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Biallelic sequence variants in *INTS1* in patients with developmental delays, cataracts, and craniofacial anomalies

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Abstract

The Integrator complex subunit 1 (*INTS1*) is a component of the integrator complex that comprises 14 subunits and associates with RPB1 to catalyze endonucleolytic cleavage of nascent snRNAs and assist RNA polymerase II in promoter-proximal pause-release on protein-coding genes. We present five patients, including two sib pairs, with biallelic sequence variants in *INTS1*. The patients manifested absent or severely limited speech, an abnormal gait, hypotonia and cataracts. Exome sequencing revealed biallelic variants in *INTS1* in all patients. One sib pair demonstrated a missense variant, p.(Arg77Cys), and a frameshift variant, p.(Arg1800Profs*20), another sib pair had a homozygous missense variant, p.(Pro1874Leu), and the fifth patient had a frameshift variant, p.(Leu1764Cysfs*16) and a missense variant, p.(Leu2164Pro). We also report additional clinical data on three previously described individuals with a homozygous, loss of function variant, p.(Ser1784*) in *INTS1* that shared cognitive delays, cataracts and dysmorphic features with these patients. Several of the variants affected the protein C-terminus and preliminary modeling showed that the p.(Pro1874Leu) and p.(Leu2164Pro) variants may interfere with *INTS1* helix folding. In view of the cataracts observed, we performed in-situ hybridization and demonstrated expression of *ints1* in the zebrafish eye. We used Clustered Regularly Interspaced Short Palindromic Repeats (CRISPR)/Cas9 to make larvae with biallelic insertion/deletion (indel) variants in *ints1*. The mutant larvae developed typically through gastrulation, but sections of the eye showed abnormal lens development. The distinctive phenotype associated with biallelic variants in *INTS1* points to dysfunction of the integrator complex as a mechanism for intellectual disability, eye defects and craniofacial anomalies.

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Introduction

The integrator complex, INT, is composed of 14 integrator subunits that associate with the C-terminal domain of Rbp1, the largest subunit of ribonucleic acid (RNA) polymerase II (RNAPII) [1, 2]. In *D. melanogaster*, the integrator complex catalyzes the endonucleolytic cleavage of nascent small nuclear RNAs (snRNAs) at their 3' ends and

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disruption of almost any integrator subunit causes snRNA misprocessing [1, 3]. There is still a paucity of information regarding the individual subunits of INT and their impact on human disease. The proteins show relatively little resemblance to other proteins, although the predicted motifs include domains that are suggestive of protein interactive surfaces, such as α -helical repeats (armadillo-like repeats (ARM), tetratricopeptide repeats (TPR), von Willebrand type A like (VWA), and HEAT domains) [1, 2, 4].

The largest integrator subunit in humans, INTS1, is composed of 2190 amino acids and has a molecular weight of 244 kDa [1]. *INTS1* is conserved amongst many metazoans, including humans, rat, mouse, zebrafish and fruit fly, suggesting a specialized and non-redundant function. Targeted disruption of *Ints1* in the mouse results in embryonic lethality at the early blastocyst stage for embryos with a pure genetic background [5]. The mutant mice have increased levels of unprocessed, primary U2 snRNA, suggestive of aberrant snRNA processing [5]. In *Oryzias latipes*, *ints1* is strongly expressed in the brain and detectable in many other tissues at 6 days post fertilization (dpf) [6]. The awabancha (awb) mutant contains a splice site loss of function variant in *ints1* (then called *kiaa1440*) and displays defective development of the thymic anlage and small eyes at 1.25 dpf and 2.5 dpf [6]. These mutants also had reduced liver and gall bladder size at 6 dpf, absence of the spleen at 7 dpf and died prior to hatching [6]. A zebrafish *ints1* mutant has not yet been reported.

Despite the murine and medaka models for *ints1* loss of function, a human phenotype associated with compromised *INTS1* function was only recently described. Three unrelated patients, all of Dutch ancestry, demonstrated moderate to severe intellectual disability, lack of speech development, impaired mobility, cataracts, and a consistent pattern of dysmorphic features comprising dolichocephaly, a prominent forehead, wide-spaced eyes, pectus deformities and irregularly implanted and overlapping toes [7]. Findings noted in single patients were hypoplasia of the cerebellar vermis, renal dysplasia, cleft lip and palate and congenital heart disease. All three patients were homozygous for the same nonsense variant in *INTS1*, c.[5351C>A];[5351C>A], predicting p.[(Ser1784*)];[(Ser1784)] (NM_001080453.2). The *INTS1* nonsense variant led to strong reduction of the mRNA expression in skin fibroblasts, consistent with a loss of function model of pathogenicity [7].

Herein we report an additional five children with biallelic sequence variants in *INTS1* who manifest a similar pattern of clinical features comprising developmental delays, juvenile cataracts and craniofacial findings. We add on detailed clinical information regarding the three previously described patients and review the phenotype. We also provide data from in-situ hybridization studies and a Clustered Regularly Interspaced Short Palindromic Repeats

(CRISPR)/Cas9 model of loss of *ints1* function in zebrafish (*Danio rerio*) that confirm the importance of this gene in eye development.

Material and Methods

Case reports

Clinical details and pedigree information were obtained from all patients as per standard clinical practice. The parents of patients 1 and 2 provided consent for publication of photographs that did not include recognizable facial features; the parents of patients 3 and 4 did not provide additional consent for publication of photographs. The parents of patient 5 provided consent for publication of facial photographs.

Exome sequencing

For patients 1 and 2, genomic DNA libraries were prepared (SeqCap EZ Human Exome Library v3.0; Roche Nimblegen, Madison, WI, USA) and sequenced on a HiSeq 2000 (Illumina, San Diego, CA, USA) for pair-end 100 cycles. Sequencing reads were aligned to the hg19 reference genome using the BurrowsWheeler Alignment tool (BWA v0.5.9) using the default parameters. Variants were subsequently called using Genome Analysis Toolkit v 1.3–21-gcb284ee [8, 9]. Variants were filtered with the recommendations listed in the Best Practice Variant Detection with GATK v3 with $\text{QD} < 2.0^1$, $\text{MQ} < 40.0^1$, $\text{FS} > 60.0^1$, $\text{HaplotypeScore} > 13.0^1$, $\text{MQRankSum} < -12.5^1$, $\text{ReadPosRankSum} < -8.0^1$ for SNPs and $\text{QD} < 2.0^1$, $\text{ReadPosRankSum} < -20.0^1$, $\text{InbreedingCoeff} < -0.8^1$, $\text{FS} > 200.0^1$ for Indels. The filtered list was then annotated with both ANNOVAR 2012Mar08 version [10] and snpEff v2.0.5 in GRCh37.64 [11]. The potential deleteriousness of novel sequence variants was assessed using PolyPhen-2 [12], Sorting Intolerant from Tolerant [13] and MutationTaster [14]. We utilized VarSifter [15] to explore the dataset and the *INTS1* sequence variants were verified using Sanger sequencing.

For patients three and four, WES sequencing was performed as part of the Care4Rare consortium as previously described [16]. Variants were filtered to those that were homozygous in both patients three and four, and had a minor allele frequency of < 0.005 in both of the 1000 Genomes or the exome variant server (EVS) databases, and absent or with a low frequency in an in-house database. The remaining variants were assessed using PolyPhen-2, SIFT, and MutationTaster. For patient five, whole exome sequencing was performed as a clinical test (GeneDx, Inc.) using the IDT xGen Exome Research Panel v1.0. Massively parallel (NextGen) sequencing was done on an Illumina

system with 100 bp or greater paired-end reads. Reads were aligned to human genome build GRCh37/UCSC hg19, and analyzed for sequence variants using a custom-developed analysis tool as previously described [17]. Exome sequencing of patient 6, 7, and 8 was previously described [7].

Clinical data and variants were submitted to the gene variant database at www.LOVD.nl/INTS1 (patient IDs 00164989–00164991).

Structure modeling

We used HHpred server [18] to identify structures of homologous proteins. HHpred identified homologs with low sequence identity of ~12% only for the C-terminal part of the protein (residues 1661–2184). We used MODELLER [19] to generate structural models for this part based on the identified homologs (PDB 2db0, 2vgl, and 4uqi).

In-situ hybridization

All animal experiments were performed under a protocol approved by the Institute for Animal Care and Use Committee (IACUC) at UCSF (AN108657–02C). EKW strain zebrafish were maintained at 28 °C with a 14 h (h) light/10 h dark cycle. We used primers to amplify the full-length *ints1* transcript from wildtype *Danio rerio* cDNA (XM_003198152.5; Table S1). Database searches showed no significant cross-hybridization between the amplified cDNA and other *Danio rerio* genes. The cDNA fragment was cloned into pCRII-TOPO (Invitrogen/Thermo-Scientific, San Francisco, CA) and cDNA orientation was verified by Sanger sequencing. Digoxigenin-labeled sense and antisense RNA probes were prepared as previously described (DIG RNA labeling kit; Roche, Indianapolis, IN) [20]. In-situ hybridization was performed according to standard protocols on wildtype, whole zebrafish embryos at 48 h post fertilization (hpf) [20].

CRISPR/Cas9 targeting of *ints1* in *Danio rerio*

We chose not to use antisense morpholinos due to the possibility of non-specific effects from this methodology on eye size and development. For CRISPR/Cas9, we used a modification of a previously described method [21]. We designed a CRISPR small guide (sg)RNA to target exon 8 of *ints1* that is upstream of an annotated domain of unknown function (DUF3677) spanning exons 8–10 of this gene (Table S2). We co-injected sgRNA and Cas9 protein into wildtype zebrafish eggs at the one cell stage and used Sanger sequencing at 24–48 hpf to assess for mutagenesis in batches of 10–20 larvae (for primers, see Table S3). FO larvae with insertion/deletion (indel) variants were raised and fin-clipped at 6–8 weeks of age in batches of 10–20 larvae for further genotyping with

Sanger sequencing to determine the exact indel variants present. We selected founders with small indels introducing a frameshift or eliminating a splice site in *ints1*, thus predicting premature protein truncation and nonsense-mediated decay (Table S4). These founders were outcrossed to wildtype *Danio rerio* and the F1 progeny were raised and genotyped to identify heterozygous fish with germline transmission of an indel variant. F1 fish that were heterozygous for indel variants in *ints1* were incrossed to obtain homozygous or compound heterozygous F2 larvae that had biallelic *ints1* variants predicting loss of function.

Larvae obtained from incrosses of heterozygous fish with *ints1* indel variants were examined for eye size and lens abnormalities at 3 to 7 dpf using light microscopy. Tails of larvae from incrosses were genotyped by Sanger sequencing and the corresponding heads of larvae with biallelic *ints1* variants were then sectioned at 6 dpf so that the eyes could be studied in more detail relating genotype to hematoxylin and eosin staining and immunohistochemistry. For hematoxylin and eosin staining, slides were immersed in hematoxylin (VWR International, Brisbane, CA) for three minutes, washed with tap water, dipped twice in Scott's water and then placed in 75% ethanol for 15 s. Slides were stained with eosin (VWR International, Brisbane, CA) for 15 s and then dipped in 75% ethanol prior to mounting. For immunohistochemistry, larval heads were fixed in 4% paraformaldehyde, washed with phosphate buffered saline (PBS) and placed in 25% (w:v) sucrose followed by 35% (w:v) sucrose. Cryosections were made according to standard techniques. Sections were dried overnight and stored at –80 °C. For staining, sections were washed three times in PBS with 0.1% Tween 20 (PBST) and incubated in blocking buffer containing 4% normal goat serum, 2% normal sheep serum and 0.2% Tween (all v-v) in PBS for 4 h at 4 °C. Slides were incubated with a 1 in 500 dilution of z1-1 primary antibody (Zebrafish International Resource Center) in blocking buffer overnight at 4 °C, washed in PBST, and then incubated overnight with a 1:500 dilution of antimouse IgG labeled with Texas red (M32017; Invitrogen, Carlsbad, CA) in blocking buffer as a secondary antibody at 4 °C. Sections were mounted with Abcam fluoroshield mounting medium containing DAPI (Abcam, Burlingame, CA) as a counterstain and imaged with a fluorescent microscope.

Lastly, in view of the lethality of the mouse model of *Ints1* loss of function at the stage of gastrulation, we used light microscopy to photograph the progeny of incrossed, heterozygous F1 adult fish at 1 h, 2 h, 4 h, and 24 hpf. After photographs were taken, DNA was extracted and the larvae were genotyped by Sanger sequencing, so that progress through gastrulation and eye morphology could be correlated to *ints1* genotype for wildtype, heterozygous, and compound heterozygous larvae.

Quantitative reverse transcriptase-polymerase chain reaction (qRT-PCR)

We used qRT-PCR to determine if there was altered expression of other INT complex genes between wildtype larvae and larvae from an incross of heterozygous, *ints1* indel fish. 1 ng cDNA was amplified using gene-specific primers (Table S5) at final concentrations of 2.5 μ M and Universal SybrGreen mastermix containing Rox (Roche, Pleasanton, CA) at a final concentration of 2.5 μ M. Reactions were run on a StepOne cycler and analyzed with StepOne™ Software and ExpressionSuite software (Thermo Fisher Scientific, San Francisco, CA) according to the $\Delta\Delta$ Ct method. We used *gapdh* (NM_001115114.1) or *eef1a1l1* (NM_131263.1) as internal control genes. All experiments were performed on three independent samples from wildtype larvae and larvae resulting from an incross of heterozygous, *ints1* indel fish. Each reaction was run in triplicate.

Results

Case reports

Clinical details from the eight cases are summarized in Table 1, Figs. 1–3, and Fig. S1. Further clinical information is available in Supplemental Data. Similar to the previously reported patients with biallelic *INTS1* variants [7], all patients demonstrated hypotonia and gait disturbances, with several patients taking independent steps at or after 2 years of age (patients 1 and 3). The mobility of the second patient deteriorated at 5 years of age after he was independently ambulant at 14 months of age, but he retained the ability to walk with assistance. All patients had significant developmental delays, with absent or minimal speech. Three were given a diagnosis of autism (patients 2, 3, and 4) and one carried a diagnosis of autism spectrum disorder (patient 5).

All patients developed cataracts between one and 3 years of age, as found in the first three reported patients, thus confirming the ocular involvement in patients with *INTS1* variants [7]. Three patients were diagnosed with myopia and the siblings from the first family had additional ophthalmological findings of microphthalmia, microcornea and colobomas (Table S6).

Four of the new patients shared dysmorphic features similar to those noted in the first three reported cases. Features described in more than half of the eight individuals with biallelic *INTS1* variants include frontal bossing, small palpebral fissures that are slanted, wide-spaced eyes, full cheeks, a broad nasal bridge and a broad or bulbous nasal tip, low-set ears, down-turned corners of the mouth and a

small chin (Table 1). Additional distinctive findings in the majority included a wide diastema between the upper incisor teeth, short neck, pectus deformities and broad halluces with overlapping toes (Table 1; Fig. 2). Other craniofacial features that were less commonly observed included brachycephaly (previous patients were reported to have dolichocephaly) [7], epicanthus and dysplastic ears (Fig. 1). Overall, the facial features and skeletal changes showed striking similarities to those of previously reported patients (Fig. 3) [7].

Brain magnetic resonance imaging (MRI) findings are summarized in Table S7 and ranged from normal (patients 7 and 8) to demonstrating non-specific abnormalities, with small anterior and hippocampal commissures with mildly reduced volume of the splenium of the corpus callosum and adenohypophysis in one child (patient 2) and hypoplasia of the inferior vermis of the cerebellum (patient 8; Table S7). Even among the three patients with the identical, homozygous nonsense variant, p.[(Ser1784*)], brain MRI abnormalities were not consistent, with only patient 6 showing hypoplasia of the inferior vermis of the cerebellum (Fig. S1). Patient 6 has also been treated for epilepsy.

The two siblings from the first family have had heights on the 5th and 1st centiles and all patients with the p.(Ser1784*) variant have significant growth retardation. The first sibling pair also demonstrated abnormal thumbs; one brother had both a multicystic, dysplastic kidney and mild prolapse of the anterior leaflet of the mitral valve. Although the second pair of siblings did not have renal ultrasound scans or echocardiograms, patient 5 had renal pelviectasis and patient 7 showed a horseshoe kidney. Patient 2 had distinctive skeletal findings, with loss of the thoracic kyphosis and lumbar lordosis, levocurvature of the lumbar spine, mild coxa valga, shortened tibiae and a leg length discrepancy. His hand radiographs demonstrated hypoplasia of the first metacarpal and phalanges of the right hand with absence of the trapezium. Congenital heart disease was only observed in patients 7 and 8 [7].

Exome sequencing

Exome sequencing showed that all patients had biallelic variants in *INTS1* (Table 2). The first sib pair were each compound heterozygous for a missense variant and a frameshift variant, c.[229C>T];[5398dup], predicting p.[(Arg77Cys)];[(Arg1800Profs*20)] in *INTS1* (NM_001080453.2). The frameshift variant was predicted to be deleterious, but the missense variant was present in 0.025% of individuals in a control database (ExAC Browser, Table 2) and has been seen in the homozygous state in one individual (<http://gnomad.broadinstitute.org/gene/ENSG00000164880>; Table 2), suggesting that it is likely

Table 1 Clinical features of patients with biallelic *INTS1* sequence variants

	Patient 1	Patient 2	Patient 3	Patient 4	Patient 5	Patient 6 ID9065364 [7]	Patient 7 ID5342723 [7]	Patient 8 ID8330971	Total
Growth parameters (gender, age) SD	(m, 14 years)	(m, 9 years)	(m, 11 years)	(f, 6 years)	(m, 21 months)	(f, 10 years)	(m, 19 years)	(m, 6 years)	
Height	Z -2.45	Z -1.44	Z - 0.01	Z -0.63	Z -3.36	-4.3 SD	-5.5 SD	-3.8 SD	
Weight	Z 0.39	Z -1.16	Z -.59	Z -1.19	Z -0.83	-4.0 SD	-5.0 SD	-1.8 SD	
Head circumference	1.21 SD	Z- 0.59	+5 SD		Z -1.41	0.5 SD	-2.5 SD	-2.3 SD	
Central nervous system									
Cognitive delays	+	+	+	+	+	+	+	+	8/8
Absent or severely reduced speech	+	+	+	+	+	+	+	+	8/8
Abnormal gait	+	+	+	+	+	+	+	+	8/8
Hypotonia	+	+	+	+	+	+	+	+	8/8
Autism	-	+	+	+	-	-	-	-	3/8
Epilepsy	-	-	-	-	-	+	-	-	1/8
Eyes									
Iris, retinal colobomas	+	-	-	-	-	-	-	-	1/8
Microphthalmia/ microcornea	+	+	-	-	-	-	-	-	2/8
Cataracts/glaucoma	+	+	+	+	+	+	+	+	8/8
Strabismus	-	-	+	-	-	+	+	+	4/8
Myopia/astigmatism	+	-	+	+	-	-	+	+	5/8
Craniofacial									
Frontal bossing	-	-	+	+	-	+	+	+	5/8
Prominent metopic ridge	-	-	-	-	-	-	+	+	2/8
Abnormal head shape	+	+	-	-	-	-	+	+	4/8
High frontal hairline	-	-	-	-	-	+	+	+	3/8
Prominent orbital ridge	-	-	-	-	-	+	+	+	3/8
Horizontal eyebrows	+	+	-	-	-	+	+	+	5/8
Deep-set eyes	NA	NA	-	-	-	+	-	+	2/6
Small palpebral fissures	+	+	-	-	-	+	+	+	5/8
Slanted palpebral fissures	+	+	+	-	-	+	+	+	6/8
Wide-spaced eyes	+	+	+	+	-	+	+	+	7/8
Epicanthus	+	+	-	-	-	-	+	+	4/8
Wide nasal bridge	+	+	+	+	-	+	+	+	7/8
Broad/bulbous nasal tip	+	+	-	-	+	+	+	+	6/8
Short nose	-	+	-	-	-	-	-	+	2/8
Full cheeks	-	-	+	-	+	+	+	+	5/8
Low-set ears	+	+	-	-	-	+	+	+	5/8
Dysplastic/prominent ears	+	+	-	-	-	+/-	+	+	4/8
Small, overfolded helices	-	-	-	-	-	+	-	+	2/8
Deep and long philtrum	+	+	-	-	-	-	-	-	2/8
Wide philtrum	-	-	-	-	-	+	+	+	3/8
Downturned corners of the mouth	+	+	+	+	-	-	+	+	6/8
Broad mouth	-	-	-	-	-	+	+	+	3/8

Table 1 (continued)

	Patient 1	Patient 2	Patient 3	Patient 4	Patient 5	Patient 6 ID9065364 [7]	Patient 7 ID5342723 [7]	Patient 8 ID8330971	Total
Microretrognathia	+	+	+	+	–	–	+	–	5/8
Short neck	+	+	–	–	–	+	+	+	5/8
Cleft lip and palate	–	–	–	–	–	–	+	–	1/8
Diastasis upper incisors/ frenula/abnormal dentition	+	+	+	–	–	+	+	+	6/8
Other findings									
Renal malformation	–	+	–	–	+	–	+	–	3/8
Pectus deformity	+	+	–	–	–	+	+	+	5/8
Thumb abnormalities	+	+	–	–	–	–	–	–	2/8
Overlapping/irregular toes	+	+	–	–	–	+	+	+	5/8
Scoliosis	–	+	–	–	–	–	–	+	2/8
Hip dysplasia	–	–	–	–	–	–	+	+	2/8
Congenital heart disease	–	–	–	–	–	–	+	+	2/8
Severe feeding problems/tube feeding	–	–	–	–	–	+	+	+	3/8
Short fingers/broad terminal phalanges	–	–	–	–	–	+	+	+	3/8
Short sternum (other than pectus carinatum)	–	–	–	–	–	+	+	+	3/8

to be a hypomorphic variant or associated with incomplete penetrance. It is possible that p.(Arg77Cys) may not lead to clinical manifestations when present in the homozygous state, but that it could have sufficient effect on the protein to cause disease when inherited with a more severe sequence variant on the second allele. Protein structure data for this variant is unavailable. It is also noteworthy that the homozygous individual in gnomAD is of Hispanic ethnicity, the same ethnicity as the siblings with this variant, and that the frequency of the variant is highest in the Hispanic population at 0.13% (<http://gnomad.broadinstitute.org/variant/7-1542657-G-A>). However, the clinical findings in these patients are similar to those in others with *INTS1* variants and we still consider this the causative gene. Both of the siblings from the second family were homozygous for a missense variant c.[5621C>T];[5621C>T], predicting p. [(Pro1874Leu)];[(Pro1874Leu)] that was also predicted to be deleterious and was absent from human genetic variation databases (Table 2). The fifth child also had biallelic *INTS1* variants—c.[5290del];[6491T>C], predicting p. [(Leu1764-Cysfs*16)];[(Leu2164Pro)]. This child's healthy brother carried the c.[6491T>C] variant only. Splicing prediction programs did not demonstrate altered splicing with any of the variants (data not shown). Several of the variants would be expected to result in loss of function of the protein, consistent with the reported homozygosity for a nonsense variant in *INTS1* in the first three reported patients [7].

Modeling studies of the C-terminal end of the protein showed that the missense variants p.(Pro1874Leu) and p.(Leu2164Pro) may interfere with helix folding (Fig. S2). Specifically, the p.(Pro1874Leu) variant is likely to disrupt the helix because it is located immediately after the helix in our model. The substitution of proline with leucine, which has a high helix propensity [22] cancels the helix break and disrupts the protein fold. The p.(Leu2164Pro) variant is expected to have an opposite effect. Leu2164 is located in the middle of the helix; the change to Pro breaks the helix. Protein structure data to model the p.(Arg77Cys) variant was not available.

In-situ hybridization

In-situ hybridization demonstrated expression of *ints1* in the developing zebrafish eye and brain at 48 hpf (Fig. S3). This is consistent with the small eyes observed in the awb medaka model of *ints1* loss of function and the eye involvement observed in the patients reported here.

CRISPR/Cas9 model of *ints1* loss of function

We successfully targeted the single, *ints1* transcript in *Danio rerio* and generated fish that were heterozygous for two separate indel variants predicting loss of function—c.1020delinsGG, predicting p.(Met341Aspfs*19), and

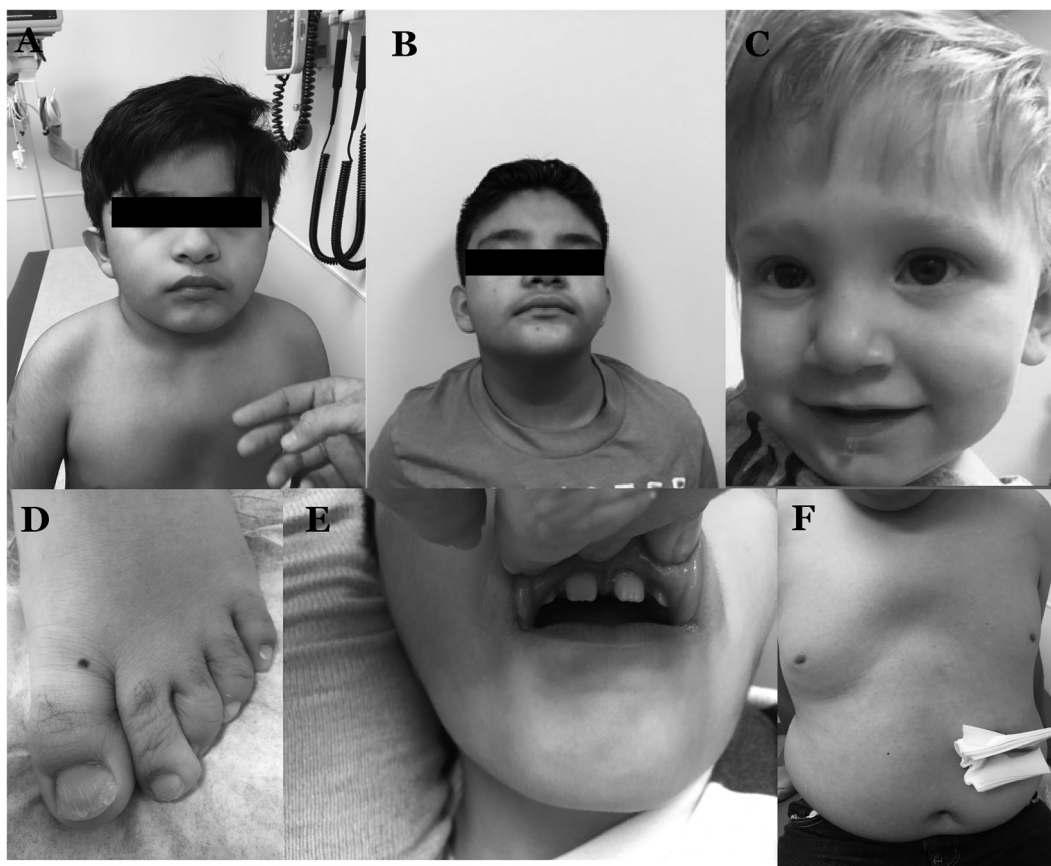


Fig. 1 Biallelic variants in *INTS1* are associated with facial dysmorphic features. **a** Front view of patient 2, the younger sibling from family 1, demonstrating a broad nasal tip, downturned corners of the mouth, low-set ears and a pectus deformity. **b** Front view of patient 1, the older sibling from family 1, demonstrating downturned corners of the mouth and low-set ears. **c** Front view of patient 5, demonstrating

frontal bossing, upslanting palpebral fissures and a wide nasal tip. **d** Left foot from patient 1, demonstrating overlapping toes with clinodactyly. **e** Oral region of patient 1, demonstrating a wide diastema between the upper two incisor teeth. **f** Chest and abdomen of patient 1, demonstrating a pectus deformity and G-tube insertion site

c.1020_1021insG, predicting p.(Met341Hisfs*18) (XM_003198152.4; Table S4; Fig. S4). After incrossing a female that was heterozygous for c.1020_1021insG in *ints1* and a male that was heterozygous for *ints1* c.1020delinsGG, genotyping at 6 dpf revealed that inheritance of the variants was present in expected Mendelian ratios, with 5/30 (16.7%) larvae wildtype for *ints1*, 8/30 (26.7%) larvae heterozygous for the c.1020insG variant in *ints1*, 10/30 (33.3%) larvae heterozygous for the c.1020delinsGG variant in *ints1*, and 7/30 (23.3%) larvae that were compound heterozygous for both the c.1020insG and c.1020delinsGG variants in *ints1*. We did not detect any external abnormalities in the larvae that were compound heterozygous for the c.1020insG and c.1020delinsGG variants in *ints1* at up to 6 dpf with light microscopy and development through gastrulation appeared to progress typically (Figs. S5 and S6).

Sectioning and staining of the eyes of larvae at 6 dpf showed grossly normal eye formation in larvae with biallelic *ints1* variants (Fig. 4g–i) compared to wildtype (Fig. 4a–c) and

heterozygous larvae (Fig. 4d–f). However, there was a subtle increase in lens thickness that was observed in the compound heterozygous larvae (Fig. 4l, m) compared to wildtype larvae (Fig. 4j) that was also detectable as an increase in staining with z1-1 antibody that labels lens fibers.

Quantitative reverse transcriptase-polymerase chain reaction (qRT-PCR)

We used qRT-PCR at 3 dpf to examine the effects of reduced *ints1* expression on genes from the INT complex on RNA obtained from wildtype larvae and RNA obtained from an incross of fish that were heterozygous for an *ints1* indel variant. A limitation of this experiment was that the incross would be expected to result in only 25% of larvae that were compound heterozygous for both *ints1* indel variants. We were unable to phenotype the larvae from the heterozygous indel incross using a phenotypic marker that was visible by light microscopy due to their typical external appearance at 3 dpf; we could also not genotype and extract



Fig. 2 Phenotype findings in patients homozygous for p.[(Ser1784*)] in *INTS1* Upper row: patient 6 at the age of 6 years; Middle row: patient 7 at the age of 19 years; Lower row: patient 8 at the age of 2

years (first two pictures) and 6 years (3rd picture). All patients share the same homozygous p.[(Ser1784*)];[(Ser1784*)] variant, but are unrelated

RNA from single larvae at this developmental stage due to larval size. We performed qRT-PCR for *ints1*, *ints3* (NM_001044932.1), *ints8* (NM_001020756.2), *ints9* (NM_001077270.1), *ints11* (NM_001020621.1) and *ints12* (NM_001002161.3) in control and incrossed larvae. The results showed significant downregulation of *ints1* as expected ($p = 0.027$; Fig. S7), but also downregulation of *ints9* ($p = 0.032$), *ints11* ($p = 0.069$), *ints8* ($p = 0.086$), *ints3* ($p = 0.163$), and *ints12* ($p = 0.176$). These findings suggest that deficiency of *ints1* could lead to dysregulation of additional genes in the INT complex. Interestingly, the results also suggested upregulation of *ints8*, although this was not statistically significant ($p = 0.086$) (Fig. S7).

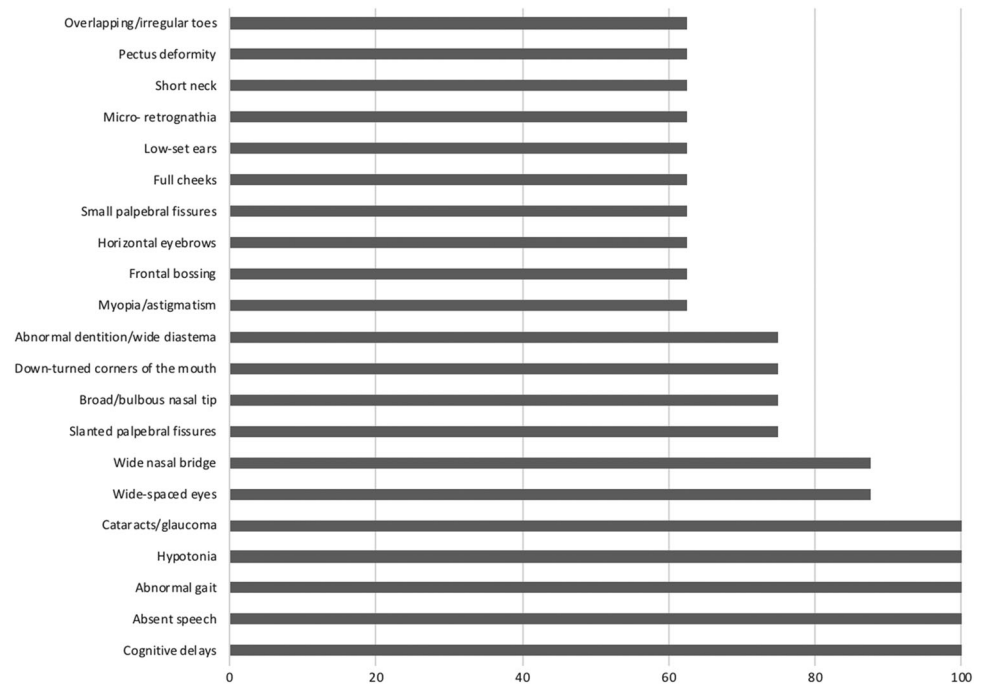
Discussion

Herein we present two sibling pairs and a fifth patient with clinical features of autism and severe speech delay, hypotonia and gait disturbances, cataracts and dysmorphic

craniofacial findings that included frontal bossing, small and slanted palpebral fissures, wide-spaced eyes, a broad nasal bridge with a broad or bulbous nasal tip, low-set ears, down-turned corners of the mouth, a wide diastema between the upper incisor teeth and a small chin. Other distinctive findings included stature at or below the 5th centile, short neck, pectus deformities and abnormal thumbs and toes. All patients had biallelic variants in *INTS1*. The overlap of clinical findings with previously reported patients who were homozygous for the p.(Ser1874*) variant in *INTS1* implicate this gene in the pathogenesis of the phenotypic features of these patients. Our data suggest that the clinical findings associated with *INTS1* variants may also include the eye findings of microphthalmia, microcornea and colobomas.

To date, there has been no clear phenotype-genotype correlation for *INTS1* variants and the first sib pair (patients 1 and 2) that have a missense variant of unclear pathogenicity, p.(Arg77Cys), share similar clinical findings to the three previously described patients (patients 6, 7, and 8) who are homozygous for the nonsense variant, p.(Ser1874*)

Fig. 3 Key phenotypic features associated with biallelic variants in *INTS1*. The frequency of the features is shown on the X-axis (%) and the features are listed on the Y-axis



(Table 2). However, the nonsense variant is also predicted to create a new splice donor site that is stronger than the endogenous splice donor site (Fig. S8) that would delete 30 nucleotides if used to result in an in-frame deletion, p.(Ser1784_Ser1793del), thus this variant may also be a hypomorphic allele (data not shown). The missense variants in patients 3 and 4, p.(Pro1874Leu), and in patient 5, p.(Leu2164Pro), were predicted to interfere with helix folding (Fig. S2) and thus may affect the ability of INTS1 to bind to other members of the integrator complex.

We used whole mount *in-situ* hybridization to show that the *ints1* gene is expressed in the developing *Danio rerio* eye (Fig. S3) and targeted the *ints1* gene using CRISPR/Cas9 in zebrafish to produce larvae with biallelic indel variants affecting *ints1* that predicted loss of function for this gene. Examination of the external appearance of larvae with biallelic *ints1* variants by light microscopy revealed the fish passed through gastrulation normally with no obvious differences in external appearance (Figs. S5 and S6). However, staining of sections of the eyes with hematoxylin and eosin at 6 dpf showed slightly thickened lenses in the larvae that were genotyped to show biallelic *ints1* variants (Fig. 4). This finding was also present with immunohistochemistry using z1-1 antibody that is expressed in lens fibers, when a broader band of z1-1 staining in larvae that were compound heterozygous for *ints1* indels was observed compared to wildtype larvae at 6 dpf. The increased staining is in accordance with early markers of adult onset cataracts that include deeper staining of fiber cells, scattered cortical opacities and fiber cell disruption [23]. These findings, together with the cataracts noted in all patients with biallelic

INTS1 variants reported to date, strongly support the involvement of this gene in eye and lens development. However, the relationship between the lens thickening and the development of cataracts is not known and the zebrafish model has limited utility to examine the other features found in patients with biallelic *INTS1* variants, such as developmental delays and dysmorphic features.

In the mouse, the *Ints1* protein has 2222 amino acids and predominantly localizes to the nucleus, with marginal cytoplasmic expression [5]. Homozygous null embryos ($-/-$) for *Ints1* demonstrated abnormal blastocyst morphology as early as embryonic day (E)3.5 [5]. In a mixed background, depletion of *Ints1* in homozygous null embryos is lethal at the morula stage [5]. After 8–10 backcrosses with the ICR strain, embryos were produced in the expected Mendelian ratios, but the blastocysts of homozygous null embryos did not expand and hatching from the zona pellucida and implantation did not occur, demonstrating developmental arrest. In vitro culture of homozygous null embryos showed increased apoptosis of the inner cell matrix and caspase-3 and caspase-7 activation [5]. Similar findings with abnormal gastrulation were not observed in our *Danio rerio* model of reduced *ints1* function, with compound heterozygous larvae passing through gastrulation with typical morphology, suggesting functional redundancy for this gene in zebrafish compared to mouse.

To date, there are few details on how *ints1* interacts with the other integrator complex subunits. INTS1 has been shown to interact with INTS12 which is involved in small nuclear (sn)RNA splicing and regulation of translation [3, 24]. In the mouse, *Ints1* may directly bind to *Ints2* [5, 25]

Table 2 Biallelic sequence variants in *INTS1* (NM_001080453.2)

Genomic Alteration (NM_001080453)	Nucleotide Alteration	Amino acid Alteration	Zygosity	Effects on Protein Domain	PolyPhen-2	Mutation Taster	1000G ^a	ExAC Browser ^b	GnomAD ^c	Conser- vation	Inter-pretation
Patients 1 and 2											
chr7:1542657G>A/	c.229C>T	p.(Arg77Cys)	Het. ^d	Unknown	PD ^e ; 0.990	DC ^f ; 1.0	-	30/120274; 0.02494	57/276876; 0.0002059	Mm ^g , Tr ^h , Dr ⁱ , Dm ^j , Ce ^k , Xt ^l	Uncertain
chr7:1515687_1515688insG	c.5398dupC	p.(Arg1800Profs*20)	Het.	-	-	DC; 1.0	-	-	1/138690; 7.21e-6	-	Deleterious
Patients 3 and 4											
chr7:1514356G>A	c.5621C>T	p.(Pro1874Leu)	HZ ^m	Abnormal helical folding	PD; 1.0	DC; 0.999	-	-	-	Mm, Gg ⁿ , Tr, Dr, Dm, Xt	Deleterious
chr7:1519188C>T	c.4207G>A	p.(Val1403Met)	HZ	-	Poss. D ^o ; 0.666	PMF ^p ; 0.999	MAF ^q 0.001597	44/17398; 0.002529	214/176252; 0.001214 1HZ	Mm, Xt	Not deleterious
Patient 5											
chr7: 1522197G>A	c.3688C>T	p.(Arg1230Cys)	Het. ^(pat)	Unknown	B ^f ; 0.34	DC; 0.999	-	3/64,838 0.00004627	7/274402; 0.0026	Mm, Dr, Dm, Ce, Xt	Not deleterious
chr7: 1515953_1515953delG	c.5290delC	p.(Leu1764Cysfs*16)	Het. ^(pat)	-	-	DC; 1.0	-	-	-	-	Deleterious
chr7:1510295A>G	c.6491T>C	p.(Leu2164Pro)	Het. ^(mat)	Abnormal helical folding	PD; 0.824	DC; 0.999	-	-	-	Mm, Gg ⁿ , Dr, Xt	Deleterious

^a1000Genomes = <http://www.1000genomes.org>

^bExAC Browser = <http://exac.broadinstitute.org/>

^cGnomAD = <http://gnomad.broadinstitute.org/>

^dHet. = heterozygous

^ePD = Probably damaging

^fDC = disease causing

^gMm = *Mus musculus*

^hTr = *Takefugu rubripes*

ⁱDr = *Danio rerio*

^jDm = *Drosophila melanogaster*

^kCe = *Caenorhabditis elegans*

^lXt = *Xenopus tropicalis*

^mHZ = homozygous

ⁿGg = *Gallus gallus*

^oD = possibly damaging

^pPM = polymorphism

^qMAF = minor allele frequency

^rB = benign

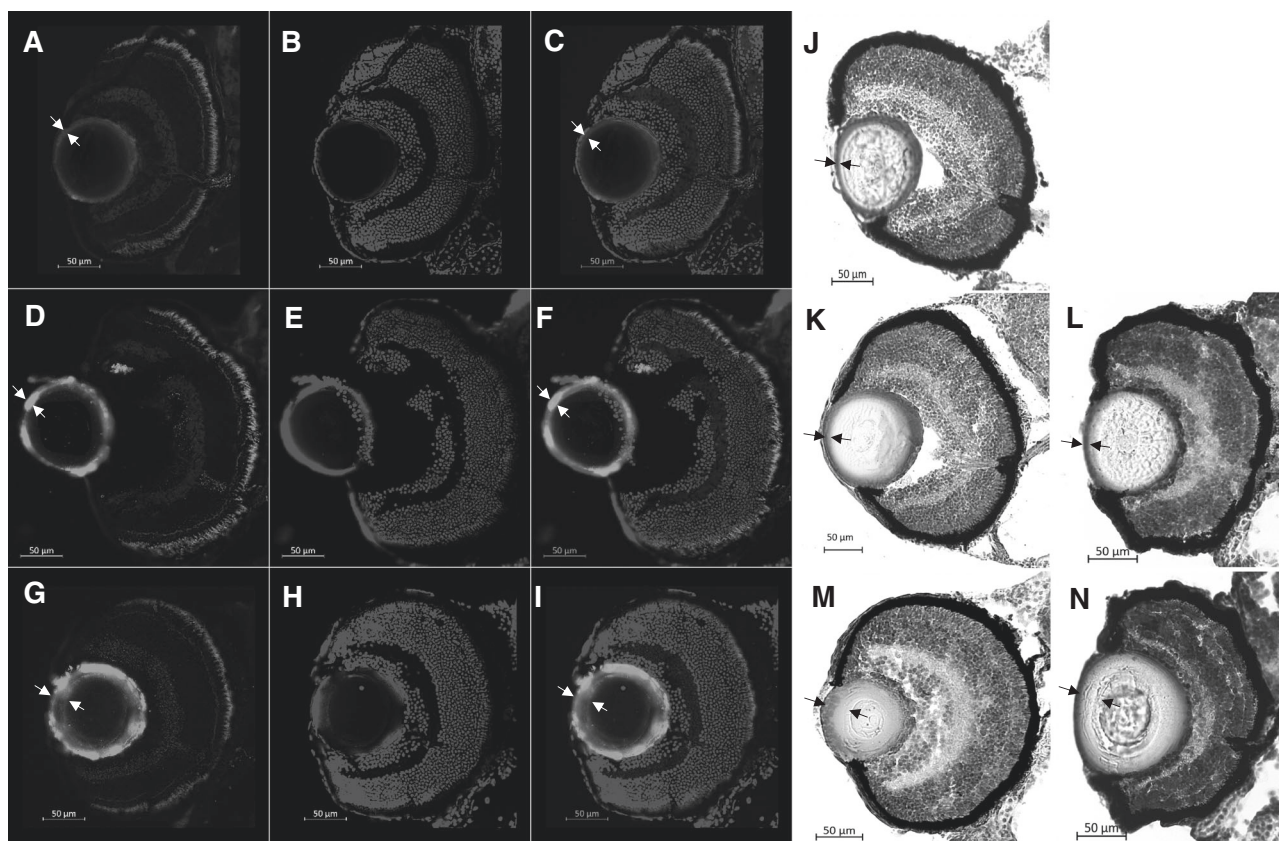


Fig. 4 Compound heterozygous *ints1* indel mutant larvae show abnormal lens morphology at 6 days post fertilization. Immunohistochemistry performed with zl-1 antibody and visualized with Texas Red demonstrates a thickened and less transparent lens appearance with reduced zl-1 staining in *ints1* compound heterozygous indel larvae compared to wildtype controls. **a–c** Larva wildtype for *ints1* (XM_003198152.4), demonstrating normal lens formation. **d–f** Larva heterozygous for *ints1* targeted allele. **g–i** Larva that are compound heterozygous for variants c.[1020delinsGG];[c.1020_1021insG],

predicting p.[(Met341Aspfs*19)];[(Met341Hisfs*18)] in *ints1*. The outer layer of the lenses is thickened and exhibits increased staining with zl-1 antibody that binds to lens fibers. Staining with hematoxylin and eosin revealed mildly thickened lenses in larvae with biallelic *ints1* variants. **j**. Larva wildtype for *ints1*. **k–l** Larva heterozygous for *ints1* targeted allele. **m–n** Larvae that are compound heterozygous for c. [1020delinsGG];[c.1020_1021insG], predicting p.[(Met341Aspfs*19)]; [(Met341Hisfs*18)] in *ints1*, showing mild thickening of the outer layer of the ocular lens

and these proteins act as scaffolding for the assembly of the entire integrator complex. Disruption to *Ints1* could thus compromise the function of the entire integrator complex [5]. *Ints11* is a predominantly nuclear protein with sequence similarity to the putative β -lactamase catalytic domain (RNA-specific endonuclease) [26]. *INTS9* and *INTS11* are paralogues of the cleavage and polyadenylation specificity factor subunits CPSF100 and CPSF73, respectively, which form the endonuclease factor responsible for cleavage of pre-mRNA at the polyA site [7]. Knockdown of *INTS1* and *INTS11* specifically abrogated the response of immediate early genes (IEGs) to epidermal growth factor (EGF) stimulation in human cells [27].

We performed qRT-PCR to determine if the genes for the remaining INT subunits were dysregulated in the larvae resulting from an incross of *ints1* heterozygous indel fish compared to an incross of wildtype fish. We found significant downregulation of *ints1* as expected, but also downregulation of *ints9*, *ints11*, *ints3*, and *ints12*,

suggesting that reduced *ints1* expression could affect the expression of other INT genes. In contrast, *ints8* may be upregulated in the incrossed larvae (Fig. S7). This is intriguing in view of the recent demonstration of biallelic variants in *INTS8* in patients with intellectual disability and neuronal migration defects [7]. In HEK293T cells stably expressing wildtype or mutant c.2917_2925(Δ EVL) *INTS8*, FLAG-*INTS8*- Δ EVL eluates contained nearly undetectable amounts of *INTS1*, *INTS12* and *RPB1*, reduced levels of associated *INTS4*, *INTS9*, and *INTS11*, but similar levels of *INTS5* and *INTS3*, compared to wildtype FLAG-*INTS8*, and thus the Δ EVL variant may impair the ability of *INTS8* to associate with members of the integrator complex, including *INTS1* [7].

In summary, we present five patients with biallelic variants in *INTS1* and provide further clinical details for three previously reported patients. All patients demonstrated severely compromised expressive speech, an abnormal gait, cataracts, and distinctive craniofacial dysmorphic features

that included frontal bossing, wide-spaced eyes, a broad nasal bridge and nasal tip and downturned corners of the mouth. We were successfully able to target the zebrafish *ints1* gene using CRISPR/Cas9. Larvae that were compound heterozygotes for deleterious *ints1* variants showed no obvious external defects, but examination of their eyes revealed abnormal lens histology with thickening of the lenses and increased staining for *zl-1* in lens fibers at 6 dpf. Our work provides additional data on the clinical features associated with biallelic *INTS1* variants in humans and confirms abnormal lens development in a zebrafish model of reduced *ints1* expression.

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Compliance with ethical standards

Conflict of interest R.E.S. is an employee of GeneDx, Inc., a wholly owned subsidiary of OPKO Health, Inc. The remaining authors declare that they have no conflict of interest.

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