Correction

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The authors wish to note: “We recently discovered an error in our calculation of fluid shear stress (FSS) in our endocytosis experiments performed in ibidi 6-chamber slides. Based on the manufacturer’s guidelines, the endocytosis studies in Figs. 2, 3, and 5 and in Figs. S3 B and C were performed using a FSS of 0.1 dyne/cm² instead of the intended 1 dyne/cm². The x axis in Fig. 2C should accordingly be scaled tenfold lower than in the original publication. This discrepancy in FSS does not alter the primary conclusions of our work. However, our suggestion based on the data in Fig. 2C that proximal tubule (PT) cells tune their internalization capacity in response to altered glomerular filtration rates (GFRs) within the normal physiologic range is now in question, as it appears that endocytosis in proximal tubule cells is activated even by exposure to very low fluid shear stress. We apologize for the error.”

The corrected Fig. 2 and its corrected legend appear below.

![Fig. 2](https://www.pnas.org/doi/10.1073/pnas.1602041113)
Shear stress-dependent regulation of apical endocytosis in renal proximal tubule cells mediated by primary cilia

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The kidney has an extraordinary ability to maintain stable fractional solute and fluid reabsorption over a wide range of glomerular filtration rates (GFRs). Internalization of filtered low molecular weight proteins, vitamins, hormones, and other small molecules is mediated by the proximal tubule (PT) multiligand receptors megalin and cubilin. Changes in GFR and the accompanying fluid shear stress (FSS) modulate acute changes in PT ion transport thought to be mediated by microrrow bending. We found that FSS also affects apical endocytosis in PT cells. Exposure of immortalized PT cell lines to physiologically relevant levels of FSS led to dramatically increased internalization of the megalin-cubilin ligand albumin as well as the fluid phase marker dextran. FSS-stimulated apical endocytosis was initiated between 15 and 30 min postinduction of FSS, occurred via a clathrin- and dynamin-dependent pathway, and was rapidly reversed upon removing the FSS. Exposure to FSS also caused a rapid elevation in intracellular Ca$^{2+}$, which was not observed in deciliated cells, upon treatment with BAPTA-AM, or upon inclusion of apyrase in the perfusion medium. Strikingly, deciliation, BAPTA-AM, and apyrase also blocked the flow-dependent increase in endocytosis. Moreover, addition of ATP bypassed the need for FSS in enhancing endocytic capacity. Our studies suggest that increased [Ca$^{2+}$], and purinergic signaling in response to FSS-dependent ciliary bending triggers a rapid and reversible increase in apical endocytosis that contributes to the efficient retrieval of filtered proteins in the PT.

Significance

The proximal tubule (PT) of the kidney is the primary site for reabsorption of ions, solutes, and filtered low molecular weight proteins. PT cells rapidly modulate ion transport capacity in response to the fluid shear stress (FSS) that accompanies changes in glomerular filtration rate. We report here that PT cells also adjust their capacity for endocytosis in response to FSS. Apical endocytosis of the megalin-cubilin ligand albumin and of fluid phase markers is markedly increased upon exposure to FSS. Moreover, Ca$^{2+}$ signaling mediated by the primary cilia on PT cells is required for this response. These studies define a novel pathway in PT cells that plays an essential role in maintaining kidney function.

Author contributions: V.R., N.M.P.-S., M.D.C., and O.A.W. designed research; V.R., Y.R., and N.M.P.-S. performed research; V.R., Y.R., N.M.P.-S., M.D.C., and O.A.W. analyzed data; and V.R. and O.A.W. wrote the paper.

The authors declare no conflict of interest.

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The kidney maintains stable efficient solute and fluid reabsorption over a wide range of glomerular filtration rates (GFRs), which is essential to preserve glomerulotubular balance (1, 2). The majority of filtered water, Na$^+$, proteins, and other solutes are reabsorbed in the proximal tubule (PT), which plays a critical role in blood volume homeostasis. Internalization of filtered low molecular weight (LMW) proteins, vitamins, hormones, and other small molecules is mediated by the PT multiligand receptors megalin and cubilin (3). Defects in the uptake of these ligands leads to LMW proteinuria, which contributes to the pathogenesis of many renal diseases including acute and chronic kidney injury, metal toxicity, cystinosis, and the X-linked disorders Lowe syndrome and Dent disease (4, 5).

Increases in GFR lead to acute changes in PT ion transport capacity. The sodium–hydrogen exchanger NHE3 rapidly accumulates at the apical surface in response to the increased fluid shear stress (FSS) on PT cells to enable increased Na$^+$ reabsorption (2, 6). Modeling studies have suggested that these flow-mediated changes in ion transport are regulated by a mechano-sensitive mechanism induced by microrrow bending (7, 8). Increases in GFR also enhance the need for megalin-cubilin-mediated uptake of filtered ligands. However, it is unknown whether or how endocytosis in PT cells responds to changes in FSS.

Here we have investigated the effect of increased flow and the accompanying FSS on apical endocytosis in PT-derived epithelial cells. We find a rapid and sustained increase in endocytic uptake of both the megalin–cubilin ligand albumin and a fluid phase marker upon exposure to physiologically relevant levels of FSS. Both basal- and FSS-stimulated uptake were inhibited by perturbants of clathrin assembly and dynamin function. Exposure to flow also triggered an increase in intracellular Ca$^{2+}$ concentration ([Ca$^{2+}$]) that required release of extracellular ATP and the presence of primary cilia. Importantly, deciliation of cells or inclusion of apyrase in the medium did not alter endocytosis under static conditions but completely abrogated the FSS-stimulated endocytic response. Our data suggest that flow sensing by mechanosensitive channels in the primary cilia modulates acute apical endocytic responses in PT cells. We discuss the impact of these results on our understanding of normal and disease kidney physiology.

Results

Exposure to FSS Stimulates Apical Endocytosis in PT Cells. A major function of the PT is to internalize solutes and LMW proteins from the glomerular ultrafiltrate. To this end, cells lining the PT express high levels of the multiligand receptors megalin and cubilin, and are specialized to maintain robust apical endocytic capacity (9–11). To confirm that immortalized cell models of the PT retain a high capacity for apical endocytosis, OK cells and LLC-PK1 cells were exposed to apically- or basolaterally added fluorescently tagged albumin (a megalin–cubilin ligand) and dextran (a marker for fluid phase endocytosis). As shown in Fig. S1, both of these cell lines internalized albumin and dextran preferentially from the apical surface. Similarly, murine S3 cells, derived from the S3 segment of the PT, also internalized albumin and dextran preferentially from the apical surface, although endocytosis was less robust than in the other PT cells (Fig. S1).
As a control, we performed similar experiments in mouse cortical collecting duct (CCD) cells. These cells express little if any megalin, and not surprisingly, little albumin was internalized from either surface of these cells. However, consistent with previous studies demonstrating more efficient basolateral internalization rates compared with apical endocytosis (12, 13), we observed considerably greater levels of dextran internalized from the basolateral surface in these cells (Fig. S1).

Healthy individuals efficiently clear LMW proteins and other freely filtered megalin–cubilin ligands despite wide fluctuations in GFR. It is unknown whether endocytic capacity in PT cells varies with changes in fluid flow and the accompanying FSS. To test this, we asked whether changes in FSS result in increased apical endocytosis in immortalized PT cells. Polarized OK and LLC-PK1 kidney cells cultured in a parallel plate flow chamber were exposed for 3 h to physiologically relevant levels of FSS (1 dyne/cm², corresponding to a GFR of 115 mL/min/1.73m² (14–16)) or maintained under static conditions. The megalin–cubilin ligand Alexa Fluor 647-albumin [40 μg/mL; the reported concentration of albumin in the PT is ∼23 μg/mL (17)] or the fluid phase marker rhodamine-dextran were included in the perfusate, and the effects on receptor-mediated and fluid phase endocytosis were quantified. We reproducibly observed a striking (typically ∼twofold) and statistically significant increase in the uptake of both albumin and dextran in both cell lines exposed to FSS compared with controls maintained under static conditions (Fig. 1). Identical results were obtained when the cells were cultured in Ibidi six-well slide chambers, which were used for most subsequent experiments. Because cells under FSS were exposed to a greater volume of medium than cells incubated under static conditions, we confirmed that fluorescent albumin and dextran were not noticeably depleted from the medium under our static incubation conditions; thus, this does not account for the different in uptake that we observed (Fig. S2). Our results are consistent with those of Ferrell et al., who recently reported similar effects of FSS on albumin uptake in OK cells (18). The increase in both receptor-mediated and fluid phase uptake in PT cells upon exposure to FSS suggests a generic stimulation in endocytosis under these conditions, rather than a selective effect on megalin–cubilin trafficking. In contrast, apical endocytosis in MDCK type II cells, which have hybrid characteristics of proximal and distal tubule cells, was unaffected by FSS (Fig. 1). Thus, the FSS-stimulated increase in endocytosis is apparently selective for cells of PT origin.

Onset, Reversibility, and FSS Dependence of FSS-Stimulated Endocytosis. To test the kinetics of FSS-stimulated endocytosis, we incubated OK cells under static conditions or exposed them to 1-dyne/cm² FSS in the presence of fluorescent albumin for various time points, then fixed and quantified uptake. As shown in Fig. 2A, uptake rates under FSS and static conditions were similar for the first 15 min and then diverged. By 30 min of exposure to flow, there was a statistically significant increase in the overall amount of albumin internalized compared with cells treated under static conditions. This increased rate of uptake continued linearly for the remaining incubation period (Fig. 2A). In separate experiments, we monitored uptake over a 1–3-h incubation period (Fig. 2A, Inset). Uptake of albumin in static and FSS-treated cells remained linear over this time period, with a roughly twofold increase in endocytosis in FSS-treated cells compared with static at each time point.

We next measured the reversibility of the FSS-stimulated increase in endocytosis. To this end, we exposed cells to FSS for 1 h in the presence (Fig. 2B, sample 1) or absence (Fig. 2B, samples 2–4) of added albumin, then stopped the flow for various times (15–60 min; Fig. 2B, samples 2–4) before adding fluorescent albumin to the cells for 1 h under static conditions. As controls, we incubated cells with fluorescent albumin for a 1-h period in the absence of flow either at the start of the experiment (at 0′; Fig. 2B, sample 5) or coincident with the last uptake period (starting at 2 h; Fig. 2B, sample 6). Albumin uptake in the presence of flow was robust compared with cells not exposed to FSS (Fig. 2B; compare sample 1 to samples 5 and 6). Strikingly, reversal of the endocytic response to FSS was essentially complete by 15 min after the cessation of FSS (Fig. 2B, sample 2). In humans, normal GFR varies between 60 and 120 mL/min/1.73m² but can reach 160 mL/min/1.73m² in diseased individuals. To test whether PT cells adjust their endocytic capacity in response to changes in FSS induced within this range, we quantified the internalization of Alexa Fluor 647-albumin in OK cells exposed to FSS within a range of 0.1–1.5 dyne/cm² for 1 h. As shown in Fig. 2C, there was a general increasing trend in the amount of Alexa Fluor 647-albumin internalized by cells exposed to this range of FSS. Albumin internalization did not reach statistical significance relative to static controls (0 dyne/cm²) at 0.2, 0.4, and 0.7 dyne/cm², but was significantly different in cells exposed to FSS of 1.0 dyne/cm² and above. Importantly, there was a statistically significant increase in albumin uptake at cells exposed to 1.0 and 1.5 dyne/cm² compared with 0.7 dyne/cm² (equivalent to a GFR of ∼60 mL/min/1.73m²). Exposure to higher FSS (1.5 dyne/cm², equivalent to a GFR of ∼150 mL/min/1.73m²) did not increase endocytic capacity above the level observed at 1.0 dyne/cm² (Fig. 2C). This suggests that PT cells tune their internalization to maximum capacity in response to altered GFR within the normal physiologic range.

FSS-Stimulated Endocytosis Occurs via a Clathrin- and Dynamin-Dependent Pathway. Megalin is internalized into clathrin-coated pits that form at the base of microvilli of PT cells (10, 19). Although some immortalized PT cell lines express caveolin, caveolae are absent in PT cells in vivo (20), suggesting that clathrin-dependent endocytosis represents the primary mechanism for internalization of...
FSS Triggers a Cytosolic Ca\(^{2+}\) Response Required for Stimulated Apical Endocytosis. Modeling studies have suggested that the flow-mediated changes in ion transport are regulated by a mechanosensitive mechanism induced by microvillar bending (7, 8). There is good evidence that primary cilia are not required for this pathway, as similar effects were observed in cells lacking mature cilia (16). In contrast, primary cilia are known to play an essential role in flow-mediated regulation of ion transport in the distal tubule (21). Genetic defects that affect cilia structure or function cause kidney disease, presumably as a consequence of aberrant FSS-dependent signaling (21, 22). Exposure to FSS is known to activate transient receptor potential channels localized on primary cilia to trigger an increase in [Ca\(^{2+}\)] and in many cell types, including kidney CCD cells (2, 21, 23). To test if exposure to FSS triggers a similar response in PT cells, polarized OK cells loaded with Fura-2 AM were perfused with Krebs buffer at an FSS of 2 dyne/cm\(^2\) and the change in [Ca\(^{2+}\)] was determined as described in Methods. Exposure to FSS caused an immediate three- to fourfold increase in [Ca\(^{2+}\)], that returned to baseline levels in 3-4 min (Fig. 4). The FSS-stimulated increase in [Ca\(^{2+}\)] was not observed when Ca\(^{2+}\) was omitted from the perfusion buffer, demonstrating a requirement for extracellular Ca\(^{2+}\) in this response (Fig. 4A).

To test the role of the primary cilia in the FSS-stimulated increase in [Ca\(^{2+}\)], we deciliated OK cells using 30 mM ammonium sulfate for 3 h. We previously showed that this treatment results in efficient and reversible removal of cilia (ref. 24 and Fig. 5A). As shown in Fig. 4B, [Ca\(^{2+}\)] in deciliated cells did not increase in response to FSS.

Previous studies conducted in collecting duct cells have shown that the FSS-stimulated, ciliary-dependent increase in [Ca\(^{2+}\)], is mediated by Ca\(^{2+}\)-stimulated Ca\(^{2+}\) release from the endoplasmic reticulum (ER) via ryanodine receptors (RyRs) (21). To assess the contribution of the Ca\(^{2+}\)-stimulated Ca\(^{2+}\) release to FSS-stimulated increase in [Ca\(^{2+}\)], we treated OK cells with the sarcoplasmic/endoplasmic reticulum Ca\(^{2+}\)-ATPase (SERCA) inhibitor tBuBHQ to deplete ER reserves of Ca\(^{2+}\) and then subjected them to FSS. Resting [Ca\(^{2+}\)] in tBuBHQ-treated cells was elevated relative to untreated cells as expected, and was unaffected upon exposure to FSS, confirming that ER stores of Ca\(^{2+}\) contribute to the FSS-stimulated rise in [Ca\(^{2+}\)] (Fig. 4C). We then depleted the RyR-sensitive pool of ER Ca\(^{2+}\) using ryanodine to test the role of RyRs in FSS-stimulated increase in [Ca\(^{2+}\)]. As shown in Fig. 4C, we observed that the flow-stimulated increase in [Ca\(^{2+}\)] was ablated posttreatment with ryanodine, confirming that release of the RyR sensitive pool of ER Ca\(^{2+}\) is requisite for the flow-stimulated increase in [Ca\(^{2+}\)]. Additionally, buffering cytosolic Ca\(^{2+}\) by incubation with the cell permeable Ca\(^{2+}\) chelator bis-(o-aminophenoxo)-N,N,N',N'-tetraacetic acid-acetoxymethyl ester time point. *P < 0.05 vs. all other conditions by ANOVA, except endocytosis measured at 1.0 vs. 1.5 dynes/cm\(^2\) are not significantly different from each other.

Fig. 2. Time course, reversibility, and FSS threshold of FSS-stimulated apical endocytosis. (A) Time course of onset of FSS-stimulated endocytosis. OK cells plated in ibidi μ-slide chambers were incubated under static conditions or exposed to 1-dyne/cm\(^2\) FSS in the presence of 40 μg/mL Alex Fluor 647-albumin for the indicated time periods, then fixed, and average internalized fluorescence quantified from 15 to 20 fields per condition. *P < 0.04 vs. paired static control by Student t test. (Inset) Albumin uptake over a 1-3-h time course. *P < 0.02 vs. static control by t test. (B) Reversibility of FSS-stimulated endocytosis. OK cells were exposed to 1-dyne/cm\(^2\) FSS for 1 h in the presence (1) or absence (2-4) of 40 μg/mL Alex Fluor 647-albumin. Cells were then fixed immediately (1) or incubated under static conditions for 15 min (2), 30 min (3), or 60 min (4) before addition of 40 μg/mL Alex Fluor 647-albumin for 1 h. As controls, Alex Fluor 647-albumin was added to cells incubated under static conditions for 1 h at the start of the time course (5) or after 2 h (6) to coincide with the uptake period for sample 4. Internalized fluorescence was quantified for five fields per condition. The average fluorescence ± range from two independent experiments is plotted. *P < 0.05 vs. static control (sample 6) by ANOVA with Bonferroni correction. All other pairwise comparisons are not significantly different. (C) OK cells were incubated with 40 μg/mL Alex Fluor 647-albumin for 1 h under static conditions (0 dynes/cm\(^2\)) or during exposure to the indicated FSS. Average internalized fluorescence was quantified from four wells for each membrane and fluid from the apical surface of these cells. To test whether the FSS-stimulated component of albumin endocytosis occurs via a mechanism similar to that of basal uptake, we asked whether perturbants of clathrin-dependent endocytosis disrupted albumin uptake under static conditions and upon exposure to FSS. To this end, we preincubated cells for 30 min with chlorpromazine (a drug that inhibits assembly of clathrin coats) before addition of fluorescent albumin under static conditions or in the presence of 1-dyne/cm\(^2\) FSS. Treatment with chlorpromazine reproducibly and significantly inhibited both basal and FSS-stimulated endocytosis (by 42% and 33%, respectively; Fig. 3A). Treatment with the dynamin inhibitor Dyno-4a also reduced cell- associated albumin (by 49% and 62% in cells exposed to static and FSS conditions, respectively; Fig. 3B).
(BAPTA-AM) inhibited the FSS-stimulated increase in \([\text{Ca}^{2+}]\) (Fig. 4C).

Bending of the cilia in distal tubule kidney cells is also known to cause extracellular release of ATP, which can trigger activation of purinergic receptors (P2YRs) causing a further increase in \([\text{Ca}^{2+}]\) (25). Therefore, we tested whether depletion of extracellular ATP alters the FSS-stimulated increase in \([\text{Ca}^{2+}]\) in PT cells. As shown in Fig. 5D, we found that inclusion of apyrase in the perfusion attenuated the FSS-stimulated increase in \([\text{Ca}^{2+}]\). The involvement of P2YRs in this signaling cascade was confirmed by addition of the pan P2YR inhibitor suramin, which also abrogated the FSS-stimulated Ca\(^{2+}\) response (Fig. 4D). Together, these data show that FSS triggers an increase in \([\text{Ca}^{2+}]\), which requires the primary cilium, extracellular \(\text{Ca}^{2+}\) influx, release of \(\text{Ca}^{2+}\) from ER stores through ryanodin receptors, and ATP-dependent activation of P2Y receptors.

**Primary Cilia and Purinergic Signaling Are Required for FSS-Dependent Modulation of Endocytosis.** To test whether primary cilia and the ATP-dependent \(\text{Ca}^{2+}\) response are also required for the endocytic response to FSS in PT cells, we deciliated OK cells as above, and measured internalization of Alexa Fluor 647-albumin in cells incubated under static conditions or exposed to 1-dyne/cm\(^2\) FSS. Indirect immunofluorescence confirmed that our deciliation protocol resulted in removal of essentially all primary cilia (Fig. 5A). Strikingly, whereas basal albumin uptake under static conditions was unaffected in deciliated cells, the FSS-induced increase in endocytic uptake was almost entirely abrogated (Fig. 5A and B). Similarly, inclusion of BAPTA-AM (Fig. 5C) or apyrase (Fig. 5D) in the medium also blocked FSS-stimulated but not basal uptake of albumin. We conclude that primary cilia and ATP-dependent P2YR signaling are both required for acute modulation of apical endocytosis in the PT in response to FSS.

Conversely, we asked whether increasing \([\text{Ca}^{2+}]\), in the absence of FSS is sufficient to trigger the downstream cascade that leads to enhanced endocytosis. As expected, addition of 100 \(\mu\text{M}\) ATP in the absence of FSS caused an acute and transient threefold increase in \([\text{Ca}^{2+}]\), whereas incubation with suramin led to a sustained elevation in \([\text{Ca}^{2+}]\), that was unchanged by FSS (Fig. S3A and Fig. 4C). Addition of ATP to cells incubated under static conditions also stimulated endocytosis by roughly 50% (Fig. S3B). Both basal and ATP-stimulated endocytosis were profoundly inhibited by suramin (Fig. S3B). Ryanodin also stimulated endocytosis in the absence of FSS, and this effect was not further augmented by exposure of the cells to FSS (Fig. S3C).

**Discussion**

PT cells are specialized to internalize and recycle large amounts of apical membrane to effectively clear LMW proteins and other molecules from the glomerular ultrafiltrate. Defects in megalin or cubilin, which mediate the uptake of these filtered ligands (3), or saturation of this pathway as can occur in diabetic nephropathy (26), lead to tubular proteinuria and eventually to renal failure. However, surprisingly little is known about how apical endocytosis is regulated in PT cells, and whether this pathway can respond acutely to variations in glomerular filtration rate to maximize uptake efficiency of filtered ligands.

Studies conducted in the past decade have underscored the importance of FSS in affecting the organization of the cytoskeleton, activity of certain transcription factors, and the trafficking and retention of transporters that regulate vectorial ion transport in PTs (2, 6). Our studies here demonstrate an additional role for FSS in the modulation of apical endocytosis in the PT. From our experiments we conclude that: (i) exposure to FSS increases apical endocytic uptake of fluid phase and membrane-
bound cargoes in immortalized PT cells in culture as well as in mouse kidney slices; (ii) the FSS-stimulated endocytic response is rapid, reversible, and is mediated by a clathrin- and dynamin-dependent pathway; (iii) FSS also stimulates an immediate spike in intracellular Ca\(^{2+}\) mediated by Ca\(^{2+}\)-dependent Ca\(^{2+}\) release from ER stores; (iv) the primary cilium of PT cells is the principal mechanotransducer mediating the spike in FSS-stimulated intracellular Ca\(^{2+}\) and the subsequent endocytic response; and (v) release of extracellular ATP triggered by the bending of primary cilia in the presence of flow is required for activation of P2YRs and for FSS-stimulated endocytic responses in PT cells. A working model for how this signaling cascade might modulate endocytic capacity is shown in Fig. 6.

Fig. 5. FSS-stimulated apical endocytosis requires cilia and extracellular ATP. (A) OK cells were treated with ammonium sulfate as indicated to deciliate cells, then incubated with Alexa Fluor 647-albumin under static conditions or exposed to FSS (1 dyne/cm\(^2\)) for 3 h. Cells were fixed and processed to detect cilia (with antiacetylated tubulin antibody; red) and internalized albumin (green); maximum projections of confocal stacks are shown. Scale bars, 10 μm. Quantitation of albumin uptake in control vs. deciliated cells ([A], mean ± SEM of three experiments), or in cells treated with 10 μM BAPTA-AM ([C], mean ± SEM of four experiments) or 1 U/mL apyrase ([D], mean ± SEM of three experiments) incubated under static conditions or exposed to 1-dyne/cm\(^2\) FSS for 1 h. *P < 0.002; **P < 0.001 by ANOVA with Bonferroni correction. Other pairwise comparisons are not significantly different.

Fig. 6. Model for FSS-regulated modulation of apical endocytosis in PT. Our data support a model in which exposure to FSS increases apical endocytic capacity in PT cells via a pathway that requires ciliary bending, and entry of extracellular Ca\(^{2+}\) via a ciliary-localized cation channel [possibly polycystin-2 (PC2)] that lead to increases in intracellular Ca\(^{2+}\) (Ca\(^{2+}\)). Bending of the primary cilium also causes release of ATP to the luminal surface (via nucleotide transporters or other mechanisms) which in turn activates P2YRs and further increases Ca\(^{2+}\). Endocytosis from the apical surface of polarized cells is known to occur exclusively at the base of microvilli via a clathrin- and dynamin-dependent pathway that is dependent on actin. We hypothesize that increased Ca\(^{2+}\) triggers a cascade that ultimately modulates actin dynamics to increase the size and volume of individual apical clathrin-coated pits.
internalized in these unevenly shaped structures, which bud from the apical membrane and fuse with a subapical network of tubules (19). We hypothesize that exposure to FSS increases the average size of these clathrin-coated structures to accommodate larger endocytic capacity. Consistent with this, there is preced-ence for modulation of clathrin-coated pit size in nonpolarized cells to accommodate larger cargoes such as virus particles (28). Unlike “traditional” clathrin-mediated endocytosis, internalization of these large cargoes requires modulation of actin dynamics at the coated pit site. We hypothesize that a similar pathway could be triggered upon FSS-stimulated [Ca2+]i increases in PT cells.

The involvement of primary cilia in the endocytic response to FSS is not well known, the first known function for cilia in PT cells and raises the possibility that defects in cilogenesis could impair the regulation of apical endocytic uptake in these cells. Genetic defects that alter ciliary function or structure lead to renal disease. To date, all disorders that result in shortened primary cilia in the kidney cause cystic disease, presumably as a consequence of aberrant flow-dependent signaling (21, 22). In contrast, transient elongation of cilia has been observed during kidney development and during recovery from acute kidney injury (29, 30). It is unclear how changes in cilia length might impact the endocytic response to FSS in the PT. Strikingly, however, cilia length has been found to be altered in cells and zebrafish depleted of OCRL1, the phosphatidylinositol 5′-phosphate kinase deficient or absent in patients with Lowe syndrome, an X-linked disorder that results in tubular proteinuria (24, 31, 32). Although the precise effects or mechanism are not yet clear, as both longer and shorter cilia were reported, these studies provide a tantalizing link that could explain how loss of OCRL1 function leads to tubular proteinuria in Lowe syndrome patients. Future studies will be required to examine whether defects in the modulation of endocytosis by FSS contribute to kidney disease in these and other patients with tubular proteinuria.

Methods

Quantitation of Flow-Dependent Endocytosis. OK cells were cultured on ibidi six-well chambers as described above. μ-Slide Luer 0.1 μm and 1 μM l-lysine-PEG rhodamine-dextran were added at the indicated FSS. Cells were perfused or maintained under static conditions for 1 h at 37 °C unless otherwise indicated. Cells were fixed with 1% paraformaldehyde or periodate-lysine-paraformaldehyde (for experiments using dextran) for 20 min at ambient temperature, the fix was removed, and 50 μL of ProlongGold (Invitrogen) was added to each well. The chambers were imaged the next day using a Leica TCS SPS confocal microscope. Five to ten confocal stacks of randomly selected fields were acquired per well. Images were exported as 8-bit tagged image file formats or Leica image files and analyzed using Fiji or Image J. Maximum intensity projections of stacks were obtained for each field and total intensity was calculated using the Measure function. Measurements for background were digitally subtracted from the total intensity of each field. No normalization was used, as raw values were generally reproducible across independent experiments. More details on other methods used are provided in SI Methods.

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