De novo 2q36.3q37.1 deletion encompassing *TRIP12* and *NPPC* yields distinct phenotypes

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**Abstract**

We report a patient with developmental delay, extremely short stature, small hands, dysmorphic facial features, hearing loss, and epilepsy carrying a de novo 2.76-Mb deletion of 2q36.3q37.1, including *TRIP12* and *NPPC*. *TRIP12* haploinsufficiency causes developmental delay with isolated dysmorphic facial features, whereas *NPPC* haploinsufficiency causes short stature and small hands. This is the first report of a unique phenotype, which is secondary to a microdeletion encompassing *TRIP12* and *NPPC*.

Thyroid hormone receptor interacting protein 12 (*TRIP12*) is a human protein homologous to the E6-AP carboxyl terminus (HECT) domain E3 ubiquitin ligase and is involved in cell cycle progression and the maintenance of chromosome integrity. *TRIP12* (MIM604506) haploinsufficiency has been reported to cause developmental delay, autism spectrum disorder (ASD), and facial dysmorphisms, which are collectively named Clark–Baraitser syndrome (MIM #617752).

C-type natriuretic peptide (CNP) plays an important role in cartilage homeostasis and endochondral bone formation and is encoded by the gene *natriuretic peptide precursor-C* (*NPPC*; MIM600296). *NPPC* variants have been reported to cause autosomal dominant idiopathic short stature with small hands. Here, we describe a case of a 4-year-old boy with a de novo 2q36.3q37.1 deletion encompassing *TRIP12* and *NPPC* who presented with severe developmental delay, extremely short stature, small hands, dysmorphic facial features, hearing loss, and epilepsy.

The boy was born to healthy nonconsanguineous Japanese parents at 40 gestational weeks. His parents had no medical history. At birth, his weight, length, and occipitofrontal circumference (OFC) were 2710 g (−0.7 standard deviation (SD)), 46.6 cm (−1.1 SD), and 33 cm (−0.4 SD), respectively. Automated auditory brainstem response detection in a neonatal hearing screening test revealed a refer outcome. Urinal cytomegalovirus DNA was not detected in the neonatal period. Hearing threshold levels were bilateral sensorineural hearing loss of 70 dB at the age of 6 months and 30–40 dB at 4 years. He showed severe developmental delay and could hold up his head at 9 months, sit alone at 14 months, and stand with support at ~2 years. At 4 years, when this report was written, he was unable to speak or stand alone. He displayed behavioral abnormalities, such as aggression, repeatedly biting his intravenous drip line during hospitalization, and frequently striking his head against the wall. His short stature became apparent by 6 months of age (Fig. 1a). His height, weight, and OFC were 76.5 cm (−4.0 SD), 8.5 kg (−3.3 SD), and 50.5 cm (+0.9 SD) at 3 years, respectively, indicating relative macrocephaly (OFC more than +1 SD compared to height). We performed several examinations to evaluate his severe short stature. Complete blood count, blood chemistry, and thyroid function tests showed normal results. Insulin-like growth factor-1 was 46 mg/mL (−1.1 SD) at 3 years. His karyotype was 46, XY. A growth hormone stimulation test was not performed because his...
parents did not provide consent. He experienced four febrile seizures and two afebrile seizures at 2 years. His electroencephalogram showed frequent spike or spike-and-wave discharges in the left parietal region during drowsy and sleep states. He was diagnosed with epilepsy, and levetiracetam was started at 3 years. Head magnetic resonance imaging showed normal results. His facial features included a prominent forehead, mild hypertelorism, epicanthal folds, short nose, low hanging columella, short philtrum, everted vermilion of the upper lip, strabismus, low-set ears (Fig. 1b, c), and his hands were small (Fig. 1d). After obtaining written informed consent, we performed array comparative genomic hybridization analysis and identified a de novo 2.76 Mb deletion (arr[GRCh37]

Fig. 1 Growth curve, images of face and hands, and deleted genes in the patient. a The patient’s growth curve. b, c The patient exhibited prominent forehead, mild hypertelorism, epicanthal folds, short nose, low hanging columella, short philtrum, everted vermilion of the upper lip, strabismus, low-set ears, and relative macrocephaly at 4 years of age. We obtained written informed consent from the patient’s family for publication of this clinical report and accompanying images. d The patient presented with small hands. e Twenty-five genes in the 2.76-Mb 2q36.3q37.1 deletion (230377789_233136164) and deleted region of our case and previously reported copy number variant (CNV) deletion cases involving TRIP12 or NPPC in Table 1.
by loss-of-function variation because the phenotype caused of being loss-of-function intolerant. Indicating that these three genes are probably intolerant to 11 patients, including four previously reported cases9,10,15 clinical features, Bramswig et al. and Zang et al. reported describe detailed patient clinical information. Regarding genes, dominant inheritance and cause developmental delay and known to affect phenotypes attributable to autosomal haploinsufficiency is not thought to confer significant survival or reproductive disadvantages. In addition, although abnormalities in SP110, ARMC9, PDE6D, and DIS3L2, which reside in the deleted region, have been reported to cause autosomal recessive inheritance diseases, our patient displayed different clinical features than those observed with these diseases.

Genetic findings related to TRIP12 mainly include rare de novo variants, which have been identified from several large ASD or ID proband cohorts by whole-exome or target sequencing. However, these reports did not describe detailed patient clinical information. Regarding clinical features, Bramswig et al. and Zang et al. reported 11 patients, including four previously reported cases9,10,15 and nine patients with inactivating single nucleotide variants or copy number variant (CNV) deletions in TRIP12, respectively, in 2017-2.3. CNP encoded by NPPC plays an important role in skeletal development. Additionally, NPPC alteration in humans was reported by Tassano et al. for the first time in a boy with short stature and several dysmorphic features; he had a 1.91 Mb 2q37 deletion, including NPPC (case-N1 in Fig. 1e and Table 1)16. Two NPPC variants were identified in six patients with short stature and small hands from only two families by Hisado-Oliva et al.6. They also showed that COP-7 cells transfected with NPPC variants detected in the families showed significantly reduced CNP-dependent cGMP synthesis even in the heterozygous state.

PSMD1 (proteasome 26S subunit, non-ATPase, 1, MIM617842) encodes a regulatory subunit of the 26S proteasome, which is a major ATP-dependent intracellular proteinase. NCL (nucleolin, MIM164035) encodes a eukaryotic nucleolar phosphoprotein and is associated with the synthesis and maturation of ribosomes. No specific phenotypes associated with PSMD1 or NCL haploinsufficiency have been reported, whereas only one patient carrying a de novo NCL variant was identified by exome sequencing of 4293 families with developmental disorders19. Given that the deletion included four genes associated with autosomal recessive inheritance diseases, the major limitation of our study is that we did not analyze a hemizygous variation of the undeleted allele.

Table 1 Summary of CNV deletions including TRIP12 or NPPC and adjacent genes.

| Case   | Deletion range chr2: | Deletion size | Gene* number | Inheritance | Phenotype                                                                 | Reference                                                                 |
|--------|----------------------|---------------|--------------|-------------|---------------------------------------------------------------------------|---------------------------------------------------------------------------|
| Our case | 230,377,789-233,136,164 | 2.76 Mb       | 25           | de novo     | Developmental delay, severe short stature, motor delay, epilepsy, bilateral hearing loss, relative macrocephaly, no words and not walk 49 months | This article                                                                |
| Case-N1 | 231,264,599-233,178,325 | 1.91 Mb       | 19           | de novo     | Development delay, cutaneous syndactyly of 2nd and 3rd toes bilaterally, peripheral hearing loss, relative macrocephaly | Tassano et al.16 Patient 1                                                 |
| Case-T1 | 230,489,478/230,513,445-231,457,431/231,508,839 | 0.98 ± 0.04 Mb | 8            | de novo     | Developmental delay, normal height, microcephaly (25 percentile), obesity, unilateral hearing loss, Autism, first words 24 months, walk 20 months | Zhang et al.3 Subject 4                                                    |
| Case-T2 | 229,076,749/229,152,599-230,801,061/230,811,273 | 1.69 ± 0.04 Mb | 4            | de novo     | Developmental delay, normal height, Autism, aggressive behaviors, 5-6 words 22 months, motor delay | Zhang et al.3 Subject 5                                                    |
| Case-T3 | no data             | 144 Kb        | 2            | Unknown     | Developmental delay, motor delay, mild short stature (6 percentile)       | Zhang et al.3 Suppl 2                                                     |
| Case-T4 | 225,043,475/225,360,778b,230,758,568/230,841,358b | 5.61 ± 0.19 Mb | 20           | de novo     | Intrauterine growth retardation, developmental delay, walk 22 months, profound speech deficit, normal height(+1 SD), large mouth, multiple renal cyst | Doco-Fenzy et al.17                                                        |
| Case-T5 | 230,778,385-230,839,009b | 61 Kb          | 2            | de novo     | Autism, first words 12 months, first phrases 24 months, 5 years 9 months: global DQ 79 | Pinto et al.21                                                             |
| Case-T6 | 230,689,285-230,976,668 | 287 Kb         | 3            | de novo     | Mild micrognathia, global developmental delay                             | DECIPHER ID 301556                                                       |
| Case-T7 | 230,020,617-231,444,802 | 1.4 Mb         | 9            | de novo     | Epicanthus, low columella, wide mouth, delayed speech and language development | DECIPHER ID 250590                                                        |
| Case-T8 | 230,724,038-230,868,128 | 144 Kb         | 2            | Unknown     | Cystic renal dysplasia, Global developmental delay                        | DECIPHER ID 281305                                                        |

DQ developmental quotient.

*Protein coding genes.

Genome coordinates differ from the original articles, which were converted from assembly NCBI36/hg18 to assembly GRCh37/hg19 by using Assembly Converter (http://grch37.ensembl.org/Homo_sapiens/Tools/AssemblyConverter).
Eight CNV deletions smaller than 6 Mb involving TRIP12 have been reported. In addition, we identified five CNV deletions with phenotype information in the DECIPHER database v9.31. None of these 13 CNVs included NPPC, NCL, or PSMD1, and five CNVs were intragenic deletions within TRIP12. The remaining eight deletions involved adjacent genes and TRIP12 (case T1–T8 in Fig. 1e and Table 1). For NPPC, case N1 was the only reported patient with a CNV deletion smaller than 6 Mb involving NPPC, and the deletion included NCL and PSMD1 (case N1 in Fig. 1e and Table 1).

The clinical features associated with TRIP12 alterations have been previously summarized. Clinical features such as developmental delay, hearing loss, and epilepsy in our patients could be attributed to TRIP12 haploinsufficiency. Our patient showed epicanthic folds, hypertelorism, short nose, low hanging columella, and low-set ears, which partly overlapped with the dysmorphic facial features observed in patients with TRIP12 alterations. However, the low-set ears, prominent forehead, strabismus, short philtrum, everted vermilion of the upper lip, and relative macrocephaly in our patient were considerably similar to those in case N1. Case N1 had 19 deleted genes, including NPPC, NCL, and PSMD1, and all of them were included within the deletion range of our case. Recognizable facial dysmorphisms in patients with NPPC alterations have not been reported. Therefore, the characteristic facies in Case N1 and in our case is not explained by NPPC deletion but by other deleted genes, including NCL and PSMD1. Given the considerable facial similarity between Case 1 and our case, we speculated that facial dysmorphism in our case would be affected more by the 19 deleted genes than by TRIP12 alteration.

Our patient showed developmental delay and short stature to a more severe extent than previously described for patients with TRIP12 alterations. He was not able to speak any words or walk independently at 49 months of age, although previously described patients with TRIP12 alterations such as case T1 and case T2 could speak their first words and walk independently at a mean age of 28 months (range 10–60, n = 15) and 20 months (range 13–29, n = 16), respectively. Their heights were variable, and the mean height SD score was −0.3 SD (range −2.2 to +1.1, n = 16). Short stature and small hands in our patient were thought to be mainly affected by the NPPC deletion. However, NPPC alterations alone do not explain his extremely short stature. The mean height SD score of the six patients with NPPC heterozygous variants (−2.9 SD, range −4.3 to −2.3) was 0.9 SD lower than that of the five patients without NPPC variants (−2.0 SD, range −4.8 to −0.4) in the two reported families. Importantly, case T4, with the largest deletion expanding in the centromeric direction, showed a normal height and moderate developmental delay (initial walk at 22 months).

Thus, genes other than TRIP12 and NPPC located in the deletion region might have contributed to the exaggerated phenotypes in terms of developmental delay and short stature. Because of extreme intolerance to loss-of-function variations, NCL and PSMD1 could be good candidate genes for the exaggerated phenotypes.

In conclusion, we report a patient with a de novo 2.76-Mb deletion of 2q36.3q37.1 encompassing TRIP12 and NPPC. This is the first report of a patient with a unique phenotype of combined TRIP12 and NPPC haploinsufficiency. The severe developmental delay and extremely short stature of the patient imply that NCL and PSMD1 are potentially disease-modulating genes.

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Conflict of interest
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