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ORIGINAL RESEARCH ARTICLE



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Zebrafish expression reporters and mutants reveal that the IgSF cell adhesion molecule Dscamb is required for feeding and survival

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ABSTRACT

Down syndrome cell adhesion molecules (DSCAMs) are broadly expressed in nervous systems and play conserved roles in programmed cell death, neuronal migration, axon guidance, neurite branching and spacing, and synaptic targeting. However, DSCAMs appear to have distinct functions in different vertebrate animals, and little is known about their functions outside the retina. We leveraged the genetic tractability and optical accessibility of larval zebrafish to investigate the expression and function of a DSCAM family member, *dscamb*. Using targeted genome editing to create transgenic reporters and loss-of-function mutant alleles, we discovered that *dscamb* is expressed broadly throughout the brain, spinal cord, and peripheral nervous system, but is not required for overall structural organization of the brain. Despite the absence of obvious anatomical defects, homozygous *dscamb* mutants were deficient in their ability to ingest food and rarely survived to adulthood. Thus, we have discovered a novel function for *dscamb* in feeding behavior. The mutant and transgenic lines generated in these studies will provide valuable tools for identifying the molecular and cellular bases of these behaviors. ARTICLE HISTORY

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Introduction

Establishing the circuits underlying perception, movement, and cognition requires the precise arrangement of intricate neuronal arbors and formation of specific synaptic contacts. Creating such complexity and precision requires diverse molecular signals functioning at the local level (Langley, 1895; Lawrence Zipursky & Sanes, 2010; Sperry, 1963). Many diseases of the nervous system manifest in debilitating neurodegenerative and psychiatric symptoms, despite the absence of major brain malformations, suggesting that molecular signals governing the local assembly of neuronal circuits may play critical roles in these conditions.

The immunoglobulin superfamily (IgSF) of cell adhesion molecules (CAMs) is one of the largest protein families in vertebrates. Their diversity, broad range of binding interactions, and abundant expression in the nervous system (Vaughn & Bjorkman, 1996), make IgSF CAMs ideal candidates for organizing neuronal circuits. One family of IgSF proteins, the Down syndrome cell adhesion molecules (DSCAMs), has attracted attention due to *DSCAM*'s location on the Down syndrome critical region (DSCR) of chromosome 21 (Yamakawa *et al.*, 1998), trisomy of which causes the disabilities associated with Down syndrome (Delabar *et al.*, 1993; Korenberg *et al.*, 1994; Korenberg, Bradley, & Disteche, 1992). DSCAMs are single-pass transmembrane

proteins, with a conserved extracellular arrangement of Ig and fibronectin repeats (Schmucker & Chen, 2009). Drosophila dscam1 has a unique mechanism of alternative splicing, which generates up to 19,008 different extracellular domains that engage in precise homophilic binding (Schmucker et al., 2000). This extreme diversity allows Drosophila DSCAMs to serve as specificity molecules mediating neurite self-recognition (Hattori, Millard, Wojtowicz, & Zipursky, 2008; Millard & Zipursky, 2008; Schmucker, 2007). Vertebrate DSCAMs, Dscam and Dscam-like 1 (Dscaml1), are not extensively alternatively spliced, but are broadly expressed in the central nervous system (CNS) and implicated in many aspects of neuronal development, including axon guidance, programmed cell death, cell migration, neurite branching and spacing, synaptic targeting, and synapse formation (Montesinos, 2014).

The functions of vertebrate DSCAMs have been best studied in the well-organized circuitry of the retina. In the mouse retina, DSCAM and DSCAML1 are expressed in non-overlapping subtypes of bipolar cells (BCs), amacrine cells (ACs), and retinal ganglion cells (RGCs). Knockout of either gene causes specific subtypes of these cells to aggregate and fasciculate, disrupting cell spacing (Fuerst *et al.*, 2009; Fuerst, Koizumi, Masland, & Burgess, 2008). In these mouse mutants, neurite branches fasciculate with other

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branches of the same neuron, as well as neighbors of the same subtype, indicating that DSCAMs are required for both self-avoidance and homotypic branch spacing. Similar to their murine counterparts, chick Dscams, and the closelyrelated Sidekick (Sdk) and Contactin (Cntn) subfamilies, are expressed in distinct retinal subtypes (Yamagata & Sanes, 2008, 2012; Yamagata, Weiner, & Sanes, 2002). Each IgSF protein is enriched in a particular sublamina of the chick inner plexiform layer (IPL) due to mutual innervation from synaptically connected BCs and RGCs, and are required for establishing those specific connections (Yamagata & Sanes, 2008, 2012; Yamagata et al., 2002). In contrast, mouse DSCAMs are broadly localized throughout the IPL (de Andrade, Kunzelman, Merrill, & Fuerst, 2014; de Andrade, Long, Fleming, Li, & Fuerst, 2014; Fuerst et al., 2009) and play a limited role in lamination that varies with genetic background (Fuerst, Bruce, Rounds, Erskine, & Burgess, 2012; Fuerst, Harris, Johnson, & Burgess, 2010; Li et al., 2015). Additional studies are required to compare the similarities and differences of DSCAM functions across vertebrate taxa.

Although DSCAMs have been best studied in the retina, they are also expressed throughout the brain and spinal cord, a feature conserved among vertebrates (Agarwala et al., 2001; Barlow, Micales, Chen, Lyons, & Korenberg, 2002; Morales Diaz, 2014; Yamakawa et al., 1998; Yimlamai, Konnikova, Moss, & Jay, 2005), but few studies have investigated roles for DSCAMs outside the retina. Analyzing other DSCAM-expressing regions could identify new roles for these proteins. Zebrafish are a promising model for investigating DSCAM functions; their external fertilization and optical transparency facilitates observation of development of the entire brain in living embryos and larvae. The zebrafish genome has three DSCAM family members: dscama, dscamb, and dscaml1. Although dscama expression and function has been investigated (Yimlamai et al., 2005), the expression of *dscamb* has only been characterized in the retina (Sun, Galicia, & Stenkamp, 2018), and its function has yet to be interrogated. In this study, we leveraged the genetic tractability of zebrafish to investigate the expression and function of dscamb. To characterize dscamb expression, we created reporters using CRISPR/Cas9-targeted enhancer traps. To determine its function, we generated dscamb lossof-function mutant lines with the TALEN targeting system. Using these tools, coupled with cellular and behavioral analyses, we identify a critical new role for dscamb in feeding and survival.

Methods

Zebrafish

Zebrafish (*Danio rerio*) were grown at $28.5 \,^{\circ}$ C on a $14 \,\text{h}/10 \,\text{h}$ light/dark cycle. Embryos were raised at $28.5 \,^{\circ}$ C in embryo water (0.3 g/L Instant Ocean Salt, 0.1% methylene blue). For live confocal imaging, embryos were treated with phenyl-thiourea (PTU) at 24 hpf to block pigmentation. All experimental procedures were approved by the Chancellor's Animal Research Care Committee at UCLA.

Supplemental Table 1 summarizes the zebrafish lines used for this study.

TALEN cloning and targeted mutagenesis

For each mutant target site in the *dscamb* locus, two 20 bp TALEN binding sites were selected (exon 1: 5'-TAG-CTTGGGGATTGAACGCA-3', 5'-TGGAGAAAGAGAAAT-GCCAA-3'; exon 2: 5'-TCTACGTTCCAGCTTATATT-3', 5'-TGGAGAACACTACCTCTTGC-3'). A restriction enzyme site between each TALEN pair was used for genotyping by RFLP. TALEN constructs were cloned using Golden Gate assembly (Cermak *et al.*, 2011) and an accompanying plasmid kit from Addgene (Addgene Kit #100000024). TALEN mRNAs were injected into 1-cell embryos to generate stable mutant lines. See Supplemental Methods for details of mutant generation and genotyping.

BAC cloning and transgenesis

The BAC(*dscamb*:Gal4) line was created by modifying Ch73-102M15, a 40.8 kb BAC containing the first exon of *dscamb*, 22.3 kb upstream of this exon, and 17.8 kb of the first intron. Using *in vitro* homologous recombination, a Gal4FF-polyA-Kan cassette was inserted in place of the *dscamb* start codon, according to a previously published protocol (Suster, Abe, Schouw, & Kawakami, 2011). See Supplemental Methods for details.

Enhancer trap creation and transgenesis

Two gRNA target sites upstream of the *dscamb* transcriptional initiation site were chosen for insertion. gRNAs were generated as previously described (Talbot & Amacher, 2014). A zebrafish codon-optimized Cas9 construct (pCS2-nCas9n, Addgene.org plasmid #47929), flanked by nuclear localization signals was used for mutagenesis (Jao, Wente, & Chen, 2013). See Supplemental Methods for details on gRNA and Cas9 mRNA synthesis.

A plasmid used to generate the enhancer trap donor construct was provided by Shin-Ichi Higashijima (pBluescript-SK-Gbait-Hsp-Gal4FF-BGHpA) (Kimura, Hisano, Kawahara, & Higashijima, 2015). After modification, this plasmid was injected into Tg(UAS:GFP), *dscamb*^{t2b} heterozygous embryos at the 1-cell stage, along with 1–5 nl of an injection mix containing Cas9 mRNA (200–300 pg/nl), donor plasmid (10–20 pg/nl), gRNA-Et1/Et2 (20–40 pg/nl), and gRNA-Mbait (20–40 pg/nl). Injected fish were screened for Gal4 expression to established stable enhancer trap lines. PCR was used to confirm enhancer trap integration and orientation. See Supplemental Methods for details.

Anatomical characterization

Details of RB and ENS characterization, ORN photoconversion, retinal dissection, sparse retinal labeling, and TEM are described in Supplemental Methods. Larvae were prepared for retinal cryosectioning according to a previously published protocol (Uribe & Gross, 2007). Immunofluorescence staining was carried out using a previously published protocol (Uribe & Gross, 2007), with some modifications, described in supplementary methods. Supplemental Table 1 summarizes antibodies used for immunofluorescence.

Behavioral assays

Touch: 2 dpf larvae were subjected to three touch trials. For each trial, a small needle was used to lightly graze the tail, and escape responses were scored. Each touch trial was separated by at least 10 s.

Background adaptation: 7 dpf fish were placed in the dark for several hours to allow pigment adaptation before being placed under bright illumination. After 30 min, larvae were categorized into either light (correct adaptation) or dark (incorrect adaptation) pigmentation groups.

Hearing: 6-8 dpf fish were placed in custom agarose chambers flooded with E3 solution and imaged with a Pike IEEE 1394b camera (Allied Vision Technologies, Stadtroda, Germany) at 544 frames per second. Startle stimuli consisted of 10 ms, 1 KHz acoustic/vibrational pulses delivered by a speaker mounted directly to the imaging platform 6 cm from the dish. Image acquisition and stimulus delivery were driven using custom software written in LabView (National Instruments, Austin, TX). For PPI testing, each dish was subjected to 15 startle-only trials and 15 PPI trials. PPI trials consisted of a non-startling prepulse followed by a startling stimulus with a 400 ms inter-stimulus interval. Startle stimulus amplitude was defined as the lowest amplitude evoking c-bend startle in at least 90% of embryos over six repetitions. Prepulse stimulus amplitude was defined as the highest amplitude evoking c-bend startle behaviors in less than 10% of embryos over six repetitions. Percent inhibition was calculated for each fish as [(% of startle trials showing shortlatency C-bends) – (% of PPI trials showing short-latency Cbends)]/(% of startle trials showing short-latency C-bends) * 100.

Optomotor response: 7 dpf fish were transferred to a 35 mm petri dish inside a circular arena lined with arrays of computer-controlled LEDs. Using MatLab software (Natick, MA), illumination of LED arrays was coordinated into bars that rotated around the chamber circumference. The angular velocity and direction of the rotating stimuli was adjusted along a 0.05 Hz sine wave. Four larvae, each in their own dish, were tested during each experiment, and their movement was recorded by a camera above the testing chamber. Each change of the stimulus direction was counted as one trial. Swimming behavior was ranked on a 1–3 scale: 3 = movement in the direction of stimulus; 2 = undirected movement; 1 = movement opposite to the stimulus.

Light and dark flash responses: 5 dpf larvae in 60 mm Petri dishes (20 fish per dish) were adapted on a white light box $(800 \,\mu\text{W/cm}^2)$ for at least 1 h prior to testing. Each larva was transferred to a well of a 4×4 testing grid (Wolman, Jain, Liss, & Granato, 2011). Responses to light and dark flash stimuli were recorded with a MotionPro Y4 high-speed

video camera (Integrated Design Tools, Tallahassee, FL) at 1000 frames per second and 512×512 pixel resolution, using a 50 mm macro lens (Sigma Corporation of America, Ronkonkoma, NY). Larvae were illuminated from above with a mounted LED light (MCWHL5 6500 K LED, powered by LEDD1B driver, Thorlabs, Newton, NJ) and below with an infrared light source (IR Illuminator CM-IR200B, C&M Vision Technologies, Houston, TX). Video images were analyzed for O-bend and routine turn response initiations and kinematics with the FLOTE software package, as previously described (Burgess & Granato, 2007a, 2007b; Burgess, Schoch, & Granato, 2010; Wolman et al., 2011). Dark flash O-bends and light flash turns were elicited and analyzed as previously described (Burgess & Granato, 2007a; Wolman et al., 2011). Once in the testing grid, larvae were acclimated to an overhead light $(85 \,\mu W/cm^2$ for dark flashes, $25 \,\mu\text{W/cm}^2$ for light flashes) for 5 min before exposure to a series of 10 flash stimuli (dark flash = lights off, light flash = light increased to $600 \,\mu\text{W/cm}^2$ within 1msec). Each stimulus lasted for 1s and stimuli was presented at a 30s interstimulus interval.

After all behavioral tests, individual fish were lysed for genotyping. Thus, the experimenter was blind to genotype during behavioral testing.

Mortality analysis

A mixture of wild-type, heterozygous, and homozygous $dscamb^{t2b}$ mutant siblings were distributed into tanks at a density of 20–40 fish/tank and raised under standard rearing conditions. At select time-points (7, 10, 13, 16, 19, 34, and 60 dpf), whole tanks were euthanized in 0.02% tricaine onice for 10 min, and individual fish were either lysed whole or fin clipped (34 and 60 dpf only) for genotyping.

Food intake assay

Analysis of food intake was adapted from a previously published protocol (Field *et al.*, 2009). See Supplemental Methods for details.

Gulping assay

At 7 dpf, Et1(dscamb^{t2b}:Gal4), Tg(UAS:GFP) heterozygous and homozygous mutant siblings were immobilized in 3% methylcellulose, with the ventral side facing up. Twenty seconds videos of spontaneous jaw movements (or 'gulps') were recorded on an AxioCam MRm CCD camera mounted on SteREO Discovery.V12 dissecting microscope with a frame interval of 0.076 frames/s. After recording, individual larvae were lysed for genotyping. Videos were analyzed by counting the total number of gulps during each video. The observer was blind to genotype while counting.

Statistical analyses

All statistical analyses were performed using the R software package (Vienna, Austria).

Results

dscamb mutations impair survival to adulthood, but produce no obvious anatomical defects

To investigate the function of Dscamb in zebrafish neuronal development, we used TALENs to generate loss-of-function mutant lines. We selected two target sites in the coding region of dscamb: (1) the translational start codon, and (2) a region in the middle of exon 2 (Figure 1(A)). We engineered a pair of TALENs flanking each target site (Cermak et al., 2011) and injected mRNAs encoding each TALEN pair into embryos at the 1-cell stage. For each target site, we used Restriction Fragment Length Polymorphisms (RFLPs) and sequencing to identify multiple founders with germline frameshift mutations that generate premature stop codons (Figure 1(B)). We established four mutant *dscamb* lines with unique mutant alleles - two mutations for each target site: $dscamb^{t1a}$, $dscamb^{t1b}$, $dscamb^{t2a}$, and $dscamb^{t2b}$. We sequenced RT-PCR products from multiple dscam^{t2a} homozygous embryos and verified that the dscamb genomic frameshift mutation was integrated into mRNA transcripts (data not shown).

Across both target sites and mutant alleles, we detected no obvious developmental or anatomical defects distinguishing wild-type and homozygous mutant siblings, at least up to 7 days post-fertilization (dpf). However, most homozygous mutants died before reaching sexual maturity (\sim 3 months old). Moreover, heterozygous siblings were significantly underrepresented relative to their wild-type siblings at the same ages (wild-type: heterozygous: homozygous sibling survival: exon 1 mutants =43:55:1, exon 2 mutants =45:52:0; wt × het exact multinomial test: exon 1 mutants p = .0317, exon 2 mutants p = .0093), suggesting that *dscamb* mutation confers a developmental deficit that impairs survival.

CRISPR/Cas9-directed enhancer trap integration reveals the Dscamb expression pattern

Closer inspection of the tissues expressing Dscamb could pinpoint developmental defects underlying mutant mortality. To identify *dscamb*-expressing cell types and image their development in live embryos, we created a dscamb enhancer trap reporter using CRISPR/Cas9-guided mutagenesis to insert a reporter transgene upstream of dscamb (Figure 1(C)) (Kimura et al., 2015). By integrating the enhancer trap upstream of the mutant allele, we could more readily identify heterozygous and mutant embryos and thus identify loss-of-function phenotypes in *dscamb*-expressing cells. We selected two guide RNA (gRNA) target sites upstream of the dscamb locus: (1) gRNA-Et1, located 69 bp upstream of the transcriptional initiation site, and (2) gRNA-Et2, located 4 bp upstream of the initiation site. dscamb-targeting gRNAs were injected into single-cell embryos along with a donor transgene plasmid containing a minimal heatshock promoter driving expression of Gal4FF (Asakawa & Kawakami, 2009), Cas9-encoding mRNA, and a second gRNA to linearize the donor plasmid (Figure 1(C)). We generated three lines with integrations upstream of the wild-type allele, and one

integrated upstream of the $dscamb^{t2b}$ mutant allele (Figure 1(D–G)). Of the wild-type enhancer trap lines, two were integrated at the gRNA-Et1 target site: Et1(dscamb^{wt}:Gal4)ⁱ, Et1(dscamb^{wt}:Gal4)^o. The remaining wild-type enhancer trap integrated at the sgRNA-Et2 site: Et2(dscamb^{wt}:Gal4). Integration upstream of the mutant allele was at the gRNA-Et1 site: Et1(dscamb^{t2b}:Gal4). With this method, the enhancer trap plasmid can integrate in either forward or reverse orientations, and multiple copies of the plasmid can integrate tandemly at the target site. Previous studies found only minor differences in enhancer trap expression between forward and reverse integration (Kimura *et al.*, 2015). All three gRNA-Et1 enhancer trap lines were integrated in the reverse orientation; the gRNA-Et2 line showed evidence of both reverse and forward integration (Figure 1(D–G)).

We crossed each enhancer trap line to Tg(5xUAS:GFP) to compare expression at 5 dpf (Figure 1(D-G)). Expression was similar regardless of target site, suggesting that integration does not disrupt critical regulatory elements and that the orientation of integration has minimal effects on expression. Integrations upstream of the wild-type and mutant alleles also showed similar expression patterns, indicating that dscamb mutations do not dramatically alter expression. UAS transgene positional effects were also minimal, as there was little difference in the expression pattern visualized with other UAS lines (UAS:KikGR, UAS:nfsb-mCherry). Since there were no obvious differences between reporter lines, we focused on the mutant enhancer trap line [Et1(dscamb^{t2b}:Gal4)] for most experiments. This allowed us to characterize the expression pattern of Dscamb while comparing development in heterozygous and homozygous mutants.

As a second approach to characterize *dscamb* expression, we generated a BAC transgenic reporter. Specifically, we replaced the start codon in a 40.8 kb BAC spanning the first exon of *dscamb* with a Gal4-containing reporter cassette and integrated this BAC into the genome to generate a stable line: BAC(dscamb:Gal4) (Figure S1(A)). Although BAC reporter expression was sparser than Et1(dscamb^{t2b}:Gal4), every region with BAC reporter expression was also observed in the enhancer trap, including the retina, otic hair cells, cranial sensory ganglia, the hindbrain, and the spinal cord, further supporting the accuracy of the enhancer trap expression pattern (Figure S1(B-G)).

The Dscamb enhancer trap is expressed throughout the central nervous system

To characterize expression of Et1(dscamb^{t2b}:Gal4), we crossed Et1(dscamb^{t2b}:Gal4), Tg(5xUAS:GFP) to a pan-neuronal line [Tg(nbt:RFP)], and imaged fish at embryonic (1 dpf) and larval (4–5 dpf) stages. At both stages, GFP was expressed throughout the brain and spinal cord (Figure 2). Although many cells were GFP+/RFP+, we frequently observed cells that were RFP+/GFP-, indicating that dscamb is not expressed in all neurons. Conversely, many cells were RFP-/GFP+, suggesting that either the nbt:RFP transgene incompletely labels neurons or that *dscamb* is also

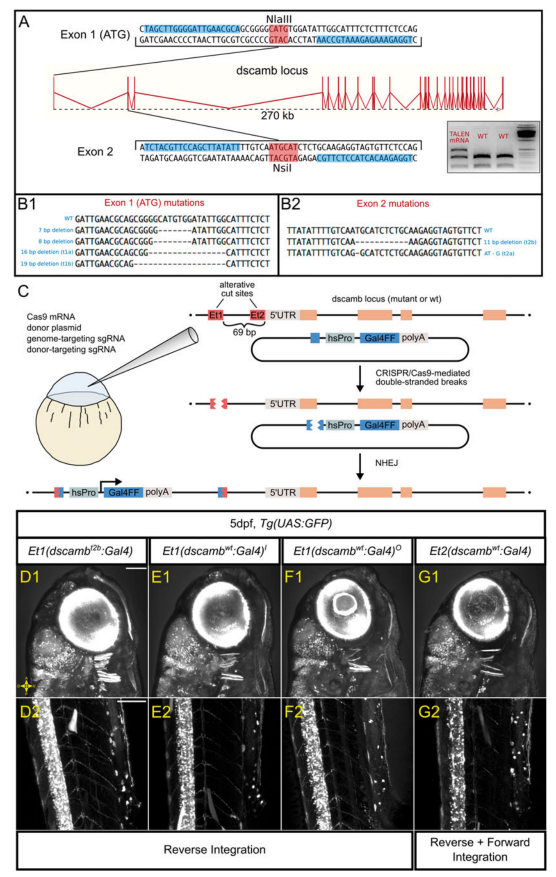


Figure 1. Using targeted mutagenesis to generate *dscamb* loss-of-function mutants and enhancer trap reporters. (A) The *dscamb* locus, including TALEN target sites in exon 1 (top) and exon 2 (bottom). Blue boxes indicate TALEN binding sites. Red boxes indicate restriction enzyme sites used for RFLP genotyping. Inset is an example of RFLP genotyping for mutations at exon 2 using genomic DNA from pooled embryos injected with TALEN-encoding mRNA or uninjected. The upper, uncut band in TALEN-injected embryos indicates mutation of the Nsil site. Gene diagram adapted from Ensembl. (B) TALEN-generated germline mutations at exon 1 (B1) and exon 2 (B2) sites. (C) CRISPR/Cas9 strategy for enhancer trap insertion. (D–G) Representative confocal images of independent enhancer trap lines, showing reporter expression in the head (D1, E1, F1, G1) and trunk (D2, E2, F2, G2). Scale bars =100 um.

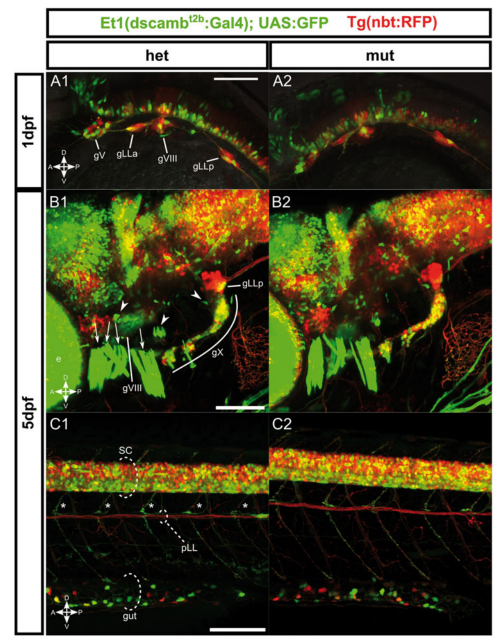


Figure 2. dscamb enhancer trap reporter is expressed broadly throughout the CNS and PNS, but homozygous mutant embryos lack obvious structural defects. (A–C) Et1(dscamb^{t2b}:Gal4) enhancer trap (green) crossed to pan-neuronal Tg(nbt:RFP) in heterozygous (left) and homozygous (right) mutants. (A1, A2) Lateral view of the head in 1dpf embryos. gV: trigeminal ganglion; gLLa: anterior lateral line ganglion; gVIII: statoacoustic ganglion; gLLp: posterior lateral line ganglion. (B1, B2) Lateral view of head in 5 dpf larvae. e: eye; gVIII: statoacoustic ganglion; gX: vagal ganglia; gLLp: posterior lateral line ganglion; arrowheads: otic vesicle hair cell patches; arrows: jaw muscles. (C1, C2) Lateral view of trunk in 5 dpf larvae. SC: spinal cord; pLL: posterior lateral line axons; asterisks: motor neuron axons; gut: intestine. Scale bars: 100 μm.

expressed in glial cells. At 4-5 dpf, the enhancer trap was broadly expressed in the forebrain, midbrain, and hindbrain (Figure 2(B)). In the forebrain, *dscamb* was expressed in the olfactory bulb and other unidentified regions (Figure 3(C); additional forebrain expression not shown). GFP + neurons were also seen in the hindbrain cerebellum and the midbrain optic tectum, where GFP + neurites spread throughout the tectal neuropil.

In the spinal cord, enhancer trap expression was more abundant ventrally (Figure 2(C)). GFP + motor axons exited the spinal cord to innervate trunk muscles from all three zebrafish primary motor neurons (the caudal (CaP), middle (MiP), and rostral (RoP) neurons, data not shown).

Somatosensory Rohon–Beard neurons (RBs) in the dorsal spinal cord also expressed the enhancer trap. Crossing the enhancer trap to a reporter for an RB subtype (BAC[TrpA1b:GFP]) (Palanca *et al.*, 2013) demonstrated that *dscamb* is expressed in a subtype of RBs partially overlapping with TrpA1b+RB neurons (Figure 3(A,B)). Since *Drosophila Dscam1* regulates self-avoidance in touch-sensing neurons (Hughes *et al.*, 2007; Matthews *et al.*, 2007; Soba *et al.*, 2007), we analyzed sensory axon morphology in the skin by injecting the BAC reporter to sparsely label RBs. We did not observe any significant changes in the frequency of axon cross-overs, suggesting that zebrafish *dscamb* does not regulate self-avoidance in RBs (Figure S2). More broadly, we

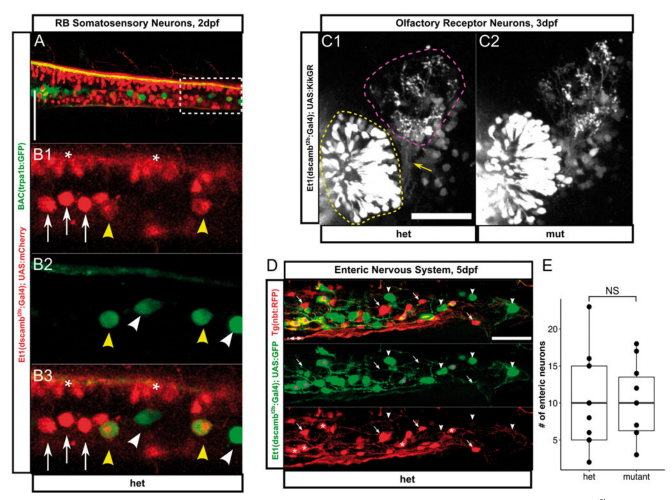


Figure 3. *dscamb* expression and function in specialized cell types. (A) Dorsal view of spinal cord in 2 dpf embryo showing Et1(dscamb^{12b}:Gal4) in red and BAC(trpa1b:GFP) in green. Scale bar: 100 μ m. Anterior is to the left. (B1–B3) Higher magnification of box in A. Yellow arrowheads: GFP+/RFP + RBs, white arrowheads: GFP+/RFP - RBs, white arrowse: GFP-/RFP + RBs, asterisks: other RFP + neurons. (C) Et1(dscamb^{12b}:Gal4) driving expression of KikGR in 3 dpf heterozygous (C1) and homozygous (C2) larvae. KikGR in ORNs (yellow dashed region) was photoconverted from green to red to delineate axon terminals in glomeruli (magenta dashed region). Yellow arrow: ORN axon coursing to the olfactory bulb. Scale bar: 50 μ m. Dorsal is up, medial is right. (D) Distal portion of gut in a 5 dpf heterozygous dscamb mutant; Et1(dscamb^{12b}:Gal4) expression in green and Tg(nbt:RFP) expression in red. Asterisks: GFP+/RFP + netroic neurons, arrowheads: GFP + only enteric neurons. Scale bar: 50 μ m. (E) Number of enteric neurons in the terminal 250 μ m of the gut. Each point represents one larvae. Mann–Whitney–Wilcoxon test: *p* = .68. Middle box line is the median; lower and upper ends of boxes are 25th and 75th percentiles.

did not detect any obvious structural differences in the brain or spinal cord of homozygous mutant animals (Figure 2). We also compared brain and spinal organization of a wildtype enhancer trap reporter [Et2(dscamb^{wt}:Gal4)] to heterozygous and homozygous enhancer trap mutants, but did not detect any obvious anatomical differences (Figure S3).

Dscamb is broadly expressed in the peripheral nervous system

Et1(dscamb^{12b}:Gal4) was also expressed broadly in the peripheral nervous system (PNS). At 1 dpf, all of the early sensory ganglia, including the trigeminal, anterior and posterior lateral line, octaval/statoacoustic ganglia, and olfactory placode, contained GFP+cells (Figure 2(A)). Compared to Tg(nbt:RFP), many cells in sensory ganglia were RFP+/ GFP-, suggesting that *dscamb* is expressed in a subtype of these sensory neurons. At larval stages (5 dpf), enhancer trap expression persisted in trigeminal, anterior lateral line, posterior lateral line, statoacoustic, and vagal ganglia (Figure 2(B)), and was also observed in the glossopharyngeal and facial ganglia (data not shown). Again, GFP expression was only detected in a subset of sensory neurons, although the size of this subset varied between ganglia. We did not detect obvious defects in the composition or organization of sensory ganglia in *dscamb* mutants (Figure 2(A,B)).

In several sensory systems, enhancer trap expression was detected in both primary sensory neurons and their downstream partners. For example, in addition to the statoacoustic ganglion, GFP+sensory hair cells were seen in all sensory hair cell (ovHCs) patches of the otic vesicle (the developing inner ear) (Figure 2(B)). GFP+axons, likely afferents from the statoacoustic ganglion, were frequently seen coursing into the ovHC patches. No pronounced changes in ovHC patches or innervating GFP+axons were detected in *dscamb* homozygous mutants.

In the olfactory system, GFP + neurons were in both primary olfactory receptor neurons (ORNs) and the downstream olfactory bulb (Figure 3(C)), suggesting that Dscamb might regulate axon guidance or synaptic coupling between these neurons. Since expression in both ORNs and the olfactory bulb made it difficult to discern the morphology of ORN axon terminals, we crossed $Et1(dscamb^{t2b}:Gal4)$ to a UAS line expressing a photoconvertible fluorescent protein (KikGR) (Figure 3(C)) (Das & Gage Crump, 2012). Using a UV laser, we selectively photoconverted KikGR from green to red in the ORNs, allowing us to distinguish their axons in the olfactory bulb. Mutant axons innervated the olfactory bulb and coalesced into glomeruli in a pattern closely resembling their heterozygous siblings, indicating that *dscamb* is dispensable for ORN axon guidance and segregation into glomeruli.

Outside of cranial sensory ganglia, Et1(dscamb^{t2b}:Gal4) was expressed in enteric neurons (ENS neurons) scattered along the length of the gut tube (Figure 2(C)). Coexpression analysis with nbt:RFP revealed that the enhancer was expressed in some, but not all, ENS neurons (Figure 3(D)). We found no difference in the median number of ENS cells between heterozygous and homozygous mutant embryos (median =10 ENS neurons in both cases; n = 10 animals per genotype) (Figure 3(E)), indicating that *dscamb* is not required for the migration of ENS neurons into the distal gut in zebrafish.

Dscamb expression in branchiomotor neurons and their target muscles

At larval stages, the dscamb enhancer trap labeled a specific population of muscle fibers, located anterior and ventral to the otic vesicle, including the levator arcus palatini (LEV-AP), dilatator operculi (DIL-OP), adductor hyomandibulae (AD-HYO), levator operculi (LEV-OP), and adductor operculi (AD-OP) (Figure 2(B)). Additional GFP + muscle fibers were located on the ventral surface of the head, including the intermandibularis anterior (INTM-A), intermandibularis posterior (INTM-P), interhyoideus (INTE), hyoideus inferior (HH-INF), hyoideus superior (HH-SUP), sternohyoideus (SH), adductor mandibulae (ADM), and the transverse ventralis (TV) (Figure 4(A)) (Diogo, Hinits, & Hughes, 2008; Schilling & Kimmel, 1997). These muscle fibers receive innervation from either the branchiomotor neurons of the V (trigeminal), VII (facial), IX (glossopharyngeal), or X (vagus) cranial nerves, located in the hindbrain. Together, these motor pools coordinate the movement of the jaw, operculum (bony gill cover), and gills during feeding and gill ventilation (Chandrasekhar, 2004; Hernandez, Patterson, & Devoto, 2005; Schilling & Kimmel, 1997). This is the first report of DSCAM family member expression in skeletal muscle, raising the possibility that dscamb may regulate the development of peripheral motor circuits.

Defects in feeding or inspiratory motor coordination provide a plausible explanation for homozygous mutant mortality, leading us to speculate that *dscamb* could be required for the development or maintenance of connections between branchiomotor neurons and their target muscle fibers. To distinguish individual hindbrain motor nuclei, we crossed the enhancer trap to Tg(isl1:GFP), which labels cranial motor neurons (Figure 4(B,C)). At 4 dpf, enhancer trap-expressing cells were not detected in the trigeminal motor nucleus (data not shown), occasionally in the facial nucleus

(Figure 4(B)), and frequently in the vagus nucleus (Figure 4(C)). Thus, at 4 dpf, dscamb is expressed in a subset of facial and vagal branchiomotor neurons. To determine if dscamb is required for establishing synapses between branchiomotor neurons and target muscles, we used alpha-bungarotoxin (aBTX) to stain neuromuscular junctions (NMJs) in 7 dpf larvae (Figure 4(D,E)). Using Et1(dscamb^{t2b}:Gal4) expression to visualize dscamb-expressing muscles on the ventral surface of the jaw, we did not detect any pronounced defects in the organization of ventral jaw muscles in dscamb mutants (Figure 4(A)). We quantified the number and volume of aBTX-labeled postsynaptic puncta on the INTM-A and INTM-P muscles of the ventral head (Figure 4(D-G)). There was no significant difference between heterozygous and homozygous mutants (Mann-Whitney-Wilcoxon test: number of puncta p = .98, volume of puncta p = .46; n = 16heterozygous and 13 homozygous), suggesting that dscamb is dispensable for forming branchiomotor NMJs, though subtle defects in synaptic development may have eluded our analyses.

Dscamb is not required for retinal lamination, cell, or branch spacing

In the mouse retina, *Dscam* is expressed primarily in RGCs, but also in subtypes of ACs and BCs, whereas Dscaml1 is expressed in rod PRs and distinct BC, AC, and RGC subtypes in rod retinal circuits (Fuerst et al., 2009). Previous studies, using transcriptomics and in situ hybridization identified dscamb expression in the ONL, INL, and RGL (Sun et al., 2018), but the extent of gene expression across the major types of retinal neurons has not been assessed in detail. We characterized *dscamb* enhancer trap expression in retinal cryosections from 5 dpf larvae - a time point at which the retina is stratified and functional (Figure 5). Although GFP expression was brightest and most abundant in PRs, the RGL was also densely labeled (Figure 5(A)). GFP expression in the inner nuclear layer (INL) was less dense, but GFP + cells were still visible throughout the INL. Based on cell body location, most GFP + INL neurons appeared to be ACs, but GFP + BCs and HCs were also observed. Thus, dscamb is expressed in all major retinal cell types, but is most broadly expressed in PRs, followed by RGCs and ACs. In the mouse retina, Dscam or Dscaml1 loss-of-function reduces programmed cell death, leading to expansion of the INL and IPL (Fuerst et al., 2008, 2009; Keeley et al., 2012). We did not observe any differences in the number of dscamb-expressing cells or thickness of the IPL in heterozygous and homozygous dscamb mutants (Figure S4), suggesting that *dscamb* does not regulate programmed cell death.

Dscamb is expressed in subtypes of ACs and RGCs but is not required for their lamination or branch spacing

We used immunofluorescence to further characterize the expression of $Et1(dscamb^{t2b}:Gal4)$ surrounding the IPL (Figure 5(B–D)). Zebrafish ACs can be located in either the INL or the RGL (as displaced ACs [dACs]). We stained 5 dpf retinal cryosections with 5E11, an antibody labeling all

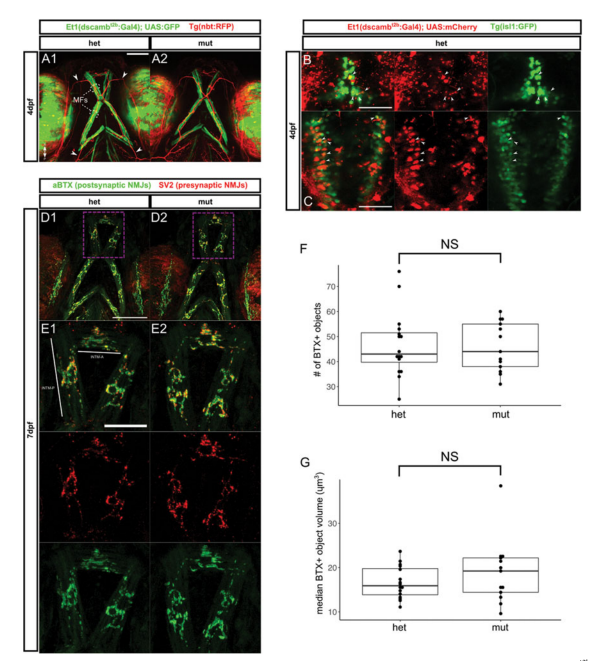


Figure 4. Jaw muscle fibers and branchiomotor innervation are intact in *dscamb* mutants. (A) Ventral view of head in 4 dpf larvae with Et1(dscamb^{12b}:Gal4) expression in jaw muscle fibers in heterozygous (A1) and homozygous (A2) mutants. Arrowheads: BMN axons, dotted lines: jaw muscle fibers (MFs). Scale bar: 100 μ m. (B, C) Dorsal images of hindbrain regions in 4 dpf larvae with Et1(dscamb^{12b}:Gal4) in red and BMNs in green [Tg(is11:GFP)]. Confocal projections of the facial (B) and vagal (C) BMN nuclei. Arrowheads: examples of mCherry+/GFP + BMNs. Scale bars: 50 μ m. (D) Confocal projections of ventral head in 7 dpf heterozygous (D1, E1) and homozygous (D2, E2) mutants with NMJs labeled presynaptically (SV2, red) and postsynaptically (aBTX, green). E1, E2 are magnifications of dashed boxes in D1, D2, showing NMJs on the INTM-A and INTM-P muscles. Scale bar D1: 100 μ m, scale bar E1: 50 μ m. (F, G) Quantification of aBTX-stained object number (F) and motion volume on the INTM-A and INTM-P of 7 dpf heterozygous and homozygous mutant larvae. *N* = 16 hets and 13 muts. Mann–Whitney–Wilcoxon test: *p* = .98 for F and *p* = .46 for G. Middle line is the median; lower and upper ends of boxes are 25th and 75th percentiles.

ACs (Figure 5(B)) (Hyatt, Schmitt, Fadool, & Dowling, 1996; Link, Fadool, Malicki, & Dowling, 2000). GFP+/5E11+ cell were present in both the IPL and RGL, confirming that *dscamb* is expressed in ACs, including dACs. However, many 5E11+ cells on both sides of the IPL were GFP-, suggesting that *dscamb* is expressed in a subpopulation of ACs. Staining retinal cryosections for Parvalbumin (Parv), a marker for an AC subtype on both sides of the IPL (Nevin, Taylor, & Baier, 2008) revealed that *dscamb*-expressing ACs sometimes also expressed Parv (Figure 5(C)). To determine if *dscamb* is expressed in RGCs, we stained for the pan-RGC marker, Hermes (Figure 5(D)) (Hörnberg *et al.*, 2013; Zearfoss, Chan, Wu, Kloc, & Etkin, 2004), which demonstrated that the *dscamb* enhancer trap labels a subtype of RGCs. There were no appreciable changes in cell number or organization of RGCs or ACs in homozygous mutant retinas (Figure S4(C-H)).

In both mice and chicks, DSCAMs regulate the segregation of neuronal processes into distinct sublamina in the IPL (Li *et al.*, 2015; Yamagata & Sanes, 2008). In the zebrafish retina, Parv + processes segregate into three distinct sublaminae: s25, s45, and s85 (Figure 5(E)) (Nevin *et al.*, 2008).

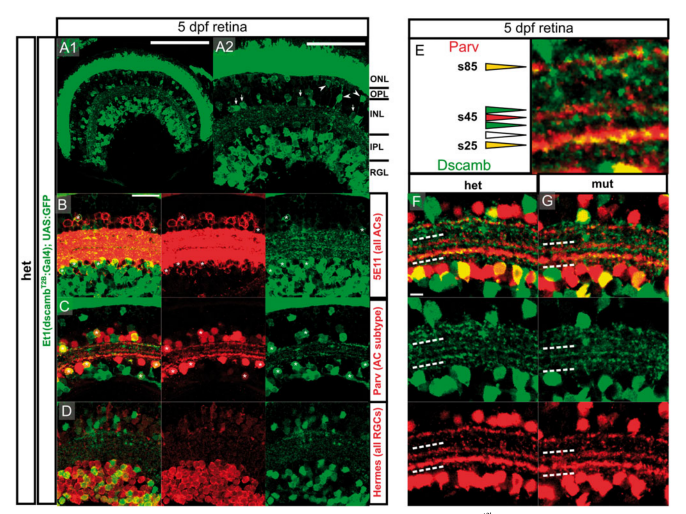


Figure 5. *dscamb* is expressed in ACs and RGCs subtypes, but is not required for IPL lamination. (A1) Et1(dscamb^{12b}:Gal4) expression in retinal section from 5 dpf larva. Dorsal is left, medial is up. Scale bars: 100 μ m. (A2) Higher magnification image of A1. Arrowheads: BCs; arrows: ACs. ONL: outer nuclear layer; OPL: outer plexiform layer; INL: inner nuclear layer; IPL: inner plexiform layer; RGL: retinal ganglion cell layer. Scale bar: 50 μ m. (B–D) Confocal images of IPL in sections from 5 dpf heterozygous larvae. Scale bar: 20 μ m. (B) Et1(dscamb^{12b}:Gal4) compared to pan-AC marker (5E11; red). Asterisks: GFP+/5E11+ ACs. (C) Et1(dscamb^{12b}:Gal4) compared to Parv + AC marker (red). Asterisks: GFP+/Parv + ACs. (D) Et1(dscamb^{12b}:Gal4) compared to pan-RGC marker (Hermes; red). (E) s25, s45, and s85 Parv + sublaminae in the IPL. Triangles indicate sublaminae that are GFP+/Parv+ (yellow), GFP + only (green), Parv + only (red), or negative for both (empty). Scale bar: 5 μ m. (F, G) Images of IPL in 5 dpf heterozygous (F) and homozygous (G) *dscamb* mutant larvae. White dashed lines outline region containing s25 and s45 Parv + sublaminae. Scale bar: 5 μ m.

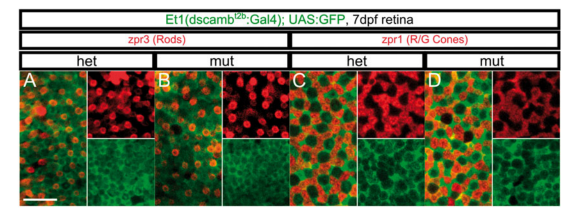
These bands corresponded to the stratification of processes from Dscamb-expressing cells (Figure 5(E)). s25, the thickest Parv + band, located closest to the RGL, often co-labeled with GFP + processes. The Parv + s45 layer was flanked by two GFP + bands. Interestingly, GFP was not expressed in s45, suggesting that this sublamina is composed primarily of processes from GFP – Parv + ACs. s85, the Parv + sublamina closest to the INL, was frequently co-labeled with GFP. Although we did not discern defects in the localization of these bands in *dscamb* homozygous mutants (Figure 5(F, G)), the absence and aggregation of Dscamb-expressing processes into consistent, identifiable sublaminae suggests that Dscamb expression is specific to certain cells types that synapse in specific regions of the IPL.

In mice, *Dscam* and *Dscaml1* mutant AC and RGC neurites fail to self-avoid, leading to extensive fasciculation and an obvious disruption of arbor morphology (Fuerst *et al.*, 2008, 2009). To determine if *dscamb* regulates self-avoidance in the zebrafish retina, we analyzed individual cells in homozygous and heterozygous mutant

retinas (Figure 6). To distinguish individual neurons, we sparsely labeled the retina by injecting low concentrations of UAS transgene into Et1(dscamb^{12b}:Gal4) embryos. At 5dpf GFP + ACs and RGCs were indistinguishable between heterozygous and homozygous mutants. Although we could not reliably resolve fine details of arbor morphology, we saw no signs of extensive fasciculation resembling the self-avoidance phenotype in mouse DSCAM mutants (Figure 6(E–H)). Thus, we found no evidence that *dscamb* is required for self-avoidance in the RGC or AC neurites, but cannot rule out that it may play a subtle role in this process.

Dscamb loss-of-function does not disrupt ribbon synapses or mosaic spacing of PRs

Both PRs and ovHCs have specialized synaptic structures called ribbon synapses. Enhancer trap expression in both these cell types raised the intriguing possibility that Dscamb may regulate ribbon synapse development. In *Dscaml1*



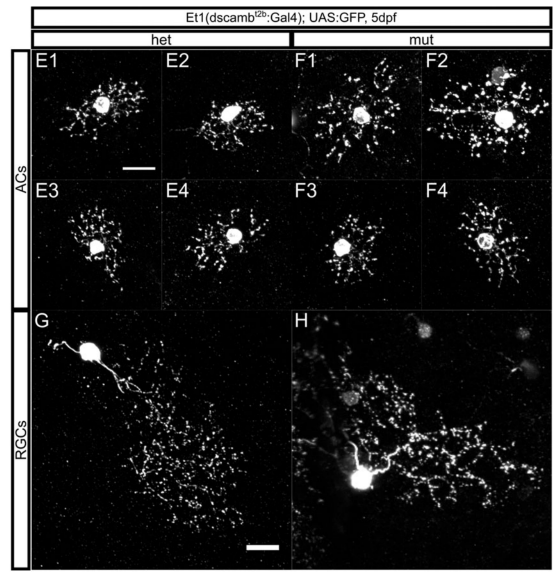


Figure 6. *dscamb* is not required for PR mosaic spacing or RGC/AC self-avoidance. (A–D) Confocal projection through the base of zpr3+ rod outer segments (red A, B) or zpr1+ R/G cone cell bodies (red C, D) in retinas from 7 dpf heterozygous (A, C) and homozygous (B, D) *dscamb* mutant larvae. Et1(dscamb^{12b}:Gal4) expression is green. Insets on the right side of each panel are single-color continuations of the left side. Scale bar in A: 10 μ m. (E–H) Sparse Et1(dscamb^{12b}:Gal4) labeling in individual ACs (E and F) or RGCs (G and H) in 5 dpf heterozygous (E and G) and homozygous (F and H) dscamb mutant retinas. Scale bars: 10 μ m.

mutant mice, rod PRs showed defects in synaptic maturation, such as an overabundance of synaptic vesicles and rudimentary synaptic ribbons (Fuerst *et al.*, 2009). Using transmission electron microscopy (TEM), we analyzed the structure of PR synaptic ribbons in 7 dpf mutant larvae (Figure S5), focusing on cone synaptic peduncles, because they are easy to identify by the presence of multiple synaptic ribbons. *dscamb* mutant cones developed synaptic ribbons that were indistinguishable from synapses in wildtype retinas.

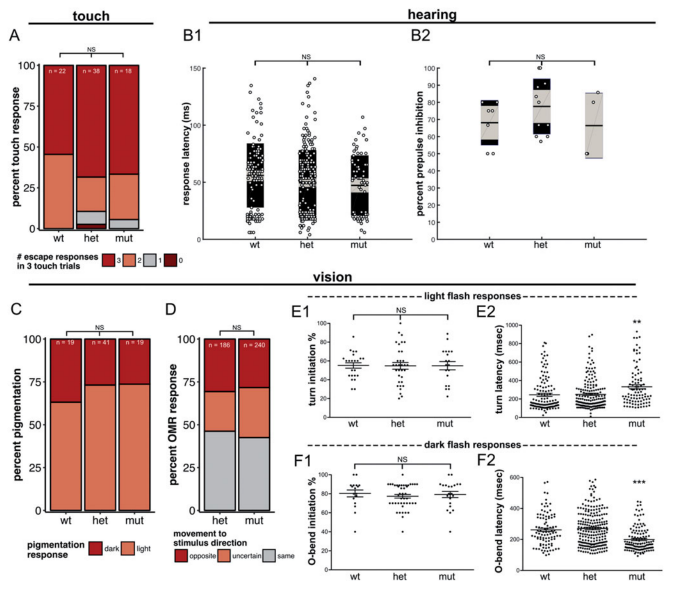


Figure 7. Sensory-evoked behavior in *dscamb* mutants. (A) Percentage of 3 dpf wild-type, heterozygous, and homozygous *dscamb* mutant larvae responding with 0, 1, 2, or 3 escape behaviors in three tail touch trials. Fisher's exact test of independence: p = .40. (B1) Auditory startle response latency in 6–8 dpf larvae. Middle bar: mean. Grey boxes: 95% confidence intervals. Outer black boxes: one standard deviation. ANOVA: p = .46. (B2) Pre-pulse inhibition (PPI) of 6–8 dpf larvae. ANOVA: p = .98. (C) Visually-mediated background adaptation. Percentage of 7 dpf larvae that adapted pigmentation (light) or failed to adapt pigmentation (dark) to a bright background. Fisher's exact test: p = .72. (D) Optomotor response (OMR). Percentage of trials in which 7 dpf heterozygous and homozygous mutant *dscamb* larvae responded to a rotating visual stimulus by moving in the same, opposite, or uncertain direction. Thirty-one hetorozygous larvae were tested in 186 trials. Forty mutant larvae were tested in 240 trials. Fisher's exact test of independence: p = .39. (E, F) Light (E) and dark (F) flash responses. Light flashes elicit turning responses and dark flashes elicit 0-bends. The percentage of trials with responses (E1, F1) and response latency during individual trials for each 5 dpf larva (E2, F2). Error = SEM. ANOVA: **p < .01, ***p < 0. $N \ge 17$ animals, ≥ 94 responses for each experiment.

In *Dscam* and *Dscaml1* mutant mice, cells of the same subtype clump together, disrupting their mosaic spacing (Fuerst *et al.*, 2008, 2009). Zebrafish PRs also have a highly regular mosaic arrangement (Bilotta & Saszik, 2001). To evaluate PR mosaic spacing, we imaged retinas from 7 dpf larvae stained for zpr3, a rod marker in embryonic and larval fish (Larison & Bremiller, 1990), and zpr1, a red and green cone marker (Figure 6(A–D)) (Schmitt & Dowling, 1996). Both zpr1 and zpr3 were completely coexpressed with Et1(dscamb^{12b}:Gal4), and GFP+cells covered the entire PR cell layer, indicating that Dscamb is expressed in all zebrafish PRs. However, the arrangement of rod and cone PRs were indistinguishable between heterozygous and

homozygous *dscamb* mutant retinas, suggesting that Dscamb is not required for PR mosaic spacing.

dscamb mutants have largely normal somatosensation and hearing, but may have subtle defects in vision

dscamb is expressed in trigeminal and RB neurons at early developmental stages, when these sensory neurons first innervate the skin. To analyze somatosensory function, we recorded the escape behavior of 2 dpf embryos in response to three trials of light tail touches (Figure 7(A)). There was no statistically significant difference in touch responses between wild-type, heterozygous, and homozygous mutant larvae (Fisher's exact test of independence p = .4), indicating that *dscamb* is not required for touch sensation in zebrafish larvae.

To characterize auditory function in larval *dscamb* mutants, we used a prepulse inhibition (PPI) assay. In PPI, an initial weak auditory stimulus inhibits behavioral response to a second stronger stimulus (Bhandiwad, Zeddies, Raible, Rubel, & Sisneros, 2013; Burgess & Granato, 2007b). 6–8 dpf larvae displayed no significant differences in PPI between wild-type, heterozygous, and homozygous mutant embryos (Figure 7(B)) (ANOVA: p = .98). The latency of the c-bend responses was also similar between genotypes (ANOVA: p = .46), indicating that *dscamb* is not required for auditory function or gross motor function.

To assess visual function in dscamb mutants, we first analyzed visually-mediated background adaptation (VBA) - a process in which larvae lighten their pigmentation in response to a bright environment and darken it in a lowlight environment (Neuhauss et al., 1999). Similar proportions of wild-type, heterozygous, and homozygous mutants adjusted their pigmentation when moved from a dark to light environment (Figure 7(C)) (light-adapted: wild-type =63%, heterozygous =73%, homozygous =74%; Fisher's exact test: p = .725). We next tested mutants for the optomotor response (OMR), a behavior in which laval fish tend to swim in the direction of a moving stimulus (Fleisch & Neuhauss, 2006). Heterozygous and homozygous dscamb mutants showed similar OMRs to multiple rounds of a rotating stimulus (Figure 7(D)). These analyses suggest that dscamb mutants detect light and that visual circuitry is sufficiently intact to process visual motion.

As a third analysis of visual function, we measured the responsiveness of mutant larvae to changes in illumination (Figure 7(E,F)). Exposing larvae to abrupt increases ('light flashes') or decreases ('dark flashes') in illumination elicits distinct turning behaviors. Dark-flash responses ('O-bends') have a larger turn angle than light-flash responses ('routine turns') (Burgess & Granato, 2007a). These responses are highly stereotyped and thus provide a sensitive measure of visual function and the circuitry underlying these behaviors. Among dscamb genotypes, there were no significant differences in the proportion of 5 dpf larvae that responded to light and dark flash stimuli (Figure 7(E1,F1)). However, homozygous mutants showed a statistically significant increase in their latency to initiate light-flash responses and a significant decrease in their latency to initiate dark-flash responses (Figure 7(E2,F2)). Other kinematic parameters of these light- and dark-elicited responses, such as distance moved or turning angle (data not shown), were indistinguishable among genotypes, suggesting that mutant behavioral changes are not secondary to motor defects. Thus, dscamb mutants likely have subtle deficiencies in initiating visually evoked behaviors.

dscamb mutants die during the first month of life and feed defectively

In the absence of marked sensory defects, a detailed characterization of *dscamb* mutant mortality might point to underlying mechanisms. By comparing the proportion of wild-type, heterozygous, and homozygous mutant siblings to the theoretical Mendelian inheritance, we found a gradual, downward trend in the survival of homozygous mutants, which first reached statistical significance at 16 dpf (Fisher's exact test: p = .0455) (Figure 8(A)). By 34 and 60 dpf there were significantly fewer mutants (p = .0013, p = .0017, respectively), and no homozygous mutants survived to 60 dpf in this experiment.

Zebrafish larvae begin acquiring nutrients from exogenous food sources at \sim 7 dpf, when the yolk sac is completely absorbed. In the absence of food, larvae perish from starvation, starting around 10 dpf, and completely succumb by 15 dpf (Versonnen, Roose, Monteyne, & Janssen, 2004; Wilson, 2012). Since dscamb mutants were under-represented at 16 dpf, we hypothesized that *dscamb* mutants may have a deficit in either finding or capturing food. To assess food intake, 7 dpf zebrafish larvae were fed larval food coated with fluorescent microspheres. After 90 min food content in the foregut (anterior intestine) was assessed (Figure 8(B–D)) (Field, Kelley, Martell, Goldstein, & Serluca, 2009) wild-type and heterozygous mutants had a similar ability to ingest food (Fisher's exact test, Bonferroni-corrected: wt \times het: p = .807), but fourfold more mutants than wild-type animals had empty foreguts (12.5% vs. 46.5%), and fivefold fewer mutants than wild-type had full foreguts (15.1% vs. 76.8%) (Fisher's exact test, Bonferroni-corrected: wt × mut: p < .001, het × mut: p < .001) (Figure 8(D)). Since Dscamb is expressed in jaw muscles, we considered the possibility that mutant jaw movements were defective, but a spontaneous 'gulping' assay demonstrated that mutant animals were capable of opening their jaws (Figure 8(E)). Alternatively, mutant animals may have an increased rate of food evacuation from the intestine, but time-lapse imaging of two animals with full guts did not reveal obvious defects in gut transit (data not shown). Thus, dscamb mutant animals may suffer from a combination of subtle defects in sensation, movement, digestion, or appetite that make their feeding behavior less efficient.

Discussion

Using targeted mutagenesis, we generated *dscamb* loss-offunction mutants and uncovered highly penetrant defects in mutant survival and food capture. Enhancer trap reporters for dscamb revealed broad expression in the central and peripheral nervous systems, and specific muscles of larval fish. In mouse embryos, Dscam also has a broad neuronal expression pattern, similar to zebrafish dscamb, including expression in the retina, ventral spinal cord, cerebellum, trigeminal ganglion, olfactory epithelium, olfactory bulb, and enteric nervous system (Agarwala, Ganesh, Amano, Suzuki, & Yamakawa, 2001; Barlow et al., 2002; Barlow, Micales, Lyons, & Korenberg, 2001; Yamakawa et al., 1998). However, the similarity between mouse and zebrafish DSCAM family members is not complete. In the retina, for example, both mouse Dscam and zebrafish dscamb are expressed abundantly in RGCs. However, unlike its murine

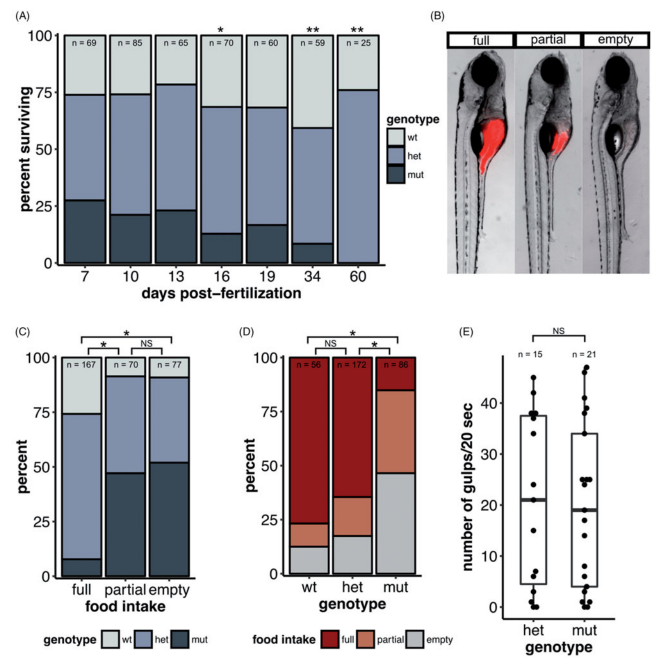


Figure 8. *dscamb* mutants have defective feeding behavior and die at 2–3 weeks of age. (A) Percentage of offspring of each genotype from heterozygous *dscamb* mutant crosses that survived to different time points. Multinomial exact test for goodness-of-fit (predicted proportions wt:het:mut =0.25:0.75:0.25) *p* values: 7 dpf =0.80, 10 dpf =0.74, 13 dpf =0.73, 16 dpf =0.046, 19 dpf =0.25, 34 dpf =0.0013, 60 dpf =0.0017. (B) Representative images of 7 dpf larvae in three categories of fluorescent food intake. (C) Percentage of each genotype in each food intake category. Fisher's exact test of independence across all groups: *p* = 1.0050. *Post hoc* Fisher's exact test with Bonferroni correction: full × partial *p* = 1.2e – 10, full × empty *p* = 4.1e – 13, partial × empty *p* = 1.0. (D) Percentage of each food intake category in each genotype. Fisher's exact test of independence across all groups: *p* = .00050. *Post hoc* Fisher's exact test with Bonferroni correction: wt × het *p* = .81, wt × mut *p* = 2.2e – 12, het × mut *p* = 1.8e – 13. (E) Quantification of number of gulps during a 20 s movie for 7 dpf heterozygous and homozygous mutants. Each data point represents one fish. Mann–Whitney–Wilcoxon test: *p* = .99. Middle line is the median; lower and upper ends of boxes are 25th and 75th percentiles.

counterpart, zebrafish *dscamb* is also expressed in photoreceptors. These distinct expression patterns suggest that DSCAM family members may have both overlapping and divergent functions across vertebrate species.

Despite broad neuronal expression and impairments in feeding and survival, we did not identify cellular or structural defects in *dscamb* mutants. Several factors could explain this observation. First, our enhancer trap reporter might not accurately reflect endogenous *dscamb* expression, and thus we did not examine the right neuronal populations. However, the fact that enhancer traps integrated at two sites, as well as a BAC reporter, had similar expression patterns argues against this possibility. Enhancer trap expression in the retina is also consistent with a recent study that identified *dscamb* mRNA expression in the ONL, INL, and RGL using *in situ* hybridization (Sun *et al.*, 2018). Second, gene expression might be incompletely knocked out in our mutant lines, or alternative splice sites or start codons could countervail frameshift mutations. Homozygous mutant mortality was highly penetrant across mutant lines with mutations at two separate target sites, providing strong evidence that the phenotype was specific to *dscamb*. In the future, generating larger genomic deletions that ablate critical domains could ensure complete loss of function. Third, dscamb may be genetically redundant with similar genes, such as dscama and dscaml1. Although the expression of zebrafish dscaml1 has not been investigated, in situ hybridization studies of dscama found broad expression throughout the brain, spinal cord, and retina, similar to dscamb (Yimlamai et al., 2005). Thus, it is possible that dscama compensates for dscamb loss-of-function. Future studies of dscama, dscamb double mutants could shed light on genetic interactions between these two genes. Nonetheless, the strong defects in homozygous mutant survival and feeding suggest that any functional redundancy is partial. Finally, dscamb may regulate subtle aspects of neuronal development that were undetectable in our anatomical analyses, similar to observations that in Dscam mutant mice the overall architecture of the brain is largely preserved (Amano *et al.*, 2009; Maynard & Stein, 2012). An accumulation of subtle developmental insults could lead to a dramatic deficit in a complex behavior such as feeding, which requires the coordinated activity of sensory, motor, and neuroendocrine systems. Although we did not detect dramatic defects with auditory, visual, or somatosensory assays, dscamb mutants did show increased latency in startle responses to a light flash and decreased latency responses to a dark flash. Thus, dscamb mutants are able to see, but may be deficient in their ability to process or respond to visual stimuli.

Vision is a critical component of larval feeding behavior, as anatomical lesions of the optic tectum abolish larval orientation towards and capture of prey paramecia (Gahtan, Tanger, & Baier, 2005), but other sensory systems likely also contribute. For example, the olfactory system regulates feeding behaviors in fish (Hamdani, Kasumyan, & Døving, 2001), and *dscamb* is expressed in both olfactory receptors neurons and the downstream olfactory bulb. Recent studies also demonstrated that molecular or anatomical ablation of branchiomotor neurons in larval zebrafish impairs food capture in a remarkably similar fashion to dscamb loss-of-function (Allen et al., 2017). dscamb is expressed in a subpopulation of larval branchiomotor neurons and their target muscles, but we did not detect obvious defects in the development of NMJs. Although mutant larvae are at least capable of opening and closing their mouths during spontaneous gulping, feeding is a complex motor behavior that requires the coordinated movement of the jaws, opercula, and the oral cavity (China & Holzman, 2014). Thus, more detailed kinematic analysis of branchiomotor-mediated movements may identify specific defects. The dscamb mutant and transgenic lines created in these studies will provide valuable tools to couple with additional behavioral analyses to unravel the cellular basis for visual and feeding impairments, thus uncovering novel functions for Dscam family proteins in vertebrate neuronal development.

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Disclosure statement

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