Loss of ICG uptake in the process of rat hepatocarcinogenesis correlates to the disappearance of glutathione-S-transferase alpha subunit.

Liu Ling*  Toshihiro Higashi†  Shigeki Tsuchida‡  
Kiyomi Sato**  Takao Tsuji††

*Okayama University,
†Okayama University,
‡Hirosaki University,
**Hirosaki University,
††Okayama University,
Loss of ICG uptake in the process of rat hepatocarcinogenesis correlates to the disappearance of glutathione-S-transferase alpha subunit.*

Liu Ling, Toshihiro Higashi, Shigeki Tsuchida, Kiyomi Sato, and Takao Tsuji

Abstract

Reduced indocyanine green (ICG) uptake is one of the functional changes of human hepatocellular carcinoma (HCC). To clarify the mechanisms of loss of ICG uptake, and determine which subunit of glutathione-S-transferase (GST), alpha or pi, plays a role in ICG transport in hepatocytes, an experimental HCC model was developed that used nodules induced by 2-acetylamino-fluorene (2-AAF) administration. Many of the ICG stained nodules, which consisted of benign and borderline lesions, were GST-alpha positive. However, the percentage of GST-alpha positive cells tended to decrease according to the disappearance of ICG staining in the process of hepatocarcinogenesis. HCCs unstained by ICG were also GST-alpha negative. GST-pi, not detected in normal rat hepatocytes, appeared in an earlier stage of hepatocarcinogenesis before the disappearance of GST-alpha, and was not observed in HCCs. No significant relationship between ICG staining and GST-pi was recognized. These results suggest that GST-alpha synthesis is disturbed in the process of hepatocarcinogenesis and results in loss of ICG uptake in HCCs, and also indicate that GST-pi may be useful for early diagnosis of preneoplastic hepatocytes showing no roles in ICG transport.

KEYWORDS: hepatocarcinogenesis, indocyanine green, glutathione-S-transferase-?, glutathione-S-transferase-?

*PMID: 7505994 [PubMed - indexed for MEDLINE]
Copyright (C) OKAYAMA UNIVERSITY MEDICAL SCHOOL
Loss of ICG Uptake in the Process of Rat Hepatocarcinogenesis Correlates to the Disappearance of Glutathione-S-Transferase Alpha Subunit

Liu Ling, Toshihiro Higashi*, Shigeki Tsuchida a, Kiyomi Sato b and Takao Tsuji

First Department of Internal Medicine, Okayama University Medical School and Second Department of Biochemistry, Hirosaki University Medical School, Aomori-036, Japan

Reduced indocyanine green (ICG) uptake is one of the functional changes of human hepatocellular carcinoma (HCC). To clarify the mechanisms of loss of ICG uptake, and determine which subunit of glutathione-S-transferase (GST), alpha or pi, plays a role in ICG transport in hepatocytes, an experimental HCC model was developed that used nodules induced by 2-acetylaminoflourene (2-AAF) administration. Many of the ICG stained nodules, which consisted of benign and borderline lesions, were GST-alpha positive. However, the percentage of GST-alpha positive cells tended to decrease according to the disappearance of ICG staining in the process of hepatocarcinogenesis. HCCs unstained by ICG were also GST-alpha negative. GST-pi, not detected in normal rat hepatocytes, appeared in an earlier stage of hepatocarcinogenesis before the disappearance of GST-alpha, and was not observed in HCCs. No significant relationship between ICG staining and GST-pi was recognized. These results suggest that GST-alpha synthesis is disturbed in the process of hepatocarcinogenesis and results in loss of ICG uptake in HCCs, and also indicate that GST-pi may be useful for early diagnosis of preneoplastic hepatocytes showing no roles in ICG transport.

Key words: hepatocarcinogenesis, indocyanine green, glutathione-S-transferase-α, glutathione-S-transferase-π

Mechanisms of uptake of organic anions (e.g., bilirubin, indocyanine green (ICG) and sulfobromophthalein (BSP) by hepatocytes have been investigated in several studies (1–6). Organic anion binding proteins (4), albumin receptors (5), or an active transport mechanism (6) are supposed to regulate the uptake. Wolkoff et al. (2) have identified a BSP receptor on the plasma membrane of hepatocytes. Other reports on the mechanism of ICG uptake indicate that glutathione-S-transferase (GST), a carrier protein of ICG, plays an important role in staining in hepatocytes (7–9).

ICG uptake has not been recognized in hepatocellular carcinoma (HCC) at the level of peritoneoscopic examination (10). In our first investigation we examined the relationship between ICG staining and cell atypism of rat liver nodules induced by administration of 2-acetylaminofluorene (2-AAF) as an experimental model of human HCC. The ICG-unstained nodules were proved to be malignant by means of histological examination and DNA ploidy pattern (11).

In this report, the existence of GST-subunits (GST-alpha and -pi) of rat liver nodules obtained from the previous experiment (11) was demonstrated by immunohistochemical technique, and the relationship between ICG staining and GST-subunits was investigated in order to clarify the role of GST-subunits on ICG uptake.

Materials and Methods

An experimental model for hepatocarcinogenesis. According to the method described by Epstein et al. (12), 36 Wistar-strain
male rats (8-week-old, 170-180g) were fed on diet containing 0.05 % 2-AAF for weeks 0-3, 4-6, or 8-13 orally (as initiation and promotion). Basal diet without 2-AAF were given for weeks 3-4 and 6-8. Surviving rats were divided into two groups and fed either a diet containing 1.22 % Syo-saiko-to, or basal diet for weeks 13-26 in order to examine at various stages of development (Fig. 1).

ICG injection and histological diagnosis. The rats were killed as previously described (11) and according to the schedule in Table 1. They were anesthetized with ether and the portal vein was cannulated with a 24-gauge thin needle. ICG (2.5 mg/ml) was injected slowly for more than 1 min, to avoid disruptive pressure until liver surface was completely stained. The inferior vena cava was then incised at the juncture with the hepatic vein and perfused with 20 ml of saline. The largest nodules, ICG stained and unstained, were fixed in 10 % neutral buffered formalin for 7 days and embedded in paraffin. The specimens were sliced 6 μm in thickness with a microtome. Formalin-fixed and paraffin-embedded sections were used for hematoxylin and eosin (H. E.) staining, and immunohistochemical analysis for GST-alpha and -π.

Estimation for specificities of anti GST-alpha and -π antibodies. According to the method of Laemmli et al. (13), GST from rat liver (Sigma Co. Ltd.) was separated by electrophoresis in 10 % polyacrylamide gels in the presence of sodium dodecyl sulfate (SDS-PAGE) and were transferred to nitrocellulose sheets (Bio-Rad Laboratories). After blocking with 3 % bovine serum albumin-0.1 M phosphate buffered saline, pH7.4 (PBS), blots were overlaid with anti GST-alpha antibody (×100) or anti GST-π antibody (×100) and were incubated for 1h at room temperature with gentle agitation. The blots were then overlaid with a 1/1000 diluted solution of peroxidase labeled swine anti-rabbit IgG for 1h. Reactivity was detected using 3, 3'-diaminobenzidine-4HCl (DAB) (Sigma Co. Ltd.) as a substrate. Preimmune rabbit serum (×100) was used as a negative control.

Immunohistochemical staining for GST-alpha and -π. Anti human GST-alpha antibody defined by K. Sato and anti human GST-π (BIO PREP Co. Ltd.) rabbit polyclonal antibody were used. Formalin-fixed and paraffin-embedded sections were deparaffinized by xylene and dehydrated by ethanol. After washing with PBS, liver sections were treated with 3 % hydrogen peroxide for 10 min and with 10 % normal goat serum for another 10 min. They were sequentially incubated with anti GST-alpha (×100) or GST-π (×1000) antibody overnight. Avidin-biotin peroxidase complex method (ABC) was used for the staining. Positive immunoreaction was detected with 3-amino-9-ethylcarbazol (Nichirei Co., Ltd., Japan). Sections were counterstained with Mayer's hematoxylin for microscopic examination. They were also stained with H.E. Preimmune rabbit serum was used as a negative antibody control. Sections were observed by light microscope, and the immunohistochemical staining was judged to be the percentage of GST positive hepatocytes per total hepatocytes as follows: 90 % < (3 +), 75 % - 90 % (2 +), 50 % - 75 % (+), 25 % - 50 % (±) and 25 % < (−).

Statistical analysis. Significance of the differences between the two groups was evaluated using χ²-test, and the mean values were compared with Student's t-test.

Results

Histological findings and ICG uptake of the nodules. Nine rats died before they could be killed. Twenty-six ICG-stained nodules (ICG(+)) and 14 ICG-unstained nodules (ICG(−)) were obtained. Histologically, 11 ICG (+) nodules were benign (42.3 %) and 15 were borderline lesions (57.7 %). ICG (−) nodules included 2 HCCs (14.3 %) and 12 borderline lesions (85.7 %) (Table 2).

Specificities of anti GST-alpha and -π antibodies. The results of Western blot analysis for anti GST-alpha and -π antibodies are shown in Fig. 1. Anti GST-alpha antibody strongly reacted with 27 kD protein, and anti GST-π antibody reacted with 29 kD protein, indicating
Table 2  ICG staining and histological finding in liver nodules examined

<table>
<thead>
<tr>
<th></th>
<th>Number of nodule lesions histologically found</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Benign</td>
</tr>
<tr>
<td>ICG (+)</td>
<td>11</td>
</tr>
<tr>
<td>ICG (-)</td>
<td>0</td>
</tr>
</tbody>
</table>

Table 3  Immunohistochemical staining pattern of GST-α and histological findings

<table>
<thead>
<tr>
<th>GST-α staining</th>
<th>Number of nodule lesions histologically found</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Benign (%)</td>
</tr>
<tr>
<td>−</td>
<td>4 (14.8)</td>
</tr>
<tr>
<td>±</td>
<td>2 (18.2)</td>
</tr>
<tr>
<td>+</td>
<td>4 (36.3)</td>
</tr>
<tr>
<td>2+</td>
<td>2 (18.2)</td>
</tr>
<tr>
<td>3+</td>
<td>3 (27.3)</td>
</tr>
</tbody>
</table>

(−) < 25%, (±) 25%−50%, (+) 50%−75%, (2+) 75%−90%, (3+) > 90%

Table 4  Relationships between GST-α staining, diameters of nodules and time from 2-AAF administration

<table>
<thead>
<tr>
<th>GST-α staining</th>
<th>Diameter of nodules (mm)</th>
<th>Time from 2-AAF administration (week)</th>
</tr>
</thead>
<tbody>
<tr>
<td>− (n=6)</td>
<td>9±3 (3−12)</td>
<td>20±5 (13−26)</td>
</tr>
<tr>
<td>± (n=7)</td>
<td>8±5 (2−15)</td>
<td>17±8 (7−26)</td>
</tr>
<tr>
<td>+ (n=10)</td>
<td>7±2 (5−11)</td>
<td>20±6 (12−27)</td>
</tr>
<tr>
<td>2+ (n=12)</td>
<td>11±5 (5−20)</td>
<td>21±5 (9−26)</td>
</tr>
<tr>
<td>3+ (n=5)</td>
<td>7±4 (2−14)</td>
<td>20±6 (10−26)</td>
</tr>
</tbody>
</table>

(−), (+), (±), (2+), (3+): See Table 3.

that each antibody was specific for GST subunits.

*Immunohistochemical staining for GST-alpha and -pi.* Early hyperplastic nodules that developed until the 10th week, 2-AAF administration were strongly positive for GST-alpha in their cytoplasm, although the surrounding tissues were negative, some showing severe fibrosis with necrosis and the destruction of the trabecular arrangement. However, the percentage of GST-alpha positive cells per total number of hepatocytes tended to decrease with the disappearance of ICG staining after 10th week of 2-AAF administration.

As shown in Table 3, the percentage of GST-alpha positive nodules (+−3+) were 81.8% (9/11) in benign tissues, 66.7% (18/27) in borderline lesions and 0% (0/2) in HCCs, respectively. The relationship between GST-alpha staining and ICG uptake was examined.

GST-alpha positive nodules consisted of 81.8% (9/11) of ICG (+) benign tissues, 93.3% (14/15) of ICG (+) borderline lesions, 33.3% (4/12) of ICG (−) borderline lesions and 0% (0/2) of ICG (−) HCCs. GST (+) nodules showed significantly larger counts of GST-alpha positive hepatocytes than ICG (−) nodules (p < 0.01), indicating that ICG uptake was closely related to GST-alpha staining (Fig. 3). No significant relationships between GST-alpha staining and the sizes of nodules or the time from 2-AAF administration were recognized (Table 4). The characteristic sections of nodules positive in GST-alpha are shown in Fig. 4.

In the process of the carcinogenesis, early hyperplastic nodules generated at week 7 were strongly positive of GST-pi as well as borderline lesions. However, 2 of HCC nodules did not show GST-pi positive hepatocytes. GST-pi was not expressed in normal rat hepatocytes and positive staining was only recognized in cholangioepithelial cells.

As the results shown in Table 5, the percentage of
Fig. 3  The relationship between GST-alpha staining and indocyanine green (ICG) uptake of the liver nodules induced by 2-AAF: Most of stained nodules were strongly stained by GST-alpha, though most of ICG unstained nodules were not stained. It might be possible that ICG stained nodules tend to be stained with GST-alpha, too. GST: See Fig. 1.

Fig. 4  Microscopic findings of liver nodules immunohistochemically stained by GST-alpha: A: Normal rat liver (7 weeks). The cytoplasm were abundant in GST-alpha. B: ICG stained benign hyperplastic nodule (10 weeks). The cytoplasm were strongly stained by GST-alpha as well as A. C: ICG unstained borderline hyperplastic nodule (26 weeks). GST-alpha was not stained in a lot of hepatocytes with fatty infiltration. D: ICG unstained hepatocellular carcinoma (HCC) (26 weeks). No hepatocytes were stained by GST-alpha. GST, ICG: See Figs 1 and 3.

GST-pi positive nodules (+ ~ 3+) were 81.8% (9/11) in benign tissues, 77.8% (21/27) in borderline lesions and 0% (0/2) of HCCs, respectively.

Table 5  Immunohistochemical staining pattern of GST-pi and histological findings

<table>
<thead>
<tr>
<th>GST-pi staining</th>
<th>Number of nodule lesions histologically found</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Benign (%)</td>
</tr>
<tr>
<td>−</td>
<td>2 (7.4)</td>
</tr>
<tr>
<td>+</td>
<td>2 (18.2)</td>
</tr>
<tr>
<td>2+</td>
<td>1 (9.1)</td>
</tr>
<tr>
<td>3+</td>
<td>6 (54.5)</td>
</tr>
<tr>
<td></td>
<td>2 (18.2)</td>
</tr>
</tbody>
</table>

(−), (+), (2+), (3+): See Table 3.

Table 6  Relationships between GST-pi staining, diameters of nodules and time from 2-AAF administration

<table>
<thead>
<tr>
<th>GST-pi staining</th>
<th>Diameter of nodules (mm)</th>
<th>Time from 2-AAF administration (week)</th>
</tr>
</thead>
<tbody>
<tr>
<td>−(n=3)</td>
<td>5 ± 3 (2–8)</td>
<td>21 ± 4 (17–26)</td>
</tr>
<tr>
<td>+ (n=7)</td>
<td>9 ± 7 (2–12)</td>
<td>18 ± 7 (7–24)</td>
</tr>
<tr>
<td>2+(n=10)</td>
<td>9 ± 3 (6–18)</td>
<td>19 ± 5 (13–24)</td>
</tr>
<tr>
<td>3+(n=5)</td>
<td>6 ± 5 (2–14)</td>
<td>17 ± 6 (10–26)</td>
</tr>
</tbody>
</table>

(−), (+), (2+), (3+): See Table 3.

The relationship between GST-pi staining and ICG uptake. GST-pi positive nodules consisted of 81.8% (9/11) of ICG (+) benign tissues, 80% (12/15) of ICG (+)
ICG Uptake and GST Staining in Liver Nodules

borderline lesions, 75% (9/12) of ICG (−) borderline lesions and 0% (0/2) of ICG (−) HCCs (Fig. 5). Hyperplastic nodules including benign tissues without histologically atypisms and borderline lesions were strongly stained by GST-pi, suggesting no relation between their ICG uptake and GST-pi staining. In addition, significant relationships between GST-pi staining and the sizes of nodules or the time from the starting of 2-AAF administration were not recognized (Table 6). The characteristic sections of the nodules stained by GST-pi are shown in Fig. 6.

Discussion

GST is a polyfunctional enzyme constituted from subunits, and involved in detoxication for various toxins. The subunits have specific roles in conjugation with glutathione, glutathione peroxidase, and participates in leukotriene and prostaglandin metabolism (14–16). Some GST-subunits act as a ligandins which bind with organic anions (e.g., bilirubin, ICG and BSP) and excrete them into bile (17). The strong expression of one of the subunits, GST-pi, in enzyme altered foci and hyperplastic nodules which are related to the preneoplastic lesions of HCCs in humans and rats has been reported (18–19). Therefore, it is supposed to be one of the available tumor markers. In our experiment, all of benign tissues were ICG (+), all of HCCs were ICG (−) and borderline lesions consisted of both ICG (+) and ICG (−). However, the fact that ICG (−) borderline lesions showed abnormal DNA ploidy patterns (11) suggest that a loss of the ability to take up ICG is one of phenotypic changes of malignant potency.

Shimada (20) and Aikawa (21) reported that GST activity and ICG staining of the liver surface observed at the level of peritoneoscopic examination correlated well with each other, and that GST proved to be the ICG binding protein in hepatocytes. However, the identification of subunits that will bind with ICG has not been sufficiently discussed.

GST-alpha was more strongly stained in ICG (+) nodules than in ICG (−) borderline lesions or HCCs, suggesting that GST-alpha acts as a ligand for ICG. And the decrease of GST-alpha, which might be due to the cellular transformation in the process of hepatocarcinogenesis, is an important reason for the low ICG staining of the preneoplastic nodules. On the contrary, the absence of significant correlations between GST-pi staining of ICG (+) and ICG (−) nodules excludes GST-pi as a ligand for ICG.

It is likely that BSP is taken up by hepatocytes through BSP receptors on the liver plasma membrane (2), however, the ICG receptor has not yet been identified. In our experiment, 4 of the GST-alpha stained nodules were ICG (−). This suggests either a qualitative or quantitative dysfunction of ICG receptors maybe also play an important role in ICG staining of the nodules in the process of hepatocarcinogenesis.

Most of the hyperplastic nodules, histologically defined as benign or borderline, were GST-alpha positive in our experiment. We interpret this to mean that the decrease of GST-alpha staining is marked at the late stage of hepatocarcinogenesis. GST-pi was positive in most of the hyperplastic nodules, defined as benign or borderline, though the normal rat liver was GST-pi negative (Figs. 4 and 6). GST-alpha stained normal rat liver tissue, but GST-pi strongly stained only hyperplastic nodules, suggesting that the increase of GST-pi staining marks an earlier stage of hepatocarcinogenesis than the decrease of ICG and GST-alpha staining. Consequently, GST-pi appears to be one of the available markers that indicate preneoplastic hepatocytes.
Sato et al. have reported that GST-pi was abundant in preneoplastic lesions or well-differentiated HCCs in spite of its scarcity in poorly-differentiated HCCs. They also mentioned that GST-pi has already appeared at the stage of enzyme altered foci, which precedes hyperplastic nodules (18). Our experiment generated 2 types of HCC. The one weakly stained (±) for GST-pi was well-differentiated HCC with acinar arrangement, and the other unstained (−) for GST-pi was a poorly-differentiated HCC. Although further study including more subjects is necessary, our results tend to support the conclusion expressed by Sao et al. Further investigations into the question of whether all GST-pi positive hepatocytes become neoplastic or not are required.

In conclusion, our findings indicate that GST-alpha acts as a possible ligand for ICG in hepatocytes and the increase of GST-pi staining appears at the earlier stage of hepatocarcinogenesis than the decrease of GST-alpha or ICG staining. These in turn suggest that GST-pi may be useful for early diagnosis of preneoplastic hepatocytes and that it does not act as ligand for ICG.

Acknowledgment. This study was supported by grants from the Ministry of Health and Welfare of Japan (Ten years strategy for cancer treatment).

References


Received September 28, 1992; accepted June 14, 1993.