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Concurrent Activation of Kras and Canonical Wnt Signaling Induces Premalignant Lesions That Progress to Extrahepatic Biliary Cancer in Mice

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ABSTRACT

Biliary cancer has long been known to carry a poor prognosis, yet the molecular pathogenesis of carcinoma of the extrahepatic biliary system and its precursor lesions remains elusive. Here we investigated the role of Kras and canonical Wnt pathways in the tumorigenesis of the extrahepatic bile duct (EHBD) and gall bladder (GB). In mice, concurrent activation of Kras and Wnt pathways induced biliary neoplasms that resembled human intraductal papillary-tubular neoplasm (ICPN) and biliary intraepithelial neoplasia (BiILN), putative precursors to invasive biliary cancer. At a low frequency, these lesions progressed to adenocarcinoma in a xenograft model, establishing them as precancerous lesions. Global gene expression analysis revealed increased expression of genes associated with c-Myc and TGFβ pathways in mutant biliary spheroids. Silencing or pharmacologic inhibition of c-Myc suppressed proliferation of mutant biliary spheroids, whereas silencing of Smad4/Tgfbr2 or pharmacologic inhibition of TGFβ signaling increased proliferation of mutant biliary spheroids and cancer formation in vivo. Human ICPNs displayed activated Kras and Wnt signals and c-Myc and TGFβ pathways. Thus, these data provide direct evidence that concurrent activation of the Kras and canonical Wnt pathways results in formation of ICPN and BiILN, which could develop into biliary cancer.

Significance: This work shows how dysregulation of canonical cell growth pathways drives precursors to biliary cancers and identifies several molecular vulnerabilities as potential therapeutic targets in these precursors to prevent oncogenic progression.

Introduction

Biliary cancer, including cholangiocarcinoma (CCA) and gall bladder carcinoma (GCC), accounts for approximately 3% of all adult malignancies (1). According to their anatomical location, CCAs are classified as either intrahepatic (iCCA), perihilar (pCCA), or distal extrahepatic biliary duct (dCCA; ref. 2). Recently, the WHO classified iCCA into two subtypes, small duct type and large duct type, based on their different etiologies and clinical behaviors (3). Risk factors for CCA include liver flukes, primary sclerosing cholangitis, hepatolithiasis, choleodochal cyst, and viral hepatitis (4). Most biliary cancer is not diagnosed until the late stages, such as the locally advanced or metastatic phases. Most of the currently available chemotherapies for biliary cancer are not sufficiently effective, and the 5-year survival rate of biliary cancer is only 5% to 15% (5, 6). To improve its poor prognosis, the development of novel diagnostic methods and therapeutic strategies is urgently needed. Accordingly, it is important to clarify the molecular mechanism underlying the development of biliary cancer and its precursor lesions.

CCA is considered to develop mainly from two distinct types of precancerous lesions, biliary intraductal papillary neoplasm (BiILN) and intraductal papillary neoplasm of the bile duct (IPNB; refs. 7–9). IPNB-like lesions in the gall bladder (GB) are commonly referred to as intracholecystic papillary-tubular neoplasm (ICPN; ref. 10). However, it is not known if these lesions are bona fide precursors of biliary cancer. Furthermore, the molecular pathogenesis of these precursor lesions is still not well understood. This is partly because these precursor lesions have only been recently defined and genetically engineered mouse models (GEMs) of these lesions have not yet been well established. GEMs, particularly those that apply the inducible CreER system, are robust tools used in the study of the molecular pathogenesis of human diseases and in the preclinical assessment of novel therapeutic strategies. To date, only a limited number of GEMs for tumors of the extrahepatic biliary system including extrahepatic biliary duct (EHBD) and GB have been...
reported, partly because there is no known tissue-specific driver CreER mouse line for the extrabiliary biliary system.

Although previous whole-genome sequencing (WGS) studies have demonstrated a complex mutational landscape of biliary cancer (11, 12), little is known about the molecular biological significance of each genetic alternation. Several recent studies of murine biliary cancer models have revealed that Pten (13), Smad4/ TgfβR2 (14), ErbB-2A (15), Kras, and p53 (16) play prominent roles in biliary tumorigenesis. However, given the heterogeneous characteristics of biliary cancer and the variety of mutations, the molecular function of other genes in biliary tumorigenesis has to be taken into consideration. WGS studies have revealed that the incidence of mutations in Kras (16.5%) and Wnt-related genes, including APC (7.1%), RNF43 (4.7%), and CTNNB (1.3%), are relatively frequent in biliary cancer (17). The incidence of Kras mutation is relatively high in both extrabiliary cholangiocarcinoma and intrahepatic cholangiocarcinoma (11). Mutation of Kras occurs in 34.2% of BilIN cases (18). According to the International Cancer Genome Consortium, biliary cancer is classified into four subgroups based on mutation status, copy number variants, gene expression, gene methylation, and prognosis. Subtype 2 biliary cancer is reported to have a high gene expression of WNT5B and Wnt-related genes, (19).

Therefore, in this study, we aimed to investigate the role of the Kras and Wnt pathways in the tumorigenesis of the extrabiliary biliary system including the EHBD and GB. We found that concurrent activation of the Kras and canonical Wnt pathways in adult biliary epithelial cells induced ICPN and BilIN, which could develop into biliary cancer, in mice. This study provides a novel GEM of ICPN and BilIN, establishing them as precursors to biliary cancer. Furthermore, mechanistically, c-Myc contributed to tumorigenesis, whereas the Tgfβ pathway inhibited it.

### Materials and Methods

#### Mice

Experimental animals were generated by crossing Hnf1b-CreERT2 mice (a gift from Jorge Ferrer, Imperial College; ref. 22), KrasG12D mice (a gift from David Tuveson, Cold Spring Harbor Laboratory, Cold Spring Harbor, NY), Ctnnb1lox(ex3)/+ mice (generated as described previously; ref. 23), Rosa26-LacZ mice (purchased from The Jackson Laboratory, JAX strain 003309), Rosa26-tdTomato mice (purchased from The Jackson Laboratory, JAX strain 007914), Dlk1CreERT2-MES-EGFP mice (24), and Bmi1-CreERT (purchased from Jackson Laboratory, JAX strain 010531). Genotyping primers are listed in Supplementary Table S1.

For the induction of Cre-mediated recombination, tamoxifen (Sigma-Aldrich) was administered by oral gavage to Hnf1b-CreERT2 mice at a dose of 400 μg/g body weight per gavage. Time courses are outlined in Figs. 1B, D, and 2A. For induction of Cre-mediated recombination, 2 mg of tamoxifen was intraperitoneally injected to Dlk1CreER, KrasG12D, Ctnnb1lox(ex3)/+ and Bmi1CreER; KrasG12D, Ctnnb1lox(ex3)/+ mice once a day for 4 consecutive days. Time courses are outlined in Supplementary Fig. S2A. All experiments involving mice were approved by the animal research committee of Kyoto University.

#### Histologic analyses and immunostaining

Mouse tissues were prepared by perfusion with 4% paraformaldehyde at sacrifice, fixed in 4% paraformaldehyde overnight, dehydrated in 70% ethanol for 2 days, embedded in paraffin, and cut into 5 μm sections. Paraffin-embedded sections were stained with hematoxylin and eosin (H&E). For X-gal staining, isolated bile ducts were incubated for 1 hour in fixative solution (4% formaldehyde, 0.2% glutaraldehyde, 0.02% Nonidet P-40, 5 mmol/L EGTA, 2 mmol/L MgCl2) at 4°C. The tissues were washed twice in PBS and then incubated in a β-galactosidase substrate (5 mmol/L K3[Fe(CN)6], 5 mmol/L K4[Fe(CN)6]×3H2O, 2 mmol/L MgCl2, 0.02% Nonidet P-40, 0.1% Na deoxycholate, and 1 mg/mL X-gal in PBS) in the dark overnight. After fixation, the sections were counterstained with Nuclear Fast Red (Sigma-Aldrich, catalog no. N3020). For IHC, antigen retrieval was performed by boiling the sections in 10 mmol/L citric acid buffer (pH 6.0) or EDTA buffer (pH 8.0) for 15 minutes at 98°C. Blocking was performed by incubating the sections with a blocking solution (Dako, catalog no. X0909). For the primary antibodies, the incubation was performed overnight at 4°C in a humidified chamber. Secondary antibody incubation was conducted for 1 hour at room temperature. Immunoperoxidase labeling was performed using the VECTASTAIN Elite ABC Standard Kit (Vector Laboratories, catalog no. PK-6100). The sections were then colored with diaminobenzidine solution (Dako, catalog no. K3468) and counterstained with hematoxylin. For immunofluorescence, sections were incubated with a primary antibody overnight at 4°C. The sections were then incubated with a fluorescence-conjugated secondary antibody (Invitrogen) for 1 hour at room temperature. The primary antibodies used in this study are listed in Supplementary Table S2.

#### Three-dimensional culture

Organoid/spheroid cultures were established from murine EHBD and GB. Mouse EHBD and GB were minced into small pieces and digested with dissociation buffer at 37°C for 15 to 20 minutes. The dissociation buffer consisted of DMEM with 10% FBS, 1 mg/mL collagenase D (Roche Diagnostics Deutschland GmbH), 0.5 mg/mL dispase (Invitrogen), and 40 μg/mL DNase (Roche Diagnostics Deutschland GmbH). The tissue suspension was passed through a 100-μm/L cell strainer and washed twice by DMEM with 10% FBS. Washed cells were mixed with 15 μL Matrigel per well on a 48-well plate. Passage was performed at 1:4 to 1:8 split ratios once per week. For passaging, organoids were digested in TrypLE (Invitrogen) for 10 minutes at 37°C. Centrifuged cells were mixed with 15 μL Matrigel per well on a 48-well plate. Expansion Medium (L-WNR Conditioned Medium-based, detailed below) was typically used. Two weeks before the extraction of total RNA or protein, organoids were incubated with Analysis Medium (without FBS, detailed below) and every organoid was passaged at 1.0 × 104 cells/well. 10048-F4 (Selleck Chem, catalog no. S7153) was used as a c-Myc inhibitor in vitro. SB431542 (Tocris Bioscience, catalog no.1614) was used as a Tgfβ inhibitor in vitro.

#### Expansion medium (L-WNR conditioned medium-based)

The Expansion Medium comprised a 50% conditioned medium of L-WNR cell line secreting Wnt3a, R-spondin3, and Noggin (L-WRN CM) and every organoid was cultured in 100 U/mL EGF, 100 U/mL FGF2, 100 μmol/L Glutamax, and every organoid was cultured in 100 U/mL EGF, 100 U/mL FGF2, 100 μmol/L Glutamax until confluence. For further expansion, organoids were passaged once a day for 4 consecutive days. Time courses are outlined in Supplementary Fig. S2A. All experiments involving mice were approved by the animal research committee of Kyoto University.

#### Analysis medium (without FBS)

The Analysis Medium (without FBS) comprised Ad-DMEM/F12 (Invitrogen) supplemented with B27 and N2, 1× Glutamax

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(Invitrogen), 10 mmol/L HEPES, 1 × penicillin-streptomycin (Invitrogen), 1 mmol/L N-acetylcysteine (Sigma-Aldrich), 10 mmol/L nicotinamide (Sigma-Aldrich), 50 ng/mL EGF (Peprotech), 1 µg/mL Rorondin-1 (Peprotech), 50 ng/mL Wnt3a (Peprotech), 100 ng/mL Noggin (Peprotech), 100 ng/mL FGF10 (Peprotech), and 10 µmol/L Y-27632 (Tocris Bioscience).

**Proliferation assay performed by PrestoBlue**

For the cell proliferation assay, spheroids (5.0 × 10^3) were seeded with 10 µL Matrigel per well in 96-well plates. Cell proliferation was evaluated using PrestoBlue Cell Viability Reagent (Invitrogen; catalog no. A13262) following the manufacturer’s protocol. On days 1, 3, and 5, the absorbance at 560 and 595 nm was measured using a plate reader.

The values were calculated as suggested by the product protocol: (absorbance at 560 nm spheroid well — absorbance 595 nm spheroid well) — (absorbance 560 nm empty well — absorbance 595 nm empty well)/(absorbance 560 nm spheroid day 1 — absorbance 595 nm spheroid well day 1) — (absorbance 560 nm empty well day 1 — absorbance 595 nm empty well day 1).

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**Figure 1.**

*Hnf1bCreER* is a suitable CreER mouse line with high efficiency to perform genetic manipulation in the extrahepatic biliary tract epithelium. A, Immunostaining for Hnf1β in the EHBD, IHBD, and GB of wild-type mice (n = 3). B, Tamoxifen administration schema for lineage tracing experiments using *Hnf1bCreER*, Rosa26-LacZ mice. C, Macroscopic (top) and microscopic (bottom) images of X-gal staining in *Hnf1bCreER*, Rosa26-LacZ mice 2 days (n = 5), 6 weeks (n = 5), and 12 weeks (n = 5) after the last tamoxifen administration. D, Tamoxifen administration schema for the experiments using *Hnf1bCreER*, Rosa26-tdTomato mice. E, Immunostaining for RFP (left) and panCK (right) in *Hnf1bCreER*, Rosa26-tdTomato mice at 2 days (n = 3) after the last tamoxifen administration. All black or white scale bars, 50 µm.
Figure 2.
Concurrent activation of Kras and Wnt pathways induced ICPN and BillN that resemble human ICPN and BillN. **A,** Tamoxifen administration schema for the experiments using Hnf1bCreER(H), Hnf1bCreER(H) KrasG12D (HK), Hnf1bCreER(H) Ctnnb1lox(ex3)/(Hb), and Hnf1bCreER(H) KrasG12D; Ctnnb1lox(ex3)/(HKb) mice (n = 5–7 mice/group). All mice were sacrificed 4 weeks after the last tamoxifen administration. **B,** Macroscopic images of the EHBD in H, HK, Hb, and HKb mice 4 weeks after the last tamoxifen administration. **C,** H&E staining (low power field) of the EHBD and GB in HKb mice 4 weeks after the last tamoxifen administration. **D,** From the top, H&E staining (high power field), immunostaining for Ki67, p-ERK, β-catenin, and coimmunostaining for DAPI (blue), GS-II (green), and CK19 (red) in the EHBD and GB in H, HK, Hb, and HKb mice 4 weeks after the last tamoxifen administration. Neoplastic changes were also observed in the peribiliary glands of EHBD in HKb mice (arrowheads). Hyperchromasia, nuclear stratification, and partial loss of nuclear polarity were observed in epithelial cells of BillN and ICPN lesions in HKb mice (insets). All black or white scale bars, 50 μm.
Proliferation and apoptosis assay by immunostaining

Immunostaining for Ki67 and cleaved Caspase3 was performed in H, HK, HJ, and HKβ spheroids. All cells, Ki67-positive cells, and cleaved Caspase3-positive cells were quantified in a low-power field.

RNA isolation and qRT-PCR

Total RNA was extracted using the RNeasy Micro Kit (Qiagen). Single-strand complementary DNA was synthesized using a ReverTra Ace qPCR RT Kit (TOYOBO). qRT-PCR was performed using SYBR Green Master Mix (Roche Diagnostics Deutschland GmbH) and LightCycler 480 (Roche Diagnostics Deutschland GmbH). The expression levels were standardized by comparing them to the levels of GAPDH or β-actin (for c-Myc). Primer sequences are listed in Supplementary Table S3.

Microarray analysis

The Clarion S Assay (Thermo Fisher Scientific) was used for microarray analysis. Sample preparation, microarray hybridization, and bioinformatics analyses were performed by Macrogen. The quality of RNA extracted from spheroids was examined with an Agilent 2100 bioanalyzer (Agilent Technologies) and a Nanodrop (Thermo Fisher Scientific). The RNA samples were hybridized to the Affymetrix GeneChip Array (Thermo Fisher Scientific). Array data export, processing, and analysis were performed using Affymetrix GeneChip Command Console Software and Transcriptome Analysis Console (Thermo Fisher Scientific). The raw data were normalized by Affymetrix Power Tools Software (Thermo Fisher Scientific). Unnamed genes were excluded. The gene expression data of the samples were analyzed using gene set enrichment analysis (GSEA) software provided by the Broad Institute of MIT and Harvard University.

Xenograft

For the xenograft model of H, HK, HJ, and HKβ biliary spheroids, every spheroid was passaged at 5.0 \times 10^5 cells/well. Spheroids were obtained from three wells 3 days after passage. Harvested spheroids were suspended in 50 µL Matrigel + 50 µL Ad-DMEM/F12 and injected subcutaneously into NOD/SCID mice (NOD.CB17-Prkdcscid/J; purchased from The Jackson Laboratory, JAX strain 001303) at one site. After 3 months, tumorigenicity was assessed.

For the xenograft model of shSmad4 and shTgfbr2 HKβ biliary spheroids, every spheroid was passaged at 1.0 \times 10^6 cells/well. Spheroids were obtained from the two wells 4 days after passage. Harvested spheroids were suspended in 50 µL Matrigel + 50 µL Ad-DMEM/F12 and injected subcutaneously into NOD/SCID mice at one site. One month later, tumorigenicity was assessed.

DNA extraction

DNA was extracted using the QIAamp DNA Micro Kit (Qiagen) according to the manufacturer’s protocol. Sample integrity and yield were assessed by the Nanodrop (Thermo Fisher Scientific) and the Agilent TapeStation. Purified DNA samples were stored at −20°C.

Library preparation and WES

The samples were prepared according to an Agilent SureSelect Target Enrichment Kit preparation guide. The libraries were sequenced with Illumina NovaSeq 6000 sequencer. Whole exome capture was carried out using Agilent’s SureSelect Mouse Kit. After DNA quality evaluation, pooled samples were sequenced on Illumina NovaSeq 6000 according to the manufacturer’s instructions for paired-end 150 bp reads. The average sequencing depth of target region and total mapped reads were summarized in Supplementary Table S4. The Whole exome sequencing was performed by Macrogen.

Exome-sequencing data analysis for SNVs and INDELs calling

Quality check in Raw data (stored as FastQ format) was performed with the FastQC (ver0.11.8). Adapter contamination and low-quality nucleotides are discarded by using Trim Galore (ver0.6.4). Read pairs were mapped to the reference genome GRCh38 using the BWA-MEM (ver0.7.17). Duplicates were trimmed from the reads using the Picard Mark Duplicates (ver2.22.9). Furthermore, we performed base quality score recalibration with Base Recalibrator and ApplyBQSR in GATK (ver4.1.9.0).

After realignment to genome, we identified and filtered variants (SNPs, Indels) using GATK Mutect2 Somatic Mutation Call, GATK Filter Mutect Calls, and GATK Select Variants to guarantee meaningful analysis. Mutations with less than 5% allele frequency, low (<5) number of read, and low (<2) number of read with mutation in xenografted tumor were excluded from analysis. More than 10bp Indel, Known SNPs and Indels registered in Mouse Genomes Project, and silent mutations were excluded from analysis. The filtered mutations were annotated with VEP (ver96). Finally, the mutation spectrum of the final variant set was analyzed with the R package maftools (ver2.6.0). The WES data analysis was performed by cBioinformatics.

Lentivirus transduction and infection

The silencing of c-Myc was achieved using pLKO-shMyc (MISSION shRNA NM_010849 TRCN0000042517, purchased from Sigma-Aldrich), the silencing of Smad4 was attained using pLKO-shSmad4 (MISSION shRNA NM_008540 TRCN0000025881, purchased from Sigma-Aldrich), and the silencing of Tgfbr2 was attained using pLKO-shTgfbr2 (MISSION shRNA NM_009371 TRCN0000294600, purchased from Sigma-Aldrich). To produce lentiviruses, HEK293T cells were transfected with the targeting plasmid, pCAG-HIVgp, and pCMV-VSV-G-RSV-Rev (donated by Dr. Hiroyuki Miyoshi, RIKEN BioResource Center) plasmids. The culture supernatants were collected 48 hours after transfection, filtered, concentrated by PEG-it (System Biosciences, catalog no. LV810A-1), and resuspended in Hanks’ balanced salt solution. Infection was performed in the presence of polybrene (8 µg/mL) for 6 hours, followed by selection with puromycin 1 µg/mL. For the proliferation analysis of the spheroids that silenced c-Myc, Smad4, and Tgfbr2, the spheroid was passaged at 2.0 \times 10^5 cells/well by puromycin.

The Cancer Genome Atlas database

The data and analysis results can be explored through The Cancer Genome Atlas (TCGA) Data Portal from the cBioPortal on the Cancer Genomics website (https://www.cibioportal.org/).

Statistical analysis

Data are presented as mean ± SEM. When n = 3 per each group, dot plots are also represented. The two-tailed Student t test was performed to analyze the statistical difference between two groups. To determine statistically significant associations between treatment groups versus a control group or between the other groups versus a control group, the data were statistically analyzed with ANOVA followed by the Dunnet test. P value of less than 0.05 was considered statistically significant. All statistical analyses were performed with either JMP 15 (SAS Institute Inc.) or GraphPad Prism, version 6.0 (GraphPad).
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Study approval
All mouse experiments were approved by the animal research committee of Kyoto University (180260) and performed in accordance with Japanese government regulations. We used 12 surgically resected ICPN tissues that were obtained from patients admitted to Kyoto University Hospital and that were stored in the hospital. The patient samples were only used for immunohistochemistry and the Ethics Committee of Kyoto University approved the use of patient samples for this experiment without requiring written informed consent. Informed consent was obtained in the form of opt-out on the website. This research conformed to the provisions of the Declaration of Helsinki. The study protocol (G1200–1 and R2904) was approved by the Ethics Committee of Kyoto University.

Data availability
All original microarray data were deposited in the Gene Expression Omnibus at National Center for Biotechnology Information (GSE191072). The whole exome sequence data were deposited in the DDBJ Sequence Read Archive (DRA) database (DRA013309). The data that support the findings of this study are included in this article and its Supplementary Table or available from the corresponding author upon reasonable request.

Results

Hnf1β was expressed in the extrabiliary biliary system including the EHBD and GB
To perform genetic manipulation in the extrabiliary biliary tract epithelium including the EHBD and GB, we first sought to determine a suitable CreER mouse line. Given that HNF1β expression has been reported to be restricted to biliary duct epithelial cells in the normal murine liver (26), we performed immunostaining (IHC) for Hnf1β to determine the expression pattern of Hnf1β in the extrabiliary biliary tract epithelium including the EHBD and GB in adult mice. Hnf1β was robustly expressed in the EHBD, IHBD and GB epithelial cells in adult wild type mice (Fig. 1A).
To assess the efficiency of Cre expression in Hnf1β positive EHBD cells in Hnf1bCreER mice, we generated Hnf1bCreER, Rosa26 LacZ mice and performed lineage-tracing experiments by the administration of tamoxifen (Fig. 1B). Macroscopically and histologically, the EHBD and GB epithelial cells were Hnf1β-lineage labeled in Hnf1bCreER, Rosa26 LacZ mice at 2 days, 6 weeks, and 12 weeks after the last administration of tamoxifen (Fig. 1C). We also generated Hnf1bCreER, Rosa26 tdTomato mice and assessed Hnf1β-lineage labeled efficiency (Fig. 1D). Almost all EHBD and GB epithelial cells, which were positive for cytokeratin (CK), were Hnf1β-lineage labeled on day 2 after the last tamoxifen administration (Fig. 1E). These results indicate that the Hnf1bCreER mouse is a suitable CreER mouse line with high efficiency for genetic manipulation in the extrabiliary biliary tract epithelium including the EHBD and GB.

Concurrent activation of the Kras and canonical Wnt pathways induced ICPN and BiliN in mice
We next investigated the functional role of Kras and canonical Wnt signal activation and the effect of concurrent activation of both pathways on tumorigenesis of the extrabiliary biliary system using Hnf1bCreER mice. In a previous report, conditional knockout of exon3 of Ctnnb1 induced the stabilization of β-catenin or constitutive activation of the Wnt/β-catenin signaling pathway in a mouse model (23). We crossed KrasG12D mice and/or Ctnnb1lox(ex3)/+ mice with Hnf1bCreER mice to generate Hnf1bCreER(H), Hnf1bCreER, KrasG12D (HK), Hnf1bCreER, Ctnnb1lox(ex3)/+ (HK), and Hnf1bCreER, KrasG12D (DK) mice. Four weeks after the last tamoxifen administration, the biliary tract was analyzed (Fig. 2A). Macroscopically, the EHBD of HKβ mice was dilated, whereas that of HK, HK, and Hβ mice appeared normal (Fig. 2B). H&E staining demonstrated an almost normal appearance of the GB and EHBD epithelial cells in H mice and a minimal neoplastic change in Hβ and HK mice. In contrast, HKβ mice (6/6 mice) displayed microscopic papillary neoplasms, which resembled human BiliN in the EHBD, and papillary neoplasms, which resembled human ICPN in the GB (Fig. 2C and D). Neoplastic changes were also observed in the peribiliary glands of the EHBD in HKβ mice. Hyperchromasia, nuclear stratification, and partial loss of nuclear polarity were observed in the epithelial cells of the BiliN and ICPN lesions in HKβ mice. Ki67 staining revealed proliferative cells in these lesions in HKβ mouse (Fig. 2D). EHBD and GB epithelial cells demonstrated increased expression of pErk in both HK and HKβ mice (Fig. 2D). Nuclear accumulation or cytoplasmic staining of β-catenin was observed in the EHBD and GB epithelial cells in both Hβ and HKβ mice (Fig. 2D). These data indicated increased activation of the Kras and Wnt signaling pathways in EHBD and GB epithelial cells in HKβ mice.

To better characterize BiliN and ICPN in HKβ mice, mucin IHC was performed. Muc1 was positive in biliary epithelial cells in H, Hβ, HK, and HKβ mice, whereas muc2 and muc5AC were negative (Supplementary Fig. S1A). Staining for Griffonia simplicifolia lectin II (GSIIL lectin), which is an alternate marker for muc6 in mice and is indicative of the pyloric gland type (27), revealed that GSIIL was partially expressed in BiliN and ICPN in HKβ mice (Fig. 2D). Staining for Ulex europaeus agglutinin I (UEA-1), an indicative marker for the gastric foveolar type in mice (28), revealed that UEA-1 was partially expressed in BiliN and ICPN in HKβ mice (Supplementary Fig. S1B). Given that both GSIIL and UEA-1 were positive in HKβ mice, these ICPN lesions resembled human gastric type ICPN.

To compare with other Cre mouse strains, we used Dclk1CreER and Bmi1-CreER mouse lines and generated Dclk1CreER; KrasG12D, Ctnnb1lox(ex3)/+ (DKβ) and Bmi1CreER; KrasG12D, Ctnnb1lox(ex3)/+ (BKβ) mice, respectively. In both DKβ and BKβ mice, BiliN-like lesions and ICPN-like lesions were formed in the EHBD and GB, respectively, 4 weeks after tamoxifen administration (Supplementary Figs. S2A–S2C).

Therefore, these data indicate that concurrent activation of the Kras and canonical Wnt pathways results in the development of gastric type ICPN and BiliN of EHBD in mice.

Establishment of spheroids derived from murine mutant GB and EHBD and characterization of their gene expression profiles
To provide insights into the underlying molecular mechanism whereby activation of the Kras and Wnt pathways cooperatively induce ICPN and BiliN in the EHBD of HKβ mice, we sought to perform global gene expression analysis of mutant biliary epithelium. However, it was difficult to isolate the extrabiliary biliary epithelium from mouse tissue. Therefore, we applied a 3D culture system of biliary spheroids in vitro. We established biliary spheroids derived from the EHBD and GB of H, HK, Hβ, and HKβ mice 4 weeks after the last tamoxifen administration (Fig. 3A). We confirmed the recombination of KrasG12D allele and Ctnnb1lox(ex3)/+ allele in H, HK, Hβ, and HKβ biliary spheroids by genotyping with PCR (Supplementary Figs. S3A–S3C). These spheroids were roundish and did not show any apparent morphologic differences between genotypes (Fig. 3B). Proliferation analysis revealed that HKβ biliary spheroids grew significantly faster than H, HK, and Hβ biliary spheroids (Fig. 3C). Moreover, we
Figure 3.
Expression of Wnt signaling pathway, c-Myc, CSC markers, and Tgfβ pathway was increased in HKβ biliary spheroids. A, Schema of biliary spheroids established from EHBD and GB in H, HK, Hβ, and HKβ mice 4 weeks after the last tamoxifen administration (n = 3 mice/group). B, Microscopic images of H, HK, Hβ, and HKβ biliary spheroids on day 1 and day 5 after passage. Scale bars, 500 μm. C, Assay of the proliferation of H, HK, Hβ, and HKβ biliary spheroids on day 1, day 3, and day 5 after passage (n = 3/group). The assay of proliferation was performed using PrestoBlue. The values were calculated as described in Materials and Methods. Data are mean ± SEM. Analyzed by one-way ANOVA with Dunnett test (vs. HKβ spheroids; *P < 0.05; **P < 0.01). D, Top, commonly high fold change (>2 log, < −2 log) expression genes on microarray analysis in HKβ biliary spheroids. Bottom, upregulated gene sets on Hallmark gene set of GSEA in HKβ spheroids. FDR, q < 0.25 (vs. HKβ spheroids). E and F, qRT-PCR analysis of H, HK, Hβ, and HKβ biliary spheroids (n = 3/group). Data are shown as individual data points and the mean ± SEM for each experimental group. Analyzed by one-way ANOVA with Dunnett test (vs. HKβ spheroids; ***P < 0.001). G, Immunostaining for Dclk1 (top) and Aldh1a1 (bottom) in the EHBD and GB in H, HK, Hβ, and HKβ mice 4 weeks after the last tamoxifen administration. Aldh1a1 was expressed at a crypt-like location between BirIL in EHBD and the inner epithelial cells of ICPN in HKβ mice (arrowheads). Scale bars, 50 μm.
performed immunostaining for Ki67 and cleaved caspase3. The ratio of Ki67 positive proliferative cells was significantly higher in HKβ biliary spheroids than H, HK, and Hβ spheroids. On the other hand, the ratio of cleaved caspase 3 positive apoptotic cells in HKβ biliary spheroids was lower than H, HK, and Hβ biliary spheroids, although it did not reach statistical significance (Supplementary Figs. S4A–S4C).

Therefore, we concluded that the high proliferation contributes to the increased growth of HKβ biliary spheroids.

Next, we determined the gene expression profile of H, HK, Hβ, and HKβ biliary spheroids by genome-wide microarray analysis. Microarray analysis revealed the molecules with high fold change (>2 log or <−2 log), which included 1,597 genes in HKβ spheroids vs. H spheroids, 1,795 genes in HKβ spheroids versus HK spheroids, and 1,454 genes in HKβ spheroids versus Hβ spheroids. Commonly high fold change (>2 log or <−2 log) expression in HKβ spheroids compared to H, HK, and Hβ spheroids included 566 genes (Fig. 3D). Upregulated genes in HKβ spheroids included Aldh1a1, Arl4c, Axin2, Bmp1, Bmp2, Bmp7, Cdh2, Cdx2, Cflg72p1, Ctgf, Serpine1, and Tgfβ3. Downregulated genes in HKβ spheroids included Hnf1α, Hnf6α, and Muc1. Other genes are listed in Supplementary Data S1. GSEA revealed gene sets corresponding to epithelial–mesenchymal transition (EMT), myogenesis, and Tgfβ, p33, and Wnt signaling as commonly enriched signatures in HKβ spheroids compared with other genotypes in a “Hallmarks” compilation (Fig. 3D). qRT-PCR analysis revealed that the expression of c-Myc, Aldh1a1, Cdh2, Snail1, Tgfβ3, Serpine1, and Dclk1 was significantly (P < 0.05) upregulated in HKβ spheroids compared to H, HK, and Hβ spheroids (Fig. 3E and F).

Downregulation of c-Myc reflects Wnt/β-catenin activity (29). High expression of Cdh2 and Snail1 indicates the tendency of EMT. Aldh1a1 and Dclk1 are known cancer stem cell (CSC) markers (24, 30). Serpine1 is known a target gene of the Tgfβ signaling pathway and encodes plasminogen activator inhibitor-1 (PAI-1; ref. 31). These results indicated that HKβ biliary spheroids harbor enhanced features of the Wnt/β-catenin signaling pathway, c-Myc, CSC, EMT, and the Tgfβ signaling pathway compared with H, HK, and Hβ biliary spheroids.

We then performed IHC analysis in murine the EHBBD and GB tissue for CSC markers, including Dclk1 and Aldh1a1. Dclk1 was expressed in a small subset of epithelial cells of the EHBBD and GB in H, HK, and Hβ mice, whereas Aldh1a1 was strongly expressed in most BilIN and ICPN in HK β mice. Aldh1a1 was not expressed in EHBBD and GB in HK, HK, and Hβ mice, whereas Aldh1a1 was expressed at a crypt-like location between BilIN in EHBBD and in the inner epithelial cells of ICPN in HKβ mice (Fig. 3G). These results, together with the microarray data above, indicated that coactivation of Kras and canonical Wnt leads to higher tumorigenic potential in mouse biliary tracts.

ICPN and BilIN have a malignant potential to progress to biliary cancer

We next investigated whether BilIN and ICPN in HKβ mice were precancerous lesions. Longer-term analysis of HKβ mice was impossible, because HKβ mice died at 6 to 8 weeks of age after tamoxifen treatment. Given that Hnf1β is also expressed in the lung, pancreatic duct, intrahepatic bile duct, and intestinal epithelium (32), we histologically analyzed the intestine, pancreas, liver, and lung in HKβ mice 4 and 6 weeks after administration of tamoxifen. We observed massive adenomas in the intestine and focal nodular adenocarcinoma in the lung, whereas pancreas and liver were almost normal in HKβ mice 4 and 6 weeks after administration of tamoxifen (Supplementary Fig. S5). HKβ mice died due to massive intestinal adenomas. Therefore, we performed xenograft experiments using biliary spheroids developed from the EHBBD and GB of H, HK, Hβ, and HKβ mice (Fig. 4A).

Notably, xenograft tumors developed in two out of 24 cases of HKβ biliary spheroids approximately 3 months after the injection of spheroids, whereas no xenograft tumors developed from H, HK, and Hβ biliary spheroids (Fig. 4B and C). Histologically, these xenograft tumors were well-differentiated adenocarcinoma expressing CK19 (Fig. 4D). Ki67 staining revealed a high proliferation of xenograft tumors in the cancer cells developed from HKβ biliary spheroids.

To investigate the mutational profiles of the xenograft tumors developed from mutant biliary spheroids, we performed WES analysis using two sets of biliary spheroids; two original mutant biliary spheroids and corresponding xenograft biliary cancer spheroids. The WES analysis revealed 10 common additional gene mutations that were observed in the xenograft biliary cancer spheroids but not in the original biliary mutant spheroids in both sets (Supplementary Figs. S6A–S6D).

These results indicated that BilIN in the EHBBD and ICPN in HKβ mice are bona fide precancerous lesions with a malignant potential to become an adenocarcinoma although at a low frequency in vivo.

C-myc is a critical mediator for HKβ biliary spheroid growth

Given that c-Myc expression was significantly increased in HKβ biliary spheroids compared to H, HK, and Hβ biliary spheroids, we subsequently focused on c-Myc (Fig. 3E). IHC analysis revealed that c-Myc was expressed in BilIN of the EHBBD and ICPN in HKβ mice (Fig. 5A). Given that c-Myc is known to be located downstream of both Kras and Wnt signaling pathways, and is important for tumor growth (33), we hypothesized that c-Myc plays a critical role in the growth of HKβ biliary spheroids. To investigate the function of c-Myc in HKβ biliary spheroids, we silenced c-Myc in HKβ biliary spheroids. The silencing of c-Myc markedly suppressed the growth of HKβ biliary spheroids compared to the controls (Fig. 5B and C). Expression of Axin2, c-Myc, and Aldh1a1 was significantly decreased by the silencing of c-Myc in HKβ biliary spheroids (Fig. 5D).

To further validate the effect of c-Myc inhibition on the growth of HKβ biliary spheroids, we performed pharmacologic inhibition of c-Myc in HKβ biliary spheroids using 10058-F4, an inhibitor of c-Myc-Max dimerization (34). Administration of c-Myc inhibitor markedly suppressed the growth of HKβ biliary spheroids compared with the nontreated controls (Fig. 5E and F). We performed Myc inhibitor experiments using normal biliary spheroids (H) as a negative control. We found that administration of 100 μmol/L 10058-F4 suppressed more effectively the growth of HKβ biliary spheroids than that of H spheroids. On the other hand, the range of suppression by administration of 50 or 150 μmol/L 10058-F4 was not different between H and HKβ spheroids (Supplementary Figs. S7A–S7C). Thus, we concluded that Myc inhibitor suppressed more effectively the growth of HKβ spheroids than H spheroids. These results defined c-Myc as a critical mediator for the growth of HKβ biliary spheroids.

Tgfβ signaling pathway suppresses the growth of HKβ biliary spheroids

Given that the Tgfβ signaling pathway was also upregulated in HKβ biliary spheroids compared with H, HK, and Hβ biliary spheroids by GSEA analysis, we next focused on Tgfβ signaling (Fig. 3D). IHC analysis revealed that p-Smad3, located downstream of the canonical Tgfβ pathway, was expressed in the nucleus of BilIN in the EHBBD and ICPN in HKβ mice (Fig. 6A). The Tgfβ signaling pathway is generally known to play an important role as a tumor suppressor at an early stage of tumorigenesis, whereas it functions as a tumor promoter at an advanced stage of tumorigenesis. We hypothesized that the Tgfβ
signaling pathway plays a role in the growth of HKβ biliary spheroids. To investigate the functional role of the Tgfβ signaling pathway in the growth of HKβ biliary spheroids, we silenced Smad4 in HKβ biliary spheroids. qRT-PCR confirmed that expression of Smad4 was 79% downregulated in Smad4 silenced HKβ biliary spheroids (Supplementary Fig. S8A). Expression of Serpine1, target gene of the Tgfβ signaling pathway was also downregulated in Smad4 silenced HKβ biliary spheroids. Silencing of Smad4 markedly increased the growth of HKβ biliary spheroids compared with the controls (Fig. 6B and C). To further investigate the functional role of Tgfβ signaling pathway in the growth of HKβ biliary spheroids, we silenced Tgfb2 in HKβ biliary spheroids. qRT-PCR analysis confirmed that the expression of Tgfb2 was 79% downregulated in Tgfb2 silenced HKβ biliary spheroids. Silencing of Tgfb2 markedly increased the growth of HKβ biliary spheroids compared with controls (Supplementary Figs. S8A–S8D).

We also performed chemical inhibition of the Tgfβ signaling pathway by administering an inhibitor of the Tgfβ signaling pathway, SB431542 (35), in HKβ biliary spheroids. We treated HKβ biliary spheroids with SB431542 and analysis of proliferation and qRT-PCR was performed. Growth of HKβ biliary spheroids was markedly increased when treated with 1 or 10 μmol/L SB431542 compared with the controls (Fig. 6D and E). qRT-PCR confirmed that expression of Dlk1, TgfbR1, Serpine1, and Cdh2 was significantly downregulated in HKβ biliary spheroids treated with SB431542 (Fig. 6F). We performed chemical inhibition experiments using normal biliary spheroids (H) as a negative control. We found that administration of 1 or 10 μmol/L SB431524 increased more effectively the growth of HKβ spheroids than that of H spheroids. Therefore, we concluded that TGFβ inhibitor increased more effectively the growth of HKβ spheroids than H spheroids (Supplementary Figs. S7D–S7G). These results further demonstrated that the Tgfβ signaling pathway functions as a suppressive mediator for the growth of HKβ biliary spheroids.

Tgfβ signaling pathway suppressed the progression of BiliIN and ICPN into biliary cancer

We hypothesized that the Tgfβ pathway also inhibits the progression of BiliIN and ICPN into biliary cancer. Accordingly, we performed xenograft experiments using Smad4-silenced HKβ biliary spheroids.
Notably, xenograft tumors developed in five out of nine cases of Smad4-silenced HKbbiliary spheroids 1 month after the injection of Smad4-silenced HKbbiliary spheroids, whereas no xenograft tumors developed from the control HKbbiliary spheroids (Fig. 6H). Histologically, xenograft tumors comprised tubular adenocarcinomas expressing CK19 and Ki67 (Fig. 6H). The xenograft tumors did not express Smad4 and were negative upon periodic acid-Schiff staining. To further investigate whether Tgfβ pathway inhibits the progression of BilIN and ICPN into biliary cancer, we performed xenograft experiments using Tgfbr2-silenced HKbbiliary spheroids. We found that xenograft tumors developed in 6 of 18 mice injected with Tgfbr2-silenced HKbbiliary spheroids 1 month after the injection, whereas no xenograft tumors developed in 18 control mice injected with HKβ biliary spheroids (Supplementary Figs. S8E and S8F). These data indicate that the Tgfβ pathway suppresses the progression of BilIN and ICPN into biliary cancer.

**Gene expression of human gastric type ICPN and biliary cancer**

To investigate whether murine ICPN in HKβ mice resembled human ICPNs, we performed IHC analysis using resected human gastric type ICPN samples. Both p-ERK and nuclear β-catenin were expressed in serial sections in 7 of 12 human gastric ICPN cases (Fig. 7A). These results suggested that the KRAS and WNT signaling pathways were activated in human ICPNs. Human ICPNs also...
Figure 6.

TGFβ signaling pathway suppresses the growth of HKβ biliary spheroids and the progression of BilIN and ICPN into biliary cancer. A, Immunostaining for p-Smad3 in the EHBD and GB in H, HK, HKβ, and HKβ mice 4 weeks after the last tamoxifen administration. B, Microscopic images of the control and the Smad4 silenced HKβ biliary spheroids on day 0, day 2, day 4, and day 6 after passage. C, Diameter (fold change) of the control and the Smad4 silenced HKβ biliary spheroids on day 0, day 2, day 4, and day 6 after passage. Data are mean ± SEM. *P < 0.05; **P < 0.01, by Student’s t test. D, Microscopic images of DMSO, 1 μmol/L, and 10 μmol/L TGFβ inhibitor-treated HKβ biliary spheroids on day 0 and day 5 after treatment. E, Diameter (fold change) of DMSO, 1 μmol/L, and 10 μmol/L TGFβ inhibitor-treated HKβ biliary spheroids on day 1, day 3, and day 5 after treatment (n = 3/group). Data are mean ± SEM. F, Relative gene expression of DMSO, 1 μmol/L, and 10 μmol/L TGFβ inhibitor-treated HKβ biliary spheroids on day 5 (n = 3/group). Data are shown as individual data points and the mean ± SEM. E and F, *P < 0.05; **P < 0.01; ***P < 0.001, by one-way ANOVA with Dunnett’s test (vs. DMSO). G, Xenograft tumors developed in five out of nine cases of ShSmad4 HKβ biliary spheroids, whereas no tumor developed in the controls (n = 9) 1 month after injection of spheroids. H, From the top, macroscopic images (one scale, 1 mm), H&E staining, immunostaining for CK19, Ki67, and Smad4, D-PAS staining of xenograft tumors derived from Smad4-silenced HKβ biliary spheroids. Scale bars, 50 μm (A and H) and 500 μm (B and D).
expressed GS-II (an alternate for MUC6), indicating the characteristics of the pyloric gland type, which was consistent with our mouse data. Moreover, in large agreement with our mouse data, ALDH1A1, C-MYC, and p-SMAD3 were also expressed in human ICPNs. These results further support our conclusion that murine BilIN and ICPN induced by biliary-specific concurrent activation of the Kras and Wnt pathways recapitulated human ICPNs.

To investigate whether ICPN of HKβ mice retained the characteristics of human biliary cancer in terms of gene expression, we performed correlative analyses using the TCGA database. Expression of MUC6, DCLK1, SNAIL1, SNAIL2, TGFβ3, TGFβRI, and TGFβR2 was positively correlated with AXIN2 expression in human biliary cancer (Fig. 7B). Expression of CTNNB1 was negatively correlated with overall survival in human biliary cancer (Fig. 7C). Therefore, these data further strengthen the human relevance of our mouse model, which provides direct evidence that concurrent activation of the Kras and Wnt pathways induces ICPN and BilIN, which could progress to biliary cancer.

Discussion

The WHO classification recently defined BilIN, IPNB, MCN, and ICPN as precancerous lesions of biliary cancer (7, 8, 10). However, it has not been demonstrated whether these lesions are bona fide
precursor lesions of biliary cancer. Moreover, the molecular mechanisms underlying the tumorigenesis of these precursor lesions and their driver mutations are not well understood. In this study, to determine the role of Kras and canonical Wnt pathways in tumorigenesis of the extrahepatic biliary tract including the EHBD and GB, we introduced endogenous expression of KRASG12D and constitutive activation of Wnt signaling in the adult mouse biliary epithelium. We revealed that concurrent activation of the Kras and canonical Wnt pathways resulted in the development of biliary neoplasms that resembled human ICPNs and BilINs in the EHBD, putative precursors to biliary cancer. Furthermore, these lesions had the capacity to progress to adenocarcinomas in a xenograft model, establishing ICPNs and BilINs as precursors to biliary cancer. Thus, together with our analyses of the human clinical samples and the datasets described above, to the best of our knowledge, we have provided the first novel GEM that recapitulates human ICPN and BilIN, which can progress to biliary cancer.

In our global gene expression analysis of HKβ biliary spheroids, expression of c-Myc and the TGFβ pathway was upregulated in HKβ biliary spheroids. c-Myc is a known oncogene that promote progression of various types of cancer. Furthermore, it is one of the common downstream targets of Kras and canonical Wnt pathways in the extrahepatic biliary system induced ICPN and BilIN, which can progress to biliary cancer.

The TGFβ pathway works as a tumor suppressor in the early stage of tumorigenesis, whereas the TGFβ pathway induces EMT and function as a tumor suppressor in the late stage of tumorigenesis including pancreatic cancer (35, 36). In this study, we demonstrated that the silencing Smad4, silencing Tgfb2, or pharmacologic inhibition of the TGFβ receptor increased the proliferation of HKβ biliary spheroids in vitro. Moreover, we showed that the silencing of Smad3 or Tgfb2 markedly increased the progression of biliary cancer in vivo. Thus, concurrent activation of the Kras and Wnt pathways in the EHBD and GB resulted in an activated TGFβ pathway, which suppresses tumorigenesis and progression to biliary cancer. Consistent with our mouse data, we also revealed p-Smad3 expression, which is indicative of the activated TGFβ pathway, in human ICPN samples. Notably, SMAD4 expression was decreased in human high-grade BilINs (37). Therefore, our data together with the previous report, suggest that the TGFβ pathway is an important barrier of ICPN and BilIN to progression to biliary cancer.

It is conceivable that co-activation of the Kras and canonical Wnt pathways upregulates TGFβ ligands, because pharmacological inhibition of the TGFβ receptor increased the proliferation of HKβ biliary spheroids in vitro. The TGFβ ligands include TGFβ1, 2, and 3. In our analysis of biliary spheroids, there were no differences in the expression of TGFβ1 and TGFβ2 among H, HK, HB, and HKβ biliary spheroids, whereas expression of TGFβ3 was significantly higher in only HKβ biliary spheroids (Fig. 3E; Supplementary Fig. S9A). Therefore, it is most likely that concurrent activation of the Kras and Wnt pathways results in activation of the TGFβ pathway via the upregulation of the TGFβ3 ligand (Supplementary Fig. S10).

We showed that both UAE-1 and GS2 were expressed in ICPN and BilIN in HKβ mice. Although Muc1 was expressed in these neoplasms via IHC, Muc1 expression in HKβ biliary spheroid was lower than that in control spheroids (Supplementary Fig. 9B and 9C). Our data suggested that concurrent activation of the Kras and canonical Wnt pathways in the EHBD and GB epithelium led to a decrease in the biliary characteristics, but increased the gastric characteristics including both the foveolar and pyloric gland types. Given that human IPNB also includes the gastric type, it would be interesting for future studies to investigate whether activation of the Wnt and Kras signaling pathways is also observed in gastric type human IPNB.

To our knowledge, we have demonstrated for the first time that Hnf1bCreER is a suitable biliary-specific CreER mouse line with high efficiency to perform genetic manipulation in the extrahepatic biliary system including the EHBD and GB. Sox9CreER (13) and CK19CreER mice (16) were previously used as a GEM in the analysis of the adult epithelium of the biliary tract. Sox9 is expressed in various organs, including chondrocytes, testis, heart, lung, pancreas, bile duct, hair follicles, retina, and the central nervous system (38). CK19 is expressed in the oral cavity, mammary gland, skin, gastrointestinal tract (from the esophagus to the rectum), and lung (39). Hnf1b is expressed in the gastrointestinal tract (from the stomach to the rectum), kidney, lung, and gonads (40). In this study, we found that HKβ mice died most likely due to massive intestinal adenomas. Therefore, massive intestinal adenomas would be also formed in Sox9CreER, KrasG12D, Ctnnb1creER33:1 and CK19CreER, KrasG12D, Ctnnb1creER33:1 mice, and thus, long-term analysis would be impossible similarly to HKβ mice. Therefore, we conclude that Hnf1bCreER line is one of the best lines for genetic manipulation in the extrahepatic biliary epithelium among several CreER mouse lines, including Sox9CreER, and CK19CreER lines.

Our lineage tracing analysis showed high efficiency of Cre expression in the epithelium of the EHBD and GB in Hnf1bCreER, Rosa26-LacZ and Hnf1bCreER, Rosa26-tdTomato mice. The difference in the efficiency of Cre recombination between Hnf1bCreER, Rosa26-LacZ and Hnf1bCreER, Rosa26-tdTomato is likely because the efficiency of reporter proteins is higher in Rosa26-tdTomato mice than in Rosa26-LacZ mice. It has been reported that the efficiency of Cre recombination of several reporter alleles varied greatly (41).

Finally, TCGA database analysis of human biliary cancer indicated a correlative relationship between the expression of AXIN2 and MUC6, DCLK1, EMT, and the TGFβ pathway. TCGA data indicated that biliary cancer in which the Wnt pathway is activated expressed MUC6. Thus, the Wnt signaling pathway could be a molecular therapeutic target for MUC6 positive biliary cancer. Future research should focus on revealing the functional role of the Kras and Wnt signaling pathways in established biliary cancer, and investigating whether Wnt signaling could be a therapeutic target for MUC6 positive biliary cancer.

In summary, concurrent activation of the Kras and canonical Wnt pathways in the extrahepatic biliary system induced ICPN and BilIN, which can progress to biliary cancer. Mechanistically, c-Myc contributed to tumorigenesis, whereas the TGFβ pathway inhibited it. Consistent with the mouse data, the Kras, Wnt signalings, c-Myc, and TGFβ pathway were activated in human ICPNs. We have provided the first novel GEM that recapitulates human ICPN and BilIN, establishing them precancerous lesions.

Authors’ Disclosures

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Authors’ Contributions

M. Nagao: Conceptualization, data curation, formal analysis, investigation, visualization, methodology, writing–original draft. A. Fukuda: Conceptualization, supervision, funding acquisition, methodology, project administration, writing–review and editing. M. Omatu: Investigation. M. Namikawa: Investigation.


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