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Tailored on demand anti-coagulant dosing: an in vitro and in

vivo evaluation of 3D printed purpose-designed oral dosage

- forms
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26 **A B S T R A C T**

27 Coumarin therapy has been associated with high levels of inter- and intra-individual variation in the required dose to reach a therapeutic anticoagulation outcome. 28 Therefore, a dynamic system that is able to achieve accurate delivery of a warfarin 29 dose is of significant importance. Here we assess, the ability of 3D printing to fabricate and deliver tailored individualised precision dosing using an in-vitro model. Sodium 30 warfarin loaded filaments were compounded using hot melt extrusion (HME) and 31 further fabricated via fused deposition modelling (FDM) 3D printing to produce capsular-ovoid-shaped dosage forms loaded at 200 and 400 µg dose. The solid 32 dosage forms and comparator warfarin aqueous solutions were administered by oral gavage to Sprague-Dawley rats. In vitro, warfarin release was faster at pH 1.2 in 33 comparison to pH 2. A novel UV imaging approach indicated that the erosion of the 34 methacrylate matrix was at a rate of 16.4 and 15.2 µm/min for horizontal and vertical planes respectively. In vivo, 3D printed forms were as proportionately effective as 35 their comparative solution form in doubling plasma exposure following a doubling of 36 warfarin dose (184% versus 192% respectively). The 3D printed ovoids showed a lower C_{max} of warfarin (1.51 and 3.33 mg/mL versus 2.5 and 6.44 mg/mL) and a longer 37 T_{max} (6 and 3.7 versus 4 and 1.5 h) in comparison to liquid formulation. This work demonstrates for the first time in vivo, the potential of FDM 3D printing to produce a 38 tailored specific dosage form and to accurately titrate coumarin dose response to an 39 individual patient.

Keywords: Rapid prototyping; Patient-centred; Personalized; Patient-specific; Three
 dimensional printing; additive manufacturing.

42

44 **1. Introduction**

45 For over 50 years now, coumarins have been the most prescribed oral anticoagulants.[1] Nevertheless, despite their wide use, coumarin therapy has been associated with a high level of 46 inter-individual variation in dose required to achieve therapeutic anticoagulation response.[2] 47 The administration of an inappropriate warfarin dose for example may place a patient in a 48 49 hypercoagulable state or increase the patient's risk of bleeding complications early in therapy. 50 As a consequence of over-anticoagulation response, there is an increased risk of major bleeding following the use of anticoagulants by 9.1% [3]. The American College of Chest Physicians 51 (ACCP) supports an "induction" dose of 2 to 5 mg per day which needs to be adjusted 52 53 according to the patient's International Normalised Ratio (INR)[4]. The pharmacodynamics and pharmacokinetics of coumarins are largely influenced by many factors such as patient age, 54 55 body weight, dietary vitamin K intake, concomitant medications, as well as various disease states.[2] Hence to ensure that the patient's INR remains within the target range, regular 56 57 coagulation monitoring and dose modification is necessary.[5]

Nevertheless, limited doses of warfarin tablets are available in the market and dose 58 modification usually requires multiple tablet ingestion or cutting or splitting of larger dose 59 60 tablets, which could lead to variations in drug content.[6, 7] An area of potential improvement to warfarin therapy would be the ability to produce flexible on-demand precision tailored dose 61 adjustments (particularly given warfarin's due to narrow therapeutic index). One technology 62 that can potentially easily benefit anticoagulant therapy is 3D printing, owing to its flexible and 63 64 precise manufacturing capability, which enables administration of the lowest effective dose of the drug to maintain the target INR. Indeed, recently, Vuddanda et al. (2017) demonstrated the 65 potential of a re-engineered thermal inkjet printer to address the challenge of warfarin dosage 66 67 personalisation, achieving highly reproducible minute warfarin dose of approximately 50 µg [8] 68

69 3D printing potential and feasibility has been revealed in several fields such as aerospace, engineering, arts, as well as in fabricating medical implants and devices. Although still at its 70 infancy in the field of personalised medicine, it is expected to revolutionise healthcare and set 71 72 an innovative platform for pharmaceutical product design and extemporaneous preparation of patient-tailored dosage forms.[9] Fused deposition modelling (FDM) 3D printing, in particular, 73 has been proposed as a platform for controlling the dose.[10] It has demonstrated its capability 74 to manufacture mechanically stable tablets fabricated from pharmaceutical grade polymers 75 without post-processing steps.[10-13] For instance, FDM 3D printing has been viably 76 established using pharmaceutical grade polymers such as PVP [9, 14], methacrylate [15] and 77 78 cellulose [12] based polymers.

79 The use of animal models is commonly used to predict formulation behaviour in humans. The use of rats in particular is favoured due to their small size, relatively low cost of breeding 80 and up-keep, as well as the presence of large databases of drug pharmacokinetic data in rats 81 and in humans.[16] Nevertheless, the testing of solid dosage forms in rats presents a challenge 82 83 in terms of ease of administration. Owing to the need to use a small dosage form size, crushed tablets filled in capsule or suspended in liquid have often been used as an inferior alternative 84 to test the in vivo performance of a tablet in rats.[17, 18] However, such approaches 85 significantly alter the nature of the dosage form. More recently, the formulation of mini-tablets 86 for animal use have been attempted [19, 20]. It is therefore important to develop strategies that 87 authentically test intact scaled down human dosage forms for animal studies to enable more 88 reliable extrapolation of human pharmacokinetic responses. 89

This work aimed to assess the suitability of FDM 3D printer technology for i) fabricating purposely designed solid dosage forms, and ii) tailoring the dose of a narrow therapeutic index drug, namely warfarin. To achieve this goal, rat-tailored FDM 3D printed warfarin ovoid tablets were printed and administered to Sprague–Dawley rats for testing to obtain their pharmacokinetics (PK) parameters.

95 **2. Materials and methods**

9697 2.1 Materials

Warfarin (sodium salt) was purchased from Arcos (UK). Eudragit E was donated from
Evonik Industries (Darmstadt, Germany). Triethyl citrate (TEC) and tri-calcium phosphate
(TCP) were supplied by Sigma–Aldrich (UK). Acetonitrile and methanol were supplied by
British Drug Houses (BDH, London, UK). Scotch Blue Painter's tape 50 mm was supplied by
3M (Bracknell, UK).

103 2.2 Preparation and optimisation of filaments

In order to fabricate drug-loaded filaments, a hot melt extrusion method was implemented 104 using a Thermo-Scientific HAAKE MiniCTW extruder (Karlsruhe, Germany). A 10 g sample 105 of Eudragit E: TEC: TCP: sodium warfarin 46.75 : 3.25 : 49:1) was accurately weighed and 106 added gradually to counter flow twin-screw hot melt extruder, HAAKE MiniCTW (Karlsruhe, 107 Germany). To allow homogeneous distribution of the powders, the molten mass was mixed in 108 the extruder for at least 5 min prior to extrusion. The specific temperature of initial feeding and 109 extrusion for the filament were 100 and 90 °C respectively. A torque control of 0.8 Nm was 110 111 used to extrude the filaments. Filaments were stored in sealed plastic bags at room temperature before 3D printing. 112

113 2.3 Design and printing of tablets

Tablets were constructed with the pre-prepared filaments using a MakerBot Replicator[®] 2X Experimental 3D Printer (MakerBot Industries, New York, USA) equipped with 0.4 mm nozzle size. The templates used to print the tablets were designed in a caplet shape using Autodesk[®] 3ds Max[®] Design 2016 software version 18.0 (Autodesk, Inc., USA). The design was saved in a stereolithography (.stl) file format and was imported to the 3D printer's software, MakerWare Version 3.9.1.1143 (Makerbot Industries, LLC., USA).

120 Two sets of 3D printed tablets were fabricated:

121 In order to establish the ability of the system to control the low dose of drug for clinical 122 use, a series of tablets with increasing volumes were then printed by increasing the dimensions 123 of the design: length × width × heights (L, H, W). The ratios between dimensions 124 (W = H = 0.4 L) remained constant. The size of the printed tablet (*M*) was changed to achieve 125 target doses of 0.5, 1, 3 or 5 mg (Table 1S).

To assess *in vivo* performance of this tablets in rats, a separate set of 3D printed ovoid shapes were manufactured with a cylindrical diameter of 2 mm and lengths of 5.5 or 11 mm to achieve a dose of 200 and 400µg respectively. Objects were printed using modified settings of the software as described earlier in our previous work at a temperature of 135 °C. [15]

130 2.4 Thermal analysis

Samples (raw materials, extruded filaments and printed tablets) were characterised using 131 differential scanning calorimetry (DSC) and thermogravimetric analysis (TGA). For DSC 132 analysis, a differential scanning calorimeter DSC Q2000 (TA Instruments, Elstree, 133 Hertfordshire, UK) with a heating rate of 10 °C/min was used. Samples were heated to 100 °C 134 for 5 min to exclude the effect of humidity then cooled to -20 °C. This was followed by a heat 135 scan from -20 °C to 300 °C. Analysis was carried out under a purge of nitrogen (50 mL/min). 136 137 The data was analysed using TA 2000 analysis software. Standard 40 µL TA aluminium pans and pin-holed lids were used with an approximate sample mass of 5 mg. All measurements 138 were carried out in triplicate. 139

For TGA analysis, raw materials, extruded filaments and 3D printed tablets were analysed using a TGA/SDTA851e Mettler Toledo (Leicester, UK). Samples (5 mg, n=3) were placed in 40 μ L aluminium pans and were then heated from 25 to 500 °C at a heating rate of 10°C/min and nitrogen gas flow of 50 mL/min. The thermal decomposition (or degradation) profile was analysed using STARE software version 9.00.

145

146 2.5 X-ray powder diffraction (XRD)

Samples (raw materials extruded filaments and printed tablets) were characterised using an X-ray diffractometer, D2 Phaser with Lynxeye (Bruker, Germany). Samples were scanned from $(2\theta) = 5^{\circ}$ to 50° using 0.01° step width and a 1 second time count. The divergence slit was 1 mm and the scatter slit 0.6 mm. The wavelength of the X-ray was 0.154 nm using Cu source and a voltage of 30 kV. Filament emission was 10 mA using a scan type coupled with a theta/theta scintillation counter over 60 min.

153

154 2.6 Characterisation of tablet properties

The hardness of six ovoid tablets was measured using a TBH 200 (Erweka GmbH, Heusenstamm, Germany). The mean crushing strength was determined, whereby an increasing force was applied to the tablet until it fractured or deformed.

In order to assess the friability of the tablets, 20 tablets were randomly selected, weighed and placed in a friability tester Erweka TAR 10 (Erweka GmbH, Heusenstamm, Germany) and the drum was then rotated at 25 rpm for 4 min. The tablets were reweighed and the differences in weight were calculated and displayed as a percentage of the original sample weight. In order to assess weight uniformity, 10 tablets were randomly selected and weighed. The average weights were measured and the percentage deviation of the individual tablets from the mean was determined.

To assess the impact of both HME and FDM 3D printing on drug content, 3 tablets from each formulation, were randomly selected and weighed. Tablets were then individually placed in a 1000 mL volumetric flask containing 0.1 M HCl and sonicated for 2 h. The solutions were filtered through 0.22 μm Millex-GP syringe filters (Merck Millipore, USA) and prepared for HPLC analysis.

170 Warfarin concentration in samples was assessed using an Agilent UV-HPLC 1260 series 171 (Agilent Technologies, Inc., Germany) equipped with Kinetex C18 column (100×2.1 mm, 172 particle size 2.6 µm) (Phenomenex, Torrance, USA) and set at temperature 26 °C. The mobile 173 phase was 4:1 mixture of methanol: pH 3 water (adjusted with orthophosphoric acid) at a flow 174 rate of 1 mL/min. The injection volume was 100 µL and the stop time was 10 min. The wavelength was set to 230 nm and the retention time of the drug was 6.3 min with a limit of detectionof 0.05 mg/L.

177 2.7 *In vitro* dissolution studies.

a.Surface dissolution imaging. A Sirius SDi2, the second generation UV imaging system, 178 designed to accommodate whole dosage forms, was used to visualize surface dissolution of 179 sodium warfarin from the 3D printed dosage forms as a whole (Fig. 1). The 3D printed tablets 180 were introduced into the SDi2's flow cell. The dissolution medium (0.1M HCl at 37°C) applied 181 at a flow rate of 8.2 mL/min. The dissolution medium was introduced into the flow cell in the 182 open loop configuration, from bottom to top, with an equivalent linear velocity of 1 cm/min. 183 Dissolution experiments were recorded for a total duration of 60 min. The two dimensional 184 185 detection area on the SDi2 is significantly larger than for the SDI (24 mm width x 28 mm height) to accommodate dissolution imaging profiling of intact whole dosage forms, with a 186 spatial resolution of 13.75 µm. The flow cell was illuminated using alternate pulses from two 187 255 and 520 nm wavelength LEDs. The dual wavelength enables two separate video captures 188 189 to be produced from a single experiment. Real-time data were then used to measure and 190 differentiate between drug release into solution and tablet erosion from the 255 and 520 nm light obtained videos, respectively. 191



¹⁹²

Figure 1. Schematic diagram of SDi2 instrument. LED's of different wavelength are em ployed to illuminate the 3D printed tablet in flow through cell filled with gastric medium. The obscuration or absorbance of the sample was recorded using an Actipix detector. The medium is pre-heated to 37°C before going through the Whole Dosage Flow Cell and is recirculated in a closed loop configuration.

197 <u>b. USP II dissolution studies.</u> The *in vitro* release of warfarin from 3D printed tablets was
 198 investigated using a USP II Erweka DT600 dissolution tester (Erweka GmbH, Heusenstamm,

199 Germany). Three tablets were randomly selected and individually placed in the dissolution 200 vessels each containing 900 mL of a fasted state simulated gastric fluid (FaSSGF) (1.75 mM 201 SLS, 0.01N HCl, 0.2% NaCl, pH 2.0) at 50 rpm and 37 ± 0.5 °C. Aliquots (5 mL) were 202 manually collected using 5 mL Leur-Lock syringes at 0, 5, 10, 15, 20, 25, 30, 40, 50 and 60 min 203 time intervals and filtered through an Agilent 0.22 µm filter. Each aliquot withdrawn was 204 replaced with 5 mL of 0.1 M HCl and analysed using the above described HPLC method.

205 2.8 *In vivo* studies

Adult heathy male Sprague–Dawley rats with an average weight of 240±15 g 206 accommodated at the University of Petra's Animal House (Amman, Jordan) under controlled 207 temperature (22 °C-24 °C), humidity (55%-65%), and a 12 hours photoperiod cycle. All rats 208 were acclimatized for 10 days before experimentation. Rats were weighed and randomized into 209 groups (n=6 rats per cage). Rats were offered standard pellet diet (Jordan Feed Company Ltd., 210 Amman, Jordan) and served clean tap water ad libitum. However, animals were fasted for 18 211 hours before the day of testing. All experiments were carried out in accordance with University 212 of Petra's Institutional Guidelines on Animal Use that adopts the guidelines of the Federation 213 of European Laboratory Animal Sciences Association (FELASA). The animal study protocols 214 were revised and approved by the Higher Research Council at the Faculty of Pharmacy and 215 216 Medical Sciences, University of Petra (Amman, Jordan).

3D printed tablets (200 or 400 µg) were administered to the rats via any oral capsule stainless 217 218 steel feeding needle. Comparison control 1 mL warfarin solutions (200 or 400 µg), equivalent to the tablet doses, were freshly prepared and administered to the rats by a stainless steel oral 219 gavage needle (Harvard Apparatus, Kent, UK). Following oral administrations, blood samples 220 221 were pooled from rat's tail (n=6 rats per group) at different time intervals namely at; 1, 2, 3, 4, 6 and 8 hours post administration. Blood was left to clot, centrifuged for 10 min at 2000G, and 222 then serum was separated and transferred directly into Eppendorf tubes, and kept in a freezer 223 at -20 °C until analysis. 224

225 2.9 Analysis of warfarin

For the analysis of warfarin an MS/MS system: API 3200 (Applied Biosystems, MDS 226 SCIEX, USA) attached to Agilent 1200 HPLC (Agilent Technologies, USA) controlled by 227 Analyst 1.6.1 software, was utilised. For the extraction of warfarin from the samples, 100 µL 228 of spiked/blank plasma were pipetted into previously labelled Eppendorf tube, 25 µL of the 229 internal standard (IS) Fenofibric acid (FFA) from 100.0 µg FFA/mL IS solution was added to 230 the tubes and vortexed for 30 sec. Afterwards, the precipitation solution, acetonitrile (400.0µL) 231 was added to the tube and vortexed for further 1 min. Samples were then centrifuged for 5 min 232 at 14,000 rpm and the supernatant was collated and transferred into an auto-sampler micro vial 233 for analysis. The mobile phase used for analysis comprised of (30:70) mixture of ammonium 234 chloride 0.001M: acetonitrile respectively eluted at a flow rate of 0.7 mL/min through a 235 Thermo BDS Hypersil C18 (50×2.1 mm, particle size 5 µm) column (Thermo Fisher Scientific, 236 Germany) at the temperature 30°C. The injection volume was 5 µL and the stop time was 237 0.7 min. The retention time of the drug was 0.3 min with a limit of detection of 10 ng/mL. 238

239 2.10 Statistical Analysis

240 Independent sample T-test was also employed using a SPSS Software (22.0.0.2) to analyse

the *in vitro* tablet characterisation results. Differences in results where $p \le 0.05$ were considered significant.

243 3. Results and discussion

In this study, we explored the adaptability of FDM based 3D printing to engineer and control 244 245 the dose of immediate release warfarin tablets. When a series of warfarin tablets with increasing dimensions were printed (Fig. 2A, Table S1), a high level of correlation was identified between 246 the theoretical volume of the tablet design and their weights ($R^2=0.9934$) (Fig. 2B). This 247 248 indicated the ability of FDM 3D printing method to achieve a sufficient control of the mass of 3D printed tablets. To establish the ability of such 3D printing method to control dosage, 249 theoretical doses based on tablet mass and measured dose of warfarin in the tablet were 250 compared. The range of dose accuracy was between 91.5% and 102.4% (Fig. 2C). The 251 coefficient of determination between target and achieved dose ($R^2 = 0.9902$) showed that it is 252 possible to fabricate tablets with desired dose of warfarin through volume modification even 253 at a minute dose of 500 µg (Fig. 2D). With the advances in 3D printers, additional safeguards 254 and quality control mechanisms can be introduced to the evolving technology [21], which are 255 expected to minimise dose variation in the near future. 256 257



Figure 2. Precision of 3D printing to control low dose sodium warfarin. (A) Images of warfarin loaded FDM 3D printed tablets with increasing dose, (B) Correlation between the theoretical volume and tablet mass, (C) warfarin dose accuracy in the 3D printed tablets and (D) correlation between theoretical volume and warfarin dose (n=3, ±SD).

Profiles from thermogravimetric analyses of warfarin and other additives as well as HME 262 processed filaments and 3D printed tablets are shown if Fig. 3A. Sodium warfarin alone or 263 incorporated in filaments did not suffer a significant weight loss at the printing temperature 264 265 135 °C. Therefore, it can be assumed that minimal or no degradation of warfarin occurs in the HME as well as in the FDM's nozzle under the utilised temperatures (Fig. 3A). The processing 266 temperatures were lower than the melting point of sodium warfarin (161 °C). Differential 267 scanning calorimetry was also conducted to examine the plasticising effect of components on 268 the methacrylic filament. As demonstrated in Fig 3B, the addition of TEC as a plasticizer 269 significantly depressed the Tg of filament to 34 °C from 54°C. However, warfarin was found 270

271 to have no significant effect on the Tg of Eudragit E. This could be attributed to the minute percentage of the drug used in the polymeric structure (1% w/w), which was insufficient to 272 significantly influence the mobility of methacrylic polymer chains within the filament matrix. 273 XRD spectra showed that β -calcium tribasic phosphate displayed peaks at 2-theta=17°, 27.8°, 274 31°, 34.4° corresponding to calcium tribasic phosphate [22], whilst warfarin drug substance 275 showed peaks at 2-theta=12.4° and 18°. XRD spectra of the warfarin filament and tablet 276 showed an absence of these specific peaks [23, 24], suggesting the warfarin is present in an 277 amorphous form within the tablet structure (Fig. 3C). 278

From determination of the mechanical properties of the 3D printed tablets, the friability of all batches was found to be zero percent. This highlights a prime advantage of FDM 3D printing in generating mechanically stable tablets over its rival technologies such as extrusion 3D printing [25] and powder-based 3D printing. [26, 27] The lack of a drying step or any postprinting finishing procedures, clearly demonstrates the potential of this technology to instantly produce a ready-to-use dosage form within minutes following a healthcare team request.



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Figure 3. Thermal analysis of Eudragit E based 3D printing filaments. (A) Thermal degradation profiles for Eudragit E, sodium warfarin, TCP, warfarin loaded filament and tablet, (B) DSC thermograph for warfarin loaded filament and tablet, (C) XRD spectra of Eudragit E, TCP, warfarin, and warfarin loaded filament and tablet.

290

The release pattern of warfarin from the methacrylic matrix was investigated using a modified FaSSGF [28] as a dissolution medium (Fig. 4). All tablets showed a release pattern of > 80% dissolution at 45 min regardless of their individual sizes. The dissolution release profile was attributed to the ionisation of the amino groups of the cationic methacrylic polymer in modified FaSSGF (pH 2.0), which leads to electrostatic repulsion between cationic polymer chains and facilitates polymer dissolution and drug release. The release was compliant with British Pharmacopeia criteria for warfarin tablets [29].



298 299 Figure 4. In vitro release pattern of sodium warfarin from 3D printed tablets of different doses from a USPII dissolution test in modified FaSSGF (pH 2.0) (n=3, ±SD). 300

To better understand the drug release from the 3D printed tablets, the dissolution behaviour 301 of the tablets at the dissolving surface in contact with the dissolution media was explored. A 302 single wavelength system has been previously used to study drug powder dissolution [30]. Here 303 we employ a UV imaging technology capable of generating visual images from simultaneous 304 spectroscopic evaluation for a complete dosage form (Fig. 5A, B). A clear advantage of using 305 such a novel UV-VIS imaging technique over the other well-established imaging techniques 306 lies in the simplicity of operation and interpretation of generated data, analogous to findings 307 by Østergaard.[31] The measurement of light intensity passing through an area of a quartz tube 308 as a function of position and time can also enable quantification of the drug substance at 309 different time intervals. During the dissolution process, drug concentration increased in the 310 first 20 min in the closed loop of the flow-through system. Simultaneously the tablet size was 311 312 eroded at a rate of 16.4 and 15.2 µm/min for horizontal and vertical planes respectively. It is worth noting that surface analysis indicated no significant swelling in the first 5 min. The 313 simultaneous drug release data suggested that under the dissolution conditions of study, the 314 majority of drug release took place by a diffusion mechanism before the erosion of the 315 methacrylic matrix within the flow-through cell is complete. 316



Figure 5. Changes in tablet height (A) and width (B) at 0, 10, 20, 30, 40, 50 and 60 min of the flow through dissolution test using Actipix SDI 2 dissolution imaging technology. (C) UV absorbance image following the illumination of follow cell containing warfarin 3D printed tablet at 255 nm wavelength.
(D) Percentage sodium warfarin release from 3D printed tablet during dissolution test (n=3, ±SD).

A prime advantage of 3D printing technologies lies in their highly flexible nature and capacity to construct dosage forms with accurate spatial distribution of ingredients compared to traditional manufacturing techniques. Therefore, constructs can now be printed to suit the anatomy of not only a particular animal but according to the weight and size of that subject. Rats are commonly considered most suitable for determining the mechanism of drug absorption and bioavailability values from powder or solution formulations [32] as well as micro- or nanoparticles [33].

Two different warfarin tablets were specially designed (Fig. 6A1) to mimic the dimensions 329 of commonly used hard capsules intended for oral delivery to rats. Tablets were successfully 330 printed (Fig. 6A2) and were orally gavaged to rats. The pharmacokinetic parameters of warfarin 331 following oral administration either as 3D printed tablets or in a solution form were evaluated 332 (Table 1, Fig. 6B, C). Warfarin plasma exposure was significantly different when an equal dose 333 was administered either as solutions or as 3D printed tablets. The solution showed a markedly 334 higher C_{max} (2.5 and 6.44 mg/mL) and shorter T_{max} (2.67 or 1.5h) for the 200 or 400 μ g/mL 335 solution respectively, in comparison to C_{max} values (1.51and 3.33 mg/mL) and T_{max} values (6 336 or 3.7 h) for 200 μ g (p<0.05) and 400 μ g (p<0.01) warfarin tablets respectively. 337

Table 1. Summary of pharmacokinetic parameters of warfarin following oral gavage of 200 or 400μg

340 from sodium warfarin solution and 3D printed tablets to adult heathy male Sprague–Dawley rats.

Dose	Cmax [*] (µg/mL)	Tmax [*] (h)	AUC₁₋₃ [*] (mg/mL.h)
Solution (200 µg)	2.5±0.3	2.67±1.15	20.64±1.9
Solution (400 μg)	6.44±0.1	1.5±0.6	39.56±7.4
3D printed tablet (200 μg)	1.51±0.09	6±1.6	10.8±2
3D printed tablet (400 µg)	3.33±0.5	3.7±1	19.93±1

^{*} C_{max} , Maximum serum concentration; T_{max} , Time at which C_{max} is observed; and AUC₁₋₈, area under curve.

343



Figure 6. (A1) Rendered images and (A2) photographs of purpose designed 3D printed tablets for oral gavage in rats, (B) Plasma concentration- time profile of warfarin following the oral dosing of 200 or 400µg from (B) warfarin solution and (C) warfarin loaded 3D printed tablets to adult heathy male Sprague–Dawley rats (n=4), error bars ±SD.

Contributing to the finding above, the additional erosion step of Eudragit E in the 3D printed 348 tablets is thought to slow down the release of warfarin from the tablets. In reality, in an in vivo 349 situation, dissolution is expected to be slower than suggested by *in vitro* dissolution techniques 350 since a significantly higher pH of the stomach contents in rats pH 3.2 (fed) and pH 3.9 (fasted) 351 [34] exists compared to the *in vitro* human simulation media conditions. Furthermore, the low 352 353 fluid volume $(3.2\pm1.8 \text{ mL})$ in the fasted rats are likely to contribute to slower dissolution rates of the methacrylate polymer *in vivo* than *in vitro*. The longer T_{max} of the tablets might also be 354 attributed to the slower transit time of the relatively large oral units in rodents as previously 355 observed to be the case for oral pellets. Such effects are likely to be minimal in healthy human 356 adults where greater volumes of gastric fluids [35, 36] and a lower pH [37] at fasted state are 357 known. In summary, when extrapolating the findings to the human situation, it should be 358 considered that such delay has been augmented by the slower erosion of cationic polymer is 359 rat gastric environments rats due to their relatively higher gastric pH and lower fluid contents 360 in comparison to humans. A key driver in the uptake and use of these polymer-rich tablets 361

(yielded by FDM 3D printing) is that they match the release from standard compressed
powdered tablets. The data we present suggests that dissolution of 3D tablets requires
acceleration. However recently, there has been reports of utilizing 3D printer geometry to
fabricate tablet with complex structure to accelerate drug release [38, 39].

On the other hand, 3D printed tablets were proportionately effective as solution 367 formulations, in that a doubling of warfarin dose from the either tablet or solution resulted in a 368 rough doubling of measured plasma exposure with AUC₁₋₈ values doubling from 20.64 ± 1.9 to 369 $39.56\pm7.4 \ \mu g/mL$ for the 200 and 400 $\mu g/mL$ solutions respectively and from 10.8 ± 2 to 370 19.93±1 µg/mL for the 200 and 400 µg 3D printed capsules, respectively (184 % versus 192% 371 372 respectively). Envisioning a future scenario, a healthcare staff member may be able to use computer software to digitally directly tailor and manufacture an individualised precision dose 373 and consequently provide plasma levels of warfarin appropriate to an individual patient's need. 374

In summary, the findings in this study clearly demonstrate the potential of 3D printing as a platform to design animal-suitable solid dosage forms and thus in principle provide a pathway for human use with the potential advantage of digitally titrating an individuals dose in response to clinical data. We have also shown the utility of a novel dissolution imaging system to give mechanistic insights into the dissolution process of a 3D-printed tablet dosage form.

381 **4.** Conclusions

This study demonstrates the flexibility of FDM 3D printers to fabricate solid dosage forms 382 to purposely suit the anatomy of an animal subject. UV imaging indicated that the erosion of 383 methacrylic matrix takes place at 16.4 and 15.2 µm/min for horizontal and vertical planes 384 385 respectively and resulted in delayed plasma exposure in comparison to warfarin solutions. Moreover, the titration of dose of a narrow therapeutic index drug, warfarin, has been 386 demonstrated in vitro and in vivo. In principle, the technology holds the promise to provide a 387 much more dynamic and responsive anticoagulant regime to suit a constantly changing 388 patient's INR profile. Such an approach can provide patients with a safer, more accurate and 389 computerised alternative to the more commonly used approach of dosing using multiple tablets 390 391 to include tablet splitting.

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- **Conflicts of interest** M A Alhnan is the innovator in patent applications WO 2016038356
- A1, WO2017072536A1 and WO2018020237A1 in the field of 3D printing of medicines.

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499 List of Figures

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501 **Figure 1.** Schematic diagram of SDi2 instrument. LED's of different wavelength are em ployed to 502 illuminate the 3D printed tablet in flow through cell filled with gastric medium. The obscuration or 503 absorbance of the sample was recorded using an Actipix detector. The medium is pre-heated to 37°C

504 before going through the Whole Dosage Flow Cell and is recirculated in a closed loop configuration.

Figure 2. Precision of 3D printing to control low dose sodium warfarin. (**A**) Images of warfarin loaded FDM 3D printed tablets with increasing dose, (**B**) Correlation between the theoretical volume and tablet mass, (**C**) warfarin dose accuracy in the 3D printed tablets and (**D**) correlation between theoretical volume and warfarin dose (n=3, ±SD).

Figure 3. Thermal analysis of Eudragit E based 3D printing filaments. **(A)** Thermal degradation profiles for Eudragit E, sodium warfarin, TCP, warfarin loaded filament and tablet, **(B)** DSC thermograph for warfarin loaded filament and tablet, **(C)** XRD spectra of Eudragit E, TCP, warfarin, and warfarin loaded filament and tablet.

Figure 4. *In vitro* release pattern of sodium warfarin from 3D printed tablets of different doses from a
USPII dissolution test in modified FaSSGF (pH 2.0) (n=3, ±SD).

Figure 5. Changes in tablet height **(A)** and width **(B)** at 0, 10, 20, 30, 40, 50 and 60 min of the flow through dissolution test using Actipix SDI 2 dissolution imaging technology. **(C)** UV absorbance image

- 517 following the illumination of follow cell containing warfarin 3D printed tablet at 255 nm wavelength.
- 518 (D) Percentage sodium warfarin release from 3D printed tablet during dissolution test (n=3, \pm SD).
- 519 **Figure 6. (A1)** Rendered images and **(A2)** photographs of purpose designed 3D printed tablets for
- oral gavage in rats, **(B)** Plasma concentration- time profile of warfarin following the oral dosing of
- 521 200 or 400µg from (B) warfarin solution and (C) warfarin loaded 3D printed tablets to adult heathy
- 522 male Sprague–Dawley rats (n=4), error bars ±SD.
- 523 List of tables

Table 1. Summary of pharmacokinetic parameters of warfarin following oral gavage of 200 or 400μg
 from sodium warfarin solution and 3D printed tablets to adult heathy male Sprague–Dawley rats.

526 Supplementary data

- **Table 1S.** Summary of length, width, height and volume of cuboid containing warfarin loaded 3D
- 528 printed tablets.

530	Tailored on demand anti-coagulant dosing: an <i>in vitro</i> and <i>in</i>
531	vivo evaluation of 3D printed purpose-designed oral dosage
532	forms
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534 535	Basel Arafat ^{1,2} , Nidal Qinna ² Milena Cieszynska ¹ , Robert T Forbes ¹ , Mohamed A Alhnan ^{1*}
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541	Supplementary Data
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546	*Corresponding author: MAlbedAlhnan@uclan.ac.uk
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Table 1S. Summary of length, width, height and volume of cuboid containing sodium warfarin loaded

552 3D printed tablets

Target dose	e Volume (mm ³)	Х	Y	Z
(µg)		(mm)	(mm)	(mm)
300	19.06	5.09	1.86	2.00
500	40.74	6.55	2.40	2.58
1000	94.93	8.68	3.18	3.42
1500	149.12	10.09	3.69	3.98
2000	203.31	11.19	4.09	4.41
2500	257.51	12.10	4.43	4.77
3000	311.70	12.90	4.72	5.08
4000	420.08	14.24	5.21	5.61
5000	528.47	15.38	5.62	6.06