New insights into cholangiocarcinoma: multiple stems and related cell lineages of origin

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Abstract
Cholangiocarcinoma (CCA) is a heterogeneous group of malignancies that may develop at any level of the biliary tree. CCA is currently classified into intrahepatic (iCCA), perihilar (pCCA) and distal (dCCA) on the basis of its anatomical location. Notably, although these three CCA subtypes have common features, they also have important inter- and intra-tumor differences that can affect their pathogenesis and outcome. A unique feature of CCA is that it manifests in the hepatic parenchyma or large intrahepatic and extrahepatic bile ducts, furnished by two distinct stem cell niches: the canals of Hering and the peribiliary glands, respectively. The complexity of CCA pathogenesis highlights the need for a multidisciplinary, translational, and systemic approach to this malignancy. This review focuses on advances in the knowledge of CCA histomorphology, risk factors, molecular pathogenesis, and subsets of CCA.

Keywords
Cholangiocarcinoma, classifications, inflammation, cells of origin, stem cells, molecular profiling

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Introduction
CCA is a heterogeneous group of malignancies emerging at any level of the biliary tree [1-3] (Fig. 1). CCA is classified into intrahepatic (iCCA), perihilar (pCCA) and distal (dCCA) based on its anatomical location [1-3]. Notably, although these three CCA subtypes have common features, they also have important inter- and intra-tumor differences that can affect the pathogenesis and outcome [4-9]. The complexity of the pathogenesis and the pronounced heterogeneity of CCAs have impeded clinical goals in their management [10].

This review focuses on advances in the knowledge of CCA classifications and histomorphology, risk factors, and molecular pathogenesis, as well as the multiple subtypes of CCA. The existence of multiple cells of origin is presented and discussed as a possible major determinant of the intertumoral heterogeneity of CCA, suggesting the possibility of a CCA classification based on cells of origin. Finally, a model of interaction between the cell of origin, the molecular alterations, and the underlying liver and biliary pathologies in different CCA subsets is proposed.

New insights into CCA classifications

In the last few years, a huge number of different classifications have been proposed [1-11]. Currently, CCA is classified as iCCA, pCCA or dCCA [1-3]. However, this classification is biased by some pitfalls. Firstly, CCA is frequently diagnosed at an advanced stage; this is especially true for pCCA, where distinguishing between an intra-hepatic or extra-hepatic location proves challenging [11]. Secondly, small bile ducts and ductules are also present in the perihilar liver parenchyma. Therefore, pCCA may originate either from these smaller ducts or from hilar ducts, though this cannot be discriminated based on gross morphology. Likewise, iCCA may originate from large or small ducts of the intrahepatic biliary tree. Thirdly, recent studies demonstrated that, from a pathological and molecular point of view, there are no differences between pCCA and iCCA that originate from large bile ducts; therefore, the distinction between these two forms of CCA is losing relevance [4,9,12-14]. Taking into consideration the macroscopic pattern of growth, iCCA has been classified as mass-forming (MF), periductal...
infiltrating (PI), and intraductal growing (IG) [2,15]. As far as pCCA and dCCA are concerned, both PI and IG patterns have been recognized. For pCCA a nodular+PI growth pattern predominates (>80%) [2,5,16-18].

Histologically, the vast majority of pCCAs and dCCAs are mucinous adenocarcinomas. In contrast, iCCAs are highly heterogeneous tumors and several classifications have been proposed [4,5,9,19]. The small bile duct type (mixed) iCCAs display an almost exclusively MF growth pattern [4,5,9,19], and are frequently associated with chronic liver diseases (viral hepatitis or cirrhosis) [4,5,9,19,20]. Notably, this subtype shares clinicopathological similarities with cytokeratin (CK) 19-positive hepatocarcinoma (HCC) [4,21]. On the other hand, large bile duct type (mucinous) iCCAs may grossly appear as MF, PI or IG types; they are more frequently associated with primary sclerosing cholangitis (PSC) and can be preceded by preneoplastic lesions, such as biliary intraepithelial neoplasm or intraductal papillary neoplasm [4,5,9,19]. Interestingly, the large bile duct type (mucinous) iCCAs share phenotypic traits with pCCA and pancreatic cancers [4]. In our opinion, this histological subtyping should be taken into serious consideration because it results in different risk factors, molecular profiles, and clinical management [3,4,9,14,22-28,52].

Multiple risk factors reveal CCA subtype-specific pathogenesis

The epidemiological profile of CCA and its subtypes (Fig. 1) displays enormous geographic differences. Although in most countries CCA is a rare cancer (incidence <6/100,000), its incidence may reach extremely high peaks in some populations of Chile, Bolivia, South Korea and North Thailand [29]. In particular, the variation in incidence rates is correlated with the different prevalence of risk factors. In regions of Thailand, for example, it is closely related to the incidence of liver flukes [30-32]. In the past century, a progressive increase in iCCA incidence was registered, whereas the incidences of both pCCA and dCCA seem to be decreasing [29]. Fig. 1 summarizes the worldwide epidemiology and incidence trends of intrahepatic and extrahepatic CCA.

In order to review the literature on risk factors associated with iCCA and/or p/dCCA, we searched for case series of iCCA

![Worldwide Incidence of CCA](image-url)

**Figure 1** Worldwide incidence (cases/100,000) of cholangiocarcinoma (CCA).
Data refer to the period 1971-2009. Green color identifies areas with lower incidence (<6/100,000 cases, rare cancer), while pink color indicates countries where CCA is not a rare cancer (>6/100,000 cases). Diagnoses have been classified according to the International Classification of Diseases (ICD-O-1, ICD-O-2, ICD-O-3, ICD-10, ICD-V9, ICD-V10, ICD-O). Where available, the more incident form (intrahepatic [IH] vs. extrahepatic [EH] CCA) and the temporal trend of incidence (↑increasing trend; §stable trend; ↓decreasing trend) have been reported
Modifed from Banales et al. Nat Rev Gastroenterol Hepatol [ref. 3]
New insights into cholangiocarcinoma origin

and p/dCCA diagnosed according to the currently recognized criteria (i.e. European RARECARE) [33], or case series with appropriate topographic classification of histologically verified CCA. The results of these studies, mostly case-control studies, are summarized in Table 1 [34-46].

Table 1 Summary of risk factors significantly associated with iCCA* and/or pCCA/dCCA# as assessed by case-control studies (Odds ratios by multivariate analyses; all odds ratios reported are significant for the 95% CIs)

<table>
<thead>
<tr>
<th>Risk factors for iCCA [references]</th>
<th>Odds ratios for increased risk</th>
<th>Risk factors for pCCA/dCCA [references]</th>
<th>Odds ratios for increased risk</th>
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<tbody>
<tr>
<td><strong>Bile duct diseases and conditions</strong></td>
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<tr>
<td>Cholecystitis [36]</td>
<td>8.5</td>
<td>Cholecystitis [36]</td>
<td>5.9</td>
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<tr>
<td>Hepatolithiasis [77,37,39,40,43]</td>
<td>50.0-4.8; 6.79§</td>
<td>Hepatolithiasis [45]</td>
<td>3.09</td>
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<tr>
<td>Choledochal cysts [36,37,44,59]</td>
<td>10.7-43.03</td>
<td>Choledochal cysts [36]</td>
<td>47.1</td>
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<tr>
<td>Choledocholithiasis [35,43,161]</td>
<td>4.17-33.35</td>
<td>Choledocholithiasis [36,45]</td>
<td>34, 9.84</td>
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<tr>
<td>Cholangitis/PSC [36,44]</td>
<td>64.2-75.23</td>
<td>Cholangitis/PSC [36]</td>
<td>45.7</td>
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<tr>
<td>Biliary cirrhosis/PBC [36,44]</td>
<td>17.08-19.8</td>
<td>Biliary cirrhosis/PBC [36]</td>
<td>11.8</td>
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<tr>
<td>Choledococtectomy [36,39]</td>
<td>3.6-5.4</td>
<td>Choledococtectomy [35,36,45]</td>
<td>5.8-12</td>
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<td><strong>Digestive diseases</strong></td>
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<tr>
<td>Inflammatory bowel diseases [36,58]</td>
<td>1.72-3.95</td>
<td>Inflammatory bowel diseases [36,58]</td>
<td>1.1-1.97</td>
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<tr>
<td>Crohn's disease [36,44]</td>
<td>1.68-2.4</td>
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<td>2.8</td>
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<tr>
<td>Ulcerative colitis [36,44]</td>
<td>3.3-4.5</td>
<td>Ulcerative colitis [36]</td>
<td>2.8</td>
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<tr>
<td>Duodenal ulcer [36]</td>
<td>3.4</td>
<td>Duodenal ulcer [36]</td>
<td>1.9</td>
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<tr>
<td>Chronic pancreatitis [36]</td>
<td>5.9</td>
<td>Chronic pancreatitis [36]</td>
<td>9.3</td>
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<tr>
<td><strong>Liver flukes</strong></td>
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<tr>
<td>Clonorchis sinensis infection [38,42]</td>
<td>8.6-13.6</td>
<td>Clonorchis sinensis infection [42]</td>
<td>6.5</td>
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<tr>
<td><strong>Endocrine disorders</strong></td>
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<tr>
<td>Thyrotoxicosis [36]</td>
<td>1.5</td>
<td>Thyrotoxicosis [36]</td>
<td>2.4</td>
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<tr>
<td>Diabetes mellitus type II [37-39,43,75,86]</td>
<td>1.8-3.2</td>
<td>Diabetes mellitus type II [35,36,45,86]</td>
<td>1.5-3.2</td>
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<tr>
<td><strong>Metabolic conditions and general risks</strong></td>
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<td></td>
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<tr>
<td>Obesity [36,44]</td>
<td>1.7-1.71</td>
<td>Alcohol intake &gt;80 g/day [39,45]</td>
<td>1.05-3.6</td>
</tr>
<tr>
<td>Alcohol intake &gt;80 g/day [37,39,75]</td>
<td>1.52-5.21</td>
<td>Smoking [36]</td>
<td>1.30</td>
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<tr>
<td>Smoking [36,44,162]</td>
<td>1.3-2.1</td>
<td>Metabolic syndrome [overall @ 44]</td>
<td>1.32-1.83</td>
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<tr>
<td>Metabolic syndrome [overall @ 44]</td>
<td>1.32-1.83</td>
<td>Family history of other cancer [45]</td>
<td>3.15</td>
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<tr>
<td>Dyslipoproteinemia [44]</td>
<td>1.65</td>
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<tr>
<td>Hypertension [44]</td>
<td>1.63</td>
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<tr>
<td><strong>Chronic liver diseases</strong></td>
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<td></td>
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<tr>
<td>Alcoholic liver disease [36,44]</td>
<td>3.1-5.69</td>
<td>Alcoholic liver disease [36]</td>
<td>4.5</td>
</tr>
<tr>
<td>Non specific cirrhosis [36,37,43,44,75]</td>
<td>18.24-28.79</td>
<td>Non specific cirrhosis [36,45]</td>
<td>5.4-6.16</td>
</tr>
<tr>
<td>Hemochromatosis [36]</td>
<td>2.6</td>
<td>Hemochromatosis [36]</td>
<td>2.6</td>
</tr>
<tr>
<td>Non alcoholic liver disease [36]</td>
<td>3</td>
<td>Non alcoholic liver disease [36]</td>
<td>3</td>
</tr>
<tr>
<td>HCV infection [77,36-40,44,75]</td>
<td>2.41-9.71; 9.7§</td>
<td>HCV infection [78]</td>
<td>1-3.05</td>
</tr>
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Table 1 was prepared to summarize findings regarding case-control studies that investigated risk factors associated

Methods for collection of case-control studies investigating risk factors and for the preparation of Table 1

Table 1 was prepared to summarize findings regarding case-control studies that investigated risk factors associated
with iCCA and/or pCCA/dCCA (extrahepatic CCA), assessed by multivariate analyses. The case-control studies were selected from papers returned by the following search terms on PubMed: (“cholangiocarcinoma” [MeSH Terms] OR “cholangiocarcinoma” [All Fields]) AND (“risk factors” [MeSH Terms] OR “risk” [All Fields] AND “factors” [All Fields]) OR “risk factors”[All Fields] OR “risk factor” [All Fields]) NOT (“review”[Publication Type] OR “review literature as topic”[MeSH Terms] OR “review”[All Fields]) AND English [lang]. The selection criteria also required that works include histologically verified case series of iCCA and/or pCCA/dCCA with appropriate topographic classification (Klatskin tumors classified as pCCA and excluded from the iCCAs).

In the 14 case-control studies selected, several putative risk factors were evaluated. The risk factors were collected based on the pathophysiological mechanisms leading to CCA (cells primarily targeted or activated by diseases and therefore involved in the carcinogenic process) and on the homogeneity of risk factors.

The investigated risk factors could be classified on the basis of the tissue or the cell primarily targeted by diseases or conditions and is therefore likely to be involved in the carcinogenic process as cell or tissue of origin. In this view, biliary diseases such as cholangitis/PSC, secondary biliary cirrhosis, cholelithiasis, hepatitis, and liver flukes are pathological conditions that primarily affect large intra- and/or extrahepatic bile ducts (Table 1) (Fig. 1). Appropriately, as shown in Table 1, these pathological conditions are risk factors for both iCCA and p/dCCA. Pancreaticobiliary maljunction, cholelithiasis, and cholecystectomy primarily affect extrahepatic bile ducts. Moreover, these conditions are recognized risk factors essentially for p/dCCA (Table 1) (Fig. 1). Parenchymal liver diseases, including chronic viral and non-viral liver diseases, recognize the interlobular bile ducts, bile ductules, and the canals of Hering as the primary targets; thus, these conditions are specific risk factors for iCCA (Table 1).

Moreover, several toxic and environmental risk factors are known or suspected to be related to CCA development; amongst them are nitrosamine-contaminated food, asbestos, dioxins, vinyl chlorides and, in the past, thorotrast [47]. However, the geographic distribution of the related CCA cases associated with these risk factors, and their combination with other identified factors or with subclinical inflammation of the liver and biliary tree (see obesity, metabolic diseases, steatosis), are largely unknown.

PSC, a disease affecting both intra- and extrahepatic bile ducts, represents the strongest independent risk factor for both iCCA and pCCA as evaluated by multivariate analysis (Table 1). Several other dated studies evaluated the cumulative risk of CCA in PSC patients, but not the discrete risk of iCCA and/or pCCA in PSC [48-51]. The lifetime incidence of CCA among PSC patients ranges from 5-10% [52-55]. Clinicopathological observations suggested that PSC is specifically associated with the development of bile-duct (mucinous) type CCA [4,56]. We have indeed found a PSC-specific pathologic spectrum involving peribiliary glands (PBGs) in large bile ducts that exhibit cell proliferation in PBGs, mucinous metaplasia, epithelial mesenchymal transition (EMT) traits, and dysplasia,
suggesting a carcinogenic progression linked with the manifestation of PSC within the bile ducts [56].

In addition, inflammatory bowel diseases (IBD) associated with or preceding PSC may significantly affect the risk of CCA. In a European multicenter study, the coexistence and duration of IBD significantly increased the risk of CCA in PSC patients [51] and the incidence of CCA was highest within the first year after diagnosis of IBD [51]. According to a recent meta-analysis of IBD patients, the estimated relative risk was 2.61 for iCCA versus 1.47 for pCCA [57]. Both Crohn's disease (CD) and ulcerative colitis (UC) were found to be associated with an increased risk of CCA, whereas CD patients seemed to have a lower risk of CCA than UC patients [57]. Recently, it has been revealed that CD and no IBD (both vs. UC) were associated with a lower risk of malignancy in PSC patients (hazard ratio, 0.77; P=0.004) [58]. In contrast, in a US study neither IBD nor its duration conferred additional risk of CCA in PSC patients [59].

In another study, Welzel et al reported that duodenal ulcer disease was significantly more common among pCCA and iCCA cases than controls [36]. Many studies have demonstrated associations between CCA and *Helicobacter pylori*, but the correlation remains controversial and a direct causal relationship has not been established [60-66].

A positive association between chronic pancreatitis and CCA has also been found. The risk is greater in pCCA than in iCCA (Table 1).

A strong association exists between liver fluke infestation (*Ophisthorchis viverrini* [OV] and *Clonorchis sinensis*) and the development of CCA, in particular in east Asia, where iCCA represents a large proportion of primitive liver cancers (Table 1) [67,68].

Some epidemiological studies have been conducted to evaluate the relationship between type 2 diabetes and CCA (Table 1) [36,69-71]. In this regard, it was demonstrated in a diabetes model and in human subjects affected by type 2 diabetes that PBGs underwent proliferation and expansion in relation to hyperglycemia [72]. Interestingly, metformin significantly reduced the risk of iCCA in diabetic patients by 60% [73,74]. A recent meta-analysis also confirmed that obesity, alcohol use, and smoking have an association with iCCA [75].

In recent years, it is becoming more evident that metabolic conditions, particularly metabolic syndrome, predispose to the development of primary liver cancers [3,44,76]. Nonalcoholic fatty liver disease/nonalcoholic steatohepatitis (NAFLD/NASH) have been found to be independent predictors of iCCA development, though with a lower strength of association than other risk factors (viral hepatitis, cirrhosis), whereas NAFLD/NASH failed to predict pCCA (Table 1). Hemochromatosis has also been identified as an independent predictor of iCCA development, though it failed to predict pCCA (Table 1).

It has long been known that the presence of cirrhosis increases the risk of iCCA [36,37,40,44,75]. Liver diseases related to hepatitis B virus (HBV) and hepatitis C virus (HCV) have been identified as definitive risk factors for CCA, with a stronger association for iCCA than pCCA [77]. A meta-

**Molecular profiling and the identification of multiple CCA subsets**

Enormous geographic and racial differences exist with CCA [3,83]. Generally, the prominent genetic alterations described in CCAs affect Tumor Protein 53 (TP53) (DNA repair) [84-86], tyrosine kinase (KRAS Proto-Oncogene, GTPase [KRAS], B-Raf Proto-Oncogene, Serine/Threonine Kinase [BRAF], Small Mother Against Decapentaplegic 4, and Fibroblast Growth Factor Receptor 2) [8,84,88], and Protein Tyrosine Phosphatase, Non-Receptor Type 3 (PTPN3) [89], deregulated Wnt Family Member/Catenin Beta 1 (WNT/CTNNB1) [90] and Notch pathways, epigenetic (Isocitrate Dehydrogenase (NADP(+)) 1, Cytoisic/IDH1 and Isocitrate Dehydrogenase (NADP(+)) 2, Mitochondrial [IDH2]) [28,84,88,91,92], and chromatin-remodeling factors [85] (AT-Rich Interaction Domain 1A, Polybromo 1, and BRCA1 Associated Protein 1) [84,86,88,91].

Since all known CCA risk factors are associated with chronic bile duct inflammation, it is understandable that molecular studies have also focused on genetic/epigenetic abnormalities involving inflammation-related genes [93-95]. In keeping with the initiation and progression of CCA in the inflammatory milieu, the enzyme cyclooxygenase (COX)-2 is induced in CCA by both bile acids and oxysterols, the oxidation products of cholesterol that are increased in the bile during biliary inflammation [96,97]. Inflammatory cytokines may also induce the expression of inducible nitric oxide (NO) synthase (iNOS) in CCA. NO promotes DNA damage directly and also by inhibiting DNA repair mechanisms, thus promoting carcinogenesis [98,99]. iNOS activation
also stimulates the expression of COX-2 [100]. Interestingly, recurrent genetic variants in the promoter of human telomerase reverse transcriptase [88] have been described in CCA that developed in the chronic inflammation milieu of the hepatitis infection [88]. This molecular alternation could be correlated with the pivotal role of telomerase in controlling stem cells, extremely challenged in conditions of chronic inflammation because the senescence of the mature hepatocytes determines the secondary stem proliferative activation [12].

The comparative evaluation of the gene expression profile (transcriptome), clinicopathological traits, and patient outcomes in iCCA cases has allowed for the identification of two main biological classes of iCCA: 1) the inflammation class (38%), characterized by activation of inflammatory signaling pathways, overexpression of cytokines, and Signal transducer and activator of transcription 3 activation; and 2) the proliferation class (62%), characterized by activation of oncogenic signaling pathways (i.e., RAS, mitogen-activated protein kinase and hepatocyte growth factor [HGF]/MET), DNA amplifications at 11q13.2, deletions at 1q42.1, mutations in KRAS and BRAF and gene expression signatures previously associated with poor outcomes for patients with HCC [7].

Technological advances have also allowed the differential characterization of genomic and genetic features of CCA epithelial and stromal compartments [101]. The tumor epithelium was defined by deregulation of the Human Epidermal Growth Factor Receptor 2 network and frequent overexpression of Epidermal Growth Factor Receptor, HGF/MET, Plastid Ribosomal Protein S6, and Kif67, whereas the stroma was rich in inflammatory cytokines [101].

The molecular mechanism of OV-associated CCA has been also studied in experimental models [32,102]. In humans, molecular studies of iCCA associated with liver flukes demonstrated an overexpression of genes involved in xenobiotic metabolism (UDP Glucuronyltransferase Family 2 Member B11, UDP Glucuronosyltransferase Family 1 Member A10, Carbohydrate Sulfotransferase 4, Sulfotransferase Family 1C Member 2), whereas, in contrast, non-OV-associated iCCA showed enhanced expression of genes related to growth factor signaling (Transforming Growth Factor β1, Placental Growth Factor, Insulin Like Growth Factor-Binding Protein). The draft genome of Clonorchis sinensis and transcriptomes of Clonorchis sinensis and OV have recently been elucidated [103,104]. In a large cohort of iCCA (n=102) associated with liver fluke infection, promoter hypermethylation in a handful of target genes was demonstrated [105]. Thus, the evaluation of the putative signature of liver flukes associated with CCA could help in screening and surveillance, with the perspective of an early diagnosis in subjects carrying the infestation [102].

CCA genetic susceptibility has been investigated in geographic areas where liver flukes are endemic. In these studies, specific haplotypes of COX2-coding gene Prostaglandin-Endoperoxide Synthase 2 (PTGS2) or Interleukin 8 Receptor Beta (IL8RB) have recently been associated with a significant risk of CCA development [106].

Different molecular signatures of the high oncogenic risk were described in PSC patients. KRAS mutations were found in the bile fluid of 30% of PSC patients without evidence of CCA [107]. Since KRAS mutations are frequently observed in CCA, this could be an early event of bile-duct carcinogenesis in PSC patients. Notably, mutational profiling can be performed in cell-free DNA of bile supernatant [108]. The inflammatory microenvironment has also been associated with an aberrant DNA methylation profile in PSC-derived CCA, which provides survival signals for the tumor [109]. Genetic susceptibility of PSC patients to CCA development was demonstrated by studies concerning the natural killer cell G2D receptor, where specific genetic variants have been described in PSC patients [110].

As far as multilevel differences between iCCA and pCCA are concerned, it has been shown that pCCA expresses higher levels of Mucin 5AC, AKT Serin/Theronin Kinase, Keratin 8, and Annexin, but less Vascular Endothelial Growth Factor. At a molecular level, distinct patterns of genetic mutations, methylation, and expression profiling may differentiate iCCA from pCCA. iCCAs were significantly more frequently B-cell Lymphoma 2+ and p16+, whereas pCCAs were more often Tumor Protein p53+ [111]. Miller et al revealed 545 genes with altered expression in p/dCCA and 2354 in iCCA [112]. Mutations in IDH1 and IDH2 were found only in iCCA (n=9), but in none of the examined p/dCCA (n=22) and gallbladder cancer (n=75) cases [113]. Recent papers confirmed that liver-fluke-negative iCCAs are enriched with IDH mutants [14,28]. A cross-platform comparison of CCA with pancreatic cancer and HCC further emphasized the presence of distinct tumor subsets, suggesting similarities between IDH-mutant CCAs and HCCs, rather than pancreatic cancers [28]. Conversely, mutations in KRAS by tumor site demonstrated a predominance of KRAS mutations in pCCAs (53.3% of hilar vs. 6.7% of peripheral type) [7]. As far as epigenetic abnormalities are concerned, the methylation of Ras Association Domain-Containing Protein 1 (RASSF1A) was more common in pCCA than in iCCA, while the opposite was demonstrated for the methylation of the Glutatione S-Transferase P (GSTP) gene [114]. Other reported alterations associated with iCCA included fibroblast growth factor receptor pathways and Ephrin Type-A Receptor 2 (EPHA2) mutations [115].

Finally, the histopathological distinction of cholangiocolocellular differentiation of iCCA has been correlated with its genome-wide molecular features and clinical significance [115]. Based on cholangiocolocellular differentiation status, iCCAs were stratified into iCCA with or without cholangiocolocellular differentiation. iCCA with cholangiocolocellular differentiation resembling an inflammation-related subtype revealed less aggressive histopathological features compared to iCCA without cholangiocolocellular differentiation resembling a proliferation subtype. Accordingly, the former showed more favorable clinical outcomes, including overall survival, than iCCA without cholangiocolocellular differentiation [116]. An updated expert review of a personalized approach based on molecular target therapies in CCA has recently been published [117].
Variable clinical presentations and diagnostic features

The clinical presentation of CCA is largely influenced by anatomic location and pattern of growth, which depend on the cells of origin. Recent studies on the origin of CCA have demonstrated that CCA comprises at least two separate entities with distinct histology, progression, and risk factors. These subtypes have recently been classified into large-bile-duct (mucinous) type CCAs and small-bile-duct or mixed CCAs. According to multiple observations, pCCAs most likely originate from mucin-secreting cholangiocytes and PBGs [118] located in hilar bile ducts, show a large-bile-duct (mucinous) type histology, and are associated with preneoplastic lesions emerging in surface epithelium [2,3] and PBGs [118]. On the other hand, iCCAs show intertumor heterogeneity and are classified into two main different histological subtypes [4,119], which probably have different cells of origin [4]: the small-bile-duct or mixed CCA and the-large-bile duct (mucinous) type iCCAs [22,119]. This last iCCA subtype displays an immunohistochemistry (IHC), gene expression and clinicopathological profile that can be superimposed on pCCA [4,120-122]. Small-bile-duct or mixed CCAs usually show a peripheral localization and an MF growth pattern. In contrast, large-bile-duct (mucinous) type CCAs usually show a periductal infiltrating and/or MF growth pattern [4]. Importantly, these separate entities differ in prognosis (with the mucin-producing iCCAs having a worse prognosis) and in associated diseases [4,10,82,123]. Indeed, parenchymal liver diseases, including chronic viral and non-viral liver diseases and liver cirrhosis, should be considered risk factors for mixed-type iCCAs [4,10,82,123]. In contrast, chronic biliary diseases or pathologies and conditions affecting the intrahepatic medium-large and extrahepatic bile ducts are risk factors for mucin-producing iCCAs and pCCAs [4,10,82,123]. These differences significantly affect the clinical presentation and modalities of diagnosis [4,10,82,123].

As far as the mixed-type MF iCCA is concerned, the clinical presentation is similar to that of other intrahepatic liver malignancies, but different from that of pCCA [4,10,82,123]. CCAs are usually asymptomatic in their early stages (20-25% of cases are incidental findings). Malaise, cachexia, abdominal pain, night sweats, fatigue and/or jaundice, whether associated or not with systemic manifestations, represent the clinical onset of symptomatic iCCA [4,10,82,123]. In contrast, a typically painless jaundice is the most frequent clinical onset in pCCA [4,10,82,123]. Regarding patients with PSC, CCA may present as the development of a rapid deterioration of clinical conditions or dominant stricture during follow up [3]. In general, the MF type represents the most frequent macroscopic presentation of iCCA (>90%), appearing on imaging as a nodule [3,123]. In the context of cirrhotic liver, it was demonstrated that, in contrast-enhanced magnetic resonance imaging (MRI), iCCAs consistently showed a lack of HCC hallmarks, though in computed tomography this occurs only in large nodules (>3 cm) [124-126]. Currently, the identification of HCC with stem-cell features (CK19+-HCC), combined HCC-CCA, cholangiolocellular carcinoma and bile-duct mixed type iCCA with imaging procedures is still an unsolved challenge [3,4,10,123,127,128]. Biopsy is therefore necessary after excluding HCC in cirrhosis, or in the context of a nodule in a non-cirrhotic liver [3,129]. From a histological point of view, the differential diagnosis of iCCA versus HCC or metastasis represents an unsolved problem [2,3,129,130], partly because of the lack of validation of specific markers.

Radiologically, pCCA usually appears as a dominant stricture; therefore, MRI+MRCP represents the imaging procedure with the highest diagnostic accuracy for localizing and sizing the stricture [3], though the definitive demonstration of malignancy is still a challenge [3]. In this respect, endoscopic retrograde cholangiopancreatography enables a number of procedures that may allow microscopic confirmation, including cytology, brushing, fluorescent in situ hybridization (FISH)-polysomy, biopsy, or further innovative techniques [3]. However, all these techniques have an unsatisfactory sensitivity [54,130-133], and even FISH-polysomy demonstrated a low sensitivity for detecting CCA in PSC patients in a meta-analysis [133]. Essentially, the diagnosis of CCA still requires a combination of clinical, radiological and non-specific histological/biochemical markers [3].

Cells of origin and cancer stem cells: inter- and intratumoral heterogeneity

The cell of origin, or cancer-initiating cell, is considered to be the normal cell that receives the first cancer-causing mutation [134,135]. On the other hand, cancer stem cells (CSCs) are the cells that sustain tumor growth and propagation [134,135]. Therefore, the phenotypes of cells of origin and CSCs may be substantially different [134,135].

CCA subtypes exhibit pronounced heterogeneity, raising the question of potentially diverse cellular origins [4,9-12]. Possible cells of origin are human Hepatic stem/progenitor cells (hHpSCs), immature neural cell adhesion molecule positive (NCAM+) cholangiocytes, mature (NCAM-) interlobular cholangiocytes, and PBGs (Fig. 1) [10]. According to different observations, pCCAs most probably originate from mucin-secreting cholangiocytes and PBGs [56,118] located in hilar bile ducts. Indeed, pCCAs are associated with preneoplastic lesions emerging in surface epithelium [2,3] and PBGs [56,118]. Moreover, the IHC and gene expression profiling of pCCA have shown a strong similarity to the cylindrical, taller, mucin-producing cholangiocytes (and PBGs) lining hilar bile ducts [4,118,136]. On the other hand, iCCAs show inter-tumor heterogeneity, leading to their classification into two main different histological subtypes [4,119], most probably with different cells of origin [4]. The large bile duct (mucinous) type iCCAs arise in larger intrahepatic bile ducts; this portion of the intrahepatic biliary tree shares anatomical and embryological similarities with the extrahepatic biliary tree and pancreatic duct system [22,119]. This iCCA subtype displays IHC, gene expression and a clinicopathological
profile that can be superimposed on pCCA [4]; in addition, it displays (together with pCCA) large similarities to pancreatic ductal adenocarcinoma [120-122]. The pattern of growth and the presence of preneoplastic lesions in cholangiocytes and PBGs lining larger intrahepatic bile ducts identify these cells as candidate cells of origin [56,136]. The small bile duct (mixed) type iCCAs show an IHC and gene expression profile similar to mucin-negative cuboidal cholangiocytes that line the smaller bile duct (interlobular bile duct and bile ductules) [4]. In addition, the phenotypic and genotypic profiles are similar to those of cholangiolocellular carcinoma thought to originate from hepatic progenitor cells [4,137,138]. A number of observations suggest that small bile duct (mixed) type iCCAs, together with cholangiolocellular carcinoma and CK19+ HCC, represent a group of primitive liver cancers originating from hHpSCs [4,21,139-141]. Thus, a unique feature of CCA is that it recognizes as origin tissues the hepatic parenchyma or large bile ducts, furnished by two distinct stem cell niches, the canals of Hering and the PBGs, respectively (Fig. 2) [12].

Based on the grade of maturation of the cell of origin within the two lineages, our suggestion is that CCAs could be classified as (Fig. 2):

- Primary liver parenchymal CCA: cholangiolocarcinoma, small-bile duct type (mixed) CCA, and combined HCC-CCA. These tumors emerge within the liver parenchyma from the canals of Hering, bile ductules and interlobular bile ducts and indeed originate from hHpSCs, immature NCAM+ cholangiocytes, or mature (NCAM-) interlobular cholangiocytes.
- Primary biliary CCA: dCCA, pCCA, and large-bile-duct (mucinous) type iCCA. These tumors emerge from the extrahepatic biliary tree and larger intrahepatic bile ducts and originate from PBGs or the surface epithelium of corresponding bile ducts.

 CSCs are defined as the cells within a tumor that possess the capacity for self-renewal and generation of heterogeneous lineages. CSCs are highly tumorigenic and are responsible for chemo-radio resistance and tumor recurrence [137,142,143]. Several CSC markers have been reported in human CCAs [137], including CD133 [144], epithelial cell adhesion molecule (EpCAM) [145], CD44 [146], CD13 [147] and CD90 [148]. In a recent study, CSCs comprised more than 30% of the tumor mass in human CCA sub-types [120]. Interestingly, the CSC profile was similar between mucinous iCCA and pCCA [120].

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**Figure 2** Classification of cholangiocarcinoma based on cell lineages of origin.

Based on the grade of maturation of the cell of origin within the two lineages, one constituted by the hepatic stem cells and non mucin-producing cuboidal cholangiocytes in Hering canals and bile ductules/interlobular bile ducts (see red points), the other constituted by peribiliary glands (PBGs) and surface epithelium of corresponding bile ducts (see blue points), our suggestion is that CCAs could be reclassified as:

- primary liver parenchymal CCA comprising combined HCC-CCA, cholangiolocarcinoma, and small-bile-duct type (mixed) CCA
- primary biliary CCA comprising dCCA, pCCA, and large-bile-duct (mucinous) type iCCA

*Modified from Cardinale et al. Nat Rev Gastroenterol Hepatol* [ref. 119]
Clinicopathological studies demonstrated the expression of cluster of designation (CD133) [34]. Epithelial cell adhesion molecule (EpCAM) [35], CD44 and SRY-related HMG-box(Sox2)[146], and S100 calcium-binding protein A4 (S100A4) EMT marker [13,149] as being significant contributors to the worsening of CCA prognosis. EpCAM and CD133 are expressed by microparticles in a liquid biopsy of CCA patients and showed significant diagnostic and prognostic potential [150]. Moreover, Sox17, a bilio-pancreatic progenitor transcriptional factor that regulates the differentiation and maintenance of the biliary phenotype, acts as a tumor suppressor in CCA and its restoration may represent a promising new therapeutic strategy [151].

Vital stem-cell signaling pathways, especially Notch, Wnt/β-catenin or Hedgehog, are crucial players in CCA pathogenesis [137,138] and could serve as molecular targets, contributing to efficient CSC target therapy [152,153]. However, CSC markers and signaling pathways are largely shared by normal and cancer stem cells, thus limiting targeted strategies specific to CSCs in primary liver cancers [137,151,153]. Moreover, CSCs display considerable crosstalk and redundancy in signaling pathways [152,153].

Cancers that follow the stem cell model are also subject to clonal evolution, as well as heterogeneity from environmental differences within tumors [154,155]. Specifically, genetically distinct subclones, together with developmental pathways and epigenetic modifications, can contribute to functional heterogeneity [155]. In CCA, the tumor microenvironment is composed of cancer-associated macrophages, fibroblasts, and vascular cells and functions as a specialized CSC niche, thus contributing to the maintenance of stemness and chemoresistance [138,149,155-157]. Many factors needed for the maintenance of CSC within its niche (cellular components, soluble factors, cytokines and growth factors) [138,157], should be considered additional potential targets for successful therapeutic strategies against CSC [152,153,157,158].

Human CCA cells expressed EMT markers both in situ and in vitro and, interestingly, subcutaneous xenografts from highly tumorigenic CD90+ or CD13+ CSCs were dominated by stromal markers [120]. By contrast, in intrahepatic xenografts in cirrhotic livers, tumors were dominated by epithelial traits reproducing the original human CCAs, suggesting that CSC subpopulations generate different types of cancer, depending on the microenvironment [120]. A similar phenomenon was described at the single-cell level in breast cancer, where metastatic cells from low-burden tissues were individuated by their increased expression of stem cell, EMT, prosurvival, and dormancy-associated genes with respect to high-burden metastasis, which is constituted by differentiated tumor cells [159]. The increased expression of stem and EMT genes in CCAs may imply a process of metastasis that could be determined by circulating tumor cells, characterized by an intermediate phenotype that is largely unknown and represents a research target with important clinical implications.

**Figure 3** We suggest a model of cholangiocarcinogenesis in which the interplay among the lineage of origin, the pathologic conditions, and the cell-etiopathology-specific somatic mutations and epigenetic modifications (+/- shared mutations) determines multiple patterns of cholangiocarcinogenesis

### Concluding remarks

Regarding the origin of CCA, a physiopathology concept that emphasizes the lineage of origin over the cell of origin should be considered (Fig. 2) [10,12,13,148]. A CCA classification based on the cell lineages of origin is more consistent with current knowledge regarding epidemiology and risk factors and may have important clinical implications in the definition of specific therapeutic targets (Fig. 2). Moreover, it highlights a lineage dependency of chronic liver diseases and related molecular carcinogenesis [12]. Since somatic mutagenesis and epigenome features are cell/lineage specific [160] and are largely driven by the preneoplastic pathologic milieu (see inflammation) (Fig. 3), the multiple lineages of origin plus the related diseases may explain the intertumoral heterogeneity observed at any level of CCA (Fig. 4). Emphasizing cell lineage in CCA development would have implications for preventive strategies in patients with clinical or subclinical underlying hepatic or biliary diseases, diagnosis and therapy (Fig. 4).

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Figure 4 Origin and multilevel repercussions of the CCA heterogeneity.

A. The anatomical-functional heterogeneity of the biliary tree may explain the intertumoral heterogeneity observed at any level in CCA. Somatic mutagenesis and epigenetic features are cell/lineage type-specific, and largely driven by the preneoplastic tissue pathologic milieu (see inflammation). These are tissue-specific. Indeed, pathologic conditions targeting liver parenchyma are chronic hepatitis, nonalcoholic steatohepatitis, and primary biliary cholangitis, while pathologic conditions targeting the bile ducts are primary sclerosing cholangitis and liver flukes. B. Cells arising from a hHpSC lineage (left) or from a biliary tree stem/progenitor cell (hBTSC) lineage (right) are considered cells of origin in cholangiocarcinoma underlie biological, pathologic, and epigenetic modifications; and associated diseases, management, prominent somatic mutations and candidate experimental models of the CCA subtypes arising within the two cell lineages are shown

**HCC, hepatocellular carcinoma;**

**CLC, cholangiolocarcinoma**

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