The World Health Organization (WHO) has reported a worldwide surge in cases of cholera caused by the intestinal pathogen *Vibrio cholerae*, and, combined, such surges have claimed several million lives, mostly in early childhood. Elevated cAMP production in intestinal epithelial cells challenged with cholera toxin (CTX) results in diarrhea due to chloride transport by a cAMP-activated channel, the cystic fibrosis transmembrane conductance regulator (CFTR). However, the identity of the main cAMP-producing proteins that regulate CFTR in the intestine and may be relevant for secretory diarrhea is unclear. Here, using RNA-Seq to identify the predominant AC isoform in mouse and human cells and extensive biochemical analyses for further characterization, we found that the cAMP-generating enzyme adenylate cyclase 6 (AC6) physically and functionally associates with CFTR at the apical surface of intestinal epithelial cells. We generated epithelium-specific AC6 knockout mice and demonstrated that CFTR-dependent fluid secretion is nearly abolished in AC6 knockout mice upon CTX challenge in ligated ileal loops. Furthermore, loss of AC6 function dramatically impaired CTX-induced CFTR activation in human and mouse intestinal spheroids. Our finding that the CFTR–AC6 protein complex is the key mediator of CTX-associated diarrhea may facilitate development of antiarrheal agents to manage cholera symptoms and improve outcomes.

Cholera is a disease characterized by severe diarrhea and resultant dehydration, hypovolemic shock, and acidosis (1). Worldwide, 172,454 cholera cases with 1,304 deaths were reported in 2015 to the World Health Organization (WHO). The WHO reports that 41% of these cases were in Africa, 37% in Asia, and 21% in the Americas. More recently, the WHO has reported major outbreaks in Somalia and Yemen with cholera cases currently exceeding one million and growing (http://www.emro.who.int/health-topics/cholera-outbreak/outbreaks.html, accessed 1st April, 2018). It is of significant public health concern that cholera is on an epidemiological rise across the world and continues to claim a significant number of lives, mostly children. Defining precise pathogenic mechanisms of cholera is critical to design preventive and disease control interventions and to prevent such outbreaks.

Cholera is caused by members of the bacterial species *Vibrio cholerae* and is transmitted via a fecal–oral route or consumption of contaminated food and water (1). When virulent cholera bacteria adhere to small bowel, they release cholera enterotoxin (CTX) formed from a B subunit pentamer, for binding to host cells, and an enzymatic A subunit. The endocytosed enzymatic A subunit ADP-ribosylates Go subunits of G proteins inside the cell, causing them to remain active. Adenylate cyclases (ACs), the membrane-bound proteins that catalyze the conversion of ATP to cAMP and pyrophosphate, are, in general, activated via Goα pathways (2). Hence, at a molecular level, CTX challenge translates into excessive cAMP production by ACs and activation of cAMP-dependent protein kinase A (PKA). Severe diarrhea in cholera occurs due to PKA-mediated phosphorylation and then hyperactivation of cystic fibrosis transmembrane conductance regulator (CFTR) protein, which continues to electrochemically drive water into the intestinal lumen coupled to its chloride transport activity (3). CFTR is a key member of the secretory epithelium in the gut and the lung that regulates overall fluid transit by means of its chloride transport function (4). Loss of CFTR function results in cystic fibrosis (CF), a fatal disease that affects multiple organs (5). The most common cause of CF is the Phe-508 deletion (F508del) mutation in CFTR that generates a clinically significant processing
defect in the protein and little to no function of CFTR at the plasma membrane. CFTR has been established as the primary driver of CTX-induced diarrhea based on studies in CF mice (6). CFTR can be regulated by multiprotein complex formation consisting of protein kinases, scaffolding proteins, ion channels, cytoskeletal elements, and other transporters (7, 8). Protein–protein interactions of CFTR have been shown to play important roles in CFTR trafficking, channel regulation, compartmentalized signaling to optimize CFTR function, and surface stabilization (7–10).

There are currently nine membrane-bound AC isoforms (Adcy1–9) identified in mammalian cells (11). The predominant AC isoform that regulates CFTR in the intestine and is relevant to secretory diarrhea has not been clearly determined (2). In this study, we elucidated that CFTR and AC6 signaling complex formation is responsible for CTX-induced diarrhea and that loss of AC6 in mice is protective against diarrhea. Our findings identify the primary diarrheagenic protein complex in cholera, presenting an opportunity to develop specific anti-diarrheal agents targeted at these complexes.

Results

**AC6 is the most abundant AC isoform in the gut**

Cellular cAMP generation is determined by the relative expression and distribution of different ACs. To determine the relative expression of ACs in gut epithelial cells, we used mRNA sequencing in mouse and human intestinal epithelial cells. We found AC6 to be the predominant AC isoform in these cells (Fig. 1A), which led us to further investigate the role of AC6 in intestinal physiology and pathophysiology. We generated an epithelium-specific knockout of AC6 using a sonic hedgehog cre recombinase (Shh Cre) driver and confirmed loss of AC6 protein expression in Adcy6f/fShh Cre/+ (Adcy6Δ/Δ) mouse ileum (Fig. 1, B–E). Immunostaining data revealed apical enrichment of AC6 in the mouse ileum (Fig. 1E). CFTR is an apical membrane-localized chloride channel in the gut and airway epithelial cells (8). CFTR is composed of two membrane-spanning domains, two nucleotide-binding domains, and a unique intracellular regulatory (R) domain of −240 amino acids (residues 590–830) (13). CFTR is activated by PKA-mediated phosphorylation of serine residues in the R domain and binding and hydrolysis of intracellular ATP at the nucleotide-binding domains. Phosphorylation of the R domain is required for opening of the CFTR channel. Based on these earlier reported data and our findings, we tested how cAMP generation by apical AC6 may regulate CFTR function.

**AC6 interacts directly with CFTR in intestinal epithelial cells**

Based on an exclusive apical localization of CFTR and AC6 in the gut epithelial cells, we considered the possibility that CFTR and AC6 might interact with each other. In a co-immunoprecipitation assay, we detected AC6 in complex with CFTR in the isolated Adcy6f/f intestinal mucosa, whereas the band corresponding to AC6 was negligibly detected in Adcy6Δ/Δ mucosa (Fig. 2A). Importantly, in this experiment, we observed that loss of AC6 did not impact CFTR protein expression (Fig. 2A). We determined, using a FRET assay, that AC6 and CFTR interact at the plasma membrane in live colon epithelial cells (HT29CL19a) (Fig. 2, B and C). In addition, we performed proximity ligation assays (PLAs) to detect in situ AC6–CFTR interactions in mice and human intestinal epithelial cells. PLA is a highly sensitive and specific immunosassay-based technology that has been designed to quantitate proteins and detect protein interactions and modifications within a spatial span of 40 nm (14). Using this method, we detected positive PLA signal corresponding to CFTR–AC6 complex at the apical membrane in Adcy6Δ/Δ intestinal epithelial cells (Fig. 2, D and E). A PLA signal was present in negligible quantities in Adcy6Δ/Δ intestine (Fig. 2, D and E). Similarly, we found a positive PLA signal corresponding to CFTR–AC6 interaction in human intestinal spheroids that was enriched at the apical or luminal side (Fig. 2F). To identify which cytosolic regions of AC6 may interact with CFTR, we generated His–S–tagged cytosolic domains of AC6 (C1 (306–672 aa) and C2 (917–1165 aa)), which together constitute the catalytic core of AC6 protein, and expressed these constructs in a bacterial protein expression system. Using S–protein pulldown, we detected an interaction of FLAG-CFTR with AC6 C2 (Fig. 2G). The interaction of AC6 C2 was maintained with FLAG-CFTRH11001, suggesting that the interaction does not require the canonical C-terminal PDZ motif in CFTR (Fig. 2G). Overall, we determined that AC6 forms a protein complex with CFTR, and, given the high abundance of the AC6 protein in intestinal epithelial cells, we hypothesized that this protein complex may potentially become the major physiological driver of CFTR function.

**cAMP generation is attenuated in Adcy6Δ/Δ intestinal epithelial cells**

Based on the abundance of AC6 in intestinal epithelial cells, we investigated how loss of AC6 affects intracellular cAMP levels. We isolated intestinal crypts from Adcy6f/f and Adcy6Δ/Δ ileum and cultured them in a Matrigel-based matrix. Subsequently, we measured cAMP levels using ELISA and detected dramatically reduced CAMP production in Adcy6Δ/Δ compared with Adcy6f/f intestinal epithelial cells in response to the AC agonist forskolin (FSK) (Fig. 3A). In addition, the CTX–induced increase in intracellular cAMP levels was significantly diminished in Adcy6Δ/Δ intestinal epithelial cells (Fig. 3B). Production of intracellular cGMP upon guanylate cyclase activation by linaclotide (15), a peptide analog of heat-stable enterotoxin, did not differ between Adcy6f/f and Adcy6Δ/Δ intestinal epithelial cells (Fig. 3C). Because CTX failed to produce diarrhea–inducing flux of intracellular cAMP in Adcy6Δ/Δ cells, we proceeded to investigate how loss of AC6 affects CFTR function upon CTX challenge.

**Adcy6Δ/Δ mice are protected against CTX-induced diarrhea**

Having determined that AC6 and CFTR interact at the plasma membrane and that loss of AC6 fails to elicit cAMP response in the presence of CTX, we next investigated intestinal fluid secretion in Adcy6Δ/Δ mice upon CTX challenge. We used the closed ileal loop model in mice to test CTX–induced fluid secretion (3, 6, 16, 17). CTX injected into the ligated loops induces hyperactivation of CFTR Cl− channel via cAMP production, causing a voluminous fluid secretion and accumulation in the closed loops. Consistently, chemical inhibition of CFTR function (e.g. using CFTRinh-172) or loss-of-function

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**Diarrheagenic complex in cholera**

Based on the abundance of AC6 in intestinal epithelial cells, we investigated how loss of AC6 affects intracellular cAMP levels. We isolated intestinal crypts from Adcy6f/f and Adcy6Δ/Δ ileum and cultured them in a Matrigel-based matrix. Subsequently, we measured cAMP levels using ELISA and detected dramatically reduced CAMP production in Adcy6Δ/Δ compared with Adcy6f/f intestinal epithelial cells in response to the AC agonist forskolin (FSK) (Fig. 3A). In addition, the CTX–induced increase in intracellular cAMP levels was significantly diminished in Adcy6Δ/Δ intestinal epithelial cells (Fig. 3B). Production of intracellular cGMP upon guanylate cyclase activation by linaclotide (15), a peptide analog of heat-stable enterotoxin, did not differ between Adcy6f/f and Adcy6Δ/Δ intestinal epithelial cells (Fig. 3C). Because CTX failed to produce diarrhea–inducing flux of intracellular cAMP in Adcy6Δ/Δ cells, we proceeded to investigate how loss of AC6 affects CFTR function upon CTX challenge.
mutation in CFTR (e.g. F508del CFTR) makes CTX less effective or ineffective in inducing secretion in ligated ileal loops in mice (16, 17). Whereas CTX induced significant fluid accumulation in Adcy6<sup>−/−</sup> mice, this phenomenon was largely absent in Adcy6<sup>−/−</sup> mice at a CTX dose of 1 μg and dramatically impaired at a dose of 10 μg (Fig. 4, A and B). Importantly, cGMP-induced fluid secretion in the ileal loops in response to linaclotide was not significantly different between Adcy6<sup>−/−</sup> and Adcy6<sup>−/−</sup> mice (Fig. 4, A and B). These findings support the conclusion that AC6 is required for CTX-induced diarrhea. Also, loss of AC6 does not impair the functional ability of cAMP-independent activation of CFTR (e.g. by cGMP in intestinal epithelial cells). To determine the effect of loss of AC6 on CFTR chloride channel function, we measured cAMP-stimulated short-circuit currents. CFTR-dependent Cl<sup>−</sup> currents were monitored in tracheal epithelial cells isolated from Adcy6<sup>−/−</sup> and Adcy6<sup>−/−</sup> mice and polarized on the transwells. As the shh cre driver is highly expressed in tracheal epithelial cells in addition to intestinal cells (Fig. 1C) and we confirmed loss of AC6 in the tracheal epithelial cells (Fig. 4C), it was appropriate to test tracheal epithelial cells to monitor CFTR-mediated Cl<sup>−</sup> transport as a function of loss of AC6. CFTR-dependent Cl<sup>−</sup> currents were reduced by 3.2-fold in Adcy6<sup>−/−</sup> versus Adcy6<sup>−/−</sup> mice (Fig. 4, D and E). Purinergic receptor stimulation by ATP that elicits calcium-dependent Cl<sup>−</sup> currents generated a similar response in Adcy6<sup>−/−</sup> and Adcy6<sup>−/−</sup> tracheal cells (Fig. 4, D and E). Hence, AC6 is the primary AC to...
mediate cAMP-dependent activation of CFTR that drives diarrhea in cholera and CFTR activation in the large airways.

Loss of AC6 impairs fluid secretion in human- and mouse-derived enterospheres

Intestinal organoids/enterospheres are cultured three-dimensional epithelial structures that contain the key intestinal cell types and recapitulate the functional features of the native intestinal tissue (18, 19). Enterosphere-based fluid secretion assay is a method that can specifically determine CFTR function (9, 20). Because of high expression of CFTR in the enterospheres, CFTR activation using forskolin causes rapid luminal area expansion that can be quantitated (9, 20). Luminal expansion is perturbed upon chemical inhibition of CFTR function.
Figure 3. cAMP generation is impaired in Adcy6<sup>Δ/Δ</sup> intestinal epithelial cells. Bar graphs represent whole-cell cAMP measurement in response to FSK (10 μM, 1 h) (A) and CTX (100 ng, 24 h) and linaclotide (LC; 500 ng, 24 h) (B) and whole-cell cGMP measurement in response to CTX (100 ng, 24 h) and linaclotide (500 ng, 24 h) (C) (same samples as used in B) in intestinal epithelial cells derived from Adcy6<sup>f/f</sup> and Adcy6<sup>Δ/Δ</sup> mice. Each dot represents a single measurement from an n = 3 intestinal cell preparation. Error bars, S.D. p value was determined by ANOVA with Bonferroni’s multiple-comparison test.

Discussion

CFTR is an important channel that controls one of the key functions of the intestinal epithelium (i.e. fluid homeostasis). Excessive activation of CFTR causes secretory diarrhea, whereas loss of CFTR function may lead to chronic constipation or to meconium ileus in some CF cases. Both of these scenarios are serious health issues.

Our results collectively demonstrate that AC6–CFTR macromolecular complex formation principally drives CFTR-dependent hypersecretion upon CTX challenge. Our findings present an avenue for development of novel anti-diarrheal agents to treat cholera that currently poses a massive health threat in many parts of the world. Small molecules that could specifically inhibit AC6 activity may prove beneficial to control cholera. Whereas administering osmolytes is the cheapest and simplest approach to prevent and treat dehydration in cholera to avert fatality, this does not control diarrhea, address the root cause of cholera, or mitigate the pathogenic effects of chronic cAMP production during cholera. Balancing cellular cAMP is necessary, as excessive cAMP can ectopically influence multiple signaling pathways and may even lead to cell toxicity (21).
This is particularly important because AC6 is such an abundant protein in the intestinal epithelial cells. Based on our data, regulation of AC6 through a small-molecule inhibitor is theoretically one of the most effective ways to control cholera. The use of a highly specific CFTR inhibitor (CFTRinh-172) as an anti-diarrheal agent could not be successful because of associative toxicity of the compound (22). Therefore, we need to not only develop safer anti-diarrheal drugs but also identify multiple targets in cholera that can be targeted with minimal side effects. Studies elucidating the roles of ACs in human diseases have been limited by the lack of specific reagents for ACs (inhibitors, activators, and antibodies) (23). We sufficiently validated that the AC5/6 antibody used in the study recognizes AC6. Many times, we resorted to using recombinant AC6 (with FLAG or fluorescent tag) for the sole purpose of validation. We also hope that defining these AC-specific roles in human diseases will incentivize related product development within pharmaceutical industries.

The relatively high abundance of AC6 in the intestine and its close proximity to CFTR through a direct interaction indicate...
why CTX-induced diarrhea becomes so exaggerated. The minimal CTX-generated fluid secretion response in \( \text{Adcy6}^{-/} \) mice suggests that other ACs may not be able to compensate for the loss of AC6 in terms of CFTR activation and formation of cellular cAMP upon CTX challenge. Only at high doses of FSK (100 \( \mu \text{M} \)) and CTX (10 \( \mu \text{g} \)) were AC6-independent mechanisms of stimulation of fluid secretion evident. We anticipate that the global cAMP generation by other ACs would stimulate CFTR-dependent fluid secretion at high doses of cAMP agonists in AC6 knockout mice; however, the fluid secretion stayed dramatically poor relative to the WT mice. There are several lines of experimental evidence to support that cAMP signaling operates in a largely compartmentalized fashion, which enables more specificity and faster kinetics of the signaling processes, underscoring the importance of cAMP compartmentalization in normal cell physiology (23, 24). Each AC isoform, although producing a common signaling molecule, cAMP, is subject to differential regulation by signaling processes (23, 25).
Diarrheagenic complex in cholera

Localization in different cellular compartments and association with specific G protein–coupled receptors and other downstream effector proteins provide means of cAMP signaling compartmentalization. Formation of macromolecular protein complexes, as in this study, between CFTR and AC6 is an elegant example of compartmentalized cAMP signaling enabling a higher degree of specificity toward CFTR activation.

It is argued that Na⁺ absorption may contribute to fluid accumulation in the enteroids at high doses of cAMP agonists, as NHE3-dependent pH activity has been detected in the enteroids (26). In the undifferentiated or proliferative state (as were the intestinal organoids used in this study), Na⁺ absorption causes minimal luminal expansion in response to FSK over a short time period (30 min). Fluid secretion in response to FSK is completely absent in F508del/F508del cftr mouse enteroids measured within 2 h of stimulant addition as previously demonstrated by us (9). This suggests that in an acute assay, fluid secretion in the enterospheres is largely CFTR-mediated. As a result, we kept the window of measurement of secretion in the enterospheres ≤ 2 h. On an important note, loss of AC6 did not inhibit the process of organoid formation and closure, and CFTR expression was not affected in Adcy6Δ/Δ enterospheres.

Our study demonstrates that AC6 is an important modifier of CFTR function. We find these studies not only relevant to cholera toxin–induced diarrhea but to CF as well. There are data suggesting that the most common CF mutation, F508del CFTR, shows diminished phosphorylation by PKA (27, 28). This effect correlated with the altered channel activity and kinetics of activation of the mutant protein at the plasma membrane. These properties of F508del CFTR are very much reflective of the way AC6 regulates normal CFTR. It would be interesting to investigate whether there is a perturbed interaction of the mutant CFTR protein with AC6. Previously, we demonstrated that F508del CFTR behaves poorly in terms of PDZ-dependent interactions compared with normal CFTR, which translates into reduced half-life and function of the rescued mutant protein at the plasma membrane (9). Understanding how peripheral interactions are orchestrated and regulate CFTR behavior is critical to determine the functional efficiency of rescued F508del CFTR. Also, it will be important to determine how the most stimulatory interactions, one of which according to our data is CFTR–AC6, can be selectively enhanced to benefit F508del CFTR function in combination with other approaches.

Materials and methods

Statistics

The statistical significance was calculated using Student’s t test or one-way ANOVA with multiple-comparison tests as applicable. p < 0.05 was considered statistically significant.

Chemicals and antibodies

Chemicals used in this study include forskolin, CPT-cAMP, isobutylmethylxanthine, and CTX from MilliporeSigma. Antibodies in this study included anti-CFTR M3A7 (Thermo Fisher Scientific) and CF3 (Abcam, Cambridge, UK), anti-AC5/6 (Santa Cruz Biotechnology, Dallas, TX), and anti-ezrin and tubulin (Cell Signaling Technologies, Danvers, MA). Linacolotide was provided by Ironwood Pharmaceuticals for research application. Ready-to-transduce GIPZ lentiviral shRNA particles (human scrambled and AC6 shRNAs) were obtained from GE Dharmacon. For lentiviral knockdown, human enterospheres were treated with the viral particles (1 × 10⁵ transducing units) for 48 h followed by 48 h of recovery and then again treated for 48 h before the experiment.

Mice

Adcy6+/− mice (Jackson Laboratory) were interbred to obtain Adcy6+/− mice. Adcy6−/− female mice were crossed with male Shh
GFP Cre mice (B6.Cg-Shh<sup>tm1T(EGFP/cre)Cjl/J</sup>, Jackson Laboratory) to finally obtain Adcy6<sup>Δ/Δ</sup> mice (Adcy6<sup>Δ/Δ</sup>/Shhm1<sup>(EGFP/cre)/wt</sup> mice). Typically, 6–12-week-old male mice were used for the study. All of the mice were maintained in a barrier facility at Cincinnati Children’s Hospital Medical Center and were fed normal chow. All procedures were performed in compliance with the Association for Assessment and Accreditation of Laboratory Animal Care and institutional guidelines and were approved by the Cincinnati Children’s Hospital Medical Center institutional animal care and use committee.

**Human biopsy tissues**

Retrieval of patient biopsies for research use was approved by the Cincinnati Children’s Hospital Medical Center institutional review board under IRB 2014-6279.

**Cell lysate preparation, protein purification, and protein-binding assays**

*Crude membrane preparation of Adcy6<sup>Δ/Δ</sup> and Adcy6<sup>Δ/Δ</sup> mouse ileum—* Small intestinal mucosa was cut into small pieces and transferred to a Dounce homogenizer. Next, 1–2 ml of cold homogenization buffer (250 mM sucrose, 1 mM EDTA, 10 mM Tris HCl buffer, pH 7.2) plus protease inhibitors (1 μM aprotinin, 1 μM leupeptin, and 1 mM phenylmethylsulfonyl fluoride) was added to the homogenizer. Tissue was homogenized using ~20 manual up and down strokes a total of three times. Samples were transferred to microcentrifuge tubes. Intact cells, nuclei, and cell debris were removed by centrifugation of the homogenate at 500 g for 10 min at 4 °C. The supernatant was the postnuclear supernatant. The postnuclear supernatant was centrifuged at 145,000 × g for 1 h at 4 °C. The supernatant following the centrifugation contained soluble proteins (cytosolic fraction). The pellet comprised the crude membrane fraction and was washed with homogenization buffer twice. The glass tube was sonicated for 20 s three times at 30-s intervals to dissolve the pellet. Protein concentration was measured using a Bradford assay or by direct absorbance at 280 nm. To perform the binding assay between AC6 C1/C2 and CFTR, 50 μg of the C1/C2 protein was added to 1 mg of protein from whole-cell BHK lysate expressing FLAG-CFTR or FLAG-CFTRHis<sup>10</sup>. Protein complexes were immunoprecipitated using S-protein–conjugated agarose in an overnight binding at 4 °C. Next day, beads were washed three times with PBS plus 0.2% Triton X-100, and samples were eluted in 5× sample buffer. Samples were subjected to traditional Western blotting and probed for CFTR using FLAG antibody.

**Intestinal fluid secretion (in vivo) measurement or closed intestinal loop experiment**

Adcy6<sup>Δ/Δ</sup>/ and Adcy6<sup>Δ/Δ</sup> mice were starved for 24 h before surgery. Mice were anesthetized under isoflurane with optimal O<sub>2</sub> and placed on a warm pad to maintain body temperature. A small abdominal incision was made to expose the distal region of small intestine. Intestinal loops (~2 cm) were exteriorized and isolated (two loops per mouse). The closed loops were then injected with 100 μl of PBS alone or PBS containing CTX (1 and 10 μg) or linacotide (1 μg). The abdominal incision and skin incision were closed with surgical sutures, and the mice were allowed to recover. After 6 h, the mice were sacrificed by CO<sub>2</sub>. Intestinal loops were removed, and loop fluid weight was measured to quantitate net fluid secretion.

**Intestinal crypt isolation and quantitation of fluid secretion in enterospheres**

Preparation of mouse intestinal crypt and quantitation were performed as described previously (19). For fluid secretion measurement with the high-content microscope (Lionheart<sup>TM</sup> FX, Biotek), average organoid area was calculated pre- and post-forskolin treatment.

**Isolation of tracheal epithelial cells and short circuit current (I<sub>sc</sub>) measurements**

The protocol for isolation of tracheal cells was followed as described previously (29). Mouse tracheal cells were polarized on Costar Transwell permeable supports (Cambridge, MA) (filter diameter, 12 mm). After 21 days of culturing, transwells were mounted in an Ussing chamber system (Physiologic Instruments) maintained at 37 °C (30). Epithelial cells were bathed in Ringer’s solution (basolateral side: 140 mM NaCl, 5 mM KCl, 0.36 mM K<sub>2</sub>HPO<sub>4</sub>, 0.44 mM KH<sub>2</sub>PO<sub>4</sub>, 1.3 mM CaCl<sub>2</sub>, 0.5 mM MgCl<sub>2</sub>, 4.2 mM NaHCO<sub>3</sub>, 10 mM HEPES, 10 mM glucose, pH 7.2, [Cl<sup>-</sup>] = 149 mM) and low-CI<sup>-</sup> Ringer’s solution (apical side: 133.3 mM sodium gluconate, 2.5 mM NaCl, 0.36 mM CaCl<sub>2</sub>).
Diarrheagenic complex in cholera

K$_2$HPO$_4$, 0.44 mm KH$_2$PO$_4$, 5.7 mm CaCl$_2$, 0.5 mm MgCl$_2$, 4.2 mm NaHCO$_3$, 10 mm HEPES, 10 mm mannitol, pH 7.2, [Cl$^-$] = 14.8. Cells were treated first with amiloride (50 μM), and, after current stabilization, CFTR was activated by adding 10 μM forskolin on the apical side. CFTRinh-172 (20–50 μM) was added to the apical side to verify CFTR dependence of the currents. Calcium-dependent currents were activated in the presence of ATP (100 μM) added to the apical side at the end of the experiment.

Immunofluorescence and proximity ligation assay

Paraffin-embedded mouse ileum segments were stained for CFTR and AC6 proteins as follows. Slides were deparaffinized in xylene three times, 5 min each, followed by dehydration with ethanol. Antigen retrieval was performed with BORG DeCloaker RTU (Biocare Medical, Concord, CA) in a pressure cooker for 5–10 min. Slides were cooled for 5 min, and tissues were permeabilized with 0.2% Triton X-100 in PBS for 10 min. Tissues were then blocked in 2.5% horse serum overnight. Slides were rinsed with 1× PBS and incubated with rabbit polyclonal AC5/6 (1:25 dilution) and mouse monoclonal CF3 (1:25 dilution) antibodies at 4 °C overnight. Human organoids embedded in Matrigel were fixed with 4% formaldehyde and permeabilized with 0.2% Triton X-100 in PBS for 30 min. Samples were incubated with rabbit polyclonal AC5/6 (1:25 dilution) and mouse CFTR antibody R1104 (1:50 dilution) for 24 h at 4 °C.

For the proximity ligation assay, anti-rabbit (plus) and anti-mouse (minus) Duo link In Situ PLA probes (MilliporeSigma) were added to the samples incubated with anti-AC5/6 and anti-CFTR antibodies as mentioned above. The next steps of PLA assay were completed as described in the manufacturer’s protocol. Slides were examined using a confocal microscope (Olympus FV1200).

FRET microscopy and data analysis

For direct sensitized emission FRET, HT29CL19A cells were transiently transfected with pAMCyan-AC6 and pEYFP-CFTR singly or in combination using Lipofectamine 3000. Single transfected cells were used to acquire cyan- or yellow fluorescent protein—only images for bleedthrough calculations. The corrected FRET (FRETc) was normalized with donor cyan fluorescent protein intensity (FRETc/cyan fluorescent protein), yielding the normalized corrected FRET (N-FRETc), and the intensity of N-FRETc images was presented in pseudocolor and monochrome mode, stretched between the low and high normalization values, according to an intensity-to-color mapped lookup table. All calculations were performed with the Channel Math and FRET modules of SlideBook software version 4.2 (Intelligent Imaging Innovations, Denver, CO).

Measurement of whole-cell cAMP and cGMP

Adcy6$^{fl/fl}$ and Adcy6$^{△/△}$ enterospheres were treated with FSK (10 μM, 1 h), CTX (100 ng, 24 h) and linaclootide (500 nM, 24 h) at 37 °C in the presence of isobutylmethylxanthine. Enterospheres were lysed in 0.1 N HCl, 0.2% Triton X-100 and centrifuged at 800 × g, and the supernatant was collected and used for cAMP- or cGMP-specific ELISA following the manufacturer’s protocol (Enzo Life Science, Farmingdale, NY).

Author contributions—K. A. and A. P. N. conceived and supervised the project. A. T., Y. R., and K. A. designed and performed experiments and together wrote the manuscript. A. P. N. edited the draft manuscript. K. M. assisted in the human enterosphere experiment.

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