Mendelsohn, Joshua B; Schilperoord, Marian; Spiegel, Paul; Balasundaram, Susheela; Radhakrishnan, Anuradha; Lee, Christopher KC; Larke, Natasha; Grant, Alison D; Sondorp, Egbert; Ross, David A; (2014) Is forced migration a barrier to treatment success? Similar HIV treatment outcomes among refugees and a surrounding host community in Kuala Lumpur, Malaysia. AIDS and behavior, 18 (2). pp. 323-334. ISSN 1090-7165 DOI: https://doi.org/10.1007/s10461-013-0494-0

Downloaded from: http://researchonline.lshtm.ac.uk/id/eprint/967532/

DOI: https://doi.org/10.1007/s10461-013-0494-0

Usage Guidelines:

Please refer to usage guidelines at https://researchonline.lshtm.ac.uk/policies.html or alternatively contact researchonline@lshtm.ac.uk.

Available under license: http://creativecommons.org/licenses/by/2.5/
Is Forced Migration a Barrier to Treatment Success? Similar HIV Treatment Outcomes Among Refugees and a Surrounding Host Community in Kuala Lumpur, Malaysia

Joshua B. Mendelsohn · Marian Schilperoord · Paul Spiegel · Susheela Balasundaram · Anuradha Radhakrishnan · Christopher C. K. Lee · Natasha Larke · Alison D. Grant · Egbert Sondorp · David A. Ross

© The Author(s) 2013. This article is published with open access at Springerlink.com

Abstract In response to a near absence of studies among refugees and host communities accessing highly active antiretroviral therapy (HAART) in urban settings, our objective was to compare adherence and virological outcomes among clients attending a public clinic in Kuala Lumpur, Malaysia. A cross-sectional survey was conducted among adult clients (≥18 years). Data sources included a structured questionnaire that measured self-reported adherence, a pharmacy-based measure of HAART prescription refills over the previous 24 months, and HIV viral loads. The primary outcome was unsuppressed viral load (≥40 copies/mL). A sample of 153 refugees and 148 host clients were recruited. Refugees were younger (median age 35 [interquartile range, IQR 31, 39] vs 40 years [IQR 35, 48], p < 0.001), more likely to be female (36 vs 21 %, p = 0.004), and to have been on HAART for less time (61 [IQR 35, 108] vs 153 weeks [IQR 63, 298]; p < 0.001). Similar proportions of those on treatment for ≥25 weeks from both groups were not virologically suppressed (19 vs 16 %, p = 0.54). The proportions in each group with <95 % adherence to pharmacy refills were 26 versus 34 %, p = 0.15. Refugee status was not independently associated with the outcome (adjusted odds ratio, aOR = 1.28, 95 % CI 0.52, 3.14). The proportions of refugee and host community clients with unsuppressed virological outcomes and sub-optimal adherence were similar, supporting the idea that refugees in protracted asylum situations are able to...
sustain good treatment outcomes, and should explicitly be included in the HIV strategic plans of host countries with a view to expanding access in accordance with national guidelines for HAART. Responding to a ausencia casi total de estudios entre refugiados y las comunidades de acogida y acceso a terapia antirretroviral de gran actividad (TARGA) en zonas urbanas, nuestro objetivo fue comparar la adherencia y resultados virológicos entre los clientes que asisten a una clínica pública en Kuala Lumpur, Malasia. Un estudio transversal se llevó a cabo entre los clientes adultos (≥18 años). Las fuentes de datos incluyen un cuestionario estructurado que midió adherencia auto-reportada, una medida farmacéutica basada en el relleno de medicamentos recetados de TARGA durante 24 meses, y la carga viral del VIH. El resultado principal fue carga viral no suprimida (≥40 copias/mL). Una muestra de 153 refugiados y 148 clientes de la comunidad de acogida fueron reclutados. Los refugiados eran más jóvenes (media de 35 años [rango intercuartil, IQR 31, 39] frente a 40 años [IQR 35, 48], p < 0.001), más probabilidades de ser mujer (36 vs 21 %, p = 0.004), y haber estado en TARGA durante menos tiempo (61 [IQR 35, 108] vs 153 semanas [IQR 63, 298], p < 0.001). Una proporción similar de las personas en tratamiento durante ≥25 semanas de ambos grupos no tuvieron supresión virológica (19 vs 16 %, p = 0.54). Las proporciones de cada grupo con <95 % de adherencia a recetas de farmacias eran 26 frente a 34 %, p = 0.15. La condición de refugiado no se asoció de forma independiente con el resultado (razón de momios ajustado, aOR = 1.28, IC del 95 %: 0.52, 3.14). Las proporciones de refugiados y de clientes de la comunidad de acogida con resultados virológicos no suprimidos y adherencia sub-óptimas fueron similares, apoyando la idea que los refugiados en situaciones de asilo prolongados son capaces de mantener buenos resultados del tratamiento, y deberían explícitamente incluirse en los planes estratégicos de VIH de los países de acogida con el fin de ampliar el acceso de acuerdo con las directrices nacionales de TARGA.

**Keywords**  Refugies · Forced migration · HIV · Antiretrovirals · Outcomes · Adherence

**Abbreviations**
- ART  Antiretroviral therapy
- CI  Confidence interval
- HAART  Highly active antiretroviral therapy
- HIV  Human immunodeficiency virus
- IDP  Internally-displaced person
- LSHTM  London school of hygiene and tropical medicine
- MSF  Médecins Sans Frontières
- NGO  Non-governmental organisation
- OR  Odds ratio
- aOR  Adjusted odds ratio

**Introduction**

Sustained excellent adherence to highly active antiretroviral therapy (HAART) is essential for achieving and sustaining suppression of HIV infection. Some estimates suggest that 6.6 million people in low and middle income-countries, or 47 % of 14.2 million eligible, are now receiving treatment [1]. Although refugees have fled across an international border and have a recognised international legal status that should enable them to receive access to medical care on an equivalent basis to host nationals in countries where they have sought asylum [2]; there are concerns as to whether refugees who are on HAART are sufficiently stable and therefore capable of sustaining optimal adherence and good treatment outcomes given potential obstacles such as language barriers, lack of employment and the risk of onwards displacement to other countries [3, 4]. In some instances, governments may be reluctant to provide treatment to refugees [5], citing concerns about stability and the prerogatives of supplying medications to their own citizens. Previous studies of adherence and treatment outcomes in other forcibly displaced and conflict-affected groups have reported high levels of adherence and acceptable outcomes suggesting that such obstacles may be overcome, but most of this work was conducted in sub-Saharan Africa or with refugees based in high-income countries [6]. There are few data available to verify the acceptability of treatment outcomes among refugees in relation to surrounding host communities in low and middle-income settings, where most of 10.6 million global refugees were situated as of 2010 [7]. In response, our objective was to study adherence and HIV treatment outcomes among refugee and host community clients accessing HAART from the same clinic in Kuala Lumpur, Malaysia. We hypothesized that refugees would exhibit inferior outcomes when compared with the surrounding host community.

**Method**

**Study Setting**

Sungai Buloh Hospital, Kuala Lumpur, Malaysia was chosen as the study setting as it met our criteria of an urban, Southeast Asian setting, with sufficient numbers of refugees...
accessing HIV treatment and care services from a single point of care. At the start of the study (April 2010), 91,985 individuals were registered by the United Nations High Commissioner for Refugees (UNHCR) as refugees or asylum seekers in Malaysia, 315 had an HIV diagnosis, and 171 were on HAART. Over 98 % of refugees on HAART were from Myanmar. By the end of 2009, the average length of a stay for an HIV-positive refugee in Malaysia was 3.7 years; 32 % were resettled to high-income countries after an average of 2.9 years (UNHCR Representation in Malaysia, Pers. Comm). Malaysia has not signed the 1951 Refugee Convention and its 1967 Protocol; however, the Ministry of Health issued a circular in 2006 that permitted refugees to access public health services, including antiretroviral therapy (ART) as part of the national HIV treatment and program. Initially not included in national strategic plans [8], refugees were formally included in the 2011–2015 Strategic Plan [9]. The Malaysian host community, comprised primarily of Malay, Chinese and Tamil groups, were fully subsidised by the national treatment program for first-line HAART (usually stavudine, lamivudine and nevirapine) and virological monitoring; second-line treatment were partially subsidised. For refugees, the national program fully subsidised first-line treatments but more expensive first and second-line drugs (e.g. efavirenz; lopinavir/ritonavir) and virological monitoring were paid for by UNHCR. Refugees did not pay out of pocket for treatment. Only refugees, meaning those who possessed documented approval of their refugee status, received subsidised treatment and support; asylum seekers were expedited through the Refugee Status Determination process in order to facilitate timely access to treatment, but did not have access to treatment until refugee status was formally confirmed.

Study Design

A 15-week (April–July 2010) cross-sectional survey, conducted at the Infectious Diseases Clinic, Sungai Buloh Hospital, aimed to recruit all refugees identified by UNHCR as recipients of HAART and a similar number of host community clients attending the same outpatient clinic. Inclusion criteria were ≥18 years of age and on HAART for ≥30 days. Refugees had routine access to the clinic one day per week, therefore we sought to recruit host community clients on only one other day per week. Those who met the inclusion criteria were recruited consecutively at the time of their regular clinic appointment and were re-contacted if they agreed but were unable to participate at the time of recruitment. In an attempt to obtain a complete sample, all eligible refugee clients on HAART who met the inclusion criteria but were not seen in the clinic during the study period were contacted by telephone or by a community representative. As attempts were made to recruit all refugees known to be on HAART, the number of eligible refugees determined the upward limit on sample size. Power calculations were initially completed using expected numbers of refugees on HAART and expected proportions virologically suppressed. Given a sample size ratio 1:1, with 150 clients per group (representing 88 % of eligible refugees) and a level of viral suppression of 70 % in the refugee group, the study had an 80 % chance of detecting a 14 % prevalence difference as statistically significant at the 5 % level. Recruitment of the host community on a 2:1 basis lowered the detectable difference to 12 % (net efficiency gain = 14 %), therefore, the 1:1 strategy was deemed sufficient for comparison. To assess representativeness of the host community sample, a sampling frame was constructed and a randomly selected comparison sample of 150 host clients was selected in order to compare basic demographic data with the recruited host community sample.

Data Sources

The primary outcome was unsuppressed viral load (≥40 copies/mL). Data sources included a structured questionnaire with self-reported adherence measures, a pharmacy-based measure of HAART prescription refills over the previous 24 months and HIV viral loads. The structured questionnaire was translated into Bahasa Malaysia, Tamil, Mandarin, Burmese, and Falam (Chin dialect), then back-translated into English. The original and back-translated English versions were reconciled, then adjusted during pretesting to enhance validity. Key self-reported adherence measures included a retrospective four-day dose-by-dose recall [10] and a retrospective one-month general recall measured on a visual analogue scale (VAS) [11]. Adherence to pharmacy refill schedule was assessed using a pharmacy-based measure of HAART prescription refills, calculated as the proportion of prescribed refills collected divided by the total required refills for up to 24 months prior to the interview date. A successful refill was determined by dividing the number of tablets claimed into the number of tablets required to avoid a personal stock-out, allowing a 14-day grace period for each collection. For all adherence measures, <95 % of doses taken as prescribed was used to signify “sub-optimal adherence”. Blood samples for HIV viral load measurement were collected using routine phlebotomy procedures and analysed using the COBAS Ampliprep/Taqman platform (Roche Diagnostics Systems, Branchburg, NJ, USA).

Statistical Methods

Socio-demographic characteristics were compared between host and refugee groups using Mann–Whitney tests,
chi-square or Fisher’s exact tests and chi-square tests for trend. Risk factors for unsuppressed viral load were evaluated using unconditional logistic regression; effect estimates were odds ratios (OR) and corresponding 95 % confidence intervals (CI). The order of entry of factors into the model was determined using a three-level, forwards, step-wise modelling approach drawing on social action theory [12] to group factors into levels representing treatment “contexts” such as socio-demographic and displacement factors; “self-change processes” such as knowledge scores and self-efficacy; and “action state” factors including the adherence measures. After univariable analyses, a “treatment context model” was fitted by adjusting for treatment context factors with \( p < 0.1 \) in univariable analyses. A “self-change processes model” was fitted by adjusting each new factor by all retained treatment context factors, then adjusting again for any additional factors with \( p < 0.1 \). An “action state (adherence) factors” model was fitted in a similar fashion but adjustment was restricted to factors from previous levels only, excluding collinear adherence measures. The final regression model was obtained by excluding factors with the highest \( p \) value, sequentially, until all remaining factors met \( p < 0.05 \). Covariates of interest retained throughout the modelling process included refugee status, age, and time on HAART. Adherence factors were retained but not included in final model building to mitigate over-adjustment bias [13–15].

Ethical Approval

Ethical approval was received by the Clinical Research Centre and the Medical Research Ethics Committee, Malaysia (Approval 3275) and the London School of Hygiene and Tropical Medicine Research Ethics Committee (Approval 5547).

Results

Study Population

We recruited 153 refugees and 148 Malaysian adults reflecting 90 % and 81 % participation rates (eligible clients who were seen or contacted and agreed to participate), respectively. The Malaysian group comprised 6 % of the target population of eligible clients (\( N = 2,870 \)) and was similar on most socio-demographic indicators to a randomly sampled host comparison group (Supplementary Table 1). Almost all (95 %) HIV-positive refugees accessing services from the study clinic were Burmese while the host community group was 61 % Chinese, 25 % Malay, and 15 % Tamil or other ethnic groups. The recruited refugee and host community groups were different on a variety of indicators (Table 1). The refugee group was younger (median age 35 vs 40 years, \( p < 0.001 \)), had a higher proportion of women (36 vs 22 %, \( p = 0.006 \)), a shorter median time on HAART (61 vs 153 weeks, \( p < 0.001 \)), a shorter time since HIV diagnosis (113 vs 315 weeks, \( p < 0.001 \)), and a lower most recent routine CD4 count (278 vs 350 cells/\( \mu L \), \( p = 0.03 \)). Among refugees, the median time of residence in Malaysia was 3.6 years (IQR 2.0, 6.2) and the median time since having received formal refugee recognition was 1.8 years (IQR 1.0, 2.9).

Virological and Adherence Outcomes

Viral load results indicated that 24 % (72/296) of clients had not achieved viral suppression (≥40 copies/mL). There was no difference between the proportions of refugees and host community clients who had not achieved viral suppression overall, or when restricting analyses to clients on treatment for ≥25 weeks (19 vs 16 %, \( p = 0.54 \); Table 2). On key measures of self-reported adherence among all surveyed clients, both groups performed similarly (Table 3). The four-day recall showed that high proportions of both groups self-reported sub-optimal adherence (8 vs 4 %, \( p = 0.20 \)), whereas the proportions who self-reported sub-optimal adherence on the one-month VAS were higher (28 vs 30 %, \( p = 0.79 \)). The pharmacy refill results were also higher but similar in both groups (26 vs 34 %, \( p = 0.15 \)). Within each group, there was evidence for ordered trends between self-reported measures of adherence and proportions not virologically suppressed among clients on treatment for ≥25 weeks. On the pharmacy refill measure, there was strong evidence for this trend among refugees, but this did not hold for the host community (see Supplementary Table 2).

Risk Factors for Unsuppressed Virological Outcomes

Unsuppressed viral load was defined as ≥40 copies/mL. In initial analyses of contextual factors (Table 4), 17 % of clients on HAART for ≥1 year were not suppressed. Among those on treatment for ≥25 weeks, 15 % of those on HAART for <1 year were not suppressed. There was no significant relationship between increasing time on treatment (over 1 year) and virological outcomes (aOR = 1.17, 95 % CI 0.69, 1.96; \( p = 0.56 \)). There was no evidence for associations between self-change process factors and the outcome (Table 5). Among exposures in the action state level (Table 6), there was a protective effect of adherence to pharmacy refill schedule (aOR = 0.47, 95 % CI 0.27, 0.83; \( p = 0.009 \)) and a harmful effect of having reported any treatment interruption in the past month (aOR = 2.77, 95 % CI 0.91, 8.43; \( p = 0.08 \)), adjusting for age group, time on HAART,
refugee status, sex, temporary travel in past year, time to clinic, time from diagnosis to HAART start and previous regimen switch.

The final multivariable model (Table 7) identified female sex (aOR = 0.39, 95 % CI 0.14, 1.05; p = 0.05), increasing time between diagnosis and treatment start (aOR = 0.64, 95 % CI 0.41, 0.99; p = 0.04) and adherence to pharmacy claim schedule (aOR = 0.47, 95 % CI 0.27, 0.81; p = 0.007) as protective, while temporary migration of ≥1 month in the past year (aOR = 4.12, 95 % CI 1.70, 9.99; p = 0.002) and average travel time to clinic of ≥1 h (aOR = 3.05, 95 % CI 1.09, 8.49; p = 0.02) were independent risk factors. There was no evidence for an association between refugee status and unsuppressed viral load (aOR = 1.28, 95 % CI 0.52, 3.14; p = 0.60) adjusting for age group, refugee status, time on HAART, sex, temporary migration in the past year, average time to clinic, and time from HIV diagnosis to HAART start.

**Discussion**

In this study, the first we are aware of that investigated adherence and treatment outcomes among both refugee and a host community in an asylum setting, a minority of both refugee (19 %) and host community clients on HAART for ≥25 weeks (16 %) did not achieve viral suppression. Only minor differences were found on self-reported and pharmacy-based adherence measures. Adherence and virological outcomes were comparable to results from other Asian
Table 2 Comparison of virological outcomes in host community and refugee clients

| Time on HAART (weeks) | Group  | ≥40 copies/mL, n (%) | Total, n (%) | p value<sup>a</sup> |
|-----------------------|--------|-----------------------|--------------|---------------------|
|                       | Host   | 32 (22)               | 144 (100)    | 0.41                |
|                       | Refugee| 40 (26)               | 152 (100)    |                     |
| <25                   | Host   | 12 (67)               | 18 (100)     | 1.00                |
|                       | Refugee| 17 (59)               | 29 (100)     |                     |
| ≥25                   | Host   | 20 (16)               | 125 (100)    | 0.54                |
|                       | Refugee| 23 (19)               | 121 (100)    |                     |

<sup>a</sup> Chi-square test

<sup>b</sup> 5 % (7/147) of client’s with a previous viral load <40 copies/mL tested in the range of 40 to 499 copies/mL. Among clients displaying this low-level viraemia, no differences were observed between the groups (Fisher’s exact test, p < 1.00)

Table 3 Proportions adhering to HAART by 4 days self-report, 1 month self-report, and pharmacy refill, in refugee and surrounding host community clients

| Dose-by-dose self-report (4 days) |   |   |   |
|----------------------------------|---|---|---|
|                                  | Host| Refugee|   |
| 0+                               | 6  | 11   | 7 |
| 80+                              | 0  | 1    | 1 |
| 95+                              | 142| 141  | 92|

| Visual analogue scale self-report (1 month) |   |   |   |
|---------------------------------------------|---|---|---|
|                                             | Host| Refugee|   |
| 0+                                           | 11 | 11   | 7 |
| 80+                                          | 33 | 32   | 21|
| 95+                                          | 104| 110  | 72|

| Pharmacy claim adherence (24 months)<sup>b</sup> |   |   |   |
|-------------------------------------------------|---|---|---|
|                                                 | Host| Refugee|   |
| 0+                                               | 14 | 9    | 7 |
| 80+                                              | 34 | 26   | 19|
| 95+                                              | 95 | 101  | 74|

<sup>a</sup> Chi-square test for trend (Cochran–Armitage test)

<sup>b</sup> Since started on HAART to a maximum of 24 months, retrospectively

HIV clinics. In a multicentre prospective cohort of 17 Asian settings, Oyomopito and colleagues found that 17 % were not virologically suppressed after 12 months on HAART [16]. We are aware of only one other report of virological outcomes among any forcibly displaced or conflict-affected groups situated in low and middle-income settings [17]. In this South African study, 24 % of “foreigners”, many of whom had emigrated from Zimbabwe but who were not explicitly identified as refugees, exhibited a study-specific measure of viral failure that included individuals with a viral load of ≥1000 copies/mL. Previous adherence data collected among other groups in low and middle-income settings have shown results that are consistent with other stable cohorts. In conflict-affected northern Uganda, Kiboneka and colleagues [18] found adherence levels of <95 % in 8 % of internally-displaced persons (IDP), as measured by a composite adherence score. In a Ugandan cross-sectional study of IDPs, mean self-reported adherence was 99.5 % [19]. In the western Equatorial province of Sudan, 12 % of refugees and IDPs on HAART for ≥6 months self-reported <95 % adherence [20]. During active conflict in the Democratic Republic of the Congo, sub-optimal adherence (measured by pill counts) was found in only 1 % of clients while CD4 gain at six months was similar to other stable cohorts [21].

Given the potential for cross-border displacement to increase the vulnerability of refugees to inferior outcomes, it was reassuring that a high proportion of refugees were virologically suppressed in the present study. In multivariable analyses, no independent association was found between refugee status and unsuppressed viral load after adjusting for age, sex, time on HAART, time from diagnosis to HAART start, temporary migration in the past year and time to clinic. Consistent with evidence from a Canadian setting showing an adverse impact of temporary migration [22], travel outside of current residence for ≥1 month in the past year (reported by 18 % of refugees and 14 % of Malaysians) led to a fourfold increase in the odds of unsuppressed viral load, a possible consequence of difficulties locating or refilling medications when personal stocks were depleted in the absence of contingency plans. Consistent with other settings, longer travel times to clinic (≥1 h) were linked to an increase in the odds of unsuppressed viral load [23–25]. By contrast, many of the obstacles thought to negatively affect treatment outcomes among refugees such as language barriers, unemployment and instability were either not associated with the outcome or were not unique to refugees. Specifically, there was no evidence for any harm contributed by the fact of a person’s employment status or language group. Language barriers in medical contexts are clearly important, but may be overcome by the effective use of interpreters and support counsellors recruited directly from refugee communities. We did not study onwards displacement to other countries directly; however, the average length of stay for an HIV-positive refugee (3.7 years) was generally indicative of stability. The finding that temporary migration (for ≥1 continuous month in the past year) was a risk factor after adjusting for refugee status suggested that this was common to the full study group. Longer times between diagnosis and HAART start were protective, even though starting HAART at a higher CD4 counts is also known to reduce mortality [26]. Longer lead-in times to routine treatment may have encouraged readiness to begin
Table 4 Association of contextual factors with unsuppressed viral load among refugees and local host community on HAART for ≥25 weeks in Kuala Lumpur, Malaysia (N = 222)

| Factor                                      | Prevalence ≥40 copies/mL, n/N (%) | p value, crude odds ratio (95 % CI) | p value, adjusted odds ratio (95 % CI) |
|---------------------------------------------|-----------------------------------|-------------------------------------|---------------------------------------|
| Age group (years)                           |                                   |                                     |                                       |
| 18--                                       | 5/25 (20)                         | p = 0.69                            | p = 0.68                              |
| 30--                                       | 18/114 (16)                       | 0.90 (0.52, 1.55)                   | 1.15 (0.60, 2.20)                     |
| 40+                                        | 13/83 (16)                        |                                     |                                       |
| Refugee status                              |                                   | p = 0.19                            | p = 0.60                              |
| Host                                        | 15/114 (13)                       | 1                                   |                                       |
| Refugee                                     | 21/108 (19)                       | 1.59 (0.77, 3.28)                   | 1.28 (0.52, 3.14)                     |
| Time on HAART (years)                       |                                   | p = 0.79                            | p = 0.56                              |
| 0--                                        | 7/46 (15)                         | 1                                   |                                       |
| 1--                                        | 9/57 (16)                         | 1.06 (0.68, 1.67)                   | 1.17 (0.69, 1.96)                     |
| 2+                                         | 20/119 (17)                       |                                     |                                       |
| Sex                                         |                                   | p = 0.04                            | p = 0.05                              |
| Male                                        | 30/155 (19)                       | 1                                   |                                       |
| Female/transgender                          | 6/67 (9)                          | 0.41 (0.16, 1.04)                   | 0.39 (0.14, 1.05)                     |
| Time from diagnosis to start (weeks)        |                                   | p = 0.07                            | p = 0.04                              |
| 0--                                        | 19/98 (19)                        | 1                                   |                                       |
| 25--                                       | 8/30 (27)                         | 0.69 (0.47, 1.03)                   | 0.64 (0.41, 0.99)                     |
| 50+                                        | 9/94 (10)                         |                                     |                                       |
| HAART regimen, dosing                       |                                   | p = 0.32                            | p = 0.13                              |
| EFV-based                                   | 21/140 (15)                       | 1                                   |                                       |
| NVP-based                                   | 12/74 (16)                        | 1.10 (0.51, 2.38)                   | 1.03 (0.44, 2.43)                     |
| Other                                       | 3/8 (38)                          | 3.40 (0.76, 15.31)                  | 6.00 (1.14, 31.74)                    |
| Current employment                          |                                   | p = 0.23                            | p = 0.21                              |
| No                                          | 13/101 (13)                       | 1                                   |                                       |
| Yes                                         | 23/121 (19)                       | 1.59 (0.76, 3.32)                   | 1.70 (0.74, 3.95)                     |
| Mother tongue                               |                                   | p = 0.19                            | p = 0.26                              |
| Bahasa Malaysia (Malay)                     | 5/39 (13)                         | 1                                   |                                       |
| Tamil                                       | 5/26 (19)                         | 1.62 (0.42, 6.27)                   | 1.56 (0.36, 6.73)                     |
| Chinese dialects                            | 3/46 (7)                          | 0.47 (0.11, 2.13)                   | 0.47 (0.09, 2.32)                     |
| Chin dialects                               | 13/54 (24)                        | 2.16 (0.70, 6.66)                   | 6.21 (0.57, 67.53)                    |
| Burmese                                     | 3/24 (13)                         | 0.97 (0.21, 4.49)                   | 2.52 (0.17, 38.58)                    |
| Other                                       | 7/33 (21)                         | 1.83 (0.52, 6.43)                   | 3.20 (0.30, 34.63)                    |
| Household size                              |                                   | p = 0.73                            | p = 0.97                              |
| 1--                                        | 9/56 (15)                         | 1                                   |                                       |
| 3--                                        | 17/112 (15)                       | 1.09 (0.66, 1.82)                   | 1.01 (0.59, 1.73)                     |
| 7+                                         | 10/54 (19)                        |                                     |                                       |
| No. dependent minors in household           |                                   | p = 0.59                            | p = 0.98                              |
| 0--                                        | 23/133 (17)                       | 1                                   |                                       |
| 1+                                         | 13/89 (15)                        | 0.82 (0.39, 1.72)                   | 1.01 (0.44, 2.33)                     |
| Temporary migration (≥1 continuous month in past year) | p < 0.001 | p = 0.002                        |                                       |
| No                                         | 23/187 (12)                       | 1                                   |                                       |
| Yes                                        | 13/35 (37)                        | 4.21 (1.87, 9.50)                   | 4.12 (1.70, 9.99)                     |
| Pathway to diagnosis                        |                                   | p = 0.50                            | p = 0.65                              |
| Voluntary test                              | 7/43 (16)                         | 1                                   |                                       |
| Mandatory test                              | 8/40 (20)                         | 1.29 (0.42, 3.94)                   | 2.01 (0.56, 7.18)                     |
| Illness/hospitalisation                     | 16/88 (18)                        | 1.14 (0.43, 3.03)                   | 1.00 (0.34, 2.93)                     |
| Other                                       | 5/51 (10)                         | 0.56 (0.16, 1.91)                   | 1.07 (0.27, 4.25)                     |
treatment while the negative impact of delaying treatment may have been confounded by delays between seroconversion and diagnosis. Specifically, clients may start HAART during acute illness when they are more motivated to get well by adhering to treatment.

The finding that women were more likely to have achieved viral suppression could have been due to gender differences proportions disclosing their status to partners (49 % of males vs 66 % of females, \( p = 0.05 \)) and in proportions with children (40 % of males vs 61 % of females, \( p = 0.004 \)). Non-disclosure of HIV status was previously shown to affect adherence to HAART [27], while having children may provide earlier pathways to care through antenatal screening [28]. This finding was consistent with results from a Chinese study [29] and a South African study that showed a tendency for men to present for treatment later and with more advanced disease [30].

Sub-optimal pharmacy refill adherence was strongly associated with lack of viral suppression, supporting the usefulness of this measure for routine monitoring especially where viral load measurement is unavailable [31, 32]. The slightly higher proportion of Malaysians not adhering optimally to the pharmacy claim schedule may have been an artefact of a system that facilitated occasional or supplementary medication collection from external pharmacies (refugees did not have similar opportunities). One-sixth of host community clients reported collecting drugs in this manner within the assessed pharmacy refill period, which supported the recommendation that multiple routine indicators could help to facilitate monitoring of adherence patterns over time [33].

Caution must be used when generalising these findings to other refugee populations given that only one setting was studied and HAART delivery systems are so often setting-specific. The HIV-positive caseload among refugees was considerably higher in Malaysia in comparison to other major programs in the region (ten cases each in Bangkok and New Delhi). Moreover, there are differences between urban, camp and rural/dispersed refugee groups in relation to service-provision challenges [34]. Socioeconomic differences between different refugee settings may be partially mitigated by individual financial assistance (distributed by UNHCR and assessed at the country-level). As with other studies that have compared different clinical

---

**Table 4 continued**

| Factor                                      | Prevalence ≥40 copies/mL, n/N (%) | \( p \) value, crude odds ratio (95 % CI) | \( p \) value, adjusted odds ratio (95 % CI)<sup>b</sup> |
|---------------------------------------------|-----------------------------------|------------------------------------------|--------------------------------------------------------|
| Average time to clinic (hours)              |                                   |                                          |                                                        |
| 0–                                          | 6/74 (8)                          | \( p = 0.01 \)                            | \( p = 0.02 \)                                         |
| 1+                                          | 30/148 (20)                       | 2.88 (1.14, 7.27)                        | 3.05 (1.09, 8.49)                                      |
| Regimen switch, ever                        |                                   |                                          |                                                        |
| No                                          | 16/120 (13)                       | \( p = 0.20 \)                            | \( p = 0.07 \)                                         |
| Yes                                         | 20/102 (20)                       | 1.59 (0.77, 3.25)                        | 2.14 (0.94, 4.85)                                      |
| Unable to refill prescription, past 3 months|                                   | \( p = 0.41 \)                            | \( p = 0.44 \)                                         |
| No                                          | 35/210 (17)                       | \( p = 0.23 \)                            | \( p = 0.41 \)                                         |
| Yes                                         | 1/12 (8)                          | 0.45 (0.06, 3.64)                        | 0.45 (0.05, 4.08)                                      |
| Any symptom or side-effect, past 4 weeks   |                                   |                                          |                                                        |
| No                                          | 6/54 (11)                         | \( p = 0.17 \)                            | \( p = 0.23 \)                                         |
| Yes                                         | 30/168 (18)                       | 1.74 (0.68, 4.44)                        | 1.51 (0.55, 4.19)                                      |
| Food security<sup>d</sup>                   |                                   |                                          |                                                        |
| Secure                                      | 10/84 (12)                        | \( p = 0.85; 0.95 \)                     | \( p = 0.64; 0.88 \)                                   |
| Insecure                                    | 26/138 (19)                       | 1.72 (0.78, 3.77)                        | 1.83 (0.67, 5.00)                                      |
| Satisfaction with primary health care       | Mean = 4.21; SD = 0.70            |                                          |                                                        |
| provider, mean score<sup>e</sup>            |                                   |                                          |                                                        |

Note: 32 clients with incomplete data were excluded (5 missing viral loads; 13 missing pharmacy claim records). Clients with missing data were not significantly different \( (p > 0.05) \) from those retained for analyses on age, sex, refugee status, and time on HAART

<sup>a</sup> Unless otherwise noted

<sup>b</sup> Adjusted for age group, sex, refugee status, travel in past year, time to clinic, time on HAART, and time from HIV diagnosis to HAART start

<sup>c</sup> Factor modelled as a linear effect (common odds ratios presented)

<sup>d</sup> Item constructed from 3 questions, each measured on a 3-point Likert scale. An endorsement of “some of the time” or “all of the time” on any of the three questions was scored as “insecure”

<sup>e</sup> Item constructed from 2 questions, each measured on a 5-point Likert scale; ascending score was consistent with greater satisfaction
settings within one national program [35], the clinic setting itself may be the primary consideration. In the present setting, the access that refugees had to HIV services from a leading reference hospital was unusual in comparison to rural, dispersed or camp-based refugee groups. As laboratory monitoring for refugees is implemented according to national protocols, any differences in access among refugees ought to have been similar to routine differences between countries.

Factors identified from these data will help to locate those who might benefit from targeted interventions. To this end, additional counselling for men on HAART, support for those HAART clients who spend lengthy periods in transit to access routine care, and those who do not consistently refill their HAART prescriptions as monitored by the pharmacy, might be beneficial. Risk assessments for clients who may travel for extended periods could be implemented to ensure that consistent medication supply is available and contingency plans are in place. Use of mobile phones, either through training in using personal alarms, or more actively through a text-message intervention, may help to mitigate some of these challenges [36, 37]. Given the importance of the pharmacy-based adherence assessment, this measure should be formalised as a routine adherence indicator, be linked to medical records, and monitored. When the reported result is poor, this should

### Table 5 Association of self-change factors with unsuppressed viral load among refugees and local host community on HAART for ≥25 weeks in Kuala Lumpur, Malaysia (N = 222)

| Factor                                      | Prevalence ≥40 copies/mL, n/N (%) | p value, crude odds ratio (95 % CI) | p value, adjusted odds ratio (95 % CI)a |
|---------------------------------------------|----------------------------------|-------------------------------------|----------------------------------------|
| Adherence self-efficacy (self-rated ability to take medications as prescribed over previous month)b |                                  |                                     |                                        |
| Excellent                                   | 16/99 (16)                       | 1                                   | 1                                      |
| Good/very good                              | 14/105 (13)                      | 1.30 (0.74, 2.26)                   | 1.02 (0.56, 1.86)                      |
| Very poor/poor/fair                         | 6/18 (33)                        |                                      |                                        |
| Serostatus disclosure to partner            |                                  | p = 0.67                            | p = 0.77                               |
| No                                         | 4/22 (18)                        | 1                                   | 1                                      |
| Yes                                        | 17/120 (14)                      | 0.74 (0.22, 2.46)                   | 1.11 (0.29, 4.23)                      |
| No partner                                  | 15/80 (19)                       | 1.04 (0.31, 3.52)                   | 1.45 (0.38, 5.53)                      |
| Serostatus disclosure to family/friends     |                                  | p = 0.23                            | p = 0.49                               |
| No                                         | 10/81 (12)                       | 1                                   | 1                                      |
| Yes                                        | 26/141 (18)                      | 1.61 (0.73, 3.53)                   | 1.37 (0.56, 3.34)                      |
| Alcohol use, past month                     |                                  | p = 0.29                            | p = 0.69                               |
| Never                                      | 24/164 (15)                      | 1                                   | 1                                      |
| One or more times                           | 12/58 (21)                       | 1.52 (0.71, 3.28)                   | 0.83 (0.33, 2.06)                      |
| Use of illegal/harmful substances, past 6 months |                                  | p = 0.23                            | p = 0.83                               |
| No                                         | 32/208 (15)                      | 1                                   | 1                                      |
| Yes                                        | 4/14 (29)                        | 2.20 (0.65, 7.45)                   | 1.18 (0.27, 5.31)                      |
| Use of traditional medicines, past 6 months |                                  | p = 0.46                            | p = 0.75                               |
| No                                         | 29/188 (15)                      | 1                                   | 1                                      |
| Yes                                        | 7/34 (21)                        | 1.48 (0.57, 3.57)                   | 1.31 (0.47, 3.70)                      |
| No. of reported barriers to adherenceb     |                                  | p = 0.46                            | p = 0.89                               |
| 0                                          | 13/82 (16)                       | 1                                   | 1                                      |
| 1+                                         | 8/67 (12)                        | 1.13 (0.82, 1.56)                   | 1.03 (0.71, 1.49)                      |
| 3+                                         | 8/36 (22)                        |                                      |                                        |
| 5+                                         | 7/37 (19)                        |                                      |                                        |
| Knowledge of HIV and AIDS (% correct of 4 questions) |                                  | p = 0.15                            | p = 0.23                               |
| 0                                          | 1/18 (6)                         | 1                                   | 1                                      |
| 50+                                        | 35/204 (17)                      | 3.52 (0.45, 27.33)                  | 3.21 (0.37, 28.05)                     |

Note: 32 clients with incomplete data were excluded (5 missing viral loads; 13 missing pharmacy claim records). Clients with missing data were not significantly different (p > 0.05) from those retained for analyses on age, sex, refugee status, and time on HAART

a Adjusted for age group, sex, refugee status, travel in past year, time to clinic, time on HAART, time from HIV diagnosis to HAART start, and previous regimen switch
b Factor modelled as a linear effect (common odds ratios presented)
alert providers and trigger more advanced and expensive testing (e.g. viral loads).

This study had important limitations. Selection bias in the host community group may have affected our findings as response rates were high in both groups, but slightly lower in the host community. The host community study sample represented 6% of the target population. As non-participants may have possessed characteristics leading to bias, we compared routine socio-demographic indicators of the study sample with a simple random sample of 150 host community clients drawn from the clinic database. The random sample was statistically similar to the study sample on all socio-demographic indicators with the exception that ethnic Chinese clients were over-represented in the study sample, which could have introduced bias as ethnic Chinese Malaysians tend to have higher household incomes than other ethnic groups in Malaysia [38]. Given that the number of refugees recruited placed an upward limit on sample size and refugees only had routine access to the clinic one day per week, we accounted for the possibility that routine appointments may not have occurred during the study period by making additional efforts (by telephone and/or community representative) to contact refugees who had not been seen in the clinic two weeks prior to the close of recruitment.

This procedure facilitated a near-complete sample, while potentially introducing bias linked to these more intensive recruitment efforts. The cross-sectional design of the study limited our ability to draw any firm causal conclusions, and to accurately measure and classify longer-term viral suppression and adherence [39]. Lastly, as only a single study viral load sample was collected, outcomes may have been subject to sporadic viral escape, or “viral blips” leading to misclassification of the outcome [40–42]. Using C⩾500 copies/mL as an indicator of viral rebound [43], we queried results falling in the 40–499 copies/mL range among clients for whom the previous viral load was suppressed (<40 copies/mL) and found no evidence for any differences between groups (Table 2).

This study excluded asylum-seekers who began HAART in their country of origin and who may have been vulnerable to inferior outcomes given the possibility that their HAART was exhausted prior to gaining refugee status and becoming eligible for the national treatment program. These cases are routinely expedited and programs should facilitate pathways to treatment for this vulnerable group in the future. Strengths of the study included detailed

| Table 6 | Association of action state (adherence) factors with unsuppressed viral load among refugees and local host community on HAART for ≥25 weeks in Kuala Lumpur, Malaysia (N = 222) |
|---|---|---|---|
| Factor | Prevalence ≥40 copies/mL, n/N (%) | p value, crude odds ratio (95% CI) | p value, adjusted odds ratio (95% CI) |
| Adherence to medication schedule, self-reported | | | |
| Never, sometimes, half of the time, | 12/62 (19) | 1 | 1 |
| most of the time | | | |
| All of the time | 24/160 (15) | 0.74 (0.34, 1.58) | 0.90 (0.39, 2.08) |
| Adherence, visual analogue scale self-report, past month (%)b | | | |
| 0− | 5/13 (39) | 1 | 1 |
| 80− | 10/46 (22) | 0.50 (0.29, 0.86) | 0.65 (0.35, 1.19) |
| 95+ | 21/163 (13) | | |
| Adherence, dose-by-dose self-report, past 4 days (%) | | | |
| 0− | 4/9 (44) | 1 | 1 |
| 95+ | 32/213 (15) | 0.22 (0.06, 0.87) | 0.32 (0.06, 1.76) |
| Adherence, pharmacy refill schedule, HAART start or 24 monthsb | | | |
| 0− | 8/22 (36) | 1 | 1 |
| 80− | 12/53 (23) | 0.45 (0.28, 0.73) | 0.47 (0.27, 0.83) |
| 95+ | 16/147 (11) | | |
| Treatment interruptions of ≥1 day, self-report, past month | | | |
| None | 27/200 (14) | 1 | 1 |
| Any | 9/22 (41) | 4.44 (1.73, 11.38) | 2.77 (0.91, 8.43) |
| Unintentional underdosing | | | |
| No | 27/180 (15) | 1 | 1 |
| Yes | 9/42 (21) | 1.55 (0.67, 3.59) | 1.66 (0.65, 4.24) |

Note: 32 clients with incomplete data were excluded (5 missing viral loads; 13 missing pharmacy claim records). Clients with missing data were not significantly different (p > 0.05) from those retained for analyses on age, sex, refugee status, and time on HAART

a Adjusted for age group, sex, refugee status, travel in past year, time to clinic, time on HAART, time from diagnosis to HAART start, and previous regimen switch

b Factor modelled as a linear effect (single common odds ratio presented)
Table 7 Final multivariate model for factors associated with unsuppressed viral load among refugees and a local host community on HAART for ≥25 weeks in Kuala Lumpur, Malaysia

| Factor | p value, crude odds ratio (95% CI) | p value, adjusted odds ratio (95% CI) |
|--------|-----------------------------------|--------------------------------------|
| Age group (years) | | |
| 18– | p = 0.69 | p = 0.68 |
| 30– | 0.90 (0.52, 1.55) | 1.15 (0.60, 2.20) |
| 40+ | 1 | 1 |
| Refugee status | | |
| Host | p = 0.19 | p = 0.60 |
| Refugee | 1.59 (0.77, 3.28) | 1.28 (0.52, 3.14) |
| Sex | p = 0.04 | p = 0.05 |
| Male | 1 | 1 |
| Female | 0.41 (0.16, 1.04) | 0.39 (0.14, 1.05) |
| Time on HAART (years) | | |
| 0– | p = 0.79 | p = 0.53 |
| 1– | 1.06 (0.68, 1.67) | 1.17 (0.69, 1.96) |
| 2+ | | |
| Time from diagnosis to start (weeks) | | |
| 0– | p = 0.03 | p = 0.04 |
| 25– | 0.61 (0.39, 0.95) | 0.64 (0.41, 0.99) |
| 50+ | | |
| Temporary migration | | |
| ≥1 continuous month in past year | p < 0.001 | p = 0.002 |
| No | 1 | 1 |
| Yes | 4.21 (1.87, 9.50) | 4.12 (1.70, 9.99) |
| Average time to clinic (hours) | | |
| 0– | p = 0.01 | p = 0.02 |
| 1+ | 2.88 (1.14, 7.27) | 3.05 (1.09, 8.49) |
| Adherence, pharmacy refill schedule, HAART start or 24 months | | |
| 0– | p = 0.002 | p = 0.007 |
| 80– | 0.45 (0.28, 0.73) | 0.47 (0.27, 0.81) |
| 95+ | | |

Note: 32 clients with incomplete data were excluded (5 missing viral loads; 13 missing pharmacy claim records). Clients with missing data were not significantly different (p > 0.05) from those retained for analyses on age, sex, refugee status, and time on HAART.

a Adjusted for all factors in table except those denoted by c. A priori factors retained: age group, refugee status, and time on HAART. Factors excluded after one iteration: previous regimen switch.

b Factor modelled as a linear effect (single common odds ratio presented).

c Factor not included in the final model process due to presumptive role as mediator; other final model factors were not adjusted for these designated factors.

In summary, the high proportion of refugee and host clients attending this public sector clinic who achieved viral suppression supports the notion that providing HAART on an equitable basis to refugee and host community groups in this urban setting is both feasible and beneficial. Given the current global reduction of funding for HIV, the future sustainability of HAART for refugees needs to be critically assessed. The Malaysian national program fully subsidises first-line treatments for refugees, however, second-line treatments and virological monitoring are paid for by UNHCR. The concern is that national treatment programs that currently include refugees may opt to exclude them if funding continues to decline. If the goal of universal access to treatment is to be reached and the public health benefits of antiretroviral therapy are to be realized, refugees and other displaced or conflict-affected persons must be fully included in country and regional proposals and planning for HIV and AIDS.

Acknowledgments The authors wish to thank the study participants, the research team, and the clinic staff at Sungai Buloh Hospital. We are grateful to Ann Burton, Chunting Wong, Nadine Cornier, Herve Isambert, Neal Alexander, UNHCR Malaysia, Ian Weller and Nathan Ford for helpful input on the protocol and draft manuscript. We thank Maria L. Vivas for translation of the abstract into Spanish. Financial support was generously provided by the Canadian Institutes of Health Research (CIHR) Grant FRN 200710IDR, the Parkes Foundation, and the Department of Infectious Disease Epidemiology (London School of Hygiene and Tropical Medicine). We acknowledge the Economic Planning Unit, Prime Minister’s Office, Malaysia, for granting permission to conduct the research. Support from the MRC Tropical Epidemiology Group at the London School of Hygiene and Tropical Medicine made open access publication possible.

Conflict of interest None.

Open Access This article is distributed under the terms of the Creative Commons Attribution License which permits any use, distribution, and reproduction in any medium, provided the original author(s) and the source are credited.

References

1. UNAIDS. UNAIDS World AIDS Day Report. Geneva; 2011.
2. UN General Assembly. Convention relating to the Status of Refugees. In: Treaty series, vol 189, p 137 United Nations; 1951.
3. International Federation of Red Cross and Red Crescent Societies. World Disaster Report 2008: Focus on HIV and AIDS. Geneva 2008; p.118–41.
4. UNHCR, Southern African HIV Clinicians Society. Clinical guidelines for antiretroviral therapy management for displaced populations. Geneva: UNHCR; 2007.

5. Trippaya V, Botswana: refugees not entitled to same services as citizens. HIV/AIDS Policy Law Rev. 2005;10(3):27–8.

6. Mendelsohn JB, Schilpvoor M, Spiegel P, Ross DA. Adherence to antiretroviral therapy and treatment outcomes in conflict-affected and forcibly displaced populations: a systematic review. Confl Health. 2012;6(1):9.

7. UNHCR. Statistical Yearbook 2010; 2011. http://www.unhcr.org/statistics. Accessed 22 May 2012.

8. Ministry of Health Malaysia. National Strategic Plan on HIV/AIDS 2006–2010. Kuala Lumpur; 2006.

9. Ministry of Health Malaysia. National Strategic Plan on HIV/AIDS 2011–2015. Kuala Lumpur; 2011.

10. Chesney MA, Ickovics JR, Chambers DB, Gifford AL, Neidig J, Zwickl B, et al. Self-reported adherence to antiretroviral medications among participants in HIV clinical trials: the AACTG Adherence Instruments. AIDS Care. 2000;12(3):255–66.

11. Walsh JC, Mandalia S, Gazzard BG. Responses to a 1 month self-report on adherence to antiretroviral therapy are consistent with electronic data and virological treatment outcome. AIDS. 2002;16(2):269–77.

12. Ewart C. Social-action theory for a public health psychology. Am Psychol. 1991;46(9):931–46.

13. Schisterman EF, Cole SR, Platt RW. Overadjustment bias and unnecessary adjustment in epidemiologic studies. Epidemiology. 2009;20(4):488–95.

14. Cole SR, Herman MA. Faliability in estimating direct effects. Int J Epidemiol. 2002;31(1):163–5.

15. Kaufman JS, Maclehose RF, Kaufman S. A further critique of the analytic strategy of adjusting for covariates to identify biologic mediation. Epidemiol Perspect Innov. 2004;1(1):4.

16. Oyomopito R, Lee MP, Panuphuk P, Lim PL, Ditangco R, Zhou J, et al. Measures of site resourcing predict virologic suppression, immunologic response and HIV disease progression following highly active antiretroviral therapy (HAART) in the TREAT Asia HIV Observational Database (TAHOD). HIV Med. 2010;11(8):519–29.

17. McCarthy K, Chersich MF, Vearey J, Meyer-Rath G, Jaffer A, Simpvalo S, et al. Good treatment outcomes among foreigners receiving antiretroviral therapy in Johannesburg, South Africa. Int J STD AIDS. 2009;20:588–62.

18. Koboneka A, Wangisi J, Nabiryo C, Tsembé J, Kusemerewera S, Olop-Olupot P, et al. Clinical and immunological outcomes of a national paediatric cohort receiving combination antiretroviral therapy in Uganda. AIDS. 2008;22(18):2493–9.

19. Garang PG, Odoi RA, Kalyango JN. Adherence to antiretroviral therapy in conflict areas: a study among patients receiving treatment from Lacor Hospital, Uganda. AIDS Patient Care STDS. 2009;23(9):743–7.

20. Salami O, Buzu A, Nzeme C. High level of adherence to HAART among refugees and internally displaced persons on HAART in western equatorial region of Southern Sudan. J Int AIDS Soc. 2010;13(Supp 1):123.

21. Calvert H, Tu D, O’Brien DP, Ellum T, Mills C, Ford N, et al. HIV treatment in a conflict setting: outcomes and experiences from Bukavu, Democratic Republic of the Congo. PLoS Med. 2007;4(5):e129.

22. Lima V, Fernandes K, Rachlis B, Druyts E, Montaner J, Hogg R. Migration adversely affects antiretroviral adherence in a population-based cohort of HIV/AIDS patients. Soc Sci Med. 2009;68(6):1044–9.

23. Charurat M, Oyegunle M, Benjamin R, Habib A, Eze E, Ele P, et al. Patient retention and adherence to antiretrovirals in a large antiretroviral therapy program in Nigeria: a longitudinal analysis for risk factors. PLoS One. 2010;5(5):e10584.

24. Ochieng-Ooko V, Ochieng D, Sidle JE, Holdsworth M, Wools-Kaloustian K, Siika AM, et al. Influence of gender on loss to follow-up in a large HIV treatment programme in western Kenya. Bull World Health Organ. 2010;88(9):681–8.

25. Wasti SP, Simkhada P, Randall J, Freeman JV, van Teijlingen E. Factors influencing adherence to antiretroviral treatment in Nepal: a mixed-methods study. PLoS One. 2012;7(5):e35547.

26. Kitahata MM, Gange SJ, Abraham AG, Merriman B, Saag MS, Justice AC, et al. Effect of early versus deferred antiretroviral therapy for HIV on survival. N Engl J Med. 2009;360(18):1815–26.

27. Sturrat MJ, Remien RH, Smith A, Copeland OQ, Dolezal C, Krieger D. The role of HIV serostatus disclosure in antiretroviral medication adherence. AIDS Behav. 2006;10(5):483–93.

28. Cornell M, Myer L, Kaplan R, Bekker LG, Wood R. The impact of gender and income on survival and retention in a South African antiretroviral therapy programme. Trop Med Int Health. 2009;14(7):722–31.

29. Dou Z, Xu J, Jiao JH, Ma Y, Durako S, Yu L, et al. Gender difference in 2-year mortality and immunological response to ART in an HIV-infected Chinese population, 2006–2008. PLoS ONE. 2011;6(8):e22707.

30. Bekker LG, Myer L, Orrell C, Lawn S, Wood R. Rapid scale-up of a community-based HIV treatment service: programme performance over 3 consecutive years in Guguletu, South Africa. S Afr Med J. 2006;96(4):315–20.

31. Bisson GP, Gross R, Bellamy S, Chittams J, Hislop M, Regensberg L, et al. Pharmacy refill adherence compared with CD4 count changes for monitoring HIV-infected adults on antiretroviral therapy. PLoS Med. 2008;5(5):e109.

32. Grossberg R, Zhang YW, Gross R. A time-to-prescription refill measure of antiretroviral adherence predicted changes in viral load in HIV. J Clin Epidemiol. 2004;57(10):1107–10.

33. Thompson MA, Mugavero MJ, Amico KR, Cargill VA, Chang LW, Gross R, et al. Guidelines for improving entry into and retention in care and antiretroviral adherence for persons With HIV: evidence-based recommendations from an International Association of Physicians in AIDS Care panel. Ann Intern Med. 2012;156(11):817–33.

34. Spiegel PB, Checchi F, Colombo S, Paik E. Health-care needs of people affected by conflict: future trends and changing frameworks. Lancet. 2010;375(9711):341–5.

35. Dahab M, Charalambous S, Karstaedt AS, Fielding KL, Hamilton R, La Grange L, et al. Contrasting predictors of poor antiretroviral therapy outcomes in two South African HIV programmes: a cohort study. BMC Public Health. 2010;10:430.

36. Horvath T, Azman H, Kennedy GE, Rutherford GW. Mobile phone text messaging for promoting adherence to antiretroviral therapy in patients with HIV infection. Cochrane Database Syst Rev. 2012;3:CD009756.

37. Lester RT, Ritvo P, Mills EJ, et al. Effects of a mobile phone short message service on antiretroviral treatment adherence in Kenya (WelTel Kenya1): a randomised trial. Lancet. 2010;376(9755):1838–45.

38. Yusof ZA, Bhattachal D. Economic growth and development in Malaysia: policy making and leadership (Working paper no. 27). Washington DC: Commission on Growth and Development; 2008.

39. Tesoriero J, French T, Weiss L, Waters M, Finkelstein R, Agins B. Stability of adherence to highly active antiretroviral therapy over time among clients enrolled in the treatment adherence demonstration project. J Acquired Immune Defic Syndr (1999). 2003;33(4):484–93.

40. Zhang S, van Sighem A, Gras L, Reiss P, Smit C, Kroon F, et al. Clinical significance of transient HIV type-1 viraemia and
treatment interruptions during suppressive antiretroviral treatment. Antivir Ther. 2010;15(4):555–62.
41. Nettles RE, Kieffer TL, Kwon P, Monie D, Han Y, Parsons T, et al. Intermittent HIV-1 viremia (Blips) and drug resistance in patients receiving HAART. JAMA. 2005;293(7):817–29.
42. Fung IC, Gambhir M, van Sighem A, de Wolf F, Garnett GP. The clinical interpretation of viral blips in HIV patients receiving antiviral treatment: are we ready to infer poor adherence? J Acquired Immune Defic Syndr (1999). 2012;60(1):5–11.
43. Grennan JT, Loutfy MR, Su D, Harrigan PR, Cooper C, Klein M, et al. Magnitude of virologic blips is associated with a higher risk for virologic rebound in HIV-infected individuals: a recurrent events analysis. J Infect Dis. 2012;205(8):1230–8.