Population Pharmacokinetics of Imipenem in Critically Ill Patients: A Parametric and Nonparametric Model Converge on CKD-EPI Estimated Glomerular Filtration Rate as an Impactful Covariate

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Abstract
Background Population pharmacokinetic (popPK) models for antibiotics are used to improve dosing strategies and individualize dosing by therapeutic drug monitoring. Little is known about the differences in results of parametric versus nonparametric popPK models and their potential consequences in clinical practice. We developed both parametric and nonparametric models of imipenem using data from critically ill patients and compared their results.

Methods Twenty-six critically ill patients treated with intravenous imipenem/cilastatin were included in this study. Median estimated glomerular filtration rate (eGFR) measured by the Chronic Kidney Disease Epidemiology Collaboration (CKD-EPI) equation was 116 mL/min/1.73 m² (interquartile range 104–124) at inclusion. The usual dosing regimen was 500 mg/500 mg four times daily. On average, five imipenem levels per patient (138 levels in total) were drawn as peak, intermediate, and trough levels. Imipenem concentration-time profiles were analyzed using parametric (NONMEM 7.2) and nonparametric (Pmetrics 1.5.2) popPK software.

Results For both methods, data were best described by a model with two distribution compartments and the CKD-EPI eGFR equation unadjusted for body surface area as a covariate on the elimination rate constant ($K_e$). The parametric population parameter estimates were $K_e$ 0.637 h⁻¹ (between-subject variability [BSV]: 19.0% coefficient of variation [CV]) and central distribution volume ($V_c$) 29.6 L (without BSV). The nonparametric values were $K_e$ 0.681 h⁻¹ (34.0% CV) and $V_c$ 31.1 L (42.6% CV).

Conclusions Both models described imipenem popPK well; the parameter estimates were comparable and the included covariate was identical. However, estimated BSV was higher in the nonparametric model. This may have consequences for estimated exposure during dosing simulations and should be further investigated in simulation studies.

1 Introduction

Because of increased antimicrobial resistance and few new antibiotics making it to market, optimization of antibiotic dosing regimens remains an important challenge to improve clinical outcomes of infections. Antimicrobial efficacy is determined by the susceptibility of a drug in vitro (usually expressed as the minimal inhibitory concentration [MIC]) and exposure to the drug in vivo, which relies on the pharmacokinetics (PK) and the dose [1]. Population pharmacokinetic (popPK) models describe the variability of exposure to a drug and are therefore used to support the optimization of dosing regimens with the objective to improve antimicrobial efficacy. During the development of new antibiotics, popPK models are recommended to support dose regimen identification and selection [2]. For already marketed antibiotics, popPK models are used in different ways to improve antibiotic dosing: individualization of dosing via therapeutic drug monitoring (TDM) software by Bayesian estimation and control; optimization of dosing regimens described in the package insert (especially for specific subpopulations); and setting clinical breakpoints [3]. Clinical breakpoints are MICs that define microorganisms as susceptible, intermediate or resistant to specific antibiotics [4].
While individual PK methods analyze the concentration-time profiles per individual subject, popPK methods analyze these profiles of a population as a whole. PopPK models describe and explain different types of PK variability, such as between-subject variability (BSV) and residual variability. PopPK modelling methods are classified as either parametric or nonparametric methods, which can each be divided in maximum likelihood or Bayesian approaches [3]. Bayesian popPK methods are used much less often than maximum likelihood popPK methods. Most published popPK models are based on parametric maximum likelihood methods (e.g. Monolix, NONMEM and Phoenix NLME), which estimate the set of PK parameters that maximize the joint likelihood of observations. Parametric methods assume that the population parameter distribution is known with population parameters to be estimated [5]. An example of nonparametric maximum likelihood software is the NPAG algorithm in the R package Pmetrics [6]. Nonparametric methods make no assumption about the shapes of the underlying parameter distributions. Another difference is that nonparametric methods use an exact likelihood function, while most parametric methods use an approximation. A disadvantage of nonparametric methods in the past was that confidence intervals about parameter estimates could not be easily determined [5, 7]. However, Pmetrics can estimate credibility intervals around median parameter estimates using a bootstrap method.

Some studies comparing parametric and nonparametric models are available in the literature. Precluding studies with currently outdated modelling software [8–11], we found eight comparison studies [12–19]. Four of these studies showed comparable parameter estimates of both models [12–15], one study showed different estimates [16], for two studies the estimates were incomparable due to a different model structure [17, 18], and in one study no parameter estimates were reported [19]. However, less similarity among methods was noticed for the BSV of parameters. The BSV is defined as the percent coefficient of variation (CV%), which is the standard deviation (SD) divided by the parameter mean. Three studies showed a higher BSV for the nonparametric model [12, 13, 16], one study showed similar BSV [14], and, for another study, the BSV of the nonparametric model was not reported [15]. Only one comparison study [18] presented goodness-of-fit (GOF) plots and visual predictive checks (VPCs) of both models. The other studies [12–17, 19] showed either GOF or VPC plots of both models, displayed the plots of only one method, or did not show any GOF or VPC plots.

Many parametric and nonparametric popPK models have been published in the literature and are often accompanied by dosing recommendations based on simulations of the model [3]. Differences in modelling results between these methods may have consequences for simulation findings and may therefore also influence dosing recommendations. Different parameter value probability distributions may influence dosing recommendations based on popPK models in TDM software.

To investigate the advantages and disadvantages of both modelling methods in practice, we developed both parametric and nonparametric popPK models using the same data, and compared the results. We used imipenem PK data from critically ill patients, given this population’s known PK variability for several antibiotics [20]. Imipenem is a carbapenem antibiotic administered in combination with cilastatin to prevent degradation by dehydropeptidase-I in the kidneys. When combined with cilastatin, approximately 70% of imipenem is recovered in the urine within 10 h and the rest is excreted as inactive metabolites via the urine [21]. The half-life is 1 h in patients with normal renal function, but is extended in patients with renal dysfunction [21]. Protein binding is reported as 10–20% [22].

Although it is known that the correlation between measured creatinine clearance (ClCr) and estimated glomerular filtration rate (eGFR) equations is weak [23], these equations are used in daily practice in many intensive care units (ICUs). In our study population, measured ClCr was unavailable, therefore we decided to test several eGFR equations during covariate model building to find the most suitable one in our population.

2 Methods

2.1 Study Population

Imipenem PK data from a previously published prospective cohort study [24] conducted between 2010 and 2013 in
the ICU of the Geneva University Hospitals (Geneva, Switzerland) were used for this popPK study. The usual dosing regimen for imipenem/cilastatin was 500 mg/500 mg four times daily, administered by intermittent intravenous infusion of 30 min. Inclusion criteria were suspected or documented severe bacterial infection and age 18–60 years, while exclusion criteria were eGFR < 60 mL/min (measured by the Cockcroft–Gault [CG] equation [25]), body mass index < 18 or > 30 kg/m², and pregnancy. The study protocol was approved by the University Hospital’s Ethics Committee (NAC 09-117). Given its observational nature, the Committee waived the requirement for informed consent from patients who were unconscious or otherwise unable to understand the study protocol.

Among the 54 critically ill patients from the Swiss study who were receiving imipenem therapy, the last 27 patients could be included because exact dosing and blood sampling times were known, in contrast to the first 27 patients, for whom levels were labelled only as trough, intermediate or peak. After excluding one subject because of missing height [26], data from the remaining 26 patients were included in the popPK study. None of these patients received probenecid, the only drug known to influence imipenem concentrations [21].

### 2.2 Study Procedures

Patients were included on their first or second day of imipenem therapy. Blood samples were planned on days 1, 2, 3, 4 and 6 of therapy, although in some patients not all planned samples were realized, e.g. due to discontinuation of therapy or problems with blood drawing. Imipenem TDM included peak (approximately 15–30 min after the end of the infusion), intermediate (midway between two sequential administrations, approximately 30 min) and trough (approximately 15 min before the next dose) concentrations. Creatinine was monitored daily.

Imipenem blood samples were drawn and immediately placed on ice, then transported to the laboratory for centrifugation. MOPS [3-(N-morpholino)propanesulfonic acid], a stabilizing buffer that protects imipenem from degradation [27], was added to an equivalent volume of plasma. Stabilized imipenem samples were subsequently stored at –80 °C for a maximum of 1 month.

Imipenem plasma concentrations were analyzed by high-performance liquid chromatography (HPLC) with ultraviolet (UV) detection at 298 nm. Ceftazidime was used as an internal standard in the HPLC-UV analysis. Acetonitrile was added to the stabilized plasma for deproteinization. The calibration curve was linear from 0.5 to 80 mg/L. Limit of detection (LOD) and limit of quantification (LOQ) were 0.2 mg/L and 0.5 mg/L, respectively [24].

### 2.3 Parametric Population Pharmacokinetic (popPK) Analysis (NONMEM)

Parametric popPK analyses were performed using nonlinear mixed-effects modelling (NONMEM version 7.2; ICON Development Solutions, Ellicott City, MD, USA). The Intel Visual Fortran Compiler XE 14.0 (Santa Clara, CA, USA) was used. The first-order conditional estimation method with interaction (FOCE-I) was used throughout the model-building process. Tools used to evaluate and visualize the model were RStudio (version 1.1.456), R (version 3.5.1), XPose (version 4.6.1) and PsN (version 4.6.0), all with the graphical interface Pirana [28] (version 2.9.4).

General model selection criteria were decrease in objective function value (ΔOFV), GOF plots and VPCs. A decrease in the OFV of 3.84 units was considered statistically significant (p < 0.05, degrees of freedom [df] = 1) in a nested model [29]. For each VPC, a set of 1000 simulated datasets was created to compare the observed concentrations with the distribution of the simulated concentrations. A numerical predictive check (NPC) of the final model was created to compare with the NPC of the final nonparametric model.

During modelling, only lower bounds (of 0) and no upper bounds were set for each parameter [29]. One-, two-, and three-compartment distribution models were evaluated [30]. Databases with untransformed and logarithmic transformed concentrations were compared by assessing GOF plots and parameter estimates. For both databases, residual unexplained variability was tested with proportional and combined (additive and proportional) error models [31]. The proportional (exponential) error model of the final NONMEM model with log-transformed data is shown in Eq. (1).

The observed concentration (OBS) consisted of the individually predicted concentration (IPRED) with added residual unexplained variability $\epsilon$ (epsilon, fixed to 1 in our model) weighted by an estimated error parameter:

$$\text{OBS} = \text{IPRED} + \sqrt{(error^2) \times \epsilon}$$  \hspace{1cm} (1)

Variability of a popPK parameter was estimated using an exponential variance model (individual popPK parameter = population popPK value × $e^\eta$). Eta ($\eta$) is a random variable drawn from a normal distribution with a mean of 0 and a variance of omega ($\omega^2$) [32]. The BSV (CV%) of a population parameter is calculated by Eq. (2) [33]. The SD is subsequently calculated by multiplying the CV% with the population parameter estimate.

$$\text{CV}(e^\eta) = \sqrt{(e^{\omega^2} - 1) \times 100\%}$$  \hspace{1cm} (2)
First, models with BSV on elimination rate constant \((K_e)\) and BSV on \(V\) were compared by assessing \(\Delta OFV\) and GOF plots. Subsequently, one-by-one addition of BSV on the other parameters was studied. A stepwise covariate model building was performed with forward addition at \(p < 0.05\) (\(\Delta OFV\) of 3.84 units, \(df = 1\)), followed by backward elimination at \(p < 0.001\) (\(\Delta OFV\) of 10.83 units, \(df = 1\)) [34]. Covariates were tested on parameters with BSV. The tested covariates are described in Sect. 2.5.

The 95% CI of each parameter in the final model was determined from a non-parametric bootstrap analysis, in which the dataset was resampled 1000 times.

2.4 Nonparametric popPK Analysis (Pmetrics)

Nonparametric popPK analysis was performed using Pmetrics version 1.5.2 (Laboratory of Applied Pharmacokinetics and Bioinformatics, Los Angeles, CA, USA) [6] in RStudio (version 1.1.456) as a wrapper for R (version 3.5.1), and the Intel Visual Fortran Compiler XE 14.0. The Non-Parametric Adaptive Grid (NPAG) program was used throughout the model-building process. The Iterative 2-Stage Bayesian (IT2B) program was used to estimate parameter ranges to pass to NPAG.

An NPAG will create a non-parametric popPK model consisting of discrete support points, each with a set of estimates for all parameters in the model plus an associated probability of that set of estimates (see Fig. 1 for an illustration of the distribution of a parameter) [6]. The sum of all probabilities is 1. There can be a maximum of 1 point for each subject in the study population [6]. Besides an overview of support points with corresponding parameter estimates, the NPAG output also contains the mean, SD and CV% of each parameter. The reported means are weighted means that are calculated by multiplying the estimate of each support point by the probability of that point and then summing up the resulting numbers. The SD is calculated from the parameter distribution. The BSV (CV%) of each parameter estimate is calculated by dividing the SD by the weighted mean.

One-, two-, and three-compartment distribution models were evaluated. A stepwise covariate model building was performed with forward addition at \(p < 0.05\) (decrease in \(- 2 \times \text{log-likelihood} [\Delta - 2\text{LL}]\) of 3.84 units, \(df = 1\)), followed by backward elimination at \(p < 0.001\) (\(\Delta - 2\text{LL}\) of 10.83 units, \(df = 1\)) [34]. Covariates were tested on parameters selected after a graphical examination of possible covariate–parameter relationships. The tested covariates are described in Sect. 2.5.

Each observation in Pmetrics is weighted by \(1/\text{error}^2\). Both gamma and lambda error models were tested (see Eqs. 3 and 4). The SD of an observation is based on the assay error polynomial (see Eq. 5) [6]; however, because the assay error polynomial was unavailable in our study, we estimated the error coefficients, with \(C_0 = 0.5 \times \text{LOQ}, C_1 = 0.1, C_2 = 0\) and \(C_3 = 0\) as a starting point.

error = SD \times \text{gamma} \tag{3}

error = \sqrt{SD^2 + \text{lambda}^2} \tag{4}

SD = C_0 + C_1 \times \text{OBS} + C_2 \times \text{OBS}^2 + C_3 \times \text{OBS}^3 \tag{5}

Model selection criteria were decrease in \(- 2\text{LL}\), bias, imprecision, GOF plots and VPCs. A decrease in the \(- 2\text{LL}\) of 3.84 units was considered statistically significant \((p < 0.05, df = 1)\) in a nested model. For each VPC, a set of 1000 simulated datasets was created to compare the observed concentrations with the distribution of the simulated concentrations. An NPC of the final model was created to compare with the NPC of the final parametric model. The raw VPC and NPC data were imported into PsN (version 4.6.0) using the Pirana interface [28] to generate plots with a similar layout as the parametric plots. VPC and NPC plots were created using XPose (version 4.6.1) within RStudio (version 1.1.456). Bias (mean weighted prediction error) and imprecision (bias-adjusted mean weighted squared prediction error) are automatically calculated by Pmetrics according to Eqs. (6) and (7), for both population and posterior predictions:

\[ \text{Bias} = \text{mean weighted prediction error} \]

\[ \text{Imprecision} = \text{bias-adjusted mean} \times \text{weighted squared prediction error} \]

\[ \text{Bias} = \frac{\sum (\text{predicted} - \text{observed})}{\sum \text{observed}} \tag{6} \]

\[ \text{Imprecision} = \frac{\sum (\text{predicted} - \text{observed})^2}{\sum \text{observed}^2} \tag{7} \]

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\[ \Delta \text{Adis} \]
\[
\text{Bias} = \frac{\sum (\frac{(\text{PRED}_i - \text{OBS}_i)}{\text{error}_i^2}) + \cdots + (\frac{(\text{PRED}_n - \text{OBS}_n)}{\text{error}_n^2})}{n}
\]

\[
\text{Imprecision} = \frac{\sum (\frac{(\text{PRED}_i - \text{OBS}_i)^2}{\text{error}_i^2}) + \cdots + (\frac{(\text{PRED}_n - \text{OBS}_n)^2}{\text{error}_n^2})}{n} - \text{bias}^2
\]

The 95% CI of each parameter in the final model was determined using a Monte Carlo simulation approach by creating 1000 samples with replacement for each support point [35], resulting in the 2.5th, 50th and 97.5th percentiles of the weighted median and the median absolute weighted deviation (MAWD). The SD was estimated by multiplying the MAWD by 1.4826 [36], while the CV% was calculated by dividing the SD by the 50th percentile of the weighted median.

### 2.5 Covariates

The tested covariates for both modelling approaches were total body weight (TBW), ideal body weight (IBW) [37], lean body weight (LBW) [37], CG eGFR [25], four-variable Modification of Diet in Renal Disease (MDRD) Study eGFR [38], Chronic Kidney Disease Epidemiology Collaboration (CKD-EPI) eGFR [39], and Jelliffe’s eGFR equation for patients with unstable renal function [40]. Per patient, one body weight measure was available, while a median of three creatinine samples per patient was drawn (see also Table 1). The MDRD, CKD-EPI and Jelliffe’s equations provide eGFR values adjusted for body surface area [BSA; mL/min/1.73 m²]. The BSA-unadjusted (absolute) values (mL/min) were also calculated by multiplying the original eGFR by the individual BSA [41] and evaluated as covariates: MDRD-abs, CKD-EPI-abs and Jelliffe-abs. The CG eGFR equation is unadjusted for BSA (mL/min) and no adjustment was made.

All covariates were evaluated by power models with normalized covariates where the median covariate value was taken as the reference value (see Eq. 8). Because multiple creatinine samples per patient were collected, which are each used to calculate CG, MDRD and CKD-EPI eGFR values, eGFR was tested as a time-varying covariate. We also tested the possible situation of reaching a maximum value of the elimination constant \(K_e\) for high (adjusted or unadjusted) eGFR values from 150, 120 and 90. For TBW, LBW and IBW, both fixed (− 0.25 for \(K_e\) and 1 for \(V\)) and estimated values of the power exponent were evaluated [42]:

\[
\text{Par}_{\text{ind}} = \text{Par} \times \left( \frac{\text{Cov}_{\text{ind}}}{\text{Cov}_{\text{median}}} \right)^{\text{power}} \times e^\eta
\]

where \(\text{Par}_{\text{ind}}\) is the individual PK parameter estimate; \(\text{Par}\) is the popPK parameter estimate (for NONMEM) or weighted median value of the Bayesian posterior distribution (for Pmetrics); Cov_{\text{ind}} is the individual covariate value; Cov_{\text{median}} is the median covariate value; power is the covariate effect; and \(\eta\) is the individual variability (\(\eta\) only for NONMEM).

See Eqs. (9) and (10) in the Results section for further clarification of the differences between equations in the final parametric and nonparametric models.

Default covariate settings were used for each modelling approach. In NONMEM, by default, the next observation is

| Parameter | Value |
|-----------|-------|
| Male [n (%)] | 18 (69) |
| APACHE II score [median (IQR)] | 22 (17–27) |
| Age, years [median (IQR)] | 51 (39–54) |
| Creatinine at inclusion, μmol/L [median (IQR)] | 59 (46–70) |
| Creatinine samples per patient [median (IQR)] | 1.5 (1.2–1.8) |
| eGFR at inclusion | 146 (123–170) |
| CKD-EPI, mL/min/1.73 m² [median (IQR)] | 116 (104–127) |
| CKD-EPI-abs, mL/min [median (IQR)] | 119 (110–139) |
| MDRD, mL/min/1.73 m² [median (IQR)] | 121 (104–159) |
| MDRD-abs, mL/min [median (IQR)] | 127 (118–162) |
| Jelliffe, mL/min/1.73 m² [median (IQR)] | 156 (132–183) |
| Jelliffe-abs, mL/min [median (IQR)] | 168 (141–202) |
| Height, cm [median (IQR)] | 175 (168–179) |
| Total bodyweight, kg [median (IQR)] | 75 (66–85) |
| Ideal bodyweight, kg [median (IQR)] | 70 (59–73) |
| Lean bodyweight, kg [median (IQR)] | 58 (46–64) |
| BMI, kg/m² [median (IQR)] | 25 (22–27) |
| BSA, m² [median (IQR)] | 1.89 (1.72–2.04) |
| Presumed infection [n (%)] | 16 (62) |
| Lower respiratory tract infection | 4 (15) |
| Intra-abdominal infection | 3 (12) |
| Surgical site infection | 1 (4) |
| Meningitis | 1 (4) |
| Gynecological infection | 1 (4) |

**Table 1** Demographic and clinical characteristics of the study population (N = 26)

**APACHE** Acute Physiology and Chronic Health Evaluation, **IQR** interquartile range, **eGFR** estimated glomerular filtration rate, **CG** Cockcroft–Gault, **CKD-EPI** Chronic Kidney Disease Epidemiology Collaboration, **MDRD-abs** absolute MDRD (i.e. MDRD multiplied by BSA), **BSA** body surface area

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carried backward (NOCB) until the time point of the previous covariate observation. For Pmetrics, covariates are applied at each dose event. By default, for missing covariate values, the last observation is carried forward (LOCF) until the last dose before the next covariate observation, when the last observation is linearly interpolated to the next observation.

### 2.6 Model Development and Comparison

Both models were developed by a medium-experienced NONMEM and Pmetrics modeler (FdV) supervised by highly experienced NONMEM (BdW) and Pmetrics (WY, MN) modelers. The development of both models occurred independently from each other according to a predefined study procedure, where the modelling workflow (e.g. model selection criteria, building of the structural model and covariate evaluation) was described. During model development, the GOF plots and VPCs were assessed in the layout of the separate programmes. During writing of the manuscript, the raw data of the GOF plots were transferred to GraphPad Prism (version 8.1.1) and the raw VPC and NPC data of Pmetrics were transferred to PsN (see Sect. 2.4) to create plots with the same layout. The $R^2$ of nonlinear regression was calculated within GraphPad Prism by the fourth equation of Willett and Singer [43] for the GOF plots as a description of the graphical fit. $R^2$ was not used during model selection.

### 3 Results

#### 3.1 Study Population

Demographic and clinical characteristics of the 26 included patients are summarized in Table 1. None of the patients received continuous renal replacement therapy (CRRT).

#### 3.2 Imipenem Samples

In total, 138 imipenem blood samples were collected from 26 patients and were subsequently analyzed. Fewer than 10% [30] of all concentrations (13/138, 9.4%) were below the limit of quantification (0.5 mg/L) and were excluded from the popPK analysis. The average number of levels per patient was 5 (range 1–11). Almost half of all samples (65/138, 47.1%) were drawn on the second day of therapy.

A graph of the analyzed imipenem concentrations ($n = 125$) plotted against the time after dose is shown in Electronic Supplementary Fig. 1. These 125 concentrations were drawn after 84 individual doses. Following 33 of these doses, two or three concentrations were taken. For the other 51 doses, one concentration was drawn. For all PK samples, a creatinine concentration was available in the 24 h before PK sampling.

#### 3.3 Parametric PopPK Model

The parametric popPK analysis using NONMEM showed that the data were best described by a model with two distribution compartments, BSV on $K_e$ and CKD-EPI-abs as a covariate on $K_e$. The parameter estimates of the final model are displayed in Table 2. Only BSV on $K_e$ was included (see also Fig. 1) because BSV on the central distribution volume ($V_c$), rate constant from the central to peripheral compartment ($K_{cp}$) and rate constant from the peripheral to central compartment ($K_{pc}$) did not significantly improve the model ($ΔOFV < 3.84$, and no improvement in GOF plots). $V_c$, $K_{cp}$ and $K_{pc}$ were the same for each subject (no BSV was included, therefore no CV% is shown in Table 2). Eta shrinkage was low (14%), and no large correlation ($>0.95$) between the parameters was detected [29]. The GOF and VPC plots (displayed in Figs. 2a, 3a and Electronic Supplementary Fig. 2a) show good predictive performance for all concentrations, for the time range after dose and for the CKD-EPI-abs range of 18–190 mL/min (except from an outlier for the 124–141 mL/min bin). The NPC did not show points outside boundaries (data not shown). As shown in Table 2, the model-based parameter estimates were similar to the bootstrap values, indicating stability of the model.

A two-compartment model described the data better than one- or three-compartment models, according to GOF plots and an OFV increase of 22.99 for a one-compartment model and 0 for a three-compartment model. The popPK parameters of a two-compartment model with logarithmic transformed data were comparable to the same model with untransformed data. The GOF plots were improved by logarithmic transformation and therefore the following analyses were performed with transformed data. An exponential (proportional) error model (see Eq. 1 in Sect. 2.3) was preferred to a combined error model due to estimation problems with the combined model. This confirmed the findings of the untransformed data, where the proportional error model had better performance than the combined error model.

Several covariates (as described in Sect. 2.5) were tested on $K_e$, the only parameter with BSV. All tested eGFR covariates on $K_e$ resulted in a significant OFV decrease ($ΔOFV 24.4$ until 38.3, $p < 0.05$) compared with the two-compartment model without covariates. However, the OFV of the model with CKD-EPI-abs as a covariate on $K_e$ was significantly better than the other eGFR models (for example, the second best eGFR was CKD-EPI on $K_e$, with an OFV increase of 4.3 compared with CKD-EPI-abs; $p < 0.05$). Due to the observation of a maximum eta in the eta-eGFR plots, implementation of a maximum $K_e$ value for eGFR values from 150, 120 and 90 was tested but this did not
further improve the model. None of the tested measures of body weight improved the model as a covariate on $K_e$ ($\Delta$OFV < 3.84) during the univariate analysis.

Equation (9) describes the calculation of individual $K_e$ ($K_{e,i}$) values in the final NONMEM model, using the population parameter estimates for $K_e$ and $K_e$(cov) from Table 2, the individual CKD-EPI-abs value at a certain time point, and $e^\eta$ (individual variability). Eta ($\eta$) is drawn from normal distribution with a mean of 0 (note Eq. 9, then $e^{0i} = 1$) and variance $\omega^2$ (estimated from the data as 0.0354). The corresponding BSV for $K_e$ (CV% 19.0%) was calculated from Eq. (2):

$$K_{e,i} = 0.637 \times \left(\frac{\text{CKD} - \text{EPI}-\text{abs}_i}{119}\right)^{0.655} \times e^\eta \quad (9)$$

A plot of the individual $K_e$ against the individual CKD-EPI-abs ($n = 86$) is shown in Electronic Supplementary Fig. 3a. Simulated concentration-time profiles for CKD-EPI-abs of 150, 120 and 90 mL/min ($n = 1000$ for each eGFR) are shown in Electronic Supplementary Fig. 4a.

The median (interquartile range) of the untransformed residuals (observed minus predicted concentrations) was $-0.159$ mg/L ($-0.960$ to $1.267$) for the population predictions and $-0.027$ mg/L ($-0.649$ to $0.698$) for the individual predictions. Residual plots are shown in Electronic Supplementary Fig. 5a.

### 3.4 Nonparametric PopPK Model

The nonparametric popPK analysis using Pmetrics resulted in the same model structure as the parametric analysis: a model with two distribution compartments and CKD-EPI-abs as a covariate on the elimination constant $K_e$. The mean parameter estimates of the final model are displayed in Table 2. For example, the mean $K_{e,\text{pop}}$ value is the mean of the support points weighted by population probabilities. This is illustrated in Fig. 1. Each individual has a Bayesian posterior (i.e. personal or individual) set of support points weighted by individual probabilities [6]. No large correlation (> 0.95) between the parameters was detected. The GOF and VPC plots (displayed in Figs. 2b and 3b and Electronic Supplementary Fig. 2b) show good predictive performance for all concentrations, for the time range after dose and for the CKD-EPI-abs range of 18–190 mL/min (except from an outlier for the 124–141 mL/min bin, similar to the parametric model). Also similar to the parametric model, the NPC did not show points outside boundaries (data not shown). As shown in Table 2, the model-based parameter estimates were similar to the bootstrap values, indicating stability of the model.

### Table 2 Population parameter estimates

| Parameter | NONMEM | Bootstrap |
|-----------|--------|-----------|
|           | Final model | Bootstrap |
|           | Parameter estimate | CV (%) | Median parameter estimate | 95% CI parameter estimate | Median CV (%) | 95% CI CV (%) |
| $V_c$ (L) | 29.6 | – | 29.4 | 22.9–34.4 | – | – |
| $K_{cp}$ (h$^{-1}$) | 0.166 | – | 0.169 | 0.092–0.436 | – | – |
| $K_{pc}$ (h$^{-1}$) | 0.192 | – | 0.192 | 0.079–0.604 | – | – |
| $K_e$ (h$^{-1}$) | 0.637 | 19.0 | 0.634 | 0.543–0.805 | 18.6 | 10.5–27.4 |
| $K_e$(cov) | 0.655 | – | 0.665 | 0.474–1.184 | – | – |
| Exponential error (mg/L) | 0.348 | – | 0.340 | 0.281–0.413 | – | – |

$V_c$, central distribution volume, $K_{cp}$ rate constant from the central to peripheral compartment, $K_{pc}$ rate constant from the peripheral to central compartment, $K_e$ elimination rate constant, $K_e$(cov) covariate effect on $K_e$, CV coefficient of variation, CI confidence interval.
A two-compartment model described the data better than one- or three-compartment models, according to GOF plots and a \(-2LL\) increase of 26.5 for a one-compartment model and 0.8 for a three-compartment model. The gamma error model was preferred to the lambda model (see Eqs. 3 and 4 in Sect. 2.4). The final values for \(C_0\) and \(C_1\) in the assay error polynomial were both 0.05, and \(C_2\) and \(C_3\) were both 0 (see Eq. 5 in Sect. 2.4). Parameter boundaries \((K_e, 0–1.5; V, 1–70; K_{cp} \text{ and } K_{pc}, 0–1)\) were set based on an IT2B run.

We evaluated several covariates (as described in Sect. 2.5) on \(K_e\) and \(V\). All tested eGFR covariates on \(K_e\) resulted in a significant \(-2LL\) decrease \((\Delta -2LL 53.3 \text{ until 59.9}; \ p < 0.05)\) compared with the two-compartment model without covariates. The \(-2LL\) value of the four models with the largest \(-2LL\) decrease (MDRD, MDRD-abs, Jelliffe-abs, and CKD-EPI-abs) did not differ significantly from each other. The model with CKD-EPI-abs had the lowest bias and imprecision compared with the three other models. Implementation of a maximum \(K_e\) value for eGFR values from 150, 120 or 90 did not improve the model. Univariate analysis resulted in a further six significant covariates \((p < 0.05)\): TBW, IBW and LBW on \(K_e\) \((\Delta -2LL 4.0 \text{ until 8.8})\), and CG, Jelliffe and Jelliffe-abs on \(V\) \((\Delta -2LL 4.6 \text{ until 5.5})\). After backward elimination at \(p < 0.001\), none of these six covariates remained in the final model.

Equation (10) describes the calculation of individual \(K_e\) values \((K_e_i)\) in the final Pmetrics model. Contrary to the parametric model with equal \(K_e\) and \(K_e(cov)\) values for each individual subject (see Eq. 9), these values are different for each individual subject in the nonparametric model. \(K_{e,med}\) is the weighted median value of the Bayesian posterior distribution for \(K_e\) in the \(i\)th individual, \(K_{e,cov,med}\) is the weighted median value of the Bayesian posterior distribution for \(K_{e,cov}\) in the \(i\)th individual, and CKD-EPI-abs\(_i\) is the individual CKD-EPI-abs value at a certain time point:

\[
K_{e_i} = K_{e,med} \times \left( \frac{\text{CKD} - \text{EPI} - \text{abs}_i}{119} \right)^{K_{e,cov,med}}
\]

\(\Delta\) Adis
A plot of the individual $K_e$ against the individual CKD-EPI-abs ($n = 86$) is shown in Electronic Supplementary Fig. 3b. This plot shows a similar $K_e$ versus CKD-EPI-abs relationship as the parametric model, although the $K_e$ distribution is wider for the nonparametric model. A wider distribution is also shown in the simulated concentration-time profiles for CKD-EPI-abs of 150, 120 and 90 mL/min (Electronic Supplementary Fig. 4b).

The median (interquartile range) of the residuals (observed minus predicted concentrations) was $-0.045$ mg/L ($-0.498$ to $1.506$) for the population predictions and $0.011$ mg/L ($-0.295$ to $0.533$) for the individual predictions. Residual plots are shown in Electronic Supplementary Fig. 5b. These are similar to the parametric plots.

4 Discussion

The structure and parameter estimates of our two independently developed parametric and nonparametric popPK models of imipenem in critically ill patients treated with imipenem/cilastatin were similar; both included two distribution compartments and CKD-EPI-abs as a covariate on $K_e$. Body weight as a covariate was not found to be a significant covariate in either model. Both models described imipenem PK well. Two main differences between the models emerged. First, the parametric model included BSV for $K_e$ only, while the nonparametric model included such variability on all popPK parameters. Second, the estimated BSV (defined as CV%) for $K_e$ was higher for the nonparametric model (34.0% vs. 19.0%). The findings of similar parameter estimates but higher BSV for the nonparametric model are in line with
two previously published studies comparing parametric and nonparametric models of other drugs [12, 13]. These BSV differences could be explained by the statistics behind the modelling methods. Both models have fixed (no BSV) and random (including BSV) parameters. However, while during parametric modelling the inclusion of BSV is examined for each popPK parameter, all popPK parameters for nonparametric models are, in principle, random parameters, while the residual error model is fixed. In addition, while parametric methods assume a normal or lognormal distribution of parameters, nonparametric methods make no assumption about the parameter distributions, which can cause a wider CI of the parameter estimates (e.g. see Fig. 1 for $K_c$, which also clearly shows the differences between the two CV% measures, and Electronic Supplementary Fig. 3). The simulated concentration-time profiles in Electronic Supplementary Fig. 4 indicate that the concentrations of the 2.5th percentile are approximately twofold lower for the nonparametric model. The consequences of this finding (e.g. higher or more frequent dosing following the nonparametric model) should be further explored by probability of target attainment calculations based on extensive Monte Carlo simulations for several dosing regimens [44].

Our finding of two distribution compartments is in accordance with other published popPK models of imipenem in critically ill patients (all with pneumoniae) [45–47]. However, our $V_c$ and clearance ($CL = \frac{V_c}{K_e}$) values were higher than previously described [45–47]. This could be explained by a higher CLcr in our population, which could be attributed to augmented renal clearance (ARC) [48, 49]. ARC is defined as increased renal elimination of circulating solutes and drugs compared with normal baseline [23] and has been reported in approximately 30–65% of critically ill patients [48]. Our $K_{cp} (0.2–0.4)$ and $K_{pc} (0.2–0.5)$ values of both models were similar to the previously published parametric model [45], but remarkably different from the two nonparametric models ($K_{cp} 3–8$ and $K_{pc}$ approximately 9) [46, 47]. Possibly, these differences could be explained by (unpublished) wider parameter boundaries of their nonparametric models. The parameter ranges in nonparametric software are strict boundaries wherein the optimal values are sought. To assist with optimal setting of parameter ranges in nonparametric popPK, we used the parametric IT2B module in Pmetrics to estimate the parameter ranges to pass to the nonparametric NPAG module. During parametric modelling with NONMEM, the parameter initial estimate is a starting point to search for the optimal value, and setting limits is not always necessary [29]. This was underlined by the fact that the results of our final NONMEM model with only lower boundaries were the same as a model with the same upper and lower boundaries as the final Pmetrics model.

Many types of standard GOF plots used for model evaluation are applied both in parametric and nonparametric modelling: observed versus population predicted concentrations; observed versus individually predicted concentrations; and weighted residual (WRES) plots. The observed versus predicted concentration plots of our popPK models are comparable. In Pmetrics, the standard layout of these plots includes the $R^2$, intercept, slope, bias (mean WRES) and imprecision. In NONMEM, these measures are not automatically calculated. A residual is the difference between an observed and a predicted concentration. WRESs are used in WRES plots, as well as for the calculation of bias and imprecision. We did not show WRES plots nor calculated weighted bias and imprecision for both methods because of two reasons. First, the weighting differs between the two modelling methods. In NONMEM, the conditional WRES (CWRES) is the residual weighted by the square root of the covariance of a FOCE model [50], while in Pmetrics, WRES is the residual weighted by the squared error [35]. Second, a WRES of logarithmic transformed data (in our NONMEM model) is not the same as a WRES of untransformed data (in our Pmetrics model). We used weighted bias and imprecision in Pmetrics only to differentiate between four covariate models with a −2LL decrease that did not differ significantly from each other. As an alternative to weighted bias, we calculated unweighted bias (the unweighted residuals of untransformed concentrations), of which the median and interquartile range were comparable for both methods.

The VPCs of both popPK models indicated a sufficient predictive performance. The means of the 95th, 50th and 5th percentiles of predictions are comparable for both models, but the 95% CIs of these percentiles differ for some bins (timeframes) of the VPCs. Most remarkably, the 95% CI of the 95th and 50th percentile of the last two bins (4.3–5.8 h after dose) is wider for the nonparametric model. This could be explained by a higher estimated BSV of $K_e$ in the nonparametric model, leading to a wider distribution of concentrations in the elimination phase. We did not show prediction-corrected VPCs (pcVPCs) because this option has been developed for parametric methods [51] and has not yet been tested for nonparametric methods. In a pcVPC, the variability coming from binning across the independent variable, e.g. due to different doses or influential covariates, is removed [51]. The pcVPCs of the NONMEM model did not show important differences from the traditional VPCs (data not shown).

Both models include CKD-EPI-abs as a covariate on $K_e$. Two of the three previous mentioned published popPK models of imipenem also included renal function as a covariate, but other measures were used, i.e. 4 h CLcr in urine [45] and the CG equation [47]. Sakka et al. did not find 12 h CLcr in urine as a significant covariate [46]. Besides eGFR, other covariates were also included in the published models, i.e. body weight [45–47], height [46, 47], BSA [46, 47], age [46, 47], sex [47] and albumin [45]. In our covariate screening...
plots, sex, age and height did not show a relationship with any PK parameters. Body weight did show a relationship in the covariate plots, but inclusion as a covariate did not improve the model. Albumin data were not available. BSA was taken into account during eGFR covariate testing. We tested both BSA-unadjusted (mL/min) and BSA-adjusted (mL/min/1.73 m²) eGFR equations because the European Medicines Agency [52] and the Kidney Disease Improving Global Outcomes (KDIGO) guideline [53] recommends to base dosing on absolute instead of BSA-normalized eGFR. The KDIGO [53] also recommends using the CKD-EPI eGFR equation. However, this guideline is based on chronic kidney disease, while our study population consisted of critically ill patients with a high median CKD-EPI eGFR of 116 mL/min/1.73 m². The correlation between measured $\text{CL}_{\text{cr}}$ and eGFR equations is known to be weak in critically ill patients [23, 54]. Nonetheless, the measurement of $\text{CL}_{\text{cr}}$ (as a surrogate for GFR) is time-consuming and is not standard practice in many ICUs. In daily practice, the eGFR is also used for dosing drugs with renal clearance, although many patients have a renal function that is not in steady state. Therefore, we decided to test several eGFR equations to find the most suitable one in our population. Minichmayr et al. performed a similar eGFR covariate analysis for meropenem in critically ill patients and found that the CG equation best described meropenem clearance [55]. We observed a maximum in the BSV–eGFR plots; however, the implementation of a maximum $K_e$ value for eGFR values from 150, 120 and 90 did not further improve the model. As already mentioned, the correlation between measured $\text{CL}_{\text{cr}}$ and eGFR equations is weak, but it is also shown that this correlation varies over the eGFR range [23]. For example, in a previous study it was shown that the CKD-EPI equation performed better for measured $\text{CL}_{\text{cr}} < 120$ mL/min than for $\text{CL}_{\text{cr}} > 120$ mL/min [23]. We did not confirm this finding in our study (see Electronic Supplementary Fig. 2), which could be explained by the different study population.

For both modelling approaches, eGFR was implemented as a time-varying covariate using a stepwise (discontinuous) approach. The default covariate settings of both methods were slightly different (i.e. NOCB without interpolation or LOCF with linear interpolation from the last dose before the next covariate value). However, the parameter estimates of both final models, developed with the default settings, were very similar to the same models using LOCF without interpolation. This is explained by the majority (85%) of patients having reasonably stable CKD-EPI-abs eGFR values around PK sampling, according to the KDIGO definition of an eGFR drop or rise of < 25% compared with the previous value [53]. This stability statement is supported by frequent creatinine monitoring. A creatinine concentration in the 24 h before TDM was available for all PK samples. Due to the stable eGFR values, a continuous covariate approach was not necessary.

We performed our study using the most used parametric (NONMEM) and nonparametric (Pmetrics) software. For approximately a decade, a nonparametric method also exists in NONMEM, and some publications [19, 56–58] regarding the evaluation and optimization of this approach are available. However, this method is seldom used in clinical practice. The parametric module IT2B in Pmetrics is used to estimate parameter ranges to pass to NPAG, but underperformed compared with other parametric algorithms [12].

One of the limitations of our study is that we used eGFR equations for a critically ill population with a high frequency of augmented clearance; however, these equations are developed for more stable patients with chronic kidney disease. Nonetheless, as measured $\text{CL}_{\text{cr}}$ was unavailable and is also not standard practice in many ICUs, we aimed to find the eGFR equation that would best describe imipenem clearance. Another drawback is that we could not compare WRESs because the weighting is different for both methods and the residuals of untransformed data (used for the nonparametric model) are different from transformed data (used for the parametric model). Other limitations are the small number of subjects and the absence of an external validation. It would be interesting to compare the predictions of both models.

## 5 Conclusions

The general structure and the parameter estimates of both models were comparable. The identical covariate results (CKD-EPI-abs on $K_e$) of the two different modelling methods strongly support the findings in this population. The nonparametric model included BSV for all parameters, while the parametric model only included BSV on $K_e$. The estimated BSV of $K_e$ was higher in the nonparametric model. The consequences of the BSV differences may affect estimated exposure during dosing simulations, and this should be further investigated in simulation studies.

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### Compliance with Ethical Standards

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**Ethical Approval** All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional research committee (Geneva University Hospital’s Ethics Committee, NAC 09-117) and with the 1964 Helsinki declaration and its later amendments.

**Informed Consent** Informed consent was obtained from individual participants included in this study, unless patients were unconscious or otherwise unable to understand the study protocol. This was approved by the abovementioned Ethics Committee, giving the observational nature of the study.

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