Abstract. Hepatitis B virus has been linked to the pathogenesis and carcinogenesis of hepatocellular carcinoma. Components of viruses have been identified within pathological specimens of hepatocellular carcinoma tissue. We characterized the in vitro response of human normal liver cells (L-02 cells) to components of infectious agents related to toll-like receptors. Immortalized human normal liver cells (L-02 cells) exhibited increased proliferation in response to exposure to CpG DNA. This molecule is a well-characterized surrogate for DNA viruses, which are common in the liver. Our experiments show that L-02 cells and some hepatoma cell lines such as HepG2, HuH7, Hep3B, express TLR 9 (CpG-specific). CpG DNA, HBV DNA, DNA of HBV middle envelope protein (MP) containing a number of CpG, supernatant of HepG2.2.15 (HepG2 cells transfected HBV) excreting HBV DNA and extraction of nucleic acids from HepG2.2.15 supernatant can all activate NF-κB. In addition, L-02 cells were less susceptible to TNF-α-induced apoptosis as measured by Annexin V-FITC staining when stimulated with CpG. mRNA of DNA methyltransferase 1 (DNMT-1) and BCL-2 was increased when L-02 cells were stimulated with CpG DNA. Our study has identified a possible novel mechanism that indicates how CpG DNA of HBV DNA may contribute to the malignant transformation of benign liver cells.

Introduction

HCC is the main cause of death in cirrhotic patients and has become a major health problem in many countries. As well, HCC is one of the most common malignancies worldwide, with an estimated annual incidence of 1 million cases (1). While development of the disease is multi-factorial including HBV, hepatitis C virus (HCV), aflatoxin, alcohol consumption and hemochromatosis (2). The toll-like receptors (TLRs) represent a primary defense against pathogens in the innate immune response (3). More than 10 mammalian Toll receptors have been identified and are involved in the recognition of microbial patterns (4). Each TLR recognizes a different, highly conserved structural motif found on many pathogens that results in the activation of transcription factors such as NF-κB. NF-κB activation is a very important reason for cell carcinization. CpG DNA is the ligand of TLR9. HBV DNA is rich in CpG. If these agents can activate NF-κB is unknown. We hypothesized that exposure of normal liver cells to components of viral pathogens might alter the character and proliferation. To test this hypothesis we used an in vitro approach employing cultured human normal liver L-02 cells treated with CpG DNA, HBV DNA, etc. We demonstrate that exposure to CpG DNA may contribute to hepatoma carcinogenesis.

Materials and methods

Cell culture and reagents. L-02 cells, HepG2, HuH7, Hep3B, HepG2.2.15 (excreting HBV DNA and transfected full-long HBV DNA) cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM, Invitrogen Corporation, Carlsbad, USA) supplemented with 10% heat-inactivated fetal bovine serum (FBS) (Hyclone Corporation, Logan, USA) and 1:100 penicillin/streptomycin at 37˚C in 5% CO2 atmosphere. HepG2.2.15 cells were supplemented with G418 (200 μg/ml). TNF-α was from PeproTech Inc (cat NO. 300-01A, Rocky Hill, NJ, USA) and 1:100 penicillin/streptomycin at 37˚C in 5% CO2 atmosphere. HepG2.2.15 cells were supplemented with G418 (200 μg/ml).

Immunocytochemistry stain for TLR9. For the immunocytochemistry staining, the cells were planted on glass coverslips in normal culture medium. The cells were fixed with 4% paraformaldehyde-phosphate-buffered solution (PBS) for 20 min at 4˚C, after which they were permeabilized with 0.5% Triton-100 for 10 min. Then blocking for 30 min the samples with 10% bovine serum album (BSA), mouse monoclonal antibody to TLR9 was applied to the coverslips. Horseradish peroxidase (HRP)-conjugated anti-mouse antibody diluted 1:500 in TBS was used to visualize the staining. At last, hematoxylin was used to stain cell nucleus. The samples were then examined using a light microscope.
Cell proliferation assay. Cell viability was evaluated via 3-[4,5-dimethylthiazol-2-yl]-2, 5-diphenyl tetrazolium bromide (MTT) reduction assay. Three thousand cells per well were plated in a 96-well plate and subjected to CpG DNA in increasing concentrations. Six parallel wells were set. The sequence of the non-methylated CpG DNA was 5'-TCGTCG TCCTGCAAGCGCTGAT-3'. The control sequence is non-CpG DNA, 5'-TCGTCGTCCTGCAAGCGCTGAT-3'. NF-κB activation is to be confirmed via immunofluorescence staining to NF-κB p65 subunit in cell nucleus, whereas NF-xB expresses in cytoplasm before activated. Cells (5x10^5) were planted on glass coverslips in 6-well plate in normal medium. After 24 h supernatant was removed and serum-free medium was added CpG (1 μM) or non-CpG (1 μM) or HBV DNA (4 μg/ml) or DNA of HBV envelope middle protein (2 μg/ml) or DNA of S envelope protein (SP) (2 μg/ml) for 4 h. In addition, supernatant of HepG2.2.15 cells or HepG2 cells (5x10^5 cells were cultured for a day) were in the place of medium, then used to stimulate human normal liver L-02 cells for 4 h. HepG2.2.15 supernatant and HepG2 supernatant (5x10^5 cells were cultured for 48 h) were used to extract nucleic acids with Phenyl hydrate/chloroform and ethanol precipitation. Extraction of nucleic acids from HepG2.2.15 supernatant and HepG2 supernatant was in 1 ml serum-free medium of L-02 cells for 4 h. Diamidino-phenyl-indole (DAPI) staining was used to locate the nucleus. The number of positive cells magnified x200 for NF-xB nuclear translocation staining and 16 fields were counted. AVONE was performed to analyze statistical significance.

Reverse transcriptase PCR for TLR9 and real-time reverse transcriptase PCR for DNMT-1 and Bcl-2. RNA was harvested using a RNA isolation kit (Tiangen, Beijing, China) acting on the manual. RT PCR was performed using the SuperScript III Cells Direct cDNA Synthesis kit (Taraka Biotechnology (Dalian) Co., Ltd., Dalian, China). TLR9 primers were as follows: upstream primer 5'-CATGCCCCTGGCGCTTCTA TTC-3', downstream primer 5'-AGCTTGGCGAGCTTG TTAGG-3'. The expected amplification size was 518 bp. Amplification size was compared to a 600 bp Plus DNA ladder (Tiangen Corporation). PCR was performed at 94˚C for 10 sec, then 40 cycles of 95˚C for 5 sec, 60˚C for 31 sec. The amplification results were analyzed using SDS·30 sec, 58˚C 30 sec and 72˚C 40 sec for 30 cycles. PCR were performed in triplicate. Paried t-test was performed subtract the average value for the blank. AVONE was performed to analyze statistical significance.

Cell proliferation assay. Cell viability was evaluated via 3-[4,5-dimethylthiazol-2-yl]-2, 5-diphenyl tetrazolium bromide (MTT) reduction assay. Three thousand cells per well were plated in a 96-well plate and subjected to CpG DNA in increasing concentrations. Six parallel wells were set. The sequence of the non-methylated CpG DNA was 5'-TCGTCG TCCTGCAAGCGCTGAT-3'. The control sequence is non-CpG DNA, 5'-TCGTCGTCCTGCAAGCGCTGAT-3'. NF-κB activity was measured by using NF-κB Activation-Nuclear Translocation Assay kit (Beyotime, Suzhou, China). NF-κB activity is to be confirmed via immunofluorescence staining to NF-κB p65 subunit in cell nucleus, whereas NF-xB expresses in cytoplasm before activated. Cells (5x10^5) were planted on glass coverslips in 6-well plate in normal medium. After 24 h supernatant was removed and serum-free medium was added CpG (1 μM) or non-CpG (1 μM) or HBV DNA (4 μg/ml) or DNA of HBV envelope middle protein (2 μg/ml) or DNA of S envelope protein (SP) (2 μg/ml) for 4 h. In addition, supernatant of HepG2.2.15 cells or HepG2 cells (5x10^5 cells were cultured for a day) were in the place of medium, then used to stimulate human normal liver L-02 cells for 4 h. HepG2.2.15 supernatant and HepG2 supernatant (5x10^5 cells were cultured for 48 h) were used to extract nucleic acids with Phenyl hydrate/chloroform and ethanol precipitation. Extraction of nucleic acids from HepG2.2.15 supernatant and HepG2 supernatant was in 1 ml serum-free medium of L-02 cells for 4 h. Diamidino-phenyl-indole (DAPI) staining was used to locate the nucleus. The number of positive cells magnified x200 for NF-xB nuclear translocation staining and 16 fields were counted. AVONE was performed to analyze statistical significance.

Cellular apoptosis analysis. Apoptosis was analyzed by fluorescence activated cell sorter (FACS) employing Annexin V-fluorescein isothiocyanate (FITC) apoptosis detection kit (Keygen Biotech. Co. Ltd. Nanjing, China). Cells Direct cDNA Synthesis kit [Taraka Biotechnology (Dalian) Co., Ltd., Dalian, China]. TLR9 primers were as follows: upstream primer 5'-AGGGAAAGGGGAGCGA-3', downstream primer 5'-AGGAAACATTCAGGTCG-3'. Bcl-2 primers were as follows: upstream primer 5'-CATGCCCTGCGCTTCATAG-3', downstream primer 5'-AGAAAACATCCAGGGTCCG-3'. Bcl-2 Activation-Nuclear Translocation Assay kit (Beyotime, Suzhou, China). Cells were treated for 24 h, after which supernatant was removed and 10% MTT serum-free medium 100 μl was added to each well and incubated at 37˚C in 5% CO2 atmosphere. For an additional 4 h, supernatant was removed carefully and 100 μl fetal dimethyl sulfoxide (DMSO) was added in each well. After shake slightly for 10 min, absorbance was read with filter in the wavelength 570/630 nm. Determine the average values from triplicate readings and subtract the average value for the blank. AVONE was performed to analyze statistical significance.

Statistical analyses. Data are presented as mean ± standard deviation (SD). Statistical analysis was performed using AVONE or paried t-test analyses using SPSS11.5 soft ware. P<0.05 was considered to be statistically significant.

Results

TLR9 are present on L-02 cells and HepG2, HuH7, Hep3B cells. RT-PCR confirmed the presence of TLR9 (Fig. 1A). TLR9 (Fig. 1B) were expressed on L-02 cells. Quantitation was accomplished by measuring the incorporation of the fluorescent dye SYBR-Green-II into the PCR product. PCR was programmed for 95˚C for 10 sec, then 40 cycles of 95˚C for 5 sec, 60˚C for 31 sec. The amplification results were analyzed using SDS2.2.1 software (Applied Biosystems) and the genes of interest were normalized to the corresponding GAPDH results. Data were expressed as fold induction relative to the control. All PCR were performed in triplicate. Paried t-test was performed to analyze statistical significance.

CpG DNA stimulate proliferation of human normal liver L-02 cells. We determined whether CpG DNA could influence live cell line L-02 cell proliferation as measured by MTT reduction assay. Different doses of CpG DNA (0.1, 1, 5 and 10 μM) or non-CpG DNA (0.1, 1, 5 and 10 μM) were used to stimulate L-02 for 24 h. CpG DNA caused an increase in cell prolifer-ation at all doses from 0.1 to 10 μM, maximum level was seen at the dose 1 μM (Fig. 2A). Non-CpG DNA did not stimulate the proliferation of L-02 cells (Fig. 2B). These results show that both human normal liver L-02 cells
are capable of a proliferative response to CpG DNA, but not to non-CpG DNA. Thus, we infer that CpG components of DNA of hepatitis virus can directly influence liver cells proliferation.

CpG DNA activates NF-κB. NF-κB activation was confirmed via immunofluorescence staining to NF-κB p65 subunit in cell nucleus. Concentrations of 1 μM of CpG DNA were used for 4 h because they demonstrated a robust proliferative response in the MTT reduction assays. Fig. 3A1 demonstrates that CpG can activate NF-κB, and the control, non-CpG DNA (1 μM), can not activate NF-κB (Fig. 3A3).

HBV DNA, MP DNA, cell supernatants and extraction of cell supernatants activate NF-κB. In order to study whether HBV DNA can activate NF-κB directly, Pblue sk-/HBV DNA containing HBV DNA full-length sequence was employed as the template to amplify HBV DNA. MP DNA and SP DNA as the control of MP DNA were amplified. HBV DNA primers were as follows: upstream primer 5'-TTT TTCACCTCTGCTAAATCA-3', downstream primer 5'-AAA AAGTTGCAATGTTGCTGG-3'. MP DNA primers were as follows: upstream primer 5'-ATGGGAGGTGGTCTTCC-3', downstream primer 5'-TCAAATGTATACCCAAAGAC-3'. SP DNA primers were as follows: upstream primer 5'-ATG GAGACCAACAA-3', downstream primer 5'-TCAAAT GTATACCCAAAGAC-3'. After 3 min of prodegeneration, for HBV DNA, PCR was performed at 94˚C for 40 sec, 65˚C for 5 min and 72˚C for 10 min for 30 cycles; for MP DNA and SP DNA, PCR was performed at 94˚C for 30 sec, 50˚C for 30 sec and 72˚C for 50 sec for 30 cycles. PCR products were separated on a 1% agarose gel and were extracted by a Gel Extract Kit. These products and the supernatant of HepG2.2.15 or HepG2 cells and extraction of nucleic acids from HepG2.2.15 supernatant and HepG2 supernatant were used to stimulate human normal liver L-02 cells. An interesting result was found that HepG2.2.15 supernatant and HBV DNA and MP DNA and extraction of nucleic acids from HepG2.2.15 can make NF-κB nuclear translocation and activate NF-κB, but the control (HepG2 supernatant and equal volume ster deionized water and SP DNA and extraction of nucleic acids from HepG2 supernatant respectively) nearly cannot (Fig. 3B1-E4). This is shown quantitatively in Fig. 3F where there is a significant increase in the number of NF-κB activated cells in each field (p<0.01). However the control groups were hardly activated.

CpG DNA inhibits TNF-α induced apoptosis. Toll-like receptor stimulation is known to up-regulate various anti-apoptotic genes; therefore, we tested the ability of CpG DNA to prevent apoptosis induced by TNF-α of human normal liver L-02 cells by measuring Annexin V-FITC and PI.
Fig. 4A-C shows apoptosis in non-stimulated and CpG DNA-stimulated and non-CpG DNA-stimulated human normal liver L-02 cells at <10%. Apoptotic cells are about 30% using TNF-α (100 ng/ml) for 24 h (Fig. 4D). Fewer cells underwent TNF-α induced apoptosis when treated with CpG DNA (Fig. 4E). Apoptotic cells are about 20% with non-CpG DNA treatment (Fig. 4F). Values are the means ± SD of three independent experiments.

**CpG DNA increased mRNA expression of DNMT-1 and Bcl-2.** Regional DNA hypermethylation and increased DNMT-1 protein expression participate in the precancerous stage, in malignant progression, and have a prognostic impact on patients with cancers (5). The proto-oncogene, Bcl-2, encodes a protein that inhibits programmed cell death (apoptosis) and play a role in cell and tissue differentiation (6). Therefore, we studied the mechanism of CpG DNA induced-carcinogenesis from these two aspects by real-time...

**Figure 3.** Different agents induce NF-κB nuclear translocation in L-02 cells. Indirect immunofluorescence analyses for cellular NF-κB localization were performed using chamber slide cultured L-02 cells. Nucleus location was marked by DAPI staining. Activated-NF-κB is localized in the nucleus and unactivated-NF-κB is localized in the cytoplasm. p65 subunit of NF-κB was stained by Cy3 representing NF-κB. NF-κB was activated by 1 μM CpG DNA (A1 and A2), supernatant of HepG2.2.15 cells (B1 and B2), 4 μg/ml HBV DNA (C1 and C2), 2 μg/ml MP DNA (D1 and D2) and extraction of Nucleic acids from HepG2.2.15 supernatant (E1 and E2). However NF-κB was not activated in the control group (respectively, in turn, non-CpG DNA (A3 and A4), HepG2 supernatant (B3 and B4), equal volume ster deionized water (C3 and C4), SP DNA (D3 and D4) and extraction of nucleic acids from HepG2 supernatant (E3 and E4). Corresponding to the above A-E groups, the number of NF-κB-activated cells in test groups are significantly increased compared with control groups (**p<0.01). The qualified analyses on the number of NF-κB activated cells in 16 fields amplified x200 (F).
RT-PCR. We found after L-02 cells were stimulated with 1 μM CpG DNA for 4 h, comparing with control group, DNMT-1 increased significantly 1.87-fold (1.63±0.27) and Bcl-2 increased significantly 3.49-fold (3.49±0.72). Thus, CpG DNA induced DNMT-1 and Bcl-2 production which possibly promoted carcinogenesis (Fig. 5).

Discussion

Inflammation and infection have been shown to be important in the pathogenesis of liver, colon, esophagus, stomach, cervical, and nasopharyngeal cancer by causing cell damage and creating a microenvironment rich in cytokines that can enhance cell replication, angiogenesis and tissue repair (7). It is well known that HBV leads to HCC. HCC incidence clearly correlates with HBV infection as 80% of HCC occur in HBV-infected patients (8). Some studies on HBV-induced HCC focus on the integration of HBV DNA (9-11), HBx (12,13), the PreS2 activators (14), and the immune system-mediated chronic inflammation of the liver (15-17). However, no study was found on HBV DNA as an inflammation element directly triggering cell cancerization in extra-hepatocytes. CpG DNA which is a major ligand of TLR9, but not the only one (18), exists in the double strands of virus and bacteria. We discovered that TLR9 is expressed on human normal liver L-02 cells and hepatoma cell lines (HepG2, HuH7 and Hep3B), and exists in human hepatocytes (19). The proliferation of L-02 cells was found with the stimulus of the different concentration of CpG DNA as the effect on the other tumor cell lines (20). This result suggests that the liver cells are capable of responding to components of pathogens by initiating cellular proliferation.

NF-κB is correlated with the HCC significantly which is bridging the action of growth factors and inflammation to hepatic oncogenesis (21). Recent findings from several laboratories have implicated constitutive activation of NF-κB as one of the early key events involved in neoplastic progression of the liver in the Mdr2-knockout mouse model. Blocking of NF-κB activity via hepatic expression of a Tet-inducible super-repressor form of IκB-a can block tumor growth (22,23). In addition, a recently published study demonstrated that NF-κB inhibited c-Myc induced activation of caspase-9 and -3 through upregulation of the anti-apoptotic target genes Bcl-XL and XIAP (24). We hypothesize that components of infectious agents may further contribute to carcinogenesis through a previously unreported method and shift the balance towards uncontrolled growth. CpG DNA activates the NF-κB and triggers a serial of downstream effect via TLR9 (25). The CpG DNA island is rich of single copy nonmethylation locus in the genome. We found that the sequence of HBV DNA is rich of CpG, and concentrated on sequence of pre S2 site encoding MP of HBV but not S site encoding SP, by online (address: http://www.bi osoft/sms) software. We employ these agents to stimulate the L-02 cells. NF-κB was remarkably activated. We infer that HBV DNA can induce directly human normal liver L-02 cell cancerization or malignant change.

Figure 4. CpG DNA pretreatment induces resistance of L-02 cells to the apoptosis induction by TNF-α. L-02 cells (5x10^5/ml) were pretreated without (A) or with 1 μM CpG DNA (B) or 1 μM (A) non-CpG DNA (C) for 24 h. L-02 cells (5x10^5/ml) were treated with TNF-α (100 ng/ml) for 24 h (D). L-02 cells (5x10^5/ml) were pretreated with 1 μM CpG DNA or 1 μM non-CpG DNA for 4 h, then stimulated with 100 ng/ml TNF-α for 24 h (E and F). The cells were stained with Annexin V-FITC and PI and subjected to FACS assay of cellular apoptosis. Values are the means ± SD of three independent experiments. Fewer cells underwent TNF-α induced apoptosis when treated with CpG DNA.

Figure 5. Effects of CpG DNA (1 μM) on mRNA expression of DNMT-1 and Bcl-2. Each experiment was conducted 3 times. Data are expressed as mean ± SD. DNMT-1 and Bcl-2 mRNA increased 1.87-fold (1.63±0.27) (mean ± SD) and 3.49-fold (3.49±0.72) respectively. Compared with control, there was a significant increase (‘p<0.05).
Studies have shown that NF-xB may be a tumor initiator by expressing anti-apoptotic genes that promote cell survival such as c-FLIP, Bcl-2, and p53 (26). As Bcl-2 appears to be involved in the turnover of stem or precursor cells, it is thought to be operational in the carcinogenesis pathways (27-29). We found increased mRNA expression of Bcl-2 with the stimulation of CpG DNA. At the same time, we showed that treatment of benign cells with CpG DNA could protect cells from TNF-α induced apoptosis, possibly altering the delicate balance between cell growth and death that leads to malignant transformation.

Recently, a study illuminated that HBV DNA is methylated in liver tissues and is non-methylated in serum from patients with high viral load. HepG2 and HepG3B cell lines can make HBV DNA methylated (30). Some studies performed almost a decade ago showed that HBV DNA can be methylated when it becomes integrated into the human genome (31). In human HCC with integrated HBV genomes, viral protein expression was found only in those cases without methylation (30). Some studies performed almost a decade ago showed that HBV DNA methylated (30). Some studies performed almost a decade ago showed that HBV DNA methylated (30).


