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Journal Scientific Reports, 13(1)

Authors

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Publication Date

2023-05-01

DOI

10.1038/s41598-023-34365-8

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A local translation program regulates centriole amplification in the airway epithelium

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Biogenesis of organelles requires targeting of a subset of proteins to specific subcellular domains by signal peptides or mechanisms controlling mRNA localization and local translation. How local distribution and translation of specific mRNAs for organelle biogenesis is achieved remains elusive and likely to be dependent on the cellular context. Here we identify Trinucleotide repeat containing-6a (*Tnrc6a*), a component of the miRNA pathway, distinctively localized to apical granules of differentiating airway multiciliated cells (MCCs) adjacent to centrioles. In spite of being enriched in TNRC6A and the miRNA-binding protein AGO2, they lack enzymes for mRNA degradation. Instead, we found these apical granules enriched in components of the mRNA translation machinery and newly synthesized proteins suggesting that they are specific hubs for target mRNA localization and local translation in MCCs. Consistent with this, *Tnrc6a* loss of function prevented formation of these granules and led to a broad reduction, rather than stabilization of miRNA targets. These included downregulation of key genes involved in ciliogenesis and was associated with defective multicilia formation both in vivo and in primary airway epithelial cultures. Similar analysis of Tnrc6a disruption in yolk sac showed stabilization of miRNA targets, highlighting the potential diversity of these mechanisms across organs.

Biogenesis of subcellular organelles requires efficient delivery and concentration of subsets of proteins to specific intracellular sites for prompt assembly^{1,2}. Targeting proteins to specific subcellular compartments with signal peptides or by translation of localized mRNAs is known to be involved in this process^{3–5}. mRNA subcellular localization and local translation are evolutionary conserved and efficient mechanisms to create functional and structural asymmetries in cells by rapidly producing encoded proteins when and where required³. The prevalence and precise subcellular localization of certain mRNAs in association with specific organelles suggest a significant contribution of local translation⁴. Mechanisms generating asymmetric distribution of proteins in polarized cells are functionally relevant in regulating multiple aspects of cell behavior, including proliferation, migration and differentiation.

Centrioles are core structures of centrosomes involved in nucleation and formation of mitotic spindles critical for cell division⁶. Centrioles also function as basal bodies for the assembly of primary cilia and multicilia^{6,7}. During multiciliogenesis, each differentiating multiciliated cell (MCC) generates hundreds of centrioles. A number of mRNAs localize and concentrate in centrioles or centrosomes^{3,8,9}. Concentration of proteins and mRNAs in centrosomes might be achieved by targeting actively translating polysomes⁸. In addition, centriolar proteins are suggested to recruit the protein translation machinery that regulates the translation of specific mRNAs¹⁰. Still, little is known about the mechanisms of mRNA localization and local translation in centriole amplification. It is also unclear how microRNAs contribute to this process.

¹The Columbia Center for Human Development, Division of Pulmonary, Allergy & Critical Care Medicine, Department of Medicine, Columbia University Medical Center, Columbia University, College of Physicians & Surgeons, 650 West 168th Street, BB 8-812, New York, NY 10032, USA. ²Division of Cellular and Developmental Biology, Department of Molecular & Cell Biology, University of California, Berkeley, CA, USA. ³Department of Biochemistry and Molecular Biology, Faculty of Medicine, University of Calgary, Calgary, Alberta, Canada. ⁴Division of Lung Diseases, National Heart, Lung, and Blood Institute, National Institutes of Health, 6705 Rockledge Drive, Room 407-J, MSC 7952, Bethesda, MD 20892-7952, USA. ⁵These authors contributed equally: Helu Liu and Huijun Li. ^{Ex}email: wvc2104@cumc.columbia.edu; jining.lu@nih.gov Trinucleotide repeat containing 6a (TNRC6A) is a direct partner of Argonaute proteins (AGOs), which play a critical role in miRNA-induced mRNA degradation¹¹. AGOs and TNRC6A have been reported in randomlydistributed cytoplasmic granules known as P-bodies, which are enriched in translation suppressors and mRNA degradation enzymes, but lack protein synthesis machinery¹². TNRC6A and AGO2 can be also detected in other cytoplasmic ribonucleoprotein granules, such as stress granules and neuronal transport granules^{13–15}. Cell culture studies show *Tnrc6a* present in most cell types, however analyses of developing and adult tissues in vivo show that levels of *Tnrc6a* expression can differ markedly^{16–18}.

In an effort to gain further insights into the tissue distribution and processes associated with *Tnrc6a* and miRNAs, here we identified *Tnrc6a* in the lung selectively expressed in MCCs of conducting airways. We found TNRC6A protein asymmetrically distributed in these cells in a not previously identified population of apical granules, which also contained AGO2. However these granules differed from P-bodies as they lacked mRNA degradation enzymes and instead were enriched or closely associated with components of the mRNA translation machinery and newly-synthesized proteins. Loss of function of *Tnrc6a* prevented these granules from forming and resulted in broad reduction of miRNA targets, including key components of the centriole biogenesis machinery required for multicilia formation. Our study suggests that this novel class of TNRC6A-containing apical granules can act as local translation hubs required to mediate large-scale production of centriolar proteins in MCCs. Our data also supports the idea that key components of the miRNA pathway can be utilized for mRNA recruitment and localization required for local translation.

Results

Thrc6a is selectively expressed in multiciliated cells of the lungs. We have previously generated a *Thrc6a*^{gt/+} reporter mouse carrying a gene trap insertion of β -galactosidase in the *Thrc6a* locus to investigate expression and function of *Thrc6a* in mouse development¹⁸. In the process of further characterizing these mutants, we performed marker analysis and X-gal staining in the adult lungs and found a striking selective pattern of β -galactosidase activity in the airway epithelium (Fig. 1a). Little to no signals were present in the alveolar epithelium and overall in the mesenchymal compartment. This pattern was similarly observed for *Thrc6a* as determined by in situ hybridization (Fig. 1b). Moreover, β -galactosidase activity co-labeled with acety-lated α -tubulin (Ac- α -tub), indicating that *Thrc6a* was selectively expressed in MCCs of the airway epithelium (Fig. 1c).

To further confirm expression and cell type specificity, we performed immunofluorescence (IF) staining of lung sections with the human index serum (termed as 18,033), widely used for detection of TNRC6A protein^{19,20}. This showed cytoplasmic signals restricted to Ac- α -tub positive cells, further corroborating TNRC6A selectivity to MCCs (Fig. 1d). To investigate when these granules are established and their association to multiciliogenesis, we analyzed airway epithelial progenitors (basal cells) from adult mouse trachea undergoing differentiation in air–liquid interface (ALI) cultures²¹. Analysis of TNRC6A (18,033) and FOXJ1, an early marker of MCC cell fate²¹ showed TNRC6A signals first detected at ALI day2 in emerging FOXJ1+ cells. By day 3–4, as these cells underwent large-scale centriole amplification to initiate multiciliogenesis²¹, TNRC6A+ cytoplasmic granules became abundant in MCCs (Fig. 1e). Signals were subsequently diminished in more mature MCCs, which by then expressed strong Ac- α -tub (Fig. 1f).

TNRC6A is enriched in a novel class of apical epithelial granules that lack mRNA degradation enzymes. To gain additional insights into the subcellular distribution of these granules in MCCs, we analyzed a series of Z-stack images of ALI day 4 cultures double-stained with the human index serum and FOXJ1. This revealed an overall population of granules heterogeneous in size, randomly distributed throughout the cytoplasm of MCCs but also identified a population of apically-localized granules of smaller size in these cells. Immunostaining using a monoclonal TNRC6A antibody (4B6) and acetylated alpha tubulin showed these apical granules underneath the multicilia of differentiating MCCs (Fig. 2a,b). Their localization in MCCs was further confirmed in ALI cultures transduced with a lentiviral vector expressing a TNRC6A-EGFP fusion protein²¹ (Fig. 2c).

The human index serum is also known to label P-bodies (processing bodies), cytoplasmic ribonucleoprotein (RNP) granules, which contain TNRC6A and various RNA pathway enzymes involved in decapping and degradation of mRNAs²². To further identify potential differences in the populations of MCC granules, we stained ALI cultures with human index serum and antibodies against components of the P-bodies. IF for 18,033 doublelabeled with EDC4, DCP1A, DDX6 and XRN1 showed these signals highly expressed in the randomly-localized population of cytoplasmic granules (P-bodies), but largely undetectable in the apical granules that we also identified using the anti-TNRC6A (4B6) antibody in the same MCC (Fig. 2b,d,e, Supplementary Fig. 1). TNRC6A is a known partner of AGO2¹². Co-IF staining of human index serum with AGO2 showed co-localization of AGO2 in both classes of granules in MCC (Fig. 3a). Notably, lentiviral gene transduction of AGO2-EGFP fusion protein showed the strong EGFP signals enriched apically underneath the cilia of differentiating MCCs (Fig. 3b).

This supported the idea that TNRC6A-containing apical granules represented a potentially novel class of granules distinct from P-bodies in size, subcellular distribution and absence of enzymes needed for mRNA degradation.

microRNAs are crucial components of the TNRC6A apical granules during multiciliogenesis. We examined the possibility that microRNAs previously reported in MCCs could be locally enriched where the TNRC6A apical granules were abundant. Indeed, ISH for miR-449a, miR-34a, miR-34b, representative MCC miRNAs²³⁻²⁶, showed strong signals apically, adjacent to the base of multicilia (Fig. 3c, Supplementary

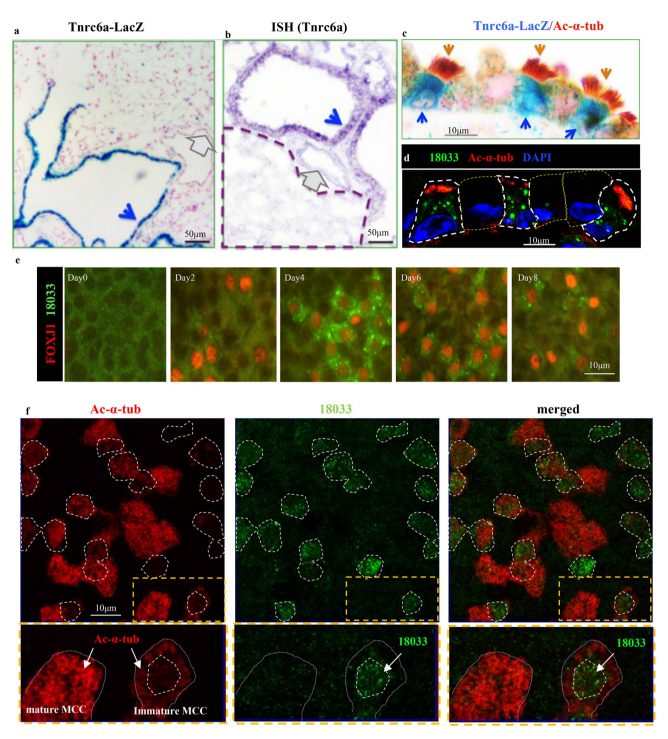


Figure 1. *Thrc6a* is selectively expressed in multiciliated cells (MCCs) of airways in the mouse lungs. (**a**) β -Galactosidase staining of *Thrc6a*^{gt/+} reporter mice showing selective expression in the airway epithelium (blue arrow), compared to vascular (gray arrow) and alveolar structures. (**b**) *Thrc6a* in situ hybridization confirming expression in airway epithelium (blue arrow), but not in vessels (red arrow) or alveoli (outlined by dashed line). (**c**) Strong β -galactosidase signals (blue arrow) in MCCs labeled by Ac- α -tub (brown arrow) compared to other airway cell types. (**d**) Immunofluorescence (IF) with human index serum (18033: green) detects TNRC6A in MCCs (Ac- α -tub: red). (**e**) Time course of detection of TNRC6A-containing granules in MCCs (FOXJ1+) differentiating from adult airway progenitors in air–liquid interface (ALI) cultures. (**f**) Distinct abundance of cytoplasmic TNRC6A expression (18033, dotted area and arrow) in immature and mature MCCs in ALI day4 cultures, as shown by strong or weak Ac- α -tub signals, respectively. Lower panels enlarged from boxed areas (yellow).

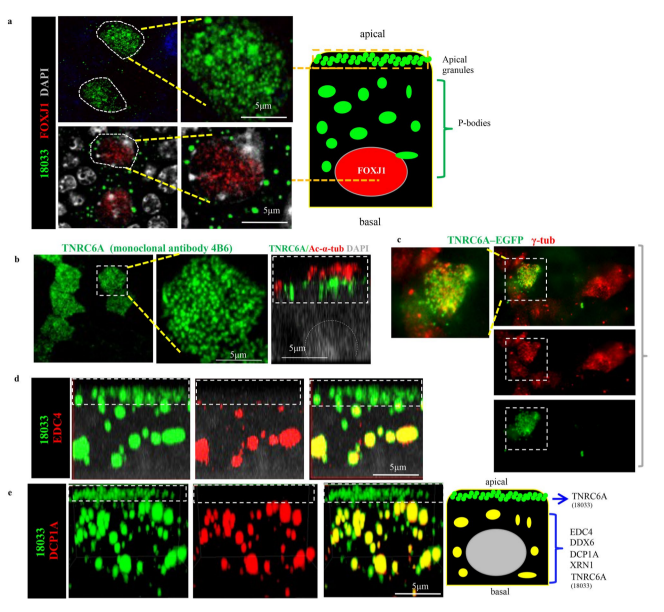


Figure 2. TNRC6A protein concentrates in apically-localized granules that lack mRNA degradation enzymes. (a) Apical granules and randomly-localized granules (P-bodies) detected by human index serum (18033) in differentiating MCC. Dashed lines depict a plane of Z-stack image from the apical or basal areas of the same MCC (diagram). (b) TNRC6A monoclonal antibody (clone#:4B6) identifies apical granules underneath the multicilia (Ac- α -tub). X–Z projection of confocal z-stack images. (c) TNRC6A-EGFP fusion protein expressed by a lentiviral vector concentrates in apical granules of MCC identified by co-labeling with γ -tub (boxed area) in ALI day3 culture. (d, e) Co-staining of human index serum with EDC4 or DCP1A in a single MCC showing double-labeling largely in P-bodies instead of the apical granules (dashed box) Diagram depicts distinct components of these granules (See also Suppl. Fig. 1).

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Fig. 2). Thus, the data suggested that the TNRC6A apical granules contained AGO2, miRNAs and presumably their targets.

We then asked whether microRNAs were necessary for formation of these apical granules during MCC differentiation. Expression of the miR-34/449 family is required for basal body maturation and apical docking during multiciliogenesis²³. We examined ALI airway epithelial cultures from miR-34/449 null mice in which all three loci encoding miR-34/449 were deleted²⁴. Analysis of ALI day 4 cultures in WT differentiating MCCs showed the typical abundant TNRC6A apical granules revealed by human index serum IF (18033 single-labeled Fig. 3d, left) and the larger randomly distributed double-labeled P-bodies (18033-DCP1A-labeled granules). By contrast, in miR-34/449 null cultures, apical TNRC6A granules were notably decreased, absent or disorganized in contrast to the P-bodies, which were largely preserved (Fig. 3d, right). Given the high abundance of miR-34/449 in MCCs (~70% of all miRNAs)²³, the data strongly suggest that miR-34/449 is a key cellular component required

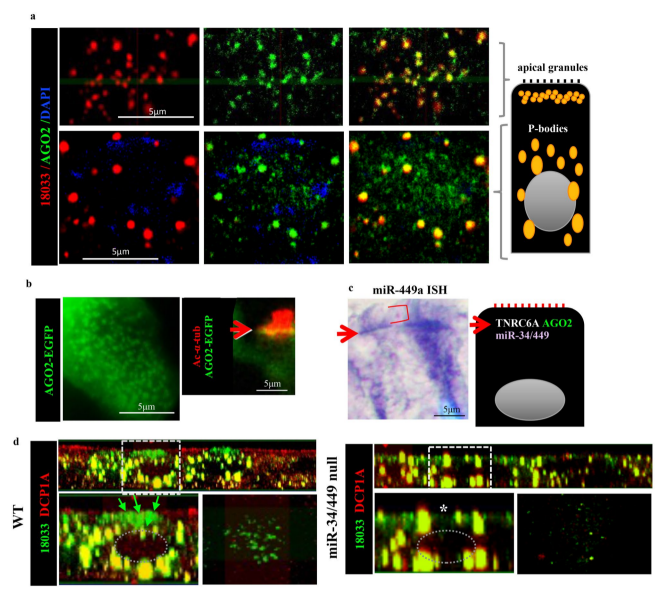


Figure 3. AGO2 and miRNAs are present in TNRC6A-containing apical granules in differentiating MCCs. (a) Co-staining of AGO2 (clone 11A9) and TNRC6A (18033, human index serum) showing double-labeling in apical granules and P-bodies within the same MCC. (b) AGO2-EGFP fusion protein produced by a lentiviral vector in differentiating MCC in ALI cultures showing apical granules underneath the emerging cilia (red, arrow points to AGO2-EGFP fusion protein underneath Ac- α -tub-labeled cilia). (c) In situ hybridization (ISH) showing miR-449a enriched apically (arrow) underneath the cilia (red bracket). (d) Co-staining human index serum (green) and DCP1A (red) in ALI day 4 cultures from WT controls (left) and miR-34/449 triple null mice (right). Boxed area in top panels enlarged in bottom panel (YZ projection and a corresponding apical Z-stack image). WT: Abundant single-labeled TNRC6A apical granules (arrows: 18033, green) and double-labeled larger granules (yellow 18033-DCP1A: P-bodies). miR-34/449 null ALI cultures: notable decrease in abundance of TNRC6A apical granules (*, green) not seen in the P-body population (yellow double-labeled granules). Nuclei represented in grey (diagrams).

for the formation of apical granules. The failure to form these granules is likely linked to events that lead to the organization of the apical domain of differentiating MCCs to initiate multicilia formation.

TNRC6A-containing apical granules are sites of newly-synthesized proteins in differentiating

MCCs. Our study identified TNRC6A, AGO2 and miRNAs in apical granules lacking enzymes for mRNA decay in differentiating MCCs. We asked whether TNRC6A was involved in mRNA localization and local translation at the apical domain in these cells, which is a site of active multicilia formation. Puromycin has been reported to label newly-synthesized proteins, since it mimics aminoacyltRNA and can enter the ribosome A site to terminate translation by ribosome-catalyzed covalent incorporation into the nascent polypeptide

C-terminus²⁷. O-Propargyl-puromycin (OPP), an alkyne analog of puromycin, has been used to detect localized translation^{27–29}.

To visualize newly-synthesized polypeptides in differentiating MCCs, ALI Day 4 mouse airway epithelial cultures were labeled with OPP for 30 min before termination and subsequently analyzed by IF-confocal microscopy. OPP staining revealed strong signals localized in apical foci underneath emerging cilia (Fig. 4a). The OPP foci were cell type-specific and dynamic, as they were not detected in mature MCCs or in other airway epithelial cell types. Notably, OPP signals were abolished by pretreatment with cycloheximide (Fig. 4b), supporting the specificity of this approach. OPP foci were similarly found in differentiating MCCs from human ALI airway epithelial cultures, suggesting conservation of this mechanism in mammalians (Fig. 4c). Importantly, OPP signals were undetected in the randomly-localized P-bodies. Co-IF of OPP with the human index serum (18033) and DCP1A in ALI day4 cultures showed OPP-18033 signals overlapping apically but no 18033 overlapped with the DCP1A positive P-bodies within the same MCC (Fig. 4d).

A closer analysis of the pattern of OPP staining showed that the OPP foci varied considerably in size and signal intensity In both human and mouse airways even within the same MCC, suggestive of a dynamic regulation (Fig. 4e). Morphometric analysis revealed OPP foci ranging from 300 to 2000 nm in diameter and signal intensity ~ 2.7 folds higher locally compared to the neighboring area (OPP mean optical density, P < 0.0001). Overall OPP staining could be identified in small and large foci in the apical region of differentiating MCCs. The small OPP foci overlapped extensively and at various degrees with the small TNRC6A-containing granules (labeled with 18033). By contrast the large OPP foci also stained apical granules but these were not 18033 (TNRC6A) positive or P-bodies (Fig. 4f,g and diagrams).

Newly-synthesized proteins and translation machinery are found in apical granules during centriole amplification. The distinct association of TNRC6A with OPP foci led us to further investigate what these sites of newly-synthesized protein represented in differentiating MCCs. Their vicinity to apically-located ciliary structures suggested that OPP could be labeling sites of local translation of centriolar components required for multicilia formation ^{30,31}. To examine this possibility we performed co-immunostaining of OPP with key centriole biogenesis markers in ALI day 3–4 cultures. This revealed the smaller OPP foci (~400–800 nm in diameter), abutting centrioles (300–400 nm in diameter), as identified by CENTRIN3, and colocalized with DEUP1, a marker of deuterosomes ^{30,31} (Fig. 5a,b).

The spatial relationship of TNRC6A, CCP110, DEUP1, and TNRC6A with the small OPP foci was examined in differentiating MCCs by double and triple-IF stainings (Fig. 5a–d). The CCP110 signals were found surrounding double positive DEUP1/OPP small foci, suggesting the co-localization of TNRC6A positive granules with procentrioles surrounding deuterosome. Interestingly, this co-IF analysis showed no overlap of the large OPP foci with CENTRIN3, CCP110, DEUP1, or TNRC6A. Instead, the large OPP foci (~800–2000 nm in diameter) colocalized with PCM1 (Fig. 5e), a component of the pericentriolar material widely reported in fibrous granules ^{30,31}.

We then asked whether components of the translation machinery were present at the sites of OPP expression. Indeed, IF analysis of ALI cultures showed the small OPP foci strongly co-labeled with eIF3B and phosphorylated RPS6 (p-RPS6) (Fig. 6a,b), indicative of active translation in the apical granules we had previously shown to contain TNRC6A (Fig. 5). Single-molecule fluorescence in situ hybridization (smFISH) revealed ribosomal protein 3 *Rpl3* mRNA (a target of mir34/449) largely expressed in the OPP small foci of TNRC6A-granules (Fig. 7a). The *Rpl3* distribution differed markedly from that of *MLS* mRNA (RNA transcribed from the mitochondrial L-strand DNA). Although known to be abundantly expressed in differentiating MCCs, *MLS* was neither co-localized nor associated with any OPP in these cultures (Fig. 7b). Interestingly, in the large OPP foci (associated with fibrous granules, Fig. 5) we detected expression of p-RPS, eIF3B and ribosomal RNAs (18S, 28S) surrounding the OPP signals (Figs. 6c,d, 7c,d). A similar subcellular arrangement has been described in the sponge bodies, cytoplasmic organelles of the developing *Drosophila* oocyte, in which the translational activator Orb and ribosomes are located at the edge or surrounding these bodies to promote local translation ³². The lack of PCM1-18033 co-labeling in the large OPP foci suggested that, in fibrous granules, newly-synthesized protein is regulated by a TNRC6A-independent mechanism. Altogether the data strongly suggested that TNRC6A-contaning apical small granules are active sites of local translation during centriole amplification in differentiating MCCs.

Disruption of *Tnrc6a* expression leads to reduced CCP110, abnormal centriologenesis and defective multicilia formation. Next, we investigated whether *Tnrc6a* expression was required for the formation of the apical granules in differentiating MCCs. For this, we examined ALI airway epithelial cultures from a *Tnrc6a* null mice carrying an EGFP reporter to identify cells committed to MCC fate (*Tnrc6a^{null}*, *FOXJ1*-*EGFP*, see "Methods"). IF staining of ALI day 4 with human index serum showed that disruption of *Tnrc6a* specifically abolished the apically-localized granules, but not the cytoplasmic P-bodies in FOXJ1-EGFP-labeled MCCs (Fig. 8a). We asked how this could affect production of CCP110, a key centriolar protein involved in ciliogenesis and known target of miR-34/449²⁴. Thus, we co-cultured airway progenitors from wild type and *Tnrc6a^{null}*; *FOXJ1-EGFP* mutants and differentiated these cells in ALI cultures. This allowed to compare potential differences in the intensity of signals between *Tnrc6a*-deficient and WT cells but expression was drastically reduced in the *Tnrc6a^{null}*; *FOXJ1-EGFP* MCCs (Fig. 8b). The presence of FOXJ1-EGFP in *Tnrc6a* mutant cells, suggested no significant defect in MCC fate determination (Fig. 8a,b). This was further supported by analysis of E18.5 lungs showing that the proportion of FOXJ1 positive cells in airways from WT and *Tnrc6a^{null}* mutants was essentially the same (Supplementary Fig. 4).

IF staining of ALI co-cultures of *Tnrc6a^{null}*, *FOXJ1-EGFP* and WT epithelial cells revealed marked differences in levels and distribution of proteins involved in centriole amplification. For example, in *Tnrc6a* null cells

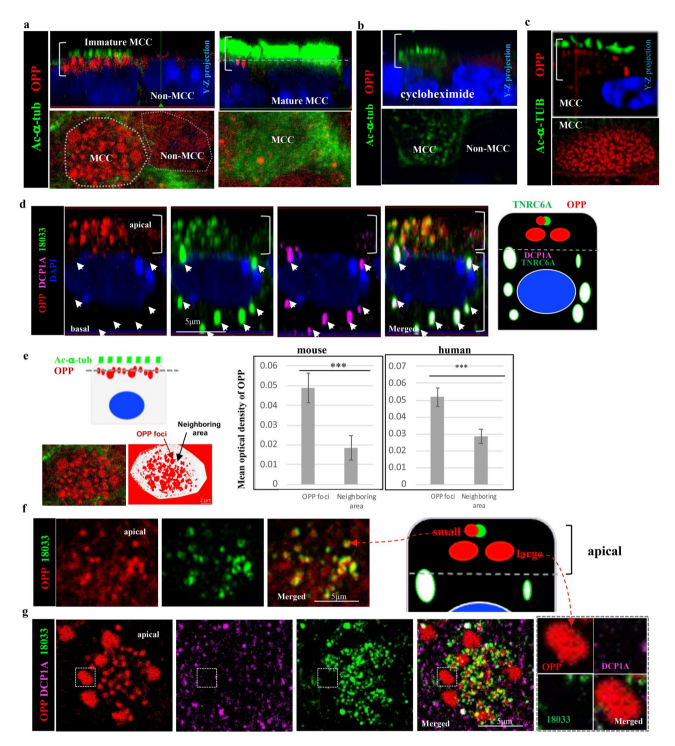


Figure 4. TNRC6A-containing apical granules are sites of newly-synthesized proteins in differentiating MCCs. (a) Strong O-propargyl-puromycin (OPP) signals of newly-synthesized proteins in the apical region (brackets) of **mouse** immature MCCs (left panels, with shorter and less cilia labeled by Ac-alpha tubulin). No OPP signals in non-MCCs (left panels) or in mature fully-differentiated MCCs (right panels: abundant tall Ac-alpha tubulin positive cilia). (b) OPP signals are abolished in **mouse** immature MCCs from ALI cultures treated with cycloheximide, a protein synthesis inhibitor. (c) Strong OPP expression and distribution in the apical region (brackets) immature MCCs from **human** ALI airway epithelial cultures. (d) OPP co-staining with DCP1A and human index serum (18033) showing OPP and TNRC6A in the apical region of differentiating mouse MCC (bracket). No OPP in DCP1A positive P-bodies of the same MCC. Diagram depicting OPP distribution in apical granules and P-bodies (compare with diagrams in Fig. 2). (e) Quantitation of mean optical density of OPP signals in apical granules compared to the neighboring area in the same plane of section (diagram on the left) using Image J. Bars and lines are mean (±sd) of measurements in 30 immature MCCs from three independent mouse or human ALI cultures ***:P<0.0001. (f, g). Heterogeneity of OPP in small and large apical foci. TNRC6A (18033-labeled) signals overlap with the smaller apical OPP foci but not with the apical large OPP foci (g). DCP1A signals do not overlap with apical TNRC6A or OPP signals. (f, g) are representative Z stack images of the apical region of an immature MCC. Boxed area in (g) is enlarged on the right panel. Diagram in (f) depicts OPP small and large foci in apical differentiating mouse MCCs.

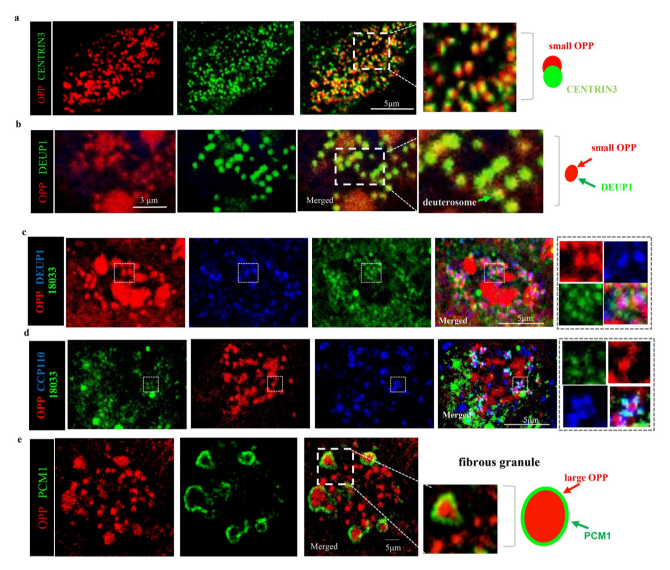


Figure 5. Sites of newly-synthesized proteins are found associated to centriolar components and fibrous granules in differentiating MCCs. (**a**, **b**) Small OPP foci abutting centrioles (CENTRIN3 positive) and in deuterosomes colocalized with DEUP1. (**c**) Small OPP foci are present in deuterosome-associated TNRC6A granules. By contrast, the large OPP foci, also found apically in differentiating MCCs, do not colocalize with TNRC6A. (**d**) TNRC6A positive granules surrounding the small OPP foci co-localize with procentrioles marked by CCP110. (**e**) The large OPP foci localize to fibrous granules (PCM1 positive). In all plates a representative plane of Z-stack image from the apical region of a single MCC is depicted. Boxed areas are enlarged on the right panels. Diagrams depict the relationship of the various components listed with the small and large OPP foci.

signals for CEP135 and PCM1 were clustered in a corner of the differentiating MCC, characteristic of an early stage of centriole amplification (Fig. 9a,b) ^{33,34}. Moreover, in mutant MCCs Centrin3 staining was drastically reduced (Fig. 9c). These changes were compatible with a significant defect of Tnrc6a mutant cells in generating the large number of normal centrioles/basal bodies required for multiciliogenesis. Consistent with a defect in centriologenesis, few cilia formed in differentiating MCCs of Tnrc6a mutants when co-cultured with WT cells (Supplementary Fig. 5a). The defect in multicilia formation was further confirmed by examining the trachea of Thrc6a null mutants at E18.5 (Supplementary Fig. 5b-d). Efficient disruption of Thrc6a was demonstrated by IF staining with the monoclonal antibody (4B6), which showed TNRC6A positive granules in tracheal MCCs of wild type but not of mutants (Supplementary Fig. 5b). Co-staining with Ac-a-tub and TNRC6A demonstrated a significantly reduced multicilia staining in *Tnrc6a* null tracheas. Scanning electron microscopy of tracheae from mutants showed the majority of MCCs (83%) with shorter and sparser cilia as compared to WT (Supplementary Fig. 5c,d). The cilia truncation was consistent with an immature MCC phenotype. We found no difference in FOXJ1 staining between WT and Tnrc6a null lungs (Supplementary Fig. 4), suggesting no defect in MCC fate determination. Transcriptome analysis of WT and Tnrc6a null lungs did not detect significant changes in expression of genes associated with proliferation or differentiation of other cell types of the lung. Moreover, no gross abnormalities were observed in these embryos or in the overall morphology of their lungs and no compensatory

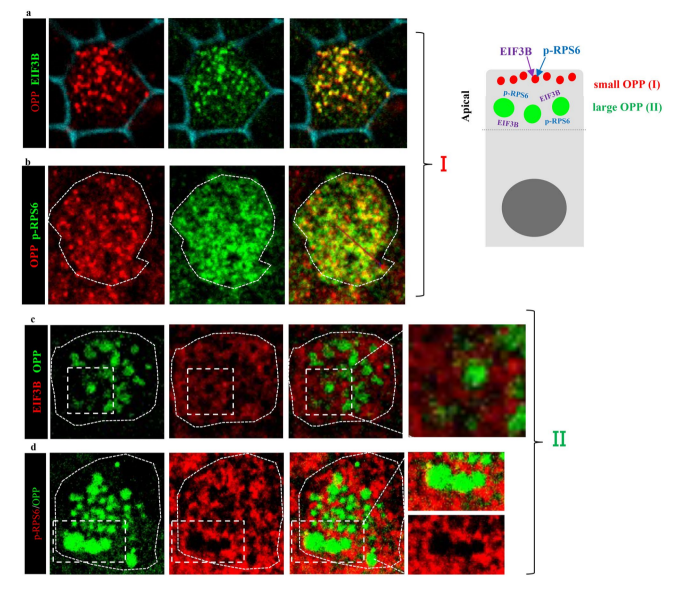


Figure 6. Translation initiation factors are co-expressed or adjacent to OPP foci in differentiating MCCs. All panels: confocal images of a single MCC in ALI day 4 culture. Diagram (right) summarizing the findings. (**a**, **b**) Translation initiation factors (EIF3B and p-RPS6) overlap with smaller OPP apical foci (I, shown to be in TNRC6A-enriched granules in Fig. 5). Phalloidin-stained cell membrane (blue) depicts apical MCC region. (**c**, **d**) EIF3B and p-RPS6 proteins are found surrounding the large OPP apical foci (II, shown to be in fibrous granules in Fig. 5). High magnification panels on the right are enlarged view of the boxed areas.

increased expression of other *Tnrc6* family member (*Tnrc6b* and *Tnrc6c*) was observed in Tnrc6A null lungs (Supplementary Fig. 6)¹⁸.

Together, these data strongly support that idea that loss of *Tnrc6a* leads to a defect selectively in MCCs, the same cell type where we found *Tnrc6a* highly expressed in WT lungs.

Disruption of Tnrc6a expression leads to reduction, rather than stabilization of miRNA targets in mutant lungs. TNRC6A has been traditionally described as a component of the RNA-induced silencing complex in microRNA (miRNA)-mediated gene suppression, repressing translation or promoting mRNA degradation. This is consistent with previous findings of defective miRNA function with increased miRNA target expression in yolk sac endoderm of Tnrc6a null mice¹⁸. We examined the subcellular localization of TNRC6A in E9.5 yolk sacs and found endodermal signals colocalized with AGO2, EDC4 and other mRNA degradation enzymes in randomly-distributed cytoplasmic granules suggestive of P-bodies¹⁷ (Supplementary Fig. 7). Intriguingly, our analysis of TNRC6A expression in the lung showed evidence of its localization in differentiating MCCs in apical granules distinct from P-bodies in size, subcellular distribution, absence of enzymes needed for mRNA degradation and enrichment in translation machinery components. Moreover we had evidence of TNRC6A apical granules as sites of active translation crucial for large-scale centriole amplification in multiciliogenesis. This suggested that TNRC6A could have a distinct role in the lung, compared to the yolk sac, where it promotes

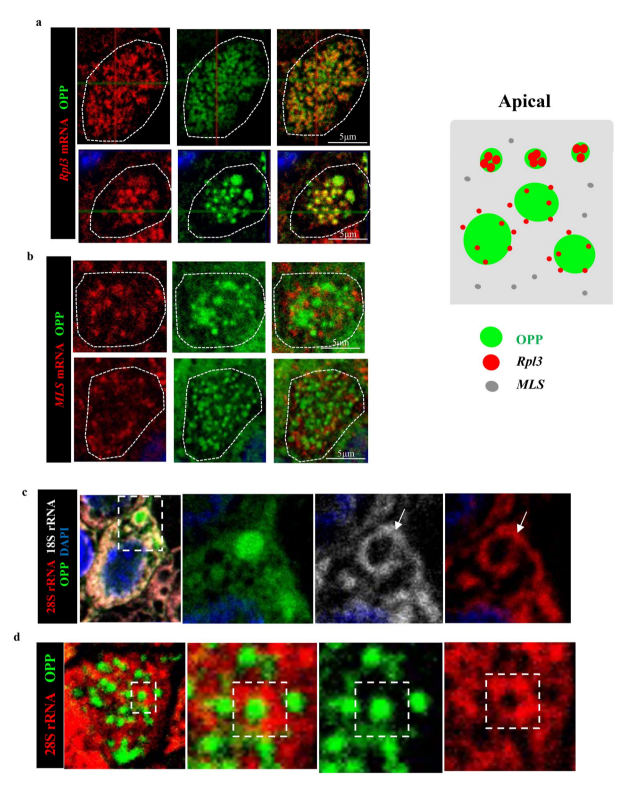


Figure 7. mRNAs and rRNAs distribution in small and large OPP foci in differentiating MCCs. RNAs detected by smFISH. Panels are representative of a plane of a Z-stack image of the apical region from a single MCC. (a) *Rpl3* mRNA strongly present in the OPP small foci (upper panel) but weaker and less concentrated in the large OPP foci (bottom panel). (b) RNA transcribed from the mitochondrial L-strand DNA does not overlap with any OPP foci. Diagram summarizing these findings (right). (c, d) 18 s and 28 s rRNA signals (arrows) strongly expressed surrounding the large OPP foci. Right panels enlarged from boxed areas.

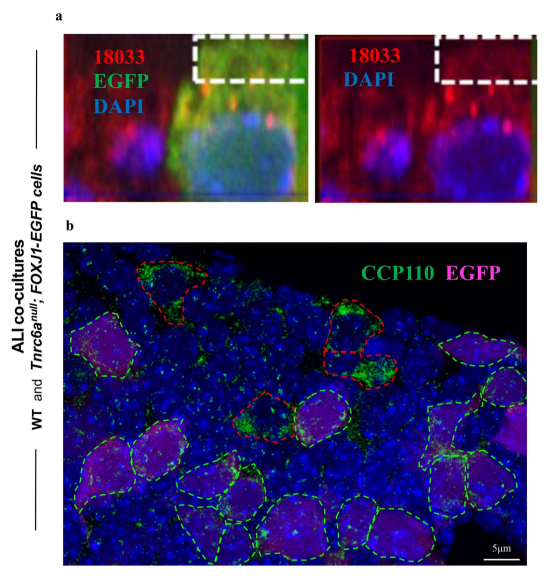


Figure 8. *Thrc6a* loss of function results in loss of apical granules and reduced CCP110 expression. ALI day 4 co-cultures of WT and *Thrc6a^{null}*; *FOXJ1-EGFP* tracheal epithelial cells. (**a**) *Thrc6a* null mutant MCC marked by FOXJ1-EGFP (green). Human index serum (red) stains both apical granules and P-bodies. However in this mutant MCC apical granules are absent due to loss of Thrc6a (boxed area). (**b**) Strong IF CCP110 signals detected in WT differentiating MCCs (outlined by red dashed line), but nearly absent in *Thrc6a* null mutant cells (marked by FOXJ1-EGFP, outlined by green dashed line).

miRNA target degradation. To investigate this possibility we examined the behavior of miRNA in the lung and yolk sac from control and Tnrc6a null mice.

We performed gene set enrichment analysis (GSEA) using a dataset containing the most evolutionary conserved targets of 83 conserved miRNAs as determined by Targetscan (http://www.targetscan.org). GSEA of these genes in yolk sac (GSE30244) showed targets of most miRNAs (78 out of 83) significantly enriched in the list of genes upregulated in *Tnrc6a* null yolk sacs (Supplementary Fig. 8). This was consistent with a role for *Tnrc6a* in the destabilization of miRNA targets^{18,35}. Strikingly, similar analysis using the *Tnrc6a* null lung dataset (GSE89327) showed that targets of most miRNAs (79 out of 83) were enriched among the genes significantly reduced in *Tnrc6a* mutant lungs, but not among the genes upregulated (Supplementary Fig. 9). This was consistent with the idea that, in the lung, *Tnrc6a* stabilizes miRNA targets instead of promoting degradation.

Indeed, analysis of miR-34/449 predicted targets in both yolk sac and lung datasets showed that, among 94 transcripts with conserved binding sites significantly altered in *Tnrc6a* null lungs, the majority (80%) was downregulated. By contrast 75% of the conserved miR-34/449 targets altered in yolk sac mutants were upregulated (72 total). Among the top 30 most downregulated genes in mutant lungs, several have been implicated in multicilia formation and 30% of these genes are predicted targets of miR-34/449, key in multiciliogenesis^{23,24} (Supplementary Table 1, Supplementary Fig. 10). For example, Spef2 and Rsph1 were strongly expressed in MCCs of the developing lung and their signals were reduced in *Tnrc6a*^{null} mutants. qRT-PCR further confirmed

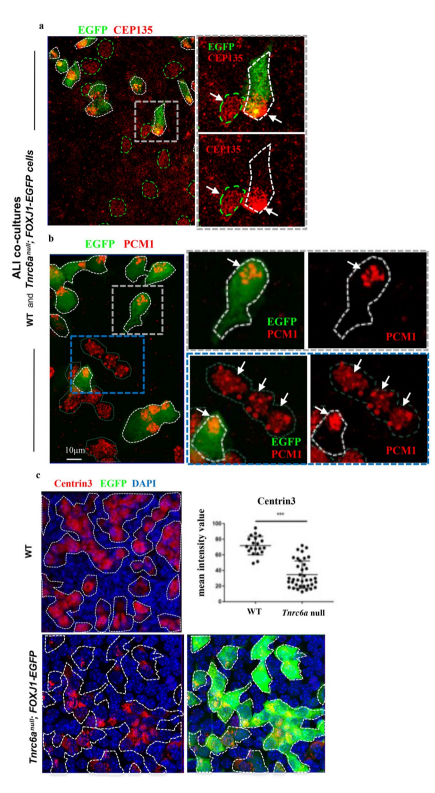


Figure 9. Defective centriole amplification in *Thrc6a*-deficient cells. (**a**, **b**) ALI day 4 co-cultures of WT and *Thrc6a^{null}*; *FOXJ1-EGFP* tracheal epithelial cells. IF for CEP135 (**a**) or PCM1 (**b**): Asymmetric distribution of signals (arrows) clustered in the corner of *Thrc6a* null defective MCCs (marked by FOXJ1-EGFP, white dashed) contrasting with the homogeneous signal distribution in WT (green dashed) MCCs. Boxed areas enlarged in the right panels. (**c**) IF of Centrin3 protein in ALI day 4 culture of WT (top panel) and *Thrc6a null*—FOXJ1-EGFP tracheal epithelial cells (bottom panels). Dashed lines depict differentiating MCCs in WT and mutants (visualized in green by FOXJ1-EGFP). Marked decrease in Centrin3 signals in mutant MCCs confirmed by quantitation of signal intensity (Image J software). Graph: dot plot (mean, ± sd of 30 MCCs from three independent experiments). Student's Test. ***P<0.0001.

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the reduced expression of additional miR-34/449 targets, such as *Rnf213*, *Tmem159* and *Calcrl*, besides *Rsph1*, *Spef2* (Supplementary Fig. 10). As demonstrated in other systems, genetic deletion of *Tnrc6a* disrupted miRNA function without affecting miRNA expression or stability¹⁸. Analysis of miR-449 (from the miR34-449 cluster) and miR-146 showed no difference in expression between WT and *Tnrc6a* null mutants (Supplementary Fig. 11).

Altogether, our data strongly suggested that, in the lung, *Tnrc6a* was not involved in degradation of miRNA targets but rather appear to be required for mRNA stabilization, and likely promoting protein synthesis.

Discussion

Here we identified TNRC6A selectively expressed in MCCs of lungs in vivo, concentrated in a novel class of apical granules containing AGO2, miRNAs and their targets, adjacent to immature centrioles. Although present in adult airways, these granules are mostly abundant in differentiating immature MCCs. Remarkably, components required for mRNA degradation are missing in these granules, while present in P-bodies within the same MCCs. Instead, they are enriched in components of the mRNA translation machinery and show evidence of concentrated newly-synthesized proteins. We found that, in the lung, loss of *Tnrc6a* leads to broad reduction instead of accumulation of miRNA targets, suggesting a role in target mRNA stabilization, rather than in degradation. Proximity-dependent biotinylation assays using centriolar and cilia proteins as baits have reported CCP110 in close proximity to TNRC6A³⁶. Here we show CCP110 and other centriolar proteins localized to the apical TNRC6A granules and no evidence of their association with enzymes involved in mRNA degradation. Notably, loss of *Tnrc6a* resulted in a drastic reduction of expression of centriolar proteins, abnormal centriologenesis and, ultimately, defective MCCs.

Cytoplasmic RNP granules in mammalian cells, including P-body, stress granule, neuronal transport granule and germ granule are generally linked to translation repression. mRNAs are not translated in stress granules where translation initiation factors are concentrated. Indeed, we found no evidence of newly-synthesized polypeptides in P-bodies in differentiating MCCs. Instead, OPP and translational machinery components were present in the apical TNRC6A-containing granules. AGO2 mainly functions as a miRNA binding protein to trigger miRNA-mediated mRNA degradation^{37,38}. Interestingly, AGO2 was originally purified and cloned as a translation initiation factor and named EIF2C2 ³⁹. It is known that AGO2 can interact with EIF1A in the biogenesis of miRNAs⁴⁰, miiRNAs are well known as a negative regulators of gene expression. However, miRNAs have been also shown to stimulate mitochondrial translation to activate translation of miRNA targets in quiescent cells and to enhance translation initiation mediated by HCV-like IRESes^{41–43}. There is evidence that AGO2 (a direct TNRC6A-binding partner) associates with ribosomes, functioning as a translation activator to mediate translation upregulation⁴⁴. Recruitment of TNRC6A to AGO2/miRNA complex typically triggers a cascade of events for translation suppression and target mRNA degradation; however evidence of TNRC6A interaction with poly (A) binding proteins (PABP) also suggests a role in translation ⁴⁵⁻⁴⁸. Our findings, together with reports of active translation associated with FUS granules ⁴⁹, point to a more prevalent role of cytoplasmic RNP granules in active local translation than previously suspected.

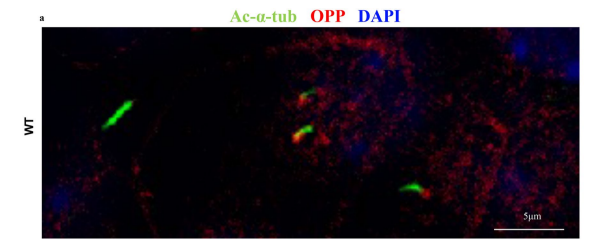
Recent studies have identified TNRC6A or AGO2 adjacent to the basal body of primary cilia or centrosomes^{50,51}, suggesting that the spatial relationship between TNRC6A/AGO2 positive granules and centrioles is conserved in both primary cilia and multicilia (as we report here). We found newly-synthesized proteins (OPP) also concentrated in granules at the base of primary cilia (Fig. 10). However, primary cilia remained intact in cultured *Tnrc6a* null cells in contrast to our findings in multicilia.

The precise mechanism by which TNRC6A promotes mRNA translation is under investigation. Based on our current observations, we propose that induction of Tnrc6a in differentiating MCCs and their accumulation in the apical granules with AGO2 and miRNAs, such as miR-34/449, lead to recruitment of target mRNAs key for multiciliogenesis, as exemplified here by those encoding centriolar proteins. The lack of mRNA degradation enzymes in the TNRC6A granules allows stabilization of these mRNA targets for local translation. Indeed, TNRC6A is known to directly bind to poly(A) binding proteins (PABP) ^{47,48}, which recruit elongation factors to initiate protein synthesis ^{52,53}. Moreover, AGO2, a binding partner of TNRC6A, directly interacts with eIF1A ⁴⁰ and presumably other elongation factors recruiting them to these apical granules. The presence of OPP and elongation factors in the TNRC6A-containing apical granules support the idea that these are site of local newly-synthesized proteins. The key role for Tnrc6a in this process was seen by the selective loss of the apical OPP-labeled foci when these granules were abolished in Tnrc6a-deficient cells (while OPP was maintained elsewhere in these cells.

In summary, our findings suggest that components of the miRNA pathways can be organized into subcellular hubs for target mRNA localization, stabilization and local translation crucial for differentiation of airway MCCs. These observations further provide strong evidence of the critical role of mRNA localization for efficient delivery and concentration of subsets of proteins to specific intracellular sites for prompt organelle biogenesis during cellular differentiation.

Material and methods

Mouse breeding, genotyping and monitoring. *Tnrc6a*^{gt/+} reporter mice carrying a gene trap insertion of the β -galactosidase gene in the *Tnrc6a* locus and Tnrc6a^{gt/gt} mutants were generated on a mixed genetic background as described¹⁸. Tnrc6a mutants were backcrossed to C57BL/6 for ten generations to overcome embryonic lethality. Mice are housed in a non-barrier animal facility at Columbia University Medical Center. The genotyping is performed as reported¹⁸. miR-34/449 null mice were bred and genotyped as described²³. Foxj1/EGFP/Tnrc6A mice were generated by breeding Tnrc6A^{gt/+} mice to Foxj1-EGFP mice (Jax, B6; C3-Tg (Foxj1-EGFP)85Leo/J) ⁵⁴. All animal studies were carried out with the approval of the Institutional Animal Care and Use Committee (IACUC), Columbia University Medical Campus (WVC IACUC #: AC-AABF2567).



Ac-α-tub EGFP DAPI

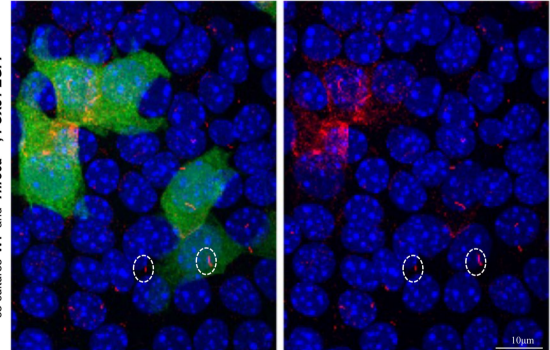


Figure 10. Disruption of *Tnrc6a* does not prevent primary cilia from forming. IF of ALI airway epithelial cultures at day 3. (**a**) WT ALI airway epithelial cultures: newly-synthesized proteins (red, label ed by OPP) are concentrated in foci at the base of primary cilia (green). (**b**) Co-culture of WT and Tnrc6a null; FOXJ1-EGFP cells at ALI day 3: *Tnrc6a* null cells marked by FOXJ1-EGFP (green) showing no significant defect in primary cilia marked by Ac-α-tub staining (circled).

Cloning and packaging of lentiviral vectors. pHAGE-EGFP-Tnrc6a vector was cloned by replacing EGFP fragment of pHAGE-CMV-eGFP-W1 vector (generous gift from Dr. Darrel Kotton, Boston University Medical School) with EGFP-Tnrc6a DNA fragment from pT7-EGFP-Tnrc6a (Addgene #25035) plasmid. To do this, we first cut pHAGE-CMV-EGFP-W1 with NotI (NEB # R0189S), followed by the treatment with DNA Polymerase I, Klenow Fragment (NEB # M0210S) to blunt the end. Then EGFP fragment was released by cutting with BamHI (NEB # R0136S), followed by separation and purification of vector fragment for ligation. To prepare EGFP-Tnrc6a fragment, P7-EGFP-Tnrc6a was cut with AgeII (NEB # R0552S), followed by the treatment with DNA Polymerase I, Klenow Fragment (NEB # M0210S) to blunt the end. Then cut again with BamHI to release and purify EGFP-Tnrc6a fragment. DNA sequencing was performed following ligation and transformation to identify and confirm the right clone. Similarly, EGFP-Ago2 fragment was released from EGFP-Ago2 plasmid (EGFP-C1 backbone, addgene, # 21981) by AgeI, blunted and followed by BamH1 digestion. The same fragment of lentiviral vector used in pHAGE-EGFP-Tnrc6A cloning was used for the ligation to generate pHAGE-EGFP-

co-cultures WT and Tnrc6anuli; FOXJ1-EGFP

Ago2. Packaging of all lentiviral vectors was performed by co-transfection of 293 T cells (80% confluent, 15 cm plate) with 2 ml of Trans-IT/DNA/media mix containing 5 plasmids encoding the lentiviral backbone (24ug) as well as tat (1.2ug), rev (1.2 μ g), gag/pol (1.2ug) and vsv-g (2.4ug) viral genes together with 90 μ l of Trans-IT (MIR #6600). Viral particles in the resulting supernatants were collected 48 h after transfection and concentrated by ultracentrifugation (90 min at 48960g on Beckman SW28 rotor). Titers used in all experiments range from 0.5–1.5 × 10⁹TU/ml. For the constructs above, lentiviral infection of basal cells isolated from mouse trachea was performed at the time of seeding and the culture were terminated at Day 3 of the differentiation, when the centriologenesis and ciliogenesis are highly active.

Histological analyses. Tissues were dissected and fixed overnight in in 4% paraformaldehyde (Ted Pella, #18505) overnight at 4 $^{\circ}$ C, then processed either for frozen section or paraffin section by following standard procedures. All the blocks were sectioned at 10 μ m either for immunostaining or for in situ hybridization.

β-Galactosidase staining. This was performed according to protocol described previously¹⁸. Briefly, adult lungs of Tnrc6a^{gt/+} mouse were perfused intratracheally with 0.25% glutaraldehyde (Electron Microscopy Science, #16220) at a constant pressure of 30 cm H₂O for 30 min, then perfused with 1XPBS for 2 h, followed with staining buffer (5 μm C6N6FeK3, 5 μm C6N6FeK4, 2 mM MgCl2, 1 mg/ml X-gal in 1XPBS) for 30 min. Then the trachea was tied and the whole lungs were incubated in the staining buffer in a 50 ml tube overnight at 37 °C. After washing in 1XPBS three times (2 h for each wash), lungs were fixed again using 4% PFA (Ted Pella, #18505) for 2 h and then processed for embedding following standard procedures. Sections of LacZ-stained lungs were either subjected to H&E staining or co-stained with antibodies specific for MCCs.

In situ hybridization (ISH). ISH detection of miR-34a and miR-449a and miR-92b or Tnrc6a mRNA expression was performed on frozen sections of mouse lungs using 5' digoxigenin-labeled LNA probes (Exiqon) or digoxigenin-labeled antisense probe of Tnrc6a. Briefly, slides were washed with DEPC-treated 1X PBS 3 times for 3 min each, followed by the treatment with 100 mM triethanolamine buffer (Sigma, Cat #90279) plus 0.25% of acetic anhydride (Sigma, #320102) for 10 min, then permeabilized by incubating in PBST (1X PBS plus 0.1% Triton X-100 (Sigma, #9002-93-1) in DEPC-treated water for 30 min, and washed 3 times for 5 min each at RT in 1X PBS. After 2 h pre-hybridization using hybridization buffer without labeled probe (50% formamide (American Bioalnalytical, # 95-12-7), 10 mM Tris-HCl pH8.0, 600 mM NaCl, 1X Denhardt's solution (Sigma, #02532), 200 µg/mL tRNA, 1 mM EDTA, 0.25% SDS (Ambion, #AM9823), 10% dextran sulfate (American Bioalnalytical, # ab427), hybridizations were carried out at 55 °C overnight in the hybridization buffer containing 25 nM of DIG-labeled miR-34a (#38487-01, ACAACCAGCTAAGACACTGCCA) or miR-449a (Exiqon, #38587-15, ACCAGCTAACAATACACTGCCA) or miR-92b (Exiqon, #610819-340, CTGCACCGCGTCCCGTC). Then, slides were sequentially washed with SSC (2X, 1X, 0.2X) at 50 °C for 30 min. followed by 0.2X SSC (Ambion, #AM9763) for 5 min at RT, and finally 1X PBS for 5 min at RT. Slides were then incubated in blocking solution (TTBS, 0.05 M Tris, pH 7.5, 0.15 M NaCl, 0.1% Tween-20, plus 5% sheep serum) and incubated with anti-digoxigenin-AP, Fab fragments (1:500, Roche, Cat. #11093274910) overnight at 4 °C. After washing in TTBS 3 times for 10 min each, signals were developed using BM purple (Roche, # 11442074001). Hybridization for Tnrc6A using digoxigenin-labeled antisense probe was performed at 70 °C by following the standard procedure. ISH of Spef2 and Rsph1 was performed by RNAscope according to manufacturer's protocol.

Immunofluorescence (IF) and immunohistochemistry (IHC). For IHC, paraffin sections were deparaffinized and rehydrated by sequentially incubating sections in three washes of xylene (5 min each), two washes of 100% ethanol (10 min each), two washes of 95% ethanol (10 min each), followed by two washes in dH_2O (5 min each). Endogenous peroxidase was quenched by incubation with 0.3% $H_2O_2/0.3$ % horse serum in 1X PBS for 5 min at RT. Then, slides were washed in TNT buffer (0.1 M Tris pH 7.5, 0.15 M NaCl, 0.05% Tween-20) 3 times (5 min each), followed by blocking with TNB buffer (0.1 M Tris pH 7.5, 0.15 M NaCl, 0.5% blocking reagent) for one hour at RT. Incubation with Ac-a-tub antibody (Sigma, #T7451, 1:10,000) diluted in TNB buffer overnight at 4 °C. Slides were then washed with TNT buffer 3 times, incubated with Horseradish peroxidase (HRP) conjugated secondary antibodies (Life Technologies) for 2 h at RT, signals were developed by using the DAB staining kit (Vector Laboratories, Cat.# SK-4100). When TNRC6A monoclonal antibody (Santa Cruz, 4B6, # sc-56314) was used for immunohistochemistry, TSA Plus Fluorescence Kit (Perkin Elmer, # NEL741B001KT) was used for signal amplification. After incubating with HRP-conjugated secondary antibody, signals of slides or ALi filters were developed by TSA Plus Working Solution following manufacture's protocol. For IF staining, cryosections or transwell filter of ALI cultures were washed in PBS, and then blocked by 5% donkey serum in PBST (1XPBS, 0.3% Triton X-100) for one hour at RT, followed by incubation with primary antibodies overnight at 4 °C (TNRC6A, human index serum, 1:3000; FOXJ1, eBioscience, # 14-9965-82, 1:100; Ac-α-tub, Sigma, # T7451, 1:10,000, or Abcam # ab125356; DCP1A, Abcam, #ab183709, 1:50; DDX6, Cell Signaling Technology, #8988S, 1:20; EDC3, Santa Cruz, # sc-55081, 1:20; EDC4/GE-1, Cell Signaling Technology, #2548S, 1:50; Xrn1, Bethyl Laboratories, #A300-443A, 1:20; y-tubulin, Abcam, #ab84355, 1:50; EIF3B, Santa Cruz, # sc-16377, 1:50). DEUP1, Dr Xueliang Zhu's lab, 1:300; PCM1, Cell Signaling, #5213, 1:50. Alexa Fluor 488 or 568-conjugated secondary antibodies (Donkey anti-Mouse IgG, or Donkey anti-Rabbit IgG or Donkey anti-Goat IgG; Life Technologies, 1:250 dilution) and Alexa Fluor 594 phalloidin (Life Technologies, #A22287) were used for the visualization of signals. Zeiss 710 confocal microscope was used for the image acquisition and analysis.

Differentiation of MCCs in Air–Liquid Interface (ALI) culture of tracheal progenitors. ALI culture of mouse tracheal epithelial cells (mTEC) were performed using basal cells isolated from adult mouse

tracheas as described previously²¹. Briefly, tracheas from adult (8- to 12-weeks old) mice were dissected and cleaned in ice-cold Ham's F-12 medium (Life Technologies, #31765-092) with 1% penicillin/streptomycin (Life Technologies, # 15140-163). To isolate epithelial cells, tracheas were digested in 1.5 mg/ml Pronase (Roche, #10165921001) in Ham's F-12 media overnight at 4 °C. The digestion was stopped by fetal bovine serum (FBS, sigma, #12133C, a final concentration of 10%). The epithelial cells including basal cells were dislodged from trachea by gentle inverting the tube, and medium containing epithelial cells was transferred to another tube. This was repeated for three time and medium containing epithelial cells were pooled and Centrifuge at 500×g, 4 °C, for 10 min. After removing the medium, epithelial cells were treated with DNase (200 µl per trachea) on ice for 5 min, and spin down again at 500×g, 4 °C. The pelleted cells were washed by pre-warmed mTEC/Basic medium containing advanced DMEM/F-12 (Life Technologies, #12634-028), 15 mM HEPES (Life Technologies, #15630-080), 1% penicillin/streptomycin, 4 mM glutamine (Life Technologies, #35050-061), 3.6 mM sodium bicarbonate (Life Technologies, #25080-094), and 0.25ug/mL Amphotericin B (Life Technologies, #15290-018), and centrifuged at 500×g for 5 min. Then cells were resuspended in mTEC/Plus medium, which contained mTEC/ Basic medium with 5% FBS, 25 ng/ml Epidermal Growth factor (EGF) (Corning, #354001), 10 µg/ml Insulin (Sigma, #I6634), 5 µg/ml Transferrin (Sigma, #T1147), 0.1 µg/ml Cholera Toxin (Sigma, #C8052), 30 µg/ml Bovine Pituitary Extract (BPE)(sigma, #P1476), and freshly added 50 nM retinoic acid (RA, Sigma, #R2625), and plated on rat tail type I collagen (Advanced Biomatrix, #5056) coated trans-wells (Corning, #3470) of 24-well plates. 10 µM Rock inhibitor Y27632 (Tocris, # 1254) was added to mTEC/Plus only once when the cells were plated. The mTEC/Plus medium was changed every other day for 7 days, until we switched the culture to air-liquid interface culture by removing the media in the apical chamber, and replacing the media in the basal chamber with mTEC/SF medium. The mTEC/SF medium was changed every other day until the culture was terminated at different time point. Cultures were terminated at different time point after the induction of differentiation and harvested for RNA isolation or Immunofluorescence. For IF, cultures filters were washed with room temperature PBS twice, then fixed by 4% PFA for 15 min, followed by three time of washes in PBS, and keep them in PBS at 4 °C.

Single-molecule fluorescence in situ hybridization (smFISH). mTEC epithelial cultures were fixed for 15 min in 4% PFA, followed by three washed in PBS then transfer to 70% ethanol for storage. In situ hybridization and imaging were performed as previously reported²⁸. Fluorescence labeled DNA oligo probes are from Itzkovitz's Laboratory. All probes are labeled with Cy5 except rRNA 18s (TMR).

O-Propargyl-puromycin (OPP) labeling and imaging. mTEC cultures were treated with either 50 μ g/ml Cycloheximide (Sigma-Aldrich, C7698) or DMSO control for 15 min, followed by 30 min incubation with 50 μ M OPP (Jena Bioscience, NU-913-5). Cultures were then washed with room temperature PBS and fixed with 4% of PFA, followed by three washes in PBS. OPP features an alkyne group that can be detected and imaged by the addition of azide-coupled fluorophores in the highly specific click reaction^{27,28}. The click reaction was performed subsequently to the IF staining by making use of the Click-iT Alexa Fluor 488 or Click-iT Alexa Fluor 647 Imaging Kits (Thermo Fisher Scientific, C10337 and C10340). The kits were used according to the manufacturer's instruction with the exception of an additional 20-fold dilution of the working solution of the Alexa Fluor azide dye in DMSO.

Immunofluorescence (IF) signal quantitation and statistical analysis. To quantify the IF staining of newly synthesized proteins (OPP) and other centriolar proteins, Z-stack of confocal images were taken for the entire cells. For OPP staining, several planes covering the entire OPP foci were chosen and the maximal projection images of these planes were acquired. Then the areas encompassing the OPP foci and background were marked using ImageJ and the mean optical intensity was calculated. For centriolar proteins, we first generated the maximal projection image for entire cells, and drew the cell boundaries. Then the mean optical intensities were measured by ImageJ. Data were presented as the mean \pm SEM and subjected to 2-tailed, paired Student's t test. P < 0.05 was considered significant. In all experiments a minimum of 3–4 replicates (ALI cultures or lung sections) were used for analyses.

Scanning Electron Microscopy (SEM). Tracheas from E18.5 WT or Tnrc6a^{gt/gt} embryonic lungs were dissected, fixed in 2.5% glutaraldehyde, 0.1 M sodium Cacodylate, 0.2 M Sucrose, 5 mM MgCl2 pH 7.4 and dehydrated through a graded series of ethanol dilutions. Samples were processed using liquid carbon dioxide in a Tousimis Samdri 795 Critical Point Drier (Rockville MD), sputter coated with chromium in a Quorum EMS 150 T ES (Quorum Technologies Ltd, United Kingdom) and examined in a Zeiss Supra Field Emission Scanning Electron Microscope (Carl Zeiss Microscopy, LLC North America), using an accelerating voltage of 2 kV. For quantitative analysis, lungs (n = 2) were examined and about 120 cells were counted for each sample.

Data availability

All procedures and experiments were performed in accordance with the relevant guidelines, regulations and protocols approved by CUIMC (Institutional Animal Care and Use Committee approval: WVC IACUC #: AC-AABF2567). All methods are reported in accordance with ARRIVE guidelines. Data referenced in this study from transcriptomics of control and Tnrc6a null mice in lung (E18.5) and yolk sac (E9.5) are available in the National Center for Biotechnology Information Gene Expression Omnibus (GEO) and are accessible through the GEO Series accession number GSE89327, GSE30244 and GSE89327. The data is available in link https://drive.google.com/drive/folders/15JZ_G_kxdZSpjS9dw1tIHfZEds-dUNwk?usp=sharin. All other relevant data are available from the corresponding author upon request.

Received: 10 January 2023; Accepted: 28 April 2023 Published online: 01 May 2023

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Acknowledgements

We thank Jun Qian for technical assistance; Dr. Andreas Moor and Dr. Shalev Itzkovitz from the Department of Molecular Cell Biology at Weizmann Institute of Science, Israel, for sharing with us probes for smFISH. We also thank the Lu and Cardoso lab members for thoughtful discussions and input. This study is supported by Grants from NHLBI (R01 HL131422 to J.L and W.V.C and R35-HL135834-01 to W.V.C).

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Competing interests

The authors declare no competing interests.

Additional information

Supplementary Information The online version contains supplementary material available at https://doi.org/ 10.1038/s41598-023-34365-8.

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