Posaconazole therapeutic drug monitoring in clinical practice and longitudinal analysis of the effect of routine laboratory measurements on posaconazole concentrations

Running title: Posaconzole TDM in clinical practice

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**Author contributions:** LFRS, DT, TSW and JWA designed the study, AGM, AV and MB collected the data, AGM and ERH performed the statistical analysis, AGM, AV, DT, ERH, TSW and JWA led the writing.

The authors confirm that the ethical policies of the journal, as noted on the journal's author guidelines page, have been adhered to and the appropriate ethical review committee approval has been received.

**Abstract**

**Background:**
Posaconazole is indicated for prophylaxis and treatment of invasive aspergillosis. Therapeutic drug monitoring (TDM) of posaconazole is used to optimize drug exposure. The aim of this study was to analyse and describe the TDM practices and exposure of posaconazole tablets.

**Materials/methods:**
Patients who received posaconazole for treatment or prophylaxis of fungal infections were included in the study. The following therapeutic window was defined: if concentration was low (<0.7mg/L for prophylaxis or <1.5 mg/L for treatment) or high (>3.75mg/L) the hospital pharmacist provided the physician with dosage advice, which implementation to patient care was analysed. A longitudinal analysis was performed to analyse if different confounding variables had an effect on posaconazole concentrations.
**Results:**
Forty-seven patients were enrolled resulting in 217 posaconazole trough concentrations. A median of 3 (IQR 1-7) samples were measured per patient. The median concentration was 1.7 mg/L (IQR 0.8-2.7) for prophylaxis and 1.76 mg/L (IQR 1.3-2.3) for treatment. Overall 78 posaconazole concentrations were out of the therapeutic window. For 45 (54%) of these concentrations a dosage change was recommended. In 54 (25%) of all measured posaconazole concentrations resulted in recommendation for dosage alteration. In the longitudinal analysis the laboratory markers and patient baseline variables did not have an effect on posaconazole concentrations.

**Conclusions:**
Adequate posaconazole exposure was shown in in 64% (affected 28 patients) of the measured concentrations. TDM practice of posaconazole can be improved by increasing the implementation rate of dose recommendation by a multidisciplinary antifungal stewardship team.

**Introduction**
Invasive fungal infections (IFIs) are still the most common infection-related causes for death among immunocompromised patients\(^1,2\). Haematopoietic stem cell transplant recipients (HSCT), solid organ transplant recipients and other immunocompromised patients are at risk for fungal infections\(^1\). According to most recent Infectious Diseases Society of America (IDSA) Aspergillosis and Candidemia guidelines, azoles (voriconazole, posaconazole, fluconazole, isavuconazole, itraconazole), liposomal Amphotericin B, micafungin, caspofungin are suggested for either treatment or prophylaxis of IFIs\(^3,4\).
Posaconazole is active against a wide spectrum of pathogens including *Candida* species, *Aspergillus* species and zygomycetes\(^5\). This has led to posaconazole being used for prophylaxis and treatment of fungal infections\(^6\)–\(^8\). However, posaconazole plasma concentrations may be influenced by other medications and diet, especially when posaconazole suspension is used\(^9\)–\(^12\). Additionally, related to the clinical condition of the patient, the physiological status of these patients can have an impact on pharmacokinetics of different drugs. For instance there can be a change in the volume of distribution during fluid therapy and metabolism or clearance of drugs during hepatic and renal function disorders\(^13\). A significant variation of posaconazole concentrations has been reported between and within patients\(^9\)–\(^10\).

Therapeutic drug monitoring (TDM) is recommended in guidelines for treatment optimization for posaconazole and other azoles like voriconazole and itraconazole\(^2\),\(^3\). TDM can be recommended based on an exposure response relationship\(^14\) and association of higher drug concentrations with better outcome in daily practice\(^6\),\(^7\),\(^15\). For posaconazole there is considered to be clinical benefit from TDM as posaconazole concentrations show large inter- and intra-patient variability, especially when the suspension is used\(^9\),\(^16\),\(^17\).

In contrast to the suspension, currently used posaconazole tablets and intravenous infusion are expected to result in more stable posaconazole concentrations\(^18\),\(^19\). TDM of posaconazole has been performed for several years\(^20\)–\(^23\), but the quality of TDM (application to clinical practice, dose alteration recommendations by pharmacists, optimal timing of measurements) and its implication to clinical practice has not been extensively addressed in studies as it has been for voriconazole\(^24\). Also, there is minimal information available on the
potential benefit of TDM in clinical practice for the newer drug formulations. Therefore, TDM of posaconazole has continued to be a subject of debate. A recent study investigated the effect of inflammation reflected by C-reactive protein (CRP) on posaconazole metabolism. It was concluded that CRP does not affect posaconazole exposure. However, other laboratory markers may be associated with altered drug exposure. For instance, due to chemotherapy, concomitant medications can cause liver function disorders which affect the pharmacokinetic processes like absorption, distribution, elimination, metabolism, which can lead to changes in posaconazole exposure. Analysing potential effect of routine laboratory markers can help defining the appropriate population for TDM of posaconazole.

The aim of this study was to evaluate the TDM practice in haematologic patients of posaconazole after the introduction of the new drug formulations and give recommendations for improvement of routine clinical practices of TDM. Additionally, we analysed if the routine laboratory measurements have effect on posaconazole concentrations.

**Material and methods**

A post-hoc analysis was performed from a prospective observational study conducted between August 2015 and June 2017 in the University Medical Center Groningen (UMCG), the Netherlands. Patients (aged ≥18 years) with haematological malignancies, who received intravenous and/or oral posaconazole for treatment, or (primary and secondary) prophylaxis of fungal infections were included in the study.
The study was reviewed by the local ethics committee and received approval (Institutional Review Board 2013-491). A written informed consent for collection of the medical data was obtained from each enrolled patient.

For every patient, information about posaconazole administration was recorded and included: posaconazole dose, indication for posaconazole (treatment or prophylaxis), route of administration (oral or intravenous), time of administration, day after treatment initiation with posaconazole and posaconazole serum concentration. In addition we collected laboratory analysis C-reactive protein (CRP), alkaline phosphatase (ALP), aspartate aminotransferase (AST), alanine aminotransferase (ALT), γ-GT (gamma-glutamyltransferase) and bilirubin values. The blood samples for measuring posaconazole serum concentrations were collected for routine care and concentrations were measured using a validated liquid chromatography-tandem mass spectrometry assay. In addition, other patient data including age, gender, height, underlying disease were collected.

During daily treatment with posaconazole, dosages were increased if pre-dose trough concentrations were too low (i.e. < 0.7mg/L for prophylaxis or < 1.5 mg/L for treatment) or decreased if pre-dose trough concentrations were too high (>3.75 mg/L, both treatment and prophylaxis), however no upper toxicity-threshold for posaconazole levels is known. For this study steady state was assumed on day 6 with a loading dose and on day 10 without a loading dose. The concentrations obtained prior to steady state were not included in the longitudinal analysis. The samples that were not at steady state were used to analyse TDM practices of our hospital.
The recommendations including dosage advice given by the clinical pharmacist if the posaconazole concentrations was out of the therapeutic range were collected from the electronic prescribing and laboratory information systems. To determine intra-patient variability in posaconazole plasma concentrations, patients who had more than one trough concentration measured were included in this subgroup analysis.

For the analysis of TDM practices it was documented if a recommendation was provided when posaconazole concentrations were out of the therapeutic window. Additionally, the overall number of recommendations provided and how many of these required a dosage change were summarized. When a recommendation to change the dose was followed by an actual dose change this was considered as a successful implementation into patient care.

For patients who received posaconazole for prophylaxis, occurrence of a breakthrough invasive fungal infection was documented. For all patients (receiving posaconazole for prophylaxis and treatment) 28-day and 12-week overall survival was documented, to analyze short and long-term survival. It was taken into account that optimum IFI treatment duration is 6-12 weeks.\(^{32}\)

Numerical variables were summarized with medians and interquartile range, while categorical variables were summarized by frequencies and percentages. The longitudinal data on posaconazole concentration was analyzed with a random intercept model for subjects. For the longitudinal analysis we included only steady state concentrations as defined in our prospective study\(^{27}\). The baseline variables gender, age, route of administration and dose, as well as the time-varying variables ALP, ALT, AST, γ-GT, bilirubin

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and CRP were included as independent variables. The Wald-type type 3 test statistic was applied to test for the null hypothesis ($\alpha = 0.05$) that the independent variables do not contribute to the posaconazole concentration. Multiple imputation, using predictive mean matching on all variables in the mixed model and 20 imputation data sets, was applied as sensitivity analysis. Pooled estimates were obtained using Rubin’s rule. The analyses were conducted with SAS version 9.4.

Results

Patient characteristics

Between August 2015 and June 2017, 47 patients with a median age of 62 (IQR 56-67) were enrolled in this study and 217 posaconazole samples were available for analysis for TDM practices and 182 samples for longitudinal analysis. Seven samples were excluded for further analysis as posaconazole was not detectable (<0.1 mg/L) because the drug was stopped before that time, and one sample for one patient because of missing start date.

Most common underlying disease was acute myeloid leukemia (AML, 61%) and the majority of patients (70%) received posaconazole for prophylaxis. Posaconazole modified release (MR) tablets were the main drug formulation used (89%) and 5 patients (11%) had treatment with both intravenous infusion which was followed by MR tablet throughout the study. Almost half (49%; 23/47) of the patients received a loading dose of 300mg two times daily on the first day of treatment. The median daily dose for all measured concentrations was 4.1 mg/kg (IQR 3.5-6.1), 2 patients were on dose 200 mg/day (prophylaxis), 33 patients were on dose 300 mg/day (26 prophylaxis, 7 treatment), 1 patient was on dose 600 mg/day (treatment) and for 11 patients (4 prophylaxis, 7 treatment) the doses varied throughout
treatment period. Other patient characteristics are described in Table 1. Figure 1 shows first and subsequent posaconazole concentrations.

**Analysis of TDM practices**

For 212 (98%) posaconazole concentrations a recommendation by a clinical pharmacist was given and made available to the physician in the electronic patient records. For 54 (25%) of these samples (31 prophylaxis, 23 curative treatment) a dosage change was recommended. However, dose recommendations were implemented in only 39% (10 prophylaxis, 11 treatment) of the cases. For 6 samples we did not have follow-up dosing.

The other dosages that were not changed (n=27) can be explained by some suggestions given on a Friday or during weekend (n=5), borderline concentrations 0.5-0.7 mg/L for prophylaxis and 1.0-1.5 mg/L for treatment (n=8), concentrations over 3.75 mg/L as there is no upper toxicity concentration confirmed (n=7), concentrations measured before day 6, the assumed steady state (n=2) and other reasons (n=5).

**Prophylaxis with posaconazole**

Thirty-three patients received posaconazole for prophylaxis (126 posaconazole samples) and 32 of them were on MR tablets only. A median of 2 (IQR 1-4) blood samples were taken per patient and the median drug concentration was 1.7 mg/L (IQR 0.8-2.7), the interpatient variance was 1.53 and standard deviation 1.24. Figure 2 shows intra- and interpatient variability for patients who had 5 or more samples measured while being on the same dose.
Overall 88 concentrations were within the therapeutic range (0.7 – 3.5 mg/L) and 32 outside (16 samples <0.7mg/L, 16 samples >3.75mg/L). Table 2 presents the samples outside the predefined therapeutic window and posaconazole therapy.

From 33 patients who received posaconazole for prophylaxis, three patients (9%) developed a probable IFI and one (3%) received posaconazole as empiric treatment for IFI (suspected breakthrough IFI). These patients had adequate posaconazole concentrations – all samples measured were over 0.7 mg/L. The detailed description of these patients are presented in Table 3.

The mortality rate in the total prophylaxis group was 6% (2 patients) after 28 days and 24% (8 patients) after 12 weeks. For the 2 patients who died after 28 days adequate posaconazole concentrations (≥0.7 mg/L) were observed. For the 8 patients who died after 12 weeks, 6 had adequate posaconazole concentrations (≥0.7 mg/L) and 2 patients both had 1 sample measured below 0.7 mg/L. Mortality was not attributed to a fungal infection.

**Treatment with posaconazole**

Fourteen patients received posaconazole MR tablets for treatment (91 posaconazole samples) and 4 received both posaconazole MR tablet followed by intravenous infusion or vice versa during the same treatment period. A median of 6 (IQR 3-9) samples were taken per patient and the median drug concentration was 1.76 mg/L (IQR 1.3-2.3), the interpatient variance was 0.5 and standard deviation 0.71. Figure 2 shows intra- and interpatient variability of patients who had 5 or more samples taken.
Forty-four posaconazole concentrations were within the therapeutic range (1.5 – 3.75 mg/L) and 46 outside (35 samples <1.5mg/L, 11 samples >3.75mg/L). Table 2 presents the samples outside the predefined therapeutic window and posaconazole therapy.

The mortality rate in this group was 14% (2 patients) after 28 days and 29% (4 patients) after 12 weeks. For the 2 patients who died after 28 days adequate posaconazole concentrations (≥1.5 mg/L) were observed. For the 4 patients who died after 12 weeks 2 had adequate posaconazole concentrations (≥1.5 mg/L) and 2 had some concentrations under the predefined therapeutic concentration (≤1.5 mg/L).

**Longitudinal analysis**

The associations of the independent variables on posaconazole concentration together with their 95% confidence interval and the Wald-type p-value are provided in Table 4. The results on the original data (with missing data) as well as the pooled estimates from the imputation are given. The original data set contains 127 measurements (from the 182 measurements) with a complete data set.

It is obvious that the dose contributed to the posaconazole concentration. In the analysis of the original data set (with missing data), ALT seemed to contribute to the posaconazole concentration, but this association seemed to disappear when multiple imputation is being used. Multiple imputation showed that subjects who had missing data on ALT had on average a lower ALT value that the subjects from whom we observed ALT data (34.0 versus 51.6). This may suggest that the associations of the independent variables in the original data are somewhat biased.

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Discussion

The objective of this study was to analyze routine TDM practices of posaconazole. Our study showed that variability in drug exposure is still present. Posaconazole concentrations might be affected by treatment setting - some patients were treated in an outpatient setting. However, some of these patients suffered from graft-versus-host-disease, which can compromise the absorption of posaconazole. Variability of posaconazole $C_{\text{min}}$ (MR tablet) was also described in a recent study on lung transplant recipients. However, we did see an increase of median posaconazole concentrations compared to a previous study done in our center with posaconazole suspension. In that study the median posaconazole concentration was 0.9 mg/L, in our study it was 1.7 mg/L (prophylaxis) and 1.76 mg/L (treatment). For most patients in van Elst et al study the patients received mostly 600 mg/day (84%) for prophylaxis and 800 mg/day (80%) for treatment. In this study 50% of the treatment group and 79% of the prophylaxis group received 300 mg of posaconazole per day. So we did see a better exposure with posaconazole tablet and intravenous formulation compared with the suspension. Lenczuk and colleagues also have shown that posaconazole concentrations are more likely to be in the therapeutic range when patients are being treated with posaconazole modified release tablet.

Posaconazole concentrations have also been described to be affected by diarrhea, body weight, male gender, use of PPIs and steroids. Our longitudinal analysis did not confirm the effect of weight and gender on posaconazole concentrations. We also did not see a change of AST levels, although posaconazole treatment is connected with liver function abnormalities. On the other hand, it has also been presented previously that liver function markers like $\gamma$-GT, ALP and ALT were not connected to higher posaconazole.
A limitation of our analysis is the fact that we did not analyse the effect of diarrhea and use of PPIs and steroids and that we included patients of a previous study, which is a part of all measured posaconazole concentrations during the study period thus does not represent the whole patient population. On the other hand, the characteristics of our dataset are somewhat similar to other studies describing posaconazole exposure in patients with haematological malignancies. The novelty of our study compared to earlier studies is the longitudinal analysis, which is taking into account the day of treatment and the time between measurements, also including all samples that have been collected for each patient. The advantage of using longitudinal analysis over univariate and multivariate analysis that have been used by earlier studies is that this type of analysis better values the effect of measurements over time.

We cannot see a relationship between low posaconazole concentrations and mortality rates. Additionally, this dataset is too small to show that low concentrations have an effect on outcomes, especially as we did not determine IFI-attributable deaths. For treatment of IFIs, higher posaconazole plasma concentrations must be obtained. In this study, over half of the concentrations measured for IFI treatment were below the therapeutic range (<1.5 mg/L). For patients receiving posaconazole as treatment significantly more samples were taken per patient compared with patients receiving posaconazole as prophylaxis. On the other hand, in this study, the defined therapeutic concentration (≥1.5 mg/L) used for treatment of IFIs was higher than previously reported (≥1 – 1.25 mg/L) to prevent antifungal resistance and to cover all strains. This caused more posaconazole concentrations to be out of the therapeutic window. If the therapeutic concentration of ≥1 mg/L was used more concentrations would have been within the range (21 samples).
Our dataset is too small to draw firm conclusions, although, most patients receiving posaconazole as prophylaxis and who had a breakthrough infection, had a therapeutic posaconazole concentration. However, three patients who received posaconazole for treatment did not have sufficient drug concentrations even when a loading dose was administered. Perhaps administering a double dose for more than 1 day when posaconazole is used for treatment of IFIs should therefore be considered.

A suggestion for dose alteration was only followed for 39% of recommendations made. The reasons behind non-implementation could have been due to borderline concentrations and samples over 3.75 mg/L as posaconazole toxic concentration has not been confirmed in literature nor by the manufacturer. Posaconazole practices were analysed before in conjunction with effect of concomitant medications, diet, concomitant chemotherapy and other variables. Additionally, in that study approximately for 20% of patients’ dosage changes were done, which led to more therapeutic concentrations. The benefit of TDM could was suggested due to also the varying posaconazole concentrations when using suspension, however not described in great detail. In this analysis we show that a quarter of posaconazole concentrations receive a suggestion for dosage change. Knowing that most of the patients received the oral formulation (tablet or suspension) we observed that TDM is still beneficial in this patient group. The overall results of this study and specific cases should be discussed in a multidisciplinary expert panel to avoid unnecessary orders and improve overall TDM practices taking into account different reasons behind non-implementation. Currently, the recommendations are documented into an electronic system and retrieved by the attending physician. To improve the communication between
physicians and pharmacists, an attending clinical pharmacist may be necessary, who would provide face-to-face consultations, thus aiding in preventing medication related errors and reducing costs\textsuperscript{39-41}.

Furthermore, antimicrobial stewardship teams are widely initiated in hospitals worldwide and it has been suggested that these teams should also include a pharmacist\textsuperscript{39}. The pharmacist could aid in choosing the best drug formulation to use, consult on appropriate empirical and prophylactic approaches, promote switching from intravenous to oral antimicrobials, analyze drug interactions and provide information about pharmacokinetics and TDM including prescribing of the new dose based on the TDM results\textsuperscript{43}. Besides this, the team should be advising appropriate antimicrobial therapy taking the specific patient and condition, documenting and analyzing resistance patterns into account\textsuperscript{40,44}.

**Conclusions**

Adequate posaconazole exposure was shown in in 64\% (affected 28 patients) of the measured concentrations. There was still an important variability present in posaconazole exposure, however in the longitudinal analysis from all the confounders only dose had a significant effect on posaconazole concentrations.

Even though posaconazole concentrations varied and recommendations were not always implemented to patient care, a large proportion of trough concentrations lied within the therapeutic range and did not need a recommendation at all. The communication between the clinical pharmacist and the attending physician should be enhanced to achieve better results in TDM practices. Close collaboration in a multidisciplinary antifungal stewardship
team and further education of medical staff is needed to increase adherence to dosage alterations.

Acknowledgements

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39. Policy statement on antimicrobial stewardship by the Society for Healthcare Epidemiology of America (SHEA), the Infectious Diseases Society of America (IDSA),

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Table 1. Patient characteristics (n=47)

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Prophylaxis</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender</td>
<td>Male</td>
<td>17 (36)</td>
</tr>
<tr>
<td>Age (years)</td>
<td>62 (57-68)</td>
<td>60 (52-67)</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>24.5 (23.5-27.7)</td>
<td>24.4 (21.7-26.6)</td>
</tr>
<tr>
<td>Underlying conditions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AML</td>
<td>19 (40)</td>
<td>11 (24)</td>
</tr>
<tr>
<td>MDS</td>
<td>7 (15)</td>
<td>1 (2)</td>
</tr>
<tr>
<td>Other*</td>
<td>7 (15)</td>
<td>2 (4)</td>
</tr>
<tr>
<td>Stem cell transplantation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Allogeneic</td>
<td>14 (30)</td>
<td>5 (11)</td>
</tr>
<tr>
<td>Autologous</td>
<td>2 (4)</td>
<td>0</td>
</tr>
<tr>
<td>No transplantation</td>
<td>17 (36)</td>
<td>9 (19)</td>
</tr>
</tbody>
</table>

Abbreviations: BMI – Body Mass Index, AML – acute myeloid leukemia, MDS – myelodysplastic syndrome

*Other includes X-linked gammaglubulinemia, T-cell prolymphocytic leukemia, follicular lymphoma, chronic myelomonocytic leukemia, Burkitt’s lymphoma, blastic plasmacytoid dendritic cell neoplasm, enteropathy associated T-cell lymphoma type 2, systemic mastocytosis, primary cutaneous T-cell lymphoma, aplastic anemia, primary myelofibrosis and acute promyelocytic leukemia.
**Table 2.** Posaconazole concentrations during prophylaxis and treatment

<table>
<thead>
<tr>
<th></th>
<th>33 (%) patients on prophylaxis</th>
<th>14 (%) patients on treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Route of administration</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oral</td>
<td>32 (68)</td>
<td>11 (24)</td>
</tr>
<tr>
<td>Intravenous and oral</td>
<td>1 (2)</td>
<td>3 (6)</td>
</tr>
<tr>
<td>Loading dose</td>
<td>17 (36)</td>
<td>6 (13)</td>
</tr>
<tr>
<td>Daily dose (mg/kg)</td>
<td>3.5 (3.4-4.3)</td>
<td>5.3 (4.2-6.8)</td>
</tr>
<tr>
<td></td>
<td>2 (1-3.5)</td>
<td>5.5 (2.75-9)</td>
</tr>
<tr>
<td>Number of samples taken per patient</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of Posaconazole samples</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Posaconazole &lt;0.7 mg/L</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Posaconazole &lt;1.0 mg/L</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Posaconazole &lt;1.5 mg/L</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Posaconazole &gt;3.75 mg/L</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Posaconazole &gt;0.7 mg/L with loading dose after/on day 6 / without loading dose after/on day 10</td>
<td>10 / 4 (# of samples)</td>
<td></td>
</tr>
<tr>
<td>Posaconazole &lt;1.5 mg/L with loading dose after/on day 6 / without loading dose after/on day 10</td>
<td>13 / 10 (# of samples)</td>
<td></td>
</tr>
<tr>
<td>120 concentrations obtained for prophylaxis</td>
<td></td>
<td>90 concentrations obtained for treatment</td>
</tr>
<tr>
<td>16 (# of samples)</td>
<td>14 (# of samples)</td>
<td></td>
</tr>
<tr>
<td>35 (# of samples)</td>
<td>11 (# of samples)</td>
<td></td>
</tr>
<tr>
<td>16 (# of samples)</td>
<td>10 / 4 (# of samples)</td>
<td></td>
</tr>
<tr>
<td>13 / 10 (# of samples)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Table 3. Clinical data of the patients who got a probable or possible breakthrough infection

<table>
<thead>
<tr>
<th>Pt</th>
<th>Demographic and clinical data</th>
<th>Initial posaconazole trough (mg/L)</th>
<th>Subsequent posaconazole troughs (mg/L)</th>
<th>IFI treatment</th>
<th>Chemotherapy/ antimicrobial therapy</th>
<th>Diagnosis of IFI</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>59-year-old woman with AML and had received a SCT</td>
<td>2.37 mg/L</td>
<td>NI</td>
<td>Amphotericin B with caspofungin</td>
<td>Cytarabine (1000 mg/m²)/ daunorubicine (60 mg/m²), prednisolone, piperacillin/tazobactam</td>
<td>HRCT: positive changes in the scan, galactomannan antigen serum index: 0.56, galactomannan antigen BAL index 0.35</td>
</tr>
<tr>
<td>2</td>
<td>46-year-old man with AML</td>
<td>1.26 mg/L</td>
<td>2.6 mg/L, 3.2 mg/L</td>
<td>Amphotericin B followed by caspofungin</td>
<td>Cytarabine (1000 mg/m²)/ daunorubicine (60 mg/m²), colistin, piperacillin-tazobactam, vancomycin</td>
<td>HRCT: positive masses in liver, galactomannan antigen serum index: 0.10</td>
</tr>
<tr>
<td>3</td>
<td>59-year-old man with systemic mastocytosis and had received a SCT</td>
<td>1.2 mg/L</td>
<td>2.4 mg/L, 1.8 mg/L</td>
<td>Amphotericin B with caspofungin</td>
<td>Ruxolitinib, cyclosporine, prednisolone, azithromycine</td>
<td>galactomannan antigen BAL index 4.90</td>
</tr>
</tbody>
</table>

Abbreviations: NI – no information

Table 4. Results of longitudinal analysis

<table>
<thead>
<tr>
<th>Variable</th>
<th>Original data set</th>
<th>Imputed data sets</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Estimate [95%CI]</td>
<td>p-value</td>
</tr>
<tr>
<td>Age</td>
<td>-0.172 [-1.341; 0.996]</td>
<td>0.766</td>
</tr>
<tr>
<td>Route of administration</td>
<td>0.032 [-0.020; 0.084]</td>
<td>0.213</td>
</tr>
<tr>
<td>Dose</td>
<td>0.322 [-0.848; 1.492]</td>
<td>0.587</td>
</tr>
<tr>
<td>ALT</td>
<td>0.006 [0.000; 0.012]</td>
<td>0.040</td>
</tr>
<tr>
<td>AST</td>
<td>0.004 [-0.013; 0.021]</td>
<td>0.648</td>
</tr>
<tr>
<td>ALP</td>
<td>-0.001 [-0.008; 0.007]</td>
<td>0.870</td>
</tr>
<tr>
<td>γ-GT</td>
<td>-0.000 [-0.003; 0.003]</td>
<td>0.939</td>
</tr>
<tr>
<td>Bilirubin</td>
<td>-0.009 [-0.035; 0.016]</td>
<td>0.467</td>
</tr>
<tr>
<td>CRP</td>
<td>0.001 [-0.004; 0.006]</td>
<td>0.597</td>
</tr>
</tbody>
</table>

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**Figure 1.** The distribution of initial posaconazole trough concentrations (n=47 panel A) and the distribution of subsequent posaconazole trough concentrations (n=170 panel B).

**Figure 2.** Intra-and interpatient variability of posaconazole concentrations in prophylaxis (A) and treatment (B) groups, x-axis presents patient number with the daily dose (mg/kg), y-axis presents number of samples.