Methylation of CAR is involved in the Suppression of CYP2C19 in

HBV-associated Hepatocellular Carcinoma

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HCC, hepatocellular carcinoma; 5-aza-dC, 5-Aza-2'-Deoxycytidine; BSP, bisulfite sequencing PCR; MSP, methylation specific PCR; HBV, hepatitis B virus; HBsAg, hepatitis B surface antigen; HBcAb, hepatitis B core antibody; HCV, hepatitis C virus; CDS, coding DNA sequence; EGF, epidermal growth factor; EGFR, epidermal growth factor receptor; CAR, constitutive androstane receptor; HNF, hepatocyte nuclear factor; GR, glucocorticoid receptor; PXR, pregnane X receptor; SAM, S-adenosyl methionine.

Abstract

Hepatocellular carcinoma (HCC), one of the most dangerous malignancies with increasing incidence and high mortality rate, represents a major international health problem. The progression of HCC is known to involve a genome-wide alteration of epigenetic modifications, leading to aberrant gene expression patterns. The activity of CYP2C19, an important member of CYP450 superfamily, was reported to be compromised in HCC, but the underlying mechanism remains unclear. To understand if epigenetic modification in HCC is associated with CYP2C19 activity change, we evaluated the expression levels of CYP2C19 and its transcriptional factors by quantitative real-time PCR using mRNA extracted from both primary hepatocytes and paired tumor vs non-tumor liver tissues ofhepatitis B virus (HBV)-infected patients. DNA methylation was examined by bisulfite sequencing (BSP) and methylation specific PCR (MSP). Results indicated that CYP2C19 could be regulated by e-box methylation of constitutive androstane receptor (CAR). The decreased expression of CYP2C19 in tumorous tissues of HBV infected HCC patients is highly correlated with the suppressed expression and promoter hypermethylation of CAR. Our study demonstrated that aberrant CAR methylation is involved in the regulation of CYP2C19 in HBV-related HCC and it may play a role during liver tumorigenesis.

Introduction

Although tremendous endeavors have been made for a long time to fight against hepatocellular carcinoma (HCC), HCC still remains the fifth most prevalent cancer worldwide and is the third leading cause of cancer mortality up to now (Jemal et al., 2011). Hepatitis B virus (HBV) infection, Hepatitis C virus (HCV) infection and alcohol abuse are considered to be the main causes, among which HBV infection accounts for 60% of HCC in Chinese population. A recent report by World Health Organization estimated that 90 million people in China - almost 7% of Chinese population - are chronically infected with HBV.

Traditionally, HCC progression was considered as a gradual multistep process of normal cells evolving into tumor cells caused by mutations which lead to altered hepatocellular phenotypes (Thorgeirsson and Grisham, 2002; Lessel et al., 2014). However, evidence accumulated in the past few decades suggest that altered epigenetic modifications also play a pivotal role in HCC development (Nishida and Goel, 2011). DNA methylation, which refers to the addition of a methyl group to the cytosine within a CpG dinucleotide, is the most intensively studied epigenetic mechanism in mammals. It has been reported that the cancerous tissues in HCC possess a unique methylation landscape which differs from adjacent tissues. In HCC, the methylome feature is roughly characterized by global hypomethylation and gene-specific DNA hypermethylation, which contribute to genomic instability and inactivation of tumor-suppressor genes, respectively (Herceg and Paliwal, 2011). Since advanced HCC is highly lethal and incurable, there is an urgent need for

understanding the tumorigenesis process as well as searching for new diagnostic biomarkers and novel therapeutic targets.

CYP2C19 is one of the most important members of CYP450 superfamily and is responsible for the metabolism of a variety of clinical important drugs like mephenytoin, diazepam and some barbitals (Griskevicius et al., 2003). The genetic polymorphisms of CYP2C19 have been well studied. Individuals can be divided into extensive metabolizers (EMs), intermediate metabolizers (IMs), poor metabolizers (PMs) and ultra-rapid metabolizers (UMs) based on the catalytic capacity of the enzyme, which is largely dependent on different alleles the individuals are carrying (Uppugunduri et al., 2012). The regulatory mechanism of CYP2C19 is currently characterized by several nuclear receptors and transcriptional factors including CAR, PXR, GR α and HNF-3 γ , among which CAR appears to play a central role and has strong association with the basal expression level of CYP2C19 (Wortham et al., 2007). It has been predicted that epigenetic might also play a role in the regulation of CYP2C19 and there exist some CpG sites in the 5'-flanking region of CYP2C19 gene, but their functional importance remains to be further elucidated (Ingelman-Sundberg et al., 2007).

The expression of CYP2C19 is reported to be decreased in advanced cancers including breast cancer, lung cancer, as well as HCC, which is also accompanied by comprised enzymatic activity (Helsby et al., 2008; Tsunedomi et al., 2005). However, regardless of the extensive effort to investigate the underlying mechanism over the past years, the altered expression of CYP2C19 in HCC does not appear to be fully

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accounted by either known polymorphisms, or inflammatory factors, or growth hormones (Helsby et al., 2008). The aim of this study is to investigate the possible epigenetic mechanism in the decreased expression of CYP2C19 in HBV infected HCC. Since CYP2C19 is also important in the disposition of a number of chemotherapeutic agents, including cyclophosphamide, thalidomide and bortezomib, elucidating the epigenetic modulation of CYP2C19 might also help understand and predict the chemotherapy outcome for HCC patients.

Materials and Methods

Cell Culture and Tissue Samples

HepG2 and HEK293 cells were grown in DMEM medium (Life Technologies, Carlsbad, CA), supplemented with 10% FBS (Life Technologies, Carlsbad, CA) and 1% PEST (Life Technologies, Carlsbad, CA), at 37°C with 5% CO₂. HK2 cells were grown in DMEM/F-12 medium (Life Technologies, Carlsbad, CA) supplemented with 18% FBS at 37°C with 5% CO. Cryopreserved primary hepatocytes (RILD Research Institute for Liver Disease, Shanghai, China) from one male and one female donor (Supplemental Table 1) were plated in InVitroGROTM CP Medium (BioreclamationIVT, Baltimore, MD) at 37°C with 5% CO₂ before drug incubation. 2 replicates were conducted for each donor.

Thirty pairs of tumorous and adjacent liver tissues were obtained from fresh surgical specimen of HBV infected HCC patients (HBV DNA>1000copies/ml or HBsAg and HBcAb test positive) and were then stored in liquid nitrogen before use. The selected patients did not receive any radiotherapy or chemotherapy before. Tissues were obtained from Zhejiang Cancer Hospital. Information of liver species is provided as Supplemental Table 2. The research protocol has been carried out in accordance with the Declaration of Helsinki and was approved by the Ethics Committee of Zhejiang Cancer Hospital.

Treatment of Primary Hepatocytes with 5-aza-dC

Primary hepatocytes were seeded at a density of 5×10^5 cells/ml in collagen-coated

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6-well plates (Life Technologies, Carlsbad, CA) with *InVitro*GROTM CP Medium (BioreclamationIVT, Baltimore, MD). Medium was changed the next day to remove the unattached cells. After 24h of plating, drug incubation started with 1 μ M and 2 μ M 5-aza-dC (Sigma, St. Louis, MO) (Christman, 2002) in *InVitro*GROTM HI Medium supplemented with 25ng/ml EGF (Sigma, St. Louis, MO) to stimulate proliferation. Control group was treated with 0.1% DMSO. Medium was changed every other day. Cells were harvested after 5 days treatment for DNA and RNA isolation.

Semi-quantitative Real-Time PCR

mRNA was extracted from tissue samples and cultured cells using the RNeasy Mini Kit (Qiagen, Valenica, CA). 500ng mRNA was used in reverse transcribed PCR reaction performed with PrimeScript RT reagents Kit (Takara, Tokyo, Japan) and thenquantitative real-time PCR was performed using the SYBR[®] Premix Ex TaqTM II (Takara, Tokyo, Japan) and the StepOnePlusTM System (Applied Biosystems, Foster City, CA). Primers for quantitative real-time PCR are listed in Table 1. Relative quantitation of gene expression was calculated with the $2^{-\Delta\Delta Ct}$ method as described previously (Schmittgen and Livak, 2008).

Table 1.

DNA Methylation Analysis

Genomic DNA from tissue samples or cells was isolated using the DNeasy Blood &Tissue Kit (Qiagen, Valenica, CA). DNA was considered qualified with a ratio of

absorbance at 260nm and 280nm (A260/A280) between 1.7 and 1.9. Samples were stored in elution buffer at -80°C before use.

Sodium bisulfite modification of DNA samples was conducted using an EZ DNA MethylationTM-Gold Kit (Zymo, Orange, CA). The location of all CpG sequences examined in this study is shown in Figure 1. PCR amplification was done with rTaq polymerase (Takara, Tokyo, Japan) in a 50µl reaction mixture with 2µl modified DNA as template. It starts with 10 cycles of touchdown PCR of 30s at 94°C, 30s at 60°C to 55° C (-0.5°C per cycle), and then 45s at 72°C and 30 cycles of 30s at 94°C, 30s at 55° C, 45s at 72°C, and finally an extension of 10min at 72°C. Amplicons were then purified and subcloned into pMD18-T vector (Takara, Tokyo, Japan) according to manufacturer's instruction. Ligation products were transformed into E. coli DH5 α cells and 10-15 random colonies with recombinant plasmids were subjected to Sanger sequencing (Sangon, Shanghai, China).

Methylation-specific PCR was also conducted using the modified DNA as template. The followingtouchdown PCR program was applied: 10 cycles of touchdown PCR of 30s at 94°C, 30s at 58°C to 53°C (-0.5°C per cycle), 20s at 72°C and 25 cycles of 30s at 94°C, 30s at 53°C, 20s at 72°C, followed by a final extension of 10min at 72°C. 10µl PCR products were shown on a 2% agarose gel containing ethidium bromide. Primers for methylation analysis are listed in table 2.

Figure 1.

Table 2.

Plasmids Construction

pCpGL-CYP2C19 (-1977~+64), pCpGL-CAR-CGI (-1446~+1971), pCpGL-CAR-2E (+175~+1971) and pCpGL-CAR-1E (+384~+1971) were constructed with a CpG-free luciferase reporter vector named pCpGL-basic (Fig. 2A) (Klug and Rehli, 2006). pCpGL-CYP2C19 contains a ~2.0kb sequence of the 5'-flanking promoter region of CYP2C19. pCpGL-CAR-CGI contains a ~3.5kb sequence with a CpG island on the 5'-flanking region and both e-box binding sites close to transcription start site and translation start site of CAR, respectively. In pCpGL-CAR-2E, the CpG island region is deleted from the sequence of pCpGL-CAR-CGI, while in pCpGL-CAR-1E, another \sim 200bp sequence which contains the first e-box binding site was deleted. Additionally, a full-length CAR CDS amplified from mRNA (primers CDS-F/R) was ligated with the 5'-flanking region as well as the untranslated exon 1 and intron 1 (amplified by primers Flanking-F/R) by overlap PCR. The ~3kb ligated fragment was then inserted into a CpG-free expression vector, pCpGfree-mcs (Fig. 2B) (InvivoGen, San Diego, CA), to construct pCpGfree-CAR. Primers used for plasmids construction are listed in Table 3. All constructs were confirmed by Sanger sequencing (Sangon, Shanghai, China).

Figure 2.

Table 3.

In vitro Methylation of Plasmid DNA

Plasmids were methylated using M. SssI (NEB, Beverly, MA) according to

manufacturer's instruction. $2\mu g$ plasmid DNA was incubated in a 40µl volume with $2\mu l$ (4U /µl) Sss I and 160µM SAM at 37°C for 6 hours. The control group of plasmid also underwent the same incubation, but in the absence of Sss I. The reaction was stopped by heating at 65°C for 20 minutes. Both methylated and control group of plasmid DNA was purified with Axygen PCR Clean-up Kit (Axygen, Hangzhou, China).

Transient Transfection Assay

HepG2, HEK293 or HK2 cells were seeded in 24-well plates at the density of 2.5×10^5 /ml. The transient transfection was carried out using Lipofectamine2000 (Life Technologies, Carlsbad, CA) when cells have reached 70~80% confluence. Reporter plasmids and expression plasmids were transfected at 500ng and 250ng per well, respectively, with 50ng pRL-TK per well as internal control. Medium was refreshed 6 hours after transfection. Luciferase activity was assessed using the Dual-Luciferase Reporter Assay System (Promega, Madison, WI) 48 hours post transfection.

Statistical Analysis

qPCR results were shown as $2^{-\Delta\Delta Ct}$ using GAPDH as housekeeping gene. The methylation percentage of BSP analysis was calculated as the number of methylated cytosines divided by the total number of cytosines on the CpG site in all amplicons. The methylation percentage of a sequence was determined as the average percentage of all CpG sites within this region. Comparison of methylation percentage and mRNA

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expression between groups was analyzed by Paired Student's t test using GraphPad Prism 5.0 (GraphPad, San Diego, CA) software. The correlation between CYP2C19 and CAR expression was determined by Pearson correlation. Difference or correlation is considered statistically significant at the level of *p < 0.05, **p < 0.01 and ***p < 0.001.

Results

Increased CYP2C19 and CAR mRNA in Primary Hepatocytes after 5-aza-dC Treatment

In order to determine whether DNA methylation is a potential mechanism of CYP2C19 regulation in the liver, a DNA methyltransferase inhibitor, 5-aza-dC, was applied to the primary cultures of human hepatocytes. Quantitative real-time PCR analysis revealed an increase of CYP2C19 and CAR expression in a dose-dependent manner, while other transcriptional factors were not significantly affected (Fig. 3). To investigate if it was the demethylation that led to the elevated expression level of CYP2C19 and CAR in primary hepatocytes, BSP was performed on the promoter regions as shown in Figure 1. BSP results revealed a moderate methylation frequency on the CpG-poor promoter of CYP2C19 (52.5%) as well as the two e-box binding sites on CAR (50% and 47.5%). After 5-aza-dC treatment, the methylation frequency of these three regions decreased to 38.3%, 13.3% and 42.5%, respectively (Fig. 4A-4C). The CpG island on CAR promoter, which was highly methylated (92.5%), also showed a decreased methylation frequency (72.5%) after exposure to 5-aza-dC (Fig. 4D). BSP results showed an overall decreased DNA methylation level on the promoter of CYP2C19 and CAR after the treatment of 5-aza-dC. These results suggested that DNA methylation is a regulatory mechanism of CYP2C19 in liver, in which CAR might also be involved.

Figure 3.

Figure 4.

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Decreased mRNA Expression of CYP2C19 and CAR in Tumorous Tissues

The expression of CYP2C19 and CAR in tumorous and adjacent tissues is shown as grouped column scatter with mean \pm S.E. in Figure 5A. The average expression of CYP2C19 in tumorous tissues (0.39 \pm 0.20) was significantly decreased compared with the normal tissues (2.4 \pm 0.40, ****p* < 0.001). In addition, the average expression of CAR also decreased significantly (0.40 \pm 0.085 *vs* 1.40 \pm 0.15, ****p* < 0.001). The alterations of expression level of CYP2C19 and CAR were correlated with a Pearson correlation coefficient of 0.31 (R²=0.09, **p* < 0.05) (Fig. 5C).

Figure 5.

Elevated Methylation Level of CpG Sites on CAR Promoter in Tumorous Tissues BSP analysis of segment 5 on CAR was performed using DNA from paired tumor and adjacent tissue samples. Representative BSP results are shown in Figure 6A and corresponding relative mRNA expression were shown in Figure 5B. Clearly, in BSP analysis, CpG 3, CpG 5 and CpG 6 in segment 5 around the transcription start site of CAR were in a higher methylated status in tumorous tissue compared with adjacent tissue. Among these three sites, CpG 3 is a potential e-box binding site, while CpG 5 and 6 are two adjacent CpG sites 121 bp downstream of CpG 3. In addition, MSP within segment 5 also revealed a higher methylation level in tumorous tissues (Fig. 6B). Interestingly, in the case of sample 246564, the expression level of CAR and CYP2C19 were increased while methylation status was decreased in tumorous tissues

(Fig. 6A). No significant methylation difference was detected between tumorous and adjacent tissues in the other BSP analyzed segments (data not shown). Figure 6.

In vitro Methylation on CAR Promoter Leads to Decreased CAR Mediated

CYP2C19 Expression in HepG2 Cells

The results of dual-luciferase reporter assay showed that the CpG-poor 2kb promoter of CYP2C19 was not affected by methylation (Fig. 7A, 7C, 7D) in all tested cell lines, suggesting that these sporadic CpG sites are irrelevant to the methylation mechanism. The unmethylated pCpGL-CAR-CGI was 2.5-fold higher in relative luciferase activity compared to the methylated group in HepG2 cells. This fold change sustained after the deletion of the CpG island as shown by the unmethylated *vs* methylated pCpGL-CAR-2E (2.2 fold), but was greatly reduced after deletion of the 200bp fragment containing the e-box binding site near the transcription start site (in the first intron), indicating that this e-box binding site was essential in the methylation regulating mechanism of CAR, whereas the e-box close to the translation start site was not involved. However, no difference between the methylated and unmethylated group was found in HK2 or HEK293 cells (Fig. 7C, 7D).

To explore if the methylation-mediated decrease of CAR could exert an effect on CYP2C19 expression, either methylated or unmethylated pCpGfree-CAR was co-transfected with pCpGL-CYP2C19 in HepG2 cells. A 2.3-fold higher luciferase activity was observed in the unmethylated pCpGfree-CAR group (Fig. 7B), which

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further supported that CYP2C19 could be suppressed through the hypermethylation

caused decrease of CAR expression.

Figure 7.

Discussion

HBV infection is still a key issue in China's health care according to the *World Cancer Report* published by the WHO in 2014. The HBV-induced hepatocarcinogenesis includes proteomic disruption, viral integrations and aberrant epigenetic modifications (Ji et al., 2014). In the present study, we demonstrated that DNA de-methylation led to elevated CAR and CYP2C19 expression in primary hepatocytes. This finding suggests that DNA methylation is a potential mechanism of CAR and CYP2C19 regulation in the liver. This notion is further supported by the suppressed CAR and CYP2C19 expression in Chinese HBV-related HCC patients.

Since CAR is a member of the nuclear receptor superfamily and a central modulator of oxidative and conjugative enzymes and transporters that are involved in drug disposition and metabolism, it is conceivable that the suppression of CAR plays an important role in the altered expression of CYP2C19 and possibly other targeting drug metabolism genes. CYPs participate in the activation and metabolism of a large number of chemotherapeutic agents such as docetaxel (Shou et al., 1998) and cyclophosphamide (Griskevicius et al., 2003). Thus, CAR suppression mediated by hypermethylation may lead to disturbance of the drug metabolizing system. Therefore, the methylation status of CAR might serve as a biomarker of altered pharmacokinetics. Indeed, it has been confirmed in other studies that several CAR-regulating CYP genes were also suppressed along with the decreased CAR expression in end-stage HCC (Chen et al., 2014), but no significant decrease of CAR and related CYP genes was found in HBV cirrhosis samples. This finding suggested that the sharp decline in the

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expression of these genes might occur when liver cirrhosis evolves into carcinoma. However, our result only offered a Pearson correlation coefficient of 0.31 between the CYP2C19 and CAR expression in contrast to a stronger correlation (Pearson correlation coefficient of 0.693) observed in healthy liver samples (Wortham et al., 2007). Since it has been demonstrated that down-regulation of metabolism related genes is preferentially linked with HBV-related HCC, rather than HCV-related HCC (Okabe et al, 2001; Iizuka et al, 2001), it is possible that the virus itself might have direct impact on the expression of certain genes. Furthermore, as the regulation of CYP2C19 is a complicated network contributed by multiple factors, the effect of CAR might be diluted in vivo. Actually, more and more mechanisms have been discovered in recent years. One example of the epigenetic mechanism is the participation of hsa-miR-29a-3p in the regulation of CYP2C19 in liver cells (Yu et al., 2015). miRNA may also participate in disrupting the correlation between CAR mRNA and protein level, thus leads to the poor correlation between CAR and CYP2C19 mRNA level. Further study is warranted to build a comprehensive map of how CYP2C19 is regulated in HCC.

Since there was a 2-fold decrease in luciferase activity after the deletion of 200 bp sequence containing the e-box at the first intron as indicated in the reporter gene assay, this sequence may act as an enhancer element of CAR. This hypothesis warrants further investigation. We performed the same *in vitro* methylation and reporter gene assay in three cell lines, HepG2, HEK293 and HK2. Interestingly, only HepG2 showed decreased luciferase activity in methylated pCpGL-CAR-2E. No difference

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between the methylated and unmethylated pCpGL-CAR-2E was detected in either HEK293 or HK2. The results indicated that certain factors specific in HepG2 cells may be involved in this methylation related regulation. Biobase (http://www.biobase-international.com/) predicted possible binding of C-myc and USF to this sequence, however, qRT-PCR results showed C-myc and USF are abundant in all three cell lines (data not shown). Since there is very limited data available regarding the transcriptional regulation of CAR, the exact function of this sequence and the candidate binding proteins remain unclear.

An e-box is a special DNA sequence (CANNTG) found in some promoter regions in eukaryotes that acts as a binding site for regulating proteins (Massari and Murre, 2000). It has been reported that methylation of the CpG dinucleotide within the e-box sequence can block the binding of N-Myc to its targeting gene promoters *in vivo*, thus contributing to the cell type-specific expression pattern of certain genes, like Caspase 8 and EGFR (Perini et al., 2005). Since two e-box sequences (CACGTG) are located near the transcription start site and translation start site of CAR respectively, methylation on the CpG dinucleotide is likely to play a role in modulating CAR expression. In ourstudy, twenty-nine liver tumor samples showed decreased CYP2C19 and CAR expression along with elevated DNA methylation around the e-box binding site compared with adjacent normal liver samples. Only in sample 246564 did the situation reverse. Further scrutiny of the diagnostic information of this patient did not reveal anything unusual. Clearly, this distinct phenomenon awaits more data to be better understood. However, the reversed correlation between

CYP2C19 and CAR expression level and DNA methylation was also observed in sample 246564, which confirms the DNA methylation status might serve as a potential marker for predicting changes in gene expression.

Nevertheless, other factors cannot be excluded that contribute to the epigenetic regulation of CYP2C19 since the distal promoter region of CYP2C19 is unidentified and more transcriptional factors might exist. However, reduced CAR expression by e-box methylation at least partially account for the suppression of CYP2C19 in the HBV-infected HCC tissue samples.

It has been well documented that DNA methylation in different genomic contexts exerts different impact on gene transcription (Jones, 2012). Much attention has been paid to those CpG-rich promoters during the past years while CpG-poor transcription start sites were overlooked. In fact, non-CpG island methylation appears to be more dynamic and therefore more changeable during development or pathogenesis. For example, the CpG-poor LAMB3 promoter was shown to be hypomethylated in primary bladder tumors. *In vitro* methylation of the promoter can directly lead to transcriptional silencing (Han et al., 2011). Furthermore, enhancers, which can reside at variable distances from promoters and are mostly CpG-poor, also tend to show a variable methylation status.

Lempiäinen et al., have demonstrated that long-term activation of CAR in mice can lead to HCC, which is accompanied by a switch in the methylome (Lempiäinen et al., 2011). However, it remains controversial whether CAR plays a direct role in human HCC. The pathogenesis of cancer is a complex process in which multiple genetic and

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epigenetic factors are regulated abnormally. This complexity poses a formidable challenge to holistic integration of information when investigation is limited to only a certain gene. Traditionally, epigenetic researches on cancer were mostly focused on tumor suppressors or oncogenes. Only in recent years has the necessity of building a comprehensive epigenome map been realized. Compared with the well-established Human Genome Project, the International Human Epigenome Consortium just launched in 2010, which aims at understanding how the epigenome has shaped human populations over generations, and how it is responded to the environment (http://ihec-epigenomes.org/). Goals of this project include generating high resolution maps of DNA methylation and informative histone modifications, as well as comparative analysis of epigenome maps between healthy and diseased status. It is highly anticipated that advances in the field of epigenetics will shed light on those intricate life processes such as tumorigenesis and aging.

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Authorship Contributions

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Footnotes

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Legends for Figures

Figure 1. Location of the sequences examined by BSP, MSP and reporter gene assay. CpG sites are shown as vertical bars with dots. Transcription start sites (TSS) are shown as curved arrows and are marked as position +1. E-box binding sites are pointed by down arrows. Double-arrow bars indicate sequences examined by BSP and the capped line indicates sequence examined by MSP. Sequence in box indicates CpG island identified by Methyl Primer Express® Software v1.0 (ABI, Foster City, CA). The sequences analyzed by reporter gene assay are shown as horizontal thick lines marked with starting and ending position.

Figure 2. Maps of the CpG-free vectors used in the reporter gene assay.

Figure 3. Expression profiles of CYP2C19 and related transcriptional factors in primary hepatocytes after 5-aza-dC treatment. The relative expression of CYP2C19, CAR, GATA-4, HNF-3 γ , PXR, GR α and HNF-4 α are shown as mean ± S.E. *p**<0.05. X-axis indicates the concentration of 5-aza-dC. Data are from 4 independent experiments of 2 donors.

Figure 4. DNA methylation of CpG sites in (a) segment 1, 2 and 3 on the promoter region of CYP2C19, (b) segment 5 near the CAR transcription start site, (c) segment 6 near the CAR translation start site, (d) segment 4 on the promoter region of CAR. CpG sites of the control group and 1µM 5-aza-dC treated group are shown as red

circles and green triangles, respectively. Data are representative of the female donor.

Figure 5. Relative expression of CYP2C19 and CAR in 30 HCC tumorous tissues compared with matched adjacent normal tissues. (a) Comparison of CYP2C19 and CAR expression in matched HCC tumorous tissues and adjacent normal tissues by Paired Student's t test. Data are shown as scatter plots with mean \pm S.E., *p****<0.001. (b) Representative CYP2C19 and CAR expression in matched HCC tumorous tissues (C) and adjacent normal tissues (N). Data are shown as mean. (c) The Pearson correlation between mRNA expression of CYP2C19 and CAR.

Figure 6 Methylation analysis of segment 5 near the CAR transcription start site. (a) Comparison of methylation status of CpG sites within segment 5 between HCC tumorous tissues (red circles) and adjacent normal tissues (green triangles). (b) MSP analysis of the e-box binding site within segment 5. DNA samples from tumorous and normal tissue are shown as C and N. Segment 5 amplified by PCR was used as either methylated control DNA (Me, M. SssI treated) or unmethylated control DNA (Un). Methylation-specific primers are shown as M and unmethylation-specific primers are shown as U.

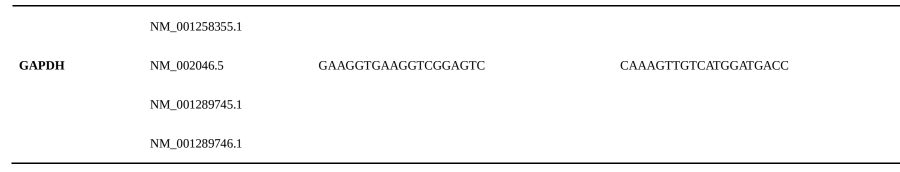
Figure 7 Effect of DNA methylation on the transcriptional activity of CYP2C19 and CAR promoter constructs. (a, c, d) The pCpGL plasmids were co-transfected with pRL-TK in HepG2, HEK293 or HK2 cells. The pCpGL plasmids were either

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methylated in the presence (methylated) or absence of (unmethylated) M. SssI. (b) The CpGfree constructs were co-transfected with pCpGL-2C19 and pRL-TK in HepG2 cells. The CpGfree constructs were either methylated in the presence (methylated) or absence of (unmethylated) M. SssI. The relative luciferase activity was calculated as ratio of firefly luciferase to Renilla luciferase. Data are shown as mean \pm S.D. with 3 replicate wells for each group. Data are representative of 3 independent experiments. $p^{**}<0.01$.

Table 1	Primer	sequences	for qRT-PCR

	Accession No.*	Forward primer (5'-3')	Reverse primer (5'-3')
CYP2C19	NM_000769.2	GAAAAATTGAATGAAAACATCAGGATTG	CGAGGGTTGTTGATGTCCATC
CAR	NM_005122.4	GCAGCTGTGGAAATCTGTCA	CAGGTCGGTCAGGAGAGAAG
GRα	NM_000176.2	GGGGAAGAGGGAGATGGA	GTGGTCAGAATGGGAGGC
PXR	NM_022002.2 NM_033013.2	CAAGCGGAAGAAAAGTGAACG	CTGGTCCTCGATGGGCAAGTC
	NM_003889.3		
GATA-4	NM_001308094.1	CTGGCCTGTCATCTCACTACG	GGTCCGTGCAGGAATTTGAGG
	NM_001308093.1		
	NM_002052.4		
ΗΝΕ-3γ	NM_004497.2	ATGCTGGGCTCAGTGAAGAT	CAGGTAGCAGCCATTCTCAA
HNF-4α	NM_178849.2	GGTCAGGTGGGGTGGATGATATAATG	TCTAGGTTAATAGGGAGGAAGGGAGG
	NM_178850.2		



*Primers were designed to amplify either all transcription variants or variants that predominate in liver.

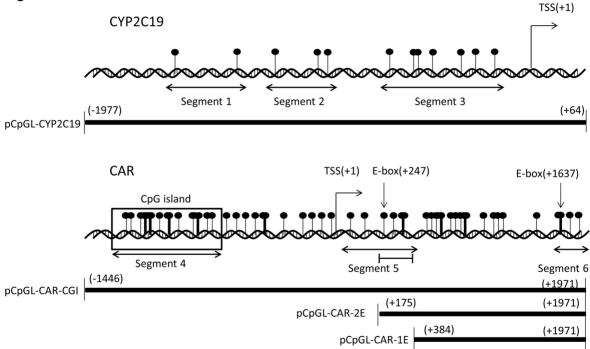
Table 2 Primer sequences for BSP and MSP

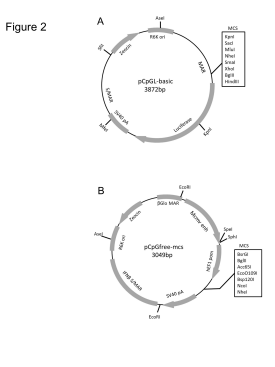
	Gene ID	Forward primer (5'-3')	Reverse primer (5'-3')
Primers for BSP			
CYP2C19 segment 1	1557	GGTAGTATATTTTTGTTAGAGTTTAAAG	AATCCACAAATATTTACTAATTATATACA
CYP2C19 segment 2	1557	GAGAATTGGAAATAATTTTATTAGG	ACCACCACAATATATAAACCTTAAC
CYP2C19 segment 3	1557	GTTTTGTTTGTTAAAATAAAGTTTTAGT	AACCCAAATAATTCCAATACACC
CAR segment 4	9970	GTAAATGAAGAAGTAGAAGATATAATAG	AATAACTCATACCTATAATCCCAACA
CAR segment 5	9970	TAGTTATTGAGAGTAATTGGAGGTTATA	CCTTAACTTCCCAAAATACTAAAATTACA
CAR segment 6	9970	TTAGAAGGGATAGAAAAGGGTTAAGG	ACTATATCCATCAAACAAACATTCAAC
Primers for MSP			
CAR-MSP-M	9970	GTTGGGGGTTAGGTATAGTGGTTTACGA	CTAAAATTACAAACATAAACTCCCGCGT
CAR-MSP-U	9970	TTGGGGGTTAGGTATAGTGGTTTATGA	TACTAAAATTACAAACATAAACTCCCACAT

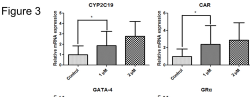
	Gene ID	Forward primer (5'-3')	Reverse primer (5'-3')
pCpGL-CYP2C19	1557	GGactagtATTATTCTAAAGAGAGCAAC	CGggatccTTAAGACAACCGTGAG
pCpGL-CAR-2E	9970	GGactagtGAAATCTGTTGGGGGACCAAAGACTAA	CGggatccCTGTGTCCATCAGACAAACATTCAG
pCpGL-CAR-1E	9970	GGactagtCCTGTAATCCCAGCACTTTGGGAAGCC	CGggatccCTGTGTCCATCAGACAAACATTCAG
pCpGfree-CAR	9970	Flanking-F	Flanking-R
		GGactcgtCTCTCTCTCTCTCTCCCAGCTTG*	TACTGGCCATGACGTCACGTGTTGGG
	NM_005122.4	CDS-F	CDS-R GAagatctCCTCAGCTGCAGATCTCCTGGA
		CGTGACGTCATGGCCAGTAGGGAAGATGAG	

*Lower-case letters indicated the added restriction sites; underlined letters indicated the overlapping part of the primers for overlap PCR.

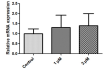


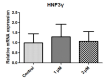




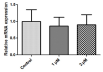


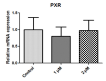


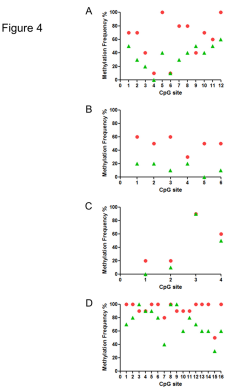


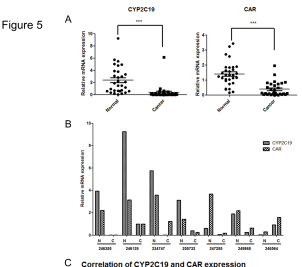














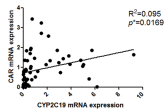
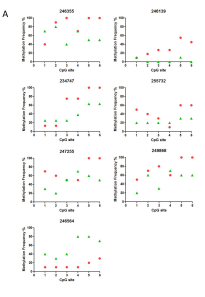


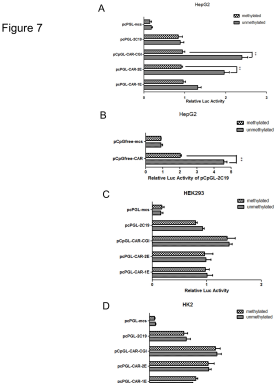
Figure 6

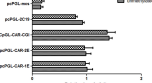


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