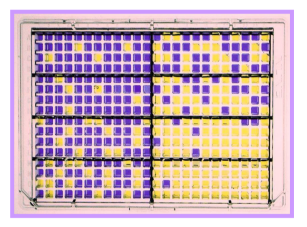


Comparison of the Performance of the Colorimetric Ames MPF Assay with the Agar Plate Method

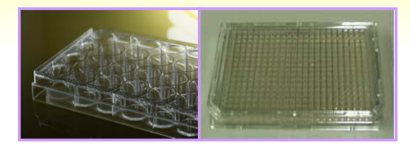


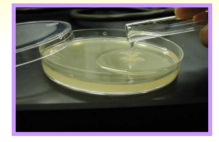
Nicole Weiland, Xenometrix AG

April 2016



Ames MPF and Ames agar plate test



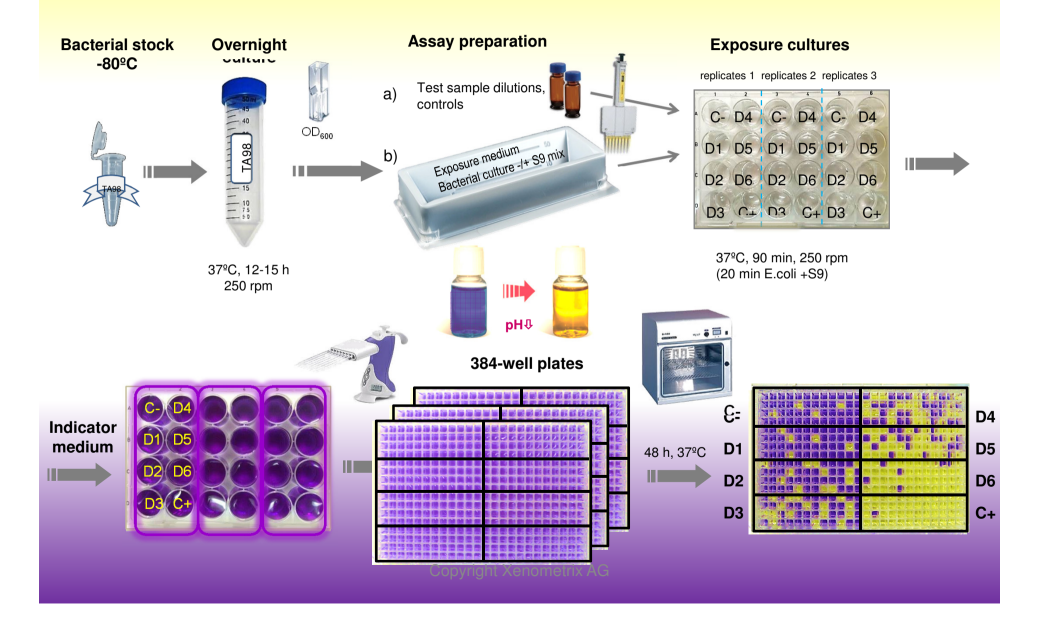


Ames MPF is based on same principle as agar plate test but

- Liquid low-volume format
- Use of microplates and multichannel pipettes
- Colorimetric read-out
- Less test sample up to 4 fold
- Less S9 up to 12 fold
- Higher throughput



Ames Microplate Assay Procedure





Measuring Points

Agar Plate test

- 1 plate 1 measuring point
- Individual handling:
 1 plate requires mixing of
 1 compound, agar and plating

liquid culture Ames MPF

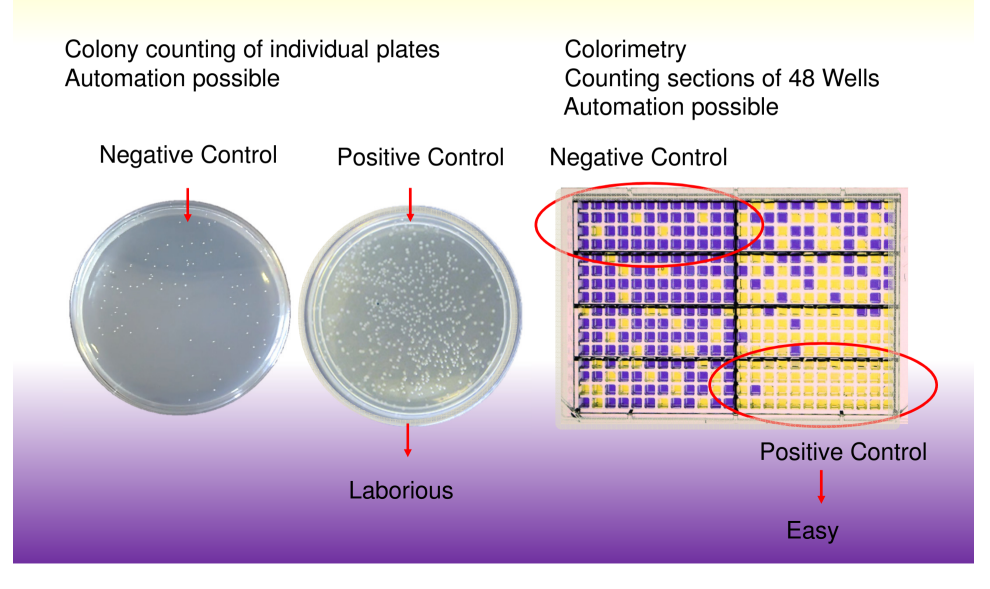
- 1 plate 24 measuring points
- Simultaneous handling of several replicates







Evaluation of Results Agar Plate Test vs Ames MPF

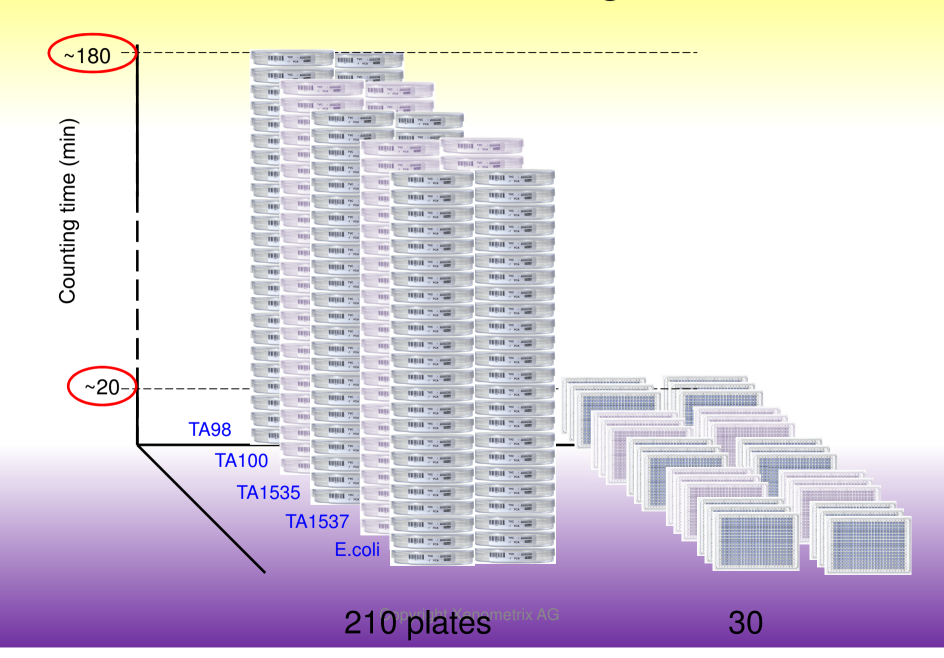


Throughput of compounds: **XENOMETRIX** Hands-on-time for 1 compound in 5 strains

1 sample, 5 concentrations, 5 strains (OECD), -/+ S9, controls, triplicates, \rightarrow Conditions: manual handling, ready-made agar plates and top agar

Ą	gar Plate / 5 Conc.	MPF / 6 Conc.	
Sample dilutions:	~5 min	~5 min	
Top agar (preparation of tubes): Addition of sample, culture, S9:	~35 min ~50 min	- ~25 min	
Plating: Transfer to 384-well plates:	~40 min -	- ~40 min	
Handling time:	~130 min	~70 min	
Counting time:	~180 min	~20 min	
Total time:	~ 5 h	~1½ h	

Visualization of Plate Counting Time



XENOMETRIX

Swiss Commitment for Bioassay

Test Sample Consumption



Minimum amount of sample needed: Agar plate test vs. Ames MPF

Setup: 5 strains (OECD 471), ½ log dilution steps, triplicates, -/+ S9

<u>Ames Agar Plate</u>:

Top dose: 5 mg/plate

Test sample: 220 mg

5 mg/ml

Ames MPF:

55 mg

Ames MPF:

⇒ 4-fold less test sample
 ⇒ Very important when compound quantity is limited!
 ⇒ Genotoxic impurities

S9 Consumption



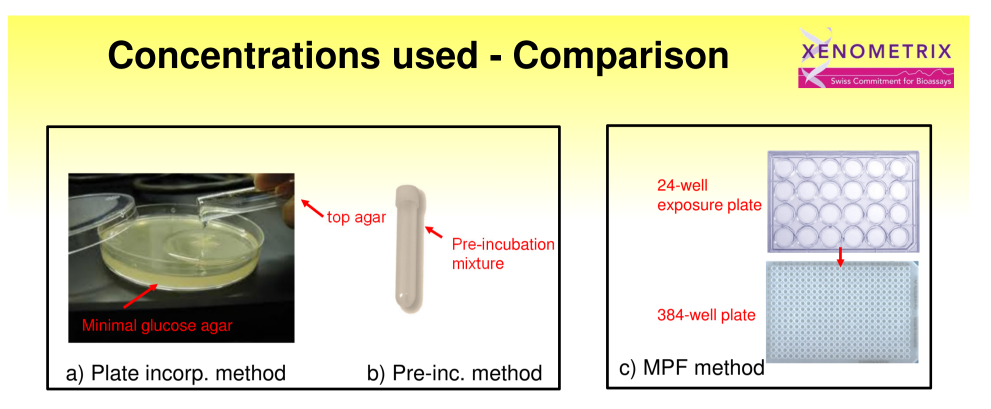
Setup: 5 strains (OECD 471), ½ log dilution steps, triplicates, S9

	Ames Agar Plate:	Ames MPF:					
S9 30%:	15.57 ml	1.35 ml - usually applied in Ames MPF					
S9 10%:	5.25 ml - usually applied in agar plate test	0.45 ml					
Ames MPF:							
 ⇒ 4-fold up to 11-fold less S9 ⇒ Reduced number of sacrificed animals ! ⇒ In line with 3Rs: Replace, Reduce, Refine ! 							



Critical Points of Ames MPF

- Comparability of concentrations used (mg/plate mg/ml)?
- 48-well limit?
- Cytotoxicity?
- Colored compounds: Interference with colorimetric read-out?



a) Plate incorporation: defined sample amount in top agar

- \rightarrow immediate pouring
- \rightarrow possible diffusion of sample and cofactors into lower agar



- \rightarrow volume not always clearly defined during exposure
- b) Pre incubation.: defined sample amount in defined volume
 - \rightarrow <u>liquid</u> pre-incubation/exposure \rightarrow dilution with top agar \rightarrow pouring
 - \rightarrow defined volume during exposure

c) Ames MPF: defined sample amount in defined volume

- \rightarrow <u>liquid</u> exposure \rightarrow dilution with indicator medium
- \rightarrow defined volume during exposure



Sample Concentration - Comparison

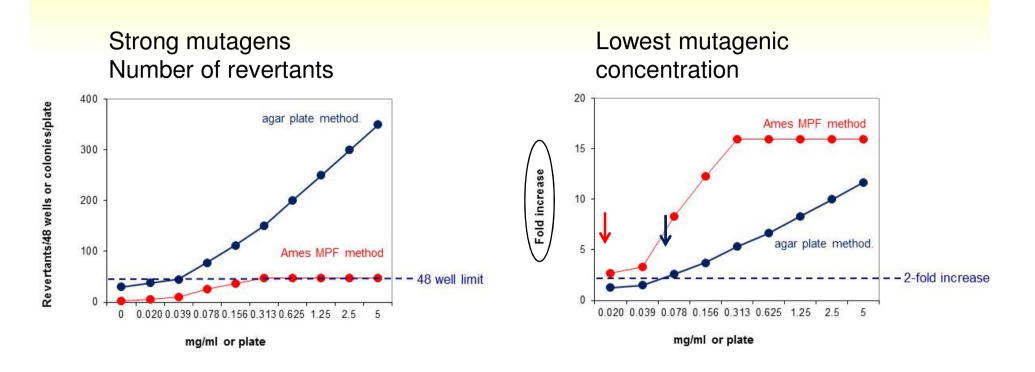
MPF method and Pre-incubation method: Both exposures performed in liquid media ⇒ Bacteria incubated with constant sample concentrations

Liquid exposure with <u>5 mg/ml</u> (MPF) or <u>5 mg/plate</u> (pre-incubation)

	Addition	Stock	Final Volume	Final concentration
MPF	10 µl	125 mg/ml	0.25 ml	5.0 mg/ml
Pre-incubation	100 μl	50 mg/ml	0.70 ml	7.1 mg/ml



48 Well Limit



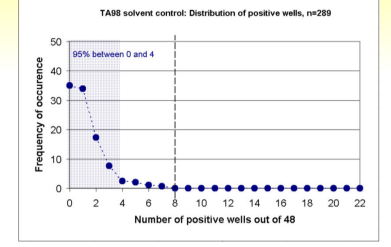
- No limits of revertants for strong mutagens in

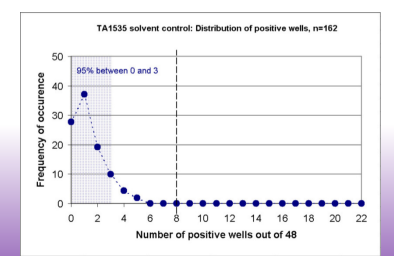
 agar test, continous increase of revertants
- Plateau of 48 wells, but: Repeated 48 revertant wells = strong mutagen in Ames MPF

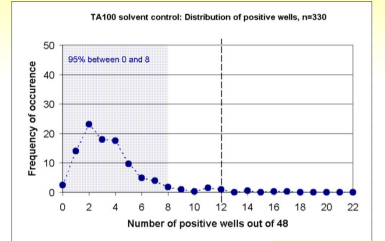
- Ames MPF detects lowest mutagenic concentration at lower dosis
- Low number of spontaneous revertants

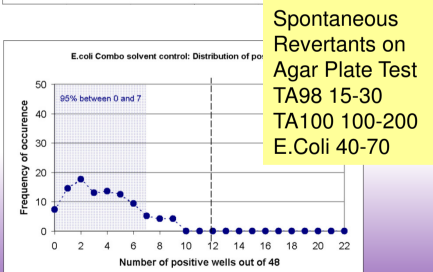
Historical Solvent Control







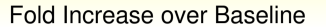


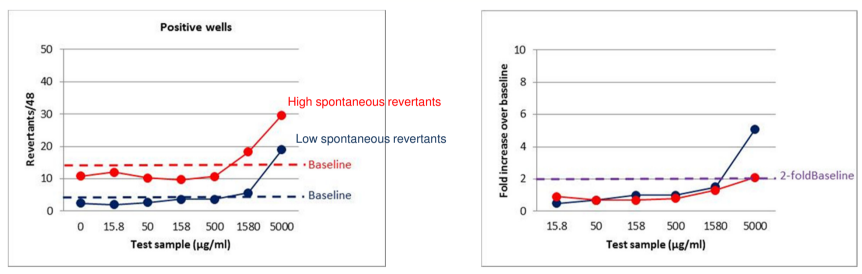




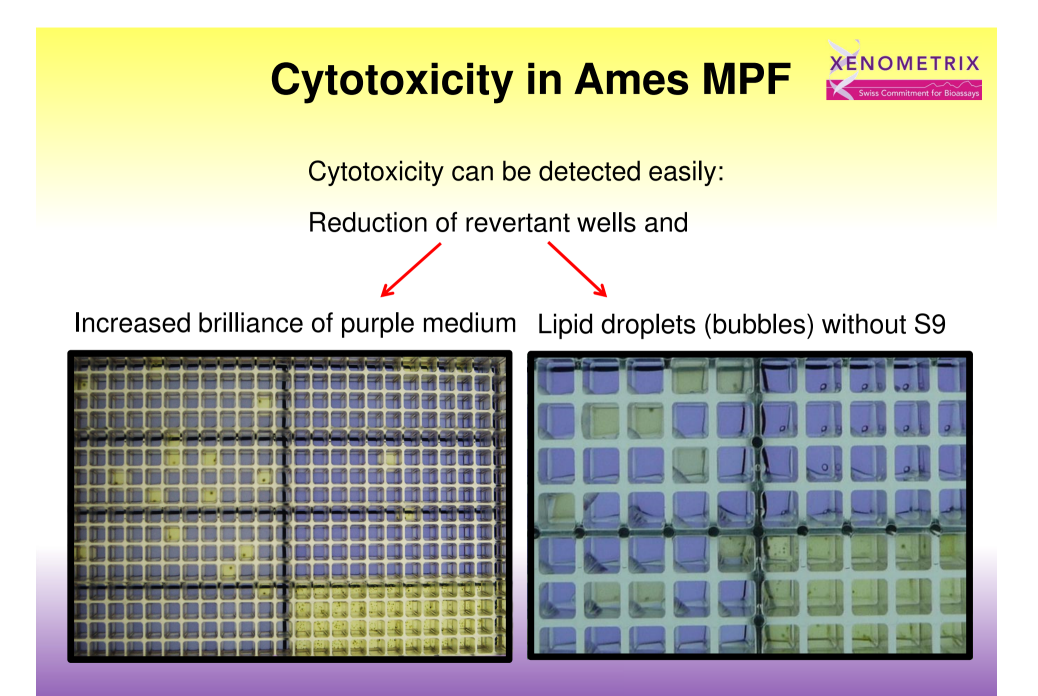
48 Well Limit in Ames MPF "Low" and "High" Spontaneous Revertants

Positive Wells



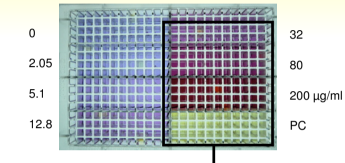


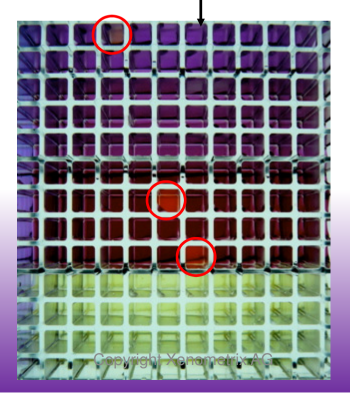
- ⇒ Pass/Fail criteria for spontaneous revertants in Ames MPF
 ⇒ Low spontanous revertants -> larger dynamic range
- Selection of cultures with low spontaneous revertant rate at Xenometrix, 2 quality controls after production





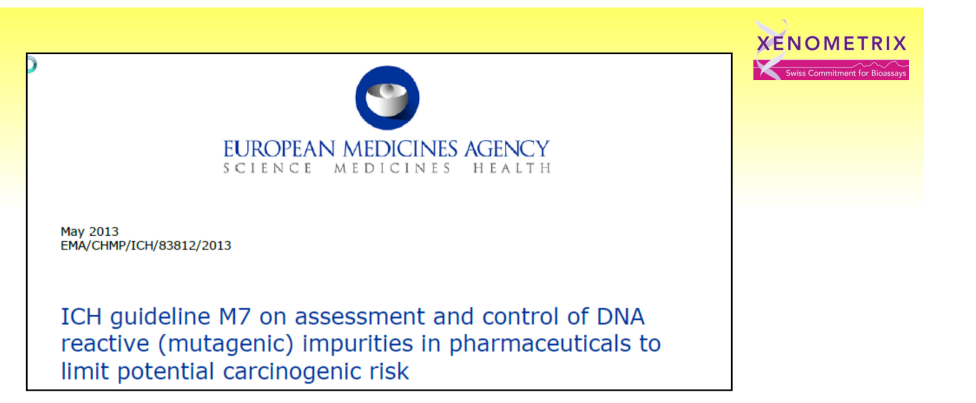
Colored compounds - colorimetric read-out





Orange instead of yellow wells

Easily detectable

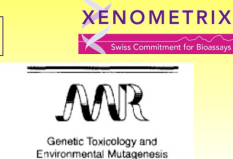


Note 2

To assess the mutagenic potential of impurities, a single bacterial mutagenicity assay can be carried out with a fully adequate protocol according to ICH S2(R1) and OECD 471 guidelines......For degradants that are not feasible to isolate or synthesize or when compound quantity is limited, bacterial mutagenicity testing could be carried out using a miniaturized assay format with proven high concordance to the ICH-compliant assay to enable testing at higher concentrations with justification.....

High concordance with agar plate test





Mutation Research 412 (1998) 115-130

Comparison of responses of base-specific Salmonella tester strains with the traditional strains for identifying mutagens: the results of a validation study

P. Gee ^{a.*}, C.H. Sommers ^a, A.S. Melick ^a, X.M. Gidrol ^a, M.D. Todd ^a, R.B. Burris ^a, M.E. Nelson ^a, R.C. Klemm ^a, E. Zeiger ^b

TA98, TA1537, TAMix compared with all strains NTP

25 chemicals tested

Overall agreement: 88%

Absuact

The ability of a TA7000 series of Salmonella his⁻ mutant tester strains to detect mutagens as classified by the traditional

High concordance with agar plate test





Available online at www.sciencedirect.com

DIRECT

Mutation Research 558 (2004) 181-197



Genetic Toxicology and Environmental Mutagenesis

www.elsevier.com/locate/gentox Community address: www.elsevier.com/locate/mutres

Assessment of the performance of the Ames IITM assay: a collaborative study with 19 coded compounds

S. Flückiger-Isler^{a,*}, M. Baumeister^b, K. Braun^c, V. Gervais^d, N. Hasler-Nguyen^e, R. Reimann^f, J. Van Gompel^g, H.-G. Wunderlich^h, G. Engelhardtⁱ

 ^a Xenometrix by Endotell GmbH, CH-4125 Allschwil, Switzerland
 ^b Boehringer Ingelheim, Department of Non-Clinical Drug Safety, Boehringer Ingelheim Pharma KG & Co. KG, D-88397 Biberach, Germany
 ^c Aventis Pharma Deutschland GmbH, Drug Innovation & Approval, Lead Optimization, Drug Safety Evaluation, D-65795 Hattersheim, Germany
 ^d Servier Group, Drug Safety Assessment, F-45403 Orléans-Gidy, France

Novartis Consumer Health, Toxicology, CH-1260 Nyon, Switzerland

^f Schering AG, Experimental Toxicology, D-13342 Berlin, Germany

⁸ Johnson&Johnson Pharmaceutical Research & Development, Department of ADME/Tox, B-2340 Beerse, Belgium
^b Federal Environmental Agency, Department for Hygiene of Drinking and Swimming Pool Water, D-08645 Bad Elster, Germany

Overall agreement standard Ames (all strains) - Ames II (TA98, TAMix): 84.2% (16/19)

Inter-laboratory consistency of 89.5% (17/19).



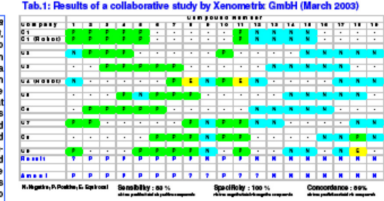
ASSESSMENT OF A SCREENING EXPERIENCE WITH THE AMES II™ TEST AND FUTURE PROSPECTS

V. GERVAIS1, D. BIJOT1 and N. CLAUDE2

¹ Drug Salety Assessment, Servier, Otléans-Gidy, France, ² IRIS, Servier, Coubevole, France

quid fluctuation version of the Salmonella mutagenicity assay, provided by Xenometrix GmbH, was used for an early compound selection in the discovery process. The ail e Ames II compared to the standard Ames test and to explore a way to reduce the required compound quantity without lowering the predictability of the test.

ETHODS of a mixture of 6 Salmonella TA7001, TA7002, TA7003, 1 TA7006, which revert to a specific base substitution n. This "mix" is used as a n, the frameshift tester strain mix and TA98 strains are lium for growth overnight at it, performed in 24-wells ws partial automation and about 60-fold less compound lard Arnes. After a 90 minith or without Aroclor-induced with solvent and positive medium lacking histidine is ch well is then aliquoted into



RESULTS

350 compounds were tested, including molecu from our own research, known non- or genoto molecules producing equivocal result concordance between the results achieve Ames II™ test and those reported in the liter the standard Arnes test ranged from 79 (Ref. (Tab.2). The concordance reached 89 collaborative study (Tab.1). No false positi were obtained with known non-mutagenic se False negative results may arise when chemi only specific strains like TA1535 or E. cdi (pKM101) which meet no equivalent in the 'mit The positive responses were randomly among the strains or the concentration range 3). In contrast, only 11% of positive results specifically in the absence of S9 (Fig.4), wh

- -----

83% Concordance Ames II vs. traditional Ames using 42 company-own chemicals (disagreement mainly with compounds that specifically revert E.coli, TA1535)

No false positive results



Mutagenesis vol. 24 no. 4 pp. 359–366, 2009 Advance Access Publication 15 May 2009 doi:10.1093/mutage/gep017

Comparison of the Ames II and traditional Ames test responses with respect to mutagenicity, strain specificities, need for metabolism and correlation with rodent carcinogenicity

Markus Kamber*, Sini Flückiger-Isler, Günter Engelhardt¹, Rudolf Jaeckh² and Errol Zeiger³

Xenometrix AG, Gewerbestrasse 25, CH-4123 Allschwil, Switzerland, ¹Experimental Toxicology and Ecology, BASF SE Product Safety, ²Regulations, Toxicology and Ecology, BASF SE Product Safety, 67056 Ludwigshafen am Rhein, Germany and ³Errol Zeiger Consulting, Chapel Hill, NC 27514, USA

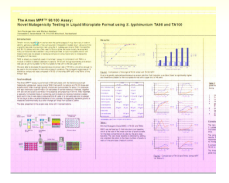
The Ames II Salmonella mutagenicity assay procedure was used to test 71 chemicals, and the results were compared with those from the traditional Ames Salmonella test using a different test method, including a simple, overall agreement or disagreement; agreement or disagreement with regard to the genetic endpoint, and whether metabolic activation is required for activity; comparisons of the active test chemical concentration ranges and with respect to the effect the test is designed to predict, i.e. cancer. Two previous studies (3,4) have compared the performance of the Ames II assay to that of the traditional Ames test procedure [i.e. the procedure with the traditional strains, as described in (5) and (6)] to validate its use as an alternative to the traditional Ames test procedure.

the NT perform format. 84% agreement between the two procedures in identifying mutagens and non-mutagens

Discordant results included chemicals requiring reductive metabolism using FMN, hamster liver S9



Xenometrix Posters: Comparison with Correspondent Traditional Strains



- TAMix vs. TA100 MPF and TA100 published traditional Ames
- TA98, TA100, TA1535, TA1537 MPF vs. TA98, TA100, TA1535, TA1537 published traditional Ames
- Ames MPF PENTA I (strains as above plus EC Combo) vs. published traditional Ames





Direct Comparison Ames MPF - Ames Pre-incubation

Mutation Research 747 (2012) 36-45



Contents lists available at SciVerse ScienceDirect Mutation Research/Genetic Toxicology and Environmental Mutagenesis journal homepage: www.elsevier.com/locate/gentox Community address: www.elsevier.com/locate/mutres

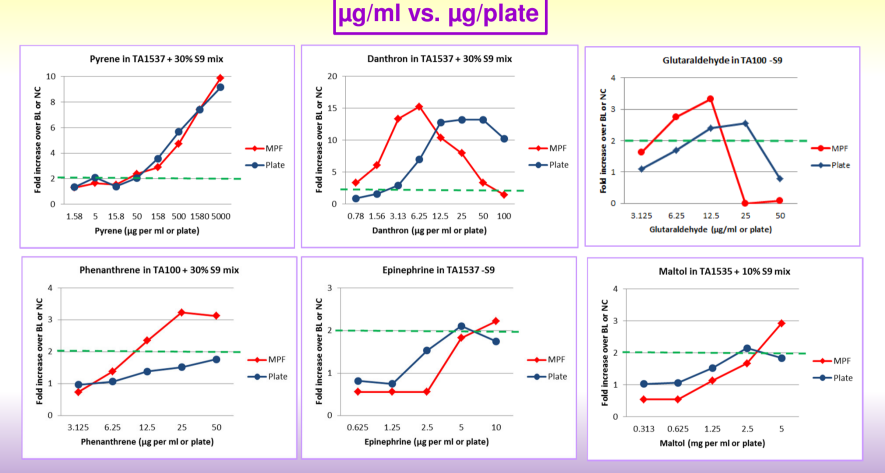


Direct comparison of the Ames microplate format (MPF) test in liquid medium with the standard Ames pre-incubation assay on agar plates by use of equivocal to weakly positive test compounds

Sini Flückiger-Isler*, Markus Kamber Xenometrix AG, Allschwil, CH-4123 Allschwil, Switzerland

- 15 equivocal to weakly positive chemicals
- Same overnight cultures, chemicals and S9 to exclude external variations, i.e. culture growth, chemical purity, weighing errors, S9 activity
- Parallel tests with most responsive strains of the NTP database (mg/plate vs. mg/ml)
- Each test was repeated at least once
- 87% concordance (13/15)
- Excellent concordance for equivocal to weak positive chemicals
- Confirms the high concordance with the ICH-compliant assay

Direct Comparison Ames MPF and Pre-incubation Method (see publication before)



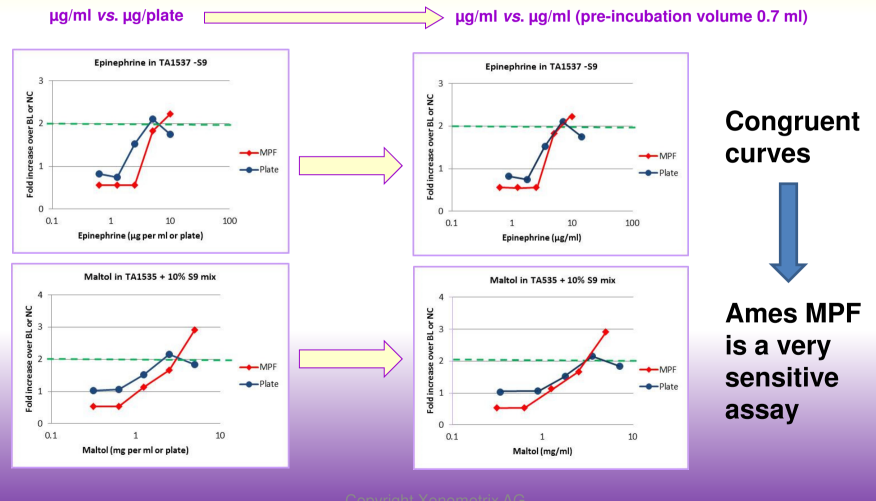
Higher sensitivity of Ames MPF with several compounds, such as Danthron, Glutaraldehyde, Phenanthrene At first glance higher sensitivity of Pre Incubation Assay with Maltol and Epinephrine, but....

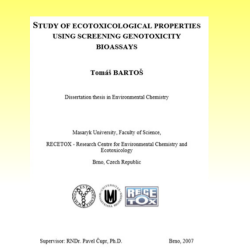
Direct Comparison Ames MPF and Pre-incubation Method – Epinephrine, Maltol

XENOMETRIX

Swiss Commitment for Big

Correction for concentration in preincubation assay (5.0 mg vs 7.1 mg)





Chapter 5

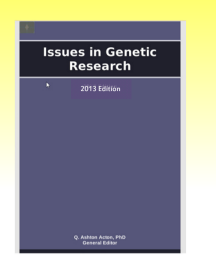
Perspectives in genotoxicity screening

"Ames II (Xenometrix, Switzerland) is a microplatebased fluctuation test version of the Ames test and probably the best Ames predictor."





Chapter 10: Ames II and Ames Liquid Format Mutagenicity Screening Assays

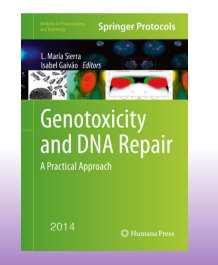


Chapter 9: Mutation Research

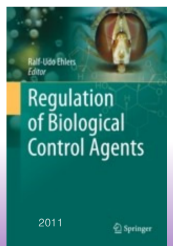


New Paradigms for the Environmental Assessment, p. 26

"Thus, the all-liquid format of the Ames II/MPF assay, which requires less test compound and allows for the use of multichannel pipettes, thus automating the pipetting steps, makes this procedure an attractive method to evaluate mutagenicity of a large number of samples at the same time - a common situation in environmental monitoring".



Chapter 2: The Ames II and Ames MPF Penta I Assay: A Liquid Microplate Format Modification of the Classic Ames Test



..it has been proposed by the European Union-funded REBECA project as a screening tool to determine whether fungal biological control agents produce genotoxic/mutagenic metabolites which require further attention in the regulatory risk assessment.

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Conclusion I - Test Performance

- Ames MPF Ames agar test: same principle, same tester strains
- Comparative studies: mean concordance of ~87%
- Comparable to the intra- and inter-laboratory reproducibility of the agar plate Ames test procedure

Conclusion II



Advantages

- 4 x less test sample necessary
- Liquid microplate format allows for less handson-time, simultaneous processing of several replicates
- Higher throughput, partly automatable
- 12 fold less consumption of S9 following 3Rs
- Quick, easy colorimetric read-out, less error prone
- Less plastic ware, reduced contaminated waste in environment
- Listed explicitly in ICH M7 Guideline
- Higher Sensitivity depending on compound



Disadvantages

- Not same large database as agar plate method
- Not listed explicitly in OECD 471



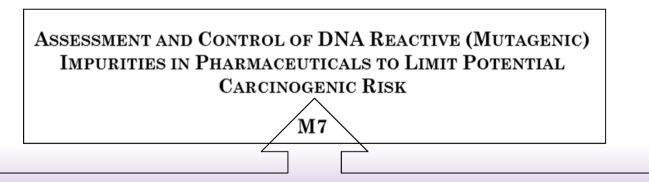
Conclusion III - ICH Guideline M7

XENOMETRIX

Swiss Commitment for F

- The Ames MPF features a miniaturized assay format with proven high concordance with the ICH-compliant assay.
- It is highly sensitive and allows testing compounds present in limited quantity.

⇒Ames MPF = Excellent tool for assessing mutagenic impurities



"For degradants that are not feasible to isolate or synthesize or when compound quantity is limited, it may not be possible to achieve the highest test concentrations recommended for an ICH compliant bacterial mutagenicity assay according to the current testing guidelines. In this case, bacterial mutagenicity testing could be carried out using a miniaturized assay format with proven high concordance to the ICH compliant assay to enable testing at higher concentrations with justification......"