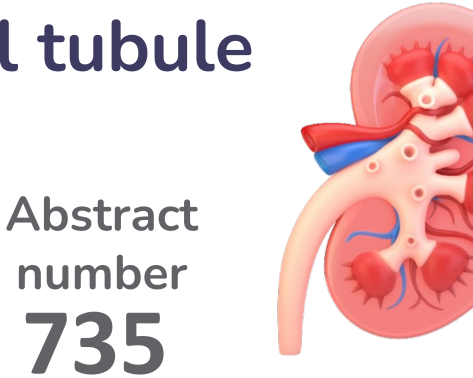


Development of highly-differentiated human primary proximal tubule MPS model (aProximate MPSTM Flow)

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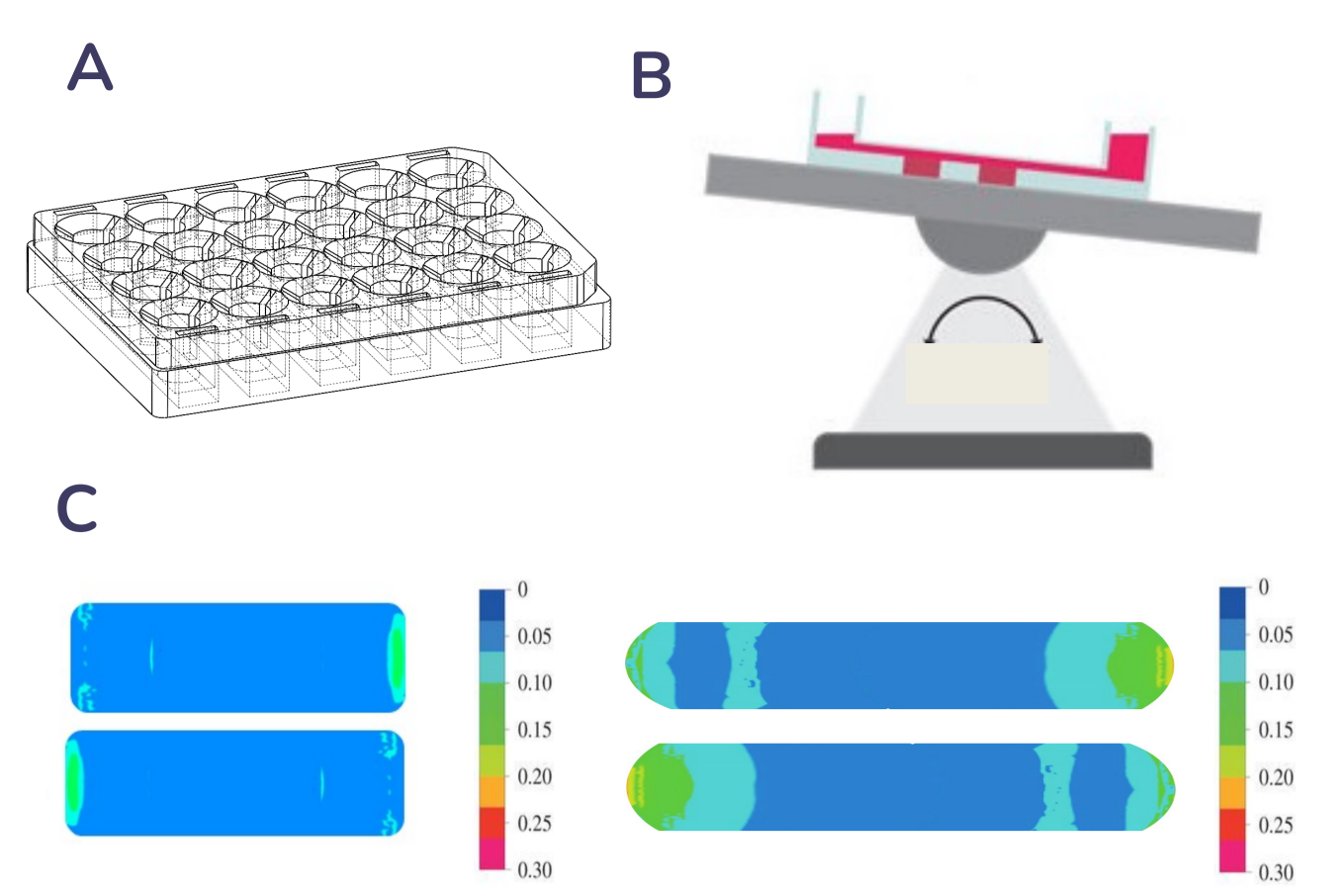
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Introduction

The proximal tubule (PT) is the key nephron segment mediating renal drug elimination and is the primary site of drug induced nephrotoxicity. However, current animal studies have proved poorly predictive of human outcome. To address this, there has been a recent upsurge in physiological relevant microphysiological systems (MPS) of PT to recapitulate differentiation and function in vitro. Here we present results from our recently developed aProximate MPS[™] human PT platform (Patent No: G001336.GB), in which primary human PT cells are subject to fluidic media flow and a shear stress between 0.1-2 dynes/cm².

Methods

Computational Fluid Dynamics (CFD) was used optimise to uniformity of the shear stress within the flow chambers. Using CAD software and 3D printing, we were able to rapidly prototype different plates with different determine format to optimal for primary cells performance within the flowplate system. Primary PT cells were seeded on underside Corning the of ThinCert™ and incubated overnight. After flipping, the flowplate was placed on a rocking platform at angle θ allowing passive liquid levelling creating lateral flow without the use of a pumping system.





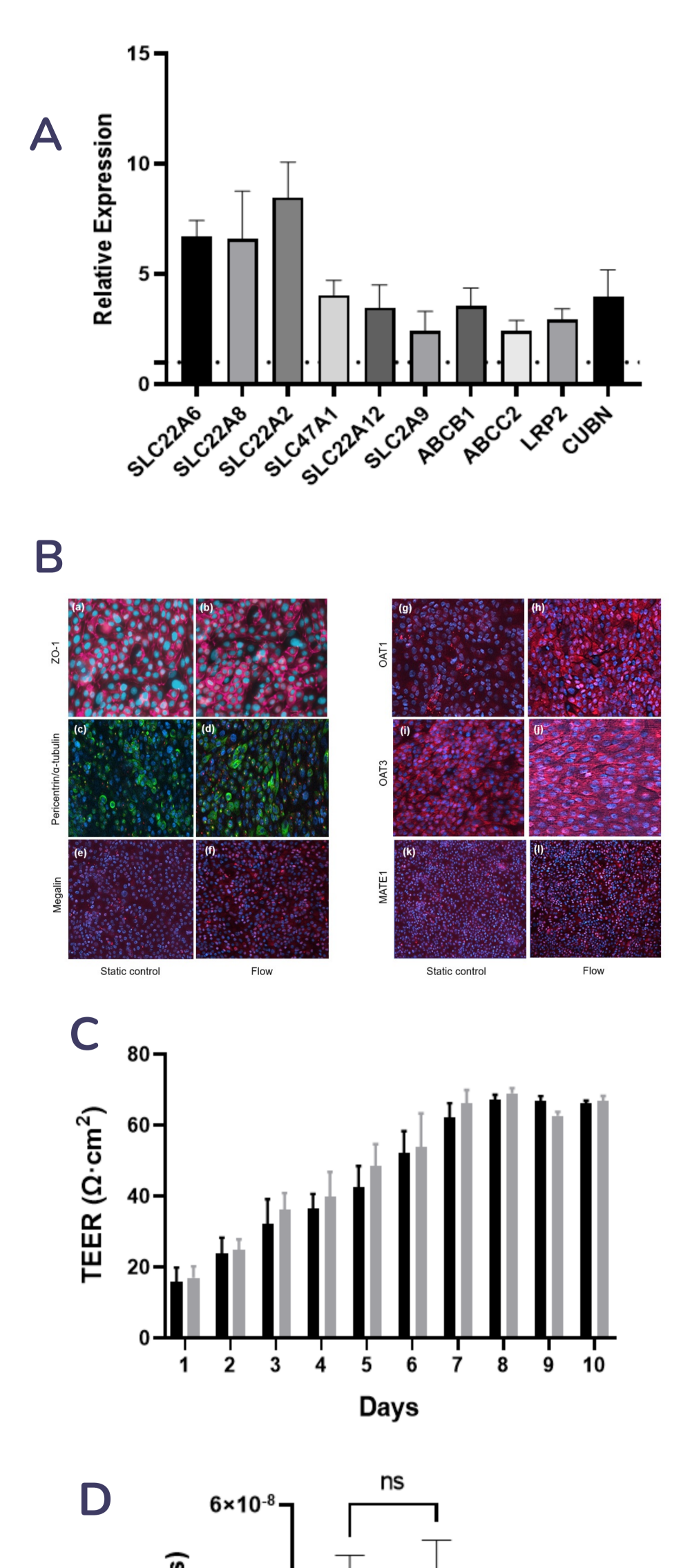
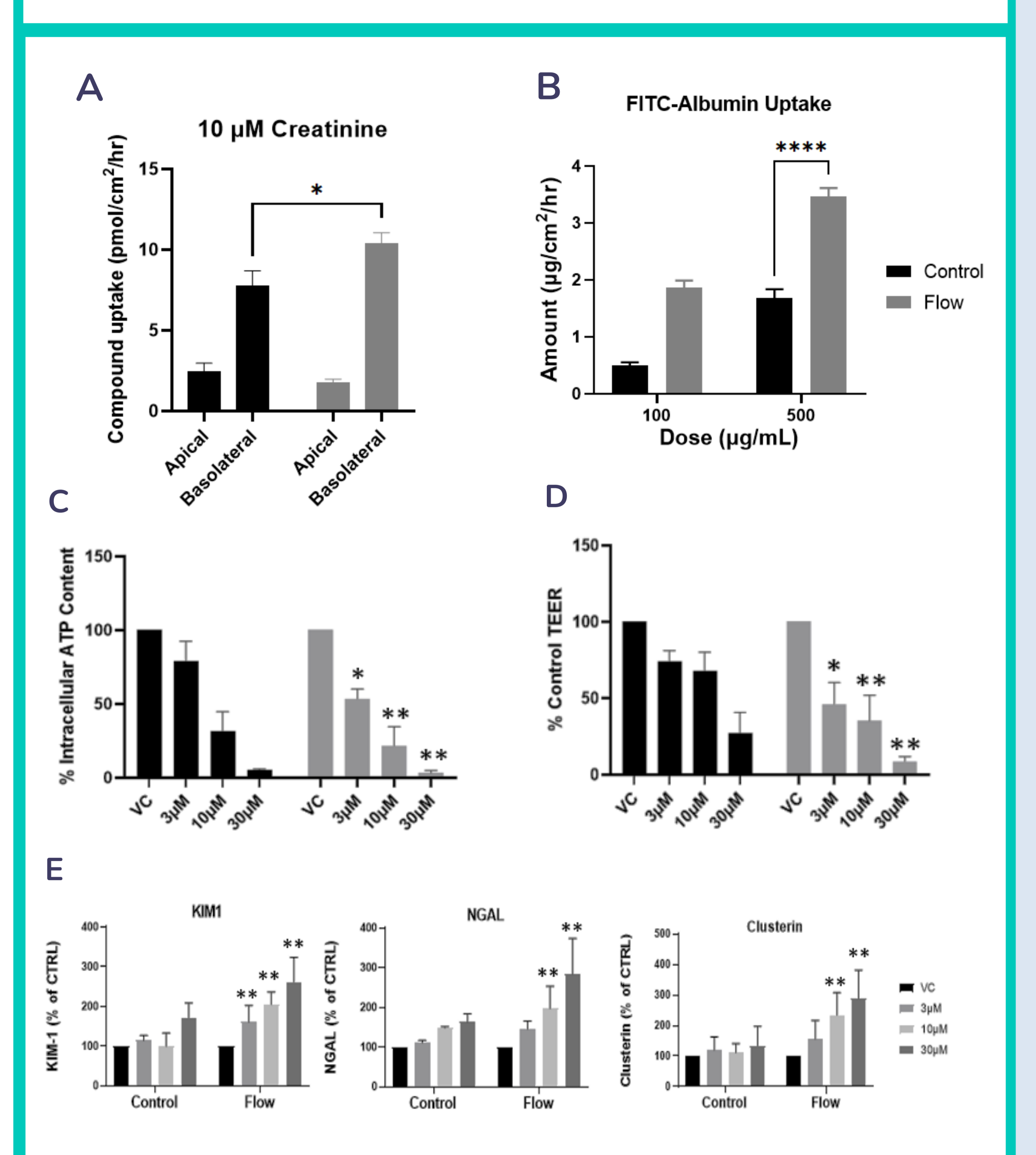
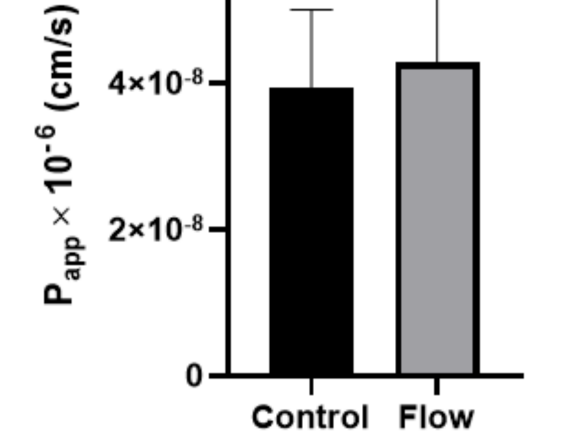


Figure 1. Design of aProximate MPSTM Flow system. A) Plate design in 24-well format, **B)** function of flowplate, **C)** CFD simulations of media flow within the channel for the 24-well connected and 24-well single channel.





Characterisation Figure function and barrier 2. assessment. A) qPCR analyses of key transporters at D7 (n=5); **B)** IF images of key proteins in human PT cells at D7; C) TEER measured from D1-D7; D) P_{app} measured using Lucifer Yellow up to 180 minutes (n=6).

Figure 5: Functional Assays on aProximate MPS[™] Flow: A) Creatinine uptake at 90min;; B) FITC-Albumin Uptake at 1hour with large dose range; Nephrotoxicity study carried out using cisplatin treatment (72hours) on stimulated cells vs control C) ATP Release measured at 72hrs following treatment. D) TEER measured at 72hrs following treatment E) Renal Injury Biomarker Release measured at 72hrs following treatment using MSD; (n=3)

Conclusions

This dataset, suggests that growing human PT cells on ThinCert[™] with media flow across the apical membrane, significantly improves phenotype and function and has significant benefit to the utility and near-physiology of the model.