

# EPHA4 signaling dysregulation links abnormal locomotion and the development of idiopathic scoliosis

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### 73 **Conflict-of-interest statement**

74 J.R.L has stock ownership in 23andMe, is a paid consultant for Regeneron  
75 Pharmaceuticals and Novartis, and is a co-inventor on multiple United States and  
76 European patents related to molecular diagnostics for inherited neuropathies, eye  
77 diseases and bacterial genomic fingerprinting. The Department of Molecular and  
78 Human Genetics at Baylor College of Medicine derives revenue from the  
79 chromosomal microarray analysis (CMA by aCGH and/or SNP arrays), clinical  
80 exome sequencing (cES) and whole-genome sequencing (WGS) offered in the  
81 Baylor Genetics (BG) Laboratory (<http://bmgl.com>).

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## 93 Abstract

94 Idiopathic scoliosis (IS) is the most common form of spinal deformity with unclear  
 95 pathogenesis. In this study, we firstly reanalyzed the loci associated with IS, drawing  
 96 upon previous studies. Subsequently, we mapped these loci to candidate genes using  
 97 either location-based or function-based strategies. To further substantiate our  
 98 findings, we verified the enrichment of variants within these candidate genes across  
 99 several large IS cohorts encompassing Chinese, East Asian, and European  
 100 populations. Consequently, we identified variants in the *EPHA4* gene as compelling  
 101 candidates for IS. To confirm their pathogenicity, we generated zebrafish mutants of  
 102 *epha4a*. Remarkably, the zebrafish *epha4a* mutants exhibited pronounced scoliosis  
 103 during later stages of development, effectively recapitulating the IS phenotype. We  
 104 observed that the *epha4a* mutants displayed defects in left-right coordination during  
 105 locomotion, which arose from disorganized neural activation in these mutants. Our  
 106 subsequent experiments indicated that the disruption of the central pattern generator  
 107 (CPG) network, characterized by abnormal axon guidance of spinal cord  
 108 interneurons, contributed to the disorganization observed in the mutants. Moreover,  
 109 when knocked down *efnb3b*, the ligand for Epha4a, we observed similar CPG  
 110 defects and disrupted left-right locomotion. These findings strongly suggested that  
 111 ephrin B3-Epha4 signaling is vital for the proper functioning of CPGs, and defects in  
 112 this pathway could lead to scoliosis in zebrafish. Furthermore, we identified two  
 113 cases of IS in *NGEF*, a downstream molecule in the EPHA4 pathway. Collectively,  
 114 our data provide compelling evidence that neural patterning impairments and  
 115 disruptions in CPGs may underlie the pathogenesis of IS.

116 **Keywords:** Idiopathic scoliosis (IS), Pathogenesis, *EPHA4* (*Ephrin type-A receptor*  
 117 *4*), Central pattern generators (CPGs).

## 118    **Introduction**

119        Idiopathic scoliosis (IS) is the most common form of spinal deformity, affecting  
 120        about 2.5% of the global population (Hresko, 2013; Luk et al., 2010). IS may have  
 121        long-term physical and mental health consequences, such as cosmetic deformity,  
 122        cardiopulmonary impairment, and even disability (Weinstein et al., 2003). All these  
 123        consequences can severely reduce the quality of life. Early intervention with  
 124        conservative treatments, such as braces, can control scoliosis progression and reduce  
 125        the need for surgical intervention (Weinstein et al., 2013). However, IS often remains  
 126        undiagnosed until malformation is evident, emphasizing the importance of  
 127        risk-prediction measurements in medical management.

128        Genetic factors are thought to play a significant role in the development of IS,  
 129        while only a few genes have been associated with the condition to date (Cheng et al.,  
 130        2015; Miller, 2007). Several single-nucleotide polymorphisms (SNPs) associated  
 131        with susceptibility to IS have been identified through genome-wide association  
 132        studies (GWASs), including SNPs linked to genes such as *LBX1*, *GPR126*, and  
 133        *BNC2*. Notably, knockdown or overexpression of zebrafish homologs of these  
 134        IS-associated genes have yielded body axis defects (Guo et al., 2016; Kou et al.,  
 135        2019; Kou et al., 2013; Ogura et al., 2015; Zhu et al., 2015). In addition to common  
 136        SNPs, rare variants with larger effect sizes or causing rare Mendelian disease traits  
 137        also contribute to IS. Rare variants in *FBN1*, *FBN2*, and other extracellular matrix  
 138        genes are associated with severe IS (Buchan et al., 2014; Haller et al., 2016).  
 139        Linkage analysis of familial IS suggests that it follows autosomal dominant (AD),  
 140        X-linked dominant, and multifactorial patterns of inheritance (Miller et al., 2005).  
 141        Variants in *CHD7* and *AKAP2* are also implicated in the pathogenesis of Mendelian  
 142        forms of IS (Gao et al., 2007; Liebeskind et al., 2016). Finally, centriolar protein  
 143        POC5 and planar cell polarity protein Vang-like protein 1 (VANGL1) were also  
 144        shown to be associated with IS. Notably, all these genes participate in a wide range  
 145        of biological processes, the mechanisms underlying IS progression still require

146 further investigation. Consequently, there is currently no consensus on the etiology  
147 of IS (Tang et al., 2021).

148 In this study, we developed a novel pipeline that combines SNP-to-gene  
149 mapping and rare variant association analysis. Using this approach, we analyzed a  
150 large Chinese population with IS and further validated our findings in East Asian  
151 and European populations. Through these analyses, we identified *EPHA4* as a novel  
152 gene associated with IS. To confirm the pathogenicity of this gene, we used a  
153 zebrafish model and found that impaired EPHA4 pathway components and resulting  
154 defects in central pattern generators (CPGs) were associated with asymmetric spinal  
155 locomotion, which may serve as a contributory factor in the development of IS.  
156 Additionally, we searched for candidate genes related to EPHA4 signaling pathways  
157 and identified mutations in the *NGEF* among IS patients. Overall, our data suggest  
158 that the impairment of the EPHA4 pathway and central pattern generators plays a  
159 previously unknown role in the development and progression of IS.

## 160 **Results**

### 161 **Enrichment analysis of rare variants in IS candidate genes**

162 To identify candidate genes associated with IS, we conducted a comprehensive  
 163 literature review of SNPs that are linked to IS from 14 published GWASs. This led  
 164 us to obtain forty-one SNPs that showed genome-wide significance ( $P < 5 \times 10^{-8}$ ) as  
 165 detailed in Table S1. We then employed positional mapping, expression quantitative  
 166 trait locus (eQTL) mapping, or chromatin interaction mapping to identify 156  
 167 candidate genes that could be potentially linked to these SNPs. Further analysis of  
 168 rare variants for the 156 candidate genes using exome data from 411 Chinese  
 169 probands and 3,800 unrelated Chinese controls identified *EPHA4* as the only  
 170 significant gene ( $P=0.045$ ,  $OR=4.09$ ) (Table S2). *EPHA4* is a member of the Eph  
 171 family of receptor tyrosine kinases which play a vital role in the development of the  
 172 nervous system (Kania and Klein, 2016). We identified three rare variants in *EPHA4*  
 173 from three IS patients, including one splicing-donor variant and two missense  
 174 variants (Table S2). The flowchart of the entire pipeline is shown in Fig. S1.

### 175 **Inheritance pattern and functional analyses of variants in *EPHA4***

176 We characterized the inheritance pattern of variants in *EPHA4* using either trio  
 177 sequencing data or Sanger sequencing for the parents. None of the parents being  
 178 tested had scoliosis based on a clinical screening test. Of the three rare variants in  
 179 *EPHA4*, two (NM\_004438.3: c.1443+1G>C and c.2546G>A [p.Cys849Tyr]) arose  
 180 *de novo* (Fig. 1A and Table 1). The heterozygous splicing variant (c.1443+1G>C)  
 181 was identified in a female IS patient (SCO2003P0846) (Fig. 1B). Scoliosis in this  
 182 patient was diagnosed in her 10s, with a main curve Cobb angle of 60°. The *in vitro*  
 183 minigene assay showed that the c.1443+1G>C variant introduced a new splicing site,  
 184 resulting in a 36-bp in-frame deletion in exon 6 (Fig. 1G, 1I, S2A). The *EPHA4*  
 185 heterozygous missense variant (c.2546G>A, p.Cys849Tyr), identified in a female IS  
 186 patient (SCO2003P2146) (Fig. 1C), is located in the protein tyrosine kinase domain

187 of *EPHA4* protein (Fig. 1A). The onset of scoliosis in this patient was in her 10s  
188 with a 70° main curve Cobb angle. Given the critical role of *EPHA4* in  
189 phosphorylating CDK5, which in turn activates downstream signaling pathways (Fu  
190 et al., 2007), a western blot analysis of *EPHA4*-c.2546G>A variant was conducted to  
191 investigate the protein expression levels of *EPHA4* and CDK5 and the amount of  
192 phosphorylated CDK5 (pCDK5) in HEK293T cells transfected with *EPHA4*-mutant  
193 or *EPHA4*-WT plasmid. Our results revealed that the missense variant resulted in a  
194 decreased phosphorylation level of CDK5 (p=0.015), suggesting a partial  
195 loss-of-function (LoF) of *EPHA4* (Fig. 1F).

196 We next investigated *de novo* non-coding variants of *EPHA4* via whole genome  
197 sequencing (WGS) in 116 trio families with IS. A *de novo* heterozygous *EPHA4*  
198 intronic variant (c.1318+10344A>G) was identified in a female patient  
199 (SCO2003P2080) in an age range of 15-20 (Fig. 1D and Table 1). She developed  
200 scoliosis in her 10s with a 70° main curve Cobb angle. This variant is predicted to  
201 affect the branch point of the fifth intron of *EPHA4* (Fig. S2B). We performed nested  
202 PCR to show that this variant induced exon 5 skipping, resulting in a 339-bp  
203 in-frame deletion (Fig. 1H, J).

204 Next, we employed an in-house gene matching approach under the framework  
205 of DISCO study, which identified a patient (SCO2003P3202) in an age range of 5-10,  
206 who had been previously diagnosed with Waardenburg syndrome caused by a 4.46  
207 Mb *de novo* deletion at 2q35-q36.2 (Li et al., 2015) (Table 1). Intriguingly, the  
208 patient also presented mild scoliosis (Fig. 1E). The deletion included the entire *PAX3*  
209 gene, which was responsible for the Waardenburg syndrome phenotype, and 36  
210 neighboring genes, including *EPHA4* (Fig. 1K). As scoliosis is not typically  
211 associated with Waardenburg syndrome caused solely by *PAX3* pathogenic variants  
212 (Tassabehji et al., 1993), we hypothesize that the deletion of *EPHA4* may be  
213 responsible for the IS phenotype in this patient. Notably, the patient IDs used in this



study are unique to our research group and will not be recognizable to anyone outside the research team, including hospital staff and the patients themselves.

Notably, the GWAS signal which we mapped to *EPHA4* (rs13398147) (Zhu et al., 2015) represents a significant eQTL in esophagus and colon tissues, with the T allele associated with decreased expression of *EPHA4*. In our East Asian GWAS cohort of 6,449 adolescent IS patients and 158,068 controls, we identified another two eQTLs in *EPHA4* associated with decreased expression of *EPHA4* in brain tissue (Table S3). In the same GWAS cohort, common SNPs in *EPHA4*, after aggregation, also showed significant enrichment ( $P=0.023$ ) in IS patients versus controls. Taken together, the convergence between rare and common variants of *EPHA4* that lead to LoF or hypomorphic effects highlights the pivotal role of *EPHA4* in the pathogenesis of IS.

# **IS-like phenotypes in zebrafish *epha4* mutants**

To investigate the role of *EPHA4* in scoliosis, we utilized zebrafish as a model system due to its versatile nature in modeling adolescent IS (Bagnat and Gray, 2020; Boswell and Ciruna, 2017; Grimes et al., 2016; Xie et al., 2022). Zebrafish have two homologs of *EPHA4*, *epha4a* and *epha4b*. Using CRISPR-Cas9, we established a stable *epha4a* zebrafish mutant line with a 63-bp deletion in exon 3, which introduced a stop codon and resulted in a truncated protein (Fig. 2A-B). The homozygous *epha4a* mutant larvae had no apparent defects in either notochord or body axis development (Fig. S3A). Consistent with previous reports, the hindbrain rhombomeric boundaries were found to be defective in both the *epha4a* homozygous mutants and morphants (Fig. S3B) (Cayuso et al., 2019; Cooke et al., 2005). Interestingly, more than 75% adult mutants showed mild scoliosis (88 of 116), and some mutants exhibited severe scoliotic phenotype (4 of 116) (Fig. 2C-D, Movie S1&S2). Remarkably, some heterozygous adult mutants also developed mild scoliosis (14 of 95), whereas none of the wild-type fish showed any signs of scoliosis (0 of 76) (Fig. 2C-D). Similarly, we further generated the *epha4b* mutants

242 with a 25-bp deletion in exon 3, which resulted in a frameshift mutation (Fig. S4A).  
 243 The *epha4b* zebrafish mutants also developed mild scoliosis (28 of 43) (Fig. S4B-C).  
 244 Intriguingly, both *epha4a* and *epha4b* mutants exhibited early onset scoliosis starting  
 245 from around 20 days post fertilization (Fig. S4D), a stage similar to that of IS  
 246 patients. Collectively, these data suggest that mutations in Epha4 proteins are linked  
 247 to the scoliotic phenotype in zebrafish.

#### 248 **Abnormal left-right swimming pattern in the absence of Epha4a**

249 While the *epha4a* mutants seemed to be grossly normal during the larvae stages,  
 250 they developed spinal curvature gradually during later development. We decided to  
 251 investigate whether these mutants exhibited any abnormalities in their behavior at  
 252 the larvae stages using EthoVision XT software. By monitoring the swimming  
 253 behavior of 8 days post-fertilization (dpf) zebrafish larvae, we found the motion  
 254 distance and swimming velocity were significantly decreased in *epha4a* mutants (Fig.  
 255 3A-B). In addition, we observed a remarkable difference in the relative turning angle  
 256 and angular velocity between these two groups. The wild-type group changed their  
 257 swimming direction randomly, showing a relative angle around 0 degrees (Fig.  
 258 3C-D). In contrast, the *epha4a* mutants favored turning to one side of their directions  
 259 (a positive angle reflects a leftward turn) (Fig. 3C-D). Moreover, the absolute turn  
 260 angles and turning speed (angular velocity) were significantly higher in these  
 261 mutants (Fig. 3E-F). To rule out potential off-target effects, we injected *epha4a*  
 262 mRNA into *epha4a* mutants, which significantly restored the swimming  
 263 coordination defects (Fig. S5).

264 Next, we compared the swimming behavior after startle response between  
 265 wild-type and *epha4a* mutants. We used a needle to touch the head or the tail of 5  
 266 days post-fertilization (dpf) zebrafish larvae to stimulate swimming behavior. In  
 267 sibling controls, the larvae responded to the tactile stimulation and swam away  
 268 quickly (Movie S3). Conversely, the *epha4a* mutants failed to respond to the initial  
 269 stimulation, and the swimming pattern was defective with an abnormal bending

270 pattern (37 of 62 from tail-stimulation group and 34 of 77 from head-stimulation  
271 group) (Fig. 3G, Movie S4). We further performed more comprehensive analysis by  
272 high-speed video microscopy. After tactile stimulation, wild-type larvae displayed a  
273 high-speed C-bend turn followed by a weaker counterbend turn after several  
274 milliseconds (Fig. 3H-I). This rhythmic left/right swimming pattern ensures that the  
275 fish swim away from the frightening stimulus. The turning angles of the control  
276 larvae after stimulation changed with a sinusoidal wave pattern (Fig. 3J). In contrast,  
277 this pattern was dramatically different in *epha4a* mutants (Fig. 3I,K). Of note,  
278 although the C-bend turning angles were similar between *epha4a* mutants and  
279 control siblings, the turning angles of the counterbend decreased significantly (Fig.  
280 3L), implying a left/right coordination defect. Taken together, both regular  
281 swimming and tactile stimulation analyses suggested that the left-right coordination  
282 swimming pattern is compromised in the absence of Epha4a.

## 283 **Defects of left-right coordination due to abnormal CPG in the absence of** 284 **Epha4a**

285 The coordinated left-right locomotion of zebrafish larvae relies on the  
286 synchronized contraction of muscle fibers, a process regulated by motor neurons  
287 situated on each side of the fish. To explore this intricate mechanism, we utilized a  
288 Tg(*elavl3*:GAL4; *UAS*: GCaMP) double transgene, allowing the expression of a  
289 genetically encoded calcium sensor in all neurons. We observed the rhythmic  
290 activation of calcium signaling in motor neurons located within the spinal cord (Fig.  
291 4A, Fig. S6A, Movie S5, Movie S6). In wild-type larvae, the calcium signals  
292 exhibited an alternating pattern between the left and right sides of the body, whereas  
293 this coordinated pattern was disrupted in both *epha4a* mutants and morphants (Fig.  
294 4A-B vs A'-B', Fig. S6A-B vs A'-B', Movie S7, Movie S8). Additionally, we found  
295 that the activation frequency of motor neurons on the left and right sides was  
296 comparable in wild-type larvae, but significantly different in the absence of Epha4a  
297 (Fig. 4C-D, Fig. S6C-D).

One well-established concept regarding left-right coordination involves the presence of central pattern generators (CPGs), which are regulated by interneuron circuitry within the spinal cord. We further examined the axon guidance of interneurons in *epha4a* mutants. First, we investigated the commissural trajectories of reticulospinal (RS) interneurons. In control larvae, the large Mauthner neurons, along with other RS neurons, were symmetrically positioned on each side of the midline and projected their axons to the contralateral sides (Fig. 5A). These bilaterally projected axons typically crossed at the midline and subsequently synapsed on motoneurons of the opposite sides, contributing to the generation of spinal cord neural circuits (Hale et al., 2016). However, in *epha4a* mutant larvae, we observed an abnormal pattern in axonal projections. Specifically, the mutant axons failed to traverse the midline and instead extended ipsilaterally (Fig. 5A). In addition, the distance between Mauthner neurons (rhombomere 4) and rhombomere 7 was significantly decreased and rhombomere 5 was scarcely visible in the mutant larvae (Fig. 5A-B). Furthermore, the sites of axon crossing between two Mauthner neurons tended to deviate to one side of the midline in the mutants (Fig. 5C-D').

Cerebrospinal fluid-contacting neurons (CSF-cNs) represent a unique type of interneuron responsible for modulating the V0 and V2a interneurons, which are integral components of locomotor CPGs (Fidelin et al., 2015; Talpalar et al., 2013; Wu et al., 2021). In wild-type larvae, we observed that the ascending axons of these neurons projected either to the right or left side from the midline, as visualized using *Tg(pkcd211:GAL4;UAS:Kaede)* double transgenic larvae (Fig. 6A). However, in the absence of *Epha4a*, the projection of these neurons exhibited notable disorganization, with numerous axons crossing the midline from one side of the trunk. This disorganized pattern was observed in both heterozygotic and homozygotic mutants (Fig. 6A-B). Additionally, we employed an optogenetic approach to activate these CSF-cNs, utilizing the *Tg(Gal4<sup>s1020t</sup>; UAS:ChR2)* double transgene. Following optical stimulation, we observed robust tail oscillations as previously described

(Wyart et al., 2009). In wild-type larvae, these tail oscillations exhibited a symmetrical left-right beating pattern (Fig. 6C-D, Movie S9). However, a striking disruption of this symmetry was observed in *epha4a* morphants, as they consistently beat towards one side of the trunk following optical stimulation (a positive angle reflects a rightward turn) (Fig. 6C'-D' and E, Movie S10). Collectively, these findings demonstrate that the left-right coordination deficiencies observed in *epha4a* mutants arise from abnormal neural circuit formation, which consequently disrupts the integrity of the CPGs.

### **Ephrin B3-Epha4 signaling regulates interneuron axon extension**

To further explore the role of Epha4a during interneuron axon extension, we examined the expression of *epha4a* during early zebrafish embryonic development. Whole-mount in situ hybridization results showed that both *epha4a* and *epha4b* were abundantly expressed in the zebrafish spinal cord (Fig. S7A). We plotted the expression of these two genes using two published single-cell transcriptome data, which showed that *epha4a* and *epha4b* were both expressed in interneurons, suggesting a role for *epha4* in interneuron function (Fig. S7B) (Cavone et al., 2021; Scott et al., 2021). Notably, the expression of *efnb3b*, encoding the ligand of Epha4, was highly enriched in the midline floor plate cells (Fig. S7B).

Ephrins, through interacting with Eph receptors, play a critical role in repulsive axon guidance during neural development (Egea and Klein, 2007; Flanagan and Vanderhaeghen, 1998). We further analyzed axon guidance in *efnb3b* morphants. Similar to those of *epha4a* mutants, the *efnb3b* morphants also displayed axon guidance defects, as well as uncoordinated calcium activation (Fig. S8A-C). In addition, morphants larvae also displayed left-right oscillations defects after optogenetic stimulation (a positive angle reflects a rightward turn) (Fig. S8D-E).

### **Candidate variants in the *EPHA4*-related genes**

Our zebrafish studies suggested that EPHA4 signaling is crucial for interneuron

353 axon guidance, hence the formation of functional CPGs. Next, we further asked  
 354 whether mutation of other components of the EPHA4 signaling can result in IS in  
 355 humans. By searching for rare variants in *EPHA4*-related genes (Fig. S9)  
 356 (Szkarczyk et al., 2019), we identified heterozygous *de novo* start-loss variant  
 357 (c.1A>G, p.Met1?) in *NGEF* in a young male (SCO2003P3332) (Fig. 7A-B, Table 1),  
 358 whose scoliosis was diagnosed in an age range of 15-20, with a main curve Cobb  
 359 angle of 60°. *NGEF* encodes the neuronal guanine nucleotide exchange factor  
 360 Ephexin that differentially affects the activity of GTPases RHOA, RAC1, and  
 361 CDC42. The activation of Ephexin is triggered by ephrin through EPHA4 (Shamah  
 362 et al., 2001).

363 Strikingly, we further identified a dominant missense variant (c.857G>A,  
 364 p.Ala286Val) in *NGEF* in a quad family with three affected members (Fig. 7C, Table  
 365 1). The proband (II:2, TSRHC01) has a 58° main curve Cobb angle. This variant,  
 366 which is located in the RhoGEF domain of Ephexin, is predicted to be highly  
 367 deleterious (CADD=29.6).

368 Altogether, our results suggest that defects of the CPGs owing to abnormal  
 369 EPHA4 signaling maybe one of the crucial factors responsible for IS.

## 370 Discussion

371 IS is a disease with diverse causes, and the underlying mechanisms can vary  
 372 even among patients with similar scoliotic phenotypes. Previous genetic studies have  
 373 highlighted the significance of the extracellular matrix (ECM) in maintaining the  
 374 balance of axial bone and supporting soft tissues in the spine, thus playing a crucial  
 375 role in IS development (Haller et al., 2016). For example, the top SNP associated  
 376 with IS maps to *LBX1*, an essential molecule for ECM maintenance and bone  
 377 homeostasis (Takahashi et al., 2011). Additionally, genetic loci in muscle  
 378 development-related genes have also been associated with the onset of scoliosis,  
 379 emphasizing the intricate interaction between bones and muscles (Ogura et al., 2015).  
 380 However, it is worth noting that the variants in these genes explain only a small  
 381 fraction of the overall heritability of IS. Consequently, it is imperative to establish  
 382 connections between the extensive genetic findings and biological mechanisms that  
 383 can elucidate the etiological landscape of IS.

384 In this study, we mapped 41 significant genome-wide loci to functional genes  
 385 through positional mapping and functional mapping such as eQTL. Then we  
 386 determined the enrichment in patient cohorts of rare variants in these genes, which  
 387 may have greater impacts compared with common SNPs. This approach revealed the  
 388 convergence of SNPs and rare variants in *EPHA4* that are enriched in patients with  
 389 IS. We also identified additional high-impact variants in *NGEF*, which is involved in  
 390 the EPHA4 pathway.

391 Our studies using zebrafish have revealed that deficiency of Epha4 can lead to  
 392 the development of scoliosis. Interestingly, we observed that even heterozygotic  
 393 *epha4a* zebrafish mutants displayed mild scoliosis (Fig. 2C), which is consistent  
 394 with the occurrence of this condition in scoliosis patients. Further analysis involving  
 395 behavior and imaging demonstrated that the absence of Epha4a resulted in defective  
 396 left-right coordination. This coordination is crucially governed by CPGs, which

397 generate rhythmic patterns of neural activity to coordinate limb movements on both  
 398 sides of the body (Kiehn, 2006; Marder and Bucher, 2001; Talpalar et al., 2013).  
 399 Previous studies have reported the involvement of ephrin and its receptors in axon  
 400 guidance during the maturation of neural circuits, including CPGs (Andersson et al.,  
 401 2012; Borgius et al., 2014; Iwasato et al., 2007). Ephrin B3, present in radial glial  
 402 cells at the dorsal midline, serves as a repulsive barrier to axons expressing *Epha4*  
 403 (Imondi et al., 2000). Axon guidance is disrupted when this repulsive pathway is  
 404 inactivated, leading to midline invasion (Butt et al., 2005; Kullander et al., 2001;  
 405 Paixão et al., 2013). In *Epha4* and *Efnb3* knockout mice, a 'hopping' gait is observed,  
 406 which is attributed to the aberrant crossing of motor neurons expressing *Epha4*  
 407 (Kullander et al., 2003). Consistent with these observations, we observed disrupted  
 408 axon guidance in the interneurons, which are integral components of CPGs, in both  
 409 *epha4a* mutants and *efnb3b* morphants. This finding suggests that CPG malfunction  
 410 may be a significant contributing factor to the scoliosis phenotype observed in these  
 411 genetic backgrounds. It is highly likely that the lack of coordinated left-right  
 412 locomotion generates imbalanced mechanical forces on the spine, gradually leading  
 413 to spinal curvature during later stages of development (Fig. 8).

414 Thus, our data provided a novel biological mechanism of IS, i.e., the  
 415 impairment of neural patterning and CPG. Previous studies have provided clues on  
 416 the role of CPGs in IS. Patients with adolescent IS showed asymmetric trunk  
 417 movement during gait, as characterized by increased relative forward rotation of the  
 418 right upper body in relation to the pelvis (Kramers-de Quervain et al., 2004; Nishida  
 419 et al., 2017). An electromyography (EMG) study also showed asymmetric activation  
 420 of paraspinal muscles between the convex and concave sides at the scoliosis curve  
 421 apex (Shimode et al., 2003). In a child with a strong family history of IS, asymmetric  
 422 hyperactivity was observed by EMG months before scoliosis was evident (Valentino  
 423 et al., 1985). These left-right locomotor coordination abnormalities indicated the  
 424 maldevelopment of CPGs as potential cause of IS. The CPG asymmetry may induce



425 an imbalance in trunk muscle strength, resulting in asymmetric rib drooping. This  
 426 leads to an abnormal vertebral rotation and then the onset of IS. Moreover, in a  
 427 companion study, Wang et al. identified a number of rare variants in *SLC6A9*, which  
 428 encodes glycine transporter 1 (GLYT1), in familial and sporadic adolescent IS cases.  
 429 The *slc6a9* mutant zebrafish also exhibited discoordination of spinal neural activities  
 430 with pronounced lateral spinal curvature, recapitulating the human IS phenotype  
 431 (Wang et al., 2024). Taken together, we propose that the dysfunction of CPGs would  
 432 cause an imbalance in the motor drive from the spinal cord and the asymmetric  
 433 transversospinalis muscle pull, eventually producing a scoliotic curve (Fig. 8).

434 In summary, our study demonstrates that both common and rare variants within  
 435 the EPHA4 pathway contribute to the genetic architecture of IS. EPHA4 pathway  
 436 dysfunction causes axon pathfinding defects, resulting in impaired coordinated  
 437 left-right locomotion by disrupting neural patterning and the function of CPGs,  
 438 thereby potentially leading to IS.

## 439 **Materials and Methods**

### 440 **Mapping of candidate genes utilizing previous association studies**

441 Through a systematic literature review, we identified IS-associated SNPs  
 442 reported in GWASs and meta-analyses of GWAS. The literature search was carried  
 443 out using MEDLINE (via Pubmed.gov) and Web of Science (via Clarivate Analytics)  
 444 and was limited to English-language articles published from January 1980 to  
 445 October 2020. The following keywords were combined to perform the search:  
 446 ‘idiopathic scoliosis’ AND ‘GWAS’ OR ‘SNP’ OR ‘single nucleotide polymorphism’  
 447 OR ‘variant’ (Table S4). The inclusion/exclusion criteria of abstracts are provided in  
 448 Table S5. After screening the titles and abstracts, we obtained from the full-text  
 449 articles the rsID, chromosome, and position for SNPs with a threshold for  
 450 genome-wide significance of  $P < 5.0 \times 10^{-8}$ . For SNPs identified in multiple studies,  
 451 we recorded the lowest P-value.

452 SNPs reported in previous studies were first pruned using Functional Mapping  
 453 and Annotation (FUMA, v1.4.1, <https://fuma.ctglab.nl/>) (Watanabe et al., 2017).  
 454 Significant SNPs were considered independent at  $r^2 < 0.6$ . All known SNPs (available  
 455 in the 1000 Genomes reference panel, <https://www.internationalgenome.org/>) were  
 456 included for further gene mapping if they were in a linkage disequilibrium (LD) block  
 457 ( $r^2 \geq 0.6$ ) with a significant independent SNP. Three SNP-to-gene strategies were  
 458 used:

- 459 1) For positional mapping of significant independent SNPs, we used annotations  
 460 obtained from ANNOVAR (<http://annovar.openbioinformatics.org>). A 10-kb  
 461 maximum distance was applied for intergenic SNPs.
- 462 2) For the eQTL mapping, we mapped significant independent SNPs and SNPs in an  
 463 LD block to eQTLs across 44 GTEx tissue types (release V8) (Consortium, 2020).  
 464 SNP-gene pairs with a false discovery rate (FDR)  $\leq 0.05$  were considered  
 465 significant.

3) For chromatin interaction mapping, we overlapped the significant independent SNPs and those in LD blocks with one end of significantly interacting regions across various tissue and cell types. Information on the interacting regions was derived from Hi-C datasets of 21 tissues and cell types provided by GSE87112. The significance of interactions for Hi-C datasets was computed by Fit-Hi-C, with an  $FDR \leq 10^{-6}$  considered significant. Genes were mapped if their promoter regions overlapped with another end of the significant interactions. The promoter region was defined as the region from -250 bp to +500 bp relative to the transcription start site.

### Cohort description

The Peking Union Medical College Hospital (PUMCH) cohort: The PUMCH cohort comprised 411 unrelated Chinese patients with severe IS (Cobb angle  $\geq 40^\circ$ ) who underwent spinal surgery in the PUMCH between October 2017 and March 2022 as part of the Deciphering disorders Involving Scoliosis and COmorbidities (DISCO) study (<http://www.discostudy.org/>). The clinical diagnosis was confirmed using standing full-spine radiographs, three-dimensional computed tomography, and magnetic resonance imaging. These patients did not show any congenital or neuromuscular defect at the time of recruitment. The control cohort consisted of 3,800 individuals without observable scoliosis and with exome sequencing or genome sequencing performed at PUMCH for clinical or research purposes. Individuals with vertebrae malformation or congenital developmental defects were excluded. A patient with a 2q35-36.2 deletion, including the *EPHA4* gene, was also recruited through in-house gene matching of the DISCO study (Li et al., 2015). This patient was subsequently evaluated for scoliosis by examination and radiography.

The East Asian cohort: The East Asian cohort includes totally 6,449 IS patients and 158,068 controls from four independent datasets (Japanese dataset 1 (Kou et al., 2013; Takahashi et al., 2011): 1,261 cases, 15,019 controls; Japanese dataset 2 (Ogura et al., 2015): 878 cases, 21,334 controls; Japanese dataset 3 (Kou et al.,

2019): 3,333 cases, 119,630 controls; Hong Kong dataset: 977 cases, 2,085 controls).  
The inclusion criteria for IS subjects was as same as our previous studies (Fan et al.,  
2012; Kou et al., 2019).

The Texas Scottish Rite Hospital for Children (TSRHC) cohort: We used a  
replication cohort of European-ancestry patients with IS from the TSRHC. Cases  
considered for inclusion in the study met criteria for a positive diagnosis of IS:  
lateral deviation from the midline greater than 15 degrees as measured by the Cobb  
angle method from standing spinal radiographs, axial rotation toward the side of the  
deviation and exclusion of relevant co-existing diagnoses.

### **Blood sample collection**

In the PUMCH cohort, genomic DNA samples were extracted from peripheral  
blood leukocytes of each subject using a QIAamp DNA Blood Mini Kit (Qiagen,  
Hilden, Germany), according to the manufacturer's protocol. Purified DNA was  
qualified by Nanodrop 2000 (Thermo Fisher Scientific, Waltham, MA, USA) and  
quantified by Qubit 3.0 using the dsDNA HS Assay Kit (Life Technologies, Carlsbad,  
CA, USA). DNA samples were stored at 4°C until used.

In the East Asian and TSRHC cohort, genomic DNA was extracted from  
peripheral blood or saliva using a standard protocol.

### **DNA sequencing and variant calling**

In the PUMCH cohort, whole exome sequencing (WES) or whole genome  
sequencing (WGS) was performed on peripheral blood DNA from all individuals and  
available family members (Table S6). A SureSelect Human All Exon V6+UTR r2 core  
design (91 Mb, Agilent) was used for exon capture. The exomes were then sequenced  
on an Illumina HiSeq 4000 (Illumina, San Diego, CA, USA) according to the  
manufacturer's instructions. For WGS, sequencing libraries were prepared using the  
KAPA Hyper Prep kit (KAPA Biosystems, Kusatsu, Japan) with an optimized  
manufacturer's protocol. We performed multiplex sequencing using an Illumina

HiSeq X-Ten sequencer (Illumina, San Diego, CA, USA). Variant calling and annotation were done in-house using the Peking Union Medical College Hospital Pipeline (PUMP) described previously (Chen et al., 2021; Zhao et al., 2021).

In the TSRHC cohort, subjects in 162 adolescent IS families were sequenced as part of the Gabriella Miller Kids First Pediatric Research Consortium (GMKF) at The HudsonAlpha Institute for Biotechnology (Huntsville, AL). In summary, DNA was normalized, sheared and then ligated to Illumina paired-end adaptors. The purified ligated DNA was amplified and exome sequencing was performed on the Illumina HiSeq X platform. The sample's sequences were aligned into GRCh38 and genotypes were joint-called per each family by the GMKF's Data Resource Center (DRC) at Children's Hospital of Philadelphia following GATK best practices as detailed here (Mukhopadhyay et al., 2020). GMKF DRC's alignment and joint genotyping pipelines are open source and made available to the public via GitHub (<https://github.com/kids-first/kf-alignment-workflow> and <https://github.com/kids-first/kf-jointgenotyping-workflow>).

### **Rare variant association analysis of the candidate genes**

To determine the contribution of rare variants in GWAS candidate genes to IS, we analyzed the gene-based mutational burden for 156 candidate genes. Only ultra-rare variants with a gnomAD population-max allele frequency  $\leq 0.01\%$  and a cohort allele count  $\leq 3$  were analyzed. Each variant was assigned a weight, and the mutational burden of a given gene was defined as the maximum weight value among all ultra-rare variants carried by the individual. The SAIGE-GENE+ package was used to determine the weighted mutational burden test for each gene (Zhou et al., 2022).

Weighting criteria for the weighted burden analysis were developed according to the variant types and in silico results. The LoF variants (canonical splicing variants, nonsense variants, or variants that cause frameshift, stop-gain, or start-loss)

were calculated together with the protein-altering variants, including missense variants and in-frame indels. Each variant was assigned a weight range from 0-1 and the mutational burden for a given gene was defined as the maximum weight value among all ultra-rare variants carried by the individual. The LoF variants annotated as ‘high confidence’ by the loss-of-function transcript effect estimator (LOFTEE) (Karczewski et al., 2020) were assigned a weight value of 1.0. The LoF variants annotated as ‘low confidence’ or unlabeled by LOFTEE, the non-canonical splicing variants with a SpliceAI score  $> 0.5$ , and the missense variants with a rare exome variant ensemble learner (REVEL) (Ioannidis et al., 2016) score of 0.8 were assigned a weight value of 0.8. The in-frame insertions/deletions (indels) with a Combined Annotation Dependent Depletion (CADD) (Kircher et al., 2014) score  $> 20$  and the missense variants with a REVEL score  $> 0.6$  and  $\leq 0.8$  were assigned a weight value of 0.6. The in-frame indels with a CADD score  $> 10$  and  $\leq 20$  and the missense variants with a REVEL score  $> 0.4$  and  $\leq 0.6$  were assigned a weight value of 0.4. The in-frame indels and the missense variants with a REVEL score  $> 0.2$  and  $\leq 0.4$  were assigned a weight value of 0.2. The remaining missense variants were assigned a weight value of 0.

### **Sanger sequencing of familial participants**

Sanger sequencing of familial participants was performed to determine the origin of variants in *EPHA4* and *NGEF*. All LoF variants and protein-altering variants identified in familial participants were validated. Variant-encoding gene regions were amplified by PCR from genomic DNA obtained from probands, as well as from parents for trios, to determine the origin of the variants. The amplicons were purified using an Axygen AP-GX-50 kit (Corning, NY, USA) and sequenced by Sanger sequencing on an ABI 3730xl instrument (Thermo Fisher Scientific, Waltham, MA, USA).

## 574 **Minigene assay**

575       The splicing variant (*EPHA4*: c.1443+1G>C) was characterized by a minigene  
576 assay. Genomic DNA from the heterozygous patient was amplified by PCR using a  
577 high-fidelity DNA polymerase. Amplicons included exon 6, intron 6, exon 7, intron  
578 7, and exon 8 of the *EPHA4* gene. PCR products were cloned into the vector via the  
579 restriction sites BamHI and MluI for pCAS2, which is based on the pcDNA3.1  
580 plasmid (Thermo Fisher Scientific, Waltham, MA, USA). Clones with wild-type or  
581 mutant genomic inserts were selected and verified by sequencing the cloned DNA  
582 fragments. The recombinant plasmids were transfected into 293T cells using  
583 Lipofectamine™ 3000 reagent (Thermo Fisher Scientific, Waltham, MA, USA).  
584 For RT-qPCR, total RNA was isolated from the transfected cells using Trizol reagent  
585 (Thermo Fisher Scientific, Waltham, MA, USA) and reverse transcription was  
586 performed using the GoScript™ Reverse Transcription System (Promega, Madison,  
587 MI, USA). PCR amplification was performed using the pCAS2-RT-F and  
588 pCAS2-RT-R primers, and the products were sequenced using pCAS2-RT-F.

589 Primers sequences for the minigene assay were as follows:

590 pCAS2-RT-F: 5'-CTGACCCTGCTGACCCTCCT-3'

591 pCAS2-RT-R: 5'-TTGCTGAGAAGGCGTGGTAGAG-3'

## 592 **Nested PCR**

593       The splicing variant (*EPHA4*: c.1318+10344A>G) was characterized by nested  
594 PCR. RNA was extracted from the whole blood of the patient using the TRIzol  
595 Reagent (CWBIO, Hangzhou, China) according to the manufacturer's guidelines.  
596 The cDNA was synthesized using HiScript II 1st Strand cDNA Synthesis Kit with a  
597 gDNA wiper (Vazyme, Nanjing, China) according to the manufacturer's guidelines.  
598 Nested PCR was performed as described previously (Yao and Tavis, 2005). Nested  
599 PCR reactions in 50 µl, included 2 µl cDNA from the reverse transcription reaction  
600 as the template for the first round of PCR or 2 µl of first-round PCR product as the

601 template for the second PCR, 1.5 µl 10 µM sense primer, 1.5 µl 10 µM anti-sense  
602 primer, 5 µl nucleotide mix (2 mM each dNTP), 5 µl 10× KOD Buffer, 1 µl 1 unit/µl  
603 Kod-Plus-Neo polymerase, 3 µl 25 mM MgSO<sub>4</sub> and 31 µl ddH<sub>2</sub>O. The PCR program  
604 was (94°C for 2 min, 98°C for 10 s, 55°C for 30 s, 68°C for 60 s) × 20 cycles and  
605 then (94°C for 2 min, 98°C for 10 s, 57°C for 30 s, 68°C for 60 s) × 30 cycles.

606 Primers sequences:

607 For the first round:

608 F: 5'-GGCTCCTGTGTCAACAACACTC-3'

609 R: 5'-GTTGGGATCTTCGTACGTAA-3'

610 For the second round:

611 F: 5'-AACTGCCTATGCAACGCTGG-3'

612 R: 5'-AGCTGCAATGAGAATTACC-3'

### 613 **Western blots**

614 HEK293T cells were transfected with *EPHA4*-C1-pEGFP plasmid or the  
615 corresponding wild type vector to detect the missense variant in *EPHA4* (c.2546G>A,  
616 p.Cys849Tyr). Cells were cultured in six-well plates and transfected with DNA (2  
617 mg/well) using Lipofectamine 3000 reagent (Thermo Fisher Scientific, Waltham,  
618 MA, USA). After 48 hours, cells were harvested, and protein extracts were prepared  
619 as described (Ding et al., 2017). The mutant proteins and wild-type proteins were  
620 fused to GFP. The expression of the two proteins was compared by western blots  
621 with an GFP antibody using the ECL detection system. The GFP antibody was  
622 purchased from Cell Signaling Technology (Cell Signaling, Danvers, MA, USA).  
623 The CDK5 antibody was purchased from Santa Cruz Biotechnology (Santa Cruz,  
624 CA, USA). A phospho-specific antibody against CDK5 phosphorylated at Tyr  
625 (GeneTex, Irvine, CA 92606 US) was purchased from GeneTex. Western blot  
626 experiments were repeated twice with similar results for the replicates.



## 627 Genotyping and imputation of GWAS

628 In the East Asian Japanese cohort, genotyping was performed by Illumina  
 629 Human610 Genotyping BeadChip, Illumina HumanOmniExpressExome and  
 630 HumanOmniExpress as our previous GWASs (Kou et al., 2019; Kou et al., 2013;  
 631 Ogura et al., 2015; Takahashi et al., 2011). For quality control (QC), subjects with call  
 632 rate  $< 0.98$ , high-degree relatedness with other subjects and outliers of East Asian  
 633 ethnicity were excluded. For variants QC, the exclusion criteria of variants were as  
 634 follows: call rate  $< 0.99$ , P-value for Hardy-Weinberg equilibrium  $< 1.0 \times 10^{-6}$   
 635 and minor allele count  $< 10$ . The reference panel for imputation, namely JEWEL7K,  
 636 was composed of 1000 Genomes Project phase 3 (v5) (Auton et al., 2015) and  
 637 Japanese whole-genome sequence data (Das et al., 2016) with 3256 high-depth  
 638 subjects ( $\geq 30$  read counts) and 4216 low-depth subjects ( $\leq 15$  read counts). Using  
 639 EAGLE 2.4.1 (<https://alkesgroup.broadinstitute.org/Eagle/>) to determine the  
 640 haplotypes, pre-phasing was conducted. Genotypes were imputed using Minimac4  
 641 (version1.0.0) (Das et al., 2016). After imputation, we excluded variants with minor  
 642 allele frequency (MAF)  $< 0.005$  and low imputation quality ( $R^2 < 0.3$ ).

643 In the East Asian Hong Kong cohort, samples were genotyped with Illumina  
 644 Human Omni ZhongHua-8 Beadchips. Illumina Genome Studio v2.0 was used to  
 645 convert raw data into PLINK format. The QC steps of samples and variants were  
 646 described in a previous study (Marees et al., 2018). Genotype phasing and imputing  
 647 were executed using SHAPEIT v2.r900 (Delaneau et al., 2012) and IMPUTE2  
 648 (Marchini et al., 2007). The imputed data was filtered using the following parameters:  
 649 INFO  $> 0.6$ , Certainty  $> 0.8$ , and MAF  $> 0.01$ . Association analysis was performed  
 650 using PLINK v1.9 logistic regression with covariates: sex, age and top 20 principal  
 651 components of the variance-standardized relationship matrix.

## 652 East Asian GWAS meta-analysis for IS

653 For the meta-analysis of the four datasets (three Japanese datasets and one  
654 Hong Kong dataset), an inverse-variance-based method was performed by METAL  
655 (version 2011-03-25) (Willer et al., 2010). SNPs in three or more of the four cohorts  
656 were used in subsequent analyses.

## 657 Gene-based common variant analysis and eQTL analysis

658 SNPs in *EPHA4* or within 20 kb flanking *EPHA4* with a significant association  
659 with IS were retrieved. SNPs were matched to potential eQTLs according to the  
660 GTEx database (v8, <https://gtexportal.org>). Gene-based common variant analyses  
661 were performed using Multi-marker Analysis of GenoMic Annotation (MAGMA)  
662 (de Leeuw et al., 2015) and FUMA (Watanabe et al., 2017) using default settings  
663 with LD information from the 1000 Genomes Project East Asian population (1KGP  
664 EAS) as a reference. SNPs located within two kb upstream and one kb downstream  
665 from *EPHA4* were included in the gene-based analysis.

## 666 Zebrafish strains, mutants, and morphants

667 Zebrafish Tuebingen (TU) strains were maintained at 28°C on a 14-hour/10-hour  
668 light/dark cycle. Embryos were raised at 28.5°C in E3 medium (5 mM NaCl, 0.17  
669 mM KCl, 0.39 mM CaCl<sub>2</sub>, 0.67 mM MgSO<sub>4</sub>) following standard protocols. Zebrafish  
670 have two homologs of *EPHA4*, *epha4a* and *epha4b*. The CRISPR/Cas9 system was  
671 used to generate zebrafish *epha4a* and *epha4b* mutants. To increase efficiency, we  
672 injected Cas9 mRNA together with multiple single guide RNAs (sgRNAs) for each  
673 gene (Table S7). The sgRNA sequences for *epha4a* and *epha4b* are listed in Table S7.  
674 Morpholino sequences for *epha4a* and *efnb3b* knockdown analysis are also listed in  
675 Table S7.

## 676 Micro CT imaging

677 Adult zebrafish *epha4a* mutants or wild-type siblings were euthanized with  
678 tricaine methanesulfonate and fixed in 4% paraformaldehyde. Micro CT images were

679 captured using a PerkinElmer Quantum GX2. Planar images acquired over 360° of  
680 rotation were reconstructed using QuantumGX. Three-dimensional renders of the  
681 skeleton were made with Analyze 12.0 software (AnalyzeDirect).

## 682 **Behavioral recordings and analysis**

683 For behavior analysis, individual zebrafish larvae were transferred into 24-well  
684 plate with fresh E3 medium at 8 dpf. Then, the plate was placed inside the  
685 Daniovision (Noldus) observation chamber for further behavior analysis.  
686 Video-tracking of swimming activity and further statistical analysis were performed  
687 using the EthoVision™ XT10 software. Behavioral data were shown as total swim  
688 distance (mm), average velocity (mm/s), average relative turn angle (degree), average  
689 absolute turn angle (degree) and average absolute angular velocity (degree/s) at a  
690 duration for 4 min. A positive angle reflects a leftward turn.

691 For tactile stimulation, zebrafish larvae were placed in a concave slide, and tactile  
692 stimulation was performed with glass capillaries to the head or tail sides. Larval  
693 startle responses were recorded using a high-speed video camera (Mikrotron, EoSens  
694 Mini1) at 1000 fps. Automated analysis of larval movement was performed using the  
695 FLOTE software package (Jain et al., 2014).

696 For optogenetic studies, the Tg(Gal4<sup>s1020t</sup>; UAS:ChR2) transgenic embryos were  
697 injected with control, *epha4a* or *efnb3b* morpholinos at one cell stage. At 5 dpf, the  
698 head of injected larva was mounted in 1% low melting-point agarose (Sigma) and the  
699 body was exposed in glass-bottom dishes (WPI). Leica M165FC fluorescence  
700 microscope was used to irradiate with a blue laser at 488nm wavelength. Larval  
701 responses were recorded using a high-speed video camera (Mikrotron, EoSens Mini1)  
702 at 500 fps. Body curvature analysis of larval movement was performed using the  
703 FLOTE software package.

## 704 **Whole-mount *in situ* hybridization and immunofluorescence**

705 The primer sequences used to amplify *epha4a*, *epha4b* and *rfng* genes were listed

in Table S7. Probe synthesis and whole-mount in situ hybridization were performed according to standard protocols. For reticulospinal neurons immunostaining, embryos were fixed in 2% trichloroacetic acid at 48 hpf for 3-4 h, washed twice in 0.5% Triton X-100 in PBS and blocked in 0.5% Triton X-100, 10% normal goat serum, 0.1% bovine serum albumin (Solarbio) in PBS for 1 h. The embryos were stained by monoclonal anti-neurofilament 160 antibody (Sigma-Aldrich) overnight at 4°C, then stained by goat anti-mouse Alexa Fluor 488 (Invitrogen) after washing.

### Analysis of neuronal calcium signals activity

Control, *epha4a* or *efnb3b* morpholinos were injected into Tg(*elavl3*:GAL4; UAS:GCaMP) transgenic embryos at one cell stage. At 24 hpf, embryos were paralyzed with 0.5 mg/ml  $\alpha$ -Bungarotoxin (AlomoneLabs), then mounted in 1% low melting-point agarose (Sigma) in glass-bottom dishes (WPI). Neuronal calcium signal images were collected with IXON-L-888 EMCCD camera equipped on Dragonfly 200 Spinning Disk Confocal Microscope using a 20 $\times$ /0.55 objective within 1 min at a frame rate of 10 fps.

Images were analyzed with imageJ software. To quantify the change in fluorescence intensity, a region of interest (ROI) was defined, and the fluorescence intensity  $F_t$  of different frames was normalized to  $F_t/F_{\min}$  based on the minimum fluorescence intensity  $F_{\min}$  in all frames. To compare the left-right alternation pattern of neuronal calcium signals, the ratio of the left-side calcium signals frequency ( $f_{\text{left}}$ ) to the right-side calcium signals frequency ( $f_{\text{right}}$ ) was logarithmically transformed (lg-transformed).

### Quantification and statistical analysis

Statistical analyses were performed in SPSS (version 15.0). Unpaired Student's t-test, Welch one-way ANOVA or two-way ANOVA followed by Turkey's multiple comparison test were applied when appropriate. All experiments were replicated at least three times independently.  $P < 0.05$  was considered statistically significant.

### 733 **Study approval**

734       Approval for the study was obtained from the ethics committee at the Peking  
 735       Union Medical College Hospital (JS-098, JS-2364), the medical ethics committee of  
 736       the Keio University Hospital (No. 20080129), the ethical committee of RIKEN  
 737       Yokohama Institute (No. H20-17(8)), and the Institutional Review Board of the  
 738       University Texas Southwestern Medical Center (protocol STU 112010□150).  
 739       Written informed consent was obtained from each participating individuals and  
 740       families in the three cohorts. For the control group, the protocols were approved by  
 741       the ethics committee at Peking Union Medical College Hospital. All zebrafish  
 742       studies were approved by the Animal Care Committee of the Ocean University of  
 743       China (Animal protocol number: OUC2012316).

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## 769 **Data and materials availability**

770 The datasets used and/or analyzed during the current study are available from  
771 the corresponding author on reasonable request.

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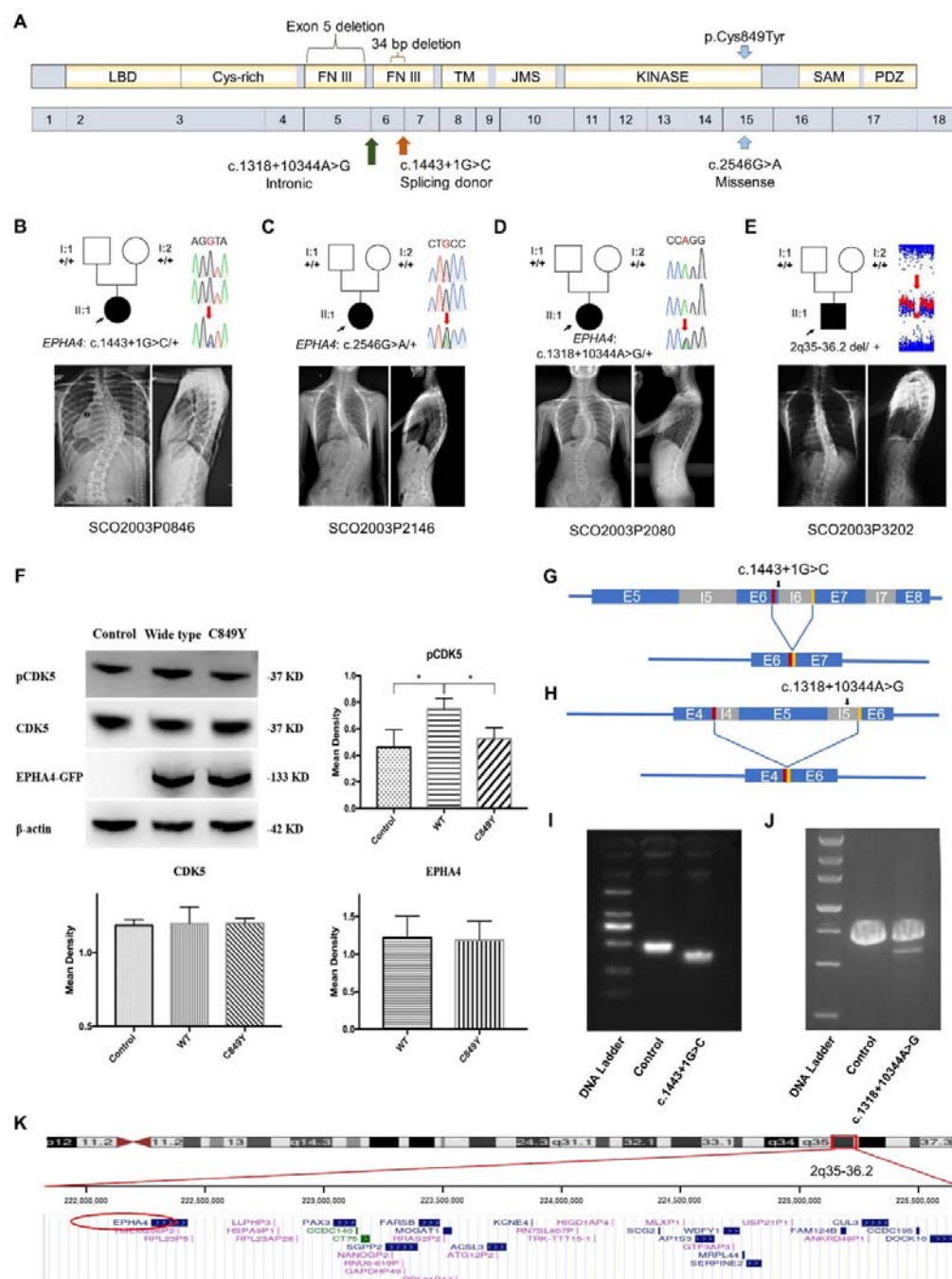
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**Table 1. Dominantly inherited variants identified in *EPHA4* and *NGEF*.**

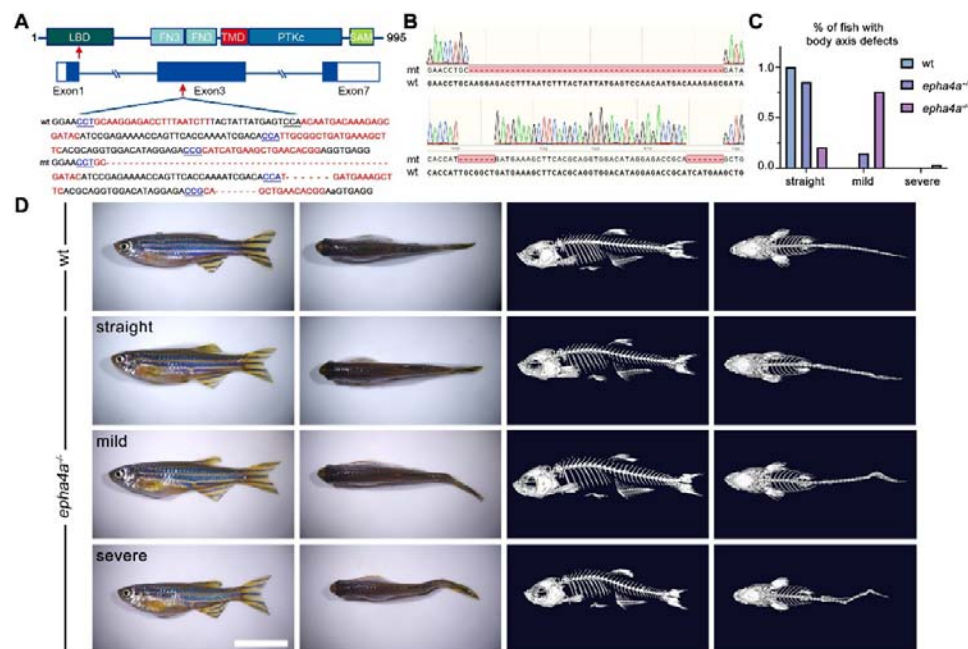
Patient ID	Chr	Gene	Ethnicity	Inheritance pattern	cDNA change	AA change	Variant type	ExAC PLI	CADD	GnomAD frequency	In-house frequency
SCO2003P0846	2	<i>EPHA4</i>	Chinese	De novo	c.1443+1G>C	NA	Splice donor	1	21.4	0	0
SCO2003P2146	2	<i>EPHA4</i>	Chinese	De novo	c.2546G>A	p.Cys849Tyr	Missense	1	28.2	0	0
SCO2003P2080	2	<i>EPHA4</i>	Chinese	De novo	c.1318+10344A>G	NA	Intronic	1	NA	0	0
SCO2003P3202	2	<i>EPHA4</i>	Chinese	De novo	2q35-36.2 4.6 Mb deletion	NA	CNV	NA	NA	0	0
SCO2003P3332	2	<i>NGEF</i>	Chinese	De novo	c.1A>G	p.Met1?	Start lost	0.95	12.8	0	0
TSRHC01	2	<i>NGEF</i>	Non-Hispanic White	AD	c.857C>T	p.Ala286Val	Missense	0.95	29.6	0.00002	0

Abbreviations: Chr (chromosomal localization); AA (amino acid); AD (autosomal dominant); CNV (copy number variant); ExAC PLI (probability of being loss-of-function intolerant from Exome Aggregation Consortium); CADD (Combined Annotation Dependent Depletion score); gnomAD (Genome Aggregation Database); NA (not applicable).

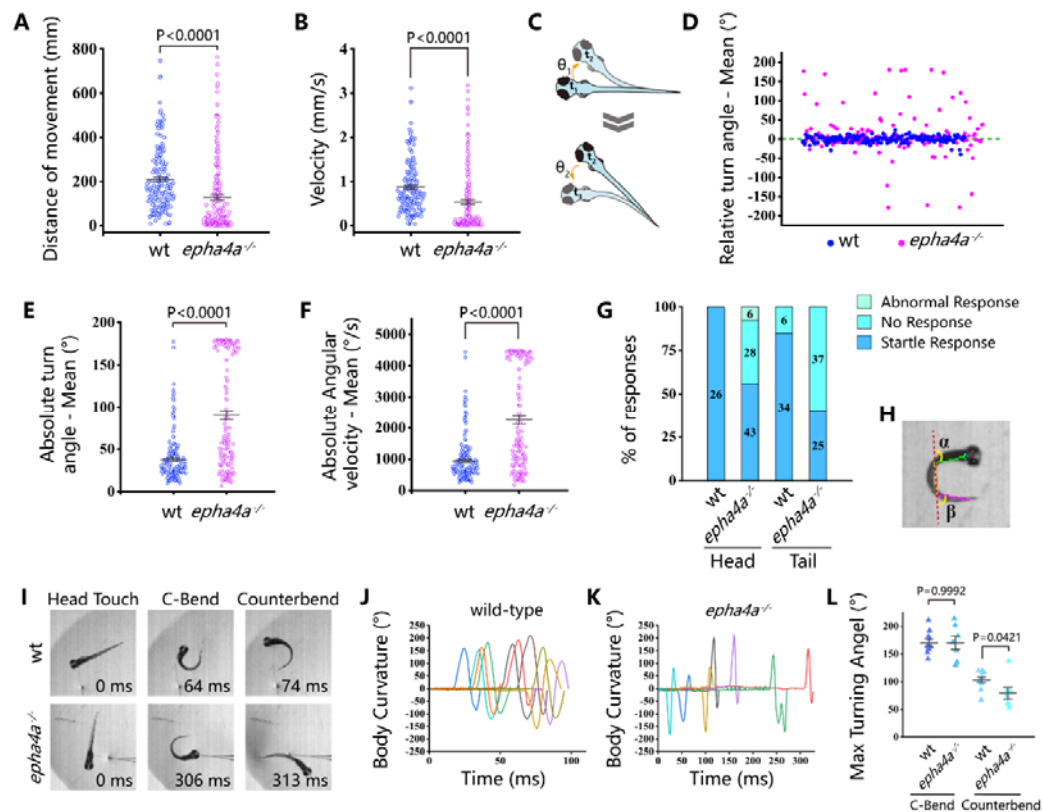


**Figure 1. Clinical and genetic information on IS patients and functional effect of *EPHA4* variants.** **A:** Locations of three *EPHA4* single nucleotide variants relative to the protein domains (top panel) and exons 1-18 (bottom panel). **B-E:** Pedigrees and spinal radiographs of four probands with dominant gene variants. Sanger sequencing confirmed the variants. The arrows indicate the probands. The term +/- denotes the wild-type, and cDNA change/+ denotes the heterozygous variant. **F:** Western blot analysis of *EPHA4*-c.2546G>A variant showing the protein expression levels of *EPHA4* and CDK5 and the amount of phosphorylated CDK5 (pCDK5) in HEK293T cells transfected with *EPHA4*-mutant or *EPHA4*-WT plasmid. WT: wild type. **G:**

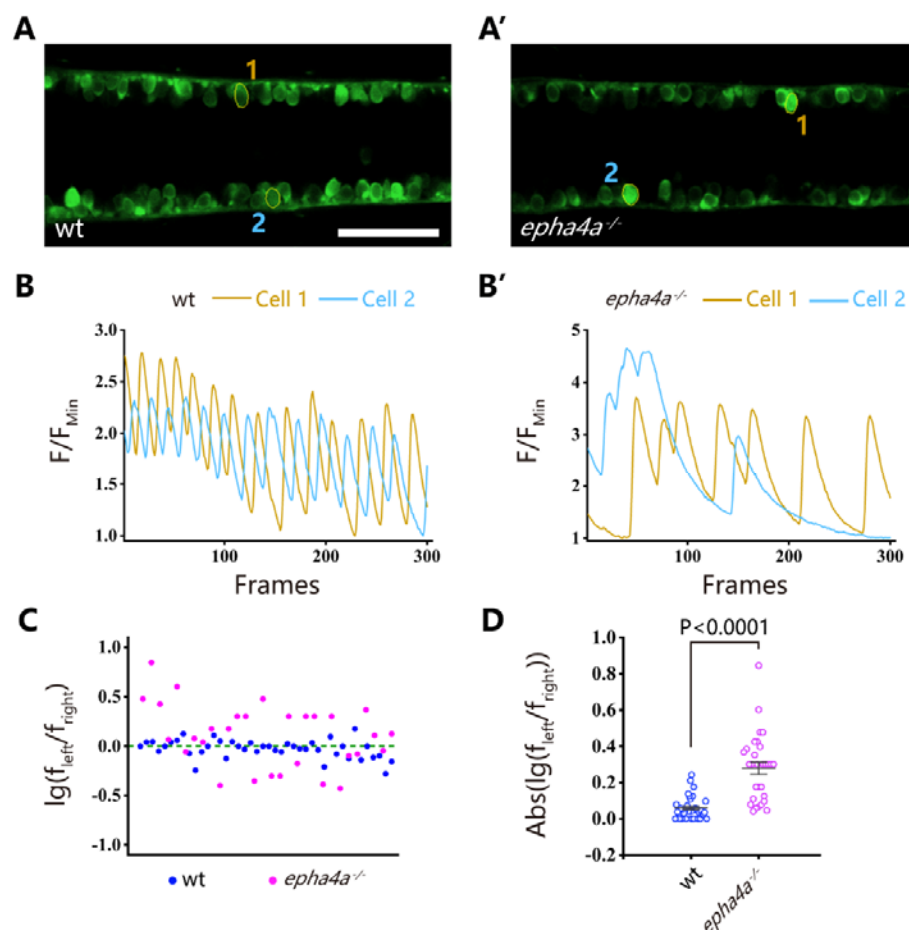
Schematic representation of the effect of the *EPHA4*-c.1443+1G>C mutation on the splicing process. This variant induced a new splicing site (red box). The yellow box indicates the splicing acceptor. **H:** Schematic representation of the effect of the *EPHA4*-c.1318+10344A>G mutation on the splicing process. This variant induced a new splicing site (red box). The yellow box indicates the splicing acceptor. **I:** The minigene assay result showed that the c.1443+1G>C variant introduced a new splicing site, resulting in a 36-bp in-frame deletion in exon 6. **J:** The nested PCR showed that the c.1318+10344A>G variant induced exon 5 skipping, resulting in a 339-bp in-frame deletion. **K:** NCBI RefSeq genes included in 2q35-q36.2 from UCSC Genome Browser. *EPHA4* is shown by the red oval.



**Figure 2. Scoliosis in zebrafish *epha4a* mutants.** **A:** Diagram of the protein domains, genomic structures, and sequences of wild-type and corresponding *epha4a* mutants. Red arrows indicate mutation sites. Blue boxes indicate open reading frames. Underlined sequences indicate the protospacer adjacent motif (PAM) region, and red fonts indicate Cas9 binding sites. LBD: ligand binding domain; FN3: fibronectin type 3 domain; TMD: transmembrane domain; PTKc: catalytic domain of the protein tyrosine kinases; SAM: sterile alpha motif. **B:** Sanger sequencing results confirmed the deletion of the target region in *epha4a* mutant transcripts. **C:** Bar graph showing the percentages of adult zebrafish with normal, mild, or severe body axis defects in wild-type (n=76), *epha4a* heterozygote (n=95), or *epha4a* homozygous (n=116). **D:** Representative images of wild-type and homozygous *epha4a* mutants. Micro CT images are shown on the right. Lateral and dorsal views are shown. Scale bar: 1 cm.

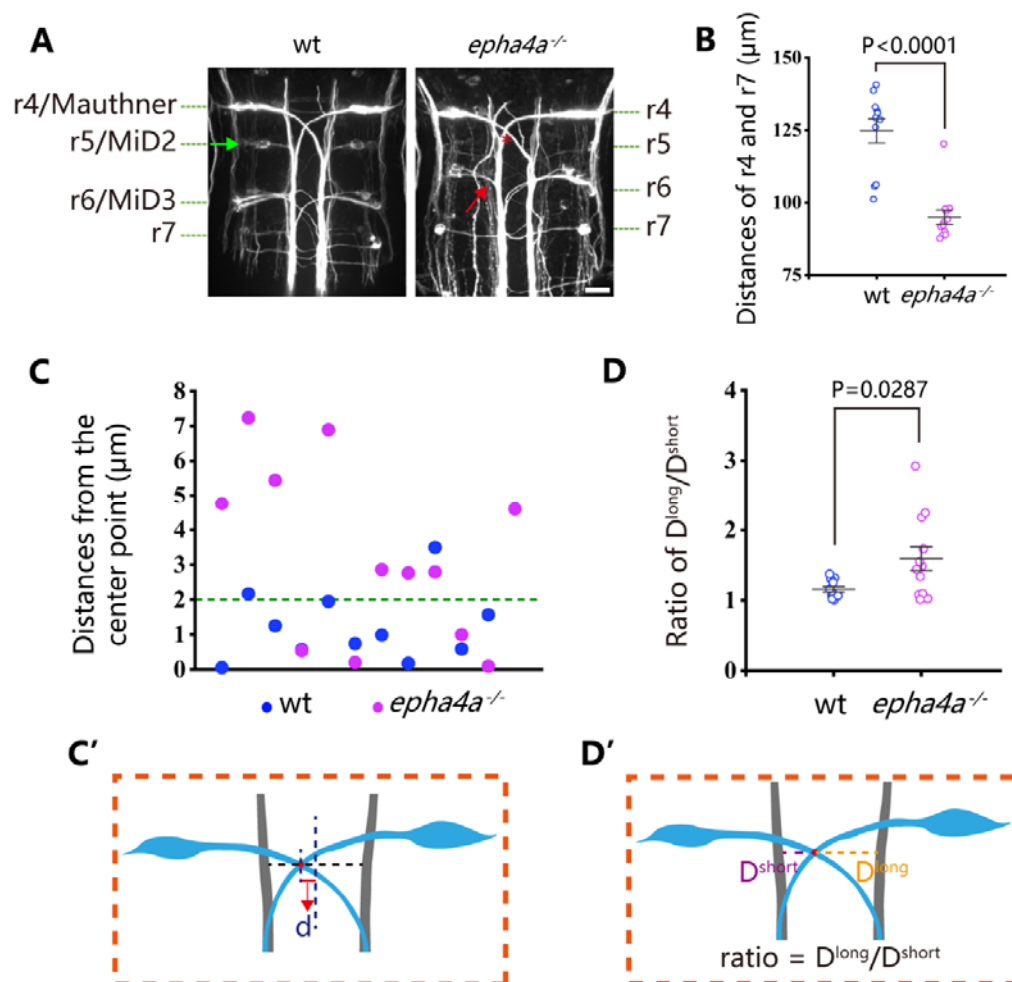


**Figure 3. Abnormal left-right swimming pattern in the absence of Epha4a. A:** Dot plots showing the swimming distance of each 8 dpf larva at a duration of 4 min (N=54 for wild-type and N=60 for *epha4a* mutants). **B:** Dot plots showing the swimming velocity of wild-type and mutant larvae as indicated. **C:** Diagram showing the turning angle ( $\theta$ ) of the larvae during swimming. **D:** Scatter plot showing the relative turning angle of wild-type and mutant larvae. The relative angles were calculated by the sum of turning angles during fish swimming with left (positive) or right (negative) turns. The *epha4a* mutants favored turning to one side of their directions compared with those of wild-type larvae. **E:** Dot plots showing the average absolute turning angle of wild-type and mutant larvae as indicated. **F:** Dot plots showing the average absolute angular velocity of wild-type and mutant larvae as indicated. **G:** Bar graph showing the percentages of 5 dpf zebrafish larvae with different reactions after tactile stimulation. N=10 for each group; the numbers of tactile stimulations are indicated in each column. **H:** Representative images of the total body curvature measurements in zebrafish larvae, with values as the sum of  $\alpha$  and  $\beta$  angles shown in the figure. **I:** Representative time-series images of 5 dpf wild-type and *epha4a* mutant zebrafish larvae after tactile stimulation to the head. Each panel represents the points of maximal body curvature for the C-bend and counterbend after the tactile startle response. **J, K:** A plot of body curvature angles as measured in panel (H) during swimming in response to tactile head stimulation in 5 dpf wild-type (Panel J, N=5 larvae, n=10 stimuli) and *epha4a* mutants (Panel K, N=5 larvae, n=7 stimuli). Each colored curve represents an independent experiment showing the response of a single larva to a stimulus. The positive angle means turning right. **L:** The maximum curvature angles during the first C-bend and counterbend after tactile stimulation in wild-type and *epha4a* mutant larvae.



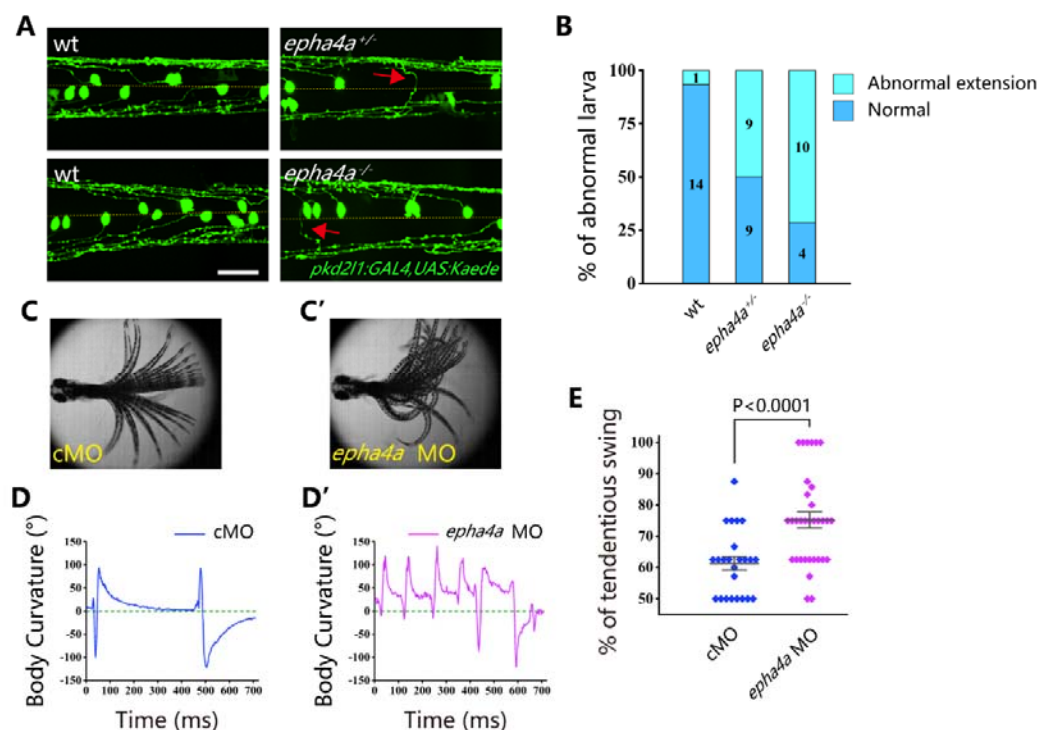
**Figure 4. Uncoordinated left-right activation of spinal cord neurons in *epha4a* mutants.** **A, A'**: Fluorescent images showing the dorsal-view of 24 hpf *Tg(elavl3:GAL4; UAS:GCaMP)* double transgenic larvae. The corresponding movies are shown in Movie S5 and S6. **B, B'**: Line charts showing the quantification of fluorescence changes of the region of interests (ROIs, circled in A, A') in wild-type larvae and *epha4a* mutants. **C**: Scatter plot showing the distribution trend of the ratio of the calcium signal frequency between left and right in wild-type (N=14 larvae, n=42 experiments) and *epha4a* mutants (N=10 larvae, n=30 experiments). **D**: Statistical graph of the ratio of the calcium signal frequency between left and right in wild-type larvae and *epha4a* mutants. Scale bars: 50  $\mu$ m in panel (A, A').



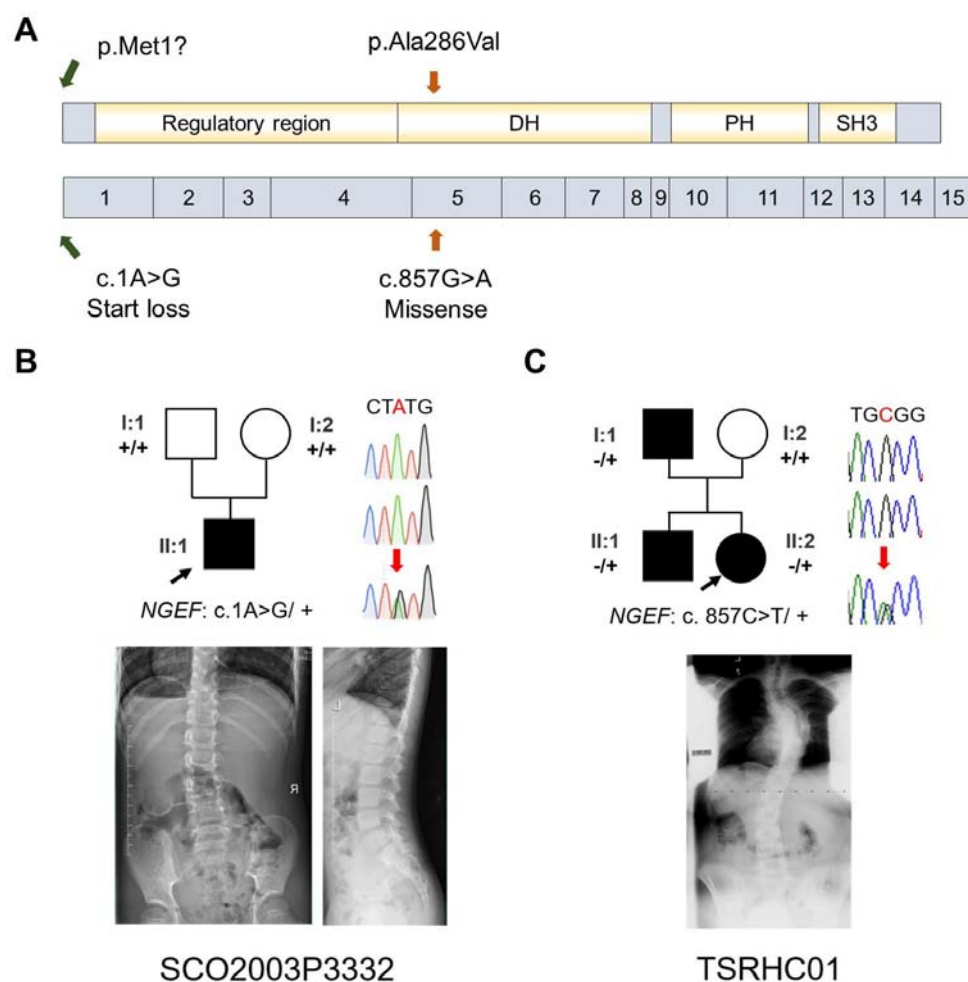


**Figure 5. Disorganized neural patterning in *epha4a* mutants.** **A:** Confocal images showing reticulospinal neuronal axons in 48 hours post-fertilization (hpf) wild-type and *epha4a* mutant larvae visualized with anti-neurofilament antibody RMO44. Asterisks indicate the cross sites of Mauthner axons. The green arrow indicates the cell body of the r5/MiD2 neuron in a wild-type larva. The red arrow points to the ipsilaterally projected axon of r6/MiD3 in the mutant larva, which is normally projected to the other side in wild-type fish. **B:** Statistical chart showing the distance between r4 and r7 of 48 hpf wild-type and *epha4a* mutants. **C:** Scatter plot showing the distance (d) between the center line and the intersection site of Mauthner axons as indicated in panel C'. **D:** The ratio of the distance between the intersection site of Mauthner axons and bilateral axon bundles in 48 hpf wild-type (N=11 larvae) and *epha4a* mutants (N=12 larvae). The ratios were calculated as in panel D'. Scale bars: 20 μm in panel (A).

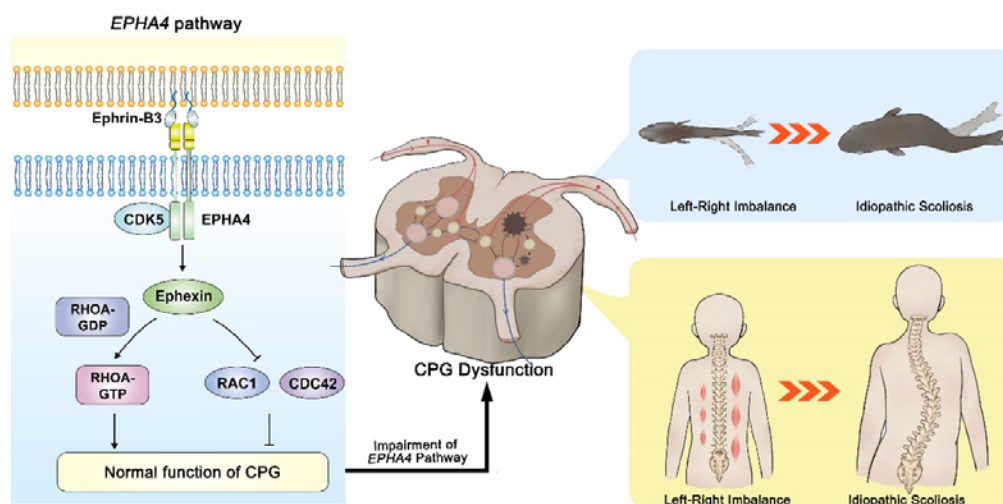




**Figure 6. Aberrant swimming as a result of abnormal extension of CSF-cNs axons.** **A:** Fluorescent images showing the distribution of ascending axons of CSF-cNs marked by Tg(*pkd2l1:GAL4,UAS:Kaede*) in 2 dpf *epha4a* mutant larvae. Yellow line indicates the midline and the red arrows indicate aberrantly extended axons in *epha4a*<sup>+/-</sup> and *epha4a*<sup>-/-</sup> larvae. **B:** Bar graph showing the percentages of abnormal extension of CSF-cNs axons in 2 dpf wild-type, *epha4a* heterozygote, and *epha4a* homozygous. The numbers of larvae are indicated in each column. **C, C':** Superimposed frames of tail oscillations in 5 dpf control and *epha4a* morphants. **D, D':** A plot of body curvature angles in panel (C) and (C'). The positive angle means turning right. **E:** Percentages of tendentious swing in control (N=8 larvae, n=24 experiments) and *epha4a* morphants (N=11 larvae, n=33 experiments). The percentages were calculated by the ratio of tendentious tail oscillation during the first eight swings. Scale bars: 50  $\mu$ m in panel (A).



**Figure 7. IS patients with potential *NGEF* variants.** **A:** Protein structure of NGEF protein with the position of potential variants. **B, C:** Pedigrees and spinal radiographs of two probands with dominant gene variants. Sanger sequencing results are shown on the right.



**Figure 8. The proposed mechanism of IS mediated by *EPHA4* dysfunction.** In healthy individuals, EphrinB3-activated *EPHA4* phosphorylates CDK5, leading to the phosphorylation of Ephexin, a protein encoded by *NGEF*. Phosphorylated Ephexin can regulate axon guidance through either activating RHOA or suppressing CDC42 and RAC1 signaling. These processes are critical to maintain the normal function of the CPG, the local neural network that provides coordinated bilateral muscle control. Impairment of the *EPHA4* pathway and CPG may cause an imbalance of the motor drive from the spinal cord during development, thus causing the uncoordinated left/right swimming behavior in zebrafish larvae and the asymmetry of the bilateral muscular pull in a young child. Although the appearance is normal in early childhood, the dysfunction produces a scoliotic curve during the growth spurt.

# Supplementary Materials for

## **Impaired central pattern generators due to abnormal EPHA4 signaling leads to idiopathic scoliosis**

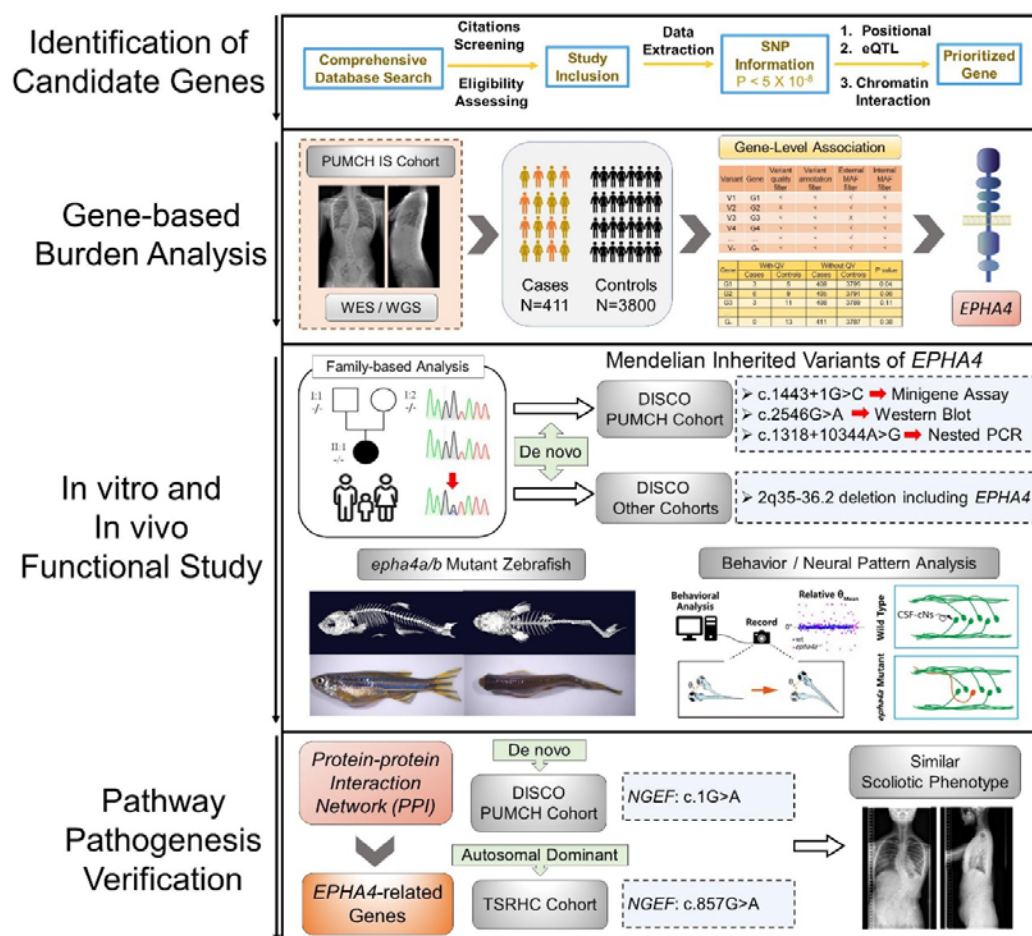
Lianlei Wang *et al.*

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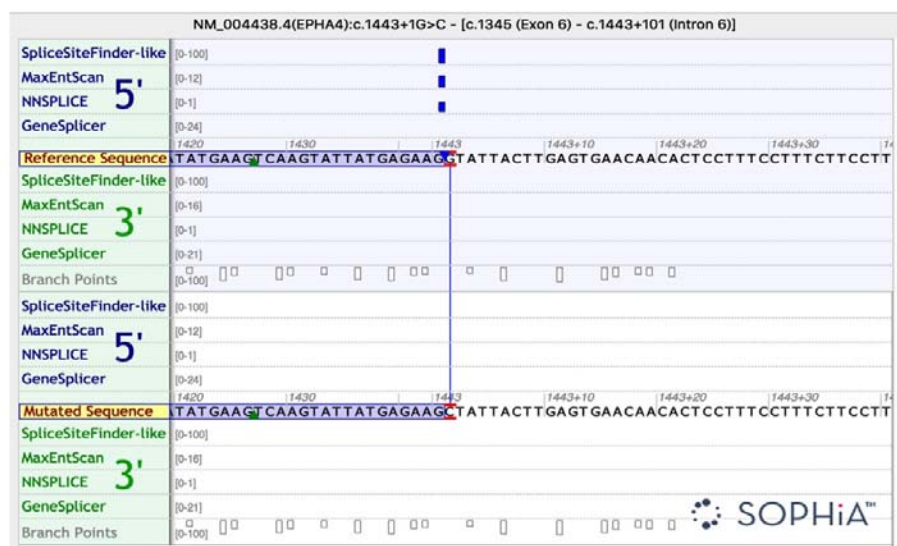
Chengtian Zhao, Ph.D., E-mail: [chengtian\\_zhao@ouc.edu.cn](mailto:chengtian_zhao@ouc.edu.cn)

Terry Jianguo Zhang, M.D., E-mail: [zhangjianguo@pumch.cn](mailto:zhangjianguo@pumch.cn)



**Supplementary Fig. 1. Flowchart for identification of causative genes.** The processes of candidate gene mapping and gene-based burden analysis were described. The variants in *EPHA4* and *NGEF* identified in each cohort were displayed. Abbreviations: SNP (single nucleotide polymorphism); eQTL (expression quantitative trait locus); IS (idiopathic scoliosis).

**A**

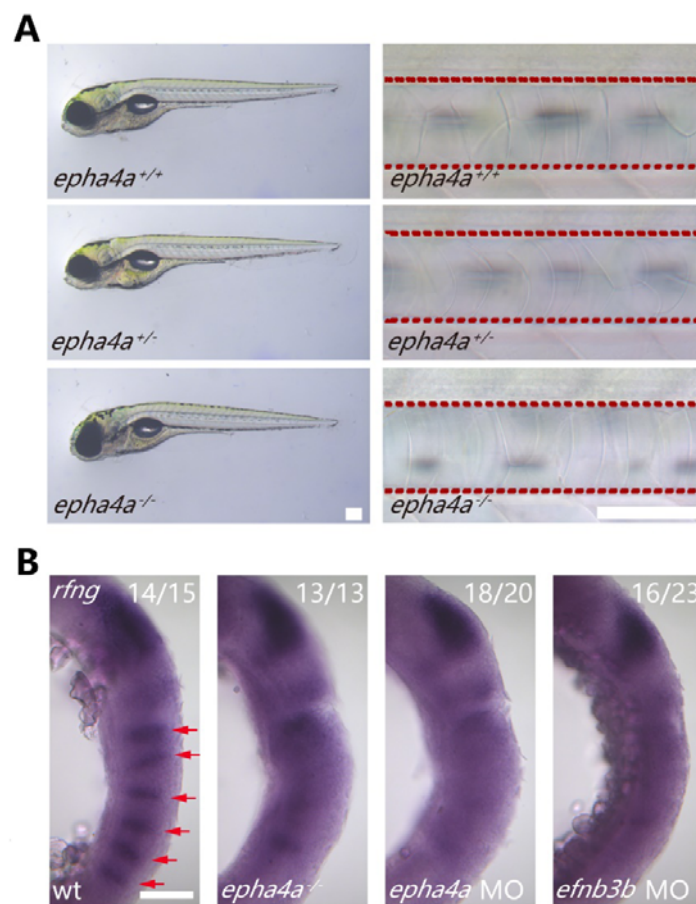


**B**

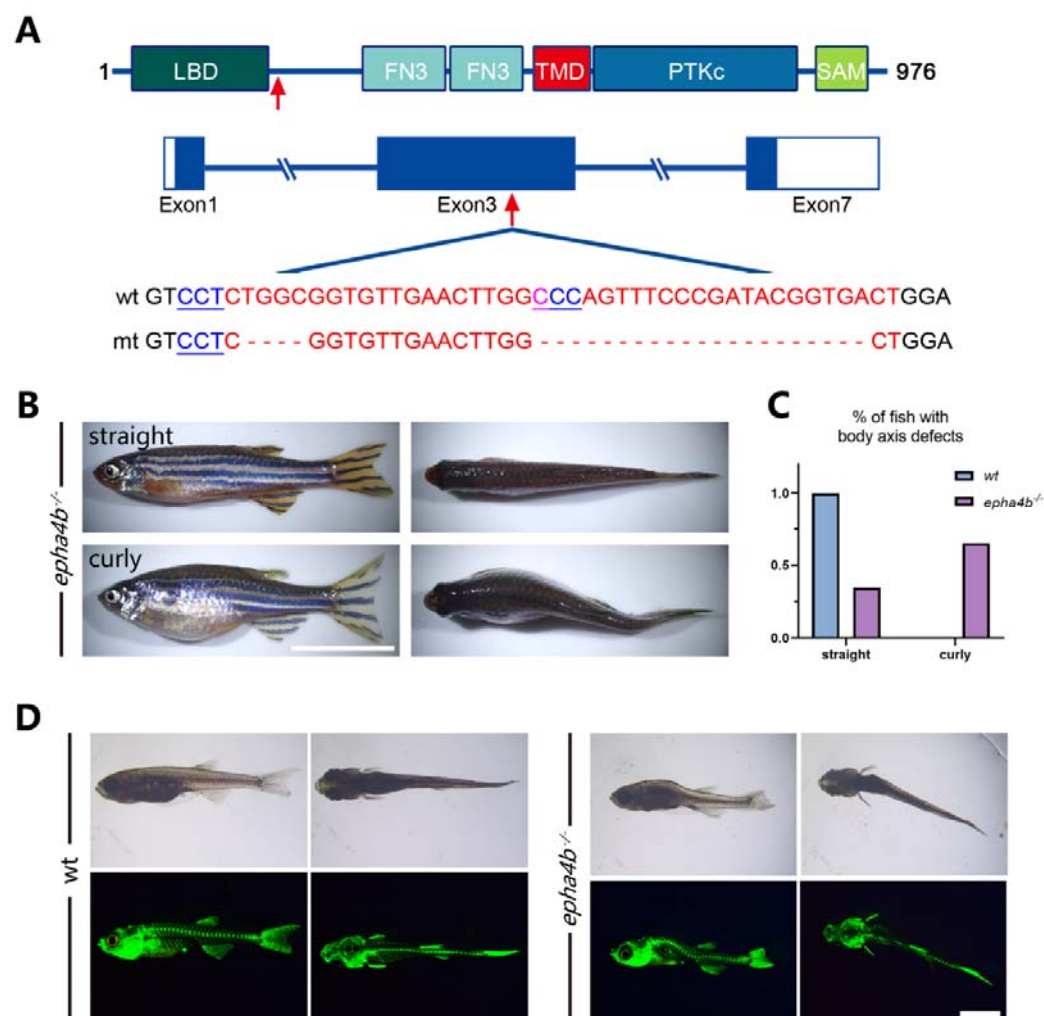


**Supplementary Fig. 2. The splicing analysis by the Alamut software.** The results of Alamut software by four algorithms (SpliceSiteFinder-like, MaxEntScan, NNSPLICE, and GeneSplicer).



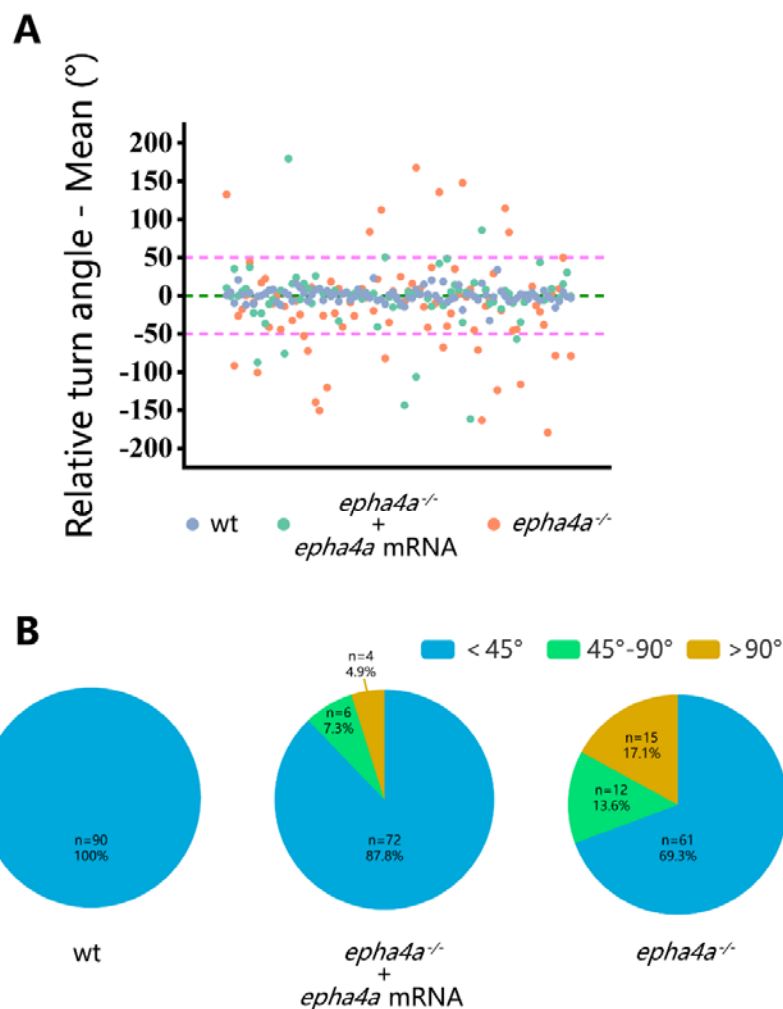


**Supplementary Fig. 3. Phenotypes of *epha4a* mutants.** **A:** Representative images showing the notochord and external phenotypes of wild-type and *epha4a* mutants. The red dashed lines indicate the area of the notochord. **B:** Whole-mount in situ hybridization results showing the expression of *rfn1* gene in 18 hpf wild type, homozygous mutant or morphant larvae as indicated. *rfn1* gene marks boundary cells in the hindbrain. The red arrows indicate boundary expression. The numbers of embryos analyzed are shown on top right. Scale bars: 200  $\mu$ m in panel (A), 100  $\mu$ m in panel (B).

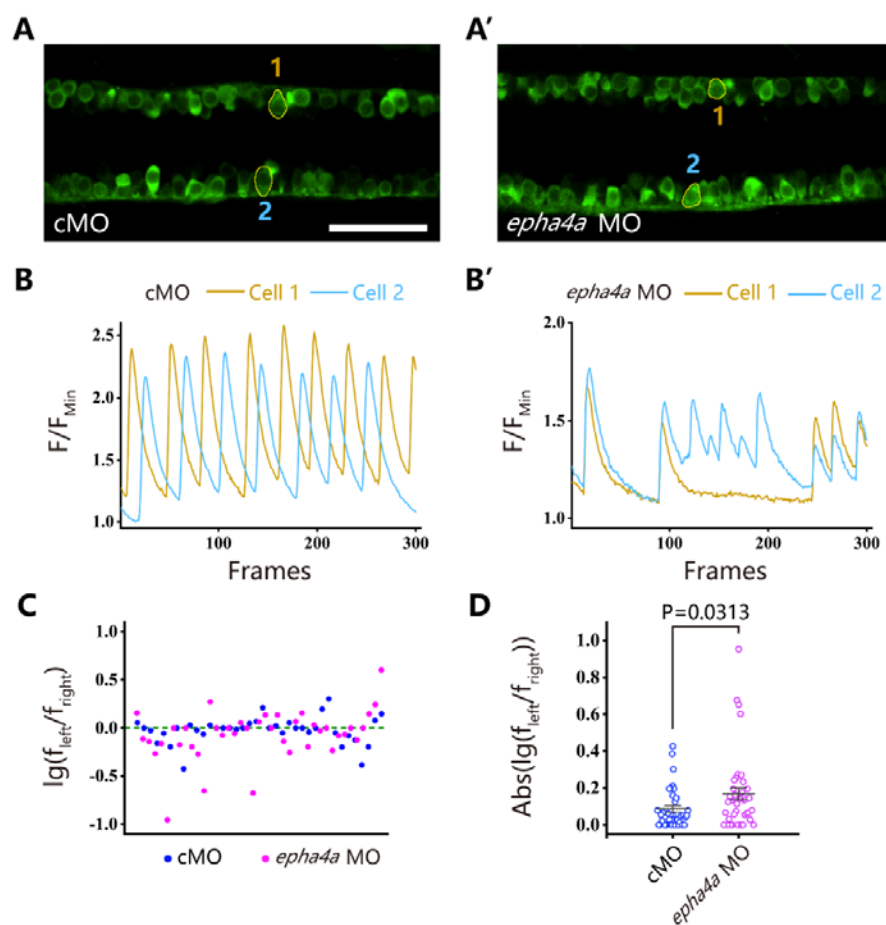


**Supplementary Fig. 4. Zebrafish *epha4b* mutants exhibited body axis defects during development.** **A:** Diagram showing the protein domains, genomic structures, and sequences of the wild-type and corresponding *epha4b* mutants. Red arrows indicate mutation sites. **B:** Representative images of *epha4b* mutants. **C:** Bar graph showing the percentages of adult fish with normal and body axis defects in wild-type (n=39) and *epha4b* mutants (n=43). **D:** Bright-field and GFP fluorescent images showing scoliosis *epha4b* mutants at 28 dpf as indicated by Tg(*Ola.Sp7:NLS-GFP*) transgene, which labels the bone skeleton. Scale bars: 1 cm in panel (B), 2 mm in panel (D).

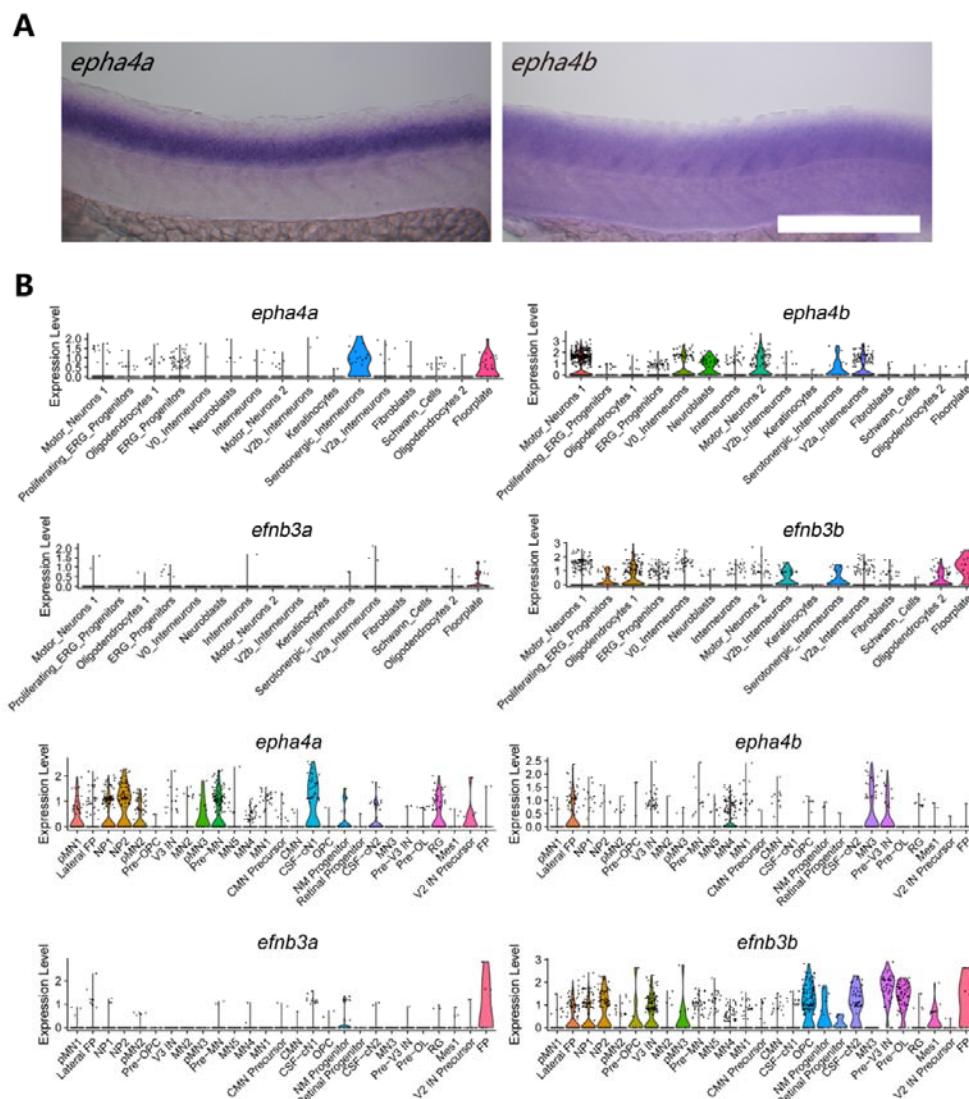




**Supplementary Fig. 5. Rescue of the *epha4a* homozygous mutant phenotype by *epha4a* mRNA injection.** A: Scatter plot showing the relative turning angle of wild-type, *epha4a* homozygous larvae and homozygous larvae injected with the corresponding mRNA. B: The pie chart shows the proportional distribution of relative turning angles in three groups of larvae. N=30 larvae, n=90 experiments. Larvae that remained stationary during the recording were excluded.

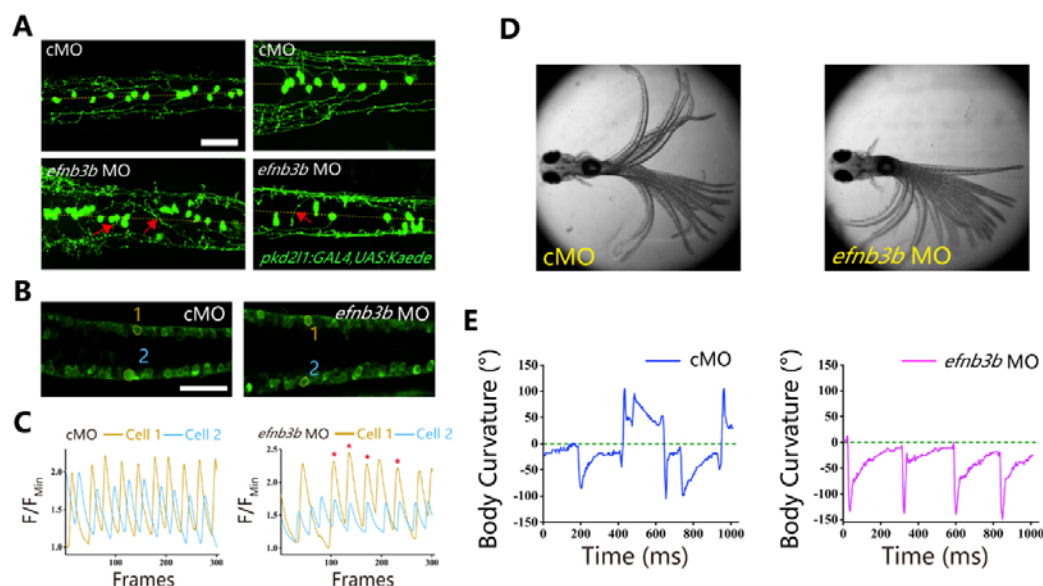


**Supplementary Fig. 6. Uncoordinated left-right activation of spinal cord neurons in the absence of EphA4a.** **A, A'**: Fluorescent images showing the dorsal-view of 24 hpf Tg(*elavl3*:GAL4; *UAS*:GCaMP) double transgenic larvae. The corresponding movies are shown in Movie S7 and S8. **B, B'**: Line charts showing the quantification of fluorescence changes of the region of interests (ROIs, circled in A, A') in control and *epha4a* morphants. **C**: Scatter plot showing the distribution trend of the ratio of the calcium signal frequency between left and right in control (N=15 larvae, n=38 experiments) and *epha4a* morphants (N=15 larvae, n=41 experiments). **D**: Statistical graph of the ratio of the calcium signal frequency between left and right in control and *epha4a* morphants. Scale bars: 50  $\mu\text{m}$  in panel (A, A').

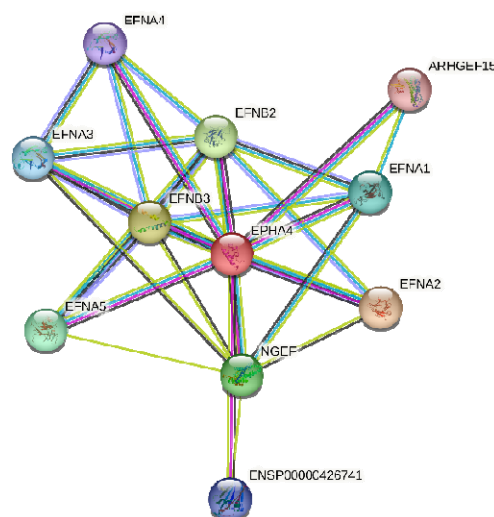


**Supplementary Fig. 7. Expression pattern of *epha4* and *efnb3* in the spinal cord.**

**A:** Whole-mount in situ hybridization results showing the expression of *epha4a* and *epha4b* genes in the spinal cord in 24 dpf zebrafish larvae. **B:** Violin plot showing *epha4a*, *epha4b*, *efnb3a*, and *efnb3b* gene expression in different cell types. Single cell transcriptome data were acquired from two different studies (top and bottom) (Cavone et al., 2021; Scott et al., 2021). Scale bars: 200  $\mu$ m in panel (A).



**Supplementary Fig. 8. Left-right coordination defects in *efnb3b* morphants.** **A:** Fluorescent images showing the distribution of ascending axons of CSF-cNs marked by *Tg(pkcd211:GAL4,UAS:Kaede)* in 2 dpf *efnb3b* morphants. The red arrows indicate aberrantly extended axons in *efnb3b* morphants. **B:** Fluorescent images showing the dorsal-view of 24 hpf *Tg(elavl3:GAL4; UAS:GCaMP)* transgenic larvae. **C:** The line chart showing the quantification of fluorescence changes of the ROIs in control morphants and *efnb3b* morphants as indicated in panel B. **D:** Superimposed frames of tail oscillations in 5dpf control and *efnb3b* morphants. **E:** A plot of body curvature angles in panel (D). The positive angle means turning right. Scale bars: 50  $\mu$ m in panel (A, B).



**Supplementary Fig. 9. Proteins in STRING that interact with EPHA4.** Protein interaction network analysis, as illustrated by STRING v11.5, was used to identify rare variants in genes related to *EPHA4*. Known and predicted protein-protein interactions are included in the STRING database. Multiple proteins, including Ephexin, are associated with EPHA4 physically or functionally. Ephexin, encoded by *NGEF*, is a neuronal guanine exchange factor.

**Supplementary Table 1. Summary of the 14 studies and their corresponding SNPs included in the candidate genes mapping.**

Citation	SNPs
Fan et al., J Hum. Genet. 2012	rs11190870 ( $p= 9.1 \times 10^{-10}$ )
Zhu et al., Hum. Mol. Genet. 2017	rs7593846 ( $p= 1.19 \times 10^{-13}$ )
	rs7633294 ( $p= 1.85 \times 10^{-12}$ )
	rs6570507 ( $p= 1.14 \times 10^{-11}$ )
	rs6047663 ( $p= 1.61 \times 10^{-15}$ )
Zhu et al., Nat. Commun. 2015	rs678741 ( $p= 9.68 \times 10^{-37}$ )
	rs4940576 ( $p= 2.22 \times 10^{-12}$ )
	rs13398147 ( $p= 7.59 \times 10^{-13}$ )
	rs241215 ( $p= 2.95 \times 10^{-9}$ )
Takahashi et al., Nat. Genet. 2011	rs11190870 ( $p= 1.24 \times 10^{-19}$ )
	rs625039 ( $p= 8.13 \times 10^{-15}$ )
	rs11598564 ( $p= 5.98 \times 10^{-14}$ )
Sharma et al., Hum. Mol. Genet. 2011	rs1400180 ( $p= 6.35 \times 10^{-8}$ )
Kou et al., Nat. Genet. 2013	rs6570507 ( $p= 1.27 \times 10^{-14}$ )
Xu et al., Spine Deform. 2018	rs169311 ( $p= 2.10 \times 10^{-8}$ )
Grauers et al., Spine J. 2015	rs11190870 ( $p= 7.0 \times 10^{-18}$ )
Ogura et al., Am. J of Hum. Genet. 2015	rs3904778 ( $p= 2.46 \times 10^{-13}$ )
Khanshour et al., Hum. Mol. Genet. 2018	rs4513093 ( $p= 1.71 \times 10^{-15}$ )
	rs1455114 ( $p= 2.99 \times 10^{-8}$ )
	rs687621 ( $p= 7.29 \times 10^{-10}$ )
	rs10756785 ( $p= 7.00 \times 10^{-10}$ )
Sharma et al., Nat. Commun. 2015	rs6137473 ( $p= 3.12 \times 10^{-8}$ )
Miyake et al., PLoS One 2013	rs11190870 ( $p= 2.80 \times 10^{-18}$ )
	rs625039 ( $p= 1.28 \times 10^{-15}$ )
	rs12946942 ( $p= 6.43 \times 10^{-12}$ )
	rs11598564 ( $p= 9.77 \times 10^{-12}$ )
	rs6570507 ( $p= 3.78 \times 10^{-8}$ )
	rs9496346 ( $p= 1.00 \times 10^{-8}$ )

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Chettier et al., PLoS One 2015	rs11190878 (p= $4.18 \times 10^{-9}$ )
Kou et al., Nat. Commun. 2019	rs9389985 (p= $3.51 \times 10^{-20}$ )
	rs7028900 (p= $2.19 \times 10^{-17}$ )
	rs144131194 (p= $1.35 \times 10^{-11}$ )
	rs6047716 (p= $1.45 \times 10^{-11}$ )
	rs141903557 (p= $9.78 \times 10^{-11}$ )
	rs11205303 (p= $1.62 \times 10^{-10}$ )
	rs12029076 (p= $2.17 \times 10^{-10}$ )
	rs1978060 (p= $3.26 \times 10^{-10}$ )
	rs2467146 (p= $5.96 \times 10^{-10}$ )
	rs11787412 (p= $1.32 \times 10^{-9}$ )
	rs188915802 (p= $1.94 \times 10^{-9}$ )
	rs658839 (p= $3.15 \times 10^{-9}$ )
	rs2194285 (p= $8.69 \times 10^{-9}$ )
	rs160335 (p= $9.10 \times 10^{-9}$ )
	rs482012 (p= $2.30 \times 10^{-8}$ )
	rs11341092 (p= $2.92 \times 10^{-8}$ )
	rs17011903 (p= $3.56 \times 10^{-8}$ )
	rs397948882 (p= $3.66 \times 10^{-8}$ )
	rs12149832 (p= $4.40 \times 10^{-8}$ )

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<b>Rank</b>	<b>Gene Symbol</b>	<b>P value</b>	<b>Variants in Cases (N = 411)</b>	<b>Variants in Controls (N = 3,800)</b>	<b>OR (95%CI)</b>
1	<i>EPHA4</i>	0.045533	3	17	4.09 (0.946-17.6)
2	<i>ARVCF</i>	0.067987	6	24	3.05 (0.807-11.6)
3	<i>TXNRD2</i>	0.092043	3	11	3.60 (0.729-17.8)
4	<i>TRIM17</i>	0.115033	3	11	3.53 (0.795-15.7)
5	<i>SLC25A1</i>	0.119166	3	9	3.39 (0.817-14.1)
6	<i>MICALL2</i>	0.199131	0	24	0.30 (0.0176-4.94)
7	<i>MTMR11</i>	0.249127	5	35	2.05 (0.663-6.36)
8	<i>MMP2</i>	0.252208	4	17	3.03 (0.613-15.0)
9	<i>BCKDHB</i>	0.293142	1	16	0.643 (0.0469-8.83)
10	<i>TTK</i>	0.293162	0	17	0.583 (0.0334-10.2)

1 **Supplementary Table 2. Summarized results of burden analysis.**



11	<i>SLC2A6</i>	0.316088	1	21	0.675 (0.049-9.3)
12	<i>SOX6</i>	0.317369	2	25	0.822 (0.0949-7.11)
13	<i>WNT3A</i>	0.329116	1	11	0.814 (0.058-11.4)
14	<i>SDR42E1</i>	0.33538	0	11	0.631 (0.0359-11.1)
15	<i>GPR126</i>	0.33628	0	12	0.699 (0.0393-12.4)
16	<i>PIGN</i>	0.338169	1	22	0.663 (0.0869-5.06)
17	<i>POLL</i>	0.348494	0	11	0.668 (0.0378-11.8)
18	<i>PAX1</i>	0.350574	0	19	0.649 (0.0368-11.4)
19	<i>INSC</i>	0.363501	0	13	0.744 (0.0416-13.3)
20	<i>IBA57</i>	0.388644	0	8	0.744 (0.0416-13.3)

2        The p-values (Fisher's Exact Test) of 20 top ranked gene were calculated on the  
3 basis of distribution differences of rare variant between the control group and the  
4 PUMCH IS cohort. Abbreviations: OR (odds ratio); CI (confidence interval).

5 **Supplementary Table 4. Search strategies for each database.**

<b>Search Number</b>	<b>Key Words</b>	<b>Results</b>
<b>MEDLINE (via Pubmed.gov)</b>		
1#	Idiopathic scoliosis, GWAS/	69
2#	Idiopathic scoliosis, SNP/	56
3#	Idiopathic scoliosis, single nucleotide polymorphism/	149
4#	Idiopathic scoliosis, variant/	129
<b>Web of Science (via Clarivate Analytics)</b>		
1#	Idiopathic scoliosis, GWAS/	42
2#	Idiopathic scoliosis, SNP/	71
3#	Idiopathic scoliosis, single nucleotide polymorphism/	198
4#	Idiopathic scoliosis, variant/	227

**6 Supplementary Table 5. Inclusion and exclusion criteria of literature review.**

	<b>Inclusion Criteria</b>	<b>Exclusion Criteria</b>
Patient	Idiopathic scoliosis	Congenital scoliosis
Recruitment	Adolescent idiopathic scoliosis	Neuromuscular scoliosis
	Early-onset idiopathic scoliosis	Syndromic scoliosis
		Secondary scoliosis
		Degenerative scoliosis
Study Type	Genome Wide Association	Linkage study
	Study (GWAS)	Candidate gene study
	Meta-analysis of GWAS	Exome sequencing
		Whole genome sequencing
Study Design	Case-control	-

7     **Supplementary Table 6. Sequencing information of PUMCH IS cohort.**

<b>Characteristics</b>	<b>Cases</b>		<b>Controls</b>	
	<b>Trio</b>	<b>Singleton</b>	<b>Trio</b>	<b>Singleton</b>
Exome sequencing	41	113	2,021	224
Genome sequencing	116	148	483	1,072
Sum	157	261	2,504	1,296
Passing QC	155	256	2,504	1,296

8     Abbreviations: QC (quality control); IS (idiopathic scoliosis).

9 **Supplementary Table 7. Sequences information of sgRNA and primers for**  
 10 **zebrafish study.**

sgRNA sequences used to generate zebrafish <i>epha4a</i> and <i>epha4b</i> mutants	
<i>epha4a</i> sgRNA-1	AAGATTAAAGGTCTCCTTGC
<i>epha4a</i> sgRNA-2	GTATCGCTCTTTGTCATTGT
<i>epha4a</i> sgRNA-3	GAAGCTTTCATCAGCCGCAA
<i>epha4a</i> sgRNA-4	CCGTGTTTCAGCTTCATGATG
<i>epha4b</i> sgRNA-1	GCCAAGTTCAACACCG-CCAG
<i>epha4b</i> sgRNA-2	AGTCACCGTATCGGGAAACT
Primers for mutant genotype	
<i>epha4a</i> -forward	GAGCTCAGCGGGTCTACATC
<i>epha4a</i> -reverse	GCGTACCTCCACTAGCGATG
<i>epha4b</i> -forward	GGTGTCTCTGAGGGTCTTTT
<i>epha4b</i> -reverse	CCTGACAGTAGTCCTTATTCTCCTC
Morpholinos used to knock down the expression of <i>epha4a</i> and <i>efnb3b</i>	
<i>epha4a</i> MO	AACACAAGCGCAGCCATTGGTGTC
<i>efnb3b</i> MO1	GTCTATTTACTCCCATCAAAGCCGT
<i>efnb3b</i> MO2	GAAATCCCGAATTCCGTCTATTTAC
Primers used for in situ hybridization analysis	
<i>epha4a</i> probe forward	AGAATACGCCAATCAGGACG

<i>epha4a</i> probe reverse	CCTTCATGTTCTTCATAGCC
<i>epha4b</i> probe forward	TTTGGTGGAGGTTAGAGGGT
<i>epha4b</i> probe reverse	GTTGTTTAGGGGTTGGGCTC
<i>rfng</i> probe forward	GGTGCTGCTTATCCCTCCTT
<i>rfng</i> probe reverse	GCGCTAAACCTCGACTGATGC

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- 11 **Supplementary Movie 1:** Video showing the swimming of three wild-type fish.
- 12 **Supplementary Movie 2:** Video showing the swimming of three *epha4a* mutants
- 13 with severe or mild scoliosis.
- 14 **Supplementary Movie 3:** High-speed video showing the startle response in a
- 15 wild-type larva at 5 dpf triggered by head tactile stimulation. Time units: ms.
- 16 **Supplementary Movie 4:** High-speed video showing the abnormal startle response of
- 17 an *epha4a* mutant larva at 5 dpf triggered by head tactile stimulation. Time units: ms.
- 18 **Supplementary Movie 5:** Alternated activation of calcium signaling in motor
- 19 neurons of 24 hpf wild type Tg(*elavl3*:GAL4; *UAS*: GCaMP) transgenic larva.
- 20 **Supplementary Movie 6:** Alternated activation of calcium signaling in motor
- 21 neurons of 24 hpf wild type Tg(*elavl3*:GAL4; *UAS*: GCaMP) transgenic larva injected
- 22 with control MO.
- 23 **Supplementary Movie 7:** Abnormal activation of calcium signaling in motor neurons
- 24 of 24 hpf Tg(*elavl3*:GAL4; *UAS*: GCaMP) transgenic larva with homozygous *epha4a*
- 25 mutant background.
- 26 **Supplementary Movie 8:** Abnormal activation of calcium signaling in motor neurons
- 27 of 24 hpf Tg(*elavl3*:GAL4; *UAS*: GCaMP) transgenic larva injected with *epha4a* MO.
- 28 **Supplementary Movie 9:** Tail oscillation after light activation of 5dpf Tg(Gal4<sup>s1020t</sup>;
- 29 *UAS*:ChR2) double transgene larva.
- 30 **Supplementary Movie 10:** Tail oscillation after light activation of 5dpf Tg(Gal4<sup>s1020t</sup>;
- 31 *UAS*:ChR2) double transgene larva injected with *epha4a* MO.