

# Measurement of Cytochrome P450 Enzyme Induction and Inhibition in human cells

Created on: 20-03-2019 - Last modified on: 28-02-2022

## Contact person

Steven Branson

## Organisation

**Name of the organisation** Vrije Universiteit Brussel (VUB)

**Department** Pharmaceutical and Pharmacological Sciences

**Specific Research Group or Service** In Vitro Toxicology and Dermato-Cosmetology

**Country** Belgium

**Geographical Area** Brussels Region

## SCOPE OF THE METHOD

<b>The Method relates to</b>	Human health
<b>The Method is situated in</b>	Basic Research
<b>Type of method</b>	In vitro - Ex vivo
<b>Specify the type of cells/tissues/organs</b>	parenchymal liver cells, stem cell-derived hepatocyte-like cells

## DESCRIPTION

### Method keywords

in vitro

hepatic cell line

luminescence

liver enzyme

viability test

## Scientific area keywords

Toxicology

Drug metabolism

drug screening

in vitro cell culture

## Method description

By the use of monolayer cultures as an *in vitro* system, the effects of drugs on CYP3A activity can be evaluated. It relies on the use of a luminogenic CYP3A substrate, namely luciferin-6'-pentafluorobenzylether (luciferin- PFBE). Upon biotransformation by CYP3A, luciferin-PFBE is converted into luciferin, which generates light when combined with a luciferin detection reagent. The normalization of the data relies on the cell number and cell viability and is evaluated by another bioluminescence reaction in which the levels of adenosine- 5'-triphosphate (ATP), the basic energy source of living cells, is measured.

## Lab equipment

Biosafety cabinet;

Luminescence plate reader;

White opaque;

96-well plates.

## Method status

History of use

## PROS, CONS & FUTURE POTENTIAL

### Advantages

Quick and easy to use.

## REFERENCES, ASSOCIATED DOCUMENTS AND OTHER INFORMATION

Coordinated by



Financed by

