Normal fat mass cannot be reliably estimated in typical pharmacokinetic studies

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Abstract
Purpose An influential covariate for pharmacokinetics is (body) size. Recently, the method of estimation of normal fat mass (NFM) has been advocated. Here, the relative contribution of fat mass, estimated as a fraction fat (F_{\text{fat}}), is used to explain differences in pharmacokinetic parameters. This concept is more and more applied. However, it remains unclear whether NFM can be reliably estimated in these typical studies.
Methods We performed an evaluation of the reliability of NFM estimation in a typical study size (n = 30), otherwise best-case scenario, by means of a pharmacokinetic simulation study. Several values of F_{\text{fat}} were investigated.
Results In a typical pharmacokinetic study, high imprecision was observed for NFM parameter estimates over a range of scenarios. For example, in a scenario where the true value of F_{\text{fat}} on clearance was 0.5, we found a 95% confidence interval of −0.1 to 2.1, demonstrating a low precision. The implications for practice are that one could conclude that fat-free mass best describes the relationship of the pharmacokinetics with body size, while the true relationship was between fat-free mass and total body weight. Consequently, this could lead to incorrect extrapolation of pharmacokinetics to extreme body sizes.
Conclusion In typical pharmacokinetic studies, NFM should be used with caution because the F_{\text{fat}} estimates have low precision. The estimation of F_{\text{fat}} should always be preceded by careful study design evaluation before planning a study, to ensure that the design and sample size is sufficient to apply this potentially useful methodology.

Keywords Normal fat mass  ·  F_{\text{fat}}  ·  Fat-free mass  ·  Population pharmacokinetics  ·  Pharmacokinetic modeling  ·  Non-linear mixed-effects modeling

Introduction

Like humans, body size descriptors come in many shapes, with total body weight (TBW) and fat-free mass (FFM) currently being most accepted in the pharmacometric community [1]. Choosing correct size descriptors, which account for the presence of peripheral fat tissue, may be especially important when dosing a drug with a narrow therapeutic index (i.e., aminoglycosides), especially in populations with extreme body size. It has been argued that “traditional” descriptors are not adequate to account for the (relative) importance of peripheral fat tissue to describe pharmacokinetic changes with increasing body size. It assumes that the pharmacokinetic parameter of interest is related to either TBW or FFM.

Recently, the method of estimation of normal fat mass (NFM) has been advocated [2]. NFM is the sum of the calculated FFM (based on height, weight, and sex) and the estimated relative contribution of fat mass. This method is increasingly employed in relatively small studies [3–14]. NFM is defined by Eq. 1, where F_{\text{fat}} is the estimate that reflects the relative contribution of fat mass. It can be used together with a standard NFM (NFM_{\text{STD}}) and an allometric coefficient (b) to scale the population estimate (\(\theta_{\text{pop}}\)) to the individual parameter (\(\theta\)) (Eq. 2) [15].

\[
\text{NFM} = \text{FFM} + F_{\text{fat}} \times (\text{TBW} - \text{FFM})
\]

(1)

\[
\theta = \theta_{\text{pop}} \times \left(\frac{\text{NFM}}{\text{NFM}_{\text{STD}}}\right)^b
\]

(2)

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It follows that when $F_{fat}$ is 1, NFM equals TBW. When $F_{fat}$ is 0, NFM equals FFM and when $F_{fat}$ is (much) larger than 1, the value of NFM is determined by the amount of fat tissue relative to FFM. In fact, an $F_{fat}$ value of $+\infty$ would represent a situation where fat tissue alone is driving changes in the parameter of interest. Finally, negative values of $F_{fat}$ are possible in a situation where, for example, clearance is impaired due to obesity. The use of the NFM method is currently reported in different ways: first, as an aid to investigate which size metric (i.e., TBW, FFM, or fat mass) is most suitable to describe the data after which one is chosen [3–10]; second, as an estimate where the value of $F_{fat}$ is reported [11–13]. And third, using a mixture of both where if an estimate is 0 or 1 then they are fixed to the respective value but when the estimates is between 0 and 1 then the actual value of the estimated is used and reported [14].

$F_{fat}$ estimates are considered drug-specific parameters that are also specific for the pharmacokinetic variable (i.e., clearance or volume of distribution) and can be used for pharmacokinetic extrapolation into populations with different body sizes or composition [2]. For this purpose, reliable estimates of $F_{fat}$ (i.e., reproducible point estimates with high precision) are needed. However, it remains unclear whether NFM, by estimation of $F_{fat}$, can be reliably determined with a representative study size seen in the clinical setting. Therefore, we investigated the reliability of the estimates of several $F_{fat}$ values in typical pharmacokinetic studies, assuming an otherwise best-case scenario.

Methods

General approach

We performed two simulation studies. The purpose of the first study was to confirm that $F_{fat}$ can be reliably estimated when your study is large enough ($n = 10,000$). This study size should not be interpreted as a realistic size but as an approximation of an infinite number of subjects. The purpose of the second study was to investigate how well $F_{fat}$ can be estimated in a typically sized, but otherwise best-case scenario, pharmacokinetic study ($n = 30$). In this report, we will refer to these studies as “large” and “typical.”

Virtual drugs

$F_{fat}$ can be defined as the contribution of fat mass to a drug clearance (CL) or volume of distribution (V) (Eq. 1). Here, we investigated 16 virtual drugs, consisting of all possible combinations of $F_{fat}$: 0, 0.5, 1, and 5, for CL and V, respectively.

Study populations

The best-case scenario for a reliable estimate of $F_{fat}$ requires a wide distribution in body size and specifically a wide distribution of the FFM and TBW difference (i.e., fat mass) [15]. We obtained realistic demographic data (weight, height, and sex) from the National Health and Nutrition Examination Survey (NHANES) database from data collected between 1999 and 2014 [16]. The FFM was predicted using weight, height, and sex according to the formula reported by Janmahasatian et al. [17]. For each virtual study, we randomly sampled per BMI group from the NHANES database to create three equally sized groups of the following: (1) normal weight (body mass index (BMI) 18.5–30 kg/m$^2$); (2) obese (BMI 30–40 kg/m$^2$); and (3) morbidly obese (BMI > 40 kg/m$^2$), with 50% of the subjects being female.

Pharmacokinetic model

All pharmacokinetic simulations and estimations were performed with NONMEM (v7.3) and Perl-SpeaksNONMEM (v4.7) [18, 19]. The virtual drug was administered as 1 mg intravenous bolus. Pharmacokinetic data were simulated using a one-compartment model with first-order elimination, CL of 0.693 L/h, and V of 1 L. Inter-individual variability (IIV) was 30% on both CL and V, and we added a proportional residual error of 15%. A 15% residual error is chosen to be in line with the most recent EMA and FDA guidelines for bioanalytical method validation, where a %CV in the precision of <15% is defined as precise [20, 21]. The sampling schedule covered three half-lives: 0.25, 0.5, 0.75, 1, 1.25, 1.5, 2, and 3 h after dosing. None of the simulated pharmacokinetic data was below the limit of quantitation. NFM was standardized around a male with a TBW of 100 kg and an FFM of 60 kg according to Eq. 2. Fixed allometric coefficients of 0.75 and 1 for CL and V, respectively, were applied. The NONMEM model code is provided in the supplemental material.

Stochastic simulation and estimation

After simulation, we re-estimated all parameters in each simulated dataset. The re-estimation was performed using the first-order conditional estimation method with interaction. For each study size, we simulated and re-estimated 1000 studies. For the typical study, we sampled new individuals from NHANES for each simulation, thereby making 1000 unique studies, to simulate...
differences in covariate distributions between studies. Figure 1 shows a flow chart of the analytical approach.

**Outcome**

The estimates from each of the 1000 studies, for each virtual drug and study size were summarized using the median and the 2.5th and 97.5th percentile. In this work, we will refer to this estimation interval as the 95% confidence interval (95% CI). Summary statistics were calculated for the pharmacokinetic parameters (CL, V, II CL, II V, and the residual error) and for the estimates of Ffat. The 95% CIs of the pharmacokinetic parameters were used to assess the reliability of the parameter estimates given the study design. To illustrate the change in precision with increasing sample size, we repeated the above method for study sizes of 60, 100, 150, 200, 250, 500, 1000, and 5000 subjects with an Ffat of 0.5 in NFM on both CL and V.

**Results**

**Reliability of pharmacokinetic parameter estimates**

Figure 2 shows a histogram of TBW and FFM distribution of a randomly sampled study (out of 1000 studies), indicating a wide range of body size. Figure S1 in the supplemental materials shows the distribution of TBW and FFM in the large (n = 10,000) study. Table 1 shows the median and 95% CI for the typical study size of a virtual drug with Ffat of 0.5 in NFM on both CL and V. All parameters were estimated with low bias (<10%) and narrow 95% CI indicating that with our typical study size it is possible to accurately estimate the primary pharmacokinetic parameters. The precision and bias of the estimates of the primary pharmacokinetic parameters in the other 15 virtual drugs were similar (data not shown).

**Reliability of Ffat estimates in large studies**

We found a narrow 95% CI for Ffat in all the 16 virtual drugs (Figs. 3 and 4 in cyan). This indicates that Ffat can be estimated accurately and precisely when a large data set is available. The 95% CI of Ffat in NFM on CL was independent of the Ffat value in NFM on V, meaning that these could be estimated independently from each other regardless the value of the other estimate of Ffat (data not shown). For example, the 95% CI the Ffat on CL with a true value of 1 was 0.11–4.4 regardless of the (true) value of the Ffat on V (i.e., whether Ffat on V was 0, 0.5, 1, or 5). Therefore, we only show combinations for the large and the typical study size where Ffat values were equal for CL and V (i.e. Ffat of 1 in NFM on both CL and V).
Reliability of $F_{fat}$ estimates in typical studies

We found a large 95% CI (Figs. 3 and 4 in red). The intervals of the different values of $F_{fat}$ were overlapping to a large extent which makes it challenging to interpret an estimated $F_{fat}$. Interestingly, the lower range of the 95% CIs of the $F_{fat}$ values of 0 and 0.5 was below zero, which indicates for CL that an increasing amount of fat mass results in a lower CL which is opposite to the relation used to simulate the data. For the $F_{fat}$ value of 5 (representing the situation where fat mass is the most important metric driving individual variability in CL or V), the 95% CIs were very wide indicating that the fact that the estimate is a high value is more informative than the exact value itself. Finally, the 95% CIs of $F_{fat}$ 0 did not contain 1 and the 95% CI of 1 did not contain 0, indicating that for our study design, size and drug characteristics using NFM as an exploratory tool should result in a model closer to the true model compared to a priori choosing a fixed value of 0 (FFM) or 1 (TBW).

Reliability of $F_{fat}$ estimates in other study sizes

The precision of the estimates of $F_{fat}$ on CL and V of other study sizes is shown in Fig. 5. As can be expected, the more subjects that are included, the narrower the 95% CI. However, even with 100 subjects, the interval for the $F_{fat}$ estimate on CL and V is relatively large including both 0.5 and 2 in its interval. Increasing the study to 250 normal weight, obese and morbidly obese subjects only slightly improves the precision.

Table 1 Pharmacokinetic parameters of the one-compartment model and the estimates in the typical study size including a virtual drug with $F_{fat}$ 0.5 in NFM on both CL and V

| Parameter                              | True value | Median estimate | 95% CI       |
|----------------------------------------|------------|-----------------|--------------|
| Clearance (L/h)                        | 0.69       | 0.71            | 0.63–0.78    |
| Volume of distribution (L)             | 1.0        | 1.0             | 0.99–1.1     |
| $F_{fat}$ on clearance                 | 0.5        | 0.51            | −0.11–2.1    |
| $F_{fat}$ on volume of distribution    | 0.5        | 0.48            | −0.06–1.6    |
| Inter-individual variability           |            |                 |              |
| Clearance (%)                          | 30         | 28              | 20–36        |
| Volume of distribution (%)             | 30         | 28              | 20–36        |
| Residual error (%)                     | 15         | 15              | 14–16        |

Median [95% CI]

- 0.0 [−0.03 – 0.03]
- 0.0 [−0.37 – 0.85]
- 0.5 [0.45 – 0.55]
- 0.51 [−0.11 – 2.1]
- 1.0 [0.93 – 1.1]
- 0.97 [0.11 – 4.4]
- 5.1 [4.7 – 5.8]
- 5.4 [1.3 – 3.3e+8]

95% CI: 95% confidence interval. Clearance and volume of distribution were standardized on a male with a total body weight of 100 kg and a fat-free mass of 60 kg.
Discussion

Although we showed that F_{fat} is identifiable in (large) pharmacokinetic studies, we found that it cannot be reliably estimated in pharmacokinetic studies with a typical study size in an otherwise best-case scenario. In our simulation study, we found values of F_{fat} in NFM on CL between −0.1 and 2.1, while the true value was 0.5.

The median value of the estimates of 1000 replicated studies of any value of F_{fat} is indeed accurate but for a single study, one never knows the true value. We show here that an incorrect value of F_{fat} will be found regularly meaning that
the estimate of \( F_{\text{fat}} \) is study-specific if the power of the estimate is not evaluated. Using an inaccurate value of \( F_{\text{fat}} \) could result in an erroneous prediction of the pharmacokinetics of a drug when extrapolated to another population.

In this simulation study, we mainly focused on two study sizes to illustrate the potential pitfall of \( F_{\text{fat}} \); hence, we did not study other study sizes or scenarios in detail. In Fig. 5, we show that increasing the sample size per study will decrease the confidence interval of \( F_{\text{fat}} \) estimates. However, a general statement on the minimal study size to estimate \( F_{\text{fat}} \) is not useful since study size is just one of the factors attributing to parameter identifiability and reliability. Other factors include the study design, number of observations, size of the interindividual variability, size of the residual variability, the distribution of the covariates (e.g., uniform, normal or log-normal), and covariate effect size [22]. In our study, we chose a best-case scenario situation, of an intravenously administered drug, with rich sampling and low residual error. Therefore, in real-life situations, we expect that NFM cannot be reliably estimated. Furthermore, Fig. 5 illustrates that even in a best-case scenario setting and a very large sample size of 100 richly sampled individuals, there would only be a marginal improvement of the precision of the \( F_{\text{fat}} \) estimates. Although our study was not designed to find the optimal design and sample size for estimation of \( F_{\text{fat}} \), considering the uniqueness of each drug and population, we nonetheless show that it is unlikely that \( F_{\text{fat}} \) can be reliably estimated in a typical study. One may argue that a poorly designed study will always result in poor estimates. From our results, it can be argued that testing fixed estimates of \( F_{\text{fat}} \) (0 for fat-free mass or 1 for total body weight) as size descriptors in a pharmacokinetic study may also result in selection of incorrect size predictors. Pharmacometricians should be aware of this phenomenon when performing pharmacokinetic studies. Future studies should investigate whether, for example, physicochemical properties of drugs may be used to predict the best size descriptor for allometric scaling in pharmacokinetic studies.

The NFM method is an elegant solution to investigate the contribution of fat mass to variability in pharmacokinetics. The NFM method can be very useful in a covariate analysis to investigate several size metrics in a single run after which the best one can be selected. The use of NFM has an advantage compared to traditional body size metrics, at the cost of only one parameter. However, its use is usually not preceded by careful study design and sample design evaluation, potentially leading to imprecise estimates. We have shown here that the point estimates of \( F_{\text{fat}} \) have low precision in realistically sized pharmacokinetic studies, even in a best-case scenario. Consequently, the estimation and reporting of NFM in these studies should be performed with caution and, if used, should always be preceded by rigorous study design evaluation.

**Conclusion**

The novel concept of NFM can be advantageous in sufficiently large studies but may have limitations in realistically sized pharmacokinetic studies, predominantly in reliability of the estimates \( F_{\text{fat}} \). This might have consequences for prediction of dosing of medications with a narrow therapeutic window in patients with extreme body sizes. We strongly advocate rigorous study design and sample size evaluation before applying this potentially very useful methodology in a pharmacokinetic study.

**Supplementary Information** The online version contains supplementary material available at https://doi.org/10.1007/s00228-020-03042-4.

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**Author contributions** All authors contributed to the study conception and design. Simulations and analysis were performed by Roeland Wasmann. The first draft of the manuscript was written by Roeland Wasmann and all authors commented on previous versions of the manuscript. All authors read and approved the final manuscript.

**Compliance with ethical standards**

**Conflict of interest** SS is an employee of AstraZeneca and may own stock or stock options. None of the other authors has a conflict to declare.

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