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4	Putative pore-forming subunits of the mechano-electrical transduction channel,
5	Tmc1/2b, require Tmie to localize to the site of mechanotransduction in zebrafish
6	sensory hair cells
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### 16 Abstract

17 Mutations in transmembrane inner ear (TMIE) cause deafness in humans; previous 18 studies suggest involvement in the mechano-electrical transduction (MET) complex in sensory 19 hair cells, but TMIE's precise role is unclear. In *tmie* zebrafish mutants, we observed that GFP-20 tagged Tmc1 and Tmc2b, which are putative subunits of the MET channel, fail to target to the 21 hair bundle. In contrast, overexpression of Tmie strongly enhances the targeting of Tmc2b-GFP 22 to stereocilia. To identify the motifs of Tmie underlying the regulation of the Tmcs, we 23 systematically deleted or replaced peptide segments. We then assessed localization and 24 functional rescue of each mutated/chimeric form of Tmie in *tmie* mutants. We determined that 25 the first putative helix was dispensable and identified a novel critical region of Tmie, the 26 extracellular region and transmembrane domain, which mediates both mechanosensitivity and 27 Tmc2b-GFP expression in bundles. Collectively, our results suggest that Tmie's role in sensory 28 hair cells is to target and stabilize Tmc subunits to the site of MET.

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#### **30** Author summary

Hair cells mediate hearing and balance through the activity of a pore-forming channel in the cell membrane. The transmembrane inner ear (TMIE) protein is an essential component of the protein complex that gates this so-called mechanotransduction channel. While it is known that loss of TMIE results in deafness, the function of TMIE within the complex is unclear. Using zebrafish as a deafness model, Pacentine and Nicolson demonstrate that Tmie is required for the localization of other essential complex members, the transmembrane channel-like (Tmc) proteins, Tmc1/2b. They then evaluate twelve unique versions of Tmie, each containing

38	mutations to different domains of Tmie. This analysis reveals that some mutations in Tmie cause
39	dysfunctional gating of the channel as demonstrated through reduced hair cell activity, and that
40	these same dysfunctional versions also display reduced Tmc expression at the normal site of the
41	channel. These findings link hair cell activity with the levels of Tmc in the bundle, reinforcing
42	the currently-debated notion that the Tmcs are the pore-forming subunits of the
43	mechanotransduction channel. The authors conclude that Tmie, through distinct regions, is
44	involved in both trafficking and stabilizing the Tmcs at the site of mechanotransduction.

45

# 46 Introduction

47 The auditory and vestibular systems detect mechanical stimuli such as sound, gravity, and 48 acceleration. These two systems share a sensory cell type called hair cells. The somas of hair 49 cells are embedded in the epithelium and extend villi-like processes from their apex into the surrounding fluid. The shorter of these, the stereocilia, are arranged in a staircase-like pattern 50 51 adjacent to a single primary cilium known as a kinocilium. Neighboring cilia are connected by 52 protein linkages. Deflection of the kinocilium along the excitatory axis tugs the interconnected 53 stereocilia, which move as a single unit called the hair bundle [1]. When tension is placed on the 54 upper-most linkages known as tip links, the force is thought to open mechanosensitive channels 55 at the distal end of the shorter stereocilia [2, 3]. These channels pass current, depolarizing the 56 cell and permitting electrical output to the brain via the eighth cranial nerve. The conversion of a 57 mechanical stimulus into an electrical signal is known as mechano-electrical transduction (MET) 58 [4]. The proteins located at the site of MET and involved in gating the MET channel are

59 collectively known as the MET complex. How the components of the MET complex, including the channel itself, are localized to and maintained at the stereocilia tips is not well understood. 60 61 To characterize the molecular underpinnings of MET and the underlying cause of 62 pathology in human patients, it is essential to examine the individual components of the 63 transduction complex in a comprehensive fashion. Thus far, only a few proteins have been 64 designated as members of the MET complex. The identity of the channel itself remains 65 contentious, but currently the best candidates for the pore-forming subunits are the 66 Transmembrane Channel-like (TMC) proteins TMC1 and TMC2. Mutations in TMC1 cause 67 human deafness [5], and double knock-outs of mouse Tmc1/2 result in the loss of MET currents 68 [6-8]. In zebrafish, overexpression of a fragment of Tmc2a generates a dominant negative effect 69 on hair-cell mechanosensitivity [9] and Tmc2a and Tmc2b are required for MET in hair cells of 70 the lateral line organ [10]. The TMCs localize to the tips of stereocilia, the site of MET, in mice 71 and zebrafish [3, 6, 8, 10-12]. A point mutation in mouse *Tmc1* results in altered channel 72 properties, suggesting direct changes to the pore [7, 13]. Likewise, in TMC2 knockout mice, 73 channel permeation properties are altered [14]. Regardless of whether the TMCs are the pore-74 forming or accessory subunits of the channel, they are essential for MET. 75 Another key component of the complex is Protocadherin-15 (PCDH15), which comprises 76 the lower end of the tip link [15, 16] and interacts with the TMCs [8, 9]. A fourth membrane 77 protein, Lipoma HMGIC fusion partner-like 5 (LHFPL5, formerly called TMHS), interacts with 78 PCDH15 and is critical for localizing PCDH15 to the site of MET [17, 18]. LHFPL5 is also 79 required to properly localize TMC1 in mouse cochlear hair cells [8]. However, loss of LHFPL5 80 in cochlear hair cells does not completely abolish MET currents, and currents can be rescued by 81 overexpression of PCDH15 [18]. This evidence suggests that LHFPL5 is not essential but rather

82 acts as an accessory protein. Another TMC1/2 interacting partner is Calcium and integrin

83 binding protein 2 (CIB2), which is a cytosolic protein that is localized in stereocilia and required

84 for MET in cochlear hair cells [19].

85 A sixth essential member of the MET complex is the transmembrane inner ear (TMIE) protein. Loss of TMIE results in deafness in all vertebrate organisms studied [20-25]. A recent 86 87 study demonstrated that TMIE is required for active MET channels in cochlear hair cells of mice 88 [26]. These authors showed that despite normal morphology of the inner ear, hair cells lacking 89 TMIE fail to label with aminoglycosides or FM 1-43, both of which are known to permeate the 90 MET channel [27, 28]. TMIE was first localized to the stereocilia of hair cells [29, 30], and then 91 to the stereocilia tips where MET occurs [25]. Zhao et al. further demonstrated that loss of TMIE 92 ablates MET currents, that TMIE interacts with both LHFPL5 and the CD2 isoform of PCDH15, 93 and that interfering with the TMIE-CD2 interaction alters MET. They proposed that TMIE could 94 be a force-coupler between the tip link and channel. However, the CD2 isoform of PCDH15 is 95 only essential in cochlear hair cells and not vestibular hair cells [31]. Zebrafish do not possess 96 the CD2 isoform [9, 32], and yet they still require Tmie for hair-cell function [21]. These 97 findings raised the tantalizing possibility that Tmie might have an additional role in MET that is 98 independent from the tip links. Here, we present an alternative role for Tmie in hair cell function. 99 We first confirmed that mechanosensitivity is absent in a zebrafish mutant of *tmie*, 100 ru1000, and demonstrated that this defect is rescued by transgenic Tmie-GFP. The localization 101 of Tmie-GFP is maintained in the absence of other transduction components, suggesting that 102 Tmie traffics independently to hair bundles. Unexpectedly, GFP-tagged Tmcs fail to localize to 103 the hair bundle in *tmie* mutants, and overexpression of Tmie leads to a corresponding increase in 104 bundle expression of Tmc2b-GFP. To determine which regions of Tmie are involved in

105	regulating the Tmcs, we performed a domain analysis of <i>tmie</i> by expressing mutated or chimeric
106	transgenes of <i>tmie</i> in <i>tmie</i> <sup><i>ru1000</i></sup> , and made three key discoveries: ( <i>i</i> ) Tmie can function without
107	its putative first transmembrane domain, (ii) the remaining helix (2TM) and adjacent regions are
108	responsible for Tmie's function in hair cells, and (iii) dysfunctional tmie constructs have reduced
109	efficacy in localizing the Tmcs, supporting the conclusion that impaired MET is due to reduction
110	of Tmc protein. Our evidence suggests that Tmie's role in the MET complex is to promote
111	localization of Tmc1/2 to the site of MET in zebrafish sensory hair cells.

112

113 **Results** 

#### 114 Gross morphology is normal in *tmie*<sup>*ru1000*</sup> mutant zebrafish

115 The literature on TMIE's role in sensory hair cells is somewhat contradictory. Earlier 116 studies proposed a developmental role for TMIE [20-22], while later studies evidenced a role in 117 MET [25, 26]. To begin our analysis and attempt to clarify the issue in zebrafish, we examined live *tmie*<sup>*ru1000*</sup> larvae at 5-7 dpf using confocal microscopy. The *ru1000* allele harbors a nonsense 118 119 mutation leading to an N-terminal truncation, L25X [21]. We observed that mature hair cells of 120  $tmie^{ru1000}$  larvae were grossly normal compared to wild type siblings in both the inner ear cristae 121 and the lateral line organ, an organ specific to fish and amphibians (Fig 1A). We noted a slight 122 thinning of the mutant hair bundles, as revealed using a transgene Actin-GFP. Thin bundles have 123 been observed in other zebrafish MET mutants, such as those carrying mutations in ap1b1 and 124 *tomt.* Both genes have been previously implicated in protein trafficking in hair cells, with *tomt* 125 having a specific role in targeting Tmc1/2 proteins to the hair bundle [11, 33].

#### 127 Tmie-deficient zebrafish are deaf due to a defect in hair cell mechanosensitivity

128 Next, we used an assay for the auditory evoked behavior response (AEBR) to quantify hearing loss in *tmie<sup>ru1000</sup>* mutants. We exposed 6 dpf larvae to a loud pure tone stimulus (157 dB, 129 130 1000 Hz, 100 ms) once every 15 seconds for three minutes and recorded their startle responses 131 (sample traces in Fig 1B). Larvae deficient in *tmie* appear to be profoundly deaf, with little to no 132 response as compared to wild type siblings (Fig 1B and 1C). We then determined basal 133 (unevoked) hair cell activity of *tmie*<sup>ru1000</sup> larvae using FM 1-43 or FM 4-64. Both are vital dyes 134 that permeate open channels, making them useful as a proxy measure of the presence of active 135 MET channels in hair cells [27, 28, 34]. A 30-second bath application of FM dye readily labels 136 hair cells of the lateral line organ, which are arranged in superficial clusters called neuromasts. We briefly exposed wild type and *tmie<sup>ru1000</sup>* larvae to FM dye and then imaged the neuromasts 137 (Fig 1D). Consistent with previous findings [21, 22, 26], *tmie*<sup>ru1000</sup> neuromasts have a severe 138 139 reduction in FM labeling, suggesting that these hair cells have a MET defect (Fig 1F). To 140 characterize mechanically evoked responses of hair cells, we recorded extracellular potentials, or 141 microphonics (Fig 1H). Using a piezo actuator, we applied a 200 Hz sine wave stimulus to 3 dpf 142 larvae while simultaneously recording voltage responses from hair cells of the inner ear. In 143 agreement with results from our FM dye assay and with microphonic recordings previously reported [21], microphonics are absent in *tmie*<sup>*ru1000*</sup> larvae (Fig 1H, gray trace). 144 145

# 146 Transgenic *tmie-GFP* rescues the functional defect in *tmie<sup>ru1000</sup>* mutants

To rescue mechanosensitivity in *tmie<sup>ru1000</sup>* larvae, we generated a construct of *tmie* tagged
with GFP on its C-terminus, then expressed this transgene using a hair cell-specific promoter, *myosin 6b (myo6b)*. Stably expressed Tmie-GFP rescued the FM labeling in *tmie<sup>ru1000</sup>* hair cells

150 (Fig 1E and 1F). Tmie-GFP also restores microphonic potentials to wild-type levels (Fig 1H, 151 orange trace). In a stable line with a single transgene insertion, we observed that Tmie-GFP 152 expression varies among hair cells, even within the same patch of neuroepithelium (lateral crista, 153 S1A Fig). Immature hair cells, which can be identified by their shorter stereocilia and kinocilia 154 (S1A Fig, bracket and arrow, respectively), consistently show a bright and diffuse pattern of 155 labeling. This high expression level in immature bundles is characteristic of transgenes expressed 156 using the *myo6b* promoter, which drives expression more strongly in young hair cells [17, 34]. In 157 mature hair cells, expression patterns of Tmie-GFP are variable. At high expression levels, Tmie-158 GFP is enriched in the bundle in a broader pattern (S1B Fig). At reduced levels, the GFP signal 159 is concentrated at the beveled edge of the hair bundle (S1C Fig). At very low levels, we can 160 observe puncta along the stereocilia staircase, consistent with localization at stereocilia tips (S1D 161 Fig). We suspect that the diffuse "bundle fill" pattern is due to overexpression, and that lower 162 levels of Tmie-GFP recapitulate the endogenous localization at the site of MET, as previously 163 observed in mice [25].

164

#### 165 **Tmie-GFP is capable of trafficking without other members of the MET complex**

Having confirmed that our exogenously expressed Tmie-GFP is functional, we used this transgene to probe Tmie's role in the MET complex. First, we characterized Tmie's interactions with other MET proteins *in vivo* by expressing transgenic Tmie-GFP in mutant *pcdh15a*, *lhfpl5a*, and *tomt* larvae (Fig 2). Because a triple knock-out of zebrafish *tmc* has not been reported, we used *tomt* mutants as a proxy for *tmc*-deficient fish based on recent studies of defective bundle localization of the Tmcs in *tomt*-deficient fish and mice [11, 35]. As in wild type bundles (Fig 2A), Tmie-GFP is detectable in the stereocilia in each of these MET mutants (Fig 2B and 2D),

even if hair bundles are splayed (Fig 2B and 2C, arrowheads). This result suggests that Tmie
does not depend on interactions with other MET components for entry into the hair bundle.

175

#### 176 Tmc1-GFP and Tmc2b-GFP fail to localize to stereocilia without Tmie

177 To determine if the loss of Tmie affects the other components of the

178 mechanotransduction complex, we expressed GFP-tagged mechanotransduction proteins

- 179 (Pcdh15aCD3, Lhfpl5a, Tmc1, and Tmc2b) in *tmie<sup>ru1000</sup>* mutants. Both Pcdh15aCD3-GFP (Fig.
- 180 3A) and GFP-Lhfpl5a (Fig 3B) showed GFP fluorescence in hair bundles with a punctate

181 distribution, similar to the pattern seen in wild type bundles. This result is consistent with the

182 intact morphology of *tmie*<sup>*ru1000*</sup> hair bundles. However, when we imaged Tmc1-GFP (Fig 3C) and

183 Tmc2b-GFP (Fig 3E), GFP fluorescence was severely reduced in the hair bundles of *tmie*<sup>*ru1000*</sup>

mutants. In mature  $tmie^{ru1000}$  hair cells, we often saw a signal within the apical soma near the

185 cuticular plate, indicative of a trafficking defect (Fig 3E, arrows; position of cuticular plate

186 denoted in Fig 1G). We quantified Tmc expression in the hair bundle region and observed a

187 striking and consistent reduction in *tmie* mutants (Fig 3D and 3F). Previously, we reported that

188 localization of transgenic Tmc-GFP is unaffected in *pcdh15a* mutants [11], demonstrating that

189 mislocalization of Tmc1/2 is not a hallmark of all MET mutants.

190

#### 191 Overexpression of Tmie increases bundle localization of Tmc2b-GFP

We hypothesized that if the loss of Tmie reduces Tmc localization in the hair bundle,
then overexpression of Tmie may have the opposite effect. To test the consequence of
overexpression of Tmie on Tmc localization, we created a second construct of *tmie* coupled with *p2A-NLS(mCherry)* driven by the *myo6b* promoter. The p2A linker is a self-cleaving peptide,

196	which leads to translation of equimolar amounts of Tmie and NLS(mCherry). Hence, mCherry
197	expression in the nucleus denotes Tmie expression in the cell (Fig 3G, lower panels). We
198	generated a stable <i>tmie</i> <sup><i>ru1000</i></sup> fish line carrying the <i>tmie</i> - <i>p2A</i> - <i>NLS</i> ( <i>mCherry</i> ) transgene and then
199	crossed it to the $Tg(myo6b:tmc2b-GFP)$ ; $tmie^{ru1000}$ line. We observed that overexpression of
200	Tmie led to a 2.5-fold increase in expression of Tmc2b-GFP in the bundles of hair cells when
201	compared to wild type siblings that carried only the <i>tmc2b-GFP</i> transgene (Fig 3G and 3H).
202	Combined with the finding that Tmc expression is lost in hair bundles lacking Tmie, our data
203	suggest that Tmie positively regulates Tmc localization to the hair bundle.
204	
205	Transgenes can effectively determine protein functionality
206	To gain a better understanding of Tmie's role in regulating the Tmcs, we characterized a
207	new allele of <i>tmie</i> , t26171, which was isolated in a forward genetics screen for balance and
208	hearing defects in zebrafish larvae. Sequencing revealed that $tmie^{t26171}$ fish carry an A $\rightarrow$ G
209	mutation in the splice acceptor of the final exon of <i>tmie</i> , which leads to use of a nearby cryptic
210	splice acceptor (S2A Fig, DNA, cDNA). Use of the cryptic acceptor causes a frameshift that
211	terminates the protein at amino acid 139 (A140X), thus removing a significant portion of the C-
212	terminal tail (S2A Fig, Protein). Homozygous mutant larvae exhibit severe auditory and
213	vestibular deficits, being insensitive to acoustic stimuli and unable to maintain balance (S2A Fig,
214	Balance). FM 4-64 labeling of <i>tmie</i> <sup>t26171</sup> mutant hair cells suggests that the effect of the mutation
215	is similar to the ru1000 mutation (S2B and S2D Figs). This finding implicates the C-terminal
216	tail, a previously uncharacterized region, in Tmie's role in MET. However, when we
217	overexpressed a near-mimic of the predicted protein product of <i>tmie</i> <sup>t26171</sup> (1-138-GFP) using the

218 *myo6b* promoter, we observed full rescue of FM labeling defects in *tmie<sup>ru1000</sup>* (S2C and S2D

Figs), as well as behavioral rescue of balance and acoustic sensitivity (n=19). These results

220 revealed that when expressed at higher levels, loss of residues 139-231 does not have a

significant impact on Tmie's ability to function.

222 This paradoxical finding highlighted an important advantage of the use of transgenes over 223 traditional mutants. There are myriad reasons why a genomic mutation may lead to dysfunction, 224 including reduced transcription or translation, protein misfolding and degradation, or 225 mistrafficking. Exogenous expression may overcome these deficiencies by producing proteins at 226 higher levels. Moreover, the use of transgenes enabled us to carry out a more comprehensive 227 structure/function study of Tmie. To test a collection of deletions and chimeras of Tmie, we 228 therefore used the *myo6b* promoter to drive exogenous expression of the constructs in hair cells 229 of the *tmie*<sup>ru1000</sup> mutant.

230 We systematically deleted or replaced regions of *tmie* to generate 12 unique *tmie* 231 constructs (Fig 4A). Earlier studies in zebrafish and mice proposed that Tmie undergoes 232 cleavage, resulting in a single-pass mature protein [21, 36]. To test this hypothesis, we generated 233 the SP44-231 construct of Tmie, which replaced the N-terminus with a known signal peptide 234 (SP) from a zebrafish Glutamate receptor protein (Gria2a). The purpose of the unrelated signal 235 peptide was to preserve the predicted membrane topology of Tmie. We also made a similar 236 construct that begins at amino acid 63, where the sequence of Tmie becomes highly conserved 237 (SP63-231). Three of the constructs contained internal deletions ( $\Delta 63-73$ ;  $\Delta 97-113$ ;  $\Delta 114-138$ ). 238 In three more constructs, we replaced part of or the entire second transmembrane helix (2TM) 239 with a dissimilar helix from the CD8 glycoprotein (CD8; CD8-2TM; 2TM-CD8). We included 240 our mimic of the zebrafish *tmie*<sup>t26171</sup> mutant, which truncates the cytoplasmic C-terminus (1-241 138). To further truncate the C-terminus, we made a construct that mimics the mouse  $sr^{J}$  mutant

(1-113). In mice, this truncation recapitulates the full-deletion phenotype [20]. Finally, we
included an alternate isoform of Tmie that uses a different final exon, changing the C-terminal
sequence (*Tmie-short*). This isoform is found only in zebrafish [21] and its function has not been
explored.

246

# Subcellular localization of mutated or chimeric Tmie reveals domains required for selflocalization to the bundle

249 We first determined the subcellular localization of each Tmie fusion protein. Plasmid 250 DNA was co-injected into *tmie<sup>ru1000</sup>* eggs with transposase to generate mosaic expression of the 251 constructs in a subset of hair cells. At 4-6 days post injection, we imaged individual hair cells 252 expressing each transgene (Fig 4B). To quantify the enrichment in the bundle versus soma, we 253 measured the integrated density of GFP fluorescence in a small central area of mature bundles 254 (Fig 4C, black oval) and separately in the plasma membrane or soma-enriched compartments 255 (Fig 4C, magenta oval). Correcting for area, we then divided the bundle values by the total 256 values (bundle/bundle + soma) and expressed this as a ratio (Fig 4D). Values closer to 1 are bundle enriched, while values closer to 0 are soma-enriched. We excluded the CD8-GFP 257 258 construct from further analyses because it was detected only in immature bundles (Fig 4B, CD8). 259 Localization fell into three broad categories: bundle-enriched, soma-enriched, and 260 equally distributed. Most of the fusion proteins were bundle-enriched, similar to full-length 261 Tmie-GFP expression (Fig 4B and 4D). Three constructs were trafficked to the bundle but also 262 expressed in the soma (SP63-231, CD8-2TM, 1-138). This result suggests that the deleted 263 regions in these constructs have some role in designating Tmie as a bundle-localized protein. 264 Also of note, the full replacement of the 2TM helix (CD8) was unable to maintain stable

265	expression in mature bundles. Half-TM replacements (CD8-2TM, 2TM-CD8) revealed that loss
266	of the first half of the helix affects trafficking, whereas alteration of the second half had no
267	effect. Only two constructs were some-enriched (Tmie-short and 1-113), suggesting an inability
268	to traffic to the bundle. These two transgenes were thus excluded from further analyses.
269	
270	FM labeling identifies functional regions in the second transmembrane domain and
271	adjacent residues of Tmie
272	To identify regions of Tmie involved in mechanosensitivity of hair cells, we measured
273	the functionality of the nine <i>tmie</i> constructs that showed hair bundle expression. As in Fig 1F, we
274	generated stable lines of each transgenic construct and quantified fluorescence in neuromasts
275	after exposure to FM 4-64 (Fig 5).
276	Of nine constructs examined, four showed wild type levels of FM fluorescence in
277	tmie <sup>ru1000</sup> neuromasts (Tmie, SP44-231, Δ114-138, and 1-138; Fig 5A and 5B). Two constructs
278	( $\Delta 97-113$ and $\Delta 63-73$ ) did not rescue above mutant levels of FM 4-64, although $\Delta 63-73$ showed
279	a non-significant increase in FM fluorescence. While residues 63-73 have not been characterized,
280	the $\Delta 97-113$ result is consistent with the findings of previous publications in humans and mice,
281	showing that mutations in this region impair hearing and hair cell function [23, 25]. Three
282	constructs were capable of partial rescue (SP63-231, CD8-2TM, and 2TM-CD8). Each one of the
283	five dysfunctional constructs altered part of a contiguous region of Tmie: the 2TM and
284	surrounding domains. These results highlight this region of Tmie as vital for function. To
285	determine whether any of the constructs also have a dominant effect on hair-cell function, we
286	compared FM label in wild type larvae with or without the individual transgenic <i>tmie</i> construct
287	(Fig 5D). SP63-231 and $\Delta 63$ -73, which had impaired rescue in <i>tmie<sup>ru1000</sup></i> , showed reduced FM

label in transgenic wild type cells (Fig 5C and 5D). Interestingly, these two dominant negative
constructs alter the extracellular region of Tmie.

290

#### 291 Recordings of mechanically evoked responses confirm that the second transmembrane

#### 292 domain and adjacent regions are required for hair cell function

293 Bath applied FM dye demonstrates the presence of permeable MET channels, but does 294 not reveal any changes in mechanically evoked responses in hair cells. Therefore, we also 295 recorded microphonics of mutant larvae expressing individual transgenes. For our recordings, we 296 inserted a recording pipette into the inner ear cavity of 3 dpf larvae and pressed a glass probe 297 against the head (Fig 6A). Using a piezo actuator to drive the probe, we delivered a step stimulus 298 at increasing driver voltages while recording traces in current clamp (Fig 6B). For each 299 transgenic *tmie* line, we measured the amplitude of the response at the onset of stimulus (Fig 6C-300 I). We limited our analysis to the lines expressing constructs that failed to fully rescue FM 301 labeling (Fig 6E-I). As positive controls, we used the full-length tmie line (Fig 65 C) and also 302 included SP44-231 (Fig 6D), encoding the cleavage product mimic. Both controls fully rescued 303 the responses in *tmie<sup>ru1000</sup>* larvae. Consistent with a reduction in labeling with FM dye, we found 304 that the microphonic responses were strongly or severely reduced in larvae expressing the SP63-231, A63-73, CD8-2TM, 2TM-CD8 and A97-113 constructs in the tmie<sup>ru1000</sup> background (Fig 6E-305 306 I). We also saw the same dominant negative effect in wild type larvae expressing transgenic 307 *SP63-231* or *∆63-73*.

# 309 Regions of Tmie that mediate hair-cell mechanosensitivity are also required for localizing 310 Tmc2b-GFP

After identifying functional regions of Tmie, we asked whether these regions are 311 312 involved in regulating Tmc localization. Therefore we quantified hair bundle expression of transgenic Tmc2b-GFP in hair cells of *tmie*<sup>ru1000</sup> mutant larvae stably co-expressing individual 313 314 transgenic tmie constructs (Fig 7B-H). As in Fig 3 G, we tagged our tmie constructs with p2A-315 NLS(mCherry) so that Tmc2b-GFP expression in the hair bundles could be imaged separately. 316 We examined SP44-231 and the five *tmie* constructs that yielded impaired mechanosensitivity. 317 Three constructs showed full rescue of Tmc2b-GFP levels in the bundle. The SP44-231 318 cleavage mimic produced highly variable levels, some in the wild type range, others increasing 319 Tmc2b-GFP expression above wild type (Figs 7B and 7H), as seen in overexpression of full-320 length Tmie (Figs 3H and 7A, right panel). We suspect that the exogenous Gria2a signal peptide 321 leads to variable processing of Tmie and thus contributes to this variability in Tmc2b-GFP 322 fluorescence. *tmie*<sup>ru1000</sup> larvae expressing the SP63-231 construct gave rise to values of Tmc2b-323 GFP fluorescence within the wild type range (Fig 7C and 7H). When we recorded microphonics 324 in these larvae, we found that co-overexpression of Tmc2b-GFP and SP63-231 resulted in better 325 functional rescue of *tmie<sup>ru1000</sup>* (S3A Fig) than when SP63-231 was expressed alone (Fig 6E). We 326 also determined that the microphonic potentials correlated with the levels of Tmc2b-GFP in the 327 bundles (S3B Fig). Likewise, the 2TM-CD8 construct also generated values of Tmc2b-GFP 328 fluorescence in the wild type range (Fig 7F and 7H). These larvae rescued microphonic 329 potentials to wild type levels (S3C Fig), unlike when 2TM-CD8 was expressed alone (Fig 6E). 330 Functional rescue again correlated with Tmc2b-GFP bundle levels (S3D Fig). These results 331 indicate that functional rescue in the SP63-231 and 2TM-CD8 lines is Tmc dose-dependent.

332	Of the three constructs with little to no functional rescue, CD8-2TM (Fig 7E and 7H) and
333	$\Delta$ 97-113 (Fig 7G and 7H) had severely reduced levels of Tmc2b-GFP in hair bundles. In
334	<i>tmie</i> <sup><i>ru1000</i></sup> expressing $\Delta 63-73$ , there was severely reduced but still faintly detectable Tmc2b-GFP
335	signal though, as with the functional rescue, this difference was not statistically significant (Fig
336	7D and 7H). The bulk of this signal was observed in immature bundles (Fig 7D, arrows, and S4
337	Fig), but there was some detectable Tmc2b-GFP signal in mature bundles (S4 Fig). Overall,
338	these results suggest that the level of functional rescue by the <i>tmie</i> constructs is correlated to the
339	amount of Tmc2b present in the hair bundle.

340

#### 341 **Discussion**

342 *TMIE* was first identified as a deafness gene in mice and humans [20, 23]. The predicted 343 gene product is a relatively small membrane protein containing a highly conserved amino acid 344 sequence near the second hydrophobic helix. Previous studies established that TMIE is required 345 for MET in hair cells [21, 25, 26] and is an integral member of the complex (Zhao et al., 2014). 346 How TMIE contributes to the function of the MET complex was not clear. Our comprehensive 347 structure-function analysis of Tmie revealed that the functional capacity of various *tmie* mutant 348 constructs is determined by their efficacy in localizing Tmc2b-GFP to the bundle, as summarized 349 in Fig 8A and modeled in Fig 8B. These findings unveil a hitherto unexpected role for Tmie in 350 promoting the localization of the putative channel subunits Tmc1 and Tmc2b to the site of MET. 351 These findings broaden our understanding of the assembly of the MET complex and point to a 352 pivotal role of Tmie in this process.

A previous study of the *ru1000* mutant suggested that Tmie's role in zebrafish hair cells was developmental, with mutant lateral line hair cells showing stunted kinocilia and the absence 355 of tip links [21]. In our hands we did not observe any gross morphological defects, and the 356 localization pattern and the levels of Pcdh15a and Lhfp15a were unaffected in *tmie<sup>ru1000</sup>* larvae. 357 This observation is consistent with intact hair bundle morphology; stereocilia that are splayed or 358 disorganized are a dominant feature of hair cells missing their tip links, as seen in *pcdh15a* or 359 *lhfpl5a* mutants [17, 32]. In TMIE-deficient mice, hair cell morphology is grossly normal up to 360 P7 [25, 26]. In agreement with a previous study in mice [25], our results indicate that  $tmie^{ru1000}$ 361 mutants are profoundly deaf due to ablation of MET in hair cells. We fully rescued this deficit in 362 zebrafish ru1000 mutants by exogenous expression of a GFP-tagged transgene of *tmie*. 363 Exogenous expression gave rise to variable levels of Tmie-GFP in hair bundles, with lower 364 levels revealing a punctate pattern expected for a member of the MET complex, and higher 365 expression levels leading to expression throughout the stereocilia. Excess Tmie-GFP did not 366 appear to cause adverse effects in hair cells, which is consistent with a previous study in the 367 circler mouse mutant [37].

368

#### 369 Tmie can localize to hair bundles independently

370 To determine the interdependence of trafficking of Tmie and the other MET components, 371 we examined the localization of Tmie-GFP in mutants of essential MET genes: the tip-link 372 protein *pcdh15a*; the accessory protein *lhfp15a*; and in *tomt* mutants (Fig 3). In both zebrafish and 373 mice, the secretory pathway protein Tomt is required for proper localization of the  $\text{Tmc}_{1/2}$  [11, 374 35]; the *tomt* mutant likely simulates the condition of a triple knockout of all three zebrafish *tmc* 375 genes (tmc1/2a/2b). Despite the absence of Pcdh15a, Lhfp15a, and Tmcs, we found that Tmie-376 GFP still traffics to the bundles of hair cells. This finding suggests that Tmie can localize 377 independently of the other proteins of the MET apparatus. This autonomy is an unusual feature

378 for membrane components of the MET complex. For example, PCDH15 largely requires 379 LHFPL5 for trafficking to the stereocilia [17, 18, 38], and depends on Cadherin 23 to maintain 380 its localization at the site of MET [17, 39]. LHFPL5 also requires PCDH15 to maintain 381 localization at the stereocilia tips [18, 38]. Thus, Tmie appears to be the exception to the rule of 382 co-dependent transport to the hair bundle. 383 Our results reveal that Tmie has distinct regions associated with self-localization and 384 function (Fig 8). Three constructs showed impaired targeting of Tmie to the bundle, namely 385 SP63-231, CD8-2TM, and the 1-138 construct, the last of which truncates the C-terminus. 386 Further manipulation to the C-terminus, either by removing more amino acids (1-113) or by 387 using an alternative final exon (*Tmie-short*), results in targeting of the protein to the plasma 388 membrane instead of the bundle. However, removing just a smaller internal segment has no 389 effect on bundle localization (Fig 4A and 4C,  $\Delta 114-138$ ). We suspect that the abundance of 390 charged residues in the C-terminus of Tmie (S2A Fig, Protein), as well as the regions altered in 391 SP63-231 and CD8-2TM, contribute to recognition by bundle trafficking machinery. 392 Mislocalization, however, did not necessarily correlate with functional rescue. Despite partial 393 mislocalization to the plasma membrane, the 1-138 construct showed full functional rescue. 394 Conversely, despite normal localization to the bundle,  $\Delta 97-113$  did not rescue function at all. 395 These results demonstrate that Tmie's functional role is separate from its ability to target to the 396 bundle. 397 398 Tmie promotes the levels of Tmc1/2 in the hair bundle

The regulatory role of Tmie with respect to the Tmcs is strongly supported by the
strikingly different effects of loss of Tmie versus overexpression of Tmie. When Tmie is absent,

401	so are the Tmcs; when Tmie is overexpressed, the level of Tmc2b in the bundle is boosted as
402	well (Fig 3C-H). These results disagree with a previous finding in mice showing that Myc-
403	TMC2 is present in hair bundles of TMIE-deficient cochlear hair cells [25]. This discrepancy
404	may be due to the use a cytomegalovirus promoter to drive high levels of expression of Myc-
405	TMC2 in an in vitro explant of cochlear tissue. Localization of TMC1 in Tmie-/- mice, which is
406	the predominant TMC protein in cochlear hair cells, was not reported. In addition, localization of
407	the TMCs in vestibular hair cells was not characterized in Tmie-/- mice. Thus, further
408	investigation is warranted to determine if the relationship between Tmie and the Tmcs uncovered
409	by our experiments is a conserved feature or is potentially dependent on the type of hair cell, as
410	MET components may vary among different cell types.
411	One important question is whether Tmie and the Tmcs can physically interact to form a
412	complex that is transported to the hair bundle. A direct interaction of the mouse $TMC1/2$ and
413	TMIE proteins was not detected in a heterologous system [25], however, our in vivo analysis
414	suggests the possibility of an indirect interaction. The deletions and chimeric forms of Tmie in
415	the present study highlight important motifs or regions of Tmie that are critical for localization of
416	the Tmcs to the hair bundle.
417	

418 The first hydrophobic helix of Tmie is dispensable

419 The membrane topology of Tmie has not been biochemically determined, however,
420 online *Phobius* software predicts an N-terminal signal peptide in mouse and human TMIE and a
421 transmembrane helix in zebrafish Tmie [40]. Interestingly, the orthologues in C. elegans or
422 Drosophila do not contain this first hydrophobic region of Tmie. Upon removal of this region,
423 we observed that SP44-231 behaved like full-length Tmie, with a comparable pattern of

localization and full functional rescue of *tmie*-deficient fish. In addition, SP44-231 rescues
Tmc2b-GFP bundle expression to wild type levels or higher. To our knowledge, these results are
the first *in vivo* evidence that Tmie can function without the putative first transmembrane
domain. Our study supports the notion that Tmie undergoes cleavage, resulting in a single-pass
membrane protein that functions in the MET complex (Fig 8B).

429

#### 430 The 2TM domain and adjacent regions of Tmie are functionally significant

431 The key functional domains of Tmie are located within and proximal to the remaining 432 transmembrane domain. We found that replacement of the entire transmembrane domain with an 433 exogenous membrane helix from the CD8 glycoprotein resulted in a protein that trafficked to the 434 bundles of immature hair cells but was not expressed in mature bundles. This finding 435 demonstrates that this domain is vital for stable localization of Tmie in mature hair cells. Half-436 chimeras of this domain revealed that the mislocalization effect is exclusive to the first half of 437 the helix, but that both halves are functionally significant (the first half more so than the second). 438 These results suggest that the transmembrane domain is critical for both Tmie's localization and 439 function in the MET complex.

Removal of the cytoplasmic amino acids 97-113, directly after the 2TM, leads to a normal localization pattern but complete loss of function. This region contains arginine residues that have previously been implicated in human deafness [23, 41-43]. Mimics of these mutations in mouse cochlear hair cells lead to altered MET currents, which has been attributed to a reduction in binding to PCDH15-CD2 [25]. Interestingly, one of the mouse mutations, R93W, resulted in loss of TMIE localization at the site of MET. In contrast to these findings, when we remove this entire intracellular region from zebrafish Tmie, it is still capable of localization in hair bundles. This result may reflect species differences in recognition sequences for traffickingmachinery.

449 The SP63-231 and  $\Delta$ 63-73 constructs both lack different segments of the extracellular 450 region of Tmie. These were the only two constructs with dominant negative effects, suggesting 451 that each construct successfully integrates into the MET complex and interferes or competes with endogenous Tmie. Both constructs only minimally rescue mechanosensitivity in *tmie*<sup>ru1000</sup> 452 453 mutants and are thus predicted to weaken the efficiency of the MET complex. Other constructs 454 such as the transmembrane chimeras also yield partial rescue but do not appear to affect the 455 function of endogenous Tmie in wild-type hair cells. These data suggest that the full 2TM 456 domain is required to produce the dominant negative effect on endogenous Tmie. Combined with 457 the finding that replacement of the 2TM with an unrelated helix causes instability of Tmie in 458 mature hair cells, we suggest that the 2TM is essential in integrating Tmie into the MET 459 complex.

460

#### 461 Impaired functionality corresponds to decreased Tmc expression

462 When co-expressed with Tmc2b-GFP, our Tmie constructs reveal a strong link between 463 function and Tmc bundle expression (Figs 7 and S3). In larvae expressing CD8-2TM and  $\Delta$ 97-464 113, both of which display little or no functional rescue, there is no detectable Tmc2b-GFP in the 465 hair bundle (Fig 7E, 7G and 7H). In addition to defects in targeting Tmcs to the hair bundle, our 466 data also suggest a role for Tmie in maintaining the levels of Tmc2b in stereocilia. The most dramatic effect on maintenance of Tmc signal in the bundle was seen in *tmie*<sup>ru1000</sup> larvae 467 468 expressing the  $\Delta 63-73$  construct. In these larvae, Tmc2b-GFP successfully traffics to the bundle 469 in immature hair cells (Fig 7D, arrows) but does not maintain strong expression in mature cells

470 (S4 Fig). Based on this data, we conclude that the first half of the transmembrane domain and the 471 intracellular residues 97-113 are involved in trafficking the Tmcs to the site of MET, while the 472 extracellular residues 63-73 stabilize Tmc expression in the MET complex (Fig 8B). 473 Surprisingly, SP63-231 and 2TM-CD8 rescue Tmc2b-GFP to wild type levels (Fig 7C, 7F and 7H), even though functional rescue of *tmie*<sup>*ru1000*</sup> by GFP-tagged versions was reduced in 474 475 both FM labeling experiments (Fig 5) and microphonic recordings of the inner ear (Fig 6). This 476 result hints at an additional role for Tmie in MET that is independent of Tmc trafficking. However, the low level of functional rescue in *tmie*<sup>*ru1000*</sup> mutants by these two constructs was 477 478 only observed in the background of endogenous levels of the Tmcs. When we co-expressed Tmc2b-GFP with either SP63-231 or 2TM-CD8, then the functional rescue of *tmie*<sup>ru1000</sup> 479 480 mechanosensitivity improved in a Tmc-dose-dependent manner (S3 Fig). Since co-expression of 481 Tmc2b-GFP can overcome the functional deficit in SP63-231 and 2TM-CD8, we propose that 482 residues 44-62 and the second half of the 2TM are important but not absolutely essential to 483 regulating Tmc bundle expression. This finding reinforces the significance of our findings with 484 the constructs  $\Delta 63-73$ , CD8-2TM, and  $\Delta 97-113$ , which still fail to rescue Tmc2b-GFP levels 485 even when Tmc2b-GFP is co-expressed.

Through a systematic *in vivo* analysis of *tmie* via transgenic expression, we identified
new functional domains of Tmie. We demonstrated a strong link between Tmie's function and
Tmc1/2 expression in the bundle. Evidence continues to mount that the Tmcs are subunits of the
MET channel, and our results implicate Tmie in promoting and maintaining the localization of
Tmc subunits at the site of MET. The precise mechanism underlying Tmie's regulation of the
Tmcs awaits further investigation.

### 493 Methods

494

#### 495 Zebrafish husbandry

- 496 Zebrafish (*Danio rerio*, txid7955) were maintained at 28°C and bred according to
- 497 standard conditions. All animal research was in compliance with guidelines from the Institutional
- 498 Animal Care and Use Committee at Oregon Health and Science University. In this study, the
- following zebrafish mutant lines were used: *tmie*<sup>*ru1000*</sup> [21], *tmie*<sup>*t26171*</sup>, *pcdh15a*<sup>*psi7*</sup> [9],
- 500 *lhfpl5a*<sup>tm290d</sup> [44],  $tmc2b^{sa8817}$  [11]. All zebrafish lines in this study were maintained in a
- 501 Tübingen or Top long fin wild type background. We examined larvae at 4-7 days post-
- 502 fertilization (dpf), of undifferentiated sex. For experiments involving single transgenes, non-
- 503 transgenic *tmie*<sup>*ru1000*</sup> heterozygotes were crossed to transgenic fish in the homozygous or
- 504 heterozygous *tmie*<sup>*ru1000*</sup> background. Mutants were genotyped by PCR and subsequent digestion
- 505 or DNA sequencing. Primers are listed in Table 1.

# **Table 1. List of primers used in this study.**

Primers f	or plasmic	l construction						
Plas	mid	Forward (5' - 3')	Reverse (5' - 3')					
pME-	Tmie	GGGGACAAGTTTGTACAAAAAAGCAGGCTCCAAAC ATGAGACGCGGGAGAAGAA	GGGGACCACTTTGTACAAGAAAGCTGGGTC TTTCTTCGCAGGCTTCTTGG					
pME-SP	44-231	GGGGACAAGTTTGTACAAAAAAGCAGGCTCCAAAC ATGATTTTGTCGGGTCTCCTTTTACCCGCGTTATGG GGACTGGCGCTCGGCCAGATACCAGACCCAGAGCT	same as pME-Tmie					
pME-SP	63-231	ACCTCAGAAACAGTGGTGTTTTGGGGA	GCCGAGCGCCAGTCCC					
pME-∆	63-73	TTATGGCAGGTTGTGGGCATTTTC	GACGGGGTCTGGCTTTTTCG					
pME-CD8	PCR 1	same as pME-Tmie	GAGAAGGACCCCACAAGTCCCGGCCAGGG GCGCCCAGATGTATCGAAGTCCCCAAAACA C					
p <b>_</b> 0.20	PCR 2	TTGTGGGGTCCTTCTCCTGTCACTGGTTATCACCCT TTACTGCAAATGCCGAATCCCAC	same as pME-Tmie					
pME-	PCR 1	ATGAGACGCGGGAGAAGAAGAGGGAAAATG	CCCACAAGTCCCGGCCAGGGGGCGCCCAGAT GTATCGAAGTCCCCAAAACACCACTGTTTC					
CD8-2TM	PCR 2	TGGCCGGGACTTGTGGGATCTTAGCAATAATAATTA CGCTCTGCTGCATCTTCAAATGCC	TTTCTTCGCAGGCTTCTTGGCACCTC					
pME-	PCR 1	same as pME-CD8-2TM	AAAGGGTGATAACCAGTGACAGGAGAAGGA CGAACATGGAGAAAATGCCCACAACCTGCC					
2TM-CD8	PCR 2	CCTTCTCCTGTCACTGGTTATCACCCTTTACTGCAA ATGCCGAATCCCACGGACG	same as pME-CD8-2TM					
pME-∆97-113		AGACTTGCTGCGAAAAATTATGCCAAC	GAAGATGCAGCAGAGCGTAATTATTATTGC					
pME-∆114-138		GCGGCAAAGGTTGAGGTGAAG	TTGCGCGTGCCGAGC					
pME-1-138		same as pME-Tmie	GGGGACCACTTTGTACAAGAAAGCTGGGTC GCCGGGCACCTCAG					
pME-1-113		same as pME-Tmie	GGGGACCACTTTGTACAAGAAAGCTGGGTC TTGCGCGTGCCGAG					
pME-Tmie-short		same as pME-Tmie	GGGGACCACTTTGTACAAGAAAGCTGGGTC AGTGCCAGGATTGGCTG					

Primers for RT-PCR				
To amplify:	Forward (5' - 3')	Reverse (5' - 3')		
t26171 cDNA	ATATGCCAACACATTGGAGACGGTGC	CCCTGAGGTGTGTGTGAGTGTTCA		
Tmie-short transcript	ATGAGACGCCCCAGAAGAAGAGGGAAAATGGCGAT G	TTAAGTGCCAGGATTGGCCGGTTCATCTTCT TCCCTG		

Primers for identifying mutants					
Mutant Forward (5' - 3') Reverse (5' - 3')					
ru1000	TGTTTCGTCCAGGCTGAAG	GGCCTCATAAAACACAAGCA			
psi7	TTGGCACCACTATCTTTACCG	ACAGAAGGCACCTGGAAAAC			
tm290d	TGGTCTTCATCCAGCCCTAC	CGATCAGCAGCAAAGAGATG			
tk256c	TGTGTATTGCAGGTCAGTGTTG	AAGCGTTTTTCTGGGTGTTG			
t26171	GCACAGCCCTAATGGATACAG	GCTTCTTTCTTTGGTGCTCCT			

#### 508 Gene accession numbers for mutants and transgenes

- 509 *tmie* (accession no. F1QA80), *tmc1* (accession no. F1QFU0), *tmc2b* (accession no.
- 510 F1QZE9), tomt (accession no. A0A193KX02), pcdh15a (accession no. Q5ICW6), lhfpl5a
- 511 (accession no. F1Q837), *actba* (accession no. Q7ZVI7).
- 512

#### 513 Transgenic lines and plasmid construction

514 The following previously published transgenic lines were used:  $Tg(-6myo6b:\beta-actin-$ 

515 *GFP-pA*) [34], *Tg*(-6*myo6b:pcdh15aCD3-mEGFP-pA*) [17], and *Tg*(-6*myo6b:GFP-lhfpl5a-pA*),

516 *Tg*(-6*myo6b:Tmc1-mEGFP-pA*), *Tg*(-6*myo6b:Tmc2b-mEGFP-pA*) [11].

517 To generate the *tmie* expression vectors, we used the Tol2/Gateway system [45]. The

518 pDestination vector contained either a *cmlc2:GFP* heart marker or α-ACry:mCherry eye marker

519 for sorting. pDESTtol2pACrymCherry was a gift from Joachim Berger and Peter Currie

520 (Addgene plasmid # 64023, [46]).

521 The 5' entry vector contained the promoter for the *myosin 6b* gene, which drives

522 expression only in hair cells. All *tmie* transgenic constructs were subcloned into the middle entry

523 vector using PCR or bridging PCR and confirmed by sequencing. The primers for each vector

are listed in Table 1. For GFP-tagging, we used a 3' entry vector with a flexible linker

525 (GHGTGSTGSGSS) followed by *mEGFP*. For *NLS(mCherry)* experiments, a p2A self-cleaving

526 peptide (GSGATNFSLLKQAGDVEENPGP) was interposed between the *tmie* construct and the

527 *NLS(mCherry)*. This causes translation of a fusion protein that is subsequently cleaved into the

528 two final proteins. The 2TM helix replacements from residues 21-43 result in the following

529 chimeric helices: CD8 (YIWAPLAGTCGVLLLSLVITLYC), CD8-2TM

530 (YIWAPLAGTCGILAIIITLCCIF), and 2TM-CD8 (LWQVVGIFSMFVLLLSLVITLYC).

531	Multisite (	Gateway	LR reactions	[47.4	481	were	performed	to	generate	the	foll	owing

532 constructs: pDest(-6myo6b:tmie-GFP-pA), pDest(-6myo6b:tmie-short-GFP-pA), pDest(-

533 6myo6b:SP44-231-GFP-pA), pDest(-6myo6b:SP63-231-GFP-pA), pDest(-6myo6b:∆63-73-

- 534 GFP-pA), pDest(-6myo6b:CD8-GFP-pA), pDest(-6myo6b:CD8-2TM-GFP-pA), pDest(-
- 535 6myo6b:2TM-CD8-GFP-pA), pDest(-6myo6b:Δ97-113-GFP-pA), pDest(-6myo6b:Δ114-138-
- 536 GFP-pA), pDest(-6myo6b:1-113-GFP-pA), pDest(-6myo6b:1-138-GFP-pA), pDest(-
- 537 6myo6b:tmie-p2A-NLS(mCherry)-pA), pDest(-6myo6b:SP63-231-p2A-NLS(mCherry)-pA),
- 538 pDest(-6myo6b: \Delta63-73-p2A-NLS(mCherry)-pA), pDest(-6myo6b: CD8-2TM-p2A-
- 539 NLS(mCherry)-pA), pDest(-6myo6b:  $\Delta$ 97-113-p2A-NLS(mCherry)-pA).

540 To generate transgenic fish, plasmid DNA and tol2 transposase mRNA were co-injected 541 into single-cell fertilized eggs, as previously described (Kwan et al., 2007). For each construct, 542 200+ eggs from an incross of  $tmie^{ru1000}$  heterozygotes were injected. To obtain stable transgenic 543 lines, >24 larvae with strong marker expression were raised as potential founders. For each GFP-544 tagged transgene, at least two founder lines were generated and examined for visible bundle 545 expression. For each *tmie* construct, we isolated a line containing single transgene insertions, 546 with the exception of the CD8-2TM construct in which we identified a single founder with high 547 transmission of the transgene (>10%) and used these offspring and their siblings for FM and 548 microphonics experiments. For NLS(mCherry) experiments, injected fish were raised to 549 adulthood and genotyped to identify *tmie<sup>ru1000</sup>* heterozygotes and homozygotes. We identified 550 founders for each construct and then crossed these founders to *tmie<sup>ru1000</sup>* heterozygotes carrying 551 Tg(myo6b:tmc2b-GFP). This generated offspring that expressed both transgenes in the tmie<sup>ru1000</sup> 552 mutant background, and we used these larvae for experiments. In SP44-231, SP63-231, and

553 *CD8-2TM*, stable transgenic lines were generated from the founder before experiments were 554 carried out.

555

#### 556 Microscopy

557 Live larvae were anesthetized with E3 plus 0.03% 3-amino benzoic acid ethylester 558 (MESAB; Western Chemical) and mounted in 1.5% low-melting-point agarose (Sigma-Aldrich cas. # 39346-81-1), with the exception of the morphology images from Fig 1A and Fig 6A in 559 560 which larvae were pinned with glass rods and imaged in E3 or extracellular solution containing 561 MESAB. The image in Fig 6A was captured at room temperature using a Hamamatsu digital 562 camera (C11440, ORCA-flash2.8), MetaMorph Advanced NX software, and an upright Leica 563 DMLFS microscope. We used differential interference contrast (DIC) with a Leica HC PL 564 Fluotar 10x/0.3 lens. For all imaging except Fig 6A, images were captured at room temperature 565 using an Axiocam MrM camera, Zeiss Zen software, and an upright Zeiss LSM700 laser-566 scanning confocal microscope. We used DIC with one of two water-immersion lenses: Plan 567 Apochromat 40x/1.0 DIC, or Acroplan 63x/0.95 W. Laser power and gain were unique for each 568 fluorophore to prevent photobleaching. We averaged 2 or 4x for each image, consistent within 569 each experiment. The Tmc1-GFP and Tmc2b-GFP transgenes are very dim, and high laser power 570 (4%) and gain (1100) were necessary. At these settings, autofluorescence from other 571 wavelengths can falsely enhance the emission peak at 488. To filter out this autofluorescence, we 572 simultaneously collected light on a second channel with an emission peak at 640 nm. 573

#### 574 Auditory Evoked Behavioral Response (AEBR)

575 Experiments were conducted as previously described [49]. Wild type and mutant larvae 576 were sorted by FM 1-43 labeling. Briefly, 6 dpf larvae were placed in six central wells of a 96-577 well microplate mounted on an audio speaker. Pure tones were played every 15 s for 3 min 578 (twelve 100 ms stimuli at 1 kHz, sound pressure level 157 dB, denoted by asterisks in Fig 1B). 579 Responses were recorded in the dark inside a Zebrabox monitoring system (ViewPoint Life 580 Sciences). Peaks represent pixel changes from larval movement. A response was considered 581 positive if it occurred within two seconds after the stimulus and surpassed threshold to be 582 considered evoked, not spontaneous (Fig 1B, green indicates movement detected, magenta 583 indicates threshold surpassed). For each larva, we used the best response rate out of three trials. 584 Response was quantified by dividing the number of positive responses by total stimuli (12) and 585 converting to a percent. If the larvae moved within two seconds before a stimulus, that stimulus 586 was dropped from the trial data set (i.e. the number of total stimuli would become 11). Each data 587 point on the graph in Fig 1C is the percent response of an individual larva. We used a two-tailed 588 unpaired t-test with Welch's correction to determine significance, \*\*\*\*p<0.0001.

589

590 FM 1-43 and FM 4-64 labeling

591 Larvae were briefly exposed to E3 containing either 3µM N-(3-

592 Triethylammoniumpropyl)-4-(4-(Dibutylamino)styryl)Pyridinium Dibromide (FM 1-43, Life

593 Technologies) or 3µM of the red-shifted N[scap]-(3-triethylammoniumpropyl)-4-(6-(4-

594 (diethylamino)phenyl)hexatrienyl)pyridinium dibromide (FM4-64; Invitrogen). After exposure

595 for 25-30 seconds, larvae were washed 3x in E3. Laser power was adjusted for each experiment

to avoid saturation of pixels but was consistent within a clutch. FM levels were quantified in

597 ImageJ [50] as described previously [9]. In brief, maximum projections of each neuromast were 598 generated using seven optical sections, beginning at the cuticular plate and moving down through 599 the soma (magenta bracket, Fig 1G). We then measured the integrated density of the channel 600 with an emission peak at 640 nm for FM 4-64, and at 488 nm for FM 1-43. This integrated 601 density value was divided by the number of cells, thus converting each neuromast into a single 602 plot point of integrated density per cell (IntDens/cell). Statistical analyses were always 603 performed between direct siblings. For Fig 5, individual values were divided by the mean of the sibling wild type neuromasts in order to display the data as a percent of wild type, making it 604 605 easier to compare across groups. Statistical significance was determined within an individual 606 clutch using one-way ANOVA.

607

#### 608 Microphonics

609 Larvae at 3 dpf were anesthetized in extracellular solution (140mM NaCl, 2mM KCl, 610 2mM CaCl<sub>2</sub>, 1mM MgCl<sub>2</sub>, and 10mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid 611 (HEPES); pH 7.4) containing 0.02% 3-amino benzoic acid ethylester (MESAB; Western 612 Chemical). Two glass fibers straddled the yolk to pin the larvae against a perpendicular cross-613 fiber. Recording pipettes were pulled from borosilicate glass with filament, O.D.: 1.5 mm, O.D.: 614 0.86 mm, 10 cm length (Sutter, item # BF150-86-10, fire polished). Using the Sutter Puller 615 (model P-97), we pulled the pipettes into a long shank with a resistance of 10-20M $\Omega$ . We then 616 used a Sutter Beveler with impedance meter (model BV-10C) to bevel the edges of the recording 617 pipettes to a resistance of 3-6 M $\Omega$ . We pulled a second pipette to a long shank and fire polished 618 to a closed bulb, and then attached this rod to a piezo actuator (shielded with tin foil). The rod 619 was then pressed to the front of the head behind the lower eye, level with the otoliths in the ear

620 of interest, to hold the head in place while the recording pipette was advanced until it pierced the 621 inner ear cover. Although it has been demonstrated that size of response is unchanged by entry 622 point [51], we maintained a consistent entry point dorsal to the anterior crista and lateral to the 623 posterior crista (see Fig 6A). After the recording pipette was situated, the piezo pipette was then 624 moved back to a position in light contact with the head. We drove the piezo with a High Power 625 Amplifier (piezosystem jena, System ENT/ENV, serial # E18605), and recorded responses in 626 current clamp mode with a patch-clamp amplifier (HEKA, EPC 10 usb double, serial # 550089). 627 Each stimulus was of 20 ms duration, with 20 ms pre- and post-stimulus periods. We used either 628 a sine wave or a voltage step and recorded at 20 kHz, collecting 200 traces per experiment. In 629 Fig 1H, we used a 200 Hz sine wave at 10V, based on reports that 200 Hz elicited the strongest 630 response [52]. In Fig 6, we used multiple step stimuli at varying voltages (2V, 3V, 4V, 5V, 6V, 631 and 10V). The piezo signal was low-pass filtered at 500Hz using the Low-Pass Bessel Filter 8 632 Pole (Warner Instruments). Microphonic potential responses were amplified 1000x and filtered 633 between 0.1-3000 Hz by the Brownlee Precision Instrumentation Amplifier (Model 440). We 634 used Igor Pro for analysis. We averaged each set of 200 traces to generate one trace response per 635 fish, then measured baseline-to-peak amplitude. These amplitudes were used to generate the 636 graphs in Fig 6. Statistical significance was determined by 2-way ANOVA comparing all groups 637 to wild type non-transgenic siblings.

638

#### 639 Quantification of Tg(myo6b:Tmc1-GFP) and Tg(myo6b:Tmc2b-GFP) in the ROI

640 Using ImageJ, maximum projections of each crista were generated for analysis (5
641 sections per stack for Tmc1-GFP in Fig 3D and Tmc2b-GFP in Fig 3F, and 13 sections per stack
642 for Tmc2b-GFP in Figs 3H and 7). Quantification of Tmc-GFP bundle fluorescence in Fig 3 was

643	achieved by outlining each bundle to encompass the entire region of interest (ROI) in a single
644	hand-drawn area (Fig 7A, right panel, black outline). In the ROI, we quantified the integrated
645	density of the channel with an emission peak at 480 nm. This was repeated in the region above
646	the bundles containing only inner ear fluid and the kinocilia in order to subtract background
647	fluorescence. Each middle crista generated one data point on the graphs in Figs 3 and 7. In some
648	cases, we saw single cells that appeared to have a GFP-fill, probably due to clipping of the GFP
649	tag. We excluded these cells from analyses, since they falsely increased the signal. Due to the 3D
650	nature of the mound-shaped cristae, it was difficult to completely exclude the apical soma region,
651	leading the signals of <i>tmie</i> <sup>ru1000</sup> to average above zero. We used the Kruskal-Wallis test for the
652	SP44-231, SP63-231, and 2TM-CD8 constructs; all others are one-way ANOVA.
653	
654	cDNA generation by Reverse Transcription Polymerase Chain Reaction (RT-PCR)
655	We sorted 30 wild type and 30 t26171 larvae by behavior (tap sensitivity and balance
656	defect at 5 dpf) and extracted RNA using the RNeasy mini kit (Qiagen). Larvae were
657	homogenized using a 1ml syringe. To generate the cDNA for the short isoform of Tmie (Tmie-
658	short) and the t26171 allele, we performed RT-PCR on these RNA samples using the RNA to
659	cDNA EcoDry Premix (Clontech, Cat # 639549). Primers are listed in Table 1. Both transcripts
660	were verified by DNA sequencing.
661	

# 662 Acknowledgements

The authors thank Cecilia Toro, Lucille Moore, and Andre Dagostin for help with
execution and analysis of the microphonics experiments, as well as Larry Trussell, Josef Trapani,
and Anthony Ricci for advice. We thank Jim Hudspeth for the *tmie<sup>ru1000</sup>* fish line. We also thank

Eliot Smith for feedback on the manuscript, and Leah Snyder and Lisa Hiyashi for laboratorysupport.

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- 822
- 823 Fig Legends
- 824

# Fig 1. Zebrafish *tmie<sup>ru1000</sup>* mutants: phenotype and functional rescue by Tmie-GFP. All 825 826 confocal images are of live, anesthetized larvae. (A) Hair cells in the lateral-line neuromasts (7 827 dpf) and inner ear cristae (5 dpf) from wild type and *tmie<sup>ru1000</sup>* larvae. A transgene (Actin-GFP) 828 was used to visualize stereocilia bundles. (B) Sample traces from an auditory evoked behavior 829 response (AEBR) assay, performed on 6 dpf larvae over the course of 3 minutes. Pure tone 830 stimuli are indicated by asterisks. Peaks represent pixel changes due to larval movements 831 (magenta indicates positive response). (C) Quantification of AEBR displayed as box-and-832 whiskers plot; significance determined by unpaired t-test with Welch's correction. (D) Top-down 833 view of neuromasts from 4 dpf larvae after brief exposure to a vital dye, FM 1-43. FM 1-43 and 834 FM4-64 permeate open transduction channels. (E) Lateral view of a neuromast from a 4 dpf 835 *tmie*<sup>*ru1000*</sup> larva expressing transgenic Tmie-GFP, after exposure to FM 4-64. (F) Quantification 836 of FM 4-64 fluorescence/cell in 5 dpf larvae; significance determined by one-way ANOVA. (G) A cartoon depiction of a group of lateral-line hair cells viewed laterally, with close-up views of a 837 838 single cell at the bundle region. The dashed green line indicates the single plane containing the 839 stereocilia bundles. The magenta bracket indicates the area used to make the maximum 840 projections that were analyzed for FM fluorescence in (F). (H) Sample traces of extracellular 841 (microphonic) recordings, evoked from the inner ear of 3 dpf larvae. A piezo actuator was used 842 to stimulate larvae with a 200 Hz sine-wave mechanical stimulus using an 8 V driver voltage. All 843 statistics are mean $\pm$ SD, \*\*\*\*p<0.0001. Scale bars: 10µm.

844

845 Fig 2. Tmie-GFP is present in the hair bundles of MET mutants. Confocal images of the 846 bundle region in hair cells of the inner-ear lateral crista in live larvae. Larvae at 6 dpf expressing 847 transgenic Tmie-GFP in the genetic backgrounds of wild type (A), and homozygous mutants for the tip link protein Pcdh15a (B, pcdh15a<sup>psi7</sup>), the accessory protein Lhfpl5a (C, lhfpl5a<sup>tm290d</sup>), and 848 849 the Golgi-localized protein Tomt (D, tomt<sup>tk256c</sup>). Tomt-deficient fish lack Tmc expression in hair 850 cell bundles [11], presumably mimicking the condition of a triple Tmc knockout. Arrowheads 851 indicate splayed hair bundles. n=8 each genotype. Scale bar: 5µm. 852 853 Fig 3. Specific loss of Tmc1 and Tmc2b in tmieru1000 larvae. Maximum projections of the 854 hair bundle region (ROI) of hair cells in the lateral crista of the inner ear, collected from live 855 larvae using confocal microscopy. (A-B) 6 dpf larvae expressing either transgenic Pcdh15aCD3-856 GFP or GFP-Lhfpl5a (n=6 each genotype). (C) 3 dpf larvae expressing Tmc1-GFP. (D) Plot of 857 the integrated density of Tmc1-GFP fluorescence in the ROI; each data point represents one 858 crista. Statistical significance determined by two-tailed unpaired t-test with Welch's correction, 859 p=0.0002. (E) 4 dpf larvae expressing Tmc2b-GFP. The arrow points to the cuticular plate/apical 860 soma region, just below the ROI. (F) Plot of the integrated density of Tmc2b-GFP fluorescence 861 in the ROI. Statistical significance determined by two-tailed unpaired t-test with Welch's correction, p=0.0005. (G) 4 dpf *tmie*<sup>ru1000</sup> larva co-expressing two transgenes, *tmc2b-GFP* and 862 863 *tmie-p2A-NLS(mCherry)*. The p2A linker is a self-cleaving peptide that results in equimolar 864 expression of Tmie and nuclear mCherry. (H) Plot of the integrated density of Tmc2b-GFP 865 fluorescence/crista in the ROI; significance determined by one-way ANOVA. All statistics are 866 mean  $\pm$  SD. Scale bars: 5 $\mu$ m.

867

868	Fig 4. Schema for a systematic domain analysis of Tmie and subcellular localization of
869	Tmie constructs. (A) A linear diagram of 12 unique constructs of tmie used in our experiments.
870	Full-length Tmie is predicted to contain two hydrophobic helices or transmembrane domains
871	(1TM and 2TM). SP44-231 and SP63-231 replace part of the N-terminus with a signal peptide
872	(SP) from the Glutamate receptor 2a (in blue). In the CD8, CD8-2TM, and 2TM-CD8 constructs,
873	all or part of the 2TM is replaced by the helix from the CD8 glycoprotein (in yellow). Tmie-short
874	is a fish-specific isoform of Tmie that contains an alternate final exon (in orange). Dotted lines
875	represent internal deletions. (B) Representative confocal images of each construct being
876	expressed as a GFP-tagged transgene in hair cells of 4-6 dpf <i>tmie</i> <sup>ru1000</sup> larvae. Expression is
877	mosaic due to random genomic insertion into subsets of progenitor cells after single-cell
878	injection. The expression of the CD8 construct is shown in a neuromast, while all others are in
879	the inner ear middle crista. (C) The localization of each GFP fusion protein was determined by
880	measuring the fluorescence/area in the bundle (b) and soma (s), and then calculating b/(b+s). (D)
881	Enrichment in the hair bundle is displayed as a ratio for each construct, with 1 being completely
882	bundle-enriched and 0 being completely soma-enriched. Scale bar in (B): 5µm.
002	

883

# Fig 5. The second transmembrane and adjacent residues of Tmie are required for rescue of FM labeling. All images are a top-down view of a representative neuromast from 6 dpf larvae collected using confocal microscopy. The left image is a single plane through the stereocilia (green dashed line in Fig 1G) with DIC + GFP fluorescence. The right image is a maximum projection of the soma region (magenta bracket in Fig 1G) showing FM 4-64 fluorescence. (A) Representative images of neuromasts in *tmie<sup>ru1000</sup>* larvae, each stably expressing an individual

890	tmie construct. FM fluorescence was normalized to wild type non-transgenic larvae generated
891	with the Tmie-GFP line. (B) Box-and-whiskers plot of the integrated density of FM
892	fluorescence/cell in transgenic <i>tmie</i> <sup>ru1000</sup> compared to non-transgenic wild type and mutant
893	siblings for each construct. (C) Representative images of neuromasts in wild type larvae with or
894	without transgene. FM fluorescence was normalized to wild type non-transgenic larvae of the
895	Tmie-GFP line. (D) Box-and-whiskers plot of the integrated density of FM fluorescence/cell in
896	wild type neuromasts with and without transgene. Significance determined within each clutch by
897	one-way ANOVA, n≥9, **p<0.01, ***p<0.001, ****p<0.0001. Scale bars in (A) and (C) are
898	10µm.
899	
900	Fig 6. The second transmembrane and adjacent regions of Tmie are required for inner ear
900 901	<b>Fig 6.</b> The second transmembrane and adjacent regions of Tmie are required for inner ear microphonics. (A) A DIC image of a 3 dpf larva anesthetized and pinned (glass fiber) for inner
901	<b>microphonics.</b> (A) A DIC image of a 3 dpf larva anesthetized and pinned (glass fiber) for inner
901 902	<b>microphonics.</b> (A) A DIC image of a 3 dpf larva anesthetized and pinned (glass fiber) for inner ear recordings. Shown are a probe attached to a piezo actuator (piezo) pressed against the head
901 902 903	<b>microphonics.</b> (A) A DIC image of a 3 dpf larva anesthetized and pinned (glass fiber) for inner ear recordings. Shown are a probe attached to a piezo actuator (piezo) pressed against the head and a recording pipette pierced into the inner ear. (B) Traces from a wild type larva. A step
901 902 903 904	<b>microphonics.</b> (A) A DIC image of a 3 dpf larva anesthetized and pinned (glass fiber) for inner ear recordings. Shown are a probe attached to a piezo actuator (piezo) pressed against the head and a recording pipette pierced into the inner ear. (B) Traces from a wild type larva. A step stimulus for 20ms was applied; 200 traces were averaged for each of the six piezo driver
901 902 903 904 905	<b>microphonics.</b> (A) A DIC image of a 3 dpf larva anesthetized and pinned (glass fiber) for inner ear recordings. Shown are a probe attached to a piezo actuator (piezo) pressed against the head and a recording pipette pierced into the inner ear. (B) Traces from a wild type larva. A step stimulus for 20ms was applied; 200 traces were averaged for each of the six piezo driver voltages: 2V, 3V, 4V, 5V, 6V, and 10V. Gray box: magnification of the onset of response in
901 902 903 904 905 906	<b>microphonics.</b> (A) A DIC image of a 3 dpf larva anesthetized and pinned (glass fiber) for inner ear recordings. Shown are a probe attached to a piezo actuator (piezo) pressed against the head and a recording pipette pierced into the inner ear. (B) Traces from a wild type larva. A step stimulus for 20ms was applied; 200 traces were averaged for each of the six piezo driver voltages: 2V, 3V, 4V, 5V, 6V, and 10V. Gray box: magnification of the onset of response in individual traces. (C-I) Same protocol as in (B). Mean amplitude of the response peak ± SD as a

910

911 Fig 7. Effect of transgenic Tmie constructs on Tmc2b-GFP bundle localization. Confocal
912 images are maximum projections of representative inner-ear lateral cristae collected from 4 dpf

913	larvae. Upper panels show the bundle region, with all larvae stably expressing transgenic
914	Tmc2b-GFP (green). Lower panels show the soma region, with some larvae expressing
915	transgenic Tmie constructs tagged with p2A-NLS(mCherry). Nuclear mCherry (magenta) is a
916	marker for equimolar translation of the indicated Tmie construct. (A) Sibling wild type,
917	<i>tmie</i> <sup><i>ru1000</i></sup> , and <i>tmie</i> <sup><i>ru1000</i></sup> expressing full-length Tmie. For the quantification in (G), Tmc2b-GFP
918	fluorescence was measured within the ROI (right panel, black line). (B-G) tmie <sup>ru1000</sup> larvae
919	expressing individual Tmie constructs tagged with p2A-NLS(mCherry), as labeled. The arrows
920	in (D) point to Tmc2b-GFP in immature hair bundles. (H) Plot of the integrated density of
921	Tmc2b-GFP fluorescence in the ROI, comparing <i>tmie</i> <sup>ru1000</sup> larvae expressing a tmie construct
922	(magenta) to wild type (black) and <i>tmie</i> <sup>ru1000</sup> (gray) siblings not expressing tmie construct.
923	Significance for SP44-231 and SP63-231 was determined by the Kruskal-Wallis test, for all other
924	tmie constructs by one-way ANOVA, n≥6, ***p<0.001, ****p<0.0001. Scale bars: 10µm.

#### 926 Fig 8. Summary of experimental results for tmie constructs and model of discrete

- 927 **functional domains of Tmie.** (A) Symbols as follows: enhanced above wild type (++),
- 928 comparable to wild type (+), partially reduced (+/-), and severely reduced or absent (-). For
- 929 *dominant negative effect*, the effect is present (+) or absent (-). Blank spaces are not determined.
- 930 Refer to Fig 4A for details on constructs. (B) A model of the protein sequence of zebrafish Tmie.
- Amino acids 1-43 are separated due to suspected cleavage as a signal peptide. Although shown
- as a TM domain, it is unclear whether the first hydrophobic region forms a helix. Note that the
- 933 extracellular region (yellow glow) was never deleted in its entirety; SP63-231 deleted
- 934 QIPDPELLPTDPPKKPDPV, and  $\Delta 63-73$  deleted TSETVVFWGLR. Also note that the TM
- domain with orange lettering only had an effect on the stability of Tmie when the entire helix

936 was substituted.

937

# 938 Supporting information

939

940 S1 Fig. Tmie-GFP shows variable expression in stereocilia. Representative images of the 941 lateral crista in a wild type larva at 6 dpf, generated using confocal microscopy. (A) The hair 942 bundle region of hair cells expressing transgenic tmie-GFP driven by the myo6b promoter. The 943 arrow and bracket show, respectively, the short kinocilium and stereocilia bundle of an immature 944 hair cell. (B) A single hair bundle with "bundle fill" expression pattern produced by 945 overexpression of Tmie-GFP. (C) A single bundle with Tmie-GFP concentrated along the 946 beveled edge of the stereocilial staircase. (D) A single bundle with punctate expression of Tmie-947 GFP suggestive of localization at the site of MET. Scale bar in (A):  $5\mu$ m, in (D):  $2\mu$ m.

## 949 S2 Fig. Differential effects on function with a genomic mutation and a transgene mimic. (A)

950 Data for a novel mutant allele of tmie, t26171. DNA: Chromatographs of the DNA sequence of 951 tmie in wild type (above) and *tmie*<sup>t26171</sup> (below) showing the genomic region where the mutation 952 occurs. An arginine is mutated to guanine in the splice acceptor (black box, above) of the final 953 exon of tmie, exon 4. The dashed black box below indicates the mutated original splice acceptor 954 site. Use of a cryptic splice acceptor (black box, below) 8 nucleotides downstream causes a 955 frameshift and an early stop codon (\*). cDNA: Chromatograph of the DNA sequence from RT-PCR of *tmie*<sup>t26171</sup> larvae bridging exons 3 and 4. Protein: The predicted protein products, shown 956 957 here as a two-pass transmembrane protein. The wild type protein has many charged residues 958 (positive in light gray, negative in dark gray) that are lost in *tmie*<sup>t26171</sup>. Balance: Photos of wild 959 type and *tmie*<sup>t26171</sup> larvae, taken with a hand-held Canon camera. Arrow points to a larva that is 960 upside-down, displaying a classic vestibular phenotype. (B) Top-down view of a representative 961 neuromast after exposure to FM 4-64, imaged using confocal microscopy. The first panel is a 962 single plane through the soma region while the second panel is a maximum projection of 7 963 panels through the soma region, beginning at the cuticular plate (as denoted by magenta bracket 964 in Fig 1G). (C) Same as (B) except that the first panel shows the bundle region so that 1-138-965 GFP can be visualized in bundles (as depicted by dashed green line, Fig 1G). The transgene is 966 driven by the myo6b promoter. (D) Plot of the integrated density of FM fluorescence per cell as a percent of wild type siblings. Displayed wild type and *tmie<sup>ru1000</sup>* data are from siblings of Tg(1-967 968 138-GFP); tmie<sup>ru1000</sup>, while tmie<sup>t26171</sup> data are from a separate experiment. Statistical significance 969 determined by one-way ANOVA, \*\*\*\*p<0.0001. Scale bar: 10µm.

# 971 S3 Fig. Functional rescue of tmieru1000 by constructs SP63-231 and 2TM-CD8 is Tmc

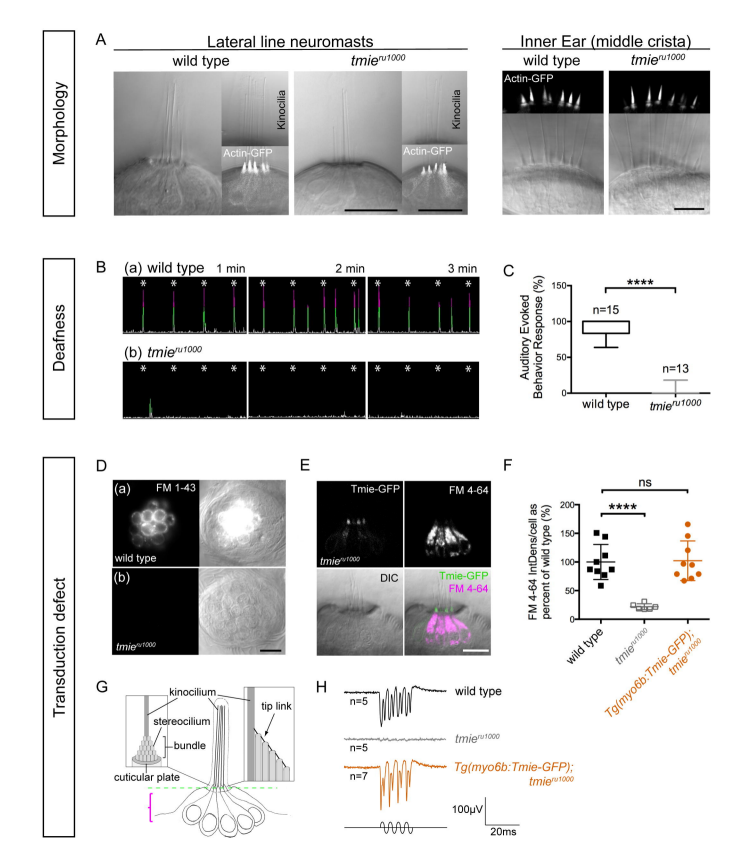
- 972 **dose-dependent.** (A) Mean amplitude of the response peak  $\pm$  SD as a function of the stimulus
- 973 intensity of the driver voltage, as described in Fig 6B. (B) XY plot of the amplitude of
- 974 microphonic response vs the integrated density of Tmc2b-GFP fluorescence in the ROI. A 10V
- 975 step stimulus was used to evoke microphonic potentials. The line is a linear regression,  $R^2$ =
- 976 0.5216. (C) Same as (A) for the 2TM-CD8 construct. (D) Same as (B) for the 2TM-CD8
- 977 construct,  $R^2 = 0.7726$ . Measurements are from 4 dpf larvae.
- 978

# 979 S4 Fig. In Δ63-73, Tmc2b-GFP traffics to bundles but does not maintain high expression in

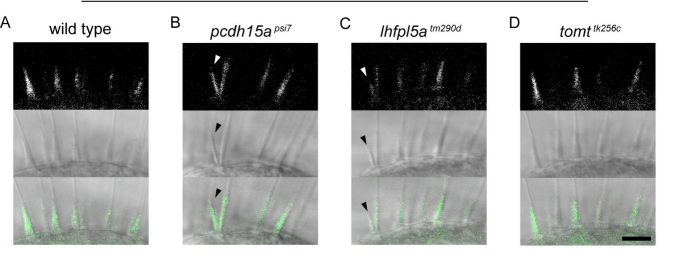
980 mature cells. Confocal images of single hair bundles from cells expressing transgenic Tmc2b-

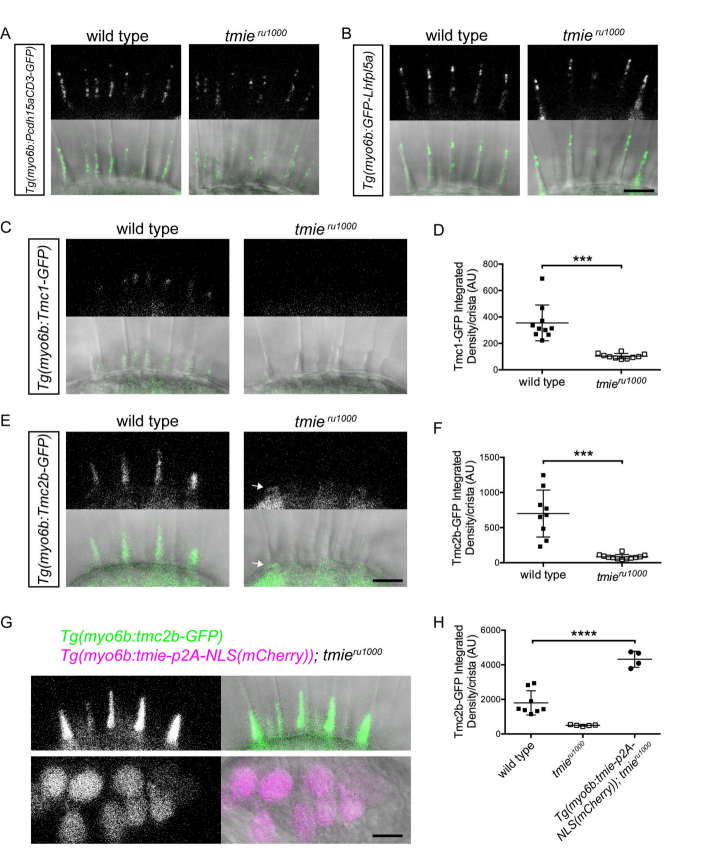
981 GFP driven by the myo6b promoter. Brackets show the stereocilia bundle, which is shorter in

- 982 immature hair cells (white brackets) and longer in mature ones (black brackets). Larvae are 4
- 983 dpf. Scale bar: 2µm.

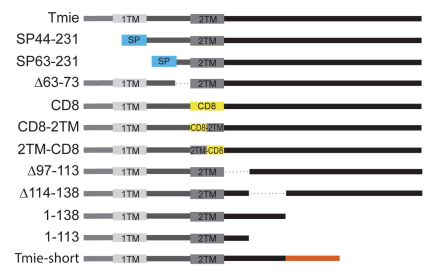


Tg(myo6b:Tmie-GFP)



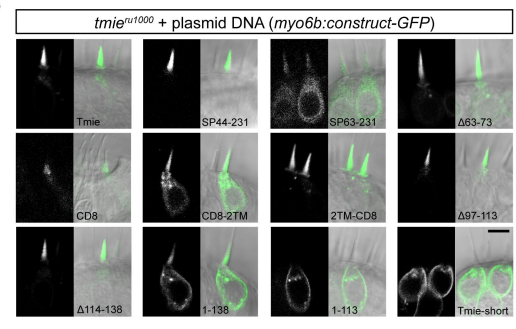




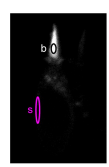


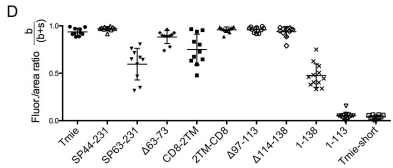
В

А

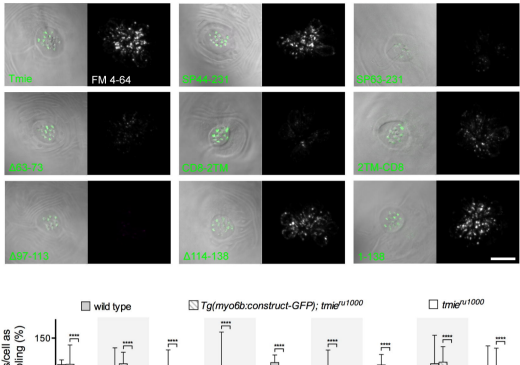


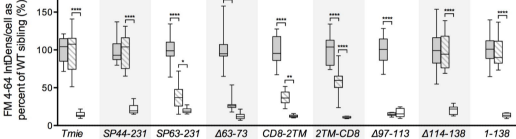






# Tg(myo6b:construct-GFP); tmie<sup>ru1000</sup>

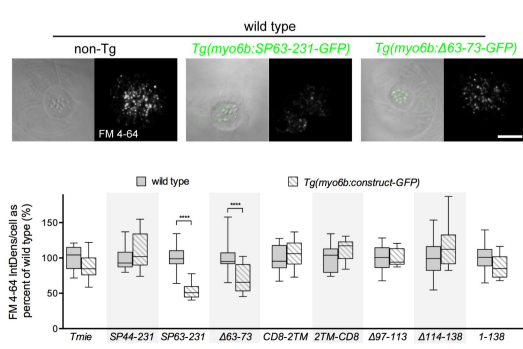




С

D

В



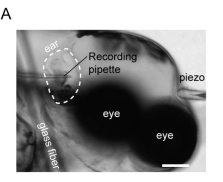
A

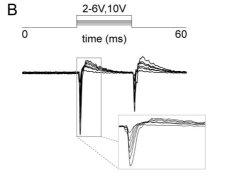
wild type

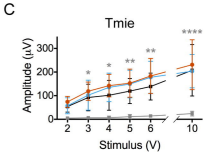
tmie ru1000 -86

Tg(myo6b:construct-GFP)

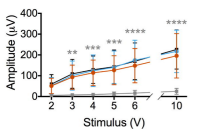
Tg(myo6b:construct-GFP); tmie<sup>ru1000</sup>

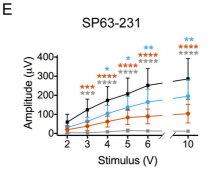








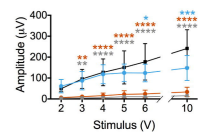






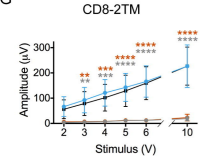
F

I

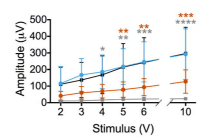


G

D

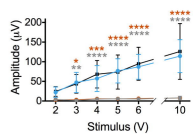


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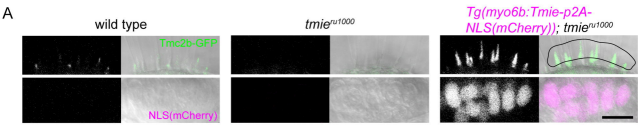


2TM-CD8

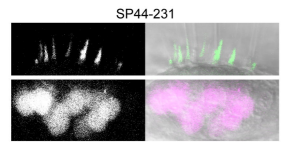




## Tg(myo6b:tmc2b-GFP)



# Tg(myo6b:tmc2b-GFP) Tg(myo6b:construct-p2A-NLS(mCherry)); tmie<sup>ru1000</sup>



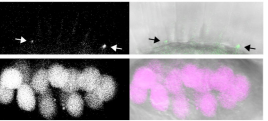
В

D

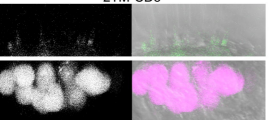
F

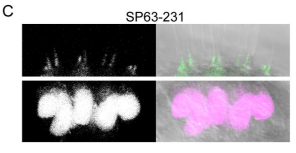
Н

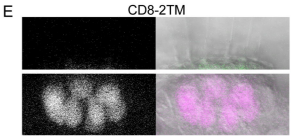
<u>∆63-73</u>



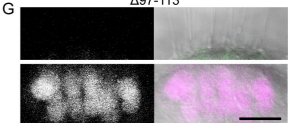
2TM-CD8

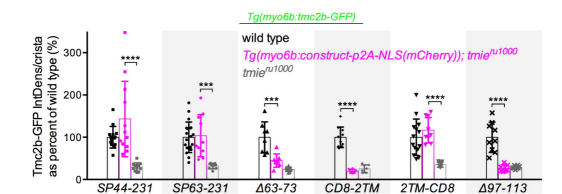


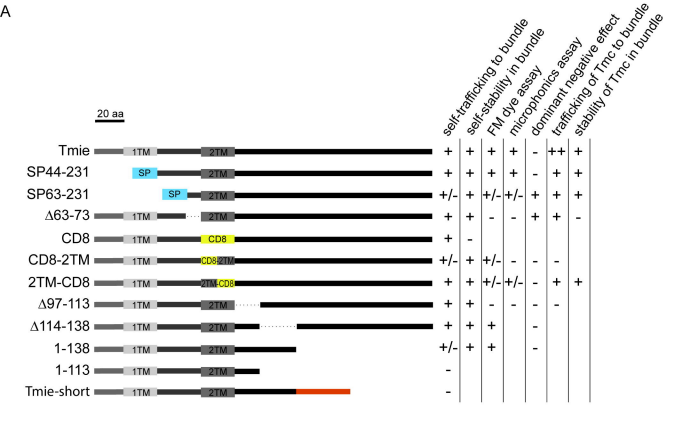












# В

