



## TABLE OF CONTENTS

Salmonella Mutagenicity Assay Kit Components .....	3
Kit Components and Additional Items Needed to Complete Assay .....	3
Introduction .....	4
The Assay .....	5
A. Experiment Planning and Set Up .....	6
B. Cell Culture .....	8
C. Treatment and Plating .....	10
D. The Results - Evaluation and Interpretation.....	15

MOLTOX®

## Salmonella Mutagenicity Assay Kit Components

Part Number	Description	Quantity
11-05L.2	Lyophilized PB/BNF-induced S9, 2 ml/vial	2 each
21-100	Nutrient Agar Plates, 20/sleeve	1 each
21-200	ST Quad PC™ Plates, 5/sleeve	1 each
21-400.2	Minimal Glucose Agar (MGA) Plates, 20/sleeve	9 each
26-503.1	Top Agar, Histidine/Biotin, 0.05mM, 100 ml/bottle	1 each
26-505.1	Oxid Nutrient Broth #2, 100 ml/bottle	4 each
60-101	ICR 191, 10 µg/vial, 5 vials/pack	1 each
60-102	Daunomycin, 60 µg/vial, 5 vials/pack	1 each
60-103	Sodium Azide, 15 µg/vial, 5 vials/pack	1 each
60-107	2-Aminoanthracene, 100 µg/vial, 5 vials/pack	1 each
60-200.4	Regensys™ "A", 40 mL (final S9 mix volume)	1 each
60-201.4L	Regensys™ "B", 123 mg/vial	1 each
71-098.5L	<i>S. typhimurium</i> TA98, 5 discs/vial	1 each
71-100.5L	<i>S. typhimurium</i> TA100, 5 discs/vial	1 each
71-1535.5L	<i>S. typhimurium</i> TA1535, 5 discs/vial	1 each
71-1537.5L	<i>S. typhimurium</i> TA1537, 5 discs/vial	1 each

### Additional items you will need to complete the assay:

Erlenmeyer flasks , ≥ 125 ml, vented, sterile (4)	Test tubes, 13 x 100 mm, sterile
Sharpie or wax pencil	Test tube rack
Micropipettes, 100 µL - 500 µL	Ice
Micropipette tips, 100 µL - 1000 µL, sterile	Ice bucket
Pipet-aid or rubber bulb	Nitrile gloves
Pipets, 1 mL - 10 mL, sterile	Incubator with shaker, 37°C
Inoculating loops, 10 µL, sterile	Vortex mixer
Microwave or boiling water bath	Dimethyl Sulfoxide (DMSO)
Water bath and/or heating block, 45°C	Swabs, sterile (optional)
Automatic colony counter or magnifying counter	Forceps, small, sterile (optional)
Water, deionized or distilled, sterile, ice-cold	Spectrophotometer capable of 650 nm
Test tubes, 15 ml, screw cap, sterile (12)	Microscope, capable of 40x

## INTRODUCTION

The materials contained in this Salmonella Mutagenicity Assay Kit include virtually all of the supplies necessary for the conduct of the “Ames Assay” as described by Maron & Ames (Maron, D. M. and B. N. Ames, Revised methods for the Salmonella mutagenicity test, *Mutation Research*, 113: 173 - 215, 1983) and Mortelmans & Zeiger (Mortelmans, K. and E. Zeiger, The Ames Salmonella/microsome mutagenicity assay, *Mutation Research*, 455: 29 - 60, 2000). This manual is a brief summary of the important aspects of testing - **We strongly recommend that you carefully read one of these papers and OECD guideline 471 before you attempt to perform the assay.**

All elements of the kit MOLTOX<sup>®</sup> are formulated and manufactured using the highest quality components and are consistent with the recommendations of Maron & Ames and Mortelmans & Zeiger (ibid.). Certain materials supplied (e.g., STDiscs<sup>™</sup> and lyophilized S9) have been specifically developed for inclusion in the assay kit by our laboratory. All of the materials contained in the kit are accompanied by GLP level Certificate of Analysis (CoA) Statements - you may be assured that each element contained within the kit has been thoroughly tested for performance in the assay.

The MOLTOX<sup>®</sup> Salmonella Mutagenicity Assay Kit is intended for use by individuals skilled in the science and art of microbiology; the use of strict aseptic technique is essential for the successful application of the materials included in the kit. While the bacterial strains included in the kit (*S. typhimurium* strains TA1535, TA1537, TA98, and TA100) are attenuated, they are potentially pathogenic and must be handled accordingly. If you have any doubts about the safe handling of the strains included in this kit do not proceed until you have consulted with us at (828) 264-9099 or have obtained the advice of a skilled biochemist or microbiologist.

The performance of the “Ames Assay” includes several distinct experimental steps; e.g., test design; S9 mix formulation; dosing and plating; phenotype confirmation; target cell titer determination; reading (counting) and analysis. The materials contained in the kit were selected so as to provide the user with considerable flexibility as regards test design. This manual was developed to assist in the utilization of the kit contents; the information provided in the manual is intended to supplement that contained in the Maron & Ames and Mortelmans & Zeiger papers. While presented in a step-by-step manner, the instructions contained in this manual are amenable to modification - if you desire assistance in any phase of the assay, please contact our Technical Service department at [TechServices@moltox.com](mailto:TechServices@moltox.com) or call (828) 264-9099 (toll-free: 800-536-7232). We will be happy to work with you to help solve any problems that may arise.

## THE ASSAY

The MOLTOX<sup>®</sup> Salmonella Mutagenicity Test kit contains four tester strains: TA1535, TA1537, TA98, and TA100. Each strain was constructed with a different lesion in the histidine operon (see Mortelmans & Zeiger, Table 1, pg 33). This mutation renders them incapable of synthesizing histidine (i.e. they are histidine auxotrophs requiring exogenous histidine). In addition, TA1535, TA1537, TA98, and TA100 have altered cell walls (*rfa*) that increase the cell's permeability to certain high molecular weight materials. These strains also share a lesion in a DNA repair-coding gene (*uvrB*) which results in increased sensitivity to a variety of mutagens. Since this lesion extends through the gene for biotin synthesis (*bio*), biotin is also required for growth]. Tester strains TA98 and TA100 carry a plasmid (pKM101) which acts to increase the activity of an error-prone DNA repair system and to confer resistance to the antibiotic ampicillin. Tester strains TA1535 and TA1537 contain no plasmids.

Because of the characteristics of the tester strains, the "Ames Assay" is uniquely suited for the detection of mutagenic activities. The several tester strains differ in their response to DNA-damaging chemical mutagens and therefore are generally employed in combination. Each strain tends to be responsive to specific classes of mutagenic chemicals due to the specific lesion in their histidine operon. However, exposure to mutagens may result in genetic reversions in the histidine operon resulting in restoration of the wild type phenotype; mutants have their histidine operons functionally "restored" and can synthesize histidine. The assay depends on the ability to distinguish between histidine auxotrophs (the tester strains) and histidine prototrophs (the mutants). Accordingly, the target cells are plated on media containing trace quantities of histidine that allows for a few rounds of cell division necessary to "fix" a mutation event; the histidine is rapidly exhausted resulting in cessation of the growth of non-mutated cells. If a mutagenic chemical is present, (comparatively) rare reversions may occur in the altered histidine operon resulting in the continuation of growth after trace histidine exhaustion. Revertant bacterial colonies that appear on histidine-limiting media plates represent prototrophs that arose either spontaneously or due to the action of a mutagen.

The sections that follow describe the procedures for the conduct of a four strain assay. For the most part, the methods described are taken directly from Maron & Ames (*ibid.*) and, subsequently, Mortelmans & Zeiger (*ibid.*). If you have no prior experience with the assay, we suggest that you follow these instructions closely; those experienced in the method will find useful information about the use of STDiscs<sup>™</sup>, QUAD PC<sup>™</sup> plates, and lyophilized S9.

## A. Experiment Planning and Set Up

1. Before setting up the assay, you should gather as much test material information as possible:
  - a. Using the available references, structural analyses or activity data, assemble as much information as possible concerning your test material or its analogues or closely related congeners. Of particular importance are questions of bacteriostatic or bacteriocidal activities, hazardous qualities and stability.
  - b. Determine the solubility of your test chemical in the appropriate solvent (water and DMSO are preferred solvents - see p. 200, Maron & Ames for listing or check the Merck Index). In many cases, you may find it necessary to dose with suspensions rather than true solutions. If an organic solvent is used (e.g., DMSO, acetone), the test material may precipitate upon addition to the aqueous top agar; e.g., 2-aminoanthracene is essentially insoluble in water and is solubilized in DMSO for dosing.
  - c. Decide on the test doses. The conventional test solution dose volume is 100  $\mu\text{L}$  per plate, but this can be increased for aqueous formulations or reduced, e.g., if a somewhat toxic organic solvent is used. The upper dose should not exceed 5 mg/plate (50 mg/mL assuming a 100  $\mu\text{L}$ /plate dosing volume). Select 5 to 7 doses separated by factors of 2, 3 (or half logs) or 5. There are 160 minimal glucose agar plates in your kit. This is sufficient for a 5 dose triplicate plate assay (including controls) conducted with and without S9.
2. To avoid difficulties on the day of the assay, design your experiment carefully and well in advance. Examples of questions you need to resolve are:
  - a. Dosimetry: Top dose? Dose intervals? Number of doses? Solvent? Volume of each dosing solution dilution required for the complete assay?
  - b. Replicates: Are you going to dose duplicate plates per condition? Triplicates?
  - c. Metabolic activation: Are you going to perform the assay with and without S9 mix?
  - d. Strains: Which strains are you going to use? Depending on your objective, you may wish to use only TA98 and TA100; e.g., if your test material is a complex mixture, a two strain assay may be desirable.

3. Assemble the supplies and equipment needed to perform the test on the day before.
  - a. Remove the QUAD PC™ plates from the refrigerator, cut off the plastic sleeve and allow to dry upright at room temperature overnight.
  - b. Label the Minimal Glucose Agar plates appropriately: Strain number(s), test material identification and dose, S9 (+/-) and date of test should be included. Writing using a wax pencil or “Sharpie” should be restricted to the dish top or side of the plate - never write on the bottom as such will interfere with scoring. Be sure to include the diagnostic positive and negative controls.
  - c. Label the Nutrient Agar plates in duplicate for the cell titer assessment: Strain number(s), final dilution factor ( $E^{-5}$  and  $E^{-7}$ ), and date of test.
  - d. Adjust the temperature of your water bath and/or dry block heater to approximately 45°C. Make sure that your incubator is adjusted to 37°C. You will need a microwave oven or a boiling water bath to melt the top agar.

## B. Cell Culture - Use of STDiscs™

Your kit is supplied with lyophilized bacterial strains in disc format. Each disc contains sufficient viable cells to serve as the inoculum for a 20 - 25 mL culture. STDiscs™ are accompanied by a CoA that describes their phenotype. The kit includes materials to confirm the phenotypes of your cultures and you may find it useful to compare your results with those described on the aforementioned CoA sheets and with the strain descriptions in the Maron & Ames (ibid.) and Mortelmans & Zeiger papers (ibid.). **Use aseptic technique when working with the strains.**

To prepare strain cultures:

### ~ THE EVENING BEFORE THE ASSAY ~

1. Remove the STDisc™ vials from the refrigerator and allow to warm to room temperature. This assists in reducing the formation of condensation within the vial upon opening. Such moisture is damaging to the lyophilized product.
2. Meanwhile, label sterile Erlenmeyer flasks with the strain number (e.g., TA1535, TA1537, TA98, and TA100) in accordance with your experimental design.
3. Using aseptic technique, carefully pipet approximately 20 - 25 mL of Oxoid #2 nutrient broth into each Erlenmeyer flask. Reserve approx. 5 mls of broth to use as a blank when measuring density.
4. Unscrew the vial closure and remove the slotted gray butyl rubber stopper from one vial (this is best accomplished by use of forceps). Do not contaminate the inner surfaces of the stopper (e.g., place the stopper in the screw cap closure or the base of a sterile, empty petri dish).
5. Using a sterile inoculating loop or small forceps, pick up one disc and drop it into the appropriately labeled flask containing nutrient broth. Aseptically replace the stopper on the STDisc™ vial and screw cap on tightly. Repeat Steps 4 and 5, using a fresh loop or new/re-sterilized small forceps, for each strain.
6. After the flasks are inoculated, transfer to a 37°C incubator and hold *stationary* overnight.

7. Early the next morning, check the density of each culture at 650nm using remainder of broth as the blank. As required, incubate with shaking [125 - 150 rpm (15mm - 25mm orbital range) - avoid foaming] at 37°C until a density of  $1 - 2 \times 10^9$  bacteria/mL is achieved (approximately 1.0 - 1.4, OD 650nm). The duration time of additional incubation/shaking is dependent on the initial density reading. These strains can double approximately every 30 minutes.
8. As the cultures achieve the appropriate density, place them in the refrigerator until shortly before the assay. Remove them approximately 1/2 hour before use to allow them to come to room temperature to avoid heat shocking the cells.

MOLTOX<sup>®</sup>

~ DAY OF THE ASSAY ~

**\*Use aseptic technique throughout this section\***

### C. Treatments and Plating

1. Melt the histidine/biotin supplemented top agar in a boiling water bath or microwave oven. Be sure to loosen the container caps - failure to do so may result in a violent explosion due to pressure build-up. Examine the melted agar carefully - if any opalescence persists, continue heating until a perfectly clear solution is obtained. After melting, place the top agar bottles into a 45°C water bath - allow at least 45 minutes for temperature equilibration.
2. As with any enzyme assay, all materials should be placed on ice prior to use and kept on ice throughout the assay. If you are using the S9 activation system, remove the tear-off seal from one or both lyophilized S9 vials. Rehydrate each vial with 2.1 mL ice-cold sterile water and mix to homogeneity. Your Regensys™ system can be used at 5% or 10% S9 concentration. For 5%, add 2.0 mL rehydrated S9 and 2.0 mL ice-cold, sterile water to the Regensys™ “A” bottle. For 10% S9, add a total of 4 mL rehydrated S9. Keep on ice. Just before use, add the contents of the Regensys™ “B” tube (NADP), mix thoroughly, and hold on ice.
3. Open the CONTROLCHEM™ packages: Nitrile gloves **must** be worn when handling these chemicals. Latex gloves **do not** provide adequate protection for chemicals dissolved in organic solvents. Remove gloves carefully to avoid skin contamination. Wash hands after use.

Add 1.0 mL of the appropriate solvent to each of the CONTROLCHEM™ tubes.

Mutagen	Amount (µg)	Strain	Solvent
Sodium Azide	15	TA1535, TA100	Water
ICR 191 Acridine	10	TA1537	Water
Daunomycin	60	TA98	Water
2-Aminoanthracene (+S9 activation control)	100	All	DMSO

4. Perform the dilutions of your test material. Remember that you will be dosing using 100 µl volumes. Therefore, your dosing solutions should be made up at 10x the desired dose. Arrange the test material dilutions so that they follow a logical sequence - e.g., solvent control, low dose to high dose.

5. Load a test tube rack with sterile 13 x 100 mm tubes with closures equal to the number of minimal glucose agar plates labeled in step A 3b. Place rack in 45°C water bath or heating block and pipet 2 mL of molten, 45°C, top agar into each tube. Remove and replace closures carefully so as to avoid contamination.
6. Arrange your previously labeled Minimal Glucose Agar plates by strain and condition (e.g., controls, +/- S9, etc.).
7. Decide which strain you are going to begin with. In the example below it is assumed that TA98 will be used first in a **duplicate** plate, + and - S9 assay. If performing a triplicate plate assay, adjust accordingly.
8. Assay your Test material:

**- WITHOUT S9 -**

- a. Add the test material doses to the tubes containing top agar. Begin with the solvent control; add 100 µL of water or DMSO (or other solvent used to solubilize your test material) to the first two tubes. Then, in ascending sequence, add 100 µL of each test material dilution to each additional pair of top agar-containing tubes.
- b. Add 100 µL of the TA98 culture to the first two tubes (solvent control tubes).
- c. Without delay, gently mix the tube contents using a vortex mixer and decant the mixture onto the surface of the appropriately labeled Minimal Glucose Agar plate. Do one tube at a time. Immediately upon decantation, gently tilt the plate and rotate so as to obtain an even distribution of the plating mixture over the surface of the bottom agar. Place onto a perfectly level surface, re-cover plate and allow to harden.
- d. Repeat steps 8b. and 8c. for each dose of the test material.

**- WITH S9 -**

- e. Repeat step 8a. using an additional set of top agar-containing tubes.
- f. Add 500 µL of the previously prepared S9 mix to the first two tubes (solvent control tubes).
- g. Without delay, add 100 µL of the TA98 culture as described in step 8b (above).
- h. Immediately mix the tube contents as before, decant onto the Minimal Glucose Agar plate, re-cover and set aside to harden. Repeat steps 8f. and 8g. for each of the test material doses.

**- REPEAT THE ABOVE PROCEDURES FOR EACH STRAIN -**

9. Treat the Positive Control Cultures:

- a. Set up 4 tubes for each strain. 2 tubes will be used for the - S9 diagnostic control and 2 will be used for the + S9 positive control.
- b. Add 2 mL of molten agar to each tube as before. Add 100  $\mu$ L of the CONTROLCHEM™ solutions according to the following scheme:

Mutagen	Strain	Dose/Plate
Sodium Azide	TA1535, TA100	1.5 $\mu$ g
Daunomycin	TA1537	6.0 $\mu$ g
ICR 191 Acridine	TA98	1.0 $\mu$ g
2-Aminoanthracene (+S9 activation control)	All (+S9)	10.0 $\mu$ g

- c. Following the methods described in Section 8, add 100  $\mu$ L of the appropriate strain and decant, spread and set aside. For 2-AA, add 500  $\mu$ L S9 mix and strains as previously described.

10. Inoculate the QUAD PC™ Plates:

- a. Using a sterile loop or swab, wet with the appropriate culture (squeeze excess liquid from swab against side of flask) and inoculate each of the four sectors of a QUAD PC™ plate using a “Z” inoculation pattern.
- b. Repeat for each strain. After all plates are inoculated, open the vial containing the crystal violet discs and, using forceps or an inoculating loop, place a single disc on the agar surface in Sector II of each of the Quad PC™ plates.

11. Determine the Titer of the Strain Cultures:

- a. Arrange the 15 ml sterile tubes with closures in sets of 3 for each strain. Pipet 9.9 mL of sterile dH<sub>2</sub>O water into each tube.
- b. Using your micropipette, inoculate the first tube with 100  $\mu$ L of the appropriate strain culture. Cap tightly and mix thoroughly using a vortex mixer at low speed or by repeated gentle inversion. This tube contains a 1:100 (10E<sup>-2</sup>) dilution of the sampled culture.

- c. Add 100  $\mu\text{L}$  of the 1:100 dilution to the second tube containing 9.9 mL sterile  $\text{dH}_2\text{O}$  water and mix as before. The second dilution is 1:10,000 ( $10\text{E}^{-4}$ ).
- d. Complete the serial dilution by adding 100  $\mu\text{L}$  of the 1:10,000 dilution to the third 9.9 mL tube and mix as before. The final dilution is 1:1,000,000 ( $10\text{E}^{-6}$ ).
- e. Arrange sets of 2 sterile tubes with closures for each strain and place in 45°C water bath or heat block. Add 2.5 or 3 mL of molten top agar to each tube.
- f. Using a micropipette, inoculate the top agar-containing tubes with 100 $\mu\text{L}$  of the 1:10,000 ( $10\text{E}^{-4}$ ) and 1:1,000,000 ( $10\text{E}^{-6}$ ) dilutions. Mix and pour onto the appropriately labeled Nutrient Agar plates (provided in the kit). The plated volumes result in final dilutions of  $1 \times 10^{-5}$  and  $1 \times 10^{-7}$  for the 1:10,000 ( $10\text{E}^{-4}$ ) and 1:1,000,000 ( $10\text{E}^{-6}$ ) dilutions in water, respectively.

## 12. Incubate the Assay

- a. Invert the plates and arrange Minimal Glucose Agar plates in stacks corresponding to each experimental condition.
- b. Place in a 37°C incubator.
- c. Incubate QUAD PC™ Plates and Nutrient Agar titer plates overnight.
- d. Incubate Minimal Glucose Agar Plates for approximately 48 h.

## 13. Read the Assay

- a. After the incubation period, remove the inverted plates and allow to come to room temperature. Turn the stacks of plates over so that the tops are up. If excessive condensate has formed on the lids while incubated in the inverted position, remove the condensate by removing the lids one at a time (keeping the plate upside down; agar should face the floor), sharply shaking the lid and replacing before turning the plates over.
- b. Colony counting can be performed manually with the aid of a magnifying counter (e.g., “Quebec” counter) or with an automatic colony counter (e.g., Neutec, Symbiosis). Depending on the activity of your test material, large numbers of colonies may develop in certain dose groups. In some cases, it may be desirable to utilize sector-counting techniques rather than full plate counts. However, sector counting is not appropriate if the distribution of colonies is nonuniform across the surface of the agar.

- c. Examine the background lawn using a microscope (40x) or similar instrument. A normal background consists of densely packed microcolonies forming a thin, somewhat granular, film. If a plate contains many very small, just macroscopic “pinpoint” colonies and reveals an absence or “thinning” of the background lawn the test material dose was toxic. Mutant colony counts from plates exhibiting toxicity should not be considered in activity determinations.
- d. After counting and recording the results for the test material treatments, the diagnostic positive control plates should be counted. The colony counts for the positive control treatments should be compared to the values described in the STDisc™ CoA sheets.
- e. Examine the cell titer (Nutrient Agar) plates. The  $1 \times 10^{-5}$  plates should be too numerous to count. In contrast, the  $1 \times 10^{-7}$  plates should contain approximately 100 - 200 colonies. Such a result indicates that the initial population (the stock culture) was in the range of  $1 - 2 \times 10^9$  viable cells per ml. Very much lower or higher initial titers may result in reduced frequencies and background or increased backgrounds, respectively.
- f. The QUAD PC™ plates (phenotypic confirmation media) should be examined and the results compared to the below table.

Sector		Strain Response	
Designation	Description	TA1535 TA1537	TA98 TA100
<b>1 (I, A)</b>	MGA + Biotin	-	-
<b>2 (II, B)</b>	MGA + Histidine/Biotin	+	+
<b>3 (III, C)</b>	MGA + Histidine/Biotin/Ampicillin	-	+
<b>4 (IV, D)</b>	MGA + Histidine/Biotin/Ampicillin/Tetracycline	-	-

All strains should exhibit a halo around the crystal violet disc (i.e., no growth in the presence of crystal violet).

## D. The Results - Evaluation and Interpretation

### 1. Negative (solvent) Control Counts

The colonies that grew on the Minimal Glucose Agar plates developed from single cells that regained their ability to grow in the absence of added histidine. The genetic reversion, from histidine auxotrophy to prototrophy, that enabled those cells to grow in the absence of exogenous histidine might have arisen spontaneously or as the result of a mutation induced by the treatments (see Maron & Ames, p. 181). It is important to realize that some of the colonies that arose in the positive control plates would have grown in the absence of treatment; they arose spontaneously. Accordingly, the negative (solvent) control colony counts constitute an important baseline in your evaluation of the test results.

Unfortunately, the spontaneous reversion frequencies for the various tester strains can be variable. Nevertheless, large deviations from the “normal” range of spontaneous reversion values may signal systematic problems with the assay. Taking into account the fact that there may be lab-to-lab differences, the following ranges for spontaneous reversion values for the included strains may be representative:

<b>Strain</b>	<b>Spontaneous Reversion Rate Values</b>
TA1535	5 - 20
TA1537	5 - 20
TA98	20 - 50
TA100	75 - 200

## 2. Diagnostic Positive Control Counts

If you used CONTROLCHEM™ chemicals as suggested, the numbers of revertants (mutants) should fall within the following guideline ranges:

<b>-S9/Direct Acting Mutagens</b>		
<b>Strain</b>	<b>Chemical</b>	<b>Number of Colonies/Plate</b>
TA1535	Sodium Azide	≥ 200 - 650
TA1537	ICR 191	≥ 40 - 200
TA98	Daunomycin	≥ 450 - 1,850
TA100	Sodium Azide	≥ 300 - 650

<b>+S9/Indirect Acting Mutagen</b>			
<b>Strain</b>	<b>Chemical</b>	<b>Number of Colonies/Plate</b>	
		<b>5% S9 Mix</b>	<b>10% S9 Mix</b>
TA1535	2-Aminoanthracene	≥ 90 - 210	≥ 75 - 150
TA1537		≥ 200 - 300	≥ 160 - 180
TA98		≥ 1280 - 1800	≥ 1200 - 2000
TA100		≥ 1500 - 1750	≥ 1150 - 1650

In general, the positive control frequencies (number of colonies per plate) should be at least 2.5 times the negative control counts (spontaneous frequency). Large deviations usually indicate problems with cell husbandry; e.g., high spontaneous frequencies (due, perhaps to culture overgrowth) often are paralleled by low induced frequencies. Such eventualities reduce the resolving power of the assay and raise questions regarding the interpretation of the results of the test material treatments.

### 3. Phenotypic Confirmation

The QUAD PC™ plates are prepared with four different media that provide basic information concerning the genotypes of the strains provided in the kit (see the CoA sheets for the specific strains). By sector, the results should be:

Sector	Observation	Genotype
<b>1 (I, A)</b>	No growth (all strains)	<i>his</i>
<b>2 (II, B)</b>	Growth. Zonal inhibition around CV disc (all strains)	<i>his, rfa</i>
<b>3 (III, C)</b>	Growth of TA98 and TA100 only	pKM101
<b>4 (IV, D)</b>	No growth (all strains )	pAQ1 (present in TA102 only)

### 4. Test Material Results

Various investigators have applied different criteria for the analysis of the assay results. In general, the 2 or 2.5 times over the background (spontaneous frequency) “rule-of-thumb” may serve as a useful way of distinguishing active mutagens from non-mutagenic materials. The presence of a dose response (not necessarily linear) is often used as an adjunct criterion for and interpretation of positive activity in the assay. We highly recommend that you read pages 195-197 in Maron & Ames (1983) or pages 49 – 50 in Mortelmans & Zeiger (2000) for additional information concerning the interpretation of the assay results.

Version FAM002.2Amesman  
12 May 2026

**MOLTOX**<sup>®</sup>  
Molecular Toxicology, Inc.

157 Industrial Park Drive Boone, NC 28607 828.264.9099 Toll Free: 800.536.7232 Fax: 828.264.0103

[www.MOLTOX.com](http://www.MOLTOX.com) [sales@moltox.com](mailto:sales@moltox.com)