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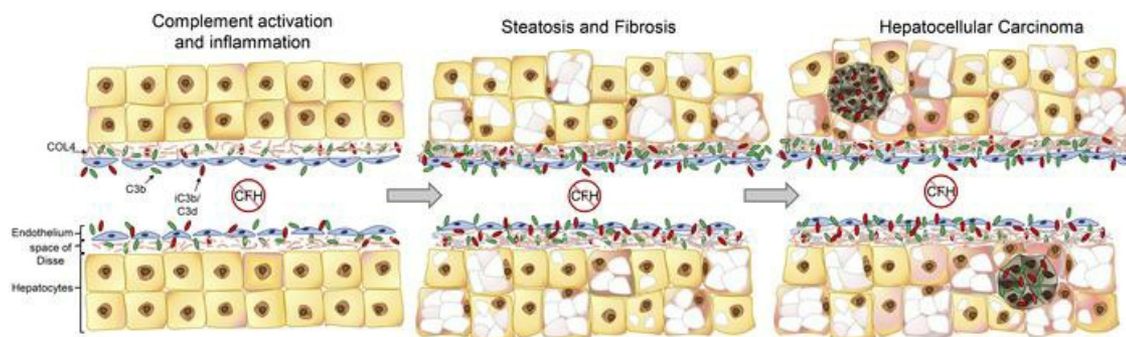
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Complement factor H-deficient mice develop spontaneous hepatic tumors

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Abstract

Hepatocellular carcinoma (HCC) is difficult to detect, carries a poor prognosis, and is one of few cancers with an increasing yearly incidence. Molecular defects in complement factor H (*CFH*), a critical regulatory protein of the complement alternative pathway (AP), are typically associated with inflammatory diseases of the eye and kidney. Little is known regarding the role of CFH in controlling complement activation within the liver. While studying aging CFH-deficient (*fH*^{-/-}) mice, we observed spontaneous hepatic tumor formation in more than 50% of aged *fH*^{-/-} males. Examination of *fH*^{-/-} livers (3-24 months) for evidence of complement-mediated inflammation revealed widespread deposition of complement activation fragments throughout the sinusoids, elevated transaminase levels, increased hepatic CD8⁺ and F4/80⁺ cells, overexpression of hepatic mRNA associated with inflammatory signaling pathways, steatosis and increased collagen deposition. Immunostaining of human HCC biopsies revealed extensive deposition of complement fragments within the tumors. Interrogation of The Cancer Genome Atlas also revealed that increased *CFH* mRNA expression is associated with improved survival in HCC patients, whereas mutations are associated with worse survival. These results indicate that CFH is critical for controlling complement activation in the liver, and in its absence, AP activation leads to chronic inflammation and promotes hepatic carcinogenesis.

Introduction

Key functions of the immune system are the ability to discriminate self from non-self, and to rapidly eliminate invasive pathogens while causing minimal injury to the host. Rapid induction of inflammation is critical for an efficient immune response and its rapid resolution prevents unintended damage. If left unresolved, however, inflammation can damage surrounding tissues, eventually leading to fibrosis, organ failure, and carcinogenesis (1, 2). The complement cascade is an important arm of the immune system and is controlled through a balance of activating and regulatory proteins (3). The classical and mannose binding lectin pathways are activated by antibodies and other pattern recognition molecules (4). The alternative pathway (AP), however, is distinct in that it is continually activated in plasma through a process called “tick-over,” indiscriminately depositing activated C3b on nearby surfaces. The deposited C3b catalyzes further AP activation, self-amplifying unless adequately controlled on the target surface by complement regulatory proteins (CRPs). CRPs selectively protect host cells but not invasive pathogens, thereby creating a system of rapid and continuous immune surveillance.

CRPs prevent autologous complement-mediated injury by cleaving and inactivating C3b (a component of the C3 and C5 convertases), and/or accelerating the decay of existing C3 and C5 convertases (3). Because the AP is constitutively activated, inadequate regulation can lead to complement-mediated injury. Congenital and acquired defects in complement regulation are associated with severe diseases, including atypical hemolytic uremic syndrome (aHUS), C3 glomerulopathy (C3G), and protein-losing enteropathy (5-7). Genetic variants of AP proteins are also associated with accelerated progression of some chronic diseases, including age related macular degeneration (AMD) and IgA nephropathy (8, 9). Complement activation has been linked with the development and spread of several cancers (10-12), raising the possibility that impaired complement regulation could be a risk factor for some types of cancer.

Factor H (CFH) is a soluble CRP that regulates AP activation in plasma and on host surfaces through its ability to bind to self-surface ligands. Regions within the CFH protein bind to anionic molecules, including glycosaminoglycans (GAGs), which are highly expressed on basement membranes and extracellular matrix (ECM), and sialic acid, a common component of cellular membranes (13, 14). Engagement of these molecules tethers CFH to host tissues, controlling AP activation at those sites (15). Although sulfated GAGs and sialic acids are ubiquitously expressed, CFH binding and regulatory activity are particularly important for specific tissues (16). For example, CFH is critical for controlling AP activation on ECM and basement membranes, as these surfaces do not intrinsically express other CRPs. Mutations in CFH binding regions are primarily associated with diseases of the kidneys and eyes (8, 17), possibly because these organs have specialized basement membranes that are highly exposed to plasma proteins.

While studying complement-mediated kidney disease, we observed spontaneous liver tumor development in *fH*^{-/-} male mice at a significantly higher rate than control mice. We hypothesized that CFH deficiency causes complement-mediated liver inflammation. Chronic inflammation can promote development of HCC (18), and complement activation can also impair the elimination of tumors by the immune system (19). Consequently, inadequate complement regulation within the liver could contribute to carcinogenesis by several mechanisms. CFH is primarily generated in the liver, as are factor B (CFB) and C3 (activating components of the AP) (20, 21). Proteins produced by hepatocytes enter the circulation by passage through the ECM-rich space of Disse and fenestrated endothelial cells lining the sinusoids, continually exposing these regions to high concentrations of complement proteins. The mechanisms of complement activation and regulation within the liver are incompletely

understood and a direct connection between CFH dysfunction and liver disease, to our knowledge, has not been previously described.

In the current study, we examined the role of CFH in regulating complement activation in the liver, and the extent to which complement activation is associated with hepatocellular injury. We examined the link between local complement activation in *fH*^{-/-} mice with changes in the hepatic immune milieu. Finally, we analyzed biopsies and clinical data from patients with HCC to look for evidence of complement dysregulation in human HCC.

Results

Mice lacking complement factor H spontaneously develop hepatocellular carcinoma

While studying aged $fH^{-/-}$ mice, we noticed a high incidence of spontaneous liver tumor development, which was especially pronounced in males (Figures 1A and 1B) compared to WT mice (Figure 1C). In male mice 15 months and older (≥ 15 months), we observed grossly visible liver tumors in 54% of $fH^{-/-}$ (n=35) compared to 10% in $fH^{-/-}fB^{-/-}$ (which are unable to activate the AP; n=10) and 7% in WT (n=29). The rate of liver tumor incidence was reduced in female mice ≥ 15 months, with 15% incidence in $fH^{-/-}$ (n=26), 3% in WT (n=29), and 0% in $fH^{-/-}fB^{-/-}$ (n=10) mice (Figure 1D). Histological evaluation of livers from WT and $fH^{-/-}$ mice from 3 different age groups (≤ 5 months, 6-14 months, and ≥ 15 months) revealed notable differences between the two strains. Hepatocyte ballooning (an indicator of hepatocyte stress or injury) was present in $fH^{-/-}$ males at all ages (Figure 1E), with multiple instances per field of view observed in over half of the ≥ 15 -month males, compared to 14% of WT males of the same age. In male mice ≥ 15 -months of age, multi-focal tumor rosettes or nests were seen in all $fH^{-/-}$ mice with visible liver tumors (Figure 1F), and in 50% of those without. Generally normal liver parenchyma was observed in WT males of the same age (Figure 1G).

Glypican-3 (GPC3) is a heparan sulfate proteoglycan that is overexpressed in HCC and has been used as a tumor biomarker (22, 23). Variation in the location of GPC3 expression has been noted in human HCC, with localization occurring on cell membranes in some and in the cell cytoplasm of other tumors (24). As further support that tumors in $fH^{-/-}$ mice are HCC, we stained livers collected from animals with and without visible tumors for GPC3 expression. We observed two distinct GPC3 expression patterns in the $fH^{-/-}$ mice, which correlated with the macroscopic findings. Livers from $fH^{-/-}$ mice with large focal tumors surrounded by otherwise

normal appearing parenchyma (Figure 1A), expressed GPC3 in a continuous, membranous pattern (Figure 1H). In contrast, GPC3 was predominantly cytoplasmic (Figure 1I) in *fH*^{-/-} livers with multi-focal tumors extending throughout the parenchyma of each lobe (Figure 1B, yellow arrows). Only membranous GPC3 deposition was seen in *WT* livers with tumors (Figure 1J).

C3 fragments are deposited throughout the livers of *fH*^{-/-} mice

During AP activation, intact fluid-phase C3 is cleaved to form C3b, which covalently attaches to nearby tissues. C3b is subsequently cleaved, generating iC3b and C3d. We immunostained liver and kidney sections from 3-5-month-old *WT*, *fH*^{-/-}, CFH heterozygotes (*fH*^{+/-}), and *fH*^{-/-}*fB*^{-/-} mice to evaluate hepatic C3 fragment deposition. In *WT* and *fH*^{+/-} mice, we found only trace amounts of punctate C3b and iC3b/C3d in the liver sinusoids (Figures 2A and Supplementary Figure S1A, respectively). In *fH*^{-/-} mice, however, C3b and iC3b/C3d were deposited extensively throughout the sinusoids (Figure 2B). C3b-iC3b/C3d deposition was linear and continuous within the sinusoidal spaces, and appeared to deposit in a gradient, with concentrated C3 in the perivenous regions adjacent to the central vein, becoming more diffuse in the periportal regions. C3b deposits colocalized with collagen IV (COL4) [a component of the basement membrane (25)] and adjacent to MECA-32 (an endothelial cell marker), further supporting that complement was activated within the sinusoidal wall (Supplementary Figures S2A and S2B). Deficiency of both CFH and CFB completely abrogated C3 deposition in livers of *fH*^{-/-}*fB*^{-/-} mice (Figure 2C). *fH*^{-/-} mice are most frequently used to study C3G which is characterized by C3 deposition localized in the glomeruli (Figure 2E), a location known to be particularly vulnerable to AP activation in the absence of CFH. C3b-iC3b/C3d deposition for each strain has been previously described in detail (26), and representative staining is shown

(Figures 2D-2F and Supplementary Figure S1A). These data demonstrate that, like the glomeruli, hepatic C3 deposition in *fH*^{-/-} mice is largely due to AP activation.

CFH is a systemic regulator; therefore, *fH*^{-/-} mice could also be susceptible to AP activation in other organs. To determine other possible locations of AP activation in *fH*^{-/-} mice, we performed whole animal PET imaging to detect localized iodinated (¹²⁴I)-C3d29 (monoclonal antibody specific for the complement fragments iC3b and C3d) (27). ¹²⁴I-C3d29 localization was only seen in the livers and kidneys of *fH*^{-/-} mice (non-specific signal in thyroid and bladder is likely due to uptake or clearance of free iodine) (Figure 2I). Bound ¹²⁴I-C3d29 was quantified (28) 96 hours after imaging (Figure 2H), which confirmed the liver and kidneys are the predominant sites of C3 deposition in *fH*^{-/-} mice.

AP activation leads to hepatocellular injury in *fH*^{-/-} mice

To assess the pathological effects of unregulated AP activation in the liver, we measured levels of alanine transaminase (ALT), aspartate transaminase (AST), alkaline phosphatase (ALP), and albumin in 3-month old *WT*, *fH*^{-/-}, and *fH*^{-/-}*fB*^{-/-} mice. ALT and AST were significantly higher in *fH*^{-/-} mice than in *WT* or *fH*^{-/-}*fB*^{-/-} mice (Figure 3A), confirming hepatocellular injury in *fH*^{-/-} mice. ALP and albumin were not different between the strains, indicating that the injury had not caused biliary obstruction or impaired synthetic function of the liver at this timepoint (29).

We stained liver sections from *fH*^{-/-} and *WT* males at 3, 9, and 18 months of age with Oil Red O, a lysochrome used to detect lipid. Lipid accumulation was mild and limited to the perivenous hepatocytes of 3-month-old *fH*^{-/-} mice but became progressively more widespread with age (Figure 3B, right panels). Positive staining was seen in the 9- and 18-month-old *WT*

mice (Figure 3B, left panels), but when quantified, was found to be significantly less than in age-matched *fH*^{-/-} mice (Figure 3C).

We next immunostained livers from 14-month old *fH*^{-/-} and *WT* mice for COL4, a structural component of basement membranes, that when deposited in excess, leads to the development of fibrosis (30). There was greater COL4 deposition in *fH*^{-/-} mice compared to *WT* (Figure 3E), occurring in a continuous pattern in the sinusoids of *fH*^{-/-} mice (Figure 3D, bottom). Only punctate deposits were seen in age-matched *WT* livers (Figure 3D, top). This suggests that the chronic inflammatory environment triggered by uncontrolled AP activation in *fH*^{-/-} livers contributes to the development of steatosis and fibrosis as the mice age.

Hepatic mRNA expression reveals a strong inflammatory signature in *fH*^{-/-} mice

We examined the mRNA expression profile of myeloid innate immune markers in the livers of 3-month-old *fH*^{-/-} mice compared to *WT*. When comparing the most differentially expressed (DE) mRNA to *WT* controls, *fH*^{-/-} livers had greater than 4.65-fold higher mRNA expression of *S100a9* and *S100a8*, the pro-inflammatory chemokine *Cxcl14*, and serum amyloid A 1 (*Saa1*) (Figures 4A and 4B). Of note, S100A9 is expressed on monocytes and neutrophils (31) and has been implicated in the development of cancer (32, 33) and tumor metastasis (34). *fH*^{-/-} mice also had increased expression of mRNA corresponding to proteins involved in the migration and activation of leukocytes, including *Itgal* (35), *Fcgr3* (36), and neutrophil cytosolic factor 2 (*Ncf2*) (Figure 4B).

Pathway analysis revealed an increase in *fH*^{-/-} liver mRNA expression in 16 of the 17 gene sets relevant to innate myeloid immune function (Figure 4C). Directed global significance (GS) scores were used to identify the most DE gene sets in *fH*^{-/-} mice as a group. Three of the

gene sets with high directed GS scores (indicating high expression compared to *WT* covariates), were the chemokine, cytokine, and Fc receptor signaling pathways. Composite scores for these three pathways (Figures 4C and 4D-4F), and the most DE genes corresponding to each of the three pathways with relative expression frequencies (Figures 4G-4I) are shown. These data provide transcript level evidence of a pro-inflammatory environment in the livers of young *fH*^{-/-} mice.

T-cell infiltration and reduced neutrophils in *fH*^{-/-} livers

To evaluate the immune milieu within the livers of *fH*^{-/-} mice, we stained liver sections for six general leukocyte markers (F4/80, CD8, CD4, CD3, B220, and Foxp3) and collected images with a Vectra quantitative pathology imaging system. Dense clusters of immune cells were observed in all *fH*^{-/-} livers near portal veins (Figure 5A, left and Figure 5B). Similar immune clusters were not seen in *WT* livers (Figure 5A, right). Significantly more F4/80⁺ and CD8⁺ T cells were seen in *fH*^{-/-} livers (Figure 5C), as well as a trend towards greater numbers of CD4⁺, CD3⁺, and CD4⁺Foxp3⁺ cells.

We assessed hepatic neutrophil populations in young *fH*^{-/-}, *WT*, and *fH*^{-/-}*fB*^{-/-} males by flow cytometry. There was greater than a 50% reduction in neutrophils (CD45⁺ CD11b⁺ Ly6G⁺) in *fH*^{-/-} livers compared to those from *WT* and *fH*^{-/-}*fB*^{-/-} (Figure 5D, left and middle). Because neutrophil infiltration within tumors may influence tumor growth (37), we assessed livers with and without tumors from 22-24-month-old *fH*^{-/-} mice. There was a trend towards a greater number of neutrophils in livers with tumors than in those without (Figure 5D, right). We stained *fH*^{-/-} liver sections containing tumor foci for neutrophils (Ly6G⁺ cells). Neutrophils were seen

within and around tumors (Figure 5E, arrows), providing evidence of neutrophil infiltration in *fH*^{-/-} liver tumors.

Skewed myeloid cell populations in *fH*^{-/-} livers

Complement fragments are chemoattractants for infiltrating monocytes, which have been shown to play an important role in shaping the tumor immune microenvironment (38). Previous studies have shown that MHC-II⁺ monocytes can inhibit tumor growth (39), whereas MHC-II⁻ monocytes, can be pro-angiogenic, suppress T cell function, and can promote tumor growth (39, 40). To examine whether the monocyte population is altered in livers of *fH*^{-/-} mice, we isolated and analyzed hepatic myeloid cells from young *fH*^{-/-}, *WT*, and *fH*^{-/-}*fB*^{-/-} males, as well as from aged *fH*^{-/-} males with and without liver tumors. Compared to *WT*, *fH*^{-/-} mice had fewer Ly6C^{high} MHC-II⁻ F4/80^{low/-} pro-inflammatory infiltrating monocytes (41) and significantly more Ly6C⁻ MHC-II⁻ F4/80^{low/-} anti-inflammatory patrolling monocytes and Ly6C^{int} MHC-II⁻ F4/80⁻ (Figure 6A). Only patrolling monocytes (Ly6C⁻ MHC-II⁻ F4/80^{low/-}) were increased in *fH*^{-/-} livers when compared to *fH*^{-/-}*fB*^{-/-} mice (Figure 6B). There were no significant differences in the monocyte populations between aged *fH*^{-/-} mice with liver tumors and those without. It is notable, however, that the Ly6C^{high} MHC-II⁻ F4/80^{low/-} infiltrating monocytes are the most numerous of these cell types in the aged *fH*^{-/-} livers, regardless of tumor status (Figure 6C), in contrast to what was seen in young *fH*^{-/-} mice.

There were no significant differences in the Ly6C⁻ MHC-II⁺ F4/80⁺ and Ly6C⁺ MHC-II⁺ F4/80⁺ macrophage populations between young *fH*^{-/-} mice and either *WT* or *fH*^{-/-}*fB*^{-/-} (Figures 6A and 6B) or between aged *fH*^{-/-} with and without liver tumors (Figure 6C). Young and aged *fH*^{-/-}

mice without tumors tend to have fewer macrophages than young *WT* or *fH^{-/-}fB^{-/-}* or aged *fH^{-/-}* with tumors, respectively.

These findings suggest that the hepatic monocyte population in young *fH^{-/-}* mice is primarily comprised of anti-inflammatory patrolling monocytes. As *fH^{-/-}* mice age and develop liver tumors, fewer pro-inflammatory Ly6C^{high} MHC-II⁺ F4/80^{low/-} monocytes transition to anti-inflammatory Ly6C⁺MHC-II⁺ F4/80^{low/-} patrolling monocytes (Figure 6D).

Recombinant murine factor H binds within the liver sinusoids and inactivates C3b

Given the pattern of C3 fragment deposition in *fH^{-/-}* livers, we examined whether CFH controls AP activation on basement membrane within the sinusoids and hepatic vasculature. We produced and functionally tested a recombinant murine CFH (rmCFH) (42), which was then conjugated with Alexa-647 and injected into male *fH^{-/-}* and *WT* mice. The mice were euthanized 24 hours later, and livers were examined for bound rmCFH. The rmCFH deposited along the sinusoids of *fH^{-/-}* livers (Figure 7A left, 7C, and 7D) in a pattern much like that of C3b deposition in unmanipulated *fH^{-/-}* mice (Figure 2B). We observed a similar pattern of rmCFH deposition in *WT* sinusoids (Figure 7A, right), although in significantly lower amounts (Figure 7B).

To assess the functionality of the rmCFH bound within the sinusoids, we stained liver tissues from the rmCFH-injected *fH^{-/-}* mice for C3b and iC3b/C3d and evaluated the sinusoids for differences in complement deposition (Figures 7C-7E). In rmCFH-reconstituted *fH^{-/-}* mice, we observed continuous deposition of iC3b/C3d in the sinusoids (Figures 7C and 7E) and a shift in C3b deposition from linear to punctate (Figures 7D and 7E). There was colocalization of rmCFH with iC3b/C3d (Figure 7C), and only sparse colocalization with C3b (Figure 7D). We quantified C3b and iC3b/C3d deposition in the reconstituted *fH^{-/-}* with that of unmanipulated *fH^{-/-}*

males of the same age. In reconstituted mice, there was a significant decrease in mean percent area C3b deposition (2% compared to 7% in unmanipulated mice) and a two-fold increase in iC3b/C3d (11% compared to 5% in unmanipulated mice) (Figure 7F). These data demonstrate that the injected rmCFH mediated cleavage of bound C3b in the sinusoids, which generated local iC3b/C3d.

Finally, to further characterize the specific location of rmCFH deposition in the sinusoids, we stained liver tissues from rmCFH-injected mice for COL4 and MECA-32. Similar to localization of C3b in the sinusoids, we observed regions of colocalization of rmCFH with both COL4 (Figure 7G) and MECA-32 (Figure 7H), indicating that rmCFH binds ECM and endothelial cells within the sinusoids and is critical for controlling alternative pathway activation at these sites.

Evidence of C3 inactivation within *fH*^{-/-} liver tumors

To examine complement regulation within liver tumors, we dual stained livers from tumor bearing mice for C3b and iC3b/C3d. In *WT* livers, we observed small tumor foci with heavy deposits of both C3b and iC3b/C3d, surrounded by normal appearing parenchyma with very little C3 deposition (Figure 8A, left panels). In contrast, *fH*^{-/-} tumors and surrounding parenchymal tissue displayed extensive C3b-iC3b/C3d deposition (Figure 8A, right panels). There were distinct regions within some of the *fH*^{-/-} tumors with minimal C3b and abundant iC3b/C3d deposition, suggesting that there was cofactor activity and inactivation of C3b within the tumor environment. To examine this further, we dual stained tumor tissues with GPC3 (to indicate tumor boundary) and either C3b or iC3b/C3d (Figures 8B and 8C). In tumors with membranous GPC3, we observed both C3b and iC3b/C3d within the tumor lesion (Figure 8B).

However, in tissues with cytoplasmic GPC3, C3b deposits were less apparent suggesting better complement regulation within these tumors (Figure 8C).

The presence of iC3b/C3d within *fH*^{-/-} liver tumors indicates that another protein serves as a cofactor for cleavage of C3b in the absence of CFH. Studies of human HCC have shown that tumors can upregulate membrane bound complement regulators, such as CD46 (membrane cofactor protein or MCP) (43, 44). Similar to CFH, CD46 is a cofactor for the cleavage of C3b, although it is reported to have limited expression in healthy mice (45). We stained *fH*^{-/-} liver tumors for CD46 and C3b and found prominent CD46 expression in some regions where C3b was absent (Figure 8D, white box). We also dual stained *fH*^{-/-} liver tumors for CD46 and iC3b/C3d. There were regions of high CD46 expression in which iC3b/C3d was not seen, as well as regions in which CD46 and iC3b/C3d colocalized. These results suggest that expression of CD46 within the HCC lesions may inhibit alternative pathway activation and it may also serve as a cofactor for the cleavage of cleavage of C3b.

C3 fragments are deposited in human HCC biopsies

We immunostained liver biopsy samples from six confirmed HCC patients for C3d/iC3b and GPC3 (Figure 9A), five of which were strongly positive C3d/iC3b. Among these five samples, there was colocalization of GPC3 and C3d/iC3b, indicating that there is complement activation within the tumors.

We next queried the TCGA through cBioPortal for Cancer Genomics (46, 47) to determine if *CFH* mutations or mRNA expression correlate with outcomes for patients with HCC. We found that *CFH* is altered in 23% (86 of 337) of the patients comprising this data set. Of those 86 patients, 50 had increased tumor *CFH* mRNA expression, seven had missense or

truncating mutations, and 29 had an amplification in *CFH* copy number (Supplementary Figure S4A). Increased expression of *CFH* is expected to help control complement activation, and we found that high *CFH* mRNA correlated with improved disease/progression-free survival as well as overall survival when compared to HCC patients with unaltered *CFH* mRNA (Figure 9B). The median disease/progression-free time for patients with high *CFH* mRNA was 55.06 months compared to 18.43 months for those with unaltered *CFH* mRNA expression. Conversely, *CFH* mutations are expected to increase complement activation in the liver and HCC tumors. Patients with *CFH* mutations had significantly worse overall survival, with a median disease/progression free time of only 9.72 months compared to 20.99 months for those without *CFH* mutations (Figure 9C). These findings demonstrate that the complement cascade is activated in the livers of patients with HCC, and that expression levels of *CFH* mRNA as well as mutations in the *CFH* gene significantly affect the prognosis for HCC patients.

To determine if these survival and prognostic outcomes relating to *CFH* mutations and increased *CFH* mRNA are unique to HCC, we queried *CFH* for all cancers in the TCGA Firehose Legacy study (Supplementary Table S1). We found that most cancer types could be grouped according to three categories: 1) worse outcome with *CFH* mutations, improved outcome with increased *CFH* mRNA, 2) worse outcomes for both *CFH* mutations and/or increased *CFH* mRNA, and 3) improved outcomes for both. We found three cancers for which the correlations were similar to HCC (worse outcomes with *CFH* mutations, improved outcomes with increased *CFH* mRNA): lymphoid neoplasm diffuse large B-cell lymphoma, lung adenocarcinoma, and esophageal carcinoma.

Discussion

Specific locations such as the eyes and kidneys, are known to be particularly dependent on CFH to control the AP. These sites are rich in ECM, have specialized basement membranes and are susceptible to AP-mediated inflammation in patients with *CFH* mutations. In the current study, we report that CFH is also critical for controlling AP activation in the liver. There is extensive complement activation throughout the sinusoids of *fH*^{-/-} mice, which is ameliorated through combined deficiency of CFH and CFB, confirming a principal role of the AP. AP activation in male *fH*^{-/-} mice was associated with hepatocellular inflammation and injury at 3 months of age, chronic liver damage and steatosis at 9 months, and the development of liver tumors for more than 50% of those aged to ≥ 15 months. This suggests that the hepatic inflammation resulting from chronic AP activation promotes a favorable environment for tumorigenesis.

C3b and iC3b/C3d were deposited in a continuous pattern along the sinusoids within the livers of *fH*^{-/-} mice. In mice reconstituted with rmCFH, the injected protein colocalized with COL4 (Figure 7G) and MECA-32 (Figure 7H) within the sinusoids and mediated cleavage of the deposited C3b to inactive iC3b/C3d. We developed a whole-body PET method to identify other sites of AP activation in *fH*^{-/-} mice and confirmed that the liver and kidneys are the primary sites of complement activation in the absence of CFH.

There is substantial experimental and clinical research linking genetic defects with *CFH* in kidney disease. Much less is known, however, regarding the role of CFH in liver disease. Although aHUS is primarily regarded as a kidney disease, it is noteworthy that abnormal liver function tests may be seen in up to 46% of patients (48). It has also been reported that complement-mediated injury can lead to hepatic failure in aHUS patients undergoing liver

transplantation (49). The presumed mechanism for this is inadequately controlled AP activation in the ischemic liver allograft, resulting from dysfunctional endogenous CFH. It is possible that the liver is a target organ in patients with aHUS associated *CFH* mutations but is overlooked due to the severe renal involvement. It is also possible that the binding mechanism of CFH to liver sinusoids and glomerular capillaries involves slightly different interactions. This is the case for AMD and aHUS, in which distinct genetic variants of *CFH* impair binding of the protein to either the eyes or the kidneys, thereby predisposing patients to one of these diseases but not the other (50).

CFH mutations have not previously been linked with the risk for HCC, although it has been reported that complement activation fragments are elevated in affected patients (51-53). In one study, complement fragments were the only independent plasma biomarker predictive of HCC by multivariate analysis (53). In the current study, we found evidence of complement activation in biopsy samples from six patients with HCC. This demonstrates that complement activation can occur within the tumors, similar to observations in mice. Furthermore, examination of the TCGA reveals that expression of *CFH* mRNA significantly correlates with survival in patients with HCC (Figure 9B). Although *fH*^{-/-} mice have complete deficiency of CFH, the TCGA data suggests that variations in CFH function and expression levels may also influence the prognosis of HCC. Future studies can examine whether CFH influences the growth of tumors by inhibiting complement within the tumor itself or through interactions in the tumor microenvironment.

It has been estimated that chronic liver inflammation may account for as much as 90% of HCCs, although the mechanisms linking inflammation with carcinogenesis are incompletely understood (54). Multiple studies have also reported that the complement cascade is activated in

the livers of patients with chronic hepatitis B (55, 56), hepatitis C (51, 52, 56), and alcoholic hepatitis (57), inflammatory diseases strongly associated with an increased risk of HCC. Based on the results of the current study, it is possible that inflammation resulting from uncontrolled complement activation within the liver is a common downstream pathway that links these chronic liver diseases with carcinogenesis. The heterogeneous pattern of complement activation within the tumors may be due to upregulation of CD46. It is also possible that additional factors promote complement activation once a cancer has formed. Antibodies may recognize tumor neoantigens, for example. Some tumor cells also produce enzymes that are capable of directly activating C3, such as cathepsins B and L (58).

Complement activation within the liver may contribute to the development of HCC by several mechanisms. Complement activation has been shown to trigger NF- κ B activation in Kupffer cells and STAT3 activation in hepatocytes (59), which may facilitate recovery of the liver after acute injury, but chronically promote hepatocyte proliferation and the development of HCC (18, 60, 61). Complement mediated inflammation in the livers of *fH*^{-/-} mice could also promote DNA damage, increasing the rate of cancer formation. Complement activation has also been shown to inhibit anti-tumor immunity in multiple types of cancer (10-12, 62, 63). For example, work has shown that C3a and C5a can induce suppressive myeloid cells or directly inhibit T cell proliferation and function (11, 63, 64). We found that the myeloid cell population is altered in the livers of *fH*^{-/-} mice, and a greater number of Ly6C^{int} MHC-II⁻ monocytes were present within the livers of these mice (Figures 6A and 6B). These cells can be immunosuppressive and may reduce anti-tumor immunity in *fH*^{-/-} livers. Thus, complement activation within the liver may both directly accelerate the development of HCC, as well as impair the ability of the immune system to eliminate these tumors once they have formed.

A limitation of our study is that it uses mice that are genetically deficient in CFH from birth. Many different signaling pathways are activated in these mice, and the development of chronic liver damage and HCC takes a relatively long time. An advantage of this model, however, is that it is characterized by spontaneous tumor formation. Most cancer models in immunocompetent mice involve genetic manipulation of tumor suppressor genes, chemical carcinogenesis, or transplant of cancer cells into recipient mice. Furthermore, in tumor transplant models, the cell lines are often derived from advanced stage tumors (65). These tumors therefore start with a high mutational burden and have already undergone immune selection. The *fH*^{-/-} mice, in contrast, provide the opportunity to study spontaneous HCCs as they develop.

In conclusion, we have found that CFH is necessary for controlling spontaneous AP activation in the liver. In the absence of CFH, dysregulated AP activation is associated with hepatocellular injury, steatosis, and development of HCC. We found evidence of complement activation in human HCC biopsies, and that *CFH* mutations and levels of *CFH* mRNA correlate with patient survival. There is clinical evidence that the complement system is activated in multiple chronic inflammatory diseases associated with increased risk of HCC. Complement activation may, therefore, be a common pathway linking viral, toxic, or autoimmune injury of the liver with carcinogenesis. Several therapeutic complement inhibitors are currently in development (66). Given the potential role of complement activation in liver carcinogenesis identified in this study, complement inhibition may represent a novel approach for treating HCC.

Materials and Methods

Antibodies, equipment, and software. Manufacture/supplier, catalog number, clone and working concentration for all antibodies is provided in Supplementary Table S2. The mouse IgG₂ isotype antibody was generated and validated using previously described methods (27). Specific details regarding equipment and software are provided in Supplementary Tables S3 and S4, respectively.

Study Design. Spontaneous hepatic tumor development was observational; therefore, group sizes were dependent on the number of animals reaching advanced age (≥ 15 months). Ethical and humane endpoints were rigorously followed. All other group sizes were determined based on previous experience (minimal phenotypic variation) and the minimum number of animals required to provide reliable statistical results. Endpoints, when applicable, were determined in advance and were based on age or a time (hours following administration of recombinant protein or imaging reagent). No data have been excluded. A minimum of three independent experiments with successful replication were performed for all studies when possible (except in cases of limited resources). All reported microscopy studies were successfully replicated in at least five independent experiments with varying conditions and equipment. Additionally, microscopy data were collected, background adjusted, converted to binary format, and evaluated in a blinded manner. All data are reported as biological replicates.

Human HCC liver tissues were provided as deidentified, pre-sectioned slides by the University of Colorado Biorepository Core facility and the University of Colorado Cancer Center Tissue Biobanking and Histology Shared Resource. The following BRISQ Tier 1 criteria were provided: Solid tissue was removed from the livers of living patients with confirmed HCC

diagnosis by surgical/clinical procedure. Both clinical and pathology diagnosis of HCC were confirmed for each sample. Resected tissues were stabilized and fixed in 10% neutral buffered formalin prior to long-term preservation in paraffin-embedded blocks. Preserved tissues were shipped and stored at ambient temperature; storage duration ranges from 10-20 years. HCC liver samples provided for this study were selected based on >20% tumor composition by the University of Colorado Biorepository Core.

Mice. Male and female C57Bl/6J wild type mice were purchased from the Jackson Laboratory. The $fH^{-/-}$ mice were created and initially provided by MCP (26). Generation of $fB^{-/-}$ mice has been previously described (67). Generation of $fH^{-/-}fB^{-/-}$ mice was achieved through interbreeding of $fH^{-/-}$ and $fB^{-/-}$ mice. All mice were bred for at least 7 generations prior to experimentation.

Liver Chemistries. Fibrin-free serum samples were analyzed using the Heska DriChem 7000 analyzer.

Light Microscopy and Evaluation of histopathological injury. Portions of the left lateral liver lobes were collected from each animal at the time of euthanasia and fixed in 10% neutral buffered formalin at 4°C. Fixed tissues were transferred to 70% ethanol and embedded in paraffin. Slides with 3 µm-thick FFPE sections were processed and stained with hematoxylin and eosin. Stained tissues were imaged at original magnifications of x100 or x200 and evaluated in a blinded fashion in groups of age- and gender-matched mice.

Oil Red O staining. Slides with 8 μm -thick frozen sections were warmed to RT and fixed in cold 10% neutral buffered formalin for 30 minutes at 4°C. Fixed sections were washed in deionized water five times and rinsed with 60% isopropanol prior to staining in freshly prepared oil red o working solution (0.5% stock diluted 3:2 with deionized water) for 15 minutes. Stained tissues were rinsed once more in 60% isopropanol followed by deionized water and mounted with a solution containing 1:1 glycerol and PBS. Slides were imaged in a blinded fashion with a Zeiss Axio Observer D1 microscope in positive phase contrast at x100 original magnification. Images were quantified with Image J (National Institutes of Health) by thresholding (for tissue segmentation) and then measuring the thresholded area. Ten separate fields of view were examined for each animal. Numerical values for percent area deposition obtained in Image J were analyzed with Prism by a two-tailed t-test with Welch's correction.

Immunofluorescent Staining. Six-micrometer-thick sagittal sections from frozen, OCT-embedded tissues were warmed to RT, fixed in absolute acetone, and washed twice with PBS. Non-specific binding and endogenous Fc receptors were blocked with 1% BSA and 5% heat-inactivated goat/rat/rabbit serum diluted in PBS for one hour at RT. Antibodies were diluted in PBS containing 2% heat-inactivated fetal bovine serum (HI-FBS) and 1% BSA, applied to blocked tissue sections, and incubated overnight at 4°C in a humidified chamber. Nuclei were stained with DAPI, washed in cold PBS, and mounted with a 1:1 solution of PBS and glycerol. Kidney autofluorescence was blocked with 0.05% Sudan Black B in 70% ethanol for 15 minutes at RT, washed twice in deionized water (10 minutes each), and mounted as described above. Slides were sealed and imaged in a blinded fashion with either an Olympus FV1000 FCS confocal (at either x100, x200, or x600 original magnification) or a Zeiss Axio Observer D1

epifluorescent microscope (x100, x200, or x400 magnification). A minimum of ten fields of view (FOV) were captured per sample. Images were converted from binary data format with either Olympus FV-10ASW (version 04.02.02.09) or Zeiss Zen Blue (version 2.6) software. Representative isotype images are shown in Supplementary Figure S1B-S1H. Image quantification was performed according to methods described in Oil Red O staining.

PET Imaging. 0.5 mCi of ^{124}I (Nordion) was added to 100 μg of mAb C3d29 (suspended in 200 μL 0.1 M PBS, pH 7.5) in a glass tube coated with 50 μg Iodogen (1,3,4,6-tetrachloro-3 α ,6 α -diphenyl glycoluril) (Pierce) to prevent antibody oxidation. After five minutes, the reaction mixture was loaded onto a P6 size exclusion column (BioRad) preequilibrated with 1% BSA/PBS, from which ^{124}I -C3d29 was eluted. Purity of ^{124}I -C3d29 was assessed by thin layer chromatography using ITLC-SG strips (Varian) and 10% trichloroacetic acid running solvent, and immunoreactivity was verified by extrapolating the binding to recombinant C3d at an infinite excess of antigen.

Male C57BL/6 and *fH*^{-/-} mice (n=4 per group) were injected intravenously with ^{124}I -C3d29 (120 μCi = 20 μg of protein in 200 μL) and underwent whole body PET/CT imaging using a Siemens Inveon microPET/CT scanner at 4, 24, 48, 72, and 96 hours post injection. Upon PET scan completion, each animal was moved automatically into the adjunct microCT scanner and low-resolution CT scans covering the entire mouse body in the axial, coronal, and sagittal directions were obtained. Images were reconstructed and analyzed with AsiProVM software (version 6.7.1.1). Animals were sacrificed after the 96-hour scan, and all major organs were collected. Organs were weighed and counted in a gamma counter along with a known amount of the radioactive injectate to determine the biodistribution of the antibody (28). Tissue

radioactivity is reported as the percentage of injected radioactivity per gram of tissue (%ID/g) corrected for background and decay.

Liver mRNA Analysis. A portion of the right medial lobe was collected, snap frozen in liquid nitrogen and stored at -80°C until RNA extraction. Liver mRNA was purified using the QIAGEN RNeasy Mini Kit (QIAGEN Sciences) according to manufacturer instructions. Purified RNA was qualitatively assessed by RNA TapeStation (Agilent) and then analyzed using the nCounter Mouse Myeloid Innate Immunity V2 panel (NanoString Technologies). Data were collected for 754 genes of interest and 22 housekeeping genes, and evaluated for quality issues, outliers, or confounders using nSolver (version 4.0) software. Genes failing to register counts above background threshold were not included. A final data analysis was performed using nCounter Advanced Analysis (version 2.0.115) and R 3.3.2 software. Briefly, probe counts were background subtracted and then normalized to the best fit housekeeping genes (Supplementary Table S5) using the GeNorm algorithm (68) within the Advanced Analysis software. The resulting data set used for analysis consisted of 414 genes. DE and pathway analyses were determined by specifying *WT* as the baseline covariate. The pathways or gene sets with the greatest differential expression were identified by directed global significance scoring, followed by analysis with composite pathway scoring. The composite pathway scoring algorithm used in nanoString advanced analysis is calculated as the first principal component of the pathway genes' normalized expression and has been described in detail (69).

Multiplex immunofluorescence of hepatic immune populations. Three-micrometer-thick, FFPE liver sections from 3-month old *fH*^{-/-} and *WT* male mice were processed and stained using the

Leica Bond-III Fully Automated IHC Autostainer. Leica ER2-pH 9 antigen retrieval buffer was used for CD3 ϵ , FoxP3, CD4, and CD8 antibodies; Dako-pH 6 was used for the F4/80 antibody; and Leica ER1-pH 6 was used for the B220 (CD45R) antibody. Primary antibodies were visualized using Opal dyes. Stained slides were scanned at low magnification with the Vectra 3.0 Quantitative Pathology Imaging System. PhenoChart (version 1.0.8) was used to review the initial scans and to specify the unbiased selection of 10 – 0.75 x 0.5 mm regions of interest (ROI) per animal by choosing the “auto” select feature. Each selected ROI was scanned again at higher magnification and resulting multispectral images (MSIs) were used for final analysis in the inForm Cell Analysis software (version 2.4.1). Briefly, each MSI was spectrally unmixed, background subtracted, and segmented by tissue, membrane, nuclear markers. Cells were phenotyped with the inForm cellular phenotyping algorithms. Quantification of data generated based on the aforementioned criteria were analyzed (approximately 7,000-7,500 cells per animal). For each animal, total cell count for each antigen was determined by taking the mean of the sum of all MSIs for each specific marker.

Liver Myeloid Cell Isolation. Liver myeloid cell isolation and flow cytometry methods were adapted from published methods (70). Male *fH*^{-/-} (with and without liver tumors), *WT*, and *fH*^{-/-} *fB*^{-/-} mice (3, 4, and/or 22 months of age) were euthanized individually by CO₂ asphyxiation. Circulating blood was removed by cardiac puncture. Liver lobes were excised, briefly washed in sterile PBS and placed on ice in 5 ml cold RPMI cell culture medium. Intact lobes were cut into \approx 1.5 mm pieces and placed in 120 U/ml collagenase 3 (Worthington) diluted in 5-10 ml of ice-cold HBSS. Samples were transferred to a 37°C water bath, placed on a culture shaker and homogenized for 45 minutes (200 rpm) at 37°C (samples were briefly vortexed twice during

digestion). Digestion was stopped with ice-cold 2% (v/v) heat-inactivated fetal bovine serum (HI-FBS) and 5mM EDTA in cold HBSS, briefly vortexed, filtered through a 70 μ m cell strainer, and centrifuged (8 minutes at 450 x g, at 4°C). The cell pellet was subject to erythrocyte lysis, followed by neutralization, filtration, and centrifugation. The cell pellet was resuspended in cold RPMI with 5% (v/v) HI-FBS and fractionated with a Lymphoprep gradient. The interphase was collected and stored in 2% (v/v) HI-FBS and 3 mM EDTA in HBSS at 4°C. The remaining lower fraction and pellet were washed and centrifuged twice, resuspended in RPMI with 5% (v/v) HI-FBS, fractionated with a 33% Percoll (EMD Millipore) gradient followed by centrifugation and 2 additional washes. The pellet was resuspended a final time in RPMI with 5% (v/v) HI-FBS, combined with fraction 1, and cell number and viability were calculated.

Flow Cytometry Analysis of Isolated Hepatic Myeloid Cells. Single cell preparations were adjusted to a concentration of 3×10^6 cells in 150 μ l. Cells were incubated on ice for 20 minutes with anti-mouse CD16/CD32 (1 μ g/ 10^6 cells) to block Fc receptors and for another 20 minutes with the conjugated primary antibody cocktail (each diluted to 0.2 μ g/ 10^6 cells). Cells were washed, centrifuged, and resuspended in 2 ml cold HBSS with 2% (v/v) HI-FBS and 3 mM EDTA. Unstained cells were used to determine cellular autofluorescence. Single-stain cells and positive/negative compensation beads were used to determine true signal, while Fluorescence Minus One (FMO) controls were used to minimize the effects of spectral overlap. Approximately 1-2 million events per sample were collected on the Cytex Aurora Multi-Spectral Flow Cytometer. SpectroFlo (version 2.1.0) was used for event acquisition and spectral unmixing. FlowJo (version 10.6.0) was used for final gating and determination of population composition. GraphPad Prism was used for statistical analysis.

Generation of recombinant murine factor H and reconstitution of mice. rmCFH was generated using the codon-optimized (*Homo sapiens*) DNA sequence for *CFH* (comprising residues 19-1234; Uniprot identifier: P06909) and has been previously described in detail (71). The resulting expression construct was stably transfected into adherent CHO cells and grown to confluency under humidified conditions at 37°C at 5% CO₂. Exhausted medium from confluent cells was collected, filtered through a 0.22 µm polyethersulfone membrane, and equilibrated for purification with the addition of stock purification buffer to a final concentration of 0.02 M sodium phosphate, 0.5 M sodium chloride, 0.02 M imidazole, and applied to a HisTrap Excel nickel sepharose column (GE Healthcare Bioscience) using a peristaltic pump. Column-bound protein was washed and eluted from the column using a 0.02 M sodium phosphate/0.5 M sodium chloride/0.2 M imidazole buffer. All purification buffers were pH-adjusted to 7.8 ± 0.1 and purification steps were done at 4° C. Eluted protein was concentrated at 4°C using 30K centrifuge filters and then buffer exchanged into Dulbecco's PBS, pH 7.8. Purified rmCFH was sterile-filtered and conjugated with Alexa-fluor 647 using the SAIWI protein/antibody labeling kit (Invitrogen) under sterile conditions. The final concentration of conjugated rmCFH was determined by Nanodrop and checked for contamination by western blot analysis. Functionality was confirmed with a guinea pig erythrocyte hemolysis assay (42).

Male mice (*fH*^{-/-} [7 months] and *WT* [9 months]; n=5 per group) selected for rmCFH reconstitution were similar in size (35 ± 1.5 g) and healthy in appearance. Mice were injected with 400 µg of Alexa 647-conjugated rmCFH by tail vein injection and observed for 20 minutes for adverse reactions. All animals were euthanized 24 hours later, and their organs were harvested for analysis. Liver sections were stained as previously discussed and imaged by

confocal microscopy. Quantification of C3b and iC3b/C3d deposition in unmanipulated and rmCFH-reconstituted *fH*^{-/-} mice was done in Image J (NIH) by thresholding (for tissue segmentation) and then measuring the thresholded area. Five separate fields of view at x600 magnification were thresholded for each animal. Numerical values for percent area deposition obtained in Image J were analyzed with Prism by a two-tailed t-test with Welch's correction.

Immunofluorescent staining of human HCC tissues. Three-micrometer-thick, de-identified FFPE sections from 6 HCC patients were subject to standard deparaffinization and rehydration methods (xylene followed by ethanol gradient). Following 2 washes in deionized water, slides were incubated in antigen retrieval citrate buffer, pH 6 at 96°C for 30 minutes, allowed to cool to RT for 20 minutes, and washed with running deionized water. Tissues were blocked with 5% HI-FBS and 1% BSA in PBS for one hour at RT and washed once with cold PBS. Primary antibodies were diluted in 2% HI-FBS and 1% BSA in PBS, applied to blocked tissues and left to incubate overnight at 4°C in a humidified chamber. Tissues were washed thrice with cold PBS prior to incubation with secondary antibody for one hour at RT, followed by a short incubation with DAPI diluted in PBS and two PBS washes. Slides were dried, mounted with 1:1 PBS and glycerol, sealed, and viewed with an Olympus FV1000 FCS confocal microscope. Images were collected at x100, x200, and x600 original magnification. Isotype images are shown (Supplementary Figure S1B-S1D).

Human cBioPortal Query. TCGA data were accessed at cBioPortal (46, 47). *CFH* mutations, copy-number alterations, and mRNA expression in patients with hepatocellular carcinoma were

queried in TCGA Firehose Legacy study. Overall survival and disease/progression-free survival curves were analyzed in Prism using the logrank (Mantel-Cox) test.

Statistical Analysis. No statistical methods were used to predetermine sample size and independent biological replicates were used for all quantitative analyses. GraphPad Prism was used for analysis, unless otherwise indicated. Quantitative data are represented as dot plots, where the center line indicates the mean and the error bars indicate the standard error of the mean (mean \pm SEM). All data points are shown. *P* values ≤ 0.05 were considered statistically significant. Equal variances were not assumed. Comparisons between two groups were analyzed by unpaired, two-tailed t-test (95% confidence) with Welch's correction and null hypothesis testing (Welch's corrected *t* and *df* [degrees of freedom] values are provided for *P* values reaching significance). Analysis of PET imaging (Figures 2H-I) was done by ordinary 2-way ANOVA ($\alpha=0.05$). Comparisons between three groups were analyzed by one-way ANOVA with post hoc Tukey's t-test. The GeNorm normalization (68) and composite pathway scoring (69) algorithms used in the NanoSting mRNA analysis are described in detail. See Supplementary Table S6 for a complete listing of experimental group sizes (delineated according to figure) categorized by gender, age, strain, and tumor status.

Study Approval. Animals were maintained under standard housing conditions and all procedures were performed in an AAALAC-accredited facility in accordance with the National Institutes of Health *Guide for the Use and Care of Laboratory Animals*. Housing conditions include a 12/12 h dark-light cycle, temperature and humidity control, and *ad libitum* access to standard rodent

chow. All experiments and protocols were approved by the University of Colorado IACUC, and guidelines establishing humane endpoints were strictly followed.

Clinical remnant archival human specimens were provided as deidentified sectioned slides in compliance with the policies and approval of the Colorado Multiple Institutional Review Board (COMIRB) under IRB-approved protocol #15-1461. Studies are conducted in accordance with the Declaration of Helsinki, the Belmont Report, and US Common Rule. The University of Colorado Biorepository Core Facility provided samples from patients with confirmed HCC status (both clinical and pathology diagnoses) and specimen tumor content \geq 20%.

Author Contributions: J.L. designed project; developed and performed experiments (except for PET imaging); collected and analyzed data; adapted flow cytometry experiment and analyzed data; prepared figures; wrote and revised manuscript; bred, maintained, and harvested mice. B.R. performed liver myeloid cell isolation and myeloid cell flow cytometry and critical review of manuscript. M.C.P. generated *fH^{-/-}* mice, provided technical advice and contributed to study design. N.J.S and P.M.S.-J. performed PET imaging and associated analyses. E.T.C. and R.A.N provided conceptual and technical guidance and critical review of manuscript. J.M.T contributed to experimental design and wrote manuscript.

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Figure Legends

Figure 1. $fH^{-/-}$ mice spontaneously develop hepatic tumors. (A-C) Necropsy images of large focal tumors (A and B, arrowheads) or multi-lobular tumors (B, arrows) in $fH^{-/-}$ livers. *WT* livers (C) are generally normal in appearance. (D) Of mice aged to ≥ 15 months, 54% of $fH^{-/-}$ males had visible liver tumors compared to only 10% of $fH^{-/-}fB^{-/-}$ and 7% of *WT* males ($*P = 0.0271$ and $****P < 0.0001$, respectively; 2-sided Fisher's exact test; $fH^{-/-}$ n=35, *WT* n=29, and $fH^{-/-}fB^{-/-}$ n=10). Tumor incidence in ≥ 15 -month-old $fH^{-/-}$ females was 15% compared to 0% in $fH^{-/-}fB^{-/-}$ and 3% in *WT* ($P = 0.5586$ and $P = 0.1777$, respectively; 2-sided Fisher's exact test; $fH^{-/-}$ n=26, *WT* n=29, and $fH^{-/-}fB^{-/-}$ n=10). (E-G) Hepatocyte ballooning (E) and tumor nests in 18-month-old $fH^{-/-}$ livers (F), and generally normal tissue in *WT* livers (G). Scale=20 μm (E-F) and 100 μm (G); n=20 males (≥ 15 months) per group. (H-J) Membranous (H) and cytoplasmic (I) GPC3 deposition in $fH^{-/-}$ livers; punctate GPC3 in *WT* livers (J). Nuclei stained with DAPI (blue). Scale=200 μm (H and J) and 100 μm (I); n=20 males (≥ 15 months) per group.

Figure 2. C3 fragments preferentially deposit in the livers and kidneys of $fH^{-/-}$ mice. (A-C) Abundant C3b and iC3b/C3d deposition in $fH^{-/-}$ livers (B), but little to no deposition in *WT* (A) or $fH^{-/-}fB^{-/-}$ (C) livers. Scale=50 μm ; n=10 males (3-5 months) per group. (D-F) Kidney C3b and iC3b/C3d staining in each strain (glomeruli indicated by arrows). Scale=100 μm ; n=5 males (3-5 months old) per group. (G) Quantification of hepatic C3b and iC3b/C3d deposition (C3b: $****P < 0.0001$ $F_{2,27}=461.1$; iC3b/C3d: $****P < 0.0001$ $F_{2,27}=584.7$; one-way ANOVA with post hoc Tukey's t-test; mean \pm SEM; n=10 per group). (H-I) PET/CT showing focal ^{124}I -C3d29 signal in the left kidney and the liver of the $fH^{-/-}$ mouse (I) and quantification of localized ^{124}I -C3d29 in

organs (**H**); **** $P = 0.0001$, $F_{5,36} = 7.067$; 2-way ANOVA interaction effect; $n=4$ males (3 months old) per group. CV: central vein; TI: tubulointerstitium; L: liver; K: kidney; Bl: bladder.

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Figure 4. Analysis of liver mRNA reveals upregulation of multiple inflammatory signaling pathways in $fH^{-/-}$ mice. (**A**) Volcano plot of $fH^{-/-}$ liver mRNA DE compared to *WT* covariate and (**B**) the fold change of the 10 most DE mRNA. Bonferroni P values are shown below. (**C**) Heatmap of myeloid innate immune pathways (data are displayed on the same scale due to Z-transformation; increasing, neutral or reduced expression, indicated by orange, black, or blue, respectively). (**D-I**) DE of mRNA in $fH^{-/-}$ livers compared to *WT*. Box-and-whisker plots (**D-F**)

showing the composite pathway scores for 3 of the most differentially expressed pathways (**** $P < 0.0001$, $t=11.01$, $df=7.954$ [D]; *** $P = 0.0002$, $t=6.777$, $df=7.816$ [E]; *** $P = 0.0006$, $t=6.033$, $df=6.752$ [F]; unpaired, two-tailed t-test with Welch's correction). For box-and-whisker plots: box = 25th-75th percentile, line = median, and whiskers = maximum and minimum values. (G-I) The most DE genes (Bonferroni-adjusted $P \leq 0.25$) within each of the three corresponding composite pathways (D-F). For all experiments, $n=5$ males (3 months old) per group. DE: differential expression; Sig: signaling.

Figure 5. Inflammation and leukocyte infiltration in $fH^{-/-}$ livers. (A-B) Large pockets of inflammatory infiltrate captured by multi-plex imaging in $fH^{-/-}$ (A, left and B) but not in WT livers (A, right). Nuclei stained with DAPI (blue). Original magnification x200; scale=50 μm (A) and 10 μm (B). (C) Increased populations of each cell type in $fH^{-/-}$ livers compared to WT (F4/80⁺: *** $P = 0.0002$, $t=6.802$, $df=7.559$; CD8⁺: ** $P = 0.0069$, $t=4.949$, $df=4.178$; CD4⁺: $P=0.1228$; CD3⁺: $P=0.1868$; Foxp3⁺/CD4⁺: $P=0.1409$; and B220⁺: $P=0.2894$). For (A-C), $n=5$ males (3 months) per group; two-tailed t-test with Welch's correction; mean \pm SEM. (D) By flow cytometry, young $fH^{-/-}$ males have fewer hepatic neutrophils compared to WT ($P = 0.2067$) and $fH^{-/-}fB^{-/-}$ (** $P = 0.0051$, $t=5.596$, $df=3.963$). Neutrophils trend higher in $fH^{-/-}$ males with liver tumors compared to those without ($P = 0.061$). Data were analyzed by unpaired, two-tailed t-test with Welch's correction; mean \pm SEM; $n=3$ per group ($fH^{-/-}$ and WT , 3 months); $n=4$ per group ($fH^{-/-}$ and $fH^{-/-}fB^{-/-}$, 4 months); $n=4$ and 2 ($fH^{-/-}$ with and without liver tumors, respectively; 22-24 months). (E) Clusters of Ly6G⁺ cells (pink) are observed in livers with multiple tumor foci (arrows). Scale=50 μm ; $n=10$ independent $fH^{-/-}$ liver tumors. Nuclei stained with DAPI (turquoise). All images are representative of 2 independent experiments.

Figure 6. Altered myeloid cell populations in the livers of $fH^{-/-}$ male mice. (A-C) The frequency of monocyte (Ly6C^{high/int/-} MHC-II⁻ F4/80^{low/-}) and macrophage Ly6C^{+/-} MHC-II⁺ F4/80⁺) populations in young $fH^{-/-}$ livers compared to *WT* (A, 3 months) and $fH^{-/-}fB^{-/-}$ (B, 4 months), and livers from aged $fH^{-/-}$ mice with and without visible liver tumors (C, 22-24 months). Data were collected by multi-spectral flow cytometry and analyzed by unpaired, two-tailed t-test with Welch's correction; mean \pm SEM; n=3 males per group ($fH^{-/-}$ and *WT*, 3 months); n=4 males per group ($fH^{-/-}$ and $fH^{-/-}fB^{-/-}$, 4 months); n=4 ($fH^{-/-}$ males with liver tumors) and 2 ($fH^{-/-}$ males without liver tumors). *P* values for each cell population are shown below each graph; (A) Ly6C^{int} MHC-II⁻ F4/80^{low/-} (t=9.896, df=3.809) and Ly6C⁻ MHC-II⁻ F4/80^{low/-} (t=5.121, df=3.474); (B) Ly6C⁻ MHC-II⁻ F4/80^{low/-} (t=3.250, df=5.568). (G) Pseudo color dot plot of representative hepatic Ly6C/MHC-II myeloid populations within CD45⁺ CD11b⁺ Ly6G⁻ cells from a young $fH^{-/-}$ mouse (4 months) compared to an aged $fH^{-/-}$ with liver tumors (22 months). Corresponding cell counts for each gate are shown.

Figure 7. Recombinant factor H binds in the sinusoids of $fH^{-/-}$ mice. (A-B) rmCFH binds within the sinusoids of $fH^{-/-}$ mice in a continuous pattern (A, left), but is reduced and punctate in *WT* (A, right); scale=20 μ m. (B) Quantification of bound rmCFH (*****P* < 0.0001, t=12.98, df=4.544; two-tailed t-test with Welch's correction; mean \pm SEM). For A and B, n=5 males per group (*WT* [9 months], $fH^{-/-}$ [7 months]). (C-E) Enlarged view of $fH^{-/-}$ image identified by dashed white box in (A). Abundant rmCFH deposition (arrowhead) near iC3b/C3d (C, turquoise) and C3b (D, red). Intermittent C3b (C and E; arrows) colocalized with linear iC3b/C3d. Scale=10 μ m. (F) Compared to unmanipulated $fH^{-/-}$, rmCFH-reconstituted $fH^{-/-}$ mice have reduced

sinusoidal C3b (**** $P < 0.0001$, $t=8.424$, $df=7.179$) and increased iC3b/C3d, demonstrating rmCFH-mediated conversion of C3b to iC3b/C3d (** $P = 0.0015$, $t=4.729$, $df=7.931$; two-tailed t-test with Welch's correction; mean \pm SEM; $n=5$ males [7 month] per group). (G-H) rmCFH colocalized with COL4 (G) and MECA-32 (H, arrows) in $fH^{-/-}$ sinusoids, but not on the large vasculature (vascular lumen denoted by asterisks). Nuclei stained with DAPI (blue). Scale=10 μm ; $n=5$. Representative images shown from 4 independent experiments.

Figure 8. Complement deposition in $fH^{-/-}$ liver tumors. (A) C3b-iC3b/C3d deposition in *WT* liver tumors (A, bottom) is sparse and focal. Generally dense C3b-iC3b/C3d deposition is seen throughout $fH^{-/-}$ tumors and the surrounding parenchyma (A, top), with distinct regions lacking C3b (white arrow). Scale=100 μm (A, *WT*), 20 μm (A, *WT* inset), and 50 μm (A, $fH^{-/-}$); $n=10$ ($fH^{-/-}$) and 2 (*WT*) males with liver tumors (≥ 15 -months). Images shown are from 20-month-old mice with visible liver tumors. (B-C) Variable patterns of C3b-iC3b/C3d and GPC3 within and surrounding individual $fH^{-/-}$ tumors. Sinusoidal C3b and iC3b/C3d deposition surrounding $fH^{-/-}$ tumors with membranous GPC3 deposition (B), and dense C3b within the tumor (B, arrowhead). Nearly absent C3b deposition (C, top) and granular iC3b/C3d deposition (C, bottom) in a $fH^{-/-}$ tumor with cytoplasmic GPC3 deposition. Scale=50 μm ; $n=10$ $fH^{-/-}$ males with liver tumors (≥ 15 -months). (D-E) CD46 expression with C3b (D) and iC3b/C3d (E) deposition in tumor-bearing $fH^{-/-}$ livers. C3b is absent from (D, white box), and iC3b/C3d (E, arrow) is reduced in regions with dense CD46. Additionally, iC3b/C3d deposition is seen colocalizing with CD46 (E, arrowhead). Scale=50 μm (D) or 20 μm (E); shown $n=5$ tumor-bearing $fH^{-/-}$ livers. Representative images from $n > 5$ (A) and 4 (B-E) independent experiments.

Figure 9. Complement activation and factor H in human HCC. (A) iC3b/C3d and GPC3 deposition in human HCC biopsies. Scale=50 μm (left panels) and 20 μm (right panels); n=6. (B-C) Kaplan-Meier plots and statistical analysis based on the work of the TCGA Firehose Legacy study, showing the effect of increased *CFH* mRNA expression (B) and *CFH* mutations (C) within the tumors of HCC patients, on survival and disease progression. Increased tumor-associated *CFH* mRNA improved both progression-free survival (B, top: $**P = 0.0020$; n=18 relapse out of 45 [high *CFH* mRNA], 161 out of 280 [unaltered]) and overall survival (B, bottom: $*P = 0.0265$; n=11 deceased out of 50 [high *CFH* mRNA] and 121 out of 326 [unaltered]). Tumor-associated *CFH* mutations reduced the number of disease/progression-free months (C, top: $P = 0.0778$; n=4 relapsed/progressed out of 5 [*CFH* mutation] and 175 out of 320 [unaltered]) and overall survival (C, bottom: $*P = 0.0308$; n=4 deceased out of 7 [*CFH* mutation] and 128 out of 369 [unaltered]). Median months survival and progression-free survival shown in parentheses for each group. Data were analyzed by logrank Mantel-Cox test.

Figure 1

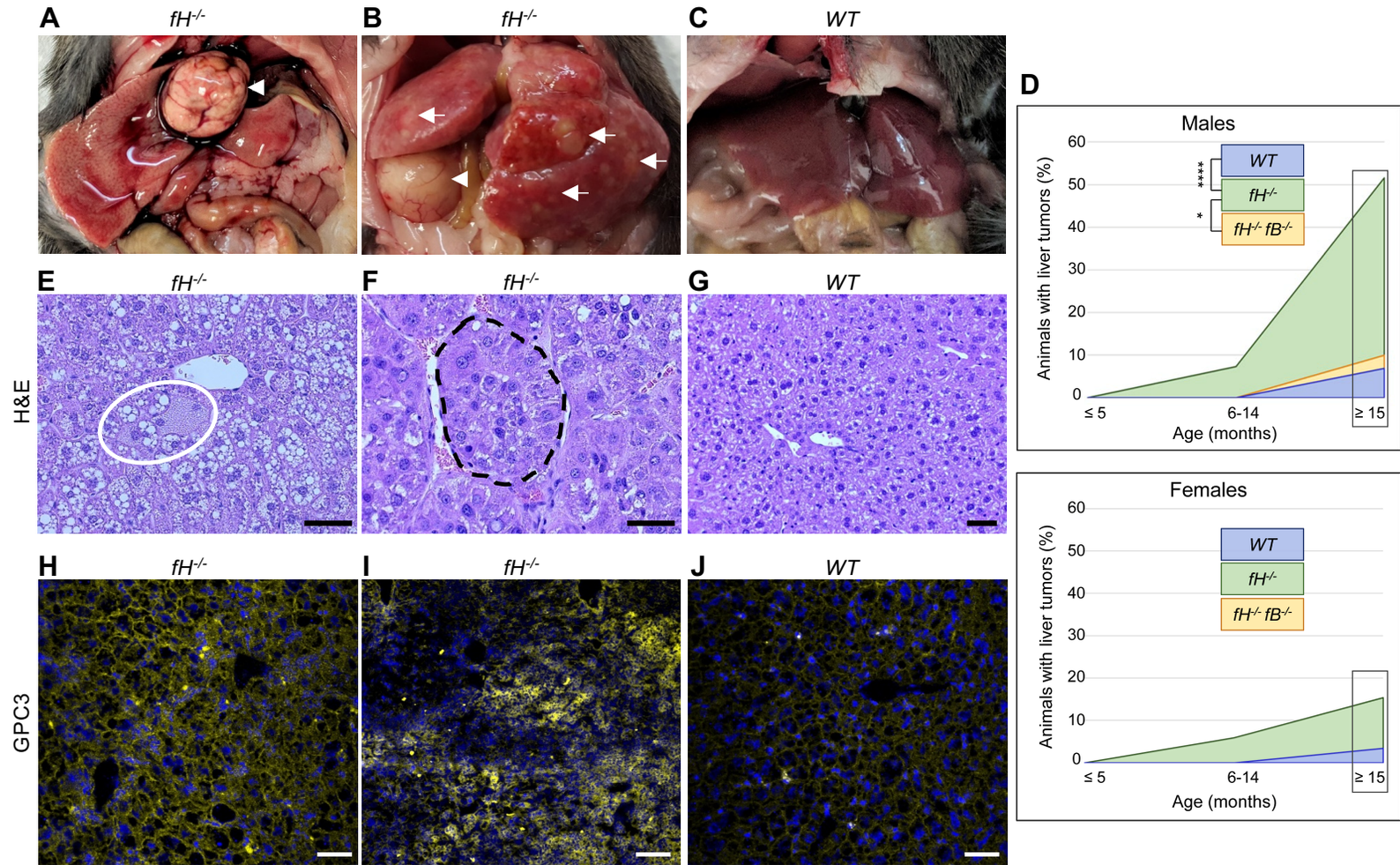


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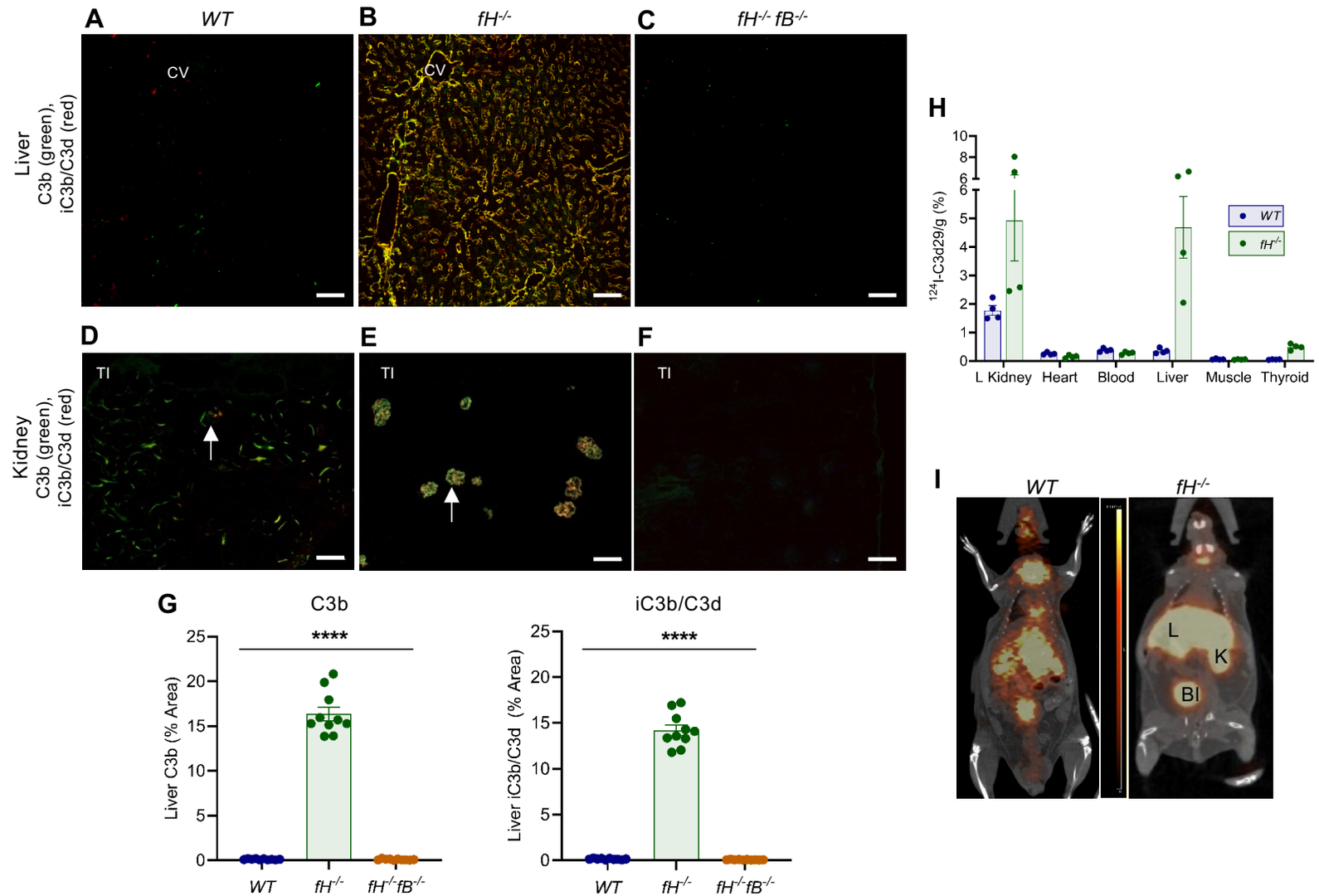


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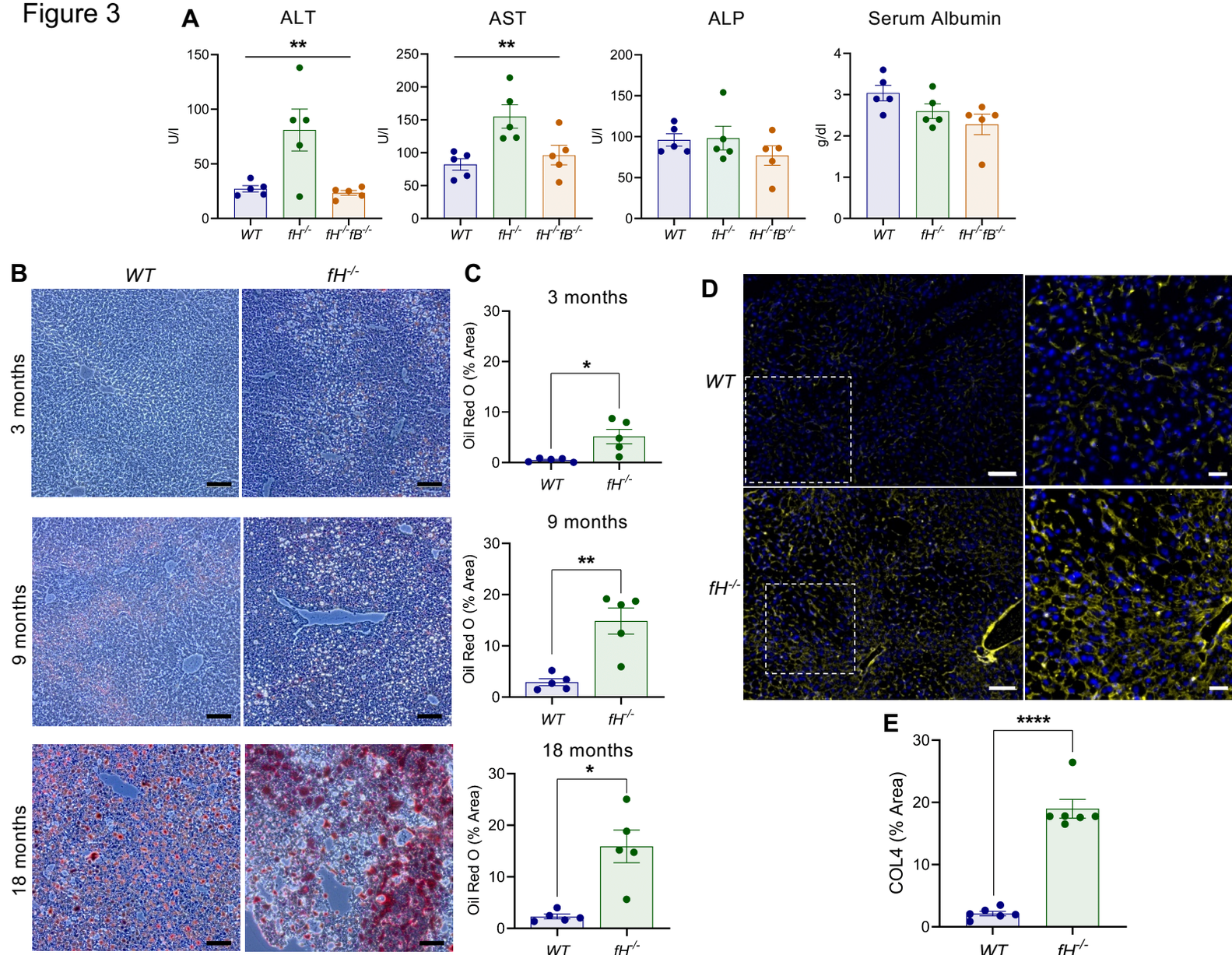


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Figure 4

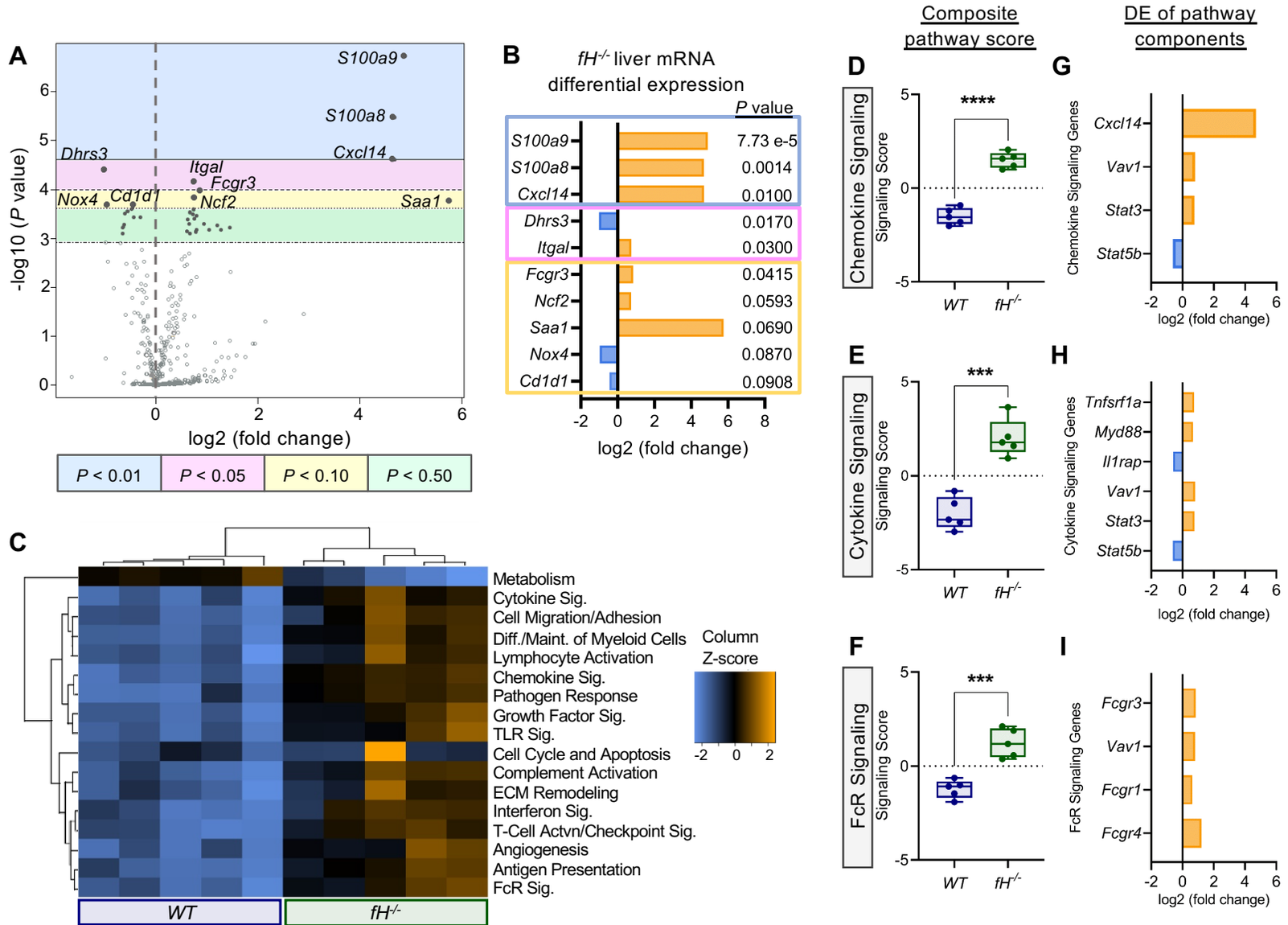


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Figure 5

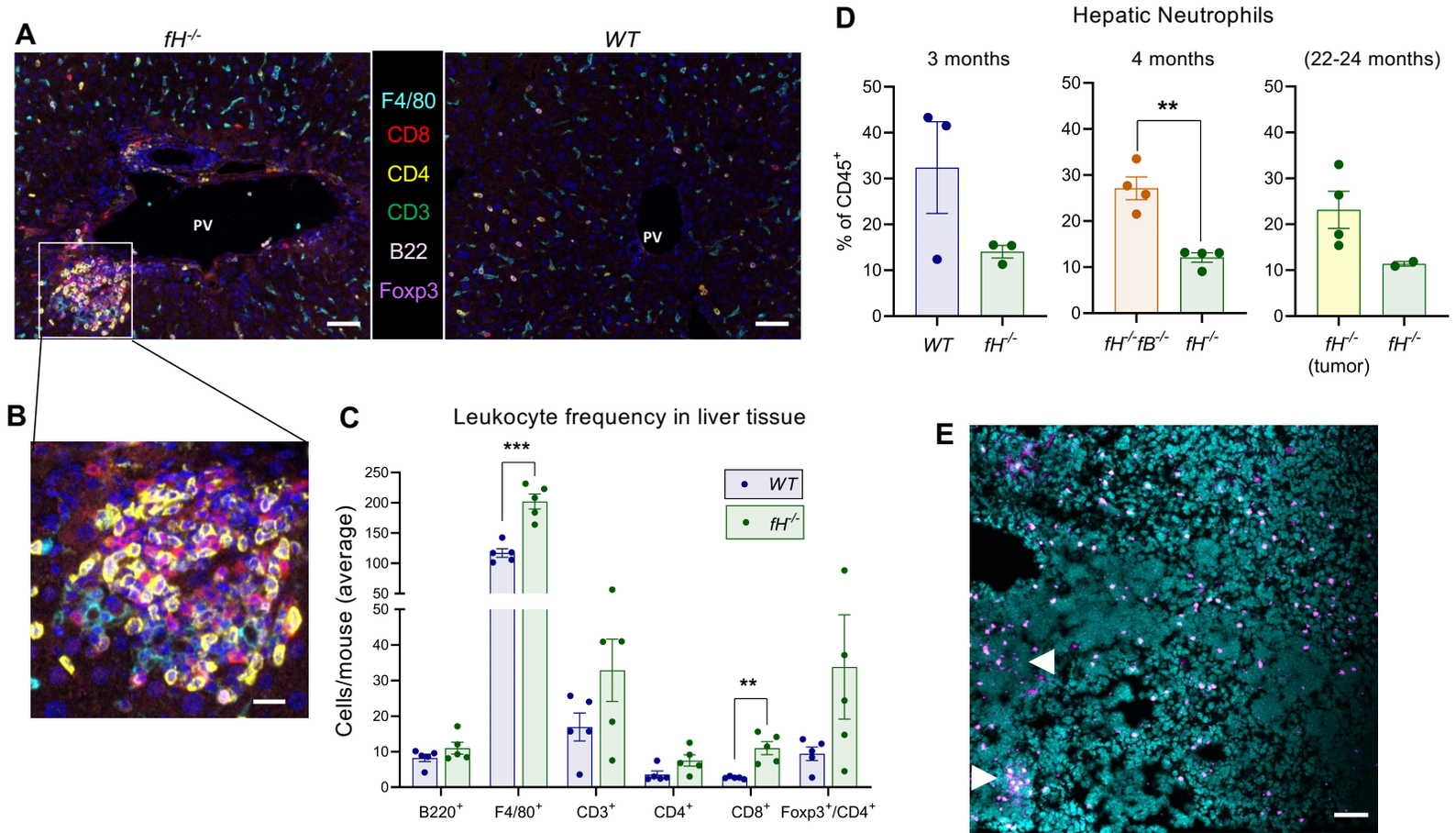


Figure 5. Inflammation and leukocyte infiltration in *fH*^{-/-} livers. (A-B) Large pockets of inflammatory infiltrate captured by multi-plex imaging in *fH*^{-/-} (A, left and B) but not in *WT* livers (A, right). Nuclei stained with DAPI (blue). Original magnification x200; scale=50 μ m (A) and 10 μ m (B). (C) Increased populations of each cell type in *fH*^{-/-} livers compared to *WT* (F4/80⁺: *** P = 0.0002, t =6.802, df =7.559; CD8⁺: ** P = 0.0069, t =4.949, df =4.178; CD4⁺: P =0.1228; CD3⁺: P =0.1868; Foxp3⁺/CD4⁺: P =0.1409; and B220⁺: P =0.2894). For (A-C), n =5 males (3 months) per group; two-tailed t-test with Welch's correction; mean \pm SEM. (D) By flow cytometry, young *fH*^{-/-} males have fewer hepatic neutrophils compared to *WT* (P =0.2067) and *fH*^{-/-}*fB*^{-/-} (** P =0.0051, t =5.596, df =3.963). Neutrophils trend higher in *fH*^{-/-} males with liver tumors compared to those without (P =0.061). Data were analyzed by unpaired, two-tailed t-test with Welch's correction; mean \pm SEM; n =3 per group (*fH*^{-/-} and *WT*, 3 months); n =4 per group (*fH*^{-/-} and *fH*^{-/-}*fB*^{-/-}, 4 months); n =4 and 2 (*fH*^{-/-} with and without liver tumors, respectively; 22-24 months). (E) Clusters of Ly6G⁺ cells (pink) are observed in livers with multiple tumor foci (arrows). Scale=50 μ m; n =10 independent *fH*^{-/-} liver tumors. Nuclei stained with DAPI (turquoise). All images are representative of 2 independent experiments.

Figure 6

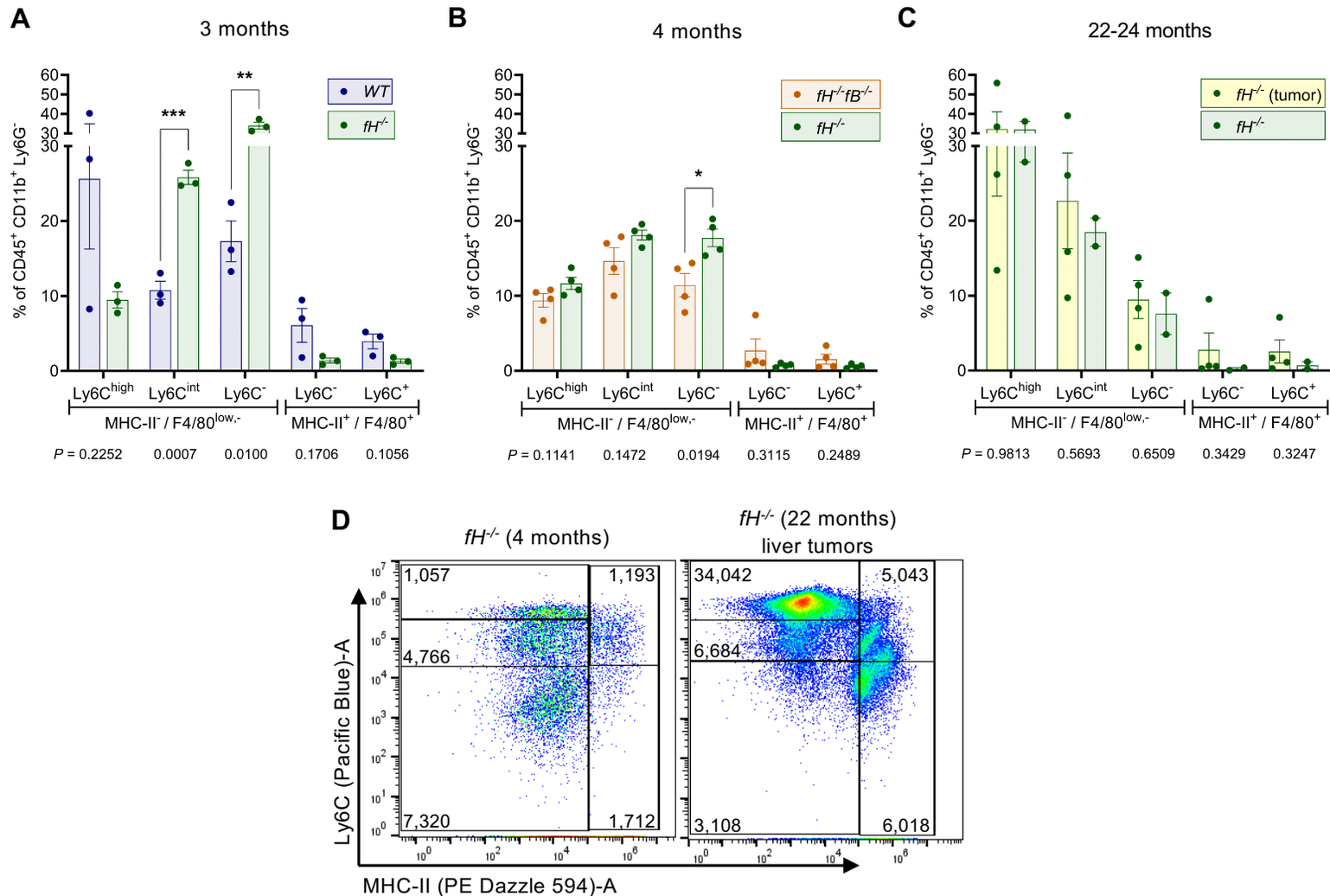


Figure 6. Altered myeloid cell populations in the livers of *fh*^{-/-} male mice. (A-C) The frequency of monocyte (Ly6C^{high/int/-} MHC-II⁻ F4/80^{low/-}) and macrophage Ly6C^{+/-} MHC-II⁺ F4/80⁺) populations in young *fh*^{-/-} livers compared to *WT* (A, 3 months) and *fh*^{-/-}*fB*^{-/-} (B, 4 months), and livers from aged *fh*^{-/-} mice with and without visible liver tumors (C, 22-24 months). Data were collected by multi-spectral flow cytometry and analyzed by unpaired, two-tailed t-test with Welch's correction; mean ± SEM; n=3 males per group (*fh*^{-/-} and *WT*, 3 months); n=4 males per group (*fh*^{-/-} and *fh*^{-/-}*fB*^{-/-}, 4 months); n=4 (*fh*^{-/-} males with liver tumors) and 2 (*fh*^{-/-} males without liver tumors). *P* values for each cell population are shown below each graph; (A) Ly6C^{int} MHC-II⁻ F4/80^{low/-} (t=9.896, df=3.809) and Ly6C⁻ MHC-II⁻ F4/80^{low/-} (t=5.121, df=3.474); (B) Ly6C⁻ MHC-II⁻ F4/80^{low/-} (t=3.250, df=5.568). (G) Pseudo color dot plot of representative hepatic Ly6C/MHC-II myeloid populations within CD45⁺ CD11b⁺ Ly6G⁻ cells from a young *fh*^{-/-} mouse (4 months) compared to an aged *fh*^{-/-} with liver tumors (22 months). Corresponding cell counts for each gate are shown.

Figure 7

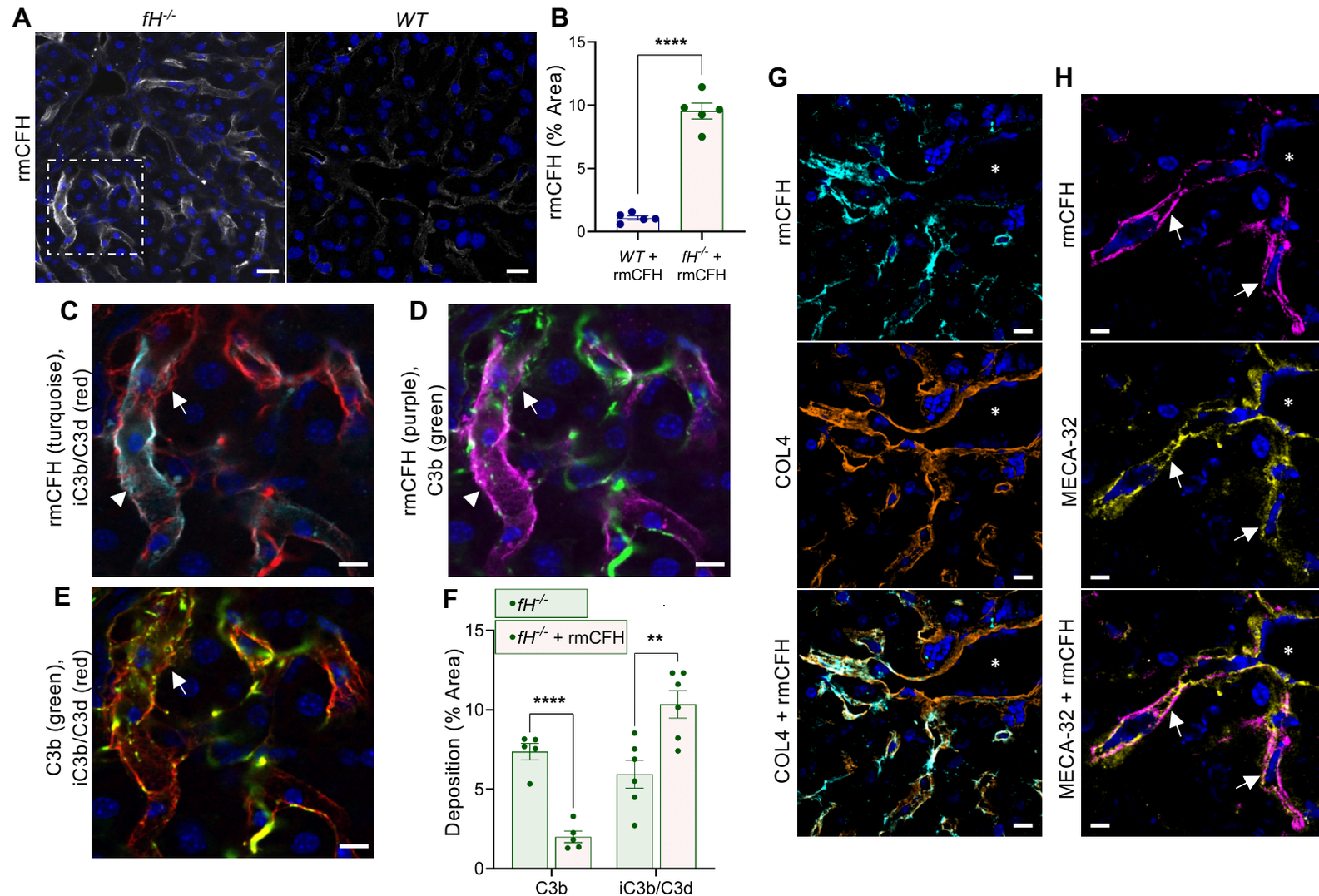


Figure 7. Recombinant factor H binds in the sinusoids of *fh*^{-/-} mice. (A-B) rmCFH binds within the sinusoids of *fh*^{-/-} mice in a continuous pattern (A, left), but is reduced and punctate in *WT* (A, right); scale=20 μ m. (B) Quantification of bound rmCFH (*****P* < 0.0001, *t*=12.98, *df*=4.544; two-tailed t-test with Welch's correction; mean \pm SEM). For A and B, *n*=5 males per group (*WT* [9 months], *fh*^{-/-} [7 months]). (C-E) Enlarged view of *fh*^{-/-} image identified by dashed white box in (A). Abundant rmCFH deposition (arrowhead) near iC3b/C3d (C, turquoise) and C3b (D, red). Intermittent C3b (C and E; arrows) colocalized with linear iC3b/C3d. Scale=10 μ m. (F) Compared to unmanipulated *fh*^{-/-}, rmCFH-reconstituted *fh*^{-/-} mice have reduced sinusoidal C3b (*****P* < 0.0001, *t*=8.424, *df*=7.179) and increased iC3b/C3d, demonstrating rmCFH-mediated conversion of C3b to iC3b/C3d (***P* = 0.0015, *t*=4.729, *df*=7.931; two-tailed t-test with Welch's correction; mean \pm SEM; *n*=5 males [7 months old] per group). (G-H) rmCFH colocalized with COL4 (G) and MECA-32 (H, arrows) in *fh*^{-/-} sinusoids, but not on the large vasculature (vascular lumen denoted by asterisks). Nuclei stained with DAPI (blue). Scale=10 μ m; *n*=5. Representative images shown from 4 independent experiments.

Figure 8

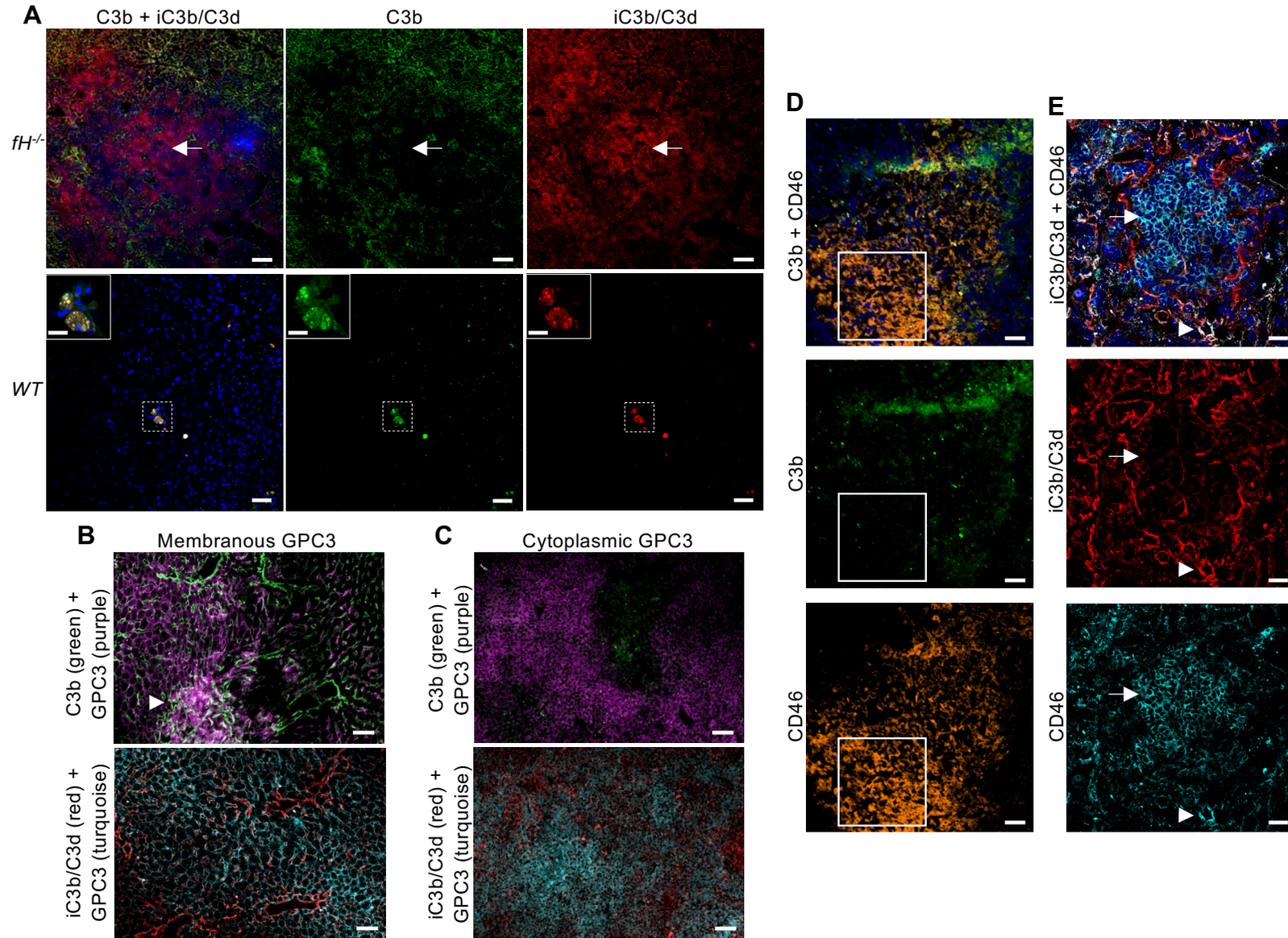


Figure 8. Complement deposition in *fh*^{-/-} liver tumors. (A) C3b-iC3b/C3d deposition in *WT* liver tumors (A, bottom) is sparse and focal. Generally dense C3b-iC3b/C3d deposition is seen throughout *fh*^{-/-} tumors and the surrounding parenchyma (A, top), with distinct regions lacking C3b (white arrow). Scale=100 μm (A, *WT*), 20 μm (A, *WT* inset), and 50 μm (A, *fh*^{-/-}); n=10 (*fh*^{-/-}) and 2 (*WT*) males with liver tumors (≥15-months). Images shown are from 20-month-old mice with visible liver tumors. (B-C) Variable patterns of C3b-iC3b/C3d and GPC3 within and surrounding individual *fh*^{-/-} tumors. Sinusoidal C3b and iC3b/C3d deposition surrounding *fh*^{-/-} tumors with membranous GPC3 deposition (B), and dense C3b within the tumor (B, arrowhead). Nearly absent C3b deposition (C, top) and granular iC3b/C3d deposition (C, bottom) in a *fh*^{-/-} tumor with cytoplasmic GPC3 deposition. Scale=50 μm; n=10 *fh*^{-/-} males with visible liver tumors (≥15-months). (D-E) CD46 expression with C3b (D) and iC3b/C3d (E) deposition in tumor-bearing *fh*^{-/-} livers. C3b is absent from (D, white box), and iC3b/C3d (E, arrow) is reduced in regions with dense CD46. Additionally, iC3b/C3d deposition is seen colocalizing with CD46 (E, arrowhead). Scale=50 μm (D) or 20 μm (E); shown n=5 tumor-bearing *fh*^{-/-} livers. Representative images from n > 5 (A) and 4 (B-E) independent experiments.

Figure 9

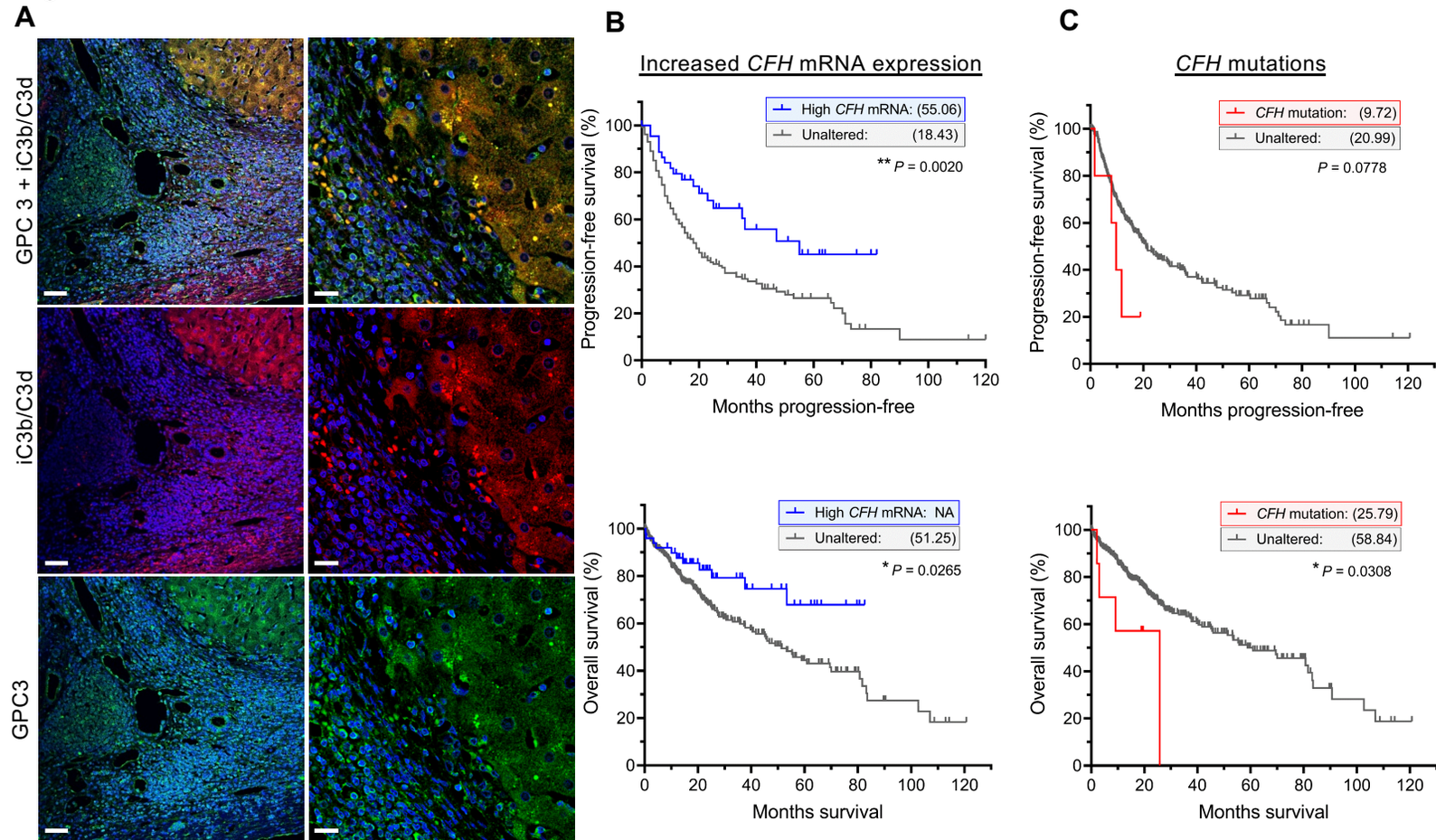


Figure 9. Complement activation and factor H in human HCC. (A) iC3b/C3d and GPC3 deposition in human HCC biopsies. Scale=50 μm (left panels) and 20 μm (right panels); n=6. (B-C) Kaplan-Meier plots and statistical analysis based on the work of the TCGA Firehose Legacy study, showing the effect of increased *CFH* mRNA expression (B) and *CFH* mutations (C) within the tumors of HCC patients, on survival and disease progression. Increased tumor-associated *CFH* mRNA improved both progression-free survival (B, top: ** $P = 0.0020$; n=18 relapse out of 45 [high *CFH* mRNA], 161 out of 280 [unaltered]) and overall survival (B, bottom: * $P = 0.0265$; n=11 deceased out of 50 [high *CFH* mRNA] and 121 out of 326 [unaltered]). Tumor-associated *CFH* mutations reduced the number of disease/progression-free months (C, top: $P = 0.0778$; n=4 relapsed/progressed out of 5 [*CFH* mutation] and 175 out of 320 [unaltered]) and overall survival (C, bottom: * $P = 0.0308$; n=4 deceased out of 7 [*CFH* mutation] and 128 out of 369 [unaltered]). Median months survival and progression-free survival shown in parentheses for each group. Data were analyzed by logrank Mantel-Cox test.