Several studies using microarrays have shown that changes in gene expression provide information about the mechanism of toxicity induced by xenobiotic agents. Nevertheless, the issue of whether gene expression profiles are reproducible across different laboratories remains to be determined. To address this question, several members of the Hepatoxicity Working Group of the International Life Sciences Institute Health and Environmental Sciences Institute evaluated the liver gene expression profiles of rats treated with methapyrilene (MP). Animals were treated at one facility, and RNA was distributed to five different sites for gene expression analysis. A preliminary evaluation of the number of modulated genes uncovered striking differences between the five different sites. However, additional data analysis demonstrated that these differences had an effect on the absolute gene expression results but not on the outcome of the study. For all users, unsupervised algorithms showed that gene expression allows the distinction of the high dose of MP from controls and low dose. In addition, the use of a supervised analysis method (support vector machines) made it possible to correctly classify samples. In conclusion, the results show that, despite some variability, robust gene expression changes were consistent between sites. In addition, key expression changes related to the mechanism of MP-induced hepatotoxicity were identified. These results provide critical information regarding the consistency of microarray results across different laboratories and shed light on the strengths and limitations of expression profiling in drug safety analysis.

Key words: methapyrilene, microarray, support vector machine, toxicogenomics, unsupervised algorithms, variability. Environ Health Perspect 112:439–448(2004). doi:10.1289/txg.6643 available via http://dx.doi.org/ [Online 15 January 2004]
minimal expression of single-cell necrosis with minimal mononuclear infiltrate without associated changes in clinical chemistry parameters (Waring et al. 2001). Thus, in the present study we chose 100 mg/kg/day as the high dose expected to elicit hepatotoxicity. A dose of 10 mg/kg/day was selected as the low dose with the expectation that no hepatotoxic effect would be observed.

Male Sprague-Dawley rats were obtained from Charles River Laboratories, Inc. (Wilmington, MA). Rats were 57 days old and weighed 233.4–274.0 g at the start of the treatment. Upon arrival to Abbott Laboratories (Abbott Park, IL), all rats were acclimated for 6 days before treatment began. The two treatment groups comprising four rats each received the test compound at a concentration of 10 or 100 mg/kg, respectively. Animals in the equally sized control group received vehicle only.

Rats were dosed once daily by gavage for 7 days. The dose volume was 10 mL/kg. Doses were milligram salt per kilogram per day and were calculated for each rat on the basis of the most recent body weight data available. Rats were fasted overnight after their last treatment, euthanized under halothane anesthesia and submitted for necropsy. Each rat received its last treatment approximately 24 hr before scheduled necropsy.

In vivo observations, pathology, and sampling. All rats were observed twice each day during the pretreatment and treatment periods for survival and general condition. Blood samples were drawn from all rats, and clinical chemistry parameters were obtained for alanine aminotransferase (ALT), aspartate aminotransferase (AST), sorbitol dehydrogenase (SDH), alkaline phosphatase (ALKPHOS), total bilirubin (TBIL), glutathione (GSH), and triglycerides (TRIG). At necropsy, liver was weighed and the percent of body weight of each organ was calculated. One part of the liver (left lateral lobe) was fixed for potential histopathology in 10% formalin and subsequently sectioned and stained with hematoxylin and eosin, while the rest of the organ was rinsed in phosphate-buffered saline, immediately flash-frozen in liquid nitrogen, and kept frozen for subsequent RNA isolation.

RNA Isolation and Distribution

Approximately 100 mg of tissue from each liver was placed into TRIzol reagent (Invitrogen, Corp., Carlsbad, CA) and homogenized. Total RNA isolation was performed exactly according to the TRizol reagent protocol. The remaining portion of the liver was retained frozen. Following isolation, the RNA was quantitated using a BioRad SmartSpec 3000 spectrophotometer (BioRad, Hercules, CA), and the integrity of the RNA was determined using an Agilent 2100 bioanalyzer (Agilent Technologies, Palo Alto, CA). The RNA from the four animals in each treatment group was then pooled using equivalent amounts from each sample. The RNA was aliquoted and precipitated in ethanol and ammonium acetate for shipment to the participating DNA microarray users. In addition, RNA from individual animals was shipped to some of the DNA microarray analysis laboratories.

DNA microarray analysis. RNA samples were analyzed independently by five different Affymetrix users: Boehringer-Ingelheim Pharmaceuticals, Novartis, Pfizer Inc, F. Hoffmann-La Roche AG, and Schering AG. Voucher samples used for the design of the microarray were derived from Build 34 of the UniGene database (http://www.ncbi.nih.gov/UniGene; created from Genbank 107/dbEST 11/18/98) and supplemented with additional annotated gene sequences from Genbank 110 (http://www.ncbi.nih.gov/GenBank/). UniGene clusters are represented by an example sequence that is the most complete and most 3’ sequence in the cluster. The oligonucleotide probes are 25mers and 16 probe pairs per sequence are used. Processing of RNA and GeneChip experiments was carried out basically as recommended by Affymetrix, with some user-specific variations (Table 1) (Lockhart et al. 1996). An initial amount of 5–20 µg total RNA was used for the synthesis of double-stranded cDNA with a commercially available kit (Superscript Choice System; Invitrogen Life Technologies or Roche Molecular Biochemicals, Mannheim, Germany) in the presence of a T7-(dT)24 DNA oligonucleotide primer. After synthesis, the cDNA was purified with phenol/chloroform/isoamylalcohol extraction and ethanol precipitation. The purified cDNA was then transcribed in vitro (Enzo Diagnostics, Inc. (Farmingdale, NY) or Ambion, Inc. (Austin, TX)) in the presence of biotinylated ribonucleotides to form biotin labeled cRNA. The labeled cRNA was then purified on an affinity resin (Rneasy: Qagen, Inc., Valencia, CA), quantified and fragmented. An amount of 10–20 µg labeled cRNA was hybridized for approximately 16 hr at 45°C to an expression probe array. The array was then washed, stained with streptavidin-R-phycocyanin (SAPE; Molecular Probes, Eugene, OR), and the signal amplified using a biotinylated goat anti-streptavidin antibody (Vector Laboratories, Burlingame, CA) followed by a final staining with SAPE. Arrays were stained using the GeneChip Fluidics Workstation 400 (Affymetrix) and then scanned twice using a confocal laser scanner (GeneArray Scanner 2500; Hewlett Packard (Palo Alto, CA) or Agilent Technologies), resulting in one average scanned image.

Data analysis. Tab-delimited files obtained from the Affymetrix Microarray Suite software, version 4.0, (*.cch files) and containing data on signal intensity [average difference (Avg Diff)] and categorical expression-level measurement (Absolute Call) were used for analysis. Data were normalized and further analyzed using Roche in-house developed software (RACE-A; F. Hoffmann-La Roche AG, Mannheim, Germany). Briefly, this software performs a normalization step on the signal intensities based on the average signal (Mean Avg Diff) of each microarray before calculating additional parameters. In the cases where biological replicates were included, RACE-A was also used to calculate the average signal (arithmetic mean), and SD for each probe set. Also, comparative analysis between control and treated was performed including fold change (Avg Diff Treated/Avg Diff Control) and a significance value (p-value), calculated using a two-tailed, unpaired t-test. Once the required statistical parameters were calculated, data were filtered and exported to MS-Excel 2002 (Microsoft Corp., Bellevue, WA) or additional software for visualization and further analysis.

In addition, methods comprising more sophisticated algorithms and designed

**Table 1. Sample preparation methods used by the contributing companies.**

| Gene expression analysis site | Acronym | Sample type | cDNA | IVT |
|------------------------------|---------|-------------|------|-----|
| Boehringer Ingelheim | BI | Pool | SSI, Invitrogen | Enzo-Affymetrix |
| Pharmaceuticals | | | | |
| Novartis Pharma AG | Nov | Pool | SSI, Invitrogen | Enzo-Affymetrix |
| Pfizer Inc | Pfi | Pool | SSI, Invitrogen | Enzo-Affymetrix |
| F. Hoffmann-La Roche | RO | Pool | AMV, Roche Molecular Biochemicals | Ambion, Inc. |
| F. Hoffmann-La Roche | RO | Individual | AMV, Roche Molecular Biochemicals | Ambion, Inc. |
| Schering AG | Sch | Pool | SSI, Invitrogen | Enzo-Affymetrix |
| Schering AG | Sch | Individual | SSI, Invitrogen | Enzo-Affymetrix |

Abbreviations: AMV, avian myeloblastosis virus; Enzo-Affymetrix, Enzo Diagnostics, Inc. and Affymetrix, Inc.; IVT, in vitro transcription; SSI, Superscript II.
specifically for multivariate data analysis such as microarray data were employed. These methods share the characteristics of reducing the dimensionality of the data to a number of dimensions (components or vectors) that explain most of the variability in the data set. They are better suited to microarray analysis and generally superior in performance than gene-by-gene analysis with conventional statistical tests because they take into account the complex data structure. Such methods are known as unsupervised (hierarchical clustering and principal component analysis (PCA)) or supervised [support vector machines (SVMs)] multivariate analysis methods. Supervised methods such as SVMs are based on algorithms that learn from a selected training data set and use this previously acquired knowledge about classes to classify unknown data. The algorithm solves the classification problem while aiming to minimize the probability of false classifications for initially unknown test data. The basic idea of the SVM method and detailed explanations are described elsewhere (Cristianini and Shawe-Taylor 2000; Schölkopf et al. 1999).

Unsupervised methods such as clustering algorithms and PCA are commonly used to determine if gene expression patterns allow the discrimination of natural subpopulations that might bear a biological meaning such as treated/untreated or healthy/diseased. PCA is a mathematical technique that reduces the dimensionality of highly multivariate data. The reduced dimensions (or components) actually describe the major part of the variation in the samples and separate natural subpopulations without a priori knowledge (Liu et al. 2002).

Cluster analysis is a method used to organize primary data. Pairwise average-linkage cluster is a form of unsupervised hierarchical clustering commonly used for the analysis of microarray data. Relationships among objects such as experimental conditions or genes are represented by a tree whose branch lengths reflect the degree of similarity between the objects as assessed by a pairwise similarity function based on correlation coefficients (Eisen et al. 1998). The clustering tools and SVM used in this analysis are modules of RACE-A, whereas PCA was performed using SIMCA-P (Umetrics, Umea, Sweden).

The complete data set is currently being submitted to ArrayExpress (EMBL-European Bioinformatics Institute, Hinxton, UK: http://www.ebi.ac.uk/arrayexpress) and will be available for public download by the second quarter of 2004. Accession numbers referencing this data set will be available on the HESI website (http://hesi.iis.org/index.cfm?pubentityid=120).

### Results

**Clinical chemistry and histopathology.** A significant change in both body weight and food consumption compared to that of control groups was seen in the high-dose, but not in the low-dose group (data not shown). Clinical chemistry values confirmed liver toxicity occurred in the high-dose rats (Table 2). There were no significant changes at the low dose. Significant increases in leakage enzyme (AST, SDH and ALKP) indicate both hepatocellular and cholangiolar injury. The dose-dependent decline in serum glucose and a trend toward a decrease in triglyceride levels might indicate compromise of hepatocellular metabolic function but may also have been influenced by reductions in food consumption.

No compound-related histopathological changes were found for the low-dose group, whereas several compound-related changes were seen in livers from rats treated at the high-dose level. These included cytoplasmic vacuolation of perisinusoid hepatocytes, minimal to mild necrosis of perisinusoid hepatocytes, increased infiltration of portal tracts by mononuclear inflammatory cells, and hyperplasia of oval cells along portal tracts.

**Comparisons across users.** After microarray analysis of the RNA, it was determined that different users obtained comparable results despite possible variation in the sample processing [from total RNA up to fragmented IVT (in vitro transcript)] and microarray hybridization protocols. In a preliminary round of analysis using rigid cutoff values to assess which genes were modulated, the number of genes detected as regulated in the pooled samples (2-fold increase or decrease) by each user were strikingly different (Table 3). All five users analyzing the pooled RNAs detected 254 genes that were regulated simultaneously, while each user recognized an excess of 1,000 genes as up- or downregulated. The data set generated at RO appeared to be a clear outlier with nearly twice the amount of modulated genes as the other users. This may be due to the modifications introduced in the sample preparation (Table 1), but no direct evidence is available in support of this.

Further microarray results demonstrated that when individual animals were analyzed, as opposed to pooled samples, the number of genes detected as induced/repressed was generally reduced. Table 3 shows the results from microarray analysis.

### Table 2. Clinical chemistry values for methapyrilene-treated rats.

| Rat no. | Dosage (mg/kg) | ALT (IU/L) | AST (IU/L) | SDH (IU/L) | ALKP(HOS) (IU/L) | TBL (mg/dL) | GLU (mg/dL) | TRIG (mg/dL) |
|---------|----------------|------------|------------|-----------|-----------------|------------|------------|------------|
| 1001    | 0              | 42         | 91         | 10.9      | 79              | 0.1        | 145        | 53         |
| 1003    | 0              | 25         | 106        | 6.6       | 170             | 0.1        | 136        | 19         |
| 1005    | 0              | 28         | 103        | 6.7       | 292             | 0.1        | 132        | 50         |
| 1007    | 0              | 30         | 90         | 7.3       | 197             | 0.1        | 121        | 48         |
| Average | 0              | 31.3       | 97.5       | 7.9       | 21.0            | 0.1        | 133.5      | 45.0       |
| SD      |                | 7.46       | 8.2        | 2.0       | 56.1            | 0.0        | 10.0       | 18.0       |
| 2001    | 10             | 51         | 168        | 17.4      | 275             | 0.1        | 102        | 28         |
| 2003    | 10             | 24         | 97         | 9.5       | 217             | 0.1        | 111        | 50         |
| 2005    | 10             | 23         | 90         | 9.1       | 235             | 0.1        | 138        | 29         |
| 2007    | 10             | 30         | 92         | 12.1      | 255             | 0.1        | 97         | 19         |
| Average | 10             | 32.0       | 111.8      | 12.0      | 245.5           | 0.1        | 112        | 31.5       |
| SD      |                | 13.0       | 37.6       | 3.8       | 25.1            | 0.1        | 18.3       | 13.1       |
| 3001    | 100            | 36         | 162        | 12.3      | 220             | 0.1        | 123        | 18         |
| 3003    | 100            | 56         | 179        | 12.9      | 283             | 0.4        | 100        | 28         |
| 3005    | 100            | 193 > 410  | 24.2       | 460       | 0.9             | 105        | 24         |
| 3007    | 100            | 51         | 200        | 19.4      | 417             | 0.3        | 88         | 15         |
| Average | 100            | 84.0       | 180.3*     | 17.2*     | 345.0*          | 0.4        | 104.0*     | 21.2       |
| SD      |                | 73.2       | 19.0       | 5.7       | 112.4           | 0.3        | 14.5       | 5.9        |

Abbreviations: ALKP, alkaline phosphatase; ALT, alanine aminotransferase; AST, aspartate aminotransferase; GLU, glucose; SDH, sorbitol dehydrogenase; TBL, total bilirubin; TRIG, triglycerides.

*Significantly different from the control group using two-tailed test (p-value < 0.05).

### Table 3. Number of genes regulated by methapyrilene across the different companies in the pooled and individual samples at the high dose.

|                  | Pooled RNA samples | Individual RNA samples |
|------------------|--------------------|------------------------|
| Common BI        | RO                 | Sch                    |
| Common RO/Sch    | All users          | RO                     |
| RO/Sch           | Sch                |                         |

**Upregulation**

| BI     | Nov | Pfi | RO | Sch | All users | RO | Sch |
|--------|-----|-----|----|-----|-----------|----|-----|
| Upregulation | 691 | 785 | 621 | 1,325 | 692 | 282 | 352 | 120 |

**Downregulation**

| BI     | Nov | Pfi | RO | Sch | All users | RO | Sch |
|--------|-----|-----|----|-----|-----------|----|-----|
| Downregulation | 480 | 283 | 728 | 292 | 405 | 130 | 75 | 262 |

Abbreviations: BI, Boehringer-Ingelheim Pharmaceuticals; Nov, Novartis Pharma AG; Pfi, Pfizer Inc; RO, F. Hoffmann-La Roche AG; Sch, Schering AG.

*Cut-off values: 2-fold change, p-value < 0.05.*
on pooled samples and individual animals conducted at RO and Sch. The inclusion of replicates very likely diminishes the influence of false signals. Nevertheless, there is still much disagreement among users when performing simple data analysis methods and defined cutoff values.

Whereas the gene expression analysis was not concordant between different laboratories, a critical question to be addressed is whether microarray results from all users reflected the observations from traditional toxicology markers and yielded similar mechanistic outcomes. When methapyrilene effects on the liver are examined, both histopathology and clinical chemistry analysis distinguished the high-dose animals from the low-dose and control animals. To determine if microarray analysis also distinguished between high and low dose, more sophisticated methods usually better suited to the analysis of highly multivariate microarray data were used. Among these methods, we chose to employ unsupervised as well as supervised approaches. Conversely, unsupervised methods are well suited to separate natural subpopulations in an unbiased manner. On the other hand, supervised methods allow incorporating knowledge obtained from the data (training set) to distinguish classes in the test data set. We analyzed the data using two unsupervised methods, namely PCA and hierarchical clustering. Using all expressed genes (4,846 probe sets), PCA analysis revealed a clear separation of the high-dose samples from controls and low-dose samples, despite the fact that the site differences are responsible for a large amount of variance. In this analysis, the second principal component (PC2; accounting for 15% of the variance) drives the treatment-related difference, as indicated by the arrows, whereas PC1 (accounting for 33% of the variance) showed a separation by site (Figure 1A, B). Excluding PC1 and relying exclusively on PC2 and PC3, a clear separation between high-dose–treated animals and the other two groups was achieved regardless of the site in which the sample processing was performed. Thus, the site-related differences do not mask the outcome of the classification. The low-dose samples could not be confidently distinguished from the vehicle-treated controls, a conclusion that accords with the clinical chemistry and histopathology findings.

To verify this latter conclusion, we grouped the data using another unsupervised clustering method, agglomerative hierarchical clustering. When we used the expressed genes employed for the PCA analysis, there was a tendency toward clustering by the site performing the microarray analysis (Figure 2A). An increase in statistical power

![Figure 1. Principal component analysis using all expressed genes (4,846 probe sets, Avg Diff Max ∼ 200, Mean Call ≥ 0.5). Abbreviations: BI, Boehringer-Ingelheim Pharmaceuticals; Nov, Novartis Pharma AG; Pfi, Pfizer Inc; RO, F. Hoffmann-La Roche AG; Sch, Schering AG; ind, individual; PC, principal component. All data points are from pooled samples unless otherwise indicated. (A) PC2 versus PC1, showing that PC1 is mainly driven by the site performing the microarray experiments. (B) PC3 versus PC2. Arrows show the separation of the high-dose samples from the vehicle and low-dose samples, mainly on the PC2.](image)

![Figure 2. Hierarchical clustering using either (A) probe sets that were detected (4,846) or (B) probe sets that showed concordant regulation in the analyses performed by the two sites using RNA from individual animals (193 probe sets). Abbreviations: BI, Boehringer-Ingelheim Pharmaceuticals; Nov, Novartis Pharma AG; Pfi, Pfizer Inc; RO, F. Hoffmann-La Roche AG; Sch, Schering AG; CONT, control; ind, individual. Each branch represents either a single chip (pooled samples) or the mean value of four replicates (individual samples). The correlation distance is represented by the distance between the samples in the dendrogram. Correlation coefficient values ranged between 0.854634 and 0.999306 in Figure 2A and between 0.490595 and 0.989861 in Figure 2B.](image)
can be achieved by including the confidence information obtained from the analysis of biological replicates (Lee et al. 2000). For most users conducting these experiments, individual replicates were not available, as the RNA had been pooled. However, replicates were available from the two sites that performed microarray analysis on individual animals. Thus, we performed hierarchical agglomerative clustering using the probe sets that were regulated in common by the high dose of MP from the individual replicates obtained by Sch and RO. With this smaller subset of genes, hierarchical clustering of the treatment groups allowed the high dose to be discriminated from the controls and low-dose–treated animals (Figure 2B). Similar to the PCA analysis, the low-dose samples could not be distinguished from the control samples.

To improve the discrimination between the groups, further analysis was performed using supervised methods. Because results from biological replicates were provided by two sites (RO and Sch), it was possible to generate a training set using the profiles obtained from the individual animals. This training set consisted of two analyses (one per site) that included 4 animals in each treatment group, amounting to 24 microarrays generated from 12 animals. This training set has the limitations of being rather small and of including in each group four biological replicates (independent) and for each of these independent replicates, two processing replicates from different sites (nonindependent). The data from individual animals were analyzed using the SVM to identify probe sets that were distinct for the three different classes of treatment (vehicle, low-dose, high-dose). The training of the SVMs and the subsequent classification were performed using all probe sets on the chip (8,799). Once the SVM was thus trained, the 15 microarrays obtained from the analysis of the pooled samples (5 controls, 5 low-dose, 5 high-dose) were used as test samples and classified. In this case, samples obtained from animals treated with low or high doses of MP were correctly classified. Classification of the control animals was relatively ambiguous, as only 2 animals were correctly classified as controls, whereas the other 3 showed no similarity to either group (Figure 3). An example of some of the genes that allow the distinction between control and treated animals (low- and high-dose) is shown in Figure 4. Thus, using supervised clustering, together with biological replicates, it was possible overall to distinguish not only the high-dose–treated group, but also the low-dose–treated group from the controls. This was not unequivocally possible using clinical chemistry, histopathology, or unsupervised clustering methods.

**Genes affected by MP.** More important than the number of regulated genes is the determination of the identity of the regulated genes, the affected cellular pathways, and their biological significance. Some genes described previously as regulated by MP or that are associated with the histopathology findings were consistently detected by all involved users. Genes associated with cell stress, cell damage or apoptosis, and

![Figure 3. Classification of the pooled samples using support vector machines. The program was trained with the individual analyses provided by RO and Sch; all five pooled analyses were used as test sets. Training and classification were performed using all probe sets on the microarray. All low-dose (blue bars) and high-dose (red bars) samples were correctly classified, as well as two of the five controls (black bars). Three additional pooled controls could not be classified in any of the training groups.](image)

![Figure 4. Examples of genes modulated at both the low and the high doses of MP. Dotted lines represent average values of the individual analyses by two users (RO and Sch) and solid lines each of the analyses of pooled samples by five users. For each panel, identification numbers in parentheses are Affymetrix probe set ID codes.](image)
| Affymetrix probe set ID | Class     | Gene                      | Max Signal | RD (individual) | Sch (individual) | Direction |
|-------------------------|-----------|---------------------------|------------|----------------|-----------------|-----------|
| X75207_s_at             | Cell cycle| Cyclin D1                  | 696        | 0.193          | 0.023           | Down      |
| D10414_g_at             | Cell cycle| Cyclin D1                  | 1,474      | 0.170          | 0.002           | Down      |
| D10414_at               | Cell cycle| Cyclin D1                  | 1,363      | 0.017          | 0.010           | Down      |
| X70671_at               | Cell cycle| Cyclin D1                  | 939        | 0.087          | 0.008           | Down      |
| E01184cds_s_at          | Cyt P450  | Cyt P450                   | 4,296      | 0.003          | 0.003           | Down      |
| M72088_at               | Cyt P450  | CYP17                      | 2,204      | 0.004          | 0.013           | Up        |
| K0241cds_at             | Cyt P450  | CYP1A2                     | 2576       | 0.005          | 0.011           | Down      |
| J04187_at               | Cyt P450  | CYP2A2                     | 11,191     | 0.007          | 0.011           | Down      |
| J02657_s_at             | Cyt P450  | CYP2C11                    | 24,686     | 0.001          | 0.001           | Down      |
| M13633cds_at            | Cyt P450  | CYP2C11                    | 7,563      | 0.012          | 0.000           | Down      |
| X79081mRNA_f_at         | Cyt P450  | CYP2C11                    | 3,775      | 0.006          | 0.001           | Down      |
| J03786_g_at             | Cyt P450  | CYP2C12                    | 7,156      | 0.021          | 0.000           | Down      |
| M33505cds_at            | Cyt P450  | CYP2C12                    | 7,758      | 0.093          | 0.000           | Down      |
| rc_AA494573_f_at        | Cyt P450  | CYP2C39                    | 16,174     | 0.009          | 0.015           | Down      |
| M30131mRNA_f_at         | Cyt P450  | CYP2C39                    | 15,025     | 0.012          | 0.000           | Down      |
| M14177_s_at             | Cyt P450  | CYP2C7                     | 17,548     | 0.000          | 0.010           | Down      |
| AB080424_s_at           | Cyt P450  | CYP2D3                     | 15,558     | 0.003          | 0.012           | Down      |
| U46118_g_at             | Cyt P450  | CYP3A49                    | 1,234      | 0.019          | 0.000           | Down      |
| M29853_at               | Cyt P450  | CYP5B1                     | 2,149      | 0.128          | 0.042           | Down      |
| D00660_at               | Glutathione| Glutathione synthetase     | 539        | 0.010          | 0.004           | Down      |
| L38615_g_at             | Glutathione| Glutathione peroxidase     | 1,169      | 0.085          | 0.023           | Down      |
| rc_AA494582_at          | Glutathione| Glutathione synthetase     | 1,049      | 0.019          | 0.010           | Down      |
| S75086_s_at             | Glutathione| Glutathione synthetase     | 2,022      | 0.004          | 0.001           | Down      |
| S52522mRNA_f_at         | Glutathione| Glutathione synthetase     | 7,568      | 0.000          | 0.000           | Down      |
| X98198_g_at             | Lipid metabolism| Acyl-CoA oxidase | 2,963  | 0.003          | 0.012           | Down      |
| AB010428_s_at           | Lipid metabolism| Acyl-CoA thioesterase 1   | 4,424  | 0.011          | 0.002           | Down      |
| Y09333_g_at             | Lipid metabolism| Acyl-CoA thioesterase 1   | 5,696  | 0.024          | 0.000           | Down      |
| Y09333_s_at             | Lipid metabolism| Acyl-CoA thioesterase 1   | 5,411      | 0.032          | 0.000           | Down      |
| D43623_g_at             | Lipid metabolism| Carnitine palmitoyltransferase | 879  | 0.053          | 0.004           | Down      |
| M26125_g_at             | Lipid metabolism| Epoxide hydrolase 1      | 18,841   | 0.023          | 0.000           | Down      |
| rc_AA883242_g_at        | Lipid metabolism| Fatty acid-CoA ligase     | 2,968  | 0.012          | 0.000           | Down      |
| M29429cds_at            | Lipid metabolism| HMG-CoA reductase         | 667      | 0.047          | 0.000           | Down      |
| X55286_g_at             | Lipid metabolism| HMG-CoA reductase         | 235      | 0.010          | 0.000           | Down      |
| J02585_s_at             | Lipid metabolism| Steroyl-CoA desaturase 1  | 3,206  | 0.017          | 0.008           | Down      |
| AB010429_s_at           | Lipid metabolism| Very long chain acyl- CoA thioesterase | 1,772  | 0.015          | 0.004           | Down      |
| L07114_g_at             | Lipid transport| Apolipoprotein B binding protein | 287  | 0.022          | 0.005           | Down      |
| AF07241_at              | Lipid transport| CD36                     | 1,077      | 0.005          | 0.000           | Down      |
| rc_AA935752_at          | Lipid transport| CD36                     | 1,446      | 0.003          | 0.000           | Down      |
| AB005743_g_at           | Lipid transport| CD36                     | 307      | 0.003          | 0.000           | Down      |
| AF072411_g_at           | Lipid transport| CD36                     | 1,979      | 0.004          | 0.000           | Down      |
| AB005743_s_at           | Lipid transport| CD36                     | 289      | 0.004          | 0.000           | Down      |
| K01190_at               | Lipid transport| Fatty acid binding protein 2 | 321  | 0.100          | 0.000           | Down      |
| U02096_s_at             | Lipid transport| Fatty acid binding protein 7 | 3,443  | 0.005          | 0.000           | Down      |
| L34649_g_at             | Lipid transport| Megalin (LRP2)            | 977       | 0.001          | 0.000           | Down      |
| U89280_at               | Phase 2 metabolism| 17-β-Hydroxysteroid dehydrogenase | 6,406  | 0.020          | 0.005           | Down      |
| AF045464_s_at           | Phase 2 metabolism| Alfafoxin b1 dehydrogenase | 9,840      | 0.016          | 0.000           | Down      |
| D30861oxon_s_at         | Phase 2 metabolism| UG77-6                 | 1,026     | 0.003          | 0.012           | Down      |
| S58538_s_at             | Phase 2 metabolism| UG77-6                 | 1,186     | 0.002          | 0.005           | Down      |
| D30861oxon_s_at         | Phase 2 metabolism| UG77-6                 | 507       | 0.003          | 0.010           | Down      |
| J02599rRNAf2 At          | Phase 2 metabolism| UG7172                | 1,299     | 0.009          | 0.000           | Down      |
| rc_N181708_s_at         | Phase 2 metabolism| UG7172                | 21,663    | 0.000          | 0.000           | Down      |
| rc_N180442_s_at         | Phase 2 metabolism| Steroid metabolism      | 1,378      | 0.008          | 0.013           | Down      |

*Continued*
| Affymetrix probe set ID | Class                  | Gene                                      | Max signal  |
|-------------------------|------------------------|-------------------------------------------|-------------|
| M95951_g_at             | Steroid metabolism    | Farensyl diphosphate synthase             | 1,040       |
| M95951_at               | Steroid metabolism    | Farensyl diphosphate synthase             | 2,011       |
| M89945mRNA_g_at         | Steroid metabolism    | Farensyl diphosphate synthase             | 3,421       |
| M89945mRNA_at           | Steroid metabolism    | Farensyl diphosphate synthase             | 4,531       |
| M81225_at               | Steroid metabolism    | Farensyltransferase                      | 1,063       |
| U33500_at               | Steroid metabolism    | Retinol dehydrogenase type II            | 1,256       |
| M19257_at               | Steroid metabolism    | Retinol-binding protein 1                | 3,941       |
| D37920_at               | Steroid metabolism    | Squalene epoxidase                       | 843         |
| U30186_at               | Stress/damage         | GADD153                                  | 1,670       |
| L23591gRNA_at           | Stress/damage         | GADD45a                                  | 1,015       |
| L23591gRNA_g_at         | Stress/damage         | GADD45a                                  | 1,154       |
| rc_A070295_g_at         | Stress/damage         | GADD45a                                  | 829         |
| rc_A179599_at           | Stress/damage         | GADD45a                                  | 1,172       |
| Y00299mRNA_at           | Stress/damage         | GADD45a                                  | 643         |
| Y00299mRNA_g_at         | Stress/damage         | GADD45a                                  | 988         |
| Y00299mRNA_g_at         | Stress/damage         | GADD45a                                  | 988         |
| Y00299mRNA_g_at         | Stress/damage         | GADD45a                                  | 988         |
| J002722c6s_at           | Stress/damage         | GADD45a                                  | 464         |
| M25157mRNA_f_at         | Sulfotransferase      | Superoxide dismutase                     | 4,406       |
| S78511_s_at             | Stress/damage         | BAX                                      | 586         |
| M60291_g_at             | Stress/damage         | B-cell translocation gene 2              | 551         |
| rc_A9A41456_s_at        | Stress/damage         | B-cell translocation gene 2              | 1,934       |
| U49729_at               | Stress/damage         | bcl12-associated X protein               | 285         |
| M33292_f_at             | Sulfotransferase      | Alcohol sulfotransferase                 | 6,794       |
| X63410cds_f_at          | Sulfotransferase      | Alcohol sulfotransferase                 | 9,853       |
| S76498_s_at             | Stress/damage         | Estrogen synthase                        | 10,020      |
| D14988_f_at             | Sulfotransferase      | Hydroxysteroid sulfotransferase          | 5,378       |
| rc_A1169695_f_at        | Sulfotransferase      | Hydroxysteroid sulfotransferase          | 5,836       |
| D14988_f_at             | Sulfotransferase      | Hydroxysteroid sulfotransferase          | 13,597      |
| D14987_f_at             | Sulfotransferase      | Hydroxysteroid sulfotransferase          | 10,861      |
| rc_AA817987_f_at        | Sulfotransferase      | Hydroxysteroid sulfotransferase          | 8,115       |
| M33133mRNA_f_at         | Sulfotransferase      | Hydroxysteroid sulfotransferase          | 14,611      |
| rc_AA818122_f_at        | Sulfotransferase      | Hydroxysteroid sulfotransferase          | 12,159      |
| rc_AA008836_at          | Sulfotransferase      | Hydroxysteroid sulfotransferase          | 602         |
| L23339_at               | Sulfotransferase      | Phenol-prefering sulfotransferase        | 2,105       |
| L23339_g_at             | Sulfotransferase      | Phenol-prefering sulfotransferase        | 15,648      |
| AB010467_s_at           | Transporter           | Myosin-like protein 2                    | 985         |
| DB0088_s_at             | Transporter           | Myosin-like protein 2                    | 4,083       |
| MB1859_at               | Transporter           | P-glycoprotein/multi-drug resistance 1   | 5,082       |
| M77479_at               | Transporter           | Sodium/bis acid con-transporter family   | 6,110       |
| rc_AL125631_at          | Unknown                | Expressed sequence tag                   | 795         |
| rc_AL172452_at          | Unknown                | Expressed sequence tag                   | 1,603       |
| rc_AA866240_f_at        | Unknown                | Expressed sequence tag                   | 8,131       |

**Table 4. Continued.**
metabolic pathway by the high dose was accompanied by downregulation of retinol dehydrogenase and retinol-binding protein 1 in the retinol metabolic pathway and of the androgen/estrogen metabolic pathways (Figure 5). In addition, MP produced a marked effect in some metabolic enzymes such as the upregulation of cytochrome P-450 (CYP)4B1, CYP2C12, and aflatoxin reductase and the downregulation of CYP1A1, CYP2A2, CYP2C11 and sulfotransferases (Ratra et al. 1998a). Additional genes involved in redox processes were affected by the treatment: glutathione S-transferase (GST)Σ-2, glutathione peroxidase, and glutathione synthetase were induced, whereas superoxide dismutase was repressed. A consistent induction of UDP-glucuronosyltransferase (UDPGT) 1–6 and a concomitant downregulation of UDPGT2B (3-hydroxyandrogen specific) were also observed. MP also seemed to have an effect on the expression levels of several transporters; MDR (P-glycoprotein), cMOAT1 (MRP2) and cMOAT2 (MRP3) were upregulated, whereas the expression of the sodium/taurocholate transporter was transcriptionally repressed.

As can be deduced from the cluster and PCA analyses (Figures 1 and 2), the effect of the low dose of MP is rather subtle, involving a small amount of regulated genes and moderate fold changes. This makes the distinction between low-dose treated animals and controls relatively difficult in a rather heterogeneous (different users, different protocols) set of samples comprising very few replicates to support statistical analysis (four biological replicates for individual sample analyses and five replicates for pooled samples). Nevertheless, some genes could be identified that are consistently modulated by the low dose of the compound. Among these genes, a dose-dependent decrease in acyl-CoA desaturase (Brunelle and Chandel 2002) and acl–acyl-CoA acyl-CoA desaturase, acyl-CoA synthase, farnesyl diphosphate FPP-transferase, FPP-synthase were upregulated, while acyl-CoA desaturase, acyl-CoA synthase, squalene epoxidase, farnesyl diphosphate synthase, mevalonate kinase, mevalonate pyrophosphate decarboxylase, 20-ketoacyl-CoA dehydrogenase, isopentenyl-diphosphate delta-isomerase, and farnesyl pyrophosphate synthase were downregulated (Ratra et al 1998b).

Figure 5. Projection of the gene expression results obtained using the individual animal data on the sterol/retinol metabolic pathways (KEGG). Red boxes indicate significant upregulation while blue boxes represent significant downregulation. Of the small colored boxes, boxes on the left represent the mean modulation by Roche (individual) and boxes on the right represent the mean modulation by Schering (individual). Nomenclature for the affected genes: farnesyl diphosphate farnesyltransferase (EC 2.5.1.21); farnesyl pyrophosphate synthase (EC 2.5.1.10); farnesyl pyrophosphate synthase (EC 2.5.1.1); 3-hydroxy-3-methylglutaryl-CoA reductase (EC 1.1.1.34); isopentenyl-diphosphate delta-isomerase (EC 5.5.3.2); mevalonate kinase (EC 2.7.1.38); mevalonate pyrophosphate decarboxylase (EC 4.1.1.33); NAD(P)H2 dehydrogenase, quinone (EC 1.6.99.2); retinol dehydrogenase (EC 1.1.1.108); squalene monoxygenase squalene (EC 1.1.108).

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inhibitor) were observed, together with a very slight decrease in betaine–homocysteine S-methyltransferase (EC 2.1.1.5) and an increase in insulin-like growth factor binding protein 1 precursor (IGFBP-1) (Mohn et al. 1991).

**Discussion**

In this study, we were able to examine differences and similarities of results from microarray analysis obtained from a common source of RNA by several users. A preliminary evaluation of the number and identity of modulated genes uncovered striking differences between the five different Affymetrix users. This is in contrast to previous studies that have shown high reproducibility with microarray chips from the same RNA source (Waring et al. 2001).

In addition, our unpublished data show that the correlation between gene expression in the liver and in the kidney from samples obtained from the same animal and prepared by the same operator following a standardized protocol is only 44%, whereas two different liver samples show a correlation of 98%. In this study some of the differences between users are likely because of different user protocols. In support of this, RO shows the most striking difference in the absolute values of regulated genes and is also the user introducing the most modifications of the sample processing and hybridization protocol (different cDNA kit, different blocking solution, etc.). Other users employing the protocols recommended by Affymetrix show less variability among them. The remaining differences are probably attributable to minor protocol deviations as well as to an expected amount of false positives. In this case the number of false positives was undoubtedly high because the samples were pooled and thus the number of replicates was low.

Further data analysis with additional tools corroborated the finding that differences between users, sample processing, and hybridization protocols affected the absolute results, but that this did not distort the major conclusion of the study. Indeed, the PCA results showed that the PC1 (accounting for 33% of the variance) was mainly site driven (Figure 1A), but this variability did not mask the effects elicited by the high dose of MP. Despite the observed differences, all users obtained similar overall results that correlated with histopathology and clinical chemistry analysis. A clear differentiation between high-dose (toxic)–treated animals and controls and low-dose–treated animals was obtained by all users, as shown using unsupervised data analysis methods (cluster analysis and PCA). Moreover, in a supervised approach it was possible to identify animals treated with high and low doses in all the pooled samples analyzed by five different users on the basis of SVM trained with data obtained from the samples processed individually. Surprisingly, the status of the pooled control samples was not clearly assigned to any treatment group. There are two possible factors that could have led to this misclassification. On one hand, the training set is very small, and this type of model has an optimal performance with large data sets. Alternatively, the effects of 10 mg/kg MP are very subtle, thus making distinction of control and low-dose–treated animals rather difficult. This is even more pronounced in a heterogeneous set of data. For very slight effects a larger number of replicates might be required for optimal performance.

As stated by Hamadeh et al. (2002), the use of unsupervised analysis tools is essential to ensure that the data contain natural subpopulations and that no preconceived bias is introduced when classes of compounds are being identified. The results obtained using cluster analysis and PCA show that gene expression profiles allow the natural classification of the high dose of MP regardless of the variation introduced by the different users. Nevertheless, these unsupervised tools mainly allow the distinction of samples showing definitive histopathological findings (high-dose) from samples without findings (controls and low-dose samples). It might be argued that this is not sensitive enough for predictive toxicogenomics studies. However, using a supervised analysis method like SVM, it was possible to correctly categorize samples into vehicle, low- and high-dose classes, which was not possible with clinical chemistry or histopathology. Thus, similar to results obtained by Burczynski et al. (2000) and Thomas et al. (2001), the ability to correctly classify compounds using toxicogenomics can be greatly improved by selecting a smaller subset of the most predictive gene sets.

A number of the genes and pathways regulated by MP toxicity were similar across users. This is particularly true for the high-dose–treated animals in which the effects were more pronounced. The genes detected as transcriptionally induced or repressed are in good agreement with results from a similarly designed study by Hamadeh et al. (2002) using cDNA spotted arrays. In-depth analysis of the genes modulated by MP sheds light on the variety of cellular processes affected.

Our results provide ample proof that gene expression analysis is a suitable method to detect effects produced by a high dose (100 mg/kg) of MP. The results presented in this article are generally in good agreement with a similar study performed by Hamadeh et al. (2002) and also show signals characteristic of the compound under investigation. The decrease in cytochrome P450 after a high dose of MP was in agreement with results of previous studies that showed this compound decreased the content of CYP2C11, CYP3A and CY2A, possibly due to suicide substrate activation (Graichen et al. 1985, Ratra et al. 1998a). MP is also known to transcriptionally induce CYP2C12 and CYP4B1 (Hamadeh et al. 2002), as was also detected by all users analyzing the samples. In addition, several of the genes detected as modulated, including the GADD family, hem oxygenase and genes related to glutathione homeostasis are indicative of the oxidative stress known to be produced by MP (Ratra et al. 1998b). Also, several of the modulated genes indicate an effect of MP on lipid metabolism, which is one of the pathways affected by MP as shown in studies using *in vitro* approaches (Iype et al. 1985) and protein analysis (Man et al. 2002). Moreover, events indicative of lipid peroxidation were observed as previously published (Hamadeh et al. 2002). The induction of mitochondrial genes (i.e., *CPT1* and acyl-CoA thioester hydrase) is also indicative of the mitochondrial proliferation that has been previously related to MP (Iype et al. 1985).

In animals treated with a low dose of MP, some genes could be identified as already being modulated after 1-week treatment with 10 mg/kg/day MP. Among these, an induction of *IGFBP-1* (Affymetrix probe set ID M58634_at) and aflatoxin B1 aldehyde reductase (Affymetrix ID AF045464_s_at), as well as the downregulation of retinol dehydrogenase type 2 (Affymetrix ID U33500_g_at) were observed. As depicted in Figure 4C, the observed downregulation of retinol dehydrogenase 2 at both the low and high doses was accompanied by the downregulation of retinol-binding protein 1 (Affymetrix ID M19257_at; Figure 4D). *IGFBP-1* (Affymetrix ID M58634_at) appeared upregulated at both doses, whereas the growth-promoting insulin-like growth factor 1 (*IGF1*, Affymetrix ID M15481_at) appeared downregulated only after rats were exposed to a high dose of MP (Figure 4A, B). The upregulation of *IGFBP-1* might be a protective mechanism for the known carcinogenic effect of MP because the levels of *IGFBP-1* regulate the mitogenic effects of IGFs (Kelley et al. 1996). In fact, *IGFBP-1* has been shown to inhibit hepatic preneoplasia in mice (Lu and Archer 2003).
An additional cell protection mechanism that appears stimulated after treatment with 10 mg/kg/day of MP is aflatoxin B1 aldehyde reductase (Affymetrix ID AF045464_s_at; Figure 4E). This detoxifying enzyme shows only a slight induction at the low dose and an extensive induction at the high dose, which is in agreement with its previously reported induction by a high dose (100 mg/kg/day) of MP (Hamadeh et al. 2002). 17β-Hydroxysteroid dehydrogenase type 2 (17βHSD2; Affymetrix ID X91234_at) shows an interesting regulation pattern, as it appears upregulated by the low dose of MP and downregulated by the high dose (Figure 4F). This enzyme is involved in the steroid conversion pathway (Akinola et al. 1996), which is one of the pathways affected by the treatment with MP (Figure 5), but the biological meaning of this finding remains unclear.

In conclusion, a high degree of user/site variability was observed with microarray analysis using the same RNA processed at different sites. Despite this, all the microarray results showed that it was nonetheless possible to distinguish toxic (i.e., histopathological findings) versus nontoxic dose levels of MP. Moreover, regardless of the user, gene expression analysis using supervised data analysis tools allowed the correct identification of the samples treated with the low dose of MP, a distinction that was not apparent from clinical chemistry or histopathology analysis.

The observed site-to-site variability did not impair the detection of molecular effects elicited by MP. In addition, crucial gene expression changes, which most likely reflect the mechanism of toxicity for MP, were observed across all user groups. These results provide critical information regarding the consistency of microarray results across different laboratories and shed light on the strengths and limitations of expression profiling in drug safety analysis.

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