Sensory primary cilium is a responsive cAMP microdomain in renal epithelia

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Primary cilia are hair-like cellular extensions that sense microenvironmental signals surrounding cells. The role of adenylyl cyclases in ciliary function has been of interest because the product of adenylyl cyclase activity, cAMP, is relevant to cilia-related diseases. In the present study, we show that vasopressin receptor type-2 (V2R) is localized to cilia in kidney epithelial cells. Pharmacologic inhibition of V2R with tolvaptan increases ciliary length and mechanosensory function. Genetic knockdown of V2R, however, does not have any effect on ciliary length, although the effect of tolvaptan on ciliary length is dampened. Our study reveals that tolvaptan may have a cilia-specific effect independent of V2R or verapamil-sensitive calcium channels. Live-imaging of single cilia shows that V2R activation increases cilioplasmic and cytoplasmic cAMP levels, whereas tolvaptan mediates cAMP changes only in a cilia-specific manner. Furthermore, fluid-shear stress decreases cilioplasmic, but not cytoplasmic cAMP levels. Our data indicate that cilioplasmic and cytoplasmic cAMP levels are differentially modulated. We propose that the cilium is a critical sensor acting as a responsive cAMP microcompartment during physiologically relevant stimuli.

The cilium is a dynamic structure that acts as a mechanosensory appendage of the cell and is involved in the pathogenesis of polycystic kidney disease (PKD)¹–⁴. The hair-like cilium extends from epithelial lining cells into the lumen and bends upon fluid shear stress, triggering an influx of calcium³–⁷. It has previously been shown that the functional efficiency of the cilium depends on the length; i.e., longer cilia can detect a lower magnitude of shear stress and elicit a larger increase in calcium influx⁸–¹⁰. Structural defects of the cilium, such as in intra-flagellar transport protein 88 (IFT88) deletion resulting in shortened cilia, cause ciliopathy phenotype in PKD mouse models¹. Likewise, functional defects of the cilium, such as dysfunctional polycystins proteins (PC1 or PC2), cause polycystic kidney disease⁷–⁴,¹¹,¹². Both PC1 and PC2 form a mechanosensory complex in the cilium, which upon bending modulates calcium fluxes¹³,¹⁴.

Similar to calcium, cyclic adenosine monophosphate (cAMP) is a second messenger involved in different signaling pathways. Studies have shown that increasing cAMP levels in cystic epithelial cells, via either adenylyl cyclase (AC) activation or addition of membrane-permeable cAMP analogs, enhances cyst formation and/or cyst enlargement. The cAMP-mediated cystogenic effect has been observed in in vitro cultures obtained from murine models and intact cysts excised from PKD patients¹⁵–¹⁷. Arginine vasopressin (AVP), the endogenous antidiuretic hormone, is also observed to aggravate the cystic phenotype as it raises cAMP levels by vasopressin activating receptor type 2 (V2R), a G protein-coupled receptor (GPCR). Activation of V2R induces an intracelluar cAMP increase, leading to the insertion of aquaporin 2 (AQP2) into the apical membrane and regulation of body fluid homeostasis¹⁸,¹⁹. Strategies focusing on blocking vasopressin action have shown reduced cystogenesis in murine models of PKD²⁰–²². Polycystic Kidney (PCK) rats develop progressive cystic enlargement of the kidneys and hepatic histologic abnormalities that resemble human autosomal dominant PKD²⁰–²⁵. Using this model, Wang et al. evaluated renal cyst development in the PCK rats without circulating vasopressin. PCK rats were crossed with Brattleboro (AVP−/−) rats and in the resulting PCK AVP−/− rats, renal cAMP levels were lowered along with a marked reduction in renal cysts²⁶,²⁷. The promising results of preclinical data have spurred interest in pursuing...
V2R antagonists as a prospective treatment for PKD^{23–25}. Clinical trials have shown that tolvaptan, a V2R antagonist, is effective in decreasing cyst growth and in slowing the decline of renal function^{26–32}.

Cyclic AMP is a ubiquitous second messenger, produced by AC activation upon binding of ligands to GPCR coupled to Gs. Thus, cAMP regulates a wide range of cellular processes, ranging from gene regulation to immune function^{33}. Previous studies have demonstrated that increasing intracellular cAMP levels, through the addition of forskolin or a cell-permeable cAMP analog, causes an increase in ciliary length of renal epithelia^{34,35}. Consistent with this finding, forskolin or a cell-permeable cAMP analog also increases ciliary length in vascular endothelia^{36}. Interestingly, forskolin or a cell-permeable cAMP has a minimal effect in serum-deprived synoviocytes^{37}. Furthermore, inhibition of ciliary-localized adenyl cyclase 3 (AC3) induces primary cilia elongation in synoviocytes^{37}. However, the effect of tolvaptan in modulating ciliary length through cAMP has never been evaluated.

Moreover, the dynamic level of cilioplasmic and cytoplasmic cAMP has not been previously examined with a targeted sensor. Given the potential treatment of tolvaptan and the role of cilia in PKD, we sought to characterize the effect of tolvaptan on ciliary length and ciliary cAMP signaling.

We show that tolvaptan increases ciliary length and enhances mechanosensitivity in response to shear stress. We also show that V2R and AC3 are localized in renal epithelial cilia. Monitoring cAMP levels in the cilioplasm after stimulation with tolvaptan revealed that the cilium is a distinct cAMP microdomain. Changes in cilioplasmic cAMP levels were found to be distinct and independent from cytoplasmic level of cAMP. Furthermore, shear stress decreased cilioplasmic, but not cytoplasmic cAMP below basal levels. We therefore hypothesize that primary cilia are chemosensitive and mechanosensitive organelles that form dynamic cAMP microdomain distinct from cell body.

**Results**

**Tolvaptan increases ciliary length and enhances shear-stress induced cytosolic calcium increase.** To determine if tolvaptan could alter the ciliary length, we measured the ciliary length distribution in epithelial and endothelial cells after treatment with different concentrations of tolvaptan (Supplementary Fig. S1). Tolvaptan treatment resulted in a dose-dependent increase in ciliary length in renal epithelial cells. The effect was maximal at 0.1 μM and reached a steady-state at higher concentrations (Fig. 1a,b). In renal epithelial cells, average ciliary length was 7.05 ± 0.15 μm. When treated with 0.1 μM tolvaptan, ciliary length increased to 9.34 ± 0.51 μm. In vascular endothelial cells, which generally have shorter cilia than epithelial cells, ciliary length increased from 3.67 ± 0.06 μm to 7.56 ± 0.14 μm after treatment with tolvaptan.

Both epithelial and endothelial cilia function as mechanosensory organelles by increasing intracellular calcium in response to fluid-shear stress^{46–47}. To test whether the increase in ciliary length translated to enhanced mechanosensitivity, cytosolic calcium levels in response to shear stress were measured with Fura-2 (Fig. 1c; Supplementary Fig. S1). After baseline measurement, cells were subjected to shear stress. Pre-treatment with tolvaptan significantly enhanced the calcium response to fluid flow in both epithelial and endothelial cells (Fig. 1d,e). These results show that tolvaptan increases ciliary length and enhances cilia mechanosensitivity.

The cilia are also intertwined with the cell cycle and are usually resorbed during cell division^{38,39}. We therefore examined if tolvaptan affected cell division using labeling with propidium iodide, a dye that binds stoichiometrically to DNA, to approximate the distribution of cells at different cell division stages. However, tolvaptan did not affect cell division stages in either epithelial or endothelial cells (Supplementary Fig. S1).

**Vasopressin receptor knockdown modulates ciliary length and function.** To further validate our results, we generated V2R-knockdown LL-CPK1 epithelial cells using lentiviral transfection. It has been reported that receptor activation or inhibition promotes receptor down-regulation^{38,39}; thus, V2R expression was examined in cells with or without vasopressin (10 μM) or tolvaptan (0.1 μM) treatment for 24 hours (Fig. 2a; Supplementary Fig. S2). V2R expression was compared between V2R-knockdown cells and the corresponding scrambled control cells (Fig. 2b). Ciliary length was determined in scrambled or V2R-knockdown treated cells in combination with addition of saline (control), tolvaptan, or vasopressin in live (Fig. 2c) and fixed (Fig. 2d) cells. Cilia in V2R-knockdown cells (5.23 ± 0.27 μm) appeared shorter than in scrambled control cells (6.11 ± 0.36 μm), but the difference was not significant (Fig. 2e). Tolvaptan treatment caused a significant increase in ciliary length of both control cells (11.01 ± 0.76 μm) and V2R-knockdown cells (7.67 ± 0.30 μm). However, the increase was significantly less in the V2R-knockdown compared to controls. Cells treated with vasopressin displayed a significant increase in ciliary length in scrambled control cells (8.13 ± 0.38 μm vs. 6.11 ± 0.36 μm in control), but not in V2R-knockdown cells (6.30 ± 0.25 μm vs. 5.23 ± 0.27 μm in V2R-knockdown control).

To investigate the role of V2R-knockdown in fluid-sensing, calcium imaging in response to fluid flow was performed. Although the differences in ciliary length were negligible between the V2R-knockdown and control cells, the V2R-knockdown failed to show the characteristic peak of calcium influx upon fluid flow (Fig. 2f; Supplementary Fig. S2). Previous work has also shown that the V2R is linked to calcium mobilization in rat IMCD cells, where activation of the V2 receptor results in a transient calcium increase, although the underlying mechanism has not been established^{40}.

Next, immunolocalization was performed to reveal whether V2R localizes to primary cilia (Fig. 3a). IMCD cells, which have been shown to contain V2R in the cilium, were used as a positive control^{41,42}. In LL-CPK1, V2R localization was observed throughout the cilium and in IMCD cilium V2R was detected in the base of cilium (Fig. 3a; Supplementary Fig. S3). V2R-knockdown cells showed very little membrane localization of V2R (Fig. S3). Because most endothelial cells did not show V2R localization to primary cilia (Supplementary Fig. S3), we only used LL-CPK1 in the remainder of our studies. Additionally, LL-CPK1 cells were grown on permeable supports and then used for immunolocalization. After scanning Z-stack images, XZ and XY planes were reconstructed which showed similar distribution of V2R in the apical and basolateral membrane (Supplementary Fig. S4).
To further confirm the presence of V2R in the cilia, primary cilia were isolated using shear stress. Isolated cilia from scrambled and V2R-knockdown cells were verified with a brightfield microscope. Due to a low concentration of total protein collected in the isolated cilia extract, a dot blot was used. The acetylated-α-tubulin blot was used to indicate the purity of our cilia lysate and the corresponding V2R blot established that the V2R was associated with the cilia.

Primary cilia are cAMP responsive microdomains. V2R is a Gs-coupled GPCR, which activates AC and, thereby, increases intracellular cAMP levels. AC3 has been reported to localize to neuronal primary cilia, whereas AC5/6 are expressed in primary cilia of mouse renal epithelial and LL-CPK1 cells. We analyzed AC expression in different cell lines and observed a differential expression pattern between the different AC isoforms. AC5/6 was not localized in cilia of LL-CPK1 cells, although it was observed in IMCD and endothelial cilia. In contrast, AC3 was localized in cilia of LL-CPK1, IMCD, and endothelial cells.

Because cAMP measurement has traditionally been performed with a solid-phase enzyme immunoassay, we used this technique to measure cAMP levels after tolvaptan treatment or induction of fluid-flow. Neither fluid flow nor tolvaptan treatment changed the cAMP levels, although a forskolin response was observed in both scrambled and V2R-knockdown cells. However, an immunoassay does not allow us to distinguish cilioplasmic from cytoplasmic cAMP changes. To monitor changes in ciliary cAMP levels, we used the...
ciliary-targeted cAMP sensor 5HT6-mCherry-cADDis. We first confirmed that the kinetics between cilia-specific (5HT6-mCherry-cADDis) and cytosolic (cAMP-cADDis) reporters did not alter (Supplementary Fig. S6).

Using imaging setup51, we were able to monitor both changes in the cytoplasmic and cilioplasmic cAMP levels. Vasopressin (10 μM) elicited an increase in cAMP levels in the cilium and the cell body (Fig. 5a; Supplementary Movie S1). In the V2R-knockdown cells, this response was absent in both the cytoplasm and the cilioplasm (Fig. 5b). Forskolin (5 μM) also increased cAMP levels in the cytoplasm and cilioplasm of scrambled control and V2R-knockdown cells (Fig. 6; Supplementary Movies S2 and S3). Tolvaptan (0.1 μM) induced an increase cilioplasmic cAMP levels, but no significant change was observed in the cytoplasm (Fig. 7a). Interestingly, tolvaptan also increased cilioplasmic cAMP in the V2R-knockdown cells (Fig. 7b). To exclude potential involvement of extracellular calcium influx and the roles of calcium-regulated ACs, cells were preincubated with calcium channel blocker verapamil (2 μM) before being challenged with tolvaptan. Blocking verapamil-sensitive calcium channels had no effect on ciliary cAMP increase nor impacted tolvaptan-induced ciliary length increase (Fig. 7c, Supplementary Fig. S7). In response to flow, the cilioplasmic cAMP underwent a significant decrease beyond the basal level (Fig. 8a; Supplementary Movie S4). The same response was not observed in the cytoplasm, where the cAMP remained at the basal level. Neither vasopressin (10 μM) nor tolvaptan (0.1 μM) affected the flow-induced repression of cilioplasmic cAMP (Fig. 8b,c). In the cytoplasm, cAMP levels remained stable during flow. Taken
together, certain stimuli modulated cAMP response differentially between cilioplasm and cytoplasm (Fig. 9), which supports the idea of the cilium being a distinct cAMP microdomain.

**Discussion**

Activation of V2R induces a GPCR signaling cascade, resulting in an intracellular cAMP increase. In turn, cAMP-dependent protein kinase (PKA) and other downstream signaling molecules are activated. Ultimately, this leads to the insertion of AQP2 into the apical membrane, increasing water reabsorption in the kidney collecting duct and enhancing urinary concentration in the thick ascending limb. Thus, the overall V2R function is to regulate body fluid homeostasis.

Using PCK rats (orthologous to human PKD) crossed with Brattleboro rats (without circulating vasopressin), Wang et al. found that vasopressin is a powerful modulator of cystogenesis. The PCK AVP−/− rats have lower renal cAMP level and show a marked reduction in renal cysts. Likewise, other animal studies show that inhibition of vasopressin-mediated signaling protects against cystogenesis. The preclinical results support the idea of pursuing V2R antagonists like tolvaptan as a prospective treatment for PKD. Clinical data have also shown positive results with tolvaptan significantly reducing the rate of total kidney volume increase and slowing the estimated glomerular filtration rate (eGFR) decline in patients. In April 2018, tolvaptan, under the trade name Jynarque was approved by the FDA to slow kidney function decline in adults with PKD. As a note, tolvaptan has been previously approved in 2008 by the FDA, marketed as Samsca, for treatment and prevention of hyponatremia. Due to the potential of liver injury caused by tolvaptan, there is an FDA approved Risk Evaluation and Mitigation Strategy (REMS) program that needs to be followed when tolvaptan is prescribed to PKD patients. The REMS include requirement of frequent tests for blood bilirubin-total (BT), alanine and aspartate aminotransferases (ALT and AST) levels to evaluate liver toxicity. Discontinuation of treatment can mitigate the hepatotoxic effects of tolvaptan when caught early hence emphasizing the requirement of frequent testing of liver function biomarkers.
In our study, we examined the effect of tolvaptan on ciliary function because defects in ciliary function have been associated with PKD. Ciliary dysfunction that prevents sensation of fluid-flow in renal epithelia i.e. calcium fluxes generated in response to fluid-flow, contributes to cyst formation. Kidney-specific inactivation of Kif3a, ...
a ciliogenic gene, in newborn mice resulted in the loss of primary cilia and produced kidney cysts\textsuperscript{61}. The loss of cilia resulted in aberrant planar cell polarity as measured by the orientation of mitotic spindles in relation to the longitudinal axis of the collecting ducts. When complemented with increased rates of cell proliferation, like in cases of acute renal injury, the aberrant planar cell polarity randomizes orientation of cell division leading to tubular dilatation and kidney cyst formation\textsuperscript{62}. Previous studies have shown that the ability of cilia to sense fluid-flow correlates with the ciliary length\textsuperscript{63}. Upon treatment with tolvaptan, ciliary length was increased in both epithelial and endothelial cells. In turn, the mechanosensitivity of the cells was increased, as shown by cytosolic calcium measurements in response to fluid-flow. Endothelial cells displayed a more pronounced amplification of the calcium signal, which could be attributed to the more pronounced increase of ciliary length after tolvaptan treatment compared to epithelial cells.

Tolvaptan is a V2R antagonist, thus a V2R-knockdown would be expected to have similar effects as tolvaptan treatment. However, our results indicate that tolvaptan increases ciliary length, whereas V2R-knockdown does not affect ciliary length. Furthermore, tolvaptan also increases ciliary length in V2R-knockdown cells. This could be due to the incomplete deletion of V2R, which allows tolvaptan to act through V2R albeit at a lower magnitude, or the action of tolvaptan is independent of V2R. To this end, we analyzed localization of V2R receptors and observed similar V2R presence in apical and basolateral membranes. We see V2R expressed throughout the cilium in LL-CPK1, as also shown by Raychowdhury et al.\textsuperscript{49}, and in the basal bodies of primary cilia in IMCD renal epithelial cells. The exact mechanism by which tolvaptan increases ciliary length and function is not clear at present. However, our data indicate that such a mechanism very likely involves ciliary cAMP. The most recent study indicates that tolvaptan can inhibit potassium (K\textsuperscript{+}) channels\textsuperscript{64}. Thus, the potential of multiple targets or mechanisms of action of tolvaptan remains a possibility.

We performed functional studies to compare cAMP levels in V2R-knockdown cells in response to V2R agonist and antagonist. In our cilia imaging experiments, vasopressin, forskolin and tolvaptan increased cilioplasmic cAMP levels. Vasopressin stimulated cilioplasmic cAMP increases were absent in V2R-knockdown cells, whereas tolvaptan elicited a cilioplasmic cAMP increase independent of V2R. We propose that while vasopressin requires the presence of V2R to engage signaling in the cilium, tolvaptan acts in a V2R-independent manner on ciliary signaling.
Previous studies have shown that the ciliary V2R is coupled with a functional AC that increases cAMP levels in cilia upon vasopressin binding. This is particularly interesting since localized cAMP responses could regulate ciliary function without stimulating global increases in cAMP that would affect cell proliferation in cystic epithelia. V2R activation stimulates AC activity and studies have shown AC isoform 5/6 to be present in the cilia. To verify this and discover if any other isoforms are present in the cilium, we immunostained for all AC isoforms. To our surprise, we found AC3 in the cilia of LL-CPK1 cells, but there was no cilia localization of AC5/6 as previously reported. AC3 localization has been reported in neuronal cilia. AC3 is one of the three CaM-sensitive AC isoforms although the effect is not as pronounced as seen in CaM activation of AC1 and AC8. Further, it is likely that AC3 is conditionally activated by calcium/calmodulin (CaM) and also by PKC but inhibited by calcium/calmodulin-dependent protein kinase II (CaMK-II). Studies have shown CaM localization at the basal body as well as throughout the cilia in a punctate pattern. In a 2011 study, Rothschild et al. showed that CaMK-II is present in zebrafish cilia and also by PKC but inhibited by calcium/calmodulin-dependent protein kinase II (CaMK-II). Morpholino mediated suppression of CaMK-II in zebrafish embryos induced hydrocephaly and pronephric cysts, and destabilized cloacal cilia. Furthermore, they establish that PKD2 and CaMK-II deficiencies are synergistic, where CaMK-II is a crucial effector of PKD2 Ca2+ required for morphogenesis of the pronephric kidney and stabilization of primary cloacal cilia. This suggests that CaMK-II might play a key role in ciliary function and response to physiological stimuli.

Cyclic AMP signaling has the potential to regulate various pathways and cell functions but when restricted in a compartmentalization model, cAMP fluxes can result in specific responses. Based on our cAMP measurements, we noted that changes in cilioplasmic cAMP is independent from cytoplasm. We also found that while vasopressin and forskolin increase both cilioplasmic and cytoplasmic cAMP, tolvaptan and fluid-flow result in a discrete change only in cilioplasmic cAMP. In response to flow, the cilioplasmic cAMP underwent a significant decrease beyond basal levels. Pre-treatment with either vasopressin or tolvaptan does not affect the flow-induced cAMP decrease in the cilia, indicating the complexity of cAMP signaling within cilioplasm. These results support the idea that the cAMP signaling occurs in a compartmentalized manner such that numerous signaling pathways can utilize this biochemical signal and still achieve distinct cellular effects (Fig. 9).
The concept of cAMP being synthesized in specific microdomains and spatially regulated is of growing interest. Studies with membrane microdomain specific sensors show that in addition to the local phosphodiesterases, which control the levels of cAMP, there are specific arrangements of the cAMP signaling components. Scaffold proteins, such as A-kinase anchoring proteins (AKAPs) and perhaps others, bring together GPCR, AC isoforms, phosphodiesterases and protein kinase A to create signaling complexes that regulate spatially constricted cAMP signaling cascades. The small cilia organelle enables enrichment of proteins with respect to the rest of the plasma membrane and may also restrict diffusion of second messengers with a transition zone that separates the cilioplasm from the cytoplasm. These features make the cilia an effective signaling microdomain with the ability to generate a high local concentration of second messengers and effectors proteins. Studies

Figure 7. Tolvaptan increases cilioplasmic cAMP. Representative time-lapse images of 1/cADDis, mCherry channels and Ratio (mCherry/cADDis) with their corresponding line graphs showing cAMP levels in response to tolvaptan (Tvp) in (a) scrambled, (b) V2R-knockdown, and (c) verapamil-treated cells. Color bar on the left indicates cAMP level from low (black) to high (green) for 1/cADDis channel. CAMP level from low (black) to high (yellow) are shown on the right color bar for the ratio images. Corresponding individual raw traces (red) and average signal (blue) are plotted in line graphs for 1/cADDis, mCherry and ratio signals in the cilioplasm and cytoplasm. Arrows indicate stimuli. N = 4 for each group.
have shown the presence of GPCR, AC isoforms and cAMP-mediated pathways present in the cilium. It is plausible to contemplate that cAMP could play a role in the dynamic modulation of the cilia, which is linked to its functional efficiency.

In conclusion, we find that tolvaptan can modulate ciliary length and function. Interestingly, tolvaptan does not fit the mold of a traditional antagonist, at least not within the ciliary structure. The prospect of tolvaptan working through an off-target effect in the cilia might open new strategies targeting the cilia and PKD relationship. In studying effects V2R/tolvaptan in cilioplasmic cAMP, our data further establish the cilium as a cAMP microdomain that responds to physiologically relevant stimulus.

**Figure 8.** Flow decreases cilioplasmic cAMP level. Representative time-lapse images of 1/cADDis, mCherry channels and Ratio (mCherry/cADDis) in response to fluid-flow (1.0 dyne/cm²) after treatment with (a) vehicle control, (b) 0.1μM tolvaptan (Tvp) or (c) 10μM vasopressin (AVP). Color bar on the left indicates cAMP level from low (black) to high (green) for 1/cADDis channel. cAMP level from low (black) to high (yellow) are shown on the right color bar for the ratio images. Corresponding individual raw traces (red) and average signal (blue) are plotted in line graphs for 1/cADDis, mCherry and ratio signals in the cilioplasm and cytoplasm. Arrows indicate stimuli. N = 4 for each group.
Materials and Methods

Reagents and antibodies. For cell culture, Dulbecco's Modified Eagle Medium (DMEM), trypsin, penicillin-streptomycin solution, phosphate buffered saline (PBS), Dulbecco's Phosphate-Buffered Saline (DPBS)
and were acquired from Corning (Manassas, VA). Fetal bovine serum (FBS) was obtained from Seradigm and Dulbecco’s phosphate buffered saline (DPBS) from HyClone (Logan, UT). Sucrose, Triton-X and Adenosine 5’-triphosphate Disodium (ATP) Salt were purchased from Fisher Scientific (Fair Lawn, NJ) and paraformaldehyde (PFA) from Electron Microscopy Services (Hatfield, PA). Propidium iodide (PI) from obtained from Biotium (Fremont, CA), forskolin from BioVision (Milpitas, CA), verapamil (Ver) from Sigma-Aldrich (Milwaukee, WI), tolvaptan (Tvp) from AK Scientific (Union City, CA) and arginine vasopressin (AVP) was purchased from Bachem (Torrance, CA). Laemmli 2X sample buffer was obtained from BioRad (Hercules, CA), cComplete Protease Inhibitor Cocktail from Roche (St. Louis, MO) and Western blot visualization kit was obtained from Thermo Scientific (Rockford, IL). Nonfat dry milk was purchased from LabScientific (Livingston, NJ). Primary antibodies, acetylated-α-tubulin was acquired from Abcam (Cambridge, MA), β-actin from Cell Biosabs (San Diego, CA) and V2R antibody from EMD Millipore (Billerca, MA). Previously validated adenyl cyclase 2, 3, 4, 5, 6, 7, 8 and 9 antibodies were obtained from Santa Cruz Biotech (Santa Cruz, CA) [80]. The secondary antibodies, fluorescein anti-mouse, texas-red anti-rabbit and mounting media with DAPI were purchased from Vector Laboratories (Burlingame, CA).

**Cell culture.** Porcine renal epithelial cells from proximal tubule (LL-CPK1), dog epithelial cells from inner medullary collecting duct (IMCD), and mouse vascular endothelial (ET) cells were cultured to a confluent monolayer in DMEM supplemented with 10% FBS at 37°C in 5% CO2. In some experiments, LL-CPK1 cells were also grown on Corning Transwell permeable supports to induce polarization for receptor localization studies and allow antibody access to the basal membrane. ET and LL-CPK1 cells have been previously described in detail [31]. Once differentiated, different concentrations of tolvaptan or vasopressin was added to culture plates. Concentration for tolvaptan was determined to be 0.1 μM based on the optimal ciliary length increase based on dose-response studies, whereas a vasopressin concentration of 10 μM was selected because it maximally increases cAMP levels [82]. Verapamil was added to cells at a final concentration of 2 μM for 10 minutes before drug treatment [83]. The drugs were mixed in starvation medium (DMEM with 2% FBS) and cells were incubated for another 20 hours. Vehicle alone (PBS containing 0.0005% DMSO) was used as a control groups to account for the DMSO concentration in both tolvaptan and vasopressin working solutions.

**FACS analysis.** Florescent-activated cell sorting (FACS) was used to investigate a possibility of an effect on cell division by tolvaptan (0.1 μM). Cells were harvested with and without drug treatment. Cells were then fixed using 70% ethanol and incubated with propidium iodide (PI), a DNA-intercalating fluorescent molecule, for 30 minutes at 37°C. Cell division analysis was carried out with flow cytometry BDFacsvsi with BD FACsuite software.

**Immunofluorescent staining.** Cells were fixed for 10 minutes (4% PFA/2% sucrose in PBS) and permeabilized for 5 minutes (10% Triton X-100). Acetylated α-tubulin (1:10,000 dilution in PBS) and fluorescein isothiocyanate (FITC)-conjugated secondary (1:1000 dilution in PBS) antibodies were each incubated with the cells for 1 hour at 37°C. For V2R visualization, V2R antibody (1:10,000 dilution in PBS) and texas-red conjugated anti-rabbit antibody were applied for 1 hour each at 37°C. All adenyl cyclase (AC) antibodies were used at a 1:1,000 dilution, and incubated with appropriate texas-red conjugated secondary antibody for 1 hour each at 37°C. Slides were then mounted with dapi hard set mounting media (VectorLabs, Burlingame, CA). Nikon Eclipse Ti-E inverted microscope or Nikon Confocal microscope with NIS-Elements imaging software (version 4.30) was used to capture images of primary cilia. Automated image acquisition was conducted in large image scan mode at 100x magnification and Z-stacks of 0.1 μm thickness. All images are shown as maximum intensity projections except confocal generated images or stated otherwise.

**Flow-induced cytosolic calcium imaging.** Cells were grown on glass-bottom plates. After treatment with the appropriate drug, the cells were incubated with 5μM Fura-2 AM (TEFLabs, Austin, TX) at 37°C for 30 minutes. After washing with DPBS, the cells were observed under a 40 × objective lens with a Nikon Eclipse Ti-E microscope. Fura-2 fluorescence images at excitation of 340/380 nm and emission of 510 nm were recorded. After equilibration under the microscope for 20 minutes, baseline calcium was recorded for 2 minutes and experimental data were acquired. Fluid-shear stress was then applied to cells with an Instech P720 peristaltic pump. The perfused fluid was pumped into the cell culture dish and retained at a shear stress of 1 dyne/cm² (for epithelial cells) or 8 dyne/cm² (for endothelial cells) with a constant flow rate of about 20 or 160 μl/sec, respectively. At the end of each experiment, the maximum calcium signal was obtained by perfusion of ATP (10 μM) to confirm cell viability. In addition to autofluorescence, both minimum and maximum fluorescence was collected as previously described [84]. Conditions for all experiments were maintained at 37°C and 5% CO2; in a stage top cage incubator (okoLab, Burlingame, CA). Calcium analysis was then followed a standard calculation as previously described [84].

**Generation of V2R-knockdown in renal epithelial LL-CPK1 cells.** Arginine vasopressin receptor type-2 (AVPR2 or V2R) targeting shRNA construct (5’-ATC GCC TTG ATG GTG TTT GTG GCA CCT GC-3; pAVP2Ra-C-shLenti) in a lentivirus backbone vector (TL513450) was ordered from Origene. Viral particles were generated in HEK cells using shRNA lentiviral packaging kit (TR30022) and passed through a 0.45-micron filter to remove cell debris. For transduction, epithelial cells were incubated with the collected viral particles and Polybrene (8 μg/ml; EMD Millipore).

For Western blot, total cell lysate was analyzed by SDS-PAGE on a 10% SDS-polyacrylamide gel. After separation, a semi-dry transfer was done using Bio-Rad system. The anti-V2R (1:1000 dilution) and anti-β-actin (1:500 dilution) antibodies were incubated with the PVDF (Polyvinylidene difluoride) membrane. Each incubation was done at 4°C overnight. For visualization, horse radish peroxidase (HRP)-conjugated secondary antibodies anti-mouse (1:1000 dilution) or anti-rabbit (1:1000 dilution) were used. For dot blot experiment, cell or ciliary
proteins were extracted in RIPA buffer with protease inhibitor. A 3 µl of sample extract was spotted onto a nitrocellulose membrane. After drying the membrane, it was blocked in 5% BSA (bovine serum albumin; Promega) for 1 hour. Incubation with V2R antibody (1:1000 dilution) was done for 30 minutes at room temperature. The membrane was then incubated with an HRP conjugated anti-rabbit antibody for 30 minutes at room temperature. For both Western and dot blots, the membrane was incubated with ECL (Enhanced Chemiluminescent; Thermo Scientific) for 1 minute. Visualization of the protein signal was done in Bio-Rad Image analysis and imager.

Cilia isolation. Cells were plated on 100 mm dishes (6 for each cell line) with 10% FBS supplemented DMEM at 37 °C in 5% CO₂. Cells were grown for 7 days and were starved overnight for differentiation as outlined above in cell culture methods. Cells were rinsed gently with PBS for a brief period and 10 ml of PBS was added. The dish was then placed on a rotary shaker (Thermo Fisher Scientific MaxQ 2508) and shaken for 4 minutes at 360 rpm, resulting in a shear stress of 10 dyn/cm². PBS was collected and transferred to a 50 ml centrifuge tube and spun for 10 minutes at 1,000 × g at 4 °C. After discarding the pellet, the supernatant was spun down in an ultracentrifuge (Thermo Fisher Scientific Sorval WX 100 + Ultracentrifuge) with a fixed angle rotor (Thermo Fisher Scientific, T-647.5) at 40,000 × g for 45 minutes at 4 °C. After discarding the supernatant, the remaining pellet containing primary cilia, was resuspended in either RIPA buffer or resuspension buffer as previously discussed84,85.

cAMP imaging and quantification. Intracellular cyclic AMP (cAMP) was quantified using ELISA and live-cell imaging. The cAMP ELISA Kit from Cayman Chemical (Cat. No. 581001) was used to measure total cAMP from cell population. Samples were lysed according to manufacturer instructions in a 96-well plate format. After primary incubation for 18 hours, the plate was developed and read at 415 nm on a spectrophotometer.

For cAMP live-cell imaging, we tested both pc3.1-SSTR3-mICNBD-FRET and 5HT6-mCherry-cADDis. The generation of pc3.1-SSTR3-mICNBD-FRET and 5HT6-mCherry-cADDis have been previously described85,86. While both constructs worked fine in response to forskolin, we chose to use 5HT6-mCherry-cADDis for the rest of our studies to obtain a relatively better transfection in our cell line. Of note is that pc3.1-SSTR3-mICNBD-FRET was introduced to the cells with chemical transfection (jetPrime; Polyplus), whereas Baculovirus transduction was utilized to express 5HT6-mCherry-cADDis (#D0211G; Montana Molecular). 2 × 10⁵ cells were plated in 6 well plates and grown in conditions described above for 3 days. Baculovirus mediated transduction was then performed using a final concentration of 5 × 10⁵ VG (viral genes)/ml and 4 mM sodium butyrate. 12 hours after initiation of viral transduction media was replaced with starvation media and incubated for another 16 hours before imaging. To enable us to monitor cilia and cell body concurrently, we used side-view imaging as previously described84. After infection, baseline fluorescence was examined for each cell. Those cells expressing the cAMP sensor in the cilia were used in our studies. In addition to cilia specific cAMP sensor, a global cytosolic sensor (#U0200R; Montana Molecular) was also used to validate our imaging system.

Fluorescence readings were taken on individually for mCherry and cADDis (cAMP difference detector in situ) at excitation/emission wavelengths of 590/610 and 490/510 nm, respectively. A set of mCherry and cADDis images were captured at 0.8 frames per second. The mCherry signal was used to correct fluorescence artifact. Because cADDis is a downward cAMP reporter with a constant mCherry fluorescence, the mCherry/cADDis ratio therefore depicted intracellular cAMP level and was used to quantify changes in cilioplastic and cytoplasmic cAMP. For ease of understanding, the cADDis signal was inverted to represent a direct relationship between cAMP level and cADDis intensity in our figures and movies (1/cADDis). Time-lapse recordings were acquired on NIS-Elements imaging software (version 4.30) and fluorescence intensities were measured for the duration of the experiment.

Data analysis. All data are reported as a mean ± standard error of mean (s.e.m). Statistical analysis was performed using ANOVA (analysis of variance) followed by Bonferroni post-hoc test. Power analysis was determined from the coefficient variant. When our coefficient variant was above 15%, the number of experimental and corresponding control groups was increased. Because in all our studies both control and experimental groups were run in parallel, our control and experimental values represented matched number of observations. In some cases, all experimental groups (including the corresponding controls) were analyzed with the post-hoc test. In other cases, only selected pairs (control vs. experimental groups) were tested. Most of our statistical analyses were performed with GraphPad Prism v.7 software. In some cases, Microsoft Excel v.15.4 was used for regression analyses. Linear regression was performed to obtain a standard calibration curve and linear equation. In this case, the analysis was done with the ordinary least squares (OLS) regression of y on x. A non-linear logarithmic regression was used to fit the sigmoidal trend-curve to show dose-response relationship. Asterisks (*) denote statistically significant differences at various probability levels (P). The P values of the significant differences are indicated in the figure legends.

When image analyses were done on fixed-specimen, images were taken at different focal planes (z-stack). If needed, a 3-dimensional image was constructed from the XY planes to validate XZ and YZ field for completion of analysis or measurement. Automated image acquisition with a thickness of 0.1 μm were taken with Nikon A1R Confocal microscope and NIS-Elements imaging software v.4.30. This system was also used for image analyses (3D object reconstruction, image segmentation, etc.). For live-imaging, Nikon Eclipse Ti-E inverted microscope and Nikon NIS Element for Advanced Research software were used for image acquisition and analyses. Our live-imaging system with controlled environmental chamber allowed more flexibility for various excitation and emission spectra from a fast wavelength exchanger DG4/5 mirror reflection system. Images were not enlarged during image analysis to avoid a false empty magnification. Ciliary length measurements were done using the measurement tool in NIS-Elements imaging software on maximum intensity projections obtained from immuno-fluorescent images. All representative images and video frames are presented with scale bars to indicate the actual image reduction size at 0.8 frames per seconds (fps). Before Fura-2 experiments, a brightfield image focused on
cell borders was obtained. Using intensity thresholds, a binary layer was created and overlaid with the Fura-2 images to create our representative images.

Free cytosolic calcium (Cyt Ca\(^{2+}\)) concentrations were calculated with the formula \(\text{Cyt Ca}^{2+} = \text{Kd} \times \frac{[\text{R} - \text{Rmin}]/(\text{Rmax} - \text{R})]}{\text{Fmax/Fmin}}\), where Kd denotes the apparent dissociation constant of the Fura-2 indicator (145 nM), R is a ratio of 510 nm emission intensity with excitation at 340 and 380 nm, and Rmax and Rmin are fluorescence intensity ratios for the calcium-bound and calcium-unbound Fura-2 with excitation at 340 and 380, respectively. We determined the Rmax and Rmin values to be stable and independent of cell type. Fmax and Fmin were the fluorescence intensity values of Fura-2 with excitation at 380 nm under the same conditions. The calcium level was radiometrically calculated. Rmin and Rmax values denote the minimum and maximum radiometric signal ratios, respectively. At the end of each experiment, the minimum fluorescence (Rmin) was obtained by incubating the cells in calcium-free solution that contained 2 mM EGTA and 10 \mu M ionomycin at pH 8.6 to optimize the ionomycin effect. After the minimum signal ratio was determined, the cells were incubated with excess calcium (10 mM) to obtain the maximum signal ratio (Rmax). Signal intensities were collected from individual cells, as well as from the whole cell population/monolayer. All the fluorescence measurements were corrected for autofluorescence. Calcium analysis was then followed a standard calculation as previously described.

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Author Contributions

R.T.S. performed the experiments, analyzed data, contributed to drafting the manuscript and oversaw the entire progress; A.M.M. contributed in generating CAMP videos; R.P. provided CAMP image capturing; D.W. provided the CAMP-FRET sensor; R.S.O. provided antibodies and CAMP expertise; S.M.N. finalized manuscript and oversaw the research project.

Additional Information

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