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SOM Text

Figs. S1 to S3

Tables S1 to S5

References (28–32)

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Evolutionarily Assembled cis-Regulatory Module at a Human Ciliopathy Locus

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Neighboring genes are often coordinately expressed within cis-regulatory modules, but evidence that nonparalogous genes share functions in mammals is lacking. Here, we report that mutation of either *TMEM138* or *TMEM216* causes a phenotypically indistinguishable human ciliopathy, Joubert syndrome. Despite a lack of sequence homology, the genes are aligned in a head-to-tail configuration and joined by chromosomal rearrangement at the amphibian-to-reptile evolutionary transition. Expression of the two genes is mediated by a conserved regulatory element in the noncoding intergenic region. Coordinated expression is important for their interdependent cellular role in vesicular transport to primary cilia. Hence, during vertebrate evolution of genes involved in ciliogenesis, nonparalogous genes were arranged to a functional gene cluster with shared regulatory elements.

Cis-regulatory modules (CRMs) provide binding sites for transcription factors that regulate the expression of neighboring genes (1). Relatively little is known about the evolution of these regulatory elements, such as how CRMs arise or how the regulated genes cofunction, other than the rare instance such as *Hox* gene clusters evolved by gene duplication and the addition of regulatory elements to regulate body patterning (2).

Joubert syndrome (JBTS) is the most common neurodevelopmental disorder among the ciliopathy spectrum, which is thought to encompass disorders of structure or function of cellular primary (nonmotile) cilia (3). Affected JBTS patients show hypotonia, ataxia, abnormal eye movement, and a distinct mid-hindbrain malformation presenting the "molar tooth" sign on brain magnetic resonance images (MTI) (fig. S1A) (4). Mounting evidence suggests that primary cilia as cellular antennae sense a wide variety of signals, including Shh signaling, and play a crucial role in vertebrate development (5).

Recently, we reported deleterious mutations of the *transmembrane protein (TMEM) 216* gene, linking to the JBTS2 locus on chromosome 11, in about half of the 10 JBTS2-linked families (Fig. 1A) (6–8). However, the remaining half of the JBTS2 families (verified by the pathognomonic MTI) were phenotypically indistinguishable (displaying optic coloboma, retinal dysplasia, nephronophthisis, and occasional occipital encephalocele) but were negative for mutations in *TMEM216* (fig. S1, A and B, and table S1). Furthermore, fibroblasts from these latter patients contained intact *TMEM216* mRNA and protein expression (fig. S2, A and B), thereby suggesting another JBTS causative gene at the JBTS2 locus.

We thus performed resequencing of all known and predicted exonic and promoter genetic elements within the minimal 17-Mb candidate interval defined by *TMEM216* mutation-negative families (9). From these data, we identified four missense mutations and one splicing homozygous deleterious mutation in evolutionarily conserved

residues of the nearby *TMEM138* gene of unknown function, thus accounting for all JBTS2-linked families (Fig. 1A, fig. S2C, and table S1). All mutations segregated according to a single recessive disease mode and were not present in 400 ethnically matched chromosomes. Among missense mutations in transmembrane domains, *TMEM138* p.H96R led to unstable protein when transfected into heterologous cells (fig. S2D), suggesting loss of function as the disease mechanism.

Although both *TMEM* genes encode transmembrane proteins (Fig. 1A), neither the genes nor the proteins demonstrated sequence homology or shared any functional domains. Phylogenetic analysis showed that they represented two distinct protein families, which have evolved separately from invertebrates (figs. S4 and S5), excluding a gene-duplication event. In all higher

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vertebrates, the two genes aligned in a head-to-tail configuration, with a conserved ~23-kilobase pair noncoding intergenic interval (Fig. 1B). A synteny map of the genomic locus in a species with deposited reference sequence indicated that they were joined by an ancient chromosomal rearrangement at the amphibian-to-reptile evolutionary transition ~340 million years ago (Fig. 1B and fig. S3). Adjacent genes often exhibit correlated expression (10) and, thus, confer a selective advantage as a genetic module for a certain biological function (11). Therefore, the arrangement of two *TMEM* genes causing indistinguishable phenotypes prompted us to test whether these genes represent co-regulated and cofunctioning genes within a CRM.

To test for coordinated expression, we examined tissue-expression patterns of human *TMEM138* and *TMEM216* using the microarray

database and in situ hybridization of human embryos. We found tight coexpression values of human *TMEM138* and *TMEM216* across the major tissues, including the brain and kidneys (fig. S6B), and similar expression patterns in various tissues, including the kidneys, cerebellar buds, and telencephalon, at 4 to 8 gestational weeks (gw) of human embryos (Fig. 2A and fig. S6C). To test whether this coordinated expression was due to the adjacent localization, we compared mRNA levels in zebrafish versus mice, representing species before and after the gene rearrangement event. Using quantitative polymerase chain reaction (qPCR), we detected tightly coordinated expression levels in mice compared with those in zebrafish (correlation coefficient $r = 0.984$ versus 0.386) (Fig. 2B and fig. S6D), which suggests that *TMEM138* and *TMEM216* might share regulatory elements

(REs) within the ~23-kb intergenic region. We further examined several experimental features and found that regulatory factor X 4, a transcription factor regulating ciliary genes, binds a RE conserved in the noncoding intergenic region to mediate coordinated expressions of *TMEM138* and *TMEM216* (see SOM text and fig. S7).

The coordinated expression and indistinguishable ciliopathy phenotype caused by either mutation of *TMEM138* and *TMEM216* suggests possible functional relations at the protein level, possibly co-regulating ciliogenesis. We noticed short cilia and failure of ciliogenesis in *TMEM138*- (p.A126T) and *TMEM216*- (p.R73L) mutated fibroblasts, respectively (Fig. 3A), as well as defects of ciliogenesis after knockdown of *TMEM138* and *TMEM216* in IMCD3 cells (fig. S8), suggesting that both *TMEM138* and *TMEM216* are required for ciliogenesis. Immunostaining of endogenous *TMEM138* with a marker (Arl13b) of the ciliary axoneme and transition zone demonstrated closely adjacent localization with *TMEM216* at the base of cilia (γ -tubulin), as reported previously (Fig. 3B and fig. S9C) (6). We also noted prominent *TMEM138* and *TMEM216* adjacent but nonoverlapping vesicular staining around the base of cilia (Fig. 3B and fig. S9, C and D).

Earlier electron-microscopic findings suggested that vesicles transporting to the cilium play a crucial role for targeted delivery of membrane proteins from the Golgi apparatus (12, 13). It is noteworthy that *TMEM216* localized to post-Golgi vesicles along microtubules, as well as the Golgi apparatus surrounding the base of cilium, whereas *TMEM138* localized to adjacent but nonoverlapping distinct vesicles, shown by immunoelectron microscopic analysis (figs. S10, S11, and S12A). This observation suggests that both *TMEM138* and *TMEM216* might mark vesicles en route to the base of cilium, which is known to be crucial for ciliary assembly (14). Using time-lapse analysis with or without fluorescence recovery after photobleaching, we found that the net flux of *TMEM216*-tagged vesicular movements is toward the centrosome in dsRed-Centrin2 (centrosomal marker) expressing cells (fig. S12B and movies S1 and S2). *TMEM138*-tagged vesicles displayed tethered vesicular movement with *TMEM216*-tagged vesicles (fig. S12C and movie S3). This observation prompted us to ask whether the two distinct vesicles differentially carry known ciliary proteins essential for ciliary assembly. We further examined several experimental features showing that *TMEM138* and *TMEM216* mark two distinct but linked vesicle pools, each associated with unique cilia-targeted proteins, including CEP290 (see SOM text and fig. S13).

To understand the adjacency of the two vesicular pools, we tested whether either was required to move vesicles containing the other. Although *TMEM216* vesicular movement was not notably affected by disruption of *TMEM138* by small interfering RNA (siRNA), the knockdown of *TMEM216* disrupted the trafficking of

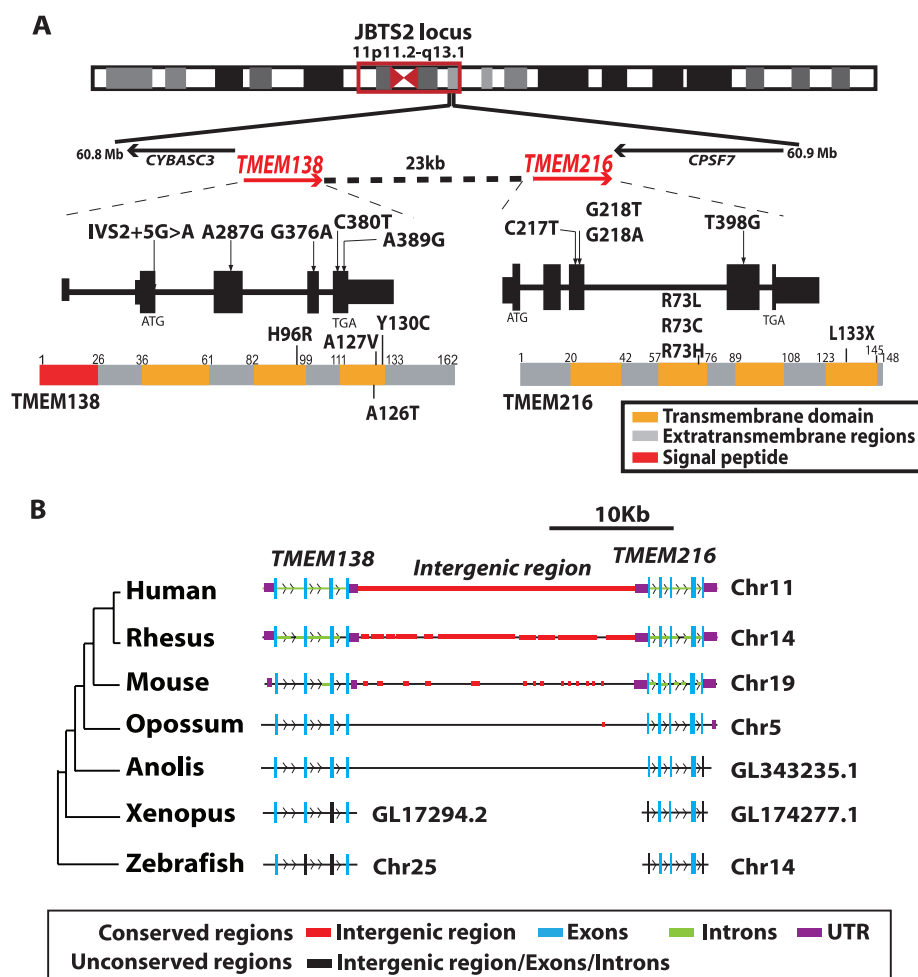


Fig. 1. Genetic heterogeneity at JBT52 locus and the evolutionary location of the *TMEM138* and *TMEM216* genes mutated in JBT52-linked families. (A) JBT52 locus (red box) on chromosome 11. *TMEM138* encoding a trispan membrane protein is aligned in a head-to-tail configuration with *TMEM216* encoding a tetraspan membrane protein. All missense and splicing mutations of *TMEM138* and *TMEM216* found in JBT52-linked families are indicated on the predicted transcript and protein. (B) Schematic synteny representation of *TMEM138*, *TMEM216*, and the intergenic region showing that the head-to-tail configuration of *TMEM138* and *TMEM216* on the same chromosome is conserved from reptiles to humans, but not in lower vertebrates, in which the two *TMEMs* are located on different chromosomes (table S2). Conserved regions have >50% sequence similarity to humans. UTR, untranslated region.

Fig. 2. Coordinated expression of adjacent *TMEM138* and *TMEM216* mediated by the noncoding intergenic region. **(A)** Similar expression patterns of *TMEM138* and *TMEM216* based on in situ hybridization at 8 gw in human embryonic tissues. *TMEM138* antisense (a and b), *TMEM216* antisense (c and d), and sense control probes (*TMEM138*) (e and f) are shown. *TMEM138* and *TMEM216* are strongly expressed in kidney, gonad (go), and adrenal gland (ad) as well as in the central nervous system, in particular in the cerebellar bud (cb), telencephalon (tel), rhombencephalon (rh), and cranial nerve ganglia such as trigeminal (V). **(B)** Real-time qPCR of *TMEM138* and *TMEM216* in selected tissues indicates tightly coordinated mRNA expressions in mouse tissues (having head-to-tail configuration), but not zebrafish (having two genes on different chromosomes). Housekeeping genes *36B4* (mouse) and *Rpl13a* (zebrafish) are used for normalization.

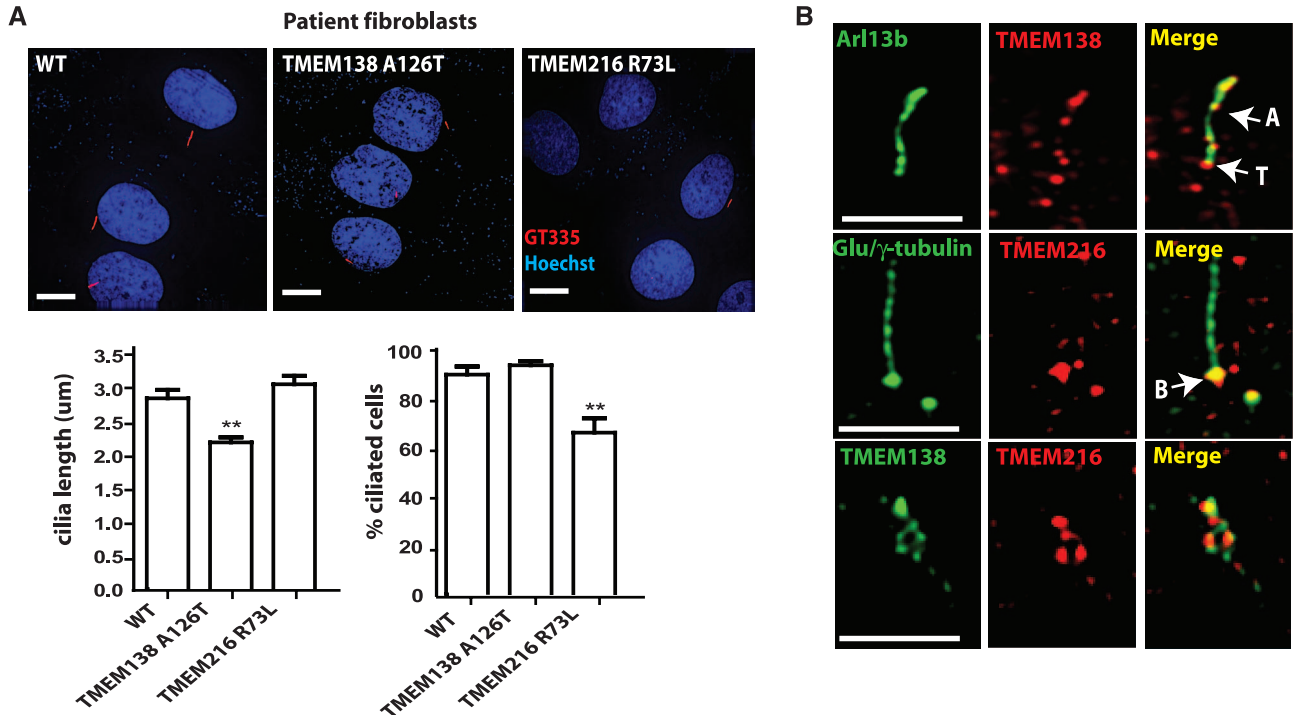
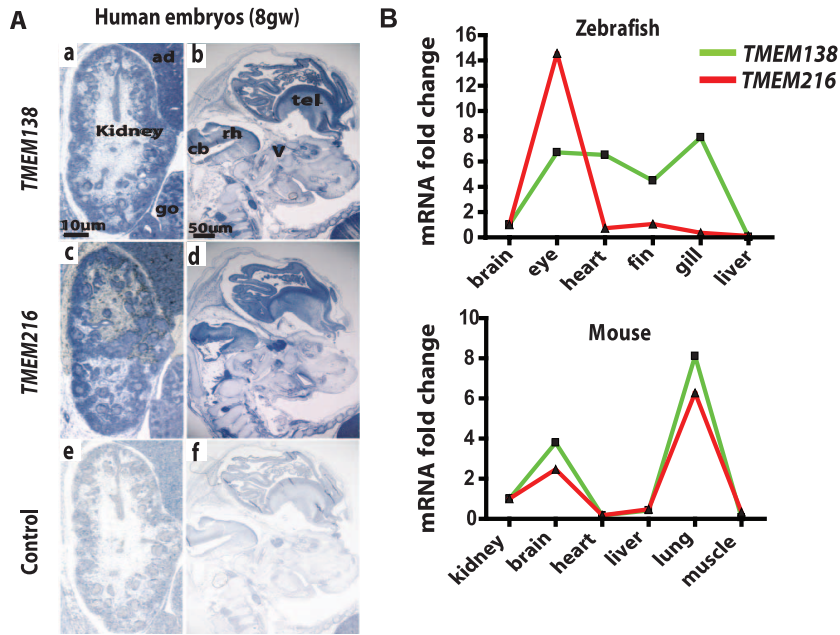


Fig. 3. Tethered vesicular trafficking of *TMEM138* and *TMEM216* to primary cilia is required for ciliogenesis. **(A)** In patient fibroblasts under 48-hour serum starvation, *TMEM138* p.A126T (MTI-656) mutations caused short cilia, and *TMEM216* p.R73L mutations disrupt ciliogenesis (defined as having cilia <1 μm long). * $P < 0.05$, ** $P < 0.01$ [versus wild type (WT) by one-way analysis of variance (ANOVA) with Bonferroni posttest, $n = 40$ to 50 cells]. Error bars indicate SEM. **(B)** In IMCD3 cells, high-resolution images of endogenous

TMEM138/216 staining show that *TMEM138* localized to ciliary axonemes and the base of cilia, whereas *TMEM216* localized primarily to basal bodies. Both *TMEMs* also show closely adjacent vesicular patterns around the base of cilia (also see fig. S9C). Anti-Arl13b (cilia), anti-Glu/γ tubulin (cilia plus centrosome), polyclonal mouse anti-*TMEM138* (fig. S9, A and B), and rabbit anti-*TMEM216* antibodies were used. A, ciliary axoneme; T, transition zone; B, basal body. Scale bars, 5 μm.

TMEM138, as well as *CEP290* (figs. S13B and S14, A and B, and movies S5 and S6), suggesting functional dependence on *TMEM216* for trafficking *TMEM138* and associated proteins.

To determine the mechanism of this functional dependence, we considered the potential role of tethering proteins in linking these two

distinct vesicular pools. Several vesicle tethering factors, including the transport protein particle (TRAPP) II complex, p115, and the conserved oligomeric Golgi complex are known to be involved in tethering Golgi vesicles (15). Among these potential tethering proteins, we found that TRAPP II mediated the tethering of *TMEM138*

and *TMEM216* vesicles crucial for ciliary assembly (Fig. 4A, SOM text, figs. S14C to S16, and movies S7 and S8).

We questioned whether the protein localization or mutant phenotype of the two *TMEMs* in species that diverged before the coordinated gene regulation should be distinct. We found the

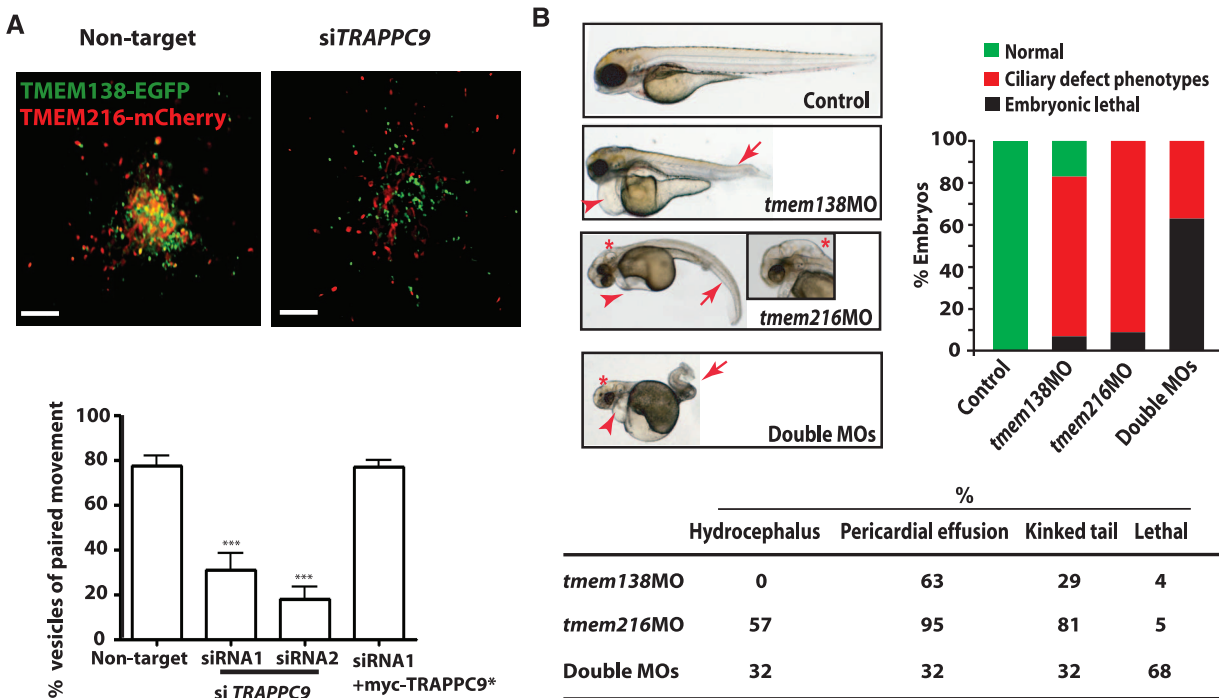


Fig. 4. Functional relatedness of TMEM138 and TMEM216. **(A)** Live-cell imaging shows that knockdown of TRAPPC9, a major subunit of the TRAPPII complex, detached the tethered TMEM138 and TMEM216 vesicles (movies S7 and S8). Defects in tethering were largely rescued by myc-TRAPPC9* (9). *** $P < 0.001$ (by one-way ANOVA with Bonferroni posttest, $n = 90$ to 110). Scale bars, 5 μ m. Error bars indicate SEM. **(B)** Injection of translation-blocking

antisense morpholinos (MOs) to *tmem138* (6 ng), *tmem216* (4 ng), or double MOs in WT (AB) zebrafish embryos leads to ciliary phenotypes of curved or kinked tail (arrows) and heart edema (arrowheads) at 3 days post-fertilization in a synergistic and dose-dependent manner (fig. S18A). Only *tmem216* morphants present hydrocephalus (*) (>50 embryos for each condition) (fig. S18B).

retention of the coordinated vesicular movement in a zebrafish cell line (ZF4) expressing the tagged version of the two TMEMs zebrafish orthologs, suggesting that the coordinated localization of proteins evolutionarily preceded the coordinated gene expression (movie S11). We next tested for conserved genetic function in zebrafish by comparing the morpholino knockdown phenotypes of each. Both knockdowns shared some typical ciliary phenotypes such as pericardial effusion, a curved or kinked tail (16) in a synergistic and dose-dependent manner, as well as gastrulation defects (Fig. 4B and fig. S18, A to D). However, only *tmem216* morphants presented hydrocephalic brains [ciliary phenotype (17)] and more severe defects of the left/right heart axis (Fig. 4B and figs. S17 and S18A). Together, the data indicate that before their adjacent genomic localization, the proteins' organismal functions were not completely congruent and were associated with distinguishable phenotypes, unlike JBTS2-linked Joubert patients.

Our findings suggest that nonparalogous genes not only can be chromosomally rearranged into a functional gene cluster during vertebrate evolution, but also can be assembled into a new CRM by evolving regulatory elements, which correlate with their coordinated expression. We are aware of few other examples in which mutations in adjacent genes lead to indistinguishable or similar human phenotypes: mutations in *EVC* or *EVC2* cause another ciliopathy, Ellis-van

Creveld Syndrome; mutations in *ABCG5* and *ABCG8* cause sitosterolemia; and disruptions of *PKD1* and *TSC2* cause renal cysts (18–20). Our results provide insight into the evolved coordinated expression and functional relatedness of adjacent nonparalogous genes as a pathogenesis of phenotypically indistinguishable genetic disorders caused by mutations at a single locus.

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Supporting Online Material

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Distinguishing Ciliopathy

Cilia were once thought to be evolutionary remnants, but structural defects reveal their importance in signaling pathways and human disease, such as Joubert syndrome. Either of the genes *TMEM138* and *TMEM216* can be found mutated in phenotypically indistinguishable ciliopathy patients. Interestingly, despite their lack of sequence homology, these genes have always been aligned in head-to-tail configuration during vertebrate evolution. The proteins expressed by these genes mark distinct tethered vesicles, which differentially carry ciliary proteins for assembly. **Lee *et al.*** (p. 966, published online 26 January; see the Perspective by **Chakravarti and Kapoor**) show that the coordinated expression of these adjacent genes depends upon a coevolved regulatory element in the noncoding intergenic region, which thus integrates the roles of both gene products. This discovery explains not only the indistinguishable pathogenesis of the patients' genotypes but also how the evolutionary clustering of genes unrelated in sequence may correlate with coordinated control of expression and function.

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