


 Cite this: *Lab Chip*, 2026, 26, 2577

## *In vitro* space of Disse model for exploration of drug induced hepatotoxicity

 Ana Mesic, \*<sup>a</sup> Antonietta Messina,<sup>b</sup> Zoe Tiprez,<sup>b</sup> Benoit Charlot, <sup>c</sup>  
 Safa Mohamed Ismail,<sup>d</sup> Nicolas Huang,<sup>d</sup> Sakina Bensalem,<sup>a</sup>  
 Jean-Charles Duclos-Vallee<sup>b</sup> and Bruno Le Pioufle<sup>a</sup>

The limited predictive power of animal models remains a major bottleneck in drug development, particularly in assessing drug-induced liver injury (DILI). To address this, we developed a novel *in vitro* liver-on-a-chip platform focused on modelling the space of Disse (sD) – a key microenvironment, yet insufficiently studied, which mediates cross-talk between endothelial and hepatic systems. The system integrates a biocompatible sodium alginate hydrogel whose mechanical properties were optimised to mimic physiological liver stiffness, enabling molecular cross-talk between human liver sinusoidal endothelial cells (LSECs) and HepaRG hepatocytes without direct contact. Under dynamic perfusion, the co-culture showed viability and polarisation for 8 days, forming organised tissues with functional bile canaliculi. The presence of LSECs markedly enhanced hepatic performance, reflected by increased albumin and urea secretion and activation of pro-regenerative secretory pathways. Prolonged acetaminophen exposure demonstrated the model's capacity to reproduce hepatotoxic responses, confirming its predictive relevance. Such versatile microphysiological platform might become a powerful tool for studying endothelial-hepatic communication, modelling liver pathologies, and improving preclinical DILI testing.

 Received 10th December 2025,  
 Accepted 13th March 2026

DOI: 10.1039/d5lc01139f

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### Introduction

The efficiency of pharmaceutical research and development (R&D) has been in continuous decline for decades, with productivity estimated to have dropped nearly 80-fold since the 1950s when adjusted for inflation.<sup>1</sup> Bringing a single drug to the market today requires on average \$1.56 billion, equivalent to 6500–15 000 person-years of work and about 10 to 12 years.<sup>2</sup> Moreover, drug-induced liver injury (DILI) is a leading cause of late-stage failure and post-marketing withdrawals.<sup>3</sup> A major driver of these inefficiencies is the limited predictive power of animal models used during preclinical testing. Despite showing apparent efficacy and safety in preclinical studies, 92% of candidate drugs ultimately fail during clinical trials.<sup>4</sup> Beyond ethical concerns, animal testing is 1.5–30 times more expensive than *in vitro* methods and often poorly predictive of human physiology.<sup>5</sup> Together, these limitations underscore the urgent need for advanced *in vitro* platforms that combine human relevance with efficiency.

Organ-on-a-chip technologies have emerged as a promising solution to the limitations of conventional preclinical models.<sup>6</sup> These platforms replicate key physiological cues using a minimal set of cells, through several complementary strategies. In the case of the liver-on-a-chip, they enable the co-culture of multiple cell types, where hepatocytes maintained alongside endothelial cells display enhanced viability and functionality, reflected in increased albumin and bile secretion.<sup>7–9</sup> Besides, the incorporation of biomimetic materials within the organ-on-a-chip structure permits the fine-tuning of extracellular matrix (ECM) properties. Indeed, hepatocytes respond through their gene expression and functionality to substrates of varying viscoelasticity, mirroring the pathological stiffening of the ECM during fibrosis.<sup>10</sup> In addition, the application of dynamic perfusion introduces physiologically relevant shear stress, which has been shown to promote the differentiation of both hepatocytes and sinusoidal endothelial cells. Finally, micro compartments allow packing of hepatocytes at high cellular density, facilitating differentiation and the formation of bile canaliculi.<sup>11,12</sup> Despite these advances, current liver-on-a-chip models fail to reproduce the space of Disse (sD) – the extracellular region bridging hepatocytes and liver sinusoidal endothelial cells – as they do not establish spatial separation between different cell types and/or replicating mechanical properties, such as elasticity or porosity.

<sup>a</sup> University Paris Saclay, ENS Paris Saclay, CNRS LUMIN, France

<sup>b</sup> Laboratory of Physio-pathogenesis and Liver Disease Treatment, INSERM U1193, France

<sup>c</sup> IES, Institute of Electronics and Systems, CNRS University of Montpellier, France

<sup>d</sup> Université Paris-Saclay, CNRS, Institut Galien Paris-Saclay, Orsay, France


The sD is a thin perisinusoidal region between liver sinusoidal endothelial cells (LSECs) and the basal membrane of hepatocytes, typically only 500–1000 nm thick.<sup>13,14</sup> Its essential function is the maintenance of liver homeostasis through exchange of nutrients, metabolites and signalling molecules, between sinusoid and hepatocytes. Additionally, it is a niche for mesenchymal stem cells involved in liver regeneration,<sup>15</sup> hepatic stellate cells responsible for the formation of scar tissue during fibrosis<sup>16,17</sup> and target localisation for activated Kupffer cells during inflammatory responses.<sup>18</sup> Being such a complex microenvironment, it is highly sensitive to pathological remodelling. In pathologies such as liver fibrosis, cirrhosis, or DILI, the sD goes through multiple cellular, structural and biochemical changes: hepatocytes experience loss of polarity, reducing their ability for molecule intake,<sup>19</sup> LSECs undergo defenestration,<sup>20,21</sup> stellate cells become activated, proliferate, and deposit excessive ECM; Kupffer cells secrete proinflammatory cytokines; the composition and mechanical properties of the ECM significantly change.<sup>22</sup>

Despite its physiological and pathological importance, modelling the sD *in vitro* remains extremely challenging. Its nanometre-scale width requires advanced microfabrication techniques for spatial separation of multiple cell types, while maintaining correct biochemical cues and functional interactions. Furthermore, replicating a continuous plasma flow, cellular diversity, and ECM complexity adds layers of technical difficulty. Currently, there are three main strategies employed in *in vitro* sD models: use of microfluidic channels, porous membranes or hydrogels:

1. Microfluidic channels allow dynamic perfusion, chemical gradient formation, and high-density hepatocyte culture, promoting bile canaliculi formation and functional alignment; however, direct exposure to shear stress and limited multicellular complexity remain challenges.<sup>23,24</sup>
2. Membrane-based systems enable the co-culture of hepatocytes and endothelial cells on opposite sides of a semipermeable barrier, allowing differential media, controlled molecular exchange, and partial replication of the sD width (1–10  $\mu\text{m}$ ), yet ECM properties tunability is limited.<sup>25,26</sup>
3. Hydrogel-based approaches provide a tuneable 3D extracellular matrix that supports hepatocyte aggregation, spheroid formation, and endothelial network integration; when combined with microfluidics, hydrogels allow perfusion and shear stress, enhancing physiological relevance and enabling vascularised hepatic constructs.<sup>27–29</sup>

Here, we report on the development of an *in vitro* sD model on a microfluidic chip with an integrated biocompatible hydrogel possessing tuneable mechanical properties. The platform integrates two cell types: hepatocytes and endothelial cells, spatially separated by a sodium alginate hydrogel barrier that reproduces the mechanical properties of the extracellular matrix while enabling molecular cross-talk. This architecture allows physiological interactions without direct cell contact, providing a more faithful representation of the *in vivo*

sinusoidal microenvironment. Furthermore, we demonstrate the advantages of this system in comparison to single cell liver-on-a-chip models, and its utility in hepatotoxicity assays using 72-hour acetaminophen exposure. This article introduces a physiologically relevant sD model that addresses a major unmet need in liver tissue engineering and toxicology testing.

## Materials and methods

### Design and fabrication of the microfluidic device

Microfluidic chip designs were created using L-Edit IC (Siemens EDA Solutions) software. Silicon master-mould was fabricated by standard photolithography. SU-8 2100 photoresist (Kayaku Advanced Materials, USA) was spin-coated onto silicon wafers at 3000 rpm for 24 s, exposed for 24 s to 365 nm UV light through a chrome mask (Toppan<sup>TM</sup>) and hard baked for 5 min at 95 °C. Uncross linked resin was removed by immersion in SU-8 Developer for 4 min. SU-8 pattern height and consistency was measured with Dektak 150 Surface Profiler (Veeco, USA).

PDMS (polydimethylsiloxane, Sylgard 184, Dow Corning) was mixed in a 10:1 base-to-curing agent ratio, degassed under vacuum for 1 h, poured over the moulds, and cured at 90 °C for 2 h. The cured PDMS replicas were peeled, cut, and punched to create inlets and outlets (1.25 mm for side channels, 0.5 mm for the middle channel). PDMS layers and glass slides were plasma-activated (Diener Atto, 50% power, 6 s,  $6 \times 10^4$  Pa O<sub>2</sub>) and bonded. Devices were finalised by a 10 min post-bonding bake at 150 °C.

### Hydrogel loading and characterisation

**Formulation.** Sodium alginate (Manucol LKX) was prepared at 20 mg mL<sup>-1</sup> and filtered for sterilisation purposes. Calcium chloride, 30 mg mL<sup>-1</sup> CaCl<sub>2</sub> (calcium chloride anhydrous granular, >93%, C1016-500G, Sigma Aldrich), was used for polymerisation.

**Loading.** Alginate solution was loaded in the chip middle channel through Hamilton syringe 100  $\mu\text{L}$  by applying manual positive pressure (which we estimated to approximately 10 kPa). Crosslinking was done by rinsing side channels with CaCl<sub>2</sub> solution for 15 minutes and subsequent rinsing and stocking with PBS 1 $\times$  until cell loading.

**Characterisation.** The swelling capacity of alginate hydrogels was evaluated using the tea bag method.<sup>30</sup> Sodium alginate mass was crosslinked by immersion in CaCl<sub>2</sub> solutions for 15 min and subsequently rinsed by immersion in Milli-Q water for 2 h. Swelling capacity was calculated using eqn (1), where  $m_0$  and  $m_t$  corresponded to the crosslinked sodium alginate mass before and after the immersion respectively.

$$\text{SC} = \frac{m_t - m_0}{m_0} \quad (1)$$

Rheological measurements were conducted at room temperature using an AR-G2 rheometer (TA Instruments)



equipped with a 40 mm plate and 1.5 mm gap. Oscillatory amplitude and frequency sweep tests were performed to determine the linear viscoelastic regime and quantify the storage,  $G'$ , and loss,  $G''$ , moduli.  $G^*$ , complex shearing modulus, was calculated from eqn (2):

$$G^* = G' + iG'' \quad (2)$$

Considering a Poisson's coefficient  $\nu$  to be 0.5 for elastomers, Young's modulus,  $E$ , was derived from eqn (3):

$$E = 2(1 + \nu)G^* \quad (3)$$

Molecular transport across sodium alginate was tested by perfusing a fluorescent molecule, 0.12 mM CDFA (5-(6)-carboxyfluorescein diacetate, MW = 445.2 Da, Thermo Fisher, C369) through the side channel at  $1 \mu\text{L min}^{-1}$  for 120 min at  $37^\circ\text{C}$ . Imaging was performed every 5 min using an EVOS M7000 microscope (Invitrogen). Relative intensity,  $I_r$ , was calculated from eqn (4):

$$I_r = \frac{I_t}{I_p} \quad (4)$$

Here,  $I_p$  was the intensity of a CDFA (5(6)-carboxy-2',7'-dichlorofluorescein) solution in the perfusion channel, and  $I_t$  is the intensity in time  $t$  on the opposite side of the alginate barrier.<sup>31</sup>

### Cell culture on a chip

**2D culture.** Undifferentiated HepaRG cells were purchased from Biopredic International (HPR101, Saint-Gregoire, France) and cultivated according to their instructions. Passages 14–19 were used for chip seeding and functional tests. The cells were seeded at a density of  $2 \times 10^4$  cells per  $\text{cm}^2$  and cultured with William's E medium supplemented with HepaRG growth medium supplement and L-glutamine (2 mM). The medium was changed every two to three days and the cells were passaged every two weeks. Primary LSEC were supplied by Biopredic International (HUV101, Saint-Gregoire, France) and cultivated according to their instructions. Passage 1 was used for chip seeding and functional tests. The cells were cultivated at a density of  $10 \times 10^3$  cells per  $\text{cm}^2$  and cultured with EBM-2 Basal Medium (CC-3156) and EGM-2 SingleQuots Supplements (CC-4176) from Lonza. The medium was changed every two days.

**Cell loading on a chip.** Chips loaded with hydrogel in the middle channel were rinsed with  $4 \text{ mg mL}^{-1}$  rat tail collagen I (Sigma Aldrich, C3867) through the side channels and incubated at  $37^\circ\text{C}$  for 1 h. The cells were trypsinised, centrifuged at 1500 rpm for 5 min and resuspended into 50  $\mu\text{L}$  of a fresh medium. 1–3  $\mu\text{L}$  of cell pellet was deposited by a pipette in the inlet of one side channel. Negative manual pressure of approximately 10 kPa was applied on the opposite end of the side channel by a Hamilton syringe

of 100  $\mu\text{L}$ . LSEC cells were loaded on the chip one day before HepaRG cells, and the chip was rotated every 30 min for  $90^\circ$  to enable cell attachment to all channel walls. The medium was perfused through the opposite side channel until the HepaRG loading, which was done on the following day (day 0).

**Cell culture on a chip.** Culture was maintained in an incubator in dynamic conditions at  $1 \mu\text{L min}^{-1}$  for 8 days. Flow was established by a syringe pump (Standard Infusion Pump 11 Elite Syringe Pump, Harvard apparatus) set at the indicated flow rate and connected to the chip through a tubing (Liquid Flow Tygon Tubing Coil 1/16" OD  $\times$  0.02" ID) inserted in the inlet of the endothelial cell channel. The medium composition consisted of HepaRG Growth Medium Supplement, HepaRG Differentiation Medium Supplement and EGM-2 medium in a ratio 1:1:2.

**Toxicity assay.** Cytotoxicity was tested in conditions of prolonged exposure to acetaminophen (APAP) at a concentration of 2 mM. APAP was diluted in a cellular medium and perfused through the endothelial channel of the chip from day 5 to day 8 of the co-culture at a flow rate of  $1 \mu\text{L min}^{-1}$ .<sup>32</sup>

### Biological characterisation of the model

**Probe staining.** Fluorescent probe staining was used for characterisation of bile canaliculi functionality (Fluobile™, SL21FL01, Starlight), hepatic viability (MitoTracker Green FM, M7514, Invitrogen) and hepatic metabolic activity (P450 Glo CYP3A4 Assay, V9002, Promega). Fluorescent probes were injected in a hepatic channel, incubated at  $37^\circ\text{C}$  and rinsed. All dilutions and rinsing were performed with William's E medium without red phenol (unless specified differently in the manufacturer's protocol). Details about probe concentrations and incubation times are provided in the SI. The images were recorded with an EVOS™ FL Auto Imaging System.

**Immunofluorescent staining.** All dilutions were performed in PBS 1 $\times$ . The lengths of steps and references of primary and secondary antibodies are detailed in the SI. After the culture period, the cells were fixed on the chip with PFA (4%). They were then permeabilised using a solution containing bovine serum albumin (BSA, 1% w/v), EDTA ( $5.37 \times 10^{-1} \mu\text{mol L}^{-1}$ ), and Triton-X-100 (0.1% v/v). They were blocked with BSA (3% w/v) and incubated overnight at  $4^\circ\text{C}$  with primary antibodies diluted in BSA (1% w/v). The cells were then washed with a solution of tween 20 (0.1% v/v), and incubated at room temperature with secondary antibodies diluted in BSA (1% w/v). They were washed again with the tween solution and finally with PBS 1 $\times$ . The images were recorded using a Leica SP8 confocal microscope.

**Nucleic acids quantification.** Quantification of nucleic acids was performed on the cells collected from the chip. To extract the cells from microfluidic channels, TRIzol solution was injected in the inlet of each side channel, and cell



suspension was collected on the outlet. DNA and RNA extraction were obtained with Ozyme, DirectZol RNA/DNA Miniprep kit (according to the manufacturer's protocol), and concentrations were measured using Nanodrop LITE spectrophotometer (ThermoFisher Scientific).

**Albumin and urea quantification.** Protein markers quantification was performed using the perfused cellular medium, collected at 24-hour intervals during the culture and stored at  $-80\text{ }^{\circ}\text{C}$  until use. Specific ELISA kits were used according to the manufacturer's instructions: Human Albumin ELISA Quantification Set, Cat N E80-129, Bethyl Laboratories and Urea Assay kit, Sigma Aldrich, MAK006.

**Proteomic profiling.** Profiling was performed on the cellular medium using the Olink Target 96 Inflammation Panel (Olink Proteomics, Uppsala, Sweden) on the Molecular Quantification platform at Institut Pasteur, following the manufacturer's protocol. Analytes were tagged with matched antibody pairs conjugated to unique oligonucleotides and detected using the proximity extension assay (PEA) technology, a multiplex immunoassay with high specificity and sensitivity. To account for technical variability, inter-plate control normalisation was applied. Data were quality controlled, and proteins with  $>15\%$  missing values were excluded. Final results were provided as  $\log_2$ -transformed Normalised Protein eXpression (NPX) values.

### Statistical and image analysis

All images were analysed in the ImageJ (Fiji) 1.54f software<sup>33</sup> (more details in the SI). All variables were expressed as means  $\pm$  standard deviations. Statistical significance between groups was evaluated by a Student's *t*-test. *P*-Value inferior than 0.05 was regarded as statistically significant (*p*-value notations: \*  $< 0.05$ , \*\*  $< 0.01$ , \*\*\*  $< 0.001$ , \*\*\*\*  $< 0.0001$ ).

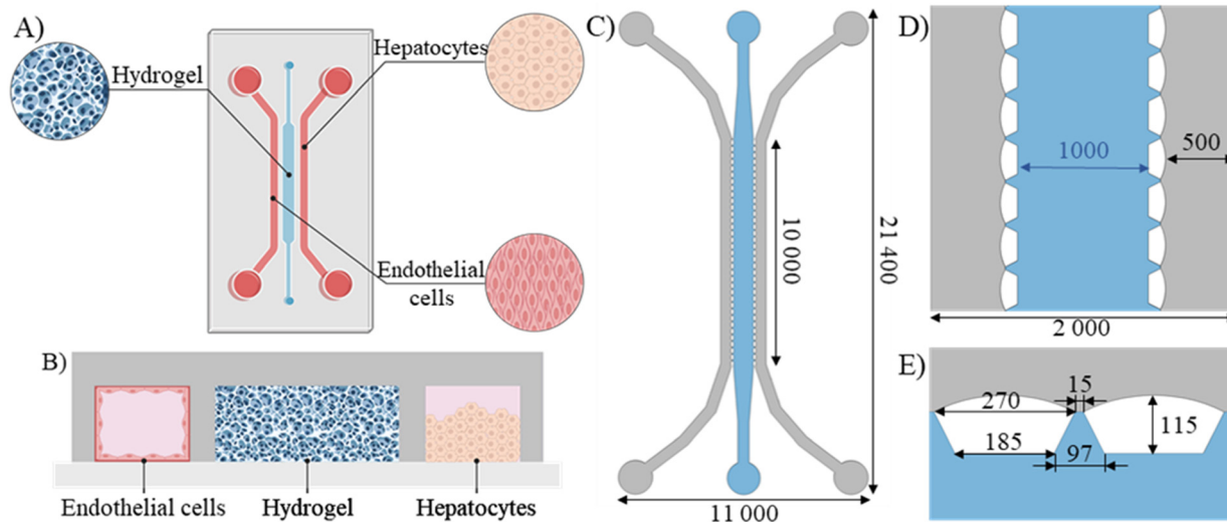
## Results

### Microfluidic chip design

The microfluidic chip was designed to replicate the spatial organisation of sinusoidal cells, hepatocytes and the sD in the liver sinusoid, based on previously published work.<sup>34,35</sup> The device consists of three parallel channels separated by two arrays of 34 polygonal pillars. This chip geometry allows mimicry a layout of the endothelial capillary, hepatocytes and space of Disse in a liver sinusoid (Fig. 1A and B). Hydrogel loaded in the middle channel mechanically mimics the ECM in the sD, while each cell type cultured in one side channel. This enables compartmentalised culture of two cell types through the mediation of a controlled molecular exchange across the middle channel. All channels have a height of  $100\text{ }\mu\text{m}$ . The central effective culturing area of the chip is  $10\text{ }000\text{ }\mu\text{m}$  long and  $2\text{ }000\text{ }\mu\text{m}$  wide (Fig. 1C and D). The hydrogel channel has a width of  $1\text{ }000\text{ }\mu\text{m}$ , while the flanking side channels measure  $500\text{ }\mu\text{m}$  each. Within the arrays, PDMS pillars are spaced  $15\text{ }\mu\text{m}$ , creating a stable retention zone for the hydrogel (Fig. 1E). Models of fluid flow in the chip were developed using finite element simulation (SI).

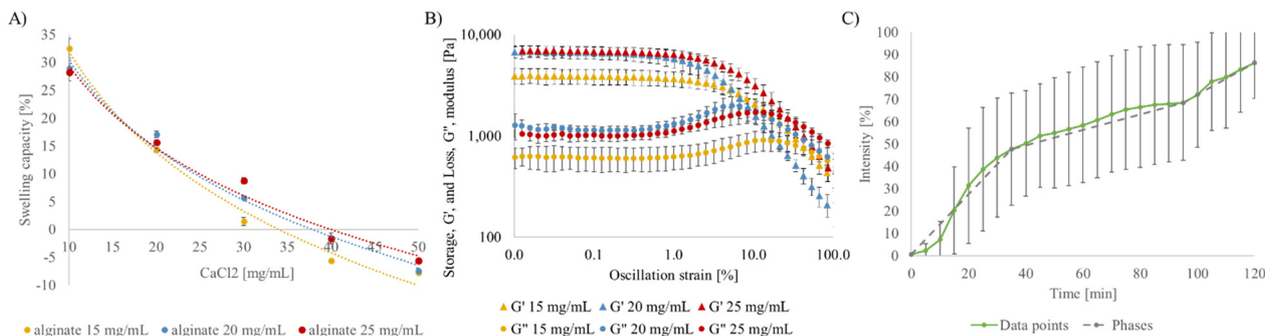
### Hydrogel characterisation

Sodium alginate is a hydrogel widely used in tissue engineering. It was chosen to mimic the sD area on the chip. It is chemically inert, derived from non-animal sources, and offers tuneable mechanical properties. Liquid sodium alginate resin was integrated in the middle channel where PDMS pillars allowed precise loading and retention. Crosslinking with  $\text{CaCl}_2$  through the side channels resulted in the formation of a stable, rigid matrix. The hydrogel remained intact after cell loading in the side channels and 8 days of culture at  $37\text{ }^{\circ}\text{C}$ ,  $5\%$   $\text{CO}_2$  saturation and  $95\%$  humidity conditions.



**Fig. 1** Microfluidic chip design. Schematics of the chip loaded with hydrogel and cells. A) Representation of the layout in the top view, *xy* axis B) representation in the cut view, *xz* axis C) dimensions of the device D) dimensions of the cell culture chamber and E) dimensions of the PDMS pillars [ $\mu\text{m}$ ].





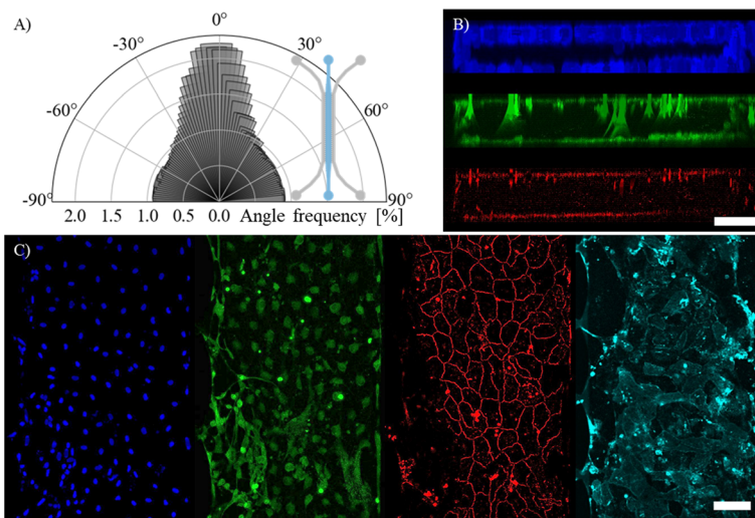
**Fig. 2** Characterisation of sodium alginate gel crosslinked with  $\text{CaCl}_2$ . A) Swelling capacity. B) Rheological testing of gross and loss modulus. C) Molecular transport of CDFA: green line – experimental points, grey line – transport kinetics phases ( $n = 3$  for each data point).

This combination of biocompatibility, structural robustness and controlled permeability encouraged us to choose the sodium alginate as hydrogel for subsequent co-culture experiments. The swelling of sodium alginate hydrogels was systematically evaluated across varying crosslinker  $\text{CaCl}_2$  concentrations to identify the formulation that maintains structural integrity of the hydrogel barrier. An optimal  $\text{CaCl}_2$  concentration of  $30 \text{ mg mL}^{-1}$  was selected, yielding a controlled swelling range of 1.4–7.8%, which ensured that the hydrogel remained stable without swelling in the adjacent channels (Fig. 2A). Rheological testing revealed that increasing alginate concentration enhanced gel stiffness. Young's modulus values varied from  $12 \pm 3 \text{ kPa}$  at  $15 \text{ mg mL}^{-1}$  to  $21 \pm 3 \text{ kPa}$  at  $20\text{--}25 \text{ mg mL}^{-1}$  (crosslinked with  $\text{CaCl}_2$   $30 \text{ mg mL}^{-1}$ ) (Fig. 2B). The  $15 \text{ mg mL}^{-1}$  formulation corresponded most closely to healthy liver tissue, while higher concentrations resembled the stiffer profiles of fibrotic or pathologically remodelled regions.<sup>36</sup> Transport of the small fluorescent molecule across sodium alginate barrier was examined by perfusion on the system with CDFA solution. CDFA molecule was selected for its fluorescent

properties, spherical structure and relatively small molecular mass comparable to glucose (445.21 Da). On-chip assays showed that 80% of total saturation was reached across the hydrogel barrier after 110 min. Molecular transport data points show three distinct phases in the transport kinetics marked with a grey dashed line on the Fig. 2C: initial diffusion-dominated transport (0–35 min), a lag phase (35–95 min) due to barrier resistance, and a new steady-state (95–120 min) after overcoming initial resistance. Slopes of these phases are 1.18, 0.3 and 0.71 respectively.

### Endothelial tube characterisation

LSEC cells were loaded in one of the side channels so that they attach to all channel walls. They were subsequently cultured in direct exposure to the shear stress induced by medium perfusion. Immunofluorescent staining of specific markers was assessed after 8 days of dynamic culture ( $1 \mu\text{L min}^{-1}$ ). F-actin staining revealed cytoskeletal alignment along the axis of medium flow. Quantitative analysis of angular fibre distribution confirmed a significant orientation of LSEC



**Fig. 3** Endothelial tube characterisation. A) Polar plot of cytoskeleton F-actin angular orientation frequency. B) 3D reconstruction of an endothelial tube cross section: blue – DAPI, green – ZO1, red – CD144 (scale bar =  $100 \mu\text{m}$ ). C) Immunofluorescent staining of endothelial specific markers: blue – DAPI, green – ZO1, red – CD144, cyan – CD1 (scale bar =  $500 \mu\text{m}$ ).



cells in the direction of shear stress. As demonstrated in Fig. 3A, the majority of cytoskeletal vectors aligned with the flow direction, with 45.4% of angles distributed within the  $-30^\circ$  to  $+30^\circ$  and 12.7% within the  $-5^\circ$  to  $+5^\circ$  range relative to the channel axis ( $0^\circ$  corresponding to medium flow).

To evaluate tissue identity and barrier integrity, immunofluorescent staining was performed for key endothelial markers. ZO1 and VE-cadherin (CD144) were localised at cell junctions, indicating the establishment of tight junctions essential for the endothelial barrier function. PECAM-1 (CD31) was strongly expressed on the cell membrane confirming differentiation status of the tissue and endothelial-specific adhesion properties (Fig. 3C). Together, these markers confirmed both the differentiation and cohesion of the endothelial sheet along the full length of the channel (full channel immunofluorescent staining in the SI). Three-dimensional reconstruction from z-stack images further demonstrated the formation of a continuous, vessel-like structure. Endothelial cells lined the bottom, top, and vertical walls of the channel, enclosing a well-defined lumen in the middle of the channel (Fig. 3B). This endothelial distribution was confirmed along the full length of the channel.

### Hepatic tissue characterisation

HepaRG cells were seeded in the hepatic channel of the microfluidic chip and cultured under dynamic perfusion ( $1 \mu\text{L min}^{-1}$ ) for 8 days. To characterise the developed tissue, cellular quantification, viability assessment and immunofluorescence staining were performed. Quantification of DNA, RNA, and protein content after cellular extraction indicated an average of 35 ng of DNA, 485 ng of RNA, and 4.37  $\mu\text{g}$  of protein per chip, corresponding to approximately 6000 hepatocytes per device.<sup>37</sup> Live-dead staining revealed high viability of the tissue with  $94 \pm 5\%$  of live cells (chips  $n = 3$ , SI). Immunofluorescent staining confirmed successful differentiation and polarisation of the hepatic tissue. Nuclear expression of HNF4 $\alpha$ , a key transcription factor for hepatocyte development, confirmed differentiation into

functional hepatocytes. Albumin, a hallmark secreted protein of hepatocytes, was detected in the cytoplasm surrounding the nuclei, highlighting active synthetic function. Further evidence of functional polarisation and bile canaliculi formation was provided by membrane expression of BSEP, a bile acid transporter, and MRP2, an efflux pump for organic anions and bilirubin (Fig. 4 and S4).

### Impact of LSEC–HepaRG co-culture on hepatic expression

LSECs have been shown to positively influence hepatocyte function, enhancing the secretion of functional markers such as albumin and urea, while at the same time improving resistance to toxicants.<sup>8,9,38</sup> To investigate these effects in our model, HepaRG cells were cultured either as monocultures or in co-culture with LSECs. Hepatic functionality was then evaluated after the culture period based on three approaches: quantification of key liver biomarkers, characterisation of hepatocyte polarisation, and probe-based assessment of protein expression.

Liver biomarkers, albumin and urea, were evaluated through ELISA quantification from the medium perfused through the chip. Albumin secretion remained stable in both monoculture and co-culture conditions, but levels were consistently higher in the LSEC–HepaRG co-culture. Indeed, average albumin production reached  $1248 \pm 214$  ng per chip in co-culture compared with  $392 \pm 168$  ng in monoculture ( $p < 0.0001$ ). Urea secretion followed the same trend, with monoculture chips secretion at  $7.84 \times 10^{-2} \pm 3.42 \times 10^{-4}$  nmol compared with  $1.69 \times 10^{-1} \pm 1.83 \times 10^{-3}$  nmol in co-culture chips. These values indicate more than a twofold increase in urea production consistent throughout the whole culture duration (Fig. 5A and B). These results demonstrate a consistent enhancement of hepatocyte metabolic activity in the presence of LSECs. Bile canaliculi formation was assessed using the live fluorescent probe Fluobile™, which marks functional canalicular networks in hepatocytes. After 8 days of culture, quantification revealed a significantly higher number and a larger total fluorescent area of bile canaliculi in LSEC–HepaRG co-culture compared to monoculture

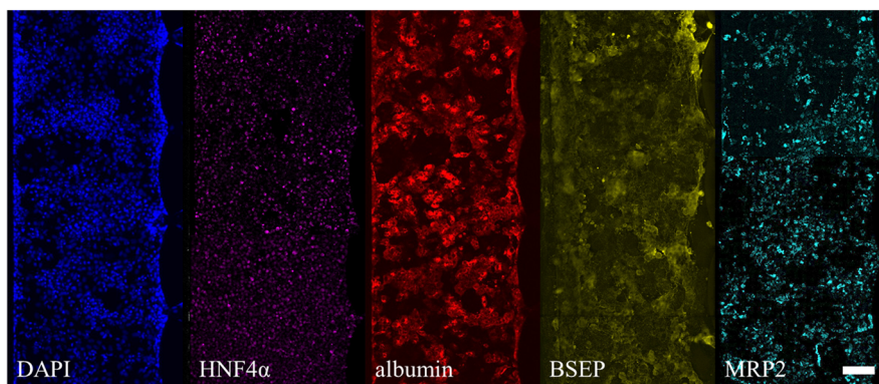
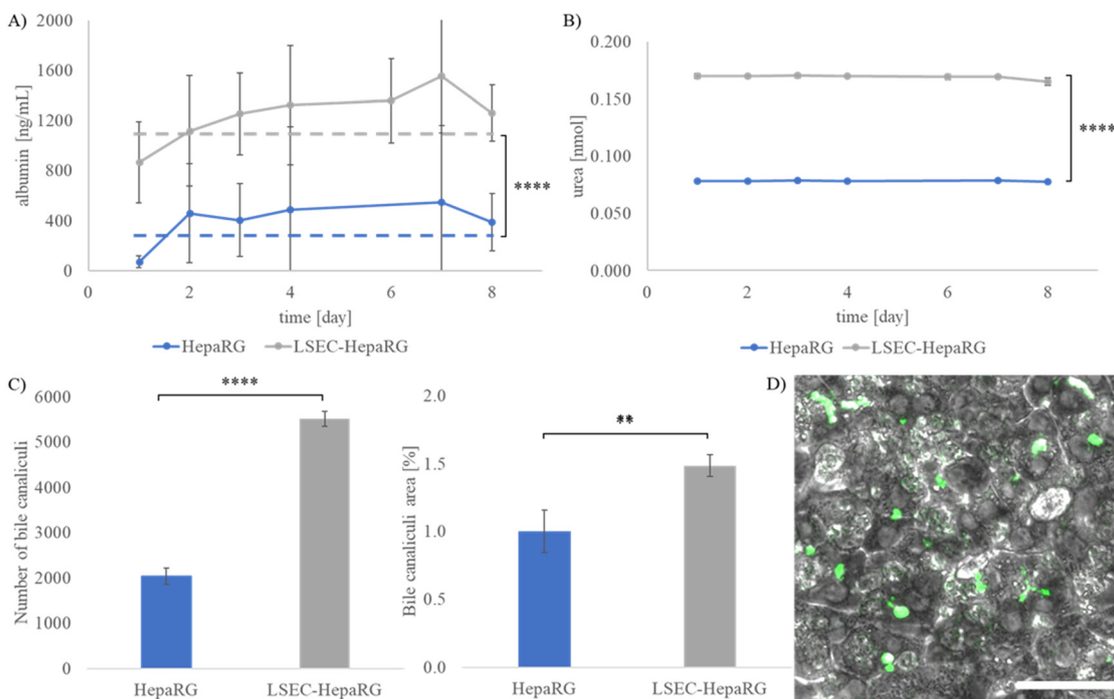


Fig. 4 Immunofluorescent staining for hepatocyte specific markers of HepaRG cells on the chip. Blue – DAPI, magenta – HNF4 $\alpha$ , red – albumin, yellow – BSEP, cyan – MRP2 (scale bar = 100  $\mu\text{m}$ ).





**Fig. 5** Characterisation of LSEC-HepaRG co-culture effect on hepatic functionality. A) Albumin and B) urea levels quantification during 8 days of co-culture (\*\*\*\* $p < 0.0001$ ). C) Hepatic polarity evaluation through quantification of number and surface area of bile canaliculi (\*\* $p < 0.001$ ; \*\*\*\* $p < 0.0001$ ). D) Image of hepatocytes stained with Fluobile: green – bile canaliculi (scale bar = 100  $\mu\text{m}$ ;  $n = 3$  for each data point).

(Fig. 5C). High-magnification imaging further showed that canaliculi formation preferentially occurred in regions of increased cellular density and within 3D structures, often extending along the membranes of adjacent hepatocytes (Fig. 5D). These observations indicate that endothelial co-culture promotes hepatocyte polarisation and the organisation of functional bile canaliculi.

To evaluate the impact of the co-culture on the hepatic secretome, proteomic profiling was performed using the Olink Target 96 Inflammation panel and analysed on  $\log_2$ -transformed NPX values.

Hierarchical clustering of  $z$ -scored NPX values (Fig. 6A) revealed a clear separation between HepaRG monocultures and LSEC-HepaRG co-cultures: co-culture samples clustered together and showed a broad, coordinated increase in signal intensity across the pre-selected analytes (selection criterion:  $p < 0.05$ ; full NPX table in the SI).

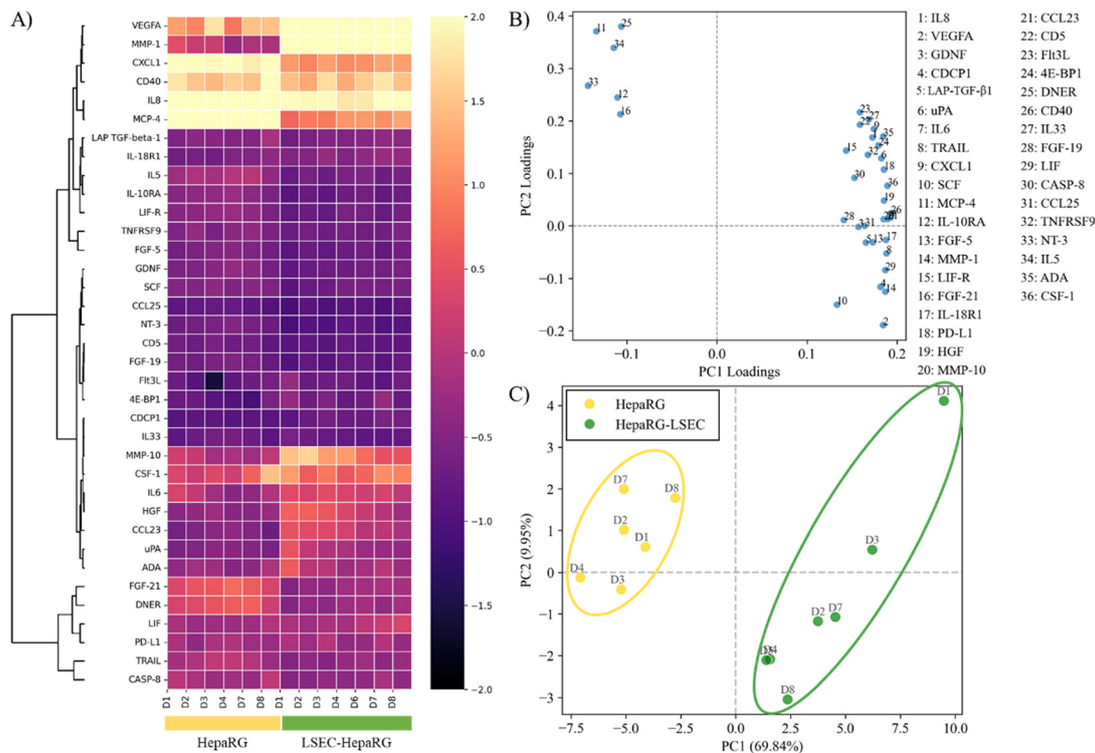
The PCA loading plot identified the analytes that most strongly contributed to sample separation. The cluster of proteins positioned on the far right of the plot – including VEGFA, CD40, MMP-1, and CSF-1 – were the major drivers along PC1, reflecting the distinction between monoculture and co-culture conditions. In contrast, proteins in the cluster located on the upper left of the plot, predominantly DNER, MCP-4, and IL-5, accounted for within-group variability along PC2, representing secondary axes of separation. Finally, principal component analysis showed robust separation of conditions along PC1 ( $\approx 70\%$  variance explained), with co-culture samples projecting positively on PC1 and

monocultures projecting negatively (Fig. 6C). Together, these results indicate that inclusion of LSECs substantially remodelled the hepatic secretome, yielding a reproducible protein signature dominated by growth/regenerative and inflammatory/remodelling signals.

### Toxicologic assay

Due to enhanced performance of the model in terms of hepatic function and polarity, the model containing LSEC-HepaRG co-culture was chosen for hepatotoxicity assay. Hepatic viability and metabolic activity were assessed under prolonged 2 mM acetaminophen (APAP) treatment by live mitochondrial and CYP3A4 probe staining. Image analysis of fluorescent MitoTracker staining images, revealed a pronounced reduction in mitochondrial activity following APAP exposure, with APAP-treated samples showing a  $77.2 \pm 5.6\%$  decrease in live mitochondrial signal compared to controls (Fig. 7A). This indicates severe impairment of mitochondrial function, consistent with oxidative stress-induced toxicity. CYP3A4 activity was also significantly affected by APAP exposure. In control tissues, CYP3A4 signal covered  $9.71 \pm 4.71\%$  of the culture area, while in APAP-treated samples it increased to  $22.81 \pm 4.93\%$  (Fig. 7B), reflecting a stress-induced alteration of drug metabolism pathways. Notably, the APAP-treated hepatic tissue displayed uneven signal distribution, fragmentation, and localised shrinkage, consistent with hepatocyte damage and detachment. These morphological alterations were





**Fig. 6** Hepatic secretome evaluation in conditions of mono- (yellow) and co-culture (green). A) Heatmap of z-scored NPX values: columns = days of culture; rows = analytes ordered by hierarchical clustering. B) PCA loading plot showing the contribution of individual analytes to PC1 and PC2. C) Principal component analysis of log<sub>2</sub>-transformed NPX value.

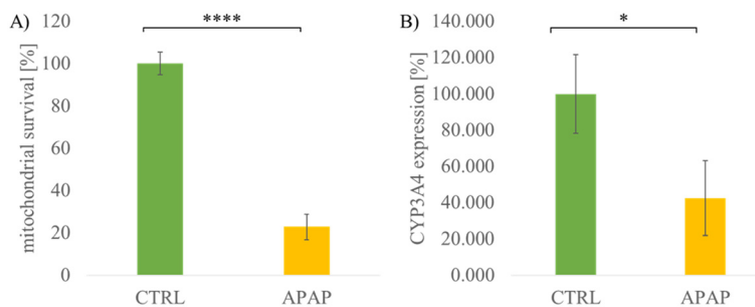
consistent with hallmark features of APAP-induced hepatocyte injury reported in previous studies.<sup>39</sup>

## Discussion

This study presents the development of a microfluidic chip used for biomimetic reproduction of the architecture and cellular interactions in a sD. The platform integrated a sodium alginate hydrogel with easily tuneable mechanical properties as an ECM analogue, enabling compartmentalised dynamic co-culture of HepaRG and LSECs. HepaRG cells were intentionally selected as a robust and well-characterised hepatic model that offers a practical compromise between physiological relevance and

experimental reproducibility for proof-of-concept microphysiological systems. Their use enabled stable long-term co-culture under dynamic conditions while minimising donor-to-donor variability and experimental complexity associated with primary human hepatocytes. The system maintained high viability and structural integrity of both tissues for at least 8 days of dynamic culture, supporting stable hepatic function and endothelial organisation. Together, these results showed the establishment of a physiological model capable of sustaining endothelial-hepatic interactions in a controlled, physiologically relevant environment.

The microfluidic design of the device directly influenced its functionality. The three parallel channel configuration



**Fig. 7** Markers of hepatic viability and function in toxicity conditions. A) Live mitochondria (MitoTracker staining, \* $p < 0.05$ ) and B) CYP3A4 expression (P450 Glo CYP3A4, \*\*\*\* $p < 0.0001$ ).



allowed spatial separation of cell types while allowing diffusional exchange through a hydrogel-filled central compartment that mimics native sD. Although the absolute width of the hydrogel channel (1000  $\mu\text{m}$ ) exceeds the *in vivo* thickness of the sD (approximately 1  $\mu\text{m}$ ), this model reproduced its physiological role as a mechanical and biochemical communication interface, while preserving structural hierarchy of the liver sinusoid. Hepatocytes and endothelial cells were allowed intercellular communication while in indirect contact. Furthermore, the design allowed a special set of conditions, such as shear stress exposure, to be applied to LSEC cells, thus closely mimicking *in vivo* situation. Current microfluidic models replicate the sD by integrating microchannels called slits, scaffolds or membranes.<sup>25,40</sup> These structures often fail to reproduce its essential mechanical characteristics, such as physiological elasticity, correct shear stress exposure and controlled porosity. The use of PDMS pillars in our chip ensured mechanical retention of the hydrogel and sustained perfusion without noticeable effect on the gel positioning within the chip. To do so, the microchannel dimensions were optimised to maintain stable hydrogel confinement and laminar flow across physiologic shear ranges.

Hydrogel optimisation was as important for the model's biomimetic performance as was the microfluidic chip design. In the majority of liver-on-a-chip models in the literature, the hydrogel of choice was typically collagen at high concentrations. These models also rely on direct embedding of hepatocytes within collagen, which often causes fibrotic phenotype due to biochemical cues of excessive ECM.<sup>41</sup> In contrast, the alginate hydrogel used here was chosen for its chemical inertness, non-animal origin, and mechanical tunability. Its porous structure enabled passive diffusion of signalling molecules while avoiding active biochemical interaction with hepatocyte or endothelial-secreted molecules, thereby preserving native intercellular communication. Crosslinking parameters were optimised to yield a Young's modulus of approximately 12 kPa, corresponding to the physiological stiffness of healthy human liver tissue.<sup>36</sup> Thereby, the non-physiological stiffening associated with fibrosis was avoided, and this allowed mechanical but not biochemical signalling from the matrix to drive cell behaviour. The hydrogel also demonstrated structural robustness and controlled permeability, with the approximate porosity of 30% estimated from tracking small-molecule transport across the barrier, thus closely resembling diffusional dynamics expected across the native sD.

Comprehensive characterisation of both endothelial and hepatic tissues confirmed successful tissue-specific organisation and functionality. Under continuous flow, LSECs aligned their actin cytoskeleton along the direction of shear stress, indicating flow-induced polarisation. While a uniform distribution of orientations would yield approximately 33.4% of vectors within  $-30^\circ$  to  $+30^\circ$  and 5.6% within  $-5^\circ$  to  $+5^\circ$ , our data showed 45.4% and 12.7%, respectively ( $p = 0.001$ ), demonstrating a significant enrichment of alignment along

the flow axis. Immunofluorescent staining for endothelial junctional markers ZO1, VE-cadherin, and PECAM-1 demonstrated the establishment of a cohesive and differentiated endothelial layer forming a tube with a lumen. The HepaRG tissue exhibited high viability ( $94 \pm 5\%$ ) and strong expression of hepatocyte-specific markers, including HNF4 $\alpha$  and albumin, confirming hepatic differentiation. The presence of apical transporters (BSEP, MRP2) indicated polarisation and bile canaliculi formation. Together, these observations demonstrated that the microfluidic environment and matrix mechanics supported physiologically relevant cell organisation and function.

Beyond single-tissue characterisation, the co-culturing of HepaRG cells with LSECs substantially enhanced hepatic performance. Co-culture resulted in a threefold increase in albumin secretion and a twofold elevation in urea synthesis compared to monocultures, consistent with previous reports of endothelial-mediated support of hepatocyte function. The presence of LSECs also promoted hepatocyte polarisation, reflected by a higher number and surface area of functional bile canaliculi, which is consistent with earlier studies.<sup>8,9,38</sup> It is likely that the enhanced hepatic functionality in the LSEC–HepaRG co-culture arises from multiple complementary mechanisms, including paracrine secretion of hepatocyte-supportive growth factors and cytokines, improved nutrient and waste exchange mediated by the endothelial layer, and direct intercellular communication through physical and biochemical signalling pathways that collectively promote polarisation and metabolic stability. Secretome profiling further revealed that co-culture triggered a coordinated remodelling of the molecular microenvironment. Biological interpretation of the analytes showed that expressed proteins were grouped into six groups based on cellular function (Table 1). Principal component and hierarchical clustering analyses showed clear separation between monoculture and co-culture samples, driven by distinct groups of proteins associated with angiogenesis, immune modulation, and extracellular matrix remodelling. Upregulation of factors such as VEGFA, HGF, SCF, and Flt3L likely contributed to enhanced hepatocyte viability and metabolic output, while increased secretion of cytokines including IL-6, IL-8, and CXCL1 suggested activation of regenerative signalling cascades. Concurrent elevation of MMP-1, MMP-10, and LAP TGF- $\beta$ 1 indicates ongoing ECM remodelling and possible low-grade activation of wound-healing pathways.<sup>42</sup> In conclusion, our secretome screening revealed activation of both pro-regenerative and pro-remodelling components, maintained through endothelial–hepatocyte crosstalk. Further application of this analytical approach should enable the identification of LSEC secreted biomarkers to determine the key signalling molecular triggers in this intercellular communication.

To demonstrate the functional relevance of the system for drug testing, a prolonged hepatotoxicity assay using acetaminophen was performed. APAP was selected as a



**Table 1** Six cellular functions of analytes with corresponding proteins

| Cellular functions                   | Analytes  |
|--------------------------------------|---|
| Angiogenesis and hepatocyte survival | VEGFA, HGF, SCF, Flt3L  |
| Inflammatory and immune response     | IL-6, IL-8, IL-10RA, CXCL1, CCL23, MCP-4, CSF-1, PD-L1, IL-18R1 |
| ECM remodelling and fibrosis         | MMP-1, MMP-10, LAP TGF- $\beta$ 1                               |
| Hepatic regeneration factors         | FGF-5, FGF-19, FGF-21, LIF, LIF-R                               |
| Apoptosis and survival regulation    | TRAIL, CASP-8   |
| Metabolic/immune modulators          | IL-33, ADA, NT-3  |

reference compound due to its established role as the gold standard in hepatotoxicity screening, with extensively characterised molecular mechanisms. Following 2 mM APAP exposure from day 5 to day 8, MitoTracker staining revealed a 77% reduction in live mitochondrial signal, directly indicating reduction in cell viability. This observation aligns with the known mechanism of APAP toxicity mediated by oxidative stress-induced mitochondrial dysfunction through the reactive metabolite NAPQI, which depletes intracellular glutathione and disrupts mitochondrial respiration.<sup>43</sup> CYP3A4 activity, assessed *via* live luminescent assay, also decreased markedly after treatment, indicating impaired hepatic metabolism. Morphological assessment confirmed tissue shrinkage and detachment; features typical of hepatocyte death. Together, these results validate the capacity of the LSEC–HepARG co-culture model to detect functional and metabolic hallmarks of drug-induced hepatotoxicity under 72-hour exposure conditions, thus serving as a proof of concept for DILI modelling.

Overall, this study introduces a liver-on-a-chip model that combines structural, mechanical, and biochemical fidelity to the native hepatic microenvironment. The integration of a tuneable alginate hydrogel enables reliable mimicking of sD mechanical properties without fibrotic phenotype induction. The model reproduces key structural and functional characteristics of the liver sinusoid, including endothelial cells organised in a vessel-like structure, hepatocyte polarisation, bile canaliculi formation, and responsiveness to drug induced toxicity. While APAP was used here for model validation, future investigation will focus on screening additional drug classes with distinct hepatotoxic mechanisms.

## Conclusion

We have developed and characterised a novel *in vitro* liver-on-a-chip platform focused on modelling of the sD. The system integrates a biocompatible sodium alginate hydrogel whose mechanical properties are optimised to closely mimic those of physiological human liver tissue. The model successfully maintains the viability and functionality of both LSECs and HepARG cells under dynamic co-culture conditions for 8 days, supporting tissue organisation and functional polarisation. Co-culturing of HepARG with LSECs significantly enhances hepatic performance, leading to increased albumin and urea secretion, improved bile canaliculi formation, and secretion of pro-regenerative and remodelling factors. The predictive capacity of

the model is further confirmed through acetaminophen exposure. Future perspectives will focus on the development of the model in two main directions. First, addition of Kupffer and stellate cells, which would increase the predictive power and reliability of the platform in conditions such as chronic inflammation. Second, screening a library of multiple drug classes and characterising the model's response in reaction to them. Altogether, this work establishes a robust and human-relevant microphysiological system for studying endothelial–hepatic interactions and drug-induced liver injury.

## Author contributions

Conceptualisation, A. M. and A. M.; methodology, A. M., A. M., Z. T. and S. M. I.; validation, A. M.; formal analysis, A. M. and S. M. I.; investigation, A. M.; resources, A. M.; data curation, A. M.; software, A. M.; writing – original draft preparation, A. M.; writing – review and editing, A. M., A. M., S. B., J. C. D. V. and B. L. P.; visualisation, A. M.; supervision, A. M., S. B., J. C. D. V. and B. L. P.; funding acquisition, J. C. D. V. and B. L. P.; project administration, A. M., S. B., J. C. D. V. and B. L. P. All authors have read and agreed to the published version of the manuscript.

## Conflicts of interest

There are no conflicts of interest to declare.

## Data availability

The data supporting this article have been included as part of the supplementary information (SI): a detailed protocol for biological characterisation and image analysis; immunofluorescent images for LSEC alignment quantification, images of hepatic tissue viability staining, NPX values table of secretome.

Supplementary information is available. See DOI: <https://doi.org/10.1039/d5lc01139f>.

## Acknowledgements

This work was supported by ENS Paris Saclay, INSERM U1193 and University Paris Saclay, Faculty of Pharmacy and ANR-21-CE-19-0025 DILI-on-chip project. AM's PhD fellowship and AM's researcher position were funded by ANR (Agence Nationale de Recherche) program DILI-on-chip through a project ANR-21-CE-19-0025.



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