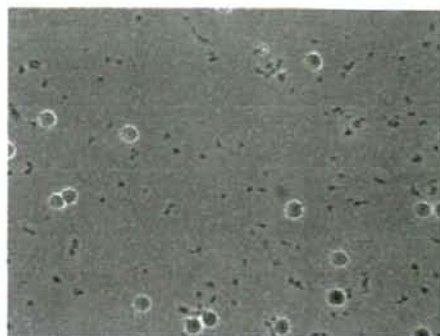


結果

分離されたPBMC

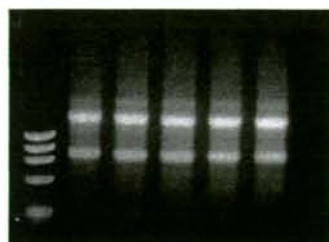


1×10^7 cells/dish \rightarrow 600 μ l で可溶化 \rightarrow total 3 μ g RNA

5×10^7 cells/dish \rightarrow 1200 μ l で可溶化 \rightarrow total 9 μ g RNA



マイクロアレイ実験に使用可能



小括

ラット血液よりPBMCを単離，培養し，遺伝子発現解析を行うことのできる実験系を準備した．血球系細胞におけるin vitroの系としてこの系は毒性メカニズム解析に有用と考えられる．

今後，各種化合物曝露による遺伝子発現解析を行い，in vivoの全血データと合わせバイオマーカー検証に用いていく予定である．

まとめ

本研究において、初年度はモデルケースとして肝臓のGSH枯渇について、メカニズムに基づくバイオマーカー候補遺伝子の抽出を行い、*in vitro*の系を立ち上げて検証を試みた。その結果、GSH枯渇時点がわからなくても*in vivo*, *in vitro*とも利用可能なマーカー候補として有望なものが得られた。今後さらに検証が必要であるが、このような特性を持つバイオマーカーはきわめて有用である。

また、PBMC初代培養系の構築を行い、血球系細胞においても*in vitro*での解析が可能となる体制を整えた。

今後これらの系を用い、阻害薬や遺伝子導入・ノックダウンなどにより、TGP2で蓄積される各種バイオマーカー候補遺伝子のメカニズム等の検証を行い、バイオマーカー確立に貢献していきたい。

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研究成果の刊行に関する一覧表

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A toxicogenomics approach for early assessment of potential non-genotoxic hepatocarcinogenicity of chemicals in rats

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ABSTRACT

For assessing carcinogenicity in animals, it is difficult and costly, an alternative strategy has been desired. We explored the possibility of applying a toxicogenomics approach by using comprehensive gene expression data in rat liver treated with various compounds. As prototypic non-genotoxic hepatocarcinogens, thioacetamide (TAA) and methapyrilene (MP) were selected and 349 commonly changed genes were extracted by statistical analysis. Taking both compounds as positive with six compounds, acetaminophen, aspirin, phenylbutazone, rifampicin, alpha-naphthylisothiocyanate, and amiodarone as negative, prediction analysis of microarray (PAM) was performed. By training and 10-fold cross validation, a classifier containing 112 probe sets that gave an overall success rate of 95% was obtained. The validity of the present discriminator was checked for 30 chemicals. The PAM score showed characteristic time-dependent increases by treatment with several non-genotoxic hepatocarcinogens, including TAA, MP, coumarin, ethionine and WY-14643, while almost all of the non-carcinogenic samples were correctly predicted. Measurement of hepatic glutathione content suggested that MP and TAA cause glutathione depletion followed by a protective increase, but the protective response is exhausted during repeated administration. Therefore, the presently obtained PAM classifier could predict potential non-genotoxic hepatocarcinogenesis within 24 h after single dose and the inevitable pseudo-positives could be eliminated by checking data of repeated administrations up to 28 days. Tests for carcinogenicity using rats takes at least 2 years, while the present work suggests the possibility of lowering the time to 28 days with high precision, at least for a category of non-genotoxic hepatocarcinogens causing oxidative stress.

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1. Introduction

Chemical carcinogenesis is a multistage process, i.e., initiation, promotion and progression (Dragan et al., 1993; Miller and Miller, 1981; Scott et al., 1984). Based on this mechanism of action, chemical carcinogens are classified as genotoxic (mutagenic) and non-genotoxic (non-mutagenic) agents (Hayashi, 1992; Melnick et al., 1996). Genotoxic agents covalently react with DNA to form DNA adducts within the cells of the target organ, contributing to the initiation process. Such chemicals could be assessed by several short-term *in vitro* and *in vivo* assays that measure DNA damage,

mutagenic effects, and chromosomal aberrations (Weisburger and Williams, 2000). In the case of non-genotoxic agents, the mechanism is much more complicated. Non-genotoxic carcinogens lack chemical reactivity with DNA and hence do not form DNA adducts, but rather induce effects that indirectly lead to neoplastic transformation or enhance the development of tumors from pre-initiated cells. Although the mechanism of action of such non-genotoxic carcinogens is not fully understood, several possibilities have been postulated in liver, such as oxidative stress, modulation of metabolizing enzymes, induction of peroxisome proliferation, alteration of intercellular communication, and disruption of the balance between proliferation and apoptosis (Butterworth and Bogdanffy, 1999; Cohen and Ellwein, 1990; Klaunig et al., 1998; Klaunig and Kamendulis, 2004; Nguyen-Ba and Vasseur, 1999; Silva Lima and Van der Laan, 2000; Williams et al., 1996). Even more complicated is the fact that many non-genotoxic carcinogens frequently cause several of these effects at once. The effects of non-genotoxic

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carcinogens in rodents are only manifested after *in vivo* exposure at high dosage levels over long periods (e.g. 2-year rodent carcinogenicity assays). Consequently, the current strategy for evaluating non-genotoxic carcinogens is not satisfactory because the test is time consuming and expensive, and it requires the use of many animals and large amounts of chemicals.

The present report is focused on the application of toxicogenomics for early assessment of potential non-genotoxic hepatocarcinogenicity of chemicals. Non-genotoxic hepatocarcinogenesis has been studied extensively, and postulated to act via a number of mechanisms: oxidative stress, increased mitogenesis, decreased apoptosis, interference with gap junction intercellular communication, and interference with tubulin polymerization (Combes, 2000; Klaunig et al., 1998). Several recent publications have described applications of microarrays and expression profiling for non-genotoxic carcinogenesis in liver (Ellinger-Ziegelbauer et al., 2005, 2008; Fielden et al., 2007; Nie et al., 2006). They attempted to extract common gene sets coordinately deregulated by several different classes of genotoxic and/or non-genotoxic hepatocarcinogenesis. It was then revealed that the modulation of extracted genes was dependent upon the class of the carcinogenesis. This strongly suggests that mechanism-based strategy should be employed in order to obtain useful biomarker gene sets for carcinogenesis. The specific aim of the present study was to develop identifiers for early assessment of non-genotoxic hepatocarcinogenicity in specific class of chemical based on gene expression profiles in reference to our large-scale database named as TG-GATES (genomics assisted toxicity evaluation system developed by Toxicogenomics Project, Japan) (Urushidani, 2007). Our strategy was to focus on common gene expression changes in livers treated with two well-known oxidative stressors, methapyri- lene (MP) (Lijinsky et al., 1980; National Toxicology Program, 2000; Ohshima et al., 1984; Ratra et al., 1998) and thioacetamide (TAA) (Becker, 1983; Diez-Fernandez et al., 1998; Duivenvoorden and Maier, 1994; Ohtsuka et al., 1998; Sanz et al., 1995) to identify a characteristic set of genes reflecting the early stage of oxidative stress-mediated non-genotoxic hepatocarcinogenesis.

2. Materials and methods

2.1. Animals and experimental design

Five-week-old male Sprague-Dawley rats were obtained from Charles River Japan, Inc. (Kanagawa, Japan). After a 7-day quarantine and acclimatization period, the animals (6-week old) were assigned to dosage groups (five rats per group) using a computerized stratified random grouping method based on individual body weight. The animals were individually housed in stainless-steel cages in an animal room that was lighted for 12 h (7:00–19:00) daily, ventilated with an air-exchange rate of 15 times per hour, and maintained at 21–25 °C with a relative humidity of 40–70%. Each animal was allowed free access to water and pellet diet (CRF-1, sterilized by radiation, Oriental Yeast Co. Ltd., Tokyo, Japan).

Table 1 lists the overview of the compounds used in this study. A total of 30 compounds (10 non-genotoxic hepatocarcinogens and 20 non-hepatocarcinogens) were available in the database when the present analysis was performed. They were subdivided in a training set, consisting of 2 non-genotoxic carcinogens (positive training set) and 6 non-hepatocarcinogens (negative training set) with the test set for additional validation consisting of 8 non-genotoxic carcinogens and 14 non-hepatocarcinogens.

According to the standard protocol in our project (Takashima et al., 2006), five rats per group were orally administered at three doses with these compounds suspended or dissolved either in 0.5% methylcellulose (MC) solution or corn oil according to their dispersibility. Traditionally, carcinogenicity studies for chemical agents have relied upon the maximally tolerated dose (MTD) as the standard method for high dose selection. In the present study, the MTD was chosen based on data derived from preliminary toxicity studies of 7 days duration.

For single-dose studies, rats were sacrificed at 3, 6, 9 and 24 h after dosing (3H, 6H, 9H and 24H, respectively). For repeated dose studies, the animals were treated daily for 3, 7, 14 and 28 days, and sacrificed 24 h after the last dosing [day 4 (4D), 8 (8D), 15 (15D) and 29 (29D), respectively]. The animals were euthanized by exsanguination from the abdominal aorta under ether anesthesia, and the liver samples

were obtained from the left lateral lobe of the liver in each animal immediately after sacrifice for examination.

The experimental protocols were reviewed and approved by the Ethics Review Committee for Animal Experimentation of National Institute of Health Sciences.

2.2. Histopathology of livers treated with MP or TAA

For light microscopic examination, the liver sample of each animal was fixed in 10% neutral buffered formalin, dehydrated in alcohol and embedded in paraffin. Paraffin sections were prepared and stained by a routine method with hematoxylin and eosin (H&E).

2.3. Microarray analysis

An aliquot of the sample (about 30 mg) for microarray analysis was obtained from the left lateral lobe of the liver in each animal immediately after sacrifice, kept in RNAlater® (Ambion, Austin, TX, USA) overnight at 4 °C, and then frozen at –80 °C until use. Liver samples were homogenized with the buffer RLT supplied in RNeasy Mini Kit (Qiagen, Valencia, CA, USA), and total RNA was isolated according to the manufacturer's instructions. Microarray analysis was conducted on three out of five samples for each group by using GeneChip® RAE230A probe arrays (Affymetrix, Santa Clara, CA, USA). The procedure was basically conducted according to the manufacturer's instructions as previously reported (Uehara et al., 2008a,b). Microarray Analysis Suite 5.0 (MAS; Affymetrix) was used to quantify microarray signals and the intensities were normalized for each chip by setting the mean intensity to 500 (per chip normalization).

2.4. Selection of persistently up/down-regulated genes in common with MP and TAA

By using statistical and clustering tools, persistently up/down-regulated genes in common with MP and TAA throughout the study periods were extracted. First, data were imported into GeneSpring 6.0 software (Silicon Genetics, Redwood City, CA), and comparisons among time-matched groups from each study of MP and TAA were performed using one-way analysis of variance (ANOVA) with Tukey's multiple comparison test for post hoc comparisons when significance was determined by ANOVA with a false discovery rate ($p < 0.05$). Probe sets exhibiting significant changes in expression by Tukey's multiple comparison test in both high- and middle-dose groups for one or more time points in each study were selected. In the next step, significant selected probe sets (452 probe sets) were divided into subsets with distinct expression profiles by *k*-means clustering using Tigr Mev 3.1 software (<http://www.tm4.org/mev.html>) (current metric: Euclidean distance; divided into nine clusters) based on logarithm (\log_2) of the ratio to control for individual gene expression. Genes not categorized in the clusters showing clear time- and dose-independent expression pattern were excluded from further analysis. Finally, a subset of 349 probe sets containing 276 up-regulated and 73 down-regulated probe sets was selected for common intersection to single and repeated studies of MP and TAA (for more information, see supplemental figures).

2.5. Class discrimination by prediction analysis of microarray (PAM)

Prediction of potential carcinogenesis was performed by an approach using PAM for R package (<http://www-stat.stanford.edu/~tibs/PAM>). PAM makes sample classification using the nearest shrunken centroid method with an automated gene selection step integrated into the algorithm (Tibshirani et al., 2002). It employs a parameter threshold Δ to select genes for class discrimination. PAM training is performed by comparing 2 positive compounds as non-genotoxic carcinogenesis (MP and TAA, high dose group only) with 6 negative compounds, i.e., APAP (Iida et al., 2005; National Toxicology Program, 1993), ASA (Giri, 1993), PhB (Meakawa et al., 1987; National Toxicology Program, 1990), RIF (Sodhi et al., 1997), ANIT (Jean and Roth, 1995; Leonard et al., 1981) and AM (Agoston et al., 2003; Delaney et al., 2004) for the ratio of expression levels of the selected 349 probe sets at various time points (a total of 64 training samples).

Ten-fold cross validation was performed to find out the optimal classifier performance, which minimized classification errors for training sets. During the validation, a threshold Δ was varied in search of the optimal classifier performance. The Δ value that settled at the lowest classification error with the fewest genes was favored as the optimal. For validation of the classifier, the optimized threshold value obtained from training was subsequently used for prediction of potential carcinogenicity for the total of 30 compounds, including training sets. PAM prediction results were expressed as a logarithm transformed score (PAM prediction score) of the ratio of positive class probability relative to negative class probability associated with the classification of each sample, i.e.,

$$\text{PAM prediction score} = \log_{10} \frac{\text{class probability : positive}}{\text{class probability : negative}}$$

Table 1
Overview of the compounds used for prediction analysis of microarrays training and/or test

| Compound | Abbreviation | CAS-number | Mode of action | Supplier | Vehicle | Dose (mg/kg) | PAM training/test |
|--|------------------|------------|----------------------------|-------------------------------|----------|--------------|----------------------------|
| Non-genotoxic hepatocarcinogens^{a,b} | | | | | | | |
| Methapyrilene | MP | 135-23-9 | Oxidative stress induction | Sigma | 0.5%MC | 10, 30, 100 | Positive training/test set |
| Thioacetamide | TAA | 62-55-5 | Oxidative stress induction | Sigma | 0.5%MC | 4.5, 15, 45 | Positive training/test set |
| Coumarin | CMA | 91-64-5 | Oxidative stress induction | Tokyo Chemical Industry | Corn oil | 150 | Test set |
| Ethionine | ET | 67-21-0 | Oxidative stress induction | Tokyo Chemical Industry | 0.5%MC | 250 | Test set |
| Carbon tetrachloride | CCl ₄ | 56-23-5 | Oxidative stress induction | Wako Pure Chemical Industries | Corn oil | 300 | Test set |
| Phenobarbital | PB | 57-30-7 | Hepatic enzyme induction | Sigma | 0.5%MC | 100 | Test set |
| Hexachlorobenzene | HCB | 118-74-1 | Hepatic enzyme induction | Tokyo Chemical Industry | Corn oil | 300 | Test set |
| Clofibrate | CFB | 637-07-0 | Peroxisome proliferation | Wako Pure Chemical Industries | Corn oil | 300 | Test set |
| Gemfibrozil | GFZ | 25812-30-0 | Peroxisome proliferation | Sigma | Corn oil | 300 | Test set |
| Wy-14,643 | WY | 50892-23-4 | Peroxisome proliferation | Tokyo Chemical Industry | Corn oil | 100 | Test set |
| Non-hepatocarcinogens^{a,b} | | | | | | | |
| Acetaminophen | APAP | 103-90-2 | - | Sigma | 0.5%MC | 600 | Negative training set |
| Aspirin | ASA | 50-78-2 | - | Wako Pure Chemical Industries | 0.5%MC | 450 | Negative training set |
| Phenylbutazone | PhB | 50-33-9 | - | Sigma | 0.5%MC | 200 | Negative training set |
| Rifampicin | RIF | 13292-46-1 | - | Wako Pure Chemical Industries | 0.5%MC | 200 | Negative training set |
| Alpha-naphthylisothiocyanate | ANIT | 551-06-4 | - | Tokyo Chemical Industry | Corn oil | 15 | Negative training set |
| Amiodarone hydrochloride | AM | 1951-25-3 | - | Sigma | 0.5%MC | 200 | Negative training set |
| Allopurinol | APL | 315-30-0 | - | Sigma | 0.5%MC | 150 | Negative test set |
| Allyl alcohol | AA | 107-18-6 | - | Tokyo Chemical Industry | Corn oil | 30 | Negative test set |
| Benzbromarone | BBr | 3562-84-3 | - | Sigma | 0.5%MC | 200 | Negative test set |
| Bromobenzene | BBZ | 108-86-1 | - | Tokyo Chemical Industry | Corn oil | 300 | Negative test set |
| Carbamazepine | CBZ | 298-46-4 | - | Sigma | 0.5%MC | 300 | Negative test set |
| Chlorpromazine | CPZ | 69-09-0 | - | Wako Pure Chemical Industries | 0.5%MC | 45 | Negative test set |
| Diclofenac sodium | DFNa | 15307-79-6 | - | Cayman Chemical Company | 0.5%MC | 10 | Negative test set |
| Diazepam | DZP | 439-14-5 | - | Wako Pure Chemical Industries | 0.5%MC | 250 | Negative test set |
| Isoniazid | INAH | 54-85-3 | - | Sigma | 0.5%MC | 200 | Negative test set |
| Nitrofurantoin | NFT | 67-20-9 | - | ICN Biomedicals | 0.5%MC | 100 | Negative test set |
| Phenytoin | PHE | 57-41-0 | - | Tokyo Chemical Industry | 0.5%MC | 600 | Negative test set |
| Propylthiouracil | PTU | 51-52-5 | - | Tokyo Chemical Industry | 0.5%MC | 100 | Negative test set |
| Sulfasalazine | SS | 599-79-1 | - | Sigma | 0.5%MC | 1000 | Negative test set |
| Valproate sodium | VPA | 1069-66-5 | - | Sigma | 0.5%MC | 450 | Negative test set |

^a Genotoxicity is based on *in vitro* genotoxicity tests (Salmonella and mammalian gene mutation tests) as reviewed in NTP (<http://ntp-server.niehs.nih.gov/>), IARC (<http://monographs.iarc.fr/>) and several published papers.

^b Carcinogenicity is based on reviews by NTP (<http://ntp-server.niehs.nih.gov/>), IARC (<http://monographs.iarc.fr/>) and several published papers.

2.6. Gene ontology (GO) analysis of PAM classifier

The identified probe sets were subjected to GO analysis by DAVID (database for annotation, visualization, and integrated discovery; <http://apps1.lniaid.nih.gov/david/>) using Fisher's exact test. Level 3 analysis was adopted.

2.7. Measurement for hepatic total glutathione contents

Hepatic total glutathione was measured in the liver of rats receiving a high dose of MP, TAA or BBZ, and their corresponding controls. Measurements were performed for three rats (gene expression was measured) per group using Glutathione Quantification Kit (Dojindo Mol. Tech, Inc., Kumamoto, Japan). In brief, the liver tissue was homogenized in 5% 5-sulfosalicylic acid and the particulate cellular debris was removed by centrifugation (8000 × g) for 10 min. The internal standards consist of serial dilutions of glutathione (1000, 750, 500, 250, 100, 50 and 0 μM). The change in absorbance at 405 nm was measured and total glutathione was calculated according to the glutathione standard curve. The results were analyzed with the use of an unpaired two-tailed Student's *t*-test or Welch's *t*-test as appropriate, and a *p*-value of <0.05 was considered statistically significant.

3. Results

3.1. Histopathology

Except for the death of one animal in the high dose group of MP on 20D, there were no other deaths in these studies of MP and TAA.

MP- or TAA-treated rats revealed typical liver damage throughout the study periods. Although the extent of the liver damage differed slightly among the animals, a similar pattern was obtained for those in the same dose group.

At high dosage of MP in the single dose study, periportal hepatocytes exhibited hypertrophy characterized by granular eosinophilic cytoplasm and enlarged nuclei with variable anisonucleosis at each time point. More striking abnormalities include mononuclear cell infiltration and hepatocellular single cell necrosis containing shrunken cells with pyknotic nuclei randomly scattered throughout the periportal region of the hepatic lobule. Associated with these lesions, increased numbers of hepatocellular mitotic figures and bile duct hyperplasia were present at each time point in the repeated dose study. At 29D, hepatocellular hyperplasia became evident, and some affected portal regions contained an increased number of oval cells arranged in clusters without a distinct lumen (Fig. 1a). In addition, for the same dose group, a pre-neoplastic altered hepatocellular focus was also observed (Fig. 1b). Middle-dose MP treatment resulted in minimal hepatocellular hypertrophy, single cell necrosis of hepatocytes, and mononuclear cell infiltration in the periportal region at 15D and 29D. Moreover, no significant histopathological alterations were observed at early time points except hepatocellular hypertrophy. In the low-dose MP-treated groups, no significant changes were observed throughout the study periods except for minimal hypertrophy of hepatocytes, observed in one animal each at 8D and 29D.

At high- and middle-dose of TAA, centrilobular hepatocytes exhibited hypertrophy with large, atypical nuclei in single and repeated dose studies (Fig. 1c). Moreover inflammatory cell infiltration and hepatocellular single cell necrosis were also observed at the centrilobular region. The degree of these lesions increased in a dose and time-dependent manner. At 15D and 29D, bile duct hyperplasia and oval cell proliferation at the periportal region became evident, and a pre-neoplastic altered hepatocellular focus was also observed (Fig. 1d). No significant histopathological alterations were observed in the low-dose groups throughout the study periods except degeneration of hepatocytes with granular and eosinophilic cytoplasm, observed in two animals at 29D.

3.2. Class discrimination by PAM in the training set

PAM training was performed using the training set to identify a minimal subset of genes expected to best characterize the early stage of non-genotoxic hepatocarcinogenesis-specific responses. Fig. 2 shows the training and cross-validation errors for different threshold values. Both the training and cross-validated errors were minimized near the threshold = 4.00, where 112 genes were selected. At this threshold, both classes of the training samples were clearly separated based on the expression pattern of these 112 genes with an overall success rate of 95%. Namely, 13 of the 16 positive sets (81%) and all of the negative sets (100%) were correctly classified (Fig. 3a). However, three positive sets (MP-3H, -4D and TAA-3H) were classified as negative, together with all of the negative sets (Fig. 3b).

The list of the genes involved in the PAM classifier is shown in Table 2 (for more information, see supplemental data). Genes were sorted according to the best prediction between the two classes. The top three important discriminators identified by PAM were "nuclear RNA helicase, DECD variant of DEAD box family (Ddx39)", "interferon-related developmental regulator 1 (Ifrd1)", and "mdm2, transformed mouse 3T3 cell double minute 2 (Mdm2)", which were highly up-regulated by MP and TAA. In the extracted 112 probe sets, 111 were prominently up-regulated in the positive training set and the remaining 1 gene (cytochrome P450 4F4) was down-regulated. Based on gene ontology, the contents of genes related to cellular metabolism including several anti-oxidative metabolism, cell proliferation, cell cycle, response to DNA damage stimulus were significantly high (Table 3). These features might reflect the cellular changes related to sustained oxidative stress in association with non-genotoxic hepatocarcinogenesis by MP and TAA.

3.3. Validation of usefulness of the PAM classifier

The 112-gene classifier generated on the training set was next applied to class discrimination for the 30 total compounds as a validation test. The classifier predicted the following samples as positive: high dose MP-6H, 9H, 24H, 8D, 15D and 29D; middle-dose TAA-29D; high dose TAA-6H, 9H, 24H, 4D, 8D, 15D and 29D; CMA-3H, 6H and 9H; ET-24H, 4D, 8D, 15D and 29D; WY-15D and 29D; BBZ-24H. All of other samples (including enzyme inducers, PB and HCB; peroxisome proliferators other than WY, such as CFB and GFZ; and other compounds) were predicted as negative.

In the present study, these prediction results were visualized as a numerical score reflecting the probabilities of class discrimination between the two classes, namely the PAM prediction score. The PAM score showed characteristic time-dependent changes by treatment with several non-genotoxic hepatocarcinogens. In the MP- or TAA-treated group, the score increased dose-dependently with a peak value at 6H for MP, 9H and 24H for TAA after single dosing, and then it markedly increased with repeated administrations (Fig. 4b, c, e, f). CMA, ET or WY treatment also resulted in an increase in the score with a peak value at 6H for CMA, 24H for ET and WY, and also showed an increase or tendency to increase with repeated dosing (Fig. 4g, h, j). Although all of the CCL4-treated groups were predicted as negative, the score showed a tendency to increase with repeated dosing (Fig. 4i). On the other hand, all of the low dose MP- or TAA-treated groups were predicted as negative without any tendency to increase in the score with repeated dosing (Fig. 4a and d). As for the enzyme inducers with carcinogenic activity, PB and HCB (Fig. 4l and m), and peroxisome proliferators other than WY, i.e., CFB (Fig. 4n) and GFZ (within Fig. 4r), showed negative scores throughout the time points. Of the non-carcinogenic samples, BBZ showed a transient increase in the score at 24H but returned to negative during repeated dosing (Fig. 4k). Other non-carcinogenic

Table 2
The list of the genes involved in the PAM classifier

| Probe ID | Accession number | Gene title | Gene symbol |
|--------------|------------------|--|----------------------|
| 1387048.at | NM.053563 | DEAD (Asp-Glu-Ala-Asp) box polypeptide 39 | Ddx39 |
| 1367795.at | NM.019242 | Interferon-related developmental regulator 1 | ifrd1 |
| 1384427.at | XM.001080981 | Transformed mouse 3T3 cell double minute 2 homolog (mouse) (predicted) | Mdm2_predicted |
| 1388986.at | - | EST | - |
| 1369921.at | NM.020540 | Glutathione S-transferase M4 | Gstm4 |
| 1368072.at | NM.019290 | B-cell translocation gene 3 | Btg3 |
| 1387060.at | NM.031642 | Kruppel-like factor 6 | Klf6 |
| 1376098.a.at | XM.001069724 | Myosin IG | Myo1g |
| 1368173.at | NM.021754 | Nucleolar protein 5 | Nol5 |
| 1373200.at | XM.001063564 | Eukaryotic translation elongation factor 1 epsilon 1 (predicted) | Eef1e1_predicted |
| 1388560.at | NM.001008771 | WD repeat domain 77 | Wdr77 |
| 1374945.at | NM.001007706 | GCD14/PCMT domain containing protein RGD1359191 | RGD1359191 |
| 1376737.at | XM.001073157 | EST | LOC686259 |
| 1388397.at | NM.001008721 | EBNA1 binding protein 2 | Ebna1bp2 |
| 1371785.at | NM.181086 | Tumor necrosis factor receptor superfamily, member 12a | Tnfrsf12a |
| 1375895.at | - | EST | - |
| 1367764.at | NM.012923 | Cyclin G1 | Ccng1 |
| 1388674.at | NM.080782 | Cyclin-dependent kinase inhibitor 1A | Cdkn1a |
| 1373499.at | NR.002704 | Growth arrest specific 5 | Gas5 |
| 1386897.at | NM.024363 | Heterogeneous nuclear ribonucleoproteins methyltransferase-like 2 (<i>S. cerevisiae</i>) | Hrmt112 |
| 1372211.at | NM.145673 | v-maf musculoaponeurotic fibrosarcoma oncogene family, protein K (avian) | Mafk |
| 1386995.at | NM.017259 | B-cell translocation gene 2, anti-proliferative | Btg2 |
| 1372510.at | NM.001047858 | Sulfiredoxin 1 homolog (<i>S. cerevisiae</i>) | Srxn1 |
| 1388900.at | XM.001076548 | RGD1566118 (predicted) | RGD1566118_predicted |
| 1370583.s.at | NM.012623 | ATP-binding cassette, sub-family B (MDR/TAP), member 1A/1B | Abcb1a/Abcb1b |
| 1398756.at | NM.012992 | Nucleophosmin 1 | Npm1 |
| 1375224.at | NM.001012206 | Plectstrin homology-like domain, family A, member 3 | Phlda3 |
| 1388155.at | NM.053976 | Keratin complex 1, acidic, gene 18 | Krt1-18 |
| 1368032.at | NM.022869 | Nucleolar and coiled-body phosphoprotein 1 | Nolc1 |
| 1388629.at | NM.199099 | Inosine 5-monophosphate dehydrogenase 2 | Impdh2 |
| 1371936.at | NM.199372 | Eukaryotic translation initiation factor 4A1 | Eif4a1 |
| 1377387.a.at | - | EST | - |
| 1374326.at | NM.001011980 | Peter pan homolog (<i>Drosophila</i>) | Ppan |
| 1367617.at | NM.012495 | Aldolase A | Aldoa |
| 1376001.at | XM.001065234 | Polymerase (RNA) I associated factor 1 (predicted) | Praf1_predicted |
| 1398832.at | NM.012749 | Nucleolin | Ncl |
| 1368121.at | NM.013215 | Aldo-keto reductase family 7, member A3 (aflatoxin aldehyde reductase) | Akr7a3 |
| 1370174.at | NM.133546 | Myeloid differentiation primary response gene 116 | Myd116 |
| 1398771.at | NM.019283 | Solute carrier family 3, member 2 | Slc3a2 |
| 1389450.at | XM.001071583 | EST | LOC360830 |
| 1371530.at | NM.199370 | Keratin complex 2, basic, gene 8 | Krt2-8 |
| 1367834.at | NM.053464 | Spermidine synthase | Srm |
| 1387282.at | NM.053612 | Heat shock 22 kDa protein 8 | Hspb8 |
| 1372043.at | XM.001071573 | EST | RGD1311709_predicted |
| 1372150.at | NM.001034146 | Ubiquitin-specific protease 10 | Usp10 |
| 1389569.at | NM.001029915 | Brix domain containing 2 | Bxdc2 |
| 1371498.at | NM.001037348 | JTV1 | MGC125271 |
| 1389815.at | NM.172045 | Protein phosphatase 1, regulatory (inhibitor) subunit 14B | Ppp1r14b |
| 1370314.at | NM.031148 | Solute carrier family 20, member 1 | Slc20a1 |
| 1372218.at | NM.199410 | WD repeat domain 12 | Wdr12 |
| 1372354.at | - | EST | - |
| 1367654.at | NM.031819 | Fat tumor suppressor homolog (<i>Drosophila</i>) | Fath |
| 1388107.at | NM.144746 | Protein phosphatase 2, regulatory subunit B, delta isoform | Ppp2r2d |
| 1372028.at | NM.001047095 | EST | RGD1305727_predicted |
| 1373767.at | NM.001008363 | Zinc finger, AN1-type domain 2A | Zfand2a |
| 1390579.at | XM.001073162 | EST | RGD1305222_predicted |
| 1388588.at | NM.001015013 | Mammary tumor virus receptor 2 | Mtvr2 |
| 1370309.a.at | NM.031330 | Heterogeneous nuclear ribonucleoprotein A/B | Hnrpab |
| 1367732.at | NM.030987 | Guanine nucleotide binding protein, beta 1 | Gnb1 |
| 1399158.a.at | NM.012992 | Nucleophosmin 1 | Npm1 |
| 1389577.at | NM.001009640 | Cirrhosis, autosomal recessive 1A (human) | Cirh1a |
| 1398757.at | NM.012992 | Nucleophosmin 1 | Npm1 |
| 1370947.at | XM.001070821 | EST | Rda279 |
| 1373677.at | XM.001061829 | Solute carrier family 39 (zinc transporter), member 10 (predicted) | Slc39a10_predicted |
| 1388244.s.at | NM.017138 | Ribosomal protein SA | Rpsa |
| 1388150.at | NM.053490 | Exportin 1, CRM1 homolog (yeast) | Xpo1 |
| 1388666.at | NM.001003401 | Ectodermal-neural cortex 1 | Enc1 |
| 1367713.at | NM.019356 | Eukaryotic translation initiation factor 2, subunit 1 alpha | Eif2s1 |
| 1386910.a.at | NM.024148 | Apurinic/apyrimidinic endonuclease 1 | Apex1 |
| 1372019.at | XM.001062474 | EST | RGD1310128_predicted |
| 1373647.at | NM.001009652 | Zinc finger protein 622 | Zfp622 |
| 1387072.at | NM.053794 | Protein kinase, lysine deficient 1 | Prkwnk1 |
| 1388754.at | - | EST | - |
| 1367870.at | NM.032614 | Thioredoxin-like 2 | Txn12 |

Table 2 (Continued)

| Probe ID | Accession number | Gene title | Gene symbol |
|--------------|------------------|--|----------------------|
| 1387950.at | NM.138847 | Nuclear import 7 homolog (<i>S. cerevisiae</i>) | Nip7 |
| 1387807.at | NM.031763 | Platelet-activating factor acetylhydrolase, isoform lb, alpha subunit 45 kDa | Pafah1b1 |
| 1371378.at | XM.001053247 | EST | LOC678808 |
| 1371735.at | – | EST | – |
| 1398791.at | NM.031614 | Thioredoxin reductase 1 | Txnrd1 |
| 1386958.at | NM.031614 | Thioredoxin reductase 1 | Txnrd1 |
| 1385616.a.at | XM.001059946 | ASF1 anti-silencing function 1 homolog A (<i>S. cerevisiae</i>) (predicted) | Asf1a_predicted |
| 1388990.at | NM.139186 | Mki67 (FHA domain) interacting nucleolar phosphoprotein | Mki67ip |
| 1388449.at | XM.001071102 | Eukaryotic translation elongation factor 1 beta 2 (predicted) | Eef1b2_predicted |
| 1373850.at | NM.001025737 | Sphingomyelin phosphodiesterase, acid-like 3B | Smpd3b |
| 1371539.at | XM.001071992 | Nucleolar protein family A, member 2 (predicted) | Nola2_predicted |
| 1387774.at | NM.013011 | Tyrosine 3-monooxygenase/tryptophan 5-monooxygenase activation protein, zeta polypeptide | Ywhaz |
| 1371980.at | NM.001034922 | ATPase family, AAA domain containing 3A | Atad3a |
| 1373075.at | XM.001061556 | EST | RGD1560888_predicted |
| 1367693.at | NM.013052 | Tyrosine 3-monooxygenase/tryptophan 5-monooxygenase activation protein, eta polypeptide | Ywhah |
| 1387973.at | NM.173123 | Cytochrome P450, family 4, subfamily f, polypeptide 4 | Cyp4f4 |
| 1390317.at | – | EST | – |
| 1371377.at | NM.001037346 | Ribosomal protein S19 | Rps19 |
| 1373380.at | NM.001010963 | Brain zinc finger protein | LOC362154 |
| 1367590.at | NM.053439 | RAN, member RAS oncogene family | Ran |
| 1370295.at | NM.138548 | Expressed in non-metastatic cells 1 | Nme1 |
| 1374632.at | NM.001012143 | Phosphatidylserine receptor | Ptdsr |
| 1388381.at | NM.001013095 | Eukaryotic translation initiation factor 3, subunit 4 (delta) | Eif3s4 |
| 1370785.s.at | NM.152935 | Translocase of outer mitochondrial membrane 20 homolog (yeast) | Tomm20 |
| 1398801.at | NM.134415 | CDK105 protein | Cdk105 |
| 1374764.at | XM.001058941 | EST | RGD1305605_predicted |
| 1374793.at | XM.001065786 | WD repeat domain 3 (predicted) | Wdr3_predicted |
| 1368106.at | NM.031821 | polo-like kinase 2 (<i>Drosophila</i>) | Plk2 |
| 1372116.at | XM.001079091 | Mitochondrial ribosomal protein S2 (predicted) | Mrps2_predicted |
| 1388507.at | NM.001037352 | Integrin beta 4 binding protein | Itgb4bp |
| 1389200.at | NM.182674 | Bystin-like | Bysl |
| 1372558.at | XM.001053949 | NMDA receptor-regulated gene 1 (predicted) | Narg1_predicted |
| 1371809.at | NM.212534 | Mitochondrial ribosomal protein S18B | Mrps18b |
| 1387911.at | NM.138708 | RAB geranylgeranyl transferase, b subunit | Rabggtb |
| 1372243.at | XM.001063411 | Calcium binding protein 39 (predicted) | Cab39_predicted |
| 1372255.at | XM.001065238 | Arginyl-tRNA synthetase (predicted) | Rars_predicted |
| 1370184.at | NM.017147 | Cofilin 1, non-muscle | Cfl1 |
| 1372461.at | NM.001012504 | EST | Set_predicted |

compounds including APL, AA, and BBz (Fig. 4o–q), and remaining 16 (Fig. 4r) were correctly predicted as negative.

3.4. Additional biological validation

In order to support the class discrimination results by PAM, hepatic total glutathione was quantified for the following selected samples: high dose MP- and TAA-treated groups, and BBz-treated groups.

Hepatic glutathione contents transiently reduced with peak values at 3H for MP, 6H for TAA and 9H for BBz after single dosing, and rapidly recovered 24H after the treatment (Fig. 5). Although hepatic glutathione content was kept at normal or higher in the BBz-treated group at all time points of repeated dose study, in the MP- and TAA-treated groups it reduced with repeated dosing (Fig. 5). These time course changes of the glutathione contents are clearly correlated with the change of the PAM score.

Of the PPAR α agonists, only WY, but not CFB and GFZ, showed positive scores at 15D and 29D. If the PAM classifier detects carcinogenesis via the activation of PPAR α and these three agonists stimulated the receptor to the same extent, all of three agonists should have been classified as positive. The dose of each compound had been determined based on a 7-day repeated preliminary study and thus the doses would not be proportional to their potency to the receptor. To assess the biological potency of each agonist *in vivo*, we compared the induction of acyl-coenzyme A oxidase 1, a gene directly regulated by PPAR α . As shown in Fig. 6, the dose of WY appeared to be too high, since enzyme induction reached its maximum by the low dose of WY. During repeated administrations,

however, the extent of the induction was almost the same as in the high dose of these three agonists. If the positive score of WY was due to its PPAR α activation, not only the high dose but also the middle and low dose should be classified as positive. We then performed PAM using the present classifier for the three doses of these three agonists, but no positive scores were obtained other than the high dose of WY at 15D and 29D (data not shown).

4. Discussion

The goal of the present study was to develop a classifier for early assessment of potential non-genotoxic hepatocarcinogenic-

Table 3
GO analysis of the PAM classifier

| Term | Count | Percentage | p-Value |
|--|-------|------------|----------|
| Cellular metabolism | 41 | 34.75 | 5.07E–03 |
| Primary metabolism | 38 | 32.20 | 1.80E–02 |
| Macromolecule metabolism | 31 | 26.27 | 8.64E–04 |
| Cell organization and biogenesis | 22 | 18.64 | 3.84E–05 |
| Biosynthesis | 14 | 11.86 | 8.19E–03 |
| Cellular localization | 12 | 10.17 | 3.03E–04 |
| Cell proliferation | 10 | 8.47 | 6.26E–03 |
| Negative regulation of physiological process | 10 | 8.47 | 1.66E–02 |
| Negative regulation of cellular process | 10 | 8.47 | 2.75E–02 |
| Protein localization | 9 | 7.63 | 7.12E–03 |
| Cell cycle | 9 | 7.63 | 1.10E–02 |
| Cell death | 8 | 6.78 | 4.27E–02 |
| Cellular morphogenesis | 7 | 5.93 | 2.60E–02 |
| Response to DNA damage stimulus | 5 | 4.24 | 2.58E–02 |
| Regulation of response to stimulus | 2 | 1.69 | 2.33E–02 |

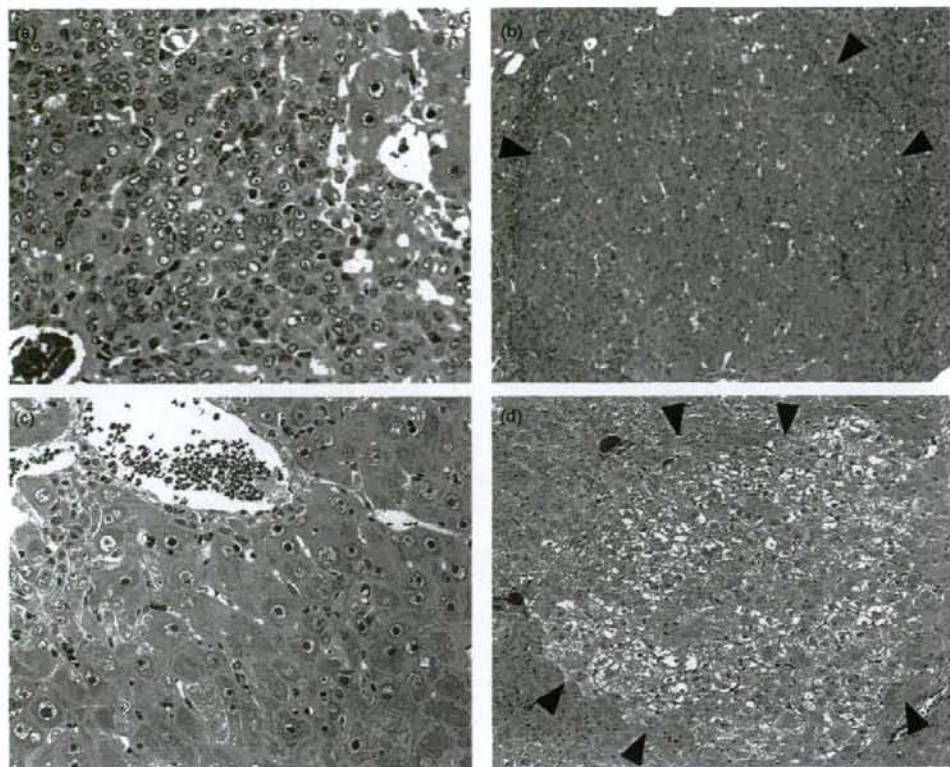


Fig. 1. Histopathology of rat liver treated with MP or TAA for 28 days. Repeated administrations of high dose of MP (100 mg/kg) for 28 days caused hepatocellular hyperplasia and some affected portal regions contained increased numbers of oval cells arranged in clusters without a distinct lumen (a), and in some cases, a pre-neoplastic altered hepatocellular focus was seen (b; arrowheads). In the centrilobular region of rat liver treated with repeated administrations of high dose of TAA (45 mg/kg) for 28 days, hepatocytes exhibited hypertrophy with large, atypical nuclei (c). As in methapyriline, a pre-neoplastic altered hepatocellular focus was also observed (d; arrowheads).

ity of chemicals based on gene expression changes stored in our database, TG-GATES. In order to utilize the classifier for practical drug development, we did not attempt to explore an original algorithm but to use a well-established one, i.e., PAM in the present case. Our advantage over the previous similar works was the quality of

the database, i.e., the quantitative gene expression data obtained in the single platform employing standardized and enriched protocol with three dose levels and eight time points (four for single and four for repeated). The enrichment of time and dose in the data has been shown to be quite powerful in toxicological analysis in various

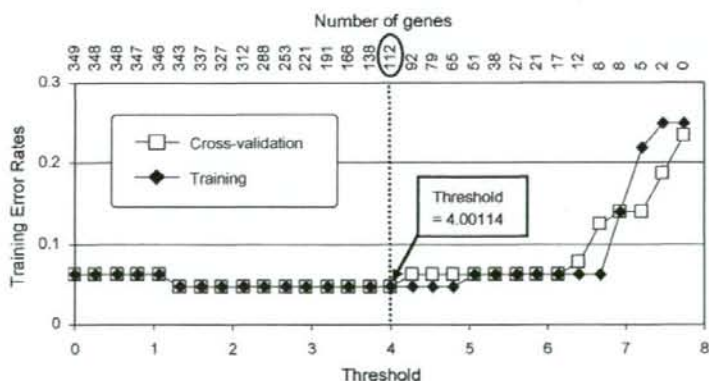


Fig. 2. PAM training and cross validation. PAM training was performed by comparing 2 positive compounds (MP and TAA, high dose group only) with 6 negative compounds (APAP, ASA, PhB, RIF, ANIT and AM) on the ratio of expression levels of the selected 349 probe sets for various time points (total of 64 training samples). Ten-fold cross validation was performed to find out the optimal classifier performance, which minimized classification errors for training sets. Both the training (black symbol) and cross-validated errors (white symbol) were minimized near the threshold = 4.00, where 112 genes (circled) were selected.

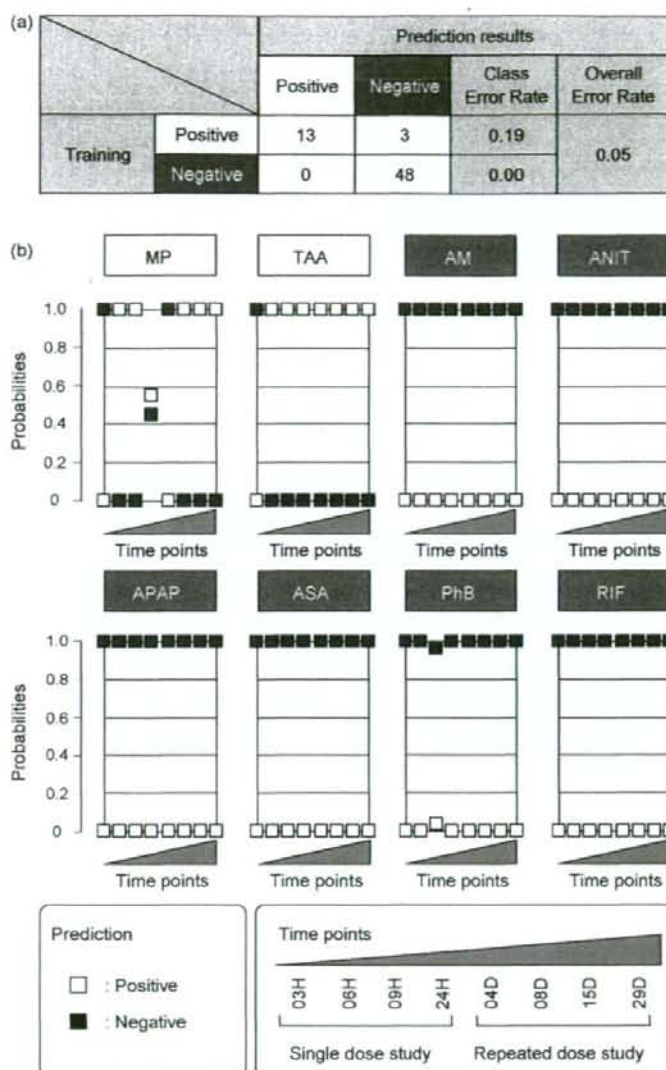


Fig. 3. Class discrimination by PAM. PAM prediction results for the condition determined in Fig. 2 are shown. (a) Prediction results of the training sets (13 positives and 48 negatives) are shown. Note that the overall success rate was 95%, i.e., 13 of the 16 positive sets (81%) and all of the negative sets (100%) were correctly classified. (b) Prediction result of individual sample. For each chemical, the samples are aligned with time as shown on the bottom. The samples predicted as positive are depicted with white and negative with black. Note that two out of three errors occurred at 3 h after single dosing.

ways (Urushidani, 2007). In the present study, genes showing clear dose- and time-dependent changes were successfully extracted by K-means clustering, and we could detect the changes of the score transient after single administration which then turned to be sustained after repeated administration. These also helped us consider the toxicological mechanism.

After PAM training, we produced a discriminator consisting of 112 of the mobilized probe sets that could discriminate between both classes with a high probability, >95%. In the training procedure, MP-3H, 4D, and TAA-3H were judged as false negatives. However, these results were considered to be reasonable because 3H of both compounds was too early for development of hepatotoxicity and 4D of MP treatment was the period when homeostatic recovery of the hepatic glutathione contents occurred.

In the present experiments, MP and TAA showed similar early morphological changes in rat liver, characterized as hepatocellular single cell necrosis with inflammatory response and hypertrophy with granular eosinophilic changes. This was confirmed by electron microscopy as proliferation and swelling of mitochondria (unpublished observations). In addition, hepatocellular altered foci were observed at 15D and/or 29D in the MP and TAA-treated groups. It is well known that this type of lesion is a pre-neoplastic transformation of the cells and is induced in the early stage of non-genotoxic hepatocarcinogenesis in the liver (Bannasch, 1976; Fischer et al., 1983). Therefore, early gene expression profiling in liver treated with these two compounds is considered to be closely related to future carcinogenesis.

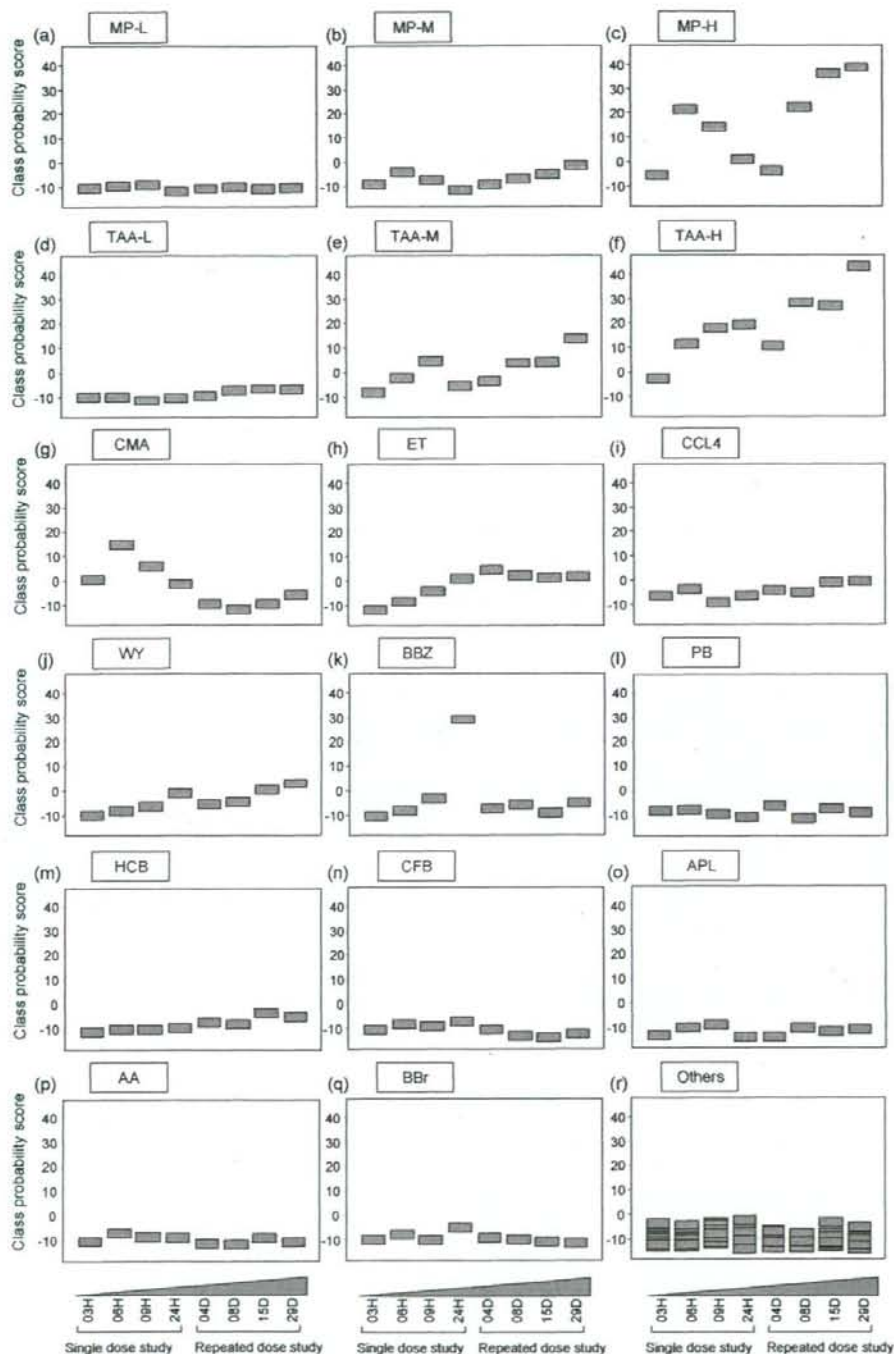


Fig. 4. PAM prediction score. The PAM class probability was converted to a score as described in Section 2 in order to enable quantitative comparison. The score is shown for MP ((a) 10 mg/kg, L; (b) 30 mg/kg, M; (c) 100 mg/kg, H), TAA ((d) 4.5 mg/kg, L; (e) 15 mg/kg, M; (f) 45 mg/kg, H), CMA ((g) 150 mg/kg), ET ((h) 250 mg/kg), CCL4 ((i) 300 mg/kg), WY ((j) 100 mg/kg), BBZ ((k) 300 mg/kg), PB ((l) 100 mg/kg), HCB ((m) 300 mg/kg), CFB ((n) 300 mg/kg), APL ((o) 150 mg/kg), AA ((p) 30 mg/kg), BBr ((q) 200 mg/kg), and (r) the other 17 chemicals. For abbreviation of the compounds, see Table 1.

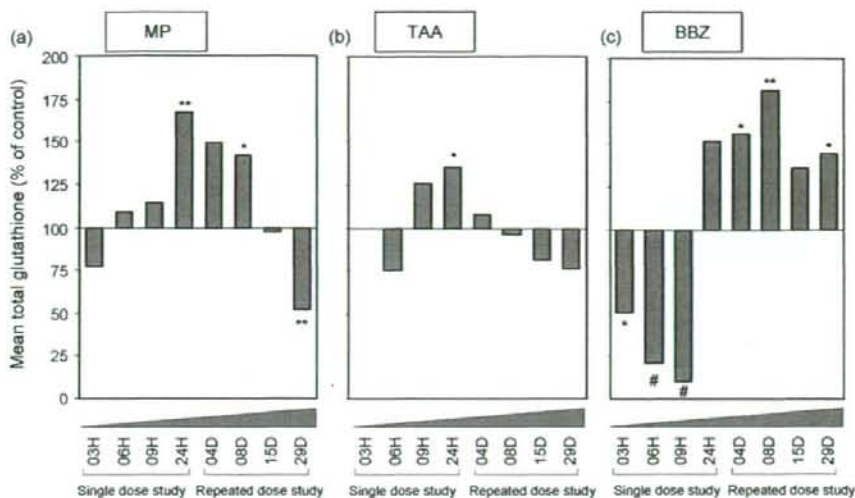


Fig. 5. Effects of MP, TAA or BBZ on glutathione contents in rat liver. Hepatic total glutathione was measured in the liver of rats receiving a high dose of MP (a), TAA (b) or BBZ (c), and their corresponding controls. Measurements were performed for five rats per group using Glutathione Quantification Kit. The results were expressed as percent of control at each time point. Statistical analysis was done by an unpaired two-tailed Student's *t*-test or Welch's *t*-test as appropriate. **p* < 0.05, ***p* < 0.01, by Student's *t*-test, #*p* < 0.05 by Welch's *t*-test.

The gene list selected as a marker for predicting hepatic carcinogenicity contained oxidant stress-, oxidative DNA damage-, and cell cycle regulation-related genes, which were changed in the early stage of administration. The oxidative stress is due to the production of reactive oxygen species more than the anti-oxidant capability of the target cells. Unregulated or prolonged production of cellular oxidants has been thought to lead to mutation as a result of oxidant-induced DNA damage, thought to participate in non-genotoxic carcinogenesis (Klaunig et al., 1998; Klaunig and Kamendulis, 2004). The observed expression changes in these genes is in accordance with previous reports that the repetitive cycle of DNA damage (initiation) and reproduction (promotion) caused by sustained oxidative stress is closely related to the carcinogenic process of non-genotoxic carcinogens. This does not mean that the classifier detects any compounds causing oxidative stress. Of the compounds used as negative sets, APAP is known as a prototypic oxidative stressor, which induces glutathione depletion in liver when overdosed (James et al., 2003; Kiyosawa et al., 2004). ASA was reported to induce some antioxidant enzymes and components

(Cai et al., 1995), and stimulates some beta-oxidation enzymes, bringing about an overproduction of H_2O_2 (Rivero et al., 1994). PhB was reported to accelerate glutathione oxidation and it induces lipid peroxidation of microsomes (Miura et al., 2002). All of these were successfully classified as negative, suggesting that the classifier discriminates non-carcinogens causing oxidative stress.

The validity of the presently developed discriminator for carcinogenesis was examined on our large-scale database, and all of the 20 chemicals except BBZ (selected as a non-carcinogen) were judged as negative at any time points. Of the eight chemicals classified as non-genotoxic carcinogens, CMA, ET, CCl₄ and WY showed positive prediction and increase in the PAM prediction scores in repeated administrations, whereas enzyme inducers such as PB and HCB, and other peroxisome proliferators were all judged as negative.

For CMA (Lake et al., 2002; National Toxicol Program, 1993), ET (Ogiso et al., 1990; Svardal et al., 1988), and CCl₄ (Castro et al., 1989; Natarajan et al., 2006), oxidative stress was reported as being involved in their hepatotoxicity and carcinogenesis. It could be con-

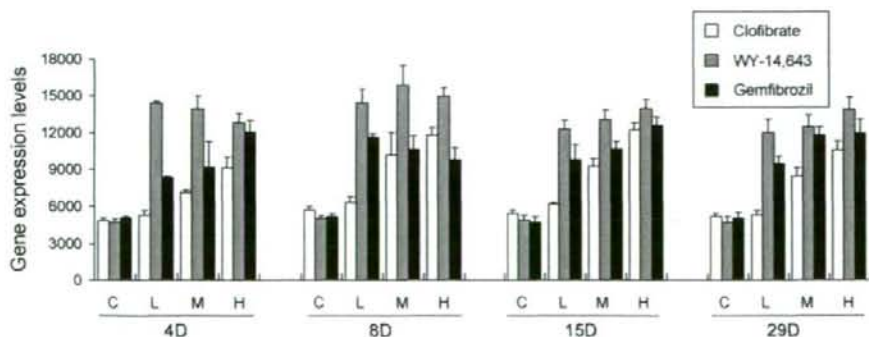


Fig. 6. Effects of repeated administration of CFB, WY or GFZ on expression of acyl-CoA oxidase-1. Expression of acyl-CoA oxidase-1, a gene directly regulated by PPAR α , was measured by GeneChip, and the intensities were normalized for each chip by setting the mean intensity to 500 (per chip normalization). The results were expressed as mean \pm S.D. (*n* = 3). For each panel, C: control, L: low dose, M: middle dose, H: high dose, for CFB: 30, 100, 300 mg/kg; WY: 10, 30, 100 mg/kg; GFZ: 30, 100, 300 mg/kg, respectively.

cluded that sustained oxidative stress plays an important role in their carcinogenesis, as in MP and TAA.

The induction of PPAR α in rodents treated with peroxisome proliferators was considered to be related to hepatocarcinogenesis (Holden and Tugwood, 1999). Moreover, increased levels of H₂O₂ generation, hydroxyl free-radical formation and lipid peroxidation were found in the liver of rats following long-term treatment with peroxisome proliferators. It was also reported that 8-hydroxydeoxyguanosine was found in the liver DNA of rats chronically treated with a PPAR α (Reddy and Lalwai, 1983; Reddy and Rao, 1989). In the present study, our discriminator designated WY as positive among the PPAR α agonists, CFB, GFZ, and WY. This result suggests that either the discriminator could predict the carcinogenesis of PPAR α agonists (although its sensitivity is relatively low) or that WY had an additional carcinogenicity differing from other PPAR α agonists. The latter would be more likely since the low and middle doses of WY (by which the induction of acyl-CoA oxidase 1 reached a maximum) did not classify as positive and since the highest doses of CFB and GFZ induce acyl-CoA oxidase 1 to almost the same extent as WY. It was also suggested that WY might share a carcinogenic mechanism with MP and TAA apart from its PPAR α agonist's activity.

The P450 enzymes generate oxygen free radicals in the process of metabolizing xenobiotic chemicals (Parke and Ioannides, 1990), including PB (Uitley and Mehendale, 1991) and HCB (Smith and De Matteis, 1990). Kinoshita et al. (2002) reported that PB-induced reversible alteration to nuclear 8-hydroxydeoxyguanosine by oxidative stress in rat liver after several days of continuous application. Furthermore, Elrick et al. (2005) provided evidence for the relationship between oxidative stress and PB-induced non-genotoxic hepatic carcinogenesis. On the other hand, HCB exposure induces long-term alterations in intercellular communication via gap junction in rat liver. This effect is thought to be a critical mechanism of HCB-induced non-genotoxic hepatocarcinogenesis and tumor promotion (Plante et al., 2002). However, these chemicals were classified as non-carcinogens based on gene expression profiling. There are likely to be numerous mechanisms involved in non-genotoxic rodent hepatic carcinogenesis. Therefore, it is thought that these chemicals induce non-genotoxic hepatocarcinogenesis through chemical-specific mechanisms.

For the evaluation of these results of prediction, we developed a PAM prediction score based on the positive/negative class probability. In the present study, we compared the score with the hepatic glutathione contents in order to examine the validity of the prediction. In association with the largest decrease of hepatic glutathione contents at 3H (MP), 6H (TAA) and 9H (BBZ), the PAM prediction score increased with the peak at 6H (MP), 9H (TAA) and 24H (BBZ). This could be explained as follows: hepatic glutathione was rapidly consumed to detoxify the oxidants produced by these toxicants, and in the subsequent glutathione-depleted state the expression of these marker genes was up-regulated. The excess production of glutathione for homeostasis tended to decrease in MP or TAA, whereas its high value was maintained in BBZ during their repeated administrations. It is known that some reactive intermediates are conjugated with glutathione to be excreted from the cell. The hepatotoxicity of the acute dose of BBZ was significantly reduced by prior sub-chronic exposure to BBZ. Therefore, the enhanced BBZ excretion by glutathione conjugation could partly explain such potential tolerance against its acute hepatotoxicity (Chakrabarti and Brodeur, 1984). It would be reasonable to speculate that BBZ, which causes transient hepatic and DNA damage by oxidative stress at the early stage of dosing, does not result in hepatic cancer since metabolic protection against oxidative stress does not allow the sustained stressful condition up to 28 days of administration, whereas a breakdown of protection occurs in the case of MP and TAA sug-

gested by the glutathione contents. There was a close correlation between the pattern of change in glutathione and PAM scores, supporting the usefulness of the present marker genes. The present scoring system also enables us to make a prediction based on important toxicological points, e.g., dose- and time-dependency and it would be a quite convenient way for evaluation of the results of discriminant analysis.

In summary, we showed that the expression profile of 112 genes selected by the PAM method could make a prediction of oxidative stress-related hepatocarcinogenicity with high precision at the early stage of administration. The possibility of non-genotoxic carcinogenicity is suggested as early as 24 h after the single dosing. Although pseudo-positives are included in the chemicals selected by the single dose experiments, these can be discriminated by the prediction based on repeated administration up to 28 days. At present, tests for carcinogenicity using rats takes at least 2 years. The present study has suggested a possibility to enable it to take as short as 28 days with high precision. Although neither a single gene nor a single pathway is sufficient to predict non-genotoxic hepatocarcinogens at present, it is evident that combinations of biomarker gene sets appeared to be useful for prediction of carcinogenesis. Further study is clearly necessary to clarify the pathophysiological roles of the genes included in the marker gene list for the process of carcinogenesis.

Conflict of interest

None.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.tox.2008.05.013.

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Original Article

Gene expression profiling of methapyrilene-induced hepatotoxicity in rat

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ABSTRACT — The present study was conducted as a model case of the toxicogenomics approach for analyzing toxicological mechanisms and toxicity assessments in the early stage of drug development by comparing with classical toxicology data. Methapyrilene (MP) 100 mg/kg produced obvious histopathological changes in liver of rats by single or repeated dose up to 28 days with significant elevation of ALT and AST. In the middle dose groups (30 mg/kg MP), no apparent changes were noted in blood biochemical data by single dosing or repeated dosing up to one week, and no obvious histopathological changes were observed except a slight hypertrophy in the hepatocytes. Comprehensive gene expression changes were analyzed using Affymetrix GeneChip® and differentially expressed probe sets were statistically extracted. These contained many genes related to “glutathione metabolism”, “apoptosis”, “MAPK signaling pathway” and “regulation of cell cycle”, which were all thought to be involved in the development of presently observed phenotypes. In the high dose groups, TGP1 scores (developed in our system in order to overview the responsiveness of drugs to multiple marker gene lists) for these categories were markedly increased from the early time point after single dose and kept their high expression throughout the repeated dose period. In the middle dose groups, the increment of the scores were noted not only at the time points when apparent pathological changes emerged, but also at the earlier stage of repeated dosing and even after single dosing. We conclude that toxicogenomics would enable a more sensitive assessment at the earlier time point than classical toxicology evaluation.

Key words: Methapyrilene; Rat; Liver; Toxicogenomics; Microarray

INTRODUCTION

The toxicogenomics approach has attracted attention in the field of drug safety assessment as a promising tool in predicting the toxicity of chemicals and analyzing the mechanism of toxicity. Classical toxicology works to characterize the emerged toxic changes, but it is not always powerful in predicting potential toxicity that has not emerged at the point of assessment but might occur in the future or to detect serious disease without apparent change

in observation of the classical phenotype. On the other hand, extraction of toxicologically meaningful information from comprehensive gene expression analysis is expected to be useful since these changes precede toxicity and occur in the lower dose range.

The toxicogenomics project was a 5-year collaborative project conducted by the National Institute of Health Sciences, the National Institute of Biomedical Innovation and 15 pharmaceutical companies in Japan that started in 2002 (Urushidani and Nagao, 2005). Its aim was to construct a

large-scale toxicology database of transcriptome for prediction of toxicity of new chemical entities in the early stage of drug development. About 150 chemicals, mainly medicinal compounds, were selected, and gene expression in the liver (also the kidney in some cases) was comprehensively analyzed by Affymetrix GeneChip®. In 2007, the project was finished and the whole system, consisting of the database, the analyzing system and prediction system, was completed and named as TG-GATEs (Genomics Assisted Toxicity Evaluation System developed by the Toxicogenomics Project, Japan).

In the present study, we selected methapyrilene, a prototypic hepatotoxicant (Lijinsky *et al.*, 1980), as a model case of the toxicogenomics approach for analyzing the toxicological mechanism and toxicity assessment in the early stage of drug development by comparing with classical toxicology data.

MATERIALS AND METHODS

Chemical

Methapyrilene (MP) was obtained from Sigma Chemical Company (St. Louis, MO, USA) and a suspension formulation was prepared by mixing with 0.5% methylcellulose (MC) solution.

Animal and experimental design

Five-week old male Sprague-Dawley rats were obtained from Charles River Japan Inc. (Kanagawa, Japan). After a 7-day quarantine and acclimatization period, the animals (6-week old) were assigned to dosage groups (5 rats per group) using a computerized stratified random grouping method based on individual body weight. The animals were individually housed in stainless-steel cages in an animal room that was lighted for 12 hr (7:00–19:00) daily, ventilated with an air-exchange rate of 15 times per hour and maintained at 21–25°C with a relative humidity of 40–70%. Each animal was allowed free access to water and pellet diet (CRF-1, sterilized by radiation, Oriental Yeast Co., Tokyo, Japan).

According to the standard protocol in our project, rats received single or repeated doses of MP by gavage at doses of 0 (vehicle only), 10, 30 or 100 mg/kg. For the single-dose study, rats were sacrificed at 3, 6, 9 and 24 hr after dosing. For the repeated dose study, the animals were treated daily for 3, 7, 14 and 28 days, and they were sacrificed 24 hr after the last dosing. The animals were euthanized by exsanguination from the abdominal aorta under ether anesthesia after blood sampling, and liver samples were obtained from the left lateral lobe of the liver in each animal immediately after sacrifice for the following exam-

inations.

The experimental protocols were reviewed and approved by the Ethics Review Committee for Animal Experimentation of the National Institute of Health Sciences.

Blood chemical examination

Blood samples were collected upon sacrifice in tubes containing heparin lithium, and aspartate aminotransferase (AST) and alanine aminotransferase (ALT) were measured using an auto analyzer (Hitachi 7080).

Histopathological examination

For light microscopic examination, liver samples were fixed in 10% neutral buffered formalin, dehydrated in alcohol and embedded in paraffin. Paraffin sections were prepared and stained by a routine method using hematoxylin and eosin (H&E).

Microarray gene expression analysis

Microarray analysis was conducted on 3 out of 5 samples for each group. Total RNA was isolated from RNAlater® (Ambion, Austin, TX, USA)-preserved samples using RNeasy kit by Bio Robot 3000 (Qiagen, Valencia, CA, USA). Homogenization was conducted by Mill Mixer (Qiagen) and zirconium beads. Purity of RNA was checked by gel electrophoresis confirming that the 260/280 nm ratio was between 2.2 and 3.0. Gene expression analysis was carried out using GeneGhip® RAE230A probe arrays (Affymetrix, Santa Clara, CA, USA) containing 15,923 probe sets. The procedure was basically conducted according to the manufacturer's instructions as previously reported. Microarray Analysis Suite 5.0 (MAS; Affymetrix) was used to quantify microarray signals and the intensities were normalized for each chip by setting the mean intensity to 500 (per chip normalization).

Microarray data analysis

To determine differentially expressed genes between high and time-matched control sample groups, Welch's *t* test was applied with a *p* cut off value of 0.01 in combination with a 1.5-fold regulation-ratio of means using GeneSpring software (Agilent Technologies, Inc., Santa Clara, CA, USA). Probe sets, which were labeled as absent by Affymetrix detection call in any of the 48 samples in single or repeated dose study, were excluded from further analysis. For the extracted probe sets, showing significant changes for at least one time point of either single or repeated administrations, pathway and GO analysis was performed using David 2.1 beta (<http://david.abcc.ncifcrf.gov/>) to identify overrepresented gene categories in

each gene list, and a *p* value of < 0.05 determined by Fisher's exact test was considered statistically significant.

Scoring of the marker gene sets (TGP1 score)

To facilitate the analytical procedures for large-scale microarray data, we developed a simple one-dimensional score, named TGP1, which is useful to overview the trend of the changes in expression of multiple biomarker gene lists all at once (Kiyosawa *et al.*, 2006). For each gene list, the signal log ratio was calculated by dividing the mean signal value of the chemical-treated group by that of the corresponding control. First, the sum of the signal log ratios for the used probe sets was calculated, and then divided by the number of probe sets used (Index 1). Next, the sum of squared signal log ratios for the used probe sets was calculated, and then divided by the number of probe sets used (Index 2). Finally, the TGP1 score was calculated by multiplying Index 1 with Index 2.

Statistical analysis

For blood chemical parameters, ANOVA followed by Dunnett's multiple comparison test or Kruskal-Wallis mean rank test followed by Mann-Whitney's U test was used as appropriate (Snedecor and Cochran, 1989).

RESULTS

Conventional toxicological parameters

Measurements of AST and ALT, representative markers of hepatotoxicity, are shown in Fig. 1 and histopathological findings are summarized in Tables 1 and 2. In the highest dose groups, both AST and ALT were elevated 3 or 6 hr after treatment, and they kept increasing with time. They also showed histopathological changes at every time point, such as hepatocellular hypertrophy and single cell necrosis in the peripheral lobular region, and obvious inflammation and anisonucleosis were noted at 24 hr after dosing (Fig. 2a). Furthermore, these lesions were aggravated and additional regenerative changes such as increased mitosis, bile duct proliferation, and hyperplasia, during repeated administrations were evident (Fig. 2b). In the middle dose groups, no apparent changes were noted in blood biochemical data by single dosing or repeated dosing up to one week, and no obvious histopathological changes were observed except a slight hypertrophy in the hepatocytes. At the time of the 15th and 29th days, single cell necrosis and infiltration of mononuclear cells were noted as in the highest group, though their severity was low. In the lowest dose groups, no obvious changes were noted either in blood biochemistry or in histopathology except for one case with hepatocyte hypertrophy in each of the 8th and

15th days.

Gene expression changes

Using the highest dose group showing obvious hepatotoxicity in terms of both blood biochemistry and histopathology, we extracted genes that showed statistically significant changes at least once in any time point in single and repeated dose experiments. For single and repeated dosing, up-regulated probe sets were 399 and 2509, respectively, and down-regulated probe sets were 235 and 876, respectively, namely, expression changes occurred in many genes in repeated dosing where obvious pathophysiological changes emerged.

Extracted genes were categorized by pathway and GO analysis and the results are summarized in Tables 3 (up-regulated) and 4 (down-regulated). Among the genes up-regulated in single dose experiments, gene ontologies or pathways related to "regulation of cell cycle", "MAPK signaling pathway" and "glutathione metabolism" were still significantly up-regulated in repeated dosing. The genes related to "apoptosis" or "ribosome" were clearly up-regulated in repeated dosing, while they were not affected by single dosing.

In single dose experiments, the number of down-regulated genes was quite a few (Table 4). Significant suppression of gene expression by repeated administration was noted in various categories, including "starch and sucrose metabolism", "steroid metabolism", "complement activation" and "complement and coagulation cascades".

We considered the pathways and GO categories of "glutathione metabolism", "apoptosis", "MAPK signaling" and "regulation of cell cycle" as most important in the toxicological mechanisms of MP. In order to overview the effects of MP on these pathways, we calculated the TGP1-score for each (Table 5). Upon calculation of the score, redundant probe sets were unified based on their reliability and dose-dependency. It is obvious from Table 5 that the marker scores of these categories markedly increased in the early stage of single dosing of the highest dose and they kept increasing during repeated dosing. In case of middle dosing, an obvious increase of scores for glutathione metabolism and apoptosis was detected not only at the 15th and 29th days with obvious pathological changes but also at the 4th and 8th days of repeated dosing and 6 hr after single dosing. In the lowest dose groups, the only detectable change was a tendency of increment in the score of apoptosis at the 8th day.