A novel nonsense SMC1A mutation in a patient with intractable epilepsy and cardiac malformation

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Abstract

Cornelia de Lange syndrome (CdLS) is a cohesinopathy caused by genetic variations. We present a female with SMC1A-associated CdLS with a novel SMC1A truncation mutation (p. Arg499Ter), transposition of the great arteries, and periodic intractable seizures from 40 months of age. A review of the literature revealed that a seizure-free period after birth of at least 15 months is required for these patients to be able to walk, irrespective of the epileptic course.

Cornelia de Lange syndrome (CdLS) [MIM: 122470, 300590, 300882, 610759, and 614701] is a congenital multisystemic disorder with widely varied characteristics ranging from mild (nonclassical phenotype) to severe (classical phenotype) that is caused by genetic variants of structural or regulatory components of the cohesin complex1,2. Classical CdLS is caused by mutations in NIPBL, while nonclassical CdLS is caused by mutations in SMC1A, SMC3A, RAD21, and HDAC83. Missense variants and small in-frame deletions in SMC1A, located at Xp11.22, account for ~5% of CdLS cases. A total of nine cases of SMC1A-associated CdLS with a male-to-female ratio of 1:23. A review of the literature revealed 60 cases of SMC1A-associated CdLS with congenital cardiac defects (CHD) have been reviewed4, although the incidence of CHD in patients with CdLS is ~30%5. SMC1A-related CdLS arises from a dominant negative effect in females2. Females with SMC1A mutations leading to protein truncation are affected by intractable epilepsy, severe developmental retardation, and few craniofacial differences4,6. Age at presentation with first epileptic seizures ranges from <1 month to 17 months6.

In our case of CdLS with an SMC1A truncating mutation, the female patient was the second child born to a healthy, nonconsanguineous couple when her mother and father were 35 and 42 years of age, respectively. There was no family history of CdLS. She was born via emergency cesarean section at 35 weeks gestation due to fetal distress. At birth, her weight was 1636 g (−1.9 SD), length was 43.5 cm (−0.43 SD), and occipitofrontal circumference (OFC) was 30.2 cm (−0.6 SD). She had been hospitalized in the neonatal intensive care unit for 3 months because of failure to thrive and transposition of the great arteries (TGA) type III congenital heart defects with a ventricular septal defect and pulmonary artery stenosis. At 11 months of age, the Blalock–Taussig shunt operation was performed, followed by home oxygen therapy. At 1 year 10 months of age, the Nissen operation was performed for gastroesophageal reflux disease. At 3 years 4 months of age, tonic seizures emerged. Hypoglycemia and high ammonia levels appeared. Hyperammonemia (405 µg/dl; normal range: 36–86 µg/dl) and repetitive hypoglycemia occurred after she suffered from bacterial pneumonia at 4 years 7 months of age. She had been taking six meals a day, two with cornstarch. She was referred to our clinic for detailed examination of...
metabolic diseases. At 4 years 9 months, she weighed 14.1 kg (−1.3 SD), her height was 100.5 cm (−1.1 SD), and her OFC was 46.8 cm (−2.2 SD). She had a prominent forehead, hypertelorism, thick eyebrows, broad nasal tip, depressed nasal bridge, full cheeks, left cupped ear, right prominent antihelix, high-arched palate, thin upper lip, crowded teeth, slender fingers, left talipes varus, and left second short toe (Fig. 1a–g). The results of all biochemical tests and gas chromatography/mass spectrometry (GC/MS) of urine and tandem mass spectrometry (MS/MS) of dry blood spots were normal. Hypoglycemia was observed during hospitalization, and occasionally episodes caused hypoglycemia irrespective of meal time. Following this, there were no episodes of hypoglycemia for 3 years. A fasting test was conducted at 7 years 3 months of age. Hypoglycemia (37 mg/dl; normal range 70–105 mg/dl) appeared 17 h after the last meal, but there were no findings of coldness, lethargy, hyperammonemia, or metabolic acidosis. Additionally, all urine GC/MS and dry blood spot MS/MS results were normal. Magnetic resonance imaging of the brain revealed normal findings. She had left sensorineural deafness (100 dB). Her developmental milestones were delayed: head control at 6 months of age, rollover at 6 months, sitting unaided at 2 years and 4 months, and walking at 2 years and 5 months. At 15 years 2 months of age, she was unable to speak meaningful words, her height was 133.0 cm (−4.6 SD), and her weight was 27.1 kg (−3.0 SD). Her menstruation started regularly at 14 years 9 months of age. At 13 years of age, six persistent deciduous teeth with no dental caries were extracted.

Three-day-long continuous intractable generalized tonic seizures (GTS) occurred every 2 weeks from 3 years to 4 months of age. At 15 years of age, moderately beneficial antiepileptic drugs (AEDs) were lamotrigine (LMT) and levetiracetam (LEV). AEDs tried without success were clobazam (CLB), potassium bromide (KBr), and carbamazepine (CBZ). She had a Jatene operation for TGA type III at 5 years 2 months of age and corrective surgery for the left talipes varus at 6 years 10 months. Chromosomal analysis showed a normal karyotype. We searched the original computerized database for possible malformation syndromes: UR-DBMS (University of the Ryukyu-Database for Malformation Syndromes: http://becomerich.lab.u-ryukyu.ac.jp) edited by Naritomi7. Suggested candidates matching over 12 signs were 4pter-p13 trisomy and Xpter-p21 monosomy, 13q14-qter trisomy and 5pter-p13 monosomy, CDLS1 (MIM 122470), CDLS3 (MIM 610759), Noonan syndrome 1 (NS1) (MIM 163950), 1q21.1 deletion syndrome, 12p trisomy, and
Table 1 Characteristics in order of age of first seizures in our patient and previously reported cases

| Feature                                      | Case 1 | Case 2 | Case 3 | Case 4 | Case 5 | Case 6 | Case 7 | Case 8 | Case 9 | Case 10 | Case 11 | Case 12 | Case 13 | Case 14 | Case 15 | Case 16 |
|----------------------------------------------|--------|--------|--------|--------|--------|--------|--------|--------|--------|---------|---------|---------|---------|---------|---------|---------|
| Age at reported (years)                      | Died aged 11M | 7 | 4 | 6 | Died aged 9YM | 4 | 3 | 8 | 6 | 10 | 46 | 6 | 3 | 14 | 14 | 15 |
| Birth OFC Z-score/ most recent               | −3.9/−2.5 | Unknown | −1.5/−4.5 | Unknown/ −2.0 | −1.6/−0.8 | −2.0/−2.5 | Unknown/ −3.0 | −1.2/−2.0 | Unknown/−2.5 | −0.8/−3.5 | −1.0/0.0 | Unknown/ −1.7 | Unknown/ −2.0 | −0.6/−2.2 |
| Most recent height Z-score/ most recent      | −2.5 | −2.6 | −5 | Unknown | 0006 | −2.6 | −3.2 | −4.5 | −2.5 | −2.3 | −0.05 | −2 | −3.7 | −4.6 |
| Developmental impairment                     | Unknown | Unknown | Moderate-severe | Severe | Unknown | Severe | Severe | Severe | Moderate-severe | Unknown | Unknown | Moderate-severe | Severe |
| Gross motor development                      | None | Unable to sit | Can take a couple of steps with support | Unable to sit without support | Non ambulant | Unable to sit | No independent mobility | Walking from 30 months | Unable to sit without support | Never crawled or walked | Run with unsteady gait | Walked at 12 months | Walked at 2 years suddenly stopped walking at 5 years | Walking from 23 years | Unsteady on feet, aged 7 |
| Speech                                       | None | None | None | Coos, laughs, cries appropriately | None | None | None | None | Lost speech aged 3 years following SE | None | None | None | None | None |
| Age at first seizure                         | <1 month | <1 month | 4 weeks | 5–6 weeks | 2 months | 4 months | 4 months | 5 months | 5 months | 6 months | 9 months | 15 months | 17 months | 2 years | 28 months | 40 months |
| Seizure types                                | Unknown | Focal with eyelid myoclonia, Focal, spasms | Bilateral tonic, focal, myoclonic, CSE | GTCS, Focal, myoclonic, CSE | Generalized tonic, focal, myoclonic, CSE | Generalized tonic, focal, myoclonic, atypical absence, focal, myoclonic | Generalized tonic, bilateral tonic, focal, myoclonic, atypical absence, focal, myoclonic, CSE | Generalized tonic, focal, myoclonic, atypical absence, focal, myoclonic, CSE | GTCS, myoclonic, atypical absence, focal, myoclonic, atypical absence, focal, myoclonic, CSE | GTCS, myoclonic, atypical absence, focal, myoclonic, atypical absence, focal, myoclonic, CSE |
| Seizure clusters                             | − | − | − | − | − | + | + | + | + | + | + | + | + | + | + | + |
| Seizure freedom                              | − | − | − | + | − | + | + | + | + | + | + | + | + | + | + | + |
| SMC1A: Amino acid change                     | C.2477delA | p.Arg825fs*48 | p.Arg1016Ter | p.Arg1016Ter | p.Glu1381Ter | p.Glu1381Ter | p.Glu1381Ter | p.Arg755Ter | p.Arg1016Ter | p.Arg1016Ter | p.Arg1016Ter | p.Arg1016Ter | p.Arg1016Ter | p.Arg1016Ter | p.Arg1016Ter | p.Arg1016Ter |
| Reference                                    | 6 | 9 | 6 | 6 | 6 | 10 | 6 | 6 | 6 | 10 | 11 | 6 | 10 | 11 | 6 | This report |

GTCS generalized tonic-clonic seizure, CSE convulsive status epilepticus, FS febrile seizure, NCSE nonconvulsive status epilepticus
16p11.2 deletion syndrome. Whole-exome sequencing was performed using the SureSelect Human All Exon V6 Kit (Agilent Technologies, Santa Clara, CA) and HiSeq2500 (Illumina, San Diego, CA). To identify disease-causing mutations, we excluded all known variants found in the 1000 Genomes database (http://www.internationalgenome.org/), Japanese Genomes database, dbSNP (http://www.ncbi.nlm.nih.gov/SNP), the genome Aggregation Database (gnomAD; http://gnomad.broadinstitute.org/), and the Human Genetic Variation Database (HGVD; http://www.genome.med.kyoto-u.ac.jp/SnpDB/). Heterozygous SMC1A (NM_006306) mutations cause CdLS, the symptoms of which fit with those of our patient. We identified a heterozygous single nucleotide variation (c. C1495T) in SMC1A exon 9 that results in a nonsense mutation (p. Arg499Ter) and a truncated protein. The p. Arg499Ter variant was not detected in her parents, suggesting that the variant was de novo. This was confirmed by Sanger sequencing (Fig. 1b). This is the first description of the p. Arg499Ter variant. Mutation Taster (http://www.mutationtaster.org) predictions indicate that this is a disease-causing variant. This study was performed in accordance with the standards of the Ethics Committee of the Ryukyus Graduate School of Medicine (Okinawa, Japan). Informed consent was obtained from her parents by Dr. Yasutsugu Chinen.

Our patient with SMC1A-associated CdLS had a non-classical CdLS type. This was determined using the clinical CdLS scoring system, which consists of 9 points, including three cardinal and three suggestive features. Her first seizures, which occurred at 40 months of age, were the latest onset recorded. Seizure onset occurred much earlier in the 15 previously reported patients with protein truncating mutations in SMC1A. When these cases, including the one presented here, were arranged in order of age of first seizures, we observed that only one (9.1%) of the 11 patients who had their first seizure at less than 9 months of age could walk. In contrast, all of the patients (100%) who had their first seizure after 15 months could walk. This observation was independent of the type of epilepsy and the truncated amino acid position in SMC1A (Table 1, modified from Symonds et al. 2017). Although the number of cases is small, this observation suggests a correlation between age of epilepsy onset and independent walking. If the first seizures develop before the patient could walk, walking without assistance may be difficult. Only one patient, case 9, developed seizures in the 5 months after birth and was able to walk at 30 months of age. This patient was seizure-free for 1 year after commencement of levetiracetam treatment, but the seizures recurred. In case 4, seizures started from 5–6 weeks of age and stopped occurring at 5 years of age after commencement of phenobarbitone treatment and adoption of a ketogenic diet. In case 8, seizures started from 5 months of age and stopped occurring at 7 years of age after commencement of gabapentin treatment and adoption of a ketogenic diet. Taken together, these cases indicate that at least the first 15 months after birth needs to be seizure-free for these patients to walk, irrespective of the epileptic course.

CHD with SMC1A-associated CdLS has been previously reported in 13 cases, including ours, yet this is the first report including a TGA diagnosis (Supplementary Table 1). In patients with CHD, the rate of substitution or mutation at an arginine position is 42% (5/12), and in patients without CHD, the rate is 50% (11/22). The post-translational modification of proteins by arginine methylation is functionally important in the SMC1A protein. Dysregulation of cohesin by the SMC1A protein may cause CHD in a zebrafish cohesinopathy model. The same variations at arginine positions, including p. Val58-Arg62del, p. Arg196, p. Arg496, and p. Arg693, revealed discordant heart defects. However, there may be no relationship between SMC1A arginine substitution and CHD, and we were unable to show whether such SMC1A variations affect arginine methylation.

Our patient’s episodes of hypoglycemia showed no remarkable causal disease and were considered clinical symptoms caused by dumping syndrome. However, we could not regularly observe reproducible events. The fasting test at 7 years 3 months of age revealed a normal reaction without hypoglycemic episodes. Infancy is considered to be a time period of low glycogen storage corresponding to age. If hypoglycemia occurs during this period, further careful investigation and dietary adjustments might be necessary.

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HGV Database
The relevant data from this Data Report are hosted at the Human Genome Variation Database at https://doi.org/10.6084/m9.figshare.hgv.2570.

Conflict of interest
The authors declare that they have no conflict of interest.

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