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Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

Smith R, Villanueva G, Probyn K, Sguassero Y, Ford N, Orrell C, Cohen K, Chaplin M, Leeflang MMG, Hine P

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[Diagnostic Test Accuracy Review]

Accuracy of measures for antiretroviral adherence in people living with HIV

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ABSTRACT

Background

Good patient adherence to antiretroviral (ART) medication determines effective HIV viral suppression, and thus reduces the risk of progression and transmission of HIV. With accurate methods to monitor treatment adherence, we could use simple triage to target adherence support interventions that could help in the community or at health centres in resource-limited settings.

Objectives

To determine the accuracy of simple measures of ART adherence (including patient self-report, tablet counts, pharmacy records, electronic monitoring, or composite methods) for detecting non-suppressed viral load in people living with HIV and receiving ART treatment.

Search methods

The Cochrane Infectious Diseases Group Information Specialists searched CENTRAL, MEDLINE, Embase, LILACS, CINAHL, African-Wide information, and Web of Science up to 22 April 2021. They also searched the World Health Organization (WHO) International Clinical Trials Registry Platform (ICTRP) and ClinicalTrials.gov for ongoing studies. No restrictions were placed on the language or date of publication when searching the electronic databases.

Selection criteria

We included studies of all designs that evaluated a simple measure of adherence (index test) such as self-report, tablet counts, pharmacy records or secondary database analysis, or both, electronic monitoring or composite measures of any of those tests, in people living with HIV and receiving ART treatment. We used a viral load assay with a limit of detection ranging from 10 copies/mL to 400 copies/mL as the reference standard. We created 2 × 2 tables to calculate sensitivity and specificity.



Data collection and analysis

We screened studies, extracted data, and assessed risk of bias using QUADAS-2 independently and in duplicate. We assessed the certainty of evidence using the GRADE method. The results of estimated sensitivity and specificity were presented using paired forest plots and tabulated summaries. We encountered a high level of variation among studies which precluded a meaningful meta-analysis or comparison of adherence measures. We explored heterogeneity using pre-defined subgroup analysis.

Main results

We included 51 studies involving children and adults with HIV, mostly living in low- and middle-income settings, conducted between 2003 and 2021. Several studies assessed more than one index test, and the most common measure of adherence to ART was self-report.

- **Self-report questionnaires** (25 studies, 9211 participants; very low-certainty): sensitivity ranged from 10% to 85% and specificity ranged from 10% to 99%.

- Self-report using a visual analogue scale (VAS) (11 studies, 4235 participants; very low-certainty): sensitivity ranged from 0% to 58% and specificity ranged from 55% to 100%.

- **Tablet counts** (12 studies, 3466 participants; very low-certainty): sensitivity ranged from 0% to 100% and specificity ranged from 5% to 99%.

- **Electronic monitoring devices** (3 studies, 186 participants; very low-certainty): sensitivity ranged from 60% to 88% and the specificity ranged from 27% to 67%.

- **Pharmacy records or secondary databases** (6 studies, 2254 participants; very low-certainty): sensitivity ranged from 17% to 88% and the specificity ranged from 9% to 95%.

- **Composite measures** (9 studies, 1513 participants; very low-certainty): sensitivity ranged from 10% to 100% and specificity ranged from 49% to 100%.

Across all included studies, the ability of adherence measures to detect viral non-suppression showed a large variation in both sensitivity and specificity that could not be explained by subgroup analysis. We assessed the overall certainty of the evidence as very low due to risk of bias, indirectness, inconsistency, and imprecision.

The risk of bias and the applicability concerns for patient selection, index test, and reference standard domains were generally low or unclear due to unclear reporting. The main methodological issues identified were related to flow and timing due to high numbers of missing data. For all index tests, we assessed the certainty of the evidence as very low due to limitations in the design and conduct of the studies, applicability concerns and inconsistency of results.

Authors' conclusions

We encountered high variability for all index tests, and the overall certainty of evidence in all areas was very low. No measure consistently offered either a sufficiently high sensitivity or specificity to detect viral non-suppression. These concerns limit their value in triaging patients for viral load monitoring or enhanced adherence support interventions.

PLAIN LANGUAGE SUMMARY

Are there good ways to find out if people living with HIV are taking their medicines every day?

The issue

For people with HIV, taking their HIV medicines every day (adherence), is vital to keep HIV under control. The best way to measure peoples' adherence to HIV medicines is with 'viral load testing', which tells us how much virus there is in the blood. Viral load testing is not available everywhere, such as in places where there is lack of funds. If we could measure adherence with a more readily available measure, this might help detect people who need more help with taking their medicines.

Aim of this review

To find out if simple measures of adherence can tell us whether people might not be taking their medication every day and might then have higher (detectable) viral loads. These people might be helped by extra viral load monitoring. This could then prevent them developing complications from HIV or passing HIV to other people.

What we found

We looked at 51 studies involving children and adults with HIV that happened between 2003 and 2021. These studies tested different ways to measure adherence, including surveys or rating scales filled out by patients, counting of patients' pills, pharmacy notes, or gadgets.

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All the measures we looked at did not help find patients who might not be taking their medications and who had higher viral loads. Different studies showed very different results. We could not explain these differences by whether the studies included children or adults, whether they were in richer or poorer areas, or what cut off they used to say if the viral load was high. This also meant that we could not combine the studies.

What are the implications of this review?

Based on the results, it is uncertain that simple measures of adherence to ART treatment can help find people living with HIV who may have a higher viral load. Still, there may be other values to trying to measure adherence that this review cannot show.

Reporting how current the evidence is

The evidence is up-to-date to 22 April 2021.



SUMMARY OF FINDINGS

Summary of findings 1. Summary of findings table 1: all index tests

Question: what is the diagnostic a suppression?	ccuracy of the different index tests	to measure adhere	ence to ART for dete	cting viral non-						
Population	HIV-positive children and adults who have been established on ART for longer than six months at the time of assessment									
Index tests	 electronic monitoring; 	tablet counts;pharmacy records or secondary database analysis, or both;								
Target condition	Viral non-suppression	Viral non-suppression								
Reference standard	Non-suppressed viral load, a 10 copies to 400 copies/mL	Non-suppressed viral load, as detected by nucleic acid testing technologies, ranging fro 10 copies to 400 copies/mL								
Action/clinical implications	<i>Low sensitivity:</i> failures to detect non-adherence.		adherent people incorrectly identified.							
	Consequences of false neg- atives: disease progression, resistance, transmission	Consequences of false neg- monitoring, patient inconvenience atives: disease progression,								
	Of greater clinical impor- tance									
Findings										
Type of index test	Studies and participants (viral non-suppression, %)	Sensitivity range (95% Cl range)	Specificity range (95% Cl range)	Certainty of the evidence (GRADE)						
Index test: self-report	25 studies	10% to 85%	10% to 99%	⊕ ###						
All participants	N = 9211	(0% to 91%)	(7% to 100%)	VERY LOW a,b,c,c						
≥95% adherence threshold	(1813, 20%)									
Any viral load threshold										
Index test: VAS	11 studies	0% to 58%	55% to 100%	⊕ ###						
All participants	N = 4235	(0% to 85%)	(46% to 100%)	VERY LOW c,d,e,f						
≥95% adherence threshold	(1479, 35%)									
Any viral load threshold										
Index test: tablet counts	12 studies	0% to 100%	5% to 99%	⊕### c,d,g,h						
All participants	N = 3466	(0% to 100%)	(2% to 100%)	VERY LOW						
≥95% adherence threshold	(504, 15%)									

Any viral load threshold

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Index test: pharmacy records or sec- ondary database All participants ≥95% adherence threshold Any viral load threshold	6 studies N = 2254 (552, 24%)	17% to 88% (11% to 92%)	9% to 95% (5% to 97%)	⊕### VERY LOW c,d,i,j
Index test: electronic monitoring	3 studies	60% to 88%	27% to 67%	⊕###
All participants	N = 186	(36% to 100%)	(11% to 80%)	VERY LOW
≥ 95% adherence threshold	(55, 30%)			k,l,m,n
Any viral load threshold				
Index test: composite measure	9 studies	10% to 100%	49% to 100%	⊕###
All participants	N = 1513	(4% to 100%)	(35% to 100%)	VERY LOW c,d,o,p
Different thresholds*	(407, 27%)			
Any viral load threshold				

^aDowngraded one level for limitations in the design and conduct of the studies due to patient selection (5 studies high risk, 7 studies unclear risk); administration and/or interpretation of the adherence measure (index test) (17 studies unclear risk); administration and/ or interpretation of the viral load test (reference standard) (6 studies unclear risk); and flow and timing of the study, including missing participant data (12 studies high risk, 2 studies unclear risk)

^bDowngraded one level for indirectness due to applicability concerns in relation to the population (2 studies high concern, 8 studies unclear concern); the measure of adherence used (index test) (3 studies unclear concern); and the viral load assessment used (reference standard) (6 studies unclear concern)

^cDowngraded two levels for inconsistency due to the extreme heterogeneity observed between the studies, both for sensitivity and specificity

^dThe evidence was not downgraded further due to imprecision as this was explained by the inconsistency observed between the studies. ^eDowngraded one level for limitations in the design and conduct of the studies due to patient selection (3 studies high risk, 3 studies unclear risk); administration and/or interpretation of the adherence measure (index test) (8 studies unclear risk); administration and/or interpretation of the adherence measure (index test) (8 studies unclear risk); administration and/or interpretation of the studies unclear risk); and flow and timing of the study, including missing participant data (6 studies high risk, 2 studies unclear risk)

^fDowngraded one level for indirectness due to applicability concerns in relation to the population (2 studies high concern, 2 studies unclear concern); the measure of adherence used (index test) (4 studies unclear concern); and the viral load assessment used (reference standard) (4 studies unclear concern)

^gDowngraded one level for limitations in the design and conduct of the studies due to patient selection (3 studies high risk, 4 studies unclear risk); administration and/or interpretation of the adherence measure (index test) (1 study high risk, 9 studies unclear risk); administration and/or interpretation of the viral load test (reference standard) (4 studies unclear risk); and flow and timing of the study, including missing participant data (7 studies high risk, 3 studies unclear risk)

^hDowngraded one level for indirectness due to applicability concerns in relation to the population (3 studies high concern, 2 studies unclear concern); the measure of adherence used (index test) (3 studies unclear concern); and the viral load assessment used (reference standard) (4 studies unclear concern).

ⁱDowngraded two levels for limitations in the design and conduct of the studies due to patient selection (3 studies high risk, 1 study unclear risk); administration and/or interpretation of the adherence measure (index test) (5 studies unclear risk); administration and/ or interpretation of the viral load test (reference standard) (2 studies unclear risk); and flow and timing of the study, including missing participant data (5 studies high risk)

jDowngraded one level for indirectness due to applicability concerns in relation to the population (3 studies unclear concern); the measure of adherence used (index test) (3 studies unclear concern) and the viral load assessment used (reference standard) (2 studies unclear concern).

^kDowngraded one level for limitations in the design and conduct of the studies due to patient selection (1 study unclear risk); administration and/or interpretation of the adherence measure (index test) (3 studies unclear risk); and flow and timing of the study, including missing participant data (1 study high risk)

^lDowngraded one level for indirectness due to applicability concerns in relation to the population (1 study high concern); and the measure of adherence used (index test) (3 studies unclear concern)

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^mDowngraded one level for inconsistency due to the heterogeneity observed between the studies, both for sensitivity and specificity ⁿDowngraded one level for imprecision due to small sample size. The evidence was not downgraded further due to imprecision as this was explained by the inconsistency observed between the studies.

°Downgraded one level for limitations in the design and conduct of the studies due to patient selection (1 study high risk, 4 studies unclear risk); administration and/or interpretation of the adherence measure (index test) (7 studies unclear risk); administration and/ or interpretation of the viral load test (reference standard) (4 studies unclear risk); and flow and timing of the study, including missing participant data (4 studies high risk, 2 studies unclear risk)

PDowngraded one level for indirectness due to applicability concerns in relation to the population (1 study high concern, 3 studies unclear concern); the measure of adherence used (index test) (3 studies unclear concern); and the viral load assessment used (reference standard) (3 studies unclear concern).



BACKGROUND

Target condition being diagnosed

Across all fields of medicine, low patient adherence is a barrier to realising the benefits of medication (Nieuwlaat 2014), and is associated with a higher mortality (Simpson 2006).

The World Health Organization (WHO) recommends provision of antiretroviral therapy (ART) to all people living with HIV, regardless of CD4 count (WHO 2016). At an individual level, ART reduces the risk of progression to AIDS or death, increases the likelihood of immune recovery, and reduces the risk of sexual transmission to seronegative partners. At a population level, widespread ART may reduce HIV incidence and offers a tool to end the HIV epidemic, as acknowledged within the Joint United Nations Programme on HIV/AIDS (UNAIDS) 90:90:90 target (UNAIDS 2014). This aims that, by 2020, 90% of all people living with HIV will know their HIV status, 90% of all people with diagnosed HIV infection will receive sustained ART, and 90% of all people receiving ART will have viral suppression.

With respect to HIV, for the individual and population level benefits of ART to be realized, patient adherence is essential. Adherence to ART is the primary determinant of viral suppression. In one metaanalysis of observational studies, only 62% of people receiving ART reported more than 90% adherence (Ortego 2011). Poor adherence increases the risk of transmission, accumulation of resistance mutations, disease progression, and death. Previous systematic reviews have identified treatments for people who achieve poor adherence to ART, thus illustrating the importance of measuring adherence in order to identify people who may benefit from such treatments (Horvath 2012; Kanters 2017; Rueda 2006).

One European consensus document defines "adherence to medications" as the process by which patients take their medication as prescribed. This term describes multiple behaviours (Vrijens 2012). There are four measurable subcategories of adherence to medications. These include:

- **initiation:** when a patient takes the first dose of a prescribed medication;
- **implementation:** the extent to which a patient's actual dosing corresponds to the prescribed dosing regimen, from initiation until the last dose;
- **persistence:** the length of time between initiation and the last dose;
- **discontinuation:** when a patient stops taking the prescribed medication.

Initiation and discontinuation are discontinuous (stop/start) measures, whereas implementation is a continuous measure. This precludes a single useful quantitative parameter to cover all three. Most research focuses on the implementation phase, that is: the extent to which a patient's actual dosing corresponds to the prescribed dosing regimen. The implementation component can be expressed via summary statistics which describe the implementation of a dosing regimen over a defined interval of time; for example, the proportion of days with the correct number of doses over a given period.

Although the implementation phase of adherence exists within a continuum from 0% to more than 100%, studies typically stratify

adherence into dichotomous variables of 'adherence' and 'nonadherence'. There are no specific consensus criteria for identifying these dichotomous categories of 'adherence' and 'non-adherence'. Traditionally, across fields of medicine, trials consider rates of less than 80% to represent non-adherence (Osterberg 2005). With respect to HIV, where non-adherence risks resistance mutations, trials have traditionally considered a threshold of greater than 95% as optimal (Paterson 2000), although more recent studies suggest a lower threshold be applied (Shilpa 2015). In practice, the level of adherence required to improve immune function and achieve viral suppression will vary by regimen and by prior history of viral suppression (Haberer 2017). For example, people with a longer-term history of viral suppression may be able to miss more doses without viral rebound (Lima 2010). Indeed, adopting lower adherence thresholds may not affect viral outcomes (Bezabhe 2016).

The definition of viral suppression is standard across guidelines, as an HIV ribonucleic acid (RNA) level below the lower limit of detection of available assays. However, the terminology for describing the absence of viral suppression is heterogeneous across the literature, incorporating concepts such as viral failure, incomplete response, viral rebound, viral blips, and low level viraemia. Table 1 summarizes the varying definitions of viral failure used internationally. Of note, the WHO definition incorporates an adherence support intervention before viral failure can be diagnosed.

Index test(s)

The index test is defined as any measures of adherence that could be utilized in resource-limited settings.

The WHO Guidelines on the use of antiretroviral drugs for treating and preventing HIV identify a need to "determine optimal ways to proactively monitor adherence and identify through simple triage those patients in greatest need of adherence support" (WHO 2016). In context, this relates to a public health approach which is "feasible on a large scale in resource-limited settings", with decentralization and integration of services such as task shifting. With respect to 'task shifting', the WHO recommends that trained and supervised community health workers can dispense ART between regular clinical visits (WHO 2016), and suggests that these workers adopt responsibility for monitoring patient's adherence (WHO 2017).

In relation to these considerations, this review focuses on measures of adherence that could be used at the 'community' or 'health centre' level as defined by a previous Cochrane Review (Kredo 2014), in a nomenclature reproduced in Table 2. As such, the measure:

- could be administered by trained volunteers, health assistants, nurse aides, and community health workers with a maximum of a few months of training;
- would not require infrastructure such as laboratories which are more commonly found at referral health centres or hospitals.

This would not preclude the use of the measure at higher levels of care. The following measures of adherence behaviour could meet these criteria:

- self-report;
- tablet counts;

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- pharmacy records or secondary database analysis, or both;
- electronic monitoring;
- composite measures of the above.

We describe these further below.

Self-report

The term 'self-report' involves a question, or set of questions, to which a patient responds. The mode of administration may be self-completion, or interviewer administered. The medium may be paper or electronic. There is no consensus taxonomy for self-report within the literature but, in broad terms, self-report questions may include:

- **behavioural questions:** questions asking patients to directly relate their adherence behaviour such as:
 - count-based questions: a specific day-by-day enquiry regarding missed doses. For example, *how many doses did you miss yesterday? The day before yesterday? Three days ago?* (Chesney 2000);
 - estimate-based questions: asking people to estimate how they took their treatment over a period of time. This might be based around a visual analogue scale (VAS), for example, mark the point along the line that most closely reflects how much of your HIV medications you have taken in the lastmonth? (Kabore 2015);
- **attitudinal questions:** these include questions asking patients about knowledge and beliefs, for example:
 - perceived barriers. *Did you ever miss a dose due to forgetfulness?;*
 - health beliefs. Sometimes, if you feel worse, do you stop taking your medications? (Knobel 2002);
 - self-efficacy. *How confident are you that you can take your medicines*? (EACS 2017).

The extent to which attitudinal questions are a valid form of assessment of adherence behaviour is unclear (Stirratt 2015), but inclusion of such questions will not preclude a questionnaire from this review. One previous systematic review covering all fields of medicine identified that the number of questions in self-report adherence measures ranged from one to 30, with a median of eight (Nguyen 2014). It is unlikely that a 30-item questionnaire could be termed 'simple triage', or be used by community health workers. Therefore, this review excludes self-report containing more than eight questions, or which the review authors deem to be prohibitively complex for use at community or health centre level.

Tablet counts

The provider counts the remaining tablets (or volume of liquid) in previously dispensed bottles and calculates an adherence percentage. This is based on expected versus actual tablets taken over a prescribed dispensing period. Counts may take place in clinic or be unannounced (in the form of telephone or home visits).

Pharmacy records or secondary database analysis, or both

Providers can use dates of prescription refills to calculate adherence measures. These can be broadly considered under three categories (Lam 2015):

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- Medication possession ratio (MPR): this measures the time for which a person possesses a supply of each medication class available, as a proportion of the time of eligibility for that medication. These measures are most commonly calculated over a three- to 12-month period but may be shorter or longer. We consider that the variability in methods used to calculate MPR will create challenges to meta-analysis. We will pool MPR data and use subgroup analyses to investigate heterogeneity introduced by different methods.
- **Tablet pick-up:** whether a person picks up all or most of their prescribed ARTs, categorizing people into either adherent or non-adherent based on specified criteria.
- **Continuous measures:** the time between prescription refills from the perspective of time gaps (periods of non-adherence) or consumption (medication availability, the days of supply/days between refills).

Electronic monitoring

Electronic monitoring devices use an embedded microprocessor to record the time and date a person opens a medication box. Health workers may access data from these devices by a cabled or cellular connection. Such devices use box opening as a proxy for medication ingestion, and as such may misclassify dose-taking behaviour. Expert opinion suggests that although devices are currently unaffordable to be used at scale in resource-limited settings, they are likely to become much cheaper in the future (Haberer 2017).

Composite measures of the above

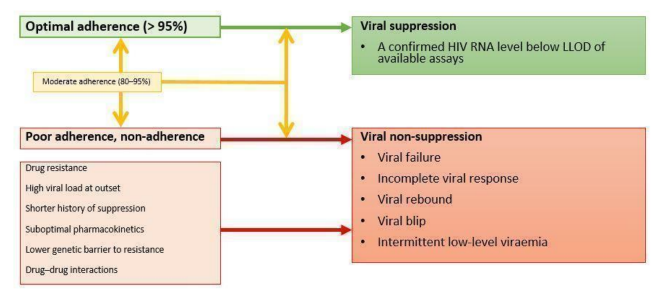
This describes the combination of two or more measures of adherence to give a more accurate impression than a single measure in isolation.

Clinical pathway

Under current national and international guidelines, when a person is diagnosed with HIV and linked to care, they are offered ART. After initiation of ART, people attend for clinical review. Clinicians may offer people more frequent clinical reviews in the months following initiation or during intercurrent illness, and less frequent clinical reviews once a person is established on and responding to therapy. Local guidelines and resources may also influence the frequency with which clinicians offer reviews. When people present for these reviews, clinicians may apply the index test (measures of adherence).

At these clinical appointments, people may also undergo viral load monitoring. This is the WHO 'gold standard' for confirmation of treatment response. The WHO also advises that viral load monitoring is the 'gold standard' for monitoring adherence (WHO 2016). Indeed, most elevated viral loads are the result of poor adherence (Bonner 2013). However, the relationship between adherence and viral load is not linear. Other patient and drug-related factors will influence viral suppression including drug resistance, viral load at outset of therapy, history of suppression, pharmacokinetics such as absorption, the genetic barrier to resistance offered by the regimen, and drug-drug interactions, as illustrated in Figure 1

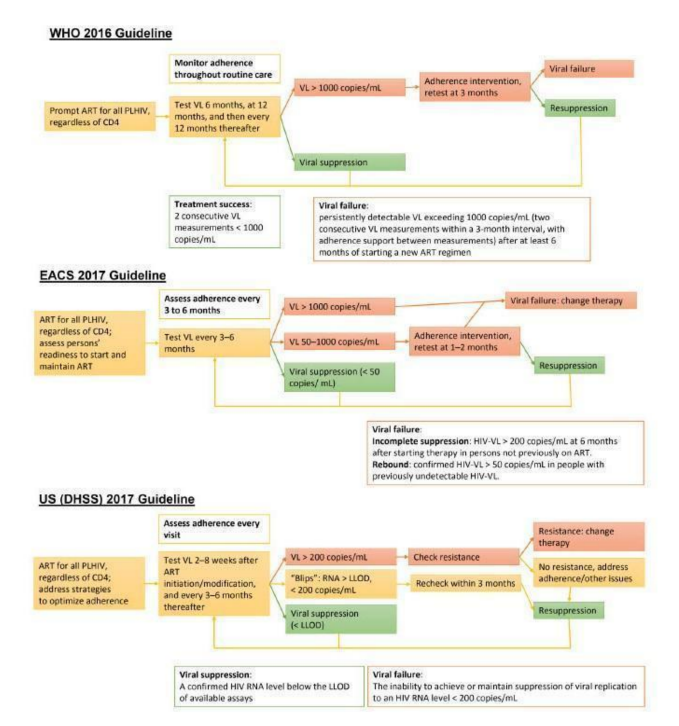
Figure 1. Patient and drug-related factors that influence viral suppression. Abbreviations: LLOD: lower limit of detection; RNA: ribonucleic acid.



The frequency with which providers offer viral load monitoring, and how the results are acted upon, will vary depending on resource availability. In resource-limited settings, a viral load measurement is recommended initially at six months, and then routinely 12 monthly thereafter, if suppressed (WHO 2016). In more resourcerich settings, viral load monitoring is more frequent, several early viral load measurements may be conducted during the first few months of ART, and routine monitoring is recommended every three to six months. Figure 2 demonstrates simplified clinical pathways as described across current guidelines.



Figure 2. Simplified clinical pathways described in current guidelines. Abbreviations: ART: antiretroviral therapy; LLOD: lower limit of detection; PLHIV: people living with HIV; VL: viral load; WHO: World Health Organization.



Prior test(s)

There are no prior tests that occur before the index test. However, elevated viral load measurements at previous visits and clinical findings may influence the decision to use measures of adherence. People may receive the index test (measures of adherence) more frequently when they have evidence of complications due to HIV. There are no differences according to age or gender.

Role of index test(s)

The index tests already in current clinical practice are used variably across the clinical pathways. If we could better understand which among all available index tests is more effective to determine viral non-suppression, this test could replace other tests within a given strategy. Additionally, an index test can be used as a triage test to enable more targeted viral load testing.



Alternative test(s)

Other measures of adherence include:

- directly observed therapy (DOTs): DOTs are often categorized as a 'direct' measure of adherence. A systematic review of DOTS showed no benefit to viral suppression of directly observed versus self-administered antiretroviral drugs (Ford 2009). We have excluded this because it does not represent 'simple triage', and there is overlap with adherence intervention;
- therapeutic drug monitoring (TDM): the absence of a drug with a long half-life gives objective evidence of recent nonadherence. We have excluded this as it is resource intensive, and generally does not give information about longer-term adherence. Other potential caveats include the following issues: serum drug levels may not reflect intracellular concentrations, therapeutic thresholds are unclear, and there is great inter- and intrapatient variability (DHHS 2017);
- pharmacological measures to quantify cumulative drug exposure: in response to the short-term nature of the information given by TDM, new measures are being evaluated to reflect drug intake and metabolism over a period of weeks to months. These include dried blood spot testing and hair testing (Castillo-Mancilla 2018). Dried blood spot testing has not yet been evaluated in relation to clinical outcomes in HIV treatment and requires deep freeze within the laboratory, which is unlikely to be viable in resource-limited settings. Hair sampling requires a person to have and be willing to part with hair, requires specialized laboratory services for processing, and is thus not likely to constitute 'simple triage'. Both these tests have future potential;
- provider clinical judgement: a small number of studies have investigated provider's subjective opinions on the likely adherence behaviour of their patient (Bangsberg 2001; Gross 2002). These represent complex qualitative assessments and are poorly amenable to meta-analysis. Therefore, we have excluded them from this review;
- **tablet identification tests:** the provider asks the patient to identify the tablets they have been prescribed from a selection of images of tablets (Parienti 2001). We have excluded these from this review because these test a patient's knowledge rather than implementation behaviour.

Adherence research has classified measures of adherence as objective and subjective, and direct and indirect. Although such terms appear in the literature, there is no formal taxonomy, and different authors may use the same term to describe different measures. Furthermore, the validity of applying these terms to HIV adherence research is questionable (Williams 2013). Therefore, we have avoided such terminology in this report.

Downstream impact of index test

The possible downstream consequences according to the four test accuracy categories, are as follows:

- **true positive (TP)** (the index test correctly identifies nonadherence to ART, and as such, detects a non-suppressed viral load): the clinician can perform additional tests (a viral load test, an increased frequency of viral load testing in future), or refer for an effective intervention (adherence support), or both;
- true negative (TN)(the index test correctly identifies adherence to ART, and as such, detects a suppressed viral load): the clinician

can continue the normal viral load testing schedule according to local practice;

- false positive (FP) (the index test misclassifies a person as nonadherent to ART, and fails to detect a suppressed viral load): the clinician may unnecessarily perform an additional test (viral load) or refer for an intervention (adherence support), or both. The blood test may cause the patient distress. The intervention may inconvenience the patient. Both test and intervention incur costs for the provider;
- false negative (FN) (the index test misclassifies a person as adherent to ART, and fails to detect a non-suppressed viral load): the patient will continue to receive the normal viral load testing schedule according to local practice. The patient has viral nonsuppression which has not been detected at that clinical review. This may lead to the consequences of transmission of HIV to other people, progression of HIV and the resultant morbidity and mortality, or development of drug resistance.

Systematic review evidence demonstrates that a number of interventions may ameliorate non-adherence, and either improve reported adherence (for example, text-messaging), or viral suppression (for example counselling or supporter interventions). The effects of interventions may be modest and wane over time (Kanters 2017).

Rationale

Although viral load testing is the reference standard measurement of treatment response, it is not universally available. In resourcelimited settings, viral load testing may either not be available or not feasible at a high frequency. In this context, the WHO has identified a demand to select through a simple triage those patients in greatest need of adherence support. This review seeks to recommend measures of antiretroviral adherence which could be used in resource-limited settings and to determine gaps in the current body of knowledge to inform future research.

OBJECTIVES

To determine the accuracy of simple measures of ART adherence (including patient self-report, tablet counts, pharmacy records, electronic monitoring, or composite methods) for detecting nonsuppressed viral load in people living with HIV and receiving ART treatment.

METHODS

Criteria for considering studies for this review

Types of studies

Inclusion criteria

- The study assesses index test(s) of interest (measures of adherence) at the time of a viral load measurement. We anticipated that most included studies would be conducted at a single time point. If studies were conducted at multiple time points, we included them if we were able to extract data from one or more specific time points, rather than aggregate or longitudinal scoring.
- The study reported data comparing the index test(s) of interest to viral load non-suppression, from which we could extract true positive, true negative, false positive, and false negative values.



• The study measured viral load using laboratory-based testing platforms.

We included observational studies (cross-sectional and prospective cohort studies) and randomized studies that provided sufficient data to create the 2 x 2 table to calculate sensitivity and specificity.

We also included studies which made within-study comparisons of the index test(s) of interest, but did not restrict inclusion only to such studies as we had anticipated few such studies existed.

Exclusion criteria

- The study did not report the lower limit of detection of the viral load assay used.
- The study used a viral load assay with a lower limit of detection greater than 400 copies/mL.
 - This is because most current laboratory assays have a lower limit of detection of less than 400 copies/mL, and there is greater clarity across literature that viral loads of less than 400 copies/mL reflect suppression.
- Studies using non-nucleic acid testing approaches.
 - An example of a non-nucleic acid approach is measurement of HIV reverse transcriptase activity; this is a surrogate for HIV viral load measurement.
- Studies using point-of-care tests.

We excluded retrospective studies or case-control study designs. These are more likely to be subject to bias, in particular, in relation to flow and timing: we anticipated that we would not be able to confirm that the timing of the adherence measure and the viral load was simultaneous, or that all patients receiving a given adherence measure would also receive a viral load.

There were no restrictions on minimal quality standard, sample sizes, or number of cases with viral non-suppression.

Participants

We included studies that recruited HIV-positive adults, adolescents, and children who had been established on ART for longer than six months at the time of assessment.

Index tests

The index tests included measures of adherence that could be utilized in resource-limited settings:

- self-report;
- tablet counts;
- pharmacy records or secondary database analysis, or both;
- electronic monitoring;
- composite measures of the above.

We categorized and analysed studies according to the above headings.

There are no specific consensus criteria for identifying adherence versus non-adherence. Studies may report different dichotomized thresholds between 'non-adherent' and 'adherent' in relation to measures of adherence that report implementation of a dosing regimen over a defined interval of time. For example:

- self-report: count- or estimate-based measures of percentage adherence over a given period;
- tablet counts: adherence percentage based on expected versus actual tablets taken over dispensing period;
- pharmacy records or secondary database analysis, or both;
- electronic monitoring: per cent of doses received as measured;
- composite measures of the above given a pooled percentage estimate.

All these measures estimate a percentage of time during which a patient takes the medication as prescribed. Typically, these studies then dichotomize 'adherence' and 'non-adherence', based on a percentage threshold.

Our definitions for the four test accuracy categories are as follows:

- true positive: the index test correctly identifies non-adherence to ART, and as such, detects a non-suppressed viral load;
- true negative: the index test correctly identifies adherence to ART, and as such, detects a suppressed viral load;
- false positive: the index test misclassifies a person as nonadherent to ART, and fails to detect a suppressed viral load;
- false negative: the index test misclassifies a person as adherent to ART, and fails to detect a non-suppressed viral load.

Target conditions

The target condition is viral non-suppression. We defined this as an HIV RNA level above the lower limit of detection of the assay used within the study in question.

Reference standards

We used a reference standard of non-suppressed viral load, as detected using nucleic acid testing technologies. This is any viral load which is above the lower limit of detection of the available assay. This varies between assays, ranging from 10 copies/mL to 400 copies/mL in those which are currently available.

Search methods for identification of studies

The Cochrane Infectious Diseases Group Information Specialists performed a comprehensive search to identify all relevant studies regardless of language or publication status (published, unpublished, in press, and ongoing).

Electronic searches

We searched the following databases from 2003 onwards, as these reflect more current ART regimens and viral load thresholds (WHO 2003).

- Cochrane Central Register of Controlled Trials (CENTRAL; the Cochrane Library);
- MEDLINE (PubMed);
- Embase (Ovid);
- Latin American and Caribbean Health Sciences Literature (LILACS);
- CINAHL (EBSCOhost);
- Africa-Wide Information (EBSCOhost); and
- Web of Science (Core Collection (Clarivate Analytics)).



We completed a preliminary search in July 2018, and adapted the search for other electronic databases. We updated the search in January 2020, and April 2021 (Appendix 1).

Searching other resources

We searched the WHO International Clinical Trials Registry Platform (ICTRP) and the ClinicalTrials.gov Clinical Study Register (www.clinicaltrials.gov).

We also screened reference lists of included studies and relevant reviews.

Data collection and analysis

Selection of studies

We merged studies identified by the keyword searches of different databases and removed duplicate reports. Review authors and collaborators from Cochrane Crowd independently scrutinized titles and abstracts from the electronic search to identify those which were potentially eligible (see summary of the protocol Appendix 2). Each study was screened by two independent authors or collaborators before inclusion or exclusion. Where there was disagreement, the first authors (PH/RS) adjudicated. We retrieved the full-text article for citations which the initial title and abstract screening identified as potentially eligible. Review authors (RS, KP and GV) independently assessed each full-text article for inclusion. We settled any discrepancies via discussion between review authors, and consultation with a third review author (PH) if further uncertainty remained. We identified studies according to the surname of the first author and year of publication.

Data extraction and management

We screened studies and extracted data independently, in duplicate. We also assessed the risk of bias and applicability concerns independently, in duplicate. We piloted the form on two studies from each adherence measure subtype, and finalized the form thereafter. We extracted data on the following characteristics.

- Author, publication year, study design (as defined by review author).
- Country of study and country income status (low-income, lower middle-income, upper middle-income, high-income), as defined by The World Bank Atlas method at the time of data extraction (World Bank 2018).
- Age and gender of included participants.
- HIV viral load assay used.
- Type of adherence assessment used, alone or as a composite measure, including:
 - for self-report: number of questions, modality (selfcompletion, interviewer-administered), question content (behavioural or attitudinal);
- for pharmacy data: MPR or tablet pick-up.
- Threshold used within the study for definition of dichotomization of optimal and suboptimal adherence.
- QUADAS-2 items (as detailed in Appendix 3).

Review authors (GV, KP, RS, YS) then extracted results and cross-tabulated data in 2×2 tables.

Assessment of methodological quality

We used the QUADAS-2 tool to appraise risk of bias and applicability (Whiting 2011). This includes four domains: patient selection, index test, reference standard, and flow and timing. To tailor the tool for our review, we changed signalling questions for each of the four domains. We have proposed an initial schema for operating the QUADAS-2 tool in Appendix 3. Review authors (KP,GV, YS and RS) independently piloted the form with two studies from each adherence measure subtype, and finalized the form thereafter. The final form for assessment of methodological quality is presented in Appendix 4 (changes to the previous form were highlighted). Risk of bias and applicability were completed for each of the included studies independently and in duplicate (GV, KP, YS). Disagreements and discrepancies were resolved by consultation between review authors, with the addition of a third author if agreement could not be reached.

We assessed the certainty of the evidence using the GRADE approach.

Statistical analysis and data synthesis

For all included studies, we used the data in the 2 x 2 tables (the binary test results cross-tabulated with the binary reference standard) to calculate sensitivity and specificity along with their 95% confidence intervals.

We have presented individual study results graphically by plotting estimates of sensitivities and specificities in a forest plot in order to facilitate visual assessment of variation in test accuracy. We used Review Manager 5 for these descriptive analyses (Review Manager 2014). For the main analysis, we used a 95% threshold or a binary (yes/ no) threshold. We chose this for the main analysis as it was commonly used and made clinical sense to the authors. However, we also conducted additional analysis using other thresholds (e.g. 80% adherence). We had planned to perform meta-analysis for each index test, but we were not able to pool the data due to the high heterogeneity among studies (see Investigations of heterogeneity). Since no pooling was not done due to heterogeneity, for the CI range we reported the lowest and the highest end of the confidence intervals across the studies that evaluated the same adherence test.

Comparing index tests

We made simple separate comparisons of summary estimates from alternative index tests. We did not encounter sufficient numbers of studies that made within-study paired comparisons of the same index tests to perform more detailed comparative analyses.

Investigations of heterogeneity

For each index test, we had planned to investigate heterogeneity by incorporating covariates to a hierarchical model in our metaanalysis. However, given that we were not able to perform meaningful meta-analysis, we instead investigated heterogeneity by subgrouping studies according to predefined categories.

These categories included:

- Setting, including income status:
 - this included the following World Bank income categories: low-income, lower- to middle-income, upper- to middleincome, and high-income economies.

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- Target population in study, as represented by child or adult.
- Lower limit of detection of viral load threshold used within the study.
- Subtype of adherence measure (e.g. by number and content of questions within self-report adherence measures).

These potential sources of heterogeneity were speculative. In addition, where stated within the results, we assessed those studies which had yielded tests with high sensitivity and specificity to assess whether there were shared characteristics.

Sensitivity analyses

We planned to conduct sensitivity analyses for each index test in which we would have excluded studies for which QUADAS-2 indicates areas of methodological concern. We also planned excluding studies in which more than four of the six QUADAS-2 domains were high risk. We also planned to assess the impact of risk of bias in relation to conduct and patient flow, and the impact of applicability in terms of whether the measure was likely to be applicable to a resource-limited setting. However, due to the high heterogeneity across/among studies, we decided against conducting sensitivity analyses as it was unlikely to influence any conclusions.

Assessment of reporting bias

We did not carry out formal assessment of publication bias because of the lack of sensitive and appropriate statistical methods for this review methodology.

Assessment of overall certainty of the evidence

We prepared a summary of findings table to present the main results and key information regarding the certainty of evidence assessed using the GRADE approach (Schünemann 2008; Schünemann 2020a; Schünemann 2020b). As recommended, we rated the certainty of evidence as high (not downgraded), moderate (downgraded by one level), low (downgraded by two levels), or very low (downgraded by more than two levels) based on four domains: risk of bias, indirectness, inconsistency, imprecision (but not publication bias). For each outcome, the certainty of evidence starts as high when there are high-quality observational studies (cross-sectional or cohort studies) that enrolled participants with

diagnostic uncertainty. When we found a reason for downgrading, we used our judgement to classify the reason as either serious (downgraded by one level) or very serious (downgraded by two levels) and recorded them in the footnotes. We applied the GRADE judgements for the GRADE domains as follows:

- Risk of bias: we used QUADAS-2 to assess the risk of bias.
- Indirectness: we used QUADAS-2 for concerns of applicability.
- Inconsistency: we carried out prespecified analyses to investigate potential sources of heterogeneity and downgraded when we could not explain the inconsistency in the accuracy estimates.
- Imprecision: we looked at the CIs of sensitivity and specificity estimates and at the unexplained heterogeneity of the results.
- Publication bias: we did not evaluate publication bias due to the lack of validated methods for diagnostic test accuracy reviews.

Equity

We did not plan our review with direct consideration of equity a priori. However, we recognize that there is a hypothetical potential for differences in adherence measure test accuracy between advantaged and disadvantaged populations. We considered the PROGRESS-Plus framework, which incorporates "Place of residence, Race/ethnicity/culture/language, Occupation, Gender or sex, Religion, Education, Socioeconomic status, Social capital and other characteristics ('Plus') such as sexual orientation, age and disability)" (Oxman 2009; Welch 2022). Our pre-planned subgroup analyses incorporated age and gender, and also income setting (which may relate to place of residence, and socioeconomic status).

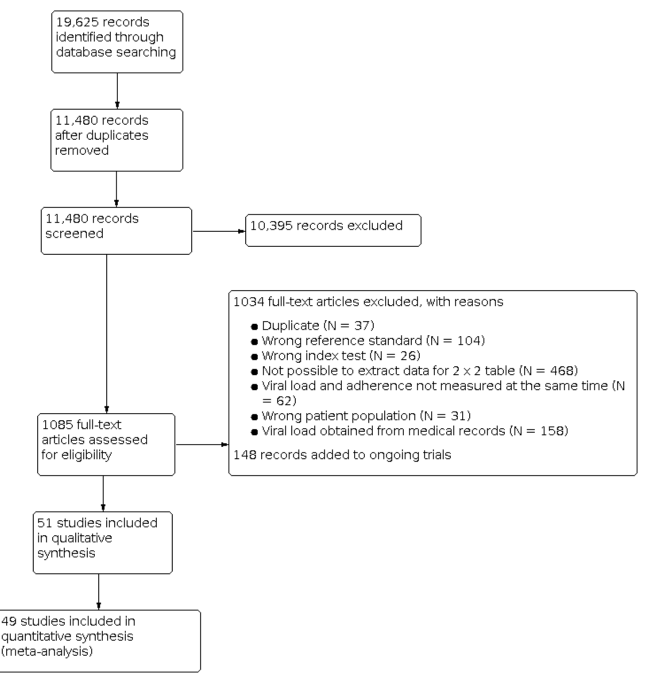
RESULTS

Results of the search

Figure 3 (Moher 2009) shows the flow of studies in the review. We identified 19,625 references from three searches: an initial search in July 2018, a repeat search in January 2020 and a last search in April 2021. From these, after removing duplicates, we identified 11,480 unique references. We considered 10,395 irrelevant to our review on initial screening. We screened 1085 references for inclusion, of which we excluded 1034 with reasons.



Figure 3. Study flow diagram



Fifty-one unique studies met our inclusion criteria and are included in the review.

Exclusions were mainly due to studies reporting duplicate data from another study (n = 37), wrong reference standard (n = 104), wrong index test (n = 26), insufficient data for the 2 x 2 table (n = 468), viral load and adherence not measured at the same time (n = 62), wrong patient population (n = 31), and viral load obtained from medical records (n = 158). We recorded the excluded studies and the reasons for their exclusion in Additional tables 3 to 9 (Table 3; Table 4; Table 5; Table 6; Table 7; Table 8; Table 9).

We also identified 148 ongoing trials (see Table 10).

Description of included studies

See Characteristics of included studies.

1. Self-report

1.1 Questionnaires

We identified 26 studies including 11,607 participants that used self-report questionnaires to estimate viral non-suppression (Avong 2015; Bajunirwe 2009; Coker 2015; Duarte 2015; Ekstrand 2010; El-Khatib 2010; Fokam 2017; Haberer 2011; Landes 2021; Mbengue 2019; McMahon 2013; Meya 2009; Mogosetsi 2018; Navarro 2014; Oette 2006; Orrell 2017; Paolillo 2017; Parker 2017; Pasquau 2018; Phillips 2019; Pulido 2009; Sangeda 2014; Segeral



2010; Segeral 2018; Tabb 2018; Zoufaly 2013). Three were RCTs (Coker 2015; Parker 2017; Pasquau 2018), nine cross-sectional (Avong 2015; El-Khatib 2010; Fokam 2017; Meya 2009; Phillips 2019; Segeral 2010; Segeral 2018; Tabb 2018; Zoufaly 2013), and the remaining 14 studies used a cohort design. A total of 9703 participants were included in the analysis, of whom 5640 had viral non-suppression. Four studies were conducted in children (Duarte 2015; Fokam 2017; Haberer 2011; Zoufaly 2013), two in mixed populations (Orrell 2017; Tabb 2018), and all others included only adults. Studies were conducted in different settings: 11 in low-income (Avong 2015; Bajunirwe 2009; Coker 2015; Haberer 2011; Landes 2021; McMahon 2013; Meya 2009; Sangeda 2014; Segeral 2010; Segeral 2018; Tabb 2018), three in lower-middleincome (Ekstrand 2010; Fokam 2017; Zoufaly 2013), five in uppermiddle-income (El-Khatib 2010; Mbengue 2019; Mogosetsi 2018; Orrell 2017; Phillips 2019), and six in high-income settings (Navarro 2014; Oette 2006; Paolillo 2017; Parker 2017; Pasquau 2018; Pulido 2009). One was conducted in a mixed setting (Duarte 2015).

Regarding the threshold for adherence, 20 studies used 100% or a binary threshold (adherent/non-adherent) (Bajunirwe 2009; Coker 2015; Duarte 2015; Fokam 2017; Landes 2021; Mbengue 2019; McMahon 2013; Meya 2009; Mogosetsi 2018; Oette 2006; Orrell 2017; Pasquau 2018; Paolillo 2017; Parker 2017; Phillips 2019; Pulido 2009; Sangeda 2014; Segeral 2010; Segeral 2018; Tabb 2018), four used 95% (Avong 2015; Haberer 2011; Ekstrand 2010; Zoufaly 2013), one used 90% (Navarro 2014), three used 80% (Haberer 2011; Phillips 2019; Segeral 2018), one used 60% (Navarro 2014).

For the viral load, two studies used 40 copies/mL (Landes 2021; Orrell 2017), 10 used 50 copies/mL (Bajunirwe 2009; Fokam 2017; Haberer 2011; Mogosetsi 2018; Navarro 2014; Oette 2006; Paolillo 2017; Pasquau 2018; Phillips 2019; Pulido 2009), four used 200 copies/mL (McMahon 2013; Parker 2017; Pasquau 2018; Zoufaly 2013), one used 250 copies/mL (Segeral 2018), and 12 used 400 copies/mL (Avong 2015; Bajunirwe 2009; Coker 2015; Duarte 2015; Ekstrand 2010; El-Khatib 2010; Mbengue 2019; Meya 2009; Phillips 2019; Sangeda 2014; Segeral 2010; Tabb 2018).

Please note that some of the studies used more than one threshold.

1.2 Visual analogue scale

We identified 14 studies including 5852 participants that used VAS to estimate viral non-suppression (Cerutti 2016; Cohen 2012; Dziva 2017; Ekstrand 2010; Gill 2010; Haberer 2011; Jiamsakul 2014; Labhardt 2012; McMahon 2013; Mbengue 2019; Meya 2009; Nelson 2010; Sangeda 2014; Segeral 2018). Seven were cohort studies, four were cross-sectional, two were RCTs and one was a prospective clinical trial. A total of 5151 participants were included in the analyses, of whom 2499 had viral non-suppression. Two studies were conducted in children (Dziva 2017; Haberer 2011), one in a mixed population (Labhardt 2012) and the remaining eleven studies in adults. Studies were conducted in different settings: seven in low-income (Dziva 2017; Haberer 2011; Labhardt 2012; McMahon 2013; Meya 2009; Sangeda 2014; Segeral 2010), three in lower-middle income (Cerutti 2016; Ekstrand 2010; Gill 2010), one in upper-middle income (Mbengue 2019), and three in mixed settings (Cohen 2012; Jiamsakul 2014; Nelson 2010).

Regarding the threshold for adherence, two studies used 100% (Sangeda 2014; Segeral 2010), 11 used 95% (Cerutti 2016; Cohen

2012; Dziva 2017; Ekstrand 2010; Gill 2010; Haberer 2011; Jiamsakul 2014; Labhardt 2012; McMahon 2013; Nelson 2010; Sangeda 2014), three used 90% (Mbengue 2019; Sangeda 2014; Segeral 2010), one used 80% (Haberer 2011), and one used a binary threshold (adherent/non-adherent) (Meya 2009).

For the viral load threshold one study used 40 copies/mL (Labhardt 2012), three used 50 copies/mL (Cohen 2012; Haberer 2011; Nelson 2010), one used 80 copies/mL (Cerutti 2016), one used 200 copies/ mL (McMahon 2013), and seven used 400 copies/mL (Dziva 2017; Gill 2010; Jiamsakul 2014; Mbengue 2019; Meya 2009; Sangeda 2014; Segeral 2010).

To note that some of the studies used more than one threshold.

2. Tablet counts

We identified 13 studies including 4899 participants that used tablet counts to estimate viral non-suppression (Apisarnthanarak 2010; Bonjoch 2006; Cerutti 2016; Coker 2015; Davies 2008; Gill 2010; Haberer 2011; Kitkungvan 2008; Mariana 2018; Moosa 2019; Okonji 2012; Orrell 2017; Sangeda 2014). Seven were cohort studies, three used a cross-sectional design, one was an RCT, and two were subanalyses of published RCTs. A total of 3808 participants were included in the analyses, of whom 2335 had viral nonsuppression. Nine studies included adults (Apisarnthanarak 2010; Bonjoch 2006; Cerutti 2016; Coker 2015; Gill 2010; Mariana 2018; Moosa 2019; Okonji 2012; Sangeda 2014), two included children (Davies 2008; Haberer 2011), and the other two were conducted in mixed populations (Kitkungvan 2008; Orrell 2017). Studies were conducted in different settings: four in low-income (Coker 2015; Haberer 2011; Okonji 2012; Sangeda 2014), four in lower-middleincome (Cerutti 2016; Gill 2010; Kitkungvan 2008; Mariana 2018), four in upper-middle-income (Apisarnthanarak 2010; Davies 2008; Moosa 2019; Orrell 2017), and one in high-income (Bonjoch 2006).

Regarding the threshold for adherence, one used 100% (Sangeda 2014), 12 used 95% (Apisarnthanarak 2010; Bonjoch 2006; Cerutti 2016; Coker 2015; Gill 2010; Haberer 2011; Kitkungvan 2008; Mariana 2018; Moosa 2019; Okonji 2012; Orrell 2017; Sangeda 2014), three used 90% (Bonjoch 2006; Davies 2008; Sangeda 2014), one used 85% (Sangeda 2014), two used 80% (Haberer 2011; Sangeda 2014), three used 75% (Apisarnthanarak 2010; Kitkungvan 2008; Sangeda 2014), one used 75% (Sangeda 2014), one used 65% (Sangeda 2014), one used 60% (Sangeda 2014), two used 55% (Kitkungvan 2008; Sangeda 2014), and one used 50% (Sangeda 2014).

For the viral load threshold, two used 40 copies/mL (Mariana 2018; Orrell 2017), four used 50 copies/mL (Apisarnthanarak 2010; Bonjoch 2006; Haberer 2011; Kitkungvan 2008), one used 80 copies/mL (Cerutti 2016), and the remaining six studies used 400 copies/mL.

To note that some of the studies used more than one threshold. We excluded Davies 2008 from the quantitative analysis because the adherence threshold was not relevant for the analysis of this review.

3. Pharmacy records or secondary databases

We identified seven studies including 2882 participants that used pharmacy records or other secondary databases to estimate viral non-suppression (Anude 2013; Hassan 2014; McMahon 2013; Messou 2011; Navarro 2014; Orrell 2017; Sangeda 2014). Six were

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cohort studies and one was a cross-sectional study. A total of 2449 were included in the analyses, of whom 1298 had viral nonsuppression. Five studies included adults (Anude 2013; McMahon 2013; Messou 2011; Navarro 2014; Sangeda 2014), and the other two studies included a mixed population. Studies were conducted in different settings: four in low-income (Hassan 2014; McMahon 2013; Messou 2011; Sangeda 2014), one in lower-middle-income (Anude 2013), one in upper-middle-income (Orrell 2017), and one in high-income (Navarro 2014).

Regarding the thresholds used to determine adherence, two studies used 100% (McMahon 2013; Sangeda 2014), six used 95% (Anude 2013; Hassan 2014; McMahon 2013; Messou 2011; Orrell 2017; Sangeda 2014), two used 90% (Navarro 2014; Sangeda 2014), one used 85% (Sangeda 2014), two used 80% (Messou 2011; Sangeda 2014), one used 75% (Sangeda 2014), one used 70% Sangeda 2014), two used 65% (Messou 2011; Sangeda 2014), two used 60% (Navarro 2014, Sangeda 2014), one used 55% (Sangeda 2014), one used 55% (Sangeda 2014), and two used 50% (Messou 2011; Sangeda 2014).

To note that some of the studies used more than one threshold.

For the viral load threshold, one used 40 copies/mL (Orrell 2017), one used 50 copies/mL (Navarro 2014), one used 200 copies/mL (McMahon 2013), one used 300 copies/mL (Messou 2011), and three used 400 copies/mL (Anude 2013; Hassan 2014; Sangeda 2014).

4. Electronic monitoring devices

We identified five studies including 475 participants that used electronic monitoring devices to estimate viral non-suppression (Evans 2016; Farley 2003; Gill 2010; Haberer 2011; Orrell 2017). All were cohort studies. A total of 392 participants were included in the analysis, of whom 92 had viral non-suppression. Two studies included children (Farley 2003; Haberer 2011), two studies included adults (Evans 2016; Gill 2010), and one study included both children and adults (Orrell 2017). Studies were conducted in different settings; one in low-income (Haberer 2011), one in lower-middle-income (Gill 2010), two in upper-middle-income (Evans 2016; Orrell 2017), and one in high-income (Farley 2003).

Regarding the thresholds for adherence, three studies used 95% (Evans 2016; Gill 2010; Haberer 2011), and four studies used 80% (Evans 2016; Farley 2003; Haberer 2011; Orrell 2017). To note that some of the studies used more than one threshold.

For the viral load threshold, one study used 40 copies/mL (Orrell 2017), one study used 50 copies/mL (Haberer 2011), and three studies used 400 copies/mL (Evans 2016; Farley 2003; Gill 2010).

5. Composite measure of adherence

We identified nine studies including 1901 participants that used composite measures of adherence to estimate viral nonsuppression (Jayaweera 2003; Mbengue 2019; McMahon 2013; Mutwa 2014; Orrell 2003; Ortega 2004; Parienti 2010; Segeral 2010; Spire 2008). Three studies were cross-sectional and six studies used a cohort design. A total of 1513 participants were included in the analysis, of whom 858 had viral non-suppression. Only one study included children (Mutwa 2014), and one study did not report on participants age (Jayaweera 2003). All the other studies included adults. Studies were conducted in different settings; four in lowincome (McMahon 2013; Mutwa 2014; Segeral 2010; Spire 2008), two in upper-middle-income (Mbengue 2019; Orrell 2003), and three in high-income (Jayaweera 2003; Ortega 2004; Parienti 2010).

Regarding the thresholds for adherence, one study used 100% (Segeral 2010), three studies used 95% (Mutwa 2014; Orrell 2003; Parienti 2010), one used 90% (Ortega 2004), one used 80% (Parienti 2010), one used 70% (Parienti 2010), and four used a binary threshold (adherent/ non-adherent of high/low) without providing exact details on percentage (Jayaweera 2003; Mbengue 2019; McMahon 2013; Spire 2008).

For the viral load threshold, two studies used 40 copies/mL (Mutwa 2014; Spire 2008), one study used 50 copies/mL (Parienti 2010), one study used 200 copies/mL (McMahon 2013), and six studies used 400 copies/mL (Jayaweera 2003; Mbengue 2019; Orrell 2003; Ortega 2004; Parienti 2010; Segeral 2010).

To note that some of the studies used more than one threshold

Methodological quality of included studies

We evaluated these studies for risk of bias in the following QUADAS-2 domains (Whiting 2011): participant selection, index test, reference standard, and participant flow. Figure 4 and Figure 5 provide a summary of the overall methodological quality for included studies.

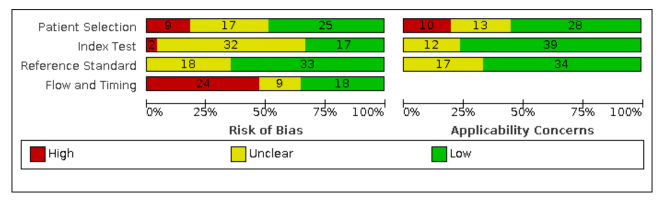


Figure 4. Risk of bias and applicability concerns graph: review authors' judgements about each domain presented as percentages across included studies

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

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	R	isk o	of Bia	is	Арр	licab	ility	Concerns
	Patient Selection	Index Test	Reference Standard	Flow and Timing	Patient Selection	Index Test	Reference Standard	
Anude 2013	?	?	Ŧ	•	•	?	Ŧ	
Apisarnthanarak 2010	•	?	?	?	•	•	?	
Avong 2015	Ŧ	?	Ŧ	•	•	•	Ŧ	
Bajunirwe 2009	Ŧ	?	Ŧ	•	•	•	Ŧ	
Bonjoch 2006	Ŧ	?	?	•	?	•	?	
Cerutti 2016	Ŧ	?	Ŧ	Ŧ	•	Ŧ	Ŧ	
C o hen 2012	?	•	Ŧ	?	•	?	Ŧ	
Coker 2015	?	•	Ŧ	•	•	Ŧ	Ŧ	
Davies 2008	Ŧ	•	?	•	•	Ŧ	Ŧ	
Duarte 2015	Ŧ	•	?	?	•	Ŧ	?	
Dziva 2017	Ŧ	?	Ŧ	•	•	Ŧ	Ŧ	
Ekstrand 2010	?	?	Ŧ	Ŧ	?	Ŧ	Ŧ	
El-Khatib 2010	•	?	Ŧ	•	?	Ŧ	Ŧ	
Evans 2016	?	?	Ŧ	Ŧ	•	?	Ŧ	
Farley 2003	?	?	Ŧ	•	•	?	Ŧ	
Fokam 2017	?	?	Ŧ	Ŧ	?	Ŧ	Ŧ	
Gill 2010	Ŧ	?	Ŧ	Ŧ	•	?	Ŧ	
Haberer 2011	Ŧ	?	Ŧ	•	•	?	Ŧ	
Hassan 2014	Ŧ	•	?	Ŧ	•	?	?	
Jayaweera 2003	?	?	Ŧ	•	•	?	Ŧ	
Jiamsakul 2014	?	Ŧ	?		•	•	?	
Kitkungvan 2008	•	?	Ŧ	?	•	•	Ŧ	
Labhardt 2012	•	?	?	•	•	•	?	
Landes 2021	?	Ŧ	Ŧ	•	?	•	Ŧ	
Mariana 2018	Ŧ	•	?	?	•	?	?	
Mbenaue 2019	Ŧ	?	?			A	?	

Figure 5. Risk of bias and applicability concerns summary: review authors' judgements about each domain for each included study



Figure 5. (Continued)

	-	-	-	i 🕶 i	I I	-	-	-		
Mbengue 2019	Ŧ	?	?	•		Ŧ	•	?		
McMahon 2013	•	?	Ŧ	•		?	•	Ŧ		
Messou 2011	•	?	Ŧ	•		?	•	Ŧ		
Meya 2009	Ŧ	Ŧ	Ŧ	Ŧ		Ŧ	•	Ŧ		
Mogosetsi 2018	•	Ŧ	Ŧ	•		?	•	Ŧ		
Moosa 2019	?	Ŧ	Ŧ	•		•	•	Ŧ		
Mutwa 2014	Ŧ	?	Ŧ	•		?	•	Ŧ		
Navarro 2014	Ŧ	Ŧ	?	•		•	•	?		
Nelson 2010	Ŧ	Ŧ	?	?		Ŧ	•	?		
0ette 2006	Ŧ	?	?	•		Ŧ	•	?		
0k o nji 2012	?	?	Ŧ	•		•	•	Ŧ		
Orrell 2003	Ŧ	?	?	?		Ŧ	•	?		
Orrell 2017	Ŧ	?	Ŧ	•		?	•	Ŧ		
Ortega 2004	Ŧ	?	?	•		?	?	?		
Paolillo 2017	?	Ŧ	?	?		Ŧ	Ŧ	?		
Parienti 2010	?	Ŧ	?	?		?	?	?		
Parker 2017	•	?	Ŧ	•		•	•	Ŧ		
Pasquau 2018	Ŧ	Ŧ	?	•		Ŧ	Ŧ	?		
Phillips 2019	Ŧ	?	Ŧ	•		Ŧ	•	Ŧ		
Pulido 2009	Ŧ	Ŧ	Ŧ	•		Ŧ	•	Ŧ		
San ged a 2014	•	?	?	•		Ŧ	•	?		
Segeral 2010	?	?	Ŧ	•		•	•	Ŧ		
Segeral 2018	Ŧ	Ŧ	Ŧ	•		Ŧ	?	Ŧ		
Spire 2008	?	Ŧ	Ŧ	•		•	Ŧ	Ŧ		
Tabb 2018	?	?	Ŧ	•		?	Ŧ	Ŧ		
Zoufaly 2013	Ŧ	?	Ŧ	•		Ŧ	•	Ŧ		
Brigh ? Unclear Brow										
• -	-					-			 	

Patient selection (QUADAS-2, domain 1)

In the patient selection domain, we considered nine studies at high risk of bias due to strict inclusion criteria with a risk for inappropriate exclusions. Seventeen studies were rated as unclear risk as it was unclear if a consecutive or random sample of patients was enrolled or they had few details on patient selection criteria, or both. The remaining studies (n = 25) were considered as low risk.

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Regarding applicability, 10 studies were rated at high concern and thirteen studies were considered at unclear concern due to patient sampling (e.g. studies only included people with a history of low adherence or viral non-suppression, people receiving support to increase adherence). The remaining studies (n = 28) were rated as low concern.

Index test (QUADAS-2, domain 2)

In the index test domain, we judged two studies at high risk of bias; in one study adherence was dichotomized as \geq 90% or < 90% as this threshold explained the largest amount of variability in the outcome (Davies 2008), and in another study there was no information on how the index test was conducted (Mariana 2018). Most studies (n = 32) were rated as unclear because there was no information on whether the index tests were interpreted without knowledge of the results of the reference standard, or if prespecified thresholds were used. The remaining studies were rated as low risk as they used a validated scale to measure adherence.

Regarding applicability, 12 studies were rated as unclear concern due to their potential complexity (e.g. long questionnaires, composite measure requiring calculations, costs of electronic monitoring devices). The remaining 39 studies were judged as low concern.

Reference standard (QUADAS-2, domain 3)

In the reference standard domain, we rated 18 studies at unclear risk of bias as there was no information to assess whether the reference standard results were interpreted without knowledge of the results of the index test or the test used to determine viral load was not described. The remaining studies (n = 33) were rated as low risk.

Regarding applicability, 17 studies were judged as unclear as there were no details on the assay used to determine viral load. The remaining studies (n = 34) were rated as low concern.

Flow and timing (QUADAS-2, domain 4)

In the flow and timing domain, we judged 24 studies to be at high risk of bias. The main reason was the high number of missing participants for the analysis (higher than 20%). Nine studies were considered at unclear concern as the interval between the adherence measure and the viral load measurement was not clear, or there was no information on the type of assay used.

Findings

The main findings are presented in Summary of findings 1. Across all included studies, the ability of measures of adherence to detect viral non-suppression showed a large variation in both sensitivity and specificity that could not be explained by subgroup analysis.

1. Self-report

1.1 Questionnaires

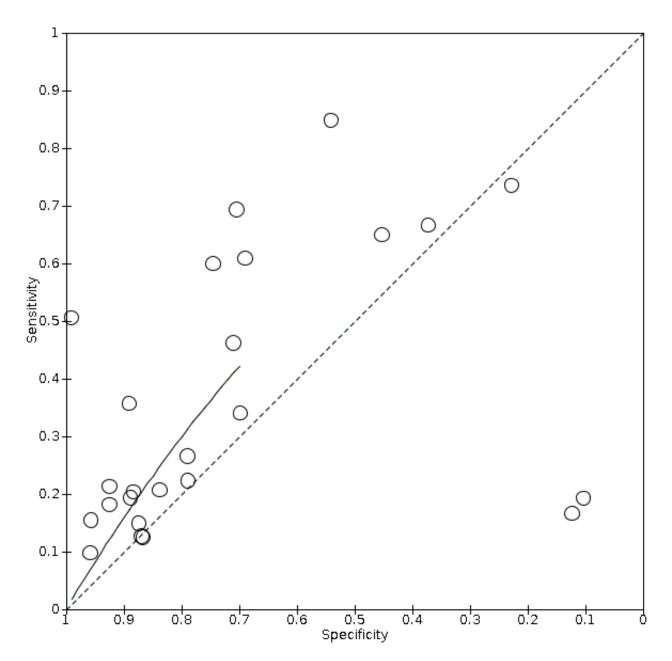
Studies using self-report questionnaires for the detection of viral non-suppression showed a large variation in both sensitivity and specificity that we could not explain by subgroup analyses or narrative review. See 'Summary of findings' table 2 in Appendix 5.

For the main analysis (Figure 6; Figure 7), we selected studies using a 100% adherence threshold (or binary yes/no) and studies using a 95% adherence threshold (25 studies with 9211 participants, of whom 1813 had viral non-suppression). The variation in point estimates for sensitivity ranged from 5% to 91% and for specificity ranged from 10% to 100%. The certainty of the evidence was assessed as very low due to risk of bias, indirectness, and inconsistency. Due to the high heterogeneity, we did not pool these studies.

Figure 6. Self-report questionnaires, various thresholds* [main analysis] *cut-off used was either ≥ 95% or 100%

Study	ТР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specifi	city (95% Cl)
Avong 2015	84	185	15	218	0.85 [0.76, 0.91]	0.54 [0.49, 0.59]	-	-
McMahon 2013	25	105	9	31	0.74 [0.56, 0.87]	0.23 [0.16, 0.31]		
Phillips 2019	34	26	15	62	0.69 [0.55, 0.82]	0.70 [0.60, 0.80]	—•·	
Fokam 2017	28	64	14	38	0.67 [0.50, 0.80]	0.37 [0.28, 0.47]		F .
Haberer 2011	13	29	7	24	0.65 [0.41, 0.85]	0.45 [0.32, 0.60]		
Parker 2017	106	458	68	1017	0.61 [0.53, 0.68]	0.69 [0.67, 0.71]		
Puli do 2009	9	27	6	79	0.60 [0.32, 0.84]	0.75 [0.65, 0.82]	-	
Coker 2015	45	2	44	224	0.51 [0.40, 0.61]	0.99 [0.97, 1.00]		
Pasquau 2018	6	51	- 7	125	0.46 [0.19, 0.75]	0.71 [0.64, 0.78]	_	-
Bajunirwe 2009	10	16	18	131	0.36 [0.19, 0.56]	0.89 [0.83, 0.94]	_ _	-
Tabb 2018	31	41	60	95	0.34 [0.24, 0.45]	0.70 [0.61, 0.77]		
Mbengue 2019	13	24	36	90	0.27 [0.15, 0.41]	0.79 [0.70, 0.86]		
Zoufaly 2013	27	23	94	86	0.22 [0.15, 0.31]	0.79 [0.70, 0.86]		
Oette 2006	13	11	48	136	0.21 [0.12, 0.34]	0.93 [0.87, 0.96]		-
Segeral 2018	28	192	107	990	0.21 [0.14, 0.29]	0.84 [0.82, 0.86]	+	
Meya 2009	10	52	39	395	0.20 [0.10, 0.34]	0.88 [0.85, 0.91]		
Paolillo 2017	46	33	192	264	0.19 [0.15, 0.25]	0.89 [0.85, 0.92]	+	-
Lan de s 2021	16	321	67	37	0.19 [0.11, 0.29]	0.10 [0.07, 0.14]		
Ekstrand 2010	10	11	45	136	0.18 [0.09, 0.31]	0.93 [0.87, 0.96]		-
Orrell 2017	1	143	5	20	0.17 [0.00, 0.64]	0.12 [0.08, 0.18]		
Duarte 2015	19	10	104	224	0.15 [0.10, 0.23]	0.96 [0.92, 0.98]		-
Segeral 2010	11	23	63	160	0.15 [0.08, 0.25]	0.87 [0.82, 0.92]		-
Sangeda 2014	7	14	48	93	0.13 [0.05, 0.24]	0.87 [0.79, 0.93]		-
Mogosetsi 2018	1	12	7	78	0.13 [0.00, 0.53]	0.87 [0.78, 0.93]		
El-Khatib 2010	10	32	92	740	0.10 [0.05, 0.17]	0.96 [0.94, 0.97]	_ 	
							0 0.2 0.4 0.6 0.8 1 0 0.2 0	40.60.81

Figure 7. Summary ROC Plot of 1 [Main analysis] Self-report, various thresholds*. *cut-off used was either ≥ 95% or 100%



We explored heterogeneity by looking at adherence threshold, type of questionnaire, population, viral load threshold used, and setting. None of these prespecified subgroups could explain the heterogeneity observed.

- Adherence threshold (Appendix 6) -100% cut-off (21 studies, N = 8204): sensitivity ranged from 18% to 85% and specificity ranged from 10% to 99%; **95% cut-off** (4 studies, N = 1007): sensitivity ranged from 18% to 85% and specificity ranged from 45% to 93%.
- Type of questionnaire (Appendix 6) 1 item only (12 studies, N = 4997): sensitivity ranged from 10% to 74% and specificity ranged from 10% to 96%; 2 to 4 items (8 studies, N = 1922):

sensitivity ranged from 13% to 69% and specificity ranged from 70% to 99%; **5 or more items** (5 studies, N = 2292): sensitivity ranged from 21% to 85% and specificity ranged from 54% to 84%.

- **Population** (Appendix 6) children (4 studies, N = 804): sensitivity ranged from 15% to 67% and specificity ranged from 37% to 96%; adults (19 studies, N = 8011): sensitivity ranged from 10% to 85% and specificity ranged from 10% to 99%.
- Viral load threshold (Appendix 6) 40 to 50 copies/mL (11 studies, N = 2290): sensitivity ranged from 14% to 69% and specificity ranged from 10% to 93%; 200 to 400 copies/mL (13 studies, N = 6664): sensitivity ranged from 10% to 85% and specificity ranged from 23% to 99%.



• Setting (Appendix 6) - low-income (11 studies, N = 4135): sensitivity ranged from 13% to 85% and specificity ranged from 10% to 99%; lower-middle-income (3 studies, N = 576): sensitivity ranged from 18% to 67% and specificity ranged from 37% to 93%; upper-middle-income (5 studies, N = 1141): sensitivity ranged from 10% to 69% and specificity ranged from 12% to 96%; high-income (5 studies, N = 2702): sensitivity ranged from 19% to 61% and specificity ranged from 69% to 93%.

In addition to prespecified subgroup analyses, we further explored the highest and lowest performing studies using self-report questionnaires, as this was the largest group of studies. We aimed to identify characteristics in common that we had not previously considered when developing the protocol. We were not able to identify any shared characteristics between those studies having or showing the highest or the lowest sensitivity estimates.

Three studies also looked at the diagnostic accuracy of a 80% adherence threshold (Haberer 2011; Phillips 2019; Segeral 2010; N = 1527; Appendix 6). Sensitivity ranged from 8% to 41% and specificity ranged from 81% to 97%.

One study was excluded from the quantitative analyses as the adherence thresholds used were not relevant for our analyses (60% and 90%) (Navarro 2014).

1.2 Visual analogue scale

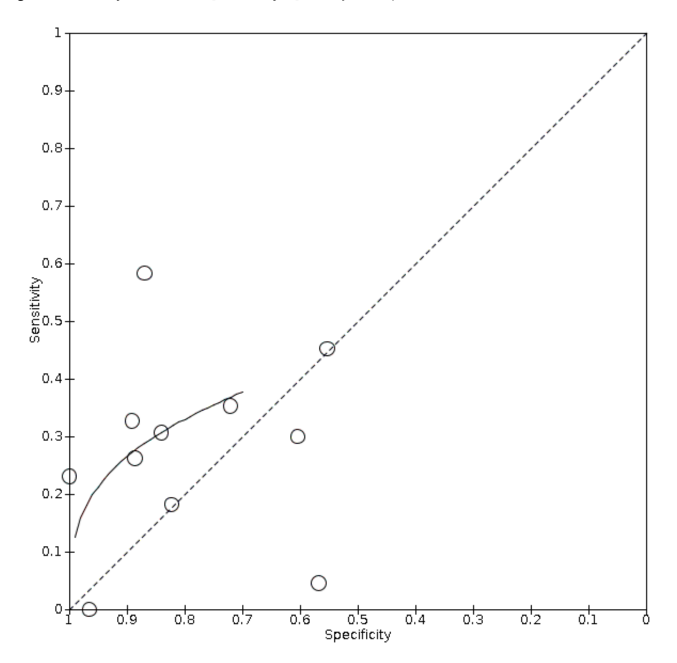
Studies using visual analogue scale questionnaires for the detection of viral non-suppression showed a large variation in both sensitivity and specificity that we could not explain by subgroup analyses See 'Summary of findings' table 3 in Appendix 5.

Eleven studies including 4235 participants (of whom 1479 had viral non-suppression) used a 95% adherence threshold (Cerutti 2016; Cohen 2012; Dziva 2017; Ekstrand 2010; Gill 2010; Haberer 2011; Jiamsakul 2014; Labhardt 2012; McMahon 2013; Nelson 2010; Sangeda 2014), and were included in the main analysis (Figure 8; Figure 9). The variation in point estimates for sensitivity ranged from 0% to 58% and for specificity ranged from 55% to 100%. The certainty of the evidence was assessed as very low due to risk of bias, indirectness, and inconsistency. Due to the high heterogeneity, we did not pool these studies.

Figure 8. Self-report using VAS; threshold: ≥ 95% adherence [main analysis]

Study	тр	FP	FN	ΤN	Sensitivity (95% CI)	Specificity (95% Cl)	Sensitivity (95%	CI)Specificity (95% CI)
Jiamsakul 2014	7	9	5	60	0.58 [0.28, 0.85]	0.87 [0.77, 0.94]		
Cerutti 2016	52	558	63	691	0.45 [0.36, 0.55]	0.55 [0.53, 0.58]		
McMahon 2013	12	38	22	98	0.35 [0.20, 0.54]	0.72 [0.64, 0.79]		
Ekstran d 2010	18	16	37	131	0.33 [0.21, 0.47]	0.89 [0.83, 0.94]		-
Nelson 2010	46	- 79	104	417	0.31 [0.23, 0.39]	0.84 [0.81, 0.87]		-
Haberer 2011	6	21	14	32	0.30 [0.12, 0.54]	0.60 [0.46, 0.74]		
Dziva 2017	16	12	45	93	0.26 [0.16, 0.39]	0.89 [0.81, 0.94]		-
Labhardt 2012	15	0	50	27	0.23 [0.14, 0.35]	1.00 [0.87, 1.00]		
San ged a 2014	10	19	45	88	0.18 [0.09, 0.31]	0.82 [0.74, 0.89]		
Cohen 2012	41	134	863	176	0.05 [0.03, 0.06]	0.57 [0.51, 0.62]	•	-
Gill 2010	0	2	8	55	0.00 [0.00, 0.37]	0.96 [0.88, 1.00]	0 0.2 0.4 0.6 0.8	1 0 0.2 0.4 0.6 0.8 1







We explored heterogeneity by looking at population, viral load threshold used, and setting. None of these prespecified subgroups could explain the heterogeneity we observed.

- **Population** (Appendix 7) **children** (2 studies, N = 239): sensitivity ranged from 26% to 30% and specificity ranged from 60% to 89%; **adults:** sensitivity ranged from 0% to 58% and specificity ranged from 57% to 96%.
- Viral load threshold (Appendix 7) 40 to 100 copies/mL (6 studies, N = 3591): sensitivity ranged from 5% to 45% and specificity ranged from 55% to 100%; 200 to 400 copies/mL (5 studies, N = 644): sensitivity ranged from 0% to 58% and specificity ranged from 72% to 96%.
- Setting (Appendix 7) low-income (5 studies, N = 663): sensitivity ranged from 18% to 35% and specificity ranged from 60% to 100%; lower-middle-income (3 studies, N = 1631): sensitivity ranged from 0% to 45% and specificity ranged from 55% to 96%.

In addition, three studies also looked at the diagnostic accuracy of a 90% adherence threshold (N = 582, Appendix 7). Sensitivity ranged from 3% to 24% and specificity ranged from 88% to 95%. Another study with 73 participants used 80% as an adherence cutoff (Appendix 7). In this study, sensitivity was 20% (ranging from 6% to 44%) and specificity was 81% (ranging from 68% to 91%).

One study was excluded from the quantitative analyses (Meya 2009) as the authors used an unclear definition for treatment adherence.



2. Tablet counts

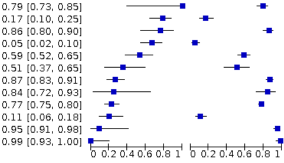
Studies using pharmacy records for the detection of viral nonsuppression showed a large variation in both sensitivity and specificity that we could not explain by subgroup analyses (See 'Summary of findings' table 4 in Appendix 5).

Twelve studies including 3466 participants (of whom 504 had viral non-suppression) used a 95% adherence threshold and were

Figure 10. Tablet counts; threshold: ≥ 95% adherence [main analysis]

Study TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)

40 0 155 1.00 [0.40, 1.00] Kitkungvan 2008 4 Sangeda 2014 43 89 12 18 0.78 [0.65. 0.88] Apisarnthanarak 2010 19 25 6 149 0.76 [0.55, 0.91] Bonjoch 2006 47 144 23 8 0.67 [0.55, 0.78] Coker 2015 25 93 22 133 0.53 [0.38, 0.68] 7 Haberer 2011 26 13 27 0.35 [0.15, 0.59] 0konji 2012 24 44 66 300 0.27 [0.18, 0.37] 9 0.25 [0.03, 0.65] Gill 2010 2 6 48 Cerutti 2016 26 275 88 941 0.23 [0.15, 0.32] 9 Orrell 2017 119 35 15 0.20 [0.10, 0.35] Moosa 2019 1 11 10 211 0.09 [0.00, 0.41] Mariana 2018 0 1 16 81 0.00 [0.00, 0.21]



included in the main analysis (Figure 10). Again, these studies showed a large variation in both sensitivity and specificity that

could not be explained. The variation in point estimates of

sensitivity ranged from 0% to 100% and the specificity ranged from

5% to 99%. The certainty of the evidence was assessed as very low

due to risk of bias, indirectness, and inconsistency. Due to the high

heterogeneity, we did not pool these studies.

We explored heterogeneity by looking at population, viral load threshold used, and setting. None of these prespecified subgroups could explain the heterogeneity we observed.

- Population (Appendix 8) children (1 study; N = 73): sensitivity was 35% (ranged from 15% to 59%) and specificity was 51% (ranged from 37% to 65%); adults (9 studies; N = 3016): sensitivity ranged from 0% to 78% and specificity ranged from 5% to 99%. The two remaining studies were conducted with mixed populations.
- Viral load threshold (Appendix 8) 40 to 80 copies/mL (7 studies; N = 2299): sensitivity ranged from 0% to 100% and specificity ranged from 11% to 99%; 400 copies/mL (5 studies; N = 1167): sensitivity ranged from 9% to 78% and specificity ranged from 17% to 95%.
- Setting (Appendix 8) low-income (4 studies, N = 942): sensitivity ranged from 27% to 78% and specificity ranged from 17% to 87%; lower-middle-income (4 studies: N = 1692): sensitivity ranged from 0% to 100% and specificity ranged from 77% to 99%; upper-middle-income (3 studies; N = 610): sensitivity ranged from 9% to 76% and specificity ranged from 11% to 95%; high-income (1 study; N = 222): sensitivity was 67% (ranged from 55% to 78%) and specificity was 5% (2% to 10%).

Two studies also looked at the diagnostic accuracy of an 80% adherence threshold (Haberer 2011; Sangeda 2014; N = 235 participants). Sensitivity ranged from 0% to 35% and specificity ranged from 69% to 100%.

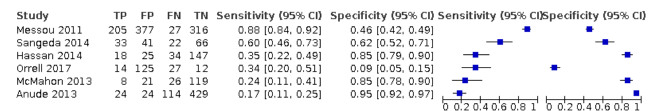
We excluded one study from the quantitative analyses as the adherence threshold used was not relevant for our analyses (90%) (Davies 2008).

3. Pharmacy records or secondary database analysis

Studies using pharmacy records for the detection of viral nonsuppression showed a large variation in both sensitivity and specificity that we could not explain by subgroup analyses (see 'Summary of Findings' table 5 in Appendix 5).

Six studies including 2254 participants (of whom 552 had viral nonsuppression) used a 95% adherence threshold (Anude 2013; Hassan 2014; McMahon 2013; Messou 2011; Orrell 2017; Sangeda 2014), and were included in the main analysis (Figure 11). The sensitivity ranged from 17% to 88% and the specificity ranged from 9% to 95%. The certainty of the evidence was assessed as very low due to risk of bias, indirectness, and inconsistency. Due to the high heterogeneity, we did not pool these studies.

Figure 11. Pharmacy records; threshold: 95% adherence [main analysis]



We explored heterogeneity by looking at population, viral load threshold used, and setting. None of these prespecified subgroups could explain the heterogeneity we observed.

- **Population** (Appendix 9) **adults** (4 studies, N = 1893): sensitivity ranged from 17% to 88% and specificity ranged from 46% to 95%. The two remaining studies were conducted with mixed populations.
- Viral load threshold (Appendix 9) 40 copies/mL (1 study, N = 178): sensitivity was 34% (ranged from 20% to 51%) and specificity was 9% (5% to 15%); 200 to 400 copies/mL (5 studies, N = 2076): sensitivity ranged from 17% to 88% and specificity ranged from 46% to 95%.
- Setting (Appendix 9) low-income (4 studies; N = 1485): sensitivity ranged from 24% to 88%; specificity ranged from 46% to 85%; lower-middle-income (1 study; N = 591): sensitivity was 17% (ranged from 11% to 25%) and specificity was 95% (ranged from 92% to 97%); upper-middle-income (1 study; N = 178): sensitivity was 34% (ranged from 20% to 51%) and specificity was 9% (ranged from 5% to 15%).

Three studies looked at the diagnostic accuracy of an 80% adherence threshold (Messou 2011; Navarro 2014; Sangeda 2014; N = 1211; Appendix 9). Sensitivity ranged from 25% to 82% and specificity ranged from 73% to 88%.

4. Electronic monitoring devices

Studies using electronic monitoring devices for the detection of viral non-suppression showed a large variation in both sensitivity and specificity that we could not explain by subgroup analyses (see 'Summary of Findings' table 6 in Appendix 5).

Three studies including 186 participants (of whom 55 had viral nonsuppression) used a 95% adherence threshold (Evans 2016; Gill 2010; Haberer 2011), and were included in the main analysis (Figure 12). Sensivity ranged from 60% to 88% and specificity ranged from 27% to 67%. The certainty of the evidence was assessed as very low due to risk of bias, indirectness, inconsistency, and imprecision. Due to the high heterogeneity, we did not pool these studies.

Figure 12. Electronic monitoring; threshold: ≥ 95% adherence [main analysis]

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)
 Sensitivity (95% CI)
 Sensitivity (95% CI)
 Gill 2010
 7
 27
 1
 30
 0.88 [0.47, 1.00]
 0.53 [0.39, 0.66]
 Image: Close and the sensitivity (95% CI)
 Image: Close and the sensitititity (95% CI)
 <t

Gill 2010	7 27	1 30	0.88 [0.47, 1.00]	
Evans 2016	20 16	76	0.74 [0.54, 0.89]	
Haberer 2011	12 17	8 35	0.60 [0.36, 0.81]	

We explored heterogeneity by looking at population, viral load threshold used, and setting. None of these prespecified subgroups could explain the heterogeneity we observed.

- Population (Appendix 10) children (1 study, N = 72): specificity was 60% (ranged from 36% to 81%) and specificity was 67% (ranged from 53% to 80%); adults (2 studies, N = 114): sensitivity ranged from 74% to 88% and specificity ranged from 27% to 53%.
- Viral load threshold (Appendix 10) 50 copies/mL (1 study, N = 72) specificity was 60% (ranged from 36% to 81%) and specificity was 67% (ranged from 53% to 80%); 400 copies/mL: (2 studies, N = 114): sensitivity ranged from 74% to 88% and specificity ranged from 27% to 53%.
- Setting (Appendix 10) low-income (1 study, N = 72): specificity was 60% (ranged from 36% to 81%) and specificity was 67% (ranged from 53% to 80%); lower-middle-income (1 study, N = 65): sensitivity was 88% (ranged from 47% to 100%); upper-middle-income (1 study, N = 49): sensitivity was 74% (ranged from 54% to 89%) and specificity was 27% (ranged from 11% to 50%).

Four studies (Evans 2016; Farley 2003; Haberer 2011; Orrell 2017) looked at the diagnostic accuracy of a 80% adherence threshold (N =327, Appendix 10). Sensitivity ranged from 24% to 89% and specificity ranged from 7% to 96%.

5. Composite measures

0.27 [0.11, 0.50] 0.67 [0.53, 0.80]

Studies using composite measures of adherence for the detection of viral non-suppression showed a large variation in both sensitivity and specificity that we could not explain by subgroup analyses (see 'Summary of Findings' table 7 in Appendix 5).

We identified nine studies including 1513 participants that used a composite adherence measure to estimate viral non-suppression (Jayaweera 2003; Mbengue 2019; McMahon 2013; Mutwa 2014; Orrell 2003; Ortega 2004; Parienti 2010; Segeral 2018; Spire 2008), of whom 407 had viral non-suppression. All studies were included in the main analysis (Figure 13). Sensitivity ranged from 10% to 100% and specificity ranged from 49% to 100%. The certainty of the evidence was assessed as very low due to risk of bias, indirectness, and inconsistency. Due to the high heterogeneity, we did not pool these studies. We explored heterogeneity by looking at adherence threshold, population, viral load threshold, and setting.

Figure 13. Composite measure; different adherence thresholds* [main analysis] *cut-off used was either ≥ 95% or 100%

Study TP_FP_FN_TN_Sensitivity (95% CI)_Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)

Parienti 2010	21	26	0	25	1.00 [0.84, 1.00]
Ortega 2004	33	43	6	54	0.85 [0.69, 0.94]
Jayaweera 2003	9	0	4	6	0.69 [0.39, 0.91]
Orrell 2003	49	69	33	91	0.60 [0.48, 0.70]
McMahon 2013	17	45	17	95	0.50 [0.32, 0.68]
Mutwa 2014	8	13	17	66	0.32 [0.15, 0.54]
Segeral 2010	20	28	54	155	0.27 [0.17, 0.39]
Mbengue 2019	9	15	40	99	0.18 [0.09, 0.32]
Spire 2008	7	9	63	267	0.10 [0.04, 0.20]

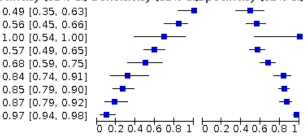
- Adherence threshold (Appendix 11) 100% adherence (6 studies, N = 1095): sensitivity ranged from 10% to 85% and specificity ranged from 56% to 100%; > 95% adherence (3 studies, N = 418): sensitivity ranged from 32% to 100% and specificity ranged from 49% to 84%.
- **Population** (Appendix 11) **children** (1 study, N = 104): sensitivity was 32% (ranged from 15% to 54%) and specificity was 84% (ranged from 74% to 91%); **adults** (7 studies, N = 1390): sensitivity ranged from 10% to 100% and specificity ranged from 49% to 97%. One study did not report on the population.
- Viral load threshold (Appendix 11) 40 to 50 copies/mL (3 studies, N = 522): sensitivity ranged from 10% to 100% and specificity ranged from 49% to 97%; 200 to 400 copies/mL (7 studies, N = 1063): sensitivity ranged from 18% to 100% and specificity ranged from 57% to 100%.
- Setting (Appendix 11) low-income (4 studies, N = 881): sensitivity ranged from 10% to 50% and specificity ranged from 68% to 97%; upper-middle-income (2 studies, N = 405): sensitivity ranged from 18% to 60% and specificity ranged from 57% to 87%; high-income (2 studies, N = 227): sensitivity ranged from 69% to 100% and specificity ranged from 49% to 100%

DISCUSSION

Summary of main results

The aim of this review was to determine the diagnostic accuracy of different adherence measures to detect non-suppressed viral load in people living with HIV. We identified 51 studies, and the main findings are presented in Summary of findings 1.

- Self-report using questionnaires: sensitivity ranged from 10% to 85% and specificity ranged from 10% to 99% (25 studies, 9211 participants; very low-certainty).
- Self-report using VAS: sensitivity ranged from 0% to 58% and the specificity ranged from 55% to 100% (11 studies, 4235 participants; very low-certainty).
- Tablet counts: sensitivity ranged from 0% to 100% and the specificity ranged from 5% to 99% (12 studies, 3466 participants; very low-certainty).
- Pharmacy records or secondary databases: sensitivity ranged from 17% to 88% and the specificity ranged from 9% to 95% (6 studies, 2254 participants; very low-certainty).
- Electronic monitoring devices: sensitivity ranged from 60% to 88% and the specificity ranged from 27% to 67% (3 studies, 186 participants; very low-certainty).



• Composite measure: sensitivity ranged from 10% to 100% and the specificity ranged from 49% to 100% (9 studies, 1513 participants; very low-certainty).

None of the methods of measure of adherence had a consistent sensitivity to detect viral non-suppression. We did not perform meta-analysis because we encountered significant heterogeneity between the studies that we could not explain by population (children and adults), viral load threshold used (above or below 100 copies/mL) or setting (low-income, lower-middle-income, upper-middle-income, high-income).

Risk of bias is presented in Figure 5 The category in which we most frequently identified high risk of bias was with regards to flow and timing. Within this category, concerns related to uncertainty regarding whether viral load and adherence were measured contemporaneously, and large amounts of missing data.

Strengths and weaknesses of the review

To our knowledge, this review represents the largest systematic collation of data in the field, including detailed assessment of the certainty of evidence provided from the field.

As stated in our protocol and background, the relationship between adherence and viral load is not linear. Other patient and drugrelated factors will influence viral suppression including drug resistance, viral load at outset of therapy, history of suppression, pharmacokinetics such as absorption, the genetic barrier to resistance offered by the regimen, and drug-drug interactions. A decrease in adherence could precede the viral non-suppression by a number of weeks. Patients who have historically had excellent adherence and viral suppression may experience 'blips' without any of these factors appearing to be at play. Such blips may correct on retesting, and not considered viral non-suppression in clinical practice. Patients may experience low-level viraemia that may be of questionable clinical significance.

Notwithstanding the non-linear relationship, a potential lag between non-adherence and viral load rise, the phenomena of 'blips', and low-level viraemia, we made a pragmatic decision to use a non-suppressed viral load as the target condition. We considered this likely to offer the best objective measure of the success of ART, and it is a clinically relevant measure. Moreover, our objective was in part to understand if there was a role for simple adherence measures in settings where viral loads were less available. We recognize that this encapsulates a broad definition but we feel this was a necessary simplification to allow for the



variation in definitions used for viral failure internationally (Figure 2). In subgroup analysis, we did not detect that changing the viral load threshold used influenced the findings. We do not feel that the target condition we chose, nor the phenomena described above including non-linear relationships, lags, blips, and low-level viraemia, could explain the low overall sensitivities seen across studies, or explain the high variation or heterogeneity seen across studies.

We included studies which had not been conceived as diagnostic test accuracy studies; rather, they were studies of other methodologies that included measures of adherence at the same time as viral load measurement and thus allowed us to extract data for 2 x 2 tables. Examples included studies aiming to describe adherence within a cohort, evaluations of adherence interventions, and randomized controlled trials comparing ART regimens. However, in an exploratory analysis of our data, we did not find that study design or objective could explain the heterogeneity we observed.

Our search identified other studies which stated they had measured adherence and viral load contemporaneously, which were similar to design studies that we included, but did not report all the data required for the 2 x 2 tables (for example, RCTs in which the text mentioned adherence had been measured, but did not report the relevant results). The author team agreed that it was not feasible to contact study authors in these instances. Given the large number of studies included in this review, we do not think that inclusion of such data would substantively influence the conclusions.

We did not perform formal subgroup analysis according to type of ART or specific ART regimens. The use of more or less 'forgiving' regimens may influence the relationship between adherence and viral non-suppression. However, the setting may to some extent reflect ART regimens used, and these did not explain heterogeneity.

In some instances, we compiled different thresholds (for example, for composite measures). This was a pragmatic decision to allow comparison of data from studies. Again, we do not think that this could explain the low sensitivities and specificities encountered, or the high variation in point estimates.

We did not make equity considerations part of the review framework and the outset, but our pre-planned subgroup analyses to some extent addressed equity considerations, including considerations on patient characteristics and income setting. Some measures of adherence might have lower accuracy in different populations due to complex influences. For example, an individual suffering from stigma due to HIV, compounded by intersectional stigma due to other characteristics (for example gender or race), might additionally fear the sense of moral judgement and labelling associated with perceived poor adherence (Eshun-Wilson 2019); this could influence their response to questions. It is beyond the scope of the methodology of this review to address complex concerns such as these in depth, and we do not consider that this would change the conclusions of the review.

Applicability of findings to the review question

The applicability domains of our QUADAS-2 assessment help determine the applicability of our findings to the review question, as described in the Methodological quality of included studies. We had low concern for patient selection applicability in most studies (55%), as these studies took place in unselected populations. We had low concern for index test applicability for most studies (74%), as they were easily implementable in all settings. For only 2% of studies, did we have high applicability concerns, as we felt the tests were too complex to administer or required expensive electronic devices. Finally, we also had low concern for reference standard applicability for most studies (67%), as study authors clearly described the viral load assay used. Overall, we feel that the findings can therefore be considered applicable to the objective of our review, which was to understand whether simple measures of adherence could be used to detect viral non-suppression in diverse settings.

Agreements and disagreements with other studies or reviews

To our knowledge, the current review represents the only systematic review of measures of adherence in HIV to formally assess diagnostic test accuracy. Almeida-Brasil 2019 is a metaanalysis including observational studies that compared adherence measures, and calculated odds ratios for a given test to detect virological failure. This, therefore, allowed for some pairwise comparison of tests without reporting sensitivity and specificity. The authors concluded that low cost measures (such as self-report) appeared equally as effective as higher cost measures such as electronic monitoring. To some extent, our review mirrors this conclusion, in so much as there was insufficient sensitivity to reliably detect viral non-suppression for all tests. Other reviews that included different approaches to measuring adherence in HIV have largely taken a narrative approach, for example, Spinelli 2020. Such reviews are valuable in appraisal of the benefits and disbenefits of using these measures clinically, but do not capture the variation and uncertainty we report.

AUTHORS' CONCLUSIONS

Implications for practice

We encountered a wide variety of adherence measures including numerous self-report measures, tablet counts, pharmacy records, electronic monitoring, and composite measures. Across groupings of similar measures, no one modality consistently offered a sufficiently high sensitivity to detect viral non-suppression. Given the variation and inconsistency between studies of the same type of adherence measure, it is not possible to recommend one type of adherence measure over another. Incorporating individual measures into composite measures do not seem to improve sensitivity above and beyond an individual method.

There is, therefore, no one adherence test that might helpfully offer an alternative to frequent viral load measurement in settings where viral load measurement was less available. This highlights the ongoing importance of viral load measurement, and ensuring access to it at individual and programmatic level. In addition, none of the adherence measures studied consistently offered a high specificity such that it might be used to identify targeted individuals who might benefit most from evidence-based adherence interventions, such as text-messaging, counselling, or supporter interventions.

Guideline and policy should recognize the high uncertainty, and probable overall limited ability of these tests to detect non-suppressed viral loads. Nevertheless, there may be other

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qualitative benefits to attempting to measure adherence that are beyond the scope of this review to detect.

Implications for research

Given the low overall sensitivity and specificity of these adherence measures to detect a non-suppressed viral load, and vast variation in point estimates, we do not consider that further adaptation and evaluation of these measures would yield significant improvements in diagnostic test accuracy. Further research might more helpfully look to reducing costs of viral load monitoring, or of reducing costs of alternative measures of objectively quantifying adherence, such as measuring drug modalities with a longer halflife in plasma, dried blood spots, or urine. The benefit of electronic devices when used to identify and act upon missed doses in realtime has yet to be fully determined.

Studies of adherence interventions should consider the uncertainty around the measures used, and may better focus on clinical outcomes including viral load, over self-reported adherence.

When considering equity, the lack of a suitable alternative to viral load assessment may have a further negative effect on people living with HIV in lower-income areas where viral load testing may be less available. Inequities could hypothetically explain some of the vast variation observed, and qualitative methodologies may be best placed to investigate this.

This review did not compare formal measures of adherence to qualitative assessment of a patient's probable adherence by healthcare providers. Future research could ask if the additive value of formal measures of adherence, as represented by the index tests in this review, above and beyond clinical assessment of adherence by healthcare providers, justifies their use in clinical practice. Qualitative research could helpfully ask whether the administration of adherence measures in practice offers clinical benefits or harms beyond predicting viral non-suppression, but this is beyond the scope of this review.

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Editorial and peer-reviewer contributions

The CIDG supported the authors in the development of this Cochrane Diagnostic Test Accuracy (DTA) Review.

The following people conducted the editorial process for this review update.

CIDG Contact Editor: Dr Lawrence Mbuagbaw; DTA Contact Editor: Ms Marta Roqué

- Sign-off Editor (final editorial decision): Professor Paul Garner
- Managing Editor (collated comments, provided editorial guidance to authors, edited the article): Dr Deirdre Walshe
- Copy Editor (copy editing and production): Anne Lethaby, Cochrane Copy Edit Support
- Peer reviewers (review stage; provided comments and recommended an editorial decision): Dr Michael McCaul, Division of Epidemiology and Biostatistics, Department of Global Health, Stellenbosch University, South Africa (clinical/ content peer review); Dr Rebecca Kuehn, Liverpool School of Tropical Medicine, UK (clinical/content peer review); Dr Jantjie Taljaard. Division of Infectious Diseases, Department of Medicine, Tygerberg Hospital and Stellenbosch University, South Africa (clinical/content peer review). Three additional peer reviewers from the DTA editorial team provided search, statistical, and general methods peer review, but chose not to be publicly acknowledged.

Dr Rebecca Kuehn is a member of CIDG, and provided peer-review comments on this article, but was not otherwise involved in the editorial process or decision making for this article. Dr Marty Chaplin, Dr Nathan Ford, and Dr Paul Hine are CIDG Editors, and Dr Mariska Leeflang is a DTA Editor, but were not involved in the editorial process of this article.

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CHARACTERISTICS OF STUDIES

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Study characteristics	
Patient Sampling	Target population: adults
	 Recruitment: a cohort of 2585 initially ART-naïve adults who started HAART between April 2008 and February 2009 were followed up for 12 months in three representative government hospitals in Nigeria: University of Abuja Teaching Hospital, Abuja (UATH), University of Benir Teaching Hospital, Benin (UBTH) and Asokoro District Hospital, Asokoro, Abuja (ADH). Inclusion criteria: ART-naïve adults who started HAART between April 2008 and February 2009 Exclusion criteria: not reported
	Study design: prospective cohort study
Patient characteristics and setting	 Country: Nigeria World Bank Income classification: low-middle-income Study setting: hospital-based (three representative government hospitals in Nigeria: University of Abuja Teaching Hospital, Abuja (UATH), University of Benin Teaching Hospital Benin (UBTH) and Asokoro District Hospital, Asokoro, Abuja (ADH)) Study dates: April 2008 to February 2009 Age of population (median, IQR): 35 years, 30 to 41
	 Gender (male %): 36.3 Participants included/analysed: 628/591 First or second-line regimen: first-line
	 Type of ART: the choice of HAART combination was both guided by national treatment protocols and the discretion of the attending physician but generally consisted of one of three first line regimens (TDF + 3TC + NVP/EFV; AZT + 3TC + NVP/EFV; stavudine + 3TC + NVP/EFV) Time on ART at enrolment: treatment-naïve

Anude 2013 (Continued)

Cochrane Database of Systematic Reviews

nude 2013 (Continued)	• Time on ART at measurement of viral load and adherence: 12 months				
Index tests	Number of index tests used: 2				
	Types of index tests: pharmacy records and self-report				
	 Blinding: no information Threshold prespecifie Adherence threshold Test 2. Self-report Validated scale: not rest 	pplicable macy refill records (no furt ion d: not reported used: 95% eported report (no further details p ion d: not reported			
	The study only reported usa	ble 2 x 2 data for this review	w for the pharmacy records.		
Target condition and reference standard(s)	 Target condition: viral non-suppression Reference standard: viral load testing was done by quantitative PCR for HIV-1 RNA in humar plasma using Roche Amplicor version 1.5, Roche Diagnostics, Basal, Switzerland. Definition of viral non-suppression: HIV viral load level > 400 copies/mL Blinded to index test: no information 				
Flow and timing	 Time interval between index and reference tests: no clear information on timing, just that it was measured at 12 months All patients received same reference standard: yes Missing data: 2585 initially included in the cohort, 805 (31%) of patients were lost to follow-up at 12 months, 628 out of the 1780 patients alive and active on the programme at 12 months were randomly selected for indepth interviews and laboratory work-up with detailed virologic and immunologic testing. Of those, 591 had data available for inclusion. Missing data > 10% 				
Comparative					
Notes	Conflicts of interest: none d	eclared			
	Cooperative Agreement Nur	nber: PS000651. Chuka Anı	ters for Disease Control and Prevention ude was funded by the US National In- Research Program (AITRP, NIH 2-D43-		
	Trial registry: not reported				
Methodological quality					
Item	Authors' judgement	Risk of bias	Applicability concerns		
DOMAIN 1: Patient Selection					
Was a consecutive or random sam- ple of patients enrolled?	Unclear				



Anude 2013 (Continued)				
Was a case-control design avoid- ed?	Yes			
Did the study avoid inappropriate exclusions?	Unclear			
Could the selection of patients have introduced bias?		Unclear risk		
Are there concerns that the in- cluded patients and setting do not match the review question?			Low concern	
DOMAIN 2: Index Test (Index test)				
Were the index test results inter- preted without knowledge of the results of the reference standard?	Unclear			
If a threshold was used, was it pre- specified?	Unclear			
Could the conduct or interpreta- tion of the index test have intro- duced bias?		Unclear risk		
Are there concerns that the in- dex test, its conduct, or inter- pretation differ from the review question?			Unclear	
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condi- tion?	Yes			
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Unclear			
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk		
Are there concerns that the tar- get condition as defined by the reference standard does not match the question?			Low concern	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval between index test and reference standard?	Unclear			

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Anude 2013 (Continued)

Could the patient flow have in-	High risk
Were all patients included in the analysis?	No
Did all patients receive the same reference standard?	Yes

troduced bias?

Apisarnthanarak 2010

Study characteristics	
Patient Sampling	 Target population: patients whose HIV level was suppressed at month 6 to < 50 copies, mL were followed up to ascertain achievement of durable HIV suppression at year 3 Recruitment: patients who were prescribed a regimen of fixed-dose, twice-daily stavudine, 3TC, and NVP and enrolled in a study at Thammasat hospital Inclusion criteria: not reported Exclusion criteria: not reported Study design: prospective cohort study
Patient characteristics and setting	 Country: Thailand World Bank Income classification: upper-middle-income Study setting: hospital- and home-based Study dates: April 2003 to January 2007 Age of population (years), median (range): 37 (15 to 61) Gender (male %): 32 Participants included/analysed: 199/199 First or second-line regimen: first-line Type of ART: stavudine, 3TC and NPV Time on ART at enrolment: not reported Time on ART at measurement of viral load and adherence: 6, 12, 18, 24, 30, 36 months
Index tests	Number of index tests used: 1 Types of index tests: tablet counts
	 Test 1. Tablet counts Validated scale: not applicable Tool description: at each routine medical encounter, the pharmacist calculated the ratio of pills taken divided by the total number of pills prescribed for the interval period. The unannounced home visits were randomly conducted by trained adherence counselling educators twice monthly and included pill counts the mean pill count ratios (based on scheduled and unannounced visits) were calculated. Blinding: no information Threshold prespecified: not reported Adherence threshold used: 95%; 75%
Target condition and reference stan- dard(s)	Target condition: viral non-suppressionReference standard: no details provided on the assay used

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

Apisarnthanarak 2010 (Continued)				
Flow and timing	 Time interval between index and reference tests: no explicit information on timing ("The unannounced home visits were randomly conducted by trained adherence counseling educators twice monthly and included pill counts the mean pill count ratios (based on scheduled and unannounced visits were calculated for each 6-month period of observation, and the HIV load was determined every 6 months") All patients received same reference standard: not reported Missing data: none. All eligible participants with viral load test and adherence measures were included in the main analysis. 			
Comparative				
Notes	Conflicts of interest: L.M.M. A.A.: no conflicts	is a consultant to WW E	pidemiology at GlaxoSmithKline, Inc.	
	Funding source: Thammasa search Fund	at University Infectious D	iseases and Infection Control Re-	
	Trial registry: not reported			
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Unclear			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate exclu- sions?	No			
Could the selection of patients have in- troduced bias?		High risk		
Are there concerns that the included patients and setting do not match the review question?			Low concern	
DOMAIN 2: Index Test (Index test)				
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear			
If a threshold was used, was it pre-speci- fied?	Unclear			
Could the conduct or interpretation of the index test have introduced bias?	Unclear risk			
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern	
DOMAIN 3: Reference Standard				

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Apisarnthanarak 2010 (Continued)			
Is the reference standards likely to cor- rectly classify the target condition?	Unclear		
Were the reference standard results inter- preted without knowledge of the results of the index tests?	Unclear		
Could the reference standard, its con- duct, or its interpretation have intro- duced bias?		Unclear risk	
Are there concerns that the target con- dition as defined by the reference stan- dard does not match the question?			Unclear
DOMAIN 4: Flow and Timing			
Was there an appropriate interval be- tween index test and reference standard?	Unclear		
Did all patients receive the same refer- ence standard?	Unclear		
Were all patients included in the analysis?	Yes		
Could the patient flow have introduced bias?		Unclear risk	

Avong 2015

Study characteristics	
Patient Sampling	 Target population: adult AIDS patients who had been treated with combination ART for at least 12 months Recruitment: at the time the study was conducted, about 11,208 AIDS patients were ever enrolled and were on ART according to programme implementation report provided by the Institute of Human Virology, Nigeria – managers of the ART clinic Inclusion criteria: participants comprised adult AIDS patients who had been treated with combination antiretroviral therapy (cART) for at least 12 months as at May 2010 Exclusion criteria: patients who were less than 18 years, critically ill or hospitalized and could not be interviewed as well as those who were not currently taking ARV Study design: cross-sectional
Patient characteristics and setting	 Country: Nigeria World Bank Income classification: low-middle income Study setting: hospital-based (tertiary level) Study dates: 2004 to 2010 Age of population (years), median (IQR): men: 42 (38 to 44); female: 36 (30 to 40) Gender (male %): 49.4 Participants included/analysed: 537/502 First or second-line regimen: HAART mixed regimens Type of ART: first-line: AZT/3TC + NVP or EFV; AZT/3TC/NVP; 3TC/NVP/d4T; TDF/FTC + EFV or NVP and second-line: TDF + 3TC + LPV/r



vong 2015 (Continued)	 Time on ART at enro 	lment: 12 months (mean du	ration of therapy was 43 months with	
	range of 16 to 70 months)			
		surement of viral load and a hs with a range of 16 to 70 m	dherence: 12 months (mean duration on the first second s	
Index tests	Number of index tests usec	1:1		
	Types of index test: self-rep	port		
	a dose, correct dose, old categories used a ent (i.e. OPTIMAL ad	Freported adherence was ass correct frequency, correct sc as cut-off for optimal adheren herence) if reported comply (5–100% and if not missing a tion ed: not reported	sessed in five different ways (not missin hedule and effective adherence).Thresh nce: a participant was considered adhe ing with the correct schedule, dose, fre ny dose in the past 3 days.	
Target condition and reference	Target condition: viral non-	-suppression		
standard(s)	 Reference standard: Roche Cobas AmpliPrep TaqMan (Cobas Amplicor; Roche Diagnostics, Switzerland) 			
	 Definition of viral non-suppression: HIV viral load > 400 copies/mL 			
	Blinded to index test: no information			
Flow and timing	 Time interval between index and reference tests: no explicit information on timing, but this is a cross-sectional study so likely to be measured simultaneously All nations received same reference standard; yes 			
	 All patients received same reference standard: yes Missing data: during the data cleaning, it was found that 28 participants had incomplete prescription refill data and one was less than 18 years. Thus, 35 participants were excluded learing 502 participants whose data was entered into the analysis. Missing data < 10% 			
Comparative				
Notes	Conflicts of interest: none of	declared		
	D43 TWO 10441 from the U	nited States' National Institu	UMB AITRP Fogarty Grant Number 5- ites of Health's Forgarty International of Human Virology, University of Mary-	
	Trial registry: not reported			
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sam- ple of patients enrolled?	Unclear			
Was a case-control design avoid- ed?	Yes			

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Avong 2015 (Continued)			
Did the study avoid inappropriate exclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the in- cluded patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Index test)			
Were the index test results inter- preted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre- specified?	Unclear		
Could the conduct or interpreta- tion of the index test have intro- duced bias?		Unclear risk	
Are there concerns that the in- dex test, its conduct, or inter- pretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condi- tion?	Yes		
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the tar- get condition as defined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Avong 2015 (Continued)

troduced bias?

Were all patients included in the Yes analysis?

Could the patient flow have in-

Low risk

Study characteristics	
Patient Sampling	 Target population: adults with HIV who received ART for at least 6 months Recruitment: all patients who received ART since December 2004, the programme's in ception, and of patients who initiated ART through December 2006 Inclusion criteria: adult patients (18 years) who received ART for at least 6 months and attended clinic at least once between April and December 2006 Exclusion criteria: no exclusions of those who met eligibility criteria Study design: prospective cohort study
Patient characteristics and setting	 Study design: prospective conort study Country: Uganda World Bank Income classification: low-income Study setting: hospital-based (Kitagata Hospital, a government-owned district hospital located in the Bushenyi district of rural southwestern Uganda) Study dates: April to December 2006 Age of population; not reported Gender (male %): 39.8 Participants included/analysed: 175/175 First or second-line regimen: first-line Type of ART: fixed-dose combination of stavudine + 3TC + NVP Time on ART at enrolment: at least 6 months Time on ART at measurement of viral load and adherence: at least 6 months
Index tests	 Number of index tests used: 2 Types of index tests: self-report and tablet counts Test 1. Self-report Validated scale: not reported Tool description: three-day recall of adherence; patients were considered nonadher ent if they missed at least 1 antiretroviral pill and 100% adherent if they had not Blinding: no information Threshold prespecified: not reported Adherence threshold used: 100% Test 2. Tablet counts Validated scale: not applicable Tool description: pill count data performed routinely by the dispenser who would asl patients to bring with them pill bottles and unused medications; percentage adher ence was calculated as the fraction of doses assumed taken among the total numbe of doses dispensed since the scheduled clinic visit. Blinding: no information Threshold prespecified: not reported Adherence threshold used: 100%

	rusted evidence. nformed decisions Setter health.			Cochrane Database of Systematic Review	
Bajunirwe 2009 (Continued)					
Target condition and refere	nce stan-	Target condition: viral nor	n-suppression		
dard(s)			04 count and plasma HIV I	RNA concentration using the Roche Am-	
		plicor v1.5 assayDefinition of viral non-suppression: HIV viral load > 50 copies/mL			
		 Blinded to index test: no information 			
Flow and timing				: no explicit information on timing; how-	
		ever, both measures at	baseline me reference standard: ye	ec.	
				viral load test and adherence measures	
		were included in the m			
Comparative					
Notes		Conflicts of interest: none	declared		
		Funding source: this study was funded, in part, by grants from the Fogarty International Center AIDS International Training and Research Program (TW00011) and the Centers For AIDS Research (AI36219).			
		Trial registry: not reported	I		
Methodological quality					
Item		Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selecti	on				
Was a consecutive or rando of patients enrolled?	m sample	Yes			
Was a case-control design a	voided?	Yes			
Did the study avoid inappro clusions?	opriate ex-	Yes			
Could the selection of pati introduced bias?	ients have		Low risk		
Are there concerns that th ed patients and setting do the review question?				Low concern	
DOMAIN 2: Index Test (Ind	ex test)				
Were the index test results i ed without knowledge of th the reference standard?		Unclear			
If a threshold was used, was specified?	s it pre-	Unclear			
Could the conduct or inter of the index test have intr bias?			Unclear risk		

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

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Bajunirwe 2009 (Continued)			
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to cor- rectly classify the target condition?	Yes		
Were the reference standard results in- terpreted without knowledge of the re- sults of the index tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the target condition as defined by the refer- ence standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval be- tween index test and reference stan- dard?	Unclear		
Did all patients receive the same refer- ence standard?	Yes		
Were all patients included in the analy- sis?	Yes		
Could the patient flow have intro- duced bias?		Low risk	

Bonjoch 2006

Study characteristics	
Patient Sampling	 Target population: adults with HIV who had been receiving an NVP-containing HAART regimen for at least 2 years
	 Recruitment: participants were identified by unselected consecutive recruitmen from outpatient clinic visits during a total period of 4 months.
	 Inclusion criteria: adult HIV-1-infected patients were included if they had been re ceiving an NVP-containing HAART regimen for at least 2 years, regardless of the reason for its initiation (first-line, salvage, or simplification).
	• Exclusion criteria: patients who discontinued treatment with NVP within the firs 2 years due to adverse events
Patient characteristics and setting	 Country: Spain World Bank Income classification: high-income
	 Study setting: 12 tertiary care hospitals in Spain
	Study dates: not reported

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

Bonjoch 2006 (Continued)	
	 Age of population (years), median (IQR): 41 (37 to 46) Gender (male %): 70.8
	 Participants included/analysed: 613/222
	 First or second-line regimen: first-line, salvage, or simplification Type of ART: regardless of the reason for initiation (first-line, salvage, or simpli-
	fication)
	 Time on ART at enrolment: at least 2 years
	 Time on ART at measurement of viral load and adherence: not reported, at least 2 years
Index tests	Number of index tests used: 1
	Types of index tests: tablet counts
	 Test 1. Tablet counts Validated scale: not applicable Tool description: the proportion of compliance was calculated by dividing the number of pills consumed during the last month by the number of pills pre-
	scribed in the same period.
	 Blinding: no information Threshold prospecified: not reported
	 Threshold prespecified: not reported Adherence threshold used: 90%; 95%
Target condition and reference standard(s)	Target condition: viral non-suppression
	Reference standard: not reported
	 Definition of viral non-suppression: HIV viral load > 50 copies/mL
	Blinded to index test: no information
Flow and timing	 Time interval between index and reference tests: no explicit information on timing, but this was a cross-sectional study so likely to be measured simultaneously All patients received same reference standard: not reported Missing data: data on adherence and viral load only available for 222/613. Missing data > 10%
Comparative	
Notes	Conflicts of interest: none declared
	Funding source: not reported
	Trial registry: not reported
Methodological quality	
ltem	Authors' judgement Risk of bias Applicability concerns
DOMAIN 1: Patient Selection	
Was a consecutive or random sample of pa- tients enrolled?	Yes
Was a case-control design avoided?	Yes
Did the study avoid inappropriate exclusions?	Unclear



Sonjoch 2006 (Continued)			
Could the selection of patients have intro- duced bias?		Low risk	
Are there concerns that the included pa- tients and setting do not match the review question?			Unclear
DOMAIN 2: Index Test (Index test)			
Were the index test results interpreted with- out knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Unclear		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Unclear		
Were the reference standard results interpret- ed without knowledge of the results of the in- dex tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Unclear risk	
Are there concerns that the target condi- tion as defined by the reference standard does not match the question?			Unclear
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between in- dex test and reference standard?	Yes		
Did all patients receive the same reference standard?	Unclear		
Were all patients included in the analysis?	No		
Could the patient flow have introduced bias?		High risk	

Cerutti 2016

Study characteristics

Cerutti 2016 (Continued)	
Patient Sampling	 Target population: patients on ART ≥ 6 months attending routine follow-up visits between May 5, 2014 and June 17, 2014 Recruitment: consecutive sample enrolled Inclusion criteria: all patients on ART ≥ 6 months attending routine follow-up visits between May 5, 2014 and June 17, 2014 and willing to participate received viral load measurement and extensive comorbidity screening, including assessment for alcohol use disorder and depressive symptoms. Exclusion criteria: being on ART < 6 months, history of treatment interruption ≥ 7 days during the last 3 months, receiving second-line ART Study design: cross-sectional
Patient characteristics and setting	 Country: Lesotho World Bank Income classification: lower-middle-income Study setting: two rural districts, Butha-Buthe and Thaba-Tseka in Lesotho Study dates: May to June 2014 Age of population (years), median (IQR): 43.6 (34.5 to 53.5) Gender (male %): 31 Participants included/analysed: 1389/1330 (tablet count); 1390/1364 (self-report) First or second-line regimen: first-line Type of ART: not reported Time on ART at enrolment: at least 6 months Time on ART at measurement of viral load and adherence: not reported; at least 6 months
Index tests	 Number of index tests used: 2 Types of index tests: Self-report and tablet counts Test 1. Self-reported (VAS) Validated scale: yes Tool description: self-reported adherence using a visual analogue scale (VAS): Adherence reports were obtained from clinical notes Blinding: no information Threshold prespecified: not reported Adherence threshold used: 95% Test 2. Tablet counts Validated scale: not applicable Tool description: the proportion of compliance was calculated by dividing the number of pills consumed during the last month by the number of pills prescribed in the same period. Blinding: no information Threshold prespecified: not reported
Target condition and reference stan- dard(s)	 Target condition: viral non-suppression Reference standard: viral RNA was prepared using an automated extractor (NucliSENS easyMAG, Biomerieux, Switzerland) Definition of viral non-suppression: HIV viral load > 80 copies/mL Blinded to index test: no information
Flow and timing	 Time interval between index and reference tests: no explicit information on timing, but this was a cross-sectional study so likely to be measured simultaneously All patients received same reference standard: yes

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Cerutti 2016 (Continued)

Missing data: self-report data available for 1330/1388 and pill count data available for 1364/1388. Missing data < 10%

Comparative			
Notes	Conflicts of interest: none	declared	
	Funding source: the Swiss through a grant to ND Labl		and Talent in Biomedical Research
	Trial registry: NCT0212669	6	
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclu- sions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Index test)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-spec- ified?	Unclear		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to cor- rectly classify the target condition?	Yes		
Were the reference standard results in- terpreted without knowledge of the re- sults of the index tests?	Unclear		

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Cerutti 2016 (Continued)	
Could the reference standard, its con- duct, or its interpretation have intro- duced bias?	Low risk
Are there concerns that the target condition as defined by the reference standard does not match the ques- tion?	Low concern
DOMAIN 4: Flow and Timing	
Was there an appropriate interval be- tween index test and reference stan- dard?	Yes
Did all patients receive the same refer- ence standard?	Yes
Were all patients included in the analy- sis?	Yes
Could the patient flow have intro- duced bias?	Low risk

Cohen 2012

Study characteristics	
Patient Sampling	Target population: treatment-naïve, HIV-1-infected adults with baseline viral load greater than or equal 5000 copies/mL and confirmed viral sensitivity to the background nucleoside/nucleotide reverse transcriptase inhibitors (N[t]RTIs)
	 Recruitment: no details reported, participants of two RCTs Inclusion criteria: treatment-naive, HIV-1-infected adults with baseline viral load greater than or equal 5000 copies per millilitre and confirmed viral sensitivity to the background nucleoside/nucleotide reverse transcriptase inhibitors (N[t]RTIs) (assessed using the vircoTYPE HIV-1 assay)
	• Exclusion criteria: documented presence of any NTI resistance-associated mutation (RAM) from a list of 39; active clinically significant disease (e.g. pancreatitis, cardiac dysfunction, active and significant psy- chiatric disorder, adrenal insufficiency, or hepatic impairment), renal impairment, pregnancy or breast- feeding
	Study design: pooled analysis of 2 RCTs
Patient characteristics and setting	 Country: each trial was conducted in 21 countries, with some overlap of countries (USA, Canada, Australia, South Africa, several countries in Europe, several in Asia, and several in Latin America). More than half the participants across the trials were from the combined USA, Canada, Europe & Australia regions. World Bank Income classification: high-income
	 Study setting: not reported
	Study dates: not reported
	Age of population (years): not reported
	Gender (male %): not reported
	Participants included/analysed:1368/1214
	 First or second-line regimen: first-line Type of ART: PI-based second-line regimen. Both groups combined: RPV 25 mg with EFV placebo once daily or EFV 600 mg with RPV placebo once daily, both in addition to (i) a fixed background N[t]RTI regimen of TDF and emtricitabine in the ECHO trial, or (ii) an N[t]RTI regimen based on the investigator's choice of TDF/FTC, zidovudine/lamivudine, or abacavir/3TC in the THRIVE trial.2: RPV

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Cohen 2012 (Continued)	 25 mg with EFV placebo once daily, in addition to (i) a fixed background N[t]RTI regimen of TDF and emtricitabine in the ECHO trial, or (ii) an N[t]RTI regimen based on the investigator's choice of TDF/ FTC, zidovudine/lamivudine (3TC), or abacavir/3TC in the THRIVE trial.3: EFV 600 mg with RPV placebo once daily, both in addition to (i) a fixed background N[t]RTI regimen of TDF and emtricitabine in the ECHO trial, or (ii) an N[t]RTI regimen based on the investigator's choice of TDF/FTC, zidovudine/lamivudine (3TC), or abacavir/3TC in the THRIVE trial o Time on ART at enrolment: treatment-naïve o Time on ART at measurement of viral load and adherence: 6 months
Index tests	Number of index tests used: 1
	Types of index tests: self-report
	 Test 1. Self-report questionnaire Validated scale: yes (M-MASRI) Tool description: Modified Medication Adherence Self-Report Inventory (M-MASRI); prescribed Blinding: no information Threshold prespecified: yes Adherence threshold used: 95%
Target condition and ref-	Target condition: viral non-suppression
erence standard(s)	 Reference standard: Amplicor HIV-1 monitor test version 1.5 (Roche, Basel, Switzerland) Definition of viral non-suppression: HIV viral load > 50 copies/mL Blinded to index test: no information
Flow and timing	 Time interval between index and reference tests: no explicit information on timing, but all data meant to be from week 48 All patients received same reference standard: yes Missing data: there were a total of 1368 patients between both trials, but only 1214 were included in
Comparativo	the analysis (686/627; and 682/587 for each trial, respectively). Overall missing data < 10%
Comparative	
Notes	Conflicts of interest: C.J.C. has received research funding from Janssen, Gilead Sciences, Bristol-Myers Squibb (BMS), Mer- ck, Tobira and ViiV Healthcare. He is on advisory boards for Gilead Sciences, Janssen, Merck, Tobira and BMS. He has received speaker honoraria from Janssen, Gilead Sciences, BMS and Merck prior to January 2011. J.M.M. has acted as a consultant, participated in advisory boards, has received speaker fees and has
	been an investigator for clinical trials for Janssen, ViiV Healthcare, Gilead Sciences, BMS, Abbott Labora- tories, Boehringer Ingelheim (BI) and Merck, Sharp, and Dohme (MSD)
	Funding source:
	C.J.C. has received research funding from Janssen, Gilead Sciences, Bristol-Myers Squibb (BMS), Mer- ck, Tobira and ViiV Healthcare. He is on advisory boards for Gilead Sciences, Janssen, Merck, Tobira and BMS. He has received speaker honoraria from Janssen, Gilead Sciences, BMS and Merck prior to January 2011.
	J.M.M. has acted as a consultant, participated in advisory boards, has received speaker fees and has been an investigator for clinical trials for Janssen, ViiV Healthcare, Gilead Sciences, BMS, Abbott Labora- tories, Boehringer Ingelheim (BI) and Merck, Sharp, and Dohme (MSD)
	Trial registry: NCT00540449 & NCT00543725



Cohen 2012 (Continued)

ltem	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selecti	ion		
Was a consecutive or ran- dom sample of patients enrolled?	Unclear		
Was a case-control de- sign avoided?	Yes		
Did the study avoid inap- propriate exclusions?	Unclear		
Could the selection of patients have intro- duced bias?		Unclear risk	
Are there concerns that the included pa- tients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Ind	lex test)		
Were the index test re- sults interpreted without knowledge of the results of the reference stan- dard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have intro- duced bias?		Low risk	
Are there concerns that the index test, its con- duct, or interpretation differ from the review question?			Unclear
DOMAIN 3: Reference Sta	ndard		
Is the reference stan- dards likely to correctly classify the target condi- tion?	Yes		
Were the reference stan- dard results interpreted without knowledge of the results of the index tests?	Unclear		

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

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Cohen 2012 (Continued)			
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the target condition as defined by the refer- ence standard does not match the question?			Low concern
DOMAIN 4: Flow and Timi	ıg		
Was there an appropriate interval between index test and reference stan- dard?	Unclear		
Did all patients receive the same reference stan- dard?	Yes		
Were all patients includ- ed in the analysis?	Yes		
Could the patient flow have introduced bias?		Unclear risk	

Coker 2015

Study characteristics	
Patient Sampling	 Target population: HIV-infected individuals initiating care and treatment from August 2006 to January 2008 Recruitment: no details reported, participants of an RCT Inclusion criteria: 18 years and older, tested positive for HIV-1 antibodies, treat ment-naive and enrolled into the US PEPFAR-funded Institute of Human Virology Nige ria's AIDS Care and Treatment in Nigeria (ACTION) programme Exclusion criteria: not reported Study design: RCT
Patient characteristics and setting	 Country: Nigeria World Bank Income classification: low-middle income Study setting: hospital-based (Aminu Kano Teaching Hospital, Northern Nigeria) Study dates: August 2006 to January 2008 Age of population (years), mean (SD): 33 (8.13) Gender (male %): 43.17 Participants included/analysed: 421/276 First or second-line regimen: first-line Type of ART: d4T-based, ZDV-based, TDF-based Time on ART at enrolment: treatment-naïve Time on ART at measurement of viral load and adherence: 9 months
Index tests	Number of index tests used: 2

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Coker 2015 (Continued)				
	Types of index tests: self-report and ph	armacy records		
	 Test 1. Self-report questionnaire Validated scale: yes (Mannheime 	ar 2006)		
	 Tool description: self-report adl 	nerence measure wa ex Questionnaire (N	as derived from an interview-ad- lannheimer 2006). Threshold for	
	• Blinding: no information			
	 Threshold prespecified: yes Adherence threshold used: 100% 			
	 Test 2. Tablet counts 	0		
	 Validated scale: not applicable 			
		y days between visit	es. This was calculated as days of s multiplied by 100. A cut-off of <	
	 Blinding: no information 			
	 Threshold prespecified: yes 			
	• Adherence threshold used: 95%			
Target condition and reference stan-	Target condition: viral non-suppressio	n		
dard(s)	 Reference standard: Roche Cobas AmpliPrep TaqMan (Copas Amplicor; tics) 			
	Definition of viral non-suppression:		copies/mL	
	Blinded to index test: no informatio	n		
Flow and timing	 Time interval between index and retime ("We assessed all variables and mL) at the end of the study using the All patients received same reference. Missing data: 70% of original sample sample. Missing data > 10 % 	d risk factors for vira e Chi-square or Fish e standard: yes	Il load suppression (< 400 copies/ er exact test and student t-test")	
Comparative				
Notes	Conflicts of interest: none declared			
	Funding source: Doris Duke Charitable to Dr. William A. Blattner and Abbott In Ndembi			
	Trial registry: not reported			
Methodological quality				
Item	Authors' judgement Risk of	bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Unclear			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate ex- clusions?	Yes			

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Coker 2015 (Continued)			
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the includ- ed patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Index test)			
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre- specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to cor- rectly classify the target condition?	Yes		
Were the reference standard results in- terpreted without knowledge of the re- sults of the index tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the target condition as defined by the refer- ence standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval be- tween index test and reference stan- dard?	Unclear		
Did all patients receive the same refer- ence standard?	Yes		
Were all patients included in the analy- sis?	No		
Could the patient flow have intro- duced bias?		High risk	

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

Davies 2008

Study characteristics	
Patient Sampling	 Target population: HIV-infected children commenced on antiretroviral triple therapy Recruitment: consecutive sample, all children that commenced therapy were eligible an agreed to participate Inclusion criteria: selection criteria for commencement of ART needed to be met, in add tion, the following limited social criteria needed to be met: having an identifiable caregive to administer medication and attend clinic appointments; resident in Cape Town for at leas 3 months; caregiver compliance with last 3 clinic appointments and caregiver willingness to comply with ongoing regular clinic attendance and monitoring. Exclusion criteria: not reported Study design: prospective cohort study
Patient characteristics and setting	 Country: South Africa World Bank Income classification: upper-middle-income Study setting: hospital-based (part of the ART program of the Red Cross Children's Hospital, a tertiary care institution in Cape Town, South Africa) Study dates: July 2002 to January 2004 Age of population (months), median (IQR): 37 (16 to 61) Gender (male %): 57.57 Participants included/analysed: 122/88 First or second-line regimen: first-line Type of ART: the majority of children were commenced on stavudine, 3TC and EFV (children > 10 kg or > 3 years) or RTV (children < 10 kg or < 3 years) as no other PI was readily available in suitable formulation and dosage in South Africa at the time Time on ART at enrolment: treatment-naïve Time on ART at measurement of viral load and adherence: 12 months
Index tests	Number of index tests used: 1
	 Types of index tests: tablet counts Test 1. Tablet counts Validated scale: not applicable Tool description: at every monthly visit for one year, caregivers were requested to return all empty medicine containers and unused medication. A dedicated programme pharma cist measured the amount of unused medication volumetrically for syrups/solutions and by pill count for tablets/capsules. The percentage adherence for each antiretroviral medication was calculated by dividing actual use (determined from returned containers and unused medication) by expected use. Blinding: no information Threshold prespecified: not reported Adherence threshold used: 90%
Target condition and reference standard(s)	 Target condition: viral non-suppression Reference standard: no details provided on the assay used ("Viral load determined using stan dard laboratory methods") Definition of viral non-suppression: HIV viral load > 400 copies/mL Blinded to index test: no information
Flow and timing	 Time interval between index and reference tests: no explicit information on timing; adherence was a composite measure of annual average percentage adherence; viral load taken at 1 months All patients received same reference standard: not reported



Davies 2008 (Continued)

• Missing data: 88/122 children included in the analysis. Missing data > 10%

Comparative	
Notes	Conflicts of interest: none declared
	Funding source: donations to fund the programme were received from Syfrets Trust Ltd, Mer- ck (Pty) Ltd, Bristol-Myers Squibb Foundation, Durbanville High School, and the University of Cape Town. Mary-Ann Davies and Andrew Boulle receive support from the International Epi- demiological Databases to Evaluate AIDS in Southern Africa (IeDEASA) collaboration which is funded by the National Institutes for Health (NIH; U01 AI069924-01).
	Trial registry: not reported

Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sam- ple of patients enrolled?	Yes		
Was a case-control design avoid- ed?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the in- cluded patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Index test)			
Were the index test results inter- preted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre- specified?	No		
Could the conduct or interpreta- tion of the index test have intro- duced bias?		High risk	
Are there concerns that the in- dex test, its conduct, or inter- pretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			

Davies 2008 (Continued)			
Is the reference standards likely to correctly classify the target condi- tion?	Unclear		
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Unclear risk	
Are there concerns that the tar- get condition as defined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Unclear		
Did all patients receive the same reference standard?	Unclear		
Were all patients included in the analysis?	No		
Could the patient flow have in- troduced bias?		High risk	

Duarte 2015

Study characteristics	
Patient Sampling	 Target population: Infants (≤ 12 months of age) who were born to women diagnosed with HIV infection either prior to or during pregnancy or within 1 month postpartum and (ii) HIV-infected infants, children and adolescents (≤ 21 years of age) Recruitment: no details reported
	 Inclusion criteria: infants (≤ 12 months of age) who were born to women diagnosed with HIV infection either prior to or during pregnancy or within 1 month postpartum and (ii) HIV-infected infants, children and adolescents (≤ 21 years of age)
	Exclusion criteria: not reported
	Study design: prospective cohort study
Patient characteristics and setting	 Country: Brazil, Mexico and Peru World Bank Income classification: Brazil (low-income and upper-middle-income during that time), Mexico (upper-middle-income), Peru (low-income) Study setting: clinic-based (14 clinical sites, 12 in Brazil, 1 each in Peru and Mexico) Study dates: 2002 to October 2007 Age of population (years), mean (range): 5.0 (< 1 to 11) Gender (male %): 50.0

Duarte 2015 (Continued)	 Participants included/analysed: 387/361 (index test 1) 387/367 (index test 2) at 6 months; 387/357 (index test 1) and 387/360 (index test 2) at 12 months.
	First or second-line regimen: second-line Turns of ADT 05% of the shidden were on combination ADT
	 Type of ART: 95% of the children were on combination ART Time on ART at enrolment: not reported
	 Time on ART at measurement of viral load and adherence: 6 and 12 months
Index tests	Number of index tests used: 1
	Types of index tests: self-report
	 Test 1. Self-report questionnaire Validated scale: yes
	 Tool description: structured questionnaire developed for use by the U.S. National Institute of Allergy and Infectious Diseases (NIAID) as part of standard practice in PACTG (Pediatric AIDS Clinical Trials Group) studies. The interview was administered in Spanish or Portuguese by a member of the clinical care or research team to the person with primary responsibility for medication administration. The participant/caregiver was asked to identify the ARV medications and number of doses (not number of pills) prescribed each day. ART adherence was derived based on the total number of doses missed during the three-day period prior to a study visit and the total number of expected doses for all of the ARVs included in the participant's treatment regimen at the time of the visit. (1) The measure was expressed in the form of a continuous measure of percent adherence calculated as binary indicator of perfect (100%) adherence. (2) Participants/caregivers were also asked to recall when they/the child last missed a dose of any ARV medication; response options included never, during the previous two weeks, during the last month, over a month ago or don't remember. This measure was dichotomized for purposes of analysis (never vs. ever). Blinding: no information Threshold prespecified: yes Adherence thresholds used: 100% (perfect adherence score: never missed a dose)
Target condition and refer- ence standard(s)	Target condition: viral non-suppression
	 Reference standard: no details provided on the assay used Definition of viral non suppression: UN/viral load > 400 conject(m)
	 Definition of viral non-suppression: HIV viral load > 400 copies/mL Blinded to index test: no information
Flow and timing	 Time interval between index and reference tests: demographic, laboratory, and clinical data were collected at enrolment and every 6 months, including HIV-1 RNA viral load, CD4 measures, CDC classification, and antiretroviral medication adherence
	All patients received same reference standard: not reported
	 Missing data: none. At 6 months, data available for 361/387 and 367/387 for index tests 1 and 2; at 12 months, data available for 357/387 and 360/387 for index tests 1 and 2. Missing data < 10%
	Demographic, laboratory, and clinical data were collected at enrolment and every 6 months, in- cluding HIV-1 RNA viral load, CD4 measures, CDC classification, and antiretroviral medication ad- herence.
Comparative	
Notes	Conflicts of interest: none declared
	Funding source: supported by NICHD Contracts N01-HD-3-3345 (2002–2007), HHSN267200800001C (2007–2012), and HHSN275201300003C (2012–2017)
	Trial registry: not reported
Methodological quality	



Duarte 2015 (Continued)

pre-specified? Could the conduct or inter- pretation of the index test have introduced bias? Are there concerns that the index test, its conduct, or in- terpretation differ from the review question? DOMAIN 3: Reference Standard Is the reference standards like- ly to correctly classify the tar- get condition? Were the reference standard results interpreted without knowledge of the results of the index tests? Unclear Could the reference stan- dard, its conduct, or its inter- pretation have introduced Unclear risk	Authors' judgement Risk of bias Ap	iem	Applicability concerns
sample of patients enrolled? Was a case-control design avoided? Ves Did the study avoid inappro- priate exclusions? Could the selection of pa- tients have introduced bias? Are there concerns that the included patients and set- ting do not match the review question? DOMAIN 2: Index Test (Index test) Unclear terpreted without knowledge of the results of the reference standard? If a threshold was used, was it Yes Could the conduct or inter- pretation of the reference standard Is the reference standard slike- Its or or the reference standard slike- Its or or or the reference standard slike- Its or	yn	OMAIN 1: Patient Selection	
avoided? Did the study avoid inappropriate exclusions? Could the selection of patters and setting do not match the review question? DOMAIN 2: Index Test (Index test) Were the index test results interrepreted without knowledge of the results of the reference standard? If a threshold was used, was it prespecified? Could the conduct or interpretation of the index test have introduced bias? Are there concerns that the index test results interpretation of the index test are suits into the reference standard? If a threshold was used, was it prespecified? Could the conduct or interpretation of the index test have introduced bias? Are there concerns that the index test into the reference standard? If a threshold was used, was it prespecified? Could the conduct or interpretation of the index test have introduced bias? Are there concerns that the index test is the reference standard like- ly to correctly classify the tar- get condition? Unclear results of the reference standard like- ly to correctly classify the tar- get condition? Unclear results interpreted without knowledge of the results of the interview question?			
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tients have introduced bias? Are there concerns that the included patients and setting do not match the review question? DOMAIN 2: Index Test (Index test) Were the index test results interpreted without knowledge of the results of the reference standard? If a threshold was used, was it pre-specified? Could the conduct or interpretation of the index test have introduced bias? Are there concerns that the review question? DOMAIN 3: Reference Standard Is the reference standards like- Unclear Were the reference standard without knowledge of the results of the result	- Yes		
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terpreted without knowledge of the results of the reference standard? If a threshold was used, was it pre-specified? Could the conduct or inter- pretation of the index test have introduced bias? Are there concerns that the index test, its conduct, or in- terpretation differ from the review question? DOMAIN 3: Reference Standard Is the reference standards like- ly to correctly classify the tar- get condition? Were the reference standard Were the reference standard Unclear results interpreted without knowledge of the results of the index tests? Could the reference stan- dard, its conduct, or its inter- pretation have introduced	ex test)	OMAIN 2: Index Test (Index tes	
pre-specified? Could the conduct or inter- pretation of the index test have introduced bias? Are there concerns that the index test, its conduct, or in- terpretation differ from the review question? DOMAIN 3: Reference Standard Is the reference standards like- ly to correctly classify the tar- get condition? Unclear Were the reference standard results interpreted without knowledge of the results of the index tests? Unclear Could the reference stan- dard, its conduct, or its inter- pretation have introduced Unclear risk	je	erpreted without knowledge f the results of the reference	
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ly to correctly classify the tar- get condition? Were the reference standard Unclear results interpreted without knowledge of the results of the index tests? Could the reference stan- dard, its conduct, or its inter- pretation have introduced	dard	OMAIN 3: Reference Standard	
results interpreted without knowledge of the results of the index tests? Could the reference stan- dard, its conduct, or its inter- pretation have introduced		to correctly classify the tar-	
dard, its conduct, or its inter- pretation have introduced		esults interpreted without nowledge of the results of the	
bias?	er-	ard, its conduct, or its inter- retation have introduced	
Are there concerns that the Unclear Unclear target condition as defined			Unclear



Duarte 2015 (Continued)

by the reference standard does not match the question?

DOMAIN 4: Flow and Timing	
Was there an appropriate in- terval between index test and reference standard?	Unclear
Did all patients receive the same reference standard?	Unclear
Were all patients included in the analysis?	Yes
Could the patient flow have introduced bias?	Unclear risk

Dziva 2017

Study characteristics	
Patient Sampling	 Target population: children aged 6 to 15 years newly diagnosed with HIV infection attending lay worker-delivered treatment support intervention to improve adherence Recruitment: all children who initiated ART were included. Inclusion criteria: children aged 6-15 years newly diagnosed with HIV infection Exclusion criteria: no exclusions reported: all children who initiated ART were included. Study design: prospective clinical trial
Patient characteristics and setting	 Country: Zimbabwe World Bank Income classification: low-income Study setting: community-based (lay worker-delivered treatment support intervention) Study dates: not reported Age of population (years), median (IQR): 11 (9 to 13) Gender (male %): 45 Participants included/analysed: 237/166 First or second-line regimen: first-line Type of ART: HIV treatment was provided according to national guidelines. Time on ART at enrolment: not reported Time on ART at measurement of viral load and adherence: 48 weeks
Index tests	 Number of index tests used: 1 Types of index tests: self-report Test 1. Self-report questionnaire Validated scale: yes (VAS) Tool description: participants completed a visual analog scale (VAS) to self-assess their adherence over the past month. The responses were given by either caregivers on behalf of the child, jointly by caregivers and children, or children alone (for older children). The authors noted that the scales routinely used to measure

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

Dziva 2017 (Continued)				
		ponding scales for childre ation fied: not reported	designed for use among adults, and n. Non-adherent = VAS score < 95%	
Target condition and reference standard(s)	Target condition: viral no	n-suppression		
	v2.0.11	suppression: HIV viral loa	oliPrep/COBAS TaqMan HIV-1 Test, d > 400 copies/mL	
Flow and timing	but both assessed at 4All patients received siMissing data: none. No	8 weeks after initiation of ame reference standard: y t all children were include		
Comparative				
Notes	Conflicts of interest: none	edeclared		
	Funding source: Wellcome Trust			
	Trial registry: not reporte	d		
Methodological quality				
ltem	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Yes			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate exclu- sions?	Yes			
Could the selection of patients have in- troduced bias?		Low risk		
Are there concerns that the included pa- tients and setting do not match the re- view question?			High	
DOMAIN 2: Index Test (Index test)				
Were the index test results interpreted without knowledge of the results of the ref- erence standard?	Unclear			
If a threshold was used, was it pre-speci- fied?	Unclear			

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Dziva 2017 (Continued)			
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correct- ly classify the target condition?	Yes		
Were the reference standard results inter- preted without knowledge of the results of the index tests?	Unclear		
Could the reference standard, its con- duct, or its interpretation have intro- duced bias?		Low risk	
Are there concerns that the target con- dition as defined by the reference stan- dard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Unclear		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	No		
Could the patient flow have introduced bias?		High risk	

Ekstrand 2010

Study characteristics	
Patient Sampling	 Target population: HIV patients, on anti-retroviral medication for at least one month in our patient Department of Medicine in a Catholic hospital in Bangalore, India Recruitment: referral from physician needed or Non Governmental Organization
	 Inclusion criteria: eligibility criteria included being at least 18 years old; capable of commun cating in English, Kannada, Tamil, or Telugu; being HIV-infected, on antiretroviral medicatio for at least one month, and willing to participate in all follow-up visits Exclusion criteria: not reported Study design: pospective cohort study
Patient characteristics and setting	 Country: India Country: India World Bank Income classification: low-middle-income Study setting: clinic-based (the outpatient Department of Medicine in a Catholic hospita)
	 in Bangalore, India) Study dates: not reported (prior to 2008)

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

Ekstrand 2010 (Continued)	 Age of population (years): mean (range): 38 (23 to 74) Gender (male %): 69 Participants included/analysed: 229/202 First or second-line regimen: not reported Type of ART: virtually all (98%) of the participants were on an NTI-based regimen, with the most common regimens being 3TC/stavudine/NVP (49%), followed by 3TC/AZT/NPV (26%), 3TC/AZT/EFV (8%), and 3TC/stavudine/EFV(7%) Time on ART at enrolment: not reported Time on ART at measurement of viral load and adherence: at least 12 months (mean: 33
Index tests	months, range: 13 to 145 months) Number of index tests used: 4
	Types of index tests: self-report
	 Test 1. Self-report questionnaire Validated scale: yes Tool description: visual analog scale in last month Blinding: no information Threshold prespecified: not reported Adherence threshold used: 95% Test 2. Self-report questionnaire Validated scale: not reported Tool description: self-reported pills missed in previous 1 month Blinding: no information Threshold prespecified: not reported Adherence threshold used: 95% Test 3. Self-report questionnaire Validated scale: not reported Adherence threshold used: 95% Test 3. Self-report questionnaire Validated scale: not reported Adherence threshold used: 95% Test 3. Self-report questionnaire Validated scale: not reported Tool description: self-reported pills missed in previous 1 week Blinding: no information Threshold prespecified: not reported Adherence threshold used: 95% Test 4. Self-report questionnaire Validated scale: not reported Adherence threshold used: 95% Test 4. Self-report questionnaire Validated scale: not reported Adherence threshold used: 95% Test 4. Self-report, % adherence in previous 4 days (detailed dose-by-dose assessment) Blinding: no information Threshold prespecified: not reported
Target condition and reference standard(s)	 Target condition: viral non-suppression Reference standard: real Time PCR assay with fluorescein labeled Taqman probe for quantitation of HIV particles
	 Definition of viral non-suppression: HIV viral load > 100 copies/mL Blinded to index test: no information
Flow and timing	 Time interval between index and reference tests: index tests and reference standard were conducted at the same study visits (at 0, 6 and 12 months). Adherence data were self-reported percentages of doses completed in previous 4 days, 1 week and 1 month. All patients received same reference standard: yes Missing data: 202 participants who attended study visits at 0 and 12 months were tested. Or 229 enrolled, 11 died and 15 were lost to follow-up, leaving 203. It is unclear what happened to be a standard to be a standard.

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Ekstrand 2010 (Continued)

one participant. Missing participants > 10%, but reasons were reported and many participants were lost to follow-up because they died.

		· · · · · · · · · · · · · · · · · · ·		
Comparative				
Notes	Conflicts of interest: none of	declared		
	Funding source: grant R01MH067513 from the National Institute of Mental Health (NIMH) (Bethesda, MD, USA). (from refID 5045 Ekstrand 2011)			
	Trial registry: not reported			
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sam- ple of patients enrolled?	No			
Was a case-control design avoid- ed?	Yes			
Did the study avoid inappropriate exclusions?	Unclear			
Could the selection of patients have introduced bias?		Unclear risk		
Are there concerns that the in- cluded patients and setting do not match the review question?			Unclear	
DOMAIN 2: Index Test (Index test)				
Were the index test results inter- preted without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre- specified?	Yes			
Could the conduct or interpreta- tion of the index test have intro- duced bias?		Unclear risk		
Are there concerns that the in- dex test, its conduct, or inter- pretation differ from the review question?			Low concern	
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condi-tion?	Yes			

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

Ekstrand 2010 (Continued)			
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the tar- get condition as defined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Could the patient flow have in- troduced bias?		Low risk	

El-Khatib 2010

Study characteristics	
Patient Sampling	 Target population: adults with HIV, being on ART for at least 12 months Recruitment: recruited through posters in two outpatient clinics. The following inclusion criteria were applied: at least 18 years old; being on ART for at least 12 months and consenting to participate in the study Inclusion criteria: at least 18 years old; being on ART for at least 12 months; and consenting to participate in the study Exclusion criteria: not reported Study design: cross-sectional
Patient characteristics and setting	 Country: South Africa World Bank Income classification: upper-middle Study setting: two outpatient clinics at the Chris Hani Baragwanath Hospital Soweto, Johannesburg Study dates: March to December 2008 Age of population (years), median: women: 41, men: 37 Gender (male %): 26.87 Participants included/analysed: 998/997 First or second-line regimen: first-line or second-line Type of ART: NNRTI-based (first-line), PI-based (second-line) Time on ART at enrolment: at least 12 months



El-Khatib 2010 (Continued)

• Time on ART at measurement of viral load and adherence: not reported; at least 12 months

Could the selection of patients have in- troduced bias?		High risk		
Did the study avoid inappropriate exclu- sions?	Unclear			
Was a case-control design avoided?	Yes			
Was a consecutive or random sample of patients enrolled?	No			
DOMAIN 1: Patient Selection				
Item	Authors' judgement	Risk of bias	Applicability concerns	
Methodological quality				
	Trial registry: not reporte	d		
	Funding source: Swedish International Development Cooperation Agency (Sida) to Z.EK. and NICD and a Karolinska Institutet faculty award (KID) to Z.EK.; African Pro- gramme for Training in HIV/TB Research Fogarty			
Notes	Conflicts of interest: none	edeclared		
Comparative				
	 All patients received same reference standard: yes Missing data: adherence/suppression reported for 997/998. Missing data < 10 % 			
Flow and timing	but this was a cross-se	ctional study so likely to	ts: no explicit information on timinន be measured simultaneously	
	Blinded to index test:			
	Basel, Switzerland)Definition of viral non-	suppression: HIV viral loa	ad > 400 copies/mL	
	-		st, v1.5 (Roche Molecular Diagnostics	
Target condition and reference standard(s)	· Target condition: viral no	n-suppression		
	 Adherence thresho 	ld used: 100%		
	Blinding: no informThreshold prespeci			
	end"		e: "Missed any pills during last week	
	 Tool description: detail about adherence measure: "adherence during the previ- ous weekend which served as a proxy for recent adherence"Threshold cate- 			
	 Test 1. Self-report Validated scale: not 	-		
	Types of index tests: self-report			
Index tests	Number of index tests use			

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



l-Khatib 2010 (Continued)			
Are there concerns that the included pa- tients and setting do not match the re- view question?			Unclear
DOMAIN 2: Index Test (Index test)			
Were the index test results interpreted without knowledge of the results of the ref- erence standard?	Unclear		
If a threshold was used, was it pre-speci- fied?	Unclear		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correct- ly classify the target condition?	Yes		
Were the reference standard results inter- preted without knowledge of the results of the index tests?	Unclear		
Could the reference standard, its con- duct, or its interpretation have intro- duced bias?		Low risk	
Are there concerns that the target con- dition as defined by the reference stan- dard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Yes		
Could the patient flow have introduced bias?		Low risk	

Evans 2016

Study characteristics	
Patient Sampling	 Target population: adult, HIV-positive patients on second-line ART, who experienced a single elevated viral load



vans 2016 (Continued)	
	 Recruitment: no details reported Inclusion criteria: Eligible patients were adult (> 18 years) HIV-positive patients at Them ba Lethu who were receiving a second-line ART regimen containing lopinavir/ritonavi or atazanavir/ritonavir and experienced a single elevated viral load (> 400 copies/mL) o second-line ART. Exclusion criteria: not reported Study design: cohort study
Patient characteristics and setting	 Country: South Africa World Bank Income classification: upper-middle-income Study setting: clinic-based (Themba Lethu Clinic in Johannesburg) Study dates: July 2011 to July 2018 Age of population (years), median (IQR): 37.6 (33.6 to 45.3) Gender (male %): 40.8 Participants included/analysed:49/49 First or second-line regimen: second-line Type of ART: second-line ART regimen containing LPV/r or atazanavir/ritonavir Time on ART at enrolment: prior to study eligibility (elevated VL on second-line), median (IQR): 48.8 months (30.4 to 68.8) Time on ART at measurement of viral load and adherence: 3 or 6 months after an elevated viral load (> 400 copies/mL) on second-line ART
Index tests	Number of index tests used: 1 Types of index tests: electronic monitoring
	 Test 1. Electronic monitoring Validated scale: not applicable Tool description: patients in the intervention cohort used an electronic adherence monitoring device (EAMD) (WisepillTM) Blinding: no information Threshold prespecified: not reported Adherence threshold used: 80% (missing ≥ 20%) or taking at least 95% (missing ≥ 5% of the prescribed medication
Target condition and reference stan- dard(s)	Target condition: viral non-suppression
	 Reference standard: no details provided on the assay used Definition of viral non-suppression: HIV viral load > 400 copies/mL Blinded to index test: yes. Viral load testing was not done by the clinic, but by a centra lab and therefore those performing the viral load tests were blinded to the study cohorts Blood samples are sent to the National Health Laboratory Service (NHLS) and viral loa and CD4 count results are uploaded directly into TherapyEdge-HIVTM from the NHLS o a daily basis.
Flow and timing	 Time interval between index and reference tests: when participants returned for the fo low-up viral load test (3 to 6 months after enrolment), the device was returned and th clinician reviewed the adherence data with the patient. All patients received same reference standard: not reported Missing data: none. All eligible participants with viral load test and adherence measure were included in the main analysis.
Comparative	
	Conflicts of interest: none declared

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Evans 2016 (Continued)

Funding source: South Africa Mission of the US Agency for International Development (USAID)

Trial registry: not reported

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate ex- clusions?	Unclear		
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the includ- ed patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Index test)			
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre- specified?	Unclear		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear
DOMAIN 3: Reference Standard			
Is the reference standards likely to cor- rectly classify the target condition?	Unclear		
Were the reference standard results in- terpreted without knowledge of the re- sults of the index tests?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the target condition as defined by the refer-			Low concern

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Evans 2016 (Continued) ence standard does not match the question?

DOMAIN 4: Flow and Timing	
Was there an appropriate interval be- tween index test and reference stan- dard?	Yes
Did all patients receive the same refer- ence standard?	Unclear
Were all patients included in the analy- sis?	Yes
Could the patient flow have intro- duced bias?	Low risk

Farley 2003

Study characteristics	
Patient Sampling	 Target population: caregivers (biologic parent, adoptive parent, foster parent, or other guardian) of perinatally HIV-infected children under the age of 13 years being treated with HAART
	 Recruitment: caregivers (biologic parent, adoptive parent, foster parent, or other guardian) of perinatally HIV-infected children under the age of 13 years being treated with HAART were invited to participate in a 3-year study of adherence involving periodic interviews and a 6- month baseline period of observation.
	 Inclusion criteria: caregivers (biologic parent, adoptive parent, foster parent, or other guardian) of perinatally HIV-infected children under the age of 13 years being treated with HAART were invited to participate in a 3-year study of adherence involving periodic interviews and a 6-month baseline period of observation. All children were receiving treatment at the University of Maryland School of Medicine.
	 Exclusion criteria: children with evidence of significant developmental delays or greater, below the mean on the Test of Nonverbal Intelligence, a severe physical handicap precluding independent ambulation, or those who were receiving all antiretrovirals in liquid formulation were not eligible to participate. Study design: prospective cohort study
Patient characteristics and setting	 Country: USA World Bank Income classification: high-income
	 Study setting: University of Maryland School of Medicine
	Study dates: recruitment from October 1998 to October 2002
	• Age of population (years), mean: 6.9
	Gender (male %): 65
	 Participants included/analysed: 31/26
	 First or second-line regimen: not reported Type of ART: defined as treatment with three different antiretroviral agents regardless of drug class
	 Time on ART at enrolment: at least 6 months
	• Time on ART at measurement of viral load and adherence: not reported; at least 6 months
Index tests	Number of index tests used: 1

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

Farley 2003 (Continued)			
	Types of index tests: electr	onic monitoring	
	 Test 1. Electronic monit Validated scale: not a 	-	
		ence rate was calculated as fo ed for the interval tion ed: yes	; System child-resistant Track Caps was llows: medication events or bottle open-
			acy refill adherence assessment and a way that data could be used.
Target condition and reference	Target condition: viral non	-suppression	
standard(s)	Reference standard: Ro	che Amplicor reverse transcr	ibed PCR method
		uppression: HIV viral load > 4	00 copies/mL
	Blinded to index test: no	DINTORMATION	
Flow and timing	 Time interval between i sured at 6 months 	ndex and reference tests: no	explicit details on timing, but both mea-
		me reference standard: yes	
	period of the study with included in this analysi	greater than 60 days of availa s. Of the 5 noncompleters, 1	nrolled; 26 completed the initial 6-month ble adherence monitoring data and were did not complete the baseline interview ng MEMS monitoring. Missing data for >
Comparative			
Notes	Conflicts of interest: none	declared	
	Funding source: R01 HD36	613 and M01 RR165001	
	Trial registry: not reported		
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sam- ple of patients enrolled?	Unclear		
Was a case-control design avoid- ed?	Yes		
Did the study avoid inappropriate exclusions?	Unclear		
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the in- cluded patients and setting do not match the review question?			Low concern

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Trusted evidence. Informed decisions. Better health.

Farley 2003 (Continued)			
DOMAIN 2: Index Test (Index test)			
Were the index test results inter- preted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre- specified?	Unclear		
Could the conduct or interpreta- tion of the index test have intro- duced bias?		Unclear risk	
Are there concerns that the in- dex test, its conduct, or inter- pretation differ from the review question?			Unclear
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condi-tion?	Yes		
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the tar- get condition as defined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Unclear		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	No		
Could the patient flow have in- troduced bias?		High risk	



Study characteristics

Fokam 2017

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Patient Sampling	Target population: adolescents living with HIV
	 Recruitment: consecutive sampling at a referral health facility for care and treat ment of HIV-infected children. Children followed up at the study site were all HIV vertically infected, except for one that was infected through unsafe blood transfu sion.
	 Inclusion criteria: eligibility criteria were: every adolescent living with HIV who: (a was aware of his HIV status, (b) was registered for ART monitoring at the study site (c) was receiving ART for at least six months, (d) was capable of responding to the study questionnaire, and (e) had provided a written consent
	Exclusion criteria: not reported
	Study design: cross-sectional
Patient characteristics and setting	 Country: Cameroon World Bank Income classification: low-middle-income
	 Study setting: National Social Insurance Fund Health Centre in Yaounde Cameroon - a referral health facility for care and treatment of HIV-infected chil dren
	Study dates: January to May 2016
	• Age of population (years), median (IQR): 13 (11 to 16)
	• Gender (male %): 48
	Participants included/analysed: 145/145
	 First or second-line regimen: mostly frst-line Type of ART: 92% were on first-line ART
	 Time on ART at enrolment: at least 6 months
	• Time on ART at measurement of viral load and adherence: at least 6 months
Index tests	Number of index tests used: 1
	Types of index tests: self-report
	 Test 1. Self-report questionnaire Validated scale: not reported
	 Tool description: poor adherence was defined as missing one dose of ART dur ing the past 14 days
	 Blinding: no information
	• Threshold prespecified: not reported
	 Adherence threshold used: 100%
Target condition and reference standard(s)	Target condition: viral non-suppression
	 Reference standard: Abbott Applied Biosystem m2000RT Real Time PCR AE m2000RT
	 Definition of viral non-suppression: HIV viral load > 50 copies/mL
	Blinded to index test: no information
Flow and timing	 Time interval between index and reference tests: index tests and reference stan dard were conducted at the same study visits
	 All patients received same reference standard: yes
	 Missing data: none. All eligible participants with viral load test and adherence mea sures were included in the main analysis.
Comparative	
Notes	Conflicts of interest: none declared

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Fokam 2017 (Continued)

Funding source: the authors received no specific funding for this work.

Trial registry: not reported

Methodological	quality
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Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of pa- tients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Unclear		
Could the selection of patients have intro- duced bias?		Unclear risk	
Are there concerns that the included pa- tients and setting do not match the review question?			Unclear
DOMAIN 2: Index Test (Index test)			
Were the index test results interpreted with- out knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Unclear		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpret- ed without knowledge of the results of the in- dex tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the target condi- tion as defined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Fokam 2017 (Continued)	
Was there an appropriate interval between in- dex test and reference standard?	Yes
Did all patients receive the same reference standard?	Yes
Were all patients included in the analysis?	Yes
Could the patient flow have introduced bias?	Low risk

Gill 2010

Study characteristics	
Patient Sampling	 Target population: adults with HIV receiving ART Recruitment: this analysis used longitudinal observational data from a three-phase adher ence study conducted among HIV-positive patients receiving ART at the Dermatology and STD clinic, Dali Second People's Hospital (DSPH) in Dali, Yunnan Province, China. Patients were eligible for participation if they were aged > 18 years and agreed to all study proce dures. Of 97 eligible patients at the clinic, 80 agreed to participate. Inclusion criteria: patients aged 18 years or older and agreed to all study procedures Exclusion criteria: not reported Study design: prospective cohort study
Patient characteristics and setting	 Country: China World Bank Income classification: low-middle Study setting: clinic-based (Sexually Transmitted disease clinic in a hospital) Study dates: June 2006 to May 2007 Age of population (years), mean (SD): 35.7 (8.1) Gender (male %): 73.9 Participants included/analysed: 69/65 First or second-line regimen: unclear Type of ART: twice-daily regimen of nevirapine or efavirenz, plus either zidovudine and lamivudine or lamivudine and stavudine Time on ART at enrolment: median 8.3 months Time on ART at measurement of viral load and adherence: median 8.3 months
Index tests	 Number of index tests used: 4 Types of index tests: self-report, tablet counts, electronic monitoring Test 1. Self-report questionnaire Validated scale: yes (VAS) Tool description: self-report/visual analog scale: indicated by where the patient marked an 'X' on the 0–100% VAS scale Blinding: no information Threshold prespecified: not reported Adherence threshold used: 95% Test 2. Tablet counts Validated scale: not applicable Tool description: [Actual number of pills in bottle]/[expected number of pills] Blinding: no information Threshold prespecified: not reported



Gill 2010 (Continued)			
	 ber of bottle opening Blinding: no informat Threshold prespecifie Adherence threshold Test 4. Electronic morito Validated scale: not a Tool description: prop 	ring pplicable portion taken: [Actual numl s] cion ed: not reported used: 95% ring pplicable portion taken within dose ti scribed time]/[expected nu cion ed: not reported	per of bottle openings]/[expected num ime: [Actual number of bottle opening umber of openings]
Target condition and reference stan- dard(s)	lands)	ganon Teknica NucliSens uppression: HIV viral load >	analyzer (BioMerieux, Boxtel, Nether 400 copies/mL
Flow and timing	 Time interval between index and reference tests: not explicitly reported, but both measured at 6 months All patients received same reference standard: yes Missing data: of 69 patients, 65 included in the analysis. Missing data < 10 % 		
Comparative			
Notes	Conflicts of interest: none c Funding source: Boston Un Trial registry: not reported		GAP/China, NIH
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate ex- clusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the in- cluded patients and setting do not match the review question?			Low concern

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



ill 2010 (Continued)			
DOMAIN 2: Index Test (Index test)			
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre- specified?	Unclear		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condi-tion?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the target condition as defined by the refer- ence standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval be- tween index test and reference stan- dard?	Unclear		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Yes		
Could the patient flow have intro- duced bias?		Low risk	

Haberer 2011

Study characteristics		
Patient Sampling	Target population: HIV-infected children initiating ART	
Accuracy of measures for	or antiretroviral adherence in people living with HIV (Review)	78



Haberer 2011 (Continued)	
	• Recruitment: the study population was drawn from the CHAPAS-1 trial, which was a randomized study of nevirapine (NVP) dose escalation among HIV-infected children initiating ART. All children were treated at the University Teaching Hospital in Lusaka, Zambia.
	 Inclusion criteria: 1. Aged 3 months to 14 years inclusive; 2. Less than 30 kg in weight; 3. Carers and children where they were appropriate, willing and able to give informed consent; 4. HIV-infected, as determined by: a. Two separate HIV-antibody enzyme-linked immunosorbent assay (ELISA) or rapid tests on the same sample in children > 18 months, b. Two positive proviral DNA tests taken on separate samples in children < 18 months; 5. Previously untreated with antiretrovirals, including any ART given to prevent mother-to-child transmission; 6. Fulfilling one of the World Health Organization (WHO) criteria for initiating treatment: a. WHO paediatric stage 4 or severe stage 3 disease regardless of CD4%, b. CD4 per cent < 15% if > 18 months of age, or < 20% if < 18 months of age, c. WHO paediatric stage 2 disease with consideration of CD4 percentage (< 15% for children > 18 months; < 20% for children < 18 months). (Note current WHO guidelines are under review and the above criteria may be changed, particularly by raising the CD4 percentage cut-off to 25% in children < 18 months; inclusion criteria would be changed accordingly for children to start ART in CHAPAS 1 trial)
	 Exclusion criteria: 1. Cannot or unwilling to regularly attend the CHAPAS clinic; 2. Severe laboratory abnormalities (contraindicating NVP-based regimen), i.e. serum creatinine > 5 times upper limit of normal (ULN) or aspartate aminotransferase or alanine aminotransferase > 10 times ULN; 3. Active opportunistic infection and/or serious bacterial infection at the time of study entry including tuberculosis (may be enrolled after the acute phase of tuberculosis); 4. Current treatment with any medication known to be contraindicated with any of the drugs prescribed for the patient's ART-therapy in this trial, including rifampicin Study design: prospective cohort substudy within an RCT
Patient characteris- tics and setting	 Country: Zambia World Bank Income classification: low-income Study setting: clinic-based (outpatients at University Teaching Hospital in Lusaka) Study dates: May 2006 to Dec 2008
	 Age of population (years), median (IQR): 6 (2 to 9) Gender (male %): 55
	Participants included/analysed: 96/73, 96/72
	 First or second-line regimen: first-line Type of ART: children randomized to initiate nevirapine (NVP) at full dose used fixed-dose combination (FDC) tablets of stavudine, 3TC, and NVP (Triomune Baby/Junior) twice daily. Children randomized to escalate their dose of NVP used Triomune Baby/Junior once daily for 14 days, together with an FDC of stavudine and 3TC (Lamivir-S) once daily. After 14 days Lamivir-S was stopped and children continued on twice daily Triomune Baby/Junior. Time on ART at enrolment: treatment-naïve Time on ART at measurement of viral load and adherence: 48 weeks
Index tests	Number of index tests used: 5
	Types of index tests: electronic monitoring, tablet counts, self-report
	 Test 1. Electronic monitoring Validated scale: not applicable Tool description: electronic monitoring with MEMS (Medication Event Monitoring System, Aardex, Switzerland) caps Blinding: no information Threshold prespecified: not reported Adherence threshold used: 95%; 80% Test 2. Tablet counts Validated scale: not applicable Tool description: home visits pill counts. Unannounced monthly home visits for further pill counts Blinding: no information Threshold prespecified: not reported
	Test 3. Tablet counts



Haberer 2011 (Continued)

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Haberer 2011 (Continued)			
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclu- sions?	Yes		
Could the selection of patients have in- troduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the re- view question?			Low concern
DOMAIN 2: Index Test	(Index test)		
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre- specified?	Unclear		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or in- terpretation dif- fer from the review question?			Unclear
DOMAIN 3: Reference	Standard		
Is the reference stan- dards likely to cor- rectly classify the tar- get condition?	Yes		
Were the reference standard results in- terpreted without knowledge of the results of the index tests?	Unclear		
Could the reference standard, its con- duct, or its interpre-		Low risk	



Low concern

Haberer 2011 (Continued) tation have introduced bias?

Are there concerns that the target condition as defined by the reference standard does not match the question?

DOMAIN 4: Flow and Timing

Was there an appro- priate interval be- tween index test and reference standard?	Unclear
Did all patients re- ceive the same refer- ence standard?	Yes
Were all patients in- cluded in the analy- sis?	No
Could the patient flow have intro- duced bias?	High risk

Hassan 2014

Study characteristics	
Patient Sampling	 Target population: HIV-infected adolescents and adults (≥ 15 years old) who had been on first-line ART for more than six months.
	• Recruitment: in the first cross-section, all consenting eligible participants were recruited between November 2008 and January 2009. At the same time, a prospective cohort was established in order to describe long-term outcomes of new clients enrolling for HIV care. All available plasma samples from participants recruited in the prospective cohort and meeting our eligibility criteria as at March 2011 were cross-sectionally retrieved.
	 Inclusion criteria: HIV-infected adolescents and adults (≥ 15 years old) who had been on first-line ART for more than six months
	• Exclusion criteria: participants with a previous history of ART exposure for prevention of moth- er-to-child transmission (PMTCT) or for post-exposure prophylaxis (PEP), and those on sec- ond-line regimens were excluded from the study.
	Study design: cross-sectional
Patient characteristics and setting	 Country: Kenya World Bank Income classification: low-income Study setting: hospital-based, rural setting Study dates: Nov 2008 to March 2011 Age of population (years), median (IQR): 36.5 (31.4 to 44.4) Gender (male %): 18.6 Participants included/analysed: 232/224

DOMAIN 1: Patient Selection			
Item	Authors' judgement	Risk of bias	Applicability concerns
Methodological quality			
	Trial registry: not reported		
	ed by Wellcome Trust fellow employees of the KEMRI/We Kenyan Ministry of Health. E was financially supported b	ships (WT089351MA and WT08 Ilcome Trust research prograr JS was funded by the Internat y the Health Protection Agency valuate Resistance (PASER), w	83579MA, respectively). SM and HN were nme while CAO was an employee of the cional AIDS Vaccine Initiative while PAC y, UK. TFRW was a member of the Phar- vhich received financial support from the
Notes	Conflicts of interest: none d		er WT089351MA). ASH and JAB were fund-
Comparative			
	 Missing data: 224 of 232 	patients included in the analys	sis. Missing data < 10 %
	 All patients received sam 		
Flow and timing	a cross-sectional study s retrieved from 12 months	o likely to be measured simul	xplicit information on timing, but this wa Itaneously. Pharmacy drug refill data wa Ition if follow-up period < 12 months) prio
	• Blinded to index test: no	information	
		d to determine virus concentra ppression: HIV viral load > 400	
	plex real time quantitativ	e probe-based assay with an	e using an inhouse assay; in brief, a multi internal control and a series of quantified
Target condition and refer- ence standard(s)	Target condition: viral non-	suppression	
	Threshold prespecifieAdherence threshold	-	
	 Blinding: no information Threshold prespecifie 		
	fore retrospectively re	etrieved pharmacy drug refill	data from 12 months (or from the date o to the date of sampling for every individua
	ART divided by the tim computed, subtracted	e between drug pickups for all d from 100% and stratified to	quivalent number of days in possession o visits. A mean MPR for each individual wa satisfactory (≥ 95%) and unsatisfactory (· conventions. Note that the authors there
	Validated scale: not aTool description: Med	oplicable icine Possession Ratios (MPRs) were calculated as proportions of the to
	Types of index tests: pharma	acy records :: Medicine Possession Ratio (N	
Index tests	Number of index tests used:		
	• Time on ART at measu	irement of viral load and adhe	erence: not reported; at least 6 months
	 First or second-line regin Type of ART: first-line Time on ART at enrolr 	regimen, not further specified	

	Cochrane
Y V	Library

Hassan 2014 (Continued)			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inappro- priate exclusions?	Yes		
Could the selection of pa- tients have introduced bias?		Low risk	
Are there concerns that the included patients and set- ting do not match the review question?			Low concern
DOMAIN 2: Index Test (Index te	est)		
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or inter- pretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or in- terpretation differ from the review question?			Unclear
DOMAIN 3: Reference Standard	d		
Is the reference standards like- ly to correctly classify the tar- get condition?	Unclear		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Could the reference stan- dard, its conduct, or its inter- pretation have introduced bias?		Unclear risk	
Are there concerns that the target condition as defined by the reference standard does not match the ques- tion?			Unclear

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

Hassan 2014 (Continued)

DOMAIN 4: Flow and Timing	
Was there an appropriate in- terval between index test and reference standard?	Yes
Did all patients receive the same reference standard?	Yes
Were all patients included in the analysis?	Yes
Could the patient flow have introduced bias?	Low risk

Jayaweera 2003

Study characteristics		
Patient Sampling	 Target population: adults with HIV, naïve to ARV Recruitment: no details reported Inclusion criteria: HIV infection, patients had to be naive to ARV, there were no limits set on HIV-1 RNA or CD4 cell counts. All patients received ritonavir 400 mg and indinavir 400 mg two times a day Exclusion criteria: not reported Study design: open-label, non-randomized, single-arm study 	
Patient characteristics and setting	 Country: USA World Bank Income classification: high-income Study setting: hospital-based, Miami city Study dates: not reported Age of population (years): not reported Gender (male %): 58 Participants included/analysed: 19/19 First or second-line regimen: first-line Type of ART: all patients received ritonavir 400 mg and indinavir 400 mg two times a day Time on ART at enrolment: treatment-naïve Time on ART at measurement of viral load and adherence: 24 weeks 	
Index tests	 Number of index tests used: 1 Types of index tests: composite measure Test 1. Self-report questionnaire and pill counts Validated scale: not applicable Tool description: patients were determined to be compliant or non-complia based on self-report with confirmation based on assessment of pill counts each study visit. No other details provided Blinding: no information Threshold prespecified: not reported Adherence threshold used: 100%; ≥ 80% 	
Target condition and reference standard(s)	Target condition: viral non-suppression	

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Jayaweera 2003 (Continued)	Reference standard: Ar	nplicor HIV-1 monitor te	st (Roche, New Jersey, USA)	
	 Definition of viral non-suppression: HIV viral load > 400 copies/mL Blinded to index test: no information 			
Flow and timing	 ment for this study wa patients has laboratory tistical analyses have b this study. All patients received sa 	s intended to be 48 wee y measurements beyond een restricted to data co me reference standard: eligible participants with	ts: Although the duration of treat- eks, only one of the noncompliant I the week-24 visit. Therefore, sta- llected during the first 24 weeks of yes viral load test and adherence mea-	
Comparative				
Notes	Conflicts of interest: none	declared		
	Funding source: Abbott La	boratories provided fina	ancial support	
	Trial registry: not reported	1		
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of pa- tients enrolled?	Unclear			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate exclusions?	Unclear			
Could the selection of patients have intro- duced bias?		Unclear risk		
Are there concerns that the included pa- tients and setting do not match the review question?			Low concern	
DOMAIN 2: Index Test (Index test)				
Were the index test results interpreted with- out knowledge of the results of the reference standard?	Unclear			
If a threshold was used, was it pre-specified?	Unclear			
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk		
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear	
DOMAIN 3: Reference Standard				

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Jayaweera 2003 (Continued)			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpret- ed without knowledge of the results of the in- dex tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the target condi- tion as defined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between in- dex test and reference standard?	Unclear		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Yes		
Could the patient flow have introduced bias?		Low risk	

Jiamsakul 2014

Study characteristics	
Patient Sampling	 Target population: HIV-infected people, treatment-experienced switching to second-line ART due to failure
	• Recruitment: patients were selected from The Therapeutics, Research, Education and AIDS Training in Asia (TREAT Asia) Studies to Evaluate Resistance Monitoring Study (TASER-M). TASER-M began recruitment in 2007 and included 12 clinical sites in Thailand, Hong Kong, Malaysia, Philippines and Indonesia.
	• Inclusion criteria: patients were either enrolled in TASER-M as treatment-naive or initiating first- line ART (not included in the analysis) or treatment-experienced switching to second-line ART due to failure.
	 Exclusion criteria: PI-minor mutations from our definition of RAMs as these minor variants may occur as common polymorphisms in HIV-1 non-B subtypes which is predominant in our cohort Study design: cohort study
Patient characteristics and set- ting	 Country: Thailand, Hong Kong, Indonesia, Malaysia and Philippines World Bank Income classification: Thailand, Indonesia, Philippines: low-middle; Hong-Kong: high; Malaysia: upper-middle
	 Study setting: 10 sites in Thailand, Hong Kong, Indonesia, Malaysia and Philippines, clin- ic-based
	Study dates: not reported (recruitment started in 2007)
	 Age of population (years), median (IQR): 36 (32 to 41)
	Gender (male %): 66
	Participants included/analysed: 105/81
	 First or second-line regimen: second-line Type of ART: not reported



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Jiamsakul 2014 (Continued)			erence: at 12 months from switch to sec-
Index tests	Number of index tests used:	1	
	Types of index tests: self-rep	ort	
	day Visual Analogue S	adherence was recorded base cale (VAS). Adherence level v be associated with virologica ssing) on d: yes	d on the WHO-endorsed self-reported 30- was categorized based on the traditional l failure: (1) always≥95%, ever <95% and
Target condition and reference	Target condition: viral non-s	uppression	
standard(s)	• Reference standard: no d	etails provided on the assay (used
		opression: HIV viral load > 400) copies/mL
	Blinded to index test: no	nformation	
Flow and timing	months after switching toAll patients received samMissing data: Of 2023 TAS) second-line ART. e reference standard: unclea SER-M participants, 105 parti	ral load and adherence was done at 12 r cipants fit inclusion criteria and were in- only available for 81 participants. Missing
Comparative			
Notes	Asia, a program of amfAR, TH Dutch Ministry of Foreign Aff tional support from amfAR a the U.S. National Institutes of International Epidemiologic Elizabeth Hospital and the Ir cil for AIDS Trust Fund. The M	sia Studies to Evaluate Resis ne Foundation for AIDS Resea airs through a partnership wi nd the National Institute of A of Health (NIH) and the Natior Databases to Evaluate AIDS (ntegrated Treatment Centre a Kirby Institute is funded by th	tance (TASER) is an initiative of TREAT irch, with major support provided by the ith Stichting Aids Fonds, and with addi- llergy and Infectious Diseases (NIAID) of hal Cancer Institute (NCI) as part of the (IeDEA) (grant no. U01AI069907). Queen are supported by the Hong Kong Coun- e Australian Government Department of dicine, UNSW Australia (the University of
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		



Jiamsakul 2014 (Continued)			
Was a case-control design avoided?	Yes		
Did the study avoid inappropri- ate exclusions?	Unclear		
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the in- cluded patients and setting do not match the review ques- tion?			Low concern
DOMAIN 2: Index Test (Index test	t)		
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpre- tation of the index test have introduced bias?		Low risk	
Are there concerns that the in- dex test, its conduct, or inter- pretation differ from the re- view question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards like- ly to correctly classify the target condition?	Unclear		
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Unclear		
Could the reference standard, its conduct, or its interpreta- tion have introduced bias?		Unclear risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			Unclear
DOMAIN 4: Flow and Timing			
Was there an appropriate inter- val between index test and ref- erence standard?	Unclear		

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Jiamsakul 2014 (Continued)

Did all patients receive the same reference standard?	Unclear
Were all patients included in the analysis?	No
Could the patient flow have introduced bias?	High risk

Kitkungvan 2008

Study characteristics	
Patient Sampling	 Target population: adults with HIV, treatment-naïve Recruitment: analysis only included patients with suppressed VL at 6 months Inclusion criteria: HIV-drug-naive patients who were ≥ 15 years old, clinically eligible for ART (CD4 count < 200 cells/mL) and subsequently prescribed GPO-VIR. Enrolled patients had confirmed HIV infection and gave written consent to study participation Exclusion criteria: patients who did not meet the criteria for ART initiation or begar an alternative regimen to GPO-VIR Study design: prospective observational study
Patient characteristics and setting	 Country: Thailand World Bank Income classification: low-middle Study setting: hospital-based Study dates: April 2003 to 31 March 2007 Age of population (years), median (range): 37 (15 to 61) Gender (male %): 64 Participants included/analysed: 205/199 First or second-line regimen: first-line Type of ART: participants were receiving fixed-dose stavudine, 3TC and NVP Time on ART at enrolment: first-line (treatment-naïve patients) Time on ART at measurement of viral load and adherence: 18 months
Index tests	 Number of index tests used: 1 Types of index tests: tablet counts Test 1. Tablet counts Validated scale: not applicable Tool description: at each routine medical encounter, the pharmacist calculated the ratio of pills taken divided by the total number of pills prescribed for the in terval period. In addition, unannounced home visits including pill counts at participants' residences were conducted randomly by trained adherence counselling educators twice monthly. Blinding: no information Threshold prespecified: not reported Adherence threshold used: 95 to 100%; 75 to 94%; 55 to 74% and 0 to 54%
Target condition and reference standard(s)	 Target condition: viral non-suppression Reference standard: no details provided on the assay used Definition of viral non-suppression: HIV viral load > 50 copies/mL Blinded to index test: no information

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

Kitkungvan 2008 (Continued)			
Flow and timing	 Time interval between index and reference tests: no detailed information about timing, but both happened at 18 months All patients received same reference standard: unclear Missing data: 199 of the 205 patients that were included in the study were included in the analysis. Missing data for < 10% participants 		
Comparative			
Notes	Conflicts of interest: none	edeclared	
		masat University Fund to	y the Thai American Physician Foun- Infectious Disease and Hospital Epi-
	Trial registry: not reported	d	
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclu- sions?	No		
Could the selection of patients have in- troduced bias?		High risk	
Are there concerns that the included pa- tients and setting do not match the re- view question?			High
DOMAIN 2: Index Test (Index test)			
Were the index test results interpreted without knowledge of the results of the ref- erence standard?	Unclear		
If a threshold was used, was it pre-speci- fied?	Unclear		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correct- ly classify the target condition?	Yes		

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Kitkungvan 2008 (Continued)			
Were the reference standard results inter- preted without knowledge of the results of the index tests?	Unclear		
Could the reference standard, its con- duct, or its interpretation have intro- duced bias?		Low risk	
Are there concerns that the target con- dition as defined by the reference stan- dard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Unclear		
Did all patients receive the same reference standard?	Unclear		
Were all patients included in the analysis?	Yes		
Could the patient flow have introduced bias?		Unclear risk	

Labhardt 2012

Study characteristics	
Patient Sampling	 Target population; patients on first-line ART for at least 6 months, aged ≥ 10 years, with viral failure
	 Recruitment: all patients aged 16 years or older who started ART with at least three drugs (two NRTIs and one NNRTI) within the two catchment areas between Janu- ary 2008 and April 2011 were included.
	 Inclusion criteria: all patients on first-line ART since at least 6 months, aged ≥ 10 years, who fulfilled clinical and/or immunological WHO-criteria for treatment fail- ure and who were followed within the study area
	 Exclusion criteria: patients taking PI-based ART were excluded from the study. The study only included patients with viral failure.
	Study design: cross-sectional
Patient characteristics and setting	 Country: Lesotho World Bank Income classification: low-income Study setting: catchment area of Seboche Hospital in northern Lesotho Study dates: October 2010 and April 2011 Age of population (years), median (IQR): 41 (33 to 49) Gender (male %): 51 Participants included/analysed: 134/92 First or second-line regimen: first-line Type of ART: not reported Time on ART at enrolment: at least 6 months Time on ART at measurement of viral load and adherence: at least 6 months (most of them 12 months)

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

Labhardt 2012 (Continued)				
Index tests	Number of index tests used: 1			
	Types of index tests: self-report			
		nurse-clinician assessed t a visual analogue scale (V ation fied: unclear	he clinical part of the score (adher- 'AS)	
Target condition and reference standard(s)	Target condition: viral no	n-suppression		
		o details provided on the suppression: HIV viral loa no information		
Flow and timing	drawn, a nurse-clinicia sured by a VAS)All patients received saMissing data: only 92	an assessed the clinical p ame reference standard:	patients could be included in the	
Comparative				
Notes	Conflicts of interest: none	edeclared		
	Funding source: no funding received			
	Trial registry: not reporte	d		
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of pa- tients enrolled?	Yes			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate exclusions?	No			
Could the selection of patients have intro- duced bias?		High risk		
Are there concerns that the included pa- tients and setting do not match the review question?			High	
DOMAIN 2: Index Test (Index test)				



abhardt 2012 (Continued)			
Were the index test results interpreted with- out knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Unclear		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Unclear		
Were the reference standard results interpret- ed without knowledge of the results of the in- dex tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Unclear risk	
Are there concerns that the target condi- tion as defined by the reference standard does not match the question?			Unclear
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between in- dex test and reference standard?	Yes		
Did all patients receive the same reference standard?	Unclear		
Were all patients included in the analysis?	No		
Could the patient flow have introduced bias?		High risk	

Study characteristics	
Patient Sampling	 Target population: post-partum mothers, screened HIV-positive at outpatient clinics ir Malawi
	 Recruitment: this is a nested study of HIV-infected mothers presenting with their 1 to 6 month-old infants at outpatient clinics in Malawi, where they were enrolled for longitudi nal follow-up in the NEMAPP study. The subset included in this study, based on regiona strata, were enrolled for intensive clinical and laboratory monitoring at 13 health facili ties across 8 districts.
	 Inclusion criteria: 1 to 6 months post-partum mothers, screened HIV-positive at outpa tient clinics in Malawi



andes 2021 (Continued)	• Exclusion criteria: not re	eported	
	Study design: prospecti		
Patient characteristics and setting	Country: Malawi		
		classification: low-income	
			ealth facilities across 8 districts)
	 Study dates: October 20 		
	_	s), median (IQR): 29 (24 to	33)
	 Gender (male %): 0 (all 		
	 Participants included/a 	=	
	 First or second-line regi Type of ART: lifelong 	men: not reported ART (i.e. tenofovir/3TC/EF	V)
	 Time on ART at enro 	lment: not reported	
	 o Time on ART at mea ≥ 24 months 	surement of viral load and	adherence: range from 6.1 months t
Index tests	Number of index tests used	d: 1	
	Types of index tests: self-re	eport	
	• Test 1. Self-report quest		
	• Validated scale: not		
			of missed ART in the last month
	 Blinding: no informa 		
	 Threshold prespecifi 	-	
	 Adherence threshold ART) 	d used: 100% (optimal adr	erence defined as 0–1 days of misse
Target condition and reference stan-	Target condition: viral non	-suppression	
dard(s)	Reference standard: Ab	bott Real-Time HIV-1 Assay	, Abbott Laboratories, Chicago, IL
	 Definition of viral non-s 	uppression: HIV viral load	> 40 copies/mL
	• Blinded to index test: no	o information	
Flow and timing			at enrolment, 12 and 24 months
		me reference standard: yes	
	 Missing data: of 590 wo in further analysis. Miss 		plete VL data at 3 visits were include
Comparative			
Notes	Conflicts of interest: not de	eclared	
	ters for Disease Control an U2GGH000721. CDC staff w ment and approval and ma the data and final responsi	d Prevention (CDC) under t vere involved as co-investig anuscript authorship. The a ibility for submission. The f	AIDS Relief (PEPFAR) through the Cen- the terms of cooperative agreement gators, assisting in protocol develop- authors acknowledge full access to al findings and conclusions in this repor sent the official position of the fund-
	Trial registry: not reported		
Methodological quality			

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Landes 2021 (Continued) **DOMAIN 1: Patient Selection** Was a consecutive or random sample Unclear of patients enrolled? Was a case-control design avoided? Yes Did the study avoid inappropriate ex-Unclear clusions? Could the selection of patients have Unclear risk introduced bias? Are there concerns that the includ-Unclear ed patients and setting do not match the review question? **DOMAIN 2: Index Test (Index test)** Unclear Were the index test results interpreted without knowledge of the results of the reference standard? If a threshold was used, was it pre-Unclear specified? Could the conduct or interpretation Low risk of the index test have introduced bias? Are there concerns that the index Low concern test, its conduct, or interpretation differ from the review question? **DOMAIN 3: Reference Standard** Is the reference standards likely to cor-Yes rectly classify the target condition? Were the reference standard results in-Unclear terpreted without knowledge of the results of the index tests? Could the reference standard, its Low risk conduct, or its interpretation have introduced bias? Are there concerns that the target Low concern condition as defined by the reference standard does not match the question? **DOMAIN 4: Flow and Timing** Was there an appropriate interval be-Yes tween index test and reference standard?

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Landes 2021 (Continued)

Did all patients receive the same refer- Yes ence standard?

Were all patients included in the analy- No sis?

Could the patient flow have introduced bias?

Mariana 2018

Study characteristics	
Patient Sampling	 Target population: adults infected with HIV, aged > 18 years, ART-naïve on ther apy for 12 months Recruitment: no details reported Inclusion criteria: adults infected with HIV, aged > 18 years, ART-naïve on ther apy for 12 months Exclusion criteria: patients with heart, renal disease and cancer were exclud ed. Study design: cross-sectional
Patient characteristics and setting	 Country: Indonesia World Bank Income classification: lower-middle-income Study setting: hospital-based (Sulianti Saroso hospital, Jakarta) Study dates: July to October 2017 Age of population (years): not reported Gender (male %): 90 Participants included/analysed: group 1: 78/78; group 2: 20/20 First or second-line regimen: first-line Type of ART: Group 1 - Fixed dose combination group: first-line treatment with tenovofir and EFV once daily; Group 2 - Free combination group: free combination of ARV Time on ART at enrolment: 12 months Time on ART at measurement of viral load and adherence: 12 months
Index tests	 Number of index tests used: 1 Types of index tests: tablet counts Test 1. Tablet counts Validated scale: not applicable Tool description: poor adherence was defined as a value of pill consumption < 95% pills. No other details reported Blinding: no information Threshold prespecified: not reported Adherence threshold used: 95%
Target condition and reference standard(s)	 Target condition: viral non-suppression Reference standard: no details provided on the assay used Definition of viral non-suppression: HIV viral load > 40 copies/mL Blinded to index test: no information

High risk

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

ariana 2018 (Continued)			
low and timing	timing, but this is a crouslyAll patients received sMissing data: none. A	ross-sectional study so l same reference standard	ith viral load test and adherence
Comparative			
lotes	Conflicts of interest: non	e declared	
	Funding source: no fund	ing received	
	NCT record number: not	reported	
1ethodological quality			
tem	Authors' judgement	Risk of bias	Applicability concerns
OMAIN 1: Patient Selection			
Vas a consecutive or random sample of patients nrolled?	Unclear		
Vas a case-control design avoided?	Yes		
id the study avoid inappropriate exclusions?	Yes		
Could the selection of patients have introduced vias?		Low risk	
are there concerns that the included patients and setting do not match the review question?			Low concern
OOMAIN 2: Index Test (Index test)			
Vere the index test results interpreted without nowledge of the results of the reference stan- lard?	Unclear		
f a threshold was used, was it pre-specified?	Unclear		
Could the conduct or interpretation of the in- lex test have introduced bias?		High risk	
are there concerns that the index test, its con- luct, or interpretation differ from the review juestion?			Unclear
OMAIN 3: Reference Standard			
s the reference standards likely to correctly clas- ify the target condition?	Unclear		
Vere the reference standard results interpret- d without knowledge of the results of the index ests?	Unclear		

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Mariana 2018 (Continued)

Trusted evidence. Informed decisions. Better health.

Could the reference standard, its conduct, or its interpretation have introduced bias?		Unclear risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			Unclear
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference stan- dard?	Unclear		
Were all patients included in the analysis?	Yes		
Could the patient flow have introduced bias?		Unclear risk	

Mbengue 2019

Study characteristics	
Patient Sampling	 Target population: HIV-positive adult (≥ 18 years of age) patients who initiated standard first-line ART Recruitment: secondary analysis of data collected from a prospective observational study Inclusion criteria: HIV infection, > 18 years, current PI-based second-line ART treatment since at least 6 months and willing to participate and consent signature Exclusion criteria: patients who transferred in on ART. While pregnant women were eligible to initiate ART, they were not included in the prospective study mainly because they were initiated using different criteria and were managed differently (e.g. transferred out to other facilities for antenatal care). Study design: prospective cohort study
Patient characteristics and setting	 Country: South Africa World Bank Income classification: upper-middle-income Study setting: Themba Lethu Clinic (TLC) in Johannesburg, South Africa (clinic visits for study visits) Study dates: treatment initiated between February 2012 and April 2016 Age of population (years): not reported Gender (male %): 33.8 Participants included/analysed: 357/163 First or second-line regimen: first-line Type of ART: standard first-line therapy included tenofovir with 3TC and efavirenz, and in April 2013 TLC introduced a single pill or fixed-dose combination which replaced the multi-pill ART regimen Time on ART at enrollment: 6 months Time on ART at measurement of viral load and adherence: not reported; at least 6 months
Index tests	 Number of index tests used: 3 Types of index tests: self-report, composite measure Test 1. Self-report VAS Validated scale: yes Tool description: VAS



6 months. Missing data > 10% Comparative Notes Conflicts of interest: DE reported grants from Health Economics and Epidemiology Research Office during the conduct of the study. The authors reported no other conflicts of interest in this work. Funding source: the American People and the President's Emergency Plan for AIDS Relief (PEPFAR) through US Agency for International Development (USAID) under the terms of Cooperative Agreements AID-674-A-12-00029 and 72067419CA00004 to HE2R0. The contents are the responsibility of the authors and do not necessarily reflect the views of PEPFAR, USAID or the United States Government. DE was supported by funding from NIH/CFAR/IAS Creative and Novel Ideas in HIV Research (CNIHR) program (sub-award with UAB Center for AIDS Research: P30AI027767) and National Research Foundation (not reported) of South Africa Thuthuka program (post-PhD track 500 TTK1206261680 Grant number 84331). Methodological quality Item Authors' judgement Risk of bias Applicability concerns	6 months. Missing data > 10% Comparative Notes Conflicts of interest: DE reported grants from Health Economics and Epidemiology Research Office during the conduct of the study. The authors reported no other conflicts of interest in this work. Funding source: the American People and the President's Emergency Plan for AIDS Relief (PEPFAR) through US Agency for International Development (USAID) under the terms of Cooperative Agreements AID-674-A-12-00029 and 72067149CA00004 to HE2RO. The contents are the responsibility of the authors and do not necessarily reflect the views of PEPFAR, USAID or the United States Government. DE was supported by funding from NIH/CFAR/IAS Creative and Novel Ideas in HIV Research (CNIHR) program (sub-award with UAB Center for AIDS Research: P30AI027767) and National Research Foundation (not reported) of South Africa Thuthuka program (post-PhD track 500 TTK1206261680 Grant number 84331). Trial registry: not reported	• /	All patients received sam	e reference standard: unclear	
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		ltem Aut	thors' judgement	Risk of bias	Applicability concerns
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DOMAIN 1. Detient Celection	DOMAIN 1. Detient Calentian				
DOMAIN 1: Patient Selection	DOMAIN 1: Patient Selection	DOMAIN 1: Patient Selection			

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Mbengue 2019 (Continued)			
Was a consecutive or ran- dom sample of patients en- rolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inap- propriate exclusions?	Yes		
Could the selection of pa- tients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Index	test)		
Were the index test results interpreted without knowl- edge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Unclear		
Could the conduct or in- terpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standa	ard		
Is the reference standards likely to correctly classify the target condition?	Unclear		
Were the reference stan- dard results interpreted without knowledge of the results of the index tests?	Unclear		
Could the reference stan- dard, its conduct, or its in- terpretation have intro- duced bias?		Unclear risk	
Are there concerns that the target condition as defined by the reference			Unclear



Mbengue 2019 (Continued) standard does not match the question?

DOMAIN 4: Flow and Timing	
Was there an appropriate interval between index test and reference standard?	Unclear
Did all patients receive the same reference standard?	Unclear
Were all patients included in the analysis?	No
Could the patient flow have introduced bias?	High risk

McMahon 2013

Study characteristics	
Patient Sampling	 Target population: adults initiating first-line ART Recruitment: consecutive initiators of ART were identified during a routine clinic visit after 11-15 months of ART. Inclusion criteria: adults initiating first-line ART who had not attended or picked up ART within 90 days of their last missed appointment. All baseline clinical and demographic data were abstracted from clinical records and ART dispensing data from pharmacy records. Exclusion criteria: patients transferred in from other sites or re-initiating ART after a treatment interruption Study design: prospective cohort study (retrospective data collection)
Patient characteristics and setting	 Country: India World Bank Income classification: low-income Study setting: clinic-based Study dates: Recruitment from October 2009 until October 2015 Age of population (years), mean (SD): 38.3 (8.7) Gender (male %): 65 Participants included/analysed: 230/170 First- or second-line regimen: first-line Type of ART: not specified. ART was provided for free in this study. Time on ART at enrolment: treatment-naïve Time on ART at measurement of viral load and adherence: 12 months
Index tests	 Number of index tests used: 5 Types of index tests: self-report, pharmacy records, composite measure Test 1. Self-report questionnaire Validated scale: yes Tool description: 30 day Self-report (5-point Likert item): "Standardized self-report adherence measures asked about adherence since; initiating ART, or the preceding 30-days". "Adherence questions were originally written in English, translated into Tamil or Telugu and independently backtranslated. Questionnaires were administered in local languages by trained staff experienced in HIV counselling and treatment." Threshold used was binary: '< Excellent' and 'Excellent'

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



McMahon 2013 (Continued)

- Blinding: no information
- Threshold prespecified: not reported
- Adherence threshold used: 100%
- Test 2. Self-report questionnaire
- Validated scale: not reported
- Tool description: self-report "Last time missed": "Standardized self-report adherence measures asked about adherence since; initiating ART, or the preceding 30-days". "Adherence questions were originally written in English, translated into Tamil or Telugu and independently backtranslated. Questionnaires were administered in local languages by trained staff experienced in HIV counselling and treatment." Threshold used was binary: '> never' and 'never'
- Blinding: no information
- Threshold prespecified: not reported
- Adherence threshold used: 100%
- Test 3. Self-report VAS
 - Validated scale: yes
 - Tool description: 30-day VAS: "An additional 30-day self-report measure was the visual analog scale (VAS) where patients indicated on a line marked from 0% to 100% the point that best corresponded to the percentage of pills taken".
 - Blinding: no information
 - Threshold prespecified: not reported
 - Adherence threshold used: 95%
- Test 4. Pharmacy records
 - Validated scale: not applicable
 - Tool description: medication possession ratio (MPR). This was calculated by dividing the days of ART dispensed by the period of time from ART start to the day of recruitment". "Patients attended monthly for medical review and picked-up ART from a pharmacy staffed by a dedicated pharmacist within the clinic".
 - Blinding: no information
 - Threshold prespecified: not reported
 - Adherence threshold used: 100 %
- Test 5. Composite measure
 - Validated scale: not applicable
 - Tool description: combined self-report and MPR: 12-month medication possession ratio (MPR) (days of ART dispensed divided by the period of time from ART start to the day of recruitment) < 100% + suboptimal adherence on either of 2 self-report measures (< excellent adherence in last 30 days, or ever reported missing ART). Threshold used was binary: 'Low adherence' and 'High adherence'.
 - Blinding: no information
 - Threshold prespecified: yes
 - Adherence threshold used: 'high versus low'

Target condition and reference standard(s)	 Target condition: viral non-suppression Reference standard: Artus HIV-1 RT-PCR (Qiagen) Definition of viral non-suppression: HIV viral load > 200 copies/mL Blinded to index test: no information
Flow and timing	 Time interval between index and reference tests: baseline characteristics and dichotomous adherence estimates after 12-months ART were compared to 12-month viral load. All patients received same reference standard: yes Missing data: 230 were included in the study, after 12 months, 177 were on ART and 174 undertook a viral load. Also, for some of the index tests 170/174 were included; unclear why additional 4 were excluded. Missing data > 10%
Comparative	

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

McMahon 2013 (Continued)

Notes	Conflicts of interest: SRL receives payment for lectures (Viiv Healthcare and Janssen), payment for educa- tional presentations (Janssen) and SRL's institution receives grant funding (Merck and Gilead). All other authors, no conflicts.
	Funding source: JM was supported by a fellowship from Tufts Medical Center Department of Geographic Medicine and Infectious Diseases, and an Australian National Health and Medical Research Council (NHM- RC) Postgraduate Scholarship. The study was supported by a Lifespan/Tufts/Brown Center for AIDS Re- search NIH grant (1P30A142853-12). AM was supported by a Fogarty International Center training grant (5D43TW000237-15). MRJ was supported by an NIH Career Development Award (5K23AI074423-04). SRL is an NHMRC Practitioner Fellow.
	Trial registry: not reported

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selec	tion		
Was a consecutive or random sample of pa- tients enrolled?	Yes		
Was a case-control de- sign avoided?	Yes		
Did the study avoid in- appropriate exclusions?	No		
Could the selection of patients have intro- duced bias?		High risk	
Are there concerns that the included pa- tients and setting do not match the review question?			Unclear
DOMAIN 2: Index Test (In	idex test)		
Were the index test re- sults interpreted with- out knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Unclear		
Could the conduct or interpretation of the index test have intro- duced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpre-			Low concern



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IcMahon 2013 (Continued) tation differ from the review question?			
DOMAIN 3: Reference Sta	andard		
Is the reference stan- dards likely to correctly classify the target con- dition?	Yes		
Were the reference standard results inter- preted without knowl- edge of the results of the index tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the target condi- tion as defined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Tim	ning		
Was there an appropri- ate interval between in- dex test and reference standard?	Unclear		
Did all patients receive the same reference standard?	Yes		
Were all patients in- cluded in the analysis?	No		
Could the patient flow have introduced bias?		High risk	

Messou 2011

Study characteristics	
Patient Sampling	Target population: HIV-infected adults who started ART
	 Recruitment: HIV-infected adults who started ART between February 2006 and May 2007 at one of three HIV outpatient clinics in Abidjan and showed up for their six-month visi were eligible for the study.
	 Inclusion criteria: HIV-infected adults who started ART between February 2006 and May 2007 at one of three HIV outpatient clinics in Abidjan and showed up for their six month visit were eligible for the study.



essou 2011 (Continued)	
	Exclusion criteria: not reportedStudy design: prospective cohort study
Patient characteristics and setting	 Country: Cote d'Ivoire World Bank Income classification: low-income Study setting: clinic-based Study dates: February 2006 to May 2007 Age of population (years), median (IQR): 36 (30 to 43) Gender (male %): 75 Participants included/analysed: 925/1206 at 12 months First- or second-line regimen: first-line Type of ART: stavudine/AZT + 3TC + NVP/EFV Time on ART at enrolment: treatment-naïve Time on ART at measurement of viral load and adherence: 6 and 12 months
Index tests	Number of index tests used: 1
	Types of index tests: pharmacy records
	 Test 1. Pharmacy records Validated scale: not applicable
	 Tool description: the medication possession ratio (MPR) was defined as the number of daily doses of antiretroviral drugs dispensed by the pharmacy to each patient, divided by that patient's total follow-up time in days since ART initiation. Blinding: no information Threshold prespecified: not reported Adherence threshold used: > 95%; > 80%; > 65%; > 50%
Target condition and reference stan- dard(s)	Target condition: viral non-suppression
	 Reference standard: ANRS real-time PCR; Biocentric, Bandol, France Definition of viral non-suppression: HIV viral load > 300 copies/mL Blinded to index test: no information
Flow and timing	 Time interval between index and reference tests: association between MPR from baseline to month 12 and virologic failure was estimated. All patients received same reference standard: yes Missing data: at 6 months, 996/1206 patients included in the analysis. At 12 months, 925/1206 were included in the analysis. Patients were defined as lost to follow-up if: (i) their last contact with study team was less than month 12; (ii) they were not known to be dead or transferred out before month 12; (iii) no further information on their vital status could be obtained within the 6 months following study endpoint (i.e. between month 12 and month 18). Missing data > 10%
Comparative	
Notes	Conflicts of interest: none declared
	Funding source: Agence Nationale de Recherches sur le SIDA et les hepatitis virales (ANot reportedS 12136, ANot reportedS 12212), National Institute of Allergy and Infectious Dis- eases (R01 AI058736, K24 AI062476) and Institute of International Education Fulbright (Fel- lowship CS)
	Trial registry: Not reported

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Messou 2011 (Continued) Item **Authors' judgement Risk of bias Applicability concerns DOMAIN 1: Patient Selection** Was a consecutive or random sample No of patients enrolled? Was a case-control design avoided? Yes Unclear Did the study avoid inappropriate exclusions? Could the selection of patients have High risk introduced bias? Are there concerns that the includ-Unclear ed patients and setting do not match the review question? **DOMAIN 2: Index Test (Index test)** Were the index test results interpret-Unclear ed without knowledge of the results of the reference standard? If a threshold was used, was it pre-Unclear specified? Unclear risk Could the conduct or interpretation of the index test have introduced bias? Are there concerns that the index Low concern test, its conduct, or interpretation differ from the review question? **DOMAIN 3: Reference Standard** Is the reference standards likely to cor-Yes rectly classify the target condition? Were the reference standard results in-Unclear terpreted without knowledge of the results of the index tests? Could the reference standard, its Low risk conduct, or its interpretation have introduced bias? Are there concerns that the target Low concern condition as defined by the reference standard does not match the question? **DOMAIN 4: Flow and Timing**



Messou 2011 (Continued)

duced bias?	9
Could the patient flow have intro-	High risk
Were all patients included in the analy- sis?	Νο
Did all patients receive the same refer- ence standard?	Yes
Was there an appropriate interval be- tween index test and reference stan- dard?	Unclear

Meya 2009

Study characteristics	
Patient Sampling	 Target population: HIV-1-positive, aged > 18 years, established on first line NNRTI-based ART for ≥ six months Recruitment: 500 patients were enrolled at a rate of approximately 10 patients per clinic day. Patients were randomly selected from the clinic reception using a list of random numbers. Inclusion criteria: patients were screened and included in the study if they were HIV-1-positive, aged > 18 years, established on first-line NNRTI-based ART for ≥ six months and did not have viral loads monitored as per routine clinic practice. Exclusion criteria: patients with acute illness were excluded from the study. Study design: cross-sectional
Patient characteristics and setting	 Country: Uganda World Bank Income classification: low-income Study setting: clinic-based - adult clinic of the Infectious Disease Institute (IDI), Mulago Hospital, Makerere University in Kampala, Uganda Study dates: not reported Age of population (years), median: 38.4 Gender (male %): 37 Participants included/analysed: 496/496 First- or second-line regimen: first-line Type of ART: first-line NTI-based ART Time on ART at enrolment: at least 6 months Time on ART at measurement of viral load and adherence: not reported; at least 6 months
Index tests	 Number of index tests used: 2 Types of index tests: self-report Test 1. Self-report questionnaire Validated scale: yes Tool description: adherence was measured by self-report, using a modified Adult AIDS Clinical Trials Group adherence questionnaire validated in the setting. Participants were asked to report adherence patterns in the three days prior to enrolment, four weeks prior to enrolment, and since the initiation of ART. A VAS, as well as a question on whether treatment had ever been interrupted for more than two days, was included to assess adherence in the four weeks prior to enrolment and since the initiation of ART; adherence measure use was the question: "Have you missed ART in the last 30 days"; responses were Yes, No. Blinding: no information Threshold prespecified: not reported

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



leya 2009 (Continued)					
	 Adherence threshold Tect 1 Solf report VAS 	used: 100%			
	 Test 1. Self-report VAS Validated scale: yes 				
	 Tool description: VAS ed for more than tw enrolment and since 	o days, was included to asse the initiation of ART; respon	ether treatment had ever been interrupt- ss adherence in the four weeks prior to ses were Yes, No.		
	 Blinding: no informa 				
	Threshold prespecifiAdherence threshold	-			
Target condition and reference	Target condition: viral non	-suppression			
standard(s)	• Reference standard: Am	plicor HIV-1 Monitor v1.5 – R	oche, Switzerland		
	 Definition of viral non-suppression: HIV viral load > 400 copies/mL 				
	Blinded to index test: no	information			
	•				
Flow and timing		ndex and reference tests: no e so likely to be measured simi	explicit information on timing, but this is Iltaneously		
	All patients received sar	=			
	 Missing data: none. All e included in the main an 		load test and adherence measures were		
Comparative					
Notes	Conflicts of interest: none of	declared			
	fectious Diseases, part of th		he National Institute of Allergy and In- th, USA. We also acknowledge support)		
	Trial registry: not reported				
Methodological quality					
Item	Authors' judgement	Risk of bias	Applicability concerns		
DOMAIN 1: Patient Selection					
Was a consecutive or random sam- ple of patients enrolled?	Yes				
Was a case-control design avoid- ed?	Yes				
Did the study avoid inappropriate exclusions?	Yes				
Could the selection of patients have introduced bias?		Low risk			
Are there concerns that the in- cluded patients and setting do			Low concern		
not match the review question?					

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

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eya 2009 (Continued)			
Were the index test results inter- preted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre- specified?	Yes		
Could the conduct or interpreta- tion of the index test have intro- duced bias?		Low risk	
Are there concerns that the in- dex test, its conduct, or inter- pretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condi- tion?	Yes		
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the tar- get condition as defined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Yes		
Could the patient flow have in- troduced bias?		Low risk	

Study characteristics • Target population: HIV-positive patients, initiating ART Patient Sampling Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

Library

Mogosetsi 2018 (Continued)

	 Recruitment: prospective cohort study among patients down referred primarily from the Phedisong 4 clinic, which offers various primary health care services to patients, including ART Inclusion criteria: age ≥ 18 years, patients initiating ART, patients who were ART-naïve and patients who had undergone the mandatory three adherence counselling classes recommended by the South African Department of Health Exclusion criteria: not reported Study design: prospective cohort study
Patient characteristics and setting	 Country: South Africa World Bank Income classification: upper-middle-income Study setting: clinic-based Study dates: November 2012 to March 2013 Age of population (years), mean (SD): viral suppression group: 37 (9.3); non-suppression group: 35 (8.0) Gender (male %): viral suppression group: 38; non-suppression group: 12 Participants included/analysed: 155/98 First- or second-line regimen: first-line Type of ART: PI-based Time on ART at enrolment: treatment-naïve Time on ART at measurement of viral load and adherence: 6 months
Index tests	Number of index tests used: 1 Types of index tests: self-report
	 Test 1. Self-report questionnaire Validated scale: yes (MMAS-4) Tool description: the adherence of each patient was assessed by means of the Morisky Medication Adherence Scale (MMAS-4). This scale is a generic self-reporting measurement of patient behaviour in taking medication. It consists of four questions with a scoring of 0 for "Yes" and 1 for "No", with a total range of 0-4 points. A score of 0 indicates high adherence, 1-2 medium adherence and 3-4, low adherence. Blinding: no information Threshold prespecified: yes Adherence threshold used: 100%; 95%
Target condition and reference standard(s)	 Target condition: viral non-suppression Reference standard: m2000 Real Time HIV-1 Viral Load System[®] supplied by Abbott Definition of viral non-suppression: HIV viral load > 40 copies/mL
	 Blinded to index test: no information
Flow and timing	 Time interval between index and reference tests: data were collected at 6 months. All patients received same reference standard: yes Missing data: 98/155 participants included in the analysis at 6 months. Missing were defaulters, deceased, transferred to another health facility, and discontinued participation. Missing data > 10%
Comparative	
Notes	Conflicts of interest: none declared Funding source: no funding received Trial registry: not reported

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

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Mogosetsi 2018 (Continued)

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclu- sions?	No		
Could the selection of patients have in- troduced bias?		High risk	
Are there concerns that the included pa- tients and setting do not match the re- view question?			Unclear
DOMAIN 2: Index Test (Index test)			
Were the index test results interpreted without knowledge of the results of the ref- erence standard?	Unclear		
If a threshold was used, was it pre-speci- fied?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correct- ly classify the target condition?	Yes		
Were the reference standard results inter- preted without knowledge of the results of the index tests?	Yes		
Could the reference standard, its con- duct, or its interpretation have intro- duced bias?		Low risk	
Are there concerns that the target con- dition as defined by the reference stan- dard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			

Could the patient flow have introduced bias?	High risk
Were all patients included in the analysis?	No
Did all patients receive the same reference standard?	Yes
Was there an appropriate interval between index test and reference standard?	Unclear
Mogosetsi 2018 (Continued)	

Moosa 2019

Study characteristics	
Patient Sampling	 Target population: HIV-tuberculosis (TB) co-infected patients receiving tuberculosis treatment Recruitment: secondary analysis of two studies. The SAPiT study is a three-arm, open-label, randomized controlled trial conducted at the CAPRISA Thekwini HIV-tuberculosis clinic with HIV-tuberculosis co-infect ed patients receiving tuberculosis treatment. After completion of follow-up in the SAPiT trial, patients were offered enrolment into a prospective observational study, TB Recurrence upon Treatment with HAART (TRuTH), investigating the rate of TB recurrence in HIV-infected adults on ART who had completed pul monary TB treatment. Inclusion criteria: SAPiT trial inclusion criteria: To be included, patients had to be independently confirmed at the Department of Medical Microbioloy, Nelson R. Mandela School of Medicine, to have AFB smear positive disease, initiated on the standard tuberculosis treatment regimen at the PCZCDC, have a CD4+ count 4500 cells/mm³ at screening and have no clinical contraindications to initiation of ART. Female participant: were required to agree to use contraception while on efavirenz. Exclusion criteria: no information on exclusion criteria for the SAPiT and TRuTH studies. In the current pape (secondary analysis), patients who never initiated ART, were lost to follow-up in the SAPiT trial, did no receive ART from the site's research pharmacy, and for whom pill count data was missing for more than 6 consecutive months in either study were excluded. Study design: three-arm, open-label, randomized controlled trial (the SAPiT study); prospective observational turdy:
Patient characteris- tics and setting	 tional study, (TRuTH study) Country: South Africa World Bank Income classification: upper-middle-income Study setting: the SAPiT trial (protocol number: CAPRISA 003) was conducted at the CAPRISA eThekwin HIV-tuberculosis clinic. This adjoins one of the largest ambulatory (outpatient) tuberculosis facilities in South Africa, the Prince Cyril Zulu Communicable Disease Centre (PCZCDC) in Durban. The TRuTH study was based in Durban, KwaZulu-Natal, South Africa, an area where it was estimated that 70% of TB patients were HIV co-infected. Study dates: SAPiT trial: June 2005 to July 2008; TRuTH observational study from November 2009 to July 2011 and follow-up was completed in 2014 Age of population (years), median (IQR): 34 (29 to 40) Gender (male %): 45.2 Participants included/analysed: 270/268 (year 1), 201 (year 2) ,166 (year 3), 243 (year 4), 233 (year 5) First or second-line regimen: first-line Type of ART: study patients were initiated on a once daily, weight-based ART regimen containing EFV or NVP plus 3TC and enteric coated didanosine (ddl) either during or after completion of tuberculosis treatment. Time on ART at enrolment: at least 6 months
	 Time on ART at measurement of viral load and adherence: different categories: 1 year, 2 years, 3 years 4 years, 5 years
Index tests	Number of index tests used: 1



loosa 2019 (Continued)	Types of index tests: tablet co	ints	
	Test 1. Tablet counts		
	 Validated scale: not app 	licable	
	 counts were conducted or 3-monthly study visit formula: (number of pill at current visit)/number days between visits) x 10 the study visits. Pill cou (time period between ex- not assessed for visits w data were missing. Blinding: no information Threshold prespecified: 	by the study pharmacist at mont s in the TRuTH study. Adherence s dispensed at previous visit - nu r of pills that should have been 00. Optimal adherence was defin nt data were not available for A kit from SAPiT study and enrolme where there was a clinician-initia n not reported	determined from pharmacy pill count data. Pill hly study visits in the SAPiT trial and at monthly percentage was calculated using the following mber of pills returned/reported remaining/lost ingested between visits (daily pill dose x no of ed as ≥ 95% of doses taken in the time between RT that was dispensed in the CAT programme ent into TRuTH study). Pill count adherence was ted treatment interruption or where pill count
	 Adherence threshold us 	ed: 95%	
Target condition	Target condition: viral non-su	ppression	
and reference stan- dard(s)	 Reference standard: SAPiT Ampliprep-Roche TaqMan 	trial: Cobas® Amplicor HIV-1 Mo	nitor, version 1.5, Roche; TRUth study: Cobas®
		pression: HIV viral load > 400 cop	ies/mL
	Blinded to index test: no in	formation	
Flow and timing	 monthly study visits in the load done at screening, rar All patients received same Missing data: 414 complete were included for the retro 	SAPiT trial and at monthly or 3 adomization and 6 monthly there reference standard: yes ad SAPiT follow-up; 379 previousl	y enrolled in SAPiT enrolled in TRuTH. Only 270 for n = 268 at year 1, n = 201 at year 2, n = 166 at
Comparative			
Notes	Conflicts of interest: none rep	orted	
	oratory and pharmacy cores w search on AIDS grant (CIPRA, g funded the care of all the SAPi cost for drugs used in the SAPi stitute, Chevy Chase, MD, USA Atlanta, GA, USA) Cooperative KwaZulu-Natal Department of ington DC, USA). KN and TG w tional Training and Research F	vere established through the Cor grant # AI51794). The US Presider T patients; the Global Fund to fig iT trial. The TRUTH study was su (grant # 55007065) and the Cent Agreement Number UY2G/ PS00 f Health and the US President's E ere supported by the Columbia U	SAPiT trial, including data management, lab- nprehensive International Program of Re- nt's Emergency Plan for AIDS Relief (PEPfAR) ght AIDS, Tuberculosis and Malaria funded the oported by the Howard Hughes Medical In- ers for Disease Control and Prevention (CDC; 1350–02. Patient care was supported by the mergency Plan for AIDS Relief (PEPFAR; Wash- Jniversity-South Africa Fogarty AIDS Interna- ogarty International Center, National Insti- d for this retrospective study.
	Trial registry: not reported		
Methodological qualit	ty		
	Authors' judgement	Risk of bias	Applicability concerns
Item	Authors judgement	NISK OF DIUS	Applicability concerns



Moosa 2019 (Continued)			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclu- sions?	Unclear		
Could the selection of patients have in- troduced bias?		Unclear risk	
Are there concerns that the included patients and setting do not match the re- view question?			High
DOMAIN 2: Index Test	(Index test)		
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre- specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or in- terpretation dif- fer from the review question?			Low concern
DOMAIN 3: Reference	Standard		
Is the reference stan- dards likely to cor- rectly classify the tar- get condition?	Yes		
Were the reference standard results in- terpreted without knowledge of the results of the index tests?	Unclear		

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Moosa 2019 (Continued)

Could the reference standard, its conduct, or its interpretation have introduced bias? Are there concerns Low concern that the target condition as defined by the reference standard does not match the question? **DOMAIN 4: Flow and Timing** Was there an appro-Unclear priate interval between index test and reference standard? Did all patients re-No ceive the same reference standard? Were all patients in-No cluded in the analysis? High risk **Could the patient** flow have introduced bias?

Mutwa 2014

Study characteristics	
Patient Sampling	 Target population: HIV-infected cART-naïve children below 15 years of age Recruitment: children were usually referred from Kigali University Teaching Hospital (which is adjacent to the TRACplus clinic), nearby district hospitals, or health centres providing 'prevention of mother to child HIV transmission services'; a few children were diagnosed at the TRACplus facility itself. All children below the age of 15 years who initiated cART at the TRACplus clinic during the study period were given the opportunity to enrol in the study. Inclusion criteria: HIV-infected cART-naïve children below 15 years of age who initiated cART Exclusion criteria: not reported Study design: prospective cohort study
Patient characteristics and setting	 Country: Rwanda World Bank Income classification: low-income Study setting: clinic- and hospital-based (TRACplus clinic, usually reffered from Kigali University Teaching Hospital which is adjacent to the TRACplus clinic), nearby district hospitals, or health centres) Study dates: March 2008 to December 2009 Age of population (years), median (IQR): 7.4 (3.2 to 11.5) Gender (male %): 43

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Low risk

DOMAIN 1: Patient Selection			
Item	Authors' judgement	Risk of bias	Applicability concerns
Methodological quality			
	Trial registry: not reported		
	Funding source: Infectious D	isease Network for Treatment a	and Research in Africa
Notes	Conflicts of interest: none de	eclared	
Comparative			
	 All patients received sam Missing data: 104/123 we pants 	-	alysis. Missing data for > 10% of partici
Flow and timing	3, 6, 12 and adherence m	easured at evey follow-up visit	d measured at enrolment and at months
	 Blinded to index test: no 		pics/illL
chee standard(s)	France	he Cobas AmpliPrcp/Cobas Ta opression: HIV viral load > 40 co	qMan HIV-I, Roche Molecular Systems
Target condition and refer- ence standard(s)	Target condition: viral non-s	uppression	
	• Adherence threshold	ised: 95%	
	Blinding: no informatiThreshold prespecifie		
	ing a structured quest the child had missed reasons for non-adhe of the medication pre counted pill dispensed	onnaire. They were asked how n during the previous 30 days and rence. Children were classified scribed in the last 30 days. In ad l and returned, unused, assuming	uestions by face-to-face interviewing us- nany doses of the prescribed medicatior at what time points this occurred, and as non-adherent if having taken < 95% dition, study nurses and pharmacy staff ng all the other pills were used.
	 Tool description: self- 	eport and pill count, and assess	ment was conducted at every clinic and
	 Test 1.Composite measure Validated scale: not approximately 		
	Types of index tests: compo	site measure	
Index tests	Number of index tests used:	1	
	 Time on ART at enroln Time on ART at measu 	ient: treatment-naïve rement of viral load and adhere	ence: 6 months
	WHO ART guidelines) adolescents less than ITT or TV, or had a seve rimoxazole prophylax (not reported TTs) and efavirenz-based or PI- been exposed to nevir two TIs and a Pl.	en: first-line nitiation in 2007, the 2007 Rwar were operational, which recom 15 years of age if they were cla ere immunodeficiency; children s; they were initiated on a first-li a nonnucleoside reverse TI. A c pased. From 2009 (revised Rwan apine in the context of PMTCT w	ndan ART guidelines (based on the 2006 mended cART initiation in children and assified as WHO paediatric clinical stage enrolled in the study received cART, cot- ine cART regimen consisting of two NRT ART regimen was defined as NVP-based dian Guidelines) children known to have vere initiated on a first-line regimen with
Autwa 2014 (Continued)			

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

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Mutwa 2014 (Continued)			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inappro- priate exclusions?	Yes		
Could the selection of pa- tients have introduced bias?		Low risk	
Are there concerns that the included patients and set- ting do not match the review question?			Unclear
DOMAIN 2: Index Test (Index te	est)		
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Unclear		
Could the conduct or inter- pretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or in- terpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard	1		
Is the reference standards like- ly to correctly classify the tar- get condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Could the reference stan- dard, its conduct, or its inter- pretation have introduced bias?		Low risk	
Are there concerns that the target condition as defined by the reference standard does not match the ques- tion?			Low concern

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DOMAIN 4: Flow and Timing	
Was there an appropriate in- terval between index test and reference standard?	Unclear
Did all patients receive the same reference standard?	Unclear
Were all patients included in the analysis?	No
Could the patient flow have introduced bias?	High risk

Navarro 2014

Study characteristics	
Patient Sampling	 Target population: adult HIV-1-infected patients who were treatment-experienced and had poor adherence to HAART
	 Recruitment: all consecutive adult HIV-1-infected patients attending the HIV unit of University Hos- pital Vall d'Hebron in Barcelona
	 Inclusion criteria: All consecutive adult HIV-1-infected patients who were treatment-experienced and had poor adherence to HAART were included in an adherence programme since its introduction in 2009. Treatment-experienced patients were defined as those who had received one or more previous HAART regimens. Patients could be receiving HAART or not at the time of entering the programme, and all of them had a detectable viral load. Patients were considered poor adherents if they had more than two consecutive detectable viral loads and admitted to having missed some doses between visits. Patients were included in the adherence program for 48 weeks. Exclusion criteria: not reported Study design: prospective cohort study
Patient characteristics and	- Country Spain
setting	 Country: Spain World Bank Income classification: high-income
	 Study setting: hospital-based
	Study dates: not reported
	 Age of population (years), median (IQR): 42.7 (36.9 to 47.2)
	• Gender (male %): 51.5
	Participants included/analysed: 136/93
	 First or second-line regimen: unclear Type of ART: present HAART regimen (treatment prescribed at the entry in the adherence programme) mainly based on a ritonavir-boosted PI regimen (n = 99, 72.8%); NNRTI and other drugs such as INSTI-integrase strand transfer inhibitors
	 Time on ART at enrolment: at least 6 months
	 Time on ART at measurement of viral load and adherence: 48 weeks
Index tests	Number of index tests used: 2
	Types of index tests: self-report
	Test 1. Self-report questionnaire
	• Validated scale: yes
	 Tool description: Simplified Medication Adherence Questionnaire (SMAQ), a validated question- naire that assesses not only the adherence to HAART or not (yes or no), but also the percentage



Vavarro 2014 (Continued)			
	 questions: (1) "Do yo taking your medicine (4) "Taking into accord "Did you not take any how many days have asses the categorical treatment over time (Blinding: no informat Threshold prespecifie Adherence threshold Test 2. Pharmacy records Validated scale: yes Tool description: base 	u ever forget to take your med s?", (3) "If sometimes you feel unt only the last week, how oft y of your medicines over the pa you not taken any medicine a variable adherence (yes or no) weeks 4, 12, 24, 36, 48, 60, 72, 8 ion ed: yes used: > 90%; > 60% s or secondary database analys ed on a drug dispensing record er of months of follow-up, assu- cion ed: yes	
Target condition and refer- ence standard(s)	 Target condition: viral non- Reference standard: not Definition of viral non-su Blinded to index test: no 	reported Ippression: HIV viral load > 50 c	opies/mL
Flow and timing	 adherence and viral load All patients received san Missing data: participan 48 of follow-up at the m were lost to follow-up wi 	I were assessed at 48 weeks ne reference standard: not reports ts who had an undetectable vi oment of the analysis were no th a viral load > 50 copies/mL, a	xplicit information on timing, just that both orted iral load but who had not yet reached week ot included in the analysis. 6 patients (4.4%) nd 7 patients had not yet reached week 48 of d missing data as well, reasons not reported.
Comparative			
Notes	Conflicts of interest: no con	npeting financial interests exist	
	Investigation Cientifica, Des		roject as part of the Plan Nacional I +D+ i, nanced by ISCIII-Subdirección General de DER)
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection	1		
Was a consecutive or ran- dom sample of patients en- rolled?	Yes		
Was a case-control design avoided?	Yes		

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Navarro 2014 (Continued)

Did the study avoid inap- propriate exclusions?	Yes		
Could the selection of pa- tients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Index	test)		
Were the index test results interpreted without knowl- edge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or in- terpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standa	ard		
Is the reference standards likely to correctly classify the target condition?	Unclear		
Were the reference stan- dard results interpreted without knowledge of the results of the index tests?	Unclear		
Could the reference stan- dard, its conduct, or its in- terpretation have intro- duced bias?		Unclear risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			Unclear
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Unclear		

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Did all patients receive the same reference standard? Were all patients included	Unclear No		
in the analysis? Could the patient flow		High risk	

have introduced bias?

Nelson 2010

Study characteristics				
Patient Sampling	 Target population: HIV-1-infected treatment-naive patients from ARTEMIS phase III trial aimed a comparing the efficacy and safety of once-daily darunavir/ritonavir (800/100 mg) versus lopinavir ritonavir (800/200 mg total daily dose), each with a fixed-dose background tenofovir and emtric itabine regimen Recruitment: participants were randomly enrolled in routine clinical practice. Inclusion criteria: ARTEMIS inclusion criteria included treatment-naive HIV-1-infected patient 			
	aged at least 18 years, with plasma HIV-1 RNA at least 5000 copies/mL. Patients coinfected with hepatitis B and/or C virus were allowed entry if their condition was clinically stable and they were not expected to require treatment during the trial.			
	 Exclusion criteria: active AIDS-defining illness; any clinically significant disease; clinical or labora tory evidence of significantly decreased hepatic function or decompensation; acute viral hepatit is at screening or calculated creatinine clearance less than 70 mL/min. Individuals with primary HIV infection or those pregnant or breastfeeding were also excluded. Patients with grade 3 or laboratory abnormalities (division of AIDS grading table) were not eligible with some exception (diabetes or asymptomatic glucose, triglyceride or cholesterol elevations) unless clinical assessment identified health risks. 			
	Study design: a randomized, phase III, open-label multicentre trial			
Patient characteristics and setting	 Country: 26 countries World Bank Income classification: all Study setting: clinic-based 			
	Study dates: September 2005 to May 2008			
	• Age of population (years), mean (SD): DRV/r: 36 (9); LPV/r: 35 (9)			
	• Gender (male %): DRV/r: 70; LPV/r: 70			
	Participants included/analysed: 689/646			
	 First or second-line regimen: unclear Type of ART: DRV/r; DPV/r; DPV/r + LPV/r 			
	 Time on ART at enrolment: at least 6 months 			
	 Time on ART at measurement of viral load and adherence: 96 weeks 			
Index tests	Number of index tests used: 1			
	Types of index tests: self-report			
	 Test 1. Self-report questionnaire Validated scale: not applicable 			
	 Tool description: patients were asked to complete a modified medication adherence self-report inventory (M-MASRI) questionnaire and report their adherence to treatment over time (weeks 4, 12, 24, 36, 48, 60, 72, 84 and 96) 			
	Blinding: no information			
	 Threshold prespecified: not reported 			

Nelson 2010 (Continued)	 Adherence threshold used: > 95% 			
Target condition and refer-	 Target condition: viral non-suppression Reference standard: VircoTYPE HIV-1 assays (Virco BVBA; Mechelen, Belgium) Definition of viral non-suppression: HIV viral load > 50 copies/mL Blinded to index test: no information 			
ence standard(s)				
Flow and timing	 Time interval between index and reference tests: no explicit information on timing, just that adherence and viral load were assessed at 96 weeks All patients received same reference standard: not reported Missing data: almost all eligible participants with viral load test and adherence measures included in the main analysis. 			
Comparative				
Notes	Conflicts of interest:			
	M. N. has received research grants and travel bursaries and served as an advisor to Johnson & Johnson/Tibotec/Janssen-Cilag.			
	PM. G. has received grants and/or fees for conferences from BMS, Gilead, GSK, Tibotec and MSD, in the previous 12 months. R. D., V. S. and L. L. are employees of Tibotec.			
	E. S. is an employee of Johnson & Johnson Pharmaceutical Services.			
	LLC. L. C. was an employee of Tibotec at the time of the study.			
M. N., P.M. G. and L. C. do not own stock in any revelant companies.			companies.	
	R. D. has been granted res	ricted stock units in J&J.		
	E. S. and L. L. own stock options in J&J.			
	V. S. owns stock and stock options in J&J.			
	Jackie Phillipson (Gardine script and collating author		provided assistance n drafting the manu-	
	Funding source: Tibotec B	/BA		
	Trial registry: NCT0025855	7		
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Unclear			

Was a case-control design avoided?	Yes
Did the study avoid inappro- priate exclusions?	Yes
Could the selection of pa- tients have introduced bias?	Low risk

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Nelson 2010 (Continued)

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Are there concerns that the Low concern included patients and setting do not match the review question? DOMAIN 2: Index Test (Index test) Were the index test results in-Unclear terpreted without knowledge of the results of the reference standard? If a threshold was used, was it Yes pre-specified? Could the conduct or inter-Low risk pretation of the index test have introduced bias? Are there concerns that the Low concern index test, its conduct, or interpretation differ from the review question? **DOMAIN 3: Reference Standard** Is the reference standards like-Unclear ly to correctly classify the target condition? Were the reference standard Unclear results interpreted without knowledge of the results of the index tests? Could the reference stan-Unclear risk dard, its conduct, or its interpretation have introduced bias? Are there concerns that the Unclear target condition as defined by the reference standard does not match the question? **DOMAIN 4: Flow and Timing** Was there an appropriate in-Unclear terval between index test and reference standard? Unclear Did all patients receive the same reference standard? Were all patients included in Yes the analysis?

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Nelson 2010 (Continued)

Could the patient flow have introduced bias?

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Unclear risk

Study characteristics	
Patient Sampling	 Target population: adults on stable HAART for more than 3 months Recruitment: participants were randomly enrolled in routine clinical practice. Inclusion criteria: unselected cohort that underwent therapeutic drug monitoring while o stable HAART for more than 3 months Exclusion criteria: not reported Study design: prospective cohort study
Patient characteristics and setting	 Country: Germany (not explicity reported) World Bank Income classification: high-income Study setting: clinic-based (university outpatient unit specialized for infectious diseases) Study dates: 2002-2004 Age of population (years), mean (SD): 44.3 (10.4) Gender (male %): 76.7 Participants included/analysed: 210/208 First- or second-line regimen: unclear Type of ART: patients were treated with the following quantifiable antiretroviral drugs used as a single or boosted agent: ritonavir-boosted amprenavir (n = 10), ritor avir-boosted atazanavir (n = 30), efavirenz (n = 24), ritonavir-boosted lopinavir (n = 78 nelfinavir (n = 17), NVP (n = 23), ritonavirboosted saquinavir (n = 4). The following combinations of compounds were applied: efavirenz and ritonavir-boosted lopinavir (n = 4 nevirapine and ritonavir-boosted lopinavir (n = 2), nevirapine and ritonavir (n = 1) Time on ART at enrolment: at least 6 months Time on ART at measurement of viral load and adherence: 24 weeks
Index tests	 Number of index tests used: 2 Types of index tests: self-reports Test 1. Self-report on self-efficacy question Validated scale: not applicable Tool description: self-efficacy evaluating the belief of the patients to be able to be compliant with the prescribed combination scheme (yes; no) Blinding: no information Threshold prespecified: not reported Adherence threshold used: sufficient; low Test 2. Self-report on medication intake question Validated scale: not applicable Tool description:correctness of medication intake was assessed by the question of having forgotten a drug dose within the last 2 days, 14 days, last weekend or never. Blinding: no information Threshold prespecified: not reported

Dette 2006 (Continued)				
Target condition and reference stan- dard(s)	Target condition: viral non-suppression			
	 Reference standard: not Definition of viral non-si 	: reported uppression: HIV viral load > 5	50 copies/mL	
	Blinded to index test: no			
Flow and timing			o explicit information on timing, but	
	 both measures were taken at 24 weeks. All patients received same reference standard: not reported Missing data: almost all eligible participants with viral load test and adherence measures were included in the main analysis. 			
Comparative				
Notes	Conflicts of interest: none of	declared		
		tory costs were financed, in coSmithKline and Hoffmann	part, by the companies Abbott, La Roche.	
	Trial registry: not reported			
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Yes			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate ex- clusions?	Unclear			
Could the selection of patients have introduced bias?		Low risk		
Are there concerns that the in- cluded patients and setting do not match the review question?			Low concern	
DOMAIN 2: Index Test (Index test)				
Were the index test results interpret-	Unclear			
ed without knowledge of the results of the reference standard?				
If a threshold was used, was it pre- specified?	Unclear			
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk		



Oette 2006 (Continued)			
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condi-tion?	Unclear		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Unclear risk	
Are there concerns that the target condition as defined by the refer- ence standard does not match the question?			Unclear
DOMAIN 4: Flow and Timing			
Was there an appropriate interval be- tween index test and reference stan- dard?	Unclear		
Did all patients receive the same reference standard?	Unclear		
Were all patients included in the analysis?	Yes		
Could the patient flow have intro- duced bias?		Low risk	

Okonji 2012

Study characteristics	
Patient Sampling	 Target population: pregnant/lactating women (32–34 weeks gestation to 24 weeks postpar- tum)
	Recruitment: not reported
	 Inclusion criteria: participants were enrolled into this substudy, if they had adherence, vira load, and CD4 data in at least 3 time points during the intervention period and agreement to exclusively breastfeed up to 24 weeks postpartum.
	Exclusion criteria: not reported
	Study design: phase IIb open-label clinical trial
Patient characteristics and setting	Country: Kenya
-	 World Bank Income classification: low-income
	 Study setting: clinic-based (outpatients)
	Study dates: July 2003 to November 2007
courses of measures for antiretroviral	adherence in people living with HIV (Review)

Methodological quality	
	Trial registry: not reported
	Funding source: the Division of HIV/AIDS Prevention, Surveillance and Epidemiology, National Center for STD, HIV and TB Prevention, Atlanta, GA and by the Kenya Medical Research Institute (KEMRI) through a cooperative agreement with the US Centers for Disease Control and Preven- tion (KEMRI Protocol number 691/CDC Protocol number 3677)
Notes	Conflicts of interest: none declared
Comparative	
	 All patients received same reference standard: yes Missing data: yes; different numbers for adherence and viral load presented based on excluding different groups from analysis
Flow and timing	• Time interval between index and reference tests: no explicit details on timing, just that both adherence and VL were tested at baseline, 14, and 24 weeks
	 Reference standard, amplicon version 1.5 monitor test standard (Roche Diagnostics Systems, Branchburg, New Jersey) Definition of viral non-suppression: HIV viral load > 400 copies/mL Blinded to index test: no information
Target condition and reference standard(s)	Target condition: viral non-suppressionReference standard: amplicor version 1.5 monitor test standard (Roche Diagnostics Systems,
	 Threshold prespecified: no Adherence threshold used: ≥ 95%
Index tests	 (3TC/ZDV) regardless of baseline CD4+ cell counts. Time on ART at enrolment: at least 6 months Time on ART at measurement of viral load and adherence: > 24 weeks Number of index tests used: 1 Types of index tests: table counts Test 1. Table counts Validated scale: not applicable Tool description: adherence was calculated as the percent of pills dispensed that were actually taken. Drug calendar and self-report data were used only to further probe adherence issues among participants. Through pill count, adherence was calculated by trained pharmacy staff over a given period, by subtracting the number of pills returned during every scheduled visit from the number dispensed. Participants were also given a simple userfriendly drug calendar to mark date, day, and time (times in day were described in pictorial forms, i.e. sunrise, sunset, etc.) when the pills were taken. Participants returned the drug calendars to the pharmacy technician for review during clinic visits. Last, self-report through standard questionnaires administered during routine study visits were used to assess adherence. Participants were asked the number of doses they missed in the past 3 days and within a specified recall period (within the last one month). Blinding: no information Threshold prespecified: no
Dkonji 2012 (Continued)	 Age of population (years), median (IQR): 24 (15 to 43) Gender (male %): 0 (all women) Participants included/analysed: 500/434 First- or second-line regimen: Type of ART: initially enrolled participants were initiated on NVP/lamivudine/zidovudine

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



DOMAIN 1: Patient Selection			
Was a consecutive or random sam- ple of patients enrolled?	Unclear		
Was a case-control design avoid- ed?	Yes		
Did the study avoid inappropriate exclusions?	Unclear		
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the in- cluded patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Index test)			
Were the index test results inter- preted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre- specified?	Unclear		
Could the conduct or interpreta- tion of the index test have intro- duced bias?		Unclear risk	
Are there concerns that the in- dex test, its conduct, or inter- pretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the tar- get condition as defined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

Okonji 2012 (Continued)

Was there an appropriate interval between index test and reference standard?	Unclear
Did all patients receive the same reference standard?	Yes
Were all patients included in the analysis?	No
Could the patient flow have in- troduced bias?	High risk

Orrell 2003

Study characteristics	
Patient Sampling	 Target population: participants from the Cape Town AIDS Cohort (CTAC), a group of HIV-positive individuals presenting to University of Cape Town HIV clinics which serve largely indigent populations Recruitment: not reported Inclusion criteria: all antiretroviral naive patients who commenced ART on any established study by December 2000 were eligible for adherence monitoring. Exclusion criteria: not reported Study design: prospective cohort study
Patient characteristics and setting	 Country: South Africa World Bank Income classification: upper-middle-income Study setting: clinic-based (University of Cape Town HIV clinics) Study dates: January 1996 to May 2001 Age of population (years), mean (SD): 33.4 (8.7) Gender (male %): 57 Participants included/analysed: 289/278 First- or second-line regimen: unclear Type of ART: all participants were antiretroviral-naive and provided written consent to participate in multicentre phase III clinical trials of combination ART. They were assigned to one of six multicentre phase III studies. Participants in two studies in 1996 were given dual therapy with an additional third concurrent, placebo-controlled and double-blinded drug (placebo versus a non-nuceoside reverse transcriptase inhibitor regimen). In the other four studies, participants were given triple therapy regimens. Time on ART at enrolment: not reported; at least 6 months Time on ART at measurement of viral load and adherence: 48 weeks
Index tests	 Number of index tests used: 1 Types of index tests: composite measure Test 1. Composite measure: tablet counts + pharmacy records Validated scale: not applicable Tool description: adherence to therapy was assessed using clinic-based pill counts and pharmacy refill data over a period of 48 weeks. Patients were instructed to return all medication bottles and unused pills at each study visit, but were not told that the returns were to be counted. All tablets of each antiretroviral medication were counted prior to dispensing and upon return. Adherence to therapy was calculated using the

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Orrell 2003 (Continued) Target condition and reference stan- dard(s)	over the 48-week stud o Blinding: no informat o Threshold prespecifie o Adherence threshold Target condition: viral non- • Reference standard: not	dy interval). ion ed: no used: > 95% suppression reported ippression: HIV viral load > 4	ets returned)/(total tablets prescribed
Flow and timing	that both were measuredAll patients received san	d at 48 weeks ne reference standard: not r eligible participants with vi	o explicit information on timing, just eported ral load test and adherence measures
Comparative			
Notes	Conflicts of interest: none d	eclared	
	Funding source: M. B. was partially funded by a grant from the BMS 'Secure-The-Future' fund. D. B. received funding from The Doris Duke Charitable Foundation.		
	Trial registry: not reported		
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate ex- clusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the in- cluded patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Index test)			
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre- specified?	Unclear		

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Orrell 2003 (Continued)			
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condi- tion?	Unclear		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Unclear risk	
Are there concerns that the target condition as defined by the refer- ence standard does not match the question?			Unclear
DOMAIN 4: Flow and Timing			
Was there an appropriate interval be- tween index test and reference stan- dard?	Unclear		
Did all patients receive the same reference standard?	Unclear		
Were all patients included in the analysis?	Yes		
Could the patient flow have intro- duced bias?		Unclear risk	

Orrell 2017

Study characteristics	
Patient Sampling	 Target population: participants of an RCT at The Hannan Crusaid Treatment Centre in Gugulethu, Cape Town
	 Recruitment: entry into the study was offered consecutively to all eligible participants presenting to the clinic.
	 Inclusion criteria: participants were eligible for the parent study if they had their own mobile phone, signed an informed consent, and had either a baseline CD4 count below 350 cells/µL or a stage 3 or 4 AIDs-defining illness in keeping with the national HIV guidelines for starting ART. All patients on the parent study with viral load data available at week 16 or week 48 were included in this substudy.
	Exclusion criteria: not reported

prrell 2017 (Continued)	Study design: cohort (substudy of an RCT)
Patient characteristics and setting	 Country: South Africa World Bank Income classification: upper-middle-income Study setting: clinic-based (public sector urban ART outpatient clinic) Study dates: July 2012 to April 2014 Age of population (years), mean (SD): 34.5 (9.1) Gender (male %): 34.8 Participants included/analysed: 230/180 First- or second-line regimen: first-line Type of ART: first-line treatment at the time of the study included tenofovir, lamivudine, and efavirenz, given as 3 separate tablets once a day. Toward the end of the study period in October 2013, a fixed-dose combination became available, but priority was given to naive patients entering care and few of the study participants were switched to the fixed-dose combination during the study. Zidovudine, stavudine, nevirapine, and lopinavir in combination with ritonavir were available as alternative agents. Time on ART at enrolment: at least 6 months Time on ART at measurement of viral load and adherence: 48 weeks (treatment-naive patients with 96 weeks follow-up in the trial)
Index tests	Number of index tests used: 4
	Types of index tests: self-report; table count; pharmacy records or secondary database analysis; elec- tronic monitoring (MEMS)
	 Test 1. Self-report questionnaire Validated scale: not applicable Tool description: study staff who were not part of the clinical team asked each participant: "Did you
	swallow your pills yesterday/2 days ago/3 days ago?"Blinding: no information
	 Threshold prespecified: yes Adherence threshold used: 100%
	 Test 2. Tablets count Validated scale: not applicable
	 Tool description: not reported
	 Blinding: no information
	 Threshold prespecified: no
	• Adherence threshold used: \geq 95%
	 Test 3. Pharmacy records or secondary database analysis Validated scale: not applicable Tool description:
	Pharmacy refill: gaps method (PR-gaps). This measure used pharmacy-dispensing quantities and refill visit dates to determine the number off medication-free days (days when the partic ipant could not have had medication in hand) in each dispensing period. The number of med- ication-free days were subtracted from the number of days in the period, and the result divid- ed by the number of days in the period (up to week 16 or up to week 48) to give an adherence percentage.
	 Pharmacy refill: average method (PR-average). An electronic dispensing system (iDART) was used at the site to record the date of ART dispensed and the quantity given to each participant Obvious errors, such as date and dispensing duplications were removed. A cumulative PR-average measure was obtained at week 16 and week 48 visits by dividing the number of days o EFV, NPV or LPV/r tablets each patient received between study randomisation date and the visit date, by the number of days they were in care over the same period.
	Blinding: no information Threshold prospecified: no
	 Threshold prespecified: no



Drrell 2017 (Continued)				
	 Test 4. Electronic monito Validated scale: not a Tool description: Wise Blinding: no informat Threshold prespecifie Adherence threshold 	pplicable epill (electronic adherence m ion ed: no	onitoring)	
Target condition and reference standard(s)	 Target condition: viral non-suppression Reference standard: HIV-1 RNA 3.0 assay[®], Bayer Healthcare, Leverkusen, Germany) Definition of viral non-suppression: HIV viral load > 40 copies/mL Blinded to index test: no information 			
Flow and timing	 Time interval between index and reference tests: no explicit information on timing. All available adherence data were used from each individual who had an HIV-RNA drawn around week 48 (weeks 32–64) in a per-protocol analysis from the time they entered care until the date of the respective viral load. All patients received the same reference standard: yes Missing data: 180/230 patients retained at 48 weeks 			
Comparative				
Notes	Conflicts of interest: none declared			
	Funding source: the Discovery Foundation supported CO through an Academic Fellowship Award in 2013 and EDCTP awarded CO a senior fellowship from 2012 to 2014: TA.2011.40200.015.			
	Trial registry: PACTR201311	000641402		
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection	on			
Was a consecutive or ran- dom sample of patients enrolled?	Yes			
Was a case-control design avoided?	Yes			
Did the study avoid inap- propriate exclusions?	Unclear			
Could the selection of pa- tients have introduced bias?		Low risk		
Are there concerns that the included patients and setting do not match the review question?			Unclear	
DOMAIN 2: Index Test (Inde	ex test)			
Were the index test re-	Unclear			

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Orrell 2017 (Continued) knowledge of the results of the reference standard?			
If a threshold was used, was it pre-specified?	No		
Could the conduct or in- terpretation of the in- dex test have introduced bias?		Unclear risk	
Are there concerns that the index test, its con- duct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Stan	dard		
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference stan- dard results interpreted without knowledge of the results of the index tests?	Unclear		
Could the reference stan- dard, its conduct, or its interpretation have in- troduced bias?		Low risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Timin	g		
Was there an appropriate interval between index test and reference stan- dard?	Unclear		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	No		
Could the patient flow have introduced bias?		High risk	



Ortega 2004

Study characteristics	
Patient Sampling	 Target population: patients attending outpatient services and receiving HAART Recruitment: all patients attending outpatient services at the Hospitals of León and Bierzo, Spain Inclusion criteria: patients treated with HAART with two NRTI and a PI not boosted with ritonavir Exclusion criteria: not reported Study design: cross-sectional
Patient characteristics and setting	 Country: Spain World Bank Income classification: high-income Study setting: hospital-based (outpatient service) Study dates: January to June 2000 Age of population (years), mean (SD): 38 (7) Gender (male %): 72 Participants included/analysed: 136/136 First or second-line regimen: unclear Type of ART: HAART with two NRTI and a PI not boosted with ritonavir Time on ART at enrolment: at least 6 months Time on ART at measurement of viral load and adherence: not reported; at least 6 months
Index tests	 Number of index tests used: 2 Types of index tests: composite (self-report + pharmacy records) Test 1. Composite measure (self-report interview + pharmacy refill records) Validated scale: not reported Tool description: patients provided information about the doses taken during the 4 days preceding the interview, and the delay in collecting drugs in the last 3 months was also registered. Blinding: no information Threshold prespecified: yes Adherence threshold used: ≥ 10% forgotten doses ≥ 9 days delay Test 2. Composite measure (self-reported interview + pharmacy refill records + other: plasmatic concentration of PIs) Validated scale: not reported Tool description: patients provided information about the doses taken during the 4 days preceding the interview, and the delay in collecting drugs in the last 3 months was also registered. Blinding: no information of PIs) Validated scale: not reported Tool description: patients provided information about the doses taken during the 4 days preceding the interview, and the delay in collecting drugs in the last 3 months was also registered. In addition, plasmatic concentration of protease inhibitor were measured. Blinding: no information Threshold prespecified: yes Adherence threshold used: ≥ 10% forgotten doses ≥ 9 days delay ≥ 10% forgotten doses ≥ 9 days delay ≥ 10% forgotten doses ≥ 9 days delay
Target condition and reference stan- dard(s)	 Target condition: viral non-suppression Reference standard: not reported Definition of viral non-suppression: HIV viral load ≥ 400 copies/mL



Drtega 2004 (Continued)	• Blinded to index test: n	o information		
Flow and timing	 Time interval between index and reference tests: no explicit information on timing, but this was a cross-sectional study so likely to be measured simultaneously All patients received same reference standard: not reported Missing data: none. All eligible participants with viral load test and adherence measures were included in the main analysis. 			
Comparative				
Notes	Conflicts of interest: none	declared		
	Funding source: Wellcome Foundation and the Biomedical Research Institute (INBIOMED of the University of León, Spain			
	Trial registry: not reported	I		
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Yes			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate exclu- sions?	Unclear			
Could the selection of patients have introduced bias?		Low risk		
Are there concerns that the included patients and setting do not match the review question?			Unclear	
DOMAIN 2: Index Test (Index test)				
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear			
If a threshold was used, was it pre-spec- ified?	Unclear			
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk		
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear	
DOMAIN 3: Reference Standard				
Is the reference standards likely to cor- rectly classify the target condition?	Unclear			
ccuracy of measures for antiretroviral adhere	ence in people living with HIV (Review)	1	

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Ortega 2004 (Continued)			
Were the reference standard results in- terpreted without knowledge of the re- sults of the index tests?	Unclear		
Could the reference standard, its con- duct, or its interpretation have intro- duced bias?		Unclear risk	
Are there concerns that the target condition as defined by the reference standard does not match the ques- tion?			Unclear
DOMAIN 4: Flow and Timing			
Was there an appropriate interval be- tween index test and reference stan- dard?	Yes		
Did all patients receive the same refer- ence standard?	Unclear		
Were all patients included in the analy- sis?	Yes		
Could the patient flow have intro- duced bias?		Low risk	

Paolillo 2017

Study characteristics	
Patient Sampling	 Target population: HIV-infected adults enrolled in NIH-funded research studies at the University of California, San Diego, HIV Neurobehavioral Research Program (HNRP) from 2003 to 2015 Recruitment: not reported Inclusion criteria: participant's baseline visit at the HNRP who were receiving ART at the time of the visit and reported drinking alcohol in the previous 30 days Exclusion criteria: non-drinkers Study design: cohort (participant's baseline visit at the HNRP)
Patient characteristics and setting	 Country: USA World Bank Income classification: high-income Study setting: clinic-based (outpatient research program) Study dates: 2003 to 2015 Age of population (years), mean (SD): 42.2 (8.6) Gender (male %): 85.6 Participants included/analysed: 535/535 First- or second-line regimen: unclear Type of ART: NRTI plus PI; NRTI plus NNRTI. Other less common ART regimen types included other combinations of the six ART drug classes. Time on ART at enrolment: at least 6 months Time on ART at measurement of viral load and adherence: median duration of exposure to ART was 63 months (IQR: 28 to 105)

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

Paolillo 2017 (Continued)				
Index tests	Number of index tests used: 1			
	Types of index tests: self-report			
	 Test 1. Self-report questionnaire Validated scale: not applicable Tool description: self-report AIDS Clinical Trial Group (ACTG), a questionnaire indicating any missed ART doses in the previous 4 days Blinding: no information Threshold prespecified: yes Adherence threshold used: 100% 			
Target condition and reference stan-	Target condition: viral non-suppression			
dard(s)	 Reference standard: not reported Definition of viral non-suppression: HIV viral load ≥ 50 copies/mL Blinded to index test: no information 			
Flow and timing	 Time interval between index and reference tests: not explicit, but seemed that both measures were taken on the same day All patients received same reference standard: not reported Missing data: none. All eligible participants with viral load test and adherence measures 			
	were included in the main analysis.			
Comparative				
Notes	Conflicts of interest: none declared			
	Funding source: Emily W. Paolillo is supported by an Institutional Ruth L. Kirschstein Na- tional Research Service Award (NRSA) T32 grant funded by the NIAAA within the National Institutes of Health (Award T32 AA013525). Data for this study was collected as part of five larger ongoing studies: 1) The CNS HIV Anti-Retroviral Therapy Effects Research (CHARTER) is supported by awards N01 MH22005, HHSN271201000036C, and HHSN271201000030C from NIH; 2) the California NeuroAIDS Tissue Network (CNTN) is supported by awards U01MH083506, R24MH59745 from NIMH; 3) the HIV Neurobehavioral Research Center (not reported) is supported by Center award P30MH062512 from NIMH; 4) the Transla- tional Methamphetamine AIDS Research Center (TMARC) is supported by Center award P50DA026306 from the National Institute on Drug Abuse (NIDA); and 5) a NIDA grant award P01DA1206507			
	Trial registry: not reported			
Methodological quality				
Item	Authors' judgement Risk of bias Applicability concerns			
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Unclear			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate ex- clusions?	Unclear			

Could the selection of patients have introduced bias?

=

Unclear risk

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Paolillo 2017 (Continued)			
Are there concerns that the includ- ed patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Index test)			
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre- specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to cor- rectly classify the target condition?	Unclear		
Were the reference standard results in- terpreted without knowledge of the re- sults of the index tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Unclear risk	
Are there concerns that the target condition as defined by the refer- ence standard does not match the question?			Unclear
DOMAIN 4: Flow and Timing			
Was there an appropriate interval be- tween index test and reference stan- dard?	Unclear		
Did all patients receive the same refer- ence standard?	Unclear		
Were all patients included in the analy- sis?	Yes		
Could the patient flow have intro- duced bias?		Unclear risk	



Parienti 2010	
Study characteristics	
Patient Sampling	 Target population: HIV-infected patients treated with ritonavir-boosted PI from two multicentre cohort studies (Etude et Surveillance par Pilulier électronique de l'Observance et de l'Incidence de la Réplication virale-ESPOIR and RE-search in Access to Care in the Homeless-REACH) Recruitment: The ESPOIR cohort selected consecutive patients receiving or starting twice-aday LPV/r-based regimens and monitored adherence in several outpatient clinics in France. The REACH cohort selected HIV-positive homeless and marginally housed individuals in San Francisco, California. Inclusion criteria: not reported Exclusion criteria: not reported Study design: analysis of data from prospective cohort studies
Patient characteristics and setting	 Country: France, USA World Bank Income classification: high-income Study setting: hospital- and community-based (the ESPOIR cohort: several outpatient clinics)
	in France; the REACH cohort: HIV-positive homeless and marginally housed individuals in San Francisco)
	Study dates: December 2006 to December 2008
	Age of population (years), mean (SD): 43.9 (7.5)
	Gender (male %): 82
	Participants included/analysed: 72/72 First or second line regiment and the second line regiment
	 First- or second-line regimen: not reported Type of ART: The ESPOIR cohort: twice-a-day lopinavir-ritonavir-based regimens; REACH cohort: ritonavir-boosted PI
	 Time on ART at enrolment: at least 6 months
	 Time on ART at measurement of viral load and adherence: at least 24 months as cohort fol- lowed for this time
Index tests	Number of index tests used: 1
	Types of index tests: composite measure
	 Test 1. Composite measure (electronic monitoring ± tablet counts, lopinavir plasma levels) Validated scale: not reported
	 Tool description: 6 caps of the Medication Event Monitoring System (MEMS) which measures patterns of missed doses with a time/date record of pill bottle opening behaviour. Average per cent dose adherence was defined as the number of MEMS events (pill bottle openings), divided by the number of prescribed doses, multiplied by 100. In addition, adherence was confirmed by measuring lopinavir plasma levels in the ESPOIR cohort and by having monthly unannounced pill counts in the REACH cohort.
	 Blinding: no information
	 Threshold prespecified: yes; (1) number of days without a dose, defined as drug discontinuation for > 24 hours and < 48 hours, (2) number of treatment interruptions lasting ≥ 48 hours, and (3) the duration of the longest treatment interruption (in days) Adherence threshold used: > 95%; > 80%; > 70%
Target condition and refer-	Target condition: viral non-suppression
ence standard(s)	Reference standard: not reported
	 Definition of viral non-suppression: HIV viral load > 50 copies/mL and > 400 copies/mL
	Blinded to index test: no information
Flow and timing	 Time interval between index and reference tests: not reported All patients received same reference standard: not reported



Parienti 2010 (Continued) Missing data: none. All eligible participants with viral load test and adherence measures were in-• cluded in the main analysis. Comparative Notes Conflicts of interest: J.-J.P. reported that he has received travel grants, honoraria for presentation at workshops, and consultancy honoraria from Abbott, Boehringer Ingelheim, and Bristol-Myers Squibb. Y.Y. reported that he has received travel grants, honoraria for presentation at workshops, and consultancy honoraria from Bristol-Myers Squibb, Boehringer Ingelheim, Gilead, GlaxoSmithKline, Merck, Pfizer, Roche, and Tibotec. All other authors: no conflicts declared Funding source: The ESPOIR cohort was supported by an Abbott Laboratories unrestricted grant (to Caen Côte de Nacre University hospital). The REACH cohort was supported by the National Institute of Mental Health (grant RO-54907) and the National Institute on Alcohol Abuse and Alcoholism (grant K-24 015287). Trial registry: not reported Methodological quality Item **Authors' judgement Risk of bias Applicability concerns DOMAIN 1: Patient Selection** Was a consecutive or random Unclear sample of patients enrolled? Yes Was a case-control design avoided? Did the study avoid inappro-Unclear priate exclusions? Could the selection of pa-Unclear risk tients have introduced bias? Are there concerns that the Unclear included patients and setting do not match the review question? DOMAIN 2: Index Test (Index test) Were the index test results in-Unclear terpreted without knowledge of the results of the reference standard? If a threshold was used, was it Yes pre-specified?



Parienti 2010 (Continued)				
Could the conduct or inter- pretation of the index test have introduced bias?		Low risk		
Are there concerns that the index test, its conduct, or in- terpretation differ from the review question?			Unclear	
DOMAIN 3: Reference Standard				
Is the reference standards like- ly to correctly classify the tar- get condition?	Unclear			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear			
Could the reference stan- dard, its conduct, or its inter- pretation have introduced bias?		Unclear risk		
Are there concerns that the target condition as defined by the reference standard does not match the ques- tion?			Unclear	
DOMAIN 4: Flow and Timing				
Was there an appropriate in- terval between index test and reference standard?	Unclear			
Did all patients receive the same reference standard?	Unclear			
Were all patients included in the analysis?	Yes			
Could the patient flow have introduced bias?		Unclear risk		

Parker 2017

Study characteristics	
Patient Sampling	 Target population: adults from AIDS Clinical Trials Group (ACTG) A5202, which randomized ART-naive, HIV- infected participants to receive placebo-controlled abacavir–lamivudine or TDF–emtricitabine with open- label ATV/r or EFV
	Recruitment: not reported
	 Inclusion criteria: age ≥ 18 years, HIV type 1 infected, had 7 days or less of ART prior to enrolment, informed consent obtained



Parker 2017 (Continued)	 Exclusion criteria: significant drug or alcohol abuse thought likely to impact adherence Study design: open-label RCT
Patient characteristics and setting	 Country: USA World Bank Income classification: high-income Study setting: not reported Study dates: 2005 to 2009 Age of population (years), median (IQR): 38 (31 to 45) Gender (male %): 83 Participants included/analysed: 1649/1857 First- or second-line regimen: unclear Type of ART: ATV/r or EFV Time on ART at enrolment: at least 6 months Time on ART at measurement of viral load and adherence: at least 6 months as RCT with 24 weeks follow-up time
Index tests	 Number of index tests used: 1 Types of index tests: self-report Test 1. Self-report single question Validated scale: not applicable Tool description: 6 potential responses to the question "When was the last time you missed any of your medications?" to assess adherence (never skip medications, more than 3 months ago, 1–3 months ago, 2–4 weeks ago, 1–2 weeks ago, and within the past week). Participants who did not provide a self-report form were considered to be not adherent for that report. Never Skip Medications (3 categories): 1. Adherent (reported never or missing their last dose more than 1 month ago on both the week 8 and week 24 reports); 2. Not adherent (reported missing their last dose within the past month on the available week 8 and week 24 reports or missing both re- ports); 3. Inconsistent (reported adherence on either the week 8 or week 24 report;); 2. Not adherent (reported missing their last dose more than 1 month ago on both the week 8 and week 24 reports); 2. Not adherent (reported never or missing both report) Missed pill more than 3 months ago or never skip medications (3 categories): 1. Adherent (reported never or missing both reports); 3. Inconsistent (reported adherence on either the week 8 ard week 24 reports or missing both reports); 3. Inconsistent (reported adherence on either the week 8 ard week 24 reports). The adherent on the other report or missing the other report) Missed pill 1.3 months ago, more than 3 months ago or never skip medications (3 categories): 1. Adherent (reported never or missing their last dose within the past month on the available week 8 and week 24 reports); 2. Not adherent (reported missing their last dose more than 1 month ago on both the week 8 and week 24 reports); 2. Not adherent (reported missing their last dose more than 1 month ago on both the week 8 and week 24 reports); 2. Not adherent (reported missing their last dose within the past month on the available week 8 ard we
	 Threshold prespecified: not reported Adherence threshold used: adherent or inconsistent, non-adherent

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

Parker 2017 (Continued)			
Target condition and reference standard(s)	Target condition: viral non-suppression		
	Reference standard: Roche Amplicor Monitor Assay Version 1.5		
	 Definition of viral non-suppression: HIV viral load > 200 copies/mL 		
	Blinded to index test: no information		
Flow and timing	• Time interval between index and reference tests: viral load measured at 24 weeks; adherence behavior through week 24		
	All patients received same reference standard: yes		
	Missing data: 89% continued on study after the week-24 visit.		
Comparative			
Notes	Conflicts of interest:		
	A. C. C reported research grants to her institution from Bristol-Myers Squibb, Merck & Co., and Roche Molec- ular Systems; Data Safety and Monitoring Board membership for Merck & Co.–sponsored clinical trials; and has developed educational presentation for International Antiviral Society-USA.		
	E. S. D. is a consultant/advisor for Bristol Myers Squibb, Gilead, Janssen, Merck, Teva, and ViiV and has re- ceived research support from Gilead, Merck, and ViiV.		
	All other authors: No reported conflicts. All authors have submitted the ICMJE Form for Disclosure of Po- tential Conflicts of Interest. Conflicts that the editors considered relevant to the content of the manuscript have been disclosed.		
	Funding source: National Institute of Allergy and Infectious Diseases of the National Institutes of Health (NIH; Al042006, Al068636, Al069481 and UM1Al068634) and the Harvard University Center for AIDS Re- search, an NIH-funded program (P30 Al060354)		
	Trial registry: AIDS Clinical Trials Group A5202		
Methodological quality	/		

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Sele	ection		
Was a consecutive or random sample of pa- tients enrolled?	Unclear		
Was a case-control de- sign avoided?	Yes		
Did the study avoid inappropriate exclu- sions?	No		
Could the selection of patients have in- troduced bias?		High risk	
Are there concerns that the included pa- tients and setting do not match the review question?			High

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Parker 2017 (Continued)

DOMAIN 2: Index Test (Index test)		
Were the index test re- sults interpreted with- out knowledge of the results of the refer- ence standard?	Unclear		
If a threshold was used, was it pre-speci- fied?	Unclear		
Could the conduct or interpretation of the index test have intro- duced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or inter- pretation differ from the review question?			Low concern
DOMAIN 3: Reference S	Standard		
Is the reference stan- dards likely to correct- ly classify the target condition?	Yes		
Were the reference standard results inter- preted without knowl- edge of the results of the index tests?	Unclear		
Could the reference standard, its con- duct, or its interpre- tation have intro- duced bias?		Low risk	
Are there concerns that the target condi- tion as defined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Ti	ming		
Was there an appro- priate interval be- tween index test and reference standard?	Unclear		



Parker 2017 (Continued)

Did all patients receive the same reference standard?	Yes	
Were all patients in- cluded in the analysis?	No	
Could the patient flow have introduced		High risk

bias?

Pasquau 2018

Study characteristics	
Patient Sampling	 Target population: adults with a positive HIV-1 antibody and/or PCR test, receiving any triple treatment containing a boosted PI
	Recruitment: not reported
	 Inclusion criteria: patients infected with HIV-1, documented with a positive HIV-1 antibodies test and/ or positive PCR confirmed for HIV-1 RNA, with an undetectable VL within the last six months and on triple antiretroviral therapy with any boosted PI. For women with childbearing potential, negative urine pregnancy test during the screening visit
	 Exclusion criteria: pregnancy or nursing, acute hepatitis, documented resistance to LPV/r or failure on a PI therapy, concomitant therapy with drugs contraindicated for use with LPV/r, known history of drug addiction or chronic alcohol consumption, current active opportunistic infection or documented infec- tion within 4 weeks of screening, renal disease with creatinine clearance < 60 mL/min, concomitant use of nephrotoxic or immunosuppressor drugs including corticosteroids, interleukin-2 or chemotherapy, prior medical history of psychiatric disorders such as depressive syndrome, schizophrenia or psychotic disease
	Study design: open-label RCT
Patient characteristics and setting	 Country: Spain World Bank Income classification: high-income
5	 Study setting: hospital-based
	Study dates: January 2010 to December 2011
	 Age of population (years), mean (SD): monotherapy group 44.5 (8); triple therapy group 45.2 (9)
	 Gender (male %): monotherapy group: 71.4; triple therapy group: 71.8
	 Participants included/analysed: 225/197
	 First- or second-line regimen: unclear
	 Type of ART: LPV/r in monotherapy or continuing combined antiretroviral triple treatment with a boosted Pl
	 Time on ART at enrolment: at least 6 months
	 Time on ART at measurement of viral load and adherence: at least 6 months as RCT with 24 weeks follow-up time
Index tests	Number of index tests used: 1
	Types of index tests: self-report
	 Test 1. Self-report questionnaire Validated scale: yes (GEEMA adherence questionnaire)
	 Tool description: this questionnaire included six individual questions. Four of the questions were qualitative ("Do you ever forget to take your medicine?", "Are you careless at times about taking your medicine?", "Sometimes if you feel worse, do you stop taking your medicine?" and "Did you not take any of your medicine over the past weekend?"); the other two questions ("Thinking about the last
	tiretroviral adherence in people living with HIV (Peview)



Pasquau 2018 (Continued)	week Hew efter heve	un un ant talkan yayı maadinin 200 a	and "Cines the last visit have many days have
		icine at all?") were independent	and "Since the last visit how many days have ly quantified and analysed.
	 Blinding: no reported 	, i	
	 Threshold prespecified 	: yes	
	 Adherence threshold u 	sed: 100% (overall GEMMA queti	onnaire)
Target condition and ref- erence standard(s)	Target condition: viral non-su	uppression	
	Reference standard: not re	•	
		pression: HIV viral load > 200 co	pies/mL and > 50 copies/mL
	Blinded to index test: no in	nformation	
Flow and timing	• Time interval between inc	lex and reference tests: 24 weeks	5
		e reference standard: not reporte	
	Missing data: 197 out of 22	25 study participants were inclue	ded in the analysis.
Comparative			
Notes	Conflicts of interest:		
		ncial grants and/or honoraria fro /iiV&Gilead as speaker for fees a	m Janssen-Cilaq, Bristol-Myers-Squibb, Ab- nd/or as Advisor fees.
	M.l. Montes has served as a s bie.	peaker for Jannsen, BMS, ViiV, A	bVie, a consultant for Janssen, BMS and Ab-
		ch grants and/or honoraria for a ′iiV, BMS, Abbott, Gildead, Janss	dvisories and/or conferences from en, Roche Farma and Merck.
			oraria from Jannsen-Cilaq, Bristol-My- beaker fees and/or as a Advisor fees.
		nent for training sessions from A nssen, Merck-Sharp & Dohme an	bbVie, Boehring Ingelheim, Bristol-Myers d ViiV Healthcare.
	funds for research from Abbv Cilag, Merck-Shap & Dohme a	ie, Boehringer Ingelheim, Bristl I and ViiV healthcare. F. Lozano ha	ultancies and educational activities, or Myers Squibb, Gilead Sciences, Janssen- s acted as a consultant for AbbVie, Janssen, Merck-Sharp & Dohme, and ViiV
		iiV healthcare and has received	Ayers Squibb, Gilead Sciences, Janssen, payment for training sessions for AbbVie,
			as sponsored by the Spciedad Andaluza de dy design, preparation of the final report
	Trial registry: NCT 01166477;	EudraCT number 2009-014430-2	25
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Select	ion		
Was a consecutive or ran- dom sample of patients enrolled?	Unclear		



asquau 2018 (Continued)			
Was a case-control de- sign avoided?	Yes		
Did the study avoid inap- propriate exclusions?	Yes		
Could the selection of patients have intro- duced bias?		Low risk	
Are there concerns that the included pa- tients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Ind	lex test)		
Were the index test re- sults interpreted without knowledge of the results of the reference stan- dard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have intro- duced bias?		Low risk	
Are there concerns that the index test, its con- duct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Sta	ndard		
Is the reference stan- dards likely to correctly classify the target condi- tion?	Unclear		
Were the reference stan- dard results interpreted without knowledge of the results of the index tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Unclear risk	
Are there concerns that the target condition as defined by the refer-			Unclear



match the question?			
DOMAIN 4: Flow and Timi	DOMAIN 4: Flow and Timing		
Was there an appropriate interval between index test and reference stan- dard?	Unclear		
Did all patients receive the same reference stan- dard?	Unclear		
Were all patients includ- ed in the analysis?	No		
Could the patient flow have introduced bias?	High risk		

Phillips 2019

Study characteristics	
Patient Sampling	 Target population: women who were enrolled in the MCH-ART study at the Midwife-Obstetric Unit, Gugulethu Community Health Centre, Cape Town Recruitment: not reported Inclusion criteria: the first 150 women between 36 and 60 months postpartum who agreed to participate and had blood drawn for ARV assays Exclusion criteria: women who were pregnant or had switched to second-line ART Study design: cross-sectional
Patient characteristics and setting	 Country: South Africa World Bank Income classification: upper-middle-income Study setting: community health centre Study dates: March 2013 to March 2017 (MCH-ART parent study) Age of population (years), mean: 33 Gender (male %): 0 (all women) Participants included/analysed: 137/137 First- or second-line regimen: first-line Type of ART: first-line regimen of TDF (300 mg), emtricitabine (200 mg) or lamivudine 300 mg (XTC), and EFV 600 mg, provided as a once-daily fixed-dose combination Time on ART at enrolment: at least 6 months Time on ART at measurement of viral load and adherence: median (IQR): 3.9 years (3.7 to 4.0)
Index tests	 Number of index tests used: 1 Types of index tests: self-report Test 1. Self-report questionnaire Validated scale: not applicable Tool description: medication adherence in the past 30 days was measured using a simple, 3-item scale. Blinding: no information



Phillips 2019 (Continued)			
	 Threshold prespecifi Adherence threshold 		
Target condition and reference	Target condition: viral non	-suppression	
standard(s)	tics, Branchburg, New J	lersey uppression: HIV viral load >	BAS TaqMan HIV-1 assay; Roche Diagnos- 400 copies/mL and > 50 copies/mL
Flow and timing		index and reference tests: a It the same day (study visit)	dherence measured and blood drawn for
			al load test and adherence measures were
Comparative			
Notes	Conflicts of interest: none	declared	
	Funding source: The President's Emergency Plan for AIDS Relief (PPFAR) through the National Institute of Child Health and Human Development (NICHD), grant number 1R01HD074558 and 1R01HD080465. The University of Cape Town (UCT) Clinical PK Laboratory is supported by the National Institute of Allergy and Infectious Diseases (NIAID) of the National Institutes of Health under award numbers UM1 AI068634, UM1 AI068636, and UM1 AI106701. Overall support for the International Maternal Pediatric Adolescent AIDS Clinical Trials Group (IMPAACT) at UCT was provided by the National Institute of Allergy and Infectious Diseases (U01 AI068632), The Eunice Kennedy Shriver National Institute of Child Health and Human Development, and Na- tional Institute of Mental Health grant AI068632. Ms Phillips receives partial funding from the South African Department of Science and Technology/National Research Foundation (DST - not reported), Centre of Excellence in Epidemiological Modelling and Analysis (SACEMA), Stel- lenbosch University, Stellenbosch, South Africa. Dr Orrell is partially supported through DAIDS grants (1R01AI122300–01, 1R34MH108393-01 and 2UM1AI0695-08).		
	Trial registry: NCT0193347	7 (parent study)	
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sam- ple of patients enrolled?	Yes		
Was a case-control design avoid- ed?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the in- cluded patients and setting do not match the review question?			Low concern

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Phillips 2019 (Continued)

DOMAIN 2: Index Test (Index test)			
Were the index test results inter- preted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre- specified?	Unclear		
Could the conduct or interpreta- tion of the index test have intro- duced bias?		Unclear risk	
Are there concerns that the in- dex test, its conduct, or inter- pretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the tar- get condition as defined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Yes		
Could the patient flow have in- troduced bias?		Low risk	

Pulido 2009	
Study characteristics	
Patient Sampling	 Target population: all patients enrolled in the OK and OK04 clinical trials randomized to receive LPV/r monotherapy Recruitment: likely consecutive Inclusion criteria: all patients enrolled in the OK and OK04 clinical trials randomized to receive LPV/r monotherapy Briefly, patients included in both trials did not have a history of virological failure while receiving a protease inhibitor, were receiving two NRTIs and LPV/r for ≥ 1 month prior to randomization and had serum HIV-1 RNA < 50 copies/mL for ≥ 6 months prior to randomization. Exclusion criteria: pregnancy, presence of serum hepatitis B surface antigen in patients treated with lamivudine, emtricitabine or tenofovir disoproxil fumarate, need for treatment with agents known to have potential major interactions with LPV/r and major psychiatric diseases as assessed by the investigator Study design: two arms from two RCTs
Patient characteristics and setting	 Country: Spain World Bank Income classification: high-income Study setting: clinic-based Study dates: OK 04 trial: December 2004 to June 2006; OK pilot trial: May 2003 to August 2004 Age of population (years), median (range): OK pilot: 42 (25-54); OK 04: 41 (28-78) Gender (male %): OK pilot: 81; OK 04: 76 Participants included/analysed: 121/121 First- or second-line regimen: unclear Type of ART: LPV/r monotherapy Time on ART at enrolment: ≥ 6 month Time on ART at measurement of viral load and adherence: at least 6 months; suppressed: median (IQR) 23 (15-32) months and non-suppressed: median (IQR): 26 (13-34) months on LPV/r
Index tests	Number of index tests used: 1
	Types of index tests: self-report
	 Test 1. Self-report questionnaire Validated scale: yes (GEEMA adherence questionnaire) Tool description: this questionnaire included six individual questions. Four of the questions were qualitative ("Do you ever forget to take your medicine?", "Are you careless at times about taking your medicine?", "Sometimes if you feel worse, do you stop taking your medicine?" and "Did you not take any of your medicine over the past weekend?"); the other two questions ("Thinking about the last week. How often have you not taken your medicine?" and "Since the last visit how many days have you not taken any medicine at all?") were independently quantified and analysed. A missed dose on ≥ 2 visits in the week prior to study visit was considered as non-adherence. Blinding: no information Threshold prespecified: yes Adherence threshold used: 100%
Target condition and refer- ence standard(s)	 Target condition: viral non-suppression Reference standard: automatized RNA extraction in an Cobas AmpliPrep instrument followed by quantification using the Cobas TaqMan HIV1 in a TaqMan 48 analyzer (Roche Molecular Systems, Inc., Branchburg, New Jersey, USA) Definition of viral non-suppression: HIV viral load > 50 copies/mL Blinded to index test: no information
Flow and timing	Time interval between index and reference tests: up to one weekAll patients received same reference standard: yes



Pulido 2009 (Continued)

• Missing data: none. All eligible participants with viral load test and adherence measures were included in the main analysis.

Comparative	
Notes	Conflicts of interest:
	FP is the recipient of a BAE grant from the Instituto de Salud Carlos III, Spanish Ministry of Health.
	JRA is an investigator from the Programa de Intensificación de la Actividad Investigadora, National Health System (I3SNS) 2008 INT07/147.
	FP and JRA have received consulting and lecture fees from Abbott Laboratories, Bristol–Myers Squibb Gilead Sciences, GlaxoSmith-Kline and Roche.
	RD has received grant support and lecture fees from Abbott Laboratories.
	MJP-E was an occasional speaker and advisor for Abbott Labora-tories, Bristol–Myers Squibb, Boer- ingher–Ingelheim, Gilead, GlaxoSmithKline, Roche and Tibotec.
	J Portilla has received lecture fees from Abbott Laboratories, Bristol–Myers Squibb, GlaxoSmithKline, Roche, Schering–Plough and Boehringer–Ingelheim.
	BC has served as a consultant on advisory boards, speakers' bureaus and in the conduct of clinical tri- als with Roche, Boehringer–Ingelheim, Abbott Laboratories, Bristol–Myers Squibb, GlaxoSmithKline, Gilead, Tibotec, Merck, Janssen, Pfizer, Siemens, Monogram Biosciences and Panacos.
	The other authors declared no competing interests.
	Funding source: Abbott Laboratories and the Fundación de Investigación Médica Mutua Madrileña
	Trial registry: NCT00114933

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection	n		
Was a consecutive or ran- dom sample of patients en- rolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inap- propriate exclusions?	Yes		
Could the selection of pa- tients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Index	x test)		



Pulido 2009 (Continued)			
Were the index test results interpreted without knowl- edge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or in- terpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standa	ard		
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference stan- dard results interpreted without knowledge of the results of the index tests?	Unclear		
Could the reference stan- dard, its conduct, or its in- terpretation have intro- duced bias?		Low risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Unclear		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Yes		
Could the patient flow have introduced bias?		Low risk	



Sangeda 2014

Study characteristics	
Patient Sampling	 Target population: adults with HIV starting ART or being on ART at a clinic Recruitment: convenient sample Inclusion criteria: HIV-infected adult patients either starting ART or being on ART who attended an HIV/AIDS Care and Treatment Centre at Amana District Hospital in Dar es Salaam, Tanzania, in 2010 Exclusion criteria: < 18 years, pregnancy, having opportunistic infections, or malignancy Study design: prospective cohort study
Patient characteristics and setting	 Country: Tanzania World Bank Income classification: low-income Study setting: clinic-based Study dates: May to July 2010 Age of population (years), median (IQR): 39 (34 to 47) Gender (male %): 36.2 Participants included/analysed: 220/162 First- or second-line regimen: not reported Type of ART: not reported Time on ART at enrolment: at least 6 months Time on ART at measurement of viral load and adherence: median (IQR): 37 months (30 to 48)
Index tests	 Number of index tests used: 3 Types of index tests: self-report, tablet counts, and pharmacy records or secondary database analysis Test 1. Self-report tool Validated scale: yes Tool description: two major sections: 1) a VAS which probed the percentage of doses taken in the previous month, and 2) two questions from the Swiss HIV Cohort Study Adherence Questionnaire (SHCS-AQ) regarding frequency of missed doses and if a patient ever missed two consecutive doses (drug holiday) in the previous month Blinding: no information Threshold prespecified: yes Adherence threshold used: VAS: 90%; 95%; 100% Two questions: 100% Test 2. Tablet count Validated scale: not applicable Tool description: at each visit, pills remaining in bottles were counted and the proportion of these pills to the dispensed pills during the previous visit was calculated based on the dose and the number of days dispensed. The pill count adherence percent was obtained by dividing the number of pills consumed by the total number of pills at the beginning of the given interval and multipiled by 100. Blinding: no information Threshold preespecified: not reported Adherence threshold used: 50%; 55%; 60%; 65%; 70%; 75%; 80%; 85%; 90%; 95%; 100% Test 3. Pharmacy records or secondary database analysis Validated scale: not applicable Tool description: Refill adherence was not calculated on a monthly basis to reduce an error of a few additional pills left over at the end of each refill period, but on the cumulative sum of the days that a patient was late for ARV pick-up appointments in each month over the year, divided by the total number of days over all periods between pick-up periods in the year of study, resulting in the percentage of time the patient was without medication over the whole year. Refill adherence was 100% if all pills left over at the end of each ref



Sangeda 2014 (Continued)	 Threshold preespecifi Adherence threshold	-	%; 75%; 80%; 85%; 90%; 95%; 100%
	Adherence measurements v cruitment (zero), one, two, a the mean of the measureme quent analyses. For self-rep	vere taken at four time points d and 12 months after recruitmer ents taken at the four time poin	luring a one-year follow-up, including at re- nt. Overall adherence for each method was ts and this mean was considered in subse- s, the 12-month visit adherence measure
Target condition and refer-	Target condition: viral non-s	suppression	
ence standard(s)	Reference standard: not		
	Definition of viral non-suBlinded to index test: no	ppression: HIV viral load > 400 information	copies/mL
Flow and timing	herence and viral load m	easures were at the same visit ole time period was used.	f-report measures and tablet count, the ad . For pharmacy refill, an overall measure c
			not included in analysis. Missing data > 10%
Comparative			
Notes	Conflicts of interest: none d	eclared	
	Vlaanderen; MRC grant in as	sociation with University of Ma	onds voor Wetenschappelijk Onderzoek nchester's Health Research Centre; au- rsity of Health and Allied Sciences (MUHAS)
	Trial registry: not reported		
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection	n		
Was a consecutive or ran- dom sample of patients en- rolled?	No		
Was a case-control design avoided?	Yes		
Did the study avoid inap- propriate exclusions?	Yes		
Could the selection of pa- tients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the			Low concern

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Sangeda 2014 (Continued)					
Were the index test results interpreted without knowl- edge of the results of the reference standard?	Unclear				
If a threshold was used, was it pre-specified?	Unclear				
Could the conduct or in- terpretation of the index test have introduced bias?		Unclear	risk		
Are there concerns that the index test, its conduct, or interpretation differ from the review question?				Low concern	
DOMAIN 3: Reference Standa	ard				
Is the reference standards likely to correctly classify the target condition?	Unclear				
Were the reference stan- dard results interpreted without knowledge of the results of the index tests?	Unclear				
Could the reference stan- dard, its conduct, or its in- terpretation have intro- duced bias?		Unclear	risk		
Are there concerns that the target condition as defined by the reference standard does not match the question?				Unclear	
DOMAIN 4: Flow and Timing					
Was there an appropriate interval between index test and reference standard?	Yes				
Did all patients receive the same reference standard?	No				
Were all patients included in the analysis?	No				
Could the patient flow have introduced bias?		High risk			

Segeral 2010

Study characteristics	
Patient Sampling	 Target population: adults with HIV/AIDS receiving HAART for at least six months Recruitment: likely consecutive (survey) Inclusion criteria: HAART prescribed according to WHO recommendations (WHO stages III and N irrespective of the CD4 cell count, or asymptomatic patients with CD4 cell counts ≤ 200/μL) bot to ARV-naive patients and to patients having previously ARV paid for themselves; all patients wh had at least one adherence assessment were included in the analysis. Exclusion criteria: not reported Study design: cross-sectional
Patient characteristics and setting	 Country: Cambodia World Bank Income classification: low-income Study setting: outpatient clinic Study dates: not reported Age of population (years), median (IQR): 35 (31 to 41) Gender (male %): 58 Participants included/analysed: 341/259 First- or second-line regimen: first-line NNRTI-based regimens Type of ART: HAART combination was (AZT or d4T)/3TC/EFV initially, but was then switched t (AZT or d4T)/3TC/NVP after July 2004, owing to EFV supply problems Time on ART at enrolment: at least 6 months Time on ART at measurement of viral load and adherence: not reported; at least 6 months
ndex tests	 Number of index tests used: 3 Types of index tests: self-report, visual analog scale, drug plasma concentration Test 1. Self-report questionnaire Validated scale: not reported Tool description: the questionnaire consisted of three questions focussing on recent drug ir take: (i) "Did you miss any HAART doses during the last four days?," (ii) "Were you late for an of your intakes by more than two hours during the last four days?," and (iii) "Did you miss an HAART doses during the last four days?," and (iii) "Did you miss an HAART doses last week-end?" Blinding: no information Threshold prespecified: yes Adherence threshold used: 100% Test 2. A visual analog scale Validated scale: yes Tool description: patients were asked to answer the question: "In general, would you say yo take your treatment? Blinding: no information Threshold prespecified: "never" (score 1) and "always" (score 10). Any answer different from 10 was considered to represent nonadherence. Adherence threshold used: score 9, score 10 Test 3. Composite measure (self-report questionnaire + drug plasma concentration) Validated scale: not reported Tool description: the questionnaire consisted of three questions focussing on recent drug ir take: (i) "Did you miss any HAART doses during the last four days?," (ii) "Were you late for an of your intakes by more than two hours during the last four days?," (ii) "Were you late for an no formation

segeral 2010 (Continued)	• Adherence threshold	used: 100%			
Target condition and refer-	Target condition: viral non-suppression				
ence standard(s)	Reference standard: ANS	second-generation (G2) real-	time RT-PCR		
		ppression: HIV viral load > 400			
	Blinded to index test: no information				
Flow and timing	a cross-sectional study s	o likely to be measured simult	xplicit information on timing, but this wa aneously		
		ne reference standard: yes			
	 Missing data: patients who did not have adherence measurements were excluded from ana 13 patients (3.8%) died, 14 (4.1%) were lost to follow-up, 9 (2.6%) were directed to other cer 12 were on an LPV/r-containing regimen, 8 were on a triple NRTI combination, and 25 could be evaluated. 				
Comparative					
Notes	Conflicts of interest: none d	eclared			
	Funding source: ESTHER pr	ogramme			
	Trial registry: not reported				
Methodological quality					
Item	Authors' judgement	Risk of bias	Applicability concerns		
DOMAIN 1: Patient Selection					
Was a consecutive or random sample of patients enrolled?	Unclear	Unclear			
Was a case-control design avoided?	Yes	Yes			
Did the study avoid inappro- priate exclusions?	Unclear				
Could the selection of pa- tients have introduced bias?		Unclear risk			
Are there concerns that the included patients and set- ting do not match the review question?			High		
DOMAIN 2: Index Test (Index te	est)				
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Unclear				
If a threshold was used, was it pre-specified?	Unclear				

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Segeral 2010 (Continued)				
Could the conduct or inter- pretation of the index test have introduced bias?		Unclear risk		
Are there concerns that the index test, its conduct, or in- terpretation differ from the review question?			Low concern	
DOMAIN 3: Reference Standard				
Is the reference standards like- ly to correctly classify the tar- get condition?	Yes			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear			
Could the reference stan- dard, its conduct, or its inter- pretation have introduced bias?		Low risk		
Are there concerns that the target condition as defined by the reference standard does not match the ques- tion?			Low concern	
DOMAIN 4: Flow and Timing				
Was there an appropriate in- terval between index test and reference standard?	Yes			
Did all patients receive the same reference standard?	Yes			
Were all patients included in the analysis?	No			
Could the patient flow have introduced bias?		High risk		

Segeral 2018

 Study characteristics

 Patient Sampling

 • Target population: adults with HIV on PI-based second-line ART regimen for at least 6 months

 • Recruitment: likely consecutive; patients were exhaustively enrolled if they were HIV-infected adults

 • Inclusion criteria: HIV infection, > 18 years, current PI-based second-line ART treatment since at least 6 months and willing to participate and consent to signature



Patient characteristics and setting	 Exclusion criteria: ongoing PI-based second-line regimen for less than 6 months at time of study intake Study design: cross-sectional
Patient characteristics and setting	
	 Country: Cambodia World Bank Income classification: low-income Study setting: 13 representative ART sites (6 in Phnom Penh and 7 in provinces) Study dates: recruitment from February 2013 to April 2014 Age of population (years), median (IQR): 42 (37 to 48) Gender (male %): 61.8 Participants included/analysed: 1348/1317 First- or second-line regimen: second-line Type of ART: PI-based second-line regimen Time on ART at enrolment: at least 6 months Time on ART at measurement of viral load and adherence: not reported; at least 6 months
ndex tests	Number of index tests used: 1
	Types of index tests: self-report
	 Test 1. Self-report questionnaire Validated scale: yes (14-item validated scale) Tool description: patients provided information about the doses taken during the days preceding the survey, about whether they respected the dose schedule during the previous 4 days and 4 weeks, and whether treatment interruption had occurrent for at least two consecutive days within the previous 4 weeks. The algorithm proposed by (Carrieri 2001). was used to calculate the adherence score corresponding to the 4 days preceding the survey, by comparing the number of pills taken with those prescribed. Blinding: no information Threshold prespecified: yes Adherence threshold used: 100%; ≥ 80%
Farget condition and reference stan- dard(s)	 Target condition: viral non-suppression Reference standard: G2 Generic HIV-1 VL ANRS kit (Biocentric, Bandol, France) conducted at National Center for HIV/AIDS, Dermatology and STD (NCHADS) laboratory Definition of viral non-suppression: HIV viral load > 250 copies/mL Blinded to index test: no information
Flow and timing	 Time interval between index and reference tests: no explicit information on timing, but this was a cross-sectional study so likely to be measured simultaneously All patients received same reference standard: yes Missing data: none. All eligible participants with viral load test and adherence measures were included in the main analysis.
Comparative	
Notes	Conflicts of interest: none declared
	Funding source: French National Agency for Research on AIDS and Viral Hepatitis (ANRS) Trial registry: NCT01801618
Methodological quality	

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Segeral 2018 (Continued)

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclu- sions?	Yes		
Could the selection of patients have in- troduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Index test)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-speci- fied?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear
DOMAIN 3: Reference Standard			
Is the reference standards likely to cor- rectly classify the target condition?	Yes		
Were the reference standard results inter- preted without knowledge of the results of the index tests?	Unclear		
Could the reference standard, its con- duct, or its interpretation have intro- duced bias?		Low risk	
Are there concerns that the target con- dition as defined by the reference stan- dard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval be- tween index test and reference standard?	Yes		

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Segeral 2018 (Continued)

Did all patients receive the same refer-Yes ence standard?

Were all patients included in the analysis? Yes

Could the patient flow have introduced bias?

Low risk

Spire 2008

Study characteristics	
Patient Sampling	 Target population: HIV-infected adults receiving ART at a hospital Recruitment: not reported Inclusion criteria: patients who had been receiving ART for 24 (± 2 months) as part of HIV program run by Médecins sans Frontières, in collaboration wth the Ministry of Health of Cambodia, at the Infectious Disease department in Khmero-Sovietic Friendship hospital in Phnom Penh Exclusion criteria: not willing to participate Study design: cross-sectional
Patient characteristics and setting	 Country: Cambodia World Bank Income classification: low-income Study setting: hospital-based Study dates: December 2004 to December 2005 Age of population (years), median (IQR): 36 (32 to 40) Gender (male %): 57.5 Participants included/analysed: 346/346 First- or second-line regimen: first-line Type of ART: ARV treatment-naïve at ART initiation (95.4%). A total of 280 patients were receiving a first-line ART regimen associating d4T, 3TC (lamivudine) and EFV while 56 were receiving d4T, 3TC and NVP and the remaining 10 another first-line regimen Time on ART at enrolment: 24 ± 2 months Time on ART at measurement of viral load and adherence: 24 months
Index tests	 Number of index tests used: 1 Types of index tests: composite Test. Composite measure (self-report questionnaire + two VAS; range 1-6) Validated scale: yes Tool description: a face-to-face interview based on a standardized questionnaire translated into Khmer was administered by an external member of staff. This questionnaire included several questions about patient's adherence to ART in the 4 days or the 4 weeks prior to the interview. Five questions regarding adherence to HAART were included in all self-administered questionnaires according to the methodology established by the AIDS Clinical Trial Group. Adherence to ART was assessed using a dichotomous score already validated in previous studies. Patients were first asked to list for each drug included in their HAART regimen the number of pills taken all of their prescribed doses in the 4 days before the visit. In addition, two visual analog scales measuring adherence in general and in the last 4 weeks were completed. Patients who reported scores < 5 were reclassified as non-adherent. Blinding: no information Threshold prespecified; yes

Spire 2008 (Continued)	• Adherence threshold	l used: 100%		
Target condition and reference	nd reference Target condition: viral non-suppression			
standard(s)	 Reference standard: real-time PCR technology which allows quantification of HIV-1 non-B subtypes including those circulating in Asia performed at HIV/hepatitis laboratoty of Necker Enfants Malades Hospital, Paris Definition of viral non-suppression: HIV viral load > 40 copies/mL Blinded to index test: no information 			
Flow and timing	 Time interval between index and reference tests: both measures taken at 24 months after initiation of ART, but this was a cross-sectional study so likely to be measured simultaneously All patients received same reference standard: yes Missing data: none. All eligible participants with viral load test and adherence measures were included in the main analysis. 			
Comparative				
Notes	Conflicts of interest: not re	ported		
	Funding source: Médecins	sans Frontières and Sidact		
	Trial registry: not reported			
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sam- ple of patients enrolled?	Unclear			
Was a case-control design avoid- ed?	Yes			
Did the study avoid inappropriate exclusions?	Unclear			
Could the selection of patients have introduced bias?		Unclear risk		
Are there concerns that the in- cluded patients and setting do not match the review question?			High	
DOMAIN 2: Index Test (Index test)				
Were the index test results inter- preted without knowledge of the results of the reference standard?	Unclear			
If a threshold was used, was it pre- specified?	Yes			
Could the conduct or interpreta- tion of the index test have intro- duced bias?		Low risk		

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Spire 2008 (Continued)			
Are there concerns that the in- dex test, its conduct, or inter- pretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the tar- get condition as defined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Yes		
Could the patient flow have in- troduced bias?		Low risk	

Tabb 2018

Patient Sampling	 Target population: HIV-positive young people who attended a monthly youth-focused HI clinic
	Recruitment: not reported
	 Inclusion criteria: HIV-positive young people who knew their HIV status attending a youth-focussed HIV clinic called 'Teen Club' (at either Kilimanjaro Christian Medical Centre or Mawer zi Regional Referral Hospital in Moshi) to receive education on topics such as stigma, adherence, and sexual reproductive health
	Exclusion criteria: not reported
	Study design: cross-sectional
Patient characteristics and setting	Country: Tanzania

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

 Study Age of Gende Partic First-o Types of Test 1 Val Too Test 1 Val Too Too Study Age of Gende Partic First-o Types of Test 1 Val Too Spectration Test 1 Val Too Spectration Test 1 Val Too Spectration Test 1 Val Too Test 1 Val Too Spectration Test 1 Val Too Spectration Test 1 Val Too Spectration Test 1 Val Too <	
 Study Age of Genda Partic First-o Types of Target condition and reference standard(s) Target c	rld Bank Income classification: low-income
 Age of Genda Partic First-o or Type one or Tin or Tin or Tin or Tin or Tin or Tin or Tin or Tin or Too spe by we oft or Test 1. or Val or Too spe by we oft or The or The	dy setting: youth-focussed HIV clinic
 Gende Partic First-o Types of Test 1 Val Too Test 1 Val Too Spe by we oft Test 1 val Too Standard(s) Target condition and reference Refere Defini Blinde Flow and timing Flow and timing Time i was a All pat Missin includ Comparative Notes Conflicts Funding an NIH-fu Fogarty 1 lows Progrand the N Scholars	dates: December 2013-December 2014 (Kilimanjaro); February to July 2015 (Moshi)
 Partic First-G Types of i Test 1 Val Too Test 1 Val Too Too Second Test 1 Val Too Too Second Test 1 Val Too Test 1 Val Too Test 1 Val Too Too Second Test 1 Val Too Second Test 1 Val Too Too Second Too Second Test 1 Val Too Second Test 1 Test 1 Test 1 Val Too Test 1 Val Too Test 1 Val Too Second Test 1 Too Second Test 1 Too Second Test 1 Too <li< th=""><th>population (years), median (IQR): 16 (14 to 18)</th></li<>	population (years), median (IQR): 16 (14 to 18)
 First-o Types of i Test 1 Val To val To val To val To or spective Blin Thi Adl Target condition and reference standard(s) Flow and timing Time i was a All pat Missin includ Comparative Notes Conflicts Funding an NIH-fu Foundary is program in the Nicholars	er (male %): almost 50%
 or Tyronover and the second sec	ipants included/analysed: 227/227
 Tindex tests Number of Types of Types of Types of Tools of the second seco	or second-line regimen: both first- and second-line regimens be of ART: first-line regimens included two NRTI and a NNRTI of either NVP or EFV; sec- d-line regimens included two NRTIs and a RTV-boosted PI of either LPV or ATV ne on ART at enrolment: at least 6 months
Types of i • Test 1. • Val • Too • page by wee oft mode na eiti • Blin • The • Adl Target condition and reference standard(s) • Reference • Defini • Blindee Flow and timing • Time i was a • All pate • Missin includ Comparative Notes Conflicts Funding s an NIH-ft Fogarty It lows Progrand the N Scholars	ne on ART at measurement of viral load and adherence: not reported; at least 6 months
 Test 1. Val Too spend by we oft model of the spectrum o	of index tests used: 1
 Val Too spe by we oft Too spe Too Too Too Selini Thin Adi Target condition and reference standard(s) Refere Defini Blinde Flow and timing Time i was a All pate Missin includ Comparative Notes Conflicts Funding : an NIH-ft Fogarty It lows Proparity It 	index tests: self-report
spe by we oft oft model na eitl o Blin o Adl Target condition and reference Target condition standard(s) Reference Flow and timing Time i Flow and timing Time i was a All pather Missin include Comparative Conflicts Notes Conflicts Funding : an NIH-ft Fogarty II Iows Progrand the N Scholars Scholars	Self-report questionnaire idated scale: not reported
 The second standard (s) Target condition and reference standard (s) Reference Definition Blinder Flow and timing Time is was a an All pather of the second standard of the second standard	ol description: a structured questionnaire was administered by trained, native Swahili- eaking, female research assistants which included queries on self-reported adherence asking dichotomously, "Have you missed any doses of your medication in the last two eks, yes or no?" and categorically, "Think about the past week (7 days); on average, how en did you miss a dose of medication?" Response options included, "(1) once a day; (2) ore than once a week, but not every day; (3) once a week; or (4) I don't miss my medicine." dequate adherence by self-report was defined as reporting any missed ART doses on her of the survey items.
 Add Target condition and reference standard(s) Refere Defini Blinde Flow and timing Time i was a All pate Missin includ Comparative Notes Conflicts Funding : an NIH-ft Fogarty It lows Progrand the Nischolars 	nding: no information
Target condition and reference standard(s) Target consistence standard(s) • Reference Definition • Definition • Blindee • Time is was a • Flow and timing • Time is was a • All path • Missinin include Comparative • Conflicts Notes Conflicts Funding so an NIH-fu Fogarty It lows Progrand the N Scholars	reshold prespecified: not reported
standard(s) Refere Defini Blinde Flow and timing Flow and timing Comparative Notes Conflicts Funding an NIH-fu Fogarty I lows Prog and the N Scholars	herence threshold used: 100%
Refere Defini Blinde Flow and timing Time i was a All pat Missin includ Comparative Notes Conflicts Funding a an NIH-fu Fogarty II lows Prog and the N Scholars	ndition: viral non-suppression
Blinde Flow and timing Time i was a All pat Missin includ Comparative Notes Conflicts Funding s an NIH-ft Fogarty II lows Prog and the N Scholars	ence standard: HIV-1 real-time PCR; the Abbott m2000, Abbott laboratories, Illinois, USA
Flow and timing	tion of viral non-suppression: HIV viral load > 400 copies/mL
was a All pat Missin includ Comparative Notes Conflicts Funding s an NIH-fu Fogarty In lows Prog and the N Scholars	ed to index test: no information
Missin includ Comparative Notes Conflicts Funding an NIH-ft Fogarty It lows Prog and the N Scholars	nterval between index and reference tests: no explicit information on timing, but this cross-sectional study so likely to be measured simultaneously
Comparative Notes Conflicts Funding s an NIH-fu Fogarty I lows Prog and the N Scholars	ients received same reference standard: yes
Notes Conflicts Funding : an NIH-fu Fogarty I lows Prog and the N Scholars	g data: none. All eligible participants with viral load test and adherence measures were ed in the main analysis.
Funding s an NIH-fu Fogarty I Iows Prog and the N Scholars	
an NIH-fu Fogarty I lows Prog and the N Scholars	of interest: none declared
	source: the current work was supported by Duke University Center for AIDS Research, inded program; International Research Scientist Development Award funded by the nternational Center and the National Institute of Mental Health; the Global Health Fel- gram of the National Institutes of Health funded by the Fogarty International Center lational Institute of Mental Health; the Infectious Diseases Society of America Medical Program. The National Institute of Allergy and Infectious Diseases at the National In- f Health funded analyses of hair samples.
Trial regi	stry: not reported
Methodological quality	- · ·

Methodological quality

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



abb 2018 (Continued)			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sam- ple of patients enrolled?	Unclear		
Was a case-control design avoid- ed?	Yes		
Did the study avoid inappropriate exclusions?	Unclear		
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the in- cluded patients and setting do not match the review question?			Unclear
DOMAIN 2: Index Test (Index test)			
Were the index test results inter- preted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre- specified?	Unclear		
Could the conduct or interpreta- tion of the index test have intro- duced bias?		Unclear risk	
Are there concerns that the in- dex test, its conduct, or inter- pretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condi- tion?	Yes		
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Unclear		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the tar- get condition as defined by the reference standard does not match the question?			Low concern

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Tabb 2018 (Continued)

DOMAIN 4: Flow and Timing	
Was there an appropriate interval between index test and reference standard?	Yes
Did all patients receive the same reference standard?	Yes
Were all patients included in the analysis?	No
Could the patient flow have in- troduced bias?	High risk

Zoufaly 2013

Study characteristics	
Patient Sampling	 Target population: HIV-1 infected children on ART at the HIV service Recruitment: all children who attended for routine follow-up and drug refill from September 2010 to August 2011 Inclusion criteria: HIV-1-infected paediatric patients on ART attending Bamenda Regional Hospital with informed consent obtained by parents or legal guardians Exclusion criteria: no plasma was available, > 18 years at enrolment, caregivers or children did not consent for participation in the study Study design: cross-sectional
Patient characteristics and setting	 Country: Cameroon World Bank Income classification: lower-middle-income Study setting: hospital-based Study dates: September 2010 to August 2011 Age of population (years), median (IQR): 8.8 (6.1 to 11.4) Gender (male %): 52.8 Participants included/analysed: 230/174 First- or second-line regimen: unclear Type of ART: ART containing two NVP in all children < 2 years irrespective of CD4+ T-cell count and thereafter according to absolute CD4+ T-cell count level. LPV/r was used when children were recently exposed to NVP in prevention of mother-to-child transmission regimens or in second-line regimens. EFV was used in case of non-tolerance of NVP. Time on ART at enrolment: median 3.4 years Time on ART at measurement of viral load and adherence: not reported; at least 6 months
Index tests	 Number of index tests used: 1 Types of index tests: self-report Test 1. Self-report number of daily doses taken Validated scale: not applicable Tool description: adherence reported by the child or caregiver was categorized according to the number of full daily doses taken in the previous 28 days, and recorded by a trained study nurse. Blinding: no information

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Zoufaly 2013 (Continued)	 Threshold prespecified: not reported Adherence threshold used: 95%; 75% 			
Target condition and reference standard(s)	Target condition: viral non-suppressionReference standard: HIV-1 real-time PCR; Abbott laboratories, Illinois, USA			
	 Definition of viral non-suppression: HIV viral load > 200 copies/mL Blinded to index test: yes 			
Flow and timing	 Time interval between index and reference tests: no explicit information on timing, but this was a cross-sectional study so likely to be measured simultaneously All patients received same reference standard: yes Missing data: none. All eligible participants with viral load test and adherence measures were included in the main analysis. 			
Comparative				
Notes	Conflicts of interest: none	e declared		
	Funding source: partly fu	nded by ESTHER Germany	1	
	Trial registry: not reported			
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Yes			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate exclu- sions?	Yes			
Could the selection of patients have in- troduced bias?		Low risk		
Are there concerns that the included pa- tients and setting do not match the re- view question?			Low concern	
DOMAIN 2: Index Test (Index test)				
Were the index test results interpreted without knowledge of the results of the ref- erence standard?	Unclear			
If a threshold was used, was it pre-speci- fied?	Unclear			
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk		



Zoufaly 2013 (Continued)

Trusted evidence. Informed decisions. Better health.

Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correct- ly classify the target condition?	Yes		
Were the reference standard results inter- preted without knowledge of the results of the index tests?	Yes		
Could the reference standard, its con- duct, or its interpretation have intro- duced bias?		Low risk	
Are there concerns that the target con- dition as defined by the reference stan- dard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Yes		
Could the patient flow have introduced bias?		Low risk	
BTC: lamivudine ACTG: AIDS Clinical Trials Group AFB: Acid Fast Bacilli AIDS: acquired immunodeficiency syndrome ANRS: National Agency for AIDS Research			

ART: antiretroviral therapy ARV: antiretroviral ATV: atazanavir AZT: zidovudine CD4: cluster of differentiation 4 CDC: centres for disease control and prevention CTAC: community training and assistance centre d4T: stavudine ddl: didanosine DNA: deoxyribonucleic acid DRV/r: darunavir/ritonavir EAMD: electronic adherence monitoring device EFV: efavirenz ELISA: enzyme-linked immunosorbent assay FDC: fixed drug combination FTC: emtricitabine G2: second generation GEEMA: Grupo Español para el Estudio Multifactorial de la Adherencia GPO: Government Pharmaceutical Organization

HAART: highly active antiretroviral therapy HIV: Human immunodeficiency virus HNRP:HIV Neurobehavioral Research Program iDART: intelligent dispensing of ART IQR: interquartile range INSTI: integrase strand transfer inhibitors ITT: intention-to-treat LPV: lopinavir LPV/r: ritonavir-boosted lopinavir MCH-ART: Maternal Child Health ART MEMS: medication event monitoring system MMAS(-4): Morisky medication adherence scale 4 M-MASRI: modified medication adherence self-report inventory MPR: medication possession ratio NA: not applicable NCT: National Clinical Trial NHLS: national health laboratory services NIAID: national institute of allergy and infectious diseases NVP: nevirapine NNRTI: nonnucleoside reverse transcriptase NRTI: nucleoside reverse transcriptase inhibitors NTI: non-structured treatment interruption NtRTI: nucleoside reverse transcriptase inhibitor PACTG: pediatric AIDS clinical trials group PCR: Polymerase chain reaction PCZCDC: Prince Cyril Zulu Communicable Diseases Centre PEP: post-exposure prophylaxis PI: protease inhibitor PIT: pills identification test PMTCT: prevention of mother to child transmissions RAM:resistance-associated mutations RCT: randomised control trial REACH: Reversing the Epidemic in Africa with Choices in HIV Prevention RNA: ribonucleic acid **RPV: ripivirine RTV: ritonavir** Rx: treatment SHCS-AQ: Swiss HIV cohort study adherence questionnaire SMAQ: simplified medication adherence questionnaire STD: standard **TB: tuberculosis** TDF: tenofovir disoproxil fumarate TRuTH: TB recurrence upon treatment with HAART ULN: upper limit of normal VAS: visual analogue scale VL: viral load WHO: World Health Organization ZDV: zidovudine

DATA

Presented below are all the data for all of the tests entered into the review.

Table Tests. Data tables by test

Test	No. of studies	No. of participants
1 [Main analysis] Self-report, various thresholds*	25	9211

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Test	No. of studies	No. of participants
2 [Subgroup analysis by adherence threshold] Self-report questionnaires; threshold: ≥ 100% adherence	21	8204
3 [Subgroup analysis by adherence threshold] Self-report questionnaires; threshold: ≥ 95% adherence	4	1007
4 [Supplementary analysis] Self-report questionnaires; threshold: ≥ 80% ad- herence	3	1527
5 [Subgroup analysis by population] Self-report questionnaires; various adher- ence thresholds*; children	4	804
6 [Subgroup analysis by population] Self-report questionnaires; various adher- ence thresholds*; adults	19	8011
7 [Subgroup analysis by setting] Self-report questionnaires; various adherence thresholds*; low-income	11	4135
8 [Subgroup analysis by setting] Self-report questionnaires; various adherence thresholds*; lower-middle-income	3	576
9 [Subgroup analysis by setting] Self-report questionnaires; various adherence thresholds*; upper-middle-income	5	1441
10 [Subgroup analysis by setting] Self-report questionnaires; various adher- ence thresholds*; high-income	5	2702
11 [Subgroup analysis by viral load] Self-report questionnaires; various adher- ence thresholds*; 40 to 50 copies/mL	11	2290
12 [Subgroup analysis by viral load] Self-report questionnaires; various adher- ence thresholds*; 200 to 400 copies/mL	13	6664
13 [Subgroup analysis by number of questions] Self-report questionnaires; various adherence thresholds*; 1-item	12	4997
14 [Subgroup analysis by number of questions] Self-report questionnaires; various adherence thresholds*; 2 to 4 items	8	1922
15 [Subgroup analysis by number of questions] Self-report questionnaires; various adherence thresholds*; 5 or more items	5	2292
16 [Main analysis] Self-report VAS; threshold: ≥ 95% adherence	11	4235
17 [Supplementary analysis] Self-report VAS; threshold: ≥ 90% adherence	3	582
18 [Supplementary analysis] Self-report VAS; threshold: ≥ 80% adherence	1	73
19 [Subgroup analysis by viral load] Self-report VAS; threshold: ≥ 95% adher- ence; 40 to 100 copies/mL	6	3591
20 [Subgroup analysis by viral load] Self-report VAS; threshold: ≥ 95% adher- ence; 200 to 400 copies/mL	5	644
21 [Subgroup analysis by population] Self-report VAS; threshold: ≥ 95% adher- ence; children	2	239



Test	No. of studies	No. of participants
22 [Subgroup analysis by population] Self-report VAS; threshold: ≥ 95% adher- ence; adults	8	3904
23 [Subgroup analysis by setting] Self-report VAS; threshold: ≥ 95% adherence; low-income	5	663
24 [Subgroup analysis by setting] Self-report VAS; threshold: ≥ 95% adherence; lower-middle-income	3	1631
25 [Main analysis] Tablet count; threshold: ≥ 95% adherence	12	3466
26 [Supplementary analysis] Tablet count; threshold: ≥ 80% adherence	2	235
27 [Subgroup analysis by viral load] Tablet count; threshold: ≥ 95% adherence; VL 40 to 80 copies/mL	7	2299
28 [Subgroup analysis by viral load] Tablet count; threshold: ≥ 95% adherence; VL 400 copies/mL	5	1167
29 [Subgroup analysis by population] Tablet counts; threshold: ≥ 95% adher- ence; children	1	73
30 [Subgroup analysis by population] Tablet counts; threshold: ≥ 95% adher- ence; adults	9	3016
31 [Subgroup analysis by population] Tablet counts; threshold: ≥ 95% adher- ence; mixed	2	377
32 [Subgroup analysis by setting] Tablet counts; threshold: ≥ 95% adherence; low-income	4	942
33 [Subgroup analysis by setting] Tablet counts; threshold: ≥ 95% adherence; lower-middle-income	4	1692
34 [Subgroup analysis by setting] Tablet counts; threshold: ≥ 95% adherence; upper-middle-income	3	610
35 [Subgroup analysis by setting] Tablet counts; threshold: ≥ 95% adherence; high-income	1	222
36 [Main analysis] Pharmacy records; threshold: ≥ 95% adherence	6	2254
37 [Supplementary analysis] Pharmacy records; threshold: ≥ 80% adherence	3	1211
38 [Subgroup analysis by viral load] Pharmacy records; threshold: ≥ 95% ad- herence; VL: 40 copies/mL	1	178
39 [Subgroup analysis by viral load] Pharmacy records; threshold: ≥ 95% ad- herence; VL: 200 to 400 copies/mL	5	2076
40 [Subgroup analysis by population] Pharmacy records; threshold: ≥ 95% ad- herence; mixed	2	402
41 [Subgroup analysis by population] Pharmacy records; threshold: ≥ 95% ad- herence; adults	4	1893



Test	No. of studies	No. of participants
42 [Subgroup analysis by setting] Pharmacy records; threshold: ≥ 95% adher- ence; low-income	4	1485
43 [Subgroup analysis by setting] Pharmacy records; threshold: ≥ 95% adher- ence; lower-middle-income	1	591
44 [Subgroup analysis by setting] Pharmacy records; threshold: ≥ 95% adher- ence; upper-middle-income	1	178
45 [Main analysis] Electronic monitoring; threshold: ≥ 95% adherence	3	186
46 [Supplementary analysis] Electronic monitoring; threshold: ≥ 80% adher- ence	4	327
47 [Subgroup analysis by viral load] Electronic monitoring; threshold: ≥ 95% adherence; VL: 50 copies/mL	1	72
48 [Subgroup analysis by viral load] Electronic monitoring; threshold: ≥ 95% adherence; VL: 400 copies/mL	2	114
49 [Subgroup analysis by population] Electronic monitoring; threshold: ≥ 95% adherence; children	1	72
50 [Subgroup analysis by population] Electronic monitoring; threshold: ≥ 95% adherence; adults	2	114
51 [Subgroup analysis by setting] Electronic monitoring; threshold: ≥ 95% ad- herence; low-income	1	72
52 [Subgroup analysis by setting] Electronic monitoring; threshold: ≥ 95% ad- herence; lower-middle-income	1	65
53 [Subgroup analysis by setting] Electronic monitoring; threshold: ≥ 95% ad- herence; upper-middle-income	1	49
54 [Main analysis] Composite measure; different thresholds*	9	1513
55 [Subgroup analysis by adherence threshold] Composite measures; 100% adherence	6	1095
56 [Subgroup analysis by adherence threshold] Composite measures; 95% ad- herence	3	418
57 [Subgroup analysis by viral load] Composite measure; various adherence thresholds*; 40 to 50 copies/mL	3	522
58 [Subgroup analysis by viral load] Composite measure; various adherence thresholds*; 200 to 400 copies/mL	7	1063
59 [Subgroup analysis by population] Composite measure; various adherence thresholds*; children	1	104
60 [Subgroup analysis by population] Composite measure; various adherence thresholds*; adults	7	1390



Test	No. of studies	No. of participants
61 [Subgroup analysis by setting] Composite measure; various adherence thresholds*; low-income	4	881
62 [Subgroup analysis by setting] Composite measure; various adherence thresholds*; upper-middle-income	2	405
63 [Subgroup analysis by setting] Composite measure; various adherence thresholds*; high-income	3	227

Test 1. [Main analysis] Self-report, various thresholds*

[Main analysis] Self-report, various thresholds*

Study	ТР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI) Sensitivity (95% CI)Specif	ficity (95% CI)
Avong 2015	84	185	15	218	0.85 [0.76, 0.91]	0.54 [0.49, 0.59]	+
Bajunirwe 2009	10	16	18	131	0.36 [0.19, 0.56]	0.89 [0.83, 0.94]	-
Coker 2015	45	2	44	224	0.51 [0.40, 0.61]	0.99 [0.97, 1.00]	•
Duarte 2015	19	10	104	224	0.15 [0.10, 0.23]	0.96 [0.92, 0.98] 🛛 🛨	-
Ekstrand 2010	10	11	45	136	0.18 [0.09, 0.31]	0.93 [0.87, 0.96] 🚽 🖛	-
El-Khatib 2010	10	32	92	740	0.10 [0.05, 0.17]	0.96 [0.94, 0.97] 🛛 🛨	
Fokam 2017	28	64	14	38	0.67 [0.50, 0.80]	0.37 [0.28, 0.47]	
Haberer 2011	13	29	- 7	24	0.65 [0.41, 0.85]	0.45 [0.32, 0.60]	
Lan de s 2021	16	321	67	37	0.19 [0.11, 0.29]	0.10 [0.07, 0.14] 🗕 💻	
Mbengue 2019	13	24	36	90	0.27 [0.15, 0.41]	0.79 [0.70, 0.86] —	
McMahon 2013	25	105	9	31	0.74 [0.56, 0.87]	0.23 [0.16, 0.31]	-
Meya 2009	10	52	39	395	0.20 [0.10, 0.34]	0.88 [0.85, 0.91]	-
Mogosetsi 2018	1	12	7	78	0.13 [0.00, 0.53]	0.87 [0.78, 0.93] —	-
Oette 2006	13	11	48	136	0.21 [0.12, 0.34]	0.93 [0.87, 0.96]	-
Orrell 2017	1	143	5	20	0.17 [0.00, 0.64]	0.12 [0.08, 0.18]	
Paolillo 2017	46	33	192	264	0.19 [0.15, 0.25]	0.89 [0.85, 0.92] 🛛 🛨	-
Parker 2017	106	458	68	1017	0.61 [0.53, 0.68]	0.69 [0.67, 0.71]	
Pasquau 2018	6	51	7	125	0.46 [0.19, 0.75]	0.71 [0.64, 0.78]	-
Phillips 2019	34	26	15	62	0.69 [0.55, 0.82]	0.70 [0.60, 0.80]	
Pulido 2009	9	27	6	79	0.60 [0.32, 0.84]	0.75 [0.65, 0.82]	-
San ged a 2014	7	14	48	93	0.13 [0.05, 0.24]	0.87 [0.79, 0.93]	-
Segeral 2010	11	23	63	160	0.15 [0.08, 0.25]	0.87 [0.82, 0.92]	-
Segeral 2018	28	192	107	990	0.21 [0.14, 0.29]	0.84 [0.82, 0.86]	
Tabb 2018	31	41	60	95	0.34 [0.24, 0.45]	0.70 [0.61, 0.77]	
Zoufaly 2013	27	23	94	86	0.22 [0.15, 0.31]	0.79 [0.70, 0.86] 🚬 👎	
-						0 0.2 0.4 0.6 0.8 1 0 0.2	0.40.60.81

Test 2. [Subgroup analysis by adherence threshold] Self-report questionnaires; threshold: ≥ 100% adherence

[Subgroup analysis by adherence threshold] Self-report questionnaires; threshold: ≥ 100% adherence

Study	ТР	FP	FN	TN	Sensitivity (95% Cl)	Specificity (95% Cl)	Sensitivity (95% CI)Specificity (95% CI)
Bajunirwe 2009	10	16	18	131	0.36 [0.19, 0.56]	0.89 [0.83, 0.94]	
Coker 2015	45	2	44	224	0.51 [0.40, 0.61]	0.99 [0.97, 1.00]	
Duarte 2015	19	10	104	224	0.15 [0.10, 0.23]	0.96 [0.92, 0.98]	-
El-Khatib 2010	10	32	92	740	0.10 [0.05, 0.17]	0.96 [0.94, 0.97]	- ·
Fokam 2017	28	64	14	38	0.67 [0.50, 0.80]	0.37 [0.28, 0.47]	
Lan de s 2021	16	321	67	37	0.19 [0.11, 0.29]	0.10 [0.07, 0.14]	•
Mbengue 2019	13	24	36	90	0.27 [0.15, 0.41]	0.79 [0.70, 0.86]	
McMahon 2013	25	105	9	31	0.74 [0.56, 0.87]	0.23 [0.16, 0.31]	- + +
Meya 2009	10	52	39	395	0.20 [0.10, 0.34]	0.88 [0.85, 0.91]	
Mogosetsi 2018	1	12	- 7	78	0.13 [0.00, 0.53]	0.87 [0.78, 0.93]	
Oette 2006	13	11	48	136	0.21 [0.12, 0.34]	0.93 [0.87, 0.96]	
Orrell 2017	1	143	5	20	0.17 [0.00, 0.64]	0.12 [0.08, 0.18]	
Paolillo 2017	46	33	192	264	0.19 [0.15, 0.25]	0.89 [0.85, 0.92]	• •
Parker 2017	106	458	68	1017	0.61 [0.53, 0.68]	0.69 [0.67, 0.71]	
Pasquau 2018	6	51	7	125	0.46 [0.19, 0.75]	0.71 [0.64, 0.78]	- - +
Phillips 2019	34	26	15	62	0.69 [0.55, 0.82]	0.70 [0.60, 0.80]	
Puli do 2009	9	27	6	79	0.60 [0.32, 0.84]	0.75 [0.65, 0.82]	_
San ged a 2014	7	14	48	93	0.13 [0.05, 0.24]	0.87 [0.79, 0.93]	+ +
Segeral 2010	11	23	63	160	0.15 [0.08, 0.25]	0.87 [0.82, 0.92]	
Segeral 2018	28	192	107	990	0.21 [0.14, 0.29]	0.84 [0.82, 0.86]	
Ta bb 2018	31	41	60	95	0.34 [0.24, 0.45]	0.70 [0.61, 0.77]	
							0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

Test 3. [Subgroup analysis by adherence threshold] Self-report questionnaires; threshold: ≥ 95% adherence

[Subgroup analysis by adherence threshold] Self-report questionnaires; threshold: > 95% adherence

Study	ΤР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)
Avong 2015	84	185	15	218	0.85 [0.76, 0.91]	0.54 [0.49, 0.59]
Ekstrand 2010	10	11	45	136	0.18 [0.09, 0.31]	0.93 [0.87, 0.96] 🗕 🗕
Haberer 2011	13	29	- 7	24	0.65 [0.41, 0.85]	0.45 [0.32, 0.60]
Zoufaly 2013	27	23	94	86	0.22 [0.15, 0.31]	

Test 4. [Supplementary analysis] Self-report questionnaires; threshold: ≥ 80% adherence

[Supplementary analysis] Self-report questionnaires; threshold: ≥ 80% adherence

Study	ΤР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% Cl)	Sensitivity (95% CI)Specificity (95% CI)
Haberer 2011	4	10	16	43	0.20 [0.06, 0.44]	0.81 [0.68, 0.91]	- -
Phillips 2019	20	3	29	85		0.97 [0.90, 0.99]	
Segeral 2018	11	31	124	1151	0.08 [0.04, 0.14]	0.97 [0.96, 0.98]	

Test 5. [Subgroup analysis by population] Self-report questionnaires; various adherence thresholds*; children

[Subgroup analysis by population] Self-report questionnaires; various adherence thresholds*; children

Study	тр	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)
Duarte 2015	19	10	104	224	0.15 [0.10, 0.23]	0.96 [0.92, 0.98] 📲
Fokam 2017	28	64	14	38	0.67 [0.50, 0.80]	0.37 [0.28, 0.47]
Haberer 2011	13	29	7	24	0.65 [0.41, 0.85]	0.45 [0.32, 0.60]
Zoufaly 2013	27	23	94	86	0.22 [0.15, 0.31]	



Test 6. [Subgroup analysis by population] Self-report questionnaires; various adherence thresholds*; adults

[Subgroup analysis by population] Self-report questionnaires; various adherence thresholds*; adults

Study	ТР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
Avong 2015	84	185	15	218	0.85 [0.76, 0.91]	0.54 [0.49, 0.59]	
Bajunirwe 2009	10	16	18	131	0.36 [0.19, 0.56]	0.89 [0.83, 0.94]	
C oker 2015	45	2	44	224	0.51 [0.40, 0.61]	0.99 [0.97, 1.00]	
Ekstrand 2010	10	11	45	136	0.18 [0.09, 0.31]	0.93 [0.87, 0.96]	
El-Khatib 2010	10	32	92	740	0.10 [0.05, 0.17]	0.96 [0.94, 0.97]	+ •
Lan de s 2021	16	321	67	37	0.19 [0.11, 0.29]	0.10 [0.07, 0.14]	- ·
Mbengue 2019	13	24	36	90	0.27 [0.15, 0.41]	0.79 [0.70, 0.86]	
McMahon 2013	25	105	9	31	0.74 [0.56, 0.87]	0.23 [0.16, 0.31]	- - - +
Meya 2009	10	52	39	395	0.20 [0.10, 0.34]	0.88 [0.85, 0.91]	
Mogosetsi 2018	1	12	- 7	78	0.13 [0.00, 0.53]	0.87 [0.78, 0.93]	
Oette 2006	13	11	48	136	0.21 [0.12, 0.34]	0.93 [0.87, 0.96]	
Paolillo 2017	46	33	192	264	0.19 [0.15, 0.25]	0.89 [0.85, 0.92]	• •
Parker 2017	106	458	68	1017	0.61 [0.53, 0.68]	0.69 [0.67, 0.71]	- · ·
Pasquau 2018	6	51	- 7	125	0.46 [0.19, 0.75]	0.71 [0.64, 0.78]	
Phillips 2019	34	26	15	62	0.69 [0.55, 0.82]	0.70 [0.60, 0.80]	
Pulido 2009	9	27	6	79	0.60 [0.32, 0.84]	0.75 [0.65, 0.82]	
San ged a 2014	7	14	48	93	0.13 [0.05, 0.24]	0.87 [0.79, 0.93]	
Segeral 2010	11	23	63	160	0.15 [0.08, 0.25]	0.87 [0.82, 0.92]	
Segeral 2018	28	192	107	990	0.21 [0.14, 0.29]	0.84 [0.82, 0.86]	0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

Test 7. [Subgroup analysis by setting] Self-report questionnaires; various adherence thresholds*; low-income

[Subgroup analysis by setting] Self-report questionnaires; various adherence thresholds*; low-income

Study	ΤР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
Avong 2015	84	185	15	218	0.85 [0.76, 0.91]	0.54 [0.49, 0.59]	
Bajunirwe 2009	10	16	18	131	0.36 [0.19, 0.56]	0.89 [0.83, 0.94]	
Coker 2015	45	2	44	224	0.51 [0.40, 0.61]	0.99 [0.97, 1.00]	
Haberer 2011	13	29	- 7	24	0.65 [0.41, 0.85]	0.45 [0.32, 0.60]	
Lan de s 2021	16	321	67	37	0.19 [0.11, 0.29]	0.10 [0.07, 0.14]	+
McMahon 2013	25	105	9	31	0.74 [0.56, 0.87]	0.23 [0.16, 0.31]	
Meya 2009	10	52	39	395	0.20 [0.10, 0.34]	0.88 [0.85, 0.91]	
Sangeda 2014	- 7	14	48	93	0.13 [0.05, 0.24]	0.87 [0.79, 0.93]	
Segeral 2010	11	23	63	160	0.15 [0.08, 0.25]	0.87 [0.82, 0.92]	
Segeral 2018	28	192	107	990	0.21 [0.14, 0.29]	0.84 [0.82, 0.86]	
Tabb 2018	31	41	60	95	0.34 [0.24, 0.45]	0.70 [0.61, 0.77]	

Test 8. [Subgroup analysis by setting] Self-report questionnaires; various adherence thresholds*; lower-middle-income

[Subgroup analysis by setting] Self-report questionnaires; various adherence thresholds*; lower-middle-inco

Study	тр	FP	FN	ΤN	Sensitivity (95% Cl)	Specificity (95% CI) Sen	sitivity (95% Cl)	Specificity (95% Cl)
Ekstrand 2010	10	11	45	136	0.18 [0.09, 0.31]	0.93 [0.87, 0.96] 🖃	-	-
F o kam 2017	28	64	14	38	0.67 [0.50, 0.80]	0.37 [0.28, 0.47]		
Zoufaly 2013	27	23	94	86	0.22 [0.15, 0.31]	0.79 [0.70, 0.86]	2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Test 9. [Subgroup analysis by setting] Self-report questionnaires; various adherence thresholds*; upper-middle-income

[Subgroup analysis by setting] Self-report questionnaires; various adherence thresholds*; upper-middle-incom

Study	ΤР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI) S	ensitivity (95% CI)Specificity (95% CI))
El-Khatib 2010	10	32	92	740	0.10 [0.05, 0.17]	0.96 [0.94, 0.97]	÷ •	
Mbengue 2019	13	24	36	90	0.27 [0.15, 0.41]	0.79 [0.70, 0.86]		
Mogosetsi 2018	1	12	- 7	78	0.13 [0.00, 0.53]	0.87 [0.78, 0.93] -	• •	
Orrell 2017	1	143	5	20	0.17 [0.00, 0.64]	0.12[0.08, 0.18] -		
Phillips 2019	34	26	15	62	0.69 [0.55, 0.82]	0.70 (0.60, 0.80) _H		

Test 10. [Subgroup analysis by setting] Self-report questionnaires; various adherence thresholds*; high-income

[Subgroup analysis by setting] Self-report questionnaires; various adherence thresholds*; high-income

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)S	pecificity (95% CI)
0ette 2006	13	11	48	136	0.21 [0.12, 0.34]	0.93 [0.87, 0.96]		-
Paolillo 2017	46	33	192	264	0.19 [0.15, 0.25]	0.89 [0.85, 0.92]	+	-
Parker 2017	106	458	68	1017	0.61 [0.53, 0.68]	0.69 [0.67, 0.71]		•
Pasquau 2018	6	51	7	125	0.46 [0.19, 0.75]	0.71 [0.64, 0.78]		-
Pulido 2009	9	27	6	79	0.60 [0.32, 0.84]	0.75 [0.65, 0.82]		0.2 0.4 0.6 0.8 1

Test 11. [Subgroup analysis by viral load] Self-report questionnaires; various adherence thresholds*; 40 to 50 copies/mL

[Subgroup analysis by viral load] Self-report questionnaires; various adherence thresholds*; 40 to 50 copies/mL

Study	ТР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% Cl)	Sensitivity (95%	CI)Specificity (95% CI)
Bajunirwe 2009	10	16	18	131	0.36 [0.19, 0.56]	0.89 [0.83, 0.94]		-
Fokam 2017	28	64	14	38	0.67 [0.50, 0.80]	0.37 [0.28, 0.47]		
Haberer 2011	13	29	7	24	0.65 [0.41, 0.85]	0.45 [0.32, 0.60]		
Lan de s 2021	16	321	67	37	0.19 [0.11, 0.29]	0.10 [0.07, 0.14]		
Mogosetsi 2018	1	12	7	78	0.13 [0.00, 0.53]	0.87 [0.78, 0.93]	-	
0ette 2006	13	11	48	136	0.21 [0.12, 0.34]	0.93 [0.87, 0.96]		-
Orrell 2017	1	143	5	20	0.17 [0.00, 0.64]	0.12 [0.08, 0.18]	-	+
Paolillo 2017	46	33	192	264	0.19 [0.15, 0.25]	0.89 [0.85, 0.92]	+	-
Pasquau 2018	6	51	7	125	0.46 [0.19, 0.75]	0.71 [0.64, 0.78]		
Phillips 2019	34	26	15	62	0.69 [0.55, 0.82]	0.70 [0.60, 0.80]		
Pulido 2009	9	27	6	79	0.60 [0.32, 0.84]	0.75 [0.65, 0.82]	0 0.2 0.4 0.6 0.8	1 0 0.2 0.4 0.6 0.8 1



Test 12. [Subgroup analysis by viral load] Self-report questionnaires; various adherence thresholds*; 200 to 400 copies/mL

[Subgroup analysis by viral load] Self-report questionnaires; various adherence thresholds*; 200 to 400 copies/ml

Study	ТР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)
Avong 2015	84	185	15	218	0.85 [0.76, 0.91]	0.54 [0.49, 0.59]
Coker 2015	45	2	44	224	0.51 [0.40, 0.61]	0.99 [0.97, 1.00]
Duarte 2015	19	10	104	224	0.15 [0.10, 0.23]	0.96 [0.92, 0.98] 📲
Ekstrand 2010	10	11	45	136	0.18 [0.09, 0.31]	0.93 [0.87, 0.96] 🗕 🗕
El-Khatib 2010	10	32	92	740	0.10 [0.05, 0.17]	0.96 [0.94, 0.97] 📲
Mbengue 2019	13	24	36	90	0.27 [0.15, 0.41]	0.79 [0.70, 0.86]
McMahon 2013	25	105	9	31	0.74 [0.56, 0.87]	0.23 [0.16, 0.31]
Meya 2009	10	52	39	395	0.20 [0.10, 0.34]	0.88 [0.85, 0.91]
Parker 2017	106	458	68	1017	0.61 [0.53, 0.68]	0.69 [0.67, 0.71]
San ged a 2014	7	14	48	93	0.13 [0.05, 0.24]	0.87 [0.79, 0.93]
Segeral 2018	28	192	107	990	0.21 [0.14, 0.29]	0.84 [0.82, 0.86] 💶
Tabb 2018	31	41	60	95	0.34 [0.24, 0.45]	0.70 [0.61, 0.77]
Zoufaly 2013	27	23	94	86	0.22 [0.15, 0.31]	0.79 [0.70, 0.86]

Test 13. [Subgroup analysis by number of questions] Selfreport questionnaires; various adherence thresholds*; 1-item

[Subgroup analysis by number of questions] Self-report questionnaires; various adherence thresholds*; 1-item

Study TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)	Study
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Bajunirwe 2009	10	16	18	131	0.36 [0.19, 0.56]	0.89 [0.83, 0.94]		-
Duarte 2015	19	10	104	224	0.15 [0.10, 0.23]	0.96 [0.92, 0.98]	-	-
Ekstrand 2010	10	11	45	136	0.18 [0.09, 0.31]	0.93 [0.87, 0.96]		-
El-Khatib 2010	10	32	92	740	0.10 [0.05, 0.17]	0.96 [0.94, 0.97]		
Fokam 2017	28	64	14	38	0.67 [0.50, 0.80]	0.37 [0.28, 0.47]		
Haberer 2011	13	29	7	24	0.65 [0.41, 0.85]	0.45 [0.32, 0.60]		
Lan de s 2021	16	321	67	37	0.19 [0.11, 0.29]	0.10 [0.07, 0.14]		+
McMahon 2013	25	105	9	31	0.74 [0.56, 0.87]	0.23 [0.16, 0.31]		
0ette 2006	13	11	48	136	0.21 [0.12, 0.34]	0.93 [0.87, 0.96]		-
Orrell 2017	1	143	5	20	0.17 [0.00, 0.64]	0.12 [0.08, 0.18]		.
Paolillo 2017	46	33	192	264	0.19 [0.15, 0.25]	0.89 [0.85, 0.92]	-	-
Parker 2017	106	458	68	1017	0.61 [0.53, 0.68]	0.69 [0.67, 0.71]		1 1 0 0.2 0.4 0.6 0.8 1

Test 14. [Subgroup analysis by number of questions] Selfreport questionnaires; various adherence thresholds*; 2 to 4 items

[Subgroup analysis by number of questions] Self-report questionnaires; various adherence thresholds*; 2 to 4

Study	тр	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% Cl)	Sensitivity (95% CI)Specificity (95% CI)
Coker 2015	45	2	44	224	0.51 [0.40, 0.61]	0.99 [0.97, 1.00]	
Meya 2009	10	52	39	395	0.20 [0.10, 0.34]	0.88 [0.85, 0.91]	
Mogosetsi 2018	1	12	- 7	78	0.13 [0.00, 0.53]	0.87 [0.78, 0.93]	
Phillips 2019	34	26	15	62	0.69 [0.55, 0.82]	0.70 [0.60, 0.80]	
San ged a 2014	- 7	14	48	93	0.13 [0.05, 0.24]	0.87 [0.79, 0.93]	+ +
Segeral 2010	11	23	63	160	0.15 [0.08, 0.25]	0.87 [0.82, 0.92]	
Tabb 2018	31	41	60	95	0.34 [0.24, 0.45]	0.70 [0.61, 0.77]	
Zoufaly 2013	27	23	94	86	0.22 [0.15, 0.31]	0.79 [0.70, 0.86]	0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

Test 15. [Subgroup analysis by number of questions] Self-report questionnaires; various adherence thresholds*; 5 or more items

[Subgroup analysis by number of questions] Self-report questionnaires; various adherence thresholds*; 5 or m

Study	ТР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% Cl)
Avong 2015	84	185	15	218	0.85 [0.76, 0.91]	0.54 [0.49, 0.59]	-	-
Mbengue 2019	13	24	36	90	0.27 [0.15, 0.41]	0.79 [0.70, 0.86]		
Pasquau 2018	6	51	- 7	125	0.46 [0.19, 0.75]	0.71 [0.64, 0.78]		-
Pulido 2009	9	27	6	79	0.60 [0.32, 0.84]	0.75 [0.65, 0.82]		
Segeral 2018	28	192	107	990	0.21 [0.14, 0.29]	0.84 [0.82, 0.86]	0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Test 16. [Main analysis] Self-report VAS; threshold: ≥ 95% adherence

[Main analysis] Self-report VAS; threshold: ≥ 95% adherence

Study	ΤР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% Cl)	Specificity (95% CI)
Cerutti 2016	52	558	63	691	0.45 [0.36, 0.55]	0.55 [0.53, 0.58]		•
C ohe n 2012	41	134	863	176	0.05 [0.03, 0.06]	0.57 [0.51, 0.62]	•	+
Dziva 2017	16	12	45	93	0.26 [0.16, 0.39]	0.89 [0.81, 0.94]		-
Ekstran d 2010	18	16	37	131	0.33 [0.21, 0.47]	0.89 [0.83, 0.94]		-
Gill 2010	0	2	8	55	0.00 [0.00, 0.37]	0.96 [0.88, 1.00]		-
Haberer 2011	6	21	14	32	0.30 [0.12, 0.54]	0.60 [0.46, 0.74]		
Jiamsakul 2014	- 7	9	5	60	0.58 [0.28, 0.85]	0.87 [0.77, 0.94]		
Labhardt 2012	15	0	50	27	0.23 [0.14, 0.35]	1.00 [0.87, 1.00]		
McMahon 2013	12	38	22	98	0.35 [0.20, 0.54]	0.72 [0.64, 0.79]		-
Nelson 2010	46	79	104	417	0.31 [0.23, 0.39]	0.84 [0.81, 0.87]	+	•
San ged a 2014	10	19	45	88	0.18 [0.09, 0.31]	0.82 [0.74, 0.89]	_ , , , , , , , , , , , , , , , , , ,	
-							0 0.2 0.4 0.6 0.8 1	0 0 2 0 4 0 6 0 8 1

Test 17. [Supplementary analysis] Self-report VAS; threshold: ≥ 90% adherence

[Supplementary analysis] Self-report VAS; threshold: ≥ 90% adherence

Study	ΤР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)	į
Mbengue 2019	12	14	37	100	0.24 [0.13, 0.39]	0.88 (0.80, 0.93) —=	
Sangeda 2014	5	10	50	97	0.09 [0.03, 0.20]	0.91 [0.83, 0.95] 💻 🗕	
Segeral 2010	2	9	72	174	0.03 [0.00, 0.09]		

Test 18. [Supplementary analysis] Self-report VAS; threshold: ≥ 80% adherence

[Supplementary analysis] Self-report VAS; threshold: ≥ 80% adherence

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)
 Sensitivity (95% CI)
 Specificity (95% CI)

Test 19. [Subgroup analysis by viral load] Self-report VAS; threshold: ≥ 95% adherence; 40 to 100 copies/mL

[Subgroup analysis by viral load] Self-report VAS; threshold: ≥ 95% adherence; 40 to 100 copies/mL

Study	ΤР	FP	FN	TN	Sensitivity (95% Cl)	Specificity (95% CI)	Sensitivity (95%	CI)Specificity (95% CI)
Cerutti 2016	52	558	63	691	0.45 [0.36, 0.55]	0.55 [0.53, 0.58]		•
Cohen 2012	41	134	863	176	0.05 [0.03, 0.06]	0.57 [0.51, 0.62]	•	-
Ekstran d 2010	18	16	37	131	0.33 [0.21, 0.47]	0.89 [0.83, 0.94]		-
Haberer 2011	6	21	14	32	0.30 [0.12, 0.54]	0.60 [0.46, 0.74]		
Labhardt 2012	15	0	50	27	0.23 [0.14, 0.35]	1.00 [0.87, 1.00]		
Nelson 2010	46	79	104	417	0.31 [0.23, 0.39]	0.84 [0.81, 0.87]	0 0.2 0.4 0.6 0.8	1 0 0.2 0.4 0.6 0.8 1

Test 20. [Subgroup analysis by viral load] Self-report VAS; threshold: ≥ 95% adherence; 200 to 400 copies/mL

[Subgroup analysis by viral load] Self-report VAS; threshold: ≥ 95% adherence; 200 to 400 copies/mL

Study	ТР	FP	FN	ΤN	Sensitivity (95% CI)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)
Dziva 2017	16	12	45	93	0.26 [0.16, 0.39]	0.89 [0.81, 0.94] —
Gill 2010	0	2	8	55	0.00 [0.00, 0.37]	0.96 [0.88, 1.00] -
Jiamsakul 2014	- 7	9	5	60	0.58 [0.28, 0.85]	0.87 [0.77, 0.94]
McMahon 2013	12	38	22	98	0.35 [0.20, 0.54]	0.72 [0.64, 0.79]
San ged a 2014	10	19	45	88	0.18 [0.09, 0.31]	0.82 [0.74, 0.89]

Test 21. [Subgroup analysis by population] Self-report VAS; threshold: ≥ 95% adherence; children

[Subgroup analysis by population] Self-report VAS; threshold: ≥ 95% adherence; children

Study	ТР	FP	FN	ΤN	Sensitivity (95% Cl)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)	1
Dziva 2017	16	12	45	93	0.26 [0.16, 0.39]	0.89 [0.81, 0.94] —	
Haberer 2011	6	21	14	32	0.30 [0.12, 0.54]	0.60 [0.46, 0.74]	

Test 22. [Subgroup analysis by population] Self-report VAS; threshold: ≥ 95% adherence; adults

[Subgroup analysis by population] Self-report VAS; threshold: ≥ 95% adherence; adults

Study	тр	FP	FN	ΤN	Sensitivity (95% CI)	Specificity (95% CI) 9	Sensitivity (95% C	I)Specificity (95% CI)
Cerutti 2016	52	558	63	691	0.45 [0.36, 0.55]	0.55 [0.53, 0.58]		•
C ohe n 2012	41	134	863	176	0.05 [0.03, 0.06]	0.57 [0.51, 0.62]	•	
Ekstrand 2010	18	16	37	131	0.33 [0.21, 0.47]	0.89 [0.83, 0.94]		
Gill 2010	0	2	8	55	0.00 [0.00, 0.37]	0.96 [0.88, 1.00]		
Jiamsakul 2014	- 7	9	5	60	0.58 [0.28, 0.85]	0.87 [0.77, 0.94]		
McMahon 2013	12	38	22	98	0.35 [0.20, 0.54]	0.72 [0.64, 0.79]		
Nelson 2010	46	- 79	104	417	0.31 [0.23, 0.39]	0.84 [0.81, 0.87]	-	
San ged a 2014	10	19	45	88	0.18 [0.09, 0.31]	0.82 [0.74, 0.89]	0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1



Test 23. [Subgroup analysis by setting] Self-report VAS; threshold: ≥ 95% adherence; low-income

[Subgroup analysis by setting] Self-report VAS; threshold: ≥ 95% adherence; low-income

Study	ТР	FP	FN	τN	Sensitivity (95% Cl)	Specificity (95% CI) Sensit	tivity (95% CI)Specificity (95% CI)
Dziva 2017	16	12	45	93	0.26 [0.16, 0.39]	0.89 [0.81, 0.94]	
Haberer 2011	6	21	14	32	0.30 [0.12, 0.54]	0.60 [0.46, 0.74] —	─ ─
Labhardt 2012	15	0	50	27	0.23 [0.14, 0.35]	1.00 [0.87, 1.00]	•
McMahon 2013	12	38	22	98	0.35 [0.20, 0.54]	0.72 [0.64, 0.79] —	• ·•
San ged a 2014	10	19	45	88	0.18 [0.09, 0.31]	0.82 [0.74, 0.89]	0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

Test 24. [Subgroup analysis by setting] Self-report VAS; threshold: ≥ 95% adherence; lower-middle-income

[Subgroup analysis by setting] Self-report VAS; threshold: ≥ 95% adherence; lower-middle-income

Study	ТР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95%	CI)Specificity (95% CI)
Cerutti 2016	52	558	63	691	0.45 [0.36, 0.55]	0.55 [0.53, 0.58]		
Ekstrand 2010	18	16	37	131	0.33 [0.21, 0.47]	0.89 [0.83, 0.94]		-
Gill 2010	0	2	8	55	0.00 [0.00, 0.37]	0.96 [0.88, 1.00]		1 0 0.2 0.4 0.6 0.8 1
							0 0.2 0.4 0.6 0.8	1 0 0.2 0.4 0.6 0.8 1

Test 25. [Main analysis] Tablet count; threshold: ≥ 95% adherence

[Main analysis] Tablet count; threshold: ≥ 95% adherence

Study	ΤР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)
Apisarnthanarak 2010	19	25	6	149	0.76 [0.55, 0.91]	0.86 [0.80, 0.90] —
Bonjoch 2006	47	144	23	8	0.67 [0.55, 0.78]	0.05 [0.02, 0.10]
Cerutti 2016	26	275	88	941	0.23 [0.15, 0.32]	0.77 [0.75, 0.80] 📲
Coker 2015	25	93	22	133	0.53 [0.38, 0.68]	0.59 [0.52, 0.65]
Gill 2010	2	9	6	48	0.25 [0.03, 0.65]	0.84 [0.72, 0.93]
Haberer 2011	- 7	26	13	27	0.35 [0.15, 0.59]	0.51 [0.37, 0.65]
Kitkungvan 2008	4	40	0	155	1.00 [0.40, 1.00]	0.79 [0.73, 0.85]
Mariana 2018	0	1	16	81	0.00 [0.00, 0.21]	0.99 [0.93, 1.00] -
Moosa 2019	1	11	10	211	0.09 [0.00, 0.41]	0.95 [0.91, 0.98] -
Okonji 2012	24	44	66	300	0.27 [0.18, 0.37]	0.87 [0.83, 0.91] 🗕 🗕
Orrell 2017	9	119	35	15	0.20 [0.10, 0.35]	0.11 [0.06, 0.18] — — — —
Sangeda 2014	43	89	12	18	0.78 [0.65, 0.88]	0.17 [0.10, 0.25]

Test 26. [Supplementary analysis] Tablet count; threshold: ≥ 80% adherence

[Supplementary analysis] Tablet count; threshold: ≥ 80% adherence

Study	ΤР	FP	FN	ΤN	Sensitivity (95% CI)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)
Haberer 2011	0	0	20	53	0.00 [0.00, 0.17]	1.00 [0.93, 1.00] 💻 🚽
San ged a 2014	19	33	36	74	0.35 [0.22, 0.49]	

Test 27. [Subgroup analysis by viral load] Tablet count; threshold: ≥ 95% adherence; VL 40 to 80 copies/mL

[Subgroup analysis by viral load] Tablet count; threshold: ≥ 95% adherence; VL 40 to 80 copies/mL

Study	тр	FP	FN	TN	Sensitivity (95% Cl)	Specificity (95% CI) Sensit	ivity (95% CI)Specificity (95% CI)
Apisarnthanarak 2010	19	25	6	149	0.76 [0.55, 0.91]	0.86 [0.80, 0.90]	
Bonjoch 2006	47	144	23	8	0.67 [0.55, 0.78]	0.05 [0.02, 0.10]	
Cerutti 2016	26	275	88	941	0.23 [0.15, 0.32]	0.77 [0.75, 0.80]	
Haberer 2011	- 7	26	13	27	0.35 [0.15, 0.59]	0.51 [0.37, 0.65]	• •
Kitkungvan 2008	4	40	0	155	1.00 [0.40, 1.00]	0.79 [0.73, 0.85]	
Mariana 2018	0	1	16	81	0.00 [0.00, 0.21]	0.99 [0.93, 1.00] =	
Orrell 2017	9	119	35	15	0.20 [0.10, 0.35]		- 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

Test 28. [Subgroup analysis by viral load] Tablet count; threshold: ≥ 95% adherence; VL 400 copies/mL

[Subgroup analysis by viral load] Tablet count; threshold: ≥ 95% adherence; VL 400 copies/mL

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)
C o ker 2015	25	93	22	133	0.53 [0.38, 0.68]	0.59 [0.52, 0.65]
Gill 2010	2	9	6	48	0.25 [0.03, 0.65]	0.84 [0.72, 0.93]
Moosa 2019	1	11	10	211	0.09 [0.00, 0.41]	0.95 [0.91, 0.98] -
0k o nji 2012	24	44	66	300	0.27 [0.18, 0.37]	0.87 [0.83, 0.91] 🚽 🗕
San ged a 2014	43	89	12	18	0.78 [0.65, 0.88]	0.17 [0.10, 0.25]

Test 29. [Subgroup analysis by population] Tablet counts; threshold: ≥ 95% adherence; children

[Subgroup analysis by population] Tablet counts; threshold: ≥ 95% adherence; children

Study	ТР	FP	FN	ΤN	Sensitivity (95% CI)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)
Haberer 2011	7	26	13	27	0.35 [0.15, 0.59]	0.51 [0.37, 0.65]

Test 30. [Subgroup analysis by population] Tablet counts; threshold: ≥ 95% adherence; adults

[Subgroup analysis by population] Tablet counts; threshold: ≥ 95% adherence; adults

Study	ТР	FP	FN	TN	Sensitivity (95% Cl)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)
Apisarnthanarak 2010	19	25	6	149	0.76 [0.55, 0.91]	0.86 [0.80, 0.90]
Bonjoch 2006	47	144	23	8	0.67 [0.55, 0.78]	0.05 [0.02, 0.10]
Cerutti 2016	26	275	88	941	0.23 [0.15, 0.32]	0.77 [0.75, 0.80]
Coker 2015	25	93	22	133	0.53 [0.38, 0.68]	0.59 [0.52, 0.65]
Gill 2010	2	9	6	48	0.25 [0.03, 0.65]	0.84 [0.72, 0.93]
Mariana 2018	0	1	16	81	0.00 [0.00, 0.21]	0.99 [0.93, 1.00] =
Moosa 2019	1	11	10	211	0.09 [0.00, 0.41]	0.95 [0.91, 0.98] -
Okonji 2012	24	44	66	300	0.27 [0.18, 0.37]	0.87 [0.83, 0.91] 🚽
Sangeda 2014	43	89	12	18	0.78 [0.65, 0.88]	0.17 [0.10, 0.25]

Test 31. [Subgroup analysis by population] Tablet counts; threshold: ≥ 95% adherence; mixed

[Subgroup analysis by population] Tablet counts; threshold: ≥ 95% adherence; mixed

Study	тр	FP	FN	TN	Sensitivity (95% Cl)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)
Kitkungvan 2008	4	40	0	155	1.00 [0.40, 1.00]	0.79 [0.73, 0.85]
Orrell 2017	9	119	35	15	0.20 [0.10, 0.35]	0.11 [0.06, 0.18]



Test 32. [Subgroup analysis by setting] Tablet counts; threshold: ≥ 95% adherence; low-income

[Subgroup analysis by setting] Tablet counts; threshold: ≥ 95% adherence; low-income

Study	ТР	FP	FN	TN	Sensitivity (95% Cl)	Specificity (95% CI) Sensitivity (9	5% CI)Specificity (95% CI)
Coker 2015	25	93	22	133	0.53 [0.38, 0.68]	0.59 [0.52, 0.65]	
Haberer 2011	- 7	26	13	27	0.35 [0.15, 0.59]	0.51 [0.37, 0.65]	
0k o nji 2012	24	44	66	300	0.27 [0.18, 0.37]	0.87 [0.83, 0.91]	•
San ged a 2014	43	89	12	18	0.78 [0.65, 0.88]	0.17 [0.10, 0.25]	

Test 33. [Subgroup analysis by setting] Tablet counts; threshold: ≥ 95% adherence; lower-middle-income

[Subgroup analysis by setting] Tablet counts; threshold: ≥ 95% adherence; lower-middle-income

Study	ΤР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)
Cerutti 2016	26	275	88	941	0.23 [0.15, 0.32]	0.77 [0.75, 0.80] 💻
Gill 2010	2	9	6	48	0.25 [0.03, 0.65]	0.84 [0.72, 0.93]
Kitkungvan 2008	4	40	0	155	1.00 [0.40, 1.00]	0.79 [0.73, 0.85]
Mariana 2018	0	1	16	81	0.00 [0.00, 0.21]	0.99 [0.93, 1.00]

Test 34. [Subgroup analysis by setting] Tablet counts; threshold: ≥ 95% adherence; upper-middle-income

[Subgroup analysis by setting] Tablet counts; threshold: ≥ 95% adherence; upper-middle-income

Study	ТР	FP	FN	TN	Sensitivity (95% Cl)	Specificity (95% Cl)	Sensitivity (95% CI)Specificity (95% CI)
Apisarnthanarak 2010	19	25	6	149	0.76 [0.55, 0.91]	0.86 [0.80, 0.90]	
Moosa 2019	1	11	10	211	0.09 [0.00, 0.41]	0.95 [0.91, 0.98]	·
Orrell 2017	9	119	35	15	0.20 [0.10, 0.35]	0.11 [0.06, 0.18]	

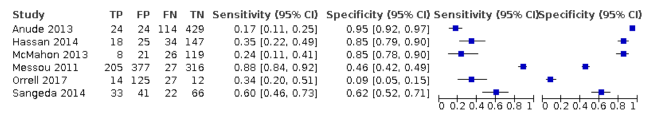
Test 35. [Subgroup analysis by setting] Tablet counts; threshold: ≥ 95% adherence; high-income

[Subgroup analysis by setting] Tablet counts; threshold: ≥ 95% adherence; high-income

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)
 Sensitivity (95% CI)

Test 36. [Main analysis] Pharmacy records; threshold: ≥ 95% adherence

[Main analysis] Pharmacy records; threshold: ≥ 95% adherence



Test 37. [Supplementary analysis] Pharmacy records; threshold: ≥ 80% adherence

[Supplementary analysis] Pharmacy records; threshold: ≥ 80% adherence

Study	ТР	FP	FN	TN	Sensitivity (95% Cl)	Specificity (95% CI) Sen	sitivity (95% CI)Spe	cificity (95% CI)
Messou 2011	148	97	84	596	0.64 [0.57, 0.70]	0.86 [0.83, 0.89]	-	
Navarro 2014	51	17	11	45	0.82 [0.70, 0.91]	0.73 [0.60, 0.83]		
San ged a 2014	14	13	41	94	0.25 [0.15, 0.39]			0.2 0.4 0.6 0.8 1

Test 38. [Subgroup analysis by viral load] Pharmacy records; threshold: ≥ 95% adherence; VL: 40 copies/mL

[Subgroup analysis by viral load] Pharmacy records; threshold: ≥ 95% adherence; VL: 40 copies/mL

Test 39. [Subgroup analysis by viral load] Pharmacy records; threshold: ≥ 95% adherence; VL: 200 to 400 copies/mL

[Subgroup analysis by viral load] Pharmacy records; threshold: ≥ 95% adherence; VL: 200 to 400 copies/mL

Study	ТР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% Cl)	Sensitivity (95%	CI)Specificity (95% CI)
Anu de 2013	24	24	114	429	0.17 [0.11, 0.25]	0.95 [0.92, 0.97]	+	•
Hassan 2014	18	25	34	147	0.35 [0.22, 0.49]	0.85 [0.79, 0.90]		+
McMahon 2013	8	21	26	119	0.24 [0.11, 0.41]	0.85 [0.78, 0.90]		-
Messou 2011	205	377	27	316	0.88 [0.84, 0.92]	0.46 [0.42, 0.49]	-	• •
San ged a 2014	33	41	22	66	0.60 [0.46, 0.73]	0.62 [0.52, 0.71]	0 0.2 0.4 0.6 0.8	1 0 0.2 0.4 0.6 0.8 1

Test 40. [Subgroup analysis by population] Pharmacy records; threshold: ≥ 95% adherence; mixed

[Subgroup analysis by population] Pharmacy records; threshold: ≥ 95% adherence; mixed

Study	ΤР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity	(95% CI)Specificity (95% CI)
Hassan 2014	18	25	34	147	0.35 [0.22, 0.49]	0.85 [0.79, 0.90]		-
Orrell 2017	14	125	27	12	0.34 [0.20, 0.51]	0.09 [0.05, 0.15]	0 0.2 0.4 0.	

Test 41. [Subgroup analysis by population] Pharmacy records; threshold: ≥ 95% adherence; adults

[Subgroup analysis by population] Pharmacy records; threshold: ≥ 95% adherence; adults

Study	ТР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI) 9	Sensitivity (95%	CI)Specificity (95% CI)
Anu de 2013	24	24	114	429	0.17 [0.11, 0.25]	0.95 [0.92, 0.97]	-	-
McMahon 2013	18	25	34	147	0.35 [0.22, 0.49]	0.85 [0.79, 0.90]		-
Messou 2011	205	377	27	316	0.88 [0.84, 0.92]	0.46 [0.42, 0.49]	-	• •
San ged a 2014	24	41	22	66	0.52 [0.37, 0.67]	0.62 [0.52, 0.71]	0 0.2 0.4 0.6 0.8	1 0 0.2 0.4 0.6 0.8 1

Test 42. [Subgroup analysis by setting] Pharmacy records; threshold: ≥ 95% adherence; low-income

[Subgroup analysis by setting] Pharmacy records; threshold: ≥ 95% adherence; low-income

Study	ТР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)
Hassan 2014	18	25	34	147	0.35 [0.22, 0.49]	0.85 [0.79, 0.90] —
McMahon 2013	8	21	26	119	0.24 [0.11, 0.41]	0.85 [0.78, 0.90] —
Messou 2011	205	377	27	316	0.88 [0.84, 0.92]	0.46 [0.42, 0.49] 🗕 🗧
San ged a 2014	33	41	22	66	0.60 [0.46, 0.73]	0.62 [0.52, 0.71]

Test 43. [Subgroup analysis by setting] Pharmacy records; threshold: ≥ 95% adherence; lower-middle-income

[Subgroup analysis by setting] Pharmacy records; threshold: ≥ 95% adherence; lower-middle-income

Test 44. [Subgroup analysis by setting] Pharmacy records; threshold: ≥ 95% adherence; upper-middle-income

[Subgroup analysis by setting] Pharmacy records; threshold: ≥ 95% adherence; upper-middle-income

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)
 Sensitivity (95% CI)
 Specificity (95% CI)

Test 45. [Main analysis] Electronic monitoring; threshold: ≥ 95% adherence

[Main analysis] Electronic monitoring; threshold: ≥ 95% adherence

Study	ТР	FP	FN	ΤN	Sensitivity (95% Cl)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
Evans 2016	20	16	7	6	0.74 [0.54, 0.89]	0.27 [0.11, 0.50]	_ - _
Gill 2010	- 7	27	1	30	0.88 [0.47, 1.00]	0.53 [0.39, 0.66]	
Haberer 2011	12	17	8	35	0.60 [0.36, 0.81]	0.67 [0.53, 0.80]	

Test 46. [Supplementary analysis] Electronic monitoring; threshold: ≥ 80% adherence

[Supplementary analysis] Electronic monitoring; threshold: ≥ 80% adherence

Study	ТР	FP	FN	ΤN	Sensitivity (95% Cl)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)
Evans 2016	11	5	16	17	0.41 [0.22, 0.61]	0.77 [0.55, 0.92]
Farley 2003	8	3	1	14	0.89 [0.52, 1.00]	0.82 [0.57, 0.96]
Haberer 2011	5	2	15	50	0.25 [0.09, 0.49]	0.96 [0.87, 1.00] —
Orrell 2017	19	95	59	7	0.24 [0.15, 0.35]	

Test 47. [Subgroup analysis by viral load] Electronic monitoring; threshold: ≥ 95% adherence; VL: 50 copies/mL

[Subgroup analysis by viral load] Electronic monitoring; threshold: ≥ 95% adherence; VL: 50 copies/mL

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)
 Sensitivity (95% CI)
 Specificity (95% CI)

Test 48. [Subgroup analysis by viral load] Electronic monitoring; threshold: ≥ 95% adherence; VL: 400 copies/mL

[Subgroup analysis by viral load] Electronic monitoring; threshold: ≥ 95% adherence; VL: 400 copies/mL

Study	ΤР	FP	FN	ΤN	Sensitivity (95% Cl)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% C	30
Evans 2016	20	16	7	6	0.74 [0.54, 0.89]	0.27 [0.11, 0.50]	
Gill 2010	7	27	1	30	0.88 [0.47, 1.00]		H

Test 49. [Subgroup analysis by population] Electronic monitoring; threshold: ≥ 95% adherence; children

[Subgroup analysis by population] Electronic monitoring; threshold: ≥ 95% adherence; children

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)
 Sensitivity (95% CI)
 Specificity (95% CI)

 Haberer 2011
 12
 17
 8
 35
 0.60 [0.36, 0.81]
 0.67 [0.53, 0.80]
 Image: Comparison of the sensitivity (95% CI)
 Image: Comparison of the sensity (95% CI)
 <td

Test 50. [Subgroup analysis by population] Electronic monitoring; threshold: ≥ 95% adherence; adults

[Subgroup analysis by population] Electronic monitoring; threshold: ≥ 95% adherence; adults

Study	ΤР	FP	FN	ΤN	Sensitivity (95% CI)	Specificity (95% Cl) Sensitivity (95% Cl)Specificity (95% Cl)
Evans 2016	20	16	- 7	6	0.74 [0.54, 0.89]	0.27 [0.11, 0.50]
Gill 2010	7	27	1	30	0.88 [0.47, 1.00]	

Test 51. [Subgroup analysis by setting] Electronic monitoring; threshold: ≥ 95% adherence; low-income

[Subgroup analysis by setting] Electronic monitoring; threshold: > 95% adherence; low-income

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)
 Sensitivity (95% CI)

Test 52. [Subgroup analysis by setting] Electronic monitoring; threshold: ≥ 95% adherence; lower-middle-income

[Subgroup analysis by setting] Electronic monitoring; threshold: ≥ 95% adherence; lower-middle-incom

Study	ΤР	FP	FN	ΤN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
Gill 2010	- 7	27	1	30	0.88 [0.47, 1.00]	0.53 [0.39, 0.66]	
							0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

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Test 53. [Subgroup analysis by setting] Electronic monitoring; threshold: ≥ 95% adherence; upper-middle-income

[Subgroup analysis by setting] Electronic monitoring; threshold: ≥ 95% adherence; upper-middle-income

Study	ΤР	FP	FN	τN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% Cl	Specificity (95% CI)
Evans 2016	20	16	7	6	0.74 [0.54, 0.89]	0.27 [0.11, 0.50]	0 0.2 0.4 0.6 0.8 1	

Test 54. [Main analysis] Composite measure; different thresholds*

[Main analysis] Composite measure; different thresholds*

Study	ΤР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specifie	city (95% CI)
Jayaweera 2003	9	0	4	6	0.69 [0.39, 0.91]	1.00 [0.54, 1.00]	_	
Mbengue 2019	9	15	40	99	0.18 [0.09, 0.32]	0.87 [0.79, 0.92]		
McMahon 2013	17	45	17	95	0.50 [0.32, 0.68]	0.68 [0.59, 0.75]		-
Mutwa 2014	8	13	17	66	0.32 [0.15, 0.54]	0.84 [0.74, 0.91]		
Orrell 2003	49	69	33	91	0.60 [0.48, 0.70]	0.57 [0.49, 0.65]		-
Ort eg a 2004	33	43	6	54	0.85 [0.69, 0.94]	0.56 [0.45, 0.66]		
Parienti 2010	21	26	0	25	1.00 [0.84, 1.00]	0.49 [0.35, 0.63]		
Segeral 2010	20	28	54	155	0.27 [0.17, 0.39]	0.85 [0.79, 0.90]		-
Spire 2008	7	9	63	267	0.10 [0.04, 0.20]	0.97 [0.94, 0.98]		4 0.6 0.8 1

Test 55. [Subgroup analysis by adherence threshold] Composite measures; 100% adherence

[Subgroup analysis by adherence threshold] Composite measures; 100% adherence

Study	ТР	FP	FN	TN	Sensitivity (95% Cl)	Specificity (95% CI) 9	Sensitivity (95% CI)Specificity (95% CI)
Jayaweera 2003	9	0	4	6	0.69 [0.39, 0.91]	1.00 [0.54, 1.00]	
MA	~		40	~~	A 1 A /A AA A A A A	A 07 (A 76 A 60)	

jayaweera 2005	9	- U	- 4	0	0.09 [0.39, 0.91]	1.00 [0.54, 1.00]	
Mbengue 2019	9	15	40	99	0.18 [0.09, 0.32]	0.87 [0.79, 0.92]	 -
McMahon 2013	17	45	17	95	0.50 [0.32, 0.68]	0.68 [0.59, 0.75]	
Ort eg a 2004	33	43	6	54	0.85 [0.69, 0.94]	0.56 [0.45, 0.66]	
Segeral 2010	20	28	54	155	0.27 [0.17, 0.39]	0.85 [0.79, 0.90]	 -
Spire 2008	7	9	63	267	0.10 [0.04, 0.20]	0.97 [0.94, 0.98]	0.20.40.60.81

Test 56. [Subgroup analysis by adherence threshold] Composite measures; 95% adherence

[Subgroup analysis by adherence threshold] Composite measures; 95% adherence

Study	тр	FP	FN	τN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)S	pecificity (95% CI)
Mutwa 2014	8	13	17	66	0.32 [0.15, 0.54]	0.84 [0.74, 0.91]		
Orrell 2003	49	69	33	91	0.60 [0.48, 0.70]	0.57 [0.49, 0.65]		-
Parienti 2010	21	26	0	25	1.00 [0.84, 1.00]	0.49 [0.35, 0.63]	0 0.2 0.4 0.6 0.8 1	

Test 57. [Subgroup analysis by viral load] Composite measure; various adherence thresholds*; 40 to 50 copies/mL

[Subgroup analysis by viral load] Composite measure; various adherence thresholds*; 40 to 50 copies/mL

Study	ΤР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)	
Mutwa 2014	8	13	17	66	0.32 [0.15, 0.54]	0.84 [0.74, 0.91]	
Parienti 2010	21	26	0	25	1.00 [0.84, 1.00]	0.49 [0.35, 0.63] — — — — —	
Spire 2008	7	9	63	267	0.10 [0.04, 0.20]	0.97 [0.94, 0.98]	

Test 58. [Subgroup analysis by viral load] Composite measure; various adherence thresholds*; 200 to 400 copies/mL

[Subgroup analysis by viral load] Composite measure; various adherence thresholds*; 200 to 400 copies/mL

Study	ΤР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
Jayaweera 2003	9	0	4	6	0.69 [0.39, 0.91]	1.00 [0.54, 1.00]	
Mbengue 2019	9	15	40	99	0.18 [0.09, 0.32]	0.87 [0.79, 0.92]	
McMahon 2013	17	45	17	95	0.50 [0.32, 0.68]	0.68 [0.59, 0.75]	
Orrell 2003	49	69	33	91	0.60 [0.48, 0.70]	0.57 [0.49, 0.65]	
Ortega 2004	33	43	6	54	0.85 [0.69, 0.94]	0.56 [0.45, 0.66]	
Parienti 2010	16	31	0	25	1.00 [0.79, 1.00]	0.45 [0.31, 0.59]	
Segeral 2010	20	28	54	155	0.27 [0.17, 0.39]	0.85 [0.79, 0.90]	

Test 59. [Subgroup analysis by population] Composite measure; various adherence thresholds*; children

[Subgroup analysis by population] Composite measure; various adherence thresholds*; children

Study	ΤР	FP	FN	ΤN	Sensitivity (95% CI)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)
Mutwa 2014	8	13	17	66	0.32 [0.15, 0.54]	

Test 60. [Subgroup analysis by population] Composite measure; various adherence thresholds*; adults

[Subgroup analysis by population] Composite measure; various adherence thresholds*; adults

Study TP FP FN TN Sensitivity (95% Cl) Specificity (95% Cl) Sensitivity (95% Cl)Specificity (95% Cl)

Mbengue 2019	9	15	40	99	0.18 [0.09, 0.32]	0.87 [0.79, 0.92]	 -
McMahon 2013	17	45	17	95	0.50 [0.32, 0.68]	0.68 [0.59, 0.75]	 -
Orrell 2003	49	69	33	91	0.60 [0.48, 0.70]	0.57 [0.49, 0.65]	 -
Ortega 2004	33	43	6	54	0.85 [0.69, 0.94]	0.56 [0.45, 0.66]	
Parienti 2010	21	26	0	25	1.00 [0.84, 1.00]	0.49 [0.35, 0.63]	
Segeral 2010	20	28	54	155	0.27 [0.17, 0.39]	0.85 [0.79, 0.90]	 -
Spire 2008	7	9	63	267	0.10 [0.04, 0.20]	0.97 [0.94, 0.98]	0 0.2 0.4 0.6 0.8 1

Test 61. [Subgroup analysis by setting] Composite measure; various adherence thresholds*; low-income

[Subgroup analysis by setting] Composite measure; various adherence thresholds*; low-income

Study	ТР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95%)	CI)Specificity (95% CI)
McMahon 2013	17	45	17	95	0.50 [0.32, 0.68]	0.68 [0.59, 0.75]		-
Mutwa 2014	8	13	17	66	0.32 [0.15, 0.54]	0.84 [0.74, 0.91]		
Segeral 2010	20	28	54	155	0.27 [0.17, 0.39]	0.85 [0.79, 0.90]		-
Spire 2008	7	9	63	267	0.10 [0.04, 0.20]	0.97 [0.94, 0.98]	0 0.2 0.4 0.6 0.8	

Test 62. [Subgroup analysis by setting] Composite measure; various adherence thresholds*; upper-middle-income

[Subgroup analysis by setting] Composite measure; various adherence thresholds*; upper-middle-income

Study	ТР	FP	FN	τN	Sensitivity (95% Cl)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI	0
Mbengue 2019	9	15	40	99	0.18 [0.09, 0.32]	0.87 [0.79, 0.92]	
Orrell 2003	49	69	33	91	0.60 [0.48, 0.70]	0.57 [0.49, 0.65]	ł

Test 63. [Subgroup analysis by setting] Composite measure; various adherence thresholds*; high-income

[Subgroup analysis by setting] Composite measure; various adherence thresholds*; high-income

Study	ТР	FP	FN	ΤN	Sensitivity (95% CI)	Specificity (95% CI) 9	Sensitivity (95% CI)Specificity (95% CI)
Jayaweera 2003	9	0	4	6	0.69 [0.39, 0.91]	1.00 [0.54, 1.00]	
Ortega 2004	33	43	6	54	0.85 [0.69, 0.94]	0.56 [0.45, 0.66]	
Parienti 2010	21	26	0	25	1.00 [0.84, 1.00]	0.49 [0.35, 0.63]	

ADDITIONAL TABLES

Table 1. Guidelines for determining viral failure

Guideline	Threshold
DHHS 2017	Persistent (> 1 reading of > 200 copies/mL) denotes viral failure after 24 weeks on an ART regimen in a person who has not yet had documented virological suppression on this regimen.
EACS 2017	Confirmed (< 1 month) HIV viral load > 50 copies/mL 6 months after starting therapy (initiation or modification) in people on ART. Depending on the HIV viral load assay, this limit could be higher or lower.
WHO 2016	Persistently detectable viral load exceeding 1000 copies/mL (i.e. 2 consecutive viral load measure- ments within a 3-month interval with adherence support between measurements) after ≥ 6 months of starting a new ART regimen.

ART: antiretroviral therapy

Table 2. Health service nomenclature

Tier	Highest cadre	Terms often used	Facility and staff	Equipment facilities		
Community	Individual with maximum of few	Family-led care	Family member	HIV tests, counselling, re- plenish drugs		
	months training, paid or unpaid	Community volun- teer	Trained volunteer; health assistants	— pienisir drugs		
		Primary care clinic	Nurse aide or community health work- ers			
Health centre	Clinical officer or nurse (≥ 2 years' training)	Health centres; dis- trict hospitals	Purpose built with ≥ 1 paramedic or nurse with some health assistants	HIV tests; antiretroviral drugs; opportunistic infec- tion medicines; point-of- care laboratories		

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

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Table 2. Health service nomenclature (Continued)

Health centre (enhanced)	Clinical officer or nurse (≥ 2 years' training)	Health centres, pri- mary health care clinics, district hos- pitals	Purpose built with ≥ 1 paramedic or nurse with some health assistants, with input from a doctor (may be via mobile support service)	HIV tests; antiretroviral drugs; opportunistic infec- tion medicines; point-of- care laboratories		
Hospital	Doctor	Health centres; dis- trict hospitals	Purpose built with ≥ 1 medical doctor with nurses/paramedics and assistants	CD4 count; medicines; not viral load		
Hospital (ad- vanced)	Specialist doctor	District hospital; re- ferral hospital	Purpose built with ≥ 2 specialist doc- tors with nurses/paramedics and assis- tants	Viral load; full investiga- tions		

CD4: cluster of differentiation

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Excluded studies: duplicate reference (N = 37)

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Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

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Matteo S, Bruno G, Astuti N, Filippo E, Valenti D, Colombo G, et al. Switching from an EFV-based STR to a RPV-based STR is effective, safe and improves HIV patients health status. 2014;17(7):A677-8



Table 5. Excluded studies: wrong index test (Continued)

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Shearer K, Evans D, Xhosa B, Hirasen K, Bracken C, Mahomed K, et al. Low prevalence of depressive symptoms among stable patients on antiretroviral therapy in Johannesburg, South Africa. PLOS One 2018;13(9):e0203797

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Strehlau R, Shiau S, Arpadi S, Patel F, Pinillos F, Tsai WY, et al. Substituting abacavir for stavudine in children who are virally suppressed without lipodystrophy: randomized clinical trial in Johannesburg, South Africa. Journal of the Pediatric Infectious Diseases Society 2018;7(3):E70-7

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Venter F, Moorhouse M, Sokhela S, Maharaj E, Akpomiemie G, Simmons B, et al. Non-inferior efficacy for darunavir/ritonavir 400/100 mg once daily versus lopinavir/ritonavir, for patients with HIV RNA below 50 copies/mL in South Africa: the 48-week WRHI 052 study. Journal of the International AIDS Society 2018;21:156-7

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Table 5. Excluded studies: wrong index test (Continued)

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Young J, Smith C, Teira R, Reiss P, Jarrin Vera I, Crane H, et al. Antiretroviral pill count and clinical outcomes in treatment-naive patients with HIV infection. HIV Medicine 2018;19(2):132-42

Table 6. Excluded studies: not possible to extract data for 2x2 table

Excluded studies: not possible to extract data for 2x2 table (N = 468)

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Achieng 2012

Achieng L, Musangi H, Ong'uti S, Ombegoh E, Bryant L, Mwiindi J, et al. An observational cohort comparison of facilitators of retention in care and adherence to anti-retroviral therapy at an HIV treatment center in Kenya. PLOS One 2012;7(3):e32727

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Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

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Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

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Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

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Table 7. Excluded studies: viral load and adherence not measured at the same time

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Table 7. Excluded studies: viral load and adherence not measured at the same time (Continued)

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Ferguson NM, Donnelly CA, Hooper J, Ghani AC, Fraser C, Bartley LM, et al. Adherence to antiretroviral therapy and its impact on clinical outcome in HIV-infected patients. Journal of the Royal Society, Interface 2005;2(4):349-63

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Kaushik V, Kalampokis I, Brown P, Finkielstein A, Chice SM, Holman S, et al. Strict adherence to highly active anti-retroviral therapy (HAART) is associated with decreased serum IgE levels and decreased viral loads among HIV-1+asthmatic women. Journal of Allergy and Clinical Immunology 2008;121(2):S229

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Malhotra A, Whitley-Williams PN, Gaur S, Petrova A. Treatment response in association with adherence patterns to highly active antiretroviral therapy in pediatric patients with perinatally acquired HIV infection. Journal of the International Association of Providers of AIDS Care 2014;13(5):461-5

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Marazzi MC, Bartolo M, Gialloreti LE, Germano P, Guidotti G, Liotta G, et al. Improving adherence to highly active anti-retroviral therapy in Africa: the DREAM programme in Mozambique. Health Education Research 2006;21(1):34-42

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Nachega JB, Hislop M, Dowdy DW, Chaisson RE, Regensberg L, Maartens G. Adherence to nonnucleoside reverse transcriptase inhibitor-based HIV therapy and virologic outcomes. Annals of Internal Medicine 2007;146(8):564-73

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Palepu A, Tyndall MW, Chan K, Wood E, Montaner JS, Hogg RS. Initiating highly active antiretroviral therapy and continuity of HIV care: the impact of incarceration and prison release on adherence and HIV treatment outcomes. Antiviral Therapy 2004;9(5):713-9

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Palladino C, Briz V, Bellon JM, Climent FJ, de Ory SJ, Mellado MJ, et al. Determinants of highly active antiretroviral therapy duration in HIV-1-infected children and adolescents in Madrid, Spain, from 1996 to 2012. PLOS One 2014;9(5):e96307

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Pasternak AO, De Bruin M, Jurriaans S, Bakker M, Berkhout B, Prins JM, et al. Modest nonadherence to antiretroviral therapy promotes residual HIV-1 replication in the absence of virological rebound in plasma. Journal of Infectious Diseases 2012;206(9):1443-52

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Petersen ML, LeDell E, Schwab J, Sarovar V, Gross R, Reynolds N, et al. Super learner analysis of electronic adherence data improves viral prediction and may provide strategies for selective HIV RNA monitoring. Journal of Acquired Immune Deficiency Syndromes (1999) 2015;69(1):109-18

Pinnetti 2015

Pinnetti C, Di Giambenedetto S, Maggiolo F, Fabbiani M, Sterrantino G, Latini A, et al. Switching to coformulated rilpivirine/emtricitabine/tenofovir in virologically suppressed patients: data from a multicenter cohort. Journal of Acquired Immune Deficiency Syndromes (1999) 2015;70(4):e147-50

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Remien RH, Exner TM, Morin SF, Ehrhardt AA, Johnson MO, Correale J, et al. Medication adherence and sexual risk behavior among HIV-infected adults: implications for transmission of resistant virus. AIDS and Behavior 2007;11(5):663-75

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Richardson LA, Kerr TH, Dobrer S, Puskas CM, Guillemi SA, Montaner JS, et al. Socioeconomic marginalization and plasma HIV-1 RNA nondetectability among individuals who use illicit drugs in a Canadian setting. AIDS (London, England) 2015;29(18):2487-95

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Shet A, Neogi U, Kumarasamy N, DeCosta A, Shastri S, Rewari BB. Virological efficacy with first-line antiretroviral treatment in India: predictors of viral failure and evidence of viral resuppression. Tropical Medicine & International Health 2015;20(11):1462-72

Simoni 2014



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Sledjeski EM, Delahanty DL, Bogart LM. Incidence and impact of posttraumatic stress disorder and comorbid depression on adherence to HAART and CD4(+) counts in people living with HIV. Aids Patient Care STDS 2005;19(11):728-36

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Tupinambas U, Ribeiro FA, Aleixo A, Greco D. Treatment switch guided by HIV-1 genotyping in Brazil. Brazilian Journal of Infectious Diseases 2006;10(2):82-8

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Zoufaly A, Jochum J, Hammerl R, Nassimi N, Raymond Y, Burchard GD, et al. Virological failure after 1 year of first-line ART is not associated with HIV minority drug resistance in rural Cameroon. Journal of Antimicrobial Chemotherapy 2015;70(3):922-5

Table 8. Excluded studies: wrong patient population

Excluded studies: wrong patient population (N = 31)

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Arrondo Velasco A, Sainz Suberviola ML, Andres Esteban EM, Iruin Sanz AI, Napal Lecumberri V. [Factors associated with adherence in HIV patients]. Farmacia Hospitalaria 2009;33(1):4-11

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Table 8. Excluded studies: wrong patient population (Continued)

Atuhaire P, Hanley S, Yende-Zuma N, Aizire J, Stranix-Chibanda L, Makanani B, et al. Factors associated with unsuppressed viremia in women living with HIV on lifelong ART in the multi-country US-PEPFAR PROMOTE study: a cross-sectional analysis. PLOS One 2019;14 (10) (no pagination)(e0219415)

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Deschamps AE, De Geest S, Vandamme AM, Bobbaers H, Peetermans WE, Van Wijngaerden E. Diagnostic value of different adherence measures using electronic monitoring and virologic failure as reference standards. AIDS Patient Care & STDs 2008;22(9):735-43

Frasca 2019

Frasca K, Morrow M, Coyle RP, Coleman SS, Ellison L, Bushman LR, et al. Emtricitabine triphosphate in dried blood spots is a predictor of viral suppression in HIV infection and reflects short-term adherence to antiretroviral therapy. Journal of Antimicrobial Chemotherapy 2019;74(5):1395-401

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Table 8. Excluded studies: wrong patient population (Continued)

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Lai HH, Kuo YC, Kuo CJ, Lai YJ, Chen M, Chen YT, et al. Methamphetamine use associated with non-adherence to antiretroviral treatment in men who have sex with men. Scientific Reports 2020;10(1):7131

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Table 8. Excluded studies: wrong patient population (Continued)

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Rosen MI, Dieckhaus K, McMahon TJ, Valdes B, Petry NM, Cramer J, et al. Improved adherence with contingency management. AIDS Patient Care and STDs 2007;21(1):30-40

Shah 2007

Shah B, Walshe L, Saple DG, Mehta SH, Ramnani JP, Kharkar RD, et al. Adherence to antiretroviral therapy and virologic suppression among HIV-infected persons receiving care in private clinics in Mumbai, India. Clinical Infectious Diseases 2007;44(9):1235-44

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Yotebieng M, Thirumurthy H, Moracco K, Edmonds A, Tabala M, Kawende B, et al. Conditional cash transfers to increase retention in PMTCT care, antiretroviral adherence, and postpartum virological suppression: a randomized controlled trial. Journal of Acquired Immune Deficiency Syndromes 2016;72:S124-9

Table 9. Excluded studies: viral load obtained from medical records

Excluded studies: viral load obtained from medical records (N = 158)

Baguso 2016

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Beckwith 2017

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Belzer 2014

Belzer ME, Naar-King S, Olson J, Sarr M, Thornton S, Kahana SY, et al. The use of cell phone support for non-adherent HIV-infected youth and young adults: an initial randomized and controlled intervention trial. AIDS and Behavior 2014;18(4):686-96

Bienczak 2017

Bienczak A, Denti P, Cook A, Wiesner L, Mulenga V, Kityo C, et al. Determinants of virological outcome and adverse events in African children treated with paediatric nevirapine fixed-dose-combination tablets. AIDS (London, England) 2017;31(7):905-15

Bisson 2008



Table 9. Excluded studies: viral load obtained from medical records (Continued)

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 Medicine 2008;5(5):e109

Blumenthal 2014

Blumenthal J, Haubrich R, Jain S, Sun X, Dube M, Daar E, et al. Factors associated with high transmission risk and detectable plasma HIV RNA in HIV-infected MSM on ART. International Journal of STD & AIDS 2014;25(10):734-41

Boarts 2006

Boarts JM, Sledjeski EM, Bogart LM, Delahanty DL. The differential impact of PTSD and depression on HIV disease markers and adherence to HAART in people living with HIV. AIDS and Behavior 2006;10(3):253-61

Bonn-Miller 2014

Bonn-Miller MO, Oser ML, Bucossi MM, Trafton JA. Cannabis use and HIV antiretroviral therapy adherence and HIV-related symptoms. Journal of Behavioral Medicine 2014;37(1):1-10

Boussari 2015

Boussari O, Subtil F, Genolini C, Bastard M, Iwaz J, Fonton N, et al. Impact of variability in adherence to HIV antiretroviral therapy on the immunovirological response and mortality. BMC Medical Research Methodology 2015;15:10

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Bradley ELP, Frazier EL, Carree T, Hubbard McCree D, Sutton MY. Psychological and social determinants of health, antiretroviral therapy (ART) adherence, and viral suppression among HIV-positive black women in care. AIDS Care 2019;31(8):932-41

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Cambiano V, Lampe FC, Rodger AJ, Smith CJ, Geretti AM, Lodwick RK, et al. Use of a prescription-based measure of antiretroviral therapy adherence to predict viral rebound in HIV-infected individuals with viral suppression. HIV Medicine 2010;11(3):216-24

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Cantudo-Cuenca MR, Jimenez-Galan R, Almeida-Gonzalez CV, Morillo-Verdugo R. Concurrent use of comedications reduces adherence to antiretroviral therapy among HIV-infected patients. Journal of Managed Care & Specialty Pharmacy 2014;20(8):844-50

Chabikuli 2010

Chabikuli NO, Datonye DO, Ansong D, Nachega J, et al. Adherence to antiretroviral therapy, virologic failure and workload at the Rustenburg Provincial Hospital: original research. South African Family Practice 2010;52(4):350-5

Chaiyachati 2011

Chaiyachati K, Hirschhorn LR, Tanser F, Newell ML, Barnighausen T. Validating five questions of antiretroviral nonadherence in a public-sector treatment program in rural South Africa. AIDS Patient Care and STDs 2011;25(3):163-70

Chander 2006



Table 9. Excluded studies: viral load obtained from medical records (Continued)

Chander G, Lau B, Moore RD. Hazardous alcohol use: a risk factor for non-adherence and lack of suppression in HIV infection. Journal of Acquired Immune Deficiency Syndromes (1999) 2006;43(4):411-7

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Chandwani S, Koenig LJ, Sill AM, Abramowitz S, Conner LC, D'Angelo L. Predictors of antiretroviral medication adherence among a diverse cohort of adolescents with HIV. Journal of Adolescent Health 2012;51(3):242-51

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Christodoulou J, Abdalian SE, Jones ASK, Christodoulou G, Pentoney SL, Rotheram-Borus MJ. Crystal clear with active visualization: understanding medication adherence among youth living with HIV. AIDS and Behavior 2020;24(4):1207-11

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Cooper RL, Brown LL, Tabatabai M, Haas DW, Shepherd BE, Myers HF, et al. The effects of perceived stress and cortisol concentration on antiretroviral adherence when mediated by psychological flexibility among Southern black men living with HIV. AIDS and Behavior 2021;25(2):645-52

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Cruz 2014

Cruz ML, Cardoso CA, Darmont MQ, Souza E, Andrade SD, D'Al Fabbro MM, et al. Viral suppression and adherence among HIV-infected children and adolescents on antiretroviral therapy: results of a multicenter study. Jornal de Pediatria 2014;90(6):563-71

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Cruz CCP, Mistro S, Mendes CMC, Schooley RT, Da Silva Badaro RJ. Monitoring of delay to pharmacy refill in assessing adherence to antiretroviral therapy. Journal of Pharmacy Practice 2018;33(2):158-63

Da 2018

Da W, Li X, Qiao S, Zhou Y, Shen Z. Evaluation of self-report adherence measures and their associations with detectable viral load among people living with HIV (PLHIV) in China. PLOS One 2018;13(8):e0203032

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Dandachi D, De Groot A, Rajabiun S, Rajashekara S, Davila JA, Quinn E, et al. Reliability and validity of a brief self-report adherence measure among people with HIV experiencing homelessness and mental health or substance use disorders. AIDS and Behavior 2021;25(2):322-9

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Domingues 2015



Table 9. Excluded studies: viral load obtained from medical records (Continued)

Domingues E, Ferrit M, Calleja M. Antiretroviral therapy, adherence and quality of life in older HIV-patients with moderate-high cardiovascular risk. European Journal of Hospital Pharmacy 2015;22(Supplement 1):A87

Enriquez 2015

Enriquez M, Cheng AL, Banderas J, Farnan R, Chertoff K, Hayes D, et al. A peer-led HIV medication adherence intervention targeting adults linked to medical care but without a suppressed viral load. Journal of the International Association of Providers of AIDS Care 2015;14(5):441-8

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Evans SD, Mellins CA, Leu CS, Warne P, Elkington KS, Dolezal C, et al. HIV treatment adherence measurement and reporting concordance in youth with perinatally acquired HIV infection and their caregivers. AIDS Patient Care and STDs 2015;29(1):43-51

Fairley 2005

Fairley CK, Permana A, Read TR. Long-term utility of measuring adherence by self-report compared with pharmacy record in a routine clinic setting. HIV Medicine 2005;6(5):366-9

Farley 2008

Farley JJ, Montepiedra G, Storm D, Sirois PA, Malee K, Garvie P, et al. Assessment of adherence to antiretroviral therapy in perinatally HIV-infected children and youth using self-report measures and pill count. Journal of Developmental and Behavioral Pediatrics 2008;29(5):377-84

Fox 2018

Fox M, Pascoe S, Huber A, Murphy J, Phokojoe M, Gorgens M, et al. Viral suppression effects of interventions for unstable ART patients in South Africa. CROI 2018;26(Supplement 1):533s

Fumaz 2009

Fumaz CR, Munoz-Moreno JA, Ferrer MJ, Negredo E, Perez-Alvarez N, Tarrats A, et al. Low levels of adherence to antiretroviral therapy in HIV-1-infected women with menstrual disorders. AIDS Patient Care and STDs 2009;23(6):463-8

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Godin 2003

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Grossberg 2004

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Gunther 2014

Gunther M, Foisy M, Houston S, Guirguis L, Hughes C. Treatment beliefs, illness perceptions, and non-adherence to antiretroviral therapy in an ethnically diverse patient population. International Journal of Clinical Pharmacy 2014;36(1):105-11

Gutierrez 2012



Table 9. Excluded studies: viral load obtained from medical records (Continued)

Gutierrez EB, Sartori AM, Schmidt AL, Piloto BM, Franca BB, De Oliveira AS, et al. Measuring adherence to antiretroviral treatment: the role of pharmacy records of drug withdrawals. AIDS and Behavior 2012;16(6):1482-90

Hersch 2013

Hersch RK, Cook RF, Billings DW, Kaplan S, Murray D, Safren S, et al. Test of a web-based program to improve adherence to HIV medications. AIDS and Behavior 2013;17(9):2963-76

Hightow-Weidman 2017

Hightow-Weidman L, LeGrand S, Choi SK, Egger J, Hurt CB, Muessig KE. Exploring the HIV continuum of care among young black MSM. PLOS One 2017;12(6):e0179688

Holstad 2010

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Holstad 2011

Holstad MM, Diiorio C, McCarty F. Adherence, sexual risk, and viral load in HIV-infected women prescribed antiretroviral therapy. AIDS Patient Care and STDs 2011;25(7):431-8

Holstad 2013

Holstad MM, Ofotokun I, Higgins M, Logwood S. The LIVE network: a music-based messaging program to promote ART adherence self-management. AIDS and Behavior 2013;17(9):2954-62

Horberg 2008

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Jeffries C, Ross P, Matoff-Stepp S, Thompson R, Harris J, Uhrig J, et al. Ucare4life: mobile texting to improve HIV care continuum outcomes for minority youth. CROI 2016;24(E-1):427

Kabore 2015

Kabore L, Muntner P, Chamot E, Zinski A, Burkholder G, Mugavero MJ. Self-report measures in the assessment of antiretroviral medication adherence: comparison with medication possession ratio and HIV viral load. Journal of the International Association of Providers of AIDS Care 2015;14(2):156-62

Kacanek 2015

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Kagee 2012

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Table 9. Excluded studies: viral load obtained from medical records (Continued)

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Kalichman 2018

Kalichman SC, Cherry C, Kalichman MO, Eaton LA, Kohler JJ, Montero C, et al. Mobile health intervention to reduce HIV transmission: a randomized trial of behaviorally enhanced HIV treatment as prevention (B-TasP). Journal of Aquired Immune Deficiency Syndromes (1999) 2018;78(1):34-42

Kapiamba 2016

Kapiamba G, Masango T, Mphuthi D. Antiretroviral adherence and virological outcomes in HIVpositive patients in Ugu district, KwaZulu-Natal province. African Journal of AIDS Research 2016;15(3):195-201

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Knafl 2010

Knafl GJ, Bova CA, Fennie KP, O'Malley JP, Dieckhaus KD, Williams AB. An analysis of electronically monitored adherence to antiretroviral medications. AIDS and Behavior 2010;14(4):755-68

Knowlton 2015

Knowlton AR, Mitchell MM, Robinson AC, Nguyen TQ, Isenberg S, Denison J. Informal HIV caregiver proxy reports of care recipients' treatment adherence: relationship factors associated with concordance with recipients' viral Suppression. AIDS and Behavior 2015;19(11):2123-9

Lampe 2010

Lampe FC, Harding R, Smith CJ, Phillips AN, Johnson M, Sherr L. Physical and psychological symptoms and risk of virologic rebound among patients with virologic suppression on antiretroviral therapy. Journal of Acquired Immune Deficiency Syndromes (1999) 2010;54(5):500-5

Langwenya 2018

Langwenya N, Phillips TK, Brittain K, Zerbe A, Abrams EJ, Myer L. Same-day antiretroviral therapy (ART) initiation in pregnancy is not associated with viral suppression or engagement in care: a cohort study. Journal of the International AIDS Society 2018;21(6):e25133

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Letourneau 2013



Table 9. Excluded studies: viral load obtained from medical records (Continued)

Letourneau EJ, Ellis DA, Naar-King S, Chapman JE, Cunningham PB, Fowler S. Multisystemic therapy for poorly adherent youth with HIV: results from a pilot randomized controlled trial. AIDS Care 2013;25(4):507-14

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Leyva-Moral JM, Loayza-Enriquez BK, Palmieri PA, Guevara-Vasquez GM, Elias-Bravo UE, Edwards JE, et al. Adherence to antiretroviral therapy and the associated factors among people living with HIV/AIDS in Northern Peru: a cross-sectional study. AIDS Research and Therapy 2019;16(1):22

Lima 2008

Lima VD, Harrigan R, Murray M, Moore DM, Wood E, Hogg RS, et al. Differential impact of adherence on long-term treatment response among naive HIV-infected individuals. AIDS (London, England) 2008;22(17):2371-80

Lima 2016

Lima VD, Hull M, McVea D, Chau W, Harrigan PR, Montaner JS. Long-term effectiveness of initiating non-nucleoside reverse transcriptase inhibitor- versus ritonavir-boosted protease inhibitor-based antiretroviral therapy: implications for first-line therapy choice in resource-limited settings. Journal of the International AIDS Society 2016;19(1):20978

MacDonell 2013

MacDonell K, Naar-King S, Huszti H, Belzer M. Barriers to medication adherence in behaviorally and perinatally infected youth living with HIV. AIDS and Behavior 2013;17(1):86-93

Machado 2013

Machado Alba JE, Vidal Guitart X. Evaluación de la respuesta y seguridada diferentes esquemas de tratamiento antirretroviral en Colombia (Response assessment and safety of different schemes of anti-HIV agents in Colombia) [Avaliação da resposta e segurança a diferentes esquemas de tratamento antirretroviral em Colombia]. Investigaciones Andina 2013;15(27):770-83

Machado-Alba 2011

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Marhefka SL, Tepper VJ, Farley JJ, Sleasman JW, Mellins CA. Brief report: assessing adherence to pediatric antiretroviral regimens using the 24-hour recall interview. Journal of Pediatric Psychology 2006;31(9):989-94

Marrone 2016

Marrone G, Mellgren A, Eriksson LE, Svedhem V. High concordance between self-reported adherence, treatment outcome and satisfaction with care using a nine-item health questionnaire in Inf-CareHIV. PLOS One 2016;11(6):e0156916

Mehta 2016



Table 9. Excluded studies: viral load obtained from medical records (Continued)

Mehta K, Ekstrand ML, Heylen E, Sanjeeva GN, Shet A. Adherence to antiretroviral therapy among children living with HIV in South India. AIDS and Behavior 2016;20(5):1076-83

Meireles 2019

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Mellins 2011

Mellins CA, Tassiopoulos K, Malee K, Moscicki AB, Patton D, Smith R, et al. Behavioral health risks in perinatally HIV-exposed youth: co-occurrence of sexual and drug use behavior, mental health problems, and nonadherence to antiretroviral treatment. AIDS Patient Care and STDs 2011;25(7):413-22

Mpawa 2017

Mpawa H, Kwekwesa A, Amberbir A, Garone D, Divala OH, Kawalazira G, et al. Virological outcomes of antiretroviral therapy in Zomba central prison, Malawi; a cross-sectional study. Journal of the International AIDS Society 2017;20(1):21623

Mugavero 2009

Mugavero MJ, Raper JL, Reif S, Whetten K, Leserman J, Thielman NM, et al. Overload: impact of incident stressful events on antiretroviral medication adherence and virologic failure in a longitudinal, multisite human immunodeficiency virus cohort study. Psychosomatic Medicine 2009;71(9):920-6

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Musiime V, Kayiwa J, Kiconco M, Tamale W, Alima H, Mugerwa H, et al. Response to antiretroviral therapy of HIV type 1-infected children in urban and rural settings of Uganda. AIDS Research and Human Retroviruses 2012;28(12):1647-57

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Table 10. Ongoing studies

Ongoing studies

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3. ChiCTR1800020357. Effect of cognitive behavioral therapy on depression and antiviral treatment efficacy of HIV/AIDs patients. who.int/trialsearch/Trial2.aspx?TrialID=ChiCTR1800020357 (first received 25 December 2018)

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27. NCT00051805. Promoting adherence to anti-HIV drug regimens [Promoting adherence to antiretroviral regimens]. clinicaltrials.gov/ct2/show/NCT00051805 (first received 17 January 2003)

28. NCT00134602. Pediatric impact: promoting adherence to medications among HIV-infected children [An intervention to promote adherence to antiretroviral medications among HIV-infected children 5-12 years of age]. clinicaltrials.gov/ct2/show/NCT00134602 (first received 25 August 2005)

29. NCT00135382. Study switching HIV-1 infected patients with an undetectable viral load on a first protease inhibitor-based regimen to an efavirenz-based regimen [A Phase 3 study switching HIV-1 infected patients with an undetectable viral load on a first protease inhibitor-based regimen to an efavirenz-based regimen]. clinicaltrials.gov/ct2/show/NCT00135382 (first received 26 August 2005)

30. NCT00160849. Lopinavir/r monotherapy as maintenance therapy after long term viral suppression [Study on the feasibility of antiretroviral therapy with a single agent - lopinavir/r - in patients treated with HAART and with viral load below 80 copies/ml]. clinicaltrials.gov/ct2/show/NCT00160849 (first received 12 September 2005)

31. NCT00194545. Effect of medication diaries on adherence to highly active antiretroviral drugs among HIV-1 infected Kenyan children [Effect of medication diaries on adherence to highly active antiretroviral drugs among HIV-1 infected Kenyan children]. clinical-trials.gov/ct2/show/NCT00194545 (first 19 September 2005)

32. NCT00196612. Once daily antiretroviral therapy in HIV infected adults treated with HAART [Phase II randomized trial comparing efficacy and safety of the maintenance of a HAART association protease inhibitor containing versus a once daily antiretroviral triple association, in HIV adult patients with undetectable viral load ANRS 099 ALIZE]. clinicaltrials.gov/ct2/show/NCT00196612 (first received 20 September 2005)

33. NCT00199979. Zidovudine/lamivudine + nevirapine twice daily, versus tenofovir + lamivudine + nevirapine once daily in ARV-naive patients [Multicenter, randomized, open-label trial, assessing the efficacy of zidovudine, lamivudine and nevirapine combination administered twice daily, versus the association of tenofovir, lamivudine and nevirapine, once daily, in antiretroviral naive HIV-1 infected patients]. clinicaltrials.gov/ct2/show/NCT00199979 (first received 20 September 2005)

34. NCT00203853. Evaluation of an intervention on adherence to Highly Active Antiretroviral Therapy (HAART) in HIV infected adults [Evaluation of an intervention (consisting of an electronic reminder device, pillboxes, and monthly telephone calls) on adherence to Highly Active Antiretroviral Therapy (HAART) in HIV infected adults]. clinicaltrials.gov/ct2/show/NCT00203853 (first received 20 September 2005)

35. NCT00224445. Boosted atazanavir and truvada given once-daily - BATON study [Boosted atazanavir and truvada given once-daily (BATON Study): a Phase 4 study of safety, efficacy & adherence in HIV infected, antiretroviral naïve subjects treated with a simple once-daily regimen]. clinicaltrials.gov/ct2/show/NCT00224445 (first received 23 September 2005)

36. NCT00234962. Study of adherence effects and clinical outcomes of kaletra based HIV antiviral therapy [Factors associated with adherence in a cohort of HIV positive subjects on a first time PI containing HAART regimen: observational study of the impact of adherence on viral load for a HAART regimen containing kaletra vs other selected PI containing HAART]. clinicaltrials.gov/ct2/show/ NCT00234962 (first received 10 October 2005)

37. NCT00273780. Highly Active Antiretroviral Therapy (HAART) adherence interventions [HAART adherence interventions in Africa: an RCT]. clinicaltrials.gov/ct2/show/NCT00273780 (first received 9 January 2006)

38. NCT00324688. Safety study of once a day ART and opiate substitute [Open-label multicenter study to assess the efficacy, the tolerability and the adherence of a once daily (QD) taken antiretroviral therapy (ART) containing the NtRTI tenofovir DF 300 mg in combination with the best suitable once a day regimen being 1 NRTI plus 1 PI or 1 NRTI plus 1 NNRTI in HIV-1-infected IVDU- patients with opiate substitution being either antiretroviral-naive or with suppressed viral load and without a history of virological failure]. clinicaltrials.gov/ct2/show/NCT00324688 (first received 11 May 2006)

39. NCT00339092. Modified directly observed therapy for improving antiretroviral therapy adherence in people with HIV [A RCT of HIV adherence case management and modified directly observed therapy]. clinicaltrials.gov/ct2/show/NCT00339092 (first received 20 June 2006)

40. NCT00408642. An enhanced adherence support programme for Highly Active Antiretroviral Therapy (HAART) [An enhanced adherence support programme for HAART]. clinicaltrials.gov/ct2/show/NCT00408642 (first received 7 December 2006)

41. NCT00602758. Effectiveness of enhanced counseling and observed therapy on antiretroviral adherence in people with HIV [ART adherence: enhanced counseling and observed therapy]. clinicaltrials.gov/ct2/show/NCT00602758 (first received 28 January 2008)

42. NCT00716040. Social-psychological intervention to improve adherence to HAART [Effectiveness of a social-psychological intervention to improve adherence to antiretroviral drug regimens for AIDS: a randomized controlled trial]. clinicaltrials.com/hiv-infections/NCT00716040/ (first received March 2008)

43. NCT00799864. A study to evaluate the pharmacokinetics, safety, tolerability, and antiviral activity of rilpivirine (TMC278) in human immunodeficiency virus infected adolescents and children aged greater than or equal to 6 years [A Phase II, open label, single arm trial to evaluate the pharmacokinetics, safety, tolerability, and antiviral activity of rilpivirine (TMC278) in antiretroviral naive HIV-1 infected adolescents and children aged >= 6 to <18 years]. clinicaltrials.gov/ct2/show/NCT00799864 (first received 1 December 2008)

44. NCT00821366. Effective AIDS treatment and support in the free state (FEATS) [Effective AIDS treatment and support in the free state (FEATS): adherence and nutritional support for effective and sustainable antiretroviral treatment in resource constrained settings]. clinicaltrials.gov/ct2/show/NCT00821366 (first received 13 January 2009)



45. NCT00959361. Evaluation of effectiveness of pharmaceutical care on the adherence of HIV-positive patients to antiretroviral therapy [Evaluation of effectiveness of pharmaceutical care on the adherence of HIV-positive patients to antiretroviral therapy - randomized clinical trial]. clinicaltrials.gov/ct2/show/NCT00959361 (first received 14 August 2009)

46. NCT01006005. Adherence-suppression-resistance relationships for atripla compared to historical antiretroviral regimens. clinical-trials.gov/ct2/show/NCT01006005 (first received 1 November 2009)

47. NCT01049568. Cell phone reminders intervention [A pilot study using cell phone interactions to improve medication adherence in adolescents who have previously failed antiretroviral therapy due to non-adherence]. clinicaltrials.gov/ct2/show/NCT01049568 (first received 14 January 2010)

48. NCT01061762. Adherence intervention for people with low-literacy [HIV treatment adherence intervention for people with poor literacy skills]. clinicaltrials.gov/ct2/show/NCT01061762 (first received 3 February 2010)

49. NCT01122186. Intervention targeting medication adherence and methamphetamine use in HIV positive men (ACE). clinicaltrials.gov/ct2/show/NCT01122186 (first received 13 May 2010)

50. NCT01347437. Improving antiretroviral medication adherence among HIV-infected youth [Improving antiretroviral medication adherence among HIV-infected youth: Phase II]. clinicaltrials.gov/ct2/show/NCT01347437 (first received 4 May 2011)

51. NCT01505660. Randomized controlled trial using patient reported outcomes and care managers to improve HIV medication adherence in routine clinical care. clinicaltrials.gov/ct2/show/NCT01505660 (first received 6 January 2012)

52. NCT01559805. Intervention to improve engagement in care among newly diagnosed HIV-positive men [Efficacy trial of a brief health enhancement intervention for newly diagnosed men]. clinicaltrials.gov/ct2/show/NCT01559805 (first received 21 March 2012)

53. NCT01641367. A5288/MULTI-OCTAVE: management using latest technologies to optimize combination therapy after viral failure [Management using the latest technologies in resource-limited settings to optimize combination therapy after viral failure (MULTI-OCTAVE)]. clinicaltrials.gov/ct2/show/NCT01641367 (first received 16 July 2012)

54. NCT01760759. Antiretroviral therapy adherence and secondary prevention of human immunodeficiency virus. clinicaltrials.gov/ct2/show/NCT01760759 (first received 4 January 2013)

55. NCT01772992. CVCTPlus: a couples-based approach to linkage to care and ARV adherence. clinicaltrials.gov/ct2/show/ NCT01772992 (first received 21 January 2013)

56. NCT02001064. Care4Today v2.0 application for improving adherence to HIV medications [Pilot study of Care4Today v.2.0 application for improving adherence to HIV medications]. clinicaltrials.gov/ct2/show/NCT02001064 (first received 4 December 2013)

57. NCT02044484. HIV clinic-based intervention to improve ART adherence and prevent HIV transmission. clinicaltrials.gov/ct2/show/ NCT02044484 (first received 24 January 2014)

58. NCT02119390. Medication adherence in human immunodeficiency virus (HIV) [Targeting enhanced adherence to medication: a pilot study in adolescents and young adults with human immunodeficiency virus (HIV)]. clinicaltrials.gov/ct2/show/NCT02119390 (first received 21 April 2014)

59. NCT02167828. Increasing social support to improve HIV care engagement and adherence in St. Petersburg, Russia [Increasing social support to improve HIV care engagement and adherence]. clinicaltrials.gov/ct2/show/NCT02167828 (first received 19 June 2014)

60. NCT02206906. Incentives to promote medication adherence among HIV-infected youth [Investigation of incentives to promote medication adherence among HIV-infected youth on antiretroviral therapy]. clinicaltrials.gov/ct2/show/NCT02206906 (first received 1 August 2014)

61. NCT02249962. Option B+: study on safety, viral suppression, and survival on second line ART [Option B+: ART safety and durability during first and subsequent pregnancies]. clinicaltrials.gov/ct2/show/NCT02249962 (first received 26 September 2014)

62. NCT02269917. Study to evaluate efficacy and safety of darunavir/cobicistat/emtricitabine/tenofovir alafenamide (D/C/F/TAF) regimen versus boosted protease inhibitor (bPI) along with emtricitabine/tenofovir disoproxil fumarate (FTC/TDF) regimen in virological-

Table 10. Ongoing studies (Continued)

ly-suppressed, HIV-1 infected participants [A Phase 3, randomized, active-controlled, open-label study to evaluate the efficacy, safety and tolerability of switching to a darunavir/cobicistat/emtricitabine/tenofovir alafenamide (D/C/F/TAF) once-daily single-tablet regimen versus continuing the current regimen consisting of a boosted protease inhibitor (bPI) combined with emtricitabine/tenofovir disoproxil fumarate (FTC/TDF) in virologically-suppressed, human immunodeficiency virus type 1 (HIV-1) infected subjects]. clinicaltrials.gov/ct2/show/NCT02269917 (first received 21 October 2014)

63. NCT02329782. Supporting treatment adherence readiness through training (START) [Controlled evaluation of the adherence readiness program for ART adherence]. clinicaltrials.gov/ct2/show/NCT02329782 (first received 1 January 2015)

64. NCT02354053. Evaluation of switching from current cART to triumeq with adherence support will enhance HIV control in vulnerable populations (TRIIADD) [A phase IV, multicentre randomized prospective open label study to evaluate whether switching from current cART to triumeq in addition to adherence support will enhance virologic control and adherence in vulnerable populations relative to adherence support alone]. clinicaltrials.gov/ct2/show/NCT02354053 (first received 3 February 2015)

65. NCT02383108. Strategy for maintenance of HIV suppression with once daily integrate inhibitor+darunavir/ritonavir in children (SMILE) [A two-arm, phase 2/3 multicentre, open-label, randomised study evaluating safety and antiviral effect of current standard antiretroviral therapy compared to once daily integrase inhibitor administered with darunavir/ritonavir (DRV/r) in HIV-1 infected, virologically suppressed paediatric participants]. clinicaltrials.gov/ct2/show/NCT02383108 (first received 9 March 2015)

66. NCT02396394. Improving ART retention and adherence in Uganda: the WiseMama study. clinicaltrials.gov/ct2/show/NCT02396394 (first received 24 March 2015)

67. NCT02464423. Improving adherence among HIV+ Rwandan youth: a TI-CBTe indigenous leader model. clinicaltrials.gov/ct2/show/ NCT02464423 (first received 8 June 2015)

68. NCT02491177. Mother and infant visit adherence and treatment engagement study (MOTIVATE!) [Maximizing adherence and retention for women and infants in the context of option B+]. clinicaltrials.gov/ct2/show/NCT02491177 (first received 7 July 2015)

69. NCT02659761. Triumeq as an integrase single tablet regimen in people with HIV who inject drugs [A prospective, single arm, openlabel 96 week observational trial of the tolerability, adherence and efficacy of a dolutegravir/abacavir/lamivudine single tablet regimen in HIV-1 antibody positive people living with HIV with a history of injection drug use switching from existing ART or starting treatment after discontinuation of ART]. clinicaltrials.gov/ct2/show/NCT02659761 (first received 20 January 2016)

70. NCT02676128. Mobile health application to improve HIV medication adherence. clinicaltrials.gov/ct2/show/NCT02676128 (first received 8 February 2016)

71. NCT02677675. Effectiveness of mobile phone technology on adherence and treatment outcomes among HIV positive patients on ART [Effectiveness of mobile phone technology in improving adherence and treatment outcomes among HIV positive patients on antiretroviral therapy (ART) in Malaysia]. clinicaltrials.gov/ct2/show/NCT02677675 (first received 9 February 2016)

72. NCT02704208. A technology-delivered peer-to-peer support ART adherence intervention for substance-using HIV+ adults. clinical-trials.gov/ct2/show/NCT02704208 (first received 9 March 2016)

73. NCT02761746. Motivational enhancement system for adherence (MESA) for youth starting ART. clinicaltrials.gov/ct2/show/ NCT02761746 (first received 4 May 2016)

74. NCT02777229. Efficacy and safety of a dolutegravir-based regimen for the initial management of HIV infected adults in resource-limited settings (NAMSAL) [A phase III randomized, open label trial to evaluate dolutegravir versus efavirenz 400 mg, both combined with tenofovir disoproxil fumarate + lamivudine for the initial management of HIV infected adults in resource-limited settings]. clinicaltrials.gov/ct2/show/NCT02777229 (first received 19 May 2016)

75. NCT02782130. Epic allies HIV ART adherence intervention [Epic allies: a gaming mobile phone application to improve engagement in care, antiretroviral uptake, and adherence among young men who have sex with men (YMSM) and trans women who have sex with men]. clinicaltrials.gov/ct2/show/NCT02782130 (first received 25 May 2016)

76. NCT02797093. Impact of ART adherence on HIV persistence and inflammation. clinicaltrials.gov/ct2/show/NCT02797093 (first received 13 June 2016)



77. NCT02797262. Measuring and monitoring adherence to ART with pill ingestible sensor system. clinicaltrials.gov/ct2/show/ NCT02797262 (first received 13 June 2016)

78. NCT02800655. Digital health feedback system for longitudinal measurement of medication adherence during anti-retroviral (ARV) therapy [A prospective single arm open label intervention study using the DHFS with HIV infected participants initiating or continuing HIV treatment]. clinicaltrials.gov/ct2/show/NCT02800655 (first received 15 June 2016)

79. NCT02878642. Adherence to dolutegravir and outcome (DOLUTECAPS) [Cohort study to assess electronic-caps defined adherence patterns - virological outcome relationship amongst HIV-1 infected subjects receiving dolutegravir-based antiretroviral therapy)]. clinicaltrials.gov/ct2/show/NCT02878642 (first received 25 August 2016)

80. NCT02888288. Integrating mental health into a HIV clinic to improve outcomes among Tanzanian youth. clinicaltrials.gov/ct2/ show/NCT02888288 (first received 5 September 2016)

81. NCT02907697. Adherence intervention for HIV-infected drug users. clinicaltrials.gov/ct2/show/NCT02907697 (first received 20 September 2016)

82. NCT02987530. National multicenter trial evaluating two treatments in patients with primary human immunodeficiency virus (HIV-1) infection (OPTIPRIM-2) [Phase III multicenter randomized trial evaluating in patients at the time of the primary HIV-1 infection, the impact on the viral reservoir of a combination including tenofovir/emtricitabine and dolutegravir or tenofovir/emtricitabine and darunavir/cobicistat]. clinicaltrials.gov/ct2/show/NCT02987530 (first received 9 December 2016)

83. NCT03076359. Traditional healers as adherence partners for persons living with HIV in rural Mozambique [Traditional healers as adherence partners for PLHIV in rural Mozambique]. clinicaltrials.gov/ct2/show/NCT03076359 (first received 10 March 2017)

84. NCT03086655. Tel-me-box: testing new, real-time strategies for monitoring HIV medication adherence in India [Tel-me-box: validating and testing a novel, low-cost, real-time monitoring device with hair level analysis among adherence-challenged patients]. clinicaltrials.gov/ct2/show/NCT03086655 (first received 22 March 2017)

85. NCT03088241. "Switch Either Near Suppression Or THOusand" (SESOTHO) [Switch to second-line versus WHO-guided standard of care for unsuppressed patients on first-line ART with viremia below 1000 copies/mL - a multicenter, parallel-group, open-label, ran-domized clinical study in rural Lesotho]. clinicaltrials.gov/ct2/show/NCT03088241 (first received 23 March 2017)

86. NCT03092115. Youth mHealth adherence intervention for HIV+ YMSM [Feasibility testing of a novel mHealth intervention to improve adherence to antiretroviral therapy among HIV+ men who have sex with men (MSM) youth]. clinicaltrials.gov/ct2/show/ NCT03092115 (first received 27 March 2017)

87. NCT03092531. Positive steps to enhance problem solving skills [Adaptive intervention strategies trial for strengthening adherence to antiretroviral HIV treatment among youth]. clinicaltrials.gov/ct2/show/NCT03092531 (first received 28 March 2017)

88. NCT03127397. Contribution of "praise messages" to HIV treatment retention and adherence among female sex workers in Ethiopia. clinicaltrials.gov/ct2/show/NCT03127397 (first received 25 April 2017)

89. NCT03149757. Connecting youth and young adults to optimize ART adherence: youTHrive efficacy trial [Connecting youth and young adults to optimize art adherence: testing the efficacy of the youth thrive intervention]. clinicaltrials.gov/ct2/show/ NCT03149757 (first received 11 May 2017)

90. NCT03195452. QDISS Stud: QD isentress as switch strategy in virologically suppressed HIV-1 infected-patient. clinicaltrials.gov/ct2/show/NCT03195452 (first received 22 June 2017)

91. NCT03198962. Use of amphetamine-type stimulants & its relationship with HIV incidence and antiretroviral adherence among MSM and TG [Use of amphetamine-type stimulants and its relationship with HIV incidence and antiretroviral adherence among Thai men who have sex with men and transgender women]. clinicaltrials.gov/ct2/show/NCT03198962 (first received 26 June 2017)

92. NCT03199027. Timing of referral to adherence clubs for antiretroviral therapy [Timing of referral to adherence clubs for antiretroviral therapy - a randomised controlled trial]. clinicaltrials.gov/ct2/show/NCT03199027 (first received 26 June 2017)

93. NCT03205566. Time to protection and adherence requirements of raltegravir with or without lamivudine in protection from HIV infection. clinicaltrials.gov/ct2/show/NCT03205566 (first received 2 July 2017)

94. NCT03256422. Antiretroviral treatment taken 4 days per week versus continuous therapy 7/7 days per week in HIV-1 infected patients [Randomized, open-label and multicentric trial evaluating the non-inferiority of antiretroviral treatment taken 4 consecutive days per week versus continuous therapy 7/7 days per week in HIV-1 infected patients with controlled viral load under antiretroviral therapy]. clinicaltrials.gov/ct2/show/NCT03256422 (first received 22 August 2017)

95. NCT03292432. Triggered escalating real-time adherence (TERA) intervention [Triggered escalating real-time adherence intervention to promote rapid HIV viral suppression among youth living with HIV failing antiretroviral therapy: the TERA study]. clinicaltrials.gov/ct2/show/NCT03292432 (first received 25 September 2017)

96. NCT03331978. A randomized controlled trial of an antiretroviral treatment adherence intervention for HIV+ African Americans. clinicaltrials.gov/ct2/show/NCT03331978 (first received 6 November 2017)

97. NCT03387397. Assessing differential adherence to medications and quality of life among people living with HIV and comorbidities. clinicaltrials.gov/ct2/show/NCT03387397 (accessed 2 January 2018)

98. NCT03394391. The effectiveness of SMS in improving antiretroviral medication adherence among adolescents living with HIV in Nigeria (STARTA) [A single-blind, randomized, parallel design study to assess the effectiveness of SMS reminders in improving art adherence among adolescents living with HIV in Nigeria (STARTA Trial-Adolescents)]. clinicaltrials.gov/ct2/show/NCT03394391 (first received 9 January 2018)

99. NCT03397576. Adherence through home education and nursing assessment, Indonesia (ATHENA-I) [A randomized controlled trial of a medication adherence intervention (ATHENA-I) to increase adherence to antiretroviral therapy among HIV-infected prisoners in Indonesia]. clinicaltrials.gov/ct2/show/NCT03397576 (first received 12 January 2018)

100. NCT03493568. Switch from dual regimens based on dolutegravir plus a reverse transcriptase inhibitor to E/C/F/TAF in virologically suppressed, HIV-1 infected patients (Be-OnE) [Open label, randomized (1:1) clinical trial to evaluate switching from dual regimens based on dolutegravir plus a reverse transcriptase inhibitor to elvitegravir/cobicistat/emtricitabine/tenofovir alafenamide in virologically suppressed, HIV-1 infected patients (Be-OnE Study)]. clinicaltrials.gov/ct2/show/NCT03493568 (first received 10 April 2018)

101. NCT03535337. Adherence interventions for HIV youth via text & cell phone - sequential multiple assignment randomized trial (SMART) [Adaptive antiretroviral therapy adherence interventions for youth living with HIV through text messaging and cell phone support embedded within the sequential multiple assignment randomized trial (SMART) design]. clinicaltrials.gov/ct2/show/ NCT03535337 (first received 24 May 2018)

102. NCT03555396. Couples ART adherence intervention for PWID in Kazakhstan [A couple-based antiretroviral therapy adherence intervention for people who inject drugs]. clinicaltrials.gov/ct2/show/NCT03555396 (first received 13 June 2018)

103. NCT03580668. Effectiveness, safety, adherence, and health-related quality of life in HIV-1 infected adults receiving bictegravir/emtricitabine/tenofovir alafenamide (BIC-STaR) [Multi-center, Canadian, non-interventional, cohort study of the effectiveness, safety, adherence, and health-related quality of life in HIV-1 infected adult patients receiving bictegravir/emtricitabine/tenofovir alafenamide (B/F/TAF)]. clinicaltrials.gov/ct2/show/NCT03580668 (first received 9 July 2018)

104. NCT03600103. Technology based community health nursing to improve combination anti-retroviral therapy (cART) adherence and virologic suppression in youth living with HIV [Technology based community health nursing to improve cART adherence and virologic suppression in youth living with HIV (TECH-N 2 CHECK-IN): a regional multi-site study]. clinicaltrials.gov/ct2/show/NCT03600103 (first received 26 July 2018)

105. NCT03618511. Interventions to improve HIV antiretroviral therapy adherence [Interventions to improve HIV antiretroviral therapy adherence in Sofala Province Mozambique]. clinicaltrials.gov/ct2/show/NCT03618511 (first received 7 August 2018)

106. NCT03665532. Youth engagement study: intervention to increase HIV treatment engagement and adherence for young people living with HIV [Unified intervention to impact HIV care continuum]. clinicaltrials.gov/ct2/show/NCT03665532 (first received 11 September 2018)



107. NCT03760458. The pharmacokinetics, safety, and tolerability of abacavir/dolutegravir/lamivudine dispersible and immediate release tablets in HIV-1-infected children less than 12 years of age [Phase I/II study of the pharmacokinetics, safety, and tolerability of abacavir/dolutegravir/lamivudine dispersible and immediate release tablets in HIV-1-infected children less than 12 years of age]. clinicaltrials.gov/ct2/show/NCT03760458 (first received 30 November 2018)

108. NCT03823261. Effects of a nurse-delivered cognitive behaviour therapy on adherence and depressive symptoms in HIV infected persons of South Korea. clinicaltrials.gov/ct2/show/NCT03823261 (first received 30 January 2019)

109. NCT03858478. Initiation of first-line antiretroviral treatment with tenofovir alafenamide - emtricitabine - bictegravir at the first clinical contact in France: trial IMEA 055 - FAST. clinicaltrials.gov/ct2/show/NCT03858478 (first received 28 February 2019)

110. NCT03978793. MyTPill: a novel strategy to monitor antiretroviral adherence among HIV+ prescription opioid users (MyTPill). clinicaltrials.gov/ct2/show/NCT03978793 (first received 7 June 2019)

111. NCT03991013. Tenofovir/lamivudine/dolutegravir combination as second line ART: a randomised controlled trial (ARTIST) [Anti-Retroviral Therapy In Second-line: Investigating Tenofovir-lamivudine-dolutegravir (ARTIST): a randomised controlled trial]. clinical-trials.gov/ct2/show/NCT03991013 (first received 19 June 2019)

112. NCT03999411. Smartphone intervention for smoking cessation and adherence to anti-retroviral therapy (ART) among people living with human immunodeficiency virus (HIV) [A novel smartphone-based intervention to support smoking cessation and adherence to antiretroviral therapy among people living with HIV: a pilot randomized clinical trial]. clinicaltrials.gov/ct2/show/NCT03999411 (first received 26 June 2019)

113. NCT04009057. Effectiveness, safety, adherence, and health-related quality of life in HIV-1 infected adults receiving bictegravir/emtricitabine/tenofovir alafenamide (B/F/TAF) [Multi-center, Israeli, non-interventional, cohort study of the effectiveness, safety, adherence, and health-related quality of life in HIV-1 infected adult patients receiving bictegravir/emtricitabine/tenofovir alafenamide (B/F/TAF)]. clinicaltrials.gov/ct2/show/NCT04009057 (first received 5 July 2019)

114. NCT04012931. A study of switching to RPV plus other ARVs in HIV-1-infected children (aged 2 to <12 years) who are virologically suppressed [A phase 2, open-label, single-arm, multicenter study to evaluate the pharmacokinetics, safety, tolerability, and effica-cy of switching to RPV plus other ARVs in HIV-1-infected children (aged 2 to <12 years) who are virologically suppressed]. clinicaltrial-s.gov/ct2/show/NCT04012931 (first received 9 July 2019)

115. NCT04024488. Group-based intervention to improve mental health and adherence among youth living with HIV in low resource settings [IMPAACT 2016 - evaluating a group-based intervention to improve mental health and antiretroviral therapy (ART) adherence among youth living with HIV in low resource settings]. clinicaltrials.gov/ct2/show/NCT04024488 (first received 18 July 2019)

116. NCT04035759. Positive affect promotion to empower optimal adherence to HIV therapy (Project APPEAL). clinicaltrials.gov/ct2/ show/NCT04035759 (first received 29 July 2019)

117. NCT04054466. Nursing counseling to the change of behavior of alcohol consumption in patients in HAART [Effectiveness of the nursing counseling to the change of behavior of alcohol consumption in patients receiving HAART]. clinicaltrials.gov/ct2/show/ NCT04054466 (first received 13 August 2019)

118. NCT04077047. ALWH: social networks, adherence and retention [Understanding and developing a network-based social support intervention to improve retention in HIV care and antiretroviral therapy adherence for adolescents living with HIV]. clinicaltrials.gov/ct2/show/NCT04077047 (first received 4 September 2019)

119. NCT04132674. Switching to a fixed dose combination of bictegravir/emtricitabine/tenofovir alafenamide (B/F/TAF) in HIV-1 infected marginalized populations who are virologically suppressed. clinicaltrials.gov/ct2/show/NCT04132674 (first received 21 October 2019)

120. NCT04222270. Caregiver peer support and ART adherence among children (CaPS) [The impact of structured caregiver peer support (CaPS) on ART adherence and viral suppression among children living with HIV in Nigeria]. clinicaltrials.gov/ct2/show/ NCT04222270 (first received 9 January 2020)

121. NTR176. The AIMS study [Theory- and evidence-based intervention to improve adherence to antiretroviral therapy among HIVinfected patients: the AIMS study]. trialsearch.who.int/Trial2.aspx?TrialID=NTR176 (first received 1 September 2005)



122. Odayar J, Malaba TR, Allerton J, Lesosky M, Myer L. Delivery of antiretroviral therapy to HIV-infected women during the postpartum period: the postpartum adherence clubs for antiretroviral therapy (PACART) trial. Contemporary Clinical Trials Communications 2019;16:100442

123. PACTR201006000222401. Children with human immunodeficiency virus (HIV) in Africa, pharmacokinetics and acceptability/adherence of simple antiretroviral regimen (CHAPAS-3) [A randomised trial to compare toxicity and pharmacokinetics of three fixeddose combination based antiretroviral regimens for treatment of human immunodeficiency virus (HIV) infected children in Africa]. pactr.samrc.ac.za/Search.aspx (first received 15 June 2010)

124. PACTR201311000641402. The TAP study [A randomised controlled Trial to explore Adherence-failure relationships in a South African antiretroviral delivery site using an electronic adherence device and sparse Pharmacokinetic sampling]. pactr.samr-c.ac.za/Search.aspx (first received 11 September 2013)

125. PACTR201405000815100. Effectiveness of an alcohol-focused intervention in improving adherence to antiretroviral therapy (ART) and HIV treatment outcomes. pactr.samrc.ac.za/Search.aspx (first received 22 April 2014)

126. PACTR201508000950148. Cell phone ringtones and antiretroviral therapy in Yaounde [Cell phone ringtones for improving adherence to antiretroviral therapy in Yaounde, Cameroon]. pactr.samrc.ac.za/Search.aspx (first received 29 November 2014)

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APPENDICES

Appendix 1. Full search strategy

The Information Specialists conducted searches on the following dates:

- First search: July 2018
- First search update: 24 January 2020
- Second search update: 22 April 2021

MEDLINE (PubMed)

(((((antiretroviral agents [Mesh] OR antiretroviral therapy, highly active [Mesh])) OR ((Antiretroviral* OR ((anti) AND (retroviral*)) OR ARV* OR ART OR "antiretroviral therapy" OR HAART OR ((highly) AND (active) AND (antiretroviral*) AND (therap*)) OR ((anti) AND (hiv)) OR ((anti)



AND (acquired immunodeficiency)) OR ((anti) AND (acquired immuno-deficiency)) OR ((anti) AND (acquired immune-deficiency)) OR ((anti) AND (acquired immun*) AND (deficienc*))))) AND (((HIV infections [MeSH] OR HIV [MeSH])) OR ((HIV OR hiv-1 OR hiv-2* OR hiv1 OR hiv2 OR hiv infect* OR human immunodeficiency virus OR human immune deficiency virus OR human immuno-deficiency virus OR human immune-deficiency virus OR ((human immun*) AND (deficiency virus)) OR acquired immunodeficiency syndromes OR acquired immune deficiency syndrome OR ((acquired immune) AND (deficiency syndrome OR acquired immuno-deficiency syndrome OR ((acquired immune) AND (deficiency syndrome OR acquired immune)) OR HIV/AIDS)))) AND (((adhere OR adherence OR adhered OR adheres OR non-adherence OR complies OR complying OR comply OR compliance OR concordance OR patient dropouts OR treatment dropouts OR treatment refusal OR "pill counting" OR "pill counting" OR "pharmacy records" OR "pharmacy recording" OR "drug counting" OR "drug counts" OR "drug count" OR dispensary OR "pharmacy recorded" OR "pharmacy recorded")) OR ((("Patient Dropouts"[Mesh]) OR ("Patient Compliance"[Mesh]) OR ("Treatment Adherence and Compliance"[Mesh]) OR ("Medication Adherence"[Mesh])))) AND (((("Viral Load"[Mesh]) OR ("Sustained Virologic Response"[Mesh]))) OR ((viral non-suppression OR viral suppression OR viral load OR virologic outcome* OR low level viraemia OR low level viremia OR viral blips OR viral failure OR viral rebound OR incomplete viral response)))) NOT ((animals [Mesh] NOT humans [Mesh])))

Embase <1946 to present

#1 *Human immunodeficiency virus/

#2 *Human immunodeficiency virus infection/

#3 (human immunodeficiency virus or human immune deficiency virus or human immuno-deficiency virus or human immune-deficiency virus).ab.

#4 (human immunodeficiency virus or human immune deficiency virus or human immuno-deficiency virus or human immune-deficiency virus).ti.

#5 (hiv-1* or hiv-2* or hiv1 or hiv2).ti. or (hiv-1* or hiv-2* or hiv1 or hiv2).ab.

#61 or 2 or 3 or 4

#7 5 or 6

8 (acquired immunodeficiency syndromes or acquired immune deficiency syndrome or acquired immuno-deficiency syndrome or acquired immuno

#9 (acquired immun* and deficiency syndrome).ti. or (acquired immun* and deficiency syndrome).ab.

#10 7 or 8 or 9

#11 (Antiretroviral* or (anti and retroviral*) or ARV* or ART or "antiretroviral therapy" or HAART or (highly and active and antiretroviral* and therap*) or (anti and hiv) or (anti and acquired immunodeficiency) or (anti and acquired immuno-deficiency) or (anti and acquired immune-deficiency) or (anti and acquired immun* and deficienc*)).mp.

#12 antiretroviral agents.mp. or antiretrovirus agent/

#13 (adhere or adherence or adhered or adheres or nonadherence or non-adherence or complies or complying or comply or compliance or concordance or patient dropouts or treatment dropouts or treatment refusal).ab. or (adhere or adherence or adhered or adheres or nonadherence or non-adherence or complies or complying or comply or compliance or concordance or patient dropouts or treatment dropouts or treatment refusal).ti.

#14 ("pill count*" or "pharmacy record*" or "drug count*" or dispensed or dispensary or "pharmacy recorded").ab.

#15 ("pill count*" or "pharmacy record*" or "drug count*" or dispensed or dispensary or "pharmacy recorded").ti.

#16 patient adherence.mp. or Patient Compliance/

#17 medication compliance/

#18 patient dropout/

#19 13 or 14 or 15 or 16 or 17 or 18

#20 11 or 12

#21 10 and 20

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#22 19 and 21

#23 ("viral non-suppression" or "viral suppression" or "viral load" or "virologic outcome*" or "low level viraemia" or "low level viremia" or "viral blips" or "viral failure" or "viral rebound" or "incomplete viral response").mp.

#24 virus load/

#25 sustained virological response.mp.

#26 23 or 24 or 25

#27 22 and 26

CINAHL (EBSCOHost)

S6 S1 AND S2 AND S3 AND S4

S5 S1 AND S2 AND S3 AND S4

S4 TX (((viral non-suppression OR viral suppression OR viral load OR virologic outcome* OR low level viraemia OR low level viremia OR viral blips OR viral failure OR viral rebound OR incomplete viral response))) OR MW Viral Load OR MW Sustained Virologic Response

S3 TX (((adhere OR adherence OR adhered OR adheres OR nonadherence OR non-adherence OR complies OR complying OR comply OR compliance OR concordance OR patient dropouts OR treatment dropouts OR treatment refusal OR "pill counts" OR "pill counting" OR "pill count" OR "pharmacy records" OR "pharmacy recording" OR "drug counting" OR "drug counts" OR "drug count" OR "drug count" OR "drug count" OR "drug count" OR "pharmacy recorded" OR "pharmacy-recorded")) OR MW Patient Dropouts OR MW Patient Compliance OR MW (Treatment Adherence and Compliance) OR MW Medication Adherence

S2 TX ((((Antiretroviral* OR ((anti) AND (retroviral*)) OR ARV* OR ART OR "antiretroviral therapy" OR HAART OR ((highly) AND (active) AND (antiretroviral*) AND (therap*)) OR ((anti) AND (hiv)) OR ((anti) AND (acquired immunodeficiency)) OR ((anti) AND (acquired immuno-deficiency)) OR ((anti) AND (acquired immune-deficiency)) OR ((anti) AND (acquired immun*) AND (deficienc*)))))) OR MW antiretroviral agents OR MW antiretroviral therapy, highly active

S1 TX ((((HIV OR hiv-1 OR hiv-2* OR hiv1 OR hiv2 OR hiv infect* OR human immunodeficiency virus OR human immune deficiency virus OR human immuno-deficiency virus OR human immune-deficiency virus OR ((human immun*) AND (deficiency virus)) OR acquired immunodeficiency syndromes OR acquired immune deficiency syndrome OR acquired immune-deficiency syndrome OR acquired immune-deficiency syndrome)) OR HIV/AIDS))) OR MW hiv OR MW hiv infection

LILACS

Search on: (anti-retroviral\$ OR antiretroviral\$ OR anti retroviral\$ OR ART OR ARV OR HAART) and (HIV\$ OR HIV/AIDS OR AIDS OR immunedeficiency OR immunedeficiency OR immune deficiency OR immuno-deficiency OR immunodeficiency OR immuno deficiency) [Words] and (adhere OR adherence OR comply OR compliance OR adherent OR compliant OR nonadherent OR non-adherent OR non-adherence OR nonadherence OR non-compliant OR non-compliance OR noncompliance OR dropout OR drop-out OR pill count OR drug count) [Words] and 2020 OR 2021 [Country, year publication]

Indexes = SCI-EXPANDED, SSCI, CPCI-S (Web of Science)

#4 #3 AND #2 AND #1

#3 TOPIC: ((((viral non-suppression OR viral suppression OR viral load OR virologic outcome* OR low level viraemia OR low level viremia OR viral blips OR viral failure OR viral rebound OR incomplete viral response OR virologic response))))

#2 TOPIC: ((((adhere OR adherence OR adhered OR adheres OR nonadherence OR non-adherence OR complies OR complying OR comply OR compliance OR concordance OR patient dropouts OR treatment dropouts OR treatment refusal OR "pill counts" OR "pill counting" OR "pill count" OR "pharmacy records" OR "pharmacy recording" OR "drug counting" OR "drug counts" OR "drug count" OR dispensed OR dispensary OR "pharmacy recorded" OR "pharmacy-recorded"))))

#1 TOPIC: ((((HIV OR hiv-1 OR hiv-2* OR hiv1 OR hiv2 OR hiv infect* OR human immunodeficiency virus OR human immune deficiency virus OR human immuno-deficiency virus OR human immune-deficiency virus OR ((human immun*) AND (deficiency virus)) OR acquired immunodeficiency syndromes OR acquired immune deficiency syndrome OR acquired immuno-deficiency syndrome OR ((acquired immun*) AND (deficiency syndrome)) OR HIV/AIDS)))) AND TOPIC: ((((Antiretroviral* OR ((anti) AND (retroviral*)) OR ARV* OR ART OR "antiretroviral therapy" OR HAART OR ((highly) AND (active) AND (antiretroviral*) OR ((anti) AND (hiv)) OR ((anti) AND (acquired immunodeficiency)) OR ((anti) AND (acquired immuno-deficiency)) OR ((anti) AND (acquired immunodeficiency)) OR ((anti) AND (acquired immunodeficiency)))))



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#1 (acquired immunodeficiency syndromes) or (acquired immune deficiency syndrome) or (acquired immuno-deficiency syndrome) or (acquired immune-deficiency syndrome) or AIDS

- #2 MeSH descriptor: [HIV] explode all trees
- #3 MeSH descriptor: [HIV Infections] explode all trees
- #4 (HIV*):ti,ab,kw
- #5 #1 or #2 or #3 or #4
- #6 Antiretroviral* or ARV* or ART or "antiretroviral therapy" or HAART
- #7 MeSH descriptor: [Anti-Retroviral Agents] explode all trees
- #8 MeSH descriptor: [Antiretroviral Therapy, Highly Active] explode all trees
- #9 #6 or #7 or #8
- #10 #5 and #9

#11 (viral non-suppression or viral suppression or viral load or virologic outcome* or low level viraemia or low level viremia or viral blips or viral failure or viral rebound or incomplete viral response)

- #12 MeSH descriptor: [Viral Load] explode all trees
- #13 MeSH descriptor: [Sustained Virologic Response] explode all trees
- #14 #11 or #12 or #13
- #15 #10 and #14

#16 adhere or adherence or adhered or adheres or nonadherence or non-adherence or complies or complying or comply or compliance or concordance or patient dropouts or treatment dropouts or treatment refusal or "pill counts" or "pill counting" or "pill count" or "pharmacy records" or "pharmacy records" or "pharmacy recorded" or "drug counting" or "drug counts" or "drug count" or dispensed or dispensary or "pharmacy recorded"

- #17 MeSH descriptor: [Patient Compliance] explode all trees
- #18 MeSH descriptor: [Patient Dropouts] explode all trees
- #19 MeSH descriptor: [Treatment Adherence and Compliance] explode all trees
- #20 MeSH descriptor: [Medication Adherence] explode all trees
- #21 #16 or #17 or #18 or #19 or #20

#22 #21 and #15

Appendix 2. Study eligibility form

Summary of the protocol	
Aim	To determine the accuracy of simple measures of adherence, including patient self-report, tablet counts, pharmacy records, electronic monitoring, or composite methods, for detecting non-sup-pressed viral load in people living with HIV
Population	HIV-positive adults, adolescents, and children who have been established on ART for \geq 6 months at the time of assessment

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(Continued)	
Target condition	The target condition is viral non-suppression. We will define this as an HIV RNA level above the lower limit of detection of the assay used within the study in question.
Index test	The index test will be measures of adherence that could be utilized in resource-limited settings, and will include: • self-report; • tablet counts; • pharmacy records or secondary database analysis, or both; • electronic monitoring; • composite measures of the above.
Reference standard	We will use a reference standard of non-suppressed viral load, as detected using nucleic acid test- ing technologies. This will be any viral load which is above the lower limit of detection of the available assay. This varies between assays, ranging from 10 copies/mL to 400 copies/mL in those which are cur- rently available. Exclude if the study: - does not report the lower limit of detection of the viral load assay used. - uses a viral load assay with a lower limit of detection > 400 copies/mL. - uses non-nucleic acid testing approaches (an example of a non-nucleic acid approach is measure- ment of HIV reverse transcriptase activity; this is a surrogate for HIV viral load measurement, but is not the reference standard). - uses point-of-care tests.
Outcomes	Our definitions for the four test accuracy categories are as follows: • true positive: the index test correctly identifies non-adherence to ART, and as such, detects a non- suppressed viral load; • true negative: the index test correctly identifies adherence to ART, and as such, detects a sup- pressed viral load; • false positive: the index test misclassifies a person as nonadherent to ART, and fails to detect a suppressed viral load; • false negative: the index test misclassifies a person as adherent to ART, and fails to detect a suppressed viral load; • false negative: the index test misclassifies a person as adherent to ART, and fails to detect a non- suppressed viral load.
Study design	Exclude: - retrospective studies or case-control study designs. There will be no restrictions on minimal quality standard, minimal sample sizes, or number of cas- es with viral non-suppression.

Appendix 3. QUADAS-2: list of signalling questions, risk of bias, and applicability

Domain	Patient selection	Index test	Reference standard	Flow and timing
Description	We will describe methods of patient selection, and the in- tended use of the adherence measure in this setting.	We will describe the mea- sure of adherence, and how the researchers interpreted it.	We will describe the method used to mea- sure viral load and the lower limit of detec- tion of the assay.	We will describe any interval between the adherence measure and the viral load measurement.
Signalling ques- tions (yes, no, unclear)	Consecutive or random sam- ple of patients?	Index test results inter- preted without knowledge of the results of reference standard?	Reference standard likely to correctly classify the target condition?	Appropriate interval between measure of adherence and viral load measurement?

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(Continued)				
used randon or consecuti No : when pa ed, for exam viously ident garding adhe Unclear: if a	Yes: if authors stated they used random patient sampling or consecutive enrolment. No: when patients were select- ed, for example, based on pre- viously identified concerns re- garding adherence. Unclear: if authors provided insufficient information.	Yes: if authors clearly reported that the measures of adherence were applied and interpreted before the viral load result was available. No: if authors reported that the measures were applied or interpreted after the viral load was available. Unclear: if authors provided insufficient information.	Yes: if authors clearly reported that a labo- ratory reference test was used at a manu- facturer recommend- ed threshold of lower limit of detection, and this was < 400 copies/ mL. No: if authors report- ed application of a post hoc threshold. Unclear: if authors provided insufficient information.	 Yes: if the measure of adherence and the measure of viral load were made on the same day. No: if time period between measure of adherence and viral load was not made on the same day. Unclear: if authors provided insufficient information.
	Did the study avoid inappro- priate exclusions?	Prespecified threshold used?	Reference standard results interpreted without knowledge of	Did all patients re- ceive a viral load, using the same as-
	Yes: if there were no inappro- priate exclusions. No: if there was evidence that	Yes: if authors reported an a priori threshold value (or values) for adherence.	the index test? Yes: if authors report- ed that viral loads	 say, and were all included in the analysis? Yes: if authors reported that all patients received a viral load using the same assay and all were included in the analysis.
	authors inappropriately ex- cluded certain patients, e.g. those deemed to have limited ability to use electronic mon- itoring devices, or excluded those with literacy concerns if self-report measures were to be self-administered. Unclear: if authors provided insufficient information.	No: if authors determined threshold values post hoc. Unclear: if authors provid- ed insufficient information.	were measured and recorded without a priori knowledge of the measure of adher- ence result. No: if authors report- ed that viral load was measured or recorded with knowledge of the measure of adherence result. Unclear: if authors	
				No: if only a selec- tion of those with ad- herence measures have viral load mea- sures, or different as- says were used.
			provided insufficient information.	Unclear: if authors provided insufficient information.
Risk of bias (high, low, un- clear)	If we answer both signalling ofIf we answer both signalling of	juestions for a domain 'yes', the juestions for a domain 'no', the juestions for a domain 'unclear questions for a domain differen the rationale for this judgemen	n we will judge risk of bias ', then we will judge the ris tly, the authors will discus	as high. sk of bias as unclear. ss this further to reach a
Applicability concerns (high, low, unclear)	Are there concerns that the in- cluded patients do not match the review question?	Are there concerns that the index test, its conduct, or interpretation differs from the review question?	_	-
	High: if some but not all in- cluded patients were concur- rently receiving interventions to improve their adherence, rather than the same standard of care, and these groups can-	the review question? High: if the measure of ad- herence was not truly ap- plicable in a resource-limit- ed setting, e.g. requiring ad- ditional remote information		
	not be separated. s for antiretroviral adherence in peop	infrastructure or analysis.		2

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

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(Continued)	Low: if all patients were re-	Low: if the measure of ad-
	ceiving the same standard of care.	herence could feasibly be applied in a resource-limit-
	care.	ed setting.
	Unclear: if there was insuffi-	U U U U U U U U U U U U U U U U U U U
	cient information to make a	Unclear: if there was insuf-
	judgement.	ficient information to make
		a judgement.

Appendix 4. QUADAS-2: list of signalling questions, risk of bias, and applicability (final version; with changes from previous version highlighted)

Domain	Patient selection	Index test	Reference standard	Flow and timing
Description	We will describe methods of pa- tient selection, and the intended use of the adherence measure in this setting.	We will describe the measure of adher- ence, and how the re- searchers interpreted it.	We will describe the method used to mea- sure viral load and the lower limit of detection of the assay.	We will describe any interval between the adherence measure and the viral load measurement.
Signalling ques- tions (yes, no, un- clear)	Consecutive or random sample of patients? Yes: if authors stated they used random patient sampling or con-	Index test results inter- preted without knowl- edge of the results of reference standard?	Reference standard likely to correctly clas- sify the target condi- tion?	Appropriate interval between measure of adherence and viral load measurement?
	secutive enrolment. No: when patients were selected, for example, based on previously identified concerns regarding ad- herence. Unclear: if authors provided in- sufficient information.	Yes: if authors clear- ly reported that the measures of adherence were applied and inter- preted before the viral load result was avail- able. No: if authors reported that the measures were applied or interpreted after the viral load was available. Unclear: if authors pro- vided insufficient infor- mation.	 Yes: if authors clearly reported that a labora- tory reference test was used at a manufacturer recommended thresh- old of lower limit of de- tection, and this was < 400 copies/mL. No: if authors report- ed application of a post hoc threshold, or use of more than one viral as- say. Unclear: if authors pro- vided insufficient infor- mation, for example, not stating which assay was used, and the low- er limit of detection for this assay. 	 Yes: if the measure of adherence and the measure of viral load were made on the same day. No: if time period between measure of adherence and viral load was not made on the same day (this is an exclusion criterion). Unclear: if authors provided insufficient information.
<i>ed?</i> Yes: no evide ent were rec adherent par No: evidence	Was case control design avoid- ed? Yes: no evidence that non-adher-			Did all patients re- ceive a viral load, using the same as-
	No: evidence that non-adher- ent were recruited different to adherent patients No: evidence that people who were poorly adherent were re-			<i>say?</i> Yes: if authors re- ported that all pa- tients received a viral

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(Continued)	cruited in a different way to those who were adherent			load using the same assay
	Unclear: not stated			No: if only a selec- tion of those with ad- herence measures have viral load mea- sures, or different as- says were used.
				Unclear: if authors provided insufficient information.
	Did the study avoid inappropri- ate exclusions?	Prespecified threshold used?	Reference standard re- sults interpreted with-	Were all included in the analysis?
	Yes: if there were no inappropri- ate exclusions.	Yes: if authors reported an a priori threshold	out knowledge of the index test?	Yes: if authors re- ported that >= 90%
	No: if there was evidence that authors excluded certain patients with factors that might influence accuracy of measurement of adherence. Examples of inappropriate exclusions:	value (or values) for ad- herence. No: if authors deter- mined threshold values post hoc.	Yes: if authors reported that viral loads were measured and recorded without a priori knowledge of the measure of adherence result.	were included in analysis. No: if only a selec- tion of those with ad- herence measures have viral load mea-
	 people with limited ability to use electronic monitoring de- vices, 	Unclear: if authors pro- vided insufficient infor- mation.	No: if authors reported that viral load was mea- sured or recorded with knowledge of the mea-	surements, or miss- ing data for > 10%. Unclear: if authors provided insufficient
	 literacy concerns if self-report measures were to be self-admin- istered 		sure of adherence re- sult. Unclear: if authors pro-	information.
	\cdot comorbidities and comedication		vided insufficient infor- mation.	
	 Viral non-suppression at base- line 			
	Unclear: if authors provided in- sufficient information.			
Risk of bias	Any answer no = high risk of bias	Any answer no = high risk of bias	Any answer no = high risk of bias	Any answer no = high risk of bias
(high, low, un- clear)	2 answers unclear = unclear	2 answers unclear = un-	2 answers unclear = un-	2 answers unclear =
	≥ 2 answers yes = low risk of bias	clear	clear	unclear
	Any answer = no, assume high risk of bias; however, author judgement may be applied.	≥ 1 answers yes = low risk of bias	≥ 1 answers yes = low risk of bias	≥ 2 answers yes = low risk of bias
]0	Any answer = no, as- sume high risk of bias; however, author judge- ment may be applied.	Any answer = no, as- sume high risk of bias; however, author judge- ment may be applied.	Any answer = no, as- sume high risk of bias; however, au- thor judgement may be applied.
Applicability concerns (high, low, un- clear)	Are there concerns that the in- cluded patients do not match the review question?	Are there concerns that the index test, its con- duct, or interpretation differs from the review question?	Are there concerns that the target condition as defined by the refer- ence standard does not	_

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(Continued)	 High: if the population is highly selected. Or if some but not all included patients were concurrently receiving interventions to improve their adherence, rather than the same standard of care, and these groups cannot be separated. Low: a general population. Or if all patients were receiving the same standard of care. Unclear: if there was insufficient information to make a judgement. 	 High: if the measure of adherence was not truly applicable in a resource-limited setting. Examples: requiring additional remote information infrastructure or analysis. Self-report of greater than 8 questions, or deemed to be of excessive complexity. MEMS that cannot be used at point of care. Low: if the measure of adherence could feasibly be applied in a resource-limited setting. Unclear: if there was insufficient information to make a judgement. 	<pre>match the review ques- tion? Low: if the viral load as- say used was clearly re- ported. Unclear: If the viral load assay used was not reported.</pre>
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Appendix 5. Additional summary of findings tables

Summary of findings table 2. Diagnostic accuracy of self-report questionnaires for the detection of viral non-suppression

Question	What is the diagnostic accuracy of	f self-report questionnaire for detecting viral non-suppression?
Population	HIV-positive children and adults who have been established on ART for longer than six months at the time of assessment	
Index test	Self-report using questionnaires	
Target condition	Viral non-suppression	
Reference stan- dard	Non-suppressed viral load, as detected by nucleic acid testing technologies, ranging from 10 copies to 400 copies/mL	
Action/clinical im- plications	<i>Low sensitivity:</i> failures to detect non-adherence.	Low specificity: adherent people incorrectly identified.
pications	Consequences of false negatives:	Consequences of false positives: increased viral load monitoring, pa- tient inconvenience
	disease progression, resistance, transmission	Of lesser clinical importance
	Of greater clinical importance	
Quantity of evi- dence	26 studies (Avong 2015; Bajunirwe 2009; Coker 2015; Duarte 2015; Ekstrand 2010; El-Khatib 2010; Fokan Haberer 2011; Landes 2021; Mbengue 2019; McMahon 2013; Meya 2009; Mogosetsi 2018; Navarro 2014; G 2006; Orrell 2017; Paolillo 2017; Parker 2017; Pasquau 2018; Phillips 2019; Pulido 2009; Sangeda 2014; S 2010; Segeral 2018; Tabb 2018; Zoufaly 2013)	
	N = 11607 participants (9703 analys	ed)

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

(Continued)	
(continued)	Total with target condition (viral non-suppression) = 5640
	Study design: 3 RCTs, 14 cohorts, 9 cross-sectional studies
	Population: 20 in adults, 4 in children, 2 in mixed population
	Setting: 11 low-income, 3 lower-middle-income, 5 upper-middle-income, 6 high-income, 1 mixed
	Adherence threshold: 20 used 100% or a binary threshold (adherent/non-adherent); 4 used 95%; 1 used 90%; 3 used 80%; 1 used 75%; 1 used 60%
	Viral load threshold: 2 used 40 copies/mL; 10 used 50 copies/mL; 4 used 200 copies/mL; 1 used 250 copies/mL; 12 used 400 copies/mL
Limitations in the ev	idence
Risk of bias	Participants: 14 studies low risk; 7 studies unclear risk; 5 studies high risk
	Index test: 9 studies low risk; 17 studies unclear risk
	Reference standard: 19 studies low risk; 7 studies unclear risk
	Flow and timing: 11 studies low risk; 2 studies unclear risk; 13 studies high risk
Applicability con-	Participants: 15 studies low concern; 8 studies unclear concern; 3 studies high concern
cerns	Index test: 23 studies low concern; 3 studies unclear concern
	Reference standard: 19 studies low concern; 7 studies unclear concern
Findings	

	Studies and participants	Sensitivity range (95% CI range)	Specificity range (95% CI range)
Main analyses			
All participants	25 studies	10% to 85%	10% to 99%
Various thresholds*	N = 9211	(0% to 91%)	(7% to 100%)
Subgroup analysis b	y adherence threshold		
All participants	4 studies	18% to 85%	45% to 93%
95% adherence threshold	N = 1007	(9% to 91%)	(32% to 96%)
All participants	21 studies	10% to 74%	10% to 99%
100% adherence threshold	N = 8204	(0% to 87%)	(7% to 100%)
Subgroup analysis b	y number of questions in the ques	stionnaire	
Single question	12 studies	10% to 74%	10% to 96%
	N = 4997	(5% to 87%)	(7% to 98%)
2 to 4 questions	8 studies	13% to 69%	70% to 99%
	N = 1922	(0% to 82%)	(60% to 100%)

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



(Continued)			
5 or more questions	5 studies	21% to 85%	54% to 84%
	N = 2292	(14% to 91%)	(49% to 86%)
Subgroup analysis b	y population		
Children	4 studies	15% to 67%	37% to 96%
	N = 804	(10% to 80%)	(28% to 98%)
Adults	19 studies	10% to 85%	10% to 99%
	N = 8011	(5% to 91%)	(7% to 100%)
Subgroup analysis b	y viral load		
40 to 50 copies/mL	11 studies	13% to 69%	10% to 93%
	N = 2290	(0% to 82%)	(7% to 96%)
200 to 400 copies/	13 studies	10% to 85%	23% to 99%
mL	N = 6664	(5% to 91%)	(16% to 100%)
Subgroup analysis b	y setting		
Low-income	11 studies	13% to 85%	10% to 99%
	N = 4135	(5% to 91%)	(7% to 100%)
Lower-middle-in-	3 studies	18% to 67%	37% to 93%
come	N = 576	(9% to 80%	(28% to 96%)
Upper-middle-in-	5 studies	10% to 69%	12% to 96%
come	N = 1141	(5% to 82%)	(8% to 97%)
High-income	5 studies	19% to 61%	69% to 93%
	N = 2702	(12% to 84%)	(64% to 96%)
Additional analysis			
All participants	3 studies	8% to 41%	81% to 97%
80% adherence threshold	N = 1527	(4% to 56%)	(68% to 99%)

Summary of findings table 3. Diagnostic accuracy of VAS for the detection of viral non-suppression

Question: what is the diagnostic accuracy of VAS for detecting viral non-suppression?		
Population	HIV-positive children and adults who have been established on ART for longer than six months at the time of assessment	

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



(Continued)			
Index test	Self-report using VAS scale		
Target condition	Viral non-suppression		
Reference stan- dard	Non-suppressed viral load, as detected by nucleic acid testing technologies, ranging from 10 copies to 400 copies/mL		
Action/clinical im- plications	Low sensitivity:failures to detect non-adherence.Low specificity:adherent people incorrectly identified.Consequences of false negatives: disease progression, resistance, transmissionConsequences of false positives: increased viral load monitoring, pa- tient inconvenienceOf greater clinical importanceOf lesser clinical importance		
Quantity of evi- dence	 14 studies (Cerutti 2016; Cohen 2012; Dziva 2017; Ekstrand 2010; Gill 2010; Haberer 2011; Jiamsakul 2014; Labhardt 2012; McMahon 2013; Mbengue 2019; Meya 2009; Nelson 2010; Sangeda 2014; Segeral 2018) N = 5852 participants (5151 analysed) Total with target condition (viral non-suppression) = 2499 Study design: 2 RCTs, 1 prospective clinical trial, 7 cohorts, 4 cross-sectional studies Population: 11 studies in adults, 2 in children and 1 in mixed population Setting: 7 low-income, 3 lower-middle-income, 1 upper-middle-income, 3 mixed settings Adherence threshold: 2 used 100%; 11 used 95%; 3 used 90%; 1 used 80%; 1 used a binary threshold (adherent/non-adherent) Viral load threshold: 1 used 40 copies/mL; 3 used 50 copies/mL; 1 used 80 copies/mL; 1 used 200 copies/mL; 7 used 400 copies/mL 		
Limitations in the e	vidence		
Risk of bias	Participants: 7 studies low risk; 4 studies unclear risk; 3 studies high risk Index test: 4 studies low risk; 10 studies unclear risk Reference standard: 9 studies low risk; 5 studies unclear risk Flow and timing: 4 studies low risk; 2 studies unclear risk; 8 studies high risk		
Applicability con- cerns	Participants: 9 studies low concern; 2 studies unclear concern; 3 studies high concern Index test: 10 studies low concern; 4 studies unclear concern Reference standard: 9 studies low concern; 5 studies unclear concern		
Findings			
Main analysis			
	Studies and participants	Sensitivity range (95% CI range)	Specificity range (95% CI range)
All participants	11 studies	0% to 58%	55% to 100%
95% adherence threshold	N = 4235	(0% to 85%)	(46% to 100%)

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)

(Continued)

Subgroup analysis by population

5 1 9	, , , ,		
Children	2 studies	26% to 30%	60% to 89%
	N = 239	(12% to 39%)	(46% to 94%)
Adults	8 studies	0% to 58%	57% to 96%
	N = 3904	(0% to 85%)	(51% to 100%)
Subgroup analysis b	y viral load		
40 to 100 copies/mL	6 studies	5% to 45%	55% to 100%
	N = 3591	(3% to 55%)	(46% to 100%)
200 to 400 copies/	5 studies	0% to 58%	72% to 96%
mL	N = 644	(0% to 85%)	(64% to 100%)
Subgroup analysis b	y setting		
Low-income	5 studies	18% to 35%	60% to 100%
	N = 663	(9% to 54%)	(46% to 100%)
Lower-middle-in-	3 studies	0% to 45%	55% to 96%
come	N = 1631	(0% to 55%)	(53% to 100%)
Additional analysis			
All participants	3 studies	3% to 24%	88% to 95%
90% adherence threshold	N = 582	(0% to 39%)	(80% to 98%)
All participants	1 study	0% to 35%	69% to 100%
80% adherence threshold	N = 73	(0% to 49%)	(59% to 100%)

Summary of findings table 4. Diagnostic accuracy of tablet counts for the detection of viral non-suppression

Question: what is the diagnostic accuracy of tablet counts for detecting viral non-suppression?		
Population	HIV-positive children and adults who have been established on ART for longer than six months at the time of assessment	
Index test	Tablet counts: adherence percentage based on expected versus actual tablets taken over dispensing period	
Target condition	Viral non-suppression	
Reference stan- dard	Non-suppressed viral load, as detected by nucleic acid testing technologies, ranging from 10 copies to 400 copies/mL	

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



Continued) Action/clinical im-	Low sensitivity: failures to detect	<i>Low specificity:</i> adherent people ir	ncorrectly identified.	
plications	non-adherence.	Consequences of false positives: in	-	
	Consequences of false negatives: disease progression, resistance,	tient inconvenience	0,1	
	transmission	Of lesser clinical importance		
	Of greater clinical importance			
Quantity of evi- dence	13 studies included (Apisarnthanarak 2010; Bonjoch 2006; Cerutti 2016; Coker 2015; Davies 2008; Gill 2010; Haberer 2011; Kitkungvan 2008; Mariana 2018; Moosa 2019; Okonji 2012; Orrell 2017; Sangeda 2014)			
	N = 4899 participants (3808 analysed)			
	Total with target condition (viral non-suppression) = 2335			
	Study design: 1 RCT, 2 sub-analyses o	of RCT(s), 7 cohorts, 3 cross-sectional s	tudies	
	Population: 9 studies in adults, 2 in c	children and 2 in mixed population		
	Setting: 4 low-income, 4 lower-midd	lle-income, 4 upper-middle-income an	d 1 high-income	
	Adherence threshold: 1 used 100%, 3 70%, 1 used 65%, 1 used 60%, 2 used	12 used 95%; 3 used 90%; 1 used 85%; d 55% and 1 used 50%	2 used 80%; 3 used 75%, 1 used	
	Viral load threshold: 2 used 40 copies/mL; 4 used 50 copies/mL; 1 used 80 copies/mL; 6 used 400 copies/mL			
Limitations in the ev	vidence			
Risk of bias	Participants: 6 studies low risk; 4 studies unclear risk; 3 studies high risk			
	Index test: 2 studies low risk; 9 studies unclear risk; 2 studies high risk			
	Reference standard: 8 studies low risk; 5 studies unclear risk			
	Flow and timing: 2 studies low risk; 3	3 studies unclear risk; 8 studies high ris	k	
Applicability con-	Participants: 8 studies low concern; 2 studies unclear concern; 3 studies high concern			
cerns	Index test: 10 studies low concern; 3 studies unclear concern			
	Reference standard: 9 studies low concern; 4 studies unclear concern			
Findings				
Main analysis				
	Studies and participants	Sensitivity range (95% CI range)	Specificity range (95% CI range	
All participants	12 studies			
95% adherence hreshold	N = 3466			
Subgroup analysis b	y population			
Children	1 study	35%	51%	
	N = 73	(15% to 59%)	(37% to 65%)	

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



(Continued)				
(continuea)	N = 3016	(0% to 91%)	(2% to 100%)	
Mixed	2 studies	20% to 100%	11% to 79%	
	N = 377	(10 % to 100%)	(6% to 85%)	
Subgroup analysis l	by viral load			
40 to 80 copies/mL	7 studies	0% to 100%	11% to 99%	
	N = 2299	(0% to 100%)	(2% to 100%)	
400 copies/mL	5 studies	9% to 78%	17% to 95%	
	N = 1167	(0% to 88%)	(10% to 98%)	
Subgroup analysis l	by setting			
Low-income	4 studies	27% to 78%	17% to 87%	
	N = 942	(15% to 88%)	(10% to 91%)	
Lower-middle-in-	4 studies	0% to 100%	77% to 99%	
come	N = 1692	(0% to 100%)	(72% to 100%)	
Upper-middle-in-	3 studies	9% to 76%	11% to 95%	
come	N = 610	(0% to 91%)	(6% to 98%)	
High-income	1 study	67%	5%	
	N = 222	(55% to 78%)	(2% to 10%)	
Additional analysis				
All participants	2 studies	0% to 35%	69% to 100%	
80% adherence threshold	N = 235	(0% to 49%)	(59% to 100%)	

Summary of findings table 5. Diagnostic accuracy of pharmacy records or secondary databases for the detection of viral non-suppression

Question: what is the diagnostic accuracy of pharmacy records or secondary databases for detecting viral non-suppression?		
Population	HIV-positive children and adults who have been established on ART for longer than six months at the time of assessment	
Index test	Pharmacy records or secondary databases	
Target condition	Viral non-suppression	
Reference standard	Non-suppressed viral load, as detected by nucleic acid testing technologies, ranging from 10 copies to 400 copies/mL	

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Continued) Action/clinical im-	<i>Low sensitivity:</i> failures to detect	Low specificity: adherent people ir	acorrectly identified		
plications	non-adherence.	Consequences of false positives: in	-		
	Consequences of false negatives: disease progression, resistance, transmission	tient inconvenience	creased viral load monitoring, pa-		
		Of lesser clinical importance			
	Of greater clinical importance				
Quantity of evi-	7 studies (Anude 2013; Hassan 2014	; McMahon 2013; Messou 2011; Navarr	ro 2014; Orrell 2017; Sangeda 2014)		
dence	N = 2882 (2449 analysed)				
	Total with target condition (viral no	n-suppression) = 1298			
	Study design: 6 cohorts, 1 cross-sec	tional study			
	Population: 5 adults, 2 mixed				
	Setting: 4 low-income; 1 lower-mide	dle-income; 1 upper-middle-income; 1	. high income		
	Adherence threshold: 2 used 100%, 70%, 2 used 65%, 1 used 60%, 1 use	6 used 95%; 2 used 90%; 1 used 85%; d 55% and 2 used 50%	2 used 80%; 1 used 75%, 1 used		
	Viral load threshold: 1 used 40 copies/mL; 1 used 50 copies/mL; 1 used 200 copies/mL; 1 used 300 copies/mL; 3 used 400 copies/mL				
Limitations in the ev	idence				
Risk of bias	Patient selection: 3 studies low risk, 1 study unclear risk, 3 studies high risk				
	Index test: 2 studies low risk, 5 studies unclear risk				
	Reference standard: 4 studies low risk, 3 studies unclear risk				
	Flow and timing: 1 study low risk, 6 studies high risk				
Applicability con-	Patient selection: 3 studies low concerns, 3 studies unclear concerns, 1 study high concerns				
cerns	Index test: 4 studies low concerns, 3 studies unclear concerns				
	Reference standard: 4 studies low concerns, 3 studies unclear concerns				
Findings					
	Studies and participants	Sensitivity range (95% CI range)	Specificity range (95% CI range)		
Main analysis					
All participants	6 studies	17% to 88%	9% to 95%		
≥ 95% adherence threshold	N = 2254	(11% to 92%)	(5% to 97%)		
Subgroup analysis by	population				
Adults	4 studies	17% to 88%	46% to 95%		
≥95% adherence threshold	N = 1893	(11% to 92%)	(42% to 97%)		
Mixed	2 studies	34% to 35%	9% to 85%		

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(Continued) ≥ 95% adherence threshold	N = 402	(20% to 51%)	(5% to 90%)
Subgroup analysis by	viral load		
All participants	1 study	34%	9%
40 copies/mL	N = 178	(20 to 51%)	(5% to 15%)
≥95% adherence threshold			
All participants	5 studies	17% to 88%	46% to 95%
200-400 copies/mL	N = 2076	(11% to 92%)	(42% to 97%)
≥95% adherence threshold			
Subgroup by setting			
Low-income	4 studies	24% to 88%	46% to 85%
≥95% adherence threshold	N = 1485	(11% to 92%)	(42% to 90%)
Lower-middle-in-	1 study	17%	95%
come	N = 591	(11% to 25%)	(92% to 97%)
≥ 95% adherence threshold			
Upper-middle-in-	1 study	34%	9%
come	N = 178	(20% to 51%)	(5% to 15%)
≥ 95% adherence threshold			
Additional analysis			
All participants	3 studies	25% to 82%	73% to 88%
≥80% adherence threshold	N = 1211	(15% to 91%)	(60% to 93%)

Summary of findings table 6. Diagnostic accuracy of electronic monitoring devices for the detection of viral non-suppression

Question: what is the diagnostic accuracy of electronic monitoring devices for detecting viral non-suppression?		
PopulationHIV-positive children and adults who have been established on ART for longer than six of assessment		
Index test	Electronic monitoring devices	
Target condition	Viral non-suppression	

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



(Continued)				
Reference standard	Non-suppressed viral load, as detect copies/mL	ted by nucleic acid testing technologi	es, ranging from 10 copies to 400	
Action/clinical impli- cations	<i>Low sensitivity:</i> failures to detect non-adherence.	Low specificity: adherent people in	-	
	Consequences of false negatives: disease progression, resistance, transmission	Consequences of false positives: increased viral load monitoring, pa- tient inconvenience		
		Of lesser clinical importance		
	Of greater clinical importance			
Quantity of evidence	5 studies (Evans 2016; Farley 2003; G	ill 2010; Haberer 2011; Orrell 2017)		
	N = 475 (392 analysed)			
	Total with target condition (viral nor	n-suppression) = 92		
	Study design: 5 cohort studies			
	Population: 2 in adults, 2 in children	, 1 in mixed population		
	Adherence threshold: 3 studies used	95%; 4 studies used 80%		
	Viral load threshold: 1 study used 40 copies/mL, 1 study used 50 copies/mL; 3 studies used 400 copies/mL			
	Setting: 1 low-income, 1 lower-midd	lle-income, 2 upper-middle-income co	ountry, 1 high-income	
Limitations in the evid	ence			
Risk of bias	Patient selection: 3 studies low risk, 2 studies unclear risk			
	Index test: 5 studies unclear risk			
	Reference standard: 5 studies low risk			
	Flow and timing: 2 studies low risk, 3	3 studies high risk		
Applicability con-	Patient selection: 3 studies low concerns, 1 study unclear concerns, 1 study high concerns			
cerns	Index test: 1 study low concerns, 3 studies unclear concerns, 1 study high concerns			
	Reference standard: 5 studies low concerns			
Findings				
	Studies and participants	Sensitivity range (95% CI range)	Specificity range (95% Cl range)	
Main analysis				
All participants	3 studies	60% to 88%	27% to 67%	
95% adherence threshold	N = 186	(36% to 100%)	(11% to 80%)	
Subgroup analysis by p	opulation			
Children	1 study	60%	67%	
95% adherence threshold	N = 72	(36% to 81%)	(53% to 80%)	
Accuracy of measures for a	ntiretroviral adherence in people living w	rith HIV (Review)	298	

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(Continued)			
Adults	2 studies	74% to 88%	27% to 53%
95% adherence threshold	N = 114	(47% to 100%)	(11% to 66%)
Subgroup analysis by v	iral load		
All participants	1 study	60%	67%
50 copies/mL	N = 72	(36 % to 81%)	(53% to 80%)
95% adherence threshold			
All participants	2 studies	74% to 88%	27% to 53%
400 copies/mL	N = 114	(47% to 100%)	(11% to 66%)
95% adherence threshold			
Subgroup by setting			
Low-income	1 study	60%	67%
95% adherence threshold	N = 72	(36% to 81%)	(53% to 80%)
Lower-middle-income	1 study	88%	53%
95% adherence threshold	N = 65	(47% to 100%)	(39% to 66%)
Upper-middle-income	1 study	74%	27%
95% adherence threshold	N = 49	(54% to 89%)	(11% to 50%)
Additional analysis			
All participants	4 studies	24% to 89%	7% to 96%
80% adherence threshold	N = 327	(15% to 100%)	(3% to 100%)

Summary of findings table 7. Diagnostic accuracy of composite measures of adherence for the detection of viral non-suppression

he diagnostic accuracy of composite measures of adherence for detecting viral non-suppression?	
Population HIV-positive children and adults who have been established on ART for longer than six months at the time of assessment	
Composite measures	
Viral non-suppression	

Accuracy of measures for antiretroviral adherence in people living with HIV (Review)



^{Continued)} Reference stan- dard	Non-suppressed viral load, as copies/mL	detected by nucleic acid testing techno	ologies, ranging from 10 copies to 400								
Action/clinical im-	Low sensitivity: failures to detect non-adherence.										
olications	Consequences of false negatives: disease progression, resistance, transmission										
	Of greater clinical importance										
	Low specificity: adherent people incorrectly identified.										
	Consequences of false positive	es: increased viral load monitoring, pat	ient inconvenience								
	Of lesser clinical importance										
Quantity of evi- dence	9 studies (Jayaweera 2003; Mb 2010; Segeral 2010; Spire 2008	engue 2019; McMahon 2013; Mutwa 20)	014; Orrell 2003; Ortega 2004; Parienti								
	N = 1901 (1513 included in the analysis)										
	Total with target condition (viral non-suppression) = 858										
	Study design: 6 cohort studies, 3 cross-sectional studies										
	Population: 7 in adults, 1 in children, 1 not reported										
	Adherence threshold: 1 used 100%; 3 used 95%; 1 used 90%; 1 used 80%; 1 used 70%; 4 used a binary thresh- old (adherent/non-adherent)										
	Viral load threshold: 2 studies used 40 copies/mL, 1 study used 50 copies/mL; 1 study used 200 copies/mL; 6 studies used 400 copies/mL										
	Setting: 4 low-income, 2 upper	-middle-income country, 3 high-incon	ne								
Limitation in the evi	dence										
Risk of bias	Patient selection: 4 studies lov	v risk, 4 studies unclear risk, 1 study hi	gh risk								
	Index test: 2 studies low risk, 7 studies unclear risk										
	Reference standard: 5 studies low risk, 4 studies unclear risk										
	Flow and timing: 3 studies low risk, 2 studies unclear risk, 4 studies high risk										
Applicability con-	Patient selection: 3 studies low concerns, 4 studies unclear concerns, 2 studies high concerns										
cerns	Index test: 5 studies low concerns, 4 studies unclear concerns										
	Reference standard: 5 studies low concerns, 4 studies unclear concerns										
Findings											
	Studies and participants	Sensitivity (95% CI range)	Specificity (95% CI range)								
Main analysis											
All participants	9 studies	10% to 100%	49% to 100%								
/arious thresholds*	N = 1513	(4% to 100%)	(35% to 100%)								
Subgroup by adhere	nce threshold										
100% adherence	6 studies	10% to 85%	56% to 100%								

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(Continued)			
	N = 1095	(4% to 94%)	(45% to 100%)
95% adherence	3 studies	32% to 100%	49% to 84%
	N = 418	(15% to 100%	(35% to 91%)
Subgroup analysis b	y population		
Children	1 study	32%	84%
Different thresh- olds*	N = 104	(15% to 54%)	(74% to 91%)
Adults	7 studies	10% to 100%	49% to 97%
Different thresh- olds*	N = 1390	(4% to 100%)	(35% to 98%)
Subgroup analysis b	y viral load		
All participants	3 studies	10% to 100%	49% to 97%%
40 to 50 copies/mL	N = 522	(4 % to 100%)	(35% to 98%)
95% adherence threshold			
All participants	7 studies	18% to 100%	57% to 100%
200 to 400 copies/ mL	N = 1063	(9% to 100%)	(31% to 100%)
Different thresh- olds*			
Subgroup by setting			
Low-income	4 studies	10% to 50%	68% to 97%
Different thresh- olds*	N = 881	(4% to 68%)	(59% to 98%)
Upper-middle-in-	2 studies	18% to 60%	57% to 87%
come Different thresh- olds*	N = 405	(9% to 70%)	(49% to 92%)
High-income	2 studies	69% to 100%	49% to 100%
Different thresh- olds*	N = 227	(39% to 100%)	(35% to 100%)

Appendix 6. Self-report questionnaire subgroup analysis

Figure 14; Figure 15; Figure 16; Figure 17; Figure 18; Figure 19

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Figure 14. Self-report questionnaires; various adherence thresholds* [subgroup by adherence threshold] *cut-off used was either \ge 95% or 100%

[Subgroup analysis by adherence threshold] Self-report questionnaires; threshold: ≥ 100% adherence

Study TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)

McMahon 2013	25	105	9	31	0.74 [0.56, 0.87]	0.23 [0.16, 0.31]	
Phillips 2019	34	26	15	62	0.69 [0.55, 0.82]	0.70 [0.60, 0.80]	
Fokam 2017	28	64	14	38	0.67 [0.50, 0.80]	0.37 [0.28, 0.47]	
Parker 2017	106	458	68	1017	0.61 [0.53, 0.68]	0.69 [0.67, 0.71]	-+ I
Pulido 2009	9	27	6	79	0.60 [0.32, 0.84]	0.75 [0.65, 0.82]	— • —
Coker 2015	45	2	44	224	0.51 [0.40, 0.61]	0.99 [0.97, 1.00]	
Pasquau 2018	6	51	7	125	0.46 [0.19, 0.75]	0.71 [0.64, 0.78]	
Bajunirwe 2009	10	16	18	131	0.36 [0.19, 0.56]	0.89 [0.83, 0.94]	
Tabb 2018	31	41	60	95	0.34 [0.24, 0.45]	0.70 [0.61, 0.77]	
Mbengue 2019	13	24	36	90	0.27 [0.15, 0.41]	0.79 [0.70, 0.86]	
Oette 2006	13	11	48	136	0.21 [0.12, 0.34]	0.93 [0.87, 0.96]	
Segeral 2018	28	192	107	990	0.21 [0.14, 0.29]	0.84 [0.82, 0.86]	+ •
Meya 2009	10	52	39	395	0.20 [0.10, 0.34]	0.88 [0.85, 0.91]	
Paolillo 2017	46	33	192	264	0.19 [0.15, 0.25]	0.89 [0.85, 0.92]	+ +
Lan de s 2021	16	321	67	37	0.19 [0.11, 0.29]	0.10 [0.07, 0.14]	
Orrell 2017	1	143	5	20	0.17 [0.00, 0.64]	0.12 [0.08, 0.18]	- - +
Duarte 2015	19	10	104	224	0.15 [0.10, 0.23]	0.96 [0.92, 0.98]	+ +
Segeral 2010	11	23	63	160	0.15 [0.08, 0.25]	0.87 [0.82, 0.92]	
Sangeda 2014	7	14	48	93	0.13 [0.05, 0.24]	0.87 [0.79, 0.93]	- -
Mogosetsi 2018	1	12	7	78	0.13 [0.00, 0.53]	0.87 [0.78, 0.93]	- -
El-Kĥatib 2010	10	32	92	740	0.10 [0.05, 0.17]	0.96 [0.94, 0.97]	. .
							0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1
IC., harava an ak	and a b		h		a shaldl Calf samet a		a shald, a OEX, a dhasanaa

[Subgroup analysis by adherence threshold] Self-report questionnaires; threshold: ≥ 95% adherence

Study	тр	FP	FN	ΤN	Sensitivity (95% Cl)	Specificity (95% CI)	Sensitivity (95% Cl)Specificity (95% Cl)
Avong 2015	84	185	15	218	0.85 [0.76, 0.91]	0.54 [0.49, 0.59]	
Haberer 2011	13	29	7	24	0.65 [0.41, 0.85]	0.45 [0.32, 0.60]	
Zoufaly 2013	27	23	94	86	0.22 [0.15, 0.31]	0.79 [0.70, 0.86]	+ +
Ekstrand 2010	10	11	45	136	0.18 [0.09, 0.31]	0.93 [0.87, 0.96]	0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

TP FP FN

15

39 395

48

7 78

62

95

86

93

44 224

34 26

10 52

1 12

45 2

31 41 60

27 23 94

11 23 63 160

7 14

Study Phillips 2019

Coker 2015

Tabb 2018

Meya 2009

Zoufaly 2013

Segeral 2010

Sangeda 2014

Mogosetsi 2018

Figure 15. Self-report questionnaires; various adherence thresholds* [subgroup by number of questions] *cut-off used was either ≥ 95% or 100%

[Subgroup analysis by number of questions] Self-report questionnaires; various adherence thresholds*; 1-item

Study TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)

McMahon 2013	25 105	9 31	0.74 [0.56, 0.87]	0.23 [0.16, 0.31]		-
Fokam 2017	28 64	14 38	0.67 [0.50, 0.80]	0.37 [0.28, 0.47]		-
Haberer 2011	13 29	7 24	0.65 [0.41, 0.85]	0.45 [0.32, 0.60]		
Parker 2017	106 458	68 1017	0.61 [0.53, 0.68]	0.69 [0.67, 0.71]		•
Bajunirwe 2009	10 16	18 131	0.36 [0.19, 0.56]	0.89 [0.83, 0.94]		-
Oette 2006	13 11	48 136	0.21 [0.12, 0.34]	0.93 [0.87, 0.96]		-
Paolillo 2017	46 33	192 264	0.19 [0.15, 0.25]	0.89 [0.85, 0.92]	-	-
Lan de s 2021	16 321	67 37	0.19 [0.11, 0.29]	0.10 [0.07, 0.14]		•
Ekstran d 2010	10 11	45 136	0.18 [0.09, 0.31]	0.93 [0.87, 0.96]		-
Orrell 2017	1 143	5 20	0.17 [0.00, 0.64]	0.12 [0.08, 0.18]		-
Duarte 2015	19 10	104 224	0.15 [0.10, 0.23]	0.96 [0.92, 0.98]	-	•
El-Khatib 2010	10 32	92 740	0.10 [0.05, 0.17]	0.96 [0.94, 0.97]		

5 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 5 0 0.2 0.4 0.6 0.8 1 [Subgroup analysis by number of questions] Self-report questionnaires; various adherence thresholds*; 2 to 4 ite

0.70 [0.60, 0.80]

0.99 [0.97, 1.00]

0.70 [0.61, 0.77]

0.79 [0.70, 0.86]

0.88 [0.85, 0.91]

0.87 [0.82, 0.92] 0.87 [0.79, 0.93]

0.87 [0.78, 0.93]

TN Sensitivity (95% CI) Specificity (95% CI)

0.69 [0.55, 0.82]

0.51 [0.40, 0.61]

0.34 [0.24, 0.45]

0.22 [0.15, 0.31]

0.20 [0.10, 0.34]

0.15 [0.08, 0.25]

0.13 [0.05, 0.24]

0.13 [0.00, 0.53]

Sensitivity (95% CI)Specificity (95% CI)



[Subgroup analysis by number of questions] Self-report questionnaires; various adherence thresholds*; 5 or mor

Study	ΤР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)
Avong 2015	84	185	15	218	0.85 [0.76, 0.91]	0.54 [0.49, 0.59]
Pulido 2009	9	27	6	79	0.60 [0.32, 0.84]	0.75 [0.65, 0.82]
Pasquau 2018	6	51	- 7	125	0.46 [0.19, 0.75]	0.71 [0.64, 0.78]
Mbengue 2019	13	24	36	90	0.27 [0.15, 0.41]	0.79 [0.70, 0.86]
Segeral 2018	28	192	107	990	0.21 [0.14, 0.29]	0.84 [0.82, 0.86]



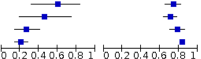




Figure 16. Self-report questionnaires; various adherence thresholds* [subgroup by population] *cut-off used was either ≥ 95% or 100%

[Subgroup analysis by population] Self-report questionnaires; various adherence thresholds*; children

Study	ΤР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% Cl)	Sensitivity (95% Cl)Specificity (95% Cl)
Fokam 2017	28	64	14	38	0.67 [0.50, 0.80]	0.37 [0.28, 0.47]	
Haberer 2011	13	29	7	24	0.65 [0.41, 0.85]	0.45 [0.32, 0.60]	
Zoufaly 2013	27	23	94	86	0.22 [0.15, 0.31]	0.79 [0.70, 0.86]	
Duarte 2015	19	10	104	224	0.15 [0.10, 0.23]	0.96 [0.92, 0.98]	

5 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 [Subgroup analysis by population] Self-report questionnaires; various adherence thresholds*; adults

Study	ТР	FP	FN	TN	Sensitivity (95% Cl)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
Avong 2015	84	185	15	218	0.85 [0.76, 0.91]	0.54 [0.49, 0.59]	
McMahon 2013	25	105	9	31	0.74 [0.56, 0.87]	0.23 [0.16, 0.31]	
Phillips 2019	34	26	15	62	0.69 [0.55, 0.82]	0.70 [0.60, 0.80]	
Parker 2017	106	458	68	1017	0.61 [0.53, 0.68]	0.69 [0.67, 0.71]	
Pulido 2009	9	27	6	79	0.60 [0.32, 0.84]	0.75 [0.65, 0.82]	
Coker 2015	45	2	44	224	0.51 [0.40, 0.61]	0.99 [0.97, 1.00]	
Pasquau 2018	6	51	- 7	125	0.46 [0.19, 0.75]	0.71 [0.64, 0.78]	
Bajunirwe 2009	10	16	18	131	0.36 [0.19, 0.56]	0.89 [0.83, 0.94]	
Mbengue 2019	13	24	36	90	0.27 [0.15, 0.41]	0.79 [0.70, 0.86]	
Oette 2006	13	11	48	136	0.21 [0.12, 0.34]	0.93 [0.87, 0.96]	
Segeral 2018	28	192	107	990	0.21 [0.14, 0.29]	0.84 [0.82, 0.86]	+ +
Meya 2009	10	52	39	395	0.20 [0.10, 0.34]	0.88 [0.85, 0.91]	
Paolillo 2017	46	33	192	264	0.19 [0.15, 0.25]	0.89 [0.85, 0.92]	· • · · ·
Lan de s 2021	16	321	67	37	0.19 [0.11, 0.29]	0.10 [0.07, 0.14]	+
Ekstrand 2010	10	11	45	136	0.18 [0.09, 0.31]	0.93 [0.87, 0.96]	
Segeral 2010	11	23	63	160	0.15 [0.08, 0.25]	0.87 [0.82, 0.92]	+ +
San ged a 2014	7	14	48	93	0.13 [0.05, 0.24]	0.87 [0.79, 0.93]	
Mogosetsi 2018	1	12	7	78	0.13 [0.00, 0.53]	0.87 [0.78, 0.93]	
El-Khatib 2010	10	32	92	740	0.10 [0.05, 0.17]	0.96 [0.94, 0.97]	0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

Librarv

Figure 17. Self-report questionnaires; various adherence thresholds* [subgroup by viral load] *cut-off used was either ≥ 95% or 100%

[Subgroup analysis by viral load] Self-report questionnaires; various adherence thresholds*; 40 to 50 copies/mL

Study	ТР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% Cl)Specificity (95% Cl)
Phillips 2019	34	26	15	62	0.69 [0.55, 0.82]	0.70 [0.60, 0.80]	
Fokam 2017	28	64	14	38	0.67 [0.50, 0.80]	0.37 [0.28, 0.47]	
Haberer 2011	13	29	7	24	0.65 [0.41, 0.85]	0.45 [0.32, 0.60]	
Pulido 2009	9	27	6	79	0.60 [0.32, 0.84]	0.75 [0.65, 0.82]	
Pasquau 2018	6	51	7	125	0.46 [0.19, 0.75]	0.71 [0.64, 0.78]	
Bajunirwe 2009	10	16	18	131	0.36 [0.19, 0.56]	0.89 [0.83, 0.94]	
0ette 2006	13	11	48	136	0.21 [0.12, 0.34]	0.93 [0.87, 0.96]	
Paolillo 2017	46	33	192	264	0.19 [0.15, 0.25]	0.89 [0.85, 0.92]	• •
Lan de s 2021	16	321	67	37	0.19 [0.11, 0.29]	0.10 [0.07, 0.14]	•
Orrell 2017	1	143	5	20	0.17 [0.00, 0.64]	0.12 [0.08, 0.18]	- -
Mogosetsi 2018	1	12	- 7	78	0.13 [0.00, 0.53]	0.87 [0.78, 0.93]	
							0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1
[Subgroup ana	lysis	by vi	ral lo	ad] S∢	elf-report questionn	aires; various adher	ence thresholds*; 200 to 400 copies/ml
.	-						
Study	TP	FP	FN	IN	 Sensitivity (95% CI) 	 Specificity (95% CI) 	Sensitivity (95% CI)Specificity (95% CI)
Study Avong 2015	1P 84		FN 15	218	,		
,		185			0.85 [0.76, 0.91]	0.54 [0.49, 0.59]	· · · · ·
Avong 2015	84	185	15	218 31	0.85 (0.76, 0.91 0.74 (0.56, 0.87	0.54 [0.49, 0.59] 0.23 [0.16, 0.31]	
Avong 2015 McMahon 2013	84 25	185 105	15 9	218 31	0.85 (0.76, 0.91 0.74 (0.56, 0.87 0.61 (0.53, 0.68	0.54 [0.49, 0.59] 0.23 [0.16, 0.31] 0.69 [0.67, 0.71]	
Avong 2015 McMahon 2013 Parker 2017	84 25 106	185 105 458	15 9 68	218 31 1017	0.85 (0.76, 0.91 0.74 (0.56, 0.87 0.61 (0.53, 0.68 0.51 (0.40, 0.61	0.54 [0.49, 0.59] 0.23 [0.16, 0.31] 0.69 [0.67, 0.71] 0.99 [0.97, 1.00]	
Avong 2015 McMahon 2013 Parker 2017 Coker 2015	84 25 106 45	185 105 458 2	15 9 68 44	218 31 1017 224	0.85 [0.76, 0.91 0.74 [0.56, 0.87 0.61 [0.53, 0.68 0.51 [0.40, 0.61 0.34 [0.24, 0.45	0.54 [0.49, 0.59] 0.23 [0.16, 0.31] 0.69 [0.67, 0.71] 0.99 [0.97, 1.00] 0.70 [0.61, 0.77]	
Avong 2015 McMahon 2013 Parker 2017 Coker 2015 Tabb 2018	84 25 106 45 31	185 105 458 2 41	15 9 68 44 60	218 31 1017 224 95	0.85 [0.76, 0.91 0.74 [0.56, 0.87 0.61 [0.53, 0.68 0.51 [0.40, 0.61 0.34 [0.24, 0.45 0.27 [0.15, 0.41	0.54 [0.49, 0.59] 0.23 [0.16, 0.31] 0.69 [0.67, 0.71] 0.99 [0.97, 1.00] 0.70 [0.61, 0.77] 0.79 [0.70, 0.86]	
Avong 2015 McMahon 2013 Parker 2017 Coker 2015 Tabb 2018 Mbengue 2019	84 25 106 45 31 13	185 105 458 2 41 24	15 9 68 44 60 36	218 31 1017 224 95 90	0.85 [0.76, 0.91 0.74 [0.56, 0.87 0.61 [0.53, 0.68 0.51 [0.40, 0.61 0.34 [0.24, 0.45 0.27 [0.15, 0.41 0.22 [0.15, 0.31	0.54 [0.49, 0.59] 0.23 [0.16, 0.31] 0.69 [0.67, 0.71] 0.99 [0.97, 1.00] 0.70 [0.61, 0.77] 0.79 [0.70, 0.86] 0.79 [0.70, 0.86]	
Avong 2015 McMahon 2013 Parker 2017 Coker 2015 Tabb 2018 Mbengue 2019 Zoufaly 2013	84 25 106 45 31 13 27	185 105 458 2 41 24 23	15 9 68 44 60 36 94	218 31 1017 224 95 90 86	0.85 [0.76, 0.91 0.74 [0.56, 0.87 0.61 [0.53, 0.68 0.51 [0.40, 0.61 0.34 [0.24, 0.45 0.27 [0.15, 0.41 0.22 [0.15, 0.31 0.21 [0.14, 0.29	0.54 [0.49, 0.59] 0.23 [0.16, 0.31] 0.69 [0.67, 0.71] 0.99 [0.97, 1.00] 0.70 [0.61, 0.77] 0.79 [0.70, 0.86] 0.79 [0.70, 0.86] 0.84 [0.82, 0.86]	
Avong 2015 McMahon 2013 Parker 2017 Coker 2015 Tabb 2018 Mbengue 2019 Zoufaly 2013 Segeral 2018	84 25 106 45 31 13 27 28	185 105 458 2 41 24 23 192	15 9 68 44 60 36 94 107	218 31 1017 224 95 90 86 990	0.85 [0.76, 0.91 0.74 [0.56, 0.87 0.61 [0.53, 0.68 0.51 [0.40, 0.61 0.34 [0.24, 0.45 0.27 [0.15, 0.41 0.22 [0.15, 0.31 0.21 [0.14, 0.29 0.20 [0.10, 0.34	0.54 [0.49, 0.59] 0.23 [0.16, 0.31] 0.69 [0.67, 0.71] 0.99 [0.97, 1.00] 0.70 [0.61, 0.77] 0.79 [0.70, 0.86] 0.79 [0.70, 0.86] 0.84 [0.82, 0.86] 0.88 [0.85, 0.91]	
Avong 2015 McMahon 2013 Parker 2017 Coker 2015 Tabb 2018 Mbengue 2019 Zoufaly 2013 Segeral 2018 Meya 2009	84 25 106 45 31 13 27 28 10	185 105 458 2 41 24 23 192 52	15 9 68 44 60 36 94 107 39 45	218 31 1017 224 95 90 86 990 395	0.85 [0.76, 0.91 0.74 [0.56, 0.87 0.61 [0.53, 0.68 0.51 [0.40, 0.61 0.34 [0.24, 0.45 0.27 [0.15, 0.41 0.22 [0.15, 0.31 0.21 [0.14, 0.29 0.20 [0.10, 0.34 0.18 [0.09, 0.31	0.54 [0.49, 0.59] 0.23 [0.16, 0.31] 0.69 [0.67, 0.71] 0.99 [0.97, 1.00] 0.70 [0.61, 0.77] 0.79 [0.70, 0.86] 0.79 [0.70, 0.86] 0.84 [0.82, 0.86] 0.88 [0.85, 0.91] 0.93 [0.87, 0.96]	
Avong 2015 McMahon 2013 Parker 2017 Coker 2015 Tabb 2018 Mbengue 2019 Zoufaly 2013 Segeral 2018 Meya 2009 Ekstrand 2010	84 25 106 45 31 13 27 28 10 10	185 105 458 2 41 24 23 192 52 11	15 9 68 44 60 36 94 107 39 45	218 31 1017 224 95 90 86 990 395 136	0.85 [0.76, 0.91 0.74 [0.56, 0.87 0.61 [0.53, 0.68 0.51 [0.40, 0.61 0.34 [0.24, 0.45 0.27 [0.15, 0.41 0.22 [0.15, 0.31 0.21 [0.14, 0.29 0.20 [0.10, 0.34 0.18 [0.09, 0.31 0.15 [0.10, 0.23	0.54 [0.49, 0.59] 0.23 [0.16, 0.31] 0.69 [0.67, 0.71] 0.99 [0.97, 1.00] 0.70 [0.61, 0.77] 0.79 [0.70, 0.86] 0.79 [0.70, 0.86] 0.84 [0.82, 0.86] 0.88 [0.85, 0.91] 0.93 [0.87, 0.96] 0.96 [0.92, 0.98]	
Avong 2015 McMahon 2013 Parker 2017 Coker 2015 Tabb 2018 Mbengue 2019 Zoufaly 2013 Segeral 2018 Meya 2009 Ekstrand 2010 Duarte 2015	84 25 106 45 31 13 27 28 10 10 19	185 105 458 2 41 24 23 192 52 11 10	15 9 68 44 60 36 94 107 39 45 104	218 31 1017 224 95 90 86 990 395 136 224	0.85 [0.76, 0.91 0.74 [0.56, 0.87 0.61 [0.53, 0.68 0.51 [0.40, 0.61 0.34 [0.24, 0.45 0.27 [0.15, 0.41 0.22 [0.15, 0.31 0.21 [0.14, 0.29 0.20 [0.10, 0.34 0.18 [0.09, 0.31 0.15 [0.10, 0.23 0.13 [0.05, 0.24	0.54 [0.49, 0.59] 0.23 [0.16, 0.31] 0.69 [0.67, 0.71] 0.99 [0.97, 1.00] 0.70 [0.61, 0.77] 0.79 [0.70, 0.86] 0.79 [0.70, 0.86] 0.84 [0.82, 0.86] 0.88 [0.85, 0.91] 0.93 [0.87, 0.96] 0.96 [0.92, 0.98] 0.87 [0.79, 0.93]	



Figure 18. Self-report questionnaires; various adherence thresholds* [subgroup by setting] *cut-off used was either ≥ 95% or 100%

[Subgroup analysis by setting] Self-report questionnaires; various adherence thresholds*; low-income

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
Avong 2015	84	185	15	218	0.85 [0.76, 0.91]	0.54 [0.49, 0.59]	
McMahon 2013	25	105	9	31	0.74 [0.56, 0.87]	0.23 [0.16, 0.31]	
Haberer 2011	13	29	7	24	0.65 [0.41, 0.85]	0.45 [0.32, 0.60]	
Coker 2015	45	2	44	224	0.51 [0.40, 0.61]	0.99 [0.97, 1.00]	
Bajunirwe 2009	10	16	18	131	0.36 [0.19, 0.56]	0.89 [0.83, 0.94]	
Tabb 2018	31	41	60	95	0.34 [0.24, 0.45]	0.70 [0.61, 0.77]	+ +
Segeral 2018	28	192	107	990	0.21 [0.14, 0.29]	0.84 [0.82, 0.86]	
Meya 2009	10	52	39	395	0.20 [0.10, 0.34]	0.88 [0.85, 0.91]	
Landes 2021	16	321	67	37	0.19 [0.11, 0.29]	0.10 [0.07, 0.14]	
Segeral 2010	11	23	63	160	0.15 [0.08, 0.25]	0.87 [0.82, 0.92]	
Sangeda 2014	- 7	14	48	93	0.13 [0.05, 0.24]	0.87 [0.79, 0.93]	
-							0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1
[Subgroup ana	lysis	by se	ettin	g] Sel	lf-report questionna	ires; various adhere	nce thresholds*; lower-middle-income
Study	ΤР	FP F	N 1	ΓN S€	ensitivity (95% CI) - Sj	pecificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
Fokam 2017	28	64 1	.4 🔅	38	0.67 [0.50, 0.80]	0.37 [0.28, 0.47]	
Zoufaly 2013	27	23 9	94 8	B6	0.22 [0.15, 0.31]	0.79 [0.70, 0.86]	+ +
Ekstrand 2010	10	11 4	5 13	36	0.18 [0.09, 0.31]	0.93 [0.87, 0.96]	
							0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1
[Subgroup ana	lysis	by se	ettin	g] Sel	lf-report questionna	ires; various adhere	nce thresholds*; upper-middle-income
Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
Phillips 2019	34	26	15	62	0.69 [0.55, 0.82]	0.70 [0.60, 0.80]	
Mbengue 2019	13	24	36	90	0.27 [0.15, 0.41]	0.79 [0.70, 0.86]	
Orrell 2017	1	143	5	20	0.17 [0.00, 0.64]	0.12 [0.08, 0.18]	- -
Mogosetsi 2018	1	12	- 7	78	0.13 [0.00, 0.53]	0.87 [0.78, 0.93]	
El-Khatib 2010	10	32	92	740	0.10 [0.05, 0.17]	0.96 [0.94, 0.97]	
							0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1
[Subgroup ana	lysis	by se	ettin	g] Sel	lf-report questionna	ires; various adhere	nce thresholds*; high-income
Study	ΤР	FP	FN	TN	I Sensitivity (95% CI) Specificity (95% Cl) Sensitivity (95% CI)Specificity (95% CI)
Parker 2017	106	458	68	1017	0.61 [0.53, 0.68] 0.69 [0.67, 0.71] 🗕 🖬
Puli do 2009	9	27	6	79	0.60 [0.32, 0.84] 0.75 [0.65, 0.82] — — — —

Parker 2017	106	458	68	1017	0.61 [0.53, 0.6
Puli do 2009	9	27	6	79	0.60 [0.32, 0.8
Pasquau 2018	6	51	- 7	125	0.46 [0.19, 0.7
Oette 2006	13	11	48	136	0.21 [0.12, 0.3
Paolillo 2017	46	33	192	264	0.19 [0.15, 0.2

o CI)	Specificity (95% CI)
.68]	0.69 [0.67, 0.71]
.84]	0.75 [0.65, 0.82]
.75]	0.71 [0.64, 0.78]
.34]	0.93 [0.87, 0.96]
.25]	0.89 [0.85, 0.92]

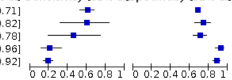


Figure 19. Self-report questionnaires; threshold: ≥ 80% adherence [supplementary analysis]

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)

11000002010	20 0	20	- 00	0.41 [0.27, 0.30]	0.97 [0.90, 0.99]		
Haberer 2011	4 10	16	43	0.20 [0.06, 0.44]	0.81 [0.68, 0.91]	-	
Segeral 2018	11 31	124	1151	0.08 [0.04, 0.14]	0.97 [0.96, 0.98]		0 0.2 0.4 0.6 0.8 1

Appendix 7. Self-report VAS subgroup analysis

Figure 20; Figure 21; Figure 22; Figure 23; Figure 24

Figure 20. Self-report using VAS; threshold: ≥ 95% adherence [subgroup by population]

[Subgroup analysis by population] Self-report VAS; threshold: ≥ 95% adherence; children

Study	ТР	FP F	N TN	Sei	nsitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
Haberer 2011	6	21 1	4 32		0.30 [0.12, 0.54]	0.60 [0.46, 0.74]	_ _
Dziva 2017	16	12 4	5 93	ł	0.26 [0.16, 0.39]	0.89 [0.81, 0.94]	
							0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1
[Subgroup ana	ilysis	s by p	opula	ation] Self-report VAS;	: threshold: ≥ 95% adh	erence; adults
Study	TP	FP	FN	ΤN	Sensitivity (95%	CI) Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
Jiamsakul 2014	- 7	9	5	60	0.58 [0.28, 0.8	85] 0.87 [0.77, 0.94]	
Cerutti 2016	52	558	63	691	0.45 [0.36, 0.5	55] 0.55 [0.53, 0.58]	
McMahon 2013	12	38	22	98	0.35 [0.20, 0.5	54] 0.72 [0.64, 0.79]	
Ekstran d 2010	18	16	37	131	0.33 [0.21, 0.4	47] 0.89 [0.83, 0.94]	
Nelson 2010	46	79	104	417	0.31 [0.23, 0.3	39] 0.84 [0.81, 0.87]	
San ged a 2014	10	19	45	88	0.18 [0.09, 0.3	31] 0.82 [0.74, 0.89]	
Cohen 2012	41	134	863	176	0.05 [0.03, 0.0	06] 0.57 [0.51, 0.62]	• •
Gill 2010	0	2	8	55	0.00 [0.00, 0.3	37] 0.96 [0.88, 1.00]	

Figure 21. Self-report using VAS; threshold: ≥ 95% adherence [subgroup by viral load]

[Subgroup analysis by viral load] Self-report VAS; threshold: ≥ 95% adherence; 40 to 100 copies/mL

Study	ΤР	FP	FN	TN	Sensitivity (95% Cl)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
Cerutti 2016	52	558	63	691	0.45 [0.36, 0.55]	0.55 [0.53, 0.58]	
Ekstrand 2010	18	16	37	131	0.33 [0.21, 0.47]	0.89 [0.83, 0.94]	
Nelson 2010	46	- 79	104	417	0.31 [0.23, 0.39]	0.84 [0.81, 0.87]	
Haberer 2011	6	21	14	32	0.30 [0.12, 0.54]	0.60 [0.46, 0.74]	- -
Labhardt 2012	15	0	50	27	0.23 [0.14, 0.35]	1.00 [0.87, 1.00]	
C o hen 2012	41	134	863	176	0.05 [0.03, 0.06]	0.57 [0.51, 0.62]	
[Subgroup ana	lysis	s by v	/iral	load]	Self-report VAS; thr	eshold: ≥ 95% adhe	rence; 200 to 400 copies/mL
Study	ТР	FP	FN	TN S	ensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
Jiamsakul 2014	7	9	5	60	0.58 [0.28, 0.85]	0.87 [0.77, 0.94]	
McMahon 2013	12	38	22	98	0.35 [0.20, 0.54]	0.72 [0.64, 0.79]	
Dziva 2017	16	12	45	93	0.26 [0.16, 0.39]	0.89 [0.81, 0.94]	
San ged a 2014	10	19	45	88	0.18 [0.09, 0.31]	0.82 [0.74, 0.89]	
Gill 2010	0	2	8	55	0.00 [0.00, 0.37]	0.96 [0.88, 1.00]	

Accuracy of measures for antiretroviral adherence in people living with HIV (Review) Copyright © 2022 The Authors. Cochrane Database of Systematic Reviews published by John Wiley & Sons, Ltd. on behalf of The Cochrane Collaboration.

Figure 22. Self-report using VAS; threshold: 95% adherence [subgroup by setting]

[Subgroup analysis by setting] Self-report VAS; threshold: ≥ 95% adherence; low-income

Study	тр	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% Cl)Specificity (95%	% CI}
McMahon 2013	12	38	22	98	0.35 [0.20, 0.54]	0.72 [0.64, 0.79]		
Haberer 2011	6	21	14	32	0.30 [0.12, 0.54]	0.60 [0.46, 0.74]	_ -	
Dziva 2017	16	12	45	93	0.26 [0.16, 0.39]	0.89 [0.81, 0.94]		-
Labhardt 2012	15	0	50	27	0.23 [0.14, 0.35]	1.00 [0.87, 1.00]		
San ged a 2014	10	19	45	88	0.18 [0.09, 0.31]	0.82 [0.74, 0.89]		-
							0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.	81
tsubgroup ana	lysis	by :	setti	ing] !	Self-report VAS; thre	shold: ≥ 95% adher	ence; lower-middle-income	
Study	ilysis TP		setti FN		Self-report VAS; thre Sensitivity (95% Cl)		ence; lower-middle-income Sensitivity (95% CI)Specificity (959	% CI)
<u> </u>	тр		FN	TN	•	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95%	% CI)
Study	тр	FP	FN 63	TN	Sensitivity (95% Cl)	Specificity (95% Cl) 0.55 [0.53, 0.58]	Sensitivity (95% CI)Specificity (95%	% CI)

Figure 23. Self-report using VAS; threshold: 90% adherence [additional analysis]

Study	ΤР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI	1)
Mbengue 2019	12	14	37	100	0.24 [0.13, 0.39]	0.88 [0.80, 0.93] —	
Sangeda 2014	5	10	50	97	0.09 [0.03, 0.20]	0.91 [0.83, 0.95] 📲	
Segeral 2010	2	9	72	174	0.03 [0.00, 0.09]	0.95 [0.91, 0.98]	1

Figure 24. Self-report using VAS; threshold: ≥ 80% adherence [supplementary analysis]

Study	ТР	FP	FN	ΤN	Sensitivity (95% CI)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)
Haberer 2011	4	10	16	43	0.20 [0.06, 0.44]	

Appendix 8. Tablet counts subgroup analysis

Figure 25; Figure 26; Figure 27; Figure 28

Figure 25. Tablet counts; threshold: ≥ 95% adherence [subgroup by population]

[Subgroup analysis by population] Tablet counts; threshold: ≥ 95% adherence; children

Study TP FF Haberer 2011 7 26 [Subgroup analysis b	13	27		0.35		[0.37, 0.65]	Sensitivity (95% CI)Specificity (95% CI)
Study	тр	FP	FN	τN	Sensitivity (95% Cl)	Specificity (95% Cl) Sensitivity (95% CI)Specificity (95% CI)
Sangeda 2014	43	89	12	18	0.78 [0.65, 0.88]	0.17 [0.10, 0.25]
Apisarnthanarak 2010	19	25	6	149	0.76 [0.55, 0.91]	0.86 (0.80, 0.90	j — - -
Bonjoch 2006	47	144	23	8	0.67 [0.55, 0.78]	0.05 [0.02, 0.10	j
Coker 2015	25	93	22	133	0.53 [0.38, 0.68]	0.59 [0.52, 0.65	j — — — —
Okonji 2012	24	44	66	300	0.27 [0.18, 0.37]	0.87 [0.83, 0.91] +
Gill 2010	2	9	6	48	0.25 [0.03, 0.65]	0.84 [0.72, 0.93] — • -•
Cerutti 2016	26	275	88	941	0.23 [0.15, 0.32]	0.77 [0.75, 0.80] — 🖛 🛛 🖷
Moosa 2019	1	11	10	211	0.09 [0.00, 0.41]	0.95 [0.91, 0.98] 🗕 🚽
Mariana 2018	0	1	16	81	0.00 [0.00, 0.21]	0.99 [0.93, 1.00	0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

Figure 26. Tablet counts; threshold: ≥ 95% adherence [subgroup by viral load threshold]

[Subgroup analysis by viral load] Tablet count; threshold: ≥ 95% adherence; VL 40 to 80 copies/mL

Study Kitkungvan 2008 Apisarnthanarak Bonjoch 2006	201	D	TP 4 19 47	FP 40 25 144	0 6 23	155 149 8	1.00 (0.40 0.76 (0.55 0.67 (0.55), 1.00] 5, 0.91] 5, 0.78]	0.79 0.86 0.05	(0.73, ((0.80, ((0.02, (0.85] 0.90] 0.10]	Sensitivity (95% CI)Specificity (95% CI)
Haberer 2011			~	26	13	27	0.35 [0.15			[0.37, (
Cerutti 2016				275	88		0.23 [0.15			[0.75, (
Orrell 2017			9	119	35	15	0.20 [0.10), 0.35]	0.11	[0.06, (0.18]	
Mariana 2018			0	1	16	81	0.00 [0.00), 0.21]	0.99	[0.93,]	1.00]	
[Subgroup anal	lysis	ь Бу	vira	al loa	d] Т	able	t count; thres	hold: ≥	95% adh	erenc	e; VL	
Study	тр	FP	FN	TN	S€	ensiti	vity (95% CI)	Specifi	city (95%)	CI)		Sensitivity (95% CI)Specificity (95% CI)
San ged a 2014	43	89	12	18		0.7	8 [0.65, 0.88]	0.17	7 [0.10, 0.2	25]		
Coker 2015	25	93	22	133		0.5	3 [0.38, 0.68]	0.59	0.52, 0.6	35]		
Ok o nji 2012	24	44	66	300		0.2	7 [0.18, 0.37]	0.87	7 [0.83, 0.9	91j		
Gill 2010	2	9	6	i 48		0.2	5 [0.03, 0.65]	0.84	ŧ [0.72, 0.9	93]		
Moosa 2019	1	11	10	211		0.0	9 [0.00, 0.41]	0.95	5 [0.91, 0.9	98]		

Figure 27. Tablet counts; threshold: ≥ 95% adherence [subgroup by setting]

[Subgroup analysis by setting] Tablet counts; threshold: ≥ 95% adherence; low-income

Study	ΤР	FP	FN	TN	I S∢	ensiti	ivity (9	95% CI)) Sp	oecifi	city (95% (CI)		Sensitivity (95% CI)Specificity (95% CI)
Sangeda 2014	43	89	12	18	3	0.73	8 [0.65	5, 0.88]	0.17	7 [0.1	0, 0.2	5]		
Coker 2015	25	93	22	133	3	0.5	3 [0.38	3, 0.68]	0.59	0.5	2, 0.6	5]		
Haberer 2011	- 7	26	13	27	7	0.3	5 [0.15	5, 0.59]	0.51	[0.3	7, 0.6	5]		
0k o nji 2012	24	44	66	300)	0.2	7 [0.18	3, 0.37]	0.87	7 [0.8	3, 0.9	1]		
															0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1
[Subgroup ana	lysi	s by	set	ting	Tal	blet o	ounts	; thre	sho	ld: ≥	95%	adhe	rence	; lov	ver-middle-income
Study	т	Р	FP	FN	τN	Sens	sitivity	/ (95%	CI)	Spec	cificit	ty (95	% CI)		Sensitivity (95% CI)Specificity (95% CI)
Kitkungvan 2008		4	40	0	155		1.00 [0	.40, 1.	00]	C).79 [0.73,	0.85]		
Gill 2010		2	9	6	48	(0.25 [0	.03, 0.	65]	C).84 [0.72,	0.93]		_
Cerutti 2016	2	6 2	75	88	941	(0.23 [0	.15, 0.	32]	0).77 [0.75,	0.80]		
Mariana 2018		0	1	16	81	(0.00 [0	.00, 0.	21]	0).99 [0.93,	1.00]		
															0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1
LSubgroup ana	lysi	s by	set	ting	Tal	blet c	ounts	; thre	sho	ld: ≥	95%	adhe	rence	;; upj	per-middle-income
Study			тр	FP	FN	TN	Sens	itivity	(95 ⁻	% CI)	Spe	cificit	y (95	% CI)	Sensitivity (95% CI)Specificity (95% CI)
Apisarnthanarak	201	0	19	25	6	149	0	.76 [0.	55,	0.91]		0.86 [0.80,	0.90]	
Orrell 2017			9	119	35	15	0	.20 [0.	10,	0.35]		0.11 [0.06, 🛛	0.18]	- - - +
Moosa 2019			1	11	10	211	0	.09 [0.	00,	0.41]		0.95 [0.91,	0.98]	
															0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1
[Subgroup ana	lysi	5 БУ	set	ting	Tal	blet o	ounts	; thre	sho	ld: ≥	95%	adhe	rence	; hig	h-income
Ctudu.	тп	гп	E.M.		6.0	n a lt li			6	a alfi a			a		Constitute INTR CICE selficity INTR CI
	ТР		FN		5e		/ity (99								Sensitivity (95% CI)Specificity (95% CI)
Bonjoch 2006	47	144	23	8		0.67	' [0.55,	0.78]		0.05	[0.0]	2, 0.10	Л		
															0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

Figure 28. Tablet count; threshold: ≥ 80% adherence [supplementary analysis]

Study	ΤР	FP	FN	τN	Sensitivity (95% CI)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI	1)
San ged a 2014	19	33	36	74	0.35 [0.22, 0.49]	0.69 [0.59, 0.78]	
Haberer 2011	0	0	20	53	0.00 [0.00, 0.17]		Ī

Appendix 9. Pharmacy records subgroup analysis

Figure 29; Figure 30; Figure 31; Figure 32

Figure 29. Pharmacy records; threshold: ≥ 95% adherence [subgroup by population]

Study	ТР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% Cl)	Sensitivity (95% CI)Specificity (95% CI)
Messou 2011	205	377	27	316	0.88 [0.84, 0.92]	0.46 [0.42, 0.49]	
San ged a 2014	24	41	22	66	0.52 [0.37, 0.67]	0.62 [0.52, 0.71]	
McMahon 2013	18	25	34	147	0.35 [0.22, 0.49]	0.85 [0.79, 0.90]	+
Anu de 2013	24	24	114	429	0.17 [0.11, 0.25]	0.95 [0.92, 0.97]	

Figure 30. Pharmacy records; threshold: 95% adherence [subgroup by viral load threshold]

[Subgroup analysis by viral load] Pharmacy records; threshold: ≥ 95% adherence; VL: 40 copies/mL

Study Orrell 2017	14	125	27	12	0.	• • •	09 (0.05, 0.15)	Sensitivity (95% Cl)Specificity (95% Cl) 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 erence; VL: 200 to 400 copies/mL
Study		тр	FP					Sensitivity (95% CI)Specificity (95% CI)
Messou 2011		205				0.88 [0.84, 0.92]		
		205	- · ·		316 66	• • •	• • •	
Sangeda 201			41			0.60 [0.46, 0.73]		
Hassan 2014		18	25	34	147	0.35 [0.22, 0.49]	0.85 [0.79, 0.90]	
McMahon 201	.3	8	21	26	119	0.24 [0.11, 0.41]	0.85 [0.78, 0.90]	
Anu de 2013		24	24	114	429	0.17 [0.11, 0.25]	0.95 [0.92, 0.97]	0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

Figure 31. Pharmacy records; threshold: ≥ 95% adherence [subgroup by setting]

[Subgroup analysis by setting] Pharmacy records; threshold: > 95% adherence; low-income

-							-				
Study		ΤР	FP	FN	TN	Sensitivity (95%	% CI)	Specificity (95% C	I) Sensitivity (95% CI)Specificity (95% CI)		
Messou 2011	2	205	377	27	316	0.88 [0.84, 0).92]	0.46 [0.42, 0.49)] 🗕 🖷		
Sangeda 2014		33	41	22	66	0.60 [0.46, 0	0.73]	0.62 [0.52, 0.7]	.] ——— ———		
Hassan 2014		18	25	34	147	0.35 [0.22, 0).49]	0.85 [0.79, 0.90)] — 🖛 —		
McMahon 2013	}	8	21	26	119	0.24 [0.11, 0).41]	0.85 [0.78, 0.90			
[Subgroup analysis by setting] Pharmacy records; threshold: ≥ 95% adherence; lower-middle-income											
Study	ΤР	FP	FN	TN	l Sei	nsitivity (95% CI)) Spe	ecificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)		
Anu de 2013	24	24	114	429)	0.17 [0.11, 0.25]		0.95 [0.92, 0.97]			
[Subgroup an	[Subgroup analysis by setting] Pharmacy records; threshold: ≥ 95% adherence; upper-middle-income										
Study 1	гР	FP	FN	ΤN	Sens	itivity (95% CI)	Spec	ificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)		
Orrell 2017]	14 1	125	27	12	C	.34 [0.20, 0.51]	0.	.09 [0.05, 0.15]			

Figure 32. Pharmacy records; threshold: 80% adherence [supplementary analysis]

Study	ТР	FP	FN	TN	Sensitivity (95% Cl)	Specificity (95% CI) 9	Sensitivity (95% CI)Specificity (95% CI)
Navarro 2014	51	17	11	45	0.82 [0.70, 0.91]	0.73 [0.60, 0.83]	
Messou 2011	148	97	84	596	0.64 [0.57, 0.70]	0.86 [0.83, 0.89]	
San ged a 2014	14	13	41	94	0.25 [0.15, 0.39]	0.88 [0.80, 0.93]	
							0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

Appendix 10. Electronic monitoring devices subgroup analysis

Figure 33; Figure 34; Figure 35; Figure 36

Figure 33. Electronic monitoring; threshold: ≥ 95% adherence [subgroup by population]

[Subgroup analysis by population] Electronic monitoring; threshold: ≥ 95% adherence; children

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)
 Sensitivity (95% CI)

[Subgroup analysis by population] Electronic monitoring; threshold: ≥ 95% adherence; adults

Study	ТР	FP	FN	ΤN	Sensitivity (95% Cl)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
Gill 2010	7	27	1	30	0.88 [0.47, 1.00]	0.53 [0.39, 0.66]	_
Evans 2016	20	16	7	6	0.74 [0.54, 0.89]	0.27 [0.11, 0.50]	

Figure 34. Electronic monitoring; threshold: ≥ 95% adherence [subgroup by viral load threshold]

[Subgroup analysis by viral load] Electronic monitoring; threshold: ≥ 95% adherence; VL: 50 copies/mL

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)
 Sensitivity (95% CI

Study	ΤР	FP	FN	ΤN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
Gill 2010	- 7	27	1	30	0.88 [0.47, 1.00]	0.53 [0.39, 0.66]	
Evans 2016	20	16	7	6	0.74 [0.54, 0.89]	0.27 [0.11, 0.50]	

Figure 35. Electronic monitoring; threshold: ≥ 95% adherence [subgroup by setting]

[Subgroup analysis by setting] Electronic monitoring; threshold: ≥ 95% adherence; low-income

TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI) Study Haberer 2011 12 17 8 35 0.60 [0.36, 0.81] 0.67 [0.53, 0.80] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 [Subgroup analysis by setting] Electronic monitoring; threshold: ≥ 95% adherence; lower-middle-income TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Study Sensitivity (95% CI)Specificity (95% CI) Gill 2010 7 27 1 30 0.88 [0.47, 1.00] 0.53 [0.39, 0.66]

[Subgroup analysis by setting] Electronic monitoring; threshold: ≥ 95% adherence; upper-middle-income

Figure 36. Electronic monitoring; threshold: ≥ 80% adherence [supplementary analysis]

Study	ΤР	FP	FN	ΤN	Sensitivity (95% CI)	Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)
Farley 2003	8	3	1	14	0.89 [0.52, 1.00]	0.82 [0.57, 0.96]
Evans 2016	11	5	16	17	0.41 [0.22, 0.61]	0.77 [0.55, 0.92]
Haberer 2011	5	2	15	50	0.25 [0.09, 0.49]	0.96 [0.87, 1.00] —
Orrell 2017	19	95	59	7	0.24 [0.15, 0.35]	

Appendix 11. Composite measures subgroup analysis

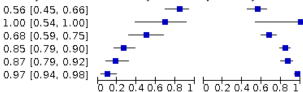
Figure 37; Figure 38; Figure 39; Figure 40

Figure 37. Composite measures; [subgroup by adherence threshold] *cut-off used was either ≥ 95% or 100%

[Subgroup analysis by adherence threshold] Composite measures; 100% adherence

Study TP FP FN TN Sensitivity (95% Cl) Specificity (95% Cl) Sensitivity (95% Cl)Specificity (95% Cl)

Ortega 2004	33	43	6	54	0.85 [0.69, 0.94]
Jayaweera 2003	9	0	4	6	0.69 [0.39, 0.91]
McMahon 2013	17	45	17	95	0.50 [0.32, 0.68]
Segeral 2010	20	28	54	155	0.27 [0.17, 0.39]
Mbengue 2019	9	15	40	99	0.18 [0.09, 0.32]
Spire 2008	- 7	9	63	267	0.10 [0.04, 0.20]



[Subgroup analysis by adherence threshold] Composite measures; 95% adherence

Study	ΤР	FP	FN	ΤN	Sensitivity (95% CI)	Specificity (95% CI)
Parienti 2010	21	26	0	25	1.00 [0.84, 1.00]	0.49 [0.35, 0.63]
Orrell 2003	49	69	33	91	0.60 [0.48, 0.70]	0.57 [0.49, 0.65]
Mutwa 2014	8	13	17	66	0.32 [0.15, 0.54]	0.84 [0.74, 0.91]

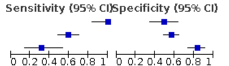


Figure 38. Composite measure; different adherence thresholds* [subgroup by population] *cut-off used was either 95% or 100%

[Subgroup analysis by population] Composite measure; various adherence thresholds*; children

Study	ΤР	FP	FN	ΤN	Sensitivity (95% CI)	Specificity (95% Cl)
Mutwa 2014	8	13	17	66	0.32 [0.15, 0.54]	0.84 [0.74, 0.91]

Sensitivity (95% CI)Specificity (95% CI)

[Subgroup analysis by population] Composite measure; various adherence thresholds*; adults

Study TP_FP_FN_TN_Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI)

· ·					, , , , , , , , , , , , , , , , , , , ,
Parienti 2010	21	26	0	25	1.00 [0.84, 1.00]
Ortega 2004	33	43	6	54	0.85 [0.69, 0.94]
Orrell 2003	49	69	33	91	0.60 [0.48, 0.70]
McMahon 2013	17	45	17	95	0.50 [0.32, 0.68]
Segeral 2010	20	28	54	155	0.27 [0.17, 0.39]
Mbengue 2019	9	15	40	99	0.18 [0.09, 0.32]
Spire 2008	7	9	63	267	0.10 [0.04, 0.20]

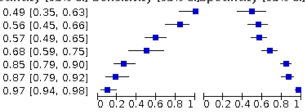




Figure 39. Composite measure; different thresholds* [subgroup by viral load threshold] *cut-off used was either ≥ 95% or 100%

[Subgroup analysis by viral load] Composite measure; various adherence thresholds*; 40 to 50 copies/mL

Study	тр	FP	FN	TN	S	ensitivity (95% CI)	Specificity (95% Cl)	Sensitivity (95% Cl	Specificity (95% CI)
Parienti 2010	21	26	0	25		1.00 [0.84, 1.00]	0.49 [0.35, 0.63]		· -∎
Mutwa 2014	8	13	17	66		0.32 [0.15, 0.54]	0.84 [0.74, 0.91]		
Spire 2008	7	9	63	267		0.10 [0.04, 0.20]	0.97 [0.94, 0.98]		
[Subgroup and	alys	is b	y vir	al lo	ad]	Composite measu	ire; various adherenc	e thresholds*; 200	to 400 copies/mL
Study	Т	ΡF	PF	N 1	ΓN	Sensitivity (95% C	I) Specificity (95% CI)	Sensitivity (95% Cl	Specificity (95% CI)
Parienti 2010	1	6 3	31	0 3	25	1.00 [0.79, 1.00	0.45 [0.31, 0.59]		_ ∎
Ortega 2004	З	3 4	13	6 !	54	0.85 [0.69, 0.94	4] 0.56 [0.45, 0.66]		-
Jayaweera 2003		9	0	4	6	0.69 (0.39, 0.9)	L] 1.00 [0.54, 1.00]		
Orrell 2003	- 4	9 6	i9 (33 9	91	0.60 [0.48, 0.70)] 0.57 [0.49, 0.65]		-
McMahon 2013	1	7 4	15 3	17 9	95	0.50 [0.32, 0.68	3] 0.68 [0.59, 0.75]		-
Segeral 2010	2	20 2	28 3	54 1	55	0.27 [0.17, 0.39	9] 0.85 [0.79, 0.90]		-
Mbengue 2019		91	.5	40 9	99	0.18 [0.09, 0.3)	2] 0.87 [0.79, 0.92]	0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Figure 40. Composite measure; different adherence thresholds* [subgroup by setting] *cut-off used was either ≥ 95% or 100%

[Subgroup analysis by setting] Composite measure; various adherence thresholds*; low-income

Study	тр	FP	FN	ты	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% Cl)Specificity (95% Cl)			
,					•					
McMahon 2013	17	45	17	95	0.50 [0.32, 0.68]	• • •				
Mutwa 2014	8	13	17	66	0.32 [0.15, 0.54]	0.84 [0.74, 0.91]				
Segeral 2010	20	28	54	155	0.27 [0.17, 0.39]	0.85 [0.79, 0.90]				
Spire 2008	7	9	63	267	0.10 [0.04, 0.20]	0.97 [0.94, 0.98]				
1							0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1			
[Subgroup analysis by setting] Composite measure; various adherence thresholds*; upper-middle-income										
	,	-,			F,					
Study	ТР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)			
Orrell 2003	49	69	33	91	0.60 [0.48, 0.70]	0.57 [0.49, 0.65]				
Mbengue 2019	9	15	40	99	0.18 [0.09, 0.32]	0.87 [0.79, 0.92]	. — — —			
	-					;	0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1			
[Suboroun ana	lysis	hv	sett	inal	Composite measure:	various adherence	thresholds*; high-income			
Loandi oah aua	., 0.0	.,	OULL		composite medodi e,		th conoldo) high hicoline			
Study	ТР	FP	FN	ΤN	Sensitivity (95% Cl)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)			
Parienti 2010	21	26	0	25	1.00 [0.84, 1.00]	0.49 [0.35, 0.63]				
Ortega 2004	33		-		0.85 [0.69, 0.94]	0.56 [0.45, 0.66]				
					• • •	• • •				
Jayaweera 2003	9	0	4	6	0.69 [0.39, 0.91]	1.00 [0.54, 1.00]				
							0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1			

HISTORY

Protocol first published: Issue 7, 2018

CONTRIBUTIONS OF AUTHORS

Rhodine Smith supported protocol development, assessed studies for inclusion, extracted data, conducted quality assessments, and drafted the review.

Gemma Villanueva assessed studies for inclusion, extracted data, conducted quality assessments, analysed the data, interpreted the analyses, and drafted the review.

Katrin Probyn assessed studies for inclusion, extracted data, conducted quality assessments, and drafted the review.

Yanina Sguassero assessed studies for inclusion, extracted data, conducted quality assessments, and drafted the review.

Nathan Ford supported protocol development, assessed studies for inclusion, and contributed to the draft manuscript.

Catherine Orrell supported protocol development, assessed studies for inclusion, and contributed to the draft manuscript.

Karen Cohen supported protocol development, assessed studies for inclusion, and contributed to the draft manuscript.

Marty Chaplin contributed to the statistical analyses and contributed to the draft manuscript.

Mariska MG Leeflang supported protocol development, assessed studies for inclusion, interpreted the analyses, and drafted the review.

Paul Hine wrote the protocol, assessed studies for inclusion, interpreted the analyses, and drafted the review.

All authors revised the draft, and agreed with its submission and publication.

DECLARATIONS OF INTEREST

Rhodine Smith has no known conflicts of interest.

Gemma Villanueva is employed by Cochrane Response, an evidence services unit operated by the Cochrane Collaboration, and has no known conflicts of interest. Cochrane Response was contracted by the CIDG to write this review.

Katrin Probyn is employed by Cochrane Response, an evidence services unit operated by the Cochrane Collaboration, and has no known conflicts of interest. Cochrane Response was contracted by the CIDG to write this review.

Yanina Sguassero is employed by Cochrane Response, an evidence services unit operated by the Cochrane Collaboration, and has no known conflicts of interest. Cochrane Response was contracted by the CIDG to write this review.

Nathan Ford has no known conflicts of interest.

Catherine Orrell was study author on the TAP study (2012-2014) and the META study (2014-2017), and has no known conflicts of interest.

Karen Cohen has no known conflicts of interest.

Marty Chaplin has no known conflicts of interest.

Mariska MG Leeflang has no known conflicts of interest.

Paul Hine has no known conflicts of interest.

The author team has no affiliations with or involvement in any organization or entity with a direct financial interest in the subject matter of the review (e.g. employment, consultancy, stock ownership, honoraria, or expert testimony).

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External sources

• Foreign, Commonwealth, and Development Office (FCDO), UK

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DIFFERENCES BETWEEN PROTOCOL AND REVIEW

We amended the title from 'Measures of antiretroviral adherence for detecting viral non-suppression in people living with HIV' (Hine 2018), to 'Accuracy of measures for antiretroviral adherence in people living with HIV'.

We added the website ClinicalTrials.gov to our search strategy after discussion with the CIDG Information Specialists.

We had planned to screen conference abstracts and contact authors in our protocol but, due to the large numbers of studies in the initial searches, we did not complete this.



At the time the protocol was drafted, we did not plan to GRADE the certainty of the evidence. We finally agreed to use the GRADE approach as it is now the recommended approach to summarize the findings from diagnostic test accuracy reviews.

There were minor changes made to the QUADAS-2 tool; these were captured and highlighted in Appendix 4.

INDEX TERMS

Medical Subject Headings (MeSH)

*Anti-Retroviral Agents [therapeutic use]; *HIV Infections [complications] [drug therapy]; Reference Standards; Sensitivity and Specificity; Viral Load

MeSH check words

Adult; Child; Humans