



Arbeitsgemeinschaft  
für angewandte  
Humanpharmakologie e.V.

## Pitfalls in BA/BE

Attempts in beating Murphy's law:  
Learnings from failures in study design,  
bioanalytics and statistics

# Helpful (?) quotations



If anything can go wrong, it will.

*Edward A. Murphy Jr.*

He who fails to plan is planning to fail.

*Winston Churchill*

You can't fix by analysis what you bungled by design.

*Richard J. Light,  
Judith D. Singer, John B. Willett*

100% of all disasters are failures of design, not analysis.

*Ronald G. Marks*

To propose that poor design can be corrected by subtle analysis techniques is contrary to good scientific thinking.

*Stuart J. Pocock*

To call the statistician after the experiment is done may be no more than asking him to perform a *postmortem* examination: He may be able to say what the experiment died of.

*Ronald A. Fisher*

If you think it's simple, then you have misunderstood the problem.

*Bjarne Stroustrup*

Whenever a theory appears to you as the only possible one, take this as a sign that you have neither understood the theory nor the problem which it was intended to solve.

*Karl R. Popper*

# Study design



- In a crossover-study the washout between treatments has to be sufficiently long
  - Avoid pre-dose concentrations which are residuals of previous period(s)
  - In order to get an unbiased estimate of treatment differences the physiological state of subjects in higher period(s) has to be the same as in the (drug-naïve) first period
  - Cave
    - Never plan the washout (generally  $\geq 5$  times the apparent half life) based on an average. Keep the distribution of half lives in mind. Some subjects might show a substantially longer half life – especially if the drug is subjected to polymorphism (poor and extensive metabolizers).
    - Think also about PD. If the drug is an auto-inducer or -inhibitor allow the body to return to its original state.

# Case 1 | Study design



- Drug A:  $t_{1/2}$  60 – 100 h (literature)
  - BA study
    - 10 mg drug A hydrochloride p.o. vs. i.v.
    - 12 subjects
    - 2×2×2 crossover, washout 35 days
    - Sampling until 312 hours post dose
    - LC/MS-MS, LLOQ 1 ng/mL (drug A base / plasma)
    - Results considered important for designing other studies
      - $t_{1/2}$  49.9 ± 13.0 h (harmonic mean ± jackknife standard deviation)
      - In none of the samples drawn at 312 h a concentration ≥LLOQ was measured
      - Extrapolated *AUC* 10.0% (median)  
3.8% – 13.9% (minimum – maximum)

# Case 1 | Study design



- Drug A:  $t_{1/2}$  60 – 100 h (literature)
  - Comparative BA study aiming to demonstrate BE
    - 10 mg drug A hydrochloride (primary target  $T_2$  vs. R, descriptive  $T_2$  vs.  $T_1$ )
    - 36 subjects
    - 3×6×3 crossover (Williams' design), washout 14 days
      - Washout planned for a worst case  $t_{1/2}$  of 66 h (covering >5 half lives)
    - Sampling until 216 hours post dose
      - No problems with extrapolated *AUC* expected (simulations)
    - GC/MS, LLOQ 0.117 ng/mL (drug A base / plasma)
  - Given that, can you imagine what happened – and why?

# Case 1 | Study design

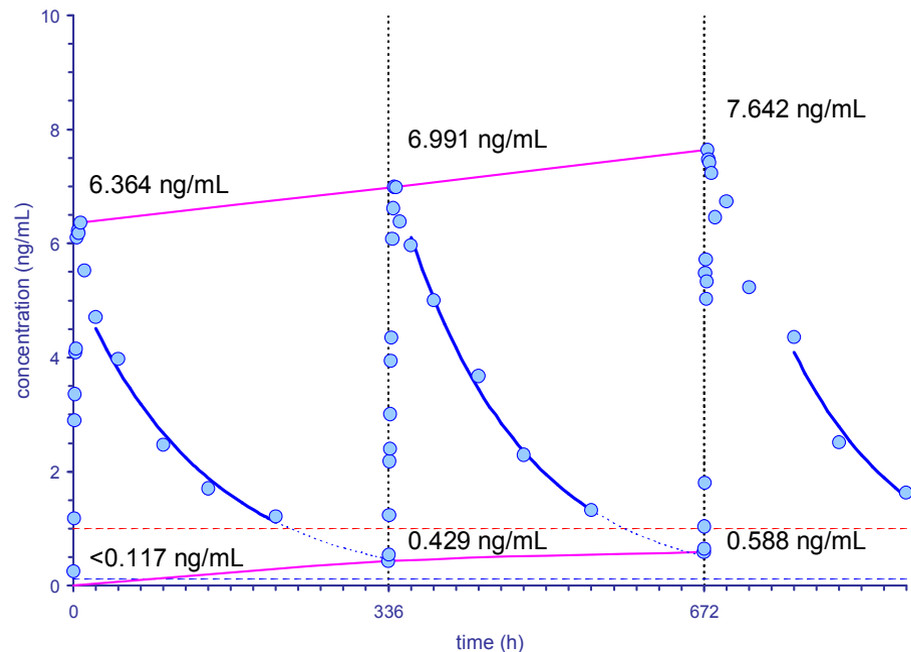


- Pre-dose concentrations  $\geq$ LLOQ: n (% of subjects, geom. means)
  - Period 1: all <LLOQ
  - Period 2: 21 (58%, 0.226 ng/mL)
  - Period 3: 18 (50%, 0.222 ng/mL) } carry-over
- Half lives (harmonic means)
  - Period 1: 51.68 h
  - Period 2: 54.20 h
  - Period 3: 63.03 h } increasing with time
- Issues
  - Improving the bioanalytical method (~9times lower LLOQ) was not a good idea
    - If we would have used the old method we would have seen not a single (!) pre-dose concentration >LLOQ
  - The shorter washout (35 days  $\rightarrow$  14) was not as well
    - Only if the estimation of  $\lambda_z$  is performed *blinded* for treatment different half lives in the periods (due to accumulation) become evident – even with the less sensitive method

# Case 1 | Study design



- Most statisticians unblind studies *before* performing NCA, which will cover potential problems
  - Half lives (harmonic means)
    - »  $T_1$ : 54.51 h
    - »  $T_2$ : 55.99 h
    - » R: 56.73 h
- Worst case  
Subject 23



- Requirements for BA/BE studies
  - Bioanalytical method developed and validated *to serve the study's purpose*
    - Calibration range
      - LLOQ  $\leq 5\% C_{max}$  in any of the subjects
      - ULOQ ideally  $\geq C_{max}$  in any of the subjects
    - (In)accuracy and (im)precision
      - 15% throughout the range (20% for ligand-binding assays)
      - 20% at LLOQ (30% for ligand-binding assays)
  - Sampling long enough to obtain reliable estimates of
    - $\lambda_z$  : at least three samples in the log/linear part
    - $AUC_{0-t}$  : covering  $\geq 80\%$  of  $AUC_{0-\infty}$  in  $\geq 80\%$  of observations
    - Note that both are *not required* if target metric is  $AUC_{0-72h}$  (IR single dose) or  $AUC_{0-T}$  (steady state)

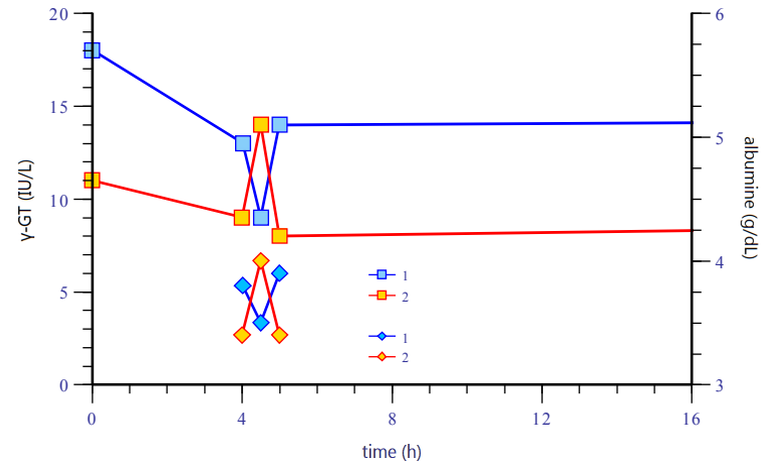
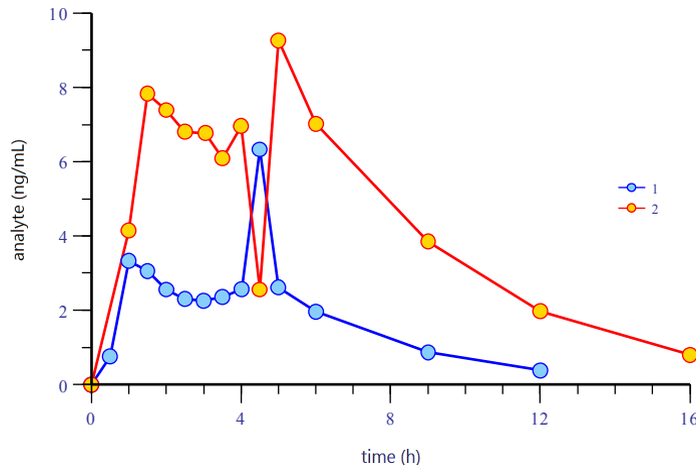
# Case 2 | Sample handling



- Clinical phase

- Biphasic modified release product of drug B
- Suspected mix-up in the transfer from sample vials after centrifugation to (plasma) sample vials

Measurable values in clin. chemistry (limited, since anticoagulant citrate)



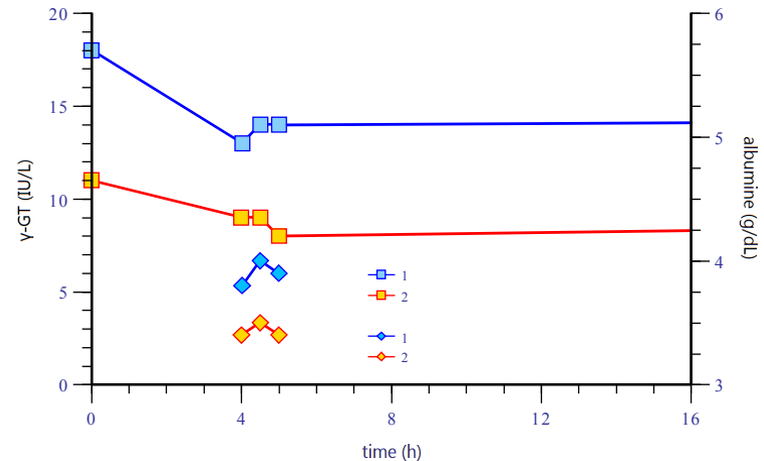
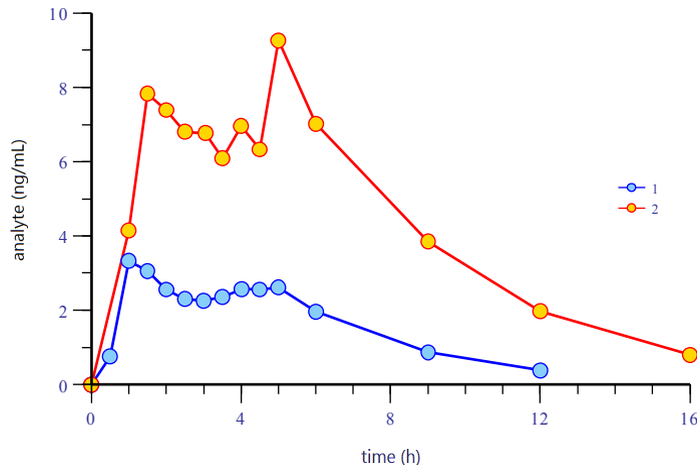
# Case 2 | Sample handling



- Clinical phase

- Biphasic modified release product of drug B
- Exploratory: values swapped (both analyte and clin.chemistry)
- Samples of subjects 1 & 2 both taken in the first period

Suspected mix-up confirmed by clin. chemistry values



# Case 2 | Sample handling



- Clinical phase
  - Barcode system failed in the first period
  - No bail-out procedure (e.g., four-eye principle)
  - Sponsor monitored plasma separation only up to two hours (when the barcode system was still working)
  - Blinded review of data for irregular profiles
    - EMA
      - According the Bioanalytical Method Validation Guideline measured results are ‘carved from stone’
        - » Exclusion of data only possible if documented error
        - » Not even repeated analysis acceptable
    - FDA
      - Acceptable
        - » Exclusion after repeated analysis possible if defined by SOP

# Case 3 | Sample handling



- Clinical phase
  - Liposome encapsulated drug C for infusion
  - Analytes
    - encapsulated drug C
    - unencapsulated drug C (*i.e.*, released from the liposomes)
    - total drug C (encapsulated + unencapsulated)
    - Metabolite (formed from unencapsulated drug C only)
  - Drug may be released from liposomes by
    - Shear forces (infusion pump, infusion needle with narrow diameter)
    - High temperature and long interval until centrifugation
    - High *g* force in centrifugation
    - Only the latter two can be avoided by proper stabilization
      - blood samples on ice
      - addition of DMSO

# Case 3 | Sample handling

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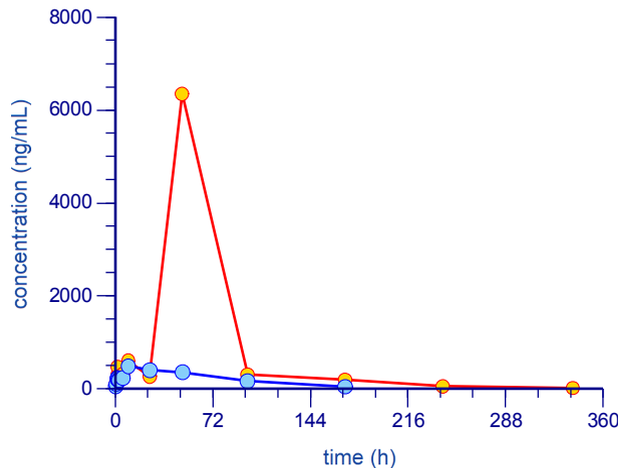


- Clinical phase
  - Multi-site study in cancer patients
  - Clinical staff educated about critical sample handling, but
    - unfamiliar procedure esp. in small clinical sites
    - necessity of following SOPs and documentation of deviations in conformity with GCP not well understood

# Case 3 | Sample handling



- Clinical phase
  - Surprises in bioanalytics
  - Extremely high concentrations of unencapsulated drug C observed in about 2% of samples
    - All values confirmed in repeated analyses

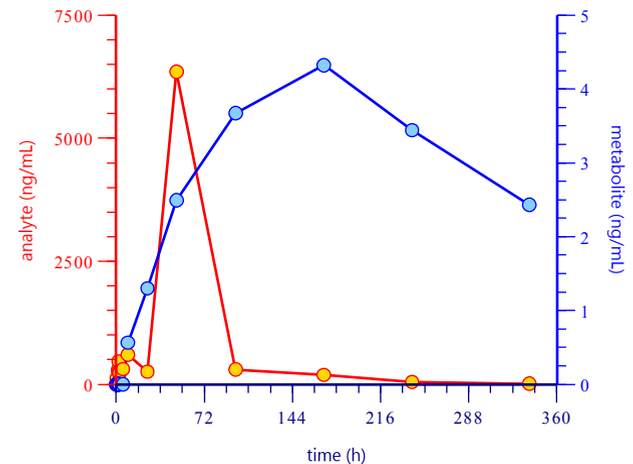
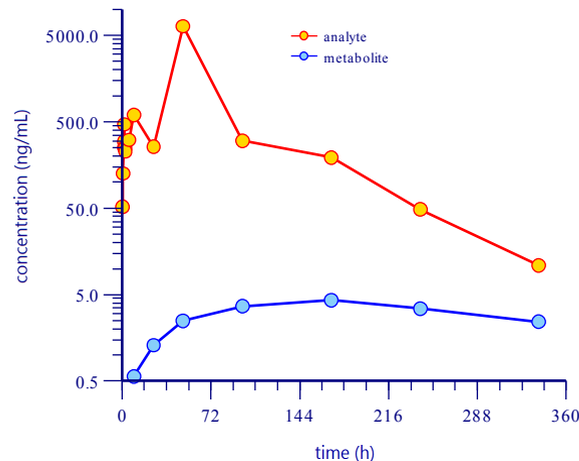


# Case 3 | Sample handling



- Clinical phase

- Extremely high concentrations of unencapsulated drug C observed in about 2% of samples
  - However, ‘normal’ concentrations of the metabolite
    - Since the metabolite can only be formed from the unencapsulated drug C, the analyte’s high concentrations were considered an artifact
    - No documentation about improper sample handling (temperature, time, stabilization)



# Case 4 | NCA

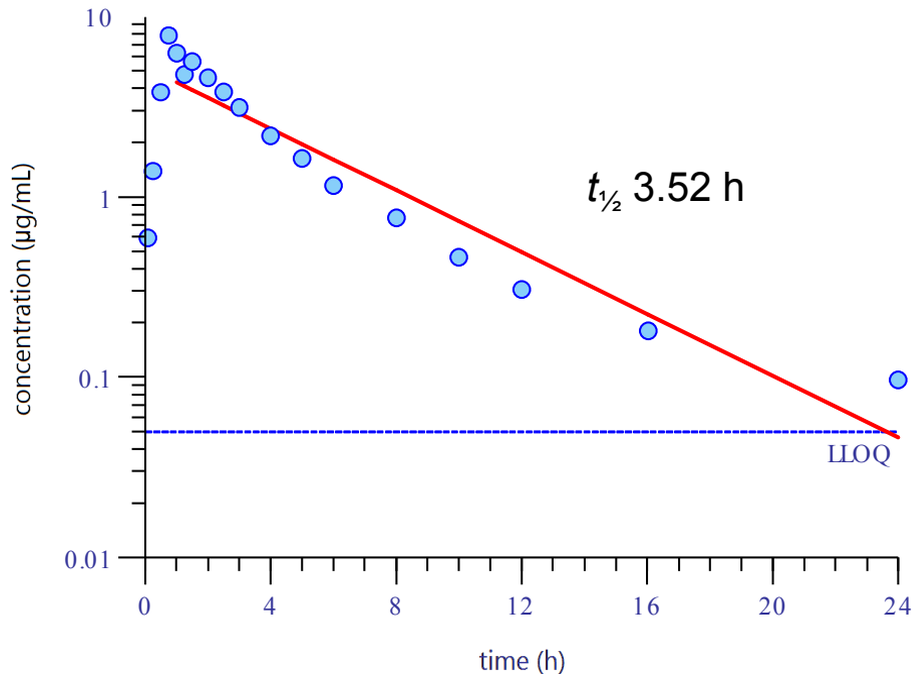


- Drug D:  $t_{1/2}$  2 – 3 h (literature)
  - BE study (500 mg D component of a 3 drug fixed dose combo)
    - liquid formulations, T vs. R
    - 27 subjects
    - TRR | RTR | RRT replicate design, washout seven days
    - Sampling until 24 hours post dose
    - LC/MS-MS, LLOQ 50 ng/mL
  - Drug D passed ABE with ease
    - $t_{1/2}$  3.92 ± 0.88 h (T), 4.98 ± 1.24 h (R)
    - Extrapolated *AUC* (median, minimum – maximum)  
T: 1.76 (0.87% – 3.61%), R: 2.42% (1.14% – 6.19%)
  - Sponsor developed a 4 drug FDC
    - Data of the BE study should be used in a PopPK model to optimize the sampling schedule for a new study

# Case 4 | NCA



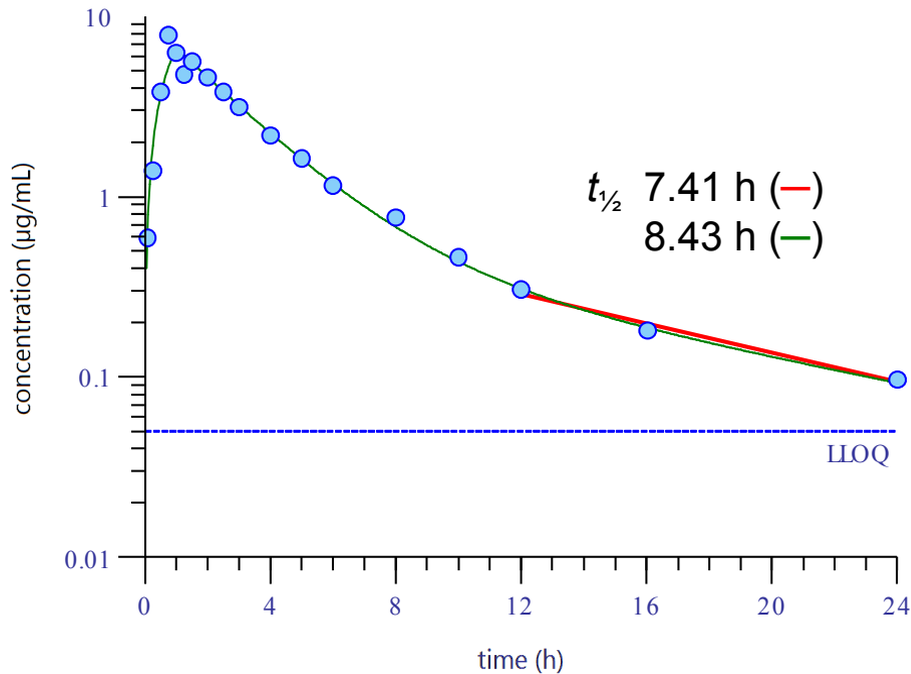
- Drug D:  $t_{1/2}$  2 – 3 h (literature)
  - No individual  $\lambda_z$  or  $t_{1/2}$  (as well as the time range used in estimation) given in the report, only  $AUC_{0-t}$  and  $AUC_{0-\infty}$
  - Reproduced the CRO's results by trial and error. Example:



# Case 4 | NCA



- Drug D:  $t_{1/2}$  2 – 3 h (literature)
  - Obviously the time range for the estimation of  $\lambda_z$  was wrong
    - Two-compartment model!
  - What I obtained in NCA (—) and by the PK model (—)



# Case 4 | NCA

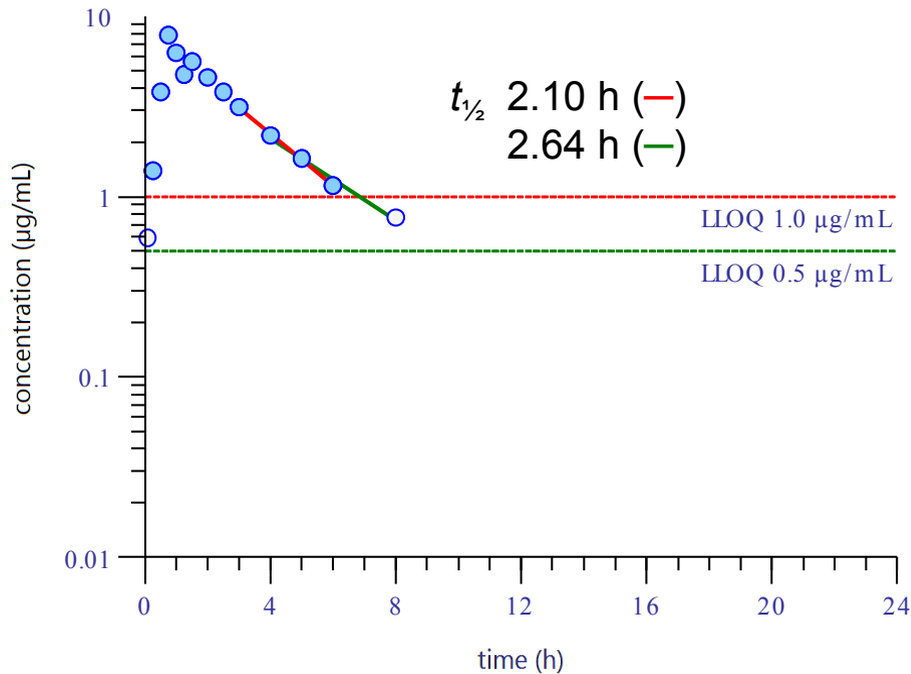


- Drug D:  $t_{1/2}$  2 – 3 h (literature)
  - Why? No problems with correct estimation of  $\lambda_z$ 
    - $t_{1/2}$  4.63 ± 1.07 h (T), 5.59 ± 1.19 h (R)
    - Extrapolated AUC (median, minimum – maximum)  
T: 2.08% (1.06% – 4.32%), R: 2.84% (1.47% – 6.19%)
  - Possible explanations
    - ‘Push-the-button-pharmacokineticist’ at work
      - Relied on an automatic algorithm?
      - No visual inspection of fits?
    - Anticipatory obedience (‘vorausseilender Gehorsam’)?
      - The bioanalytical method was at least 10times more sensitive than ones used in the past (drug D approved in 1955)
      - Maybe the CRO wanted to avoid a single sentence in the discussion section explaining why a second phase is apparent – explaining a half live longer than the one known from literature

# Case 4 | NCA



- Drug D:  $t_{1/2}$  2 – 3 h (literature)
  - Estimation of  $\lambda_z$  by bioanalytical methods with a LLOQ of 1.0 or 0.5  $\mu\text{g}/\text{mL}$  explains short half lives given in 'old' literature



# Case 4 | NCA



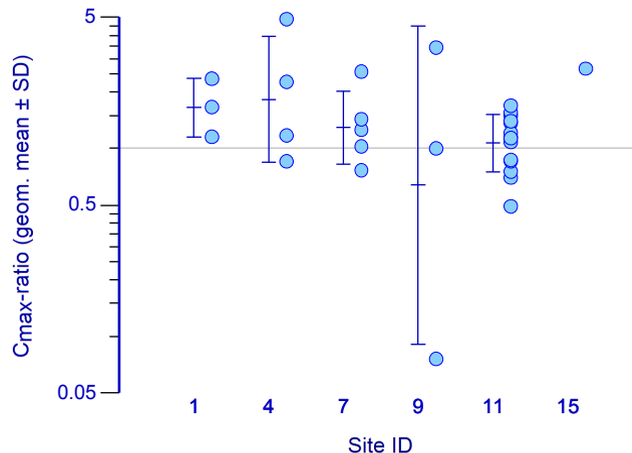
- Drug D:  $t_{1/2}$  2 – 3 h (literature)
  - Lessons learned
    - Insist that the (draft!) PK report allows independent assessment
    - Mandatory<sup>1,2</sup>
      - All raw data
      - $\lambda_z$  and/or  $t_{1/2}$  as well as time ranges used in estimation
      - All derived PK metrics
    - Desirable
      - Data in a machine-readable format (CSV, SAS transport, CDISC)
    - Unacceptable
      - A 500+ page PDF generated by SAS
      - As above but a scanned print

1 Schulz H-U, Steinijans, VW. *Striving for standards in bioequivalence assessment: a review.* Int J Clin Pharm Ther Toxicol. 1991;29(8):293–8. [PMID 1743802](#).

2 Sauter R, Steinijans VW, Diletti E, Böhm E, Schulz H-U. *Presentation of results from bioequivalence studies.* Int J Clin Pharm Ther Toxicol. 1992;30(Suppl.1):S7–30. [PMID 1601535](#).

- Requirements for BA/BE studies
  - Design should allow accurate (unbiased) assessment of the treatment effect
  - EMA (2010)
    - The study should be designed in such a way that the formulation effect can be distinguished from other effects.
    - The precise model to be used for the analysis should be pre-specified in the protocol. The statistical analysis should take into account sources of variation that can be reasonably assumed to have an effect on the response variable.

- Continuing Case 3
  - Extreme  $C_{max}$ -ratios of unencapsulated drug C observed only in small clinical sites



- Pooling data of sites *only* if
  - similar variances
  - no treatment-by-site interaction

# Case 5 | Statistics



- Planned evaluation of unencapsulated drug C
  - Statistical model *did not* take the multi-site nature of the study into account (*i.e.*, data of all sites were naïvely pooled);
    - Failed: PE 121.01% (90% CI: 96.13 – 152.33%),  $CV_w$  55.6%
- Sensitivity analysis
  - Statistical model suggested by the FDA including the site-by-treatment interaction
    - Highly significant ( $p$  0.00063)
    - Hence, pooling of sites is *not* justified
  - Therefore, analysis of largest site #11 only
    - Passed: PE 103.80% (90% CI: 89.87% – 119.90%),  $CV_w$  21.4%

- Adaptive Two-Stage Sequential Design in BE

- EMA (2010)

It is acceptable to use a two-stage approach [...]. If this approach is adopted appropriate steps must be taken to preserve the overall type I error of the experiment [...]. For example, using 94.12% confidence intervals for both the analysis of stage 1 and the combined data from stage 1 and stage 2 would be acceptable, but there are many acceptable alternatives and the choice of how much alpha to spend at the interim analysis is at the company's discretion.

- The 94.12% CI ( $\alpha$  0.0294) preserves the patient's risk in simulation-based methods *only* if

- GMR 0.95 and
    - Target power 80%

# Case 6 | Statistics



- Adaptive Two-Stage Sequential Design in BE
  - GMR 0.90, target power 85%,  $\alpha$  0.0294
  - Stage 1:  $n_1$  24
    - Failed: PE 89.00% (94.12% CI: 77.24 – 102.54%)
    - Since interim power 37.7%, stage 2 with 54 subjects initiated
  - Analysis of pooled data:  $n_1+n_2$  78
    - Passed: PE 91.00% (94.12% CI: 82.16 – 100.79%)
  - Inflated patient's risk (5.23%)
  - The study's conditions would require *more* adjustment (at least an  $\alpha$  of 0.0279 or a 94.42% CI)
    - *Post hoc* assessment
      - Passed: PE 91.00% (94.42% CI: 82.05 – 100.92%)
      - Type I Error 4.99%
      - Wider CI but conclusion agrees with the original analysis

- Adaptive Two-Stage Sequential Design in BE
  - However, correct would have been to find a suitable  $\alpha$  (0.0278) already *before* and implement it in the study
  - Stage 1:  $n_1$  24
    - Failed: PE 89.00% (94.44% CI: 77.09 – 102.75%)
    - Since interim power 36.6%, stage 2 with 54 subjects initiated
  - Analysis of pooled data:  $n_1+n_2$  78
    - Passed: PE 91.00% (94.44% CI: 82.05 – 100.93%)
  - Controlled patient's risk (4.99%)