

# Cholangiocarcinoma stem-like subset shapes tumor-initiating niche by educating associated macrophages

Chiara Raggi<sup>1,†,\*</sup>, Margherita Correnti<sup>1,†</sup>, Antonio Sica<sup>3,4</sup>, Jesper B. Andersen<sup>5</sup>, Vincenzo Cardinale<sup>6</sup>, Domenico Alvaro<sup>6</sup>, Giovanna Chiorino<sup>7</sup>, Elisa Forti<sup>1</sup>, Shannon Glaser<sup>8</sup>, Gianfranco Alpini<sup>8</sup>, Annarita Destro<sup>9</sup>, Francesca Sozio<sup>10,11</sup>, Luca Di Tommaso<sup>9,12</sup>, Massimo Roncalli<sup>9,12</sup>, Jesus M. Banales<sup>13</sup>, Cédric Coulouarn<sup>14</sup>, Luis Bujanda<sup>13</sup>, Guido Torzilli<sup>15</sup>, Pietro Invernizzi<sup>1,2,\*</sup>

<sup>1</sup>Center for Autoimmune Liver Diseases, Humanitas Clinical and Research Center, Rozzano, Italy; <sup>2</sup>Program for Autoimmune Liver Diseases, International Center for Digestive Health, Department of Medicine and Surgery, University of Milan-Bicocca, Italy; <sup>3</sup>Laboratory of Molecular Immunology, Humanitas Clinical and Research Center, Rozzano, Italy; <sup>4</sup>Department of Pharmaceutical Sciences, University of Piemonte Orientale "Amedeo Avogadro" Novara, Italy; <sup>5</sup>Biotech Research and Innovation Centre, University of Copenhagen, Copenhagen, Denmark; <sup>6</sup>Department of Medico-Surgical Sciences and Biotechnologies, Sapienza University of Rome, Rome, Italy; <sup>7</sup>Cancer Genomics Laboratory, Fondazione Edo ed Elvo Tempia, Biella, Italy; <sup>8</sup>Research, Central Texas Veterans Health Care System, Scott & White Digestive Disease Research Center, Scott & White, Department of Medicine, Texas A&M Health Science Center, College of Medicine, Temple, TX, United States; <sup>9</sup>Pathology Unit, Humanitas Research Hospital, Rozzano, Italy; <sup>10</sup>Leukocyte Migration Laboratory, Humanitas Clinical and Research Center, Rozzano, Italy; <sup>11</sup>Department of Molecular and Translational Medicine, University of Brescia, Brescia, Italy; <sup>12</sup>University of Milan Medical School, Milan, Italy; <sup>13</sup>Department of Liver and Gastrointestinal Diseases, Biodonostia Research Institute, Donostia University Hospital, University of the Basque Country (UPV/EHU), CIBERehd, Ikerbasque, San Sebastián, Spain; <sup>14</sup>Inserm U991, Hôpital Pontchaillou, 35033 Rennes cedex, France; <sup>15</sup>Department of Hepatobiliary and General Surgery, Humanitas Research Hospital, Rozzano, Italy

**Background & Aims**: A therapeutically challenging subset of cells, termed cancer stem cells (CSCs) are responsible for cholangiocarcinoma (CCA) clinical severity. Presence of tumorassociated macrophages (TAMs) has prognostic significance in CCA and other malignancies. Thus, we hypothesized that CSCs may actively shape their tumor-supportive immune niche.

**Methods**: CCA cells were cultured in 3D conditions to generate spheres. CCA sphere analysis of *in vivo* tumorigenic-engraftment in immune-deficient mice and molecular character-ization was performed. The *in vitro* and *in vivo* effect of CCA spheres on macrophage precursors was tested after culturing healthy donor cluster of differentiation (CD)14<sup>+</sup> with CCA-sphere conditioned medium.

**Results**: CCA spheres engrafted in 100% of transplanted mice and revealed a significant 20.3-fold increase in tumor-initiating fraction (p = 0.0011) and a sustained tumorigenic potential through diverse xenograft-generations. Moreover, CCA spheres were highly enriched for CSC, liver cancer and embryonic stem cell

<sup>&</sup>lt;sup>†</sup> These authors contributed equally as joint first authors.



markers both at gene and protein levels. Next, fluorescenceactivated cell sorting analysis showed that in the presence of CCA sphere conditioned medium, CD14<sup>+</sup> macrophages expressed key markers (CD68, CD115, human leukocyte antigen-D related, CD206) indicating that CCA sphere conditioned medium was a strong macrophage-activator. Gene expression profile of CCA sphere activated macrophages revealed unique molecular TAM-like features confirmed by high invasion capacity. Also, freshly isolated macrophages from CCA resections recapitulated a similar molecular phenotype of in vitro-educated macrophages. Consistent with invasive features, the largest CD163<sup>+</sup> set was found in the tumor front of human CCA specimens (n = 23) and correlated with a high level of serum cancer antigen 19.9 (n = 17). Among mediators released by CCA spheres, only interleukin (IL)13, IL34 and osteoactivin were detected and further confirmed in CCA patient sera (n = 12). Surprisingly, a significant association of IL13, IL34 and osteoactivin with sphere stem-like genes was provided by a CCA database (n = 104). In vitro combination of IL13, IL34, osteoactivin was responsible for macrophage-differentiation and invasion, as well as for in vivo tumor-promoting effect.

**Conclusion**: CCA-CSCs molded a specific subset of stem-like associated macrophages thus providing a rationale for a synergistic therapeutic strategy for CCA-disease.

**Lay summary**: Immune plasticity represents an important hallmark of tumor outcome. Since cancer stem cells are able to manipulate stromal cells to their needs, a better definition of the key dysregulated immune subtypes responsible for cooperating in supporting tumor initiation may facilitate the development of new therapeutic approaches. Considering that human

Keywords: Cholangiocarcinoma; Cancer stem cells; Tumor-associated macrophages.

Received 7 June 2016; received in revised form 17 August 2016; accepted 18 August 2016; available online 1 September 2016

<sup>\*</sup> Corresponding authors. Addresses: Center for Autoimmune Liver Diseases, Humanitas Clinical and Research Center, via Manzoni 113, 20089 Rozzano (Milan), Italy. Tel.: +39 (02) 8224 5127 (C. Raggi), or Program for Autoimmune Liver Diseases, International Center for Digestive Health, Department of Medicine and Surgery, University of Milan-Bicocca, Via Cadore 48, 20900 Monza (MB), Italy. Tel.: +39 02 6448 8052 (P. Invernizzi).

*E-mail addresses:* chiara.raggi@humanitasresearch.it (C. Raggi), pietro.invernizzi@ unimib.it (P. Invernizzi).

cholangiocarcinoma represents a clinical emergency, it is essential to move to predictive models in order to understand the adaptive process of macrophage component (imprinting, polarization and maintenance) engaged by tumor stem-like compartment.

© 2016 European Association for the Study of the Liver. Published by Elsevier B.V. All rights reserved.

### Introduction

Neoplastic transformation of intra- and extra-hepatic biliary epithelial cells (cholangiocytes) gives rise to cholangiocarcinoma (CCA), a major form of primary liver cancer along with hepatocellular carcinoma (HCC). Due to its aggressiveness, late diagnosis and treatment-refractory nature, CCA represents a clinical emergency [1,2]. CCA incidence and mortality rates are increasing worldwide and therapeutic options are limited.

Due to its desmoplastic nature, CCA is associated with massive presence of stromal cells [3,4]. Among various immunesubgroups (e.g., T cells, neutrophils, granulocytes) [5–8], CCAassociated macrophages represent a poorly defined but very intriguing immune-subset. Indeed, high tumor-associated macrophage (TAM) density significantly correlated with poor prognosis, unfavorable survival and metastasis tendency in CCA patients, suggesting a major role of macrophages in cholangiocarcinoma progression (Reviewed in [4]).

Recently, cancer stem cells (CSCs) have been proposed as a driving force of tumor initiation, dissemination and drug-resistance thus representing a primary therapeutic target.

Importantly, the view that CCA is traditionally considered to originate either from cholangiocytes or progenitor cells, has been recently challenged. Indeed, it is proven that CCA can also originate from mature hepatocytes in mouse models [9].

Although it has been shown that HCC progression is driven by CSCs [10,11], identification of CCA stem-subset is still largely unexplored and limited to classical antigenic approaches (Reviewed in [4]). Similar to normal stem cells, CSCs are believed to reside in specialized microenvironment ("CSC-niche") within tumor context [12–14] that supports self-renewal and drug-resistance [15–19].

Tumor stroma co-evolves together with cancer cells and plays a critical part in both malignancy initiation as well as key steps of growth and metastasis [20]. Since macrophages show a remarkable degree of plasticity during tumor development, we assumed that, depending on CSC tumorigenic-state, CSCs might actively release soluble mediators that engaged circulating monocytes to tumor-initiating niche and prompted macrophagedifferentiation towards an exclusive subset of TAMs, the CSCassociated TAMs. Although bulk tumor cells support TAMs recruitment and activation, specific effects of CSCs on TAMs phenotype are still unexplored in human CCA.

The current study sought to 1) identify a functional CSCsubset in human CCA and 2) verify the existence of a peculiar TAM-compartment as result of a bioactive CCA stem-like cells.

#### Materials and methods

#### Mice

## JOURNAL OF HEPATOLOGY

Laboratory) were used in all experiments. The mice were used at 6 weeks of age, in accordance with the guidelines and with the approval of the local Experimental Animal Committee [16].

### Sphere assay

The cells were grown in anchoring-independent conditions with selective serumfree Dulbecco's modified Eagle's medium (DMEM)/F12 medium supplemented with 1X B27 supplement minus vitamin A (Life Technologies), human recombinant epidermal growth factor (hrEGF) (R&D System) (20 ng/ml), and bFGF (R&D System) (20 ng/ml) for 15 days.

#### Monocyte isolation and macrophage-differentiation

Human monocytes were obtained from healthy blood donor buffy coats by gradient centrifugation using a Ficoll gradient (GE Healthcare) and further purified from peripheral blood mononuclear cells (PBMC) by magnetic-activated cell sorting (MACS) using CD14 microbeads (Miltenyi Biotech). After MACS purification, two fractions were obtained: the CD14<sup>+</sup> fraction and the eluate (composed of all PBMC CD14<sup>-</sup> cells). The purity of CD14<sup>+</sup> cells was >90%. CD14<sup>+</sup> cells were cultured in presence of 30% CCA sphere- or monolayer conditioned medium (CM).

#### In vivo study

In vivo experiments were performed in accordance with the guidelines and approval of the local Experimental Animal Committee. The following materials were used: 1000 sphere- or monolayer-derived cells to measure the tumorigenic potential of CCA cells and 1000 monolayer-derived cells and 300 macrophages for TAMs + CCA monolayer co-injection experiments. The cells were dissociated into single-cell suspensions and resuspended in 100  $\mu$ l of DMEM and reduced Matrigel<sup>®</sup> growth factor (BD Bioscience) (1:1), and the mixture was subcutaneously (s.c.) injected into the right flank of 6-week-old NSG mice. Tumor growth was monitored, and the diameter of the growing tumors was measured in millimeters every week using a caliper. The animals were sacrificed when the xenografts reached 2.0 cm in diameter [15,16,26]. The limiting dilution analysis was performed by sorting 1000/100/10 alive cells from dissociated CCA monolayer and spheres. Web-based extreme limiting dilution analysis (ELDA) statistical software (http://bioinf.wehi.edu.au/software/limdil/index. html) was used [28].

#### Patient samples

All patients gave written, informed consent. The study was approved by local institutional review boards. CCA samples and peritumoral non-cancerous liver were obtained from patients submitted to surgical resections (Department of Hepatobiliary Surgery, Istituto Clinico Humanitas). Distinctions between intrahepatic CCA and perihilar CCA were based on clinical records including surgical reports.

Further details may be found in the Supplementary material.

#### Results

#### Human CCA Spheres retain stem-like tumor-initiating features

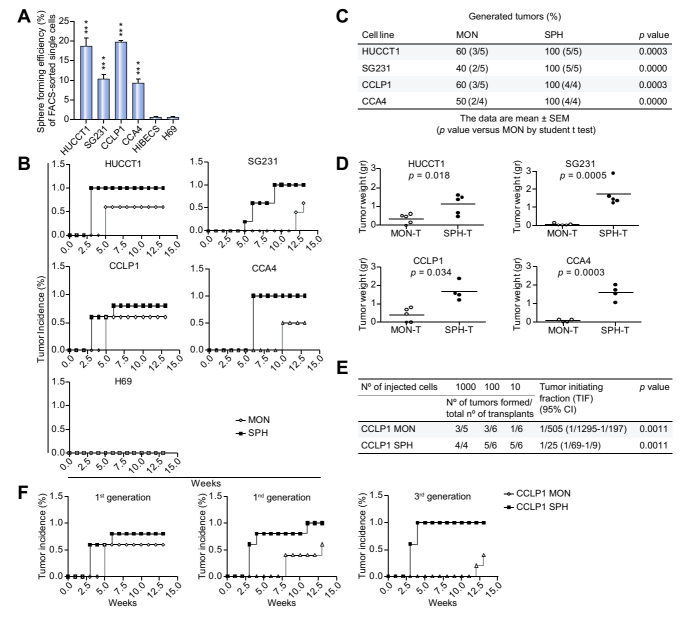
CCA stem-like cells were identified by using a functional tool of 3D culture system based on defined serum-free medium, as shown in many patient-derived tumors [21,22]. First, established (SG231, HUCCT1, CCLP1) [23–25] and primary human intrahepatic CCA-derived (CCA4) cell lines [26] were tested for sphereforming efficiency (SFE) as a representation of tumor stem-like subset. To this end, CCA fluorescence-activated cell sorting (FACS)-sorted single cells were cultured in 3D non-adherent condition in a 96-well plate. Importantly, single-cell sorting prevents formation of cell-aggregation and avoids large fraction of nonstem cells to form clusters. As expected, only malignant cholangiocytes retained sphere-forming potential in contrast to normal

The Humanitas Animal Care and Research Advisory Committee approved housing and experimental animal procedures. NOD/SCID/IL2r $\gamma$  null (NSG) mice (Jackson

counterparts such as normal immortalized cholangiocytes (H69) and human intrahepatic biliary epithelial cells (HiBECs) (Fig. 1A). However, CCA spheres (SPH) revealed a high grade of heterogeneity in terms of efficiency (HUCCT1 and CCLP1, approximately 20%; SG231 and CCA4, 10%), size and morphology (Supplementary Fig. 1A).

Next, an extensive characterization, including evaluation of *in vivo* tumorigenicity in immune-deficient NOD-SCID/*IL2Rg*<sup>null</sup> (NSG) mice, drug-responsiveness and gene expression analysis of CCA-SPH, was performed.

To assess CCA-SPH ability to generate tumors, spheres formed by HUCCT1, SG231, CCLP1 and CCA4 were dissociated and 1000 cells injected s.c. into NSG mice (Fig. 1B–D) [27]. Consistently with their *in vitro* SFE (Fig. 1A), CCA-SPHs were highly tumorigenic and engrafted 100% of transplanted mice in all examined cell lines albeit with differences in tumor size (Fig. 1B–D). In contrast to CCAs grown in monolayer condition (MON), including H69-MON, CCA-SPHs were responsible for early tumor initiation and formation (3 weeks for HUCCT1, CCLP1; 5 weeks for SG231; 6 weeks for CCA4) during 13 weeks of observation. In addition,



**Fig. 1. CCA self-renewal capacity** *in vitro* **and tumorigenic potential** *in vivo.* (A) Sphere-forming efficiency (SFE) of CCAs (HUCCT1, SG231, CCLP1, CCA4) and normal cholangiocytes (HiBECs, H69) calculated by dividing the sphere number by number of single cells seeded and expressed as a percentage. Mean  $\pm$  SEM (n = 4, *p* value vs. H69 and human intrahepatic biliary epithelial cells (HiBECS) by Student's *t* test, \*\*\**p*  $\leq$  0.001). Sphere tumorigenic capacity in NSG mice: (B) Tumor growth kinetic (n = 5), (C) frequency and (D) weight of generated tumors at 13 weeks after subcutaneous injection into NSG mice of 1000 SPH/MON isolated cells as monitored by weekly palpation. Mean  $\pm$  SEM (*p* value vs. MON-T by Student's *t* test). (E) Self-renewing CCLP1 cells calculated by ELDA program (see Methods section) (*p* value vs. MON tumor-initiating fraction (TIF) by Student's *t* test). (F) Serial transplantations of 1000 CCLP1 SPH/MON cells into flanks of NSG mice (n = 4).

Journal of Hepatology 2017 vol. 66 | 102-115

SPH-derived tumors (SPH-T) showed a large difference in terms of tumor weight and size compared to CCA-MON derived tumors (MON-T) (Fig. 1D), indicating that SPH-T retained high tumorigenic potential when xenotransplanted in animal hosts and represent an important source of de novo tumor growth. The most common method to determine frequency of self-renewing cells within tumors is a limiting dilution cell transplantation assay. Thus, due to slight differences in tumor potential between SPH and MON in CCLP1, we specifically performed limiting dilution transplantation assay in NSG mice. After s.c. injection of 1000, 100 or 10 cells of both CCLP1-MON and SPH conditions, data analysis using ELDA software [28,29] revealed a significant 20.3-fold increase in the absolute number of tumor-initiating fraction (TIF) within CCLP1-SPHs (TIF = 1/25) compared to MONs (TIF = 1/505) (*p* = 0.0011, Fig. 1E). Indeed, as few as 10 SPH cells produced tumors, whereas 1000 MON cells gave rise to fewer tumors at 13 weeks (Fig. 1E) confirming that SPHs were remarkably enriched in self-renewing CCA-propagating cells. Additionally, since self-renewing stem-like cancer cells have an unlimited ability to promote tumor growth, capability for serial transplantation was also tested (Fig. 1F). Cells were re-isolated from CCLP1-T established from both 1000 SPH and MON cells, propagated in short-term cultures and re-transplanted into secondary recipient mice. Notably, SPHs not only sustained tumorigenic potential in serial transplantations but also progressively improved in tumor frequency. Conversely, corresponding MONs showed either dramatic increase in tumor latency and decrease in tumor incidence at later generations (Fig. 1F). Together these data indicated that, in accordance with long-term self-renewal potential, SPHs significantly maintain tumorigenic potential through diverse xenograft-generations while MONs reduced it.

Furthermore, because drug-resistance represents a key feature of cancer stem/initiating cells, 3D cultures were tested for susceptibility to chemotherapeutic agents. Indeed, in presence of common anti-CCA drugs such as cisplatin, 5-fluorouracil, oxaliplatin and gemcitabine, CCA-SPH revealed a better survival rate compared to parental CCA-MONs, as measured by proliferation MTT assays and indicated by the 50% inhibitory concentration values (IC50) (Supplementary Fig. 1B). Although a remarkable heterogeneity in terms of drug-sensitivity, SPH-resistance was a common phenomenon observed across all CCA lines.

Next, to corroborate our in vivo and in vitro findings, an extensive molecular characterization of CCA-SPH was achieved by PCR arrays specific for CSC, liver cancer and embryonic stem cell (ESC) pathways. In accordance with the observed heterogeneity in SFE, tumor potential and drug-resistance, clustering analysis of differentially expressed genes for each pathway revealed SPH geneenrichment as compared to respective MONs, specific for each cell line (Fig. 2A-C; Supplementary Fig. 2A). Indeed, CCLP1, HUCCT1 and CCA4 SPHs presented augmented expression for CSC- (55-69%) and liver cancer-related (61-77%) genes, whereas SG231-SPH retained the highest fraction of ESC-related (37%) genes (Supplementary Fig. 2A). However, a core of 30 common dysregulated genes in all CCA-SPHs was identified. Specifically, expression of 23 CSC-genes, including key molecules for pluripotency and self-renewal (SOX2, POU5F1, KLF4, BMI1, NOTCH1, MYC, KITLG, LIN28A, MAML1, YAP1), drug-resistance and survival (ABCG2, LATS1, NFKB1, NOS2), metastasis (TGFBR1, BPM7, FGFR2) and stem-like surface markers (CD24, CD44, PROM1, THY1, EPCAM), resulted enhanced. Moreover, this common gene-set revealed overexpression of 6 hepatic oncogenic drivers (CDKN1A,

## JOURNAL OF HEPATOLOGY

*BCL2L1, CTNNB1, IGF2, ITGB1, LEF1*) and key ESC factors such as *HNF4* (Fig. 2B), in addition to the well-described liver CSC markers CD13 and LGR5 (not included in the arrays) (Fig. 2C). Accordantly, further confirmation of stem cell content was considered at protein level by FACS-analysis. Indeed, superior expression of CD13, CD44, THY1 and EPCAM were verified in CCA-SPHs from four cell lines (Fig. 2D; Supplementary Fig. 2B) thus suggesting that 3D cultures were effectively enriched for tumorigenic cells endowed with stem-like molecular traits.

Particularly, presence of the most significant CSC surface markers, such as *CD44*, *PROM1*, *THY1* and *EPCAM* was validated by CCA transcriptome of 104 patient database [1] (Supplementary Fig. 2C).

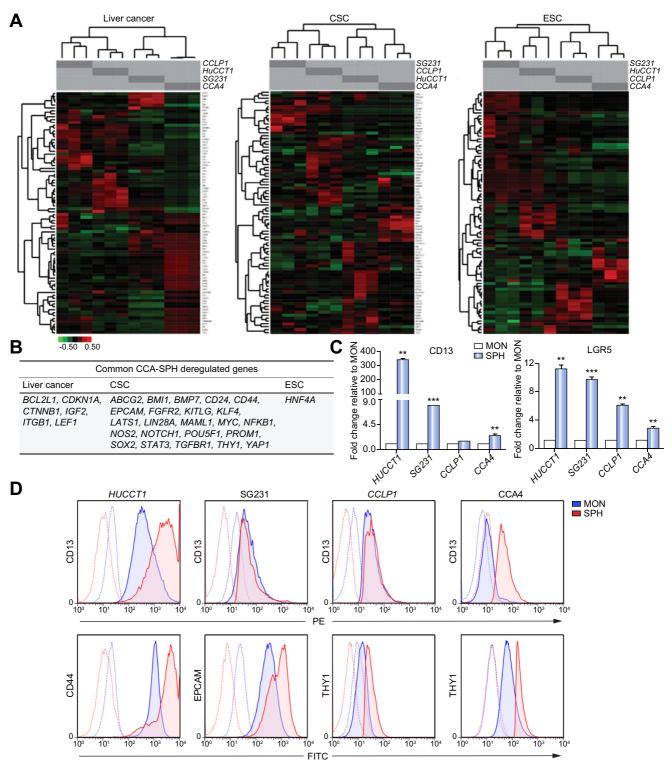
Whereas gene expression profile was consistent with stem-like traits of SPH-condition, we also tested the strength of 3D culture system compared to antigenic approach. To this end, primary CCA4 was immune-sorted for standard stem-related marker THY1 and verified for in vitro sphere-ability. However, no significant differences in in vitro self-renewing capability were found between thymocyte differentiation antigen (THY)<sup>+</sup> and parental cells as control (Supplementary Fig. 2D). Next, we combined expression of stem-like markers to sphere-forming capacity, thus THY<sup>+</sup> cells were grown in 3D condition (THY<sup>+</sup> SPH) and tested for tumorigenic potential. Notably, in vivo analysis revealed similar TIF subset between  $THY^+$  SPH (TIF = 1/47.8) and parental SPH (TIF = 1/35.7) (*p* = 0.2028; Supplementary Fig. 2D, right) also confirmed by expression of liver cancer, CSC and ESC-like markers (Supplementary Fig. 2D). These results indicated that 3D culture system potentially represents a reliable tool and valid alternative to the antigenic approach for CCA stem-like cell selection.

Altogether, this broad functional and molecular characterization proposed the existence of a stem/progenitor-like compartment in human CCA identified by 3D culture system. Indeed, CCA-SPHs exhibited typical stemness hallmarks, such as upregulated stem cell biomarkers, drug-resistance and enhanced capability to initiate malignancy.

CCA stem-like compartment educates macrophage precursors towards acquisition of CSC-associated TAM phenotype

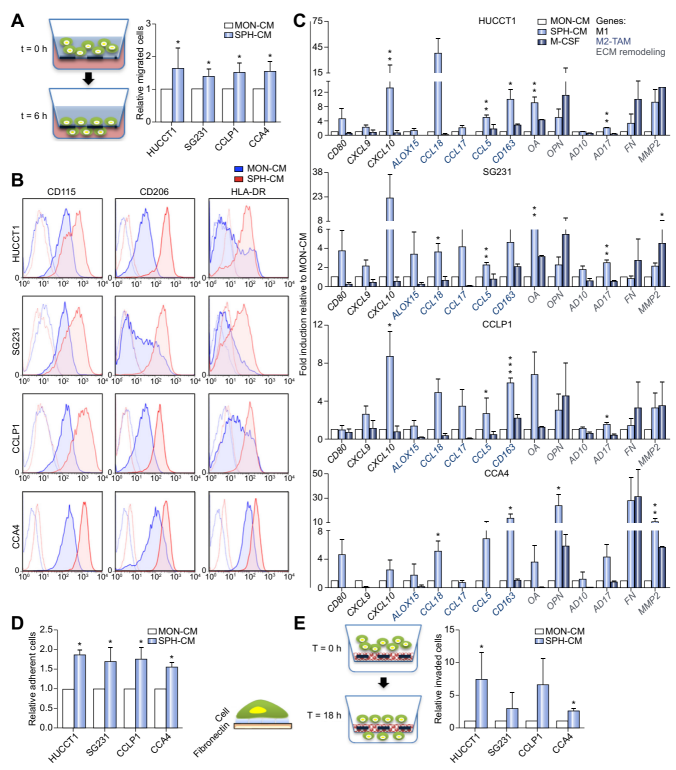
In addition to diverse cellular components and extracellular matrix (ECM), CSC-milieu is enriched with tumor cell products such as growth factors, cytokines, prostaglandins and diverse protein factors [30]. Thus, to mimic a bioactive CSC-niche, freshly isolated healthy donor circulating monocytes (CD14<sup>+</sup>) were exposed to CCA-SPH conditioned medium (SPH-CM) and tested for 1) chemotactic recruitment based on migration properties and 2) macrophage-differentiation by analysis of marker expression and functional properties (Fig. 3; Supplementary Fig. 3A). First, by using a chemotaxis chamber, a higher number of CD14<sup>+</sup> migrating towards SPH-CM gradient than MON-CM was recorded within 6 h, suggesting that SPH-CM acted as a strong monocyte attractor (Fig. 3A). Hence, greater amount of recruited mononuclear cells by CCA-SPHs maybe find a potential explanation in creating a tumorigenic-niche supporting CSCmaintenance.

It is widely accepted that macrophages constitute an extremely heterogeneous population. Most macrophages originate from blood monocytes, which differentiate into distinct macrophage types, schematically identified as M1 (or classically activated) and M2 (or alternatively activated). It is now generally



**Fig. 2. Molecular properties of CCA spheres.** (A) Pathway-focused qRT-PCR arrays. Hierarchical clustering distinguished each CCA-SPH type based on significant gene expression compared to CCA-MON. Data first centered and normalized, then clustered using centered correlation metrics with complete linkage. Dendrograms depict similarity of individual genes and cases. Right side indicates clusters of coordinately expressed genes with higher expression levels in CCA-SPH than CCA-MON. Relative gene expression level depicted according to the scale bar. (B) List of commonly upregulated genes in SPH of all CCA cell lines. Genes divided according to pathway of belonging. (C) Relative expression of *CD13* and *LGR5* transcript-encoding markers. glyceraldehyde 3-phosphate dehydrogenase (*GAPDH*) as internal control. All mRNA levels are presented as fold changes normalized to 1 (mean expression of monolayer). Mean  $\pm$  SEM (n = 3, p value vs. MON by Student's t test, \*p  $\leq 0.05$ , \*\*p  $\leq 0.01$ , \*\*\*p  $\leq 0.001$ , (D) FACS-profile of CD44, CD13, THY, EPCAM.

Journal of Hepatology 2017 vol. 66 | 102-115



**Fig. 3. Effect of CCA stem-like cells on macrophage precursors.** (A) Migrated CD14<sup>+</sup> towards SPH-CM (6 h). Relative migrated CD14<sup>+</sup> cells normalized to migrated CD14<sup>+</sup> in presence of MON-CM. Mean ± SEM (n = 3, p value vs. MON). (B) FACS-profile of CD115, HLA-DR and CD206 expression in macrophages obtained by culture of CD14<sup>+</sup> with SPH- and MON-CM. Histograms represent three independent experiments with macrophage from three different healthy donors. (C) Relative expression of M1/M2 and matrix remodeling-related genes. As control macrophage-differentiated with M-CSF. *GAPDH* as internal control. All mRNA levels presented as fold changes normalized to 1 (n = 3). (D) Adhesion assay using FN-supports. Cells counted and normalized to MON macrophages (n = 3). (E) SPH macrophage-invasion assay using Matrigel-coated transvells. Cells counted and normalized to migrated MON macrophages (n = 5). Mean ± SEM (p value vs. MON macrophage by Student's t test, \* $p \in 0.05$ , \* $^*p \in 0.01$ , \*\*\* $p \in 0.001$ ).

recognized that TAMs show an M2-like cancer-promoting phenotype [31–33].

To determine the effect of the CCA stem-like component on macrophage precursors, healthy donors CD14<sup>+</sup> were cultured in presence of CCA-SPH-CM, and after 6 days, investigated for macrophage-differentiation and polarization. Consistently with observed changes in morphology (data not shown), by using a panel of conventional markers typically used for M0 classical differentiation (CD115, macrophage colony-stimulating factor receptor or M-CSF receptor), M1 (HLA-DR, human leukocyte antigen and major histocompatibility complex class II) and M2-TAM (CD206, macrophage mannose receptor) polarization (Supplementary Fig. 3B), FACS-analysis revealed that in presence of CCA-SPH-CM, in vitro-educated CD14<sup>+</sup> expressed higher CD115 level in contrast to CD14<sup>+</sup> exposed to MON-CM or M-CSF, which acted as control for M0 classical differentiation (Fig. 3B; Supplementary Table 1). As expected and regardless culture conditions, CCA-CM induced macrophage-differentiation of human monocytes as indicated by CD14<sup>+</sup> CD68<sup>+</sup> cells in both MON and SPH-CM (Supplementary Fig. 3C and D). Notably, more than 94% of SPH-educated CD14<sup>+</sup> expressed CD14, CD68 and CD115 markers (Fig. 3B; Supplementary Fig. 3C and D). Moreover, differentiation towards neutrophil lineage and dendritic cells was excluded by CD66b and CD1a staining [34,35], also confirmed in MOmacrophages (Supplementary Fig. 3E and F). Furthermore, CCA-SPH activated-macrophage (SPH macrophage) expressed greater levels of CD206 and HLA-DR TAM-like markers in contrast to CCA-MON activated-macrophage (MON macrophage) (Fig. 3B; Supplementary Table 1). All these data indicated that both MON- and SPH-CM induced macrophage-differentiation, but SPH-CM was responsible for macrophage-acquisition of a specific CSC-dependent phenotype.

Likewise, in *in vitro*-educated macrophages, expression genes belonging to M1 (e.g., *CD80*, *CCL5*, *CXCL9*, *CXCL10*), M2 (*ALOX15*, *CCL18*, and *CCL17*) and TAM-like (*CD163*) categories, including genes involved in ECM remodeling and adhesion such as osteoactivin (*OA*), fibronectin (*FN*), osteopontin (*OPN*), metalloproteinase ADAM (*AD10*, *AD17*), and matrix metalloproteinase-2 (*MMP2*), were determined (Fig. 3C). Notably, SPH macrophages retained unique molecular features with concomitant expression of TAM-like (e.g., *CD163*), M1-like (e.g., *CCL5*, *CXCL9*, *CXCL10*), M2like (e.g., *CCL17*, *CCL18*) and matrix remodeling-related (e.g., *OA*, *AD17* and *MMP2*) genes (Fig. 3C).

Consistent with our gene expression data, SPH macrophages possessed greater adhesion ability on FN-support (Fig. 3D) and better invasion capacity as shown by *in vitro* Matrigel-coated transwell assays (Fig. 3E) [36] compared to MON macrophages. Both these features are likely associated with dynamic properties of TAM-like cells within tumor-initiating niche because of their incessant and deregulated activity regarding ECM-reorganization and turnover. Owing to their potential ability to remodel tumor stroma, SPH macrophages may invade and at the same time adhere to the ECM-component, thus providing sustenance to tumorenvironment. This result showed concordance with the evidence that mononuclear phagocytes support stem cell functions, thus contributing to tissue repair and remodeling [37].

CCA humanized-mice recapitulate in vitro-educated-macrophage traits

To translate our *in vitro* data in a human-like setting, we analyzed the *in vivo* immunological response of human mononuclear cells

to CCA stem-like component at tumor site. Thus, a cotransplantation of human PBMC (hPBMC) and patient-derived CCA4-SPHs into NSG mice was performed (Fig. 4A-B; Supplementary Fig. 4A). As already extensively demonstrated, NSG mice bearing a targeted mutation in *IL2r* gene (responsible for a severe defective murine immune component) support considerably high levels of human hematopoietic and lymphoid cell engraftment providing an exciting substitute for human immunobiology studies [16]. Therefore, CCA4-SPH-T bearing mice (when tumor size reached 50 mm<sup>3</sup>, 51 days after s.c. injection) were engrafted with three successive doses of hPBMC ( $15 \times 10^6$ /dose) intravenously (i.v) in the retro-orbital sinus. Efficiency of human engraftment was tested 6 days after first hPBMC dose and presence of human circulating mononuclear cells (mCD45<sup>-</sup> hCD45<sup>+</sup> hCD14<sup>+</sup>) was confirmed in mouse peripheral blood by FACSanalysis. In agreement with published data [16], frequency of mCD45<sup>-</sup> hCD45<sup>+</sup> hCD14<sup>+</sup> circulating cells were almost 0.7% indicating an effective human cell engraftment (Supplementary Fig. 4B and C).

At the end of experiment, both CCA4-SPH-T and CCA4-MON-T were removed and FACS-analyzed for human mononuclear cells presence. Well-matched with *in vitro* data, CCA4-SPH-T retained higher number of human CD45<sup>+</sup> CD68<sup>+</sup> (83.55%) in comparison with CCA4-MON-T (70%) (Fig. 4A). Again, infiltrated human CD45<sup>+</sup> CD14<sup>+</sup> in CCA4-SPH-T displayed enhanced expression of M1-like (e.g., *CD80*, *CXCL9*), M2-like (e.g., *CCL18*, *CCL17*) and matrix remodeling-related genes (e.g., *OPN*, *AD10*, *AD17*, *MMP2*, full-length FN and oncofetal FN (*ED-B*)) (Fig. 4B).

Collectively, our *in vitro* and *in vivo* observations indicated that SPHs recruit higher number of mononuclear cells at tumor site and CCA-SPH macrophages retain a unique phenotype compared to MON macrophages.

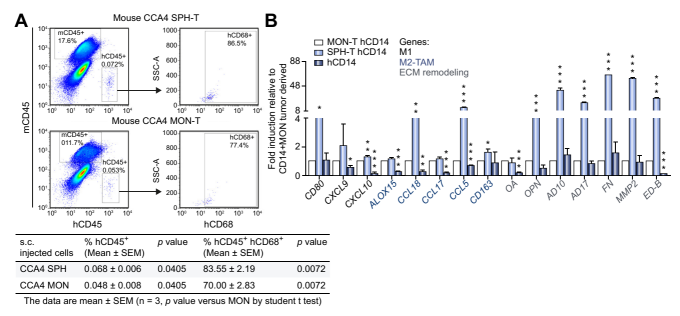
### *Mixed phenotype of infiltrating macrophages in CCA patients*

Since a broad analysis of human CCA-TAMs has not been provided yet, we corroborated our in vitro data with a global investigation of CCA patient-associated macrophages. Abundance, phenotypes and distribution of macrophages residing in CCA intratumoral and peritumoral tissue were evaluated via immunohistochemistry (IHC). Paraffin-embedded tissue samples from 23 CCA patients were tested for tumor-promoting macrophages (CD163) presence. An immunohistochemical analysis of tumor sections showed that CD163<sup>+</sup> cells were more highly expressed in tumor than peripheral tumor region (Fig. 5A; Supplementary Table 2). Remarkably, analysis of macrophage-distribution within tumor lesion revealed that the largest proportion of CD163<sup>+</sup> was located in the tumor front (Fig. 5A, lower). On further evaluation of relationships between TAM-marker density and clinical pathological features, we found that CD163 progressively increased along with tumor grade (G2/G3 > G1) and was significantly associated with CCA pathological grade (p = 0.006) (Fig. 5B, Supplementary Table 2) as well as CA19.9 serum marker (Fig. 5C).

Likewise, FACS-profiling of mononuclear subsets from fresh resected CCA samples confirmed CD115, CD206 as well as HLA-DR expression of infiltrating CD68<sup>+</sup> cells (Fig. 5D; Supplementary Fig. 5A–D). In addition, isolated CD14<sup>+</sup> cells from fresh CCA tissue [16] revealed expression of M1-like (e.g., *CD80, CXCL9*) and M2-like (e.g., *CCL18*) markers as well as genes involved in ECM remodeling including *OA, FN* and *ED-B* (Fig. 5E).

Similarly, transcriptome of laser capture micro-dissected epithelium (EPI) and stroma (S) from 23 CCA patients revealed

Molecular and Cell Biology



**Fig. 4. Evaluation of TAM-infiltration in mouse model.** (A) FACS-profile of mouse CD45<sup>-</sup>, human CD45<sup>+</sup> CD68<sup>+</sup> cells in isolated mononucleate subset from CCA4-SPH-T and CCA4-MON-T. Representative dot plots and table with percent of human CD45 CD68 double positive cells determined by FACS. Data are mean  $\pm$  SEM (n = 3, *p* value *vs.* MON-injected mice by Student's *t* test). (B) Relative gene expression of human CD14<sup>+</sup> isolated from CCA4-T. As control, human CD14<sup>+</sup>. Human *GAPDH* as internal control. All mRNA levels presented as fold changes normalized to 1 (mean expression of CCA4 MON-T CD14<sup>+</sup> cells). Mean  $\pm$  SEM (n = 3, *p* value *vs.* MON-T CD14<sup>+</sup> cells). Student's *t* test, \**p*  $\leq 0.05$ , \*\**p*  $\leq 0.01$ , \*\*\**p*  $\leq 0.001$ .

a significant deregulation of CD115, HLA-DR, CXCL9, MMP2 and CD206 in CCA-S compared to CCA-EPI compartment (Fig. 5F; Supplementary Fig. 5E). Further validation was provided by tumor (T) transcriptome *vs.* surrounding liver tissue in an independent data set of CCA patients (n = 104) (Supplementary Fig. 5F).

Furthermore, IHC analysis of CCA sections revealed a colocalization of both CD163 positive elements together with expression of CSC-related markers (CD44 or EPCAM) strongly supporting the potential relationship of infiltrating TAMs with CSC-niche (Fig. 5G).

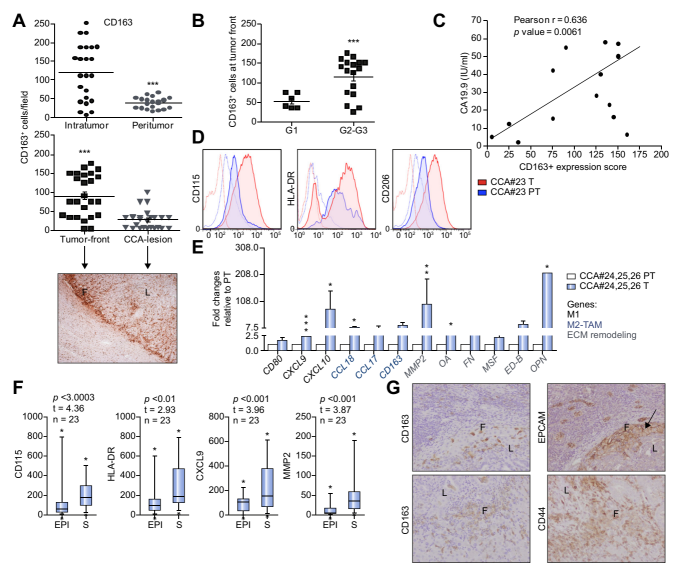
Therefore, our comprehensive characterization showed that phenotype of CCA patient macrophages retained mixed M1-M2 features, further strengthening our *in vitro* results.

### CCA-SPH secretory profiling specifically involved in macrophagedifferentiation

Intriguingly, IHC data revealed that SPH-derived tumors retained a prominent and well-defined stromal compartment attested by  $\alpha$ -SMA staining, massive collagen presence (Sirius Red) and abundance of tumor-associated CD31-positive vessels as well murine F4/80 tumor-macrophages (Fig. 6A). This evidence suggested that SPH cells acted differentially in creating their associated surrounding microenvironment.

Furthermore, we tested the diverse functionality of CCA cells on the surrounding niche, in particular lymphocytes as well as endothelial cells. Thus, CellTrace<sup>TM</sup> 5(6)-Carboxyfluorescein Nhydroxysuccinimidyl ester (CFSE) labeled CD4 peripheral blood lymphocytes (CD4<sup>+</sup> T) were cultured in presence of both MON and SPH-CM. It was found that SPH-CM were not able to induce CD4<sup>+</sup> T cell proliferation, unlike MON-CM did (Fig. 6B) suggesting a more immunosuppressive properties of SPH-associated microenvironment [38,39]. Since angiogenesis is important hallmark of tumor progression, we evaluated the biological activity of CCA-CM on the induction of *in vitro* capillary-like structures [40]. By using the well-known Matrigel angiogenesis assay, the effects of SPH-CM and MON-CM on human umbilical vein endothelial cells (HUVEC) cells in term of capillary tube-like formation were examined *in vitro*. In accordance with IHC data of SPH tumor-derived tissues (Fig. 6A), HUVEC cells treated with SPH-CM for 24 h revealed the highest neovascular response in comparison to MON-CM treatment (Fig. 6C; Supplementary Fig. 6A).

Hence, based on this result, we investigated the presence of specific soluble mediators released by SPHs using enzymelinked immunosorbent assay (ELISA) profiling of conditioned media. Among 37 tested molecules (chemokines, cytokines, interleukins) specifically involved in macrophage-differentiation, activation and recruitment, several factors were secreted at extremely low levels (<50 pg/ml), whereas 16 showed significant amount in both CCA-MON and SPH-CM (Fig. 6D, Supplementary Table 3). A 3-level analysis identified 1) common SPH and MON, 2) MON-specific, 3) SPH-specific released factors. First, we found that both SPH and MON produced high vascular endotheial growth factor (VEGF) concentrations. Therefore, VEGF role in driving macrophage-differentiation was in vitro investigated. For that reason, healthy donor CD14<sup>+</sup> were cultured for 6 days in presence of VEGF added to fresh monocyte medium, not tumor conditioned (Supplementary Fig. 6B). Clearly, FACSprofile reveals enriched expression of CD68 and CD115 as classical macrophage markers (Supplementary Fig. 6B) suggesting its contribution in CCA macrophage-differentiation instead of classically described M-CSF. Notably, both SPH and MON macrophages overexpressed corresponding VEGF receptors (VEGFR1-3) (Supplementary Fig. 6C).

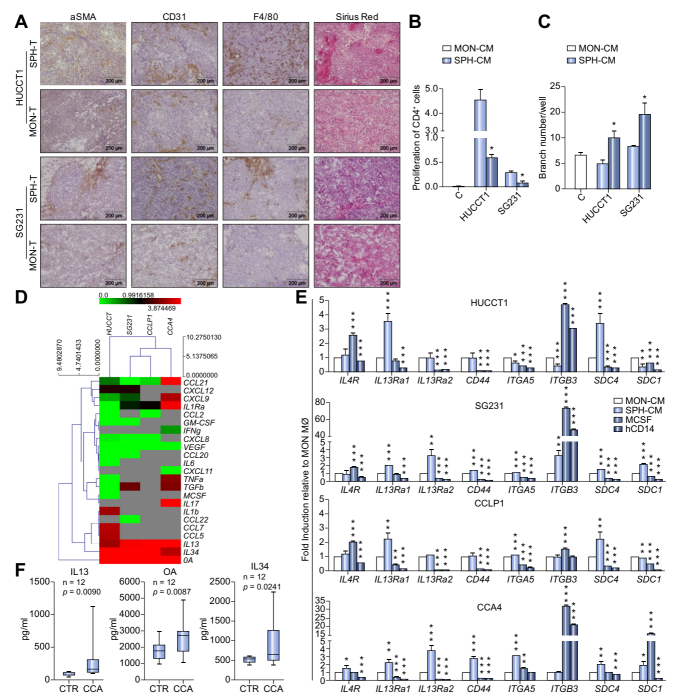


**Fig. 5. Evaluation of TAM-infiltration in human CCA samples.** (A) Quantification of CD163<sup>+</sup> cells in both CCA intratumoral (T) and peritumoral (PT) regions. Mean  $\pm$  SEM (n = 23, *p* value vs. PT). Distribution of CD163<sup>+</sup> cells in CCA lesion (L) and tumor front (F) shown by a representative image and corresponding quantification. Mean  $\pm$  SEM (n = 23, *p* value vs. F). (B) Correlation of % CD163<sup>+</sup> cells tumor grade (G) (n = 25, Supplementary Table 2, one-sided Student's *t* test applied to log ratios in order to compare G1 to G2-G3). (C) Cancer antigen 19.9 serum levels in CCA patients (n = 17, Supplementary Table 2, Pearson correlation between two parameters was calculated using R and the cor.test function, yielding correlation coefficients and *p* values). (D) Expression of CD115, HLA-DR and CD206 in CCA-infiltrated macrophages by FACS included both CCA T and PT regions. Representative histograms of CCA#23 patient. (E) Relative expression of M1/M2 and matrix remodeling-related genes. *GAPDH* as internal control. All mRNA levels displayed as fold changes normalized to 1 (mean expression of PT-macrophages). Histograms represent the average of three different CCA patients (#24, #25, #26 patients). Mean  $\pm$  SEM (n = 3, *p* value vs. PT). (F) Gene expression evaluated in a set of CCA patients (n = 23) where paired intratumoral epithelial (EP1) and stromal compartments (S) obtained by laser micro-dissection. Student's *t* test, \**p*  $\leq 0.01$ . (G) Representative images for IHC analysis of CD163 and CSC-related markers (CD44, EPCAM) in CCA sections (20x) (CCA lesion (L) and tumor front (F)).

Nevertheless, a cell line-specific MON secretory profile was identified, with altered supernatant levels of GM-CSF, M-CSF, TNF $\alpha$ , transforming growth factor  $\beta$  (HUCCT1), GM-CSF, CCL22, CCL20, CXCL8 (SG231), CCL2, and CXCL18 (CCLP1 cells), CXCL11 (CCA4) (Supplementary Table 3). However, release of soluble mediators was cell line-dependent and no common molecules among CCA-MONs were recognized.

Crucially, a specific secretory SPH-profile consisting of IL13, IL34 and OA was identified in all tested cell lines (Fig. 6D; Supplementary Fig. 6D) and more importantly SPH macrophages overexpressed receptors for IL13 (*IL13Ra1*, *IL13Ra2*, *IL4R*), OA (*CD44*, *ITGA5*, *ITGB3*, *SDC1*, *SDC4*) (Fig. 6E) as well as for IL34 (CD115) already shown in Fig. 3B. Therefore, these data proposed the potential implication of IL13, OA and IL34 in the macrophage-acquisition of CSC specific phenotype.

Surprisingly, elevated levels of circulating IL13, OA and IL34 were significantly found in CCA patient group (n = 12) compared to healthy control subjects (n = 12), proposing a potential association with CCA-disease (Fig. 6F, Supplementary Table 4). However, due to limited availability of human CCA specimens, important question such correlation between serum level and abundance of TAMs as well clinical pathological features of CCA



**Fig. 6. SPH-specific production of bioactive molecules.** (A) Representative images show immunohistochemistry of  $\alpha$ -SMA, CD31, F4/80 and Sirius Red staining for collagen on SPH and MON tumor sections. Scale bar: 200 µm. (B) Effect of SPH- and MON-CM on CD4<sup>+</sup> cells proliferation. CD4<sup>+</sup> without stimulation used as control. Data presented as % of proliferating CD4<sup>+</sup> cells. Mean  $\pm$  SEM (n = 3, *p* value vs. MON-CM by Student's *t* test,  $\pm p \approx 0.05$ ,  $\pm p \approx 0.001$ ,  $\pm p \approx 0.001$ , (C) Effect of SPH- and MON-CM on CD4<sup>+</sup> cells proliferation. CD4<sup>+</sup> without stimulation ability of HUVEC tube formation. HUCCT1 and SG231 MON- and SPH-CM were added to the medium of HUVEC to assay their effect on the tube formation ability of HUVEC. Data presented as number of branches/well. Mean  $\pm$  SEM (n = 3, *p* value vs. MON-CM by Student's *t* test,  $\pm p \approx 0.05$ ,  $\pm p \approx 0.001$ ). (D) Heat map representation of soluble mediators released by SPH and MON (ELISA). Concentration as pg/ml. Molecules clustered with names shown on right of heat map. Each raw corresponds to a single compound, and each column represents an independent condition. Heat map color scale corresponds to relative molecule expression (on the top, minimum and maximum of all values). Results are average of three independent experiments. (E) Relative expression of transcript-encoding receptors for IL4 (*IL4-R*), IL13 (*IL13Ra1*, *IL13Ra2*), OA (*CD44*, *ITGA5*, *ITGB3*, *SDC4*) and IL34 (*SDC1*) in SPH and MON macrophage. CD14<sup>+</sup> cells as well M-CSF derived macrophage, as controls. GAPDH as internal control. All mRNA levels are mean  $\pm$  SEM (p value vs. MON macrophage by Student's *t* test,  $\pm p \ll 0.001$ ,  $\pm p \ll 0.001$ ). MØ, macrophage.

Table 1. Correlation of SPH stem-like genes and OA, IL34 and IL13 in clinical CCA (n = 104 patient tumors) using a microarray database (1).

	OA	
SPH genes	R	p value
BMI1	0.3504	0.0003
CD44	0.3844	0.0001
CTNNB1	0.4419	0.0000
KITLG	0.3975	0.0000
KLF4	0.3603	0.0001
LEF1	0.6321	0.0000
THY1	0.5118	0.0000
	IL34	
MAML1	0.3423	0.0004
THY1	0.2966	0.0022
	IL13	
LIN28A	0.4906	0.0000
POU5F1	0.2685	0.0059
SOX2	0.3121	0.0013

Pearson correlation between gene pairs was calculated using R and the "cor.test" function, yielding correlation coefficients and *p* values.

patients was not addressed. Nonetheless, correlative data were provided by a well-described transcriptome database of 104 CCA patients [1]. Amazingly, Pearson correlation coefficient test showed significant associations of specific CCA-SPH released molecules (*IL13, OA, IL34*) with SPH stem-like genes such as *BMI1, CTNNB1, CD44, KITGL, KLF4, LEF1, LIN28A, MAML1, POUF5, SOX2 and THY1* (Table 1) reinforcing the importance of CCA stem-secretory profiling in tumor context in CCA patients. Furthermore, intimate connection between CSCs and tumormacrophages in human CCA was also denoted by meaningful correlations between stem-like SPH and SPH macrophage-molecular traits (Supplementary Table 5).

IL13, IL34, and OA are required for acquisition of CSC-like TAM identity

To confirm the contribution of IL13, OA, IL34 in shaping molecular and functional aspect of SPH macrophages, inhibition of these molecules by addition of single or combination neutralizing antibodies to SPH-CM (previously in vitro tested, to define an effective and non-toxic concentration, Fig. 7; Supplementary Fig. 7A), was tested in vitro and in vivo. Strikingly, our results indicated a downregulation of CD115, CD206 and HLA-DR proteins (Fig. 7A; Supplementary Table 6) as well as gene expression (CXCL9, CXCL10, CCL18, CCL5, CD163, OPN and AD17) (Fig. 7B) in addition to loss of functional properties such as migration, macrophage-invasion and adhesion (Fig. 7C) suggesting a shift to MON macrophage phenotype. Particularly, each single IL13, IL34, or OA seemed to be independently involved in CD115 (Fig. 7A). More importantly, only OA molecule is accountable for CD206 and HLA-DR expression (Fig. 7A) as well as macrophage-invasion and adhesion (Fig. 7C), whereas combination of neutralizing antibody has a major impact in diminishing the expression of M1, M2 and ECM remodeling genes (Fig. 7B).

Well-matched with SPH-CM inhibition, the addition of these molecules to MON-CM reproduced same SPH macrophage differentiative impact. Indeed, IL13, IL34, and OA were able to restore SPH macrophage-like profile with overexpression of CD115, HLA-DR and CD206 markers and presence of an equal amount of CD68 protein (Fig. 7A; Supplementary Fig. 7B and C, Supplementary Table 6). Remarkably, our data displayed the reexpression of specific gene-set (*CXCL9, CXCL10, CCL18, CCL5, CD163, OPN* and *AD17*) (Fig. 7B; Supplementary Fig. 7D) as well as re-acquisition of functional properties such as monocyte recruitment, macrophage-invasion and adhesion (Fig. 7C; Supplementary Fig. 7E and F). Pertinently, all these results pointed to a peculiar contribution of these molecules in driving SPH macrophage-differentiation. Thus, we concluded that although the underlying mechanism need to be explored, combination of IL13, OA, IL34 directly trigger differentiation of monocytes into stem-like TAM-subtype.

Furthermore, to investigate the significance of diverse macrophage-subsets in supporting tumorigenicity, three diverse *in vitro*-educated macrophage-subsets such as 1) SPH macrophage, 2) MON macrophage, 3) MON (+ IL13 + OA + IL34) macrophage, 4) SPH (+ single or combination of antibodies against IL13, OA, IL34) macrophage were co-injected with CCA-MONs into NSG mice. As expected, MON macrophages enhanced tumor growth of MON cells injected alone as already described in diverse study for TAMs. Outstandingly, SPH macrophages boosted tumorigenic potential compared to MON macrophages, indicating that CSC-associated TAMs increased *in vivo* tumor-promoting effect of CCA-CSCs. As confirmation, co-injected MON with (+ IL13 + OA + IL34) macrophages reproduced similar SPH macrophage tumorigenic effect (Fig. 7D), proving direct functional relation-ship between CSC TAMs and tumor outcome.

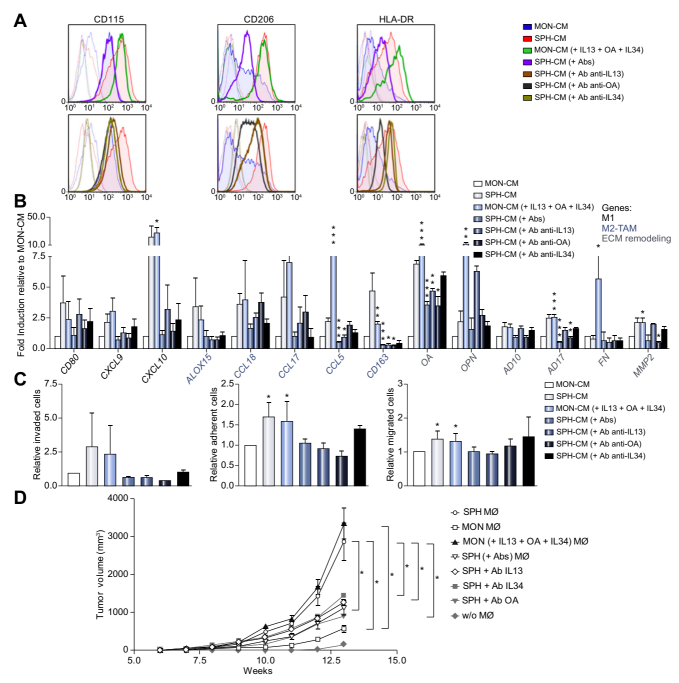
### Discussion

Current CCA treatment strategies are largely ineffective. Therefore, understanding the pathogenesis of this disease is essential for identifying potential curative targets. Concept of stemnessdriven carcinogenesis highlights the existence of a therapeutically challenging subset responsible for tumor initiation, dissemination and drug-resistance termed CSCs. Our study mainly focused on exploration of stem-compartment in human intrahepatic CCA by using a 3D culture system as a well-known functional tool to select and enriched for tumor stem-like cells. Indeed the absence of CCA-CSC specific markers limits the use of classical antigenic approach. Thus, by using *in vitro* spherogenicity and *in vivo* tumorigenicity as criteria of stemness properties, for the first time we provided extensive molecular and functional evidence for CCA stem-like compartment identified by 3D cultures.

Secondly, here we proposed that tumor stem-like cells might model their immunopathological-niche according to their necessities. Indeed, we focused on CSC-secretome responsible for infiltrating-monocyte recruitment and macrophage-priming. Pertinently, the aim of this study was the exploration of CSC impact on TAM traits in CCA. Indeed, we showed that TAMs acquire a peculiar phenotype dependent on bioactive CSCsecretome. Further studies are required to deepen SPH macrophage properties on tumorigenic potential and drug-resistance of SPH cells.

A variety of modulators released by stromal and tumor cells recruit circulating monocytes to tumor sites, cause their differentiation into macrophages and severely affect their functions [20,41–43]. Our results showed profound molecular and functional differences among macrophage-activated by CCA-SPH in comparison to MON macrophages suggesting that different TAM-compartments within same tumor may reflect diverse

Molecular and Cell Biology



**Fig. 7. IL13, OA, IL34 combination mimics SPH-like effects on macrophage-differentiation and monocyte recruitment.** (A) FACS-profile of CD115, HLA-DR and CD206 expression in macrophages obtained by CD14<sup>+</sup> cultured in presence of IL13 (80 pg/ml), OA (0.9 ng/ml) and IL34 (80 pg/ml), added to the MON-CM (green). Inhibitory impact of human antibodies anti-IL13 (800 ng/ml, 10,000x), OA (2700 ng/ml, 3000x), IL34 (800 ng/ml, 10,000x) alone (in brown, dark grey and dark green, respectively) or in combination was shown (violet). Effects of both SPH- (red) and MON- (blue) CM also shown. Histograms represent three independent experiments using macrophage from three different healthy donors. (B) Relative expression of M1/M2 and matrix remodeling-related genes. *GAPDH* as internal control. All mRNA levels displayed as fold changes normalized to 1 (mean expression of MON macrophage) (n = 3). Mean ± SEM (*p* value *vs.* MON- or SPH-CM by Student's *t* test, \**p*  $\leq$ 0.01, \*\*\**p*  $\leq$ 0.001). (C) Invasion and adhesion assay with fibronectin supports. Cells counted normalized to MON macrophages (n = 5). Migration assay of monocytes counted and normalized to monocyte migrated in presence of MON-CM (n = 5). Mean ± SEM (*p* value *vs.* MON- or SPH-CM by Student's *t* test, \**p*  $\leq$ 0.01, \*\*\**p*  $\leq$ 0.001). (D) macrophage-role in supporting *in vivo* tumorigenicity. 1000 MONs (SG231) co-injected with 300 *in vitro*-educated macrophages into NSG mice. Tumor growth evaluated. Mean ± SEM (n = 5, *p* value *vs.* MON or SPH macrophage.

responses to divergent local signals. Indeed, tumor stem-like associated-TAMs displayed mixed M1-M2 molecular traits including high invasion and adhesion capability. Hence, a mixed

state of macrophage-subsets reinforced the concept of TAM plasticity. It is important to note that CSC ability to modulate a macrophage dominated inflammatory response is similar to the

reactive phenotype displayed by cholangiocytes of dysplastic bile ducts in a mouse model of congenital hepatic fibrosis [44], a disease with high risk of CCA development.

Of relevance, our findings indicated that CCA-SPH released molecules, such as IL13, OA and IL34, model the macrophage component associated to CSCs. Interestingly, IL34 has recently been identified as a second ligand for hematopoietic colony-stimulating factor-1 receptor (CSF-1R) together with the cytokine CSF1, also known as M-CSF [45]. CSF1-R activation establishes signaling cascades leading to differentiation and functionality of monocytes, tissue-macrophages and antigen-presenting dendritic cells [31,41]. Thus, macrophage-differentiation may be possibly driven by IL34 or alternative mechanisms (e.g., VEGF) proposing new clues for CCA-immunotherapy strategies [42].

Consistently, IL13 has been described as a typical T helper 2 cell type cytokine that, together with IL4, generated alternatively activated M2-macrophages [31,41]. This clearly suggested that SPH induced-macrophages resemble M2-polarized cells.

Lastly, OA (also known as glycoprotein non-metastatic melanoma protein b [gpnmb]), a type 1 transmembrane glycoprotein, is produced by embryonic nervous system, developing nephrons, osteoblasts and osteoclasts. Remarkably, OA is overexpressed in patients with glioblastoma multiform and significantly correlated with poor outcome [46]. Moreover, in breast cancer cells, OA high expression levels are required for their enhanced invasiveness and osteolytic bone metastases formation [47].

Together our observations pointed to a stem-like secretome (IL13, OA, IL34 production) and resulting-macrophage phenotype in human CCA. Although, additional studies are warranted to formally dissect molecular mechanisms underlying differentiation of recruited monocytes towards SPH macrophage-subset, we provide a rationale for IL13, OA and IL34 targeting. Indeed, extra analyses are essential for demonstrating the base knowledge for effective prevention and/or treatment of malignant SPH macrophage-subset, thus leading to exploration of combination therapies in CCA patients.

Notably, despite several limitations, our humanized xenograft (s.c. patient CCA-SPH and i.v. hPBMC) highlights SPH biological relevance in priming associated macrophages. In this respect, we know that orthotopic allograft models for human CCA represent a potential tool to recapitulate key clinical, cellular and molecular features of this aggressive tumor [48]. While diverse animal models of CCA (including numerous new genetic models) do exist, an effective patient-like CCA animal model remains to be established [48]. Thus, this lack represents a limitation in the understanding the CCA pathogenesis. Nevertheless, we presented a human CCA-like setting as preclinical platform to investigate mechanisms bridging CCA-disease to CCA-infiltrating macrophages for synergistic therapies. To this end, in the contest of nanotechnology medicine, in vivo administration of IL13, OA and IL34 antibodies combined with chemotherapies, potentially represents a versatile biomedical tool to improve CCA outcome.

### **Financial support**

The study was supported partly by the Italian Foundation of Cancer Research award (MFGA17588) to Dr. Raggi, partly by the Dr. Nicholas C. Hightower Centennial Chair of Gastroenterology from Scott & White, a VA Research Career Scientist Award, VA Merit award (5101BX000574) to Dr. Alpini and a VA Merit award (5l01BX002192) to Dr. Glaser. Dr. Raggi was supported by U. Veronesi Foundation Post-Doctoral fellowship.

#### **Conflict of interest**

The authors who have taken part in this study declared that they do not have anything to disclose regarding funding or conflict of interest with respect to this manuscript.

### Authors' contributions

Study design: CR; Data acquisition/analysis: MC JBA AD FS LDT EF; Manuscript drafting: CR; Critical revision: AS GA DA; Statistical analysis: GC; Obtained funding: CR GA PI; Technical support: VC SG MR JMB CC GT CC JMB LB

### Acknowledgements

We thank Dr. A.J. Demetris (University of Pittsburgh, Pittsburgh, PA) for HUCCT1, CCLP1 and SG231 cells; Dr. G. J. Gores (Mayo Clinic, Rochester, MN) for human immortalized non-malignant cholangiocyte cell line H69; Dr. E. Morenghi for assistance with the statistical analysis; Dr. E. Trombetta for human anti-CD44 and EPCAM antibodies. This material is the result of work supported by resources at the Central Texas Veterans Health Care System. The views expressed in this article are those of the authors and do not necessarily represent the views of the Department of Veterans Affairs.

### Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.jhep.2016.08. 012.

## References

Author names in bold designate shared co-first authorship

- [1] Andersen JB, Spee B, Blechacz BR, Avital I, Komuta M, Barbour A, et al. Genomic and genetic characterization of cholangiocarcinoma identifies therapeutic targets for tyrosine kinase inhibitors. Gastroenterology 2012;142:1021–1031 e1015.
- [2] Blechacz B, Gores GJ. Cholangiocarcinoma: advances in pathogenesis, diagnosis, and treatment. Hepatology 2008;48:308–321.
- [3] Raggi C, Invernizzi P, Andersen JB. Impact of microenvironment and stemlike plasticity in cholangiocarcinoma: molecular networks and biological concepts. J Hepatol 2015;62:198–207.
- [4] Banales JM, Cardinale V, Carpino G, Marzioni M, Andersen JB, Invernizzi P, et al. Expert consensus document: Cholangiocarcinoma: current knowledge and future perspectives consensus statement from the European Network for the Study of Cholangiocarcinoma (ENS-CCA). Nat Rev Gastroenterol Hepatol 2016;13:261–280.
- [5] Lin G, Liu Y, Li S, Mao Y, Wang J, Shuang Z, et al. Elevated neutrophil-tolymphocyte ratio is an independent poor prognostic factor in patients with intrahepatic cholangiocarcinoma. Oncotarget 2016;7:50963–50971.
- [6] Sabbatino F, Villani V, Yearley JH, Deshpande V, Cai L, Konstantinidis IT, et al. PD-L1 and HLA class I antigen expression and clinical course of the disease in intrahepatic cholangiocarcinoma. Clin Cancer Res 2016;22:470–478.

Molecular and Cell Biology

- distribution in cholangiocarcinoma. World J Gastroenterol 2015;21:4961–4968.[8] Sasaki M, Tsuneyama K, Ishikawa A, Nakanuma Y. Intrahepatic cholangio-
- carcinoma in cirrhosis presents granulocyte and granulocyte-macrophage colony-stimulating factor. Hum Pathol 2003;34:1337–1344.
  [0] Fan B. Malato Y. Cubici DE Navi S. Pazumilava N. Pibback S. et al.
- [9] Fan B, Malato Y, Calvisi DF, Naqvi S, Razumilava N, Ribback S, et al. Cholangiocarcinomas can originate from hepatocytes in mice. J Clin Invest 2012;122:2911–2915.
- [10] Marquardt JU, Raggi C, Andersen JB, Seo D, Avital I, Geller D, et al. Human hepatic cancer stem cells are characterized by common stemness traits and diverse oncogenic pathways. Hepatology 2011;54:1031–1042.
- [11] Raggi C, Factor VM, Seo D, Holczbauer A, Gillen MC, Marquardt JU, et al. Epigenetic reprogramming modulates malignant properties of human liver cancer. Hepatology 2014;59:2251–2262.
- [12] Kreso A, Dick JE. Evolution of the cancer stem cell model. Cell Stem Cell 2014;14:275–291.
- [13] Hanahan D, Coussens LM. Accessories to the crime: functions of cells recruited to the tumor microenvironment. Cancer Cell 2012;21:309–322.
- [14] Plaks V, Kong N, Werb Z. The cancer stem cell niche: how essential is the niche in regulating stemness of tumor cells? Cell Stem Cell 2015;16:225–238.
- [15] Jinushi M, Chiba S, Yoshiyama H, Masutomi K, Kinoshita I, Dosaka-Akita H, et al. Tumor-associated macrophages regulate tumorigenicity and anticancer drug responses of cancer stem/initiating cells. Proc Natl Acad Sci U S A 2011;108:12425–12430.
- [16] Su S, Liu Q, Chen J, Chen F, He C, Huang D, et al. A positive feedback loop between mesenchymal-like cancer cells and macrophages is essential to breast cancer metastasis. Cancer Cell 2014;25:605–620.
- [17] Wan S, Zhao E, Kryczek I, Vatan L, Sadovskaya A, Ludema G, et al. Tumorassociated macrophages produce interleukin 6 and signal via STAT3 to promote expansion of human hepatocellular carcinoma stem cells. Gastroenterology 2014;147:1393–1404.
- [18] Lu H, Clauser KR, Tam WL, Frose J, Ye X, Eaton EN, et al. A breast cancer stem cell niche supported by juxtacrine signalling from monocytes and macrophages. Nat Cell Biol 2014;16:1105–1117.
- [19] Zhou W, Ke SQ, Huang Z, Flavahan W, Fang X, Paul J, et al. Periostin secreted by glioblastoma stem cells recruits M2 tumour-associated macrophages and promotes malignant growth. Nat Cell Biol 2015;17:170–182.
- [20] Biswas SK, Sica A, Lewis CE. Plasticity of macrophage function during tumor progression: regulation by distinct molecular mechanisms. J Immunol 2008;180:2011–2017.
- [21] Roberts PE. Isolation and establishment of human tumor stem cells. Methods Cell Biol 2008;86:325–342.
- [22] Pollard SM, Yoshikawa K, Clarke ID, Danovi D, Stricker S, Russell R, et al. Glioma stem cell lines expanded in adherent culture have tumor-specific phenotypes and are suitable for chemical and genetic screens. Cell Stem Cell 2009;4:568–580.
- [23] Miyagiwa M, Ichida T, Tokiwa T, Sato J, Sasaki H. A new human cholangiocellular carcinoma cell line (HuCC-T1) producing carbohydrate antigen 19/9 in serum-free medium. In Vitro Cell Dev Biol 1989;25:503–510.
- [24] Han C, Wu T. Cyclooxygenase-2-derived prostaglandin E2 promotes human cholangiocarcinoma cell growth and invasion through EP1 receptor-mediated activation of the epidermal growth factor receptor and Akt. J Biol Chem 2005;280:24053–24063.
- [25] Shimizu Y, Demetris AJ, Gollin SM, Storto PD, Bedford HM, Altarac S, et al. Two new human cholangiocarcinoma cell lines and their cytogenetics and responses to growth factors, hormones, cytokines or immunologic effector cells. Int J Cancer 1992;52:252–260.
- [26] Cardinale V, Renzi A, Carpino G, Torrice A, Bragazzi MC, Giuliante F, et al. Profiles of cancer stem cell subpopulations in cholangiocarcinomas. Am J Pathol 2015;185:1724–1739.
- [27] Sirica AE, Zhang Z, Lai GH, Asano T, Shen XN, Ward DJ, et al. A novel "patientlike" model of cholangiocarcinoma progression based on bile duct inoculation of tumorigenic rat cholangiocyte cell lines. Hepatology 2008;47:1178–1190.

- [28] Saeed AI, Sharov V, White J, Li J, Liang W, Bhagabati N, et al. TM4: a free, open-source system for microarray data management and analysis. Biotechniques 2003;34:374–378.
- [29] Quintana E, Shackleton M, Sabel MS, Fullen DR, Johnson TM, Morrison SJ. Efficient tumour formation by single human melanoma cells. Nature 2008;456:593–598.
- [30] Raggi C, Mousa HS, Correnti M, Sica A, Invernizzi P. Cancer stem cells and tumor-associated macrophages: a roadmap for multitargeting strategies. Oncogene 2015;35:671–682.
- [31] Sica A, Schioppa T, Mantovani A, Allavena P. Tumour-associated macrophages are a distinct M2 polarised population promoting tumour progression: potential targets of anti-cancer therapy. Eur J Cancer 2006;42:717–727.
- [32] Steidl C, Lee T, Shah SP, Farinha P, Han G, Nayar T, et al. Tumor-associated macrophages and survival in classic Hodgkin's lymphoma. N Engl J Med 2010;362:875–885.
- [33] Bingle L, Brown NJ, Lewis CE. The role of tumour-associated macrophages in tumour progression: implications for new anticancer therapies. J Pathol 2002;196:254–265.
- [34] Schmidt T, Brodesser A, Schnitzler N, Gruger T, Brandenburg K, Zinserling J, et al. CD66b overexpression and loss of C5a receptors as surface markers for staphylococcus aureus-induced neutrophil dysfunction. PLoS One 2015;10 e0132703.
- [35] Solinas G, Schiarea S, Liguori M, Fabbri M, Pesce S, Zammataro L, et al. Tumor-conditioned macrophages secrete migration-stimulating factor: a new marker for M2-polarization, influencing tumor cell motility. J Immunol 2010;185:642–652.
- [36] Chen J, Yao Y, Gong C, Yu F, Su S, Liu B, et al. CCL18 from tumor-associated macrophages promotes breast cancer metastasis via PITPNM3. Cancer Cell 2011;19:541–555.
- [37] Pollard JW. Trophic macrophages in development and disease. Nat Rev Immunol 2009;9:259–270.
- [38] Biswas SK, Mantovani A. Macrophage plasticity and interaction with lymphocyte subsets: cancer as a paradigm. Nat Immunol 2010;11:889–896.
- [39] Sica A, Mantovani A. Macrophage plasticity and polarization: in vivo veritas. J Clin Invest 2012;122:787–795.
- [40] Dang W, Qin Z, Fan S, Wen Q, Lu Y, Wang J, et al. MiR-1207-5p suppresses lung cancer growth and metastasis by targeting CSF1. Oncotarget 2016;7:32421–32432.
- [41] Mantovani A, Allavena P. The interaction of anticancer therapies with tumor-associated macrophages. J Exp Med 2015;212:435–445.
- [42] Franklin RA, Liao W, Sarkar A, Kim MV, Bivona MR, Liu K, et al. The cellular and molecular origin of tumor-associated macrophages. Science 2014;344:921–925.
- [43] Van Overmeire E, Laoui D, Keirsse J, Van Ginderachter JA, Sarukhan A. Mechanisms driving macrophage diversity and specialization in distinct tumor microenvironments and parallelisms with other tissues. Front Immunol 2014;5:127.
- [44] Locatelli L, Cadamuro M, Spirli C, Fiorotto R, Lecchi S, Morell CM, et al. Macrophage recruitment by fibrocystin-defective biliary epithelial cells promotes portal fibrosis in congenital hepatic fibrosis. Hepatology 2016;63:965–982.
- [45] Felix J, Elegheert J, Gutsche I, Shkumatov AV, Wen Y, Bracke N, et al. Human IL-34 and CSF-1 establish structurally similar extracellular assemblies with their common hematopoietic receptor. Structure 2013;21:528–539.
- [46] Kuan CT, Wakiya K, Dowell JM, Herndon 2nd JE, Reardon DA, Graner MW, et al. Glycoprotein nonmetastatic melanoma protein B, a potential molecular therapeutic target in patients with glioblastoma multiforme. Clin Cancer Res 2006;12:1970–1982.
- [47] Rose AA, Pepin F, Russo C, Abou Khalil JE, Hallett M, Siegel PM. Osteoactivin promotes breast cancer metastasis to bone. Mol Cancer Res 2007;5:1001–1014.
- [48] De Minicis S, Kisseleva T, Francis H, Baroni GS, Benedetti A, Brenner D, et al. Liver carcinogenesis: rodent models of hepatocarcinoma and cholangiocarcinoma. Dig Liver Dis 2013;45:450–459.