SLC26A3 inhibitor identified in small molecule screen blocks colonic fluid absorption and reduces constipation

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SLC26A3 (downregulated in adenoma; DRA) is a Cl⁻/anion exchanger expressed in the luminal membrane of intestinal epithelial cells, where it facilitates electroneutral NaCl absorption. SLC26A3 loss of function in humans or mice causes chloride-losing diarrhea. Here, we identified slc26a3 inhibitors in a screen of 50,000 synthetic small molecules done in Fischer rat thyroid (FRT) cells coexpressing slc26a3 and a genetically encoded halide sensor. Structure-activity relationship studies were done on the most potent inhibitor classes identified in the screen: 4,8-dimethylcoumarins and acetamide-thioimidazoles. The dimethylcoumarin DRA\textsubscript{inh}-A250 fully and reversibly inhibited slc26a3-mediated Cl⁻ exchange with HCO\textsubscript{3}⁻, I⁻, and thiocyanate (SCN⁻), with an IC\textsubscript{50} of ~0.2 μM. DRA\textsubscript{inh}-A250 did not inhibit the homologous anion exchangers slc26a4 (pendrin) or slc26a6 (PAT-1), nor did it alter activity of other related proteins or intestinal ion channels. In mice, intraluminal DRA\textsubscript{inh}-A250 blocked fluid absorption in closed colonic loops but not in jejunal loops, while the NHE3 (SLC9A3) inhibitor tenapanor blocked absorption only in the jejunum. Oral DRA\textsubscript{inh}-A250 and tenapanor comparably reduced signs of constipation in loperamide-treated mice, with additive effects found on coadministration. DRA\textsubscript{inh}-A250 was also effective in loperamide-treated cystic fibrosis mice. These studies support a major role of slc26a3 in colonic fluid absorption and suggest the therapeutic utility of SLC26A3 inhibition in constipation.

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SLC26A3 inhibitor identified in small molecule screen blocks colonic fluid absorption and reduces constipation

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Introduction

SLC26A3, originally named downregulated in adenoma (DRA), is an anion exchanger expressed at the lumen-facing plasma membrane of enterocytes in the colon and small intestine (1–4). SLC26A3 facilitates exchange of various monovalent anions, including Cl−, HCO3−, and thiocyanate (SCN−), as well as the divalent anion oxalate (5–9). SLC26A3-mediated Cl−/HCO3− exchange is involved in electroneutral NaCl absorption, which drives water absorption in the colon and small intestine (4, 8, 10–14). Loss-of-function mutations in SLC26A3 cause congenital chloride-losing diarrhea (CLD) in humans, which is characterized by severe diarrhea that starts in utero (15, 16). In mice, knock-out (KO) of slc26a3 recapitulates CLD, producing diarrhea and metabolic alkalosis (10). SLC26A3 inhibition is, thus, a potential approach to treat constipation by blocking intestinal fluid absorption, which may be effective in the major types of constipation, including opioid-induced constipation (OIC), and in constipation associated with cystic fibrosis (CF) in which the pro-secretory CFTR Cl− channel is defective (17).

SLC26A3 is a member of the SLC26 gene family of anion channels and transporters. The closest SLC26A3 homolog, SLC26A4 (pendrin; 45% amino acid identity), is an anion exchanger in the inner ear, thyroid, inflamed airways, and kidney (6, 9, 18). We previously identified small-molecule pendrin inhibitors that increased airway surface liquid depth in airway epithelial cell cultures and produced a diuretic response in mice, suggesting potential therapeutic utility in inflammatory lung disorders, such as CF and asthma, and in volume-overload edema (19, 20). The second most closely related SLC26A3 homolog, SLC26A6 (putative anion transporter-1 [PAT-1]; 34% amino acid identity), is also an anion exchanger expressed at the apical membrane in the small intestine that, together with SLC26A3, facilitates electroneutral NaCl absorption (21, 22).

We report here the identification, by high-throughput screening, of nanomolar-potency small molecule SLC26A3 inhibitors, their optimization by structure-activity analysis, and their characterization in cellular models. Inhibition of intestinal fluid absorption was demonstrated in closed intestinal loops in mice, and
efficacy was demonstrated in an experimental model of constipation. Inhibition of slc26a3 reduced manifestations of constipation with comparable efficacy to a blocker of intestinal Na⁺ absorption, the NHE3 inhibitor tenapanor, and when coadministered, slc26a3 and NHE3 inhibitors fully reversed constipation.

Results

High-throughput screen to identify slc26a3 inhibitors. Screening was done using a Fischer rat thyroid (FRT) cell line stably expressing murine slc26a3 and a halide-sensitive yellow fluorescent protein (FRT-YFP-slc26a3 cells). FRT cells were used because of their low intrinsic permeability to anions that are transported by slc26a3 and because of their good adherence and rapid growth on uncoated plastic. As diagrammed in Figure 1A, slc26a3-mediated transport was assayed from the kinetics of YFP fluorescence quenching in response to extracellular addition of I⁻ to drive the Cl⁻/I⁻ exchange; transport inhibition reduces the rate of fluorescence decrease. Figure 1B shows representative fluorescence data from inactive and active compounds tested in screening, together with data from vehicle (DMSO) control and cells not expressing slc26a3. For screening, the nonselective chloride channel blocker niflumic acid was used as a positive control, which strongly inhibited slc26a3 at 350 μM. The Z-factor for the slc26a3 screening assay was ~0.7.

To test whether second messengers regulate slc26a3 activity in the transfected FRT cells, experiments were done in which FRT-YFP-slc26a3 cells were preincubated with forskolin (20 μM; to elevate cAMP), 8-bromoguanosine 3’;5’-cyclic monophosphate (100 μM; 8-Br-cGMP; a cell permeable cGMP analog), ATP (100 μM; to elevate cytoplasmic Ca²⁺), phorbol 12-myristate 13-acetate (100 nM; PMA), or 20 mM potassium chloride (high K⁺; to depolarize membrane potential). None of these maneuvers significantly altered electroneutral slc26a3-mediated Cl⁻/I⁻ exchange (Figure 1C). Therefore, inhibitors identified in the screen are likely to inhibit slc26a3 exchange directly rather than acting on upstream signaling mechanisms.

Screening of 50,000 synthetic small molecules at 25 μM identified 136 active compounds that inhibited slc26a3 by >75%. The compounds clustered in several compound classes, including 4,8-dimethylcoumarins, acetamide-thioimidazoles, triazoleamides, and thiazole-thiophenesulphonamides (Classes A–D, respectively) (Figure 2, A and B). A scatter plot of inhibition percentage for a subset of screened compounds, shown in the order as screened (Figure 2A, left), revealed clusters of active compounds as a consequence of grouping of chemical scaffolds in compound plates. On retesting, 36 compounds inhibited slc26a3 by more than 50% at 10 μM (Figure 2A, right), including multiple Class A and Class B compounds.
Figure 2C shows concentration-dependence measurements for inhibition of slc26a3-mediated Cl⁻/I⁻ exchange for a selected Class A (IC₅₀ ~ 4 μM) and Class B (IC₅₀ ~ 9 μM) compound discovered in the primary screen, together with data for niflumic acid (IC₅₀ ~ 60 μM).

Structure-activity relationship studies. Based on potency data and structural considerations, compounds of Classes A and B were prioritized for further study. A total of 176 commercially available Class A analogs and 175 Class B analogs were tested. Figure 3, A and B, summarizes the structural determinants of activity for Class A and Class B compounds (left panels), with concentration-dependence data of selected analogs provided (right panels). Class A compounds are coumarins that are substituted with methyl group on the 4 and 8 positions. Structure-activity analysis showed that the acetic acid moiety at the 3 position (R²) is required for activity, as the corresponding acetate ester was inactive. Replacement of the acetic acid with propanoic acid greatly reduced activity, and nonacid substituents including alkyl and benzyl were inactive. Greatest potency was found for hydroxylation at the 7 position on the coumarin with a benzyl ring (R¹), with 3-chloro, 3-fluoro, and 3-bromo substituents giving greatest potency.

Class B compounds are acetamide-thioimidazoles with substituents on the imidazole (R¹) ring and acetamide linker (R²). For the R¹ group, phenyl gave the best activity, with benzofuran, benzyl, methyl, and acetate reducing activity. Unsubstituted phenyl gave the greatest potency, although 4-methoxy and 4-chloro were also active, while substituents on the 2 position reduced activity. For R², several substituted anilines were active, whereas nonaniline, alkyl, benzyl and allyl groups reduced activity. Anilines substituted with 3-chloro-4-methyl were most active. Other substituents on the aniline ring — including 4-halide, 4-methyl, and 4-methoxy — were active. In general, 3,4-disubstituted and 4-substituted anilines with electron-neutral or donating groups were best, whereas 4-NO₂ and 3,4-disubstituted anilines with electron-withdrawing groups reduced activity.

Supplemental Tables 1 and 2 (supplemental material available online with this article; https://doi.org/10.1172/jci.insight.121370DS1) list IC₅₀ values for selected active Class A and Class B compounds, respectively. DRAₙₐₙ-A250 was the most potent Class A compound with an IC₅₀ of 0.15 μM (Figure 3A, right panel). Several other Class A analogs were also found to be more potent than DRAₙₐₙ-A01, the original compound identified in the primary screen. For Class B, a number of analogs showed greater potency.
than the original active compound (DRA inh-B01, IC\textsubscript{50}~9 \textmu M), with DRA inh-B601 having the lowest IC\textsubscript{50} (~1.5 \textmu M; Figure 3B, right panel). For further studies, DRA inh-A250 was resynthesized (Figure 3C) by Pechmann condensation of 2-methylresorcinol with dimethyl acetylsuccinate to yield 7-hydroxycoumarin (intermediate 1), which was further alkylated with 3-bromobenzyl bromide to give coumarin ester (intermediate 2) (23, 24). Hydrolysis of the coumarin ester gave the corresponding 3-acetic acid-coumarin DRAinh-A250 in good yield (Figure 3C).

**DRAinh-A250 characterization.** Concentration-dependence measurements for inhibition of the slc26a3-mediated exchange of Cl\textsuperscript{-} and HCO\textsubscript{3}\textsuperscript{-}, the physiologically relevant activity of slc26a3, was measured using slc26a3-expressing FRT cells labeled with BCECF as a cytoplasmic pH sensor. Labeled cells were initially incubated in a Cl\textsuperscript{-}-containing HCO\textsubscript{3}\textsuperscript{-} buffered solution, and HCO\textsubscript{3}\textsuperscript{-} influx/Cl\textsuperscript{-} efflux with cytoplasmic alkalinization was produced by addition of a gluconate-containing HCO\textsubscript{3}\textsuperscript{-} buffered solution. The increase in cytoplasmic pH (~0.05 pH units/min) was inhibited by DRA inh-A250 with IC\textsubscript{50} ~0.1 \textmu M (Figure 4A). Concentration-dependence measurements for inhibition of slc26a3-mediated Cl\textsuperscript{-}/SCN\textsuperscript{-} exchange gave an IC\textsubscript{50} of ~0.3 \textmu M (Figure 4B). The kinetics of DRA inh-A250 inhibition, measured by incubation of FRT-YFP-slc26a3 cells with 1 \textmu M DRAinh-A250 for different times prior to initiation of the Cl\textsuperscript{-}/I\textsuperscript{-} exchange, showed a half-time (t\textsubscript{1/2}) for inhibition of ~30 seconds (Figure 4C). To confirm full reversibility, FRT-YFP-slc26a3 cells were incubated with 1 \textmu M DRA inh-A250 for 10 minutes and then washed 3 times prior to assay Cl\textsuperscript{-}/I\textsuperscript{-} exchange (Figure 4C, inset). Inhibition of (human) SLC26A3 by DRAinh-A250 was also tested, using a HEK cell model expressing SLC26A3 and YFP. SLC26A3-mediated Cl\textsuperscript{-}/I\textsuperscript{-} exchange was inhibited by DRA inh-A250 with IC\textsubscript{50} ~0.25 \textmu M, with complete inhibition at higher DRAinh-A250 concentrations (Figure 4D).

**Figure 3.** Structure-activity relationships for Class A and Class B slc26a3 inhibitors. (A) Class A. Left panel: core structure of the Class A inhibitor is boxed; active and inactive substituents are listed. Right panel: concentration-dependence measurements for selected Class A compounds, including DRA\textsubscript{inh}-A250 (mean ± SEM, n = 3). (B) Class B, showing core structure (boxed) with active and inactive substituents (left panel) and concentration-dependence measurements (right panel; mean ± SEM, n = 3). (C) Route for synthesis of DRA\textsubscript{inh}-A250.
DRAinh-A250 selectivity. To investigate DRAinh-A250 selectivity, transport assays (full experimental details in Supplemental Methods) were done on a panel of SLC26-family homologs and other relevant epithelial ion transporters and channels. Preincubation of cells with DRAinh-A250 at 10 μM did not alter Cl–/I–-exchange activity of (murine) pendrin (slc26a4), PAT-1 (slc26a6), or slc26a9, as shown in original fluorescence time course data (Figure 5A) and summary data (Figure 5B). DRAinh-A250 at 10 μM also did not alter the activity of the Cl– channels CFTR and TMEM16A. In separate studies, to investigate possible action of DRAinh-A250 on multiple ion transporters and channels in a human epithelium, human bronchial epithelial (HBE) cells were studied by short-circuit current (Isc), with successive additions of amiloride (to block the epithelial sodium channel; ENaC), forskolin, CFTR inh-172 (to block CFTR), and ATP (to activate the Ca2+-activated chloride channel; CaCC) (Figure 5C). Pretreatment for 20 minutes with 10 μM DRAinh-A250 did not alter Isc responses, indicating that DRAinh-A250 did not affect ENaC, CFTR, or CaCC activity, or the activity of other transporters — including K+ channels and the Na-K-Cl cotransporter NKCC1 — that are required to support their activity. In a preliminary toxicity study, incubation of T84 cells with 10 μM DRAinh-A250 for 24 hours did not cause toxicity (Supplemental Figure 2).

DRAinh-A250 inhibits fluid absorption in the distal colon in mice. Closed intestinal loop studies in mice were done to study effects of DRAinh-A250 on fluid absorption. DRAinh-A250 administration directly in isolated distal colonic loops completely prevented the reduction in the loop weight/length ratio at 60 minutes, a direct measure of fluid absorption (Figure 6A, top panel). Changes in luminal pH were also determined as a semiquantitative measure of slc26a3-facilitated HCO3– secretion/Cl– absorption, with slc26a3 inhibition predicted to reduce luminal fluid alkalinization. For these experiments, a phosphate-buffered solution (PBS) without HCO3–/CO2 was used to avoid confounding effects of luminal carbonic anhydrases. DRAinh-A250 reduced alkalinization of luminal fluid in the distal colonic loops.
at 60 minutes, consistent with inhibition of Cl⁻/HCO₃⁻ exchange (Figure 6A, bottom panel). Administration of the NHE3 (Na⁺/H⁺ exchanger 3) inhibitor tenapanor (25) had no effect on the weight/length ratio or luminal pH in distal colonic loops. Results with coadministration of DRA inh-A250 and tenapanor were similar to those with DRA inh-A250 alone. These results support the conclusion that slc26a3 is the predominant proabsorptive transporter in murine distal colon, whereas NHE3 does not contribute significantly to fluid absorption in this segment.

DRA inh-A250 does not alter jejunal fluid absorption in mice. Administration of DRA inh-A250 to midjejunal loops did not produce significant changes in the loop weight/length ratio, whereas fluid absorption was abolished in tenapanor-treated loops (Figure 6B, top panel). DRA inh-A250 did not alter acidification of jejunal fluid at 30 minutes, with tenapanor preventing acidification and producing slight alkalinization (Figure 6B, bottom panel). Loops coadministered DRA inh-A250 and tenapanor showed a loop weight/length ratio and luminal fluid pH that were similar to treatment with tenapanor alone. These results provide evidence against a significant role of slc26a3 in fluid absorption in mouse jejunum and support the conclusion that NHE3 is a predominant proabsorptive transporter in this segment.

DRA inh-A250 reduces loperamide-induced constipation in WT and CF mice. Orally administered DRA inh-A250 at 5 mg/kg partially prevented the loperamide-induced reduction in stool weight, pellet number, and water content in WT mice (Figure 7, A and B). Tenapanor given orally at 5 mg/kg also partially prevented loperamide-induced reduction in stool weight, pellet number, and water content. The magnitude of these responses is similar to that seen with very high doses of the approved drug linaclotide (26, 27). Remarkably, coadministration DRA inh-A250 and tenapanor completely prevented loperamide-induced constipation, suggesting an additive effect of slc26a3 and NHE3 inhibition.
on stool hydration (Figure 7B) (see Discussion and Figure 8 for proposed mechanisms). Importantly, DRAinh-A250 was also effective in a loperamide-induced constipation model in CF mice having loss of function of the prosecretory Cl⁻ channel CFTR (Figure 7C). To study reversibility in vivo, WT mice were treated with DRAinh-A250 as in Figure 7A, and loperamide was administered at 24 hours. The loperamide effect was similar to that in mice without preexposure to DRAinh-A250 (stool weight 15 ± 5 mg, pellet number 1.5 ± 0.6, stool water content 31% ± 4%, mean ± SEM, n = 4 mice). The anticonstipation effect of DRAinh-A250 is, thus, not the consequence of a toxic effect on the gastrointestinal tract.

**Discussion**

The goal of this study was to identify potent and selective small molecule inhibitors of slc26a3 in order to test their utility for treatment of constipation and to investigate the role of slc26a3 in intestinal fluid absorption. High-throughput screening identified several classes of slc26a3 inhibitors, which — following structure-activity analysis and optimization — produced the 4,8-dimethylcoumarin DRAinh-A250 with an IC₅₀ of 100–200 nM for inhibition of slc26a3-mediated anion exchange. Importantly, DRAinh-A250 did not inhibit homologous slc26a-family anion exchangers or relevant intestinal transporters, and it was nontoxic in cell culture (Supplemental Figure 2). Studies in mice demonstrated DRAinh-A250 efficacy in a loperamide model of constipation and provided novel data that clarified the mechanisms of intestinal fluid absorption.

Constipation is a common problem, with an estimated prevalence of ~15% in the US population (28, 29). The most common types of chronic constipation include chronic idiopathic constipation (CIC), OIC, and constipation-predominant irritable bowel syndrome (IBS-C). Current treatment approaches for chronic constipation include lifestyle and dietary changes, over-the-counter laxatives (e.g., stimulants of intestinal contraction, osmotic agents), and recently approved prescription drugs that promote intestinal fluid secretion.
Three prosecretory drugs that activate Cl– channels have been approved: lubiprostone, a prostaglandin derivative that indirectly activates CFTR and chloride channel 2 (ClC-2); linaclotide, a peptide agonist of the guanylate cyclase C receptor that indirectly activates CFTR; and plecanatide, a uroguanylin analog that also acts as an agonist of the guanylate cyclase C receptor (30). Our group recently reported CFTR activators with marked pro-secretory action and improved efficacy over lubiprostone and linaclotide in experimental animal models of constipation (26, 27, 31).

Inhibition of intestinal fluid absorption represents an alternative approach to increase stool hydration in constipation. Recent evaluation of the NHE3 inhibitor tenapanor in a Phase 3 clinical trial for IBS-C showed limited efficacy, with 27% combined pain and stool pattern response rate for the treated group versus 18.7% for placebo (30). NHE3 is expressed in the small intestine and proximal colon, but not in distal colon (12, 32). The limited clinical efficacy of NHE3 inhibition for constipation might be due to unimpaired downstream fluid absorption in the distal colon. As an alternative antiabsorptive approach, we targeted SLC26A3, the major Cl–/HCO3– exchanger in the colon, and demonstrated comparable efficacy to tenapanor in a murine model of constipation. Theoretically, SLC26A3 inhibition might be more effective than NHE3 inhibition for constipation therapy, as it blocks absorption in the distal colon where stool is dehydrated to its final form.

Remarkably, we found additive actions of NHE3 and slc26a3 inhibitors in preventing loperamide-induced constipation. Our results suggest the efficacy of SLC26A3 inhibitors alone for various types of constipation and their combination with tenapanor in patients that do not respond adequately to monotherapy.

In CF subjects, gastrointestinal-related problems are common, including meconium ileus (~15% occurrence in neonates), constipation (up to 47% lifetime prevalence), and distal intestinal obstructive syndrome (~15% lifetime prevalence) (17, 33). Impaired function of the prosecretory CFTR Cl– channel is believed to be the cause of these disorders. We showed previously that an orally administered

Figure 7. Oral DRAinh-A250 reduces loperamide-induced constipation in WT and cystic fibrosis mice. (A) Mouse model of constipation produced by loperamide. (B) Effects of DRAinh-A250 (5 mg/kg) and tenapanor (5 mg/kg), individually and together, on 3-hour stool weight, number of pellets, and stool water content in loperamide-treated WT mice (mean ± SEM, n = 4–7 mice per group). Vehicle control shown for comparison. (C) Study as in A done in cystic fibrosis mice lacking functional CFTR (mean ± SEM, n = 4–5 mice per group). *P < 0.05, **P < 0.01, ***P < 0.001, one-way ANOVA with post-hoc Newman-Keuls multiple comparisons test.
CFTR activator was effective in experimental mouse models of constipation, and — as expected — CFTR activation was ineffective in CF mice lacking functional CFTR (26). Here, we demonstrated efficacy of a slc26a3 inhibitor in CF mice in a loperamide model of constipation, which supports the conclusion that intestinal slc26a3 function is not impaired in CF. SLC26A3 inhibition therapy may, thus, be beneficial in promoting stool hydration and treating constipation in human CF subjects — and potentially in preventing or treating meconium ileus in the neonatal period.

There are at least 3 pathways for absorption of Na⁺ and Cl⁻ in the intestine: nutrient-coupled Na⁺ absorption, electroneutral NaCl absorption, and electrogenic Na⁺ absorption (13). Nutrient-coupled Na⁺ absorption in the small intestine involves sodium-glucose transporters (SGLTs) and Na⁺-amino acid cotransporters in which Cl⁻ is absorbed through a paracellular pathway. Electroneutral NaCl absorption in the small intestine and proximal colon involves Na⁺/H⁺ exchangers (NHE2/SLC9A2 and NHE3) and Cl⁻/HCO₃⁻ exchangers (SLC26A3 and SLC26A6), which are thought to function in tandem to absorb NaCl.
Electrogenic Na⁺ absorption in the distal colon involves ENaC. The major absorptive pathways in the jejunum and distal colon are diagrammed in Figure 8A. NHE3 and NHE2 are expressed in the small intestine and proximal colon; NHE3 (SLC9A3) is the dominant Na⁺/H⁺ exchanger, as it can compensate for loss of NHE2 (34). SLC9A3 mutations cause congenital Na⁺ diarrhea with mild metabolic acidosis, and slc9a3−/− mice have mild diarrhea with luminal fluid alkalinization (35). However, compensatory increased ENaC activity in the distal colon of slc9a3−/− mice is thought to limit the severity of diarrhea. SLC26A3 is mainly expressed in the distal colon and duodenum, with low expression in the jejunum and ileum (3, 4), whereas SLC26A6 is mainly expressed in the small intestine (36). In agreement with these findings, we found — in mouse jejunum — that slc26a3 inhibition had no effect, while NHE3 inhibition blocked fluid absorption, whereas — in mouse distal colon — slc26a3 inhibition blocked absorption while NHE3 inhibitor had no effect. These results provide pharmacological evidence that slc26a3 plays little or no role in NaCl absorption in murine small intestine where SLC26A6 is likely the dominant Cl⁻/HCO₃⁻ exchanger, in agreement with earlier studies (36, 37).

Though much of the current understanding of intestinal NaCl absorption comes from studies using KO mice, it is recognized that KO mice can produce confounding secondary phenotypes because of altered gene expression patterns or organ development. For example, with slc26a4 (pendrin), we found remarkable differences in renal physiology with pharmacological inhibition, as compared with slc26a4−/− mice (20). SLC26A3 loss of function in humans or KO mice produces CLD with marked diarrhea, whereas the inhibitor studied here had only partial efficacy in a mouse model of constipation. This difference could be due to partial efficacy of the inhibitor or, perhaps more likely, to secondary phenotypes in the KO mouse. Our finding that slc26a3 inhibition blocked fluid absorption only in the distal colon supports the latter possibility, as does the additive effect seen on coadministration of slc26a3 inhibitor and tenapanor. An additional contributing factor may be regulatory interactions among NHE3, Slc26a3, and CFTR that may involve the NHE-regulatory factor (NHERF), SLC26 sulfate transporter antisigma (STAS) domain, and the CFTR regulatory domain (13). As many CLD-associated SLC26A3 mutations occur in the STAS domain, the severe CLD phenotype might be a consequence of both the loss of SLC26A3 absorption and dysregulation of NHE3 and CFTR activities. Prior studies in subjects with CLD and slc26a3−/− mice showed impaired fluid absorption in the jejunum (4, 38), whereas closed-loop studies here suggested that slc26a3 is of minimal importance in fluid absorption in this segment. Further studies will be required to test these ideas.

The closed-loop method used in Figure 6 to measure intestinal fluid absorption is an extension of the commonly used closed-loop method to measure intestinal fluid secretion in which fluid transport is deduced from the weight/length ratio of closed intestinal loops at a specified time after fluid injection. Compared with the conventional method involving in vivo intestinal perfusion and use of a volume marker to deduce fluid transport, the closed-loop method is technically more simple and robust, and it provides data with relatively low variance. Validation studies reported in Supplemental Figure 1 show the expected effect of sodium and chloride ion substitution in preventing fluid absorption, and the expected effect of glucose in increasing fluid absorption in jejunum. We note that use of this method requires an initial time course study to select an appropriate time window and, for quantitation of absolute fluid transport rates, determination of the relationship between loop weight/length ratio and luminal fluid volume.

Though the results here provide proof-of-concept for the therapeutic utility of SLC26A3 inhibition in promoting stool hydration, several caveats are noted. There is a theoretical relationship between SLC26A3 inhibition and tumorigenicity based on the original findings of SLC26A3 downregulation in adenoma and manifestation of an expanded colonic epithelial proliferative zone and surface mucosal hyperplasia in slc26a3−/− mice (1, 10). Notably, however, CLD subjects with SLC26A3 mutations do not manifest an increased incidence of gastrointestinal cancers (39, 40). Reduced SLC26A3 activity is observed in ulcerative colitis, and single-nucleotide polymorphisms in SLC26A3 may be a risk factor for development of ulcerative colitis (41, 42); however, loss of SLC26A3 function does not produce intestinal inflammation, as only 6% of CLD subjects report symptoms of Crohn’s disease or unspecified colitis (15, 40). Renal insufficiency is observed in ~25% of CLD subjects, although this likely results from delayed diagnosis and consequent chronic volume and salt depletion (15). SLC26A3 mutations are also associated with male subfertility, although available evidence suggests that impaired SLC26A3-CFTR interaction and consequent impaired CFTR activity are responsible (43). In general, with appropriate salt-replacement therapy, the long-term health in CLD is good, suggesting the absence of significant tumor and nongastrointestinal problems with SLC26A3 mutation (40).
There have been a few prior reports on the biological properties of the 7-benzoxy-4,8-dimethyl-3-acetic acid-coumarin scaffold, as found in DRAinh-A250. Similar coumarins were reported as inhibitors of *Bacillus anthracis* and *Staphylococcus aureus* replicative DNA helicase (24). Another 3-acetic acid-coumarin was reported to inhibit the oncogene signal transducer and activator of transcription (Stat3) activation by inhibiting the nuclear translocation of phosphorylated Stat3 (44). In general, coumarins are plant-derived natural products with diverse inflammatory, antitumor, antibacterial, and antifungal properties (45). The physico-chemical properties of DRAinh-A250 include the presence of multiple hydrogen bond acceptors, a molecular mass of 417 Da, a logP value of 4.6, and a topologic polar surface area of 73 Å², all of which are consistent with drug-like properties, including oral bioavailability. In addition, the 3-acetic-acid-coumarin scaffold does not belong to promiscuous binders known as panassay interference compound molecules. Synthetically, the 3-acetic-acid-coumarin scaffold can be prepared in 3–5 steps from commercially available starting materials, which allows facile synthesis of targeted analogs.

In summary, we found that slc26a3 is a druggable target in which small molecule inhibitor therapy may be beneficial for antiabsorptive therapy of the common types of constipation, as well as in constipation associated with CF.

**Methods**

**Molecular biology.** Vectors containing complementary DNA (cDNA) for slc26a3, SLC26A3, slc26a6, and slc26a9 were purchased from Origene and manipulated using standard techniques (see Supplemental Methods). To generate a cell line for screening, slc26a3 was subcloned into pIRESpuro3 (Clontech). Human SLC26A3 was subcloned into pLVX-IRES-mCherry (Clontech), and other slc26 family members were subcloned into vectors that were generated to coexpress the halide sensitive EYFP-H148Q/I152L/F46L (YFP).

**Cell culture and transfections.** FRT epithelial cells (UCSF Cell Culture Facility), HEK cells (UCSF Cell Culture Facility), and HBE cell cultures (UCSF Cystic Fibrosis Research Development Program) were cultured using standard methods (19). For slc26a3 inhibitor screening, an FRT cell line virally transduced to express YFP (46) was transfected with pIRESpuro3-slc26a3, selected using 0.15 μg/ml puromycin, and a clonal cell line (termed FRT-YFP-slc26a3) was isolated. To generate a cell line enriched for expression of SLC26A3, HEK293 cells were transduced with lentiviral particles generated from the pLVX-IRES-mCherry-SLC26A3 vector and then transfected to express YFP. FRT cells expressing YFP and slc26a4 (pendrin) or CFTR were previously described (20, 26). FRT cells were transfected with Lipofectamine 2000 (Thermo Fisher Scientific), and HEK cells were transfected with NanoFect (Alstem).

**Screening and synthetic chemistry.** High-throughput screening to identify slc26a3 inhibitors using FRT-YFP-slc26a3 cells was done with a semiautomated Beckman Coulter platform with FLUOstar OMEGA plate readers (BMG Labtech), essentially as described for discovery of pendrin inhibitors (19). Initial screening was done of ~50,000 drug-like synthetic small molecules (ChemDiv) at a concentration of 25 μM using a Cl⁻/I⁻ exchange protocol to assay slc26a3 function. Following initial screening, compound analogs were purchased (ChemDiv) to generate structure-activity data. The lead slc26a3 inhibitor was resynthesized and confirmed by 1H and 13C NMR, and >95% purity was confirmed by liquid chromatography–mass spectrometry (LC-MS).

**Selectivity studies.** YFP-based assays of CFTR, pendrin, and TMEM16A were as described (19, 26, 47, 48). HEK cells transfected to express YFP and SLC26A3, slc26a6, or slc26a9 were functionally assayed using a Cl⁻/I⁻ exchange protocol. Assays of the Cl⁻/HCO₃⁻ exchange were done using BCECF as an intracellular pH indicator as described for pendrin (19). Iₛₑₑ measurements in well-differentiated HBE cell cultures were done as described (19).

**Murine models.** Closed-loop models and the loperamide-induced model of constipation were done in mice as described (26, 27). Mice were housed and bred in the UCSF Laboratory Animal Resource Center.

**Statistics.** Details of statistical methods are provided in individual figure legends. One-way ANOVA was used for multiple comparisons, with Newman-Keuls post hoc multiple comparisons test. Student's *t* test (two-tailed) was used to test significance for comparisons of 2 data sets. *P* < 0.05 was considered significant.

**Study approval.** Animal experiments were approved by the UCSF IACUC. Procurement and use of HBE cell cultures was approved by the UCSF Human Research Protection Program IRB. For further details, see online Supplemental Methods.
Author contributions
PMH, OC, SL, JAT, AAR, and PWP performed in vitro and in vivo experiments. PMH, OC, JAT, and PWP analyzed data. PMH, OC, SL, and ASV designed experiments. PMH, OC, SL, PWP, and ASV wrote the manuscript. PMH and ASV developed the original concept for this study.

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