Effect of Ginkgo biloba extract on the expressions of Cox-2 and GST-Pi in rats with hepatocellular carcinoma risk

Chao Ou 1 *, Hai-Ping Zheng 2 *, Jian-Jia Su 1*, Ji Cao 1, Guo-Jian Li 2, Le-Qun Li 3

1. Department of Experimental Pathology, The Affiliated Tumor Hospital of Guangxi Medical University, Nanning, Guangxi Province 530021, China
2. Department of Medical Oncology, The First Affiliated Hospital of Guangxi Medical University, Nanning, Guangxi Province 530021, China
3. Department of Hepatobiliary Surgery, The Affiliated Tumor Hospital of Guangxi Medical University, Nanning Guangxi Province 530021, China

*These authors contributed equally.

Abstract
Background: Hepatocellular carcinoma (HCC) is one of the most common and aggressive cancers worldwide, and the pathogenesis is complicated at present. There are few effective therapeutic measures, and novel therapeutic strategies are urgently required to improve clinical outcome. Ginkgo biloba extract (EGb) is reported to have an anti-cancer activity.

Objectives: To explore the effect of EGb on expressions of cyclooxygenase-2 (Cox-2) and glutathione S-transferase Pi (GST-Pi) in the pathogenesis of HCC.

Methods: 120 Wistar rats were divided into three groups at random: normal control group (control group), HCC risk group without treatment (HCC risk group), HCC risk group treated with EGb (EGb group); n=40, respectively. The HCC risk in rats was induced by aflatoxin B1 injection. At the end of 13-week, 33-week, 53-week and 73-week, 10 rats in each group were killed and the relevant samples were collected.

Results: The mRNA and protein expressions of Cox-2 and GST-Pi were measured by real-time reverse transcription polymerase chain reaction, immunohistochemical analysis and western-blot. When compared with those in the control group in 73-week, the mRNA and protein expressions of GST-Pi in EGb group were weaker than those in HCC risk group in 73-week. However, the mRNA and protein expressions of Cox-2 in HCC risk group were increased than that of control group, and there was no statistical difference for mRNA and protein expressions of Cox-2 between HCC risk group and EGb group.

Conclusion: EGb can regulate the expression of GST-Pi, but it does not seem to have an effect on Cox-2 expression in the liver of HCC risk rats.

Keywords: Hepatocellular carcinoma (HCC); Ginkgo biloba extract (EGb); Cox-2; GST-Pi

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Introduction
Hepatocellular carcinoma (HCC) is one of the most lethal and prevalent liver [1, 2]. HCC is highly prevalent in many Asian countries, accounting for 80% of victims worldwide [3, 4]. Most HCC patients show chronic hepatitis or cirrhosis caused by persistent infection with hepatitis B virus (HBV) or hepatitis C virus (HCV) [5-7]. Aflatoxin B1 (AFB1) is also very important in the onset of HCC. Hepatectomy is regarded as the optimal curative treatment for HCC at present; however, the postoperative outcome remains unsatisfactory [8]. Its underlying biological mechanism remains unclear and no well-documented drug and targeting has been approved to date [9].

Cyclooxygenase 2 (Cox-2) is expressed highly in cancer and it is found to be implicated in tumor progression, and its inhibition can reduce tumor growth and augment therapy [10, 11]. Glutathione S-transferase Pi (GST-Pi) is a subgroup of GST family, which provides cellular protection against free radical and carcinogenic compounds due to its detoxifying function [12].
was no report studying the expressions of Cox-2 and GST-Pi in the HCC rats induced by AFB1. In this study, we investigated the association of Cox-2 and GST-Pi expressions with HCC risk in rats induced by AFB1.

Ginkgo Biloba is one of the oldest living tree species and it has been described as a living fossil. Its beneficial effects were known 5000 years ago in traditional Chinese medicine. The study of the biological activities of Ginkgo Biloba extract 761 (EGb 761), a standardized extract of Ginkgo Biloba with a well-defined mixture developed more than 20 years ago, has gained great popularity in European countries and the USA [13]. EGb 761 has been standardized to ensure the consistency of its composition and a reliable, safe, efficacy profile [13]. The EGb 761 extract exhibits important properties, such as anti-oxidation, anti-apoptosis and anti-inflammation. At present, some studies [14-16] reports that EGb can take an anti-cancer role. Whether there was an association between EGb and Cox-2 / GST-Pi expression in the HCC risk rats induced by AFB1, there was no any report. In our study, we investigated the alteration of mRNA and protein expressions of Cox-2 and GST-Pi in liver tissue in a rat with HCC risk, treated with EGb to explore the effect of EGb on expressions of Cox-2 and GST-Pi in the pathogenesis of HCC risk.

Materials and methods

Animal model

120 healthy male rats, 180g to 200g, of Wistar backgrounds were purchased from the Experimental Animal center of Guangxi Medical University, Nanning, China. All procedures were approved by the animal ethics committee of Guangxi Medical University. The rats were divided into three groups at random: normal control group (control group), HCC risk group without treatment (HCC risk group), HCC risk group treated with EGb (EGb group); n=40, respectively. The risk of HCC in rat was induced by intraperitoneal injection using aflatoxin B1 (AFB1) as follow: 4-7 weeks and 9-12 weeks, 200μg/kg, thrice a week; 14-17 weeks, 19-22 weeks, 24-27 weeks, and 29-32 weeks, 100μg/kg, twice a week; 34-73 weeks, 100μg/kg, once a week. The rats in EGb group were treated with EGb761 from the first week to the end (2g/kg.d; Sitexin, co.; Guilin, China). At the end of 13-week, 33-week, 53-week and 73-week, 10 rats in each group were killed and the relevant samples were collected for molecular biology determination.

Real time reverse transcription polymerase chain reaction to detect Cox-2 and GST-Pi mRNA expression in liver tissue

Renal tissue was homogenized and total RNA was extracted with TRIzol (Beijing Tiangen, Co., China). Ultraviolet spectrophotometer measuring absorbance, agarose gel electrophoresis confirmed that there had been no degradation of RNA by visualizing the 18S and 28S RNA bands under ultraviolet light [17, 18]. Primers of were designed according to primer design principles by Primer Premier 5.0. The primers for Cox-2, GST-Pi and internal control β-actin were as follows: F 5’- CCTCGTCCAGATGCTATCTT-3’ and R 5’-GAAGGTGCTAGGTTTCCAG-3’ for Cox-2; F 5’-AGATGTCTGGCTTCAAGGCT-3’ and R 5’- ATTTGCATCGAAGGTCTCC-3’ for GST-Pi; F 5’- GCCCCCTGAGGAGCACCCTGT-3’ and R 5’- ACGCTCGGTCAAGGATCTTCA-3’ for β-actin. One microgram total RNA from the liver tissue of each rat was reverse transcribed into cDNA with an ExScript RT reagent kit (Fermentas). Cox-2, GST-Pi and β-actin were amplified with SYBR Premix Ex Taq (Beijing Tiangen, Co., China). Gene expression of β-actin was also measured in each sample and used as an internal control for loading and reverse transcription efficiency. The analysis for each sample was performed in triplicate. The average threshold cycle (Ct, the cycles of template amplification to the threshold) was worked out as the value of each sample. The data for fold change was analyzed using 2−ΔΔCt [19, 20].

Immunohistochemical analysis of Cox-2 and GST-Pi

The operation was implemented using the streptavidin-peroxidase immunohistochemical method. Renal tissue samples were fixed in 10% neutral formaldehyde, dehydrated with ethanol, and embedded in paraffin. Serial 4 μm sections were collected sequentially on glass slides. The paraffin was removed from the sections with xylene and rehydrated in graded ethanol. In order to retrieve antigenicity from formalin fixation, we incubated the sections for 10 min in 10 mmol/l sodium citrate buffer using a microwave oven. Endogenous peroxidase activity was blocked by further pretreatment with 3% hydrogen peroxide and methanol. Finally, the sections were incubated with antibody against GST-Pi (Thermo Fisher Scientific, Co., Runcorn, UK), Cox-2 (Thermo Fisher Scientific, Co., Runcorn, UK) overnight at 4°C. The sections were washed thoroughly in phosphate-buffered saline (PBS) solution and incubated with rabbit anti-mouse biotinylated second antibody immunoglobulin (Maixin Bio, Co., Inc., China) for 30 min. Finally, the sections were stained with diaminobenizidine (Maixin Bio, Co., Inc., China).
We obtained negative controls by replacing specific antisera with PBS solution. Brownish yellow granular or linear deposits in the cells or matrix were interpreted as positive areas. Semi-quantitative evaluation was performed by computer-assisted image analysis (Leica Co., Germany). The positive staining of Cox-2 or GST-Pi was measured at 400-fold original magnification selected from coded sections of each rat at random.

Western-blot analysis
The total proteins of liver tissue were extracted using lysing buffer (Beyotime Institute of Biotechnology, China), following a centrifugation at 12000g for 10 minutes at 4°C, and protein content in the supernatant was detected using the BCA protein assay. Equal amounts of extract (30 μg) were then separated on an 8% sodium dodecylsulfate (SDS)-poly-acrylamide gel and transferred by electroblotting to PVDF membranes. The membranes were incubated in 5% skimmed milk for one hour and were washed with 0.1% TBST at room temperature, and followed by overnight incubation at 4°C with primary antibodies: anti-GST-Pi (Thermo Fisher Scientific, Co., Runcorn, UK), anti-Cox-2 (Thermo Fisher Scientific, Co., Runcorn, UK) and anti-β-actin (1:5000, Epit Mics). Primary antibodies were diluted using 0.1% TBST. Alkaline phosphatase conjugated secondary antibodies were incubated with blots for one hour at room temperature. After washing, blots were developed by the ECL Western Blotting detection system, and then exposed to X-ray film for visualization of the protein bands, and membranes were semi-quantified using the Quantity One image analysis system (4.3.1, Bio-Rad, USA). The expression level of GST-Pi or Cox-2 was corrected by comparison with β-actin, and the final data were expressed used the fold chance.

Statistical analysis
The data are shown as mean ± standard deviation (SD). To compare the groups in relation to parameters with normal distribution, One-way analysis of variation (ANOVA) with post-hoc Fisher’s LSD (least significant difference) was used. For parameters without normal distribution, Kruskal-Wallis with post-hoc Mann-Whitney (only for the weight parameter) was used (21, 22). A value of $P < 0.05$ was accepted as statistically significant. Statistical analysis was performed using the Statistical Package for Social Sciences (SPSS, version 13.0; Chicago, IL, USA).

Results
The Cox-2 and GST-Pi mRNA expression in liver tissue
The mRNA expression of Cox-2 in HCC risk group was similar to that of control group in 13-week or 33-week, and the difference was not statistically significant (all $P > 0.05$; Figure 1). Interestingly, the mRNA expression of Cox-2 in HCC risk group was notably increased than that of control group in 53-week and 73-week, and the difference was marked (all $P < 0.01$; Figure 1). However, there was no statistical difference for mRNA expression of Cox-2 between HCC risk group and EGb group in all the time points (all $P > 0.05$; Figure 1).
When compared with that in control group in all time points, the mRNA expression of GST-Pi in HCC risk group was increased (all $P<0.01$; Figure 2). Interestingly, the difference of mRNA expression of GST-Pi between HCC risk group and EGb group was no notably in 13-week, 33-week and 53-week. However, the mRNA expression of GST-Pi in EGb group was marked attenuated than that in HCC risk group in 73-week ($P<0.01$; Figure 2).
Figure 2 The mRNA expression of GST-Pi in three group. (: $P<0.01$ compared with control group; ): $P>0.01$ compared with AFB1 group; §: $P>0.05$ compared with AFB1 group.

**Protein expressions of Cox-2 and GST-Pi in liver tissue using immunohistochemical analysis**

Immunohistochemical staining for Cox-2 was performed. The staining for Cox-2 was markedly enhanced in HCC risk group in 53-week or 73-week when compared with those in control group (all $P<0.01$; Figure 3), but the difference was not observed in 13-week or 33-week (all $P>0.05$; Figure 3). Furthermore, the difference of Cox-2 expression between HCC risk group and EGB group was not observed in 13-week, 33-week, 53-week or 73-week. The typical immunohistochemical staining figure of for Cox-2 in 73-week was shown in Figure 3.
Immunohistochemical staining for GST-Pi was also conducted. The staining for GST-Pi was markedly enhanced in HCC risk group in 13-week, 33-week, 53-week or 73-week when compared with those in control group (all $P<0.01$; Figure 4). The GST-Pi expression in EGB group in 73-week was marked reduced when compared with that in HCC risk group ($P<0.01$; Figure 4). However, the difference of GST-Pi expression between HCC risk group and EGB group was not observed in 13-week, 33-week, or 53-week (all $P>0.05$; Figure 4). The typical immunohistochemical staining figure of for Cox-2 in 73-week was shown in Figure 4.
Protein expressions of Cox-2 and GST-Pi in liver tissue using western-blot analysis

There was no statistical difference for the protein expression of Cox-2 between HCC risk group and control group in 13-week and 33-week (all $P>0.05$; Figure 5). Interestingly, the protein expression of Cox-2 in HCC risk group was markedly increased than that of control group in 53-week and 73-week, and the difference was notably (all $P<0.05$; Figure 5). There was no statistical difference for protein expression of Cox-2 between HCC risk group and EGb group in all the time points (each $P>0.05$; Figure 5). The typical blot bands for Cox-2 were presented in Figure 5.

Figure 4 Statistical parameters of the protein expression of GST-Pi and the typical immunohistochemical staining figure for GST-Pi in three group. (): $P<0.01$ compared with control group; (): $P>0.01$ compared with AFB1 group; §: $P>0.05$ compared with AFB1 group.
The protein expression of GST-Pi in HCC risk group was increased than that of control group in all time points (all P<0.01; Figure 6). Interestingly, the difference of protein expression of GST-Pi between HCC risk group and EGb group was no notably in 13-week, 33-week and 53-week. However, the protein expression of GST-Pi in EGb group was marked attenuated than that in HCC risk group in 73-week (P<0.01; Figure 6). The typical blot bands for GST-Pi were presented in Figure 6.
Discussion

In this study, we found that Cox-2 expression in the HCC risk group was notably increased when compared with control group. Increased Cox-2 was associated with the risk of HCC in rats. The results from real time reverse transcription polymerase chain reaction, immunohistochemical analysis and western-blot analysis were consistent. Cox-2 might be taking part in the pathogenesis of HCC. Over-expression of Cox-2 has been reported to be associated with hepatocarcinogenesis. Ogunwobi et al. [23] reported that upregulation of Cox-2 could promote epithelial-mesenchymal transition and carcinogenesis in vitro, and could take part in the development of HCC. Lee et al. [24] reported that the activation of Cox-2 signaling pathway had particular relevance to HCV-associated HCC in patients with HCV-associated HCC. He et al. [25] reported that over-expression of Cox-2 in noncancerous liver regions was an independent and significant indicator predictive of early recurrence of HCC in patients with HBV-related cirrhosis. Those results mentioned above indicated that over-expression of Cox-2 was associated with the onset of HCC. It was similar to the conclusion from our study. Cox-2 might take part in the pathogenesis of HCC.

GST-Pi expression was also increased in the HCC risk group than that in control group. In our study, we found that the expression of GST-Pi was increased step by step in the HCC risk group when compared with in control group. The results from real time reverse transcription
polymerase chain reaction, immunohistochemical analysis and western-blot analysis were consistent. Elevation of GST-Pi was associated with the risk of HCC. There were also some reports finding that there might be a relationship between the expression of GST-Pi and the HCC risk. Yang et al. [26] reported that higher GST-pi expression mediated resistance to cisplatin in HCC cell line. Morsi et al. [27] reported that HCC patients had significantly higher serum GST-Pi. Niu et al. [28] found that the increased GST-Pi might be a marker enzyme for immunohistochemical detection of human HCC and its preneoplastic lesions. The conclusion from those reports mentioned above was consistent with ours. So, increased GST-Pi might be associated with the onset of HCC.

In our study, we found that the EGb had no effect on the expression of Cox-2 in rats. Jang et al. [29] found that Cox-2 expression in the cambial meristematic cells was almost completely suppressed by EGb. Bao et al. [30] reported that EGb could inhibit the production of Cox-2 in myocardial ischemia/reperfusion rats. Park et al. [31] found that EGb could inhibit the LPS-induced Cox-2 in macrophage cell line. Our result was inconsistent with that from the studies mentioned above. However, there was no other report to study the relationship between EGb treatment and Cox-2 expression in progress of HCC. More studies on HCC risk should be performed in the future.

Interestingly, we found that EGb could take a protective role and regulate the expression GST-Pi. The GST-Pi expression was weakened when compared with that in HCC risk group in 73-week. There was only one study reporting the relationship between EGb treatment and GST-Pi expression. Sasaki et al. [32] reported that EGb significantly increased the protein level of GST-Pi in the mouse liver. In our study, we didn’t found that EGb could induce the GST-Pi expression in the rats with HCC risk. Its conclusion was similar to ours. We speculated that the EGb could take the protective role against HCC risk in rats by reducing the expression of GST-Pi, but not Cox-2.

**Conclusion**
The expressions of Cox-2 and GST-Pi were increased in the rats with HCC risk, and EGb could play a protective role against HCC risk in rats induced by AFB1 by reducing the expression of GST-Pi, but not Cox-2, although the detailed mechanisms were not fully understood. At the moment, EGb is already used in clinical treatment for various diseases, such as thyroid carcinoma, Alzheimer's disease and so on. In this investigation, we have demonstrated that EGb can protect against HCC risk in rats treated with AFB1. This observation might offer some new insights to prevent the onset of HCC. However, cells culture and inhibition of signaling pathway for EGb are needed to be conducted to be sure of its effect and to explore the detailed mechanism.

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**Competing Interests:**
The authors have declared no competing interest.

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