Public comment

Improved stratum corneum sampling *in vivo* delivers <u>added value</u> for topical bioequivalence assessment

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SC sampling: Added value for BE assessment

Translational methodology for *in vitro* (IVPT) observations

- Drug/formulation specific in vitro-in vivo correlation (IVIVC)
- Simpler than PK; available when plasma levels are too low for PK
- Simpler than open flow microperfusion/microdialysis
- Measures drug delivery rate from SC
 - Measure mass of drug in SC after period of clearance (¥)
 - Compare to mass of drug in SC at end of uptake (
- Calculate the average flux from the SC to deeper tissues

Average Flux =
$$\frac{\left(M_{Up} - M_{Clear}\right)/A}{t_{Clear} - t_{Up}}$$



SC sampling in vivo: Example 1

DICLOFENAC SODIUM

Compare 3 products (all Q1 different)

- 2% solution (Pennsaid) 10 mg/cm² (contains DMSO)
- ♦ 3% gel (Solaraze) 20 mg/cm²
- 1% gel (Voltaren) 10 mg/cm²
- 17 h clearance after 6 h uptake
- 14 subjects

SC sampling: Mass and BE assessment

DICLOFENAC SODIUM



Uptake: Closed symbols Clearance: Open symbols

Error bars, 90% CI of the log mean

SC sampling: Average clearance flux

DICLOFENAC SODIUM



IVPT data
■ △ Bath pig (n=4)
■ △ Yucatan mini-pig (n=4)

Error bars, 90% CI

In vitro permeation test (IVPT) 30 Cumulative Penetration, mcg cm⁻² Diclofenac Bath pig (n=4) 20. Ρ 10 -0 10 20 40 50 30 0 Time post drug application, h Calculate from mass permeated over comparable interval (8 - 24 h)

Error bars, 1 SD

SC sampling in vivo: Example 2

ACYCLOVIR

Compare 3 creams (5%) in 2 trials

- Trial 1
 - US Zovirax (US)
 - UK Zovirax (UK)
- Trial 2
 - Aciclovir 1A Pharma (AT)
 - US Zovirax (US)
- 15 mg/cm²
- 17 h clearance after 6 h uptake
- 10 subjects/trial

SC sampling: *Mass and BE assessment*

ACYCLOVIR





SC sampling: BE assessment compared to dOFM

Open Flow Microperfusion (dOFM)*



Uptake: Closed symbols Clearance: Open symbols

*Bodenlenz M et al. *Clin Pharmacokinet,* 56:91-98 (2017)

Error bars, 90% CI of the log mean

BE Ratio

SC sampling: Average clearance flux

ACYCLOVIR





Flux from SC sampling similar for US, UK & AT

Flux from IVPT for US & UK also similar



Time post drug application, h

Average flux from mass permeated over comparable interval (8 - 24 h)

Drug removed in SC sampling but not in IVPT may explain quantitative differences

Error bars, 90% CI

SC sampling in vivo: Example 3

DRUG Z

- 3 gel products with the same concentration of Z
 - Ref Commercial product
 - Test1 Q1 & Q2 equivalent to Ref
 - Test2 more gelling agent; otherwise Q1 & Q2 equivalent
- Identical amounts of each formulation applied
- 12 h clearance after 6 h uptake
- 14 subjects

SC sampling: *Mass and BE assessment*

DRUG Z



Uptake: Closed symbols Clearance: Open symbols

Error bars, 90% CI of the log mean

SC sampling in vivo: Valuable tool to assess BE

- Measured in humans in vivo
- Improved SC sampling protocol demonstrated to be robust and reliable across labs and operators
 - Demonstrated for 4 drugs, 3 formulations/drug, 3 labs, 5 operators (including econazole presented in presentation by Dr. Richard Guy)
 - Technically accessible and economical method
- Complementary to other surrogate assessment methods
 - IVPT, open flow microperfusion/microdialysis, plasma PK
 - <u>Obvious value</u> for drugs acting on or in the stratum corneum
 - <u>Added value</u> for drugs acting deeper in the skin

Can assess clinically-relevant topical bioavailability (BA)

 Formulation effects on skin barrier function after repeat dosing (see two slides at end of this presentation)

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SC sampling in vivo: Assess repeat dose effect

Gel vs. lotion at same strength



Nathalie Wagner PQRI, Rockville, MD March 13, 2013



Dry clean; first 2 tapes discarded; 22 tapes (n = 4) <u>One application</u>: Drug mass in SC "plateaued" by ~2 h <u>Daily application</u>: Sampled 2 h after application

SC sampling in vivo: Assess repeat dose effect

Gel vs. lotion at same strength



- Different "steady state" after 1 and multiple applications
- Measurements after a few applications on the recommended clinical schedule might be appropriate for formulations containing ingredients that affect the SC
- Multiple applications more representative of the clinical intended use