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Data in Brief

Gene expression profiling of the tumor microenvironment in human intrahepatic cholangiocarcinoma

Laurent Sulpice a,b,c, Mireille Desille a,b,d, Bruno Turlin a,b,d,e, Alain Fautrel a,b, Karim Boudjema a,b,c, Bruno Clément a,b, Cédric Coulouarn a,b,*

a Inserm, UMR991, Liver Metabolisms and Cancer, F-35033 Rennes, France
b Université de Rennes 1, F-35043 Rennes, France
c CHU Rennes, F-35033 Rennes, France
d CHU Rennes, Centre de Ressources Biologiques Santé, F-35033 Rennes, France
e CHU Rennes, Service d’Anatomie et Cytologie Pathologiques, F-35033 Rennes, France

ABSTRACT

Intrahepatic cholangiocarcinoma (ICC) is the second most common type of malignant primary tumors in the liver. ICC is an aggressive cancer with a poor survival and limited therapeutic options. At the histological level, ICC is characterized by an abundant stroma (i.e. the tumor microenvironment that notably includes components of the extracellular matrix, stromal cells and soluble factors). Tumor microenvironment is known to play a key role in tumor onset and progression but it is poorly characterized at the molecular level. Thus, this study was specifically designed to identify genes that are significantly deregulated in the tumor microenvironment of human ICC. Here we provide a detailed description of the experimental design and methods used to acquire the genomic data deposited into Gene Expression Omnibus (GEO) under the accession number GSE45001.

Our genomic dataset provides insights on the molecular pathways altered in the microenvironment of ICC and allows the identification of novel ICC biomarkers, as exemplified previously in Hepatology (PMID: 23775819).

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2. Experimental design, materials and methods

2.1. General objective and experimental design

The aim of the study was to identify genes significantly deregulated in the microenvironment of human ICC. For this purpose, we applied a gene expression profiling approach using Agilent pangenomic microarrays and total RNA isolated from human ICC samples after laser capture microdissection (LCM) of the normal vs. tumoral stroma from 10 patients with ICC.

2.2. Patients

Gene expression profiling was performed from fresh frozen tissues of 10 patients with ICC [1]. Fresh frozen tissues and formalin-fixed paraffin-embedded (FFPE) tissues were provided by the biobank of the hospital-university (Centre de Ressources Biologiques [CRB] Santé de Rennes; BB-0033-00056). Patients underwent liver resection at Rennes university hospital between Jan. 1997 and Aug. 2011. Only mass-forming types ICC were included, as defined by the Liver Cancer Study Group of Japan. Written informed consent was obtained from all patients. The study protocol fulfilled national laws and regulations and was approved by the local ethics committee and the Institutional Review Board (IRB00003888) of Inserm (IORG0003254).

http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE45001

* Corresponding author at: Inserm, UMR991, Liver Metabolisms and Cancer, F-35033 Rennes, France.
E-mail address: cedric.coulouarn@inserm.fr (C. Coulouarn).

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2.3. Laser capture microdissection (LCM) and RNA extraction

The integrity and the quality of all tissue sections were first validated at the histological level after hematoxylin and eosin (H&E) staining by an experienced liver pathologist (Fig. 1A–B). LCM was then performed to isolate the fibrous tissue from tumoral and non-tumoral ICC samples. LCM was performed using the Arcturus Veritas™ microdissection system (Applied Biosystems, Carlsbad, CA). From frozen tissues,
serial sections of 10 μm were prepared using a Leica 3050 S cryostat (Leica Microsystems, Wetzlar, Germany) and mounted onto a PEN membrane glass slide (Applied Biosystems). Tissue sections were dehydrated by successive immersions (30 s. twice) in 70%, 90% and 100% ethanol solutions. Enzymatic activity was locked by the immersion in a xylene solution (1 min. Twice) before performing LCM (Fig. 1C). LCM was performed within 1 h to limit RNA degradation. The average microdissected area was 5.07 ± 1.42 mm² (6.01 ± 1.02 mm² for tumor stroma and fibrous tissue in the adjacent non-tumor tissue, respectively) (Fig. 1D-E). Total RNA from laser capture microdissected tissues (Fig. 1E) was extracted and purified using an Arcturus Picopure RNA isolation kit according to the manufacturer’s instructions (Applied Biosystems, Carlsbad, CA). RNA was qualified using a Nanodrop 2000 spectrophotometer (Thermo Fischer Scientific, Waltham, MA). The average RNA concentration was 6.18 ± 3.05 ng/μL and the 260/280 ratio was above 2.

2.4. Gene expression analysis

Genome-wide expression profiling was performed using a low-input QuickAmp labeling kit and human SurePrint G3 8x60K pan genomic microarrays (Agilent Technologies, Santa Clara, CA) as previously described [2]. Total RNA, in presence of an external spike-in RNA mixture, were amplified and labeled with Cy3 fluorescent dye using Agilent one color low-input QuickAmp labeling kit following the manufacturer’s instructions. Starting from 50 ng total RNA purified from LCM tissue samples, amplification yield was 1.8 ± 0.7 μg cRNA and specific activity was 5.8 ± 3.4 pmol Cy3 per μg cRNA.

The same amount of Cy3-labeled cRNA (600 ng) was fragmented at 60 °C for 30 min in a reaction volume of 25 μl containing 1 × Agilent fragmentation buffer and 2 × Agilent blocking agent following the manufacturer’s instructions. Following the fragmentation reaction, 25 μl of 2 × Agilent hybridization buffer (HI-RPM) was added to the fragmentation mixture. Forty μl of the mixture were hybridized to Agilent SurePrint G3 Human GE 8x60K for 17 h at 65 °C in a rotating Agilent hybridization oven. After hybridization, microarrays were washed 1 min at room temperature with GE Wash Buffer 1 (Agilent) and 1 min with GE Wash buffer 2 at 37 °C (Agilent).

Microarray were scanned immediately after washing with the Agilent DNA Microarray C Scanner (G25655) using one color scan setting for 8x60k array slides. The scanned images were analyzed with Feature Extraction Software 10.7.3.1 (Agilent Technologies) using default parameters (protocol GE1_107_Sep09_sssSurrogates and Grid: 028004_D_F_20101102). Quality control showed a broad distribution of background subtracted signals, suggesting that RNA with various abundances (from low to high) could be analyzed from the genomic dataset (Fig. 2A). The analysis also demonstrated...
an expected correlation between the coefficient of variation and signal intensity (Fig. 2B). The use of Agilent spike-In RNA demonstrated a linear relation between signal intensity and the relative abundance or concentration of the RNA (Fig. 2C). All 20 microarrays from the genomic dataset were qualified and further analyzed using GeneSpring software (Agilent Technologies).

Microarray data were first normalized by using the quantile normalization algorithm. As shown in Fig. 3 this normalization method corrected putative technical variations between samples. Without initial gene filtration, Principal Component Analysis (PCA) using expression values of all genes in all 20 samples mostly separated tumor from non-tumor samples (Fig. 4). Then, a filtration by “flag” and “signal intensity” was applied. Were retained only the entities in which at least 50% of the values in any of the two conditions (Non Tumoral vs. Tumoral) had a “detected flag” (i.e. a positive and significant feature as defined by GeneSpring). For the filtration by signal intensity, were retained the entities in which at least 50% of the values in any of the two conditions (non tumoral, NT vs. tumoral, T) were within the range of interest (i.e. 20–100th percentile). Differentially expressed genes were identified by a two-sample univariate t-test and a random variance model as described [1]. Permutation P-values for significant genes were computed based on 10,000 random permutations.

In conclusion, our genomic dataset is relevant to provide insights on the molecular pathways altered in the microenvironment of ICC and to identify novel ICC biomarkers [1,3].

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