Rifampicin Alters Metformin Plasma Exposure but Not Blood Glucose Levels in Diabetic Tuberculosis Patients

Lindsey H.M. te Brake1,2,†, Vycke Yunivita3,†, Resvi Livia4, Nanny Soetedjo5, Eleonora van Ewijk-Beneken Kolmer1, Jan B. Koenderink2, David M. Burger1, Prayudi Santoso6, Reinout van Crevel7, Bachti Alisjahbana4, Rob E. Aarnoutse1, Rovina Ruslami3 on behalf of the TANDEM Consortium

The pharmacokinetic (PK) and clinical implications of combining metformin with rifampicin are relevant to increasing numbers of patients with diabetic tuberculosis (TB) across the world and are yet unclear. We assessed the impact of rifampicin on metformin PKs and its glucose-lowering effect in patients with diabetic TB by measuring plasma metformin and blood glucose during and after TB treatment. Rifampicin increased metformin exposure: plasma area under the plasma concentration-time curve from time point 0 to the end of the dosing interval (AUC0–\(\tau\)) and peak plasma concentration (Cmax) geometric mean ratio (GMR; during vs. after TB treatment) were 1.28 (90% confidence interval (CI) 1.13–1.44) and 1.19 (90% CI 1.02–1.38; \(n=22\)). The metformin glucose-lowering efficacy did not change (\(\Delta\)glucose − Cmax; \(P=0.890; n=18\)). Thus, we conclude that additional glucose monitoring in this population is not warranted. Finally, 57% of patients on metformin and rifampicin, and 38% of patients on metformin alone experienced gastrointestinal adverse effects. Considering this observation, we advise patients to take metformin and rifampicin with food and preferably separated in time. Clinicians could consider metoclopramide if gastrointestinal adverse effects occur.

Tuberculosis (TB) remains a leading cause of morbidity and mortality in developing countries. In 2017, an estimated 10.0 million people developed active TB and 1.6 million patients died.1 At the same time, ~415 million people had diabetes mellitus (DM). The DM prevalence is growing rapidly, especially in low-income and middle-income countries, where TB is endemic. Furthermore,
DM increases the risk of developing active TB. Overall, there is an increasing number of TB cases attributable to DM, namely 10% in 2010, and 15% in 2013. Patients with concurrent TB and DM face a higher risk of TB treatment failure, relapse after cure, and death. DM management in patients with TB is also problematic. The TB drug rifampicin may affect blood glucose concentrations and induce hyperglycemia by augmenting intestinal absorption of glucose or reducing insulin sensitivity. More importantly, rifampicin increases the clearance of most oral antidiabetic drugs that are commonly used in low-income to middle-income countries. For example, sulphonylureas are metabolized in the liver by cytochrome P450 enzymes, of which rifampicin is a very potent inducer. To overcome the effects of rifampicin on treatment and maintenance of glycemic control, metformin has been proposed as a good alternative to other oral antidiabetic drugs, as it is not metabolized in the liver. Moreover, metformin is the first choice antidiabetic according to type 2 DM treatment guidelines. The drug is relatively cheap, widely available, and not associated with weight gain or hypoglycemia. The main disadvantage of metformin is the risk for lactic acidosis and the high frequency of gastrointestinal adverse effects. The oral absorption, hepatic uptake, and renal excretion of metformin are largely mediated by organic cation transporters (OCTs) and multidrug and toxin extrusion protein 1 and 2K (MATE1 and MATE2K), all members of the solute carrier family. It is eliminated unchanged in the urine with a half-life of ~5 hours. Its renal clearance is greater than that of creatinine, indicating that tubular secretion contributes to its elimination.

There is very limited data on co-administration of metformin and rifampicin and the need for dose adjustments in patients with TB-DM. Rifampicin is an agonist of the pregnane X receptor (PXR), a transcription factor that upregulates a large number of genes involved in xenobiotic detoxification, including drug-metabolizing enzymes and drug transporters. In rats, the PXR agonist pregnenolone-16-carbonitrile upregulated the expression of OCT1 in the liver and OCT2 in the kidneys, which significantly reduced metformin plasma exposure. In healthy volunteers, rifampicin caused altered metformin absorption kinetics, leading to higher exposure levels and enhanced glucose-lowering action. Extrapolation of these results to complex patients with TB-DM should be made cautiously, because disease status can alter transporter expression levels, and patients with diabetes have altered glucose regulation compared with healthy volunteers.

In summary, the pharmacokinetic (PK) and clinical implications of combining metformin with rifampicin are relevant to increasing numbers of patients with TB with type 2 DM across the world and are yet unclear. We, therefore, assessed the effect of rifampicin on the steady-state PK parameters and glucose-lowering effect of metformin in patients with TB-DM.

RESULTS

Subjects

From the TANDEM TB-DM cohort, 57 patients were eligible and asked for informed consent, of which eventually 24 patients (12 women and 12 men) participated in the study. For further details on eligibility, inclusion, and exclusion of patients see Supplementary Files S1–S4. Nineteen patients were on 500 mg of metformin (test dose during PK day), of which two patients were once daily, nine patients were twice daily, four patients were thrice daily, and three patients were thrice daily in combination with 850 mg doses. One subject switched from twice daily 500 mg to thrice daily 500 mg in between the PK sessions. This was corrected for by comparing area under the plasma concentration-time curve (AUC) across the entire dosing interval. Five patients were taking 850 mg of metformin (test dose during PK day) all thrice daily. Median fasting blood glucose during the first screen was 152 (range: 91–329) mg/dL and estimated glomerular filtration rate (eGFR) amounted to 101 (range: 65–141) mL/min. Median age at both sessions was 51 years (range: 24–63 years). Body mass index remained comparable between sampling sessions 1 and 2, with 23 (14–34) kg/m² and 24 (16–35) kg/m², respectively, supporting within-subject comparability of the two sessions (P = 0.339).

Effect of rifampicin on metformin PKs

Steady-state metformin PKs were assessed for subjects who underwent PK sampling in the first (n = 24) and second session (n = 22, Table 1). Rifampicin increased metformin exposure: for rifampicin + metformin (during TB treatment) relative to metformin alone (after TB treatment) the geometric mean ratio estimates of area under the plasma concentration-time curve from time point 0 to the end of the dosing interval (AUC₀₋₉₆) and peak plasma concentration (C₉₀₀) were 1.28 (90% confidence interval (CI) 1.13–1.44) and 1.19 (90% CI 1.03–1.38), respectively. Tubular secretion of metformin remained comparable (P = 0.777; Table 1 and Figure 2a–c). Creatinine clearance was slightly reduced (~4 mL/min) after rifampicin discontinuation.
Table 1 Comparison of steady-state metformin pharmacokinetic parameters and renal elimination parameters with and without co-administration of rifampicin

<table>
<thead>
<tr>
<th>Parameter</th>
<th>With rifampicin (n = 24)</th>
<th>Without rifampicin (n = 22)</th>
<th>P value or GMR estimate (90% CI) (n = 22)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AUC_{0–τ} (mg/L × hour)</td>
<td>15.4 (7.8–32.8)</td>
<td>12.1 (8.1–21.6)</td>
<td>1.28 (1.13–1.44)^a</td>
</tr>
<tr>
<td>C_{max} (mg/L)</td>
<td>2.3 (1.25–5.22)</td>
<td>2.0 (1.10–3.86)</td>
<td>1.19 (1.03–1.38)^b</td>
</tr>
<tr>
<td>T_{max} (hour) – median</td>
<td>1.5 (1–4)</td>
<td>1.5 (0–6)</td>
<td></td>
</tr>
<tr>
<td>CL/F (L/hour)</td>
<td>36 (21–69)</td>
<td>47 (34–73)</td>
<td>&lt;0.001^b</td>
</tr>
<tr>
<td>V_{d}/F (L)</td>
<td>332 (201–681)</td>
<td>310 (87–477)</td>
<td>0.586^b</td>
</tr>
<tr>
<td>t_{1/2} (hour)</td>
<td>6.4 (4.2–17.2)</td>
<td>4.6 (1.6–6.6)</td>
<td>0.002^b</td>
</tr>
<tr>
<td>AUC_{0–8 hour} (mg/L × hour)</td>
<td>13.4 (7.8–32.8)</td>
<td>10.9 (7.4–11.2)</td>
<td>0.083 (n = 14)</td>
</tr>
<tr>
<td>AUC_{0–τ} (mg/L × hour) –excluding sessions with ≥ 20% extrapolation</td>
<td>15.1 (7.8–32.8)</td>
<td>12.2 (8.1–21.6)</td>
<td>0.005^b</td>
</tr>
<tr>
<td>CL_{creatinine} (mL/min)^c</td>
<td>110 (88–155)</td>
<td>106 (74–148)</td>
<td></td>
</tr>
<tr>
<td>CL_{0–8 hour, renal} (mL/min)^d</td>
<td>309 (90–625)</td>
<td>332 (126–659)</td>
<td>0.836^b</td>
</tr>
<tr>
<td>CL_{0–8 hour, secretion} (mL/min)^d</td>
<td>205 (1–494)</td>
<td>219 (52–533)</td>
<td>0.777^b</td>
</tr>
<tr>
<td>CL_{0–τ, secretion} (mL/min)^d</td>
<td>212 (27–494)</td>
<td>216 (52–533)</td>
<td>0.856^b</td>
</tr>
</tbody>
</table>

Metformin plasma concentrations were extrapolated from 8–12 or 24-hours postdose to calculate AUC_{0–τ}, for patients on once or twice daily metformin, respectively. Pharmacokinetic parameters are expressed as geometric mean (range), unless stated otherwise. AUC_{0–τ} area under the plasma concentration-time curve from time point 0 to the end of the dosing interval; CI, confidence interval; CL, clearance; C_{max}, maximum plasma concentration; F, bioavailability; GMR, geometric mean ratio; t_{1/2}, half-life; T_{max}, time to maximum plasma concentration; V_{d}, volume of distribution.

^aEvaluation of a pharmacokinetic interaction by means of the bioequivalence approach. ^bPaired-samples t test on log-transformed pharmacokinetic parameters.

Figure 2 Individual changes in steady-state metformin pharmacokinetic parameters, AUC_{0–τ} (a), C_{max} (b) and tubular secretion (c), with and without co-administration of rifampicin. Metformin plasma concentrations were extrapolated from 8 until 12–24 hours to calculate AUC_{0–τ} for patients taking metformin once or twice daily. Tubular secretion was based on observed urine collections up to 8 hours after metformin intake. Data were assessed with paired-samples t test on the log-transformed parameters. AUC_{0–τ} area under the plasma concentration-time curve from time point 0 to the end of the dosing interval; C_{max}, maximum plasma concentration. ETH, ethambutol; IC, informed consent; INH, isoniazid; MTF, metformin; PK-GC, pharmacokinetic-glucose curve sampling session; PZA, pyrazinamide; RIF, rifampicin.

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The PK parameters of rifampicin are shown in Supplementary File S2 and demonstrate significant exposure to the perpetrator drug.

Effect of rifampicin on the glucose-lowering efficacy of metformin

Blood glucose concentrations for both sessions are displayed in Supplementary File S5. Fasting blood glucose concentrations differed slightly (15% reduction) during and after rifampicin treatment, but this did not reach statistical significance ($P = 0.081$, $n = 18$; Table 2). Similarly, the ability of metformin to reduce maximum blood glucose levels ($G_{\text{max}}$) and the AUC of glucose was not significantly different when measured during and after TB treatment (Table 2 and Figure 3).

Drug adherence and adverse events

Patients’ adherence to metformin and TB drugs was good. Mean percentages of drug adherence prior to the first session, according to physician assessment, pill count, self-assessment, and diary were 100% ($n = 24$), 99% ($n = 24$), 100% ($n = 24$), and 92% ($n = 19$), respectively. Adherence was 100% ($n = 22$), 96% ($n = 22$), 100% ($n = 22$), and 100% ($n = 17$) for the second session, respectively. We expect any nonadherence to be of minor importance, because we actively stimulated drug intake during the final 3 days before PK sampling by personally contacting each patient.

Of the 24 patients who underwent PK sampling, 13 patients experienced gastrointestinal adverse event(s) when they took both metformin and rifampicin (57%; during TB treatment), and 8 patients experienced gastrointestinal adverse event(s) when they took only metformin (38%; after TB treatment; Table 3). These additional gastrointestinal events could be attributed to more cases of nausea and vomiting (Table 3). Data do not include the cases that were excluded from the analysis due to vomiting on the PK-glucose curve (GC) days (Supplementary File S4). This occurred mainly prior to the addition of metoclopramide and separation of intake of metformin and TB drugs; four out of the first six patients vomited during PK-GC sampling session 1 (67%), with $n = 2$ during PK-session 1 and $n = 4$ during GC-session 1. All adverse events were grades 1 or 2.

DISCUSSION

This study was the first to evaluate the effect of rifampicin on metformin exposure and activity in patients with TB-DM. We observed an increase in metformin exposure, with an AUC0–$\tau$ GMR (exposure during vs. after TB treatment) of 1.28 (90% CI 1.13–1.44), suggesting a PK interaction (bio-inequivalence) when metformin is co-administered with rifampicin.23 This interaction did not result in a statistically significant change in the glucose-lowering effect of metformin.

Strikingly, renal clearance and tubular secretion were unaffected by rifampicin in our study. Because metformin is also not metabolized, we hypothesize that the higher metformin total and peak exposures are the result of increased metformin absorption. Metformin has relatively poor oral absorption with a bioavailability of 55%, 14 leaving substantial space for absorption increases. OCT1, OCT3, and plasma membrane monoamine transporter (PMAT) are the relevant metformin transporters in the small intestine, with PMAT being the most abundant.14,24 Rifampicin-induced upregulation of these transporters is probably responsible for the increased absorption assumed in our study. Indeed,
rifaximin increased OCT1 expression in blood cells in healthy volunteers. Further study is needed to evaluate the inductive effect of rifampicin on tissue (e.g., intestinal) expression of metformin transporters to confirm if this mechanism explains the rifampicin-metformin interaction.

The effect of rifampicin on systemic exposure to metformin is consistent with the findings of Cho et al. in healthy volunteers, although the PK interaction in our study seems larger (14–18% vs. 17–30%). This difference may be explained by the enhanced renal clearance and tubular secretion in the healthy volunteer study, whereas we did not identify such an elimination effect. Disease status may have altered expression levels of transporters important for metformin elimination, such as OCT1, MATE1, or MATE2K. Information on the transcriptional regulation or inducibility of the MATE family in humans is lacking.

Importantly, increased metformin exposure in our study was not associated with any clinically relevant or statistically significant increase in the glucose-lowering effect of metformin. Several explanations can be brought forward. First, rifampicin may have differential effects on expression of the various drug transporters in the intestine, liver, and kidneys. In humans, the liver has a central role in the regulation of systemic glucose, but also the kidneys contribute to glucose uptake, gluconeogenesis, and glucose utilization. Theoretically, altered OCT1 and OCT2 expressions influence local metformin exposure and, thus, overall drug efficacy. Unaffected expression would be in contrast with findings in rats, where the PXR agonist pregnenolone-16-carbonitrile upregulated the expression of both OCT1 in the liver and OCT2 in the kidneys. Second, PXR itself is known to influence glucose homeostasis, and rifampicin has been reported to induce early phase hyperglycemia possibly by enhancing glucose absorption. These parallel processes may have counteracted or dampened a pharmacodynamic interaction.

To reduce the risk of gastrointestinal adverse effects, intake of metformin and rifampicin was separated in time and patients were provided with metoclopramide after a high incidence of vomiting in the initial patients (see Supplementary Files S4 and S6). Without these preventive measures we would not have been able to complete the study. Nonetheless, (mild) gastrointestinal adverse events remained extremely common, especially among patients with TB-DM on TB treatment and metformin (57%), with more nausea and vomiting compared with patients using metformin alone. Preventive actions should be considered when treating patients with TB-DMM with metformin. Normally, it is advised to take TB drugs on an empty stomach, as food is known to limit their bioavailability, but they can also be taken with food if gastrointestinal adverse effects occur.

A limitation of this study is its sequential design; a clinically significant interaction might have led to adjustment of the metformin dose by the attending physician. Indeed, in three of the first seven subjects, the dosing interval of metformin had been changed after the first PK-GC session. Two of these patients could be measured in a third session, after switching back to the dosing schedule of their first PK-GC session. Furthermore, bias as a consequence to the fixed sequence design could have been introduced by differences in comedication and food within-patient between sampling sessions. To avoid such interference, we standardized the timing and content of meals, as well as fixed the timing of intake of comedication (if any) to assure patients were sampled under comparable conditions throughout the study. Overall, we feel that our measures have been sufficient to prevent any confounding of the metformin-rifaximin drug-drug interaction and were, most importantly, mimicking the real-life clinical situation.

Another limitation is the use of creatinine clearance as estimation for the fraction metformin eliminated from the body by glomerular filtration (eGFR). It is well known that creatinine is freely filtered by the glomerulus, however, active secretion by the proximal tubule also occurs. This possibly explains the slight change in creatinine clearance after stopping rifampicin (−4 mL/min;
The impact of this finding is minor; creatinine secretion is only a small contributor to creatinine clearance, ~10–20% of the GFR, whereas renal clearance of metformin can be up to 4–11 times as great as creatinine clearance.

As another limitation of our study, we cannot exclude an effect of isoniazid on the PKs and efficacy of metformin, as it was always co-administered with rifampicin in our study. However, to our knowledge, interactions of isoniazid with drug transporters have not been reported in literature. Similarly, we do not anticipate interactions with metformin drug transporters caused by other concomitant drugs used by patients in this study.

Finally, we would like to note here that the average percentage of extrapolation for twice-daily dosing (from $AUC_{0–8\text{ hour}}$ until $AUC_{0–\tau}$ was 21%; range: 15–26%), which is within acceptable limits for reliable extrapolation. Average percentage of extrapolation for once-daily dosing was 29% (range: 14–36%), which may be considered bordering/exceeding reliable extrapolation. However, this only concerned two patients in the entire study. The comparison for nonextrapolated total exposures ($AUC_{0–8\text{ hour}}$ i.e., with vs. without rifampicin), was similarly significant ($P = 0.001; n = 22$) as for $AUC_{0–\tau}$ (Table 1, Figure 2). The comparison for $AUC_{0–\tau}$, excluding subjects with an AUC extrapolation > 20% for at least one sampling session, showed less significance ($P = 0.083$), probably because of a loss in power ($n = 14$), because geometric means and ranges were similar to $AUC_{0–\tau}$ in the entire group (Table 1).

To summarize, we found that rifampicin exposure increases plasma metformin concentrations, without affecting its glucose lowering effect, suggesting that additional monitoring of glycemic control may not be necessary when treating patients with TB-DM. We observed a high incidence of gastrointestinal adverse effects, and, therefore, we advise patients to take metformin and rifampicin with food, and preferably separated in time. In addition, physicians could consider combining TB-DM treatment with metoclopramide as an anti-emetic when patients are experiencing gastrointestinal adverse effects.

**METHODS**

**Subjects**

Study subjects were Indonesian patients with pulmonary TB and type 2 DM that had been included in the TANDEM study (ClinicalTrials.gov: NCT02106039) at the University of Padjadjaran, Bandung, Indonesia. Patients were enrolled if they were between 18 and 65 years of age; were taking metformin, irrespective of other blood glucose control drugs; were treated with the standard Indonesian continuation phase TB regimen containing rifampicin and isoniazid thrice weekly; had a stable renal function, classified as an eGFR of > 60 mL/min; and signed an informed consent. Patients were excluded if they had an alanine aminotransferase > 3 × upper limit of normal; had inadequately controlled blood glucose concentrations, classified as an average fasting blood glucose and 2-hour postprandial glucose > 300 mg/dL; and were pregnant or lactating. The study was approved by the Independent Ethics Committee, Faculty of Medicine, University of Padjadjaran, Bandung, Indonesia. All procedures were in accordance with the Helsinki Declaration of 1975 (as revised in 1983) and Good Clinical Practice.

**Blood and urine collection during sampling sessions**

Each of the two sampling sessions consisted of 2 days, one PK day and one GC day. During the PK days, metformin and rifampicin/isoniazid were initially taken together, and with food ($n = 6$), but this resulted in a high incidence of vomiting ($n = 4$ in total; 2/6 patients during PK1 and 4/6 patients during PK2).

**Figure 4** Schematic overview of the study design in weeks from the start of tuberculosis (TB) treatment (wk 0). The TB treatment period (week 0–24) is colored dark gray. Patients were enrolled if they were continuation phase of TB treatment. Daily intake of TB drugs (for > 7 days) is indicated by dashed lines. In between sessions, there was at least a 1-month washout period, after which any induction caused by rifampicin was expected to have dissipated. $AUC$, area under the plasma concentration-time curve; $C_{\text{max}}$, maximum plasma concentration.
and another 2/6 during GC1). Subsequently, patients were provided with oral metoclopramide (10 mg) 1 hour before taking metformin, and the TB drugs were ingested 3 hours after metformin. Patients were ordered to first take their breakfast and immediately afterward their metformin. Metoclopramide was the preferred anti-emetic, as there may be a drug–drug interaction between rifampicin and domperidone and there is less experience with the use of other anti-emetics in this setting. Eight serial blood samples were drawn at 0 (predose), 1, 2, 3, 4, 5, 6, and 8 hours after witnessed ingestion of metformin with a standardized breakfast. During the PK day, cumulative urine from all subjects was collected for 8 hours in 2-hour intervals. Patients taking metformin once or twice daily received a urine container to collect their urine at home between the 8-hour until the 12–24-hour interval. Additional blood was withdrawn to assess plasma creatinine concentrations for the estimation of creatinine clearance.

On the second morning of each session, metoclopramide and metformin were taken once more, this time on an empty stomach. A total of 75 g of glucose was ingested on an empty stomach, at 2.5 hours after metformin intake when metformin concentrations are expected to be at their maximum ($C_{\text{max}}$). Glucose concentrations were measured just before and at $\frac{1}{2}$, 1, 1½, 2, 2½, and 3 hours after ingestion of glucose. See Supplementary File S6 for a schematic overview of the PK-GC sampling days.

### Study drugs

All patients received 500 or 850 mg metformin tablets (Glucophage) from PT Merck (Jakarta, Indonesia). TB drugs (rifampicin (450 mg) and isoniazid (300 mg)) were from PT Kimia Farma (Bandung, Indonesia), formulated in separate tablets. The bioequivalence of the rifampicin tablets and an international reference standard was established previously. Metoclopramide (Metolon) 10 mg was provided by PT Bernofarm (Sidoarjo, Indonesia). Drug adherence was monitored from 1 week prior to the first PK session until the end of the study, using physician assessment, pill count, self-assessment, and a patient-kept diary.

### Bio-analysis and PK evaluation

Plasma (metformin and rifampicin) and urine (metformin) concentration analyses were performed with validated ultra-performance liquid chromatography assays (Supplementary Files S1 and S2). Noncompartmental PK analyses were applied to calculate metformin and rifampicin PK parameters using Phoenix WinNonlin version 6.3 (Pharsight, Mountain View, CA; Supplementary Files S2 and S3). The net tubular secretion of metformin was calculated by subtracting creatinine clearance from metformin renal clearance (total amount excreted divided by $AUC_{\text{inf}}$; Supplementary File S3 for full details).

To assess changes in the glucose lowering effect of metformin, basal/fasting blood glucose and $C_{\text{max}}$ were determined directly from the GC. The glucose $AUC$ until 3 hours ($\text{G-AUC}_{0-3\text{ hours}}$) was calculated using the Linear Trapezoidal Linear Interpolation rule using Phoenix WinNonlin.

### Statistical analysis

Patient characteristics, metformin PKs, glucose data, and adverse events were presented descriptively for all patients included in the study. A bioequivalence approach was used to evaluate the metformin–rifampicin interaction. To conclude the absence of an interaction, or bioequivalence, the 90% CI of the GMRs of AUC-session 1: AUC-session 2 and $C_{\text{max}}$-session 1: $C_{\text{max}}$-session 2 should be between 0.80 and 1.25. Further conclusions with regard to the presence of an interaction were drawn as described by Williams et al. To assess differences between sampling sessions in patient characteristics, metformin elimination, and glucose exposures, either Related-Samples Wilcoxon Signed Rank tests or paired-samples t tests on log-transformed plasma/blood parameters were performed. All statistical analyses were performed with IBM SPSS Statistics 22 for Windows. The P values < 0.05 were judged significant.

### SUPPLEMENTARY INFORMATION

**Supplementary File S1.** Metformin concentration analyses.  
**Supplementary File S2.** Rifampicin concentration analysis and pharmacokinetic evaluation.  
**Supplementary File S3.** Pharmacokinetic evaluation of metformin.  
**Supplementary File S4.** Eligibility, inclusion, and exclusion of patients.  
**Supplementary File S5.** Blood glucose curves.  
**Supplementary File S6.** Schematic overview of the PK-GC sampling days.

### FUNDING

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### CONFLICTS OF INTEREST

The authors declared no competing interests for this work.

### AUTHOR CONTRIBUTIONS


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