Original Article

Mutations in the NOG gene are commonly found in congenital stapes ankylosis with symphalangism, but not in otosclerosis

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Human noggin (NOG) is a responsible gene for multiple synostosis syndrome (SYNS1) and proximal symphalangism (SYM1), two conditions that are recently known to be within a wider range of clinical manifestations of stapes ankylosis with symphalangism. This study was performed to determine the range of phenotype caused by NOG mutations, using Japanese patients with various phenotypes including sporadic inherited SYM1, dominantly inherited SYM1, stapes ankylosis with broad thumb and toes (Teunissen and Cremer syndrome). In addition, 33 patients with typical otosclerosis (without symphalangism) were studied. Direct sequencing analysis disclosed three novel mutations of the NOG gene in three SYM1 families. None of the otosclerosis patients without symphalangism had NOG mutations, indicating that NOG mutations may be restrictively found within patients with various skeletal abnormalities. These results together with the literature review indicated that there are no clear genotype-phenotype correlations for NOG mutations. With regard to surgical outcome, most of the patients in these three families with NOG mutations showed remarkable air-bone gap recovery after stapes surgery. Molecular genetic testing is useful to differentiate syndromic stapes ankylosis from otosclerosis, and even mild skeletal anomalies can be a diagnostic indicator of NOG-associated disease.

Conflict of Interest

The authors declare no conflict of interest.

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Human noggin (NOG) is a responsible gene for a wide range of clinical manifestations of stapes ankylosis with symphalangism. Proximal symphalangism (SYM1: MIM #185800) (1) is known as an autosomal dominant disorder with high penetrance. The most common clinical features are the immobility of the proximal interphalangia (PIP) joints of the hands and toes, and congenital conductive hearing loss due to stapes ankylosis. Multiple synostosis syndrome (SYNS1: MIM #186500) (1) is characterized by a more severe phenotype of the proximal symphalangism, such as progressive and expanded bony fusion of joints and unique facial manifestations. In addition, mutations in NOG have been identified in Tarsal–Carpal Coalition syndrome (TCC: MIM #186570) (2), stapes ankylosis with broad thumb and toes (MIM #184460) (Teunissen and Cremer syndrome) (3), and Brachydactyly type B2 (BDB2: MIM #611377) (4).

Otosclerosis (MIM #166800) is known as the single most common cause of progressive conductive hearing loss, characterized by abnormal bone remodeling in the otic capsule. Although there are a small number of familial cases that are likely to be monogenic, the majority of cases are sporadic. A series of studies has
suggested that this condition involves both genetic and environmental factors (5).

Typical otosclerosis was included in this study because it is an interesting question as to whether some of the typical otosclerosis is a continuum of a category of disease caused by NOG mutations. We thought this may be true because (i) within SYM1, NOG mutations were found in patients with minor skeletal anomalies without symphalangism (3), and (ii) stapes ankylosis is an important phenotype of the animal model for NOG+/− mice (6).

To date, no detailed survey was available for NOG mutations in the stapes ankylosis patients with symphalangism in Asian populations. Therefore this study was undertaken to address whether NOG mutations are also causative and commonly found in those populations and if so, whether there is a different mutation spectrum.

In addition, previously reported NOG mutations were reviewed to determine their spectrum as well as whether there are any particular genotype–phenotype correlations caused by NOG mutations.

Materials and methods

Subjects

We ascertained three Japanese families to be associated with conductive hearing loss and symphalangism, including an autosomal dominant SYM1 family, a sporadic SYM1 case with normal parents, and an autosomal dominant stapes ankylosis with broad thumb and toes (Teunissen and Cremer syndrome) family. Thirty-three Japanese otosclerosis patients, who underwent stapes surgery, were also screened for mutations in the NOG gene. Their clinical symptoms, including ages at surgery (36–77 years old: average 54.4 years old), onset age (15–57 years old: average 37.3 years old), gender (10 male and 23 female), laterality (9 unilateral and 24 bilateral), and hearing threshold (average 63.1 dB), are summarized in Table 1. Average onset age, was hearing threshold, was evaluated using pure-tone audiometry classified by a pure-tone average over 250, 500, 1000, 2000, and 4000 Hz. By detailed anamnestic and medical examination, no patients had any associated skeletal abnormalities. All of the patients were sporadic cases and no similar condition was observed within their familial members. Satisfactory outcomes after stapes surgery were obtained in all 33 subjects.

We obtained informed consent for participation in this project from each subject and also from 192 normal control subjects who were unrelated Japanese individuals without any noticeable hearing loss evaluated by auditory testing. Oto logic examination, audiometric analysis, and radiologic imaging were carried out for each patient.

Family 1

As shown in the pedigree (Fig. S1a), patient #991 was diagnosed with symmetric conductive hearing loss of 50 dB (Fig. 1b) at the age of 6 years. Tympanography indicated type A sclerosis and absence of the stapedius reflex, whereas otomicroscopy results were normal. Temporal bone computed tomography (CT) scan revealed no inner or middle ear malformations. Her hearing level was stable and non-progressive, and she received hearing aids in both ears. At the age of 17, exploratory tympanotomy of the left ear showed bony fixation of the footplate without any other deformities in the middle ear and stapedotomy using a Teflon piston and wire was performed, resulting in a remarkable improvement in hearing. One year later, stapedotomy was undertaken in her right ear also. After the surgery, the postoperative hearing levels showed 20–30 dB and she did not use her hearing aids. The X-ray presented in Fig. 1 shows symphalangism in the PIP joints of the second to fifth fingers of both hands and in the distal interphalangeal (DIP) joints of the left second and fifth fingers and of the right fifth finger. There was symphalangism in both hands, resulting in limited mobility of the fingers. Symphalangism (fixation of the proximal interphalangeal joint) in both feet was also found.
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Fig. 1. (a) Photograph with arrowheads indicating symphalangism in the hands of patient #991. (b) Audiograms from patient #991 showing conductive hearing loss.

Fig. 2. (a) Photograph with arrowheads indicating symphalangism in the hands of patients #3925 and #3926. (b) Audiograms from patients #3925 and #3926 showing conductive hearing loss.

The ankylosis was confirmed by X-ray examination (Fig. 1a). Congenital hyperopia (only in this patient within the family) was also present.

Family 2

As shown in the pedigree (Fig. S2a), a girl and her father (patients #3925 and #3926) visited our hospital due to bilateral hearing loss. Audiograms indicated bilateral mixed hearing loss (Fig. 2b). Anamnestic evaluation suggested that the hearing loss was non-progressive without any associated symptoms such as ear fullness, tinnitus or vertigo. Patient #3926 underwent stapedotomy at the age of 42, achieving significant recovery of his hearing. There was symphalangism in the PIP joint of both fifth fingers, resulting in limited mobility of the fingers. Fixation of the proximal interphalangeal joint was not found in either foot.
Stapes ankylosis with NOG mutation

The ankylosis was confirmed by X-ray examination (Fig. 2a). Congenital hyperopia was not present in this family.

Family 3

The pedigree shows hearing loss was inherited in four generations, indicating autosomal dominant inheritance (Fig. S3a). In addition to conductive hearing loss (Fig. 3b), the family members were associated with the following clinical phenotype in various degrees: hyperopia, broad thumbs and first toes, symphalangism, syndactyly, and fused cervical vertebrae (Fig. S3a). The clinical diagnosis was therefore stapes ankylosis with broad thumb and toes (3) or Teunissen and Cremer syndrome (7). Patient #4351 had conductive hearing loss, hyperopia, broad thumbs and first toes, symphalangism, and syndactyly. She noted her hearing loss around age 10. Stapedotomy was performed when she was 37 (right) and 38 (left) years old, achieving significant recovery of hearing. Patient #4106 had conductive hearing loss, hyperopia, broad thumbs and first toes, and fused cervical vertebrae. His hearing loss was noted around age 3 and was diagnosed at the age of 8. Stapedotomy was performed when he was 9 (right) and 10 (left) years old, achieving significant recovery of hearing.

Mutation identification

Human NOG gene coding is constituted of one single exon, in which an open reading frame of 696 nucleotides encodes a NOG polypeptide of 232 amino acids. A sequence obtained from GeneBank U31202 was used to design primers containing the entire coding region of NOG. Two fragments to entirely cover the coding region of NOG were amplified with polymerase chain reaction (PCR) and two specific primer pairs, as follows: F1, 5'-CTTGTGCTTCTCCTCCGC-3'; R1, 5'-TACTGGATGGGAATCCAGCC-3'; and F2, 5'-TACGACCCAGCTTATGAGC-3'; and R2, 5'-TAGCAGCGACCTTCGACT-3'.

PCR reactions were carried out in 25 μl total volume containing 40 ng of genomic DNA, 10 pmol of each primer, 2 mM dNTPs, ×10 PCR buffer and 0.2 U of ExTaq polymerase (Takara, Tokyo, Japan). PCR conditions were denaturing at 94°C for 2 min; 35 cycles at 96°C for 30 s, 60°C for 30 s, 72°C for 1 min extension, with a final extension step at 72°C for 5 min in a Perkin-Elmer 9600 thermal cycler (Perkin-Elmer, Foster City, CA). PCR products were purified with a Suprec filter (Takara) and sequenced directly, using four primers (F1, R1, F2, and R2) and ABI BigDye terminators, on an ABI 3100 sequencer (Perkin-Elmer).

Results

Three novel mutations of the NOG gene were found by direct sequencing analysis in three families, whose common clinical features were compatible with SYM1, i.e. immobility of the PIP joints of the hands and toes, and congenital conductive hearing loss due to stapes ankylosis. Patient #991 of family 1 had a heterozygous G>T transversion at nucleotide 551 (Fig. S1b), predicting a cysteine (C) for phenylalanine (F) substitution at amino acid 184 (C184F) in the coding region of NOG. Since the C184F mutation was not found in either parent and was found only in the proband (patient #991), it was suggested that the mutation arose de novo in only the affected individual. Patients #3925 and #3926 of family 2 had a heterozygous T>A transversion at nucleotide 463 (Fig. S2b), predicting a cysteine (C) for serine (S) substitution at amino acid 155 (C155F) in the coding region of NOG. Since the C184F mutation was not found in either parent and was found only in the proband (patient #991), it was suggested that the mutation arose de novo in only the affected individual. Patients #3925 and #3926 of family 2 had a heterozygous G>T transversion at nucleotide 463 (Fig. S2b), predicting a cysteine (C) for serine (S) substitution at amino acid 155 (C155F) in the coding region of NOG. Patients #4106 and #4351 had a heterozygous C215X mutation.

None of the otosclerosis patients had NOG mutations. These three mutations were not observed in any of the other family members nor in the 192 unrelated Japanese controls (384 chromosomes).

Discussion

This study identified three novel mutations in the NOG gene in families with symphalangism, being consistent with the previous work showing that NOG is the responsible gene for SYM1 and stapes ankylosis with
Table 2. NOG mutations reported in SYM1, SYNS1, TCC, BDB2 and TCS families

| Nucleotide change | Amino acid | Family information | Phenotype | Evolutionary conservation + | Domain/structure/motif ++ | Authors |
|-------------------|------------|--------------------|-----------|-----------------------------|---------------------------|---------|
| c. 53delC         | Frameshift | Japanese, AD       | SYNS1     | —                           | Finger/clamp region interface of NOG and BMP? | Takahashi (8) |
| c. 103C>G         | p. R33A    | German, AD         | BDB2      | Conserved                   | Finger/clamp region interface of NOG and BMP? | Lehmann (4) |
| c. 103C>T         | p. R33S    | Turkish, AD        | BDB2      | Conserved                   | Finger/clamp region interface of NOG and BMP? | Hirschorn (9) |
| c. 103C>T         | p. R33S    | Italian, AD        | SYM1      | Conserved                   | Finger/clamp region interface of NOG and BMP? | Margino (10) |
| c. 104C>T         | p. R34R    | NI, sporadic       | SYM1      | Conserved                   | Finger/clamp region interface of NOG and BMP? | Leiden (8) |
| c. 106G>C         | p. G35E    | Danish, AD         | BDB2      | Almost conserved            | Finger/clamp region interface of NOG and BMP? | Debeer (11) |
| c. 110C>T         | p. R37R    | Belgian, AD        | TCC       | Conserved                   | Finger/clamp region interface of NOG and BMP? | Debeer (12) |
| c. 124C>G         | p. A42A    | Belgian, de novo   | TCC       | Conserved                   | Finger/clamp region interface of NOG and BMP? | Debeer (12) |
| c. 125C>G         | p. A42R    | NI, AD             | SYNS1     | Conserved                   | Finger/clamp region interface of NOG and BMP? | Oxley (13) |
| c. 129-130dup     | Frameshift | Dutch, AD          | SABTT     | —                           | Finger/clamp region interface of NOG and BMP? | Weikamp (14) |
| c. 142G>A         | p. E48K    | Japanese, sporadic | SYM1      | Conserved                   | Finger/clamp region interface of NOG and BMP? | Kassaki (19) |
| c. 142G>A         | p. E48K    | Iranian, AD        | BDB2      | Conserved                   | Finger/clamp region interface of NOG and BMP? | Leiden (8) |
| c. 149C>G         | p. K42R    | Belgian, de novo   | TCC       | Conserved                   | Finger/clamp region interface of NOG and BMP? | Debeer (12) |
| c. 252-253insC    | Frameshift | Dutch, AD          | SABTT     | —                           | —                           | Brown (3) |
| c. 30delG         | Frameshift | Dutch               | SYM1      | —                           | —                           | Thomeer (16) |
| c. 328C>T         | p. G110X   | Italian, AD        | SABTT     | Conserved                   | —                           | Brown (3) |
| c. 38T>A          | p. L130X   | Japanese, AD       | SYM1      | Almost conserved            | —                           | Takahashi (8) |
| c. 591C>T         | p. Q171X   | Dutch               | SYM1      | Almost conserved            | —                           | Thomeer (16) |
| c. 481T>A         | p. G175S   | Japanese, AD       | SYM1      | Conserved                   | —                           | Leiden (8) |
| c. 499C>G         | p. R175G   | North American, sporadic | BDB2     | Conserved                   | —                           | Present study |
| c. 551G>A         | p. C184Y   | Japanese, sporadic | SYM1      | Conserved                   | —                           | Present study |
| c. 559C>T         | p. P175S   | British, AD        | BDB2      | Conserved                   | —                           | Leiden (8) |
| c. 561T>C         | p. I183C   | Belgian, AD        | SYNS1     | Conserved                   | —                           | Oxley (13) |
| c. 608T>C         | p. L203P   | Dutch, AD          | SABTT     | Conserved                   | β-Sheet 3 of NOG structure | Weikamp (14) |
| c. 611G>T         | p. R204L   | NI, AD             | SABTT     | —                           | β-Sheet 3 of NOG structure | Dixon (2) |
| c. 644G>A         | p. W205X   | sporadic           | SYNS1     | Conserved                   | β-Sheet 4 of NOG structure | Dawson (17) |
| c. 655G>C         | p. W205C   | Belgian, AD        | SYNS1     | Conserved                   | β-Sheet 4 of NOG structure | Declau (18) |
| c. 655G>C         | p. W205C   | American, sporadic | SABTT     | Conserved                   | β-Sheet 4 of NOG structure | Enemy (19) |
| c. 645C>A         | p. C215X   | Japanese, AD       | SABTT     | Disulphide bounds in cysteine knot motif to stabilize finger 2 structure | Present study |
| c. 649T>G         | p. W217G   | Hawaiian, AD       | SYNS1     | Conserved                   | β-Sheet 4 of NOG structure | Gong (1) |
| c. 659-660TC>A    | p. I230N   | Belgian, AD        | SYM1      | Almost conserved            | —                           | Leiden (8) |
| c. 665T>A         | p. Y222D   | Belgian, AD        | SYM1      | Almost conserved            | —                           | Leiden (8) |
| c. 666A>G         | p. Y222C   | American, AD       | SYM1      | Conserved                   | —                           | Leiden (8) |
| c. 666A>G         | p. Y222C   | NI, AD             | TCC       | Conserved                   | —                           | Leiden (8) |
| c. 688C>T         | p. P222L   | Belgian, AD        | SYM1      | Conserved                   | —                           | Leiden (8) |
| c. 696G>C         | p. C222W   | German, AD         | SYNS1     | Conserved                   | —                           | Rudnik-Schöneborn (20) |

+ residues are conserved across mammals except for zebrafish.

+ Residues are conserved across mammals except for zebrafish and chicken.
broad thumb and toes/Teunissen and Cremer syndrome. One mutation was a nonsense mutation (C215X), leading to a truncated protein, and was likely to be a pathologic mutation. The other two mutations are also likely to be pathologic rather than functionally neutral polymorphic changes because: (i) none were found in any of the controls, (ii) the alignment of NOG sequences from human, mouse, chicken, Xenopus laevis and zebrafish showed that C155 and C184 are well-conserved amino acids in all species (data not shown), and (iii) all affected subjects showed similar phenotypes.

To date, 36 NOG mutations have been reported in SYM1, SYNS1, TCC, BDB2 and TCS families (Table 2). Although the NOG mutations have been reported mainly in dominant families (Table 2), de novo NOG mutations have also been reported in sporadic SYM1 (8) and sporadic SYNS1 (1). Therefore, genetic investigation may be needed for determining pathogenesis of congenital stapes ankylosis with stiffness of the PIP joints, even in sporadic cases. A milder phenotype (3) as well as the present case with minor joint anomalies in family 2 indicated that it may be clinically important to check such skeletal abnormalities when diagnosing and treating patients with stapes ankylosis, because it may be difficult to differentiate congenital stapes ankylosis from otosclerosis when conductive hearing loss is delayed to adulthood.

Whether NOG mutations can be found more frequently in sporadic conductive hearing loss patients is an interesting question. In this study, mutations were not found in any otosclerosis patients who did not have any associated abnormality. Therefore, typical otosclerosis is not a continuum of the category of diseases associated with NOG mutations. These results, together with the previous literature, indicate that the NOG mutations are restrictively found within patients with various skeletal abnormalities regardless of severity. It is noted that the reported NOG mutation in mild cases (patients with stapes ankylosis without symphalangism) have minor skeletal abnormalities such as broad thumbs and great toes (3), but these cases had symphalangism in the little fingers only.

A review of the reported 41 mutations showed that, interestingly, the majority of NOG mutations are located in the evolutionally well conserved and therefore functionally critical region (Table 2), suggesting that this region might be functionally relevant in NOG polypeptides. This study added three novel NOG mutations in conserved cysteine residue within the cystine knot motif and confirmed that NOG is a causative gene for this category of disease. In addition, there was no particular racial-specific founder mutation within this gene (Table 2). With regard to a genotype–phenotype correlation, phenotypes seem to be independent of the location of the mutation and type of mutations (Table 2). Other genetic factors and/or interacted proteins may also be involved in determining clinical phenotypes.

With regard to surgical outcome, stapes surgery for conductive hearing loss due to NOG mutations may be a good therapeutic option for most cases. In fact, two of these seven patients who underwent stapes surgery (#991 and #4106) had hearing deterioration 3–10 years after the initial surgery, in accordance with a previous report (3, 22), hypothesizing that excessive bony overgrowth and refixation of the ossicle chain may occur after initially successful surgery. The other cases in this study maintained good hearing even after long-term follow-up periods (more than 10 years). Therefore, surgical outcome should be carefully evaluated after long-term observation. Careful explanation of possible limitations of surgical treatment and alternative treatment options such as a bone-anchored hearing aid may be appropriate for such patients with this genetic background.

The identification of the causative genes responsible for various middle/inner ear diseases will enable us to classify new congenital deafness groups in the future, and lead to clinical application in the diagnosis of middle ear disorders and better counseling for the selection of ideal intervention.

Supporting Information

The following Supporting information is available for this article: Fig. S1. (a) Pedigree of family 1. Filled symbol represents the affected individual. (b) Sequence analysis of noggin (NOG). Arrow indicates a G to T change at nucleotide 551 in patient #991. This substitution causes codon 184 to change from TGC (cysteine: C) to TTC (phenylalanine: F).

Fig. S2. (a) Pedigree of family 2. Filled symbols represent the affected individuals. (b) Sequence analysis of noggin (NOG). Arrow indicates a T to A change at nucleotide 463 in patients #3925 and #3926. This substitution causes codon 155 to change from TGC (cysteine: C) to AGC (serine: S).

Fig. S3. (a) Pedigree of family 3. Filled symbols represent affected symptoms (conductive hearing impairment, hyperopia, and finger malformation. (b) Sequence analysis of noggin (NOG). Arrow indicates a C to A change at nucleotide 645 in patients #4106 and #4351. These two patients had a heterozygous C215X mutation. This nonsense mutation (C215X) leads to a truncated protein.

Additional Supporting information may be found in the online version of this article.

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