

ORIGINAL ARTICLE

PKPD Modeling of Predictors for Adverse Effects and Overall Survival in Sunitinib-Treated Patients With GIST

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A modeling framework relating exposure, biomarkers (vascular endothelial growth factor (VEGF), soluble vascular endothelial growth factor receptor (sVEGFR)-2, -3, soluble stem cell factor receptor (sKIT)), and tumor growth to overall survival (OS) was extended to include adverse effects (myelosuppression, hypertension, fatigue, and hand-foot syndrome (HFS)). Longitudinal pharmacokinetic–pharmacodynamic models of sunitinib were developed based on data from 303 patients with gastrointestinal stromal tumor. Myelosuppression was characterized by a semiphysiological model and hypertension with an indirect response model. Proportional odds models with a first-order Markov model described the incidence and severity of fatigue and HFS. Relative change in sVEGFR-3 was the most effective predictor of the occurrence and severity of myelosuppression, fatigue, and HFS. Hypertension was correlated best with sunitinib exposure. Baseline tumor size, time courses of neutropenia, and relative increase of diastolic blood pressure were identified as predictors of OS. The framework has potential to be used for early monitoring of adverse effects and clinical response, thereby facilitating dose individualization to maximize OS.

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A challenge of pharmacological interventions for the treatment of cancer is to provide the most effective drug at the most efficient dosing regimen to increase the probability of successful outcomes. If objective (and early) prognostic measures of the pharmacodynamic activity for a given anticancer drug can be established, identification of the patients most likely to respond to treatment, in addition to patients with a higher risk of experiencing adverse effects, can be envisioned.

Tyrosine kinase inhibitors comprise a group of anticancer drugs where effectiveness may be improved through use of a biomarker-guided dose individualization approach. Currently, these drugs are typically dosed as a fixed regimen (dose amount and frequency) across all patients despite the high between-patient variability in pharmacokinetics (PK). Such regimens increase the potential for suboptimal exposures being achieved in particular patients.¹

Sunitinib malate (SUTENT, Pfizer) is an oral, small-molecule, multitargeted tyrosine kinase inhibitor currently approved in, e.g., the United States and Europe for the treatment of metastatic renal cell carcinoma, imatinib-resistant gastrointestinal stromal tumor (GIST), and pancreatic neuroendocrine tumor.

In a companion article, we present an overarching modeling framework that links drug exposure, soluble biomarkers, tumor growth dynamics, and overall survival (OS) with the aim of identifying robust predictors of clinical response.² A soluble form of the vascular endothelial growth factor (VEGF) receptor, soluble vascular endothelial growth factor receptor (sVEGFR)-3, was identified as the most effective predictor of OS in sunitinib-treated GIST patients. sVEGFR-3 was found to be a more effective predictor of OS than the relative change in tumor size at week 7, which was previously proposed for other cytotoxic treatments.³ An additional question

to address was whether sVEGFR-3 or any of the other previously investigated candidate biomarkers (VEGF, sVEGFR-2, or sKIT) were predictive of adverse effects, which could assist in determining an individualized dose. Additionally, the adverse effects themselves may be “useful” alternative early indicators of pharmacodynamic activity as they could be more practical to use in the clinical setting. Hypertension is a commonly observed adverse effect for VEGF(R) inhibitors⁴ and has been reported as a biomarker of treatment efficacy in metastatic renal cell carcinoma patients treated with sunitinib.^{5,6} A positive correlation has also been identified between the severity of neutropenia and both time to progression and OS⁷ in sunitinib-treated GIST.

Relationships between drug exposure, candidate biomarkers (VEGF, sVEGFR-2, sVEGFR-3, and sKIT), and the sunitinib-related adverse effects—fatigue, hand-foot syndrome (HFS), neutropenia, and hypertension—were assessed in sunitinib-treated GIST using nonlinear mixed-effects pharmacokinetic–pharmacodynamic modeling. Adverse events were also evaluated as potential predictors of OS.

RESULTS

The analysis included data pooled from four clinical trials^{8–11} in phases I–III, which comprised 303 patients with imatinib-resistant malignant GIST treated with sunitinib and/or placebo (**Table 1**). Available data were biomarker candidates (VEGF, sVEGFR-2, sVEGFR-3, and sKIT), OS, and the most commonly reported treatment-related adverse effects, fatigue, HFS, neutropenia, and hypertension (diastolic blood pressure, dBP).

Longitudinal information on dose, sunitinib daily area under the concentration–time curve (AUC), and biomarkers

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Table 1 Data summary of the analyzed studies

Study number	Study 1004	Study 1047	Study 1045	Study 013
Reference	Demetri <i>et al.</i> ¹²	George <i>et al.</i> ¹³	Shiara <i>et al.</i> ¹⁴	Maki <i>et al.</i> ¹⁵
N	202 active 47 placebo	13	36	52
Study design	Double-blind, randomized, placebo-controlled, phase III	Nonrandomized, evaluating continuous treatment regimen, phase II	Nonrandomized, dose-escalating study in Japanese patients, phase I/II	Nonrandomized, dose-escalating study, phase I/II
Dosing schedule (weeks on/weeks off)	0, 50 mg q.d. 6-week cycles (4/2)	37.5 mg q.d. 4-week cycles, continuous treatment	25, 50, 75 mg q.d. 6-week cycles (4/2)	25, 50, 75 mg q.d. 3-week cycles (2/1) 4-week cycles (2/2) 6-week cycles (4/2)
Fatigue (%) ^a	0: 30 1: 33 2: 25 3: 11 4: 1	0: 54 1: 23 2: 23 3: 0 4: 0	0: 31 1: 39 2: 19 3: 8.3 4: 2.7	0: 11 1: 48 2: 33 3: 5.7 4: 1.9
Hand-foot syndrome (%) ^a	0: 83 1: 5.0 2: 6.9 3: 5.4 4: 0	0: 100 1: 0 2: 0 3: 0 4: 0	0: 14 1: 20 2: 34 3: 31 4: 0	0: NA 1: NA 2: NA 3: NA 4: NA
Absolute neutrophil counts ($\cdot 10^9/l$) ^b , median (range)	3.1 (0.080–20)	1.8 (0.010–7.5)	2.1 (0.28–12)	2.6 (0.16–15)
Diastolic blood pressure (mmHg) ^b , median (range)	80 (20–120)	78 (40–120)	79 (40–120)	80 (50–130)
Survival (weeks), median (range)	61 (4–226)	31 (81–15)	37 (27–48)	39 (4–96)

NA, not available; q.d., once daily.

^aHighest observed severity score within an individual (%). ^bObserved median (range) response during the treatment period.

were evaluated as predictors for adverse effects. The time courses of the biomarker [BM(t)] concentrations were characterized by indirect response models, as described in the study by Hansson *et al.*² The predicted value of the different biomarkers for adverse effects was assessed by investigating how descriptive measures of the individual model-predicted biomarker time courses were, alone or in addition to drug exposure. The model-predicted baselines (BM_0), absolute and relative change from baseline (BM_{REL}), or the complete biomarker time courses were evaluated as predictors.

Myelosuppression model

A semiphysiological myelosuppression model¹² using Box-Cox-transformed data adequately described the extent and time course of the change in absolute neutrophil count (ANC) following sunitinib treatment. A more pronounced reduction in ANC was observed during the first treatment cycle, with partial recovery to baseline levels during the subsequent off-treatment period. A smaller decline in ANC was subsequently observed during the second treatment cycle and thereafter. Placebo-treated patients did not show any systematic alterations in ANC levels.

All of the investigated biomarkers were statistically significantly correlated with the changes in ANC levels when assessed at a fixed time point (landmark). However, the longitudinal model-predicted relative change in sVEGFR-3 from baseline ($sVEGFR-3_{REL}$) was the better descriptor of the myelosuppression time course (change in the objective function value, $\Delta OFV = 170.8$, compared with AUC). No further improvement in the description of the data was observed when any of the other biomarkers or sunitinib AUC was added to the univariate model.

An E_{max} function most appropriately characterized the biomarker-ANC relationship. A separate baseline parameter (ANC_0) was estimated to account for lower ANC levels in Study 45, which was conducted in Japanese patients. The final model included interindividual variability in ANC_0 , mean transit time (MTT) through the nonsensitive compartments, E_{max} and EC_{50} (Table 2), with a correlation between ANC_0 and E_{max} of 90%. For a typical patient receiving a daily 50-mg sunitinib dose (4/2 schedule) and an ANC_0 of 4.94 ($\cdot 10^9/l$), the model predicted a 62% decrease in ANC corresponding to a nadir of 1.9 ($\cdot 10^9/l$).

The predictive performance of the final myelosuppression model, as illustrated by a prediction-corrected visual

Table 2 Final model parameter estimates (relative SE, %)

Parameter	Estimate	RSE, %	IIV CV, %	RSE, %
Myelosuppression model				
ANC ₀ (-10 ⁹ /l)	4.94	2.8	42	5.6
ANC ₀ : Study 45 (-10 ⁹ /l)	3.69	6.9	42	5.6
MTT (hours)	248	3.6	17	19
ANC E _{max}	0.520	9.1	13	36
ANC EC ₅₀ (pg-hour/l)	0.552	17	46	16
γ	0.362	7.4	—	—
Residual error ^a	0.406	4.3	—	—
Blood pressure model				
dBP ₀ (mmHg)	71.8	1.0	12	6.7
dBP ₀ placebo (mmHg)	77.6	1.6	12	6.7
MRT (= 1/k _{out}) (hours)	361	17	83	12
dBP _{slope} (l/mg-hour)	0.119	9.4	65	11
Residual error (mmHg)	6.24	16	—	—
Residual error (%)	6.97	24	—	—
Survival model				
λ (/week)	0.0079	55	—	—
α	1.15	9.1	—	—
α _{cens}	1.27	44	—	—
β ₁ ANC (l·10 ⁹)	4.76	31	—	—
β ₂ dBPREL	-1.29	27	—	—
β ₃ Tumor base (/mm)	-0.00172	46	—	—
λ _{cens} (/week)	0.0019	6.6	—	—

ANC₀, baseline absolute neutrophil count; CV, coefficient of variation; dBP₀, baseline diastolic blood pressure (dBP); dBP_{slope}, parameter relating drug exposure to the change in dBP; IIV, interindividual variability; MRT, mean residence time; MTT, mean transit time; RSE, relative standard error; α, shape factor in Weibull probability density function; α_{cens}, shape factor in Weibull probability density function for censoring; β₁ANC, parameter relating ANC(t) to the hazard; β₂dBPREL, parameter relating the relative change in dBP to hazard; β₃Tumor base, parameter relating observed baseline tumor size to the hazard; γ, feedback factor; λ, scale factor in the Weibull probability density function; λ_{cens}, scale factor in the Weibull probability density function for censoring.
^aResidual error on Box-Cox-transformed scale.

predictive check (VPC; **Figure 1**), shows a good description of the ANC data.

Blood pressure model

An indirect response model with stimulation of the production rate (K_{in}), as proposed by Keizer *et al.* for another antiangiogenic drug,¹³ described the observed elevated dBP during sunitinib treatment periods and the return to near baseline during off-treatment periods (Eq. 1). No increase in dBP could be identified for placebo patients. However, this group had a significantly higher baseline dBP (dBP₀) when estimated separately.

$$\frac{ddBP}{dt} = K_{in} \cdot (1 + dBP_{Drug\ effect} \cdot AUC) - K_{out} \cdot dBP(t) \quad (1)$$

$$K_{in} = dBP_0 \cdot K_{out}$$

Sunitinib AUC was linked to the production rate (K_{in}) by a linear slope factor (dBP_{Drug effect}). None of the evaluated biomarkers were found to be significantly related to the dBP time course. Interindividual variability was estimated for dBP₀, the mean residence time MRT (=1/k_{out}), and dBP_{Drug effect} and a combined additive and proportional residual error model was used. A correlation between dBP₀ and dBP_{Drug effect} (65%) was estimated. The final model predicted a drug-induced increase in dBP by 10 mmHg for the typical patient with a baseline dBP of 71.8 mmHg treated with 50 mg sunitinib receiving a 4/2 schedule.

The relative standard errors (RSEs, %) for the estimated parameters were low to intermediate, showing an adequate precision in the estimates (**Table 2**). The prediction-corrected VPC shows a good predictive performance of the final blood pressure model (**Figure 2**).

Fatigue and HFS models

The data were treated as ordered categorical (grades 0, 1, 2, ≥ 3), and an extension of the proportional odds model was used to describe the probability and severity of fatigue and HFS over time.^{14,15} The extension included a first-order Markov model to condition the probability of transition between different severities based on the preceding one, thereby accounting for that the severity of the adverse effects is not independent from one time point to another. Logit transformations were used to constrain the estimated probabilities to values between 0 and 1, and the function describing the probability of transition from grade *a* to grade *b* for the *i*th patient at the *j*th observation was given the structure shown in Eq. 2.

$$\text{Logit}(P_{ijb|a}) = \ln \frac{P_{ijb|a}}{1 - P_{ijb|a}} = f_{b|a} + \eta_i \quad (2)$$

$$f_{b|a} = B_{b|a} + g(x_i)$$

where $f_{b|a}$ is a function of baseline transition probabilities ($B_{b|a}$), and $g(x_i)$ is a linear function on the logit scale relating explanatory factors, such as time, drug exposure (dose and AUC), and absolute/relative changes in biomarker concentrations over time, to the probability of developing fatigue and HFS. The interindividual random effect for patient *i* (η_i) was assumed to be normally distributed with a mean of zero and a variance of ω^2 .

All of the biomarkers were significantly related to the probability and severity of fatigue and HFS. However, the relative change over time for sVEGFR-3 (sVEGFR-3_{REL}) showed the most profound relationship (Δ OFV = -103 for fatigue, Δ OFV = -159 for HFS, compared with AUC), and no further improvement in the description of the data was observed when AUC or the other biomarkers were also incorporated. No significant trend over time was identified. Increasing sVEGFR-3_{REL}, i.e., a more pronounced reduction in sVEGFR-3, was associated with increased probability and severity of fatigue and HFS. Incorporation of an effect compartment into the model [$k_{e0} = 0.424/\text{hour}$ (fatigue) and

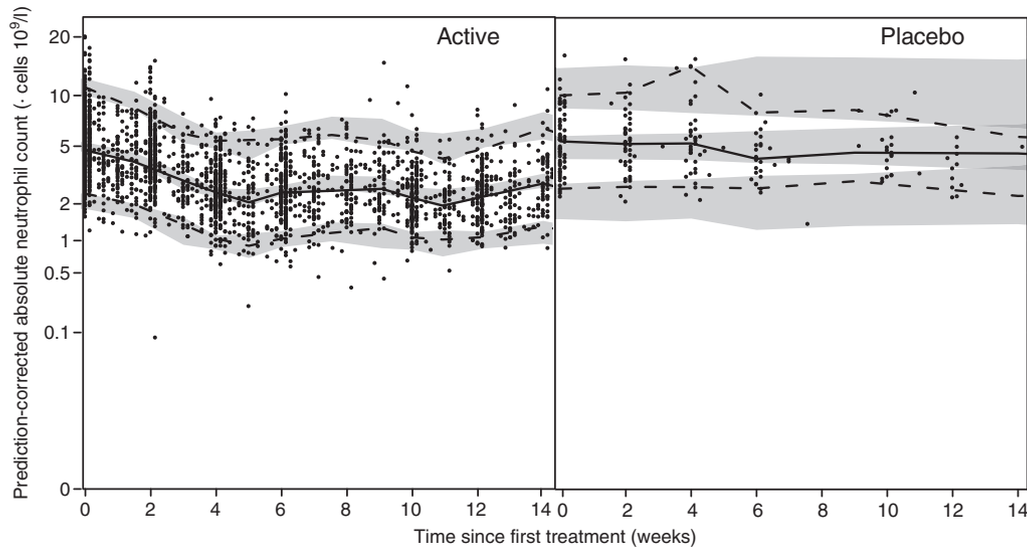


Figure 1 Prediction-corrected visual predictive checks of the final semimechanistic myelosuppression model for sunitinib-treated (left) and placebo-treated (right) patients using soluble vascular endothelial growth factor receptor (sVEGFR)-3_{REL} as descriptor. Solid circles (●) represent observed absolute neutrophil counts, solid lines (—) represent the median of the observed data, and dashed lines (---) represent the 5th and 95th percentiles of the observed data. Shaded areas are the 95% confidence intervals based on the simulated data ($n = 500$) for the corresponding percentiles.

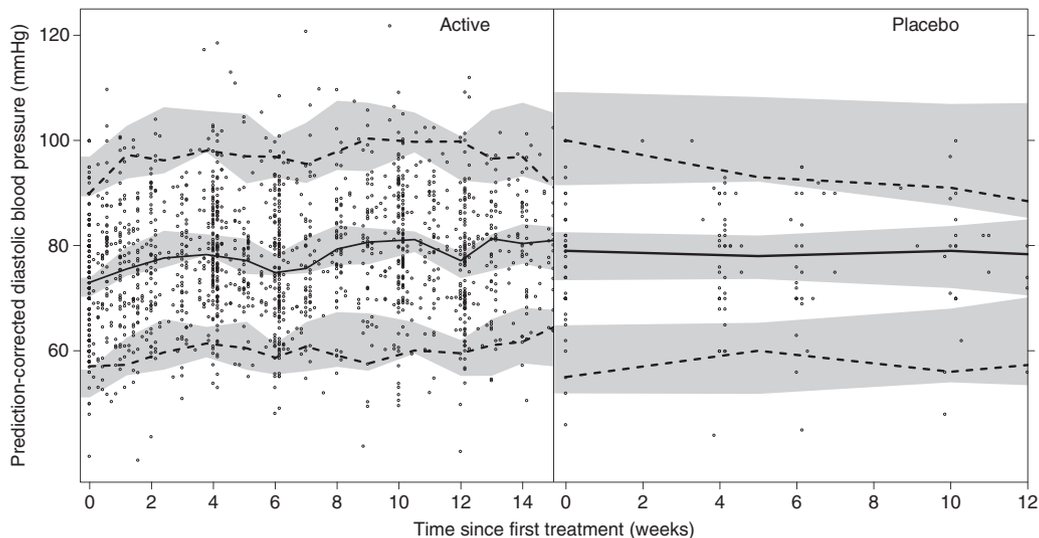


Figure 2 Illustration of the predictive performance of the final blood pressure model for sunitinib-treated (left) and placebo-treated (right) patients by prediction-corrected visual predictive checks. Solid circles (●) represent observed diastolic blood pressure, solid lines (—) correspond to the median of the observed blood pressure data, and dashed lines (---) correspond to the 5th and 95th percentiles of the observed data. The shaded areas are the 95% confidence intervals based on the simulated data ($n = 500$) for the corresponding percentiles.

0.347/hour (HFS)] significantly improved both the fatigue and HFS models (Table 3).

The parameters were estimated with acceptable precision and are reported on the logit scale in Table 3. The simulation-based model evaluation described well the time course for the probability and severity of fatigue and HFS (Table 3, Figure 3). Furthermore, the simulated numbers of transitions between different severity grades were consistent with the observed values (results not shown).

OS model

A Weibull model described the underlying baseline hazard for the observed survival data (λ ; hazard coefficient, α ;

shape factor; Eq. 3, Table 2). The time course of neutropenia [ANC(t)] was the most significant predictor of OS ($\Delta\text{OFV} = -42.6$). Additionally, $\text{dBP}_{\text{REL}}(t)$ was significantly related to OS ($\Delta\text{OFV} = -25.4$), whereas fatigue and HFS were not related. A more pronounced decrease in ANC over time and/or a larger relative increase in dBP decreased the hazard risk of death.

To be able to compare our model with the previously reported survival model from that in our companion article,² which included sVEGFR-3_{REL} and tumor size at start of treatment as predictors, these two predictors were also included and reevaluated for significance, in addition to the presence of the adverse events. The model including ANC and dBP

Table 3 Time course for the probability and severity of HFS and fatigue

Parameter	HFS model		Fatigue model	
	Estimate	RSE, %	Estimate	RSE, %
$B_{1 0}$	-10.4	11	-5.85	3.0
$B_{2 0}$	-0.974	13	-1.14	7.8
$B_{3 0}$	-1.59	19	-1.60	14
Slope $_{x 0}$	-8.00	14	-1.93	22
$\omega_{x 0}$	3.07	67	1.06	20
$B_{1 1}$	2.29	14	2.63	10
$B_{2 1}$	-9.53	5.0	-10.7	3.1
$B_{3 1}$	-1.33	24	-1.77	22
Slope $_{x 1}$	-6.00	18	-4.62	17
$\omega_{x 1}$	0.902	54	1.25	30
$B_{1 2}$	3.04	15	2.86	12
$B_{2 2}$	-0.747	14	-0.427	20
$B_{3 2}$	-9.09	5.1	-11.6	5.0
Slope $_{x 2}$	-3.23	43	-4.64	22
$\omega_{x 2}$	0.270	118	1.30	24
$B_{1 >3}$	3.4	21	3.06	23
$B_{2 >3}$	-1.65	23	-0.090	115
$B_{3 >3}$	-0.453	37	-0.636	33
Slope $_{x >3}$	-4.75	32	-3.32	51
$\omega_{x >3}$	NE	NE	0.841	71

$B_{0|a}$ intercept for the probability of transition from grade *a* to grade *b*; HFS, hand-foot syndrome; NE, not estimated; RSE, relative standard error; Slope $_{x|b}$ parameter relating soluble vascular endothelial growth factor receptor (sVEGFR)-3 to the probability of the severity score *x* given the previous score *b*; $\omega_{x|b}$ interindividual random variability.

resulted in a similar OFV as sVEGFR-3 $_{REL}$ and could thereby be an alternative to the model developed earlier.² A summary table of differences in OFV for the significant predictors is provided in **Supplementary Table S1** online. The final model using adverse effects as predictors was parameterized as in Eq. 3.

$$h(t) = \lambda \cdot \alpha \cdot t^{\alpha-1} \cdot e^{(\beta_1 \cdot (ANC(t)C-5)/5 + \beta_2 \cdot dBP_{REL(t)} + \beta_3 \cdot \text{Tumor base})} \quad (3)$$

The predictive properties of the survival model are adequate as shown in the VPC (**Figure 4**). The Kaplan–Meier plot of the observed survival data is well within the 95% confidence interval calculated from simulations of 200 data sets. The median number of simulated events ($n = 151$, range: 126–176) was in accordance with the number of observed events ($n = 163$).

Evaluation of how well the model can predict survival using the model-predicted time course of ANC and dBP based on data from only the first treatment cycle (<6 weeks, and an average of five (ANC) and four (dBP) measurements), compared with the use of all data, resulted in a similar model fit ($\Delta\text{OFV} = 9.5$). This indicated that a small number of ANCs and blood pressure measurements may be needed during the first treatment cycle to predict survival given the model. The predictive properties of the model when using data from only

the first treatment course are illustrated by the Kaplan–Meier plot in **Figure 4**.

DISCUSSION

This study described the association between sunitinib exposure, selected biomarkers, myelosuppression, hypertension, fatigue, and HFS through the development of longitudinal pharmacokinetic–pharmacodynamic models, and relationships were added to the modeling framework described for sunitinib in GIST (**Figure 5**). Predictors of the toxicity dynamics were assessed, and the soluble protein sVEGFR-3 was identified as being most predictive of neutropenia, fatigue, and HFS. The link between the longitudinal adverse effects data and OS indicated that dBP, myelosuppression, and baseline tumor size were most predictive. The relationship between AUC and adverse effects has previously partially been characterized in this population without considering the time aspect.¹⁶ In this study, the complete time course of the adverse effects were characterized to evaluate the relationships.¹⁷

A semimechanistic model¹² was successfully applied to describe sunitinib-induced myelosuppression. sVEGFR-3 was statistically the best predictor of the neutropenic time course. All of the investigated biomarkers were, however, significantly related to the time course of myelosuppression when evaluated one by one, reflecting the inhibitory effect of sunitinib on KIT (stem cell factor receptor) and the role of VEGF in myelopoiesis.⁴ These biomarkers may also have provided more longitudinal information on drug exposure because they were monitored for a longer period than drug concentrations. A longer MTT in the myelosuppression model (248 hours) was estimated for sunitinib than that typically reported for traditional intravenously administered cytotoxic drugs in short treatment schedules.^{23,31} Van Kesteren *et al.* reported on longer MTT for prolonged schedules, which may be the result of a decreased effect of the endogenous growth factors following longer periods of low number of circulating cells.¹⁸

The elevated dBP as a result of sunitinib treatment was described by an indirect response model with a linear relationship between AUC and the increased production rate (K_{in}). The mechanism mediating the elevated blood pressure in antiangiogenic treatment is not clear. One hypothesis is that decreased production of nitric oxide and vascular rarefaction due to VEGF blockade induces vasoconstriction and hypertension.⁴ The developed model has no direct mechanistic link between exposure and appearance of hypertension, which limits extrapolation outside the range of dosing schedules studied. A contrasting result to the other investigated adverse effects was that none of the evaluated biomarkers were significantly related to the model-predicted time course of dBP.

The developed first-order mixed-effects Markov models characterized the dynamics of longitudinal fatigue and HFS data. The model provides an alternative approach to traditional analysis of toxicity data, which often only reports the highest severity during the study for a patient and thereby discards the evolution of toxicity over time. An increased sunitinib exposure or sVEGFR-3 response was related to a higher

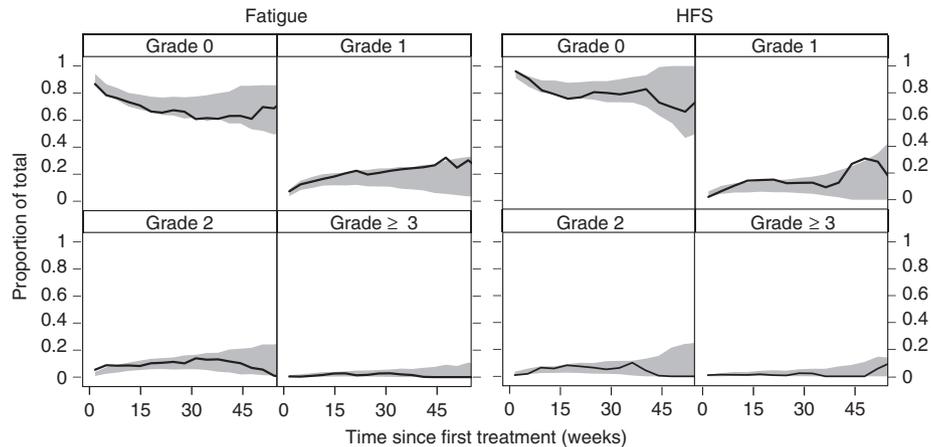


Figure 3 Visual predictive check of the final model for the time course of the probability and severity of fatigue (left) and hand–foot syndrome (HFS) (right) for actively treated patients stratified by severity grade. The solid lines (—) represent the observed time course of each severity grade, and the shaded areas are the 95% confidence intervals generated from simulations.

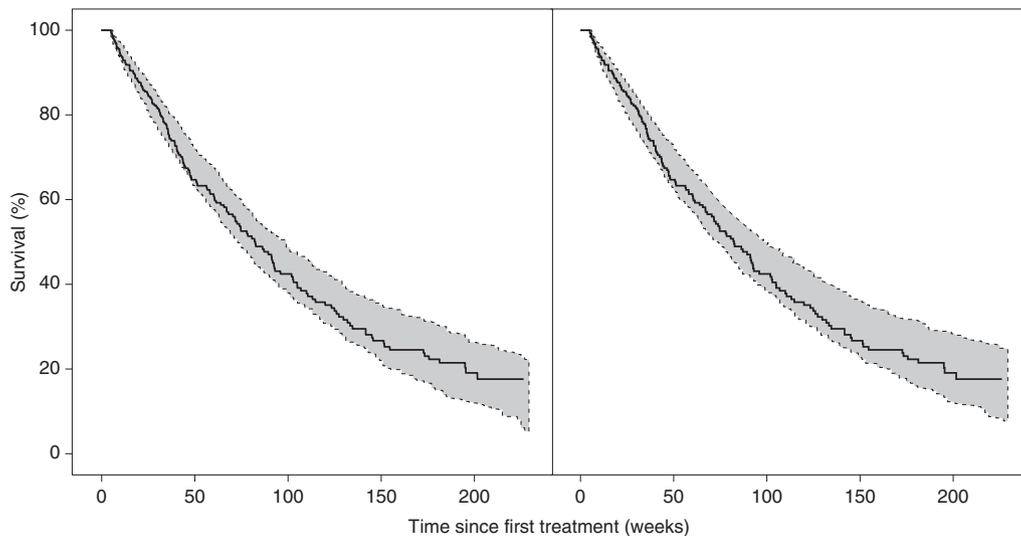


Figure 4 Kaplan–Meier plot of the observed survival data (solid line) and the 95% confidence interval (shaded area) based on the simulated data ($n = 200$) for the final survival model (left). A Weibull model ($\lambda = 0.0019$, $\alpha = 1.3$) was applied in the simulations to describe censoring. The right panel illustrates the predictive properties of the model when only using data from the first 6 weeks of treatment.

risk and severity of adverse effects. The proposed models with sVEGFR-3 as a descriptor are empirical and provide limited contribution to the understanding of the mechanism for development of HFS and fatigue but could be of prognostic value.

The relative change in sVEGFR-3 over time is predictive of OS (ref 2) and the adverse event dynamics for myelosuppression, fatigue, and HFS. Therefore, monitoring sVEGFR-3 has the potential to identify the patients at highest risk of toxicity and to enhance dose optimization. Efficient dose individualization could minimize the occurrence of severe adverse events and improve the treatment efficacy. A maintained dose intensity is of importance because higher exposures have been shown to be associated with longer OS and time to progression in a previously reported exposure–response analysis.¹⁶

The relative change in dBP, myelosuppression time course, and tumor size at start of treatment were predictive

of OS. The model predicted that patients with a greater relative change in blood pressure and ANC, together with a smaller tumor size at baseline, displayed the longest OS. Cutoff values of blood pressure and neutropenia have previously been identified as predictors in sunitinib treatment using traditional statistical analysis. ANC $< 1.5 \times 10^9/l$ or dBP > 90 mmHg at any time during treatment were associated with longer OS.^{5–7}

ANC and dBP measurements from the first treatment cycle can predict OS. The developed survival model may guide inpatient dose escalation based on dBP and neutropenia and explore the effectiveness of alternate dosing strategies. The potential impact on the augmented incidence and severity of HFS and fatigue due to the increased dose adjustments can be considered. A future simulation study will evaluate a dose individualization approach to ultimately optimize the use of sunitinib in GIST. These factors will require confirmation in larger prospective trials. In

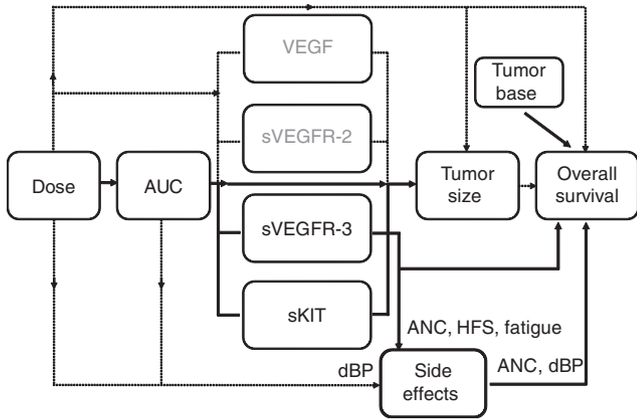


Figure 5 Relationships evaluated in the framework for sunitinib in gastrointestinal stromal tumor. Solid lines indicate relationships included in the final models, and dashed lines indicate relationships investigated but not included in the final models. In contrast to the other adverse effects investigated, blood pressure was, however, better predicted by sunitinib daily area under the concentration–time curve (AUC) than soluble vascular endothelial growth factor receptor (sVEGFR)-3. Tumor size before treatment initiation (baseline) was a predictor in the final model of overall survival, whereas the relative change in sVEGFR-3 over time was a better predictor than other evaluated metrics of tumor size after start of treatment.² The current analysis showed that absolute neutrophil count (ANC) and diastolic blood pressure (dBP) are, in combination, as good predictors of overall survival as is sVEGFR-3. HFS, hand-foot syndrome; sKIT, soluble stem cell factor receptor.

conclusion, this analysis proposed the relative change in sVEGFR-3 over time as a predictor of the occurrence and severity of myelosuppression, fatigue, and HFS following sunitinib treatment. Furthermore, sunitinib-induced hypertension and neutropenia were identified as predictors of OS in GIST. The developed framework, including adverse effects as biomarkers, has a potential to be used for early monitoring of response, thereby facilitating effective dose individualization.

METHODS

Patients and study design. The analyzed data were from four clinical trials^{8–11} in phases I–III, which comprised patients with imatinib-resistant malignant GIST treated with sunitinib. Only patients with biomarker and survival data reported were included in the analysis, totaling 303 patients. Sunitinib was administered in one of four different treatment schedules, including the 4/2, 2/2, and 2/1 (weeks on/weeks off) schedule, and continuous treatment, with doses ranging from 0 to 75 mg orally once daily (Table 1). Patients randomized to receive placebo treatment ($n = 47$) in the placebo-controlled trial (study 1004) were offered sunitinib on disease progression as defined by the Response Evaluation Criteria in Solid Tumors (RECIST).¹⁹ The studies were approved by local ethics committees and were performed in accordance with the Declaration of Helsinki. Written informed consent was obtained from all patients.

Data analysis. This population PK/PD analysis was performed sequentially using the nonlinear mixed-effects modeling

approach. The first-order conditional estimation method with interaction (FOCEI) and the Laplacian estimation method implemented in the software NONMEM (version 7.2, gFortran version 4.5.0, Gnu compiler collection; **Supplementary Data**) were used for parameter estimation. The R-based software Xpose (version 4)²⁰ was used for model diagnostics and graphical visualization of the results, and the PsN toolkit (version 3)^{21,22} was used for the postprocessing of results.

Model selection was based on assessment of graphical diagnostics and comparison of the OFV, provided by NONMEM, in the log-likelihood ratio test. The difference in OFV for two nested models is proportional to minus twice the log likelihood of the data and is approximately χ^2 distributed. A significance level of $P < 0.01$ was used in this analysis, which corresponds to a decrease in OFV of at least 6.63 for the addition of one extra parameter. Evaluation of model robustness was based on relative standard errors (RSEs, %) of the model parameter estimates determined by nonparametric bootstrapping ($n = 200$). Prediction-corrected VPCs²³ were also assessed to judge the predictive performance of the developed models.

Pharmacokinetics. Dose or daily area under the concentration–time curve (AUC) calculated as $\text{Dose}/(\text{CL}/F)$, with $\text{AUC} = 0$ during off-treatment periods, was used as the exposure measure for sunitinib (no data for the equipotent metabolite SU1266 were available). Total oral plasma clearance (CL/F) was described by the individual parameter estimates or population values [(when no PK data were available ($n = 57$)) obtained from a population PK model.²⁴

Biomarker models. The time courses of the biomarkers were characterized by individual parameter estimates obtained from previously developed biomarker models.² The models described the changes in biomarker concentrations [BM(t)] through indirect response models assuming a decreased production of the zero-order rate constant (K_{in}) for sVEGFR-2, sVEGFR-3, and sKIT and an inhibited degradation of VEGF (k_{out}). A linear disease progression model characterized the increase of VEGF and sKIT over time. Data on sVEGFR-3 were not available for two of the studies ($n = 69$), but the high correlation between sVEGFR-2, sVEGFR-3, and VEGF to sVEGFR-3 enabled derivation of information on sVEGFR-3 in individuals with missing data.²

Myelosuppression model. The time course of sunitinib-related changes in ANC was described by a semimechanistic model for myelosuppression.¹² The model is composed of a compartment representing drug-sensitive proliferating progenitor cells in the bone marrow, a compartment of systemic circulating neutrophils, and a link between them through three transit compartments reflecting cell maturation. The model also includes a feedback function mimicking the effect of endogenous growth factors, e.g., granulocyte colony stimulating factor (G-CSF), which increases the proliferation rate when neutrophil levels in the blood are low. The half-life of circulating neutrophils in blood was fixed to the literature value of 7 hours²⁵ to enhance the physiological interpretation of the model. Estimated system-specific parameters were

ANC_0 , MTT, and the feedback factor (γ). Furthermore, the drug effect—which is assumed to act by reducing the proliferation rate and inducing cell loss—was estimated. Herein, linear, E_{max} , and sigmoid E_{max} models were evaluated to link drug exposure (dose and AUC) and biomarkers (baseline, absolute, and relative change) to cell death.

Modeling was performed on Box–Cox-transformed neutrophil data ($ANC_{transformed} = (ANC^\lambda - 1)/\lambda$), with $\lambda = 0.2$, as this previously has been shown to result in approximately normally distributed residuals.^{26,27} Interindividual variability was assumed to be log-normally distributed with a mean of zero and a variance of ω^2 and was evaluated for ANC_0 , MTT, and the drug effect parameters. Residual variability was described by an additive (on Box–Cox scale) error model. The predictive performance of the myelosuppression model was evaluated by prediction-corrected VPCs (500 simulations).²³

Blood pressure model. The sunitinib-induced hypertension was reported in terms of increase in dBP. The actual times of the day for blood pressure measurements were not available and were therefore assumed to occur in the morning for all observations. The increase in dBP following sunitinib treatment was described using (i) an indirect response model with stimulation of K_{in} or (ii) the alternative model with inhibition of k_{out} . dBP_0 and k_{out} (reported as $MRT = 1/k_{out}$) were estimated, and K_{in} was derived as $dBP_0 \times k_{out}$. Linear, E_{max} , and sigmoid E_{max} drug effect relationships to sunitinib exposure (dose and AUC) and biomarkers (baseline, absolute, and relative changes over time) were evaluated. Interindividual variability was assumed to be log-normally distributed with a mean of zero and a variance of ω^2 . Additive, proportional, or combined additive and proportional residual error models were evaluated.

The predictive properties of the final blood pressure model was evaluated using prediction-corrected VPCs ($n = 500$).²³

Fatigue and HFS model. Fatigue and HFS were assessed daily throughout treatment. They were reported according to the National Cancer Institute common toxicity criteria (version 3.0) as different grades, where grade 0 corresponds to no adverse event and grade 4 refers to a life-threatening event. However, grade 4 was only reported in a few patients (for fatigue: 1%; and for HFS: 0%), and these occurrences were consequently grouped with grade 3 into a single category.

Models for ordered categorical data (grades 0, 1, 2, and ≥ 3) with a first-order Markov model were applied. Transitions between all the different severity grades were considered in the analysis, totaling 16 different possible transitions. The sum of the associated probabilities for each grade ($P_{x|0}$, $P_{x|1}$, $P_{x|2}$, and $P_{x|\geq 3}$) is one, and therefore three probabilities for each grade were directly estimated and the fourth probability ($P_{0|0}$, $P_{1|0}$, $P_{2|0}$, and $P_{3|0}$) was expressed as one minus the sum of the associated probabilities. Linear and nonlinear models for the explanatory factors were assessed, and the addition of an effect compartment to account for a delay in the drug effect was tested.²⁸

The fatigue and HFS models were evaluated by categorical VPCs, where 95% confidence intervals were generated from 500 simulations and overlaid with the observed time course of fatigue and HFS stratified by severity. In addition, predictive checks were created by comparing the simulated and observed numbers of transitions between different toxicity grades.

OS model. A parametric survival (time-to-event) model was developed to evaluate whether any of the studied treatment-related adverse effects were predictive of OS. The underlying distribution of the observed survival data was evaluated by exponential, Weibull, log-logistic, extreme value, and Gompertz probability density functions. The individual predicted time courses, using AUC as predictor, for neutropenia and hypertension were extrapolated based on the developed models assuming dosing and schedule according to protocol until time of censoring/death. For neutropenia and hypertension baseline levels, absolute time course and absolute and relative change from baseline over time were evaluated as predictors of OS. For fatigue and HFS, the observed severity scores (last observation carried forward) were evaluated as predictors of OS by including each observed score as a predictor. To be able to compare our model with the previously reported survival model, which included sVEGFR-3_{REL} and tumor size at start of treatment as predictors,² these two predictors were also included and reevaluated for significance, in addition to the presence of the adverse events.

The predictive properties of the survival model were assessed by Kaplan–Meier plots of the observed survival data compared with a 95% confidence interval generated from simulations of 200 replicates of the data sets. Censoring because of, e.g., a short follow-up period was described by a Weibull model, which was applied in the simulations.

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Conflict of interest. G.M., M.A.A., and P.A.M. are employees of Pfizer Ltd. At the time the work was carried out, G.M. was supported by a research grant from Pfizer Ltd. and J.F. was an employee at Pfizer Ltd. M.O.K. and L.E.F. have acted as paid consultants to Pfizer Ltd. As Deputy Editor-in-Chief of *CPT: Pharmacometrics & Systems Pharmacology*, L.E.F. was not involved in the review or decision process for this article. E.K.H. declared no conflict of interest.

Study Highlights

WHAT IS THE CURRENT KNOWLEDGE ON THE TOPIC?

- ✓ Sunitinib-induced change in the angiogenic biomarker soluble vascular endothelial growth factor receptor (sVEGFR)-3 was a predictor of overall survival (OS) in a modeling framework evaluating exposure–biomarker–tumor size–OS relationships for gastrointestinal stromal tumor (GIST).

WHAT QUESTION DID THIS STUDY ADDRESS?

- ✓ In a population PKPD analysis, exposure and the changes in angiogenic biomarkers over time were investigated as predictors for sunitinib-induced adverse effects. The time courses of the adverse effects were tested for their predictive capacities for OS.

WHAT THIS STUDY ADDS TO OUR KNOWLEDGE

- ✓ The framework was expanded to integrate adverse effects. The relative change in sVEGFR-3 over time was a predictor of myelosuppression, fatigue, and hand–foot syndrome. The relative increase in diastolic blood pressure (dBP_{REL}) was positively correlated with sunitinib AUC. Neutrophil count and dBP_{REL} were identified to predict OS equally well as sVEGFR-3.

HOW THIS MIGHT CHANGE CLINICAL PHARMACOLOGY AND THERAPEUTICS

- ✓ Models of four common adverse effects of sunitinib were added to the developed framework on integrated PKPD models. Neutrophil counts and blood pressure measurements show promise to be applied in feedback individualization to increase OS in GIST.

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