Correlation between lamivudine plasma concentrations and patient self-reported adherence to antiretroviral treatment in experienced HIV patients

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Background: Adherence to antiretroviral treatment (ART) is important to achieve treatment success in human immunodeficiency virus (HIV)-infected patients. Most HIV clinics apply the patient self-report (PSR) method. However, the reliability of this method in experienced HIV patients remains questionable.

Purpose: To validate the PSR method for measuring adherence to ART using lamivudine (3TC) plasma concentrations in experienced HIV patients.

Methods: The study was conducted in Dar Es Salaam and involved 220 patients who were receiving ART services at HIV clinics for more than 12 months. Self-reported adherence information to ART was obtained on the day of HIV clinic visit. The patients were asked to mention the number of doses missed within the past 7 days. In addition, blood samples (2 mL) were collected from each patient on the same day. The blood samples were determined for 3TC plasma concentrations. The target 3TC plasma concentration as indicator concentration for adherent patients was determined in 20 patients who took their evening dose of antiretrovirals under supervision. The blood from these patients was drawn 3 hours after drug administration.

Results: Complete drug levels of 3TC and self-reported adherence data was obtained in 200 treatment-experienced HIV patients. Lamivudine plasma concentrations obtained in these patients ranged between 0.02–17.36 µg/mL. The mean time from dose administration to blood drawing was 3.1 ± 1.2 hours with coefficient of variation 39%. The mean 3TC plasma concentration obtained in 20 patients who took their antiretroviral dose under supervision was found to be 0.67 ± 0.46 µg/mL, range 0.25–2.33 µg/mL. As many as 82.5% of experienced HIV patients had PSRs in agreement with their 3TC plasma concentrations.

Conclusion: PSR adherence is still a valid method for ascertaining adherence to ART in treatment-experienced HIV patients.

Keywords: patient self-report, adherence, experienced HIV patients, lamivudine plasma concentrations

Introduction

Adherence to antiretroviral treatment (ART) has been demonstrated to reduce human immunodeficiency virus (HIV)-1 RNA, elevate CD4 cell counts, decrease morbidity and mortality due to acquired immunodeficiency syndrome (AIDS), and improve quality of life.1 Studies have shown that for successful ART, adherence to dosage regimen should be ≥95%.1-3 However, there are many factors which contribute to poor ART adherence, such as complexity of dosing schedule, strict dietary restrictions, drug toxicity, pill burden, forgetting, and poor drug supply chain.4-8 Nonadherence to ART may lead...
to suboptimal drug levels with possible therapeutic failure, deterioration of the immune system, and/or emergence of medicine-resistant HIV strains.1–3

Estimates of nonadherence to ART have varied widely, depending on the method of measurement, definition of non-adherence, the population studied, and the time over which adherence was monitored. There are various approaches to measure adherence, ranging from patient self-report (PSR), physician assessments, electronic monitoring, pill count, and prescription-refill days.9–11 Determination of antiretroviral drug level has been recently exploited as an alternative method for measuring adherence to ART.9,10 Other methods include visual analog scale; viral load HIV-1 RNA; and macrocytosis, defined as mean corpuscular volume measurement, in patients undergoing stavudine- and zidovudine-containing regimen.12–14

Most HIV clinics in low- and middle-income countries, apply PSR as a routine method for measuring adherence in ART patients. The PSR method is simpler and less costly compared with other subjective and nonsubjective methods. PSR does not require expensive bioanalytical techniques and highly skilled personnel. However, it has been reported that the PSR method can lead to an overestimation of adherence in treatment-experienced HIV patients.5–10 In addition to lacking a standardized instrument for universal use in routine clinical practice,15–19 the validity of PSR in experienced HIV patients in developing countries may be questionable. This study reports the extent of validity of the PSR method in measuring adherence to ART in experienced HIV patients in Tanzania.

Materials and methods

Study site

This was a cross-sectional study involving 220 HIV-infected patients receiving ART at Mwananyamala and Muhimbili HIV clinics located in Dar Es Salaam. The study took place between July and December 2010. Approval to conduct the study was granted by Muhimbili University of Health and Allied Sciences (MUHAS) Ethics Committee.

Enrollment of subjects

Nurse counselors working at the HIV clinics were trained as research assistants. They were also involved in the identification of files of eligible patients for inclusion in the study. Initially, 400 patients were purposely selected and the major inclusion criterion was the receiving of lamivudine (3TC)-based ART for more than 12 months and continued attendance at the HIV clinics for care and treatment. Nurses and counselors at each HIV clinic were involved in identification of those patients who were assessed by HIV clinicians and found to be eligible for the study. Patient selection was based on the “first-in, first-taken” principle, as long as the patient met the inclusion criteria. Out of 400 patients, 220 patients who consented to participate to the study on the day of clinic visitation were selected. The drug combination given to patients was zidovudine/stavudine + 3TC + nevirapine/efavirenz. The research assistants documented all available data from patients’ files in line with routine measurements.

Collection of adherence information from patients

Eligible patients were asked by the research assistants to report the exact number and date of the doses they had missed in the past 7 days, as is done routinely in HIV clinics in Tanzania. They were also asked to indicate the hour at which they ingested the last ARV dose before presenting to the clinic. In addition, blood samples were collected from each patient. A single venous blood sample (2 mL) was drawn into an ethylenediaminetetraacetic acid tube. The time interval between the last ARV dose intake and blood sampling was documented. The patients were not aware of the study before attending their consultation and therefore did not know that they were having blood drawn to measure 3TC plasma levels. All blood samples were single samples obtained just after undergoing medical examinations. The blood samples were centrifuged immediately at 1000 g to obtain plasma. The obtained plasma samples were kept at −80°C until drug assay.

The study determined the indicator (target) 3TC plasma concentration in 20 separate patients who had also been on ART for more than 12 months. Since on average, a patient spent 3 hours at the HIV clinic before blood withdrawal, analogously, blood was drawn from these patients 3 hours after taking their ARV evening dose under supervision. The blood samples were processed to obtain plasma and stored until drug assay, as described above.

Determination of ARV in blood samples

Lamivudine plasma concentrations were determined by using a high-performance liquid chromatography method published by Zhou and Sommadossi with minor modifications.19 Sample analysis was carried out in the MUHAS–Sida (Swedish International Development Cooperation Agency) Bioanalytical Laboratory, Unit of Pharmacology and Therapeutics, School of Pharmacy, MUHAS in Dar Es Salaam, Tanzania. Prior to running the samples from the study patients, the method was validated with respect to intra- and interday

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precision and accuracy. The intra- and interday precision rates were always \(<10\%\) and sample recovery was consistently between 90\%–110\% of the spiked samples, thus indicating the accuracy and suitability of the analytical method used.

**Data analysis**

To verify the accuracy of the PSR method in treatment-experienced HIV patients, the proportion of patients whose self-reported responses were in agreement with the determined plasma concentrations was calculated. Table 1 reflects the plasma concentrations obtained in 20 patients who took their evening ARV dose under supervision. The mean 3TC plasma concentration determined from blood samples drawn 3 hours after drug intake in these supervised patients was \(0.67 \pm 0.46 \, \mu g/mL\) (0.21–1.13 \(\mu g/mL\)) and ranged from 0.25 to 2.33 \(\mu g/mL\). Therefore, all patients with 3TC plasma concentrations \(\geq 0.21–2.33 \, \mu g/mL\) were taken as adherent as long as they stated so when interviewed by the research assistant. To confirm the existence of a correlation adhering to blood drawing was 3.1 \(\pm\) 5 years.

The mean time (data not shown) from dose administration to blood drawing was 3.1 \(\pm\) 1.2 hours with coefficient of variation (CV) 39\%. Some demographic data, such as age, sex, weight, height, and concomitant therapy, were included in patient files, but in most files these data were missing.

Table 1 reflects the 3TC plasma concentrations obtained from 20 patients who took 3TC-based antiretroviral therapy (ART) in the evening under supervision and had their blood drawn 3 hours after drug intake. The mean 3TC plasma concentrations obtained from these patients was \(0.67 \pm 0.46 \, \mu g/mL\) (0.21–1.13 \(\mu g/mL\)) and ranged from 0.25 to 2.33 \(\mu g/mL\) with CV 69\%.

Complete drug levels of 3TC and PSR-adherence data were obtained in 200 out of 220 study patients. Figure 1 shows the distribution of 200 patients with respect to 3TC plasma concentrations. The concentrations have been categorized into three groups: (1) those below the cut-off point of \(<0.21 \, \mu g/mL\); (2) those within the target predetermined range \(-0.21 \text{ to } 2.33 \, \mu g/mL\); (3) and those \(>2.33 \, \mu g/mL\).

Lamivudine plasma concentrations obtained in these patients ranged between 0.02 and 17.36 \(\mu g/mL\) (CV \(>60\%\)). Self-reported adherence indicated that, out of 200 patients whose plasma samples were analyzed for 3TC concentrations, 5\% (10/200) reported missing two to three doses in the past 7 days, including the day they visited the clinic, and their plasma concentrations were \(<0.21 \, \mu g/mL\). There were six other patients who reported not missing any dose in the past 7 days but had 3TC plasma concentrations below the level required for adherent patients. Three patients reported taking two doses of ARV in the morning and had 3TC plasma concentrations \(>2.33 \, \mu g/mL\).

The number of patients whose self-reported responses were in agreement with the determined plasma concentrations was calculated as follows:

1. Patients whose 3TC plasma concentrations were found to lie within the predetermined target concentration (0.21–1.13 \(\mu g/mL\)) obtained in 20 supervised patients.
2. Patients who reported not missing any dose and their 3TC plasma concentrations were found to be >1.13 µg/mL but <2.33 µg/mL (1.13–2.33 µg/mL).
3. Patients who reported missing ARV doses and also had 3TC plasma concentrations <0.21 µg/mL.

Table 2 summarizes the distribution of the 200 study patients in relation to PSR adherence and 3TC plasma concentration. The proportion of HIV-treated patients whose self-reported responses were in agreement with the determined plasma concentrations is also demonstrated in Figure 2. As many as 82.5% of patients (165) had PSR responses in agreement with their 3TC plasma concentrations (Table 2).

Discussion
The study has determined the validity of PSR adherence to ART in experienced HIV patients in a low-income country. Lamivudine plasma concentrations were used as indicator of adherence in the study patients. The quantified 3TC plasma levels are grouped as reflected in Figure 1. Lamivudine plasma concentrations obtained in these patients ranged between 0.02 and 17.36 µg/mL (CV > 60%). There was high variation of 3TC plasma concentrations obtained among the study patients. The results are consistent with those obtained by Moore et al and Panhard et al.20,21

The large variation of 3TC plasma concentrations among patients may be partly explained by the different hours at which the patients took their morning ARV doses before blood was drawn by the phlebotomist. It could also be explained by variability of body weights and heights (body mass index) and renal conditions among patients. Unfortunately, the study clinic did not register the body mass index of most patients, and none of the patients had a serum creatinine assessment to determine renal performance. The study used only the data available in patient files in line with what is done routinely in Tanzanian clinics. Lamivudine is a prodrug that undergoes phosphorylation when catalyzed by intracellular kinases to form 3TC-5′-triphosphate, the active metabolite that prevents viral replication.23 Variation in 3TC disposition and, subsequently, plasma concentrations among patients could therefore depend on a patient’s individual capacity to phosphorylate the parent drug. Nevertheless, 3TC is renally excreted and, recently, a low 3TC renal clearance was demonstrated in patients who had impaired renal functions.24,25 Lamivudine has very short elimination half-life of 3–4 hours21,22 and this may also have contributed to the large variation in the drug plasma concentrations among individual patients.

In this study, it was shown that there was discrepancy between the obtained drug plasma concentrations and

Table 2 Proportion of patients (number in brackets) in relation to patient self-report (PSR) and lamivudine (3TC) plasma concentration agreement (n = 200)

<table>
<thead>
<tr>
<th>3TC plasma concentration in µg/mL</th>
<th>Proportion of patients with agreement between 3TC concentration and PSR</th>
<th>Proportion of patients with disagreement between 3TC concentration and PSR</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;0.21 µg/mL</td>
<td>5 (10)</td>
<td>3 (6)</td>
</tr>
<tr>
<td>0.21–1.13 µg/mL</td>
<td>51 (102)</td>
<td>3 (6)</td>
</tr>
<tr>
<td>&gt;1.13–2.33 µg/mL</td>
<td>25 (50)</td>
<td>5 (10)</td>
</tr>
<tr>
<td>&gt;2.33 µg/mL</td>
<td>1.5 (3)</td>
<td>6.5 (13)</td>
</tr>
<tr>
<td>Total</td>
<td>82.5 (165)</td>
<td>17.5 (35)</td>
</tr>
</tbody>
</table>
Patient self-reported drug intake in about 18% of patients. Although the proportion of nonadherent patients in the study was found to be low, it is still important to validate the PSR method regularly to reinforce its effectiveness, particularly in treatment-experienced HIV patients in low- and middle-income countries. Overall results show that there was good adherence in the patients studied. Several factors could explain the good adherence obtained during the study; for instance, free access to ARV drugs, the then-effective ARV procurement system, and adequate counseling of eligible patients.

The study applied 3TC plasma concentrations as a validity marker of PSR. However, it is still debatable whether the best drug of choice, as a marker of adherence, should have short or long half-life. Those in support of short half-life drugs argue that missing a single dose of such a drug will lead to a detectable change in drug plasma concentration, as opposed to drugs with long half-life. As for drugs with a long half-life, a single missed dose may not be detectable, especially if the steady state plasma concentration had already been attained. Recently, Segeral et al applied plasma concentrations of efavirenz and nevirapine (both have long half-lives) to determine adherence in Cambodian patients undergoing first-line World Health Organization-recommended HAART; the results obtained were consistent with those obtained by other methods.12

Figure 2 Proportion of patients whose determined lamivudine (3TC) plasma concentrations was in agreement with patient self-report.

**Conclusion**

This study has demonstrated good agreement between PSR adherence and 3TC plasma concentrations in over 80% of the study patients. It can be concluded that the PSR-adherence method is still reliable for measuring adherence in patients undergoing ART, including experienced HIV patients in HIV clinics in Tanzania.

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**Disclosure**

The authors report no conflicts of interest in this work.

**References**


