Cholangiocytes derived from human induced pluripotent stem cells for disease modeling and drug validation

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The study of biliary disease has been constrained by a lack of primary human cholangiocytes. Here we present an efficient, serum-free protocol for directed differentiation of human induced pluripotent stem cells into cholangiocyte-like cells (CLCs). CLCs show functional characteristics of cholangiocytes, including bile acids transfer, alkaline phosphatase activity, γ-glutamyl-transpeptidase activity and physiological responses to secretin, somatostatin and vascular endothelial growth factor. We use CLCs to model in vitro key features of Alagille syndrome, polycystic liver disease and cystic fibrosis (CF)-associated cholangiopathy. Furthermore, we use CLCs generated from healthy individuals and patients with polycystic liver disease to reproduce the effects of the drugs verapamil and octreotide, and we show that the experimental CF drug VX809 rescues the disease phenotype of CF cholangiopathy in vitro. Our differentiation protocol will facilitate the study of biological mechanisms controlling biliary development, as well as disease modeling and drug screening.

Cholangiocytes are the main site of dysfunction in cholangiopathies, a diverse group of bile duct disorders that includes inherited diseases such as CF-associated cholangiopathy, developmental diseases such as Alagille syndrome and autoimmune diseases such as primary biliary cirrhosis, as well as drug- and toxin-induced conditions. Cholangiopathies carry significant morbidity and mortality, accounting for up to a third of adult and more than 70% of pediatric liver transplantations. However, research on their pathophysiology has been limited by poor access to primary biliary tissue, difficulties in culturing primary cholangiocytes in vitro and inadequate animal disease models.

The capacity of human induced pluripotent stem cells (hiPSCs) to proliferate indefinitely in culture and differentiate into a broad spectrum of cell types makes them well suited to in vitro disease modeling. Early methods for deriving cholangiocytes from hiPSCs were based on spontaneous differentiation and had limited characterization of the resulting cells. Despite recent advances toward guided differentiation of hiPSCs to CLCs, existing protocols show poor differentiation efficiency (<31%), and the derived cells differ considerably from primary biliary tissue in their transcriptional profiles. Furthermore, in vitro-generated cholangiocytes have not been shown to reproduce key functions of bona fide cholangiocytes, such as enzymatic activity (for example, alkaline phosphatase (ALP) and γ-glutamyl transferase (GGT)), responses to hormonal stimuli (secretin and somatostatin) and chloride transfer through cystic fibrosis transmembrane regulator (CFTR) activity. Demonstration of these properties is essential for recapitulating cholangiopathies and studying the effects of therapeutic agents. Finally, current systems diverge from the physiological pathways controlling biliary development in vivo, limiting their value for developmental studies.

Here we report a stepwise method for cholangiocyte differentiation that recapitulates native biliary development (Fig. 1a). The quality, functionality and purity of the resulting CLCs are substantially higher than those of cells generated by previous methods (see Supplementary Note, Supplementary Methods and Supplementary Fig. 1 for detailed comparison).

RESULTS
Cholangiocyte progenitors generated from hiPSCs
We focused first on the generation of bipotent hepatoblasts, the common progenitor of hepatocytes and cholangiocytes. To achieve this goal,

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we adapted our established hepatic hiPSC differentiation protocol\(^{11,12}\). Cells generated with the adapted protocol after 12 days of differentiation expressed hepatoblast markers, including α-fetoprotein (AFP), HNF4A, HNF1B, TBX3 and CK19 (KRT19) (Fig. 1b,d), and had the potential to differentiate toward both the hepatic ( Supplementary Fig. 2a–c) and biliary (Fig. 1c,d and Supplementary Table 1) lineages. To differentiate these hepatoblast-like cells into cholangiocyte progenitors (CPs), we interrogated pathways reported to control early biliary specification\(^{10}\) (Supplementary Fig. 3a–c and data not shown) and found that activin in combination with retinoic acid suppressed the expression of the hepatoblast markers AFP, HNF4A and TBX3 (Supplementary Fig. 3c). Addition of fibroblast growth factor (FGF)-10 along with activin and retinoic acid induced the expression of the early biliary specification markers SOX9, HNF1B and CK19 (Fig. 1c,d)\(^{10}\), resulting in a population in which 75.1% of cells were CK19\(^{+}\)Sox9\(^{+}\) (Supplementary Fig. 4a). Flow cytometry analyses identified the majority of the remaining cells as Sox9\(^{−}\)AFP\(^{+}\) hepatoblasts (Supplementary Fig. 4a), explaining the presence of reduced but detectable levels of AFP in our culture (Fig. 1d). Mature biliary markers such as secretin receptor (SCR), also known as \textit{SCTR}, somatostatin receptor 2 (SSTR2), aquaporin1 (AQP1) and anion exchanger 2 (AE2, also known as \textit{SLC4A2}) were not expressed (Fig. 1d). Thus, activin, retinoic acid and FGF10 promote the differentiation of hepatoblast-like cells into early CLCs or CPs \textit{in vitro}. To promote maturation of the CPs, we used three-dimensional culture conditions known to induce cholangiocyte maturation through organoid formation\(^{7–9}\). CPs grown in these conditions proliferated rapidly, organized into ring-like structures after 48–72 h and within 5–7 d gave rise to cystic organoids and branching tubular structures (Fig. 2a,b) bearing primary cilia (Fig. 2c) similar to those of primary cholangiocytes. The organoids expressed biliary markers, including CK7 (KRT7), CK18 (KRT18), CK19, HNF1B, γ-glutamyl transferase (GGT1), Jagged1 (JAG1), NOTCH2, CFTFR, SCR, SSTR2, AQP1 and AE2 (Figs. 1d and 2d) at levels similar to those in primary cholangiocytes (Fig. 1d and Supplementary Fig. 5). Transcriptomic analyses of common bile duct primary tissue and cells at key stages of our differentiation protocol (Fig. 2e and Supplementary Table 2) using Euclidean hierarchical clustering revealed that CLCs are distinct from earlier developmental stages (Fig. 2e and Supplementary Fig. 6), cluster closely with primary common bile duct cholangiocytes (Pearson correlation coefficient for CLCs versus primary cholangiocytes \(r = 0.747\), CLCs versus hepatoblasts \(r = 0.576\), CLCs versus hiPSCs: \(r = 0.474\)) and express both mature (SSTR2, ALPL, KRT7) and fetal (SOX9) biliary markers, as shown by immunofluorescence and quantitative PCR (QPCR) analyses (Fig. 2e and Supplementary Fig. 6). These results confirm that CPs can differentiate into cells resembling biliary epithelial cells when grown in three-dimensional culture.

**Figure 1** Generation of CPs from hiPSCs.
(a) Overview of the protocol used to differentiate hiPSCs to CLCs. DE, definitive endoderm; FP, foregut progenitors; HB, hepatoblasts; HC: hepatocytes; RPMI, Roswell Park Memorial Institute medium; SB, SB-33542; HGF, hepatocyte growth factor; RA, retinoic acid; EGF, epidermal growth factor; FGF, fibroblast growth factor.
(b) Immunofluorescence analyses demonstrating the expression of key hepatoblast markers, as indicated, in day 12 hepatoblasts. DAPI, 4’,6-diamidino-2-phenylindole dihydrochloride. Scale bars, 100 μm. (c) Immunofluorescence analyses demonstrating the expression of early biliary markers by immature CPs on day 16. Scale bars, 100 μm. (d) Gene expression profile of hiPSC-derived cells at key stages of biliary differentiation and of primary cholangiocytes (PCs). \(n = 4\) biological replicates for each stage of differentiation, \(n = 3\) independent samples for PCs. Center line, median; box, interquartile range (IQR); whiskers, range (minimum to maximum). IQR is not calculated for PCs, as \(n = 3\). Asterisks represent statistical significance of differences between HBs, CPs and CLCs; \# indicates lack of statistical significance (\(P > 0.05\)) (one-way ANOVA with Tukey correction for multiple comparisons). Values are relative to the housekeeping gene porphobilinogen deaminase (\textit{PBGD}, also known as \textit{HMBS}).

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The efficiency of our differentiation protocol can be estimated from the observation that, by day 26, 1 × 10⁶ hiPSCs produced 74.4 × 10⁶ cells, of which 57.28 × 10⁶ expressed mature biliary markers (74.4 × 10⁶ cells generated × 77% (average differentiation efficiency across three lines, s.d. = 6.5%) = 57.28 × 10⁶ CLCs) (Supplementary Fig. 4a, b). More specifically, 74.5% of the resulting cells expressed both the biliary marker Sox9 and the mature biliary marker CK7 (Supplementary Fig. 4a). A further 7.5% of the cells, consistent with immature cholangiocytes, expressed Sox9 but not CK7 (Supplementary Fig. 4a); 15% of the cells expressed both AFP and albumin (Supplementary Fig. 4a), indicating the presence of a small fraction of hepatocytes in our culture conditions and explaining the detection of low AFP levels by QPCR analyses (Fig. 1d). The remaining 3% of the cells were not characterized further. These results were confirmed on three independent hiPSC lines (Supplementary Fig. 4a).

Next we characterized the functionality of the generated organoids. In vivo, cholangiocytes reabsorb bile acids and modify the composition of canalicular bile through a series of secretory and resorptive processes regulated by intracellular calcium signaling. Native biliary epithelial cells have ALP and GGT activity and proliferate in response to stimuli such as vascular endothelial growth factor (VEGF). The secretory potential of CLCs generated in vitro was confirmed using rhodamine 123, a fluorescent substrate for the cholangiocyte surface glycoprotein multidrug resistance protein-1 (MDR1). Rhodamine 123 was actively secreted in the lumen of CLC organoids, and luminal dye accumulation was prevented by the MDR1 inhibitor verapamil (Fig. 3a-c), confirming MDR1-dependent transfer of rhodamine 123. The capacity of CLCs to interact with bile acids through the apical salt and bile transporter (ASBT) was demonstrated by showing active export of the fluorescent bile acid cholyly-syl-fluorescein (CLF) from the lumen of CLF-loaded organoids compared to controls loaded with fluorescein isothiocyanate.

Figure 2 Generation of CLCs from CPs. (a,b) Immunofluorescence (left) and light microscopy (right) images of CLC organoids (day 26) demonstrating the formation of cystic (a) and branching (arrows) tubular structures (b). Scale bars, 100 µm. (c) Transmission electron microscopy images (right) and immunofluorescence analyses (left) for acetylated α-tubulin demonstrating the presence of cilia (arrowhead). The immunofluorescence image was acquired from the bottom of a large cystic organoid using confocal microscopy. Scale bars, 500 nm and 100 µm, respectively, L, lumen. (d) Immunofluorescence analyses demonstrating the expression of early and mature biliary markers in day 25 CLCs as indicated. Scale bars, 100 µm. (e) Euclidean hierarchical clustering analysis focusing on the genes that define the transcriptional signature of CLCs and HBs (4,963 genes differentially expressed in CLCs versus hiPSCs or in HBs versus hiPSCs). For each probe, standard scores (z-scores) indicate the differential expression measured in number of s.d. from the average across all samples. CLCs cluster closer to and present higher correlation coefficient with common bile duct cholangiocytes (CBD) used as a primary control, rather than to HBs or hiPSCs (Pearson, CLCs versus PCs r = 0.747, CLCs versus HBs r = 0.576, CLCs versus hiPSCs r = 0.474). Representative genes are indicated. Data correspond to biological triplicates.
CLCs model development of the human biliary system

To investigate potential applications of our system for developmental studies, we characterized signaling pathways that control organoid formation in vitro as compared with native duct development. First, we interrogated activin–TGF-β signaling in view of its pivotal role in physiological biliary specification and tubulogenesis. We blocked the activity of TGF-β, which is normally present in Matrigel, using the activin receptor inhibitor SB-431542. SB-431542 completely abrogated organoid formation, confirming that activin–TGF-β signaling is a key regulator of organoid formation in our system.

We performed similar analyses for Notch signaling during biliary specification of hepatoblasts to CPs and CLC organoid formation in vitro. Deregulation of Notch signaling is associated with Alagille syndrome, characterized by a paucity of bile ducts. To explore the hypothesis that blocking Notch signaling would impair organoid formation in vitro, we first characterized the activity of the Notch pathway in our system. Notch activation results in cleavage and nuclear translocation of its intracellular domain. Immunofluorescence analysis with antibodies specific to the cleaved Notch intracellular domain (NICD) confirmed the presence of active NICD, with increased nuclear localization in CPs and CLC organoids. The expression of NOTCH2, as well as its ligand JAG1 and its downstream target HES1, were also increased in both stages compared to hepatoblasts, consistent with pathway activation (CPs versus hepatoblasts: NOTCH2, P < 0.001; JAG1, P < 0.001; HES1, P < 0.05; CLCs versus CPs or hepatoblasts: P < 0.0001, one-way ANOVA with Tukey correction for multiple comparisons). Inhibition of Notch signaling in three-dimensional culture conditions using the γ-secretase inhibitor L-685,458 blocked cleavage of the NICD, suppressed HES1, NOTCH2 and JAG1 expression in CPs and CLCs.}

![Supplementary Videos 1 and 2](image-url) ASBT expression was confirmed by QPCR and immunofluorescence analyses (Supplementary Fig. 7a,b). Furthermore, CLCs responded to acetycholine and ATP stimuli by increasing intracellular calcium (Fig. 3g and Supplementary Videos 3 and 4), demonstrated increased proliferation in response to VEGF stimulation (51% increase in expansion, P < 0.0001, two-tailed t-test) (Fig. 3h,i) and exhibited GGT and ALP activities similar to those of primary controls (GGT activity: 160% of human serum activity, P < 0.0001, one-way ANOVA with Dunnett correction for multiple comparisons) (Fig. 3j,k). Together, these observations confirm that our hiPSC-derived CLCs display a range of functions of the native biliary epithelium.
expression (Fig. 4c) and blocked organoid formation (Fig. 4e,f), confirming the importance of Notch signaling to the generation of organoids incorporating a luminal space in our system.

These results reinforce previous findings obtained in mice by demonstrating the importance of activin–TGF-β and Notch signaling pathways in human cholangiocyte specification24. They underline the potential of our culture system for studying human biliary tree development in vitro.

CLCs validate drugs for polycystic liver diseases

Polycystic liver diseases are characterized by multiple cystic lesions in the liver arising from fetal cholangiocytes25,26. Intraluminal fluid secretion and cholangiocyte proliferation result in cyst expansion and liver impairment owing to space-occupying effects27,28. We explored the use of CLC organoids to identify compounds that might reduce cyst size in polycystic liver disease. The secretory activity of cholangiocytes is increased by the hormone secretin and reduced by the hormone somatostatin, resulting in changes in duct size. Octreotide, a synthetic analog of somatostatin, is used clinically to restrict cyst size in polycystic liver disease28–32. CLC organoids express both SCR and STTR2 (Figs. 1d and 5a,b), suggesting that these pathways might be functional in our cells. Accordingly, secretin increased organoid size (6.1% average diameter increase, P < 0.01, one-way ANOVA with Dunnett correction for multiple comparisons), whereas somatostatin and octreotide decreased organoid size, compared to untreated controls (7.9% and 4.9% average diameter decrease, respectively; P < 0.001 and P < 0.05, respectively; one-way ANOVA with Dunnett correction for multiple comparisons) (Fig. 5c–d and Supplementary Videos 5–7).

Furthermore, octreotide negated the effects of secretin and decreased intracellular cyclic AMP (45% of somatostatin response; P = 0.001, one-way ANOVA with Dunnett correction for multiple comparisons), in keeping with previous studies26,30 (Fig. 5e). To further test the effects of octreotide on disease-specific CLCs, we differentiated hiPSCs derived from a patient with polycystic liver disease (Supplementary Fig. 9) to CLCs (Fig. 5f). Octreotide treatment reduced organoid size (4.86%, P < 0.0001, one-way ANOVA with Dunnett correction for multiple comparisons) (Fig. 5g,h), reproducing the effects of the drug in vitro.

CLCs model CF liver disease

The autosomal recessive disorder CF is caused by mutations in the CFTR gene, encoding a cell-surface chloride transporter33,34. CFTR mutations in cholangiocytes result in reduced intraluminal chloride secretion, increased bile viscosity and focal biliary cirrhosis secondary to bile plugs occluding the intrahepatic bile ducts35–37. To model CF biliary disease in vitro, we generated hiPSCs from skin fibroblasts of a patient homozygous for the most common CF mutation, ΔF508 (CF-hiPSCs)38, and differentiated them into CLCs. CF-hiPSC-derived CLCs (CF-CLCs) expressed markers (Fig. 6a) and displayed functional characteristic (Fig. 6b) of biliary epithelial cells. Transcription of the CFTR gene was confirmed using QPCR (Fig. 6a), and immunofluorescence analyses detected minimal CFTR protein expression (Fig. 6c), in agreement with studies reporting very rapid endoplasmic reticulum degradation of the misfolded protein39. We also used the fluorescent chloride indicator N-(6-methoxyquinolinyl)acetoethyl ester (MQAE)40 to monitor intracellular and intraluminal chloride secretion and CFTR function (Fig. 6d–e, Supplementary Fig. 10a–c). We also used an assay to monitor intracellular chloride secretion (Supplementary Fig. 10d) and a fluorescent chloride indicator N-(6-methoxyquinolinyl)acetoethyl ester (MQAE)40 to monitor intracellular and intraluminal chloride secretion (Supplementary Fig. 10d) and a fluorescent chloride indicator N-(6-methoxyquinolinyl)acetoethyl ester (MQAE)40 to monitor intracellular and intraluminal chloride secretion.
concentration. Wild-type (WT) CLC organoids appropriately modified intracellular chloride in response to media with varying chloride concentrations, whereas no change was observed in CF-CLCs (Fig. 6d,e), confirming the absence of functional CFTR in these cells.

Next we investigated the effects of the experimental CF drug VX809 (ref. 41) in the context of biliary disease. VX809 stabilizes CFTR, corrects folding defects in patients with the ΔF508 mutation and increases CFTR functionality in lung cells42. Incubation of CF-CLCs with VX809 for 48 h increased CFTR function analyzed by MQAE to a level similar to that of WT-CLCs (Fig. 6d,e). This effect was negated by CFTR inhibitor 172, confirming that the phenotypic rescue of CF-CLCs by VX809 depended on improved CFTR function (Fig. 6d,e).

Given the association between chloride and fluid secretion in cholangiocytes33, we studied the impact of VX809 on organoid size. CF-CLC organoids treated with VX809 increased in size compared to their untreated counterparts (5.6% mean diameter increase, P = 0.001, two-tailed t-test) (Fig. 6f,g). This observation confirmed that VX809 increases CFTR function and improves intraluminal fluid secretion, suggesting a previously unreported therapeutic effect for this drug in the context of CF liver disease.

DISCUSSION

We present a protocol for the generation of CLCs from hiPSCs that addresses the limitations of previously published differentiation methods by more closely recapitulating natural bile duct development (Fig. 1a). Our results confirm the importance of FGF10 and activin–TGF-β for early biliary specification, as previously described in vitro10,19,43, and reveal a role for retinoic acid in this process, at least in vitro. We show that the combination of these factors under chemically defined, serum-free conditions is sufficient to promote the specification of hiPSC-derived hepatoblasts into CPs with high efficiency. Production of CPs, mimicking physiological biliary development, has not been demonstrated previously, to our knowledge, and likely explains the efficiency of our culture system in generating CLCs that closely resemble bona fide cholangiocytes at the transcriptional (Supplementary Fig. 1 and Supplementary Tables 3 and 4) and functional (Supplementary Table 3) levels.

Our protocol will enable various applications, such as developmental studies, disease modeling, therapeutic target validation and drug screening. Accordingly, we used patient-derived hiPSCs to model polycystic and CF liver disease and applied these models to reproduce the effects of the therapeutic compounds verapamil, octreotide and VX809. VX809 has already completed phase 2a clinical trials42 for CF, but its effects on CF-associated cholangiopathy have not been described, to our knowledge. Our results suggest that CLCs provide a suitable system for identifying new therapeutic agents, which is particularly
important given the lack of high-throughput drug screening platforms for cholangiopathies. CLCs may also contribute to tissue engineering of livers or liver organoids that incorporate a biliary system for the treatment of end-stage disease.

METHODS

Methods and any associated references are available in the online version of the paper.

Accession codes. ArrayExpress: E-MTAB-2965.

Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

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AUTHOR CONTRIBUTIONS

E.S.: design and concept of study, execution of experiments and data acquisition, development of protocols and validation, generation, collection and interpretation of data, production of figures, manuscript writing, editing and final approval of manuscript. M.C.d.B., F.A.C.S.: technical support, execution of experiments. P.M.: bioinformatics and statistical analyses. A. Bertero: bioinformatics analyses. K.S.-P., E.S., E.M.: primary tissue provision. T.H.K., J.A.B., W.T.H.G., S.D., A. Baker, A.K., G.J.A.: critical revision of the manuscript for important intellectual content. N.R.F.H.: design and concept of study, study supervision, generation
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COMPETING FINANCIAL INTERESTS
The authors declare competing financial interests: details are available in the online version of the paper.

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ONLINE METHODS

Generation of hiPSC lines. All hiPSC lines used here were derived and characterized previously by our lab.11,12,44 Briefly, the lines used were generated from human skin fibroblasts and peripheral blood (ethics reference no. 08/H0311/201 for Hertfordshire Regional Ethics Committee (REC) and 09/H0304/77 for National Research Ethics Service (NRES) Committee East of England, Cambridge East), using the Yamanaka approach.6,9 The CF fibroblasts were obtained from the Coriell cell repository. The lines were authenticated using SNIP arrays and regularly tested negative for mycoplasma contamination.

Culture of hiPSCs. Human iP cells were maintained in defined culture conditions as previously described11,12,44, using activin-A (10 ng/ml) and b-FGF (12 ng/ml).

Differentiation of hiPSCs into cholangiocyte progenitors. hiPSCs were differentiated into foregut progenitor cells (FPs) as previously described12,44. Bipotent hepatoblasts were generated by culturing FPs in RPMI (Gibco, Invitrogen) plus B27 supplemented with SB-431542 (10 µM, Tocris Bioscience) and BMP4 (50 ng/ml) for 4 d. To induce biliary specification, we cultured hepatoblasts for another 4 d in the presence of RPMI (Gibco, Invitrogen) plus B27 supplemented with FGF10 (50 ng/ml, Peprotech), activin-A (50 ng/ml) and retinoic acid (3 µM, Sigma-Aldrich).

Maturation of cholangiocyte progenitor cells to CLCs and organoid formation in 3D culture. Human CPs were passaged using Cell Dissociation Buffer (Gibco, Life Technologies, catalog number 13151-014) and suspended at a density of 8 × 104 cells/ml, in a mixture of 40% Matrigel (BD Biosciences, catalog number 356237) and 60% William’s E medium (Gibco, Life Technologies) supplemented with 10 mM nicotinamide (Sigma-Aldrich), 17 mM sodium bicarbonate (Sigma-Aldrich), 0.2 mM 2-phospho-l-ascorbic acid trisodium salt (Sigma-Aldrich), 6.3 mM sodium pyruvate (Invitrogen), 14 mM glucose (Sigma-Aldrich), 20 mM HEPS (Invitrogen), ITS+ premix (BD Biosciences), 0.1 µM dexamethasone (R&D Systems), 2 mM Glutamax (Invitrogen), 100 U/ml penicillin, 100 µg/ml streptomycin and 20 ng/ml EGF (R&D Systems). A 50-µL droplet of the cell suspension was added in the center of each well of a 24-well plate; the gel was allowed 1 h at 37 °C to solidify and then overlayed with William’s E medium with supplements. The medium was changed every 48 h and the cells were cultured for a total of 10 d.

Using the same methodology, we have been able to culture CLC organoids in formats ranging from 6- to 96-well plates. To generate large numbers of CLC organoids, multiple 50-µL droplets were added in a well of a 6-well plate or a 10-cm dish. To provide a large number of wells compatible with high-throughput screening and large-scale experiments, 30-µL droplets were added per well of a 96-well plate. In both cases, the gel was allowed 1 h at 37 °C to solidify and then overlaid with William’s E medium with supplements.

Inhibition of activin and Notch signaling in 3D culture and assessment of organoid formation. Human CPs were suspended at a density of 8 × 104 cells/ml, in a mixture of 40% Matrigel and 60% William’s E medium (Gibco, Life Technologies) with supplements as described above. The cell suspension was distributed in three aliquots of equal volume. Each aliquot received no further supplementation and was used as a positive control. The second aliquot was further supplemented with SB-431542 for the inhibition of activin–TGF-β signaling. 50 µM L-685,458 (Tocris Biosciences) was added in the third aliquot for the inhibition of Notch signaling. Each aliquot was distributed in three aliquots of equal volume. One aliquot received no further supplementation and was used as a positive control. The second aliquot was added to the medium overlaying the Matrigel on a daily basis. After a total of 10 d in 3D culture, the total number of cysts in four random wells of a 24-well plate was counted for each condition by a researcher blinded to sample identity. Error bars represent s.d.

Flow cytometry analyses. hiPSCs, FPs, HBs and CPs were dissociated to single cells using Cell Dissociation Buffer (Life Technologies). The cells were then counted using a hemocytometer and fixed using 4% PFA for 20 min at 4 °C. Cell staining and flow cytometry analyses were performed as previously described44.

Primary cholangiocytes. Frozen primary human cholangiocytes derived from common bile duct were obtained from Celprogen (catalog number 36755-11). The cells were derived from donors negative for hepatitis B surface antigen, HIV 1 and 2, syphilis, hepatitis B virus, human T lymphotropic viruses 1 and 2, hepatitis C virus, West Nile virus and Trypanosoma cruzi. Each frozen ampule (1.2 × 106 cells) of 1 × 106 viable cells was thawed with gentle agitation in a 37 °C water bath. The cells were transferred to a sterile centrifuge tube and 1 µl of prewarmed Human Cholangiocyte Cell Culture Complete Growth Medium (Celprogen, catalog number M36755-115) was added. The ampule was washed twice with Human Cholangiocyte Cell Culture Complete Growth Medium and the cells were centrifuged at 100g for 7 min. The pellet was resuspended in 500 µl of Human Cholangiocyte Cell Culture Complete Growth Medium. The cells were distributed equally to two wells of a 6-well plate precoated with Human Cholangiocyte Cell Culture Extracellular Matrix Celprogen, catalog number M36755-11-6-wells) with 2 µl of Human Cholangiocyte Cell Culture Complete Growth Medium per well. The cells were incubated at 37 °C in a 5% CO2, humidified incubator for 48 h and the medium was changed once. After another 24 h, when the wells were confluent, the cells were lysed and RNA was extracted as previously described11,12, or the cells were fixed with 4% PFA as previously described11,12 for ALP staining. QPCR analyses for biliary markers (Fig. 1d) and ALP staining (Fig. 3k) confirmed the nature and cholangiocyte properties of the cells.

Primary biliary tissue. Primary biliary tissue (bile duct) was obtained from an organ donor from whom organs were being retrieved for transplantation. A section of the bile duct was excised during the multi-organ retrieval operation after obtaining informed consent from the donor’s family (REC reference number: 09/H0306/73, NRES Committee East of England–Norfolk). The tissue was homogenized using a tissue homogenizer and RNA was extracted as previously described41.

Immunofluorescence, RNA extraction and quantitative real-time PCR. Immunofluorescence, RNA extraction and QPCR were performed as previously described9. A complete list of the primary and secondary antibodies used is provided in Supplementary Table 5. A complete list of the primers used is provided in Supplementary Table 6. All QPCR data correspond to four independent biological replicates, with the exception of the primary CDB cholangiocytes from Celprogen, where three independent samples were used.

For immunofluorescence in 3D Matrigel cultures, the organoids were fixed in Matrigel with 4% PFA for 20 min at room temperature, to avoid Matrigel liquefaction. The samples were permeabilized and blocked with 0.1% Triton X-100 and 10% donkey serum respectively for 30 min and incubated with primary antibody in 1% donkey serum overnight at 4 °C. The following day the samples were washed three times with PBS for 45 min per wash, incubated with secondary antibody in 1% donkey serum for 60 min at room temperature and washed again three times in PBS. Hoechst 33258 was added to the first wash. All immunofluorescence images were acquired using a Zeiss Axiovert 200M inverted microscope or a Zeiss LSM 700 confocal microscope. ImageJ 1.48k software (Wayne Rasband, NIH, USA, http://imagej.nih.gov/ij) was used for image processing. Changes in brightness or contrast during processing were applied equally across the entire image.

For RNA extraction in 3D Matrigel cultures, the organoids were washed once with PBS and 1 µl of ice-cold dispaee was added per well of a 24-well plate.
The Matrigel was mechanically dissociated, transferred to a Falcon tube and kept on ice to allow the combination of low temperature and disperse digestion to liquefy the Matrigel. After 10 min the cells were centrifuged at 1,600 r.p.m. (44g) for 3 min and the supernatant was aspirated. The pellet was washed once with PBS and the centrifugation step repeated. Finally, the supernatant was aspirated and 350 µl of RNA lysis buffer were added to the pellet. RNA was extracted from the lysate using a kit (GenElute Mammalian Total RNA Miniprep Kit, Sigma-Aldrich, catalog number RTN350) according to the manufacturer's instructions.

**Microarrays.** 500 ng of total cellular RNA was amplified and purified using the Illumina TotalPrep-96 RNA Amplification kit (Life Technologies) according to the manufacturer's instructions. Three biological replicates for each condition were analyzed. Biotin-labeled RNA was then normalized to a concentration of 150 ng/µl and 750 ng were hybridized to Illumina Human-12 v4 BeadChips for 16 h (overnight) at 58 °C. Following hybridization, BeadChips were washed and stained with streptavidin-Cy3 (GE Healthcare). BeadChips were then scanned using the BeadArray reader and image data were processed using Genome Studio software (Illumina). The raw and processed microarray data are available on ArrayExpress (accession code E-MTAB-2965).

**Microarray analysis.** Probe summaries for all arrays were obtained from the raw data using the method “Making Probe Summary” in Genome Studio (Illumina). These values were transformed (variance stabilized) and quantile normalized using the R/Bioconductor package lumi. Standard lumi QC procedure was applied and no outliers were identified.

Differential expression between pairs of conditions was evaluated using the R/Bioconductor package limma. A linear model fit was applied, and the procedure was applied and no outliers were identified.

Differential expression between pairs of conditions was evaluated using the method of Benjamini and Hochberg to correct the P values. Probes that failed to fluoresce above background in both conditions were removed. Differentially expressed probes were selected using a cutoff of adjusted P < 0.01 and absolute fold-change > 2.

Probes differentially expressed between hiPSCs and CLCs or hiPSCs and HBs (representing the aggregate transcriptional signature of CLCs and HBs) were selected for Euclidean hierarchical clustering using Perseus software (MaxQuant). Standard scores (z-scores) of the log2-normalized probe expression values across the different conditions were calculated and used for this analysis.

**Rhodamine 123 transport assay.** CLC organoids were incubated with 100 µM rhodamine 123 (Sigma-Aldrich) for 5 min at 37 °C and then washed with William’s E medium three times. Fresh William’s E medium with supplements was added following the third wash. The organoids were incubated at 37 °C for another 40 min. To demonstrate that rhodamine 123 transfer indeed increased in absorbance over 10 min by 353. Error bars represent s.d. Multiple mean absorbance comparisons (CLCs versus substrate, CLCs versus MEFs, CLCs versus human serum) were performed using one-way ANOVA with Dunnett correction for multiple comparisons.

**Fluorescence intensity comparisons were performed using a two-sided Student’s t-test.**

**Cholyl-lysyl-fluorescein transport assay.** CLC organoids were loaded with 5 µM cholyl-lysyl-fluorescein (CLF, Corning Incorporated) for 30 min at 37 °C and then washed with Leibovitz’s medium (Life Technologies) three times. Following completion of the third wash, time-lapse images were taken using a confocal microscope for 10 min. To demonstrate that the changes in CLF fluorescence intensity observed were secondary to active export of CLF from the organoid lumen, the experiment was repeated with 5 µM unconjugated fluorescein isothiocyanate (FITC) (Sigma-Aldrich) as a control. Around 1,000 fluorescence measurements were made between the organoid interior and exterior.

Fluorescence in the organoid lumen was normalized to background measured in the surrounding external area. Each experiment was repeated in triplicate. Error bars represent s.d. Mean fluorescence intensity comparisons were performed using a two-sided Student’s t-test.

**Measurement of intracellular calcium.** Intracellular calcium signaling, regulated by stimuli such as acetylcholine and ATP, constitutes a key second messenger for cholangiocytes. CLC organoids were incubated with 25 µM of the calcium indicator Fluo-4 a.m. (Life Technologies) for 60 min at 37 °C and washed three times with William’s E medium. Fresh William’s E medium with supplements was added following the third wash. The organoids were stimulated with 1 µM acetylcholine (Sigma-Aldrich) or 30 µM ATP (Sigma-Aldrich), while time-lapse images were taken. Each measurement was repeated in triplicate. To calculate the number of cells responding to stimulation, the number of cells loaded with Fluo-4 a.m. was counted by two different researchers before the start of the experiment. Following stimulation with ATP or acetylcholine the number of responding cells (increase in fluorescence) was also counted and responsiveness was expressed as the ratio of responding cells to the total number of cells loaded with Fluo-4 a.m. For the statistical approach for smoothing the data and plotting bands for the confidence limits, please see “Statistical analyses.”
Supplementary Videos 5–7 were made by taking images before and after treatment at 2-min intervals until organoid size stabilized.

cAMP levels. cAMP levels were measured in triplicate using the cAMP-Glo assay kit (Promega) according to the manufacturer’s instructions and a P450-Glomax 96 microplate luminometer (Promega). Error bars represent s.d. Statistical significance was calculated using one-way ANOVA with Dunnett correction for multiple comparisons.

CFTR activity. CFTR activity was measured as previously described. Briefly, MQAE is a fluorescent dye quenched by the presence of chloride but not affected by other anions or pH changes. Chloride transfer across the cell membrane is mainly regulated by CFTR in cholangiocytes. Therefore, cells with a functional CFTR will respond to a chloride challenge by rapidly increasing intracellular (and intraluminal, in the case of organoids) chloride concentration, thereby quenching MQAE fluorescence. Chloride depletion using a nitrate solution will have the opposite effect.

Cells were incubated with 8 mM MQAE (Life Technologies) and 5 µM forskolin for 4 h at 37 °C. Standard Ringer’s solution containing NaCl, KCl, CaCl2, MgCl2 glucose and HEPES was used to provide a chloride challenge expected to increase intracellular chloride in the presence of functional CFTR. Modified Ringer’s solution consisting of NaNO3, KNO3, CaNO3, MgNO3, glucose and HEPES was used to promote chloride efflux and deplete intracellular chloride. Live pictures were captured every minute as each solution was added. To demonstrate the effect of VX809 on CFTR functionality, CLC organoids were incubated with 30 mM VX809 (Selleck) for 48 h. The assay was repeated as described above in the presence and absence of 7 µM CFTR inhibitor 172 (Sigma-Aldrich) to confirm the specificity of the compound for CFTR. Intracellular fluorescence intensity was measured in three random areas from the wall of each organoid using ImageJ software and normalized to the minimum fluorescence value for each area. Error bars represent s.d.

Cytochrome p450 activity. Cyp3A4 activity was measured using the p450-Glo assay kit (Promega) according to the manufacturer’s instructions and a P450-Glomax 96 microplate luminometer (Promega).

Timing of experiments on CLC organoids. All the experiments and characterization with regards to CLCs were performed on CLC organoids after 10 d of 3D culture unless stated otherwise.

Statistical analyses. All statistical analyses were performed using GraphPad Prism 6 or the R statistical environment. For comparison between two mean values, a two-sided Student’s t-test was used to calculate statistical significance. For comparison between multiple values, one-way ANOVA was used with Tukey correction for multiple comparisons when comparing multiple values to each other (for example, QPCR plots) or Dunnett correction for multiple comparisons when comparing multiple values to a single value (for example, functional assays where the values are compared to a negative control). The normal distribution of our values was confirmed using the Kolmogorov-Smirnov test where appropriate. Further information on the statistical analysis of our data is provided in Supplementary Table 7 (test used for each experiment or analysis, test statistic, degrees of freedom, P value).

To smooth our data for generating the curves in Figure 3g, we used functional data analysis theory implemented in the R package “Ida” (http://cran.r-project.org/web/packages/Ida/index.html). First, we represented our data values (3 replications at each fluorescence intensity measurement) using 60 equidistant B-spline basis functions, and roughness penalties in the second derivative (λ = 1). We used the functions create.bspline.basis and smooth.fd over the interval 1–100 s. Then we evaluated the mean and the s.d. of the functional data objects using the R functions mean.fd and sd.fd.

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