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Fractional laser-assisted topical delivery of bleomycin quantified by LC-MS and visualized by MALDI mass spectrometry imaging


ABSTRACT

Bleomycin exhibits antiproliferative effects desirable for use in dermato-oncology but topical use is limited by its 1415 Da molar mass. Ablative fractional laser (AFL)-assisted drug delivery has been shown to enhance drug uptake in skin. The aim of this study was with AFL to deliver bleomycin into skin, quantify uptake, and visualize biodistribution with mass spectrometry. In a Franz diffusion cell study, pig skin samples (n=66) were treated with AFL (λ=10,600 nm), 5% density, and 0, 5, 20, or 80 mJ/microbeam (mb) pulse energies before exposure to bleomycin for 0.5, 4, or 24 h. Bleomycin was quantified in biopsy cryosections at depths of 100, 500, and 1500 μm using high-performance liquid chromatography-mass spectrometry (LC-MS), and drug biodistribution was visualized for 80 mJ/mb samples by matrix assisted laser desorption/ionization mass spectrometry imaging (MALDI-MSI). The pulse energies 5, 20, and 80 mJ/mb resulted in microscopic ablation zones (MAZs) reaching superficial, mid, and deep dermis respectively. Bleomycin was successfully delivered into the skin and deeper MAZs and longer exposure time resulted in higher skin concentrations. After 24 h, AFL exposure resulted in significant amounts of bleomycin throughout all skin layers (>510 μg/cm², p≤0.002). In comparison, concentrations in intact skin exposed to bleomycin remained below limit of quantification. MALDI-MSI supported the quantitative LC-MS results by visualizing bleomycin biodistribution and revealing high uptake around MAZs with delivery into surrounding skin tissue. In conclusion, topical drug delivery of the large and hydrophilic molecule bleomycin is feasible, promising, and should be explored in an in vivo setting.

Introduction

Pretreatment with ablative fractional lasers (AFL) shows promise to improve topical therapies by enhancing drug delivery through the skin (Haedersdal et al., 2016). AFL targets water and in precise microbeams ablates tissue, resulting in laser channels termed microscopic ablation zones (MAZ), with surrounding vertical coagulation zones (CZ) of variable thickness (Hantash et al., 2007). Drug properties such as high molecular weight (>500 Da) and water solubility pose challenges to uptake through the lipid bilayers in the stratum corneum of intact skin (Morrow et al., 2007; Aulton, 2013). AFL-assisted drug delivery has been shown to facilitate topical drug uptake of anticancer agents including 5-FU (130 Da) (Wenande et al., 2017), cisplatin (300 Da) (Wenande et al., 2018), vismodegib (431 Da) (Olesen et al., 2018), and methotrexate (455 Da) (Taudorf et al., 2015), all of which have substantially smaller masses than bleomycin (1415 Da and 1426 Da for bleomycin A2 and B2, respectively).

Bleomycin is a cytotoxic glycopeptide used in the treatment of a variety of malignant diseases. The drug's mechanism of action includes intercalation in DNA and chelation with transition metals to induce reactive oxygen species (Saitta et al., 2008; Povirk et al., 1979). With chemical properties including molar masses of 1415–1426 Da and a predicted logP of −7.5 (Kim et al., 2016), bleomycin cannot cross the skin barrier unassisted to any significant degree.

Bleomycin is approved for systemic use in the treatment of squamous cell carcinoma, lymphoma, and testicular cancer (Saitta et al., 2008), and has been used off-label with intraleisional administration for a wide variety of malignant and nonmalignant conditions in dermatology, e.g. basal-cell carcinoma (Glass et al., 1997) and keloids (Berman et al., 2017). AFL-assisted drug delivery of bleomycin may be an alternative to intravenous and intraleional administration, to enable a homogenous biodistribution in skin, and minimize systemic effects.

We aim to explore topical delivery of the large and hydrophilic drug bleomycin after AFL delivery, quantify skin uptake with high-performance liquid chromatography-mass spectrometry (LC-MS), and visualize drug biodistribution in superficial (100 μm), mid (500 μm), and deep (1500 μm) dermis.
using matrix assisted laser desorption/ionization mass spectrometry imaging (MALDI-MSI).

Materials and methods

Study design

Delivery of bleomycin by AFL treatment, reaching three different laser channel depths, was examined in an in vitro Franz diffusion cell model. Pig skin samples were exposed to bleomycin for 0.5, 4, or 24 h. A total of eleven interventions were tested in six repetitions (n = 66) including non-AFL bleomycin interventions and saline controls (Table 1). LC-MS was used to quantitate bleomycin in cryosections from superficial, mid, and deep dermis (100, 500, and 1500 μm depths) along with receiver and donor fluids. MALDI-MSI imaging visualized cutaneous bleomycin biodistribution (100, 500, and 1500 μm depths) in skin samples treated with the deepest laser channels.

Skin preparation

Full-thickness skin samples from the flanks of two pigs (female, 46 and 44 kg) were utilized in the study. Skin was excised immediately after euthanasia, trimmed for hair, and relieved of subcutaneous fat. The skin was cut into square pieces of approximately 3 x 3 cm, stored for up to a month at −80°C and thawed to room temperature at study initiation.

Ablative fractional laser

AFL treatment was performed with a fractional CO₂-laser (10,600 nm, DeepFx handpiece, Lumenis Inc., Santa Clara, CA, USA). Laser settings comprised single pulses of 5, 20, or 80 mL/microbeam (mb) pulse energy at 5% density, generating MAZs reaching superficial-, mid- and deep dermis.

Franz diffusion cell model

Immediately after AFL treatment, full-thickness skin samples were mounted on a Franz cell model (PermeGear Inc., Hellertown, PA, USA) consisting of a donor and receiver compartment. Skin sample stratum corneum (0.64 cm²) faced donor compartments, while receiver compartments contained 5.5–5.8 mL phosphate buffered saline (pH 7.4, 37°C) and a magnetic stir bar. At baseline, 750 μL of bleomycin sulfate (Baxter, Deerfield, Chicago, USA) (15,000 IU/mL = 7.06 mg/mL) was topically applied to skin samples via donor compartments.

At 0.5, 4, or 24 h diffusion time, skin samples were dismounted, padded dry, and biopsied using an 8-mm punch. A biopsy from each sample was then mounted on Tissue-Tek (Sakura, Alphen, Netherlands) for LC-MS quantitation and laser channel morphology microscopy. The biopsies for MALDI-MSI were mounted on Milli Q water (ELGA CENTRA R200, ELGA LabWater, Buckinghamshire, United Kingdom). Biopsies were sectioned at skin depths of 100-, 500- and 1500 μm to produce samples of 30 μm thickness for LC-MS and 10 μm for MALDI-MSI. Samples were stored at −80°C prior to analysis. For laser channel morphology, cryosections were stained with hematoxylin-eosin and photo documented with ProgRes Capture Pro v. 2.8.8 (Jenoptik, Jena, Germany). Laser channel and coagulation-zone dimensions were digitally measured with ImageJ software v. 1.52c (National Institute of Health, Maryland, USA).

LC-MS

Before LC-MS quantitation, donor and receiver fluids were collected directly from the Franz cells. For skin samples, bleomycin was initially extracted from cryosections for 2 h in 1 mL PBS, rotating vials every 30 min. All samples (250 μL) were then precipitated with 500 μL 2% ZnSO₄·7H₂O in 25% methanol, centrifuged at 20,000 rcf for 5 min at 4°C and the supernatant was analyzed by LC-MS. Standards underwent equivalent analysis.

Quantitative analysis of bleomycin in skin sections, donor, and receiver compartments was performed using an Agilent
Laser channel morphology and dimensions in HE-stained cryosections. (A) MAZ-Supf, depth: 195 μm, width: 89 μm (80–101 μm), and coagulation zone: 34 μm (28–35 μm), n = 17 channels. (B) MAZ-Mid, depth: 602 μm (513–665 μm), width: 140 μm (118–160 μm), and coagulation zone: 47 μm (43–51 μm), n = 21 channels. (C) MAZ-Deep, depth: 1496 μm (1324–1599 μm), width: 251 μm (197–286 μm), and coagulation zone: 65 μm (60–76 μm), n = 21 channels. The coagulation-zones bordering the laser channels are visualized by more intense hematoxylin staining than the surrounding tissue. Dimensions are shown as median with interquartile range. Magnification ×40. MAZ Supf: Microscopic ablation zone reaching superficial dermis; MAZ Mid: Microscopic ablation zone reaching mid dermis; MAZ Deep: Microscopic ablation zone reaching deep dermis.

**Bleomycin imaging in skin**

Qualitative determination of the distribution of bleomycin was performed on a Thermo QExactive Orbitrap mass spectrometer equipped with an AP-SMALDI-10 ion source (TransMIT, Giessen, Germany). MALDI imaging was performed on 12 samples and 4 control tissue sections. Prior to analysis, 10 μm sections of frozen tissue, both horizontal at 100, 500 and 1500 μm depth and selected vertical sections, were thawed in a vacuum desiccator and spray-coated with a 30 mg/mL solution of 2,5-dihydroxybenzoic acid in 50:50 methanol/water with 1% trifluoroacetic acid (Hsieh et al., 2007). 300 μL of matrix solution was pneumatically sprayed at a pressure of 2.5 bar, a flow rate of 30 μL/min and a distance between the sample and spray tip of 110 mm to form a homogenous layer of matrix crystals covering an area of approximately 18 mm in diameter. While applying the matrix solution, the sample was rotated at 600 rpm. The mass spectrometer was operated in positive ion mode (scan range m/z 400–1600) at mass resolving power of 14,000 @ m/z 200. A mass accuracy of 1 ppm was ensured by using a matrix peak as lock-mass. The samples were imaged with pixel sizes of 40 μm for horizontal tissue sections and 35 μm for vertical tissue sections. The control tissue sections were analyzed using a pixel size from 100–120 μm, depending on the size of the section. Raw data was converted to imzML (Imaging mass spectrometry markup language) using an imzML converter (Schramm et al., 2012). Images were generated in MSiReader 0.06 using a bin width of 0.02 Da (Robichaud et al., 2013). The image analysis was performed using accurate masses of the two forms of bleomycin in positive ion mode (m/z 1415.52609 and 1425.56323) and the tissue was localized on basis of the localization of the endogenous phospholipid PC(34:1) (m/z 798.54092).

**Units and calculations**

Skin bleomycin concentrations (μg/cm³ and IU/cm³) were calculated based on cryosection dimensions of a halved biopsy with a surface area of 0.25 cm² and thickness of 0.003 cm. Receiver concentrations (μg/cm³) were calculated based on the exposed skin area of the Franz cell setup (0.64 cm²). Percentage of applied concentration present in a cryosection was based on the donor chamber concentration of bleomycin (7.06 mg/mL). In the result section, all concentrations mentioned are median values of 6 repetitions and p-values represent the difference between the intervention and control (non-AFL bleomycin treated skin for 24 h).

**Figure 1.** Laser channel morphology and dimensions in HE-stained cryosections. (A) MAZ-Supf, depth: 195 μm (158–219 μm), width: 89 μm (80–101 μm), and coagulation zone: 34 μm (28–35 μm), n = 17 channels. (B) MAZ-Mid, depth: 602 μm (513–665 μm), width: 140 μm (118–160 μm), and coagulation zone: 47 μm (43–51 μm), n = 21 channels. (C) MAZ-Deep, depth: 1496 μm (1324–1599 μm), width: 251 μm (197–286 μm), and coagulation zone: 65 μm (60–76 μm), n = 21 channels. The coagulation-zones bordering the laser channels are visualized by more intense hematoxylin staining than the surrounding tissue. Dimensions are shown as median with interquartile range. Magnification ×40. MAZ Supf: Microscopic ablation zone reaching superficial dermis; MAZ Mid: Microscopic ablation zone reaching mid dermis; MAZ Deep: Microscopic ablation zone reaching deep dermis.
2,5-dihydroxybenzoic acid and trifluoroacetic acid were purchased from Sigma-Aldrich. Methanol was from ChemSolute/TH Geyer. Milli Q water was produced on a Millipore Direct-Q3 (Merck, Darmstadt, Germany).

**Statistical analyses**

Descriptive statistics are presented as medians with interquartile ranges. Testing for statistical significance (5% level), unpaired samples were compared using the non-parametric Kruskal–Wallis and Mann–Whitney U tests. If concentrations were below LOQ (75 ng/ml) the LOQ value was used in the calculations. All p-values are exact values and two-tailed. Statistical tests were computed in SPSS 24 (IBM Corporation, Armonk, NY, USA). Graphical presentation of data was plotted using GraphPad Prism 7 (GraphPad Software Inc., San Diego, CA, USA).

**Results**

**Laser channel dimensions**

Pulse energies of 5, 20 and 80 mJ/microbeam resulted in median MAZs reaching the superficial- (MAZ-Supf; 195 μm depth), mid- (MAZ-Mid; 602 μm depth) and deep dermis (MAZ-Deep; 1496 μm depth). Histological evaluations of MAZs are shown in Figure 1.

**LC-MS quantified bleomycin concentrations**

Overall, drug uptake depended on MAZ depth and exposure time, resulting in higher bleomycin concentrations using deeper MAZs and longer exposure times (Figure 2 and Table 2).

After 24 h of drug exposure, all three MAZs depths provided significantly increased bleomycin concentrations in all skin layers (p ≤ .002). Thus, MAZ-Supf, MAZ-Mid and MAZ-Deep delivered concentrations of 722, 1107, and 1658 μg/cm³ bleomycin in superficial dermis, 670, 1104, and 1918 μg/cm³ in mid dermis, and 510, 680, and 1100 μg/cm³ bleomycin in deep dermis, respectively (Table 2 and Figure 2(C)). In non-AFL-exposed skin, bleomycin concentrations remained below LOQ in all skin layers.

At shorter exposure times, MAZs similarly facilitated significant concentrations of bleomycin in the superficial and mid dermis, though not all MAZ depths resulting in bleomycin in the deep dermis. After 0.5 h of drug exposure, bleomycin reached the superficial dermis with MAZ-Supf (109 μg/cm³, p=n.s.), MAZ-Mid (204 μg/cm³, p=.026) and MAZ-Deep (302 μg/cm³, p=.002). As seen in Table 2 and Figure 2(A), after short-term 0.5 h of drug exposure, only MAZ-Deep delivered bleomycin into mid dermis (150 μg/cm³, p=.041), and none of the MAZ depths facilitated delivery into deep dermis.

After 4 h of topical drug exposure, further increases in bleomycin were observed in the superficial dermis with MAZ-Supf (287 μg/cm³, p=.004), MAZ-Mid (484 μg/cm³, p=.002) and MAZ-Deep (695 μg/cm³, p=.002). Bleomycin was also detected in the mid dermis with MAZ-Supf (211 μg/cm³, p=.041), MAZ-Mid (434 μg/cm³, p=.002) and MAZ-Deep (547 μg/cm³, p=.026). In deep dermis, MAZ-Mid and MAZ-Deep delivered 163 μg/cm³ (p=.015) and 183 μg/cm³ (p=.093) bleomycin, respectively (Table 2 and Figure 2(B)), while MAZ-Supf delivery was below LOQ (data not shown).

Bleomycin concentrations in the transdermal receiver chambers were below LOQ or negligible after 0.5 and 4 h of drug exposure with all MAZs. After 24 h, bleomycin reached quantifiable concentrations (Table 2). For all interventions, donor compartments contained more than 75% of the originally applied bleomycin concentration at examined time points (data not shown). Normal saline controls were negative for bleomycin by LC-MS analysis.

![Figure 2](image-url)
In Figure 3, enhancement ratios depending on MAZ depth and exposure time are shown as a heatmap of investigated skin layers. As illustrated, skin pretreatment with AFL enabled topical delivery of bleomycin with deeper MAZs resulting in enhanced delivery. Longer drug exposure times further enhanced concentrations. Four hours drug exposure time using MAZ-Deep thus offered the same enhancement ratio in superficial skin as 24 h exposure with MAZ-Supf.

### Table 2. Quantification and comparison of bleomycin in intervention and control skin.

<table>
<thead>
<tr>
<th>Intervention</th>
<th>n</th>
<th>Sample</th>
<th>Quantified bleomycin concentration (median)</th>
<th>Percentage of applied conc.</th>
<th>Compared with control</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>LC-MS(^a)</td>
<td>per cm(^2) skin(^b)</td>
<td>Receiver(^c)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>ng/mL (IQR)</td>
<td>µg/cm(^3)</td>
<td>IU/cm(^3)</td>
</tr>
<tr>
<td>No MAZ</td>
<td>6</td>
<td>100 µm</td>
<td>LOQ (LOQ–114.0)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>500 µm</td>
<td>LOQ (LOQ–101.0)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>1500 µm</td>
<td>LOQ (LOQ)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Receiver</td>
<td>LOQ (LOQ–85.0)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>MAZ-Supf</td>
<td>0.5 h</td>
<td>100 µm</td>
<td>82.0 (LOQ–101.0)</td>
<td>109</td>
<td>232</td>
</tr>
<tr>
<td></td>
<td>500 µm</td>
<td>LOQ (LOQ)</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>1500 µm</td>
<td>LOQ (LOQ)</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Receiver</td>
<td>LOQ (LOQ)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>4 h</td>
<td>100 µm</td>
<td>215.5 (176.0–234.0)</td>
<td>287</td>
<td>610</td>
</tr>
<tr>
<td></td>
<td>500 µm</td>
<td>158.5 (151.0–164.0)</td>
<td>211</td>
<td>449</td>
<td>3.0</td>
</tr>
<tr>
<td></td>
<td>1500 µm</td>
<td>LOQ (LOQ)</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Receiver</td>
<td>LOQ (LOQ)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>24 h</td>
<td>100 µm</td>
<td>541.5 (537.0–545.0)</td>
<td>722</td>
<td>1533</td>
</tr>
<tr>
<td></td>
<td>500 µm</td>
<td>503.0 (438.0–604.0)</td>
<td>671</td>
<td>1424</td>
<td>9.5</td>
</tr>
<tr>
<td></td>
<td>1500 µm</td>
<td>382.5 (254.0–427.0)</td>
<td>510</td>
<td>1083</td>
<td>7.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Receiver</td>
<td>LOQ (LOQ–972.0)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>MAZ-Mid</td>
<td>0.5 h</td>
<td>100 µm</td>
<td>153.0 (148.0–173.0)</td>
<td>204</td>
<td>433</td>
</tr>
<tr>
<td></td>
<td>500 µm</td>
<td>LOQ (LOQ–114.0)</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>1500 µm</td>
<td>LOQ (LOQ)</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Receiver</td>
<td>LOQ (LOQ)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>4 h</td>
<td>100 µm</td>
<td>363.5 (281.0–424.0)</td>
<td>489</td>
<td>1029</td>
</tr>
<tr>
<td></td>
<td>500 µm</td>
<td>325.5 (303.0–355.0)</td>
<td>434</td>
<td>922</td>
<td>6.1</td>
</tr>
<tr>
<td></td>
<td>1500 µm</td>
<td>122.3 (97.0–167.0)</td>
<td>163</td>
<td>347</td>
<td>2.3</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Receiver</td>
<td>LOQ (LOQ)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>24 h</td>
<td>100 µm</td>
<td>830.5 (759.0–924.0)</td>
<td>1107</td>
<td>2352</td>
</tr>
<tr>
<td></td>
<td>500 µm</td>
<td>828.0 (702.0–933.0)</td>
<td>1104</td>
<td>2345</td>
<td>15.6</td>
</tr>
<tr>
<td></td>
<td>1500 µm</td>
<td>510.5 (402.0–582.0)</td>
<td>681</td>
<td>1446</td>
<td>9.6</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Receiver</td>
<td>LOQ (LOQ–2001.0)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>MAZ-Deep</td>
<td>0.5 h</td>
<td>100 µm</td>
<td>227.0 (215.0–261.0)</td>
<td>303</td>
<td>643</td>
</tr>
<tr>
<td></td>
<td>500 µm</td>
<td>112.5 (104.0–185.0)</td>
<td>150</td>
<td>319</td>
<td>2.1</td>
</tr>
<tr>
<td></td>
<td>1500 µm</td>
<td>LOQ (LOQ)</td>
<td>–</td>
<td>–</td>
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<td></td>
<td></td>
<td>Receiver</td>
<td>LOQ (LOQ)</td>
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<td>–</td>
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<tr>
<td></td>
<td>4 h</td>
<td>100 µm</td>
<td>521.0 (476.0–621.0)</td>
<td>695</td>
<td>1427</td>
</tr>
<tr>
<td></td>
<td>500 µm</td>
<td>410.0 (309.0–552.0)</td>
<td>547</td>
<td>1161</td>
<td>7.7</td>
</tr>
<tr>
<td></td>
<td>1500 µm</td>
<td>137.5 (LOQ–196.0)</td>
<td>183</td>
<td>389</td>
<td>2.6</td>
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<td></td>
<td></td>
<td>Receiver</td>
<td>LOQ (LOQ)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>24 h</td>
<td>100 µm</td>
<td>1244.0 (1010.0–1400.0)</td>
<td>1659</td>
<td>3522</td>
</tr>
<tr>
<td></td>
<td>500 µm</td>
<td>1438.5 (1035.0–1635.0)</td>
<td>1918</td>
<td>4073</td>
<td>27.2</td>
</tr>
<tr>
<td></td>
<td>1500 µm</td>
<td>825.0 (676.0–883.0)</td>
<td>1100</td>
<td>2336</td>
<td>15.6</td>
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<tr>
<td></td>
<td></td>
<td>Receiver</td>
<td>4380.5 (3775.0–5178.0)</td>
<td>383</td>
<td>–</td>
</tr>
</tbody>
</table>

MAZ: Microscopic ablation zone.
MAZ-Supf: Microscopic ablation zone reaching superficial dermis.
MAZ-Mid: Microscopic ablation zone reaching mid dermis.
MAZ-Deep: Microscopic ablation zone reaching deep.
LOQ: Below limit of quantification (75 ng/mL).
IQR: Interquartile range (Q1-Q3).
\(^a\)Concentration as measured with LC-MS.
\(^b\)Cryosection volume is 0.00075 cm\(^3\) (surface area 0.25 cm\(^2\) \times thickness 0.003 cm).
\(^c\)Surface area is 0.64 cm\(^2\).

**Enhancement ratios**

In Figure 3, enhancement ratios depending on MAZ depth and exposure time are shown as a heatmap of investigated skin layers. As illustrated, skin pretreatment with AFL enabled topical delivery of bleomycin with deeper MAZs resulting in enhanced delivery. Longer drug exposure times further enhanced concentrations. Four hours drug exposure time using MAZ-Deep thus offered the same enhancement ratio in superficial skin as 24 h exposure with MAZ-Supf.

**MALDI mass spectrometry imaging**

Qualitative imaging revealed bleomycin concentrating primarily in CZs, though also present in surrounding tissue (Figure 4(A)). Supporting quantitative LC-MS data, a trend of higher concentrations with increased drug exposure time was seen. The highest concentrations of bleomycin were noted in the superficial dermis, with decreasing intensity down through the skin layers.

Drug imaging of the superficial dermis (Figure 4(Ba–Bc)) indicated increasing concentrations of bleomycin over time.
Despite poorly defined cryosection outlines, imaging also revealed visible laser channels in analyzed samples.

Mid dermis imaging (Figure 4(Bd–Bf)) showed well-defined skin sample outlines with an organized grid of MAZs. Peak bleomycin concentrations were detected corresponding to CZs and a trend for higher concentrations is seen with increasing drug exposure time.

Deep dermis imaging (Figure 4(Bg–Bi)) showed no visible bleomycin in the center of the 0.5-hour image but the drug is present peripherally (Figure 4(Bg)). This corresponds to the quantitative analysis by LC-MS being below LOQ. The 4 and 24 h images (Figure 4(Bh,Bi)) show well-defined outlines and grid-organized laser channels with a trend of higher bleomycin concentrations with increasing drug exposure time.

No bleomycin was detected in negative control samples with normal saline (data not shown).

Discussion

The present study is the first assessment of AFL-assisted topical delivery of bleomycin quantified with LC-MS and visualized with MALDI-MSI. Bleomycin was successfully delivered into the deep dermis by AFL while drug detection in non-AFL exposed skin was negligible even at superficial depths (Table 2 and Figure 2(C)). Bleomycin concentrations depended on MAZ depth, and higher concentrations of drug in tissue were achieved using deeper MAZs and longer drug exposure times. As expected, a non-linear relationship between drug uptake and MAZ depth as well as exposure time was observed, indicating a saturation level of bleomycin in the skin similar to data published for other drugs previously delivered by AFL in vitro, e.g. 5-FU (Wenande et al., 2017) and methotrexate (Taudorf et al., 2015).

The quantitative LC-MS results are supported by the qualitative MALDI-MSI. Furthermore, the qualitative imaging data provided insight in biodistribution and confirms high uptake of bleomycin in CZs as well as substantial delivery in the surrounding skin tissue. It has been hypothesized that AFL-induced CZs function as a barrier against drug uptake (Erlandsson et al., 2016). However, in the present study, we utilized MALDI imaging to demonstrate substantial biodistribution beyond CZs, similar to what has previously been shown with smaller drugs than bleomycin, e.g. 5-FU (Wenande et al., 2017). Thus, our data is in line with previous research which shows high uptake in CZs with other topically-applied compounds (Haak et al., 2017).

AFL-assisted delivery of bleomycin appears to be a promising candidate for a new topical treatment with uniform and reproducible biodistribution. The AFL modality is associated with less interphysician variance than might be expected with current local treatment methods, e.g. injection delivery. AFL-assisted drug delivery may also enable precision delivery to specific skin depths and might also limit transdermal access to the vascular system and systemic effects; an advantage of local therapy. Although cutaneous biodistribution is highly dependent on drug concentrations and characteristics, a recent in vivo study investigating AFL-assisted cisplatin (300 Da) and 5-FU (130 Da) delivery did not find any quantifiable transdermal uptake (Wenande et al., 2018), which indicates that systemic access might be limited depending on the drug and treated area.

AFL-assisted drug delivery in a clinical setting can reduce the need for long drug exposure times. After just 30 min, deep laser channels yielded almost the same drug uptake enhancement ratio in superficial skin as superficial channels after 4 h drug exposure. Thus, increasing MAZ-depth may dramatically reduce drug exposure time.

Compared with reported IC50 test results, our data suggests that AFL-assisted drug delivery of bleomycin can provide clinically relevant concentrations with all tested MAZs. Two studies listed in the Genomics in Drug Sensitivity Cancer (GDSC) project database differ greatly in their IC50 test results for bleomycin on a wide range of cell lines. For the often-used SCC-25 cell line, reported values were 0.97 μg/cm³ (0.69 μM) and 219 μg/cm³ (155 μM), respectively (Yang et al., 2012).

Another study on the same cell line suggests 15.0 μg/cm³ (10.7 μM) after 24 h (Olesen et al., 2017a). Importantly, all three values can easily be reached using AFL under in vitro conditions.

We used the standard vehicle for bleomycin for our Franz diffusion cell study, an aqueous saline solution. Topical application in the clinic may benefit from a more viscous
formulation in order to adhere securely on the skin for the duration of the treatment, e.g. a gel-based vehicle. In a recent study, we compared different types of vehicles for use with laser-assisted delivery and found that a gel-based vehicle enters the skin almost as effectively as a liquid solution (Olesen et al., 2017b). Future in vivo studies might investigate bleomycin’s efficacy and compatibility with different vehicles compared with normal saline. Other factors including drug loading in skin (Karadzovska et al., 2012), closing of laser channels over time (Banzhaf et al., 2017), and vascular wash-out effects also represent areas of further investigation, though best studied in vivo (Oni et al., 2012; Ibrahim et al., 2018).

A limitation of this in vitro study represents the almost infinite supply of bleomycin dissolved in normal saline available to the skin during exposure. This does not necessarily translate to clinical settings, but may emulate continuous application under

Figure 4. Mass spectrometry imaging. MALDI-MSI of bleomycin B2 (m/z 1425.56323) with MAZ-Deep laser channels. (A) Vertically cut skin cryosection after 24 h of topical drug exposure. Laser channels are easily seen with high concentrations of bleomycin (yellow) in the coagulation zones and drug dissemination into the surrounding tissue. (B) Horizontally cut skin cryosections. A trend towards higher concentrations is seen along the x-axis of time, and lower down the y-axis of skin depth. The depicted intensity values for each image are based on maximum bleomycin detection within the individual skin sample and thus cannot be compared inter-individually. MALDI-MSI: Matrix assisted laser desorption/ionization mass spectrometry imaging; MAZ-Deep: Microscopic ablation zone reaching deep dermis.
occlusion during a similar time-frame in the clinic. Furthermore, Figures 4Bg-Bi suggest that punch biopsy can cause depression of peripheral tissue from the skin surface downwards, resulting in cryosections from deep skin compartments displaying tissue from more superficial layers in their outer rims. Higher drug concentrations would in consequence be visualized in the periphery of the cryosections, a phenomenon similarly observable in earlier studies (Wenande et al., 2017). Finally, a few samples of non-AFL-exposed skin were shown to contain bleomycin. However, this finding most likely reflects inadvertent contamination during handling and these low concentrations near the LOQ had a limited impact on our findings.

In conclusion, we show for the first time that AFL-assisted topical drug delivery of the large hydrophilic molecule bleomycin is feasible. Cutaneous bleomycin concentrations can be increased by using deeper MAZs and longer exposure times. Imaging further reflected the quantitative LC-MS data by visualizing bleomycin distributed in CZs and surrounding tissue, as well as higher drug concentrations associated with increased exposure time. AFL-assisted bleomycin delivery appears to be a promising candidate for future in vivo studies.

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