Screening of known disease genes in congenital scoliosis

Kazuki Takeda1,2 | Ikuyo Kou1 | Shuji Mizumoto3,4 | Shuhei Yamada3 | Noriaki Kawakami5 | Masahiro Nakajima1 | Nao Otomo1,2 | Yoji Ogura1,2 | Noriko Miyake6 | Naomichi Matsumoto6 | Toshiaki Kotani7 | Hideki Sudo8 | Ikuho Yonezawa9 | Koki Uno10 | Hiroshi Taneichi11 | Kei Watanabe12 | Hideki Shigematsu13 | Ryo Sugawara14 | Yuki Taniguchi15 | Shohei Minami7 | Masaya Nakamura2 | Morio Matsumoto2 | Japan Early Onset Scoliosis Research Group | Kota Watanabe2 | Shiro Ikegawa1

1Laboratory of Bone and Joint Diseases, Center for Integrative Medical Sciences, RIKEN, Tokyo, Japan
2Department of Orthopaedic Surgery, Keio University School of Medicine, Tokyo, Japan
3Department of Pathobiochemistry, Faculty of Pharmacy, Meijo University, Nagoya, Japan
4Department of Women’s and Children’s Health, Dunedin School of Medicine, University of Otago, Dunedin, New Zealand
5Department of Orthopaedic Surgery, Meijo Hospital, Nagoya, Japan
6Department of Human Genetics, Yokohama City University Graduate School of Medicine, Yokohama, Japan
7Department of Orthopaedic Surgery, Seirei Sakura Citizen Hospital, Sakura, Japan
8Department of Advanced Medicine for Spine and Spinal Cord Disorders, Hokkaido University Graduate School of Medicine, Sapporo, Japan
9Department of Orthopaedic Surgery, Juntendo University School of Medicine, Tokyo, Japan
10Department of Orthopaedic Surgery, National Hospital Organization, Kobe Medical Center, Kobe, Japan
11Department of Orthopaedic Surgery, Dokkyo Medical University School of Medicine, Mibu, Japan
12Department of Orthopaedic Surgery, Niigata University Hospital, Niigata, Japan
13Department of Orthopaedic Surgery, Nara Medical University, Kashihara, Japan
14Department of Orthopedics, Jichi Medical University, Shimotsuke, Japan
15Department of Orthopaedic Surgery, Faculty of Medicine, The University of Tokyo, Tokyo, Japan

Correspondence
Shiro Ikegawa, Laboratory of Bone and Joint Diseases, Center for Integrative Medical Sciences, RIKEN, Minato-ku, Tokyo, Japan.
Email: sikegawa@ims.u-tokyo.ac.jp
Kota Watanabe, Department of Orthopaedic Surgery, Keio University School of Medicine, Shinjuku-ku, Tokyo, Japan.
Email: watakota@gmail.com

Abstract

Background: Congenital scoliosis (CS) is defined as a lateral curvature of the spine due to the vertebral malformations and has an incidence of 0.5–1/1,000 births. We previously examined TBX6 in Japanese CS patients and revealed that approximately 10% of CS was caused by TBX6 mutations. However, the genetic cause of remaining CS is unknown.

Methods: We recruited 78 CS patients without TBX6 mutations and major comorbidities, and investigated the genes previously reported to be associated with CS and congenital vertebral malformations by whole-exome sequencing.

The authors wish to be known that, in their opinion, the last two authors should be regarded as joint corresponding authors.

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1 INTRODUCTION

Congenital scoliosis (CS) is defined as a lateral curvature of the spine due to vertebral malformations; its incidence is 0.5–1/1,000 births worldwide (Giampietro, 2012; Giampietro et al., 2013; Liljenqvist, 2004; McMaster & Ohtsuka, 1982). Segmentation of the vertebrae occurs between 20 and 35 days after conception in human embryonic development (Turnpenny et al., 2007). Somites, the precursors to the ribs, striated muscle, and dermis of the back, are also formed during this period. This process is called somitogenesis. Disruptive perturbation of somitogenesis contributes to congenital vertebral malformations leading to CS. CS is well known to be a heterogeneous disorder ranging from nonsyndromic to possibly syndromic CS who has various major comorbidities (Turnpenny et al., 2007).

Perturbation of somitogenesis causes not only vertebral malformations but also rib malformations. Therefore, many CS patients also have rib malformations. Such a severe form of CS associated with rib malformations is referred to as spondylocostal dysostosis (SCD) (Bulman et al., 2000; Sporrong et al., 2015; Sparrow, Guillem-Navarro, Fatkin, & Dunwoodie, 2008; Whittock et al., 2004). While CS is the view from the orthopedic aspect of the disorder, SCD considers developmental abnormalities of the skeleton (Supporting Information Figure S1). Therefore, some SCD have only mild scoliosis due to the balanced abnormality of the spine even though they have severe vertebral abnormalities. SCD is included in the nosology and classification of genetic skeletal disorder: 2015 revision as the group 35, “Dysostoses with predominant vertebral with and without costal involvement” (Bonafe et al., 2015). Severe spine and rib malformations of CS and SCD usually affect the thoracic growth and function adversely, and lead to thoracic insufficiency syndrome (Campbell et al., 2003; Flynn et al., 2013). Early diagnosis and intervention are mandatory to keep the deformity under control.

RESULTS

We identified the compound heterozygous missense variants in LFNG in one patient. No likely disease-causing variants were identified in other patients, however. LFNG encodes a GlcNAc-transferase. The LFNG variants showed loss of their enzyme function.

CONCLUSIONS

A LFNG mutation is reported in a case of spondylocostal dysostosis (SCD), a skeletal dysplasia with severe malformations of vertebra and rib. The CS patient with LFNG mutations had multiple vertebral malformations including hemivertebrae, butterfly vertebrae, and block vertebrae, and rib malformations. LFNG mutations may cause a spectrum of phenotypes including CS and SCD. The current list of known disease genes could explain only a small fraction of genetic cause of CS.

KEYWORDS

congenital scoliosis, LFNG, spondylocostal dysostosis, whole-exome sequencing

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Perturbation of somitogenesis resulting in congenital vertebral malformations is etiologically heterogeneous (Giampietro et al., 2013). Several genes associated with congenital vertebral malformations have been identified in human subsequent to identification of model organisms; the examples include PAX1 (MIM# 167411), SLC35A3 (MIM# 605632), and T (MIM# 601397) (Ghebranious et al., 2008; Giampietro et al., 2005; Thomsen et al., 2006). Whole-exome sequencing (WES) has identified DYNC1H1 (MIM# 600112) in a patient with CS and spinal atrophy with lower extremity predominance (Punetha et al., 2015). Notch signaling pathway genes, Dll3 (MIM# 602768), Mesp2 (MIM# 605195), LFNG (MIM# 602576), Hes7 (MIM# 608059), Ripply2 (MIM# 609891), and Tbx6 (MIM# 602427), are associated with somitogenesis and their mutations have been identified in SCD (Bulman et al., 2000; Giampietro et al., 2006; McInerney-Leo et al., 2015; Sparrow et al., 2006, 2008, 2013; Summer et al., 2004). It was reported that compound heterozygosity for null mutations and the common hypomorphic risk haplotype composed by three SNPs in Tbx6 caused CS (Wu et al., 2015). Recently, several reports suggested that Tbx6-associated CS and SCD may represent a spectrum of a disease caused by the compound heterozygosity model (Lefebvre et al., 2017; Takeda et al., 2017; Wu et al., 2015).

We previously examined Tbx6 in undiagnosed Japanese CS patients and found that approximately 10% of CS was caused by Tbx6 mutations (Takeda et al., 2017). The incidence of Tbx6-associated CS was similar worldwide (Lefebvre et al., 2017; Takeda et al., 2017; Wu et al., 2015). However, the genetic cause of CS other than Tbx6 remains unknown. Therefore, we investigated the mutations in genes associated with the development of spine in Japanese CS by WES. We identified the compound heterozygous missense mutations in LFNG in one CS patient. These mutations in LFNG lead to loss of its enzyme function.
2 MATERIALS AND METHODS

2.1 Subjects

We recruited 196 Japanese CS patients who were seen at participating hospitals. They received clinical and radiological examinations by expert spinal surgeons and pediatricians. Patients who have known syndromic scoliosis, including Alagille syndrome, Goldenhar’s syndrome, Jarcho-Levin syndrome, Klippel-Feil syndrome, SOTOS syndrome, and VACTERL association, were excluded. Genomic DNA was extracted from peripheral blood leukocytes using standard methods or from saliva using Oragene DNA collection kit (DNA Genotek, Ottawa) according to the manufacturer’s protocol. Informed consent was obtained from the patients and their parents. The study was approved by the ethical committee of all participating hospitals and RIKEN.

2.2 WES

TBX6 mutation was examined as previously described (Takeda et al., 2017). We conducted WES as previously described (Guo et al., 2017; Wang et al., 2017). Briefly, approximately 3 μg DNA was sheared and used for a SureSelect Human All Exon V5 library (Agilent Technologies, Santa Clara, CA) according to the manufacturer’s instructions. Patients’ DNA samples were sequenced on a HiSeq2000 (Illumina, San Diego, CA) with 101-bp paired-end reads.

2.3 Analysis of WES data

Although all patients were sporadic, the mode of inheritance in CS remains still unclear. So, we considered both autosomal dominant and autosomal recessive inheritance. We excluded variants within exons ±30 bp from exon–intron junction that minor allele frequency was above 0.03 in dbSNP137, the National Heart Lung and Blood Institute Exome Sequencing Project Exome Variant Server (NHBLI-ESP 6500; https://evs.gs.washington.edu/EVS/), Exome Aggregation Consortium (ExAC; https://exac.broadinstitute.org/), Integrative Japanese Genome Variation Database (iJGVD; https://ijgvd.megabank.tohoku.ac.jp/), and our in-house database from 575 Japanese individuals since the incidence of CS is 0.5–1/1,000 births and SCD is more rare disease than CS worldwide. We conducted the pathogenicity prediction using SIFT (https://sift.bii.a-star.edu.sg/) (Adzhubei et al., 2010), Polyphen-2 (https://genetics.bwh.harvard.edu/pph2/) (Kumar, Henikoff, & Ng, 2009), and MutationTaster (https://www.mutationtaster.org/) (Schwarz, Cooper, Schuelke, & Seelow, 2014). We defined a variant as “deleterious” when it was predicted as damaging in the three prediction tools. We investigated the variants satisfying the above-mentioned conditions in previously reported CS and SCD genes: PAX1, SLC35A3, T, DYNC1H1, DLL3, LFN, HES7, DLL1, MESP2, and RIPPL2.

2.4 DNA sequencing of LFNG

The genomic region of the LFNG gene including two missense variants (c.467T>G and c.856C>T) was PCR-amplified using KOD Fx and primers 5′-CCAGTGCGGCC- GCCTGGTGCTGTTGCTAATAAGGTT-3′ (forward) and 5′-CCAGTGAAATTCATACCCGTAGCTACGCTACCCTG-3′ (reverse). The PCR amplicons were cloned into the NotI and EcoRI sites of the pBluescript SK(-) cloning vector (Agilent Technologies). After ligation, the vectors and insert DNA fragments were transformed to E.coli competent cells. The bacterial colonies were collected and verified by Sanger sequencing. LFNG variants were detected by direct sequencing of PCR products using KOD Fx (Toyobo, Tokyo) and primer sets using a 3730xl DNA analyzer (Applied Biosystems, Foster City, CA). Nucleotide numbering uses +1 as the A of the ATG translation initiation codon in the reference sequence, with the initiation codon as codon 1.

2.5 Expression of the soluble form of the LFNG proteins and enzyme assay of their GlcNAc activity

The expression vector of human LFNG was constructed as described previously with slight modifications (Rampal et al., 2005). A truncated form of LFNG (wild-type) was amplified from human placenta cDNA (BioChain, Newark, CA) by two-round PCR. The first PCR was performed with a forward primer, 5′-GCTGCT CGCCTGCGCTG CTGTTGCTCA-3′, and a reverse primer, 5′-CTCCA- CAGCA GAACGTCCCAGCAGCCT-3′, followed by nested PCR with nested primers: a forward primer containing an in-frame EcoRI site located 259-bp downstream from the initial codon, 5′-CCGGAAATTCA-GATGCCGGCCCGCCGCCGC-3′, and a reverse primer containing a KpnI site located 3-bp downstream from the stop codon, 5′-CGGGGTACCGATGGGTCTCGACGTC-CTGCCC-3′. Each PCR was conducted with KOD-Plus DNA polymerase (Toyobo). A DNA fragment, which encoded the human LFNG lacking the first 86 amino acids including the predicted transmembrane domain, was subcloned into a p3xFLAG-CMV8 (Sigma, St. Louis, MO) vector, resulting in the fusion of LFNG (wild-type) to the preprotrypsin leader sequence and the 3xFLAG tag sequence at the N-terminus present in the vector.

3 Site-specific mutagenesis of LFNG to produce the p.Leu156Arg and p.Arg286Trp mutants was performed using two rounds of PCR. The first PCR was performed
with a 5′-primer containing an in-frame EcoRI site described above and a 3′-internal mutagenic oligonucleotide primers (5′-GGCCGTTGCTCTGCGCCCG GCCCT CATTTTTCCC-3′ or 5′-GGGATCAGTGCAAGGCCAGA TCCGCTC AGCCGT-3′) for p.Leu156Arg or p.Arg286Trp, respectively, or 5′-internal mutagenic oligonucleotide primer (5′-GGGAAAGATGAGGCCGGCCAGGCACAG GC-GCC′ or 5′-ACGGCTAGCAGTCTGGCTGCTT GATGACTG-3′) for p.Leu156Arg or p.Arg286Trp, respectively) and 3′-primer containing an EcoRV site described above, and p3xFLAG-CMV8/LFNG (wild-type) as a template. The second PCR was performed with a 5′- and 3′-primers containing a HindIII and an EcoRV sites described above and KOD Plus polymerase, and the first PCR products as a template. The amplified fragments were digested with EcoRI and KpnI, inserted into p3xFLAG-CMV8, and sequenced using a 3730xl DNA Analyzer.

The expression plasmid was transfected into HEK293T cells on a T-75 flask using FuGENE 6 HD (Promega, Madison, WI). Three days after transfection, 3 ml each of the culture medium and the cells were collected. The cells were lysed with phosphate-buffered saline containing 0.5% TritonX-100 and protease inhibitor cocktail (Roche, Basel). Those conditioned media and cell lysates were incubated with 10 µl of anti-FLAG M2 agarose resin (Sigma) for 2 hr and overnight, respectively, at 4°C. The beads were washed with 1 ml of 50 mM Tris-HCl, pH 7.5 containing 150 mM NaCl, 0.02% Tween-20, and then analyzed on a 10% sodium dodecyl sulfate (SDS)-polyacrylamide gel, transferred to a nitrocellulose membrane (BioRad), and incubated for 1 hr with anti-FLAG M2 antibody (Sigma). The bound antibody was detected with anti-mouse IgG conjugated with a fluorescent dye, IRDye 800IRD (Li-Cor, Lincoln, NA) using ODYSSEY CLx (Li-Cor). The amount of recombinant LFNG protein was estimated by standard curve from fluorescent intensity of 3xFLAG-tagged bovine alkaline phosphatase (Sigma) using Image Studio (ver 4.0, Li-Cor).

GlcNAc-transferase activity was examined as described previously (Rampal et al., 2005). Briefly, the GlcNAc-transferase assay mixture contained 10 µl of enzyme-bound anti-FLAG affinity resins, 50 mM 2-(N-morpholino) ethanesulfonic acid-NaOH (pH 6.5), 10 mM MnCl2, 0.1 mM UDP-GlcNAc (ultrapure grade, Promega) as the sugar donor substrate, and 1 mM p-nitrophenyl-α-L-fucose (pNP-Fuc, Sigma) as the sugar acceptor in a total volume of 50 µl. The reaction mixture was incubated at 37°C for 2–4 hr. The reaction product, UDP moiety released from UDP-GlcNAc, was mixed with UDP detection reagent, which contains an enzyme converting UDP to ATP, in UDP-Glo™ Glycosyltransferase Assay kit (Promega). The newly synthesized ATP to be measured using a luciferase/luciferin reaction and the luminescent signals were detected using a luminometer, Victor X4 (PerkinElmer, Waltham, MA).

3 | RESULTS

3.1 | Clinical features of the CS patients

We found TBX6 mutations in 15 out of 196 CS patients. From the patients without TBX6 mutations, we selected 78 nonsyndromic CS patients (Supporting Information Table S1) and examined them by using WES. Their mean age was 12.0 ± 5.4 years old. Ten patients had family histories of scoliosis; two had CS in their families. The mean Cobb angle of the main curve was 43.0° ± 13.4°. Eight patients had various comorbidities. Various types of rib malformations were found in 23 patients; one irregular shaped, 11 fused, six missing, and five additional ribs.

3.2 | WES

The mean depth of coverage for reads was 74.3x, and 94.3% of the targeted bases had more than 20 reads on average (Supporting Information Table S2). By evaluation of the variants for their pathogenicity, we identified a CS patient (S1289) who had two likely disease-causing variants in LFNG: c.467T>G (p.Leu156Arg) and c.856C>T (p.Arg286Trp) (Table 1). The sequences of the variants were confirmed by Sanger sequencing (Figure 1a,b). We did not identify any likely disease-causing variants in other genes (PAX1, SLC35A3, T, DynC1H1, DLL3, HES7, DLL1, MESP2, and Ripply2). The coverages of the exons of these genes which had more than 10 reads on average were over 90% (Supporting Information Table S3).

3.3 | Evaluation of LFNG variants

c.467T>G and c.856C>T in LFNG had been annotated in ExAC and iJGVD; allele frequency of c.467T>G was 3.0 × 10−4 in iJGVD and that of c.856C>T was 2.5 × 10−5 in ExAC. Both variants were predicted to be disease causing by SIFT, Polyphen-2, and MutationTaster (Table 1). Previously, only one LFNG mutation (c.564C>A; p.Phe188Leu) has been reported in a SCD patient.

Because the parents’ DNAs were unavailable, we examined the haplotype of the two missense variants in S1289 by a long-range PCR followed by sequencing the subcloned amplicons. We found that S1289 was compound heterozygotes for the two missense variants.

3.4 | Functional analysis of LFNG variants in vitro

To examine whether the LFNG missense variants, p.Leu156Arg and p.Arg286Trp cause a reduction of the GlcNAc-transferase activity, the LFNG proteins in soluble
forms with a 3xFLAG epitope were generated by replacing the first 86 amino acids of LFNG with a cleavable pre-trypsin signal sequence. The soluble 3xFLAG-tagged LFNG proteins were expressed in HEK293T cells. The enzyme activity for wild-type LFNG (Supporting Information Figure S2b). These data indicate that the two LFNG variants lead to loss of the enzyme function. By contrast, the expression of wild-type and a mutant enzyme, p.Arg286Trp-LFNG, but not p.Leu156Arg-LFNG, was detected in the conditioned medium (Supporting Information Figure S2a). These observations suggest that the secretion pathway of p.Leu156Arg-LFNG might be impaired. The GlcNAc-transferase activity of p.Arg286Trp-LFNG from the conditioned medium was also drastically decreased compared to that of the wild-type LFNG (Supporting Information Figure S2b).

### 3.5 Clinical features of the patient with LFNG mutations

S1289 was a 16 years old male who was born as a child of nonconsanguineous healthy parents. He presented with mild short stature (−2.1 SD) and had no comorbidity (Supporting Information Table S1). He had severe scoliosis and was diagnosed with CS. The main Cobb angle was 70° at T9-L3 (Figure 3a). He had multiple vertebral malformations: hemivertebrae at T5, L1, and L4 (one segmented type and two nonsegmented type), block vertebra at T8-T9, and butterfly vertebrae (nonsegmented type) at T7, T12, and L5 (Figure 3b). Multiple rib malformations, including hypoplasia of bilateral 2nd ribs and left 6–7th, 8–9th, right 5–6th, 7–8th, and 9–10th fused ribs, were also found.

### 4 DISCUSSION

Our WES identified a CS patient who is compound heterozygous for two novel missense variants of LFNG. Only one LFNG mutation (c.564C>A, p.Phe188Leu) has previously been reported, which causes SCD3 (MIM# 609813) in the homozygous state (Sparrow et al., 2006). Both of the variants we found were rare and were predicted deleterious in all bioinformatic tools examined. LFNG encodes an O-fucosylpeptide 3-β-N-acetylgalcosaminyltransferase that adds N-acetylgalcosamine (GlcNAc) residues to O-fucose on the EGF-like repeats of Notch receptors (Bruckner, Perez, Clau sen, & Cohen, 2000; Moloney et al., 2000). Our variants and the previously reported mutation are all located in highly conserved region that encodes a glucosaminyl-transferase; particularly, the Arg286 residue where we found the mutation was conserved in all known fringe proteins from Drosophila melanogaster to human (Correia et al., 2003). Although these in silico evaluation data strongly suggested that the two variants are disease-causing mutations, it has been reported that to predict the deleterious effect of missense variants in silico is sometimes difficult (Majithia et al., 2016; Takeda et al., 2017). Therefore, we examined the effect of the variants in vitro and revealed that both lead to
loss of the GlcNAc-transferase activity. Based on these observations, the two LFNG variants are loss of function mutations and hence are pathogenic to the CS phenotype. p.Leu156Arg was suggested to cause mislocalization of the mutant protein in addition to its loss of enzymic activity (Supporting Information Figure S2a). It is reported that p.Phe188Leu mutation was also mislocalized and enzymatically inactive by a functional assay (Sparrow et al., 2006).

**FIGURE 2** Protein level and GlcNAc-transferase activity of recombinant LFNG expressed in HEK293T cells. (a) Western blot analysis of the recombinant LFNG proteins: wild-type, L156R (p.Leu156Arg), and R286W (p.Arg286Trp). The purified recombinant LFNG from cell lysate was detected with the anti-FLAG and fluorescence-conjugated anti-mouse IgG antibodies. Asterisks indicate heavy and light chains of the anti-FLAG antibody from the anti-FLAG agarose resin for purification. (b) GlcNAc-transferase activity of the recombinant LFNG in the cell lysate. Both L156R and R286W showed significantly decreased enzyme activities. Values are the means ± SE (n = 3). *p < 0.0001 versus the wild-type was calculated by the Student's t test.

**FIGURE 3** Radiographic phenotype of the congenital scoliosis patient with LFNG mutations. (a) The standing spinal antero-posterior radiograph. The patient has severe scoliosis with a Cobb angle of 70° at T9-L3. (b) Three-dimensional CT (left: front view, right: back view). Multiple congenital vertebral malformations including hemivertebrae, block vertebrae, and butterfly vertebrae are noted. Multiple rib malformations including hypoplasia and fused ribs are also found.
The CS patient who carried the LFNG mutations (c.467T>G; p.Leu156Arg and c.856C>T; p.Arg286Trp) had severe scoliosis due to interspersed vertebral malformations from thoracic to lumbar spine together with multiple rib malformations, while the SCD patient who carried the LFNG mutation (c.564C>A; p.Phe188Leu) had nonprogressive scoliosis but more extensive and a large number of congenital vertebral malformations like “pebble beach” from cervical to lumbar spine as well as severe short trunk and finger abnormalities (Sparrow et al., 2006; Turnpenny et al., 2003). Thus, LFNG mutations are considered to cause a spectrum of vertebral malformation recognized as CS and/or SCD (Supporting Information Figure S1). Such phenotypic variation including CS and SCD is also known in vertebral malformation caused by TBX6 mutations (Lefebvre et al., 2017; Takeda et al., 2017). Further, accumulation of the patients with LFNG mutations is mandatory to conclude the phenotypic range of the LFNG mutation and genotype–phenotype association.

In the screenings of SCD genes, known gene mutations have been found in 20%–25% of the patients examined; 60% of SCD is caused by DLL3 mutations (Bonafe, Giunta, Gassner, Steinmann, & Superti-Furga, 2003; Turnpenny et al., 2003; Turnpenny, Sloman, & Dunwoodie, 1993; Whittock et al., 2004). In contrast, our screening of known disease genes for CS and vertebral malformation could only find the causal genes in <10% of nonsyndromic CS patients who had no major comorbidities; CS in our cohort did not have the mutations in known genes causing CS and SCD other than TBX6 and LNFG. CS is a very heterogeneous disorder; the type, location, number of vertebral malformations, and comorbidities are different between the patients (Giampietro et al., 2013). On the other hand, CS caused by the TBX6 mutation has been reported to have specific abnormality of the vertebra (Takeda et al., 2017; Wu et al., 2015). The vertebral anomaly along the whole length of the spine with the vertebrae of rounded shape (“pebble beach” sign) is a feature of SCD1 (MIM# 277300), which is caused by DLL3 mutations (Turnpenny et al., 2003). Their clinical differences suggest considerable genetic heterogeneity of CS. In the current study, we investigated the limited number of genes that are considered to be associated with the development of spine. Most of them are related to the Notch signal. Somitogenesis of vertebrate species relies on the intersecting gradients and cross-regulatory activities of various signal transduction pathways, not only Notch signaling but also Fgf, Wnt, and retinoic acid signaling pathways (Gibb, Maroto, & Dale, 2010; Wahi, Bochter, & Cole, 2016). It was reported that mutations in Wnt3a identified as a negative regulator on Notch signaling and somitogenesis associated with congenital vertebral malformations (Aulehla et al., 2003; Hayward, Kalmar, & Arias, 2008). Therefore, further comprehensive approach might identify the causal genes for CS. However, our preliminary screening of unknown CS gene(s) for the exome data did not present a possible causal gene that had likely deleterious mutation in more than two cases. To discover the unknown causal genes for the rare diseases by WES is still challenging and tends to come up against the N = 1 problem (Akawi et al., 2015). It would be mandatory to increase the sample size and evaluate the phenotype well.

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CONFLICT OF INTEREST

The authors declare no conflict of interest associated with this manuscript.

ORCID

Kazuki Takeda http://orcid.org/0000-0003-3857-4985

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.

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