Background and Objective

Characterisation of exposure-response for therapies where the clinical endpoint is a bivariate outcome generally involves two assumptions: (i) the random effect determining the dose-exposure relation is not related to any parameter of the exposure-response relation, and (ii) the drug effect is linear on the logit scale. This study aims at assessing the possibility for randomised dose controlled trials (DCT) and concentration controlled trials (CCT) to characterise such phenomena, with a particular focus on drugs with narrow therapeutic index.

Methods

PKPD model: A simulation-based study was performed using NONMEM VI considering a hypothetical immunosuppressant agent with rejection as main efficacy endpoint. The PK-model was described by equations:

\[ C_t = \frac{C_{L_t} - C_{L_e}}{C_{L_t}} \]

and the PD-relationship with a regression logistic model:

\[ \text{Logit}_j = \theta_{d_j} + \theta_{s_j} \cdot C_i \]

\[ P_x = \frac{e^{\text{Logit}_i}}{1 + e^{\text{Logit}_i}} \]

Two alternative PD-models were considered:

a) PKPD covariance between CL and baseline or slope parameter:

\[ \text{Logit}_i = \theta_{d_{ij}} + \theta_{s_{ij}} \cdot C_j \]

b) Nonlinear exposure-response relationship:

\[ \text{Logit}_i = \theta_{d_{ij}} + \theta_{s_{ij}} \cdot C_i \]

Simulation setup: the default considered as typical value for clearance (\( C_{L_t} \)), 20 L/h and 45% of IV. As baselines and slopes: \( \text{BINF}_i = -3.5 \), \( \text{BINF}_i = 15 \), \( \text{BREJ}_i = -1 \) and \( \text{BREJ}_i = -12 \).

As \( \delta \) and \( \gamma \) the values shown in the equations. When \( \gamma \) took the values 1, 2, 3, 4, the \( \text{BREJ}_i \) took values of -12, -120, -1200 and -12000, respectively. So that, in all cases the covariance/non-linearity was inside the range of study and could be detected a priori.

Study design: 3 randomized, cross-over designs with two dose/exposure levels were considered:

(i) DCT with two dose level targets;

(ii) TCCT with exposures that reflect the expected typical exposure in the corresponding DCT;

(iii) VCCT targeting the two exposure levels that result in equal exposure variability as the corresponding DCT.

A study size of 500 subjects was considered and four different ranges of target levels were explored.

When assessing the non-linearity, the CCTs designs targeted three levels of exposures inside the same range as the corresponding DCT, so that the VCCT still was equip-variable to the DCT.

Results and Discussion

With regards to precision and bias in parameter estimates: DCT and VCCT were superior for a (+) and (-) PKPD covariance between CL and baseline, respectively (Figure 1 & 2).

When the PKPD covariance existed between CL and slope, the VCCT design was the more precise regardless of the sign of the correlation (Figure 3). The VCCT and TCCT showed highest power to detect the correlation in all cases (results not shown).

Acknowledgments

R.L.G. was supported by a grant from Pfizer, Sandwich, U.K.

References


Conclusion

For drugs with narrow therapeutic index either a VCCT or DCT have shown to be more informative designs to describe the exposure-response relationship when there is a PKPD covariance in the parameters, whereas a DCT seems to be more informative when describing non-linear relationships between exposure and response. However, typical studies of this type would not have enough power to reliably detect such relationships regardless of design.