Non-combustible Next Generation Products induce lower biological activity than combustible tobacco on a human cardiovascular model on-a-chip

SOT 62nd Annual Meeting March 19-23, 2023 Abstract 3459/ P583

Fiona Chapman^{1*}, Luuk de Haan², Linda Gijzen², Edgar Trelles Sticken³, Roman Wieczorek³, Sarah Jean Pour³, Liam Simms¹, Matthew Stevenson¹ ¹Imperial Brands PLC, 121 Winterstoke Road, Bristol, BS3 2LL, UK; ²Mimetas B.V., Biopartner Building 5, De Limes 7, 2342 DH Oegstgeest, the Netherlands; ³Reemtsma Cigarettenfabriken GmbH, An Imperial Brands PLC company, Albert-Einstein-Ring-7, D-22761, Hamburg, Germany *Corresponding author's e-mail: Fiona.Chapman@uk.imptob.com

INTRODUCTION

Cigarette smoking is a cause of serious diseases in smokers, including heart diseases. Atherosclerosis progresses through a pathway of endothelial dysfunction, lipid infiltration, macrophage recruitment and vascular remodelling, and can be driven by exposure to exogenous agents such as some of the harmful compounds (HPHCs) present in cigarette smoke. However, the development of next generation nicotine delivery products (NGPs), such as electronic nicotine delivery systems (ENDS) and heated tobacco products (HTPs), offers adult smokers, uninterested or unwilling to quit smoking, potentially reduced harm alternatives to continued cigarette smoking, potentially reduced harm alternatives to continued cigarette smoking. levels of HPHCs compared to cigarettes [1, 2].

AIM

Using the OrganoPlate[®] 2-lane chip (Mimetas BV) (Figure 1) [3], this study aimed to assess the impact of exposure to cigeratte smoke and NGP aerosol extracts on a Human Coronary Artery Endothelial Cell (HCAEC) and THP-1 monocyte co-culture, to model and assess early (inflammatory) processes involved in atherosclerosis development [4]

METHODS

Test Articles

- 1R6F Reference Cigarette (University of Kentucky)
- NGP: HTP, Pulze with iD stick (Balanced Tobacco)
- NGP: ENDS product, myblu EU Tobacco 1.6% Nicotine

Smoke/ aerosol extract generation

Smoke/ NGP aerosol was generated using a VITROCELL VC-10-S smoking machine (VITROCELL Systems GmbH). Smoke/ NGP aerosol was bubbled through three in-line impingers each containing 10ml (combined final stock volume: 30ml) of phosphate buffered saline (PBS) solution to generate extracts for addition to the experimental model (Figure 2). 1R6F stock concentration: 1.8 puffs/ml; NGP stock concentrations: 4.8 puffs/ml.



Figure 2: Smoke/ aerosol extraction schematic (e-cigarette = ENDS)

Nicotine and select carbonyls were quantified within the smoke/ aerosol bubbled PBS (bPBS) samples (Table 2). Nicotine was quantified using liquid chromatography with tandem mass spectrometry (LC-MS/MS) with an AB Sciex API 6500 QTRAP (SCIEX) using nicotine-d4 as the internal standard. For the analysis of carbonyls, bPBS samples were diluted with 2,4-dinitrophenylhydrazine (DNPH) The carbonyl-DNPH derivates were then quantified using high performance liquid chromatography with a diode-array detector (HPLC-DAD; Agilent Technologies 1100 Series). Table 1 details the subsequent in vitro testing methodology.

Table 1: Overview of the *in vitro* testing methodology

Day -1	Collagen I (Cultrex) extracellular matrix (ECM) gel seeded into OrganoPlate
Day 0	HCAECs seeded onto ECM gel
Day 3	Test article extracts added to cell culture medium (containing THP-1 cells) (pre-conditioning) (negative control = PBS)
Day 4	OrganoPlate culture exposed to: - Preconditioned medium - Test article extracts (compound only controls) - Positive control, TNFα - Positive control (glutathione (GSH) assay), ethacrynic acid
	4h analysis: Glutathione (GSH) assessment (monochlorobimane added to cultures)
	Fresh THP-1 monocytes added for adhesion assay
Day 5	24h analyses: ICAM-1 expression (immunofluorescent readout) Monocyte adhesion (immunofluorescent readout) Inflammatory mediators (in medium samples)

Data analyses

REFERENCES

Nine chips/test condition were used across 2 biological replicates (n=9, N=2). Data was normalised to respective controls for visualisation, and error bars plotted were standard deviation (SD) about the mean. Statistical significance was determined using a one-way analysis of variance (ANOVA). Statistical significance is marked as follows: p < 0.05, p < 0.01, p < 0.001 and p < 0.0001.

- CONCLUSIONS
- reduced risk potential of NGPs [2]





Medium preconditioning (with THP-1 monocytes and the test article extracts) combined with the HCAECs elicited clear dose- and test article-dependent responses. Inclusion of compound only controls confirmed the effect of preconditioning Across the endpoints assessed, 1R6F elicited the greatest changes, when compared on a nicotine concentration basis. The outcomes reflect the proposed relative risk (of exposure to toxicants) of the test articles under the conditions of the test: 1R6F >> HTP > ENDS and add to the weight of evidence of the

- The results suggest that cigarette has the greater potential effect on atherosclerotic processes compared to the NGPs. Future work will involve the addition of further endpoints related to atherosclerotic processes including cell migration

[1] Rudd, K., et al. (2020). Chemical Composition and In Vitro Toxicity Profile of a Pod-Based E-Cigarette Aerosol Compared to Cigarette Smoke. AIVT, https://doi.org/10.1089/aivt.2019.0015. [2] Chapman, F., et al. (2023). Multiple endpoint in vitro toxicity assessment of a prototype heated tobacco product indicates substantially reduced effects compared to those of combustible cigarette.



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Figure 6: Heatmap of Inflammatory cytokine levels (log 2 transformed; normalised to respective control values) following exposure of the model to

[4] Poussin, C., et al. (2020). 3D human microvessel-on-a-chip model for studying monocyte-to-endothelium adhesion under flow – application in systems toxicology. ALTEX, https://doi.org/10.14573/altex.1811301. [5] Simms, L, et al. (2022). Use of Human Induced Pluripotent Stem Cell-Derived Cardiomyocytes to Predict the Cardiotoxicity Potential of Next Generation Nicotine Products. Front Toxicol, https://doi.org/10.3389/ftox.2022.747508.

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Figure 1: OrganoPlate[®] 2-lane and the cell model used in this study

