

# MAINTAIN CYP ACTIVITY & INDUCIBILITY

### INTRODUCTION

A relevant *in vitro* model should display CYP inducibility that mimics in vivo metabolic responsiveness to known modulators. This response of CYPs to induction is a critical parameter for predicting adverse drug reactions.

#### **Materials required**

- BIOMIMESYS®Hepatocyte
- Cryopreserved human hepatocytes
- HCM BulletKit (Lonza)
- Rifampicin, Omeprazole (inducers)
- Salicylamide (phase 2 enzymes inhibitor)

by

- > Testosterone, Buproprion, Phenacetin (substrate)
- > DNA quantitation kit, fluorescence assay using bisBenzimide (Sigma Aldrich)
- ➢ LC-MS/MS

### Hydroscaffold properties

Translucent and porous

### Method

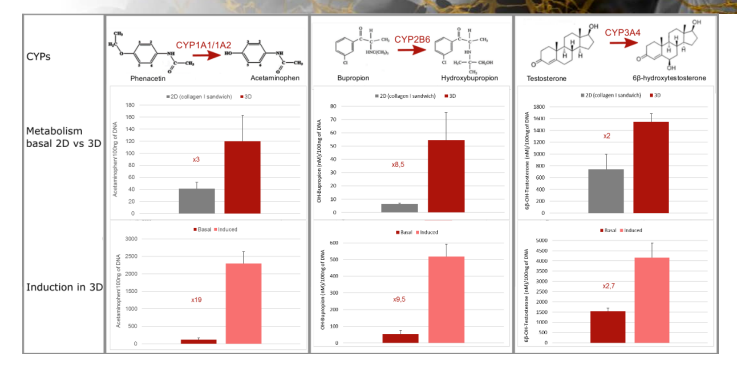
- Expose hepatocytes to CYP-inducers for 48h with daily medium exchange
- Add specific substrates for 24h
- Measure the DNA quantity and quantify the metabolites by LC-MS/MS at day 6 for 2D culture (collagen I sandwich) and day 10 for 3D culture

#### RESULTS

Measurement of basal and induced activities of CYP1A1/1A2, CYP2B6 and CYP3A4 in cryopreserved human hepatocytes (n=3), with BiIOMIMESYS<sup>®</sup>Liver



# **Biofunctionalized** hydroscaffold for 3D culture



Pharma

## CONCLUSION

Basal and induced CYP1A1/A2, CYP2B6, CYP3A4 activities are higher in cryopreserved human hepatocytes using BIOMIMESYS<sup>®</sup>Liver compared to 2D culture conditions. BIOMIMESYS<sup>®</sup>Liver is a robust 3D culture system for DILI studies.

## **Contact Information**

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