In vitro mammalian cell micronucleus test

Commonly used acronym: In vitro MN test

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SCOPE OF THE METHOD

<table>
<thead>
<tr>
<th>The Method relates to</th>
<th>Human health</th>
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</thead>
<tbody>
<tr>
<td>The Method is situated in</td>
<td>Basic Research, Regulatory use - Routine production</td>
</tr>
<tr>
<td>Type of method</td>
<td>In vitro - Ex vivo</td>
</tr>
<tr>
<td>This method makes use of</td>
<td>Animal derived cells / tissues / organs</td>
</tr>
<tr>
<td>Species from which cells/tissues/organs are derived</td>
<td>Hamster</td>
</tr>
<tr>
<td>Type of cells/tissues/organs</td>
<td>CHO-K1 cells</td>
</tr>
</tbody>
</table>

DESCRIPTION

Method keywords
DNA damage
micronuclei
chromosomal damage
OECD TG 487
structural and numerical aberration chromosomal aberration
• automated scoring

Scientific area keywords
in vitro toxicology
Method description

The *in vitro* micronucleus test is a genotoxicity test for the detection of micronuclei in the cytoplasm of interphase cells and has been described in detail in OECD TG 487. Micronuclei may originate from acentric chromosome fragments (i.e. lacking a centromere), or whole chromosomes that are unable to migrate to the poles during the anaphase stage of cell division. Consequently, this *in vitro* method can detect both structural and numerical chromosome aberrations in cells that have undergone cell division during or after exposure to the test chemical. Within our lab, the *in vitro* micronucleus test with the actin polymerisation inhibitor cytochalasin B is applied. Cytochalasin B allows for the identification and selective analysis of micronucleus frequency in cells that have completed one mitosis, because such cells are binucleate. Scoring of the micronuclei in the binucleated cell is done in an automated way using the Metafer system. Other imaging software or flow cytometry can also be used as well as manual scoring.

Lab equipment

- Standard equipment for working with cell cultures;
- Fluorescence microscope;
- Analysis software for automated scoring (e.g. metafer 4).

Method status

Published in peer reviewed journal
Validated by an external party (e.g. OECD, EURL ECVAM,...)

PROS, CONS & FUTURE POTENTIAL

Advantages

- Simple and easy to identify endpoint.

Challenges
- Distinction between structural and numerical chromosome aberrations requires combination with other techniques (e.g. Fluorescence In Situ Hybridization - FISH);
- Detection of artefacts during automated scoring, manual verification needed;
- Automated flow cytometric detection for faster scoring of more test concentrations with more cells (benchmark dose approach) Flow Cytometry for automated cytotoxicity detection along with micronuclei scoring.

**Modifications**

- Use of other cell types, preferably p53-competent human-derived cell lines;
- Combination with Fluorescence In Situ Hybridization (FISH) to distinguish between structural and numerical chromosome aberrations;
- Use of flow cytometry for bench mark dose approach and simultaneous detection of cytotoxicity and MN scoring.

**REFERENCES, ASSOCIATED DOCUMENTS AND OTHER INFORMATION**

**References**


**Associated documents**

OECD in vitro mammalian cell micronucleus test.pdf

**Links**

OECD Test guideline

**PARTNERS AND COLLABORATIONS**

**Organisation**

Name of the organisation Sciensano

Department Chemical and physical health risks

Specific Research Group or Service Risk and health impact assessment
Country: Belgium
Geographical Area: Brussels Region