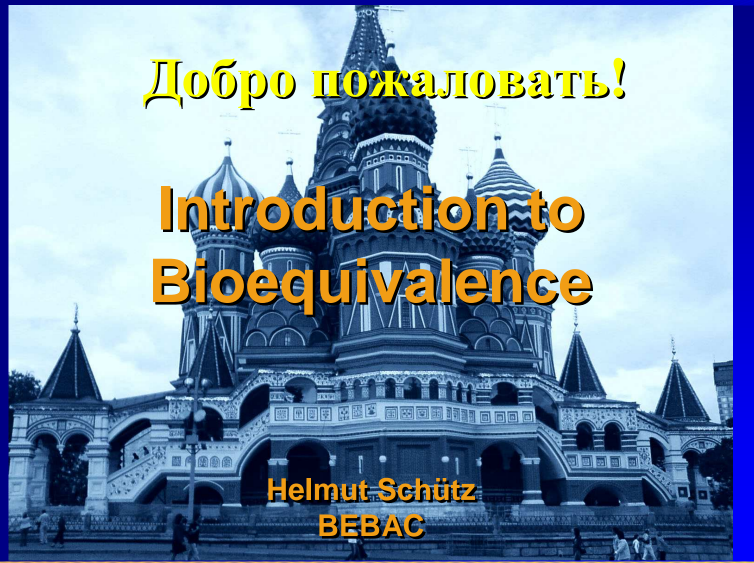


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# Answering the Question: What is Enlightenment?

"Enlightenment is man's emergence from his self-imposed immaturity for which he himself was responsible. Immaturity and dependence are the inability to use one's own intellect without the direction of another. One is responsible for this immaturity and dependence, if its cause is not a lack

Beantworfung ber Frage ; Bas ift Aufflärung ?

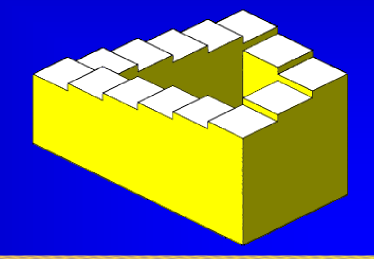
"Unfklärung ift der Ausgang des Mene ichen aus feiner felbst verschuldeten Unmundigkeit. Unmundigkeit ist das Unvermde gen, sich seines Verstandes ohne Leitung eines andern zu bedienen. Selbst verschuldet ist diese Unmuns digkeit, wenn die Ursache berfelben nicht am Mangel des Verstandes, sondern der Entschließung und des Muthes liegt, sich seiner ohne Leitung eines andern zu bedienen. Sapere auch habe Muth, dich deines eiges nen Verstandes zu bedienen! ift also der Wahlspruch der Aufklärung.

of intelligence, but a lack of determination and courage to think without the direction of another. **Sapere aude!** Have courage to use your **own** understanding! is therefore the slogan of Enlightenment." *Immanuel Kant (1784)* 

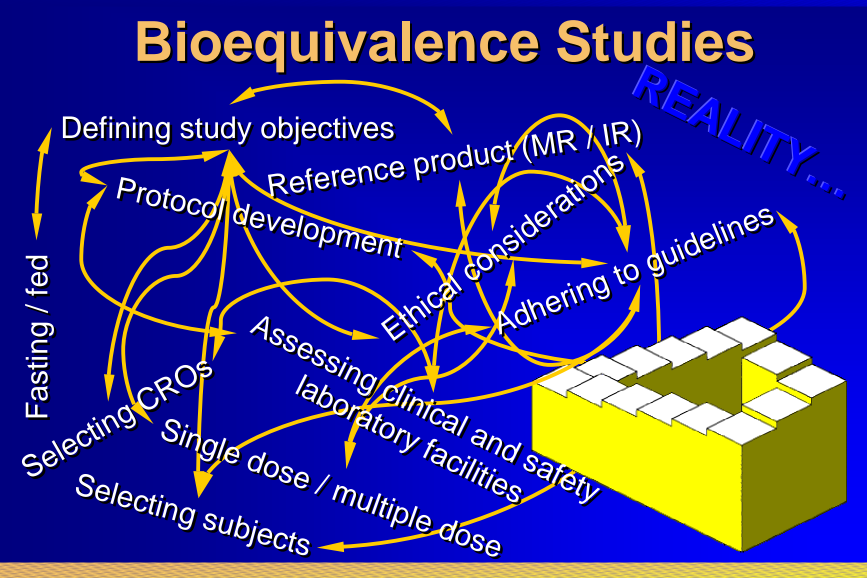


# **Bioequivalence Studies** DREAM

- Defining study objectives  $\overline{}$
- Fasting / fed
- Single dose / multiple dose
- Reference product (MR / IR)
- Selecting CROs
- Protocol development
- Ethical considerations
- Assessing clinical and safety laboratory facilities
- Selecting subjects
- Adhering to guidelines









Bioequivalence
Surrogate of clinical equivalence or
Measure of pharmaceutical quality?
Types of studies
Pharmacokinetic (PK)
Pharmacodynamic (PD)
Clinical (equivalence and/or safety/efficacy)



Types of studies (cont'd)
Healthy Subjects
Patients
Single dose
Multiple dose
Cross-over, replicate
Parallel

Reference product (MR, IR, solution)



 Types of studies (cont'd) Food effect PK interaction Design Issues Dose regimen Fasted / fed state Type of standard meals Bioanalytics (not GLP!) Parent drug / metabolite(s) / enantiomers / pro-drugs Validation / routine application



•Ethics (GCP!) Dose levels / number of administered doses Number / volume of blood samples Drug and/or adverse effects Clinical performance (GCP!) CRO selection Responsibilities of sponsor / investigator Audits / monitoring



### •NCA / PK (PD)

Sampling schedule
 Metrics (*AUC*, *C<sub>max</sub>*; *AUEC*, *Ae<sub>max</sub>*,...)
 Design, methods, evaluation

### Sample size

 Estimation from previous and/or pilot studies, literature

Highly variable drugs

### Biostatistics

- Models & assumptions
- Protocol, evaluation, report

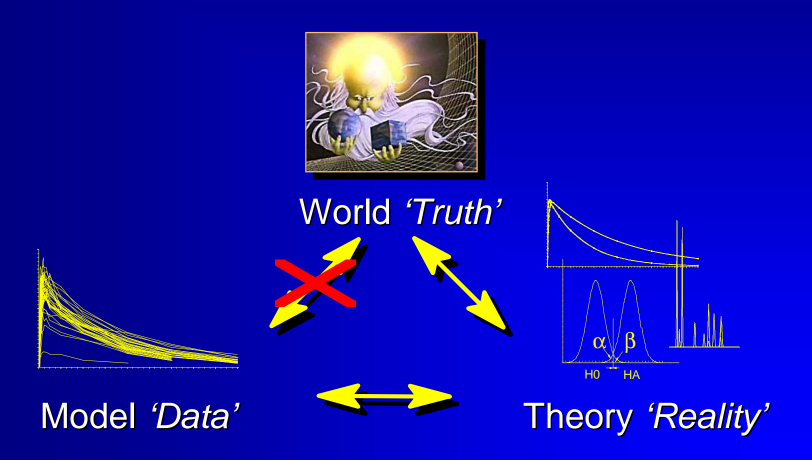




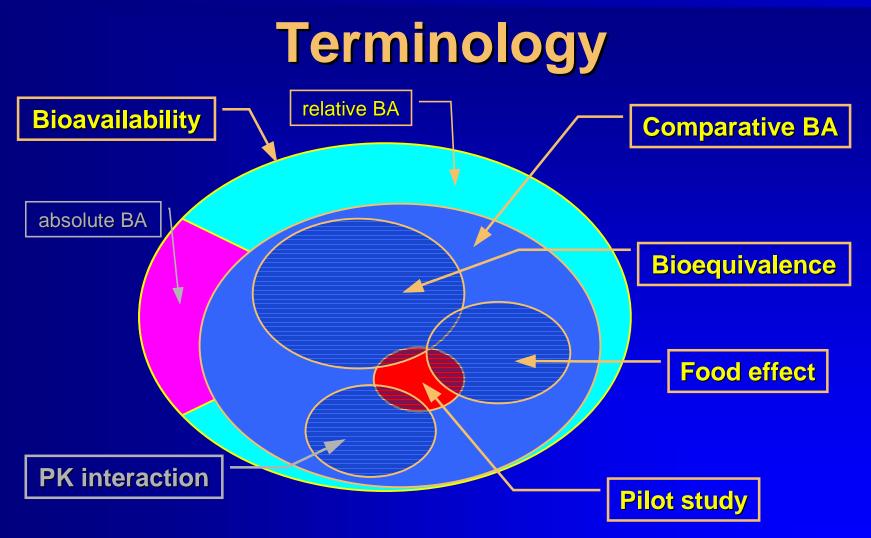
'What if'-scenarios
Common pitfalls
Blind review
'Failed' studies
Deficiency letters



## Assumptions









## **Definitions**

### •EMEA Guideline on BE (2010)

- A bioequivalence study is basically a comparative bioavailability study designed to establish equivalence between test and reference products.
  - Comparative BA,
  - designed to demonstrate BE,
  - reference = innovator's product.

### Russian BE Guideline (2008)

Two drug preparations are considered to be bioequivalent if bioavailability of drug substance is the same.

#### EMEA Human Medicines Evaluation Unit / CPMP Guideline on Investigation of Bioequivalence(2010) http://bebac.at/Guidelines.htm - EU Ministry of Health and Social Development Russian Federation Drugs Bioequivalence Evaluation (2008) http://bebac.at/Guidelines.htm - RU



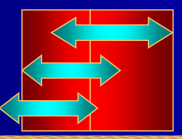
## **Bioequivalence...**

Comparative BA

- true experiment; no bibliographic comparison
- Designed to demonstrate BE
  - variability,
  - deviation of test from reference,
  - drop-out rate,...

to be able (statistical power!) to demonstrate BE

•Reference = Innovator's product



#1: BE [90%–125%]
#2: BE [80%–110%]
#3: not BE [76%–103%]; (but 'BE' to #2)



## **Bioequivalence...**

### •EMA GL on BE (2010)

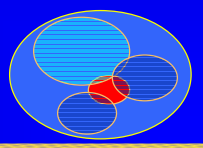
Two medicinal products containing the same active substance are considered bioequivalent if they are pharmaceutically equivalent or pharmaceutical alternatives and their bioavailabilities (rate and extent) after administration in the same molar dose lie within acceptable predefined limits. These limits are set to ensure comparable in vivo performance, i.e. similarity in terms of safety and efficacy.



## **Bioequivalence...**

### Russian GL on BE (2008)

Two drug preparations are considered to be bioequivalent if bioavailability of drug substance is the same. Bioavailability – percentage amount of the drug substance entered systemic blood flow (extent of absorption) and the rate of this process (rate of absorption).





## **Global Harmonization?**

- In almost all regulations two PK metrics are necessary to demonstrate BE, namely
  - extent  $(AUC_t \text{ or } AUC_{\infty})$  and
  - rate ( $C_{max}$ ) of exposure.
- One exception: US-FDA (where AUC, and AUC, must demonstrate extent of exposure)
  - Although stated in the GL, such a requirement is statistically flawed.
    - Multiplicity issues (what is the patient's risk?)
    - Impossible *a*-adjustment (interdependence)

There can be only one!





# **History of BE**

### Bioequivalence

- Problems first noticed with NTIDs (Narrow Therapeutic Index Drugs) in the late 1970s
- Intoxications (and even some fatallities!) were reported (warfarin, digoxin, phenytoin)
  - Warfarin, digoxin: Patients switched between formulations which were got approval solely based on *in vitro* data (innovator ↔generic)
  - Phenytoin: The innovator's API was changed from a microcrystalline to an amorphous form resulting in 10times higher plasma concentrations in steady state



# **History of BE**

#### Bioequivalence

- Surrogate of clinical equivalence (1980+)
  - Studies in steady state in order to reduce variability
  - Studies based on active metabolite
  - Wider acceptance range if clinical justifiable (not FDA!)

Measure of pharmaceutical quality (2000+)

- Single dose studies preferred
- Generally parent drug
- Widening of acceptance range exceptional (except FDA HVDs and EMA C<sub>max</sub> of HVDs)





## Early 1980s

#### First method

 FDA's 75/75 Rule BE, if 75% of subjects show ratios of 75%-125%. Not a statistic, variable formulations may pass by chance...

#### BE Cabana

Assessment of 75/75 Rule: FDA Viewpoint J Pharm Sci 72, 98-99 (1983) JD Haynes FDA 75/75 Rule: A Response J Pharm Sci 72, 99-100 (1983)

		R	T/R	75%-125%
1	71	81	87.7%	yes
2	61	65	93.8%	yes
3	80	94	85.1%	yes
4	66	74	89.2%	yes
5	94	54	174.1%	no
6	97	63	154.0%	no
7	70	85	82.4%	yes
8	76	90	84.4%	yes
9	54	53	101.9%	yes
10	99	56	176.8%	no
11	83	90	92.2%	yes
12	51	68	75.0%	yes
				75.0%



## Mid 1980s I

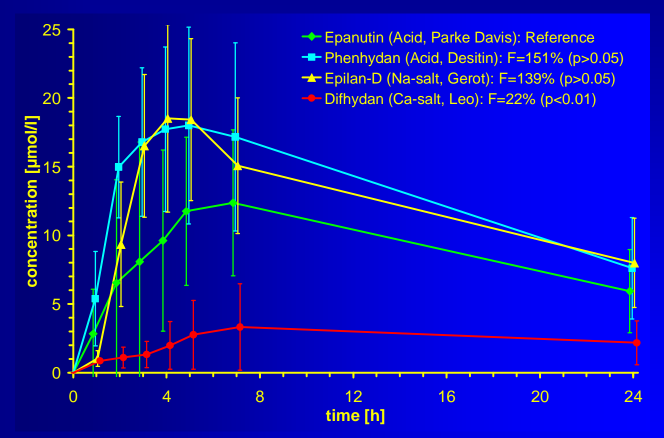
### Early method

- Testing for a significant difference (*t*-test) at α 0.05 Problem:
  - High variability in differences → formulation will pass ( $p \ge 0.05$ )
  - Low variability in differences
    - $\rightarrow$  formulation will fail (*p* < 0.05)
  - This is counterintuitive and the opposite of what we actually want!

	Т	R	T–R
1	71	81	-10
2	61	65	-4
3	80	94	-14
4	66	74	-8
5	94	54	+40
6	97	63	+34
7	70	85	-15
8	76	90	-14
9	54	53	+1
10	99	56	+43
11	83	90	-7
12	51	68	-17
mean	75	73	+2
SD	16	15	23
CV%	21.4%	20.6%	940%
		t-table	2.2010
		t-calc	0.3687
			n.s.



## Example



#### Nitsche V, Mascher H, and H Schütz

Comparative bioavailability of several phenytoin preparations marketed in Austria Int J Clin Pharmacol Ther Toxicol 22(2), 104-107 (1984)



## Mid 1980s II

### Later method FDA's 80/20 rule At least 80% power to be able to demonstrate a 20% difference (*t*-test) at $\alpha$ 0.05 Essentially the 75/75 rule in more statistical terms. Power 71.5% < 80! (not BE)</p> In any study (even at 'true' T=R) with variability $s_{1}/2/n > 6.44$ it is impossible to show BE!

	Т	R	T–R
1	71	81	-10
2	61	65	-4
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4	66	74	-8
5	94	54	+40
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10	99	56	+43
11	83	90	-7
12	51	68	-17
mean	75	73	+2
SD	16	15	23
		t-table	2.2010
		t-calc	0.3687
			n.s.
		power	71.59%



## Late 1980s

### TOST (Two One-Sided Tests)

- First formulation of the problem based on equivalence rather than a difference
  - Two One-Sided t-tests
  - Bioequivalent if p(<80%) + p(>120%) ≤0.05
  - Equivalent to a 90% confidence interval within an acceptance range of 80% – 120%

#### **DA Schuirmann**

A Comparison of the Two One-Sided Tests Procedure and the Power Approach for Assessing the Equivalence of Average Bioavailability J Pharmacokin Biopharm 15, 657–680 (1987)

	т	R	T–R
1	71	81	-10
2	61	65	-4
3	80	94	-14
4	66	74	-8
5	94	54	+40
6	97	63	+34
7	70	85	-15
8	76	90	-14
9	54	53	+1
10	99	56	+43
11	83	90	-7
12	51	68	-17
	p(<80%)		0.0069
	p(>	>120%)	0.0344
		p(total)	0.0414
		T/R	103.32%
	90%	5 CI (lo)	88.35%
	90%	5 Cl (hi)	118.30%



## Human Guineapigs I

- BE studies as a surrogate for clinical efficacy / safety ('essential similarity')
  - We want to get unbiased estimates, *i.e.*, the point estimate from the study sample ...

$$PE = \frac{\hat{X}_{Test}}{\hat{X}_{Reference}}$$



should be representative for the population of patients.

$$F_{Pop} = \frac{\mu_{Test}}{\mu_{Reference}}$$





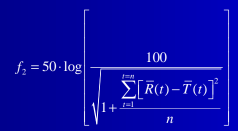
# Human Guineapigs II

- BE studies as a special case of documented pharmaceutical quality
  - The *in vivo* release in the biostudy ...

$$PE = \frac{\hat{X}_{Test}}{\hat{X}_{Reference}}$$



should be representative for the *in vitro* performance.







## Science $\rightarrow$ Regulations

- We can't compare bioavailabilities in the entire population of patients
  - Scientific Reductionism (based on assumptions)
    - Similar' concentrations in healthy subjects will lead to 'similar' effects in patients.

Equal doses and inter-occasion clearances!

$$AUC_{T} = \frac{D_{T} \cdot F_{T}}{CL_{T}}, AUC_{R} = \frac{D_{R} \cdot F_{R}}{CL_{R}}$$
$$\begin{bmatrix} D_{T} \cong D_{R}, CL_{T} \cong CL_{R} \end{bmatrix}$$
$$F_{rel}(BA) = \frac{F_{T}}{F_{R}} \cong \frac{AUC_{T}}{AUC_{R}}$$

Introduction to Bioequivalence



## Models vs. Reality





## **A Reminder**

Rose is a rose is a rose is a rose.



Gertrude Stein (1913)

Guidelines are guidelines are guidelines.

Henrike Potthast (ca. 2004)

In advanced engineering, you expected failure; you learned as much from failures as from successes – indeed if you never suffered a failure you probably weren't pushing the envelope ambitiously enough.

Stephen Baxter; Transcendent, Chapter 36 (2006)



### Thank You! Introduction to Bioequivalence Open Questions?



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