3D-Printed Franz cells – update on optimization of manufacture and evaluation

B. C. Sil*, R. G. Belgrave†, M. P. Alvarez‡, L. Luo‡, M. Cristofoli*†, M. R. Penny‡, D. J. Moore§, J. Hadgraft†, S. T. Hilton§, and M. E. Lane‡

*London Metropolitan University, 166-220 Holloway Road, London N7 8DB, †UCL Chemical Engineering, Torrington Place, London WC1E 7JE, ‡UCL School of Pharmacy, 29-39 Brunswick Square, London WC1N 1AX and §GSK Consumer Healthcare, Skin Health R&D, St George’s Ave, Weybridge KT13 0DE, UK

Received 4 March 2020, Accepted 26 March 2020

Keywords: 3D printing, Franz cells, SEM analysis, In vitro, permeation studies

Abstract
OBJECTIVES: Laboratory in vitro permeation processes require the use of modified Franz type diffusion cells which are conventionally fabricated from glass. Fragility and high cost are frequently associated with this type of laboratory apparatus. The purpose of our present research was to develop a simple, economical and versatile approach to manufacture Franz type cells using additive manufacturing (AM).

METHODS: Graphical Franz diffusion cell designs were reproduced with a stereolithography (SLA) 3D printer and assessed over a minimum period of 24 h. The surface morphology of AM printouts was analysed before and after compatibility studies using scanning electron microscopy (SEM). Comparative permeation studies in both glass and AM Franz type diffusion cells were conducted using a caffeine solution (1.5 mg mL⁻¹), applied to a model silicone membrane.

RESULTS: Testing of the 3D printed scaffolds confirmed similar recovery of the permeant when compared to glass cells: 1.49 ± 0.01 and 1.50 ± 0.01 mg mL⁻¹, respectively, after 72 h. No significant differences were visible from the SEM micrographs demonstrating consistent, smooth and non-porous surfaces of the AM Franz cells’ core structure. Permeation studies using transparent 3D printed constructs resulted in 12.85 ± 0.53 µg cm⁻² caffeine recovery in the receptor solution after 180 min with comparable permeant recovery, 11.49 ± 1.04 µg cm⁻², for the glass homologues.

CONCLUSION: AM constructs can be considered as viable alternatives to the use of conventional glass apparatus offering a simple, reproducible and cost-effective method of replicating specialised laboratory glassware. A wider range of permeants will be investigated in future studies with these novel 3D printed Franz diffusion cells.

Résumé
OBJECTIF: les processus de perméation in vitro en laboratoire nécessitent l’utilisation de cellules de diffusion de type Franz modifiées, fabriquées traditionnellement en verre. La fragilité et un coût élevé sont fréquemment associés à ce type d’appareil de laboratoire. L’objectif de nos travaux de recherche actuels était de développer une approche simple, économique et polyvalente pour fabriquer des cellules de type Franz à l’aide de la fabrication additive (FA).

MÉTHODES: les conceptions des cellules de diffusion Franz graphiques ont été reproduites avec une imprimerie 3D stéréolithographie (SLA) et évaluées sur une période minimum de 24 h. La morphologie de surface des impressions FA a été analysée avant et après des études de compatibilité à l’aide de la microscopie électronique à balayage (MEB). Des études comparatives de perméation des cellules de diffusion de type Franz en verre et FA ont été réalisées à l’aide d’une solution de caféine (1.5 mg ml⁻¹) appliquée à un modèle de membrane en silicone.

RÉSULTATS: les tests des supports imprimés 3D ont confirmé une récupération similaire du perméant par rapport aux cellules de verre : 1,49 ± 0,01 et 1,50 ± 0,01 mg ml⁻¹, respectivement, après 72 h. Aucune différence significative n’a été observée sur les micrographiques MEB, montrant des surfaces cohérentes, lisses et non poreuses de la structure centrale des cellules Franz FA. Les études de perméation utilisant des constructions transparentes imprimées en 3D ont conduit à une récupération de la caféine de 12,85 ± 0,53 µg cm⁻² dans la solution de récepteur après 180 min avec une récupération de perméant comparable, 11,49 ± 1,04 µg cm⁻², pour les homologues de verre.

CONCLUSION: les constructions FA peuvent être considérées comme des alternatives viables à l’utilisation d’appareils de verre conventionnels offrant une méthode simple, reproductible et rentable de réplication de la verrerie de laboratoire spécialisée. Une gamme plus large de perméants sera étudiée dans de futures études avec ces nouvelles cellules de diffusion Franz imprimées en 3D.

Introduction

The evaluation of permeation profiles from cosmetic formulations is considered to be a crucial component in both the development and quality assurance of any new product [1,2]. Data gathered from such studies allow researchers to assess the viability of delivering different materials to and through biological membranes. To date, laboratory in vitro permeation processes require the use of modified Franz type diffusion cells, conventionally fabricated from glass, which are available in different formats that can be customized to experimental requirements [3]. We have previously reported on the development of a novel low-cost approach to additive
manufactured (AM) Franz diffusion cells and compared the permeation of different active ingredients between glass cells and these newly developed 3D-printed constructs. Results suggested a possible interaction of the permeants with the transparent resins used to produce the diffusion apparatus [4]. In the present work, the permeation of caffeine (CAF), a widely studied hydrophilic permeant, was investigated in both AM cells and glass cells [5,6]. Microscopic analysis of these 3D Franz cells was also conducted in order to assess physical integrity of the scaffolds [7].

Caffeine (CAF) was purchased from Sigma Aldrich, Dorset, UK. Phosphate-buffered saline (PBS) tablets were purchased from Oxoid Limited, Basingstoke, UK. Polydimethylsiloxane (PDMS) membrane with a thickness of 250 \( \mu \)m was supplied by Shielding Solutions Limited, Braintree, UK. High-performance liquid chromatography (HPLC) grade water and methanol were obtained from Fisher Scientific, Loughborough, UK. Stereolithography resin GPCL04 for printing of transparent materials was purchased from Formlabs, Massachusetts, USA.

AM Franz type diffusion cells were produced using typical glass homologues measurements which were collected using a Vernier callipers (RS Components Ltd., Corby, UK) and drawn in silico, using an online computer-aided design (CAD) program – TinkerCAD\textsuperscript{TM} (Autodesk\textsuperscript{®}, California, USA). Graphical designs were then imported to the Preform\textsuperscript{TM} software tool (version: 2.14.0, Formlabs, Massachusetts, USA) prior to printing. Stereolithography (SLA) 3D printing was carried out using a Form2 printer (Formlabs, Massachusetts, USA). The 3D-printed receptor compartment was reproduced with the following specifications: outer diameter (O.D.) = 30 mm, inner diameter (I.D.) = 10 mm, height (\( h \)) = 16 mm and aliquot collection arm (length (\( l \)) = 54 mm with I.D. = 3 mm) as shown in Fig. 1. The inner object volume was 2.33 ± 0.03 cm\(^3\) (weight (\( w \)) = 499.23 ± 0.41 mg) for the receptor compartment of the AM Franz cell. The donor compartment was printed with O.D. = 30 mm, I.D. = 10 mm, \( h \) = 10.7 mm (Fig. 1) which gave a total inner object volume of 1.15 ± 0.01 cm\(^3\) (\( w \) = 309.22 ± 0.30 mg). Printing was carried out using 100 \( \mu \)m resolution and a resin tank temperature of 28 \( ^\circ \)C. Preform\textsuperscript{TM} generated supports (point size: 0.50 mm, point density: 1:0) were used for all printed scaffolds. Post-curing of AM constructs was achieved by exposing the 3D-printed Franz type cell compartments to UV light (405 nm) at 60\(^\circ\)C for 15 min using Form Cure (Formlabs, Massachusetts, USA) [8]. All transparent 3D-printed Franz cells were tested for leaks by filling both compartments with a PBS (pH 7.3 ± 0.1) solution. These printouts (compartments) were clamped together using in-house manufactured metallic clamps. The AM cells were examined for leaks over a minimum of 24 h, and the printouts were considered successful if no aqueous media was present on the outer wall after this period.

Caffeine analysis was conducted using a HPLC (Agilent Technologies 1200 series) equipped with an Agilent G1322A degasser, G1311A quaternary pump, G1329A auto sampler and G1316A thermostat column compartment (Agilent Technologies, Cheadle, UK). The analysis was performed using a Phenomenex Luna Phenyl Hexyl column fitted with a guard column (Phenomenex, Cheshire,
The length, internal diameter and particle size were 250 mm, 4.6 mm and 5 µm, respectively. The mobile phase consisted of water:methanol (40:60). Prior to its use, the mobile phase was degassed using an ultrasonicator (VWR International, Lutterworth, UK) to remove air bubbles. The flow rate of the mobile phase was 1 mL min\(^{-1}\), and the column temperature was set at 23°C. The chromatogram was acquired at a wavelength of 278 nm. A sample volume of 10 µL was injected for a total run time of 10 min. A known amount of CAF was dissolved in PBS (pH 7.3, 0.1), and a stock solution (1000 µg mL\(^{-1}\)) was prepared. The stock solution was diluted to prepare various concentrations of CAF. The CAF peak was evident at 3.9 min. The calibration curve was constructed in the concentration range of 0.05–500 µg mL\(^{-1}\). A linear relationship was found between concentration and peak area with regression coefficient values (\(r^2\)) of greater than 0.999. The LOQ and LOD values were 0.5 and 0.05 µg mL\(^{-1}\), respectively [9].

Caffeine compatibility studies with AM Franz type diffusion cells were conducted by filling the diffusion cell receptor compartment with a 1.5 mg mL\(^{-1}\) CAF solution prepared in PBS (pH 7.3 ± 0.1). The AM Franz cell was then sealed with Parafilm (Bemis NA, 02 4 4 8 7 2 1.35 1.40 1.45 1.50 1.55 1.60 278 μm Figure 3 Amount in mg mL\(^{-1}\) of CAF recovery from glass and 3D-printed receptor compartments at 0, 24, 48 and 72 h (\(n = 4\), mean ± SD).

Figure 5 Amount of CAF permeation from initial dose per cm\(^2\) of PDMS membrane at 0, 15, 30, 45, 60, 90, 120 and 180 min using glass and 3D-printed Franz type diffusion cells (\(n = 4\), mean ± SD).

© 2020 The Authors. *International Journal of Cosmetic Science* published by John Wiley & Sons Ltd on behalf of Society of Cosmetic Scientists and Société Française de Cosmetologie

*International Journal of Cosmetic Science*, 42, 415–419
Neenah, USA) and placed in a JB Nova thermostatically controlled water bath (Grant, London, UK) set to 32 ± 1°C equipped with a HP 15 stirring system (Varimag, Florida, USA). About 200 µL aliquots were taken from the cell receptor compartment at different time points, 0, 24, 48 and 72 h and replaced with the same volume of fresh CAF solution. The samples were appropriately diluted to be in the range of the calibration curve and analysed using HPLC.

The surface morphology of the AM Franz type diffusion cells from pre- and post-compatibility studies was analysed using an EVO MA 10 (Carl Zeiss SMT GmbH, Oberkochen, Germany) scanning electron microscope (SEM). Constructs were cut in different cross-sectional directions and sputter coated with gold for 45 s using an Emitech 550 (Emitech Ltd., Ashford, UK). The 3D-printed materials were then attached onto adhesive carbon disks (Fig. 2), and SEM micrographs were taken of different longitudinal and cross-sections. SEM was conducted at low magnification, with an electron accelerating voltage of 10–20 kV under vacuum. Images of the surface structure of the samples were captured and collected using the Everhart-Thornley detector image acquisition and processing software (Carl Zeiss SMT GmbH, Oberkochen, Germany).

In vitro permeation studies in both glass and AM Franz type diffusion cells were conducted using 1 mL CAF (1.5 mg mL⁻¹) solution, applied to a PDMS model membrane. Freshly prepared PBS (pH 7.3 ± 0.1) was used as the receptor solution. The temperature of the PDMS membrane was equilibrated to 32 ± 1°C, and the CAF solution applied. The donor compartment was not occluded after application of the CAF solution. Samples of 200 µL of receptor solution were removed from the receptor compartment at various time intervals (0, 5, 10, 15, 30, 45, 60, 90, 120 and 180 min) and replaced with fresh temperature equilibrated PBS solution. The samples were appropriately diluted to be in the range of the calibration curve and analysed using HPLC.

Glass diffusion cells are conventionally used in dermal permeation studies given their lack of interaction with permeants [10]. However, high-costs are associated with this type of glass apparatus and they are not mechanically robust. We previously reported the development of 3D-printed Franz diffusion cells in different permeation experiments with results showing chemical incompatibility between the electron rich domains of permeants and the chemical structure of the polymer in the printed constructs [4]. In this study, CAF was chosen as a model molecular permeant because of its hydrophilic nature (Table 1) and wide use in skin permeation studies [6,9,11,12]. With a log P value of −0.07 and an aqueous solubility of 21.6 mg mL⁻¹, CAF was initially screened for its AM Franz diffusion cell compatibility prior to conducting in vitro permeation studies. Stability testing of CAF with the Formlabs clear resin GPCLO4 confirmed comparable outcomes as for glass diffusion cells, with results demonstrating similar recovery between the two constructs (1.49 ± 0.01 and 1.50 ± 0.01 mg mL⁻¹, respectively) after 72 h as reported in Fig. 3 (Two-way ANOVA Tukey’s multiple comparisons). This suggested that no interaction occurred between CAF and the transparent 3D-printed scaffold indicating the suitability of these AM constructs to be used for permeation studies.

The morphology of the AM Franz type diffusion cells surfaces was characterized using SEM. As shown in Fig. 4, different cross-sectional images of the 3D-printed constructs using 54 to 256 × magnifications confirmed the structural similarity between AM scaffolds studied prior (Fig. 4A-C) and post-compatibility (Fig. 4D-F). No significant differences were visible from the SEM micrographs demonstrating consistent, smooth and non-porous surfaces of the 3D-printed Franz cells’ core structure. Additionally, shape and size uniformity across all printed samples was consistent for both compatibility and permeation studies with no evident swelling of the polymeric constructs.

In vitro permeation studies showed that similar permeation of CAF was observed in AM diffusion cells when compared with the Franz type glass cells as shown in Fig. 5. All experiments were conducted using the same type of PDMS membrane, confirming the suitability of CAF as a model molecular permeant with the GPCLO4 AM resin. Permeation studies using transparent 3D-printed constructs resulted in 12.85 ± 0.53 µg cm⁻² CAF recovery in the receptor solution after 180 min with a comparable CAF permeation, 11.49 ± 1.04 µg cm⁻², for the glass homologues.

Similar to their glass counterparts, 3D-printed Franz type diffusion cells were shown to be structurally solid with SEM analysis displaying no printout degradation during the AM resin/model active compatibility studies. Also, subsequent comparative permeation analysis conducted in typical glass diffusion cells and 3D-printed scaffolds using CAF showed consistent results. Physical and/or chemical interactions between printing resins and other model active ingredients were previously described as a limitation for the use of AM technology for Franz cell fabrication [4]. Here, we have demonstrated the relationship of using a more hydrophilic molecular entity as an alternative for the testing of these 3D-printed transparent constructs. This choice relied on our previously reported methodology that showed improved compatibility between the polymeric scaffold and the model tested compounds when less lipophilic permeants were used (i.e. nicamamide, terbinaline hydrochloride and diclofenac free acid) [4]. Hence, the increase in compound recovery translates into a robust analytical process suitable to be used in real-life training scenarios. With a price range of 2–3 USD per set (i.e. Franz cell receptor and donor compartments) and a printing time of ~3.5 h to reproduce four full sets, AM technology may now be considered as a viable alternative to the use of conventional glass cells.

Acknowledgments
This research was supported by GSK Consumer Healthcare. We thank our colleagues from the UCL Skin Research Group and Hilton Laboratory who provided insight and expertise that greatly assisted the research. We also like to acknowledge the expertise and support given by Toby Neville, Professor Paul Shearing and Professor Dan Brett from UCL Department of Chemical Engineering, Faculty of Engineering Science.

References
1. Clowes, H.M., Scott, R.C. and Heylings, J.R. Skin absorption: Flow-through or static diffusion cells. Toc. in Vtro. 8, 827–830 (1994).
2. Abd, E., Yousef, S.A., Pastore, M.N. et al. Skin models for the testing of dermal drugs. Clin. Pharmacol. 8, 163–176 (2016).
3. Franz, T.J. Percutaneous absorption. On the relevance of in vitro data. J. Inv. Derm. 64, 190–195 (1975).
4. Sil, B.C., Alvarez, M.P., Zhang, Y. et al. 3D-printed Franz type diffusion cells. *Int. J. Cosm. Sci.* **40**, 604–609 (2018).

5. Alex, A. Physicochemical profiling (Solubility, Permeability and Charge State). *Curr. Top. Med. Chem.* **1**, 277–351 (2001).

6. Luo, L. and Lane, M.E. Topical and transdermal delivery of caffeine. *Int. J. Pharm.* **490**, 155–164 (2015).

7. Zaleski, R., Stefaniak, W., Maciejewska, M. and Goworek, J. Porosity of polymer materials by various techniques. *J. Por. Mat.* **16**, 691–698 (2008).

8. Formlabs. *A guide to post-curing formlabs resins*. Formlabs, USA 2018 (Accessed 05/06/2019).

9. Potard, G., Laugel, C., Baillet, A., Schaefer, H. and Marty, J.P. Quantitative HPLC analysis of sunscreens and caffeine during in vitro percutaneous penetration studies. *Int. J. Pharm.* **189**, 249–260 (1999).

10. Skelly, J.P., Shah, V.P., Maibach, H.I. et al. FDA and AAPS report of the workshop on principles and practices of in vitro percutaneous penetration studies: relevance to bioavailability and bioequivalence. *Pharm. Res.* **4**, 265–267 (1987).

11. Trauer, S., Patzelt, A., Otberg, N. et al. Permeation of topically applied caffeine through human skin—a comparison of in vivo and in vitro data. *Br. J. Clin. Pharmacol.* **68**, 181–186 (2009).

12. Gajewska, M., Paini, A., Sala Benito, J.V. et al. In vitro-to-in vivo correlation of the skin penetration, liver clearance and hepatotoxicity of caffeine. *Food Chem. Toxic.* **75**, 39–49 (2015).

13. Hansch, C., Leo, A. and Hoekman, D. *Exploring QSAR - Hydrophobic, Electronic, and Steric Constants*. American Chemical Society, Washington, DC (1995).

14. Yalkowsky, S.H., He, Y. and Jain, P. *Handbook of Aqueous Solubility Data*. 2nd ed. CRC Press, Boca Raton, FL (2010).