

An Overview of In Vitro BE Studies

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Disclaimer



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Learning Objectives



- Understand why an in vitro study may be used to establish bioequivalence (BE)
- Identify resources to aid with in vitro BE study selection and development
- Describe types of in vitro BE studies

Why In Vitro?

- In vivo BE studies
 - Expensive
 - Time-consuming
- In vitro BE studies
 - Can reduce risk of harm
 - May be the best method to determine BE



Key Resources

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Product-Specific Guidances (PSG)



- <u>PSGs</u> provide recommendations to support ANDA drug development
- Over 2000 PSGs available

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Reference Standard

Identify the appropriate reference listed drug (RLD) from the <u>Orange</u>
 <u>Book</u>

– See also: <u>Purple Book</u>

 Can use a different reference standard in some cases Approved Drug Products with Therapeutic Equivalence Evaluations | Orange Book

Purple Book Database of Licensed Biological Products

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Types of In Vitro BE Studies

Common In Vitro BE Studies

- In vitro permeability testing (IVPT)
- In vitro release testing (IVRT)
- In vitro binding testing
- Size distribution studies
 - In vitro globule size distribution study
 - Particle size distribution/determination (PSD) study
 - In vitro liposome size distribution study
- In vitro aerosol studies (5- or 6-test battery)

Other In Vitro Studies

- FDA
- In vitro dissolution testing for BE determination
- BCS dissolution testing
- BCS solubility testing
- BCS permeability testing
- In vitro NG/G tube study
- In vitro microbial kill rate study

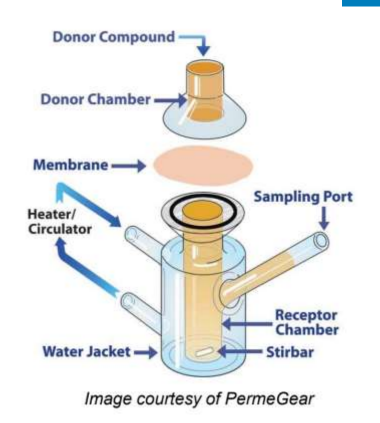
BCS-based biowaiver



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- Semi-solid topical dermatological drugs
 - In vitro BE approaches:
 IVRT and IVPT
 - In vivo BE approach:
 clinical endpoint
 - IVRT can be used for other formulations





IVRT

- Synthetic membrane
- Consistent
- Infinite dose
- Release rate
- Not expected to correlate or predict vivo BA/BE

IVPT

- Human skin
- Variable
- Finite dose
- Flux profile
- Expected to have in vitro-in vivo correlation



- Both
 - Method development
 - Method validation
- IVRT
 - Study

Α	В	А	В	А	В
В	A	В	А	В	А

- IVPT
 - Pilot study
 - Pivotal study



- Example: acyclovir 5% topical cream
 - In vitro
 - Q1 and Q2 sameness
 - Q3 physical and structural tests
 - IVRT <u>and</u> IVPT
 - In vivo
 - Clinical endpoint



- Resources
 - United States Pharmacopeia (USP) General Chapter
 <1724>, Semisolid Drug Products Performance
 Tests
 - FDA/CRCG 2021 workshop
 - In Vitro Bioequivalence Data for a Topical Product: Bioequivalence Review Perspective (Dr. Suman Dandamudi, 2017)



In Vitro Binding Testing

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In Vitro Binding Testing

- 21 CFR §320.23(b)(2)
- Phosphate or bile acid-binding drugs (GI)
- Equilibrium (pivotal) and kinetic testing
- Measure unbound analyte(s) in filtrate

In Vitro Binding Testing

Equilibrium

- ± acid pre-treatment*
- 8+ concentrations of phosphate/bile salts
- Incubate till equilibrium

Kinetic

- or ± acid pre-treatment*
- 2 concentrations of phosphate/bile salts
- 8+ lengths of time

In Vitro Binding Testing

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- Example: sucralfate oral suspension
 - Only in vitro recommended
 - Equilibrium binding study with bovine or human serum albumin
 - Equilibrium binding study with bile salts
 - Kinetic binding study with bile salts
 - In vitro enzyme (pepsin) activity study



In Vitro Aerosol Studies

In Vitro Aerosol Studies

Inhaled

- 1. Single actuation content (SAC)
- 2. Aerodynamic particle size distribution (APSD)
- 3. Spray pattern
- 4. Plume geometry
- 5. Priming and repriming

Nasal

1. SAC

- 2. Droplet size distribution by laser defraction
- 3. Drug in small particles/droplets
- 4. Spray pattern
- 5. Plume geometry
- 6. Priming and repriming

Draft Guidance



In Vitro Aerosol Studies

- Both in vitro <u>and</u> in vivo commonly recommended
- <u>Example</u>: albuterol sulfate, aerosol, metered; inhalation
 - In vitro: 5-test battery for inhaled aerosols
 - In vivo: PK study



Size Distribution Studies



Size Distribution Studies

- Globule, particle, or liposome size distribution studies
- Help ensure uniformity and consistent dosing
- Different formulation types
- Varying methods

Size Distribution Studies

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- <u>Example</u>: Cyclosporine ophthalmic emulsion
 - In vitro
 - Q1 and Q2 sameness
 - Q3 comparable
 - Globule size distribution, viscosity, pH, zeta potential, osmolality, surface tension
 - In vivo
 - Clinical endpoint
- Resource: <u>Assessment of Complex Drug Product Physicochemical</u> <u>Characteristics to Support In Vitro BE Studies</u> (Dr. Asif Rasheed, 2020)



Challenge Questions

Challenge Question #1



Which of the following statements is <u>NOT</u> true?

- A. Reference standards can be identified using the Yellow Book.
- B. Acceptable study types are described in the product specific guidances.
- C. If a reference listed drug is unavailable, FDA may select a new one to serve as a reference standard.
- D. The Purple Book details licensed biological products.



Challenge Question #2

Which of the following are components of in vitro aerosol studies?

- A. Single actuation content
- B. Spray pattern
- C. Plume geometry
- D. All of the above

Summary



- In vitro BE studies
 - can be conducted with or instead of in vivo BE studies
 - can vary greatly and are highly dependent upon the drug and formulation

Closing Thought



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Questions?

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