at amino acid position 98 into a trytophan (p. Arg98Trp). This missense mutation affects the stable expression of PEX26. The PEX6 binding domain is destroyed and the ability to participate in the interaction between PEX6 and PEX1 is reduced (27,28). However, the effect of recurrent c.292C> T mutation was relatively minor. It has been reported not only in NALD but also in IRD. The same mutation suggests that different clinical phenotypes, which may be related to the temperature sensitivity of PEX26 cell lines (14). Since this mutation has not yet been identified in ZS, we infer that the missense mutation c.292C>T causes residual protein levels and function, and thus exhibits a relatively mild clinical phenotype.

Mutations of PEX26 are distributed across the entire gene, but 70% (14/20) of the deleterious mutations of ZS are located in the N-terminal part (Figure 3), including nonsense, frameshift and splicing mutations, that theoretically causes a truncation of functional domain

| Variants | ZS (N) | NALD&IRD (N) | Total (N) |
|----------|--------|--------------|-----------|
| Missense | 6      | 22           | 28        |
| Deleterious | 12    | 4            | 26        |
| Total    | 18     | 26           | 44        |

P = 0.0005

N>40; Tmin=16x18/44=6.5>5; Z=3.477. P=0.0005. N, number of cases; ZS, zellweger syndrome; NALD, neonatal adrenoleukodystrophy; IRD, infantile Refsum disease.
and/or large domain deletion in peroxins, resulting in dysfunction. The closer the mutation site is to the N terminal, the greater the damage to the structure and homeostasis of peroxisomes. ZS patient-derived PEX26 mutants seriously impaired peroxisomal matrix protein import, which progressively influence the energy GTP generated and provided for the fission of peroxisomes (29). Thus, these mutations are total or almost total loss of protein levels and function, and are most severe in patients with ZS. In contrast, missense variants that retain partial function have less effect on peroxisome assembly (30,31). Patients with NALD or IRD, mainly composed of missense mutations, showed moderate or mild clinical phenotypes. The severity of the mutation depends on the effect on the function of PEX26. Our result showed that the proportion of deleterious mutation in ZS is significantly higher than NALD/IRD (\(P<0.05\)). Thus, this may explain why the ZS phenotype is more severe than other phenotypes. The type of PEX26 mutation can help predict the severity of individual cases.

In conclusion, we reported a ZS neonate developed facial deformities, hypotonia, feeding difficulties, and seizures. The new pathogenic mutation c.34del was found in the PEX26 gene. We also summarized the manifestations of reported ZS neonates with PEX26 mutation, including characteristic craniofacial dysmorphism, hypotonia, cerebral dysfunctions (cerebral dysfunctions), liver dysfunction. Elevated C26 and/or C24 in VLCFA is usually shown in blood plasma analysis. However, seizures and renal cysts are less obvious for neonatal presenting in newborns with PEX26 mutation. In addition, the c.292C>T mutation occurred at a relatively high frequency in ZSD with the PEX26 mutation, which resulted in a relatively mild clinical phenotype. Finally, we compared the occurrence of deleterious variants, associated with a disease severity, between ZS groups and NALD/IRD groups, ZS accounting for the significant higher percentage and therefore demonstrated the most severe clinical manifestation.

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**Footnote**

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