TITLE: Mixed hepatocellular-cholangiocarcinoma tumors: cholangiolocellular carcinoma is a distinct molecular entity.

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**ABBREVIATIONS:**

HCC-CCA: Hepatocellular cholangiocarcinoma  
HCC: Hepatocellular Carcinoma  
iCCA: Intrahepatic Cholangiocarcinoma  
WHO: World Health Organization  
CLC: Cholangiolocellular Carcinoma  
TGF-beta: Transforming Growth Factor, Beta 1  
TERT: Telomerase Reverse Transcriptase  
KRAS: Kirsten Rat Sarcoma Viral Oncogene Homolog  
IDH1/2: Isocitrate Dehydrogenase 1/2  
SNP: Single Nucleotide polymorphism
WES: Whole Exome Sequencing
SALL4: Spalt-Like Transcription Factor 4
IRB: Institutional Review Board
FFPE: Formalin-Fixed, Paraffin-Embedded
TACE: Transarterial Chemoembolization
FF: Fresh-Frozen
OCT: Optimal Cutting Temperature compound
KRT7: Keratin 7
KRT19: Keratin 19
HEP1: Hep Par 1
CK7: Cytokeratin 7
CK19: Cytokeratin 19
NCAM: Neural Cell Adhesion Molecule 1
GPC3: Glypican 3
EpCAM: Epithelial Cell Adhesion Molecule
CNV: Copy Number Variation
CCND1: Cyclin D1
FGF19: Fibroblast Growth Factor 19
EMT: Epithelial to Mesenchymal Transition
INF: Interferon
IPA: Ingenuity Pathway Analysis
IGF: Insulin-like Growth Factor
IGF1R: Insulin-Like Growth Factor 1 Receptor
MYC: V-Myc Avian Myelocytomatosis Viral Oncogene Homolog
BRAF: B-Raf Proto-Oncogene, Serine/Threonine Kinase
CTNNB1: Catenin Beta 1
TP53: Tumor Protein P53
FGFR2: Fibroblast Growth Factor Receptor 2
CN-LOH: Copy Neutral Loss of Heterozygosity
NTP: Nearest Template Predictions
GSEA: Gene Set Enrichment Analysis
FDR: False Discovery Rate
SD: Standard Deviation
RNA: Ribonucleic acid
DNA: Deoxyribonucleic acid
dsDNA: Double Stranded Deoxyribonucleic Acid
PCR: Polymerase Chain Reaction
qRT-PCR: Quantitative Real Time Polymerase Chain Reaction
HBV: Hepatitis B virus
HCV: Hepatitis C virus
PSC: Primary Sclerosing Cholangitis

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interpretation of data), MES (study concept and design), JML (study concept and design; drafting of the manuscript; critical revision of the manuscript for important intellectual content; obtained funding; study supervision).
ABSTRACT

Background and aims: Mixed hepatocellular-cholangiocarcinoma (HCC-CCA) is a rare and poorly understood type of primary liver cancer. We aimed to perform a comprehensive molecular characterization of this malignancy.

Methods: We performed gene expression profiling, DNA copy number detection, and exome sequencing using formalin-fixed samples from 18 patients with mixed HCC-CCA encompassing the whole histological spectrum of the disease. Comparative genomic analysis was performed with independent datasets of HCC (n=164) and iCCA (n=149).

Results: Integrative genomic analysis of HCC-CCAs revealed that cholangiolocellular carcinoma (CLC) represents a distinct biliary-derived entity compared with the stem-cell and classical types. CLC tumors were NCAM positive (6/6 vs 1/12, \( P<0.001 \)), chromosomally stable (mean chromosomal aberrations 5.7 vs 14.1, \( P=0.008 \)), showed significant upregulation of TGF-beta signaling and enrichment for inflammation-related and immune response signatures (\( P<0.001 \)). Stem-cell tumors were characterized by SALL4 positivity (6/8 vs 0/10, \( P<0.001 \)), enrichment of progenitor-like signatures, activation of specific oncogenic pathways (i.e. MYC and IGF), and signatures related to poor clinical outcome. Regarding classical type, a significant correlation in the copy number aberrations of the iCCA and HCC components suggested a clonal origin. Exome sequencing revealed an average of 63 non-synonymous mutations per tumor (mean driver mutations:2). Among those, \( TP53 \) was the most frequently mutated gene (6/21, 29%) in HCC-CCAs.

Conclusions: Mixed HCC-CCA represents a heterogeneous group of tumors, with stem-cell type characterized by features of poor prognosis and classical type with common lineage for HCC and iCCA components. CLC stands alone as a distinct biliary-derived entity associated with chromosomal stability and TGF-beta signaling.
LAY SUMMARY

The molecular characterization of mixed hepatocellular cholangiocarcinoma (HCC-CCA) defined that cholangiolocellular carcinoma (CLC) is a distinct molecular entity with biliary-derived origin and no traits of HCC. On the other hand, within the mixed HCC-CCA, the stem-cell type shared aggressive phenotype and poor outcome whereas the classic type shows a common cell lineage for both the HCC and the iCCA component. These data supports re-defining the pathological classification of mixed HCC-CCA in light of the novel molecular data provided.
INTRODUCTION

Liver cancer is the second leading cause of cancer-related deaths, with more than 850,000 new cases annually worldwide [1]. Mixed hepatocellular-cholangiocarcinoma (HCC-CCA) is a rare type of primary liver cancer accounting for less than 1% of all primary liver malignancies [2,3]. Diagnosis is based on histological examination and requires unequivocal presence of both hepatocellular carcinoma (HCC) and intrahepatic cholangiocarcinoma (iCCA) elements intimately admixed [2]. Due to its low incidence and the lack of an established consensual pathological diagnosis, the demographic features and clinical behavior of these tumors remain ill-defined. Roughly, the age, sex specific incidence and geographical distribution are similar to those for HCC [2,4,5]. Median overall survival rates of HCC-CCA are similar to iCCA [3,6–8]. To date, clinical practice guidelines do not include a specific treatment recommendation for HCC-CCA and surgical resection, when feasible, remains the standard of practice.

Histologically, mixed HCC-CCA is a heterogeneous group of primary liver tumors. According to the 2010 World Health Organization (WHO) classification [2], they are divided in two main categories: the classical type and the stem-cell features type. The classical type is characterized by areas of typical HCC and iCCA with an intermediate transition which holds mixed features of both entities. The category of stem-cell features is further subdivided into typical, intermediate and cholangiolocellular carcinoma (CLC). Subtypes with stem-cell features are composed of tumor cells with intermediate histological features between hepatocytes and cholangiocytes. In addition, recent studies have suggested the presence of distinct properties for each subtype of HCC-CCA with stem-cell features given their association with different clinicopathological factors [9,10]. Unlike HCC or iCCA, there is no genome-wide characterization of mixed HCC-CCA tumors. Indeed, it is unclear whether histologic subtypes have a well-defined correlate at
the molecular level. Gene expression profiling on a small series of HCC-CCA samples suggested that HCC-CCA might share common characteristics with poorly differentiated HCC and iCCA with stem-cell traits [11–15]. Furthermore, WNT/beta-catenin and TGF-β signaling were reported to be significantly activated in mixed HCC-CCA when compared to progenitor-like HCC[13]. Mutational analysis has suggested common recurrent driver mutations in HCC and HCC-CCA in comparison to iCCA, such as larger frequency of TERT promoter mutations and a lower frequency of KRAS and IDH1/2 mutations [14]. On the other hand, genome wide allelotyping analyses of classical HCC-CCA suggested a closer genomic proximately to iCCA than to HCC [16].

Herein, we provide a comprehensive molecular characterization of mixed HCC-CCA including histological characterization, whole-genome expression profiling, single-nucleotide polymorphism (SNP) array, and whole-exome sequencing (WES). Integrated analysis to evaluate the genomic overlap with a large independent set of HCC and iCCA samples was also performed. Overall, integrative genomic analysis indicates that CLC is a distinct entity with a biliary molecular profile, low chromosomal instability, and enrichment of TGF-β and immune-related signaling. The other mixed tumors can be molecularly distinguished in two main subclasses: the stem-cell subclass characterized by the presence of the early progenitor marker (SALL4) and signatures of more aggressive phenotype, and the classical subclass, constituted by components of both HCC and iCCA with a clonal origin. Thus, we propose a molecular classification that encompasses two groups within the mixed HCC-CCA tumors (stem-cell and classical). In addition, our data suggest that CLC stands alone as an independent biliary-derived entity not sharing any molecular traits of HCC.
MATERIAL & METHODS

Human samples and nucleic acid extraction

For the purpose of this study, we evaluated 4728 consecutive patients who underwent surgery for primary liver cancer between 1994 and 2013 at the Icahn School of Medicine at Mount Sinai [HCC (4307, 91%), iCCA (360, 7.7%), and mixed HCC-CCA (61, 1.3%)], following local Institutional Review Board (IRB) approval. Among the 61 mixed HCC-CCA cases, 43 cases were excluded due to several reasons: a) pre-treated with locoregional therapies (such as transarterial chemoembolization, TACE), b) lack of sample availability or c) low tumor cell viability. The diagnosis of mixed HCC-CCA was confirmed by two expert hepato-pathologists (MIF and ST). A final set of 18 patients with available fixed paraffin-embedded (FFPE) samples and clinical data were selected and classified according to the latest WHO classification [2], as classical (n=4) or with stem-cell features (n=14). The stem-cell features subgroup included Typical (n=2), Intermediate (n=6), and CLC (n=6). For the purpose of molecular profiling in the case of the classical mixed tumors, nucleic acids were extracted separately from the HCC-like and CCA-like components (4 patients, 8 tumor samples total). Table 1 summarizes the main clinico-pathological features of the 18 patients included in the study. For the purpose of integrative genomic analysis, molecular data previously reported by our group on HCC (n=164) and iCCA (n=149) was used [17,18]. In addition, for the identification of driver mutations in stem-cell features subtype, fresh frozen optimal cutting temperature compound (FF-OCT) embedded tumor tissues and corresponding normal tissue (n=6 pairs including 3 overlapping cases with above) were provided by the Mount Sinai Institutional Biorepository after IRB committee approval. For detailed description of nucleic acid extraction see supplementary Material and Methods section.
Immunohistochemistry, Whole-genome gene expression profiling, Genome-wide analysis of DNA copy number alteration, Whole-exome sequencing, and Statistical analyses

See Supplementary Materials and Methods section.
RESULTS

CLC, stem-cell and classical types are distinct entities.

In order to understand if the different histological subtypes of mixed HCC-CCA represent distinct subgroups of the disease, we performed gene expression-based unsupervised clustering. The unsupervised clustering analysis (Figure 1, Supplementary Figure 1) revealed three distinct clustered groups: 1) CLC tumors in cluster C, \( P=0.0013 \), 2) stem-cell feature tumors in cluster D \( P=0.0062 \), 3) Classical tumors depending on its components (HCC-component in clusters A and B, \( P=0.024 \)). In addition, the fact that the iCCA-like components of the classical subtype co-clustered with either stem-cell feature tumors or CLC suggest the presence of common molecular traits among these (Figure 1A). Differential molecular profile of CLC with respect to other stem-cell feature tumors was further confirmed by integrative genomic analysis with an independent set of HCC (n=164) and iCCA (n=149) samples (Figure 2). CLC tumors significantly co-clustered together suggesting high genomic similarity among them in comparison to other primary liver tumors \( P<0.001 \). Significant genomic proximity was also observed for stem-cell HCC-CCA \( P<0.001 \). Moreover, CLC tumors co-clustered with iCCA from the proliferation class, whereas the stem-cell mixed tumors co-clustered with HCC with progenitor-like traits (Figure 2, \( P<0.001 \)). Subsequent analysis of cell lineage with specific marker genes further corroborated the observation that CLC may represent a separate entity, as indicated by the expression of a biliary phenotype with significant up-regulation of biliary-specific genes (e.g. \( KRT7, KRT19, ITGB4 \)) and down-regulation of hepatocyte-related genes (e.g. \( ADH1A, ALB, APOB, HNF1A \))[19] (Figure 1). These findings were in concordance with the immunostaining profile (Figure 1B and Figure 3), which defined CLC tumors as negative for the hepatocyte marker HepPar1 (0/6 in CLC vs 10/12 in others, \( P=0.015 \), Supplementary table 1), but positive for biliary markers (CK7, CK19).
and more specifically the progenitor-like marker NCAM (6/6 in CLC vs 1/12 in others, \( P<0.0001 \), Supplementary table 1).

The stem-cell molecular subclass was characterized by the expression of both hepatocyte and biliary markers (Figure 1B and Figure 3), and the early progenitor cell marker SALL4 (6/8 vs 0/10 in rest of mixed tumors, \( P=0.0004 \), Supplementary Table 1). On the other end of the spectrum, the HCC-like and iCCA-like components of the classical subtype showed a biphenotypic profile with simultaneous expression of hepatocytic and biliary markers (Figure 1B) despite their distinct histological features (Figure 3). Comparison of gene expression levels and immunostaining grading scores showed significant correlation for all markers used (Supplementary Figure 2).

The different genomic profile of CLC tumors was further confirmed by the DNA copy number variation (CNV) analysis which revealed significantly higher chromosomal stability in CLC compared to the non-CLC tumors (5.7 mean alterations in CLC vs 14.1 for others, \( P=0.008 \), Figure 4). In contrast, classical and stem-cell subclasses presented frequent broad chromosomal aberrations, recapitulating those previously reported in both HCC and iCCA, including gains of 1q, and 8q, and losses of 4q, 8p, 9q, 16q and 17p (Figure 4). High-level amplifications of 11q13, harboring the oncogenes \( CCND1 \) and \( FGF19 \), were detected in 3 cases of mixed HCC-CCA (Supplementary Table 2). Thus, the above findings support that mixed HCC-CCA tumors can be classified into 2 distinct molecular subclasses (stem-cell and classical). CLC tumors can be defined as a separate entity with biliary phenotype and no traits of HCC. We, then, further characterized the molecular traits of each of these tumor subtypes.

**CLC subclass: characterized by TGF-β signaling and immune-related response signaling**
When evaluating previously reported prognostic gene signatures in liver cancer, CLC showed a significant enrichment of S1 subclass [20] characterized by higher TGF-β activation, and Late TGF-β induced signature [21] (Figure 5A, Supplementary Table 3). Furthermore, unsupervised hierarchical clustering with independent set of iCCA and HCC samples, suggested a shared molecular profile of CLC and progenitor-like iCCA tumors (Figure 2, P<0.001). On the other hand, gene set enrichment analysis (GSEA) of canonical and hallmark oncogenic pathways showed that the CLC subclass was significantly enriched with TGF-β signaling, epithelial to mesenchymal transition (EMT), inflammation and immune response related signaling (i.e. TNF-α, INF-γ, IL2/STAT5, IL6/STAT3, T cell response) (Supplementary Table 4). Notably, significant enrichment of immune cells [22] including T cells (i.e. effector memory, CD8+ PD1 high), cytotoxic lymphocytes, T cell helper 1 (Th1), natural killer (NK) and neutrophils was also detected in CLC (Figure 5A-middle panel, P<0.05). In addition, specific chemokines (e.g. CXCL12, CCL2, CCL21) and several cytokine receptors (i.e. CXCR4, IL8RB, IL10RA, IL17RC) were also identified to be significantly up-regulated in CLC (Supplementary Figure 3, Supplementary Figure 4). Furthermore, the Ingenuity Pathway Analysis (IPA) of the top deregulated genes in CLC (Supplementary Table 5), predicted TGF-β as the major activated upstream effector molecule in these tumors (Supplementary Figure 5A, z-score>2, p<0.0001). Specifically, the ligand TGFβ2 was found to be significantly up-regulated among the different TGF-β superfamily of ligands (Supplementary Figure 3). Other significantly deregulated candidate upstream effectors and top bio-functions in CLC included activation of TP53 (Supplementary Figure 5A), and DNA damage response checkpoint regulation and Ataxia telangiectasia mutated (ATM) signaling (Supplementary Figure 5B), respectively. Overall, TGF-β and pro-inflammatory response related signaling pathways were found to be specifically up-regulated in CLC tumors.
**Stem-cell subclass: associated with progenitor-like phenotype and proliferative signaling pathways**

We next sought to assess dominant genomic traits in the stem-cell subclass. GSEA of the SALL4 positive stem-cell tumors showed enrichment of progenitor-like liver cancer subtypes (i.e Stem cell [12], CK19 [17], EpCAM [23]) and more aggressive HCC—including HCC Proliferation subclass [24], cell-cycle deregulated G3 subclass [25], poor survival cluster A [26], and S2 subclass [20]—together with activation of IGF1R [27] and NOTCH [28] signaling (**Figure 5A, Supplementary Table 3**). Consistently, MYC, mTORC and NOTCH signaling were identified as the main canonical pathways associated with stem cell subclass (**Supplementary Table 6**). IGF2 signaling was further confirmed as one of the main deregulated signaling pathway networks in these tumors (**Figure 5C, Supplementary Table 7**). Another core node up-regulated in stem cell subclass included genes implicated in the hepatic specification of liver progenitor cells (i.e. PROX1, HNF1B, FOXA1, FOXA3), suggesting a more hepatocyte committed lineage in contrast to the biliary phenotype observed in CLC. In addition, consistent with SALL4 expression, the pluripotent embryonic related OCT4 signaling was found as one of the top activated canonical pathways (**Supplementary Figure 6**). On the other hand, in regards to CNV profiling (**Figure 4**), the most frequent chromosomal alterations reported in HCC showed an enrichment trend in stem-cell subclass, including 1q gains (6/8 vs 2/10), 8q gains (5/8 vs 1/10), 1p losses (4/8 vs 1/10), and 4q losses (6/8 vs 2/10). Altogether, these data suggest that the stem-cell subclass shares common molecular features with more aggressive and progenitor-like HCCs.

**Classical subclass: HCC and iCCA components are derived from the same clone**

The classical subclass showed enrichment of the poor prognosis iCCA subclass [29] and the chromosome 7 polysomy HCC subclass [24]. By performing separate analysis of the
classical iCCA and HCC components, we found enrichment of several gene sets in the iCCA component whereas none reached our pre-specified statistical threshold in the HCC component (Figure 5A, Supplementary Table 3). Moreover, the classical iCCA component seemed to have an intermediate molecular profile to stem-cell and CLC subclasses, as it showed association with liver derived gene signatures enriched in either stem-cell (i.e. HCC Proliferation, G3, S2, cluster A, CK19, IGF1R) or CLC (i.e. Late TGF-β induced signature). GSEA of canonical signaling pathways in the iCCA component (Supplementary Table 8) and HCC component (Supplementary Table 9) of the classical tumors further suggested a more aggressive phenotype for the iCCA component. Pathways enriched in the iCCA component included pro-mitotic DNA replication related signaling, proliferative signals such as MYC and mTOR, and pro-inflammatory pathways such as INF-γ and downstream IL2/STAT5 signaling (Supplementary Table 8). These data indicate that the iCCA component of the classical subclass may have a more aggressive molecular profile within classical mixed tumors.

Clonality analysis based on CNV profiling of the HCC-like and iCCA-like components of the classical HCC-CCA showed a significant correlation and remarkable similarity (mean 51%, $P<0.001$) between both components in 3 out of 4 cases (Supplementary Figure 7 and 8, Supplementary Table 10). Specifically, focal chromosomal aberrations harboring known driver genes (i.e. 11q13 gain and 9p21 deletion), were identified in both components in 1 classical subclass case (Supplementary Table 2, Supplementary Figure 8). These findings suggest that the HCC and iCCA components of the classic subtype may share a common cell of origin that later undergoes clonal divergent expansion.
Landscape of mutations in mixed HCC-CCA

Next, we evaluated the mutational landscape of mixed HCC-CCA tumors (4 CLC and 2 stem-cell, including 3 overlapping cases with above analysis) using exome sequencing. Somatic substitutions were predominantly G>A and C>T transitions in both CLC and stem cell tumors (Supplementary Figure 9A), as previously reported in iCCA [30,31], HCC [32,33] and other cancers [34]. Globally, an average of 63 non-synonymous mutations (range 10-129) and 3 small insertions and deletions (indels, range: 1-5, Supplementary Table 11) were identified per mixed HCC-CCA tumor (Supplementary Figure 9B). Non-synonymous mutations included 93% missense (average 59 per tumor, range: 9-115) and 7% nonsense (average 4 per tumor, range: 1-14,) mutations. Overall, both CLC and stem-cell subclasses presented similar percentage of missense, silent, and nonsense mutations (Figure 6A). Among the non-synonymous mutations, 10 affected known cancer driver genes [35,36] (average of 2 driver mutations per tumor), which were confirmed by independent PCR and sequencing in each tumor (Figure 6B, Supplementary table 12).

We further explored the incidence of hot-spot mutations identified by exome sequencing in known oncogenic drivers (BRAF, DNMT3A, IDH1) together with recurrent mutations reported in HCC (TERT promoter, CTNNB1, TP53) and iCCA (KRAS, FGFR2-BICC1, FGFR2-PPHLN1, IDH2) in the remaining FFPE samples. TP53 mutations emerged as the most frequent alteration (6/21, 29%), regardless of the subclass (Figure 6C). The mutational profile of CLC included TP53 and IDH1, while the stem-cell subclass (i.e. TERT promoter, TP53, AXIN1, BRAF, FGFR2-BICC1) seemed to recapitulate those characteristic of both HCC and iCCA (Figure 6B and 6C). Interestingly, the most frequent HCC driver mutation, TERT promoter mutation, co-occurred with TP53 mutation in 2 cases (1 classical case and 1 stem-cell SALL4 negative case). The co-occurrence of TERT promoter and TP53 mutations (Figure 6C), in both the HCC and iCCA component
of the classical subclass case further supported the model of single cell of origin suggested by CNV profiling. Other screened mutations such as \textit{CTNNB1}, \textit{KRAS}, \textit{FGFR2-PPHLN1} and \textit{IDH2} were absent in our cohort. In summary, mutational profiling of mixed HCC-CCA revealed common drivers in typical HCC and iCCA tumors. However, the CLC subclass seems to have a distinct mutational profile in comparison to stem-cell subclass.

**External validation confirms CLC as a distinct entity from stem-cell HCC-CCA**

We validate the molecular subclasses in a publicly available dataset including 20 HCC-CCAs described with CLC features [13]. The CLC main molecular characteristics, such as biliary committed phenotype and activation of TGF-beta and immune-response signaling, were successfully reproduced. (Supplementary Figure 10A). In addition, a CLC-derived 156-gene signature (Supplementary Table 13) was significantly enriched in ~60% (11/20) of these CLC tumors. Subclass mapping approach further reinforced the notion that CLC tumors have a sole biliary-like phenotype, with no traits of HCC [37] (Supplementary Figure 10B). Moreover, stem-cell HCC-CCAs shared molecular traits of HCC but not of CLC tumors (Supplementary Figure 10B) suggesting a hepatocyte lineage. These results confirm that CLC and stem-cell mixed tumors represent distinct molecular entities.
DISCUSSION

Mixed HCC-CCAs are a heterogeneous group of primary liver cancers. The current histological classification describes two main subtypes, classical and with stem-cell features. Our study provides molecular evidence confirming a biphenotypical fingerprint for stem-cell and classical types, while defining CLC as a separate biliary-derived entity with no genomic features of HCC. We were able to characterize these differences based on integrated genomic analysis of gene expression, DNA copy number alteration, signaling pathway deregulation and mutational profiling (Figure 7).

CLC was characterized by chromosomal stability, up-regulation of TGF-beta signaling, prominent enrichment in immune-related pathways and defined biliary features. Histologically, CLC were characterized by being strongly embedded in fibrous stroma and by presenting positive NCAM immune-staining, as previously described [5,11,38,39]. Characteristically, TGF-β signaling was activated in CLC tumors as opposed to other mixed HCC-CCA. TGF-β is a pleiotropic cascade with different functions depending on the cellular context. It exerts pro-tumorigenic effects by enhancing tumor growth and invasion through the induction of EMT, activation of myofibroblasts and collagen deposition [40]. TGF-β has also been associated with the transformation of hepatic progenitor cells (HPCs) into tumor initiating cells [41] and biliary differentiation during early development [42]. From the signaling stand point, CLC tumors also showed up-regulation of EMT-related markers, down-regulation of hepatocyte specific gene markers and enrichment in immune cells (T cells, cytotoxic lymphocytes, Th1, NK) and immune-response related gene signatures. Particularly, the CXCL12-CXCR4 axis, which has been described to be instrumental for the chemoattraction of myeloid and lymphoid cells in to the tumor [43], was also shown to be significantly up-regulated in CLC. All these findings -coordinated Th1 cell and cytotoxic immune infiltration- are consistent with those reported in other
tumor types [44–46]. It is intriguing to understand how activation of TGF-β signaling, which is a known immune-suppressor, coexists with activation of T cells in the same tumor, a finding that requires further studies. In our view, all these specific molecular traits support the concept that CLC should stand alone as a molecularly differentiated subtype of intrahepatic biliary carcinomas.

Tumors belonging to the stem-cell subclass share completely distinct molecular traits. First, they showed enrichment for signatures defining activation of proliferative signals such as MYC, IGF2, mTOR and NOTCH and HCC poor prognosis. Histologically, they were characterized by the presence of SALL4 – an early progenitor-like marker- in 75% (6/8) of cases compared to none of other mixed subclasses. Of note, re-expression of the oncofetal SALL4 transcription factor has also been described in a subset of HCCs with progenitor features and poor prognosis [47].

The classical subtype represents a completely distinct entity, since it shares features from both typical HCC and iCCA. Our findings support a single cell of origin model for these tumors based on similarities in CNV aberrations in HCC and iCCA components in 3 out of 4 paired cases. These results are aligned with previous findings based on LOH analysis where 70% (8 out of 11) of the cases studied showed significant similarities [48]. Moreover, the presence of a characteristic transition area between the HCC and iCCA components with biphenotypic features [2], further supports the model of a single clonal process in which genetic divergence within the tumor parallels the histological diversity.

From the molecular standpoint, each component retained genomic and biomarker traits resembling either HCC or iCCA. Similarly, mutational profiling data showed common oncogenic driver mutations characteristic of either HCC or iCCA in classical HCC-CCA subclass. Finally, we did not identify WNT/β-catenin signaling activation or presence of
*CTNNB1* mutations in our classical HCC-CCA samples, in agreement with previous studies [13,16].

After thoroughly exploring all genomic results, we can speculate on the diverse cellular lineage of this heterogeneous group of tumors (Supplementary Figure 11). Certainly, our results support the possible existence of multiple cells of origin [49]. Mixed HCC-CCA may share a common ancestor, the HPCs, but also might derive from more mature progenitor cells. Our gene expression profiling data suggested a biliary committed precursor in the CLC cases and a biphenotypic progenitor-like precursor in the stem-cell and classical subtypes. First, CLC characteristically express NCAM, and has a loss of hepatocyte markers only retaining cholangiocyte markers, both features suggesting a more mature biliary progenitor ancestor. In addition, IDH1 mutation, a known inhibitor of hepatocyte differentiation and gatekeeper for iCCA generation [19], was only detected in CLC. Conversely, the stem cell subclass might derive directly from HPCs, or from transdifferentiation of more mature ancestors. Specifically, those tumors were enriched with stem-cell signatures and SALL4 positive staining and, thus are the logical candidates to derive from HPC lineage. Finally, the classical subtype retained markers of both HCC and iCCA, and since it seems to have a clonal origin, the cells of origin should be mature enough to have lost early progenitor markers (SALL4), albeit retaining biphenotypical markers.

In summary, our study provides a comprehensive molecular characterization of mixed HCC-CCA. First, from the molecular standpoint both the stem-cell and classical types retain biliary and HCC components, and thus fit within the HCC-iCCA definition. Conversely, our data supports defining CLC as a distinct biliary-derived molecular entity with no HCC traits. These results provide the rationale for re-defining the current pathological classification [2], and for establishing more precise therapeutic approaches.
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### Table 1: Clinico-pathological characteristics of mixed HCC-CCA cohort according to mixed molecular subclass.

<table>
<thead>
<tr>
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<td>33 (1-1573)</td>
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</table>

CLC: cholangiolocellular carcinoma. Stem cell subclass includes the histological stem cell feature typical and intermediate subtypes.

\(^1\) Not available in 5 patients

\(^2\) Not available in 1 patient

p-value corresponds to statistical analysis of the 3 molecular subclasses. Fisher Exact Probability Test was used for categorical variables and The Kruskal-Wallis Test for the continuous factors.
FIGURE LEGENDS

Figure 1: Molecular classes of mixed HCC-CCA and correlation with histopathological features. **A)** Non-negative matrix factorization–based clustering analysis of 22 tumor samples from 18 patients with mixed HCC-CCA. **B)** Heat-map representing expression of specific cell lineage gene markers (upper panel) and immunostaining profiling data (lower panel) of HCC-CCA tumors. The iCCA and HCC components of classical subclass were analyzed separately.

Figure 2: Integrative genomic analysis of HCC-CCA with HCC and iCCA. **A)** Unsupervised hierarchal clustering analysis was performed on merged gene expression profile cohorts of HCC-CCA (6 CLCs and 8 stem-cell subclass), HCC (n=164) and iCCA (n=149). Integration of the datasets was based on the Z-score transformation of the differentially expressed gene in each independent cohort. Nearest prediction method was used for the association with liver cancer derived gene signatures. **B)** Schematic representation of overlapping molecular profile and genomic proximity of CLC and stem-cell subclass mixed tumors with iCCA and HCC, respectively, based on unsupervised clustering analysis. Statistically significant association was calculated by Fisher Exact Probability Test (p<0.05).

Figure 3: Immunostaining profile of mixed HCC-CCA subclasses. Representative morphological and positive immunohistochemical staining features observed for each mixed HCC-CCA subclass.

Figure 4: Broad chromosomal alterations detected in mixed HCC-CCA subclasses. Chromosome arms are displayed in descending order along the vertical axis. Detected broad chromosomal gains, losses, and copy neutral loss of heterozygosity (CN-LOH) per tumor sample have been highlighted. The iCCA and HCC components of each classical
case were analyzed separately. Bars indicate the total number of broad chromosomal gains and losses. CLC showed higher chromosomal stability in comparison to other mixed HCC-CCAs (p<0.01, two-sided T-test).

**Figure 5: Whole-genome gene expression analysis of different mixed HCC-CCA subclasses.** A) Heat-map representing prediction of liver cancer derived molecular classification and gene signatures (upper panel), immune-related gene signatures for cell type and activated signaling (lower panel). B) Network analysis of deregulated genes in CLC showed TGF-β signaling as one of the main activated signaling nodes. C) Network analysis of deregulated genes in SALL4 stem-cell subclass showed activation of IGF2 signaling and hepatic specification of progenitor cells. In the network analyses, a node symbolizes a gene or gene product, and direct and indirect interactions are indicated by solid lines and dotted lines, respectively. Statically significant associated features in CLC are highlighted with (*) and in stem-cell subclass with (¥) using Fisher Exact Probability Test for categorical variables and two-sided T-test for continuous variables. *p<0.05, **p<0.01, ***p<0.001. The iCCA and HCC components of each classical case were analyzed separately.

**Figure 6: Mutational profile of mixed HCC-CCA.** A) Histogram of the number of mutations in each primary tumor sample (upper panel) and pie chart representation of the percentage of non-synonymous somatic mutations for CLC and stem-cell subclasses (lower panel). B) Heat-map representing the individual mutation in known cancer driver genes identified by exome-sequencing. C) Heat-map representing the validation of exome-sequencing results and screening of the most prevalent oncogenic mutations reported in HCC and iCCA in the study cohort. Overlapping cases from which fresh-frozen
and FFPE samples were analyzed separately by exome-sequencing and PCR validation are highlighted with an asterisk.

**Figure 7:** Summary of molecular characterization of mixed HCC-CCAs and CLC tumors as distinct entities. CLC only share biliary-derived features, as opposed to HCC-iCCA tumors. Specific cell lineage markers, liver cancer derived gene signatures, pathway signaling, chromosomal stability and driver mutations are depicted for each entity.
Figure 1

A

![Tumor type diagram with clusters A, B, C, D]

Histological tumor type
- CLC
- Classical (iCCA-part)
- Classical (HCC-part)
- Stem-cell Intermediate
- Stem-cell Typical

Cluster
- A
- B
- C
- D

B

<table>
<thead>
<tr>
<th>Expression of cell lineage marker genes</th>
<th>CLC</th>
<th>Stem-cell</th>
<th>Classical</th>
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<tr>
<td>Hepatocyte</td>
<td>ADH1A</td>
<td>ALB</td>
<td>APOB</td>
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<tr>
<td>Biliary</td>
<td>KRT7</td>
<td>KRT19</td>
<td>ITGB4</td>
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<tr>
<td>Stem cell</td>
<td>EpCAM</td>
<td>NCAM</td>
<td>SALL4</td>
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Immunohistochemistry
- Hep-like
- other
- Biliary
- SC-marker

Gene expression
- High (score 3-4)
- Low (score 1-2)
- Negative
- High
- Low
Figure 2

A

Non-proliferation liver cancer (124/327, 38%)

Proliferation liver cancer (204/327, 62%)

B

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<tr>
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<th>Proliferation-like HCC</th>
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<td>S2 class</td>
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<td>Stem-cell HCC-CCA</td>
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Legend:
- Positive
- Not applicable
Figure 4
Figure 5

A

B

C

Figure 5
Figure 7

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<thead>
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<th>Biliary-derived tumor</th>
<th>Mixed HCC-CCA tumors</th>
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<td>CLC</td>
<td>Stem-cell</td>
</tr>
<tr>
<td>Classical</td>
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**Histological markers**

- NCAM+  
- SALL4+  
- SALL4-  
- CK7+  
- CK19+  
- HEP1+  
- GPC3 +

**Gene expression**

- Biliary-like  
  - Biphenotypic (hepatocyte and biliary marker genes)  
  - Hepatocyte-like  
  - Biphenotypic

**Gene signatures enrichment**

- S1 (TGF-WNT)  
  - Late TGF-beta

**Copy Number Variation**

- Chromosomal stability  
  - (Gains: 1q, 8q; Losses: 4q, 8p, 9q, 16q, 16p)

**Common HCC or iCCA driver mutations**

- IDH1  
- TP53  
- FGFR2-BICC1  
- TP53  
- BRAF  
- TERT promoter TP53  
- TERT prom