NGLY1 Deficiency Zebrafish Model Manifests Abnormalities of the Nervous and Musculoskeletal Systems

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Background: NGLY1 is an enigmatic enzyme with multiple functions across a wide range of species. In humans, pathogenic genetic variants in NGLY1 are linked to a variable phenotype of global neurological dysfunction, abnormal tear production, and liver disease presenting the rare autosomal recessive disorder N-glycanase deficiency. We have ascertained four NGLY1 deficiency patients who were found to carry a homozygous nonsense variant (c.1294G > T, p.Glu432*) in NGLY1.

Methods: We created an ngly1 deficiency zebrafish model and studied the nervous and musculoskeletal (MSK) systems to further characterize the phenotypes and pathophysiology of the disease.

Results: Nervous system morphology analysis has shown significant loss of axon fibers in the peripheral nervous system. In addition, we found muscle structure abnormality of the mutant fish. Locomotion behavior analysis has shown hypersensitivity of the larval ngly1(−/−) fish during stress conditions.

Conclusion: This first reported NGLY1 deficiency zebrafish model might add to our understanding of NGLY1 role in the development of the nervous and MSK systems. Moreover, it might elucidate the natural history of the disease and be used as a platform for the development of novel therapies.

Keywords: NGLY1 deficiency, zebrafish, nervous system, musculoskeletal system, abnormalities

1 INTRODUCTION

N-glycanase deficiency (OMIM # 615273) is a rare autosomal recessive disorder linked to a variable phenotype in the patients (Enns et al., 2014) and caused by pathogenic variants in NGLY1 (Need et al., 2012; Lam et al., 2017). Currently, there are fewer than 100 patients reported worldwide, and possibly many more undiagnosed, who suffer from this devastating disease with no effective treatment (Tickotsky-Moskovitz, 2015). One of the major manifestations in patients with N-glycanase deficiency is severe global developmental delay (motor and cognitive), neurological dysfunction and damage to both the central nervous system (CNS) and peripheral nervous system (PNS), leading to significant locomotion disability (Jones et al., 2012). Furthermore, early onset liver disease and elevated liver transaminases are also part of...
2 MATERIALS AND METHODS

2.1 Clinical Data
The patients were recently reported (Kalfon et al., 2021). The guardians of the affected individuals signed an informed consent form for participating in this study. The Israeli Supreme Helsinki committee approved the study; GMC 03-04-2006.

2.2 Cell Culture
Primary fibroblast cell cultures were derived from skin biopsies taken from one NGLY1 deficiency patients and maintained in a 37°C and 5% CO2 incubator.

2.3 Western Blotting
Proteins were extracted from fibroblast cells of healthy donors and affected individuals as well as from amputated fish tails (3mpf) of WT and mutant fish. Electro-blotting was done using a power supply with the conditions of 60 V/180 mA, for 1.5 h. NGLY1 primary antibody, at 1:500 concentrations, was added to the membrane for overnight incubation at 4°C, and then underwent 1 h of incubation with horseradish peroxidase (HRP) secondary antibody Goat-anti-rabbit at 1:2000 concentration. Finally, the membrane was filmed using G:BOX (Syngene, Frederick, WA, United States) with a chemiluminescence light.

2.3.1 Antibodies Used
Anti NGLY1 antibody (Rabbit anti Human), Sigma-Aldrich, HPA036825 (Sigma-Aldrich, St Louis, MO, United States).
Secondary antibody IgG-HRP Goat anti-Rabbit, Santa Cruz Biotechnology (Dallas, Texas United States; Cat. No. SC 2004).
Anti β-Actin antibody, anti-Mouse monoclonal, cell-signaling technology (Danvers, MA, United States) Cat. No. 8H10D10.
Secondary antibody, IgG-HRP Goat anti-Mouse, Invitrogen (Paisley, United Kingdom; Cat. No. A16072).

2.4 Immunofluorescence Staining
The cells were fixed with 4% formaldehyde, permeabilized with 0.5% Triton X-100 in PBS, and blocked with blocking solution (10% FBS, 1% BSA, and 0.05% Triton X-100 in PBS). Immunofluorescence staining was performed with 1:100 Anti NGLY1 antibody (Rabbit anti-Human), Sigma-Aldrich, HPA036825 in blocking solution overnight at 4°C. Next, the cells were incubated with 1:700 Alexa 488-labeled secondary antibody (A-21202 by Invitrogen, excitation at ~495 nm, emission ~519 nm). Finally, cover slips were laid on slides containing 4 µl of mounting medium with DAPI nucleus staining (Vector Laboratories, Burlingame, CA, United States). Fluorescent images were obtained by Nikon Eclipse Ti and Nikon C2 confocal scanner (Nikon Instruments Inc., Melville, NY, United States). Fluorescence images were analyzed with ImageJ software.

2.5 Bioinformatic Tools and Online Databases
Bioinformatic software and databases were used throughout the research, as follows:
Primer design tools: PRIMER3, NCBI primer BLAST.
Genome databases: UCSC, ENSEMBLE genome browser, NCBI PUBMED, OMIM (Online Mendelian Inheritance in Man), GeneCards, HGMD (Human Genome Mutation Database), dbSNP (Database of Single Nucleotide Polymorphism).
TABLE 1 | Heterozygous fish with ngly1 mutant alleles.

| Fish number | Allele 1 | Allele 2 |
|-------------|----------|----------|
|             | Nucleotide change | Amino acid change | Nucleotide change | Amino acid change |
| 1           | c.1759-1777del | p.517fs | - | - |
| 2           | c.1553-1557del | p.443fs | - | - |

Analysis tools: Nudieotide primer BLAST, reverse complement design tool, Mutation Taster, Alamut Visual. Restriction enzymes design tools: NEB Cutter. CRISPR design (fish) tool https://crispr.cos.uni-heidelberg.de/.

2.6 Generating Zebrafish Model
All animal research followed a protocol approved by BIU IACUC- 023_b15280_80.

2.7 Ngly1 Mutant Zebrafish Established by Targeting Ngly1 Exon 9 Using CRISPR-Cas9
In order to create a premature stop codon in exon 9 similar to the genetic variant in our ngly1 patients, we used CRISPR/Cas9 mediated ngly1 targeting in ZF embryo (Figure 2A). A guide RNA (gRNA) was designed to target exon 9 of ngly1 (NM_001020601.1). We analyzed ngly1 CRISPR/Cas9 mediated gene targeting in G0 generation by extracting DNA from 24 hpf injected embryos to determine mutagenesis efficiency. Cas9 protein and gRNAs (Sigma-Aldrich) targeting ngly1 exon 9 were injected to one cell stage zebrafish embryo. Twenty-four hours after injection a sample of injected embryos was genotyped by PciI restriction enzyme which has a recognition site at the Cas9 nuclease cleavage site. In order to obtain ngly1 knockout zebrafish, F0 founder fish harboring ngly1 mutation were mated with WT. F1 heterozygotes (3mpf) were genotyped. Two different ngly1 mutant alleles were found (Table 1; Figure 2B). We crossed F1 del 19bp fish with a desired wild-type (WT) fish (outcrossing) in order to reduce off-target mutations effect. F2 heterozygotes were crossed with each other to create ngly1<sup>+/−/−</sup> and ngly1<sup>+/+(+/−)</sup> fish.

2.8 Zebrafish Maintenance
Zebrafish (Danio rerio) of AB strain were maintained at 28°C under 14 h light: 10 h dark cycles. For the experiments, adults and larval fish were euthanized by immersion in methane sulfonate (MS222), Tricaine 0.4% with subsequent placing on ice.

2.9 The Guide RNA Design
The website (https://crispr.cos.uni-heidelberg.de/) was used to design gRNAs for targeting exon 9 of zebrafish ngly1 gene. The gRNA designed contained recognition site for PciI restriction enzyme near the PAM sequence which allows screening for CRISPR’s success in mutating exon 9 of ngly1.

2.10 Embryo Injection
A mix of ngly1 gRNA and Cas9 mRNA was injected directly into one-cell-stage embryos, using a pneumatic Pico Pump (WPI, Worcester, MA, United States). Each embryo was injected with 500 µl of solution containing ~300 ng/µl of gRNA and ~300 ng/µl of Cas9 protein.

2.11 Zebrafish Genotyping
2.11.1 Tail Clipping (Adults)
Fish at 3 months post fertilization (mpf) were anesthetized by immersion in fish system’s water containing 0.016 mg/ml Tricaine (Sigma-Aldrich) and tail clipping was performed.

2.11.2 DNA Extraction From zebrafish Tail and PCR Amplification
DNA was extracted from fish tail (3mpf) by using a standard protocol.

2.12 cDNA Synthesis
2.12.1 RNA Purification From ZF Tail
RNA and cDNA production was performed from fish tail (3mpf) using standard protocol.

2.12.2 mRNA Levels Evaluation
cDNA amplicons for ngly1 and rpl32 were evaluated using RT-PCR. rpl32 was used as a reference gene and normalized to healthy controls.

2.12.3 Fluorescent In-Situ Hybridization
Fluorescent probe was designed in order to label the ngly1 mRNA in zebrafish larvae tissues (Fried and Sherma, 2010) (Integrated DNA Technologies, Coralville, IA, United States). At 6dpf the larvae were fixed in 4% paraformaldehyde (Sigma-Aldrich). Next the larvae were dehydrated with 25%, 50%, and 75% ethanol for 15 min each. Larvae were incubated with 10 µg/ml Proteinase K for 45 min to elevate the permeability of the tissues. Finally, the larvae were incubated with 200 ng/µl ngly1 probe: (5’CGCTTCACCACATAGACCGTGCTTGTCTCCCGCGAAGGAAACACCCTA) and visualized under Nikon Eclipse Ti and Nikon C2 confocal scanner (Nikon Instruments Inc.). In order to validate the ngly1 expression in the nervous system.

2.13 Establishing a Stable Ngly1 Mutant Transgenic Zebrafish Line
Tg (HUC:gal4 uas:mem YFP-mito CFP) co-expresses memYFP and mitoCFP (Zada et al., 2014) and was kindly provided by Prof. Lior Appelbaum (Bar-Ilan University, Ramat-Gan, Israel). In order to establish a ngly1 mutant transgenic fish we crossed F0 Tg (HUC:gal4 uas:mem YFP-mito CFP) to homozygote ngly1<sup>+/−</sup> and selected heterozygotes from F1. F1 heterozygotes whose inbreeding produced homozygous ngly1<sup>+/−</sup>/Tg (HUC:gal4 uas:
mem YFP-mito CFP) and ngly1(+/+)Tg (HUC:gal4 uas:mem YFP-mito CFP) siblings (F2). The PNS morphology was imaged by confocal Nikon eclipse.

2.14 Phalloidin Staining
Larvae at 6 dpf were fixed in 4% paraformaldehyde (Sigma-Aldrich). Next the larvae were dehydrated with 50% and 70% ethanol and then stained with Phalloidin fluorescent dye that stains F-actin in fixed cells, stabilizes actin filaments in vitro and visualizes them (Wulf et al., 1979). Finally, we focused on larval muscle by visualization under fluorescent microscope- Nikon Eclipse Ti and Nikon C2 confocal scanner (Nikon Instruments Inc.). Images were converted in 12 bit using software and exported in tiff format. Fluorescence was measured using Nikon Eclipse Ti software; data obtained were represented as the ratio between total pixel intensity and the number of pixels. Mean size of the somite was measured in μm using Nikon Eclipse Ti software. Muscle morphology and intensity were compared between ngly1(−/−) and ngly1(+/+) siblings (Mignani et al., 2020).

2.15 Motor Assay
Zebrafish larvae at 6 dpf were placed in 96-wells plate, and behavior was recorded by DanioVision (Noldus Information technology, Wageningen, Netherlands) system. The results were analyzed by the Ethovision XT -11 locomotion tracking software (Noldus). We performed a basic locomotor assay that consisted of three different phases: 15 min lights-on, then 15 min lights-off (stress stimulus) and finally 15 min of lights-on (recovery). ngly1(−/−) and ngly1(+/+) siblings were video tracked, and swimming distance patterns were measured and analyzed.

2.16 Pain Sensitivity Test
A pain sensitivity test based on sensitized acid aversion assay developed by Steenbergen and Bardine (2014) was performed; the premise of this experiment is based on the increase in movement presented by larvae exposed to acidic environment. 6dpf zebrafish larvae were placed in 96 wells plate containing an insert of 96 wells with a 100-um mesh bottom (Millipore, Massachusetts United States); all larvae could be simultaneously moved from the water with a neutral pH to acetic acid 0.01% water (pH = 3.7–3.9) where they stayed for 10 min. Fish behavior was recorded by DanioVision (Noldus) system for 10 min, and swimming distance patterns of ngly1(−/−) and ngly1(+/+) siblings were calculated.

3 RESULTS

3.1 Fibroblast Cell Culture Analysis
In order to investigate the studied genetic variant effect on NGly1 protein level in humans, proteins were extracted from fibroblasts derived from a patient (NH-130) and a healthy control (NH-94). NGly1 protein was absent in the patient with NGly1 deficiency (Figures 1A,B).

3.2 Zebrafish mRNA and Protein Expression
A high percentage (>90%) of embryos harbored mutations at ngly1 exon 9, indicating the efficiency of CRISPR-Cas9 in inducing mutations with the ngly1 gRNA designed. Out of 62 F1, 14 fish were heterozygotes to a deletion of 19bp (c.1759–1777) resulting in p.met 517fs (Figures 2A,B). This mutation causes the appearance of a premature stop codon creating a 535 amino acid protein instead of the 644 amino acid full length of ngly1 zebrafish protein. The 14 F1 heterozygote fish were crossbred with each other to obtain the F2 generation. F2 were in-bred to generate F3 ngly1 homozygote, heterozygote and WT offspring for phenotype-genotype correlations analysis (Figure 3A). A significantly lower ngly1 mRNA expression was detected in the ngly1(−/−) fish compared to the ngly1(+/+) fish as expected (p < 0.001) (Figure 3B). The positive control rpl32 mRNA showed similar expression levels in both genotypes. Similarly, the western blot analysis demonstrated a significant reduction in protein level of ngly1 in the ngly1(−/−) fish compared to the ngly1(+/+) fish (p < 0.01) (Figures 3C,D).

3.3 Ngly1 Expression Pattern
To study the expression pattern of ngly1 in the CNS and PNS, we performed in-situ hybridization analysis in larvae nervous system. Ngly1 is significantly expressed in the brain, spinal cord and peripheral nervous system of WT ZF larvae (Figures 4A–D). In order to validate the ngly1 expression in the nervous system, the green fluorescent regions were compared to a Tg (HUC:gal4 uas:mem YFP-mito CFP)- fish that constantly expresses a fluorescent protein which labels the nervous system as a positive control (Figures 4E,F).

3.4 Nervous System Live Imaging Analysis
Nervous system morphology was analyzed at 6dpf in ngly1(−/−) Tg (HUC:gal4 uas:mem YFP-mito CFP) and ngly1(+/+) Tg (HUC:gal4 uas:mem YFP-mito CFP) sibling. The results demonstrate that ngly1(−/−) display a significant reduction (p < 0.001) of peripheral axon fibers in comparison to control group (Figures 4G–I).

3.5 Musculoskeletal Morphology
Muscle structure was analyzed at 6dpf in ngly1(−/−) fish and WT siblings. The result confirmed that ngly1(−/−) fish demonstrate a reduction in muscle mass (p < 0.01) and somite size (p < 0.001) in comparison to the control group (Figure 5).

3.6 Motor Assays
To investigate the functionality of the MSK system in ngly1 mutant fish, we tested ZF swimming. The ngly1(−/−) larval ZF displayed significantly less swimming distance with respect to the control ngly1(+/+) (p < 0.001). Under stress stimulus, when lights were turned off, both groups showed an increase in the distance moved compared to lights-on phase (Figures 6A–C). However, ngly1(−/−) fish showed a significantly higher increase in swimming distance than the control ngly1(+/+) fish (p < 0.01).

3.7 Sensitized Acid Aversion Assay
To further characterize the neural phenotype, we performed a pain insensitivity test by sensitized acid aversion assay. The premise of this experiment is based on the nociceptive response presented by zebrafish larvae. Steenbergen and
Bardine (2014), demonstrated that zebrafish larvae show behavioral responses to a noxious chemical stimulus and that the sensory nervous system receptors mediate a response to pain by acid aversion. The two fish groups—ngly1(−/−) and ngly1(+/+)
were compared at the larval stage (6 dpf). The ngly1(−/−) larvae reacted to the pain with a significant increase in swimming distance compared to ngly1(+/+) after the treatment by acetic acid (pH 3.8) (p < 0.05) (Figures 6D,E).

4 DISCUSSION

In the present study we established a viable ngly1 mutant zebrafish model and studied the phenotypic consequences of the perturbed ngly1. The chosen fish model offers unique opportunities for gene modification by CRISPR/Cas9 technologies followed by gross morphological examination, live imaging of the dynamic behavior and anatomic characterization of the whole organism, from the embryonic to adult stages. As far as we know, no zebrafish model for NGLY1 deficiency had been generated to date.

Our research was based on a study of four patients from two related Druze families who presented with clinical features of NGLY1 deficiency (Kalfon et al., 2021).

The literature describes several attempts to create animal models that mimic the disease in mice (Fujihira et al., 2017), Drosophila (Galeone et al., 2017) and rats (Asahina et al., 2020).
However, there was little evidence for the causative role of NGLY1 disruption in the pathophysiology of the disease.

We created a nonsense genetic variant in exon 9 from 11 exons in a zebrafish model, similar to the genetic variant in our patients that disrupts the normal gene sequence and leads to a premature termination codon in position 534 of the protein (ngly1’s full protein length is 644 amino acid). The AA sequence of the mutant fish is not identical to that of human patients, however it is very similar as both the human and fish mutation are a nonsense mutation in exon 9 from 11 exons. qPCR analysis of mRNA levels

![Figure 3](image-url)

**FIGURE 3** | Mutation characterization in zebrafish. (A) Genotype screening in DNA extracted from zebrafish. Representative electrophoregrams indicating the sequences of WT, heterozygote and homozygote. (B) qPCR analysis quantification of ngly1 mRNA levels. t-test (ngly1+/+) n = 3, ngly1−/− n = 3, ***p < 0.001. (C) WB analysis quantification of ngly1 protein levels. t-test (ngly1+/+) n = 3, ngly1−/− n = 3, **p < 0.01. (D) WB analysis with an antibody against ZF ngly1; β-actin was used as a loading control.
in the zebrafish revealed significant reduction in the levels of ngly1 mRNA in the ngly1<sup>−/−</sup> fish compared to the ngly1<sup>+/+</sup>. We also observed massive decrease in the levels of ngly1 protein in the ngly1<sup>−/−</sup> fish compared to the ngly1<sup>+/+</sup>. However, we demonstrated a residual amount of the ngly1 protein in the mutant fish. Indeed, Smits et al. (2019) described this phenomenon of residual protein expression, that is, characteristic to the CRISPR-Cas9 system, supporting the validity of our data. Another possible explanation for the residual protein in the mutant fish, is alternative splicing variation in different tissues of the fish model. However, since exon 9 of ZF codes for 63 amino acids (~7 KD) and the WB analysis showed a normal size of the mutant protein, the mechanism of exon skipping is less likely to be the cause for the presence of residual protein. The residual protein in the mutant fish has size similar to the normal ngly1 protein, therefore we assume that there is minimal catalytic activity in the mutant fish. This enzymatic activity may explain the fact that mutant larva present with “mild” phenotype compared to the human patients carrying the same genetic variant.

A number of loss-of-function pathogenic variants in human NGLY1 have been reported, including nonsense, missense, frameshift, and splice site variants. These occur all through the gene, with no obvious hot spots (Gene reviews NGLY1). The p.Arg401Ter nonsense variant is the most common, accounting for approximately one third of pathogenic alleles (Enns et al., 2014). Affected individuals harboring at least one copy of c.1201A>T (p.Arg401Ter) tend to have a more severe clinical course with higher scores on the Nijmegen Pediatric CDG Severity scale (Lam et al., 2017). A sib pair with the cryptic pathogenic c.930C>T splice site variant (predicted as a silent p.Gly310=) and a p.Gln208Ter nonsense variant exhibited relatively mild impairment in all domains (Lam et al., 2017). Nevertheless, no clear phenotype genotype correlation has yet been established.

In order to compare between the phenotypes of NGLY1 deficiency fish model and the human patients, we explored and characterized two systems: the nervous system and the MSK system.
The expression pattern of ngly1 in ZF tissue has not been reported before. Therefore, we focused on the anatomic regions defined by ngly1 expression in 6dpf larvae by RNA *in situ* hybridization. In the ZF larvae we observed significant expression of ngly1 mRNA in the nervous system (brain and the spinal cord) as hypothesized (Figures 4A–D), suggesting the importance of ngly1 in ZF nervous system development and maintenance. Furthermore, we demonstrated a distinct neural phenotype of the PNS morphology showing a loss of axon fibers; this finding supports a causative correlation of the NGLY1 nonsense genetic variant and the neurological phenotype in ZF and possibly in humans. Furthermore, we analyzed ZF locomotion behavior during startling by light/dark switch (Hamilton et al., 2017). The general motor test showed a reduction of movement in ngly1<sup>−/−</sup> larvae compared to ngly1<sup>+/+</sup> with light on (ideal conditions) similar to the phenotype reported in a rat model and in patients who universally present with delayed or lost motor milestones. In contrast, under stress stimulus, when lights were turned off, both groups showed an increase in swimming distance (Figure 6B) compared to lights-on phase (Figures 6A–C). However, ngly1<sup>−/−</sup> fish increased the swimming distance significantly compared to the control ngly1<sup>+/+</sup> fish. This result revealed a surprising observation, by which ngly1<sup>−/−</sup> fish have an impairment of the sensory regulation system—hypersensitivity to a stress trigger unlike the human patients and the rat model. Clinical case reports described pain insensitivity in patients with NGLY1 deficiency. In the case of the ZF model, the ngly1<sup>−/−</sup> larvae reacted to a pain trigger with an increase in swimming distance compared to ngly1<sup>+/+</sup> after treatment with acetic acid (pH 3.8). Since the increase in fish movement correlates to pain response (Curtright et al., 2015), we conclude that the ngly1<sup>−/−</sup> larvae displayed pain hypersensitivity reactions (Figures 6D,E) unlike the human patients and the rat model. This result provides additional evidence of unexpected phenotype of sensory hypersensitivity reaction in junior mutant ZF. Finally, our study shows an increase in the sensitivity of the mutant fish model to stress conditions. The mechanism of this phenomenon is yet to be determined.

We speculate that this phenomenon can be explained by the fact that the loss of peripheral axons is part of complex peripheral neuropathy, which can lead to hyperesthesia. Moreover, in order to add understanding about the mechanism underlying the paradoxical behavior of the mutant fish, it is important to further investigate the fish brain, that is, known to have a crucial role in responding and regulation of stress stimuli (Maldonado and De Jesus, 2021).

Another phenotype found in patients is MSK abnormalities that are described as one of the secondary phenotypes of NGLY1 deficiency and occur as a result of the loss of motor control (Need et al., 2012). In order to explore the MSK system phenotype and understand in which stage of the disease the muscle phenotype starts taking place, we studied the muscle morphology of 6dpf larvae. Similar to the human patients

![FIGURE 5](image-url)
(Panneman et al., 2020), we observed a significant difference in the muscle structure and morphology between the two ZF genotypes—ngly1<sup>−/−</sup> and ngly1<sup>+/+</sup>.

In general, the ZF model for NGLY1 deficiency does not display the anticipated dramatic phenotype such as global developmental delay (He et al., 2015). This might be explained by the fact that there is residual protein level in ngly1<sup>−/−</sup> fish. Also, another possible explanation is based on the study that has described an effective compensation mechanism in ZF as a response for nonsense-mediated mRNA decay (NMD) in case of premature stop codon in mRNA (El-Brolosy et al., 2019) as we created in our fish model.

In summary, we have created a ZF model for NGLY1 deficiency and characterized a neural phenotype of loss of axon fibers in PNS, a MSK phenotype of reduction in muscle mass and somite size, and a surprising sensory hypersensitivity to external triggers such as darkness and acetic acid-related pain. These multi-organ phenotypes of the mutant fish, suggest—a versatile function of NGLY1—and possible tissue specific mechanisms underlying the complexed phenotype of NGLY1 deficiency. Therefore, future studies should focus on several specific tissues in paralleled to studying the entire fish to accurately unveil the mechanism of this disorder. Further characterization of the brain, liver, skeleton, and eyes in adult fish might enable documentation of the natural history and the progression of the disease in comparison to the human phenotype. Our results further support previous reports showing that zebrafish are highly amenable to mutagenesis and germline transmission. Continued investigations of the NGLY1 deficiency ZF model might lead eventually to resolving the molecular details and mechanisms behind this rare systemic disorder and might serve as a platform for experimental treatments to rescue/medicate this devastating disease in humans.

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/supplementary material.
AUTHOR CONTRIBUTIONS

AM: generated the zebrafish model, designed and performed the experiments, designed the microscope analyses and behavioral analyses of the fish wrote the first draft. GN: participated in design of experiments, executed experiments, developed the microscope imaging and prepared figures. Participated in writing the original draft. CS: participated in design and generation of the fish model, Participated in writing the original draft. LK: supervised the experiments, wrote and edited the figures. Participated in writing the manuscript. AK: participated in the design and generation of the fish model. DK: supervision of fish experiments, wrote and edited the manuscript. TCF-Z: conceived and designed the project. Supervised the experiments. Wrote the paper.

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