A Homozygous Truncating Mutation in NALCN Causing IHPRF1: Detailed Clinical Manifestations and a Review of Literature

Amir Hossein Karimi1
Mohammad Reza Karimi1
Poopak Farnia2,3
Farshid Parvini1
Majid Foroutan4

1Department of Biology, Faculty of Basic Sciences, Semnan University, Semnan, Iran; 2Mycobacteriology Research Centre (MRC), National Research Institute of Tuberculosis and Lung Disease (NRITLD), Shahid Beheshti University of Medical Sciences, Tehran, Iran; 3Department of Biotechnology, School of Advanced Technologies in Medicine, Shahid Beheshti University of Medical Sciences, Tehran, Iran; 4Department of Internal Medicine, Semnan University of Medical Sciences, Semnan, Iran

Abstract: Infantile hypotonia, with psychomotor retardation and characteristic facies 1 (IHPRF1), is a rare disorder characterized by global developmental delay and dysmorphic features. This syndrome is caused by genetic anomalies within the NALCN gene. The current report examines a 9-year-old female IHPRF1 patient. Our objective was to contribute to the delineation of the underlying factors influencing this rare condition. Whole exome sequencing (WES) was utilized to identify the disease-causing mutation in the affected individual. Subsequently, Sanger sequencing was performed for the patient, her parents, and two close relatives in order to confirm the detected mutation. Moreover, detailed clinical examinations including EEG, echocardiography, and biochemical/physical tests were carried out to elucidate the effects of the mutation. WES identified a homozygous nonsense mutation in the NALCN gene (c.2563C>T p.R855X). This mutation was confirmed by Sanger sequencing in the patient and her family members and segregated with the autosomal recessive inheritance pattern of IHPRF1. Moreover, genotype-phenotype correlation analysis confirmed the disease-causing nature of this mutation. The current report provides the first detailed description of a patient with this homozygous nonsense mutation (c.2563C>T p.R855X) and expands the clinical spectrum of IHPRF1 disease. Possible influences of sex and other factors on this disease are discussed and a review of the literature is also provided.

Keywords: global developmental delay, dysmorphism, intellectual disability, motor retardation, cognitive delay

Plain Language Summary
Infantile hypotonia with psychomotor retardation and characteristic facies 1 (IHPRF1), is a rare neurological disorder caused by mutations in the NALCN gene. The product of this gene is the main component of a conserved ion transporting channel involved in determining the resting membrane potential of different cell types including neurons and is involved in many physiological processes such as respiration, intestinal pacemaking, and regulation of circadian cycle. Individuals affected by IHPRF1 are most prominently characterized by global developmental delay and dysmorphic features. Due to our poor understanding of the NALCN ion transporting channel and the limited number of reported IHPRF1 patients so far, there is no developed comprehensive perspective on the underlying factors affecting this disorder and the variability of the affected systems and processes. Herein, we provide a detailed description of an IHPRF1 patient with a homozygous truncating mutation, manifesting a wide range of severe abnormal symptoms. We also provide a thorough review of the literature and propose sex and other elements as possible factors influencing this rare syndrome. Moreover, we draw the attention of the future investigators to a described
activator of the NALCN ion transporting channel known as 
substance P, as a potential participant in many of the anomalies 
observed in the diseases associated with NALCN ion transport-
ing channel.

**Introduction**

Sodium leak channel, non-selective (NALCN) is the sole 
non-selective member of the 24-TM ion channel 
superfamily. This channel is an essential component in 
the regulation of the resting membrane potential of several 
cell types including neurons by conducting a sodium back-
ground conductance. NALCN along with UNC80 and 
UNC79 and NLF-1/FAM155A are the four main subunits 
of a channelosome which is known to be involved in 
rhythmic behaviors including circadian cycle, respiration, 
and intestinal pacemaking.

The activity of the NALCN channelosome is regulated 
by different types of mediators. This includes its activation 
via neurotensin, acetylcholine, and substance P (SP) and 
its inhibition by GABA, dopamine, and extracellular calcium.

Several diseases have been shown to be associated with 
the NALCN channelosome. While no disease-causing 
mutation has been reported for the UNC79 gene, mutations 
in the NALCN and UNC80 genes can cause infantile 
hypotonia with psychomotor retardation and characteristic 
facies 1 (IHPRF1, OMIM 615,419) and IHPRF2 (OMIM 
616,801), respectively. Both IHPRF1 and IHPRF2 are 
shown to follow the autosomal recessive inheritance pattern 
of Mendelian disorders. It is worth mentioning that 
mutations in NALCN could also lead to autosomal domi-
nant congenital contractures of the limbs and face, hypo-
tonia, and developmental delay syndrome (CLIFAHDD, 
OMIM 616,266). A recent study indicates that IHPRF1 
and CLIFAHDD are caused by loss and gain of function 
mutations, respectively.

Here, we report a 9-year-old female IHPRF1 patient 
from a highly consanguineous Afghan family residing in 
Iran. The patient manifests a wide range of symptoms 
including severe hypotonia, severe global developmental 
delay, lack of speech, severe muscle atrophy, hypersalivation, 
and severe constipation (Table 1). The patient has 
also had an older male sibling with similar abnormalities 
who has died undiagnosed at the age of 3.5 months. 
Parents sought genetic counseling during the mother’s 
third pregnancy after their gynecologist suspected genetic 
defects in the proband. Our examinations revealed 
a homozygous nonsense mutation in the NALCN gene 
(NM_052867:c.2563C>T) as the cause of the defects in 
the proband and the heterozygosity for this mutation was 
confirmed prenatally for the third sibling. The present 
study further expands the clinical symptoms associated 
with IHPRF1 by describing the wide range of abnormal 
phenotypes manifested in the affected individual. A review 
of the literature is provided and a number of issues that 
might prove beneficial to be taken into account in future 
reports and studies are also discussed.

**Materials and Methods**

This study was approved by the ethics committee of 
Semnan University of medical sciences, Semnan, Iran. 
Informed consent was obtained from the legal guardians 
and Karyotype test was performed for the patient. Since 
no chromosomal abnormalities were detected, genomic 
DNA samples were prepared with the QIAamp DNA 
Blood Mini Kit (Germany) and whole exome sequencing 
(WES) was carried out for about 100 million reads, using 
the Illumina Hiseq2000 sequencer platform. Obtained 
results were analyzed by open-access bioinformatics 
tools, namely BWA aligner, GATK, and annovar. In 
order to inquire about the novelty of the mutation, 
public resources including ClinVar, gnomAD, Kaviar, 
and GME were utilized. Moreover, the local 
population database with more than 3000 unrelated indi-
viduals (BayanGene) was examined in order to deter-
mine the frequency of the mutation in the local 
population.

Online bioinformatics tools such as MutationTaster and 
CADD_phred were used in order to predict the probable 
effects of the mutation. Subsequently, Sanger sequencing 
was performed for the patient, her parents, and two close 
relatives to confirm the presence of the identified mutation 
and analyze its segregation. The primers used for this test 
were: F-5' AGCAAGAGGACGCATGAT 3' and R-5' 
CTTCATTGGGAAAATGGC 3'. Chromas software 
was applied to analyze the resulting data from Sanger 
sequencing. To determine the status of the mutation in the 
fetus, genomic DNA isolated from the amniotic fluid cells 
was sequenced. Furthermore, quantitative fluorescence poly-
merase chain reaction (QF-PCR) was carried out to investi-
gate short tandem repeat (STR) markers on chromosomes X, 
Y, 13, 18, and 21 of the fetus, using “3130 Genetic Analyzer” 
and fragment analysis software.

Clinical examinations including echocardiography and 
electroencephalography (EEG) as well as physical exam-
inations were performed for the patient. Since any further
Table 1 Clinical Characteristics of the Proband and Her Deceased Sibling

| Characteristics       | Patient | Sibling |
|-----------------------|---------|---------|
| Sex                   | Female  | Male    |
| Cognitive delay       | Severe  | Severe  |
| Height                | 68 cm   | Na      |
| Weight                | 6.8 kg  | Na      |
| Head circumference    | 46 cm   | Na      |
| Hypotonia             | Severe  | Severe  |
| Microcephaly          | +       | + (Severe) |
| Micrognathia          | +       | +       |
| Smooth philtrum       | –       | –       |
| Low set ears          | –       | –       |
| Large ears            | –       | –       |
| Strabismus            | +       | +       |
| Nystagmus             | –       | –       |
| Wide mouth            | +       | +       |
| Persistently open mouth | +     | +       |
| Pectus carinatum      | +       | +       |
| Pes varus             | +       | +       |
| Scoliosis             | Severe  | Severe  |
| Triangular face       | +       | +       |
| High arched palate    | +       | +       |
| Hypersalivation       | +       | +       |
| Eye contact           | Poor    | Na      |
| Muscle atrophy        | Severe  | Severe  |
| Seizure               | –       | –       |
| Sleep disturbance     | +       | +       |
| Lack of speech        | +       | Na      |
| development           | Cardiovascular malformation | ASD, TR |
|                       | ASD     | ASD     |
| Pain sensation        | +       | Na      |
| Spastic tetraplegia   | –       | Na      |
| Recurrent infection   | + (Respiratory tract) | + |
| Constipation          | Severe  | Na      |
| Optic atrophy         | –       | Na      |
| Apnea                 | +       | +       |
| Neuroimaging          | Frontal and temporal lobe atrophy | Na |
| Electroencephalography| Severe abnormal (hypsarrhythmia) | Na |
| Early death           | –       | + (3.5 months) |

Abbreviations: Na, not available; ASD, atrial septal defect; TR, tricuspid regurgitation.

Results

The WES analysis identified a stop gain mutation in the NALCN gene on chromosome 13 (NM_052867; chr13:101107503G/A c.2563C>T p.R855X (rs376152742). The mutation was predicted to be pathogenic by MutationTaster and CADD_phred. Our review of the public resources, local population database, and the literature revealed that there is no previous publication describing this mutation in homozygous form. However, a single ClinVar submission confirmed the pathogenic effects of this genetic anomaly but merely provided an overview of the resulting manifestations with no detailed information. Furthermore, it failed to distinguish between IHPRF1 and CLIFAHDD and resulted in confusion by referring to several publications on CLIFAHDD as supporting evidence (SCV000742166.1). Sanger sequencing confirmed the presence of the mutation and its homozygosity in the proband and segregated with the autosomal recessive inheritance pattern of IHPRF1 in all tested individuals (Figure 1).

Clinical examination results of the proband included severe muscle atrophy, severe scoliosis (improved by bracing), constipation, poor eye contact, and dysmorphic features such as a triangular face, long thin fingers, strabismus, pectus carinatum, and pes varus. Echocardiography results diagnosed the patient with atrial septal defect (ASD) and tricuspid regurgitation (TR).

Additionally, our examination of the patient’s clinical history revealed that severe global developmental delay was evident from the first months of life. She never gained the ability to roll over or sit without support. Neuroimaging showed frontal and temporal lobe atrophy, particularly around the Sylvian fissure region, and cortical perisylvian polymicrogyria was suspected. EEG was performed for the patient during sleep and included generalized slow activity and attenuation with superimposed beta range activity. Asymmetry was present and the results were compatible with hypsarrhythmia. The complete blood count and biochemistry profile included a high level of white blood cells, low hematocrit, low levels of creatinine and blood ammonia, a very high level of aspartate aminotransferase (AST), and a normal level of alanine aminotransferase (ALT). Plasma amino acid profile included low levels of glycine, alanine, alpha-amino butyric acid, tryptophan, valine, and leucine. Abdominopelvic ultrasound was unremarkable. The TSH and T4 levels were within the normal range.
range. The Patient also manifested recurrent and prolonged respiratory tract infections (Table 1).

The results of the QF-PCR performed prenatally for the third sibling showed a female fetus with no chromosomal abnormalities. Sanger sequencing revealed that the fetus was heterozygote for the identified mutation in the NALCN. Echocardiography at 26 weeks of gestation diagnosed the fetus with ASD. She passed away at 3 months of age due to an unknown cause.

**Discussion**

Here, we report an IHPRF1 patient with a homozygous mutation c.2563C>T p.R855X in the NALCN gene. The wild type NALCN protein contains four homologous repeats, each of which comprises six transmembrane segments with the pore-forming region of each domain located between the fifth and the sixth transmembrane segments. Due to the truncating effect of the mutation, loss of 884 out of 1738 amino acids of the wild type protein is predicted. Hence, the resulting protein would contain only two out of the four above-mentioned domains (Figure 1).

Although the same mutation was previously described in a compound heterozygote individual (pat: p.R855X, mat: p. R1094Q), there are no previous publications of this mutation in the homozygous form. Hence, the current report provides the first description of the direct effects of this truncating homozygous mutation. Interestingly, the patient described by Campbell et al (2018) is probably the least affected individual among all the reported IHPRF1 patients, so far. They suggested that the milder manifestation of the disease phenotypes might

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**Figure 1** (A) Patient's dysmorphic phenotypes. Pectus carinatum, severe muscle atrophy, pes varus, and scoliosis are evident. (B) Patient's pedigree along with the results of Sanger sequencing for the respective individuals. All tested individuals, except for the proband, were revealed to be heterozygote for the detected mutation in the NALCN gene. (C) The NALCN protein structure. This protein consists of four transmembrane domains (shown by D1 to D4), each of which contains six transmembrane segments. Twenty-one of the described missense, frameshift, and nonsense mutations are marked with green, orange, and red, respectively. The homozygous nonsense mutation detected in our patient (Arg855Ter) (distinguished by an ellipse) is located on the cytoplasmic loop, between the second and the third transmembrane domains. The hypothetical truncating effect of this mutation on the protein is illustrated.
be due to the fact that the missense mutation of the compound heterozygote patient is located on the third transmembrane domain of the NALCN protein. However, it is noteworthy that a missense mutation in the fourth transmembrane domain of the NALCN also resulted in a milder manifestation of the disease.\textsuperscript{20} Moreover, there are reports of several truncating mutations located both in and after the third transmembrane domain of the NALCN protein, which resulted in much more severe manifestations of the disease.\textsuperscript{20–22} Considering these, the exhibition of milder clinical symptoms in the patients reported by Campbell et al (2018), the patients in the family 2 described by Al-Sayed et al (2013), and individuals 16 and 17 described by Bramswig et al (2018) might also be attributed to the missense nature of the mutation. Three points, however, should be taken into account: First, hypothetically, mRNAs produced from the mutated genes with stop gain mutations can undergo nonsense-mediated decay (NMD), thus, the more severe exhibition of the disease phenotypes in patients with stop gain mutations in or after the third transmembrane domain of the NALCN might be due to the general downregulation of the truncated protein by this mechanism. Second, to the best of our knowledge, there are only four reported missense mutations in the NALCN gene which are all located in the third and the fourth transmembrane domains of NALCN protein (Figure 1).\textsuperscript{19,20,23} Third, in a recent study, no sodium background current was detected in cells expressing p.Trp1287Leu missense mutation which might suggest underlying factors other than residual channel activity for the milder exhibition of the symptoms in the individuals affected by this mutation.\textsuperscript{10} With that being said, further accumulation of the IHPRF1 reports can help in the delineation of this matter. Beyond any doubt, rapid progress in the field of next-generation sequencing allows for a more accurate and less expensive diagnosis and study of genetic defects.\textsuperscript{24–26}

Considering the extremely high rate of constipation in all the IHPRF1 patients reported so far (present in 31 out of 34 examined individuals), constipation can be considered as one of the most prominent symptoms of this disease, along with global developmental delay and dysmorphic features (for a summary of the symptoms manifested in all the 36 patients reported so far, please see the Supplementary table). Cardiovascular malformation, however, is seldom associated with IHPRF1. To the best of our knowledge, other than the current case, there is only one other reported individual diagnosed with ASD,\textsuperscript{27} and there is no previous report of TR in IHPRF1 patients.

The patient described in this report had an older male sibling with similar phenotypes who passed away genetically undiagnosed at the age of 3.5 months. Interestingly, the male sibling also exhibited seizures which is a common phenotype in IHPRF1 patients (present in more than 47% of the described individuals). This made us suspect that sex might play a role in the extent of the disease severity. We specifically compared the disease status in all the seven opposite-sex IHPRF1 siblings described in the previous reports.\textsuperscript{19,20,22,28,29} Our interpretation of the information provided by the authors in their respective manuscripts is that there seems to be a bias towards the more severe exhibition of the IHPRF1 phenotypes in the male individuals. In two of the described siblings (individuals 2 and 3, and individuals 11 and 12 described by Bramswig et al (2018)), we were not able to identify any major difference between the individuals. In all the other siblings described, the male individuals seem to present a more severe phenotype compared to their female siblings. Although the number of reported siblings is not enough to draw conclusions, we suggest that the sex of the affected individual might have an influence on the extent to which IHPRF1 phenotypes manifest. In agreement with this hypothesis, the study of the effects of estrogen and progesterone hormones on the regulation of the NALCN expression in human myometrial smooth muscle cells suggests a sex-dependent biology for the NALCN.\textsuperscript{30}

NALCN has been identified as a cation channel involved in the conductance of substance P-activated cation channel current.\textsuperscript{4} Considering that it has been shown that SP plays important roles in a wide variety of physiological processes including cardiovascular system activity, cell growth and proliferation, inflammation, pain sensation, respiration, and pacemaking activity in the gastrointestinal tract,\textsuperscript{31–33} some of the phenotypes associated with IHPRF1 disease can be attributed to the disability of the mutated NALCN in the conductance of the SP-activated cation channel current. Indeed, it has been shown that NALCN is an important component in the SP-mediated modulation of gastrointestinal motility through interstitial cells of Cajal and also is essential in respiratory system response to SP.\textsuperscript{34,35} Furthermore, the study of IHPRF1 from the perspective of the other known regulators of the NALCN channelosome, namely neuropeptide, acetylcholine, GABA, extracellular calcium, and dopamine, would be essential for a better understanding of this disorder.\textsuperscript{5–7}

**Conclusion**

We utilized WES and Sanger sequencing to identify a disease-causing mutation in the NALCN gene in
a patient with dysmorphic features and severe global developmental delay. Our main limitation was the patient’s intolerance for further clinical examinations. Our results further expand the clinical spectrum of IHPRF1. We propose sex-based differences as possible influencers of this disease. Further study of the biology of NALCN is essential for a better understanding of this disease. Regarding the importance of SP in a wide variety of physiological activities, the future study of the role of NALCN in these processes might prove beneficial.

Abbreviations
ALT, alanine aminotransferase; AST, aspartate aminotransferase; ASD, atrial septal defect; CLIFAHD, congenital contractures of the limbs and face, hypotonia, and developmental delay syndrome; EEG, electroencephalography; IHPRF, infantile hypotonia with psychomotor retardation and characteristic facies; NALCN, sodium leak channel, non-selective; NMD, nonsense-mediated decay; QF-PCR, quantitative fluorescence polymerase chain reaction; STR, short tandem repeats; SP, substance P; TR, tricuspid regurgitation; WES, whole exome sequencing.

Ethics Approval
This study was conducted under the approval of the ethics committee of Semnan University of medical sciences, Semnan, Iran. Written informed consent was obtained from the legal guardians prior to all the relevant clinical tests and the preparation and submission of the manuscript. Consent for publication of patient’s data and any identifiable features is also available.

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Author Contributions
All authors made substantial contributions to conception and design, acquisition of data, or analysis and interpretation of data; took part in drafting the article or revising it critically for important intellectual content; agreed to submit to the current journal; gave final approval of the version to be published; and agree to be accountable for all aspects of the work.

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