

# Mouse *hitchhiker* mutants have spina bifida, dorso-ventral patterning defects and polydactyly: identification of Tulp3 as a novel negative regulator of the Sonic hedgehog pathway

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The mammalian Sonic hedgehog (Shh) signalling pathway is essential for embryonic development and the patterning of multiple organs. Disruption or activation of Shh signalling leads to multiple birth defects, including holoprosencephaly, neural tube defects and polydactyly, and in adults results in tumours of the skin or central nervous system. Genetic approaches with model organisms continue to identify novel components of the pathway, including key molecules that function as positive or negative regulators of Shh signalling. Data presented here define Tulp3 as a novel negative regulator of the Shh pathway. We have identified a new mouse mutant that is a strongly hypomorphic allele of Tulp3 and which exhibits expansion of ventral markers in the caudal spinal cord, as well as neural tube defects and preaxial polydactyly, consistent with increased Shh signalling. We demonstrate that Tulp3 acts genetically downstream of Shh and Smoothed (Smo) in neural tube patterning and exhibits a genetic interaction with Gli3 in limb development. We show that Tulp3 does not appear to alter expression or processing of Gli3, and we demonstrate that transcriptional regulation of other negative regulators (Rab23, Fkbp8, Thm1, Sufu and PKA) is not affected. We discuss the possible mechanism of action of Tulp3 in Shh-mediated signalling in light of these new data.

## INTRODUCTION

Birth defects remain the greatest cause of infant mortality in the Western world. These developmental anomalies can affect any organ in the body and are often severely disabling or fatal. In the UK, over 90 000 pregnancies per year (~2.3%) are affected by a major congenital malformation, and amongst the most common anomalies are defects of the nervous system and limbs, which are observed in 10 and 16% of affected fetuses, respectively (1). Reducing the clinical burden from birth defects requires a more complete understanding of the causative factors, including knowledge of the genetic, cellular and molecular mechanisms involved, in order to permit the develop-

ment of appropriate preventative therapies. Studies with model organisms are essential in helping to unravel the complex network of processes involved in normal development, and the identification of mutant animals provides key insights into the pathways that are involved in these disorders. Mutagenesis of mice with ethylnitrosourea (ENU) has proven to be a powerful method for the generation of new mutations that provide good models of human developmental disorders (2–4). Investigation of these mutants promises to shed new light on the causes of these defects as well as identifying novel components of known signalling pathways.

Development of the mammalian nervous system involves the initial formation of the neural tube, the precursor of the brain and

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spinal cord, and the subsequent differentiation of the neural tissue into spatially appropriate neuronal cell types. The rolling up and closure of the neural plate to form the neural tube constitutes the process of neurulation, and this occurs with a characteristic sequence of events and varying neural fold morphology in different regions of the embryo (5–7). Initiation of neural tube closure occurs at the level of the cervical/hindbrain boundary (Closure 1) at E8.5 in mouse, and continues from here both cranially into the hindbrain and caudally along the developing spine. Two further sites of closure initiation are observed in the cranial region, at the forebrain–midbrain boundary (Closure 2) and at the most rostral extent of the forebrain (Closure 3). Neurulation in the head is completed with closure of the intervening regions, termed the anterior neuropore and hindbrain neuropore, by E9.5. Neural tube closure continues until E10.5 in a cranio-caudal direction from the site of Closure 1 along the developing spine. The region immediately caudal to the closed neural tube, in which the neural folds are elevating and about to close, is termed the posterior neuropore (PNP). The morphology of the neural folds within the PNP varies along the spinal axis, and this has led to the characterization of three modes of spinal neurulation (6). In mode 1 (E8.5–E9), the neural plate exhibits pronounced bending at the ventral midline, the median hinge point (MHP), but straight lateral folds, whereas in mode 2 (E9–E9.75) the neural folds bend both at the MHP and also at paired dorso-lateral hinge points (DLHPs). Closure of the upper sacral level involves mode 3 neurulation (E9.75–E10.5), in which the neural plate bends solely at the DLHPs.

The Sonic hedgehog (Shh) signalling pathway is essential for the normal growth and patterning of many organs including the nervous system and limbs (8,9). Disruption of Shh signalling results in a range of congenital defects, including holoprosencephaly, craniofacial abnormalities and skeletal malformations (8,9). Conversely, inappropriate activation of the pathway leads to neural tube defects, polydactyly, basal cell carcinomas and tumours of the central nervous system (7,10,11). Shh plays a key role in regulating closure of the neural tube since Shh expression in the notochord is necessary and sufficient to inhibit formation of DLHPs, through a mechanism involving antagonism of Bmp signalling (12,13). Shh signalling from the notochord and, subsequently, the floor plate is also important for correct dorso-ventral (DV) patterning of the nervous system (14–18). The spatially restricted specification of neuronal subtypes is essential for the subsequent assembly of functional neuronal circuits. High concentrations of Shh induce floor plate markers, whereas lower concentrations specify increasingly more dorsal interneuron subtypes within the ventral neural tube. Absence of Shh signalling results in loss of ventral markers and complete dorsalization of the spinal cord, whereas ectopic Shh signalling leads to expansion of ventral markers at the expense of more dorsal marker domains. Patterning along the anterior–posterior (AP) axis of the limb is also dependent on the Shh pathway (19–21). Shh is expressed in the zone of polarizing activity (ZPA) in the posterior domain of the limb buds, and the spatial and temporal gradients of Shh activity across the limb determine digit number and identity. Increased or ectopic Shh activity leads to polydactyly, whereas decreased Shh activity results in narrowed limbs and reduced number of digits.

The vertebrate Shh pathway shares fundamental similarities with that of *Drosophila* (22–24). Hh ligand binds to Patched

(Ptc) receptor, releasing the repressive activity of Ptc on Smoothed (Smo) and, thereby, allowing Smo to initiate intracellular signalling. Downstream of Smo the signal culminates with the activation of the Ci/Gli transcriptional activators and removal of the Ci/Gli repressors. Vertebrates express three Gli family members, Gli1–3; Gli1 functions as a transcriptional activator, whereas Gli2 and Gli3 can act both as transcriptional activators and, following proteolytic removal of the C-terminal region, as transcriptional repressors. The relative importance of the Gli activator and repressor forms differs in the patterning of the neural tube and limb. Neural tube patterning is mostly dependent on ventral Gli1 and Gli2 activators, with less important roles of Gli3 repressor dorsally (15,25–29). In contrast, limb patterning is generated predominantly by Gli3, with opposing gradients of Gli3 activator and Gli3 repressor (30–32). Although vertebrate and *Drosophila* hedgehog pathways are grossly similar, the intermediate steps between the release of Smoothed and the activation or processing of Ci/Gli are distinct between species and, indeed, are not yet fully understood (9,22,33,34).

Forward genetic screens in mouse have successfully identified additional components of the vertebrate Shh pathway, with the advantage of no prior bias on the type of molecule that may be involved (35,36). A number of positive regulators of Shh signalling have recently been discovered, based on loss of ventral cell types in the neural tube, including proteins involved in the formation/maintenance of the primary cilia or intraflagellar transport (37). Several Shh pathway components localize within these subcellular organelles (38–42), and the general consensus is that some aspects of vertebrate Smo-mediated Gli processing occur within cilia. Specific proteins have been revealed as negative regulators of the pathway, since disruption of their function leads to activation of Shh signalling, as observed by the expansion of ventral marker domains within the neural tube. The known negative regulators include Ptc1, Sufu, protein kinase A (PKA), Rab23, Fkbp8 and Thm1 (42–51). PKA phosphorylates Gli3 prior to processing by the ubiquitin-proteasome pathway (52), whereas Sufu inhibits the transcriptional activity of Gli activators in the nucleus and may also sequester Gli in the cytoplasm (49,50,53,54). Thm1 is involved in intraflagellar transport (42). The functions of Rab23 and Fkbp8 are not clear, although they may have roles in vesicular trafficking (55,56). Clearly, the vertebrate Shh pathway is complex, and our incomplete understanding of the processing and regulation of intermediate steps suggests that additional components remain to be discovered.

Here, we report a new ENU-induced mouse mutant that we have named *hitchhiker* (*hhkr*), which exhibits a range of developmental abnormalities including neural tube defects and polydactyly. We reveal that *hitchhiker* carries a strongly hypomorphic mutation in *Tulp3*, and we show that mutants have ventralization of the caudal spinal cord. We demonstrate that *Tulp3* functions genetically downstream of *Shh* and *Smo*, to give activation of Shh pathway targets in the absence of Shh and Smo; complementary results are reported in an accompanying paper (Norman *et al.*, submitted for publication). In addition, we show that the *hitchhiker* mutant exhibits a genetic interaction with *Gli3* in limb development, but does not substantially affect Gli3 processing or expression.

We demonstrate that transcriptional regulation of other negative regulators (Rab23, Fkbp8, Thm1, Sufu and PKA) is not affected in *hitchhiker* mutants. We propose that Tulp3 functions as a novel negative regulator of the Shh pathway and we discuss the possible mechanism of action of Tulp3 in Shh-mediated signalling in light of these new data.

## RESULTS

### *hitchhiker* is a novel mutant with defects in neural tube and limb development

*hitchhiker* (*hhkr*) arose during an on-going screen for recessive ENU-induced mutations that affect the morphology of mid-gestation embryos. The mutant was initially identified through the presence of neural tube defects, with exencephaly and/or lumbosacral spina bifida aperta, and mutants also commonly exhibited oedema (Fig. 1A–D). Histological sections revealed abnormal morphology of the lower spinal neural tube in mutants in which the neural tube had closed, with an expanded lumen and reduced thickness of the neuroepithelium, particularly in the ventral half of the spinal cord (Fig. 1F, compare with E). Mutants also demonstrated oedema in the surrounding tissue, and small, dorsally misplaced dorsal root ganglia (Fig. 1F). Sections through other mutants demonstrated the widely splayed neural folds characteristic of spina bifida (Fig. 1G). Some *hitchhiker* fetuses exhibited a closed thoracolumbar neural tube but which was not enclosed within the bony vertebral column, either with or without lumbosacral spina bifida (Fig. 1I, compare with H), and this phenotype may be similar to the human defect of spina bifida occulta. Histological sections through E15.5 *hitchhiker* embryos with this phenotype confirmed the presence of a closed neural tube, but with a grossly enlarged spinal canal, a thin dorsal covering of surface ectoderm and flanked by the widely spaced cartilage primordia of the developing vertebrae (Fig. 1K, compare with J). Skeletal preparations of E18.5 fetuses with the exposed thoracolumbar neural tube (without spina bifida) clearly demonstrated splayed vertebrae in the thoracolumbar region (Fig. 1M, compare with L) and also revealed abnormal morphology of the ribs, commonly with bifurcations (Fig. 1O, compare with N). Since the ribs, vertebrae and dorsal dermis all derive from the somites (57), the observation of defects in these tissues might suggest possible abnormalities in somitic development in *hitchhiker*.

The *hitchhiker* line was named after a limb defect, in which the first digit exhibited an abnormal backward-bent position (Fig. 1P and Q). More often, mutants exhibited preaxial polydactyly on both fore- and hindlimbs, with six or occasionally seven digits (Fig. 1R and S). Skeletal preparations revealed a spectrum of defects, including apparent bifurcation or duplication of digit one, or full or partial duplication of another digit (Fig. 1T–W and data not shown).

The observed phenotypes were all partially penetrant and occurred in any combination: 68% of fetuses exhibited polydactyly ( $n = 73$ ;  $\geq$ E12.5); 65%, spina bifida aperta; 58%, oedema; 37%, exencephaly; 8%, spina bifida occulta and 5% no detectable defect ( $n = 126$ ,  $\geq$ E10.5). Approximately

6% of heterozygotes exhibited a mild head misshaping or slight caudal oedema.

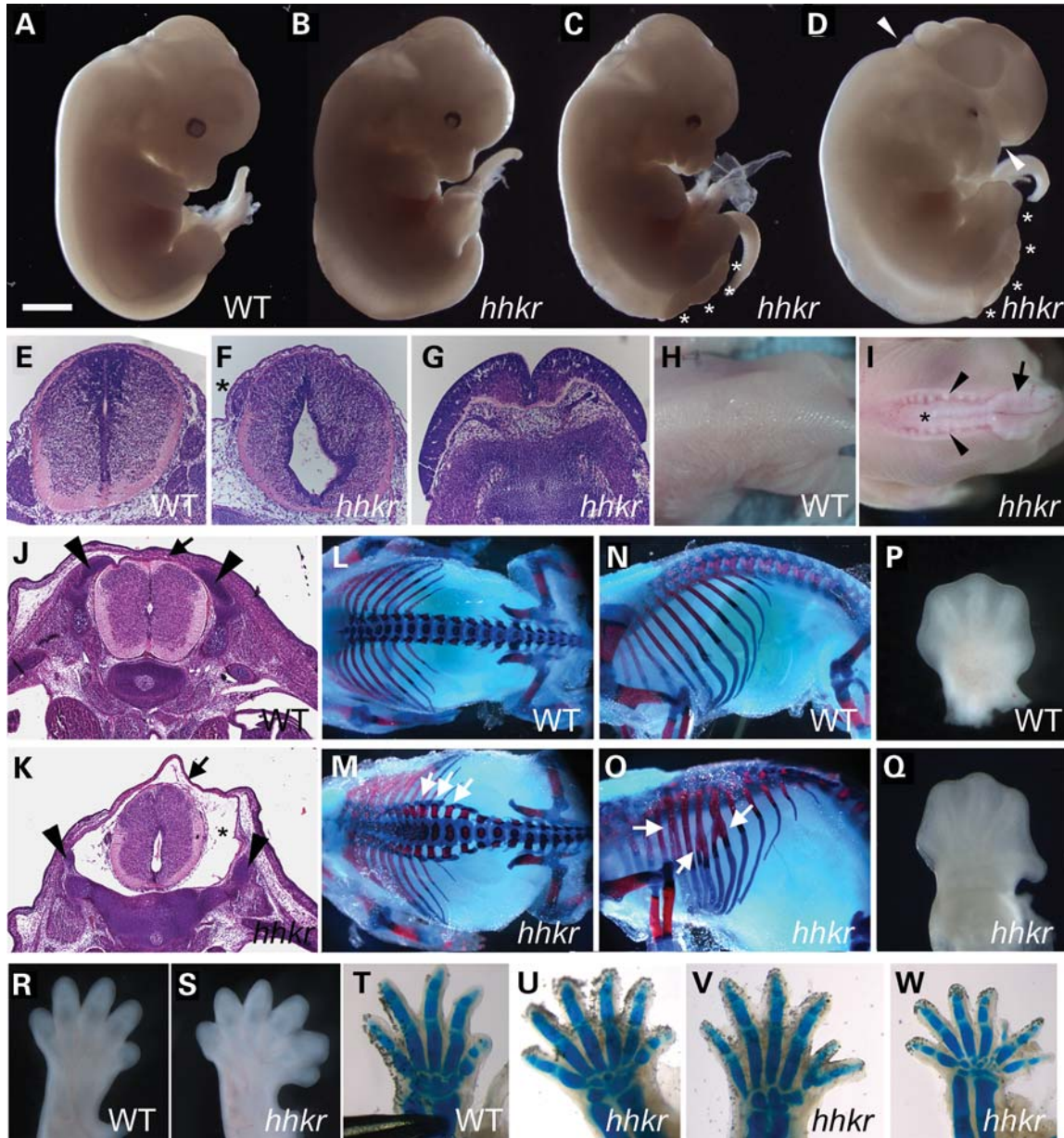
### *hitchhiker* carries a splice site mutation in *Tulp3* causing a strongly hypomorphic function

Genetic mapping using a genome-wide marker panel and nine affected individuals revealed linkage of *hitchhiker* to Chromosome 6 (Fig. 2A), and analysis with additional markers and individuals refined the interval to 12 Mb (Fig. 2B). Within this region lies *Tulp3*; since a targeted allele of *Tulp3* exhibits neural tube defects similar to *hitchhiker* (58), we examined *Tulp3* in our mutant. RT-PCR analysis revealed a reduced size for one *Tulp3* amplicon (Fig. 2C), and sequencing revealed a deletion of 52 bp corresponding precisely to exon 2 (data not shown). Genomic DNA analysis identified a single nucleotide substitution at the highly invariant position 2 of the splice donor site immediately 3' to exon 2 (Fig. 2D and E) (59).

Deletion of exon 2 creates a frameshift that is predicted to cause premature truncation of the protein (Fig. 2F) and a null phenotype. Western blots confirmed a dramatic reduction in Tulp3 protein in *hitchhiker* homozygotes, although long exposures detected Tulp3 at  $\sim$ 4% of the wild-type level (Fig. 2G and H), suggesting that *hitchhiker* is a strong hypomorph of *Tulp3* rather than a complete null. Tulp3 protein was reduced to 60% wild-type level in heterozygotes (Fig. 2G and H). To further verify that *Tulp3* mutation is responsible for the *hitchhiker* phenotype, we intercrossed *hitchhiker* heterozygotes with mice heterozygous for the *Tulp3*-targeted allele, *Tulp3*<sup>tm1Jng</sup> (58), with both strains on a C3H background. This yielded embryos exhibiting the range of phenotypes displayed by either homozygous mutant (Fig. 2I and J;  $n = 8$ ), indicating a failure of complementation. The phenotype of both the targeted allele homozygotes and the double heterozygous embryos appears to be slightly more severe than that of *hitchhiker* homozygotes, with more extensive oedema and defects in eye development. Together, these data indicate that *hitchhiker* is a strong hypomorph of *Tulp3*.

### Neural tube defects in *hitchhiker* result from a failure of brain and PNP closure

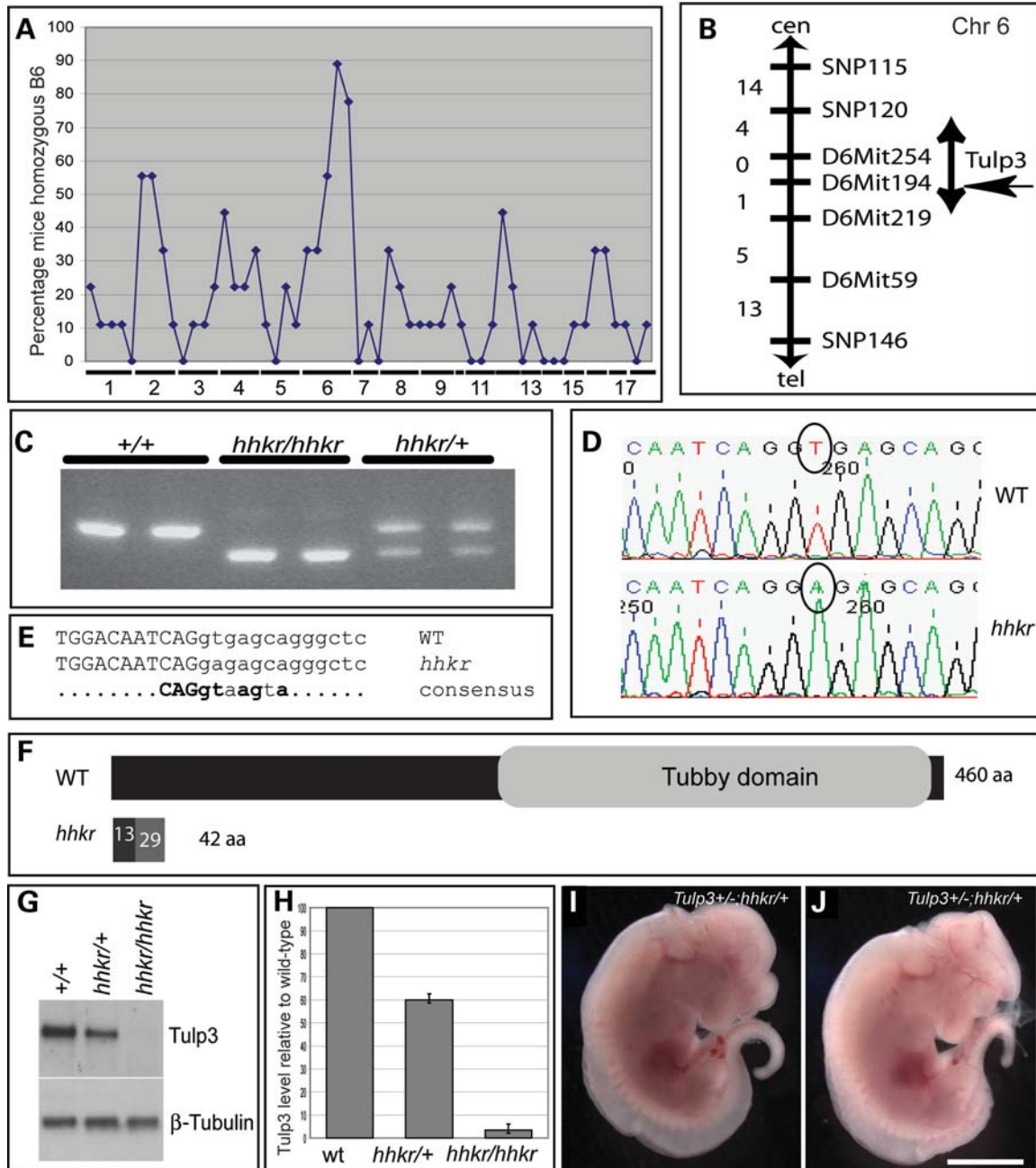
Following identification of the mutant gene, we conducted detailed morphological and histological examination of mutant embryos during the stages of neurulation in order to determine the cause of the neural tube defects. In the cranial region, mutant embryos exhibited normal *de novo* closure at the future cervical region (Closure 1) and the rostral extent of the forebrain (Closure 3) (data not shown). In contrast, the occurrence of Closure 2, at the forebrain–midbrain boundary, was delayed or failed: 22% of *hitchhiker* embryos exhibited failure of Closure 2 even at the 24–27 somite stage, while a further 15% had achieved Closure 2 but retained an open hindbrain neuropore at this stage, in contrast to the complete head closure seen in wild-type embryos by the 21 somite stage (Fig. 3A and B). The exencephalic phenotype observed in 37% of fetuses at later stages, therefore, corresponds precisely to the earlier failure of either Closure 2 or hindbrain neuropore closure.



**Figure 1.** *hitchhiker* mutant mouse embryos exhibit defects in neural tube, skeleton and limb development. (A–D) E13.5 wild-type (A) and *hitchhiker* embryos (B–D) with a range of phenotypes, including oedema (B), oedema with spina bifida (C) and oedema with spina bifida and exencephaly (D). Asterisks mark region with spina bifida and exencephalic region is marked with arrowheads. (E–G) Transverse sections stained with haematoxylin and eosin (H&E) through spinal neural tissue in wild-type (E), mutant with caudal oedema and dorsally positioned dorsal root ganglia (asterisk) (F) and mutant with spina bifida (G). (H and I) Dorsal views of E18.5 wild-type (H) and *hitchhiker* mutant (I) fetuses demonstrating splayed vertebrae (arrowheads) and visible neural tube (asterisk) adjacent to a region of open spina bifida (arrow) in mutants. (J and K) H&E-stained sections through the thoracolumbar region of E15.5 wild-type (J) and *hitchhiker* (K) fetuses. The mutant exhibited the splayed vertebrae phenotype, and sections demonstrate a closed but abnormally shaped neural tube within a grossly enlarged spinal canal (asterisk), covered dorsally with a thin layer of surface ectoderm (arrow) and flanked by widely spaced vertebral primordia (arrowheads), compared with wild-type. (L–O) Skeletal preparations of E18.5 wild-type (L and N) and *hitchhiker* mutant (M and O) fetuses stained with alcian blue for cartilage and alizarin red for bone, demonstrating splayed vertebrae (M; arrows) and rib bifurcations (O; arrows) in mutants. (P and Q) Forelimbs from wild-type (P) and *hitchhiker* mutants (Q) at E13.5, demonstrating malposition of digit 1 in mutants. (R and S) E15.5 wild-type (R) and *hitchhiker* (S) limb buds showing preaxial polydactyly with six digits. (T–W) E16.5 skeletal preparations for wild-type forelimb (T) and *hitchhiker* mutants (U–W) demonstrating preaxial polydactyly with a range of phenotypes. Limbs are oriented with anterior to right. Scale bars: A–D and H: 2.5 mm; E–G: 0.5 mm; H and I: 5 mm; J and K: 1 mm; L–O: 4 mm; P and Q: 1 mm; R and S: 1.1 mm; T–W: 1.2 mm.

In the spinal region, embryos displayed a defect in PNP closure. From around the 17-somite stage, *hitchhiker* mutants exhibited a significantly enlarged PNP, compared with stage-matched wild-type and heterozygous littermates

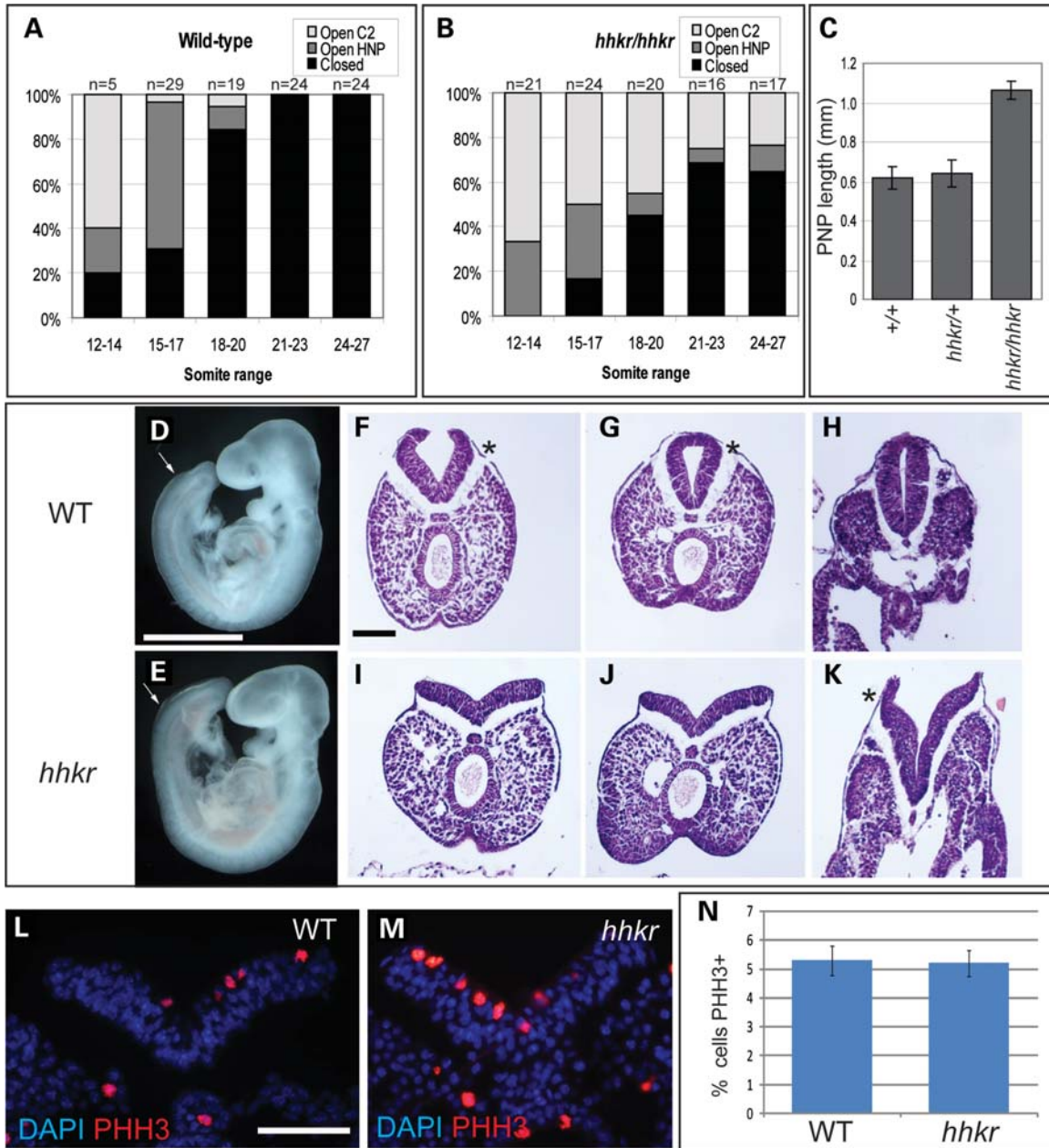
(Fig. 3C–E). In wild-types and heterozygotes, the PNP remained open in only 25% of embryos by the 27 somite stage ( $n = 4$  of 20), whereas 72% of *hitchhiker* mutants exhibited an open PNP at this stage ( $n = 13$  of 18). These figures



**Figure 2.** *hitchhiker* carries a splice site mutation in *Tulp3* resulting in severely reduced protein expression and function. (A) Genome-wide scan with 55 SNP markers on nine affected fetuses, showing percentage of mice typing as homozygous C57BL/6 at each marker; *hitchhiker* shows linkage to Chromosome 6. (B) Genetic map following analysis with 67 affected fetuses; numbers on left indicate genetic separation of markers (in centimorgan). *hitchhiker* maps to 12 Mb (5 cM) region between SNP120 and D6Mit219, a region containing *Tulp3*. (C) Gel of RT-PCR products of *Tulp3* amplified between exons 1 and 3, from wild-type, homozygous and heterozygous mutants, showing reduced amplicon size in *hhkr*. (D) Sequence trace from genomic DNA revealed single nucleotide substitution, T in wild-type to A in *hhkr* (circled). (E) Comparison with consensus splice sequence (59) reveals *hhkr* mutation occurs in the splice junction immediately 3' to exon 2. Uppercase: exon, lowercase: intronic sequence, bold: invariant. (F) Schematic representation of Tulp3 protein in wild-type and predicted truncation following exon 2 deletion in *hitchhiker*. (G) Western blot analysis on total cell lysates from whole E9.5 embryos demonstrating reduced Tulp3 protein in heterozygous embryos and almost complete absence in homozygous mutant embryos, relative to wild-type littermates. (H) Quantitation of Tulp3 expression in heterozygous and mutant embryos, normalized to  $\beta$ -tubulin, shown as a percentage of wild-type;  $n = 3$  blots, E9.5 or E11.5 whole embryos. (I and J) Complementation test between *hitchhiker* and the *Tulp3*-targeted allele, *Tulp3*<sup>im1.3mg</sup> (58) generated double-mutant embryos with exencephaly, spina bifida and oedema, E12.5. Scale bar: I and J: 2.5 mm.

correspond closely to the spina bifida observed in later embryos, both in the incidence (65%) and the region affected; at later stages, *hitchhiker* embryos exhibited an open neural

tube corresponding to the region between somites 24 and 34. More caudally, the spinal neural tube appeared normal, suggesting that secondary neurulation proceeds unhindered



**Figure 3.** Exencephaly and spina bifida in *hitchhiker* result from a failure of neural tube closure. (A and B) Proportion of embryos demonstrating completely closed cranial neural tube (black bar), initiation of head closure at Closure 2 with open hindbrain neuropore (dark grey) or open cranial region with failure of Closure 2 (light grey), within the somite ranges marked, in wild-type (A) and *hitchhiker* (B) embryos. Number of embryos within each group is shown at the top of each column. (C) Length of PNP in wild-type, heterozygote and *hitchhiker* mutant embryos, at 22–24 somite stage, in millimetre; mean  $\pm$  SEM. (D and E) Lateral view of E9.5 (20 somite) wild-type (D) and *hitchhiker* (E) embryos demonstrating enlarged PNP in mutants; anterior limit of PNP is marked with arrows. (F–K) Transverse H&E-stained sections through the caudal end of 24 somite wild-type (F–H) and *hitchhiker* (I–K) embryos, within the PNP (F, G, I and J), or closed neural tube (H). Wild-type embryos exhibit Mode 2 neurulation with clear DLHPs (asterisks), whereas *hitchhiker* embryos have reduced DLHPs that are apparent only adjacent to the closed neural tube (K, asterisk). (L and M) Immunostaining for phosphorylated histone H3 to examine cell proliferation in the caudal neuroepithelium of E9.5 wild-type (L) and *hitchhiker* (M) embryos. Red; phosphorylated histone H3; blue, DAPI. (N) Quantitative analysis of cell proliferation in the neuroepithelium, displayed as percentage of cells stained for phosphorylated histone H3, revealing no significant difference in percentage proliferation in *hitchhiker* caudal ends. Scale bar: D and E: 1 mm; F–M: 100  $\mu$ m.

in *hitchhiker*. Thus, in the spinal region as in the cranial region, we have demonstrated that neural tube defects occur as a primary defect in neural tube closure.

Previous detailed analysis of spinal neural tube closure has documented three ‘modes’ of closure, characterized by the

precise morphology of the neural folds (6). Histological sections through the PNP revealed abnormal morphology in *hitchhiker* embryos. At the 24 somite stage, wild-type and heterozygous embryos exhibited prominent DLHPs, characteristic of mode 2 neurulation (Fig. 3F–H). In contrast,

*hitchhiker* mutants exhibited a reduction in this dorso-lateral bending, with small DLHP seen only in the sections immediately adjacent to the closed neural tube and straight neural folds observed more caudally (Fig. 3I–K). The rates of cell proliferation and cell death in the neuroepithelium of the caudal neural tube at E9.5 were not significantly different between *hitchhiker* mutants and wild-type littermates (Fig. 3L–N and data not shown). These results suggest that the abnormal neural fold morphology may be causative for the delay in PNP closure in *hitchhiker* mutants.

#### ***hitchhiker* mutants exhibit increased Shh pathway activity during spinal neural tube closure**

To investigate the cause of the neuroepithelial defects, we examined the expression of *Shh*, a key molecule involved in the regulation of neural tube closure. Strong expression of *Shh* in the notochord represses DLHP formation during mode 1 neurulation, whereas weaker *Shh* activity during mode 2 neurulation permits DLHPs to form (13). By whole mount *in situ* hybridization, *Shh* expression appeared comparable between *hitchhiker* mutants and wild-type embryos at E9.5 (Fig. 4A and B), and transverse sections through the caudal region revealed similar intensities of expression in the notochord (Fig. 4C and D). At this stage and axial level, *Shh* expression is not yet induced in the wild-type floor plate, and *Shh* is similarly absent from the *hitchhiker* floor plate (Fig. 4C and D). To investigate the activity of the Shh pathway, as opposed to the expression of Shh ligand, we examined the expression of the transcriptional target of Shh signalling, *Ptch1*. Whole mount *in situ* hybridization revealed a dramatic up-regulation of *Ptch1* expression in the caudal region of the neural tube in *hitchhiker* mutants (Fig. 4E and F). Transverse sections through these embryos demonstrated an expanded domain of *Ptch1* expression throughout the DV extent of the caudal neural tube in *hitchhiker*, compared with wild-type (Fig. 4G and H). Quantitative analysis by real-time RT–PCR (qRT–PCR) detected a 1.65-fold increase in *Ptch1* expression in the caudal third of E9.5 embryos, compared with wild-type (Supplementary Material, Fig. S1A). In addition, a similar qRT–PCR analysis of *Gli1*, another transcriptional target of the Shh pathway, detected a 1.5-fold increase in *Gli1* expression in *hitchhiker* (Supplementary Material, Fig. S1B;  $P < 0.05$ ). These data suggest that the activity of the Shh pathway is increased in *hitchhiker* mutants, at E9.5, independently of a change in *Shh* expression. We suggest that this may inhibit the formation of DLHPs.

Further analysis at E10.5 revealed continued overexpression of *Ptch1* in *hitchhiker* mutants, compared with wild-type littermates. At this stage as at E9.5, the most dramatic up-regulation of *Ptch1* expression was observed in the caudal neural tube, but ectopic expression was also evident in the caudal somites (Fig. 4I–L). Although *Shh* expression looked normal at E9.5, by E10.5 *Shh* is ectopically expressed in *hitchhiker* mutants, in approximately one-third of the spinal cord anterior to the hindlimb bud (Fig. 4M and N; asterisks). Transverse sections through the lumbar neural tube demonstrated a dramatic increase in the *Shh* expression domain. *Shh* expression extends throughout the ventral half of

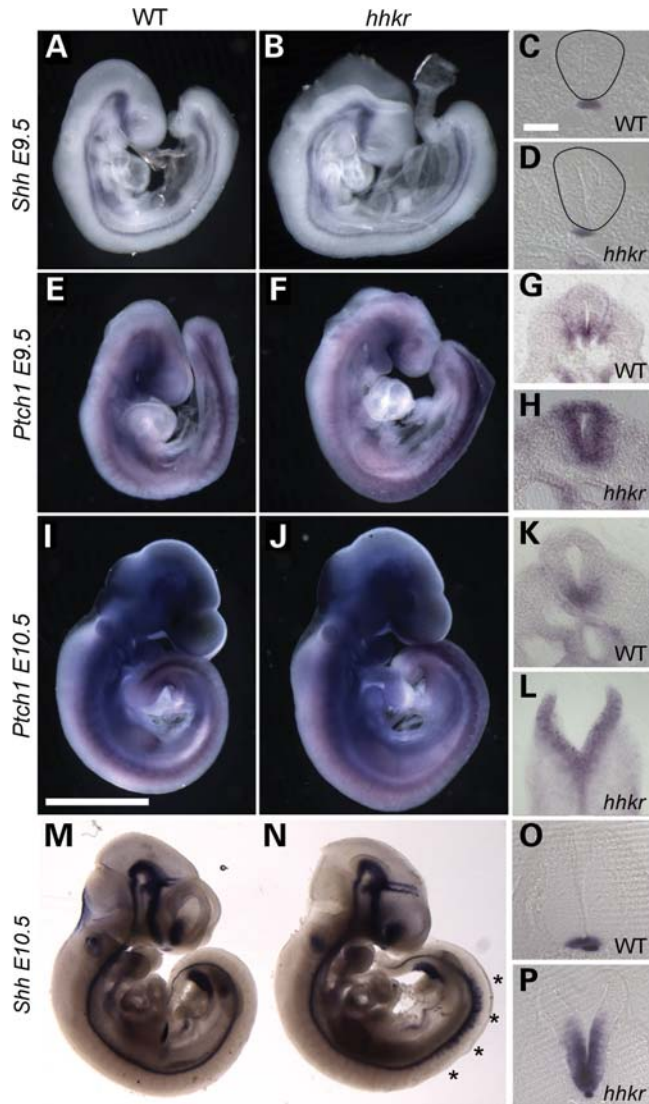
the neural tube in *hitchhiker* mutants, whereas it is restricted to the floor plate in wild-type embryos (Fig. 4O and P). The spatial and temporal relationship between the onset of *Ptch1* and *Shh* overexpression suggest that in *hitchhiker* mutants, activation of the *Shh* pathway occurs independently of increased Shh ligand, which then subsequently (and indirectly) induces the ectopic expression of *Shh* within the neural tube.

#### ***hitchhiker* mutants exhibit ventralization of the caudal spinal cord**

Activity of the Shh pathway is a key regulator of DV patterning within the neural tube, and the observed changes in *Ptch1* and *Shh* expression in *hitchhiker* mutants prompted us to examine the expression of markers of DV patterning at E10.5. As anticipated from the expansion of *Shh* mRNA expression, we detected dorsal expansion of both *Shh* and *Foxa2* protein expression, markers of floor plate, in the lumbar neural tube of *hitchhiker* mutants (Fig. 5A–D), although the region that exhibits the columnar morphology characteristic of floor plate appears to be unaltered. In addition, *Nkx2.2*, which marks the progenitors of the V3 ventral interneurons adjacent to the floor plate, and *Nkx6.1*, and *Olig2* which mark other ventral and ventro-lateral neurons, are all expanded dorsally in *hitchhiker* mutant embryos (Fig. 5E–J). *HB9/MNR2* and *Islet1/2*, markers of differentiating motoneurons, were also ectopically expressed in dorsally extended domains in *hitchhiker* mutants (Fig. 5K–N). Correspondingly, markers of dorso-lateral and dorsal neurons, *Pax6*, *Pax7* and *Msx*, exhibited smaller expression domains in *hitchhiker* mutants, with a dorsal shift in their ventral limit of expression (Fig. 5O–T). Patterning of the neural tube at more cranial spinal levels was largely unaffected in *hitchhiker* mutants (Supplementary Material, Fig. S2). Thus, our data reveal ventralization of the caudal neural tube in *hitchhiker* mutants, consistent with *Tulp3* acting as a negative regulator of the Shh pathway.

#### **Genetic analysis reveals Shh pathway activation in *hitchhiker* occurs independently of Shh ligand**

The expression changes observed at E9.5 suggested that up-regulation of the Shh pathway in *hitchhiker* mutants occurs independently of a change in *Shh* expression. We hypothesized that mutation of *Tulp3* leads to loss of repression of the Shh pathway, downstream of Shh ligand. To test this genetically, we intercrossed *hitchhiker* with the *Shh*-targeted null allele (60). Doubly heterozygous (*Shh*/+;*hhkr*/+) mice were viable and fertile, and intercrosses yielded E10.5 embryos with the expected Mendelian ratio of genotypes. Although *hhkr* homozygotes often exhibited exencephaly and caudal oedema (Fig. 6B), *Shh* null mutants were characterized by holoprosencephaly and reduced growth (Fig. 6C, compare with A), as described previously (60). Compound homozygous *Shh/Shh;hhkr/hhkr* embryos exhibited exencephaly, caudal oedema and spina bifida (Fig. 6D). These phenotypes are characteristic of *hitchhiker* and distinct from those of the *Shh* null, demonstrating that *hitchhiker* is epistatic to *Shh*.



**Figure 4.** *hitchhiker* mutants exhibit increased *Ptch1* expression prior to increased *Shh* expression. E9.5 (A–H) and E10.5 (I–P) embryos following wholemount *in situ* hybridization with riboprobes for *Shh* (A–D and M–P) or *Ptch1* (E–L). (A–D) At E9.5, *Shh* expression is comparable in wild-type (A) and *hitchhiker* (B) embryos, and transverse sections reveal expression is confined to the notochord (C and D). (E–H) *Ptch1* expression at E9.5 is increased in *hitchhiker* mutants (F) compared with wild-type (E), in the caudal half of the embryo. Transverse sections demonstrate up-regulated and expanded expression in the caudal neural tube of mutant (H) compared with wild-type (G). (I–L) At E10.5, *Ptch1* expression remains greatly enhanced at the caudal end of *hitchhiker* embryos (J and L) compared with wild-type (I and K). (M–P) At E10.5, ectopic *Shh* expression is evident in *hitchhiker* embryos along approximately one-third of the spinal cord, anterior to the hindlimb (N, asterisks, compare with M). Transverse sections reveal greatly expanded *Shh* expression domain within the caudal neural tube of *hitchhiker* (P) compared with wild-type (O). Scale bars: A, B, E, F, I, J, M and N: 1 mm; C, D, G, H, K, L, O, P: 100  $\mu$ m.

To investigate this further, we assessed the DV patterning in the caudal neural tube of double mutant embryos. Immunostaining on transverse sections through the caudal spinal cord of *Shh* mutants revealed a dorsalized phenotype, distinct from both the wild-type expression pattern (Fig. 6E–H) and

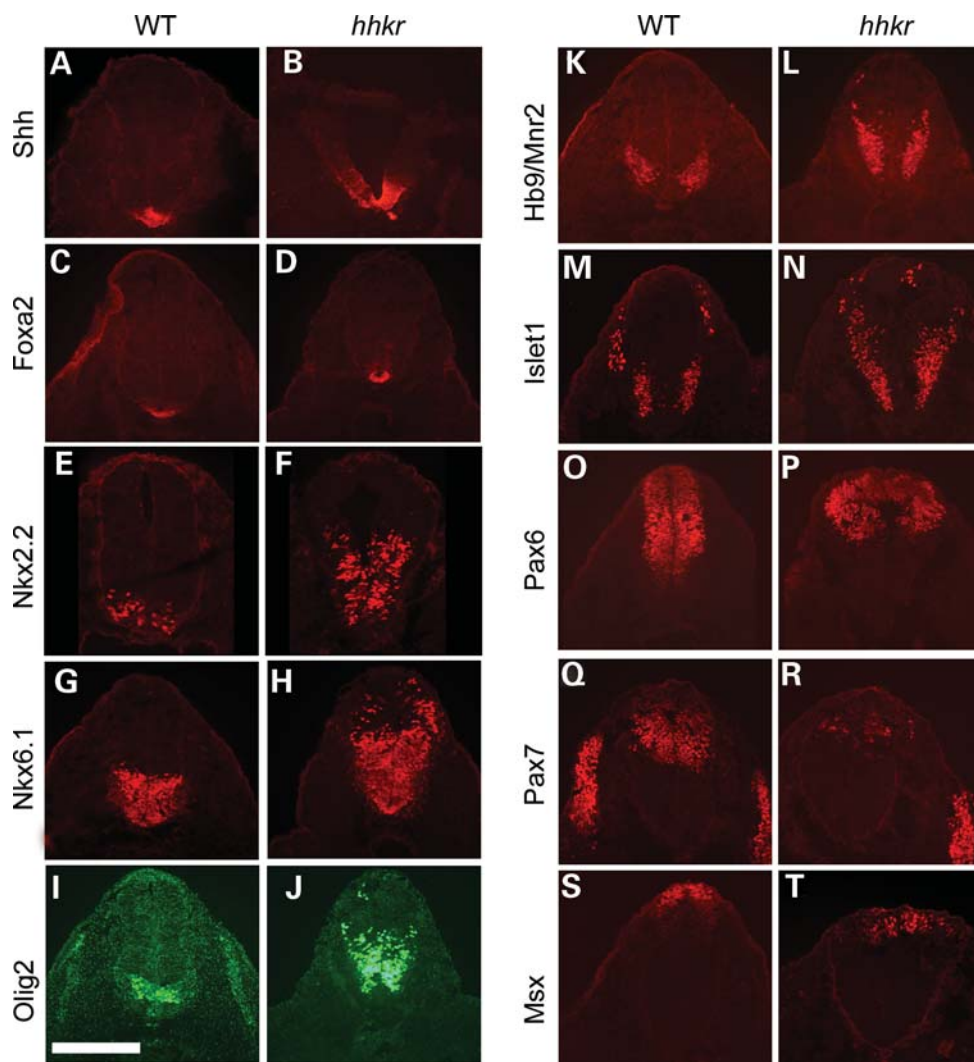
ventralized phenotype of *hhkr* embryos (Fig. 6I–L). In *Shh* mutants, *Nkx2.2*, *Nkx6.1*, *Olig2*, *HB9* and *Islet1* are absent while *Pax6*, *Pax7* and *Msx* are expanded ventrally (Fig. 6M–P and data not shown), as reported previously (45). In the *Shh/Shh;hhkr/hhkr* double mutants, *Nkx2.2*, *Nkx6.1*, *Olig2*, *HB9* and *Islet1* were expressed throughout a dorsally extended domain in double mutants, in a pattern similar to *hhkr* (Fig. 6Q–S and data not shown). The ventral limits of *Pax6*, *Pax7* and *Msx* expression domains were shifted dorsally (Fig. 6T and data not shown), similar to *hitchhiker*. Thus, doubly homozygous *Shh/Shh;hhkr/hhkr* embryos revealed ventralization of the neural tube similar to that in *hitchhiker* mutants, and dissimilar to the dorsalization seen in *Shh* mutants. In addition, analysis of *Ptch1* and *Gli1* expression demonstrated increased expression in *Shh/Shh;hhkr/hhkr* embryos, similar to that observed in *hhkr* (Supplementary Material, Fig. S3), and in striking contrast to the absence of *Ptch1* expression in *Shh* mutant embryos. These data further support our conclusion that *Tulp3* acts downstream of *Shh*, and indicates that mutation of *Tulp3* leads to activation of *Shh* pathway targets in the absence of *Shh* ligand.

#### *hitchhiker* is genetically epistatic to *Smoothened*

Since the hedgehog pathway can be activated by ligands other than *Shh*, we sought to test whether the pathway activation observed in *hitchhiker* mutants could be caused by increased expression of another ligand, such as Indian hedgehog or Desert hedgehog. Immunostaining with the 5E1 antibody that was raised against *Shh* (61), but which detects all hedgehog ligands (62), detected no observable signal in the notochord, neural tube or adjacent tissue of *Shh/Shh;hhkr/hhkr* double mutants, suggesting this is not the case (Fig. 7B, compare with A). For a more formal test, we intercrossed *hitchhiker* with the *Smo*-targeted null allele (63) to test for dependence on *Smoothened*, an obligate component of all hedgehog signalling. Doubly heterozygous (*Smo/+;hhkr/+*) mice were viable and fertile, and intercrosses yielded E10.5 embryos with the expected Mendelian ratios of genotypes. *Smo* null mutants were characterized by cyclopia, holoprosencephaly, retarded growth and pericardial oedema (Fig. 7E), as described previously (63). Compound homozygous *Smo/Smo;hhkr/hhkr* fetuses exhibited a phenotype more closely reminiscent of *hhkr* homozygotes than *Smo* null embryos, with expansion of the caudal spinal cord and a relatively normal head and growth, although pericardial oedema was also observed (Fig. 7F, compare with C–E).

Immunostaining on transverse sections through the caudal spinal cord of doubly homozygous (*Smo/Smo;hhkr/hhkr*) embryos revealed ventralization of the neural tube similar to that seen in *hitchhiker* mutant embryos, and distinct to the dorsalized phenotype observed in *Smo* mutants (Fig. 7O–R). *Nkx2.2* and *Islet1* were expressed throughout a dorsally extended domain (Fig. 7S and T), whereas the ventral limits of *Pax6*, *Pax7* and *Msx* expression domains were shifted dorsally (Fig. 7U and V and data not shown), similar to *hitchhiker* (Fig. 7K–N). These data reveal that *Tulp3* acts downstream of *Smo*, and indicates that mutation of *Tulp3* leads to activation of the *Shh* pathway independently of *Smo* receptor.





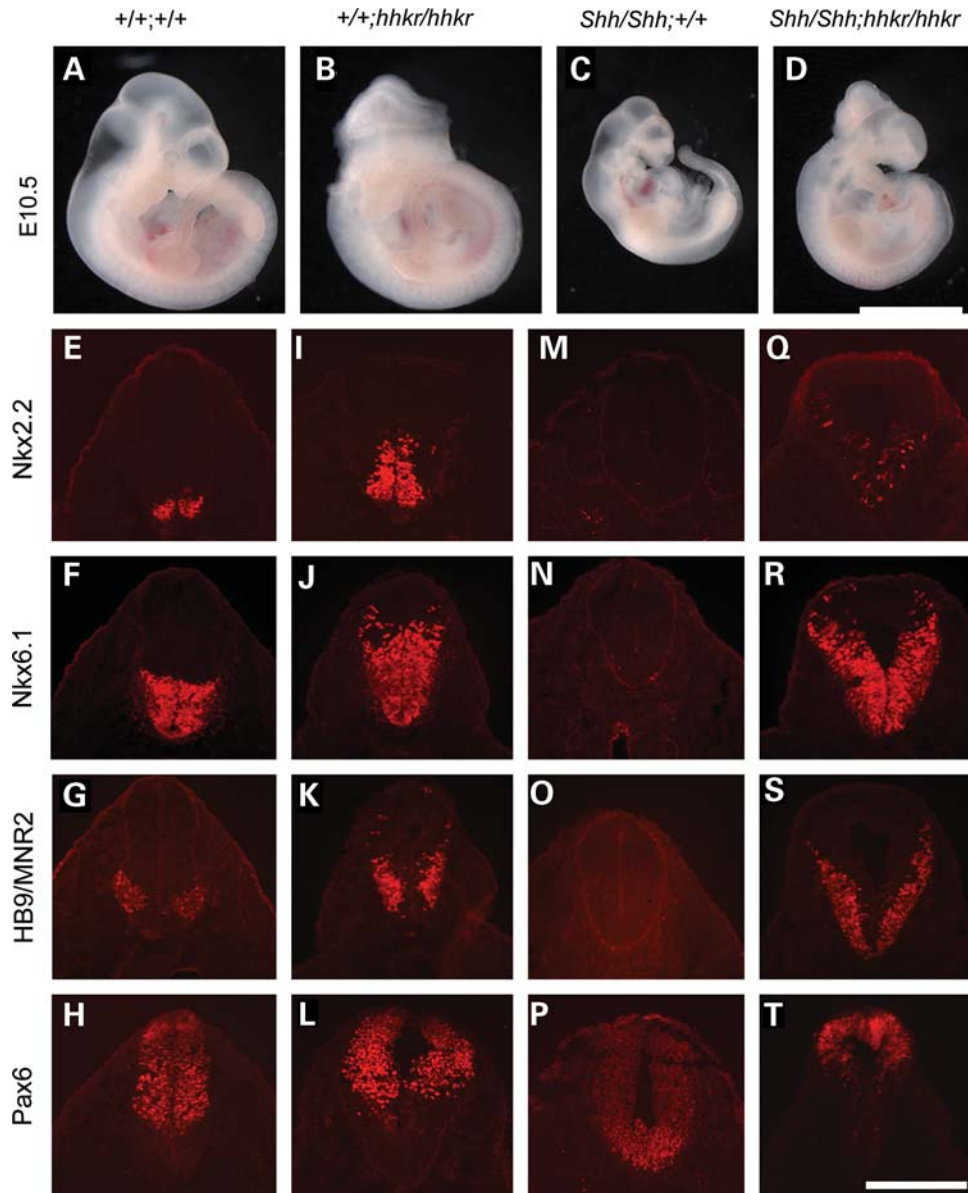
**Figure 5.** *hitchhiker* mutants exhibit ventralization of the caudal spinal cord. Immunostaining of transverse sections through the neural tube immediately anterior to the hindlimb bud with antibodies against Shh (A and B), Foxa2 (C and D), Nkx2.2 (E and F), Nkx6.1 (G and H), Olig2 (I and J), HB9/MNR2 (K and L), Islet1 (M and N), Pax6 (O and P), Pax7 (Q and R) and Msx (S and T), in wild-type (A, C, E, G, I, K, M, O, Q and S) and *hitchhiker* (B, D, F, H, J, L, N, P, R and T) embryos at E10.5. Ventral and ventro-lateral markers (Shh, Foxa2, Nkx2.2, Nkx6.1, Olig2, HB9/MNR2 and Islet1) exhibit expanded expression domains in *hhkr* while dorsal markers (Pax6, Pax7 and Msx) are reduced. Scale bar: 300  $\mu$ m.

### ***hitchhiker* limb buds exhibit ectopic anterior activation of Shh signalling and a genetic interaction with Shh and Gli3**

The *hitchhiker* mutants exhibit preaxial polydactyly, and we hypothesized that this might be caused by ectopic activation of the Shh pathway, similar to that observed in the neural tube. To address this, we examined the expression of targets of the Shh pathway in the developing limb buds. *Shh* expression appeared normal in *hitchhiker* limb buds, with a single domain of expression in the ZPA at E11.0 (Fig. 8A). In contrast, ectopic expression of *Ptch1* was observed in the anterior region of mutant limb buds at E11.0, whereas expression appeared normal in the posterior region (Fig. 8B). We also observed ectopic expression of *Gli1* in the anterior domain (Fig. 8C), although this was less robust than for *Ptch1*, and *Hoxd13* (Fig. 8D) another target of Shh pathway activation. Thus, *hitchhiker* mutants show normal expression patterns of markers in the posterior region of the

limb buds but appear to have ectopic activation of Shh pathway targets in the anterior region of the limb, in the absence of ectopic Shh ligand.

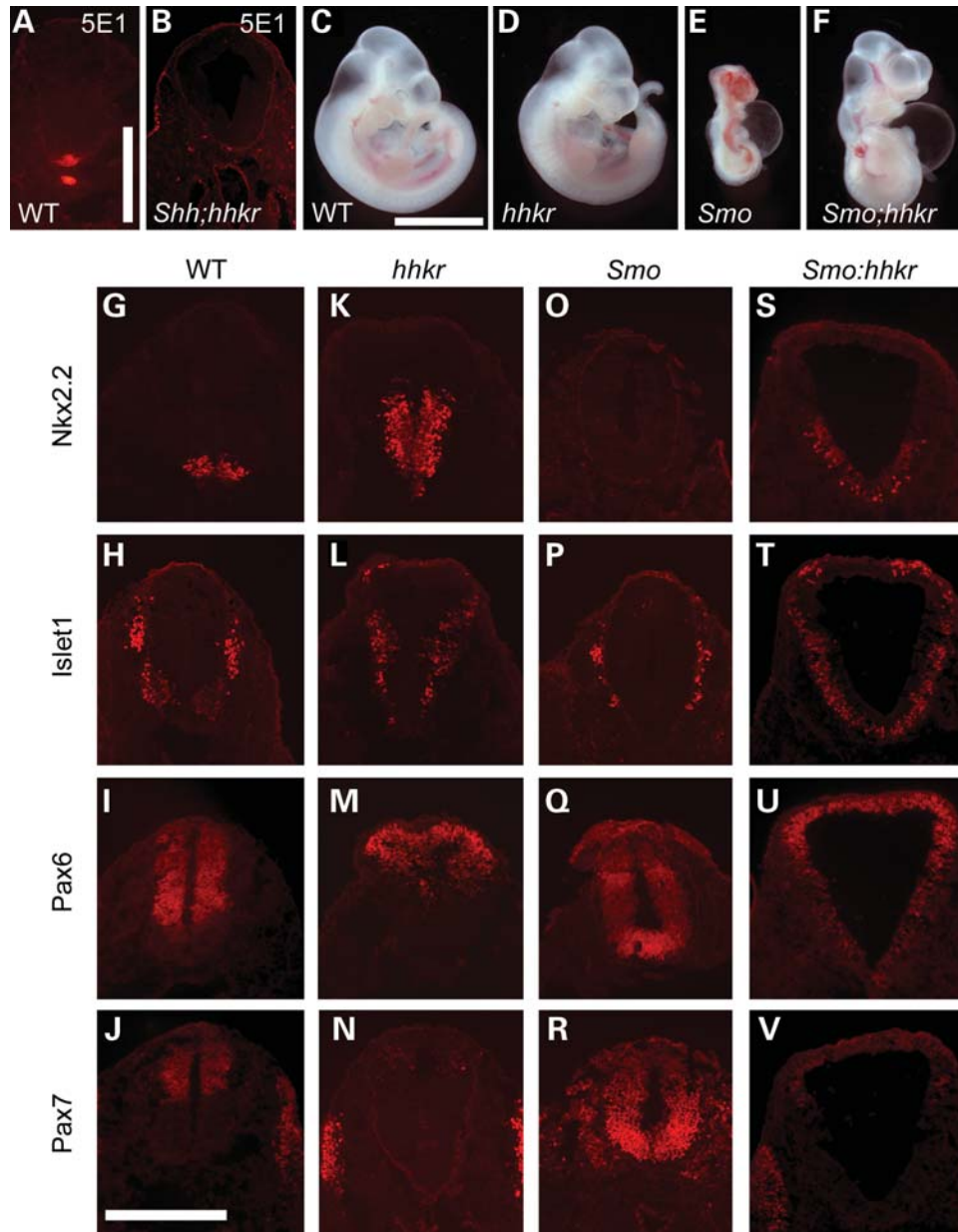
To determine whether *Tulp3* acts downstream of Shh in the limb (as well as in the neural tube), we examined limb morphology in *Shh/Shh;hhkr/hhkr* double mutants. Although *hitchhiker* mutants exhibited preaxial polydactyly with six digits on both forelimbs and hindlimbs (Fig. 9B, compare with A), *Shh* mutants are characterized by shortened and narrowed limbs bearing only a single digit (Fig. 9C), as described previously (64). The *Shh/Shh;hhkr/hhkr* double homozygotes reproducibly demonstrated an intermediate phenotype, with two digits on the forelimbs and four digits on hindlimbs (Fig. 9D). Although the limbs of double homozygotes do not resemble either single mutant, the mutation of *Tulp3* can partially rescue the severe limb defects of *Shh* mutants, consistent with *Tulp3* acting genetically downstream of Shh.



**Figure 6.** Genetic analysis of epistasis places *Tulp3* downstream of *Shh*. (A–D) E10.5 embryos generated from intercrossing *Shh*<sup>+/+</sup>;*hhkr*<sup>+/+</sup> doubly heterozygotes, demonstrating phenotypes of wild-type (A), *hitchhiker* (B), *Shh* (C) and *Shh/Shh*;*hhkr/hhkr* doubly homozygous (D) embryos. Double mutants grossly resemble *hitchhiker*. (E–T) Immunostaining of transverse sections through the neural tube immediately anterior to the hindlimb bud with antibodies against Nkx2.2 (E, I, M and Q), Nkx6.1 (F, J, N and R), HB9/MNR2 (G, K, O and S) and Pax6 (H, L, P and T) in wild-type (E–H), *hitchhiker* (I–L), *Shh* (M–P) and *Shh/Shh*;*hhkr/hhkr* doubly homozygous (Q–T) embryos. Immunostaining of double mutants is similar to *hitchhiker*. Scale bar: A–D: 2.5 mm; E–T: 300  $\mu$ m.

The downstream effectors of the *Shh* pathway are the three Gli-family transcription factors, with Gli3 playing the major role in limb patterning (30–32). To investigate the relationship between *Gli3* and *Tulp3* genetically, we intercrossed *hitchhiker* mutants with mice heterozygous for the *Gli3* null allele, *Gli3*<sup>XtJ</sup> (65). Doubly heterozygous *hhkr*<sup>+/+</sup>;*Gli3*<sup>+/+</sup> mice were viable and fertile, and intercrosses yielded embryos of all the expected genotypes except for surviving doubly homozygous mutants, at E15.5. Analysis of digit number in the different genotypic classes revealed an additive genetic interaction between

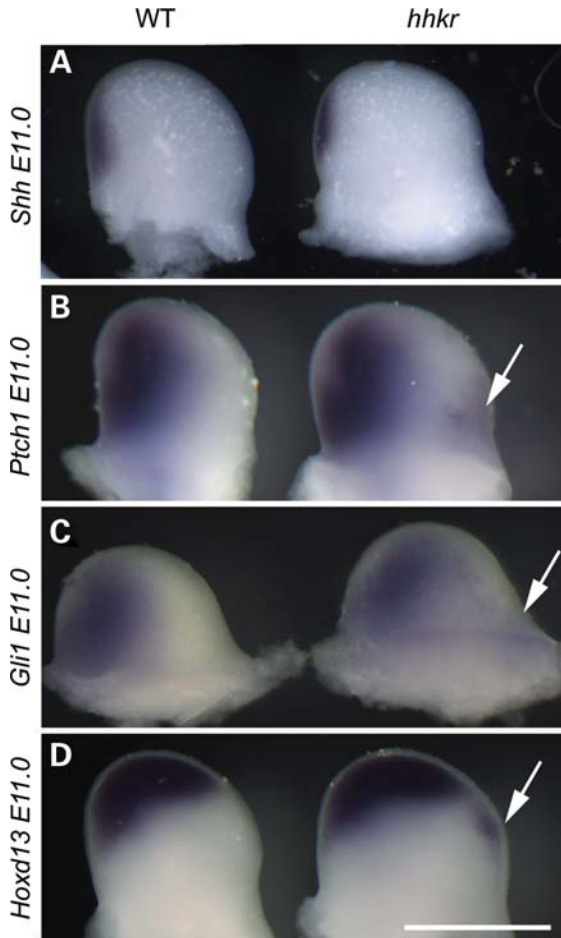
*hhkr* and *Gli3* (Table 1). Heterozygous *hitchhiker* mice exhibited normal limbs (Fig. 9F, compare with E), whereas *Gli3*<sup>+/+</sup> fetuses exhibited an extra preaxial digit (Fig. 9G). The *hhkr*<sup>+/+</sup>;*Gli3*<sup>+/+</sup> double heterozygotes appeared almost phenotypically indistinguishable to single *Gli3*<sup>+/+</sup> heterozygotes, with most limbs exhibiting a single extra digit (Fig. 9H). The *hhkr/hhkr* single homozygotes exhibited preaxial polydactyly with six or seven digits (Fig. 9I). Introduction of one *Gli3* mutant allele onto the homozygous *hhkr/hhkr* genotype appears to exacerbate the *hitchhiker* phenotype, with more severe polydactyly than



**Figure 7.** Ventralization of the spinal cord in *hitchhiker* occurs independently of *Smo*. (A and B) Immunostaining of wild-type and *Shh/Shh;hhkr/hhkr* double homozygous embryos with 5E1 antibody to detect all Hedgehog ligands. (C–F) E10.5 embryos generated from intercrossing *Smo/+;hhkr/+* double heterozygotes, demonstrating phenotypes of wild-type (C), *hitchhiker* (D), *Smo* (E) and *Smo/Smo;hhkr/hhkr* doubly homozygous (F) embryos. Double mutants more closely resemble *hitchhiker* than *Smo*. (G–V) Immunostaining of transverse sections through the neural tube immediately anterior to the hindlimb bud with antibodies against Nkx2.2 (G, K, O and S), Islet1 (H, L, P and T), Pax6 (I, M, Q and U) and Pax7 (J, N, R and V) in wild-type (G–J), *hitchhiker* (K–N), *Smo* (O–R) and *Smo/Smo;hhkr/hhkr* (S–V) doubly homozygous embryos. Scale bar: A and B: 500  $\mu$ m, C–F: 2.5 mm; G–V: 300  $\mu$ m.

observed in either *hhkr/hhkr* single mutants or *Gli3/+* heterozygotes, with often seven or eight digits (Fig. 9J; compare with I and G; Table 1). Homozygous *Gli3/Gli3* embryos exhibited severe polysyndactyly with often seven to nine digits (Fig. 9K), as described previously (65). Introduction of one *hhkr* mutant allele onto the *Gli3/Gli3* genotype (*hhkr/+;Gli3/Gli3*) appears to make no difference to the limb defect, which is almost indistinguishable to that observed in *Gli3/Gli3* single mutants, with seven to nine digits formed (Fig. 9L, compare with K; Table 1). Doubly

homozygous *hhkr/hhkr;Gli3/Gli3* embryos did not survive to E15.5. Attempts to collect embryos at E12.5 yielded only one double mutant in which limb morphology was abnormal, but retarded development meant digit number could not be reliably ascertained. The absence of doubly homozygous mutant data precludes a formal assessment of epistasis. However, the analysis of intermediate genotypes demonstrates that there is a genetic interaction between the two loci, and suggests that *Tulp3* and *Gli3* may act in concert during limb morphogenesis.



**Figure 8.** *hitchhiker* mutants exhibit ectopic expression of Shh pathway targets in the anterior region of the limb buds. (A–D) Forelimb buds from E11.0 wild-type (left) or *hitchhiker* (right) embryos following wholemount *in situ* hybridization with riboprobes for *Shh* (A), *Ptch1* (B), *Gli1* (C) and *Hoxd13* (D); anterior to right. *Shh* expression looks similar in wild-type and mutant limbs, whereas *Ptch1*, *Gli1* and *Hoxd13* show normal posterior expression but a small region of ectopic expression in the anterior domain (arrows). Scale bar: 1 mm.

### Gli2 and Gli3 abundance and Gli3 processing appear unaltered in *hitchhiker* mutants

Shh pathway activity is mediated by the stimulation of Gli activators and inhibition of Gli repressors. Since we have demonstrated that Tulp3 acts genetically downstream of Shh and Smo, and genetically interacts with Gli3, we sought to determine whether Gli2 and Gli3 expression are altered in *hitchhiker* mutants. By qRT-PCR, the expression of *Gli2* and *Gli3* was not detectably altered in the caudal end of E9.5 mutant embryos, compared with wild-type (Fig. 10A). Other mutants that affect Shh signalling demonstrate dramatic differences in Gli3 processing, as seen, for example, in the *ftm*, *IFT88/polaris* and *Dnchc2* mutants (66–69), prompting us to look for similar changes in the abundance of the Gli3 isoforms in *hitchhiker* mutants. Since the Gli3 Ab we used detects multiple bands, we performed western blots with protein extracts from *Gli3*<sup>X<sub>1</sub>/X<sub>1</sub></sup> mutants, which lack Gli3 protein (65,70), in order to confirm the correct band identity (Fig. 10B). Exper-

iments with forelimb buds isolated from E9.5 embryos, thus at a stage before changes in gene expression are apparent, revealed no observable change in the abundance of either the full length or cleaved Gli3 isoforms compared with wild-type or heterozygous littermates (Fig. 10B). In addition, Westerns performed with nuclear extracts from E10.5 embryos detected no change in the abundance of either full length or cleaved isoforms, in *hitchhiker* mutants, suggesting that Gli3 entry into the nucleus is not affected (Fig. 10C). From these data, we conclude that the action of Tulp3 does not appear to be mediated through regulation of Gli expression, processing or nuclear localization, although we cannot rule out the possibility that small differences might exist that would not be easily detectable due to experimental variability.

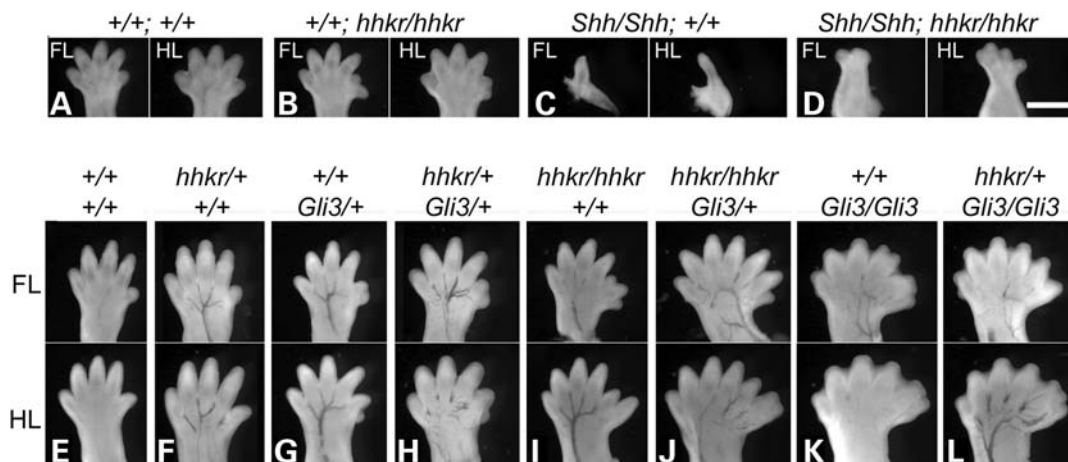
### Expression of *Rab23*, *Fkbp8*, *Thm1* and Sufu, activity of PKA and morphology of cilia are unaltered in *hitchhiker*

Since tubby family proteins may function as transcription factors (71), we hypothesized that Tulp3 might act by regulating the expression of another negative regulator. By qRT-PCR, we detected no significant change in *Rab23*, *Fkbp8* or *Thm1* (*Ttc21b*) expression in *hitchhiker* homozygotes compared with wild-type or heterozygous littermates, in the caudal thirds of E9.5 embryos, where the phenotype is apparent (Fig. 11A). Western blotting detected no change in Sufu expression in *hitchhiker* mutants compared with wild-type embryos at E9.5 (Fig. 11B). Assays for PKA activity in protein extracts from the caudal third of E9.5 embryos detected no significant difference in PKA activity between *hitchhiker* mutants and wild-types (Fig. 11C). We conclude that Tulp3 does not regulate the expression of *Rab23*, *Fkbp8*, *Thm1* or Sufu, nor the activity of PKA.

Recent evidence indicates that cilia play a major role in Shh pathway activity. Although most mutants affecting ciliary morphology or function lead to a decrease in Shh pathway function, such as seen in the *IFT52*, *IFT57*, *IFT88/polaris*, *IFT172* and *Kif3a* mutants (66–69,72,73), at least one mutant that affects retrograde intraflagellar transport results in activation of the Shh pathway (42). This prompted us to examine ciliary morphology in *hitchhiker* mutants. Scanning electron microscopy (SEM) on limb buds from E11.5 embryos revealed the presence of cilia on both wild-type and *hitchhiker* mutant cells, with no overt difference in their length or morphology (Fig. 11D–I). Immunofluorescence with anti-acetylated tubulin and anti-polaris on cultures of mouse embryonic fibroblasts or limb bud cells confirmed the presence of cilia in both *hitchhiker* mutant and wild-type cells with no detectable difference in polaris distribution (Fig. 11J–M and data not shown). Thus, no gross defect in ciliary morphology was observed in *hitchhiker* homozygotes.

## DISCUSSION

In this study, we have identified a novel ENU-induced strongly hypomorphic allele of Tulp3 which exhibits ventralization of the caudal neural tube, neural tube defects and polydactyly. We demonstrate ectopic expression of Shh pathway targets, independently of a change in Shh expression, and show



**Figure 9.** Genetic analysis shows *hitchhiker* acts downstream of *Shh* and interacts with *Gli3* in limb development. (A–D) E12.5 embryos generated from intercrossing *Shh/+;hhcr/+* double heterozygotes, demonstrating phenotypes of the forelimbs (FL) and hindlimbs (HL) of wild-type (A), *hitchhiker* (B), *Shh* (C) and *Shh/Shh;hhcr/hhcr* double mutant fetuses (D); anterior to right. (E–L) FL and HL of E15.5 embryos generated from intercrossing *hhcr/+;Gli3/+* double heterozygotes; positioned with anterior to the right. Wild-type (E) and *hhcr/+* (F) embryos have normal limbs with five digits; *Gli3/+* (G) and *hhcr/+;Gli3/+* (H) display six digits with a small additional preaxial digit; *hhcr/hhcr* (I) demonstrates six or seven digits; *hhcr/hhcr;Gli3/+* (J) exhibits seven digits; *Gli3/Gli3* (K) has polysyndactyly with seven or eight digits, plus an extra postaxial nubin; *hhcr/+;Gli3/Gli3* (L) exhibits polysyndactyly with seven or eight digits, plus an extra postaxial nubin. Scale bar: 1 mm.

genetically that *Tulp3* functions downstream of *Shh* and *Smo*. We also demonstrate that *Tulp3* genetically interacts with *Gli3* in regulating limb morphogenesis, but does not noticeably affect *Gli3* expression or processing. We propose that *Tulp3* functions to negatively regulate the *Shh* pathway in the caudal neural tube and limb buds.

### **Tulp3 functions as a negative regulator of *Shh* signalling in neural tube and limb**

The targeted allele of *Tulp3* has previously demonstrated a requirement for *Tulp3* in neural tube closure (58); however, *Tulp3* has not previously been associated with DV patterning of the neural tube nor with limb defects. In this study, we have demonstrated that *Tulp3* functions as a negative regulator of the *Shh* pathway in the developing neural tube and limbs. This conclusion is based on three sets of experimental results, which are the established criteria for identifying novel components of the *Shh* pathway (45,47,74). First, we observed increased/ectopic expression of genes activated by *Shh* signalling in the *hitchhiker* mutants, in the caudal spinal cord and anterior limb domains. *Shh* is necessary and sufficient for the expression of ventral neural tube cell types including floor plate, motor neurons, V2 and V3 interneurons (60,61,75), and markers of these neurons/progenitors were expressed in expanded domains in the caudal neural tube of *hitchhiker* mutants. Neural tube patterning involves both the ventralizing signal of *Shh* from the notochord and floor plate, and the dorsalizing signal of BMPs and Wnts from the roof plate and surface ectoderm, with modulation of patterning by FGFs and retinoic acid (14–18,76). Although antagonism of BMP signalling in neural explants leads to a dorsal-to-ventral switch in the neuronal progenitor cell identity induced by a set concentration of *Shh* (77), embryos in which BMP or Wnt signalling is abrogated exhibit disruption of the dorsal interneuron cell populations yet no defects in ventral

neuronal patterning (78–80) arguing against a role for *Tulp3* in activating BMP or Wnt signalling. Moreover, four of the overexpressed genes (*Foxa2*, *Nkx2.2*, *Ptch1* and *Gli1*) are direct transcriptional targets of *Shh* signalling (81,82). Secondly, genes that are normally repressed by *Shh* signalling are shifted dorsally in the *hitchhiker* neural tube. Thirdly, analysis of *Shh/Shh;hhcr/hhcr* and *Smo/Smo;hhcr/hhcr* double mutants shows that the activation of *Shh* target genes in *hitchhiker* mutants is independent of both *Shh* and *Smo*, and that substantial rescue of neural tube patterning and limb development occur in double mutants. Although BMP and Wnt signals can modulate the cellular response to *Shh*, the specification of ventral cell fates does not occur in the absence of hedgehog signalling, even without a ‘repressive’ effect of the dorsalizing signals (77,83). These defects observed in *hitchhiker* are established criteria for identifying negative regulators of the *Shh* pathway, as used in the characterization of *Fkbp8* and *Rab23* (45,47).

We have shown that the DV patterning of the neural tube in *Shh/Shh;hhcr/hhcr* and *Smo/Smo;hhcr/hhcr* double mutants is more closely similar to *hitchhiker* homozygotes than either *Shh* or *Smo* mutants, indicating that the DV markers are expressed independently of *Shh* and *Smo*. However, the expression patterns are not identical between double mutants and *hitchhiker* homozygotes. *Nkx2.2* and *Islet1* are expressed throughout the most ventral (floor plate) domain in double mutants, whereas they are excluded from this region in wild-type and *hitchhiker* homozygote. *Foxa2* is expressed in an expanded domain in *hitchhiker* homozygotes, but appears to be absent from double mutants (Supplementary Material, Fig. S3). In addition, low-intensity *Pax6* expression was evident throughout the ventral neural tube of *Smo/Smo;hhcr/hhcr* mutants, while it is excluded from this region in wild-types and *hitchhiker* homozygotes. These results indicate that the activation of *Foxa2* expression and the repression of *Nkx2.2*, *Islet1* and *Pax6* from the ventral domains remain

**Table 1.** Genetic interaction between *hitchhiker* and *Gli3*, determined by analysis of digit number

<i>hhkr</i> genotype	+/+	+/+	+/+	<i>hhkr</i> /+	<i>hhkr</i> /+	<i>hhkr</i> /+	<i>hhkr</i> / <i>hhkr</i>	<i>hhkr</i> / <i>hhkr</i>
<i>Gli3</i> genotype	+/+	<i>Gli3</i> /+	<i>Gli3</i> / <i>Gli3</i>	+/+	<i>Gli3</i> /+	<i>Gli3</i> / <i>Gli3</i>	+/+	<i>Gli3</i> /+
Number of fetuses examined	10	10	5	21	24	7	7	9
Average number of digits	<b>20.0</b>	<b>24.0</b>	<b>28.8</b>	<b>20.0</b>	<b>24.2</b>	<b>29.0</b>	<b>23.9</b>	<b>27.0</b>
Percentage of limbs with no extra digits	100	0	0	100	0	0	14	0
Percentage of limbs with one extra digit	0	100	20	0	96	14	75	36
Percentage of limbs with two extra digits	0	0	45	0	2	50	11	53
Percentage of limbs with three extra digits	0	0	25	0	2	32	0	11
Percentage of limbs with four extra digits	0	0	10	0	0	4	0	0

dependent on functional Shh and Smo activity. *Foxa2* induction and *Nkx2.2*, *Islet1* and *Pax6* repression require very high levels of Shh activity (84–86), suggesting that down-regulation of *Tulp3* in *hitchhiker* mutants is not sufficient to induce high-level activation of the pathway. A similar situation seems to occur in the developing limbs, whereby the intermediate phenotype of double mutants suggests that, in the absence of Shh, mutation of *Tulp3* is not sufficient to induce full activation of the Shh pathway. Intriguingly, *Shh*<sup>-/-</sup>;*Gli3*<sup>+/+</sup> mice typically form limbs with two to four digits (31), similar to the phenotype observed in *Shh*/*Shh*;*hhkr*/*hhkr* double homozygotes. Although *Tulp3* mutation can lead to activation of some markers in a Shh-independent manner, the level of Shh pathway activation does not reach the highest levels that normally occur and, therefore, some residual Shh-dependent patterning remains.

Some regulators of Shh activity are also transcriptional targets of the Shh pathway. By western blotting, we found no substantial change in the abundance of *Tulp3* protein in *Shh*, *Smo* or *Gli3* mutants (data not shown), suggesting that *Tulp3* is not itself a transcriptional target of the Shh pathway. Involvement of *Tulp3* as a negative regulator of the Shh pathway suggests that down-regulation of *Tulp3* activity might be required for Shh signalling. *Tulp3* has several putative sites for phosphorylation, myristoylation and glycosylation (87). It is possible that regulation of *Tulp3* activity might occur through effects on post-translational modification.

### Possible functions of *Tulp3* in the Shh pathway

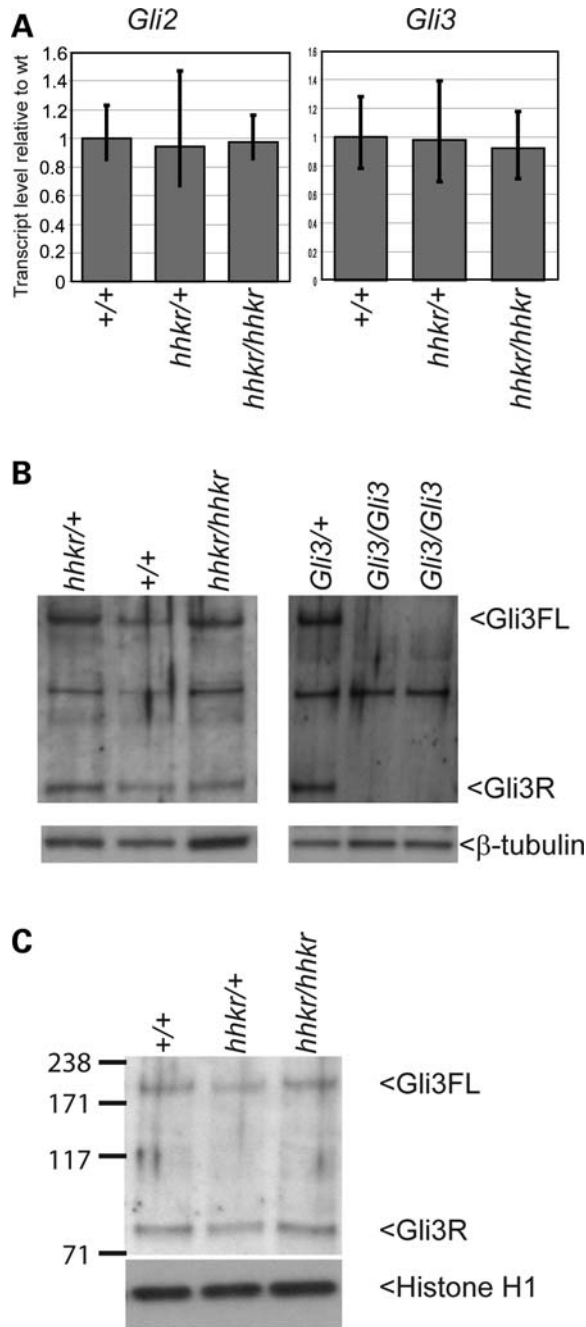
The tubby family consists of four proteins, Tubby and *Tulp1*–*3*, which are characterized by the C-terminal ‘tubby’ domain (88,89). Disruption of other tubby proteins has been associated with a range of phenotypes, including retinal and cochlear degeneration following progressive cell apoptosis, and maturity-onset obesity (90–94). No tubby member has previously been linked to Shh signalling.

We have demonstrated that *hitchhiker* mutants exhibit a phenotype consistent with *Tulp3* acting as a negative regulator of Shh signalling, and we have positioned *Tulp3* genetically downstream of Shh and Smo. Many regulators of the Shh pathway have recently been discovered, and several have been shown to act genetically downstream of Shh and Smo. These include proteins involved in the formation or maintenance of cilia and intraflagellar transport, such as *Arl13b*/*hennin*, *IFT88*/*polaris*/*flexo*, *IFT172*/*wimple*, *Kif3a*, *Rpgrip11*/*Fantom* and *Thm1* (38,42,66,68,69,72,95). Disruption of IFT

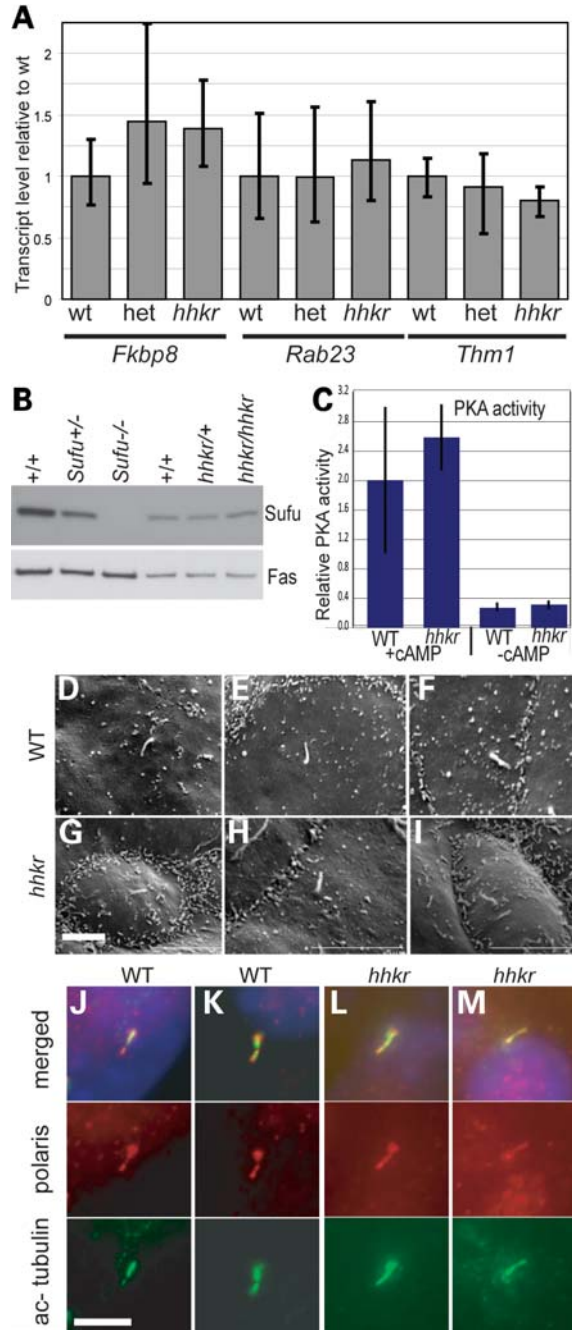
or cilia usually results in reduced Shh pathway activity in the neural tube, evidenced by the absence of Shh-dependent ventral cell types, as seen in mutants for *IFT88*/*polaris*/*flexo*, *IFT172*/*wimple*, *Kif3a*, *Rpgrip11*/*Fantom*, *Dync2h1*/*Dnchc2*, *IFT52*/*Ngd5*, *IFT57*/*hippi* and *Ofdl* (67,68,72,96,97). However, mutation of *Arl13b*/*hennin* results in constitutive activation of the Shh pathway at an intermediate level (95), whereas loss of *Thm1* causes increased Shh activity and expansion of ventral markers (42). The Gli-mediated transcriptional response to Shh and the effects of the negative regulators Sufu and PKA are abrogated in cells mutant for IFT components (98). The negative regulators *Rab23* and *Fkbp8* function genetically between Shh/Smo and *Gli2* (45,47,99,100), and their activity is dependent on functional IFT (66,100). *Rab23* is a member of the Rab-GTPase family of vesicular transport proteins and may regulate vesicular transport within (or dependent on) cilia, although the targets of *Rab23* trafficking have not been identified (55,99,101). *Fkbp8* is a member of the FK506-binding protein family (immunophilins), and may also play a role in membrane-associated proteasome function and trafficking (56,102).

The importance of cilia is further underlined by the finding that several Shh pathway components are localized within the cilia, including Smo, Ptch, Sufu and Gli proteins (38–41,67,98). Smo becomes enriched in cilia following treatment of cells with Shh (39,40,98), and this ciliary localization is necessary (39) but not sufficient (98) for pathway activation. *Tulp3* protein can be detected at the tips of cilia (37), and evidence from other members of the tubby family suggests that *Tulp3* might play a role in vesicular trafficking. *Tulp1* interacts with several proteins implicated in the formation and movement of vesicles including F-actin, dynamin-1, clathrin heavy chain, dynein intermediate chain and tubulin (103,104). *Tulp1* mutants exhibit accumulation of rhodopsin-containing vesicles in the interphotoreceptor matrix, consistent with loss of polarized trafficking between photoreceptor segments (91,105). However, examination of ciliary morphology in *hitchhiker* mutants has detected no overt ciliary defect, whereas immunofluorescence studies revealed no apparent difference in localization of *polaris*/*IFT88*. Thus, there is no evidence to suggest that ciliary morphology or transport is grossly disrupted in *hitchhiker* mutants, although more subtle structural or functional defects would not have been detected.

An alternative hypothesis for the role of *Tulp3* is as a transcriptional regulator. The tubby proteins are characterized by the C-terminal ‘tubby’ domain that binds to double-stranded DNA, while the N-terminal region can function as a



**Figure 10.** *Gli2* and *Gli3* expression and *Gli3* processing are not altered in *hitchhiker* mutants. (A) Quantitative RT-PCR analysis with primers for *Gli2* and *Gli3* on RNA from caudal ends of E9.5 wild-type, heterozygous and *hitchhiker* mutant embryos, showing no change in expression. (B) Western blot analysis on total cell lysates from E9.5 limb buds demonstrate no apparent change in abundance of the full length 190 kDa *Gli3* isoform (*Gli3FL*) nor the 83 kDa cleaved repressor isoform (*Gli3R*), in *hitchhiker* embryos compared with wild-type or heterozygous littermates. Western blots with extracts from *Gli3*/+ and *Gli3/Gli3* mutants confirm correct *Gli3* band identify.  $\beta$ -Tubulin was used as a loading control. (C) Western blots with nuclear extracts from E10.5 caudal ends demonstrate similar abundance of the 190 and 83 kDa *Gli3* isoforms in *hitchhiker* embryos compared with wild-type or heterozygous littermates; histone H1 was used as a loading control.



**Figure 11.** Evaluation of changes in downstream molecules and cilia structure. (A) Quantitative RT-PCR analysis with primers for *Fkbp8*, *Rab23* and *Thm1* (*Ttc21b*) on RNA from caudal ends of E9.5 wild-type, heterozygous and *hitchhiker* mutant embryos, showing no significant change in expression. (B) Western blot analysis on total cell lysates from E10.5 limb buds for *Sufu* expression revealed no change in *hitchhiker* mutants; fatty acid synthase (*Fas*) was used as a loading control. (C) PKA activity with or without cAMP addition, in protein extracts from E9.5 caudal ends of wild-type and *hitchhiker* embryos; no significant difference is detected. (D–I) SEM of E11.5 forelimb buds from wild-type (D–F) and *hitchhiker* mutant (G–I) embryos revealing cilia of normal length and morphology. (J–M) Immunostaining of MEFs from wild-type (J and K) and *hitchhiker* embryos (L and M) with anti-acetylated tubulin (green), anti-polaris (red) antibodies and DAPI (blue) demonstrate similar staining in mutants and wild-type cilia. Scale bars: 2  $\mu$ m.

transcriptional activator (71). The tubby proteins also encode nuclear localization signals and can be detected in the cell nucleus (87,93,106). Experiments suggest tubby proteins translocate from the plasma membrane into the nucleus in response to  $G\alpha_q$ -coupled receptor signalling (107). The phenotype of *hitchhiker* mutants might be consistent with transcriptional down-regulation of another negative regulator of the Shh pathway. Indeed, it is striking that *hitchhiker* bears many similarities with the *Rab23<sup>opb</sup>* mutant, including gross phenotypic effects (exencephaly, dorsal expansion of the caudal spine, small and dorsally misplaced dorsal root ganglia, polydactyly, rib bifurcations and splayed vertebrae) (108,109), similar patterning defects in the caudal neural tube (45,99,110) and positioning genetically downstream of *Shh* and *Smo* (45,99). The disruption of *Fkbp8* or partial loss of function of PKA (*Pka-plof*) also yield partial ventralization of the neural tube, similar to that seen in *hitchhiker* (46,47,100). However, we have shown that expression of *Rab23*, *Fkbp8*, *Thm1*, *Gli3* and *Sufu* and activity of PKA are not detectably altered in *hitchhiker* mutants. Thus, although we cannot rule out a transcriptional role for *Tulp3* on some other yet undiscovered negative regulator, we have excluded transcriptional modulation of the known negative modulators of Shh signalling.

Modulation of Gli protein activity is the downstream consequence of the Shh pathway, since the presence of Shh ligand acts to inhibit cleavage of Gli3 (30) and suppress the processing and degradation of Gli2 (111). Many mutants that exhibit down-regulation of the Shh pathway show impaired Gli3 processing, with a decrease in the amount of Gli3 repressor relative to the amount of full-length Gli3 protein, as seen in mutants for *Dync2h1/Dnchc2*, *IFT88/polaris*, *IFT172/wimble*, *Kif3a*, *Rpgrip11/fantom* and the chicken *talpid<sup>3</sup>* mutant (38,67–69,72,73,112). Gli3 processing is also disrupted in the *Thm1* mutant, which has activation of Shh signalling (42). The negative regulator *Sufu* binds to the Gli proteins (113,114) and may act in part by sequestering Gli1 and Gli2 in the cytoplasm (49,53,114,115), a function that involves CRM1-dependent nuclear export of Gli1 but not Gli2 (54). We have shown that *Tulp3* genetically interacts with Gli3 in regulating limb development. However, we observed no change in the expression levels of Gli2 or Gli3, and we detected no substantial change in the abundance of the Gli3 isoforms in *hitchhiker*. This suggests that *Tulp3* function is not associated with Gli expression or processing. In addition, we observed no alteration in abundance of Gli3 within the nucleus, suggesting that *Tulp3* does not act to regulate entry into the nucleus.

It is possible that *Tulp3* may function as a co-factor in Gli activity or may affect the interaction of Gli with other regulators. *Sufu* can inhibit the transcriptional activator function of Gli1 and Gli2, independently of cytoplasmic sequestration (54). *Sufu* mutant MEFs exhibit constitutive Gli activity yet do not show accumulation of Gli1 in the nucleus (50). *Sufu* interacts with SAP18, Galectin3 and other nuclear proteins (116) as well as with Gli1 bound to DNA (53), and *Sufu* represses Gli-mediated transcription by recruiting the histone deacetylation machinery through interaction with SAP18 (117). Although we detected no change in *Sufu* expression level, it is possible that loss of *Tulp3* might affect the interaction of *Sufu* with Gli; recent data have revealed that

*Cdc211* can act as a pathway activator, through binding to *Sufu* and relieving its inhibition on Gli (118). Alternatively, *Tulp3* might act to regulate the expression or activity of other factors that regulate Gli protein function, such as MED12 (119), MIM/BEG4 (120), CBP (81), Ski (121), Dyrk1 (122), DYRK2 or MAP3K10 (123). The precise functional and molecular relationship between *Gli3* and *Tulp3* requires further analysis.

It is intriguing that the *hitchhiker* limb buds exhibit a localized domain of ectopic gene expression, at the anterior limit of the limb buds. This is distinct from the ectopic expression patterns observed in other mutants, in which there is an anterior expansion of the normal expression domains. Many other mutants with polydactyly have impaired processing of Gli3, and this may cause the limb defect. It is possible that *Tulp3* acts in regulation of specific targets of Gli3 function, which then subsequently results in the region-specific activation of other molecules. Microarray analysis of transcriptional changes in *Gli3* mutant limb buds identified a number of targets, including some that are expressed in localized anterior domains and which are down-regulated in *Gli3* mutants; these include *Pax9*, *Cdo* and *Id2* and novel transcript *BG074838* (124). We might hypothesize that *Tulp3* has a role in co-regulation of specific Gli3 targets, resulting in loss of an anterior-specific factor, rather than a general change in Gli3 effects in the limb.

#### ***hitchhiker* mutants exhibit spina bifida as a result of a failure of neural tube closure**

We have demonstrated that spina bifida in *hitchhiker* results from a primary failure of spinal neural tube closure. Closure of the PNP occurs with three ‘modes’ of closure, at different gestational ages and, therefore, at different axial levels, typified by the formation of MHPs (mode 1), DLHPs (mode 3) or both (mode 2) (6). The regulation of DLHP formation is modulated by Shh expression, with strong Shh expression from the notochord during mode 1 neurulation inhibiting DLHPs (12), through the noggin-mediated antagonism of Bmp signalling (13). Formation of DLHPs is critical for spinal closure, and failure to form DLHPs results in severe spina bifida as seen in the *Zic2* mutant (13). In *hitchhiker* mutants, the formation of DLHPs appears to be partially suppressed, although they are not inhibited completely, and correspondingly, the spina bifida in *hitchhiker* is less severe than observed in *Zic2* mutants. We have demonstrated increased activation of the Shh pathway at the caudal end of *hitchhiker* mutants, and this would be anticipated to repress DLHP formation. Indeed, the loss of other negative regulators of the Shh pathway can also result in spina bifida, including *Sufu*, *Rab23*, *PKA* and *Ptch1* (45,46,48,109). However, we also observed defects in DV patterning of the neuroepithelium at E9.5 (data not shown). It is likely that PNP closure is inhibited by a combination of these effects.

#### **Regional differences in the role of *Tulp3***

Our data reveal a role for *Tulp3* in DV patterning of the neural tube, with a more important role in caudal than cranial regions of the spinal cord. *Tulp3* is expressed ubiquitously (58),



suggesting that the AP difference is not attributable to differential expression. It is notable that the role of other molecules in DV patterning also varies along the AP axis. For example, mutation of *Rab23*, *Fkbp8* or PKA-deficiency affect only caudal spinal cord patterning (45–47,110), whereas the *Dync2h1/Dnchc2* mutant exhibits a more severe patterning disruption in rostral than caudal regions of the spinal cord (72) and retinoic acid modulates patterning in the rostral spinal cord of quail embryos (125). *Smo*<sup>-/-</sup>;*Gli3*<sup>-/-</sup> embryos show reduced rescue of the ventral markers more caudally, and this is taken to indicate an AP difference in the requirement for Hh/Gli3 interactions in the specification of similar cell identities (75). The molecular basis of the regional differences and the relationship between the pathway regulators has yet to be fully elucidated.

The data presented here led to similar conclusions as a study performed with the *Tulp3*-targeted allele (Norman *et al.*, submitted). However, there are minor variations between the phenotypes, such as the absence of *Foxa2* expression in the *Shh/Shh;hhkr/hhkr* double mutant, compared with expression of this floor plate marker in double mutants with the *Tulp3* null allele, and no significant change in proliferation in the caudal neural tube of *hitchhiker*, compared with decreased proliferation in the *Tulp3* knockout. These qualitative differences likely reflect the difference in abundance of *Tulp3* protein, in the two alleles, since the targeted allele is likely to be a true null, while we have demonstrated in *hitchhiker* that a small amount (~4%) of wild-type protein remains.

In conclusion, correct regulation of the activity of the Shh pathway is essential for normal development and continued health in adults. Misregulation during embryonic development generates a spectrum of congenital abnormalities, including holoprosencephaly, neural tube defects and polydactyly. The continued identification of new components of the Shh pathway, such as *Tulp3*, coupled with further research to understand their molecular function, holds significant promise for the diagnosis and treatment of human disease.

## MATERIALS AND METHODS

### Mice and embryos

*hitchhiker* was identified during a three-generation (G3) recessive mutagenesis screen (details to be described elsewhere). Briefly, C57BL/6 males were injected with ENU and outcrossed to C3H/HeH; male F1 offspring were mated to C3H/HeH and F2 females backcrossed to their father. G3 embryos were examined at E13.5 for developmental abnormalities. Affected fetuses were used for genetic mapping with a 55-marker genome-wide SNP panel (sequences available on request), and additional microsatellite markers. The mutant line was maintained by backcrossing to C3H/HeH and, following gene identification, genotyped for the mutation by pyrosequencing. The MGI accession ID for this allele is MGI:3662473. Mice heterozygous for *Shh*<sup>tm1Chg</sup> (60) or *Smo*<sup>tm1Amc</sup> (63) were maintained on 129S6/SvEv and genotyped by PCR (N. Warr, personal communication). Mice heterozygous for the *Tulp3*<sup>tm1Jng</sup>-targeted allele (58) on C3Heb/FeJ were obtained from J. Eggenschwiler (Princeton University). Mice heterozygous for *Gli3* deletion (*Gli3*<sup>XtJ/+</sup>) on a

mixed C57BL/6/C3H background were obtained from T. Theil (Edinburgh University) and genotyped as described (126). Animals were maintained following guidelines of the Medical Research Council and in accordance with the Animals (Scientific Procedures) Act, 1986.

Mice were maintained routinely on a 12 h light–dark cycle (dark from 19:00 to 07:00 h). Embryos were generated by overnight matings, with the day of finding a copulation plug designated as embryonic day (E) 0.5. Some mice were maintained on a reverse light–dark cycle (dark from 10:00 to 22:00 h), and litters from these animals were designated as E1.0 on the day of plugging. Embryos were dissected in PBS with 10% newborn calf serum and processed according to downstream application.

### Sequencing

RNA was extracted from E13.5 fetuses using GenElute (Sigma) and reverse transcribed with MMLV-RT (Invitrogen). DNA and cDNA were amplified with intron or exon-specific *Tulp3* primers (sequences available on request), purified using Qiaquick (Qiagen) then sequenced with BigDye reagent (ABI) and ABI3700.

### Histology and *in situ* hybridization

Histology and skeletal preparations used standard protocols. Wholemount *in situ* hybridization was performed essentially as described (127), using digoxigenin-labelled riboprobes for *Shh*, *Ptch1*, *Gli1*, and *Hoxd13*. At least three embryos of each genotype were analysed with each probe and processed under identical conditions. Embryos were photographed on a Leica MZ16 stereomicroscope and vibratome sectioned at 35–50 μm as described (128). Slide *in situ* hybridization was performed using 10 μm cryosections as described elsewhere (129), with digoxigenin-labelled riboprobes for *Ptch1* and *Gli1*.

### Protein extraction, western blots and PKA assay

Total cell lysates were generated in RIPA buffer (PBS with 1% Nonidet P40, 0.5% sodium deoxycholate, 0.1% SDS). Nuclear fractions were generated using the Proteoextract Subcellular Fractionation kit (Calbiochem). Proteins were quantitated using the DC assay (Biorad). Westerns used 1–5 μg protein per lane on 7% or 3–8% Tris-Acetate NuPAGE gels with SeeBluePlus2 or HiMark ladders (Invitrogen). Proteins were transferred onto Hybond ECL (GE Healthcare) and detected with antibodies against *Tulp3* (1:300; gift from J. Eggenschwiler), *Gli3* (1:1000; Santa Cruz sc-20688), *Sufu* (1:200; sc-10933),  $\beta$ -tubulin (1:5000; sc9104), histone H1 (1:1000, sc-8030) or fatty acid synthase (1:1000; sc-55580) with HRP-conjugated secondary antibodies (1:12 000; DAKO) and detection with ECL Advance (GE Healthcare). Extracts from *Gli3*<sup>XtJ/XtJ</sup> (65) and *Sufu*<sup>-/-</sup> (50) mutants were used to verify *Gli3* and *Sufu* band identity, respectively, and the antibodies used have been published previously (70,130). Bands were quantitated from scanned images using Adobe Photoshop CS2; data are expressed as means  $\pm$  standard errors, and significance tested using two-sample equal

variance, two-tailed distribution Student's *t*-test. PKA activity was measured on extracts from E9.5 caudal thirds using a non-radioactive assay (Calbiochem), following Bradford assay quantitation.

### SEM, cell culture and immunofluorescence

Embryos were processed for SEM as described (38) and imaged on an Hitachi S-530 scanning electron microscope. Mouse embryonic fibroblast primary cultures were immunostained for N-acetylated tubulin (1:10 000; Sigma) or Polaris (1:1000; gift from B. Yoder) as described (38). Immunofluorescence on 10  $\mu$ m cryosections was performed as described (131), using antibodies against Nkx2.2, Shh, Islet1, Msx1/2, Foxa2, HB9/MNR2, Nkx6.1, Pax6 and Pax7 (1:10; Developmental Studies Hybridoma Bank) and Olig2 (1:500, Abcam ab33427). Cell proliferation rates were determined by staining with antibody against phospho-Histone H3 (pHH3, Upstate Cell Signalling). Primary antibody staining was detected with appropriate secondary antibodies conjugated with AlexaFluor-488 or AlexaFluor-594 (1:250 dilution; Invitrogen), mounted in Vectashield with DAPI (Vector Laboratories) and imaged with a Zeiss Axiophot microscope. Quantitation of proliferation rates was performed by counting pHH3+ cells and total (DAPI+) cells within the caudal neuroepithelium of E9.5 (17–19 somite) embryos; the mitotic index was calculated as the average from 12 sections from each of six embryos of each genotype, with analysis from the region immediately posterior to the caudal extremity of the vitelline artery thus within the PNP.

### Quantitative real-time RT-PCR

Total RNA was extracted from caudal thirds of E9.5 embryos and DNase treated using RNeasy plus mini kit (Qiagen), quantified with a Nanodrop ND1000 and the integrity assessed by gel electrophoresis. Double-stranded cDNA was then synthesized from 1  $\mu$ g RNA using High Capacity cDNA Archive kit (Applied Biosystems). Quantitative real-time PCR was performed with 10 ng cDNA using TaqMan<sup>®</sup> gene expression assays on a 7500 Fast Real-Time PCR System (Applied Biosystems). The assays used were: Mm00494645\_m1 (Gli1); Mm00487418\_m1 (Fkbp8); Mm01293116\_m1 (Gli2); Mm00492333\_m1 (Gli3); Mm01545399\_m1 (Hprt); Mm00436209 (Rab23); Mm00436026 (Ptch1) and Mm01270405 (Ttc21b). Samples were analysed in quadruplicate, using at least three embryos of each genotype, and calculations performed using the comparative C<sub>T</sub> method. The values were normalized to the endogenous reference gene (Hprt) and the transcript levels are presented as fold change relative to the wild-type sample in relative quantification (RQ) units. Error bars indicate the calculated maximum (RQ<sub>Max</sub>) and minimum (RQ<sub>Min</sub>) expression levels, with a 95% confidence level. Statistical significance was tested using two-sample equal variance, two-tailed distribution Student's *t*-test and is also represented as non-overlapping error bars if the samples are significantly different ( $P < 0.05$ ).

### SUPPLEMENTARY MATERIAL

Supplementary Material is available at *HMG* online.

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