Modified Ames test using a strain expressing human sulfotransferase 1C2 to assess the mutagenicity of methyleugenol

Hiroshi Honda1*, Kazuyuki Minegawa2, Yurika Fujita1, Noriko Yamaguchi2, Yoshihiro Oguma2, Hansruedi Glatt3, Naohiro Nishiyama1 and Toshio Kasamatsu1

Abstract

Introduction: Several alkenylbenzenes, including methyleugenol (ME), are present in a wide range of botanicals and exhibit carcinogenic and mutagenic properties. Negative results are generally obtained for alkenylbenzenes in standard in vitro genotoxicity tests, including the Ames test. A lack of mutagenicity observed in such tests is thought to result from impaired metabolic activation of alkenylbenzenes via hydroxylation, with subsequent sulfocoujugation to its ultimate mutagenic or carcinogenic form. Although recent studies have reported the mutagenicity of hydroxylated ME metabolites in the Ames test using modified TA100 strains expressing human sulfotransferases (SULTs), to our knowledge, the detection of ME mutagenicity has not yet been reported.

Findings: Using strain TA100-hSULT1C2, which expresses human SULT1C2, we optimized the protein content of S9 Mix and the pre-incubation time required to promote metabolic activation in the Ames test. This procedure enabled us to obtain a positive response with ME.

Conclusions: We established Ames-test conditions enabling the detection of ME-induced mutagenicity, using a strain expressing human SULT1C2 in the presence of induced-rat S9 Mix. This simple approach will help assess the mutagenicity of other alkenylbenzenes and related chemicals.

Keywords: Ames test, Methyleugenol, Alkenylbenzene, Sulfotransferase, S9

Findings

Introduction

Alkenylbenzenes are present in a wide range of botanicals, including basil, nutmeg, and fennel, which are used in herbal teas, food flavorings, and food supplements. However, several alkenylbenzenes such as methyleugenol (ME), estragole, safrole, and β-asarone are known to exhibit mutagenicity [1] and hepatocarcinogenicity in rodents [2–4]. Findings from mechanistic studies indicate that alkenylbenzene-induced liver tumors result from the metabolism of these compounds to DNA-reactive intermediates. For example, ME is metabolized to 1'-hydroxymethyleugenol (1'-HME) by cytochrome P450 (CYP). Subsequent sulfoconjugation of 1'-HME by sulfotransferases (SULTs) leads to the production of highly reactive electrophiles that can form DNA adducts and thereby induce mutations (Fig. 1) [5, 6].

Human health-related concerns regarding the use of botanicals or botanical ingredients containing alkenylbenzenes have been raised; therefore, hazard assessments of alkenylbenzenes are of high priority [1]. However, the mutagenic properties of alkenylbenzenes are difficult to measure with standard in vitro genotoxicity assays, such as the bacterial reverse-mutation assay (Ames test) [7], due to a lack of specific enzyme activities in the exogenous metabolic system (rat S9 Mix) that is generally used in these assays. Although several studies have shown that alkenylbenzenes are mutagenic [8–12], the experimental systems used require considerable effort or expertise when compared to standard genotoxicity assays (e.g., the Ames test). Therefore, a simple approach for utilizing standard
genotoxicity assays, or a minor modification of such assays, is necessary.

Recently, Herrmann et al. demonstrated that hydroxylated metabolites of ME, including 1′-HME, are mutagenic in the Ames test when using Salmonella typhimurium TA100 strains expressing human SULTs [13]. However, detection of mutagenicity using the parent compound, ME, has not yet been reported in the Ames test. To assess the mutagenicity of alkenylbenzenes, we sought to establish Ames test conditions enabling the detection of mutagenicity of ME. As described above, ME is bioactivated in two steps, involving hydroxylation by CYP and sulfoconjugation by SULT. Employing the S. typhimurium strain TA100-hSULT1C2, which expresses human SULT1C2, we optimized the protein content of the S9 Mix and the pre-incubation time for co-incubating bacteria with test chemicals and the S9 Mix before the main incubation step, which promoted the generation of oxidized ME metabolites by CYP.

### Materials and methods

#### Test strain
We used strain S. typhimurium TA100-SULT1C2, established by transforming the S. typhimurium TA100 strain with an expression vector, pKKneo-SULT, for the human SULT1C2 gene [EMBL/DDBJ: AF186263] [14]. pKKneo-SULT was developed from cloning of human SULT cDNA into pKKneo (pKK233-2 [Stratagene] with the ampicillin-resistance marker replaced by a neomycin resistance marker) [14].

Several known mutagens exhibited strong mutagenic activity in TA100-hSULT1C2 but were inactive in the parental strain TA100 [13–16]. Among these mutagens, sulfoconjugation in furfuryl alcohol and 5-hydroxymethylfurfural has been demonstrated, and enzyme kinetic data have been established in cytosolic preparations from hSULT1C2 expressed in S. typhimurium [17].

#### Chemicals/reagents
JIS special grade dimethyl sulfoxide (negative-control solvent), 2-(2-furyl)-3-(5-nitro-2-furyl)-acrylamide (AF-2; 99.6 % purity), and benzo[a]pyrene (B[a]P; 99.7 % purity) were used as positive-control reagents. These three reagents, as well as magnesium sulfate (>98 % purity), were obtained from Wako Pure Chemical Industries, Osaka Ltd. (Japan). Nutrient Broth No. 2 Culture Medium was purchased from OXOID Ltd. Hampshire (UK). Minimum-glucose medium was obtained from Kyokuto Pharmaceutical Industrial Co., Ltd. Tokyo (Japan). The protein contents were adjusted by changing the amount of the S9 fraction, prepared from 5,6-benzoflavone and phenobarbital-induced Sprague–Dawley rats, and water (total volume: 5 mL), and the mixtures were added to 10-mL vials of Cofactor-I (MgCl$_2$: 16.3 mg; KCl: 24.6 mg; glucose-6-phosphate: 17.1 mg; Na$_2$HPO$_4$: 119.6 mg; NaH$_2$PO$_4$/2H$_2$O: 24.7 mg) (also see the Optimization of the S9 Mix protein content section).

#### Ames test
We modified the pre-incubation method [18], as described previously [13]. Briefly, 30 mM magnesium sulfate was used instead of 0.1 mM sodium phosphate buffer (pH 7.4), and optimization of metabolic activation conditions was explored in terms of the protein content in the S9 Mix and pre-incubation times. Inorganic sulfate is required for the bacteria to synthesize 3′-phosphoadenosine-5′-phosphosulfate, the cofactor for sulfo transferases. Magnesium ions are activators of some enzymes involved in the synthesis of 3′-phosphoadenosine-5′-phosphosulfate. To grow bacteria, 10 mL of Nutrient Broth No. 2 Culture Medium was placed into a sterilized, 40-mL capacity L-shape test tube, to which 10 μL of a bacterial suspension was added. Bacterial cultures were incubated at 37 °C with shaking at 100 rotations/min for 9 h before starting the experiment.
Ten microliters of test formulation, solvent, or positive-control reagent (AF-2 or B[a]P) solution was added to sterilized tubes. Next, 0.5 mL of 30 mM magnesium sulfate was added to the tubes for experiments not involving metabolic activation, or 0.25 mL of 60 mM magnesium sulfate and 0.25 mL of S9 Mix were added to promote metabolic activation. Subsequently, 0.1 mL of bacterial culture was added to each tube. Immediately after stirring, the tubes were pre-incubated for the indicated durations at 37 °C, with shaking at 80 rotations/min. Top agar (2.0 mL; 45 °C) was added and the resulting homogeneous mixtures were overlaid on agar plates containing Minimum-Glucose Medium. After the top agar solidified, the minimum-glucose agar plates were inverted in an incubator and cultured for 48 h at 37 °C. After incubation, the colonies were enumerated using an automatic colony counter (Colony Analyzer CA-11D systems, System Science Co., Ltd.). The bacteria were inspected under a stereomicroscope for the presence/absence of bacterial growth inhibition.

Optimization of the S9 Mix protein content
Length of the pre-incubation time in this experiment was 1 h. For preliminary examinations, the S9 Mix was prepared with a total protein content of 4.0, 1.2 (common content for Ames tests), 0.3, or 0.1 mg protein/plate. Based on the results of the preliminary examinations, the main experiments were conducted using S9 Mix containing 2.4, 1.2, 0.6, or 0.3 mg of protein/plate.

Optimization of the pre-incubation time
The concentration of the S9 fraction was set at 10 % (total protein content: 1.2 mg/plate), as normally used for Ames tests. The pre-incubation time was varied at 0 min, 20 min (ordinary time for the liquid pre-incubation version of the Ames tests), 1 h, and 2 h.

Evaluation of results
We evaluated whether the number of revertant colonies in the presence of ME were increased by at least twice above that of the solvent control in a dose-dependent manner, which is a commonly used criterion for a positive response.

Results and discussion
Positive controls
To confirm that the experimental conditions other than the activity of hSULT1C2 work properly in the TA100-hSULT1C2 strain, we tested AF-2 without metabolic activation and B[a]P with metabolic activation under the same conditions as ME. As a result, both AF-2 and B[a]P yielded clear positive responses, indicating that the experimental conditions were valid (data not shown).

Optimization of S9 Mix protein content
In preliminary experiments, slight increases in the number of revertant colonies were observed in ME treatment groups when the S9 Mix protein content was above 0.3 mg/plate (data not shown). Next, we prepared S9 Mix for subsequent experiments such that the protein content ranged from 0.3 to 2.4 mg/plate. The number of revertant colonies increased similarly regardless of protein content, but these increases were not sufficient to be judged as positive responses (2-fold increase over the solvent control; Fig. 2). Thus, our results suggested that the protein content offers diminishing returns above a threshold level.

Optimization of the pre-incubation time
Effects of varying pre-incubation time on the mutagenicity of ME are shown in Fig. 3. When the pre-incubation time was 2 h, the number of revertant colonies increased in a time-dependent manner to at least twice that of the negative-control group. Then, we conducted an Ames test for ME under the optimized condition (protein content: 1.2 mg/plate, pre-incubation time: 2 h), and a positive response was confirmed under the metabolic-activation condition (Fig. 4). Although the response was just above a 2-fold increase compared to the negative control group, the reproducibility of the positive response was confirmed (Figs. 3 and 4).

![Fig. 2 Optimizing the protein content in the S9 Mix. Based on the results of the preliminary investigation, the total protein contents in the S9 Mix were set at 0, 0.3, 0.6, 1.2, or 2.4 mg/plate. The pre-incubation time was set at 1 h. The plots indicate the ratio of the average number of revertant colonies observed (treated: solvent control) in two plates and the number of spontaneous revertants. The number of spontaneous colonies per plate ranged from 108 to 151, varying slightly in response to the different protein levels used](image-url)
Implication for evaluating the mutagenicity of alkenylbenzenes
To our knowledge, this is the first report describing the detection of the ME mutagenicity using the Ames test. Because 1'-HME mutagenicity has been detected previously using the TA100-hSULT1C2 strain [13], prolonging the pre-incubation time may be effective in enhancing the production of mutagenic metabolites from ME by CYP and SULT. It should be noted that the approach used in the present study is primarily suited for hazard assessment, rather than risk assessment. Human SULT1C2, expressed in the TA100-hSULT1C2 cells used in this study, is a fetal enzyme that is rarely expressed in adult tissues [19]. Therefore, using the TA100 strain expressing human SULT1A1, which is abundantly expressed in many tissues in adult humans and shows considerable 1’-HME-sulfoconjugation activity [13], would be more relevant to human risk assessment. Indeed, genetic manipulation of the SULT1A status (knockout of the endogenous SULT1A gene or overexpression for the human SULT1A1/2 gene cluster) drastically affected the formation of hepatic DNA adducts in mice treated with ME or 1’-HME [20].

Furthermore, alkenylbenzene hydroxylation may vary with CYP expression levels or isoforms. Gardner et al. reported that the highest enzyme activities in human subjects were equivalent to that observed in untreated rats [21]. Cartus et al. found somewhat higher activities in human liver microsomes than in rat liver microsomes, in particular at low ME concentrations [22]. As we employed induced rat S9 Mix obtained from a rat administered 5,6-benzoflavone and phenobarbital, our system could potentially be optimized to promote ME hydroxylation and would be a very conservative approach in view of risk assessment. Safrole, another known mutagenic alkenylbenzene, is primarily 1’-hydroxylated by CYP2A6. CYP1A2 shows the highest activity in hydroxylating ME among the CYP enzymes [23]. Thus, careful attention is required when selecting a metabolic-activation system with an appropriate origin, taking into consideration structure-metabolism relationships. Nevertheless, the optimization of metabolic-activation conditions should be universally applicable.

Conclusion
We established Ames test conditions that enable detection of a positive mutagenic response with ME. This simple approach will help assess the mutagenicity of alkenylbenzenes or other related chemicals.

Abbreviations
1’-HME: 1’-hydroxymethyleugenol; AF-2: 2-(2-furyl)-3-(5-nitro-2-furyl)acrylamide; B[a]P: benzo[a]pyrene; CYP: cytochrome P450; ME: methyleugenol; SULT: sulfotransferase.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
KM, NY, and YO carried out the experiments. HH, KM, YF, TK, and NN drafted the manuscript. HH, YF, TK, and KM participated in the analysis and interpretation of the data. HH, TK, and HG participated in the design of the study, and HH and TK participated in its coordination. All authors read and approved the final manuscript.
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Author details
1R&D Safety Science Research, Kao Corporation, 2606 Akabane, Ichikai-Machi, Haga-Gun, Tochigi 321-3497, Japan. 2Tokyo Laboratory, BoZo Research Center Inc., 1-3-11, Hanegi, Setagaya-Ku, Tokyo 156-0042, Japan. 3Department of Nutritional Toxicology, German Institute of Human Nutrition (DIfE) Potsdam-Rehbruecke, Arthur-Scheunert-Allee 114-116, D-14558 Nuthetal, Germany.

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