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


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Loss of c-Jun N-terminal Kinase 1 and 2 Function in Liver Epithelial Cells Triggers Biliary Hyperproliferation Resembling Cholangiocarcinoma

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Targeted inhibition of the c-Jun N-terminal kinases (JNKs) has shown therapeutic potential in intrahepatic cholangiocarcinoma (CCA)-related tumorigenesis. However, the cell-type-specific role and mechanisms triggered by JNK in liver parenchymal cells during CCA remain largely unknown. Here, we aimed to investigate the relevance of JNK1 and JNK2 function in hepatocytes in two different models of experimental carcinogenesis, the diethylnitrosamine (DEN) model and in nuclear factor kappa B essential modulator (NEMO)^{hepatocyte-specific knockout (Δ hepa)} mice, focusing on liver damage, cell death, compensatory proliferation, fibrogenesis, and tumor development. Moreover, regulation of essential genes was assessed by reverse transcription polymerase chain reaction, immunoblottings, and immunostainings. Additionally, specific *Jnk2* inhibition in hepatocytes of NEMO ^{Δ hepa}/JNK1 ^{Δ hepa} mice was performed using small interfering (si) RNA (si*Jnk2*) nanodelivery. Finally, active signaling pathways were blocked using specific inhibitors. Compound deletion of *Jnk1* and *Jnk2* in hepatocytes diminished hepatocellular carcinoma (HCC) in both the DEN model and in NEMO ^{Δ hepa} mice but in contrast caused massive proliferation of the biliary ducts. Indeed, *Jnk1/2* deficiency in hepatocytes of NEMO ^{Δ hepa} (NEMO ^{Δ hepa}/JNK ^{Δ hepa}) animals caused elevated fibrosis, increased apoptosis, increased compensatory proliferation, and elevated inflammatory cytokines expression but reduced HCC. Furthermore, si*Jnk2* treatment in NEMO ^{Δ hepa}/JNK1 ^{Δ hepa} mice recapitulated the phenotype of NEMO ^{Δ hepa}/JNK ^{Δ hepa} mice. Next, we sought to investigate the impact of molecular pathways in response to compound JNK deficiency in NEMO ^{Δ hepa} mice. We found that NEMO ^{Δ hepa}/JNK ^{Δ hepa} livers exhibited overexpression of the interleukin-6/signal transducer and activator of transcription 3 pathway in addition to epidermal growth factor receptor (EGFR)-rapidly accelerated fibrosarcoma (Raf)-mitogen-activated protein kinase kinase (MEK)-extracellular signal-regulated kinase (ERK) cascade. The functional relevance was tested by administering lapatinib, which is a dual tyrosine kinase inhibitor of erythroblastic oncogene B-2 (ErbB2) and EGFR signaling, to NEMO ^{Δ hepa}/JNK ^{Δ hepa} mice. Lapatinib effectively inhibited cystogenesis, improved transaminases, and effectively blocked EGFR-Raf-MEK-ERK signaling. **Conclusion:** We define a novel function of JNK1/2 in cholangiocyte hyperproliferation. This opens new therapeutic avenues devised to inhibit pathways of cholangiocarcinogenesis. (*Hepatology Communications* 2020;4:834-851).

Bile duct hyperplasia and aberrant cholangiocyte growth can result in hepatic cystogenesis, differentially diagnosed on the basis of cholangioma, cholangiofibrosis, intrahepatic cholangiocarcinoma (CCA), and oval cell hyperplasia.^(1,2) CCA, a malignancy that arises in the setting of chronic inflammation of biliary epithelium cells, has an increasing incidence and is the second most common primary

Abbreviations: α -SMA, alpha smooth muscle actin; Δ hepa, hepatocyte-specific knockout; A6, Notch-1; ALT, alanine aminotransferase; AST, aspartate aminotransferase; BW, body weight; CAF, cancer-associated fibroblast; CC3, cleaved caspase 3; CCA, cholangiocarcinoma; CK, creatine kinase; CLD, chronic liver disease; Coll1A1, collagen type I alpha 1; DEN, diethylnitrosamine; dmbt1, deleted in malignant brain tumors 1; ECM, extracellular matrix; EGFR, epidermal growth factor receptor; EMT, epithelial-mesenchymal transition; ERBB, erythroblastic oncogene B; ERK, extracellular signal-regulated kinase; f/f, floxed mice; gabrrp, gamma-aminobutyric acid A receptor; pi, GAPDH, glyceraldehyde 3-phosphate dehydrogenase; HCC, hepatocellular carcinoma; HER, human epidermal growth factor receptor; HNF, hepatocyte nuclear factor; HPF, high-power field; IHC, immunohistochemistry; IL, interleukin; JAK, Janus kinase; JNK, c-Jun N-terminal kinases; LoxP, locus of X-over P1; LPC, liver parenchymal cell; LW, liver weight; MAPK, mitogen-activated protein kinase; MEK, mitogen-activated protein kinase kinase; mRNA, messenger

liver cancer globally. Unfortunately, survival beyond a year of diagnosis is less than 5% and therapeutic options are scarce.⁽³⁾

Several *in vivo* and *in vitro* models as well as research with human tissue samples help to elucidate the main pathways implicated in CCA formation. However,

none of these studies recapitulates the human disease, and translation into improved patient outcome has not been achieved. In addition, the pathophysiology of CCA remains poorly understood. Thus, there is an urgent need for new models to improve the management of this insidious and devastating disease.

RNA; *MUC*, mucin; *NEMO*, nuclear factor kappa B essential modulator; *NF-κB*, nuclear factor kappa B; *OSM*, oncostatin M; *p*, phosphorylated; *PCNA*, proliferating cell nuclear antigen; *qRT-PCR*, quantitative reverse-transcription polymerase chain reaction; *Raf*, rapidly accelerated fibrosarcoma; *RIPK*, receptor-interacting serine/threonine-protein kinase; *si*, small interfering; *SOCS3*, suppressor of cytokine signaling 3; *SOX-9*, transcription factor *SOX 9*; *STAT*, signal transducer and activator of transcription; *TKI*, tyrosine kinase inhibitor; *TNF*, tumor necrosis factor; *TUNEL*, terminal deoxynucleotidyl transferase-mediated deoxyuridine triphosphate nick-end labeling.

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The c-Jun N-terminal kinases (JNKs) are evolutionarily conserved mitogen-activated protein kinases (MAPKs) and play an important role in converting extracellular stimuli into a wide range of cellular responses, including inflammatory response, stress response, differentiation, and survival.⁽⁴⁾ In tumorigenesis, JNK has been shown to have tumor suppressive function in breast,⁽⁵⁾ prostate,⁽⁶⁾ lung,⁽⁷⁾ and pancreas⁽⁸⁾ cancer. However, the pro-oncogenic role for JNK has also been well documented.⁽⁹⁻¹¹⁾ Importantly, JNK has lineage-determinant functions in liver parenchymal cells (LPCs) where it not only favors proliferation of biliary cells but also directly biases biliary cell-fate decisions in bipotential hepatic cells. It has been reported that JNK inhibition delays CCA progression⁽¹²⁾ by impeding JNK-mediated biliary proliferation. These data indicate that JNK modulation would be of therapeutic benefit in patients with CCA. Nevertheless, little is known about the cell-type-specific role and mechanism of JNK in biliary overgrowth in order to have a targeted and definite therapy against CCA.

In the present study, we investigated the implications of hepatocyte-defective JNK signaling in experimental carcinogenesis. Unexpectedly, loss of *Jnk1/2* in LPCs inhibited hepatocellular carcinoma (HCC) but triggered biliary epithelium hyperproliferation and features compatible with CCA. Overall, our data uniformly suggest that hepatocytic JNK is pivotal for biliary epithelial hyperproliferation resulting in ducto/cystogenesis.

Materials and Methods

GENERATION OF MICE AND ANIMAL EXPERIMENTS

Albumin (*Alb*)-*Cre* and *Jnk2*-deficient mice in a C57BL/6J background were purchased from the Jackson Laboratory (Bar Harbor, ME). *Jnk1*^{locus of X-over P1(LoxP)/LoxP}/*Jnk2*^{-/-} (Hepatocyte-specific knockout of *Jnk1* [JNK1^{Δhepa}]) mice were created as reported.⁽¹³⁻¹⁵⁾ We used male mice for all experiments. For *in vivo* experiments, mice were treated with a daily dose of lapatinib (150 mg/kg weight; n = 7 mice per group) or vehicle (0.5% hydroxypropylmethylcellulose/1% Tween 80) (n = 6) by oral gavage starting at 6 weeks of age over a period of 6 weeks. For small interfering (si)RNA-mediated knockdown experiments, 8-week-old nuclear factor kappa B (NF-κB) essential

modulator (NEMO)^{Δhepa}/JNK1^{Δhepa} were injected with a dose of 0.2 mg/kg body weight (BW) *siJnk2* or small interfering luciferase (*siLuc*) once per week over a period of 4 weeks. In parallel, lapatinib was given orally to *siJnk2*-treated NEMO^{Δhepa}/JNK1^{Δhepa} mice on the same day of the first *siJnk2*-injection. Induction of tumorigenesis was performed by intraperitoneal injection of 25 mg/kg BW of diethylnitrosamine (DEN; Sigma-Aldrich, Munich, Germany) at 14 days of age. Mice were killed 24 weeks later. Vehicle-injected (saline) male mice served as controls.

Animal experiments were carried out according to the German legal requirements and animal protection law and approved by the authority for environment conservation and consumer protection of the state of North Rhine-Westphalia (LANUV; Germany). All strains were crossed on a C57BL/6 background. The mice were housed in the Institute of Laboratory Animal Science at the University Hospital RWTH-Aachen University according to German legal requirements (Animal Welfare Act [Deutsches Tierschutzgesetz], Federation of European Laboratory Animal Science Associations [FELASA], Society of Laboratory Animal Science [GV-SOLAS]) under a permit of the Veterinäramt der Städteregion Aachen. All animals received humane care according to the criteria outlined in the Guide for the Care and Use of Animal Models. All organ explants and animal experiments were approved by the local authority for environment conservation and consumer protection of LANUV on the following animal grants: 30034G (AZ-84-02.04.2016.A080) and TVA-11324GZ (AZ-84-02.04.2016.A490).

INTERFERENCE RNA AGAINST *Jnk2* (*siJnk2*)

The siRNA molecules were purchased from Axolabs GmbH (Kulmbach, Germany) and were chosen due to their ability to specifically target *Jnk2* in mice with mismatches to *Jnk1* (2-18 nucleotides) to increase *in vivo* stability and suppression of the immunostimulatory properties, as described.⁽¹⁶⁾

DATA AND SOFTWARE AVAILABILITY

Affymetrix Microarray was performed as described,⁽¹⁷⁾ and data were deposited with the National Center for Biotechnology Information Gene

Expression Omnibus (<http://www.ncbi.nlm.nih.gov/geo/>) under accession number GSE140498.

STATISTICAL ANALYSIS

All data are expressed as mean \pm SEM. Statistical significance was determined by two-way analysis of variance (ANOVA) followed by a Student *t* test or by one-way ANOVA followed by a Newman-Keuls multicomparison test. *P* < 0.05 was considered significant.

Results

COMBINED LOSS OF *Jnk1/2* FUNCTION IN HEPATOCYTES TRIGGERS BILIARY HYPERPROLIFERATION AND DUCTO/CYSTOGENESIS IN AN EXPERIMENTAL MODEL OF CHRONIC LIVER DISEASE

Mice lacking NEMO in LPCs spontaneously develop HCC, rendering these animals an ideal model that perfectly mimics progression of chronic liver disease (CLD) as observed in humans.⁽¹⁸⁾ We previously established that *Jnk1* and *Jnk2* have specific roles for the progression of NEMO ^{Δ hepa}-dependent chronic liver injury, indicating that the MAPK genes expressed in liver cells have pivotal functions in cell death and inflammation.⁽¹⁹⁾ We then found that combined activities of *Jnk1* and *Jnk2* specifically in hepatocytes protected against toxic liver injury (CCl₄ and acetaminophen).⁽¹⁵⁾ However, *Jnk1/2* deficiency in hepatocytes increased tumor burden in an experimental model of HCC.⁽¹⁰⁾ Therefore, in the present study we aimed to investigate the definitive contribution of JNK genes to liver cancer.

For this purpose, we generated NEMO ^{Δ hepa}/JNK ^{Δ hepa} and their respective controls NEMO ^{Δ hepa} and NEMO^{floxed (f/f)} mice (Supporting Fig. S1) and examined the progression of liver disease. At 13 weeks of age, histologic evaluation of NEMO ^{Δ hepa} unveiled the presence of dysplastic nodules, steatohepatitis, and cell death (Supporting Fig. S2A,B). All NEMO ^{Δ hepa}/JNK ^{Δ hepa} livers with no external signs of nodules spontaneously showed hyperproliferation of biliary epithelium translated into hepatic ducto/cystogenesis and lymphoid aggregates surrounding these structures (Supporting Fig. S2A,B).

Development of HCC is characteristic of 1-year-old NEMO ^{Δ hepa} mice, visible both macroscopically and histologically. However, most 52-week-old NEMO ^{Δ hepa}/JNK ^{Δ hepa} mice displayed no signs of HCC. Here, yellowish coloration of the liver parenchyma was associated with hyperproliferation of biliary epithelial cells and lymphoid cell accumulation. This was much more pronounced than at earlier stages of CLD. Interestingly, histopathological examination of these specimens in two different institutes (Utrecht and Graz) revealed that approximately 33% of 52-week-old JNK ^{Δ hepa} livers presented small cysts while an increased frequency and a higher number of ductular cells were evident in the hepatic parenchyma of all NEMO ^{Δ hepa}/JNK ^{Δ hepa}, with atypia compatible with CCA (Fig. 1A,B; Table 1).

Moreover, combined JNK1/JNK2 deletion in hepatocytes of NEMO ^{Δ hepa} mice triggered significantly reduced BW and liver weight (LW) compared with NEMO ^{Δ hepa} mice but a similar LW/BW ratio as hepatocyte-specific NEMO-deficient mice (Fig. 1C). Notably, at 13 weeks of age, NEMO ^{Δ hepa}/JNK ^{Δ hepa} animals had a significantly increased hepatosomatic ratio compared with NEMO ^{Δ hepa} mice, albeit no differences in BW or LW (Supporting Fig. S2C).

Surprisingly, 1 year-old NEMO ^{Δ hepa}/JNK ^{Δ hepa} livers exhibited reduced HCC compared with NEMO ^{Δ hepa} animals (Fig. 1D). In contrast, JNK1/2-deleted NEMO mice displayed ducto/cystogenesis that was much more pronounced than at earlier stages of CLD (Fig. 1B; Supporting Fig. S2D). NEMO ^{Δ hepa}/JNK ^{Δ hepa} livers exhibited significantly elevated glutamate dehydrogenase, total bilirubin, alkaline phosphatase, aspartate aminotransferase (AST), and alanine aminotransferase (ALT) levels compared with NEMO ^{Δ hepa} mice, already detectable at 13 weeks (Fig. 1D,E; Supporting Fig. S2D,E). Altogether, these data indicated that deletion of *Jnk1/2* in an experimental model of CLD has pivotal implications in cell death, cholestasis development, and ductular proliferation of cholangiocytes.

ANALYSIS OF THE MICROENVIRONMENT DRIVING MASSIVE BILE DUCT PROLIFERATION IN NEMO ^{Δ hepa}/JNK ^{Δ hepa} ANIMALS

Lack of NF- κ B activity in hepatocytes triggers spontaneous HCC. In turn, 52-week-old

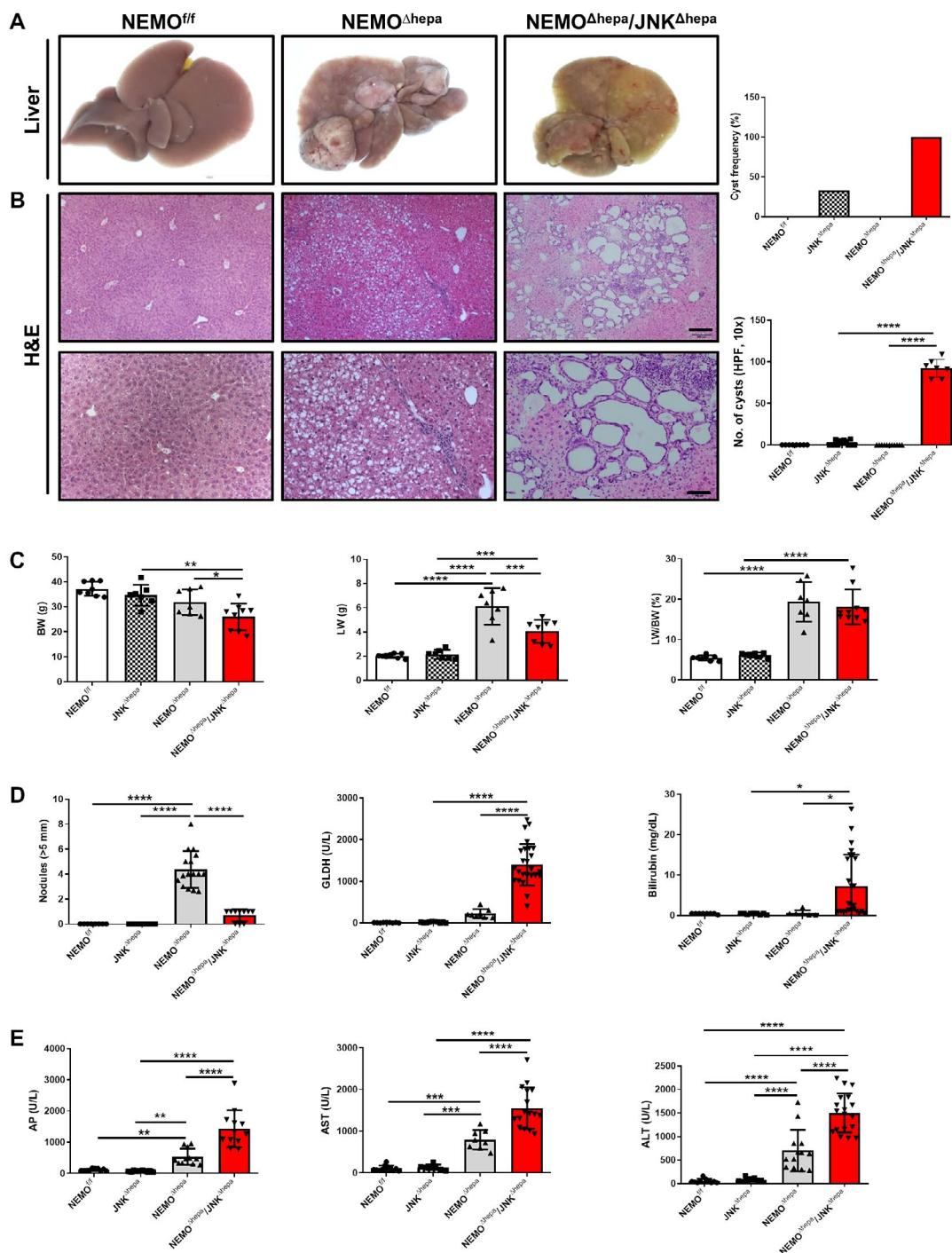


FIG. 1. Deletion of *Jnk1/2* in 52-week-old NEMO^{Δhepa} livers triggers cyst formation. (A) Macroscopic view of livers from 52-week-old NEMO^{fl/fl} (wild type), NEMO^{Δhepa}, and NEMO^{Δhepa}/JNK^{Δhepa} mice. (B) Representative H&E staining of liver sections of NEMO^{fl/fl}, NEMO^{Δhepa}, and NEMO^{Δhepa}/JNK^{Δhepa} livers at 52 weeks of age. Different magnifications were used (left). Scale bar 200 μm. Cyst frequency and number of visible microscopic cysts per 10× view field were calculated and graphed (right), magnification is 10× for upper and magnification is 20× for lower. (C) BW (left); LW (center); LW/BW ratio (right). (D) Tumor burden for each individual mouse was characterized by calculating total number of visible tumors >5 mm in diameter per mouse (left); serum levels of GLDH (center); bilirubin (right). (E) Serum levels of AP (left), AST (center), and ALT (right), in 52-week-old NEMO^{fl/fl}, JNK^{Δhepa}, NEMO^{Δhepa}, and NEMO^{Δhepa}/JNK^{Δhepa} mice. Data are presented as mean ± SEM. **P* < 0.05; ***P* < 0.01; ****P* < 0.001; *****P* < 0.0001. Abbreviations: AP, alkaline phosphatase; GLDH, glutamate dehydrogenase; H&E, hematoxylin and eosin.

TABLE 1. HISTOPATHOLOGICAL CHARACTERISTICS OF THE DIFFERENT MOUSE GROUPS (SCALE, 0-4)

	NEMO ^{Δhepa}	NEMO ^{Δhepa} /JNK ^{Δhepa}
Neoplasia	HCC	Cystic cholangioma Mucinous CCA
Anisokaryosis	3.5	2.0
Altered foci	2.0	1.8
Mitosis/HPF (40×)	2.5	1.0
Cellular hypertrophy	3.0	2.5
Dysplasia	3.0	2.5
Oval cell proliferation	1.5	2.5
Portal inflammation	1.0	1.0
Overall inflammation	3.0	3.5
Ductular reaction	1.5	2.0
Apoptosis	1.0	3.0
Fibrosis	3.0	4.0
Steatosis	1.4	0.0
Others	Pale cytoplasm hepatocytes	Massive bile duct proliferation

NEMO^{Δhepa}/JNK^{Δhepa} mice displayed significantly increased bile duct proliferation. We next investigated liver damage associated with the phenotype of these animals. We first applied terminal deoxynucleotidyl transferase-mediated deoxyuridine triphosphate nick-end labeling (TUNEL) staining, which detects several types of cell death, including necrosis, apoptosis, and necroptosis.⁽²⁰⁾ NEMO^{Δhepa} livers displayed a high percentage of TUNEL-positive cells, while NEMO^{Δhepa}/JNK^{Δhepa} livers showed significantly increased levels of cell death (Fig. 2A).

Apoptosis and necroptosis are two relevant forms of cell death in the pathogenesis of human and murine liver disease.^(21,22) To discriminate between these two types of cell death, we performed immunohistochemistry (IHC) analyses, which revealed significantly higher levels of the apoptosis marker cleaved caspase 3 (CC3) in NEMO^{Δhepa}/JNK^{Δhepa} livers, especially in cholangiocytes but also in immune infiltrates of these livers (Fig. 2B). We subsequently studied proteins involved in necroptosis, e.g., receptor-interacting serine/threonine-protein kinase 3 (RIPK3) by western blot and IHC, which was found overexpressed in NEMO^{Δhepa}/JNK^{Δhepa} livers, while no difference in RIPK1 protein expression was found (Fig. 2E; Supporting Fig. S3A).

Cell-cycle dysregulation is characteristic of biliary overgrowth, resulting in CCA. Increased Ki-67-positive and proliferating cell nuclear antigen (PCNA)-positive cells were observed in NEMO^{Δhepa}/JNK^{Δhepa} frozen

and paraffin sections, respectively, the latter further confirmed by immunoblot analysis (Fig. 2C,E; Supporting Fig. S3B). Moreover, the number of transcripts for PCNA and cyclin D1 was up-regulated in NEMO^{Δhepa}/JNK^{Δhepa} livers (Supporting Fig. S3C). P21 has been shown to induce cell-cycle arrest and promote the DNA repair gene, thus acting as a tumor suppressor.^(23,24) Interestingly, we observed p21 overexpression in *Jnk1/2*-deficient NEMO^{Δhepa} mice (Fig. 3E). Moreover, oxidative stress in the liver microenvironment has a definitive role in CCA development. We thus measured lipid peroxidation and the antioxidant defense of these animals. Interestingly, strong 4-hydroxynonena immunostaining and catalase depletion (Supporting Fig. S3D,E) in NEMO^{Δhepa}/JNK^{Δhepa} livers confirmed elevated reactive oxygen species (ROS) production associated with cholangiocellular proliferation.

Hepatic stellate cells (HSCs) are important cells shaping the hepatic microenvironment after liver damage. Hence, HSCs are involved in the high number of α -smooth muscle actin (α -SMA)-positive myofibroblasts and extracellular matrix (ECM) deposition associated with CCA development.⁽²⁵⁾ Sirius red staining and quantification confirmed strong collagen deposition in NEMO^{Δhepa}/JNK^{Δhepa} livers (Fig. 2D; Supporting Fig. S4A). Furthermore, collagen type I alpha 1 (Col1A1) and α -SMA were dramatically overexpressed in NEMO^{Δhepa}/JNK^{Δhepa} liver protein lysates (Fig. 2E) and were associated with significantly increased transcripts of *col1A1*, matrix metalloproteinase (*mmp*)7/9/12, and tissue inhibitor of metalloproteinase (*timp*)1 (Supporting Fig. S4B,C). Cytokine-mediated tissue fibrosis was also measured, and levels of monocyte chemoattractant protein 1 (*mcp1*), tumor necrosis factor (*tnf*), interleukin (*il*)1 β , and regulated upon activation, normal T cell expressed, and secreted (*rantes*) were significantly up-regulated in NEMO^{Δhepa}/JNK^{Δhepa} mice (Supporting Fig. S4D,E). Altogether, these data suggest that fibrogenesis and unresolved inflammation are involved in CCA development in the NEMO^{Δhepa}/JNK^{Δhepa} context while HCC-related tumorigenesis is diminished as confirmed by low glutamine synthase expression (Supporting Fig. S5A).

NEMO^{Δhepa}/JNK^{Δhepa} LIVERS EXHIBIT TYPICAL FEATURES OF HUMAN CCA

Immunohistochemical analysis revealed that many ductules were composed of creatine kinase

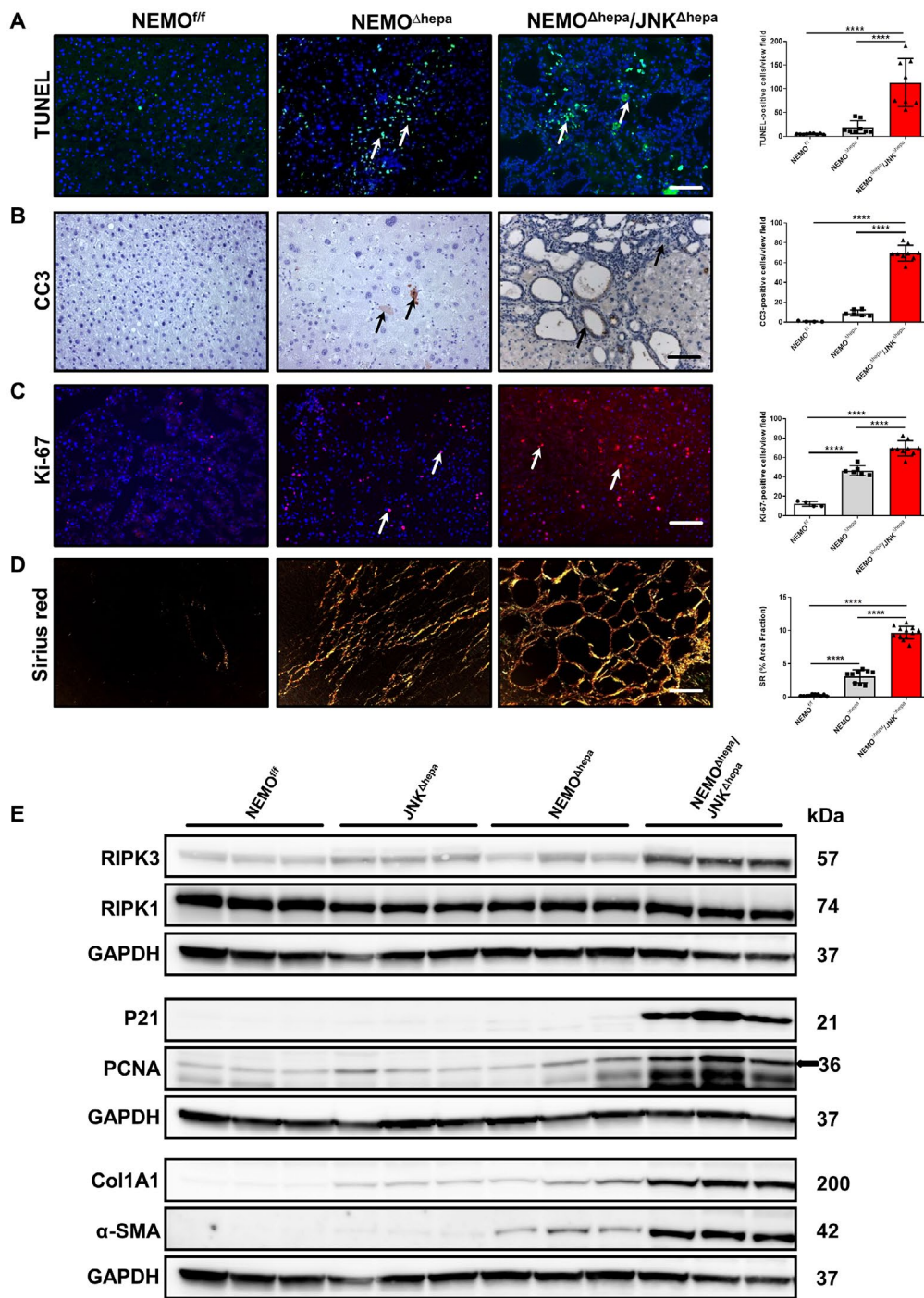


FIG. 2. Characterization of cell death, cell proliferation, and collagen deposition in 52-week-old NEMO^{Δhepa}/JNK^{Δhepa} mice. (A) Representative TUNEL staining of liver sections of NEMO^{fl/fl}, NEMO^{Δhepa}, and NEMO^{Δhepa}/JNK^{Δhepa} livers at 52 weeks of age (left). TUNEL-positive cells were quantified and graphed (right). (B) Representative IHC staining for CC3 of the same livers (left). CC3-positive cells were quantified and graphed (right). (C) Immunofluorescent staining for Ki-67 of liver cryosections from the same livers (left). Ki-67-positive cells were quantified and graphed (right). (D) Representative sirius red staining of paraffin sections from the indicated genotypes (left). Quantification of the positive sirius red area fraction was performed with Image J and graphed (right). (A-D) Scale bars, 200 μm. Arrows (→) indicate positive cells. Data are presented as mean ± SEM; *****P* < 0.0001. (E) Protein levels of α-SMA, Col1A1, PCNA, p21, RIPK1, and RIPK3 from whole-liver extracts of 52-week-old NEMO^{fl/fl}, JNK^{Δhepa}, NEMO^{Δhepa}, and NEMO^{Δhepa}/JNK^{Δhepa} mice were analyzed by western blot with the indicated antibodies. GAPDH was used as a loading control.

(CK)19-positive cells. In fact, $NEMO^{\Delta hepa}/JNK^{\Delta hepa}$ cystic and CCA-like structures exhibited CK19-positive staining that was corroborated by dramatically

elevated CK7/19 messenger RNA (mRNA) expression in these livers (Fig. 3A,D). $NEMO^{ff}$ had regular small bile ducts, and $NEMO^{\Delta hepa}$ exhibited mild to

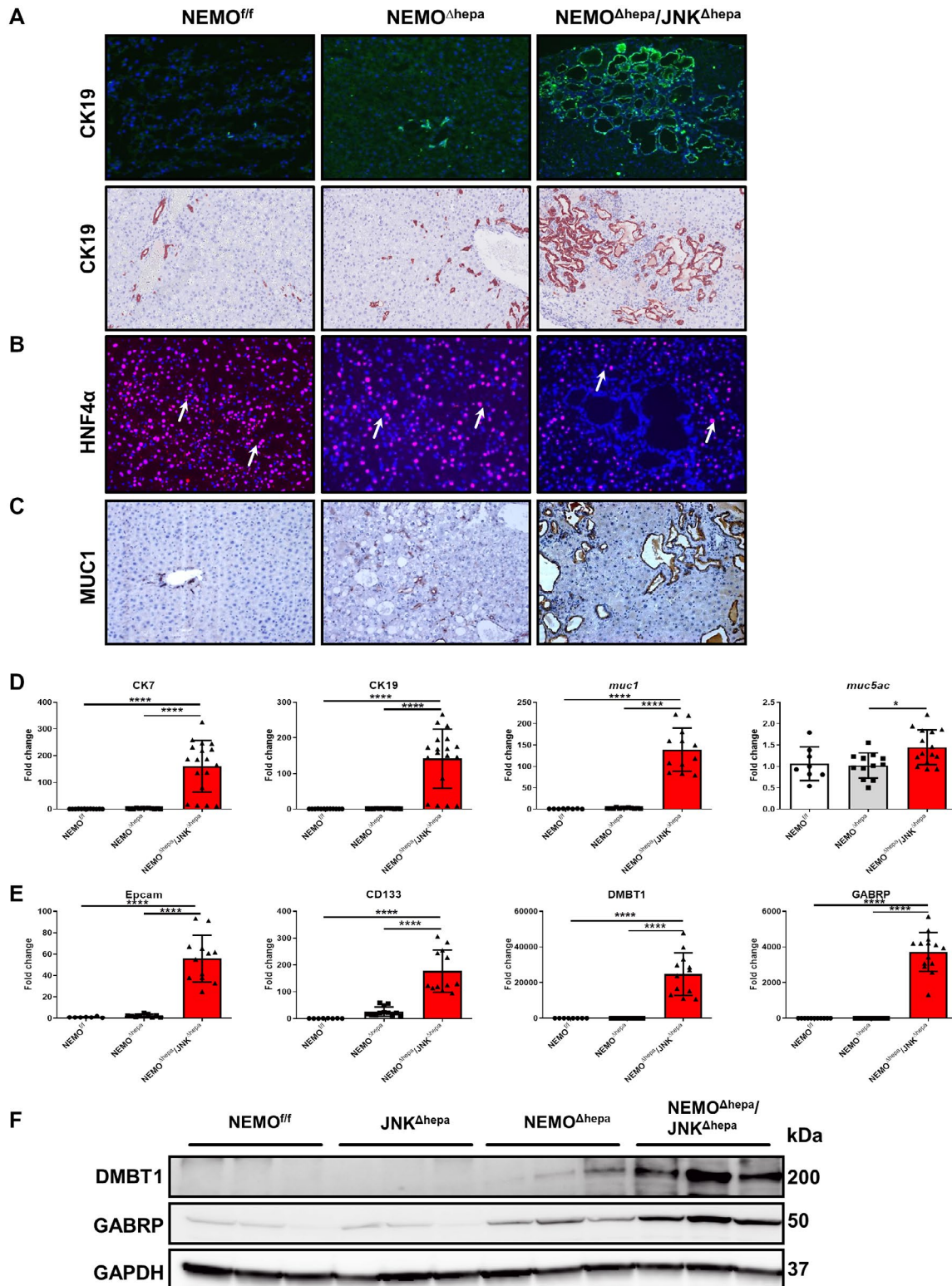


FIG. 3. Loss of *Jnk1/2* in NEMO^{Δhepa} hepatocytes triggers cholangiocellular proliferation. (A) Representative IF for CK19 of liver cryosections was performed in 52-week-old NEMO^{fl/fl}, NEMO^{Δhepa}, and NEMO^{Δhepa}/JNK^{Δhepa} livers (upper panel). IHC staining for CK19 reveals densely packed cholangiocellular proliferates with a tubular growth pattern mimicking morphologic features of well-differentiated human CCA (lower panel), magnification is 20×. (B) Representative IF staining for HNF-4α of the same livers, magnification is 20×. (C) Representative IHC staining for Muc1, magnification is 20×. (D) mRNA expression analysis of CK7 (left), CK19, and *muc1* (center) and *muc5ac* (right) was quantified by qRT-PCR in the same livers. (E) mRNA expression analysis of Epcam (left), CD133 and DMBT1 (center), and GABRP (right) was quantified by qRT-PCR of samples taken from NEMO^{fl/fl}, NEMO^{Δhepa}, and NEMO^{Δhepa}/JNK^{Δhepa} livers killed at 52 weeks. (F) Protein expressions of GABRP and DMBT1 from whole-liver extracts of 52-week-old NEMO^{fl/fl}, JNK^{Δhepa}, NEMO^{Δhepa}, and NEMO^{Δhepa}/JNK^{Δhepa} mice were analyzed by western blot with the indicated antibodies. GAPDH served as loading control. Data are presented as mean ± SEM. **P* < 0.05; *****P* < 0.0001. Abbreviations: CD, cluster of differentiation; Epcam, epithelial cell adhesion molecule; IF, immunofluorescence.

moderate ductular proliferates similar to a ductular reaction in humans. Noticeably, livers of NEMO^{Δhepa}/JNK^{Δhepa} mice displayed cholangiocellular proliferates with a tubular architecture resembling the morphology of human well-differentiated CCA, highlighted by a strong-positive CK19 signal. In contrast, hepatocyte nuclear factor (HNF)-4α staining was characteristic of pericyclic areas and hepatocytes of NEMO^{Δhepa} and NEMO^{fl/fl} livers (Fig. 3B). Cells containing cytoplasmic mucin (MUC) granules are common in CCA tissues.⁽²⁶⁾ Thus, we assessed the expression of MUC1 and MUC5AC, which are closely related to dedifferentiation, infiltrative growth pattern, and patient survival.⁽²⁶⁾ Whereas MUC1 staining was limited to areas of ductogenesis or oval cell reaction in NEMO^{Δhepa} livers, it was very strong in cysts of NEMO^{Δhepa}/JNK^{Δhepa} livers, which were associated with significantly increased *muc1* and *muc5ac* mRNA levels in these mice (Fig. 3C,D).

High expression of cholangiocyte markers, including cluster of differentiation (*cd*)133 and expression of CCA/tumor-enriched markers, including epithelial cell adhesion molecule (*epcam*), deleted in malignant brain tumors 1 (*dmbt1*), and gamma-aminobutyric acid A receptor, pi (*gabbrp*),⁽²⁷⁾ together with the accumulation of transcription factor SOX 9 (SOX-9)-positive cells (Fig. 3E,F; Supporting Fig. S5B) suggest that loss of *Jnk1/2* function in hepatocytes promotes the shift from HCC to CCA in this experimental model of CLD.

ADMINISTRATION OF DEN TO JNK^{Δhepa} MICE TRIGGERS HEPATIC DUCTO/CYSTOGENESIS WITHOUT HEPATOCARCINOGENESIS

To validate the relevance of *Jnk1/2* in mediating the shift from HCC to CCA, we applied a second model of carcinogenesis, the DEN model. Previously, Das and colleagues,⁽¹⁰⁾ using mice with compound

deficiency of *Jnk1/2* in hepatocytes, demonstrated that the JNK genes possess tumor-suppressing roles in liver carcinogenesis that depend on the cell types of the liver. Additionally, we showed that hepatocyte-specific *Jnk1/2* knockout female and male mice have no phenotype affecting the correct function of the liver.⁽¹⁵⁾ These previous results are confirmed in the present study in a larger pool of animals ranging from 13 to 52 weeks of age (Supporting Figs. S6A-D and S7A-D). However, approximately 33% of these mice from week 30 of age histologically displayed tumors resembling human CCA in their liver parenchyma that did not affect liver function (Fig. 1A,B; Supporting Fig. S7A-D).

JNK^{Δhepa} mice were challenged with the carcinogen DEN. Interestingly, these mice exhibited jaundice and the liver was yellowish albeit with decreased tumor load (Supporting Fig. S8A). The LW/BW ratio was significantly decreased in JNK^{Δhepa} compared with JNK^{fl/fl} mice (Supporting Fig. S8B). Histologic evaluation performed by two blinded pathologists demonstrated the presence of cystogenesis and cholangioma-like structures in liver parenchyma accompanied by strong infiltration of immune cells (Fig. S8C). The analysis of serum transaminases in JNK^{Δhepa} demonstrated decreased ALT and AST levels in these animals (Fig. S8D,E). These results indicated that JNK1 and JNK2 might influence cell fate during liver tumorigenesis.

To further analyze the differences between both animal models, age progression versus chemically induced HCC, we performed a microarray analysis (Supporting Fig. S9A,B). Interestingly, *dmbt1*, *muc1*, *gabbrp*, and immunoglobulin heavy chain (gamma polypeptide) (*ighg*)2b were commonly up-regulated in *Jnk1/2*-deficient NEMO^{Δhepa} mice and *Jnk1/2*-deficient mice challenged with DEN, indicating that epithelial–mesenchymal transition (EMT) might be modulated through a JNK-dependent mechanism.

Next, we analyzed total common up-regulated and down-regulated genes in NEMO^{Δhepa}/JNK^{Δhepa} versus JNK^{Δhepa} + DEN, which were the majority compared to specific changes of each experimental model (Supporting Fig. S9C,D). Heat map analysis of these genes showed up-regulation of genes related to CCA, like *dmbt1*, *muc1*, *gabbrp*, and matrix deposition, including *timp1* and *mmp12*, and down-regulation of serpins, which are often implicated in HCC development (Fig. 4A). Altogether, these data reinforce the notion that the JNK signaling pathway modulates cell fate during liver carcinogenesis.

OVEREXPRESSION OF THE IL-6/SIGNAL TRANSDUCER AND ACTIVATOR OF TRANSCRIPTION 3 PATHWAY IS CHARACTERISTIC FOR NEMO^{Δhepa}/JNK^{Δhepa} LIVERS

Cytokines play an important role during carcinogenesis by shaping the inflammatory microenvironment toward malignant transformation. In particular, IL-6 has been demonstrated to have an integral role in CCA biology and other cancers as a growth and survival factor.⁽²⁸⁾ Under physiologic conditions, IL-6 through a Janus kinase (JAK)/signal transducer and activator of transcription (STAT) 3 pathway induces expression of suppressor of cytokine signaling 3 (SOCS3), accelerating inflammation, cell growth, and tumor formation. We found overexpression of *il6*, *il6* receptor (*il6r*), and *socs3* mRNA as well as other IL-6 family members, including LIF IL-6 family cytokine (*lif*) and oncostatin M (*osm*) (Fig. 4B,C; Supporting Fig. S10A). OSM has recently been identified as an important regulator of the EMT/cancer stem cell plasticity program that promotes tumorigenic properties.⁽²⁹⁾ *Osm* was significantly increased in NEMO^{Δhepa}/JNK^{Δhepa} livers.

Concomitantly, high phosphorylated STAT3 (pSTAT3) levels were evident in NEMO^{Δhepa}/JNK^{Δhepa} compared with NEMO^{Δhepa} livers (Fig. 4D; Supporting Fig. S10B), indicating that this pathway is strongly activated in our murine model.

THE EGF-Raf-MEK-ERK1/2 PATHWAY IS OVEREXPRESSED IN NEMO^{Δhepa}/JNK^{Δhepa} ANIMALS

Activation of Notch and Wnt signaling pathways stimulates proliferation of the hepatic progenitor

cell compartment. In particular, NOTCH signaling is implicated in the commitment toward the cholangiocyte fate, while WNT can trigger differentiation toward the hepatocyte lineage.⁽³⁰⁾ Therefore, we first assessed the relevance of Notch-1 (A6) in NEMO^{Δhepa}/JNK^{Δhepa} livers. Interestingly, A6 staining was positive in clusters of immune cells in NEMO^{Δhepa} livers. NEMO^{Δhepa}/JNK^{Δhepa} livers exhibited increased Notch-1 expression throughout the liver parenchyma (Fig. 5A). In fact, other family members, including *Notch2*, were significantly up-regulated or had a tendency toward increased mRNA levels in NEMO^{Δhepa}/JNK^{Δhepa} livers (Fig. 5C; Supporting Fig. S11A-F). However, no differences in β-catenin expression between NEMO^{Δhepa} and NEMO^{Δhepa}/JNK^{Δhepa} livers were observed (Fig. 5D). Of note, JNK1/2-knockout mice had reduced phosphorylation of β-catenin, indicating that JNK is necessary for β-catenin phosphorylation, as suggested.⁽³¹⁾

The epidermal growth factor receptor (EGFR) family includes erythroblastic oncogene B (ERBB)1, 2, 3, and 4, with ERBB1/EGFR and ERBB2/human epidermal growth factor receptor 2 (HER2) (Neu in rodents) being frequently implicated in the multi-step carcinogenesis of CCA.⁽³²⁾ Specifically, HER2 is a well-described predictive biomarker for positive anti-HER2 therapy response in breast and gastric cancer and, lately, in CCA.^(33,34) Our first results showed dramatically increased ErbB2 protein and mRNA levels in livers of NEMO^{Δhepa}/JNK^{Δhepa} compared with NEMO^{Δhepa} hepatic tissue (Fig. 5B-D). Interestingly, we also found increased expression of the EGFR ligand *egf* (Fig. 5C). Consistently, the levels of EGFR phosphorylation (pEGFR) were markedly elevated in the livers of NEMO^{Δhepa}/JNK^{Δhepa} mice (Fig. 5D).

The rapidly accelerated fibrosarcoma (RAF)-mitogen-activated protein kinase kinase (MEK)-extracellular signal-regulated kinase (ERK) transduction pathway is a key signaling cascade that regulates cellular proliferation, differentiation, and apoptosis and is frequently dysregulated in HCC⁽³⁵⁾ and in biliary tract cancer.⁽³⁶⁾ Binding of EGF to EGFR triggers its tyrosine kinase activity downstream signaling leading to the end phosphorylation of MEK1/2 and ERK1/2, with translocation of ERK to the nucleus and expression of genes related to proliferation. Increased phosphorylation (i.e., activation) of RAF, MEK1/2, and ERK/2 was

found in NEMO^{Δhepa}/JNK^{Δhepa} mice (Fig. 5D) associated with biliary overgrowth. These results were further corroborated in livers of 13-week-old mice

(Supporting Fig. S12), indicating that the RAF-MEK-ERK pathway was constitutively activated at early stages of cholangiocarcinogenesis.

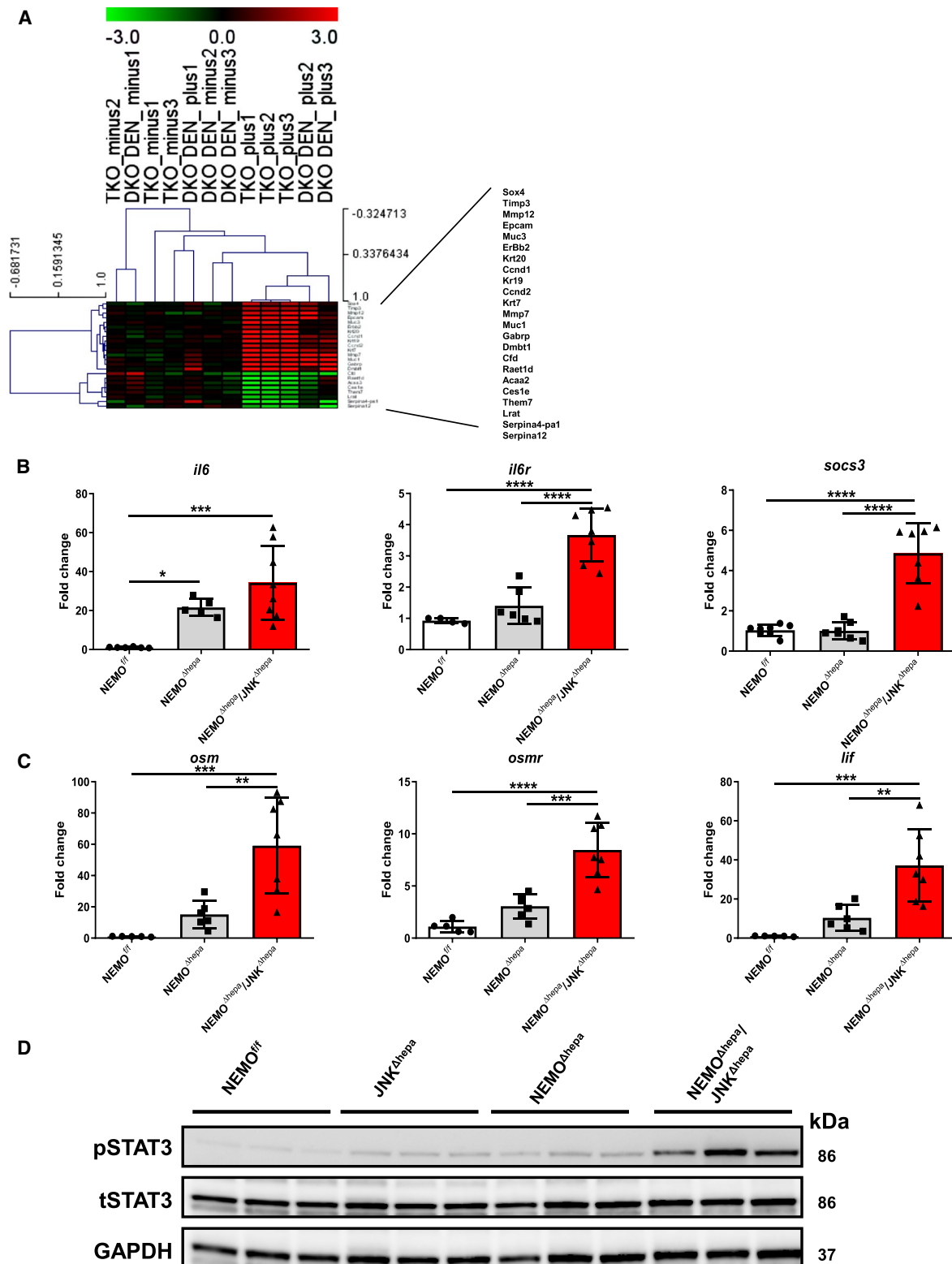


FIG. 4. The IL-6/STAT3 pathway is pivotal in biliary cell proliferation during liver carcinogenesis. (A) Gene array analysis was performed in 8-week-old NEMO^{Δhepa}/JNK^{f/f} and NEMO^{Δhepa}/JNK^{Δhepa} livers in addition to 26-week-old wild type and JNK^{f/f} and JNK^{Δhepa} livers challenged with DEN. Correlation of the fold induction of gene expression in liver is shown. Log₂ expression values of the individual mice were divided by the mean of the NEMO^{Δhepa}/JNK^{Δhepa} mice. Log ratios were saved in a .txt file and analyzed with the Multiple Experiment Viewer. Top up-regulated and down-regulated target substrates are shown (red, up-regulated; green, down-regulated; n = 3, 3.0 < fold change > 3.0). (B) mRNA expression analysis of *il6* (left); *il6r* (center), and *socs3* (right). (C) mRNA expression analysis of *osm* (left), *osmr* (center), and *lif* (right) was quantified by qRT-PCR of samples taken from NEMO^{f/f}, NEMO^{Δhepa}, and NEMO^{Δhepa}/JNK^{Δhepa} livers killed at 52 weeks. (D) Protein expression levels of STAT3 and pSTAT3 from whole-liver extracts of 52-week-old NEMO^{f/f}, JNK^{Δhepa}, NEMO^{Δhepa}, and NEMO^{Δhepa}/JNK^{Δhepa} mice were analyzed by western blot with the indicated antibodies. GAPDH served as loading control. Abbreviations: DKO, JNK1^{f/f}/JNK2^{-/-} + DEN; LIF, leukemia inhibitory factor; osmr, oncostatin M receptor; TKO, NEMO^{Δhepa}/JNK^{Δhepa}.

LAPATINIB TREATMENT PREVENTS BILIARY EXPANSION AND DUCTO/CYSTOGENESIS IN NEMO^{Δhepa}/JNK^{Δhepa} MICE

Our data strongly indicate the EGFR-HER2 pathway being involved in biliary cell hypergrowth and cholangiocarcinogenesis in NEMO^{Δhepa}/JNK^{Δhepa} livers. Lapatinib is a dual EGFR2/HER2 tyrosine kinase inhibitor (TKI) targeting both EGFR and HER2.^(37,38) It successfully inhibited the growth of HER-overexpressing breast cancer cells in culture and in tumor xenografts.⁽³⁹⁾ Taking into consideration the promising therapeutic option of lapatinib, we subsequently explored the potential beneficial effect of this dual TKI in our model of cholangiocellular proliferation.

Lapatinib was administered daily by oral gavage for 6 weeks in NEMO^{f/f}, NEMO^{Δhepa}, and NEMO^{Δhepa}/JNK^{Δhepa} mice (Fig. 6A). The TKI did not cause any relevant histopathological or serum biochemistry alterations to NEMO^{f/f} or NEMO^{Δhepa} animals (Fig. 6B). However, lapatinib treatment of NEMO^{Δhepa}/JNK^{Δhepa} mice resulted in a significant reduction of cyst-like structures (Fig. 6B). Lapatinib also significantly reduced serum AST and ALT levels in NEMO^{Δhepa}/JNK^{Δhepa} compared with NEMO^{f/f} and NEMO^{Δhepa} animals (Fig. 6C).

Next, the impact of lapatinib on EGFR/HER2 signaling was tested by investigating the downstream RAF-MEK-ERK pathway below EGFR/HER2. Decreased activation of EGFR and abrogation of RAF, MEK, and ERK signaling pathways were found in lapatinib-treated livers compared with vehicle-administrated NEMO^{Δhepa}/JNK^{Δhepa} livers (Fig. 6D). These results suggest that lapatinib successfully decreases EGFR-HER2 signaling and functionally links inhibition of this pathway with biliary overgrowth in NEMO^{Δhepa}/JNK^{Δhepa} livers.

COMBINED JNK1/JNK2 DELETION IS ESSENTIAL FOR HYPERPROLIFERATION OF BILE DUCTS

Our data were generated in global JNK2^{-/-} mice. Hence, we aimed to distinguish if hepatocytic or nonparenchymal JNK2 function is essential to direct bile duct proliferation. We applied our recently developed hepatocyte-specific *Jnk2* siRNA (*siJnk2*) protocol using lipid nanoparticles⁽¹⁶⁾; *siLuc* served as controls. We included 6- to 8-week-old untreated floxed NEMO^{f/f}/JNK^{f/f}, *siLuc*-treated NEMO^{Δhepa}/JNK1^{Δhepa}, and *siJnk2*-challenged NEMO^{Δhepa}/JNK1^{Δhepa} mice in this analysis. In parallel, animals were treated with vehicle or lapatinib.

Interestingly, *Jnk2* knockdown in hepatocytes triggered massive biliary cyst formation in livers of vehicle-treated NEMO^{Δhepa}/JNK1^{Δhepa} animals compared with *siLuc*-NEMO^{Δhepa}/JNK1^{Δhepa} and untreated floxed NEMO^{f/f}/JNK^{f/f} mice (Supporting Fig. S13A). These results demonstrate that combined *Jnk1/2* deletion in hepatocytes is responsible for directing biliary hyperproliferation in this model. Additionally, lapatinib treatment successfully reduced liver cystogenesis and significantly ameliorated serum transaminases in NEMO^{Δhepa}/JNK1^{Δhepa} + *siJnk2* livers, confirming the efficacy of this TKI in experimental biliary cystogenesis (Supporting Fig. S13B-D).

Discussion

Hyperplasia of the biliary epithelia with variable atypia in cystic bile ducts may give rise to malignant transformation leading to intrahepatic CCA for which an enormous unmet clinical and research need exists. CCA is an epithelial neoplasm derived from

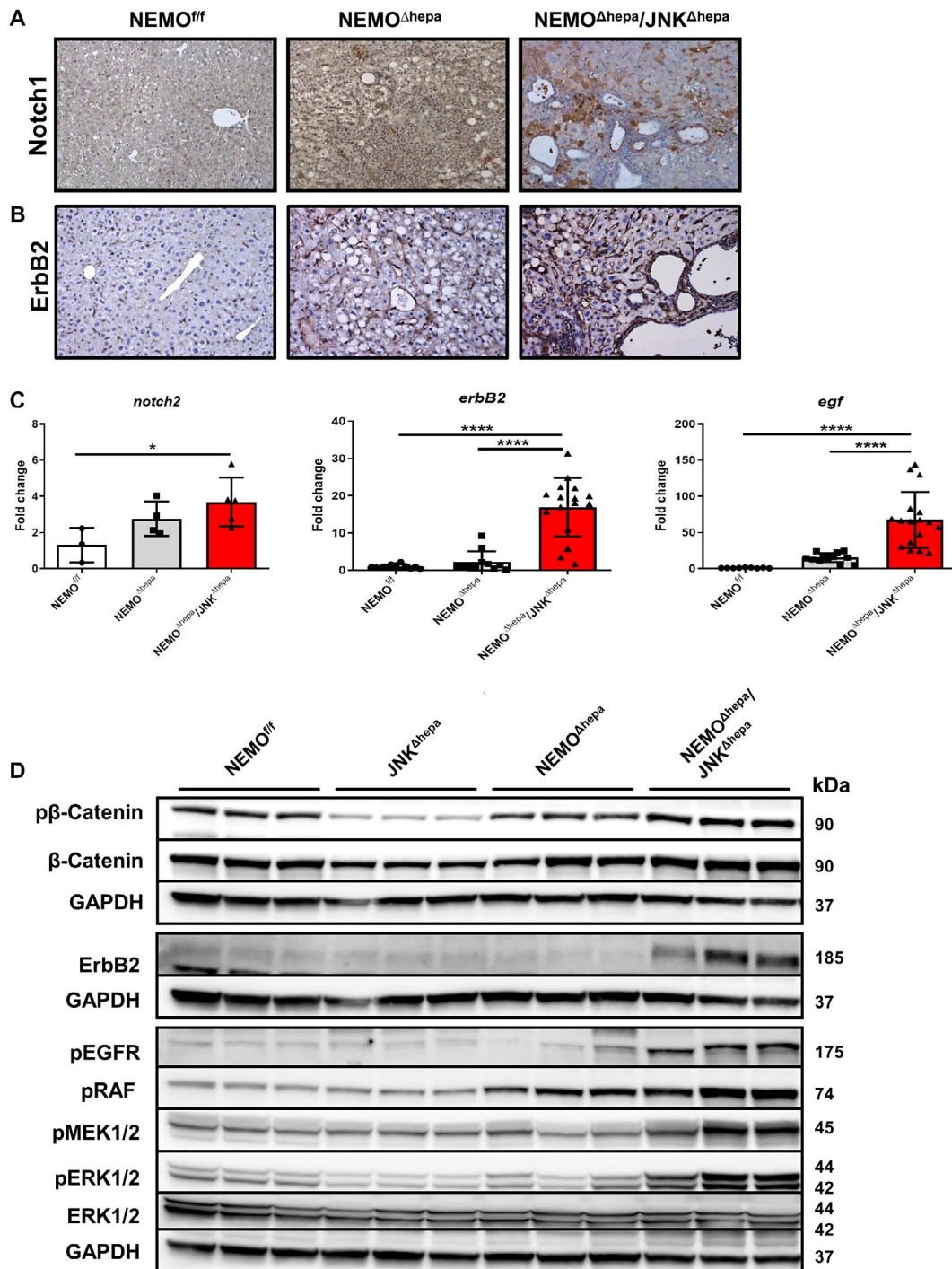


FIG. 5. Activation of EGF-EGFR-RAF-MEK1/2-ERK1/2 is distinctive of NEMO^{Δhepa}/JNK^{Δhepa} mice. (A) Representative IHC for NOTCH-1 staining of liver sections of 52-week-old NEMO^{fl/fl}, NEMO^{Δhepa}, and NEMO^{Δhepa}/JNK^{Δhepa} livers, magnification is 20×. (B) Representative IHC for ErbB2 staining of liver sections of the same livers. (C) mRNA expression analysis of *notch-2* (left), *erbB2* (center), and *egf* (right) was quantified by qRT-PCR of samples taken from NEMO^{fl/fl}, NEMO^{Δhepa}, and NEMO^{Δhepa}/JNK^{Δhepa} mice killed at 52 weeks. Data are presented as mean ± SEM. **P* < 0.05; *****P* < 0.0001. (D) Protein expressions of phospho-β-catenin, β-catenin, ErbB2, phospho-EGFR, phospho-RAF, phospho-MEK1/2, phospho-ERK1/2, and total ERK1/2 from whole-liver extracts of 52-week-old NEMO^{fl/fl}, JNK1^{Δhepa}/JNK2^{-/-}, NEMO^{Δhepa}, and NEMO^{Δhepa}/JNK^{Δhepa} mice were analyzed by western blot with the indicated antibodies. GAPDH served as loading control. Abbreviation: phospho, phosphorylated.

primary and secondary bile tracts and accounts for 5%-10% of primary liver cancer, the incidence and mortality of which are steadily increasing. The 5-year

survival of patients with CCA remains unacceptably low, and survival has not dramatically improved in the past 20 years.⁽⁴⁰⁾ This is in part due to a lack of

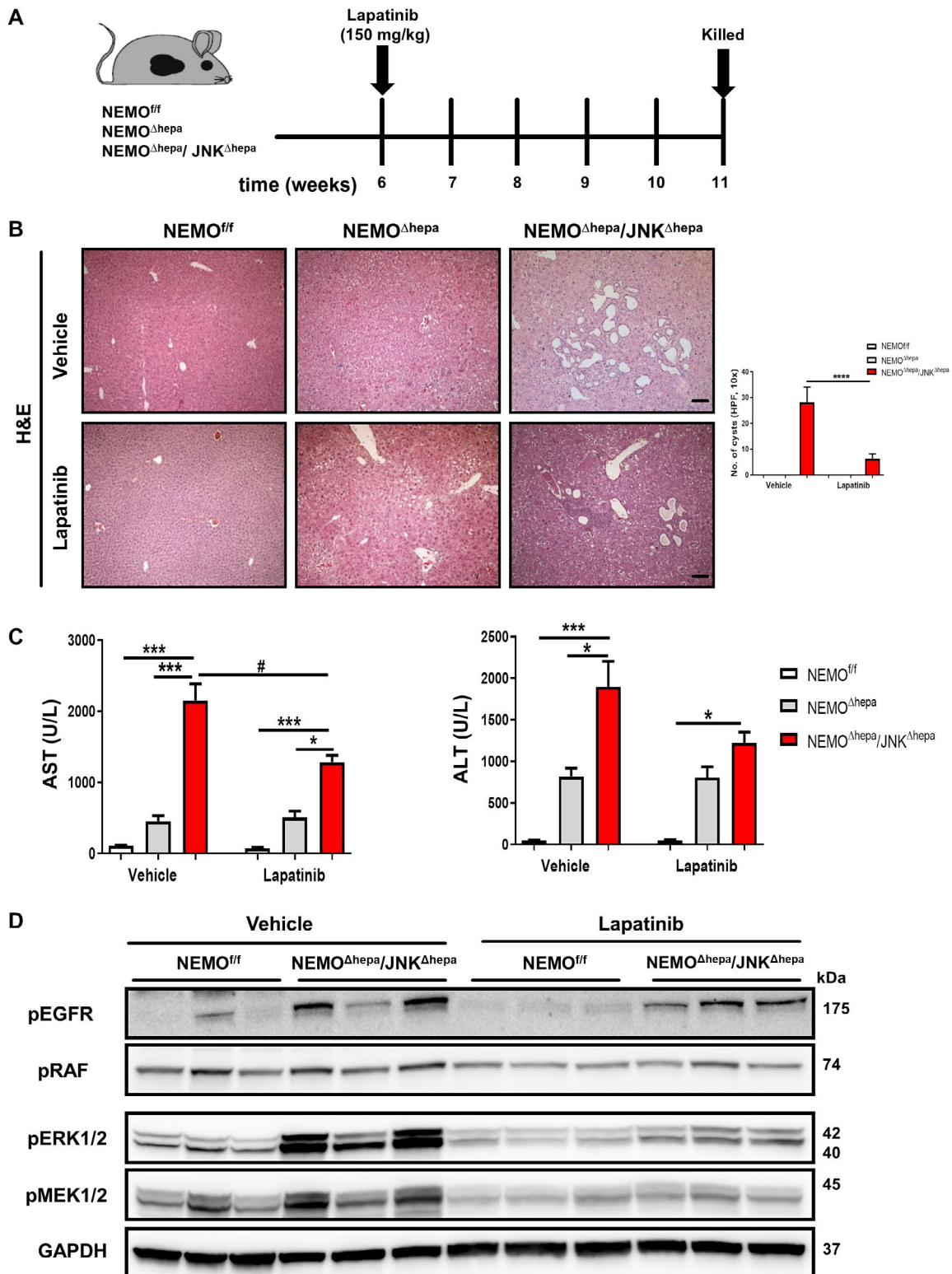


FIG. 6. Lapatinib, a dual tyrosine kinase inhibitor, protects against hyperbiliary proliferation in 52-week-old NEMO^{Δhepa}/JNK^{Δhepa} animals. (A) Experimental design of lapatinib (150 mg/kg BW) or vehicle administration to 6-week-old NEMO^{Δhepa}, NEMO^{Δhepa}/JNK^{Δhepa}, and NEMO^{Δhepa}/JNK^{Δhepa} mice. (B) Representative H&E staining of liver sections of NEMO^{Δhepa}, NEMO^{Δhepa}/JNK^{Δhepa}, and NEMO^{Δhepa}/JNK^{Δhepa} mice treated with either vehicle or lapatinib for 6 weeks (left), magnification is 10×. Number of visible microscopic cysts per 10× view field were calculated and graphed (right). (C) Serum levels of AST and ALT, in NEMO^{Δhepa}, NEMO^{Δhepa}/JNK^{Δhepa}, and NEMO^{Δhepa}/JNK^{Δhepa} mice after 6 weeks of lapatinib treatment. Data are presented as mean ± SEM. **P* < 0.05, ****P* < 0.001, #*P* < 0.05 (intergroup). (D) Levels of phospho-ERK1/2, phospho-MEK1/2, phospho-RAF, and phospho-EGFR from whole-liver extracts of the indicated genotypes treated with vehicle or lapatinib were analyzed by western blot with the indicated antibodies. GAPDH served as loading control. Abbreviations: H&E, hematoxylin and eosin; phospho, phosphorylated.

understanding of the pathophysiologic mechanisms underlying CCA. Because biliary tract cancer is often diagnosed late, the success of the only curative procedure (surgical resection) is limited, and the lack of biomarkers or diagnostic tools that would lead to early diagnosis is a matter of concern.⁽⁴¹⁾

A recent study⁽¹²⁾ opened a Pandora's box for therapeutic options targeting the JNK signaling pathway, a major regulator of cell proliferation, against CCA. Earlier, several studies confirmed the critical role of the JNK signaling pathway in liver cancer.^(10,19,42-44) At present, little is known on the role of JNK in directing differentiation of LPCs and more specifically of bipotential hepatic cells not only in liver homeostasis but also following liver injury.

We first focused on the specific roles of the *Jnk* genes in the progression of experimental chronic liver disease using NEMO mice and showed how *Jnk1* or *Jnk2* tipped the balance toward HCC or necroinflammation, respectively.⁽¹⁹⁾ We later demonstrated that combined *Jnk1/2* deletion in hepatocytes triggers more severe liver injury, inflammation, and progression after toxic liver injury.⁽¹⁵⁾

Thus, we next sought to investigate the consequences of hepatocytic *Jnk1/2* ablation in liver parenchymal proliferation and growth in experimental HCC. For this purpose, we generated NEMO^{Δhepa}/JNK^{Δhepa} mice, which displayed reduced tumor burden, despite the fact that they had signs of jaundice and hyperbilirubinemia, compared to NEMO-deficient mice developing HCC.⁽¹⁸⁾ Remarkably, *Jnk1/2*-deleted NEMO^{Δhepa} livers exhibited hyperproliferation of the biliary epithelium, forming cyst-like structures compatible with cholangioma or malignant CCA, as assessed by two independent pathologists (Table 1).

These livers were characterized by cell death, inflammatory microenvironment, and ECM deposition. Necroptosis-associated hepatic cytokine microenvironment induces the shift from HCC to CCA development.⁽²²⁾ NEMO^{Δhepa}/JNK^{Δhepa} livers display

considerable CC3 staining and RIPK3 protein levels, triggering exacerbated compensatory proliferation of LPCs, which are associated with increased ROS production and failure of the antioxidant defense.

Moreover, deposition of ECM and periductal/pericyclic scar formation was a prominent feature in NEMO^{Δhepa}/JNK^{Δhepa} livers. A unique characteristic of CCA is the presence of cancer-associated fibroblasts (CAFs) surrounded by numerous immune cells.⁽⁴¹⁾ CAFs promote the secretion of chemokines/cytokines including EGF in CCA cell lines.⁽⁴⁵⁾ Moreover, TNF might be another important culprit in CCA development, as suggested.⁽¹²⁾

In parallel with proliferating hepatocytes and a marked expansion of the biliary epithelium, the liver parenchyma of NEMO^{Δhepa}/JNK^{Δhepa} animals was positive for CK19 and SOX-9 and negative for HNF-4α. Moreover, we found increased expression of mucin genes. Considering that CCA tissues are characterized by the presence of mucin-secreting cells, this finding further supports the CCA diagnosis.⁽⁴⁶⁾ Interestingly, analysis of gene expression showed the occurrence of CCA/epithelial-transformed neoplasia-enriched markers, including *Dmbt1* and *Gabrp*, not only in NEMO^{Δhepa}/JNK^{Δhepa} liver but also in a second model of liver carcinogenesis, the DEN model. Confirmation in the two models suggests that CCA development relies on *Jnk1/2* combined function in hepatocytes but is independent of NF-κB activity in LPCs. Moreover, microarray studies highlighted the pivotal role of *Jnk1/2* in modulating cell fate by promoting hepatocarcinogenesis and under-regulating cascades linked with cholangiocarcinogenesis. Our data undoubtedly indicate that loss of *Jnk1/2* function promoted CCA in both experimental CLD and chemically induced HCC.

Jnk1/2-deleted NEMO livers exhibited strong expression of NOTCH-1/A6 and *Notch-2* and a clear tendency toward increased expression of Notch signaling pathway effectors. These results are consistent

with evidence from mouse studies, suggesting that NOTCH and WNT/ β -catenin are key drivers of CCA development. Specifically, NOTCH-1, -2, and -3 were shown to be overexpressed in human cholangiocellular injury.⁽⁴⁷⁾ In contrast, no differences in β -catenin expression between NEMO ^{Δ hepa} and NEMO ^{Δ hepa}/JNK ^{Δ hepa} were observed, indicating that hepatocyte differentiation in our model is inhibited in favor of ductular reaction and cholangiocyte differentiation. Of note, JNK1/2-knockout mice have reduced phosphorylation of β -catenin, suggesting that JNK is necessary for β -catenin phosphorylation, as suggested.⁽³¹⁾

Our protein and mRNA data overwhelmingly indicated that overexpression and activation of the EGFR/ErbB2 family was implicated in multistep cystogenesis toward CCA in NEMO ^{Δ hepa}/JNK ^{Δ hepa} livers. Indeed, EGFR overexpression occurs in 11%-27% of human CCA, whereas HER2 overexpression is less frequent but very characteristic in transgenic mouse models.^(36,48) It has been reported that EGF is released by tumor-associated macrophages (TAM).⁽⁴⁹⁾ Binding of EGF to its receptors induces their homodimerization or heterodimerization, which in turn activates downstream signaling pathways that regulate cell differentiation, migration, angiogenesis, and survival.⁽⁵⁰⁾ In our study, we specifically focused on RAF-MEK1/2-ERK2 and JAK/STAT signaling. Both pathways were dramatically induced in NEMO ^{Δ hepa}/JNK ^{Δ hepa} livers, suggestive of the strong proliferative and inflammatory microenvironment within the hepatic parenchyma.

Therefore, we tested the possibility of blocking EGFR/HER2 signaling as a novel strategy in treating hyperproliferation of the biliary epithelium. We used lapatinib, a dual TKI that efficiently inhibits both EGFR and HER2. Treatment not only prevented RAF-MEK-ERK activation but also inhibited biliary cyst formation. However, its efficacy in clinical translation may highly depend on enrollment of patients with EGFR/HER2 hyperactivation or overexpression.

Recently, Heikenwalder's group showed the therapeutic relevance of inhibiting JNK in CCA both *in vivo* and in cell lines.⁽¹²⁾ LPC-specific *Jnk1/2* knockout mice were used in two different CCA models. In contrast to our observations, they found reduced cholangiocellular injury in both models. This apparent discrepancy between both studies may be reconciled considering that adeno-associated virus-mediated *Cre* expression in hepatocytes might not exactly resemble

Alb-Cre excision since birth, as employed in our study. To further confirm the implications of *Jnk1/2* ablation in hepatocytes, we blocked *Jnk2* specifically in hepatocytes using a liposome-delivery system coupled to an siRNA that was recently reported by our laboratory.⁽¹⁶⁾ Our data undoubtedly indicated that *Jnk1/2* in hepatocytes prevents biliary cell hyperproliferation. In addition, lapatinib successfully prevented activation of the EGFR-RAF-MEK1/2-ERK1/2 pathway in *Jnk1/2* mice. However, more studies (e.g, assessment of JNK levels *in vitro* and in organoids) need to be performed to better define the specific role of JNK for CCA initiation.

Overall, our results show that complete inhibition of JNK signaling in hepatocytes in an experimental HCC model triggers binding of EGF (most likely released by the increased TAM-derived environment) to its receptor activating EGFR-RAF-MEK1/2-ERK1/2 signaling. This cascade is essential to drive EMT transdifferentiation of oval cells into biliary cells and massive ducto/cystogenesis, which shares the molecular features of CCA. Here, CAF-derived cytokines, including *tnf* or transforming growth factor beta (*tgfb*), might further contribute to exacerbated proliferation of biliary cells (Supporting Fig. S14).

Our study better delineates the pathogenesis of CCA by describing a novel function of JNK in cholangiocyte hyperproliferation. It also defines new therapeutic options to inhibit pathways involved in cholangiocarcinogenesis.

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