



Government of Malawi Ministry of Health

Integrated HIV Program Report July – September 2015

- *Integrated HIV Program Supervision*
- *HIV Testing and Counselling / Early Infant Diagnosis*
- *Blood Safety*
- *Post Exposure Prophylaxis*
- *HIV Exposed Child Follow-Up*
- *Pre-ART*
- *Prevention of Mother to Child Transmission /
Antiretroviral Therapy*
- *TB / HIV*
- *Sexually Transmitted Infections*
- *Supply of HIV Program Commodities*

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1 Executive Summary

A summary of the key achievements between **July and September 2015** is provided below:

- Scale-up of integrated HIV services had reached the following number of sites:
 - **724** static (579 within and 145 outside of health facilities) and 188 outreach HTC sites
 - **717** (static) ART sites
 - **619** PMTCT sites (Option B+, all included in ART sites above)
 - **663** Pre-ART sites
 - **650** sites with HIV-exposed child follow-up
- **625,803** persons were tested for HIV and received their results; **190,051 (30%)** accessed HTC for the first time; **435,752 (70%)** were repeat testers and **13,093 (3%)** of these received confirmatory testing (after having tested positive in the past). This is equivalent to **51%** confirmatory testing coverage among 25,737 patients initiating ART this quarter. **34,966 (6%)** clients received a positive result for the first time.
- **16,645 (94%)** of 17,773 blood units collected were screened for (at least) HIV, hepatitis B and syphilis.
- **142,701 (91%)** of 156,851 women at ANC had their HIV status ascertained; **11,113 (8%)** of these were HIV positive. **129,887 (97%)** of 133,931 women at maternity had their HIV status ascertained **9,868 (8%)** of these were HIV positive.
- **25,737** patients started ART this quarter.
- **585,660** patients were alive and on ART by end of September 2015. This means that **59%** of the estimated 1 million HIV positive population was on ART. ¹ Estimated ART coverage among people in need for treatment¹ was **50%** (50,533 / 101,000) for children (<15 years) and **68%** (531,990 / 779,000) for adults.
- **78%** of adults and **79%** of children were retained alive on ART at 12 months after initiation. Actual retention rates are thought to be about **10%** higher due to misclassification of 'silent transfers' as defaulters in clinic-based survival/retention analysis. (see section 15.4)
- **510,798 (93%)** of 574,610 patients on first line adult ART were on regimen 5A (TDF/3TC/EFV).
- **11,823² (79%)** of an estimated **14,926¹** HIV infected pregnant women in Malawi were on ART this quarter. **7,031 (59%)** of these were already on ART when getting pregnant and **4,792 (41%)** started ART during pregnancy/delivery.
- An additional **1,555²** breastfeeding women started ART due to **Option B+** (in WHO stage 1/2)
- **78%, 71%, 67%** and **66%** of women started under **Option B+** were retained on ART at **6, 12, 24** and **36 months** after initiation, respectively.
- **9,019 (7%)** of infants discharged alive from maternity were known to be HIV exposed, **8,414 (93%)** of these received ARV prophylaxis (nevirapine). **7,558 (84%)** were enrolled in exposed child follow-up before age 2 months.
- A total of **11,357** HIV exposed children and **6,929** pre-ART patients were enrolled for follow-up in *HIV Care Clinics (HCC)* during this quarter.

¹ 2015 Spectrum estimates based on 2014 definition of eligibility for ART in Malawi (CD4<500, Option B+, UT for U5).

² Adjusted for double counting due to patient transfers / 'failed ART initiations' among women lost to follow-up within 6 months of ART registration.

2 Integrated HIV Program Overview

Malawi implemented a revised HIV Program in all health facilities following the release of the **2011 Malawi Integrated Clinical HIV Guidelines**. The second edition of these guidelines was published in March 2014 and implementation of revised policies commenced in April 2014. Key policies include:

- **PMTCT Option B+**: Universal life-long ART for all HIV infected pregnant and breastfeeding women regardless of clinical or immunological stage.
- Standard **HIV exposed child follow-up** to age 24 months. This program aims to improve early infant diagnosis and ART initiation using DNA-PCR testing for all infants from age 6 weeks; rapid antibody testing is considered diagnostic from age 12 months and repeated at 24 months. HIV exposed child enrolment and follow-up should be integrated with maternal ART follow-up (Option B+) to improve retention and adherence.
- **Early ART initiation**: universal ART for children under 5 years (confirmed HIV infection, CD4% no longer required), children over 5 years and adults with a **CD4** count ≤ 500 , patients with HIV and hepatitis B co-infection.
- Transition to a **new first line ART regimens** for adults (Malawi regimen 5A: tenofovir / lamivudine / efavirenz) and children (Malawi regimen 2: zidovudine / lamivudine / nevirapine), including provision of paediatric ARV formulations for all children under 25kg. Transition to 5A was completed by end 2013.
- Standardized **pre-ART services** for all HIV-infected persons not yet eligible for ART. The pre-ART program aims to reduce the incidence of HIV-related diseases and to enable early ART initiation based on the new CD4 cell threshold (500) through scheduled CD4 count monitoring.
- Provider-initiated provision of **contraceptives and condoms** for all adults in pre-ART and ART clinics to reduce the rate of unwanted pregnancies among HIV-infected adults and to reduce HIV-transmission between sexual partners.
- Isoniazid preventive therapy (**IPT**) for pre-ART patients to reduce the incidence of TB and intensified TB case finding (**ICF**) for all patients in pre-ART and ART follow-up to enable early diagnosis and treatment of TB and to reduce TB transmission in HIV clinics.
- Roll-out of scheduled **viral load monitoring** to improve early detection of treatment failure and initiation of second line ART.

Implementation of PMTCT Option B+ requires provision of ART services at all health facilities with Maternal and Child Health services. This required a massive acceleration of the **decentralization of ART services** from 303 (static) sites in June 2011 to currently 717 sites.

3 Supportive Site Supervision

3.1 Methods

The Department for HIV and AIDS has coordinated quarterly supportive supervision visits to all health facilities with ART services since the start of the national treatment program in 2004. Supervision teams are composed of: experienced HIV clinicians; nurses and M&E staff from health facilities in the public and private sector; district and zonal PMTCT and ART coordinators; program officers and technical staff from the Department for HIV and AIDS; technical staff from implementing partners. The TB and HIV programs have fully integrated their respective site supervision exercises since April 2015.

Each quarter, a one-day pre-supervision meeting is organised for all supervisors participating in the upcoming round to share program updates, discuss observations from the previous round, distribute materials and organise logistics, transport and accommodation.

Standard supervision forms are used to guide implementation of the supervision protocol, to update site information and collect M&E reports. Custom forms with previous data for each site are printed from the HIV Department database. The supervision forms include:

- Contact details of HIV service providers at each site
- Quality of service checklist
- Follow up on action points noted during the previous visit
- Next visit date
- M&E reports from HTC, ANC, maternity, exposed child and pre-ART follow-up, ART and TB
- Physical drug stock-level assessment
- Identification of sites in urgent need of clinical mentoring
- Semi-structured feedback and performance rating for the supervision teams by facility staff

One copy of the supervision form is returned to the Department for HIV and AIDS, where data are entered in a custom MS Access database to produce national reports and to manage program logistics and the commodity supply chain. A second copy of the supervision form is left at the sites.

The supervision protocol includes a systematic review and verification of primary records (patient cards and registers) at all sites. This effectively provides a quarterly quality audit for M&E records, which has resulted in exceptional accuracy and completeness of HIV Program data in Malawi. At the same time, the systematic chart review helps to identify complex cases or deviations from clinical protocol, allowing the supervision team to provide targeted mentoring and clinical advice. The quarterly supervision exercise also aims to boost staff morale and motivation through *Certificates of Excellence* that are awarded by MOH to sites with an excellent score on the quality of service checklist. A growing number of health workers from sites all over the country participate as supervisors in this quarterly exercise and this has strengthened the national HIV Program identity and has greatly facilitated communication between program staff at the national, zonal, district and facility level.

The HTC Program usually conducts a separate supportive site supervision exercise each quarter, targeting a sample of HTC sites both within and outside of health facilities. Supervision teams consist of district, zonal and national level HTC coordinators, supported by implementing partners.

3.2 Supervision Outcomes

727 public and private sector facilities were visited for **clinical HIV program supervision** between 5th and 16th October 2015.

The large number of sites was covered by **166** supervisors working in **32** teams that spent a total of **1,989 working hours** at the sites. Each site visit lasted on average **2.8** hours, but up to 2 days were spent at the busiest sites. **330 (45%)** sites were awarded a *Certificate of Excellence* for **excellent performance**. The

number of sites with excellent performance is lower than previous quarter 372. **68 (9%)** sites had significant weaknesses and were rated to require **intensive mentoring**. The capacity to provide site mentoring will need to be further expanded.

Table 1: Outcomes of integrated HIV services supervision for 2015 Q3

Zone	Total facil. visited*	Supervision hours spent at facilities		Performance (# and % of sites)	
		Total	Average per site	Excellent perform.	Mentoring needed
NZ	132	347	2.6	64 48%	16 12%
CEZ	101	303	3	57 56%	11 11%
CWZ	163	397	2.5	63 39%	9 6%
SEZ	164	502	3.1	82 50%	23 14%
SWZ	167	440	2.7	64 38%	9 5%
Malawi	727	1,989	2.8	330 45%	68 9%

* includes facilities that were visited for assessment of readiness, but that may have not (yet) been designated to provide integrated HIV services.

Table 1 provides a summary of the supervision outcomes by zone. Most facilities were using the standard national M&E tools. **137** sites had cumulatively registered more than 2,000 ART patient and **49** of these had registered more than 5,000. **51 (37%)** of these high burden sites were using electronic data system for ART (EDS). Some NGO supported sites were using custom tools

compatible with the national standard reporting requirements.

4 Inventory of Sites and Services

4.1 Sites and Services

There were **724** static and **188** outreach HTC sites in Q3 2015; **145** of these were outside of health facilities.

Table 2: Facilities with integrated HIV services in the 5 Zones. Availability of services defined by performance (at least 1 patient enrolled) during 2015 Q3

Zone	Total fac.(1)	Facilities providing HIV services				CD4 count machines (2)			Results
		Exp. child	Pre-ART	PMTCT B+	ART	Installed	Functional	Results	
NZ	134	118 88%	117 87%	109 81%	130 97%	34 25%	28 82%	2,811	
CEZ	101	95 94%	93 92%	87 86%	98 97%	16 16%	14 88%	1,289	
CWZ	164	132 80%	132 80%	132 80%	160 98%	30 18%	27 90%	3,454	
SWZ	169	146 86%	163 96%	137 81%	166 98%	39 23%	37 95%	6,093	
SEZ	164	159 97%	158 96%	154 94%	163 99%	51 31%	45 88%	5,866	
Malawi	732	650 89%	663 91%	619 85%	717 98%	170 23%	151 89%	19,513	

(1) Total facilities in the public / private sector designated to provide integrated HIV services in this quarter. Individual site selection is reviewed and may change each quarter.

(2) CD4 machines that have produced at least 1 result during the reporting period are defined as functional.

Table 2 shows the distribution of the **732** sites designated to provide clinical HIV services in Q3 2015, by zone. At the national level, there were **717** (static) sites with at least one patient on ART, **619** sites had enrolled women under PMTCT Option B+; **663** sites were providing pre-ART services. **650** had enrolled HIV exposed children for follow-up. ART services were now available at almost all designated sites in the 5 zones.

The SEZ had reached 99% of designated sites with ART services. SWZ and CWZ had reached 98% of designated sites with Option B+.

CD4 count machines (including 'point of care' machines) were installed at **170** sites, and **151** (89%) of these had produced at least 1 result during Q3 2015. The total number of CD4 results produced decreased from 20,977 in Q1 2015 to **19,513** during Q3 2015. 31% of these outputs were generated by 37 machines in the SW zone, implying that many CD4 machines continued to experience down-time or to be running considerably below capacity. The raised CD4 count threshold for ART eligibility may have also resulted in a decrease in the number of pre-ART patients requiring CD4 count monitoring as a large proportion will now be started on ART after an initial CD4 test.

4.2 Staffing of HIV Services

4.2.1 HTC Services

The Department for HIV and AIDS has maintained a dedicated system for professional registration and performance tracking for HTC Providers since 2011. This separate registration system is needed because HTC providers include lay persons with HTC training who are not registered with any other professional body. All HTC providers are issued with a unique ID and a professional logbook for documentation of duty stations, HTC trainings, sit-in observation and proficiency testing results. Logbook holders are requested to record the total number of tests done at the end of each month. Logbooks are routinely reviewed during quarterly supervision and key performance data for each provider are summarized on the site supervision forms.

	2014 Q4	2015 Q1	2015 Q2	2015 Q3
Sites visited	718	717	718	727
Sites with any tests done	660 92%	666 93%	674 94%	684 94%
Sites with registered HTC staff	659 92%	670 93%	671 93%	669 92%
Total HTC staff at visited sites	3,574	3,692	3,830	3,933
Staff with any test done	2,195 61%	2,249 61%	2,495 65%	2,287 58%
Staff with 300+ tests done this quarter	286 11%	268 10%	326 11%	474 17%
Logbooks reviewed	2,526 71%	2,559 69%	2,870 75%	2,856 73%
HTC staff participating in PT this quarter	170 7%	1,651 65%	931 32%	209 7%
Total tests (HTC register)	464,292	480,249	494,006	625,803
Tests accounted for by individual staff	331,981 72%	321,858 67%	380,159 77%	443,193 71%
Source: logbooks	303,046 91%	291,525 91%	359,042 73%	420,985 95%
Source: HTC register	28,935 9%	30,333 9%	21,117 6%	22,208 5%
Total tests by staff with 300+ tests	145,300 44%	134,272 42%	166,291 44%	263,234 59%

669 (92%) of the 727 visited facilities had registered HTC providers and **684** (94%) sites had performed at least one test during Q3 2015. **2,856 (73%)** of **3,933** HTC providers had their logbooks available for review.

According to the 2,856 reviewed logbooks, **209 (7%)** HTC providers had participated in proficiency (panel) testing (PT) this quarter. This is lower than the participation rate from the previous quarters. Only 73% of providers at the visited sites had their logbooks available for review and documentation of PT may have been incomplete. The national HIV reference laboratory is aiming to organize six monthly PT rounds for all practising HTC providers. However, the distribution of PT panels from the reference lab was delayed this quarter and district lab supervisors did not receive adequate PT supplies for their sites.

443,193 (71%) of all 625,803 tests conducted this quarter (according to HTC register reports) were accounted for by individual HTC staff working at the visited sites. **420,985 (95%)** of these tests were documented in the

reviewed logbooks and an additional **22,208 (5%)** could be attributed to individual providers from staff codes in the HTC registers. **474 (17%)** of 2,287 providers with documented activity had tested 300 or more clients this quarter. A dedicated full-time HTC provider is expected serve 300 clients per quarter (average of 5 clients per day for 60 working days per quarter). The 474 HTC providers who met or exceeded this target produced **263,234 (59%)** of the total number of tests accounted for by individual staff this quarter.

4.2.2 ART/PMTCT

Integrated HIV program supervision has included a staffing census for ART clinics since Q3 2014. This census is implemented during the site visits, indicating all staff members who actually worked at the ART clinic on the most recent clinic day. The census is designed to provide an accurate snapshot of the actual staffing of ART services each quarter. The numbers collected may be slightly lower than longer term averages, because around 100 service delivery staff are themselves participating in the supervision exercise and will not be counted as having worked in their ART clinic during the supervision period. The table below shows that total staffing levels have been fairly consistent over the last 3 quarters.

In October 2015, **701** clinicians (physicians, clinical or medical officers); **983** nurses and **967** auxiliary staff (health surveillance assistants, clerks, etc.) were working in ART clinics in Malawi.

	2014 Q4		2015 Q1		2015 Q2		2015 Q3	
Clinicians	617	26%	604	27%	652	27%	701	26%
Nurses	855	37%	814	36%	892	36%	983	37%
Pharmacy staff	11	0%	14	1%	12	0%	15	1%
Auxiliary Staff	847	36%	837	37%	891	36%	967	36%
Total	2,330		2,269		2,447		2,666	

An estimated 2.8 million ART patient visits are currently managed at the 717 ART sites per annum, based on approximately 585,000 patients alive on ART and an average dispensing interval of 2.5 months. With 260 working days per year, an average of 10,800 patient visits are therefore managed by the ART sites per working day. At current staffing levels, this translates into an average of **15** ART patient visits per clinician and **11** per nurse per day. This approximate HRH capacity assessment does not take account of site-specific differences in patient burden and staffing levels and there are several medium and high burden sites with sub-optimal staffing. However, the national treatment program is fully decentralized to the health centre level and the program continues to devolve the growing patient burden to peripheral facilities. Since 2011, the steepest increase in ART patient numbers has been recorded at the 300 small peripheral sites that have the largest collective staffing capacity (see Figure 4 on page 20).

5 HIV Testing and Counselling Program Outputs

HTC protocols have been revised in 2013 and a new HTC register was implemented in the course of a national re-training campaign for all HTC providers between May and November 2013. Protocol revisions include:

- Clear recommendations for re-testing based on the client’s test result and risk assessment
- Proper documentation of confirmatory testing for clients with a prior positive result (usually performed at enrolment into care).

The full national HTC data are presented in the **Appendix**.

5.1 HTC data

625,803 people³ were tested and counselled for HIV between July and September 2015. Testing outputs increased by **27%** from the previous quarter. This was most likely due to the deployment of new dedicated staff (*HIV Diagnostic Assistants, HDAs*) at about 200 facilities. HDAs are currently hired by PEPFAR implementing partner organizations and seconded to public sector facilities, primarily to boost routine provider-initiated HIV testing for patients.

586,926 (94 %) of all tests were performed at health facilities, **14,563 (2 %)** were done in stand-alone HTC sites and **24,314 (4%)** were done outside of facilities / in the community. Out of a total of **34,966** people newly diagnosed with HIV this quarter, **32,541 (93%)** were tested at health facilities, **1,064 (3%)** at stand-alone HTC sites and **1,361 (4%)** in a community-based testing.

5.2 HTC access type

361,626 (58%) of people tested were patients receiving provider-initiated testing and counselling (PITC); **261,272 (42%)** accessed voluntary counselling and testing, door-to-door, community-based testing, etc.; and **2,905 (<1%)** came for testing with a *Family HTC Referral Slip* (FRS) that was issued to a family member at a prior HTC encounter. Based on a total of 16,934 FRS issued to index clients this quarter, the successful referral rate for family members was **17%** (2905 / 16934). This is lower than in previous quarter (9%). Referral slips have remained under-utilized.

5.3 Age and sex distribution among HTC clients

Out of **625,803** people tested and counselled, **37%** were males and **63 %** were females. **44%** of females were pregnant. The proportion of males (51%) to non-pregnant females (49%) was similar, implying gender balanced access to HTC services. Pregnant women have to be excluded from this comparison because testing among pregnant women is almost entirely provider-initiated and there is no comparable access route targeting males.

143,254 (23%) of all people tested accessed HTC with their partners (as a couple).

51% of all people tested and counselled were 25 years and above, **39 %** were between 15-24 years and **10%** were children below 15 years. **3,228 (<1%)** of rapid tests done were among infants.

5.4 First time, repeat and confirmatory test results

The 2011 and 2014 Malawi Clinical HIV Guidelines stipulate: *All patients need a confirmatory HIV test to rule out any possibility of mix-up of test results or fraudulent access to ART: either at enrolment into pre-ART follow-up, or before starting ART if the test to confirm was not done in pre-ART. Children under 12 months starting ART with a positive DNA-PCR do not need another confirmatory test before starting ART, but all need a confirmatory rapid antibody test at age 12 and 24 months.*

190,051 (30%) accessed HTC for the first time and **435,752 (70%)** were repeat testers. Based on the cumulative number of people who accessed HTC for the first time, a total of **6,063,479** people have been tested since introduction of the *first time HTC access* indicator in July 2007.

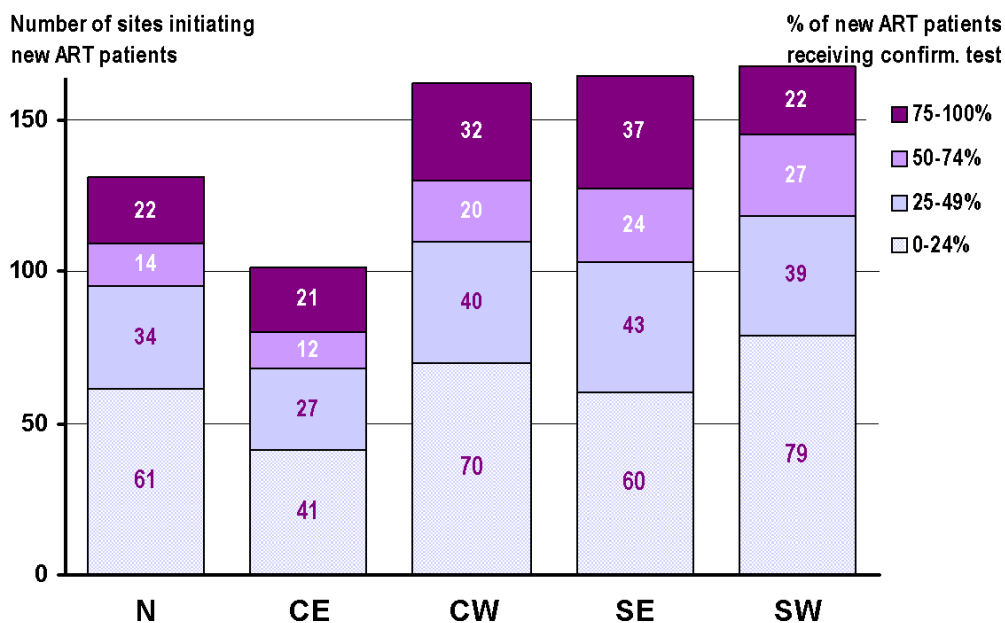
34,966 (6%) out of all clients received a positive result for the first time. Positive rapid test results among infants (**1,122**) and inconclusive test results (**2,392**) both accounted for **<1 %** of new results given to clients.

³ Reports from the HTC register are based on client encounters. It is not possible to de-duplicate people who access HTC multiple times in the reporting period. However, very few individuals come for repeat testing in less than 3 months and the number of HTC encounters in one quarter is therefore assumed to represent individuals.

419,284 (96%) of 435,752 repeat testers reported a *last negative* result. **13,904 (3%)** were reported as *previous positives* and all of these should have been classified as receiving a confirmatory test. For most of the **13,904** *previous positives*, testing was probably initiated by a health worker before enrolment into care. *Confirmatory test results* accounted for **12,513 (90%)** of *previous positive* clients. The remainder (1,391) may have been misclassified as *new positive* or *new inconclusive* because they were among clients who independently sought confirmation of their positive status. **12,513 (96%)** of 13,093 confirmatory tests were

Figure 1: Confirmatory HIV testing coverage at ART sites in the 5 zones

Num.: total confirmatory HIV tests documented in HTC registers. Denom.: total new patients initiating ART at the site



concordant positive and **580 (4%)** were classified as *confirmatory inconclusive*. This category includes parallel concordant negative and discordant test outcomes (Determine HIV1/2 and Uni-Gold HIV1/2 are used in parallel for confirmatory testing). This relatively high proportion of clients who did not have a concordant positive confirmation may be explained by selective confirmatory testing among clients with doubts about their previous positive status, but it underscores

the importance of both routine confirmatory testing before ART initiation as well as the need to continue strengthening HTC quality assurance processes.

The 13,093 confirmatory test results documented this quarter indicate that only **51%** of the 25,737 patients initiating ART this quarter received confirmatory testing and **Figure 1** shows that confirmatory testing remained low in all 5 zones. Only **134 (19%)** of facilities throughout the country had performed confirmatory testing for $\geq 75\%$ of patients newly initiated on ART. Implementation of the confirmatory testing policy will be further reinforced over the next quarters.

6 DNA-PCR testing for Early Infant Diagnosis of HIV (EID)

DNA-PCR testing is performed at 8 labs (Mzuzu Central Hospital, Mzimba District Hospital, Kamuzu Central Hospital, Queen Elizabeth Central Hospital, DREAM Blantyre, DREAM Balaka, Zomba Central Hospital and Partners in Hope, Lilongwe). EID counsellors collect infant blood samples as dried blood spots on filter paper. Health facilities are requested to maintain a standard EID DNA-PCR logbook to document EID samples and to track results. The logbook includes the dates of collection, dispatch, receipt of result from the lab and communication of the result to the mother. Supervision teams were asked to collect basic data from these logbooks.

514 (80%) of 650 sites with HIV exposed children in follow-up had collected and recorded at least 1 DNA-PCR sample during Q3 2015. A total of **8,066** DNA-PCR samples were collected and recorded. By the time the logbooks were reviewed (between 1 and 3 weeks after the end of the quarter), results had been received at the sites for **2,607 (32%)** of these specimens and **1,252 (48%)** of these results had been communicated to the mother. The proportion of results received at the sites was **52%, 37%** and **9%** for samples collected in July, August and September, respectively. A total of **110 (4%)** results received at the sites were positive.

The **8 laboratories** registered the **receipt of 7,647** DNA-PCR samples that were collected during Q3 2015. This represents 95% of the 8,066 samples recorded in the logbooks at the sites. 6,200 (81%) of the 7,647 registered samples arrived in the same quarter.

A total of **6,050** valid DNA-PCR results were dispatched from the labs in Q3 2015. **3,150 (52%)** of the dispatched results were from samples collected in Q3 2015, while 2,900 (48%) were from samples collected in the previous quarters. The median time between sample collection and dispatch of the result was **44 days**; 75% of results were dispatched between 27 and 70 days after sample collection.

3,343 (55%) of all results were from infants under 2 months old at the time of sample collection. 2,005 (33%) were 2-5 months, 558 (9%) were 6-11 months and 17 (<1%) were 12-17 months. 12 results were from older children or adults, presumably from samples sent to the lab as ‘tie-breaker’ for inconclusive rapid test results. The date of birth was missing for 115 samples.

Age at sample collection	Tot. Results	Positives	
<2 months	3,343	68	2.0%
2-5 months	2,005	113	5.6%
6-11 months	558	77	13.8%
12 months +	29	4	13.8%
(missing)	115	5	4.3%

267 (4.4%) of all results dispatched were positive. The age-specific number (%) of positive results is shown on the left. Receipt of the DNA-PCR result at the health facility is a prerequisite to updating of patient records and for appropriate clinical management. Considering the delays between

sample collection and dispatch of the test result from the lab, the child’s age at the time of dispatch of the result from the lab is a useful indicator for the process of early infant diagnosis and early initiation of ART for confirmed infected infants. The table below shows the distribution of ages when results were dispatched from the lab.

Age when result disp. from lab	Tot. Results	(Col %)	Positives	(Col %)
<2 months	293	5%	3	1%
2-5 months	4,459	74%	125	47%
6-11 months	1,090	18%	116	43%
12 months +	93	2%	18	7%
(missing)	115	2%	5	2%
Total	6,050	100%	267	100%

Out of 267 positive results dispatched, only 3 (1%) were sent before the child was 2 months old. A total of 128 (48%) positive results were sent before the child was 6 months old and 244 (91%)

were sent before the child was 12 months old. A total of 103 infants were started on ART in WHO stage 1 or 2 on the basis of confirmed HIV infection (see ART section below). This is equivalent to **42%** of the number of positive DNA-PCR results dispatched for children <12 months this quarter.

7 Blood Safety

The Malawi Blood Transfusion Service (MBTS) is striving to provide safe blood products for the entire country using voluntary non-remunerated donors and quality assured screening for transfusion transmissible infections (TTIs). For the last years, MBTS has not been able to meet the national demand and several hospitals continue to supplement or rely entirely on blood units collected from replacement donors. Complete reports from MBTS have been available throughout, but blood safety reports from health facilities have not been consistently available and it has been challenging to compile national reports relying on the data passively submitted by the sites. Therefore, the PMTCT/ART supervision teams were tasked with active collection of blood donor and cross-matching data from all visited health facilities. Some of the visited laboratories were not using the standard MOH registers and the aggregation of data for reporting may have been affected by incomplete documentation at some sites.

A total of **17,773** blood units were collected in Malawi during Q3 2015. MBTS collected **10,454 (58%)** of these, **100%** of which were screened comprehensively for the relevant TTIs (HIV, Hepatitis B, Hepatitis C, syphilis, malaria). In addition, **54** hospitals in Malawi collected a total of **7,319** units from replacement donors. **6,191 (85%)** of these units were screened for at least the 3 key TTIs (HIV, HepB and syphilis) and

3,368 (54%) of these were also screened for HepC and malaria. This means that a total of **16,645 (94%)** of all 17,773 units collected by MBTS and from replacement donors this quarter were screened at least for HIV, HepB and syphilis. Based on the blood donor registers at the sites that collected blood from replacement donors, 17 units were screened for HIV and HepB only and 2 were screened only for HIV. 1,109 were screened with any other combination of tests for TTIs.

A total of **10,685** potential replacement donors were documented in the blood donor registers at the facilities and 7,319 (68%) of these ended up donating. Facilities may have used different screening algorithms and potential donors may have been excluded on the basis of different criteria, including TTIs, blood group, haemoglobin concentration and/or clinical conditions. Testing for less prevalent TTs may have only been carried out for donors who passed the screening for more common conditions. In total, 80% of potential donors were tested for HIV, 80% for HepB, 80% for syphilis, 67% for malaria and 48% for HepC. Detailed data on outcomes of individual tests among all potential blood donors are presented in the Appendix.

8 Post Exposure Prophylaxis (PEP)

A total of **1,593** persons received PEP during Q3 2015. This is a 13% increase from the previous quarter (1,408).

9 Provider-Initiated Family Planning (PIFP)

The Integrated Clinical HIV Guidelines encourage health workers to routinely provide condoms to all adults in pre-ART and ART clinics. Women should also be offered at least the standard injectable contraceptive (Depo-Provera) during any pre-ART or ART visit. This policy aims to address the significant unmet need for family planning that had been observed among HIV patients in Malawi and to reduce the number of unwanted pregnancies among HIV-infected women (*PMTCT Prong 2*). HIV program reporting on PIFP is limited to women who received an injection of Depo-Provera in pre-ART and ART clinics during the last quarter. The report does not account for family planning need nor does it include women who accessed family planning services outside of HIV clinics.

Table 3: Number and % of women retained in HIV care * who were on injectable contraceptives (Depo) by the end of 2015 Q3.

Zone	Pre-ART		ART		Both patient groups	
	Tot. women	On Depo	Tot. women	On Depo	Tot. women	On Depo
NZ	568	61 11%	33,122	7,934 24%	33,691	7,995 24%
CEZ	388	59 15%	26,850	4,364 16%	27,238	4,423 16%
CWZ	3,224	1,099 34%	68,425	19,072 28%	71,649	20,171 28%
SEZ	2,950	1,051 36%	105,140	39,415 37%	108,090	40,467 37%
SWZ	5,098	814 16%	110,555	22,467 20%	115,653	23,280 20%
Malawi	12,229	3,084 25%	344,093	93,252 27%	356,322	96,336 27%

* estimated from the total number of patients retained in pre-ART and ART, multiplied by the proportions of females and adults registered

Table 3 shows that **96,336 (27%)** of 356,322 women in care received Depo-Provera from HIV clinics in Q3 2015. The SE Zone had achieved the highest coverage among women in pre-ART and ART. Patient coverage and stock availability has slightly decreased in this quarter. 589 (82%) of ART/PMTCT sites had stocks of Depo-Provera in October 2015 from 76% in July 2015.⁴ The HIV Program is no longer supplementing FP supplies through procurement and distribution of additional Depo-Provera to sites.

⁴ Many Mission hospitals do not provide family planning.
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10 Cotrimoxazole Preventive Therapy (CPT)

All patients in HIV care are universally eligible for CPT in order to reduce the frequency and severity of several HIV-related diseases. Patients with confirmed HIV infection are provided lifelong CPT in pre-ART and ART clinics. CPT is also given to HIV exposed children until exposure to breast milk has stopped and HIV infection has been ruled out (usually around age 24 months). Fewer than 5% of patients are expected to require stopping of CPT due to toxicity.

Table 4 shows that **651,717 (91%)** of 718,058 all patients in care were on CPT at the end of Q3 2015.

Table 4: Number and % of patients retained in HIV care who were on cotrimoxazole and isoniazid preventive therapy (CPT, IPT) by the end of 2015 Q3.

Zone	CPT									IPT	
	Exp. child		Pre-ART		ART		All patient groups		Pre-ART		
	Tot. pat.	On CPT	Tot. pat.	On CPT	Tot. pat.	On CPT	Tot. pat.	On CPT	Tot. pat.	On IPT	
NZ	8,723	6,918 79%	2,276	2,217 97%	58,532	52,549 90%	69,531	61,684 89%	2,276	1,978 87%	
CEZ	7,882	6,303 80%	1,811	1,803 100%	46,826	43,623 93%	56,519	51,729 92%	1,811	1,644 91%	
CWZ	16,356	13,670 84%	10,336	7,408 72%	118,393	110,819 94%	145,085	131,898 91%	10,336	6,169 60%	
SEZ	30,751	26,992 88%	11,945	10,884 91%	169,473	155,986 92%	212,169	193,862 91%	11,945	9,687 81%	
SWZ	28,745	25,803 90%	16,849	15,798 94%	189,160	170,942 90%	234,754	212,544 91%	16,849	13,996 83%	
Malawi	92,457	79,687 86%	43,217	38,111 88%	582,384	533,919 92%	718,058	651,717 91%	43,217	33,474 77%	

10.1 Intensified TB Case Finding (ICF)

TB is one of the most important HIV-related diseases in Malawi and a considerable proportion of (mainly early) deaths on ART are attributed to undiagnosed TB. ICF is carried out using a standard symptom checklist at every HIV patient visit. ICF outcomes are documented on HIV exposed child, pre-ART and ART patient cards, but routine M&E reporting is currently limited to ART patients in order to reduce the burden of reporting secondary cohort outcomes. It is assumed that implementation of ICF is similar in pre-ART and exposed child follow-up.

577,441 (98%) of all patients retained on ART were screened for TB at their last visit before end of September 2015. As of that visit, **3,270 (1%)** patients were new TB suspects and had presumably been referred for examination by a clinician and for TB investigations. **802 (<1%)** patients had confirmed TB (clinical or lab based). Out of these, **729 (91%)** were confirmed to be on TB treatment and **73 (9%)** had not yet started or had interrupted TB treatment. An excerpt from the data in the **Annex (Cumulative ART outcomes)** is shown below.

Current TB status among ART patients (ICF)

ICF not done (Current TB status unknown/ not circ)	4,943	1%
ICF done	577,441	99%
TB not suspected	573,369	99%
TB suspected	3,270	1%
TB confirmed	802	0%
TB confirmed, not on treatment	73	9%
TB confirmed, on TB treatment	729	91%

10.2 Isoniazid Preventive Therapy (IPT)

All pre-ART patients with a negative screening outcome for TB symptoms are eligible for IPT. The first (large scale) distribution of isoniazid and pyridoxine for the HIV programs reached the sites during July 2012. **33,474 (77%)** of 43,217 patients retained in pre-ART were on IPT by the end of September 2015. Isoniazid was in stock at 621 facilities during the October 2015 supervision visit.

11 HIV-Related Diseases

Table 5 shows the number of patients treated for key HIV-related indicator diseases. **4,346** patients were started on TB treatment this quarter and HIV status was ascertained for **3,973 (91%)**. **2,230 (56%)** of these were HIV positive and **1,573 (71%)** of all HIV positives were already on ART when starting TB treatment. The Diflucan (fluconazole) donation program has received renewed attention and significant stocks of fluconazole were distributed to all sites in November 2012. In Q3 2015, **525** and **808** patients received Diflucan for acute cryptococcal meningitis and oesophageal candidiasis, respectively. **323** patients with Kaposi sarcoma were registered for ART in this quarter.

Table 5: Number new cases of key HIV-related diseases registered per quarter (KS = Kaposi Sarcoma, CM = cryptococcal meningitis, OC = oesophageal candidiasis).

	TB				KS *	CM *	OC *
	Tot. cases	HIV status asc.	HIV positive	Already on ART	Tot. cases	Tot. cases	Tot. cases
2014 Q4	4,204	3,989 95%	2,103 53%	1,546 74%	336	475	663
2015 Q1	4,158	3,765 91%	1,954 52%	1,408 72%	260	865	610
2015 Q2	4,288	4,074 95%	2,200 54%	1,513 69%	265	459	599
2015 Q3	4,346	3,973 91%	2,230 56%	1,573 71%	323	525	808

12 HIV-Exposed Child Follow-Up

12.1 Methods and Definition of Indicators

There are multiple entry points into HIV exposed child follow up: children of HIV infected mothers may be enrolled at birth at maternity / postnatal ward; they may be found at Under 1 or Under 5 Clinics through active screening for HIV exposure; they may be identified when presenting sick to OPD; or they may be seen with their mothers in ART follow-up. Although the targeted enrolment age is below 2 months, children may theoretically be enrolled up to 23 months of age (when HIV infection can be ruled out by rapid antibody test and breast milk exposure is likely to have stopped).

Initial registration data and details for every visit are recorded on an *Exposed Child Patient Card* and a subset of the registration data is copied in the *HIV Care Clinic (HCC) register* (one record per patient). Registration data are reported from the HCC register on a quarterly basis. Follow-up outcomes are reported monthly, selecting children who were **2, 12 and 24 months** old in the respective reporting month. Outcomes are determined from the latest visit details recorded on each card. HIV infection status is evaluated as **known negative** if a negative DNA-PCR or rapid test result was available at the last visit; HIV infection status is evaluated as **known positive** if a positive DNA-PCR result was available at any age or a positive rapid antibody test was available from age 12 months; HIV infection status is counted as **unknown** if HIV infection has not been confirmed and/or a negative test result pre-dated the last visit (assuming on-going HIV exposure through breast milk). All children under 24 months with confirmed HIV infection and those under 12 months with confirmed HIV infection through DNA-PCR or HIV antibody and symptoms of *presumed severe HIV disease* are **eligible for ART**.

The main outcome indicator for the HIV exposed child follow-up program is **HIV-free survival at 24 months of age**. This is defined as the proportion of children who were discharged as confirmed HIV uninfected by the age of 24 months.

12.2 HIV Exposed Child Registration Data

11,357 HIV exposed children were newly enrolled into follow-up during Q3 2015; **7,558 (67%)** of these were under the age of 2 months. This represents timely enrolment for **84%** of the 9,019 known HIV exposed children discharged from maternity this quarter. The total number of new enrolments (11,357) exceeds by 2,338 (26%) the total number of known HIV exposed children discharged from maternity (9,019). This apparent discrepancy may be explained by delayed enrolment of infants born in previous quarters; by double-counting of infants who transferred between sites; or by identification and enrolment of additional HIV exposed infants after birth. Overall, enrolment into follow-up for known HIV exposed infants appears to be almost complete.

The documentation of follow-up outcomes, particularly the updating of DNA-PCR results on patient cards, remained incomplete at several sites. This has led to an underreporting of ascertainment of HIV status among the 2-month old cohort.

12.3 Birth Cohort Outcomes

There were **7,800** infants in the **2-month age cohort**. **2,362 (30%)** had received a DNA-PCR result. **43 (2%)** of these were confirmed HIV infected. An additional **20** infants were diagnosed with *presumed severe HIV disease*, which means that a total of **63** infants were eligible for ART. **33 (52%)** of these had started ART. The proportion of positives starting ART is slightly lower than the previous quarter (57%). Out of the entire 2-month age cohort, **7,020 (92%)** were retained in exposed child follow-up, **33 (<1%)** had started ART and **34 (<1%)** were discharged confirmed uninfected⁵. **19 (<1%)** were known to have died and **526 (7%)** had been lost to follow-up.

There were **8,434** children in the **12-month age cohort**. Current HIV infection status was known for **4,056 (48%)** children (DNA-PCR or rapid antibody test) and **170 (4%)** of these were confirmed HIV infected. **20 (<1%)** additional children had been diagnosed with *presumed severe HIV disease*, which means that a total of **190** children were eligible for ART. **175 (92%)** had started ART. Out of the entire age cohort, **6,148 (75%)** were retained in exposed child follow-up, **175 (2%)** had started ART and **73 (<1%)** were discharged confirmed uninfected.⁵ **1,747 (21%)** were lost to follow-up and **74 (<1%)** were known to have died.

There were **8,057** children in the **24 month age cohort**. Current HIV infection status was known for **3,930 (49%)** children (DNA-PCR or rapid antibody test) and **225 (6%)** of these were confirmed HIV infected. **10** additional children had been diagnosed with *presumed severe HIV disease*, which means that a total of **235** children were eligible for ART. **212 (90%)** of these had started ART. Out of the entire age cohort, **842 (11%)** were retained in exposed child follow-up, **212 (3%)** had started ART and **3,540 (45%)** were discharged confirmed uninfected. **3,132 (40%)** were lost to follow-up and **103 (1%)** were known to have died.

Confirmed HIV-free survival at age 24 months in this quarter remained implausibly low at **45%**. This was related to the fact that only 49% in this cohort had a known HIV status. 4,127 (51%) children were classified as 'current HIV infection status unknown' and many of these may be among the 3,132 children lost to follow-up and the 103 children who had died. However, 842 (11%) were retained in follow-up beyond age 24 months and a final rapid test was not available for these children, possibly due to continued breast feeding.

⁵ A small number of children may be rightfully discharged as 'confirmed uninfected' by 2 or 12 months of age, provided that HIV exposure through breast milk has definitely stopped (e.g. maternal death) and a negative HIV test was obtained at least 6 weeks thereafter.

There are still problems with scheduled HIV testing (and documentation of test results) at 6 weeks, 12 and 24 months of age.

13 Pre-ART

13.1 Pre-ART Registration Data

A total of **6, 929** patients were newly registered for pre-ART follow-up in Q3 2015. **587 (5%)** of these were children aged 5-14 years. The number of new pre-ART enrolments slightly increased from the previous quarter (6,077 total, 472 children). Several sites had already established pre-ART services before July 2011 and the cumulative number of pre-ART patients ever registered was **201,804**.

13.2 Cumulative Pre-ART Follow-up Outcomes

43,217 (22%) of all patients ever registered were retained in pre-ART follow-up by the end of September 2015; **102,297 (53%)** had started ART; **46,939 (24%)** had been lost to follow-up; **1,917 (1%)** were known to have died. The proportion of patients starting ART is bound to increase in the cumulative pre-ART cohort analysis over time. Based on a subtraction of cumulative outcomes from the previous quarter, **4,411** pre-ART patients started ART during Q3 2015; **2,803** were lost to follow-up and **368** died.

CPT coverage among pre-ART patients was **88%** in Q3 2015 and IPT coverage declined slightly to **77%**. **3,084 (25%)** of 12,229 women had received Depo-Provera from their pre-ART clinic. Further details on CPT, Depo-Provera and IPT are presented in **Tables 3** and **4** in the sections above.

14 PMTCT / ART

The implementation of **PMTCT Option B+** has effectively integrated PMTCT and ART services. The program aims to initiate lifelong ART for all HIV infected women as early as possible in pregnancy. ART may be started and continued at ANC, labour and delivery, and at ART clinics. All infants born to HIV-infected women are supposed to start daily nevirapine prophylaxis for the first 6 weeks of life. Nevirapine syrup is given to women at ANC at the earliest opportunity to take home with instructions how to give it to the new-born.

14.1 Data Sources and Reporting Methods

New standard M&E tools for ANC and maternity were implemented in January 2010 and revised in Q2 2012 to reflect the Option B+ policy. ANC and maternity clinic registers and reporting forms include patient management information and all relevant data elements for the maternal and child health and HIV programs. The ANC register was specifically designed to avoid data duplication that previously affected PMTCT reports from ANC due to the inability to account for individual women's outcomes in the course of multiple visits. The cohort reporting system is designed to aggregate women's outcome data after they have completed their ANC visits. The outcome report is completed for women who started ANC 6 months before the reporting period.

From **Q2 2015**, the PMTCT data elements (HIV ascertainment and ART status) were also added to the first section of ANC reporting form that captures women's status at their first (booking) visit. The ANC report now includes the HIV and ART status at the first visit for women starting ANC in the reporting period and the final HIV and ART status of women who had completed ANC by the end of the reporting period. This addition aims to monitor PMTCT service implementation more closely in time, allowing for corrective action in the course of subsequent visits.

Data from ANC and maternity are collated and presented separately because records do not allow identification of individual women and hence are subject to double counting if not separated.

All patients starting ART are recorded using standard program monitoring tools (ART patient treatment cards and ART clinic registers). **ART baseline data** for all patients registered are reported each quarter from ART
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clinic registers. **ART outcomes** of all patients ever registered are reported after reviewing the cards of all new patients and of those who were on ART at the end of the previous quarter, updating the status of patients who have subsequently died, stopped or been lost to follow-up. Secondary outcomes such as current regimen, CPT status, side effects, adherence and TB status are reported for all patients retained on ART.

ART scale-up has resulted in a growing proportion of HIV-infected women who are already on ART when getting pregnant. Implementation of *Option B+* will further increase ART coverage in this group. **Maternal ART coverage** is estimated from the number of pregnant women who were already on ART when getting pregnant (**maternity reports**) *plus* those who newly started ART when pregnant (**ART reports**).

Maternity reports capture ART status at the time of delivery (up to the time of discharge from the postnatal ward). The timing of ART initiation is categorized into: (any time) before pregnancy; during 1st / 2nd trimester; during 3rd trimester; during labour. About 97% of pregnant women in Malawi attend ANC, but only 83% of women in the general population deliver at a health facility in Malawi. Maternity reports therefore have the potential for undercounting the number of mothers and infants receiving ARVs. However, there is evidence from ANC and maternity reports that almost all of the known HIV infected women deliver at health facilities. ARV coverage among known positives is therefore reliably calculated from maternity reports. Women admitted at maternity who are referred to another facility before / after delivery are double-counted in aggregated maternity data. Assuming the probability of referral is independent of ART status, the number of women already on ART when getting pregnant is therefore **adjusted** by the overall proportion of referrals among women admitted to maternity.

ART program reports capture pregnancy (and breastfeeding) status at the time of *ART initiation*, providing information on the number of new women starting ART while pregnant (or while breastfeeding). ART reports do not capture women who become pregnant after starting ART. For the estimation of maternal ART coverage, the number of women starting ART in pregnancy is **adjusted for**:

a) Double-counting of women starting ART in pregnancy and subsequently transferring to another site. These women are counted multiple times as 'pregnant at the time of starting ART' in the quarterly ART cohort reports because the disaggregation of age, sex and reason for starting ART applies to all patients newly registered in the quarter, including transfers in. Separate *ART 'survival' analyses* are collected each quarter for women started under Option B+. The proportion of women transferred within 12 months of registration is used to adjust the quarterly number of pregnant women starting ART for transfers.

b) Failed ART initiation is thought to be the main underlying reason for early loss to follow-up among the Option B+ cohort. Patients are recorded on patient cards and in clinic registers when the first supply of ARVs is dispensed and all new entrants are counted as ART initiations in the quarterly ART cohort report. Recent operational studies indicate that most pregnant women lost to follow-up within the first 6 months never return after this first dispensing visit and many of these may have never actually started taking ART. The proportion of women lost to follow-up in the 6-month survival analysis is therefore used to adjust the number of pregnant women starting ART in the quarterly ART cohort reports for *failed initiations*.

Infant PMTCT coverage is estimated from maternity reports, based on the number of infants born to known HIV-infected women and discharged alive who started nevirapine prophylaxis.

Coverage is calculated by dividing the number of patients served by population denominators. The denominators are derived from expected pregnancies based on population projections and HIV prevalence from epidemiological surveillance (source: 2015 Spectrum model for Malawi). There are an estimated 14,926 HIV infected pregnant women in the population⁶ per quarter (1/4 of 59,704 in 2015).⁶

⁶ 2015 Spectrum estimates based on current definition of eligibility for ART in Malawi (CD4<500, Option B+, UT for U5).
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14.2 ARV Coverage among Pregnant / Breastfeeding Women and Exposed Infants

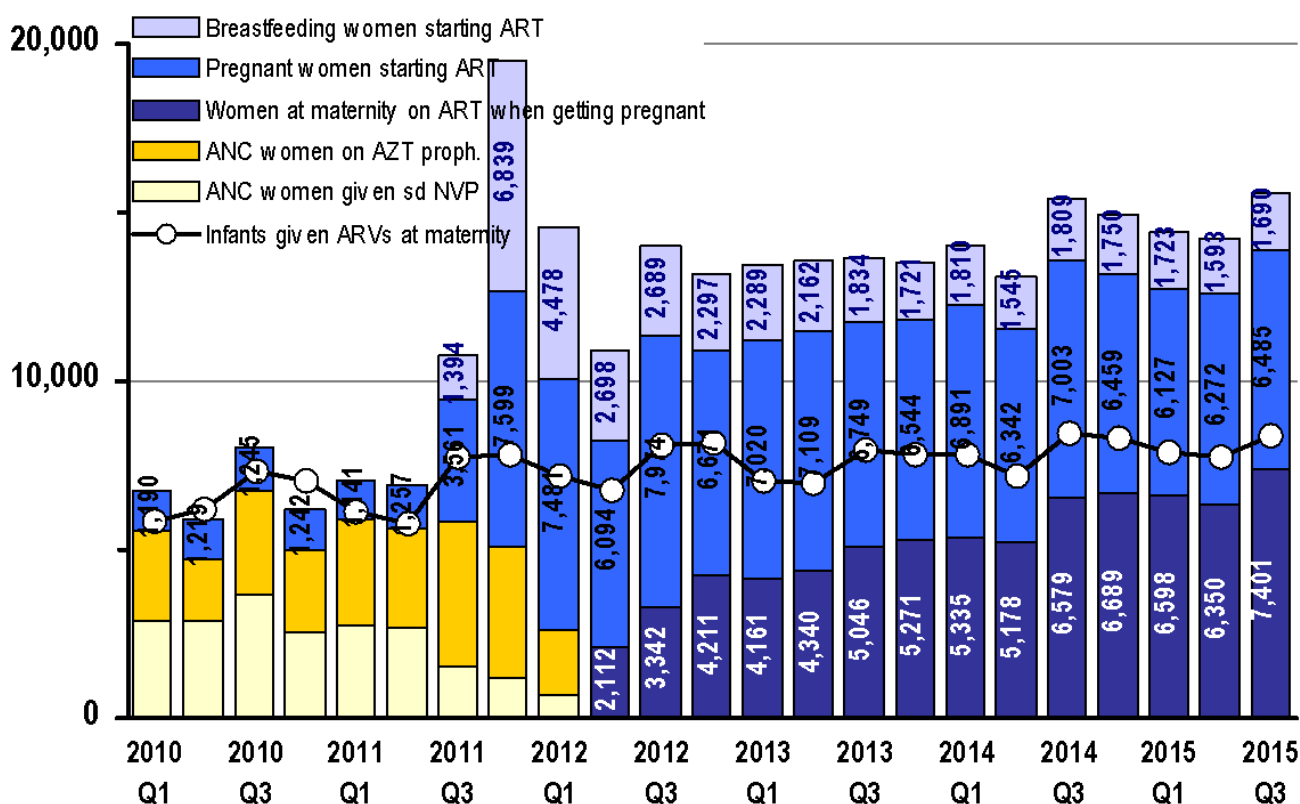
11,823 (79%) of the estimated 14,926 HIV infected pregnant women in Malawi this quarter were on ART. This is based on **7,031**⁷ women at maternity who were already on ART when getting pregnant and **4,792**⁸ women who newly initiated ART in pregnancy. This is a slight increase in ART coverage from 70% in the previous quarter.

An additional **1,555**⁹ breastfeeding women started ART due to **Option B+** (in WHO clinical stage 1 or 2), bringing the total number newly started on ART under **Option B+** to **6,347**. Most women starting ART while breastfeeding were probably identified late in maternity or early in the postnatal period, but this group may also include some women who re-initiated after interrupting ART in pregnancy. **8,414** infants were confirmed to have started NVP prophylaxis at maternity.

Figure 2 shows the transition from prophylactic ARV regimens for HIV infected mothers to universal ART under **Option B+** (registration data; not adjusted as above). The (less effective) single dose NVP regimen and AZT combination prophylaxis had been phased out by April 2012. The average number of pregnant women registered for ART each quarter **increased almost 6-fold** from **1,221** in the 12-month period before introduction of Option B+ to an average of around **6,500** since Q4 2011.

Figure 2: Transition from prophylactic ARV regimens for PMTCT to Option B+ in Malawi

Women who moved to Option B+ from sdNVP / AZT were double counted between Q3 2011 - Q1 2012. It is likely that <12,000 total women were on ARVs during these quarters. Data on women already on ART when getting pregnant are only available from Q2 2012.



⁷ 7,401 women who started ART before pregnancy admitted at maternity; reduced by 5% to adjust for double-counting of 7,272 referrals among 132,255 total admissions.

⁸ 6,485 women registered at ART clinics who were pregnant at the time of starting ART; a) 8% are discounted to adjust for double-counting of transfers based on 752 of 9,082 women who transferred within 12 months of registration (12 month Option B+ survival analysis); b) 19.7% are discounted to account for presumed failed ART initiations based on 1,560 of 7,928 women lost to follow-up within 6 months of registration (6 month Option B+ survival analysis).

⁹ 1,690 women registered at ART clinics who were breastfeeding at the time of starting ART; reduced by 8% to adjust for double-counting of transfers based on 752 out of 9,082 women who transferred within 12 months of registration (12 month Option B+ survival analysis). Failed ART initiations are thought to be less common among this group, so no further adjustment is made.

14.3 HIV Services at ANC

The full national data from ANC are presented in the **Appendix**.

14.3.1 HIV Ascertainment and ART Coverage

Booking cohort:

150,456 women attended ANC for their first visit between July and September 2015. This is 91% of the estimated 166,000 pregnant women in the 2015 population during one quarter.¹⁰ **135,411 (90%)** of women in the booking cohort had their HIV status ascertained at the first visit. Out of these, **12,636 (9%)** presented with a valid previous test result and **122,775 (91%)** received a new test. A total of **10,062 (7%)** of women were found HIV positive: **5,368 (53%)** of these from a documented previous test and **4,694 (47%)** from a new test. **9,043 (90%)** of all positives were on ART: **4,849 (54%)** of these were already on ART when starting ANC and **4,194 (46%)** newly started ART at their first ANC visit. Out of these, **3,469 (83%)** were in their 1st or 2nd trimester and **725 (8%)** were in the 3rd trimester of pregnancy.

Outcome cohort:

156,851 women had started ANC between January and March 2015 and their outcomes were reported between July and September 2015. Only **38,323 (25%)** of women in this cohort attended the recommended minimum of 4 focussed ANC visits.

142,701 (91%) of the outcome cohort had their HIV status ascertained at least once in the course of ANC. This is similar to the previous quarter (91%). **12,197 (9%)** presented with a valid documented previous HIV test result and **130,504 (91%)** received a new HIV test result at ANC. A total of **11,113 (7.8%)** women were found HIV positive. This is slightly lower than the latest Spectrum projections (9.0% HIV prevalence among pregnant women in 2015).⁶

10,099 (91%) of (known) HIV infected women were on ART by the end of ANC. This represents **68%** coverage of the estimated 14,926 HIV positive pregnant women per quarter at the population level. Of the **10,099** ANC women who were known to receive ART, **5,045 (50%)** were already on ART when starting ANC, **4,047 (40%)** initiated before 28 weeks of pregnancy and **1,007 (10%)** initiated during the last trimester of pregnancy. **10,317 (93%)** of HIV infected women at ANC were on Cotrimoxazole Preventive Therapy. **9,433 (85%)** of known HIV infected women attending ANC received the infant dose of ARVs (nevirapine syrup) to take home.

14.3.2 Syphilis Screening

30,074 (19%) of women in the outcome cohort were tested for syphilis and **612 (2%)** were syphilis positive. The low testing rate probably explains the higher (2%) than expected proportion (<1%) of positives as the testing was likely selective of those suspected to be positive.

14.4 HIV Services at Maternity

The full national data from maternity are presented in the **Appendix**.

Between July and September 2015, **124,983** women were admitted for delivery to maternity; **7,272** of these were referred to another facility before delivery, resulting in **132,255** total admissions to maternity during Q3 2015. Out of all admissions, **124,100 (96%)** delivered at health facilities, while **4,867 (4%)** had already delivered before reaching a facility. The **124,100** facility deliveries represent **75%** of the estimated 166,000

¹⁰ Estimated as ¼ of 664,000 births projected for 2015 (Demographic Proj Spectrum 2015).

Malawi National Statistics Office. (2008). Malawi Population Projections 2008-2030.

Retrieved from http://www.nsomalawi.mw/images/stories/data_on_line/demography/census_2008/MainReport/ThematicReports/Population Projections Malawi.pdf

quarterly deliveries in the population in 2015 which is less than the 83% reported in the Integrated Household Survey Report of 2010-2011.

A total of **121,450 (96%)** deliveries were conducted by skilled birth attendants, **604 (<1%)** by paramedical staff and **4,605 (4%)** were not attended by any of the above (probably mainly among women who delivered before reaching maternity). **16,310 (12%)** of women developed obstetric complications. The most common leading complications were obstructed / prolonged labour (**5,637** cases) and post-partum haemorrhage (**1,811** cases). A total of **128,967** babies were born, **124,549 (97%)** were singletons and **4,418 (3%)** were twins/multiples. There were **126,776 (98%)** live births and **2,191 (2%)** stillbirths. **125,653 (99%)** of babies born alive were discharged alive and **1,123 (1%)** died before discharge. **126,556 (>99%)** of women were discharged alive and **103 (<1%)** women died before discharge, which is equivalent to a maternal mortality ratio of **81 per 100,000** live births among women attending maternity.

14.4.1 HIV Ascertainment at Maternity

129,887 (97%) women had their HIV status ascertained at maternity. Out of these, **126,231 (97%)** presented with a valid previous HIV test result and **3,656 (3%)** received a new HIV test result. A total of **9,868 (8%)** women were HIV positive and **120,019 (92%)** were negative. The **129,887** women whose HIV status was ascertained at maternity represent **78%** of the expected 166,000 women delivering in the population.

HIV exposure status was ascertained for **122,547 (98%)** out of 125,653 babies born and discharged alive. **9,019 (7%)** of these were born to a known HIV positive mother.

14.4.2 ARV Coverage at Maternity

A total of **9,659 (98%)** of known HIV infected women admitted to maternity received ART. Out of these, **7,401 (77%)** had started ART before pregnancy, **1,207 (12%)** initiated ART during the 1st or 2nd trimester, **890 (9%)** initiated during the 3rd trimester and **161 (2%)** initiated ART at maternity.

A total of **8,414 (93%)** of 9,019 infants who were known HIV exposed and discharged alive started daily NVP prophylaxis at maternity. This represents **56%** coverage of the estimated 14,926 HIV exposed infants born in the population in this quarter.

15 ART Access and Follow-Up Outcomes

The full national data from the ART Program are shown in the **Appendix**.

15.1 New ART Registrations during Q3 2015

By the end of September 2015, there were **724 static ART sites** in Malawi, managed by government, mission, NGOs and the private sector. Out of these, **92** were ART facilities in the private sector, charging a nominal MK500 per monthly prescription of drugs per patient.

Similar to previous quarter, the electronic reporting system supported by MSF in Chiradzulu district experienced a prolonged technical fault and accurate ART cohort reports for the 10 affected facilities could not be obtained in time for this report. MSF therefore provided estimates for Q3 2015, for these 10 sites. This has led to some inconsistencies in the data presented.

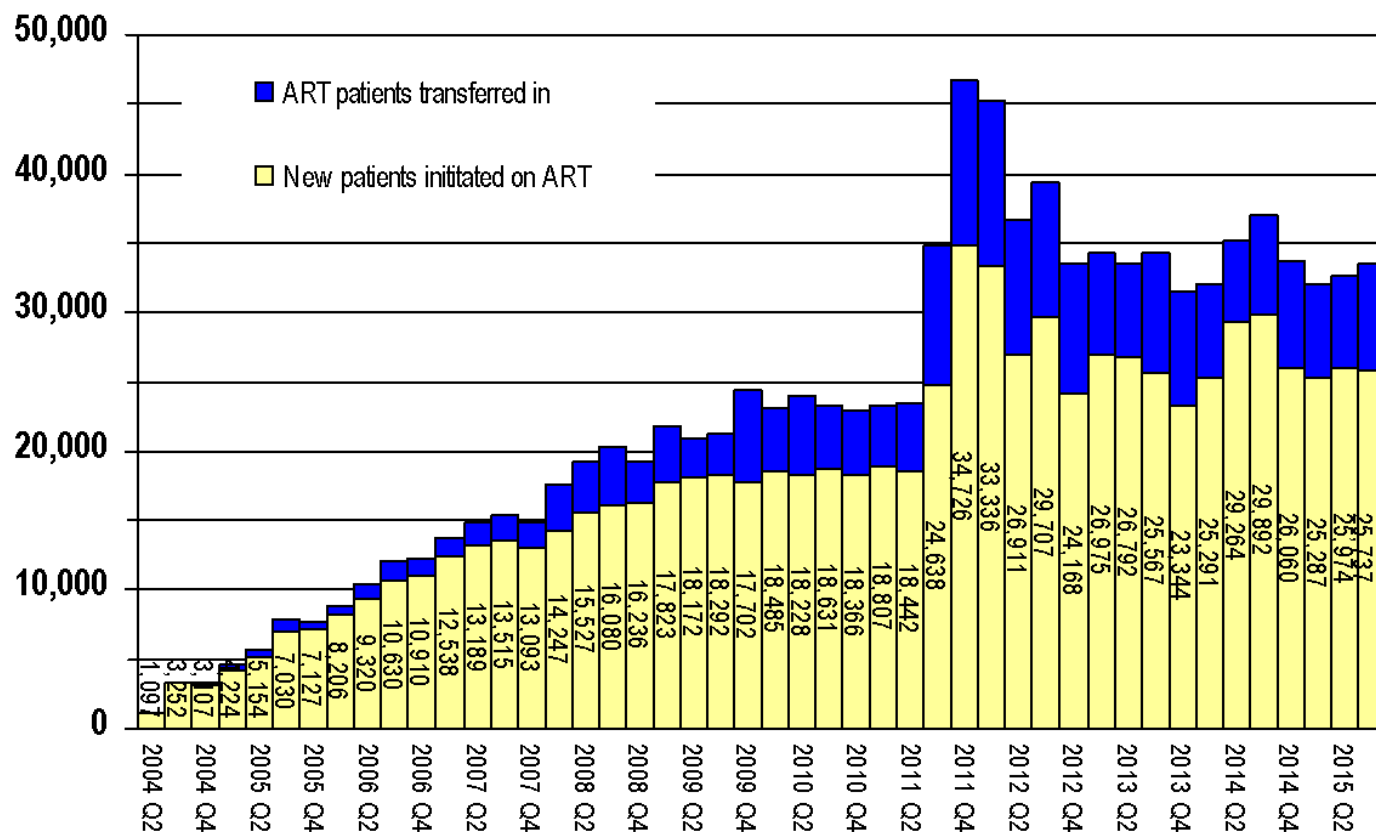
Implementation of the Malawi Integrated Clinical HIV Guidelines started in July 2011, triggering a massive surge in new ART initiations (see **Figure 3**). **25,737** patients initiated ART in Q3 2015 and **7,328** patients were registered as a transfer in (already on treatment; 22% out of all 33,474 clinic registrations). These numbers are similar to previous quarter.

Among all new registrations **37%** were males, **63%** were females. **6,485 (31%)** of females were pregnant. **6,475 (99.8%)** of pregnant women were started under **Option B+** (in WHO stage 1 or 2 with unknown CD4

or CD4 above 500), while 10 were in more advanced stage of HIV infection. An additional **1,690** women in WHO stage 1 or 2 were started because of breastfeeding, bringing the total number of women registered as started under **Option B+**¹¹ to **8,165**.

Figure 3: Patients newly initiated on ART and total ART clinic registrations per quarter

Total ART clinic registrations include patients who transferred between sites. This results in double counting of patients at the national level. For 'patients newly initiated on ART' every patient is only counted once.



A total of **21,603 (65%)** of all patients registered started in WHO stage 1 or 2 and **12,723 (59%)** of these started due a low CD4 count. **10,008 (30%)** of patients registered started in WHO stage 3 and **1,401 (4%)** started in stage 4.

2,607 children were registered at ART sites in Q3 2015. **672 (26%)** of these were registered under the expanded policy of universal ART for children aged 12-59 months in WHO stage 1 or 2, independent of CD4 count. **104 (4%)** of children started ART with presumed severe HIV disease, is similar to the previous quarter (113). **103** infants in WHO stage 1 or 2 started due to confirmed HIV infection through DNA-PCR, which is higher than the previous quarter (87). Early paediatric ART access has remained below expectations, but the relatively low number is consistent with reduced transmission rates due to Option B+: considering that 9,019 HIV exposed infants were identified at maternity and assuming a 2% transmission rate among the 98% of HIV positive mothers at maternity who received ART (and 20% transmission in the 2% who did not receive ART)¹², only about 235 of these known HIV exposed infants may have been infected perinatally during Q3 2015. However, considering the projected 1,560 new infant HIV infections in the 2015 population per quarter⁶, early infant treatment coverage remains low at an estimated **7%** (103 / 1,560). The most significant bottleneck for early infant treatment remains the identification of HIV infected pregnant / breastfeeding women.

¹¹ Universal ART for all HIV infected pregnant and breastfeeding women in WHO stage 1 or 2, independent of CD4 count

¹² UNAIDS Reference Group on Estimates Modelling and Projections (2011). Working paper on mother-to-child-transmission rates for use in Spectrum. Geneva, UNAIDS.

982 (3%) out of all ART clinic registrations were patients with TB: **649 (2%)** had a current and **333 (1%)** a recent history of TB. **323 (1%)** of patients registered had Kaposi's sarcoma.

15.2 Cumulative ART Registrations up September 2015

By the end of September 2015, there were a cumulative total of **1,060,580** clinic registrations, representing **849,368 (80%)** patients who newly initiated ART and **199,346 (19%)** patients who transferred between clinics. **11,866 (1%)** out of all clinic registrations were patients who re-initiated ART after treatment interruption. Out of all registrations, **36%** were males and **64%** were females, **91%** were adults and **9%** were children (<15 years). Private sector clinics accounted for **31,759 (3.0%)** of total patient registrations.

15.3 ART Outcomes

585,660 patients were alive on ART by the end of September 2015. This is equivalent to **59% ART coverage** among the estimated 1 million HIV positive population in Malawi in 2015. The number of patients on ART includes an estimated 3,137 patients in transit between sites (50% of the 6,273 patients newly registered as transferred out at sites across the country).

Out of the **849,368** patients ever initiated on ART, **585,660 (69%)** were retained alive on ART, **79,280 (9%)** were known to have died, **195,982 (23%)** were lost to follow-up and **3,448 (<1%)** were known to have stopped ART. An estimated **531,990** adults and **50,533** children (<15 years) were alive on ART by the end of September 2015.

Figure 4 Patients alive on ART at the end of each quarter, stratified by size of facility (number of patients alive on ART)

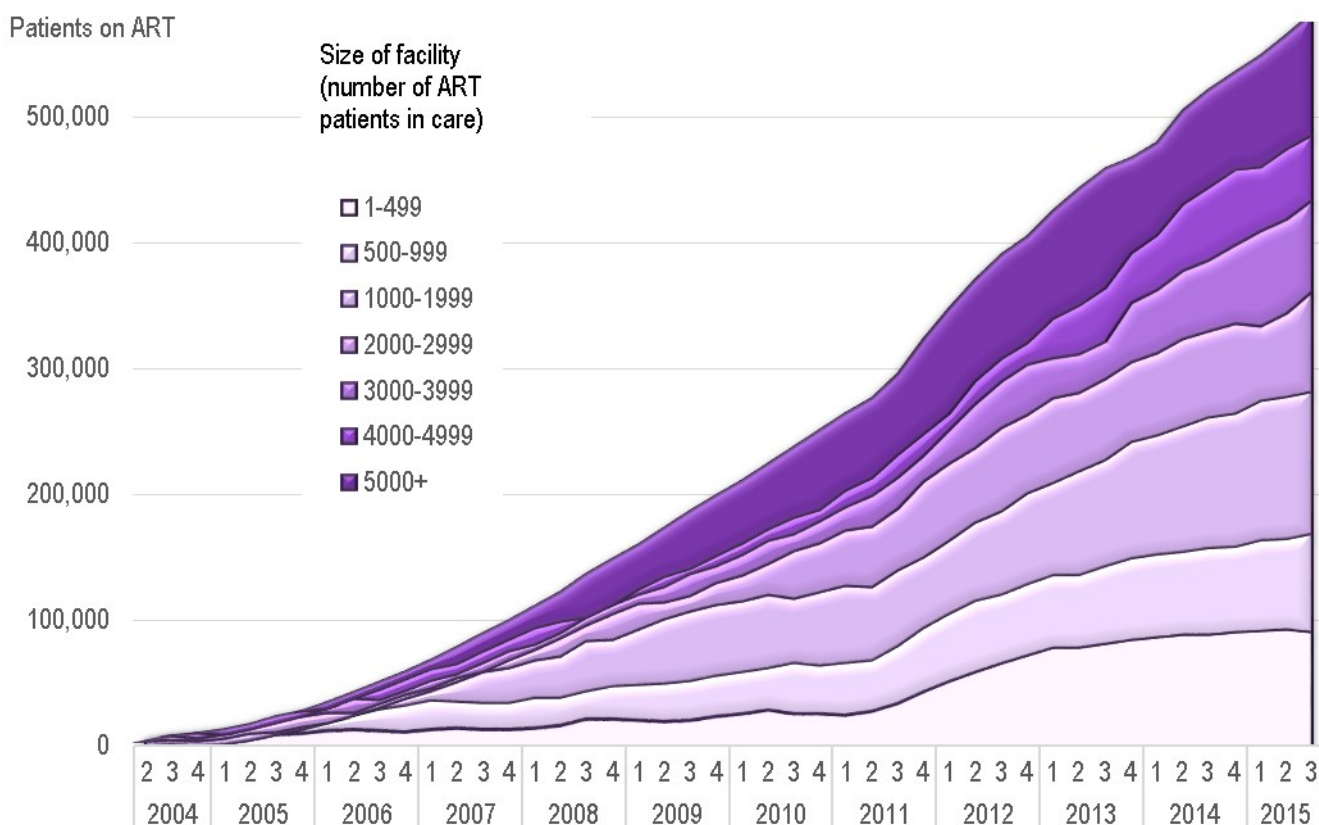


Figure 4 shows the increase of patients alive on ART by the end of each quarter. The number of patients alive on ART **increased by 17,190** in Q3 of 2015. **Figure 4** also illustrates the ongoing decentralization of Malawi's ART program. From Q3 2011, the greatest increase in ART patient numbers was seen at sites with fewer than 500 patients alive on ART. By the end of September 2015, **48%** of the national ART patient cohort was in care at sites with fewer than 2,000 patients.

Figure 5: Quarterly rates of ART drop out (ART stop, defaulters and deaths)

Numerator: new ART stops, new defaulters and new deaths in the respective quarter

Denominator: total patients retained alive at the end of the previous quarter plus new patients registered in the respective quarter)

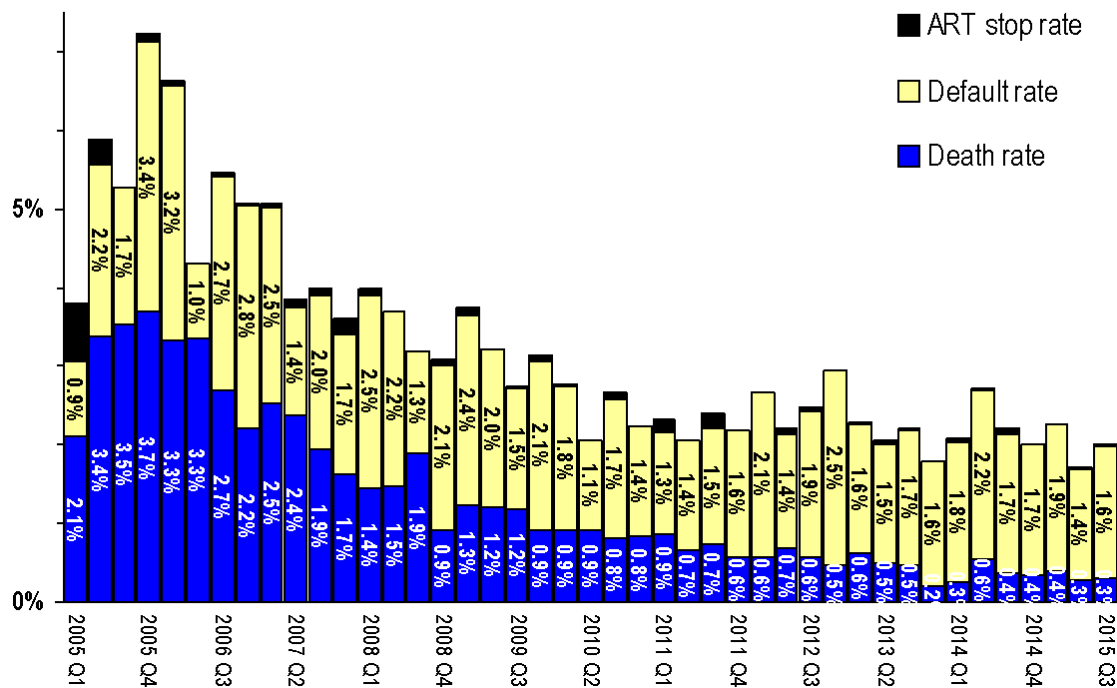


Figure 5 shows the considerable decrease of ART drop-out rates since the start of the national program. There were **1,938** new deaths, **9,908** new defaulters and **82** new stops in Q3 2015. This translates into a quarterly death rate of **0.3%** and a defaulter rate of **1.6%** among the patients alive and on treatment in this quarter. These rates were similar to the previous quarter. Based on previous operational studies, about half of the patients in stage 3 and 4 who are classified as lost to follow-up are thought to have died. There is also an indication that 10-15% of pregnant women who were registered as ‘initiated on ART’ under Option B+ may have never actually started taking ARVs due to inadequate preparation at ANC. Importantly, the ascertainment of loss to follow-up requires updating of patient treatment cards after analysis of the most recent dispensing visit. Any lack in rigour in this process will lead to a misclassification of patients who have been lost to follow-up as ‘retained alive on ART’.

By end of September 2015, a cumulative **79,280 (9%)** patients were known to have died **195,982 (23%)** were lost to follow-up and **3,448 (<1%)** were known to have **stopped ART**.

Figure 6: Patients starting ART in WHO stage 4 and deaths in the first 3 months after ART initiation. (Shown as proportions among new patients registered each quarter)

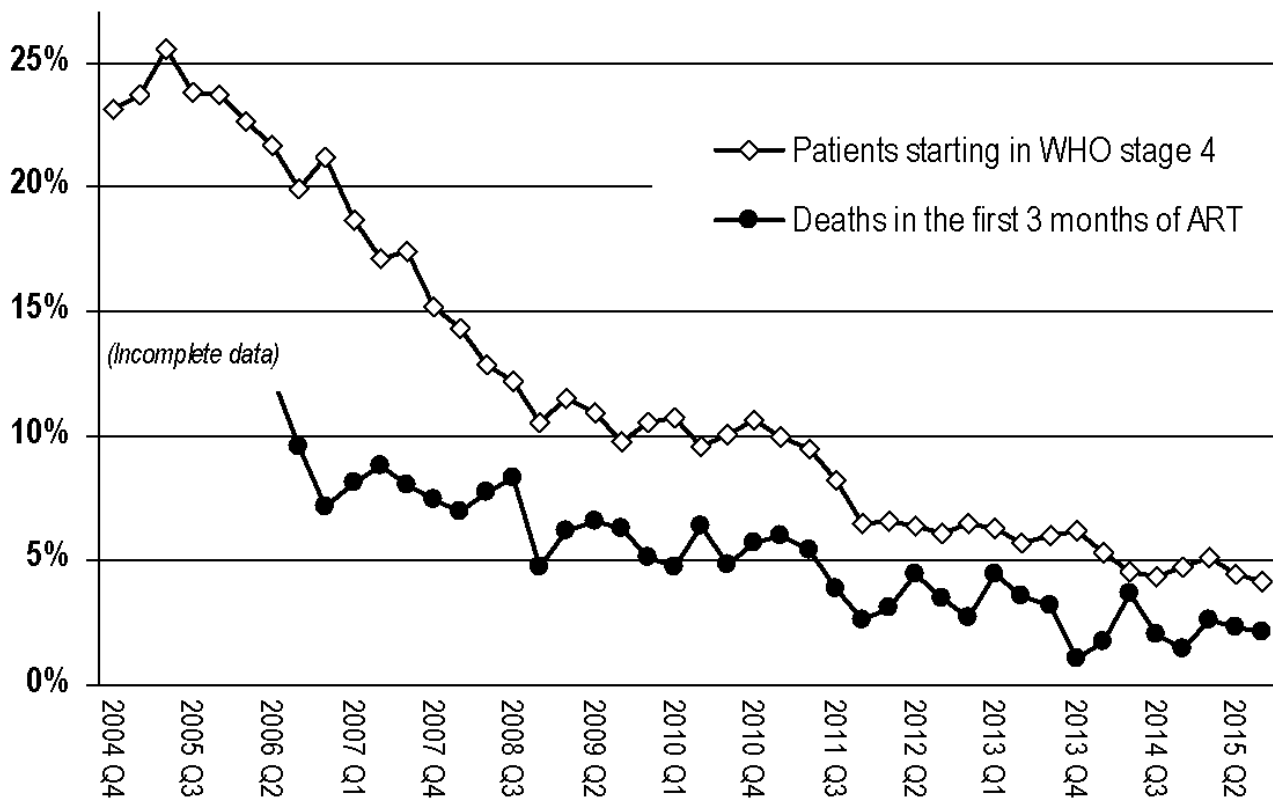


Figure 6 shows the considerable decline in **early mortality** since the start of the program. In Q2 of 2006 11% of new patients died within the first 3 months of ART initiation. There has been a remarkable decline in early mortality and the lowest point thus far has been reached in Q4 2013. The decrease in early mortality is probably mainly due to earlier ART initiation (patients in WHO stage 2 with a CD4 count below the threshold or in stage 3). It correlates well with the decline in the proportion of patients starting ART in WHO clinical stage 4 from 25% in 2005 Q2 to **4%** in Q3 2015. Slight fluctuations in the calculated early mortality rates are mainly due to inconsistent classification of month of death at the sites with electronic patient record systems. The 2014 guidelines have led to further reduction in early mortality, as more patients are started in WHO stage 1 and 2 (CD4 threshold for eligibility <500; universal ART for HIV infected pregnant and breastfeeding women and children under 5 years).

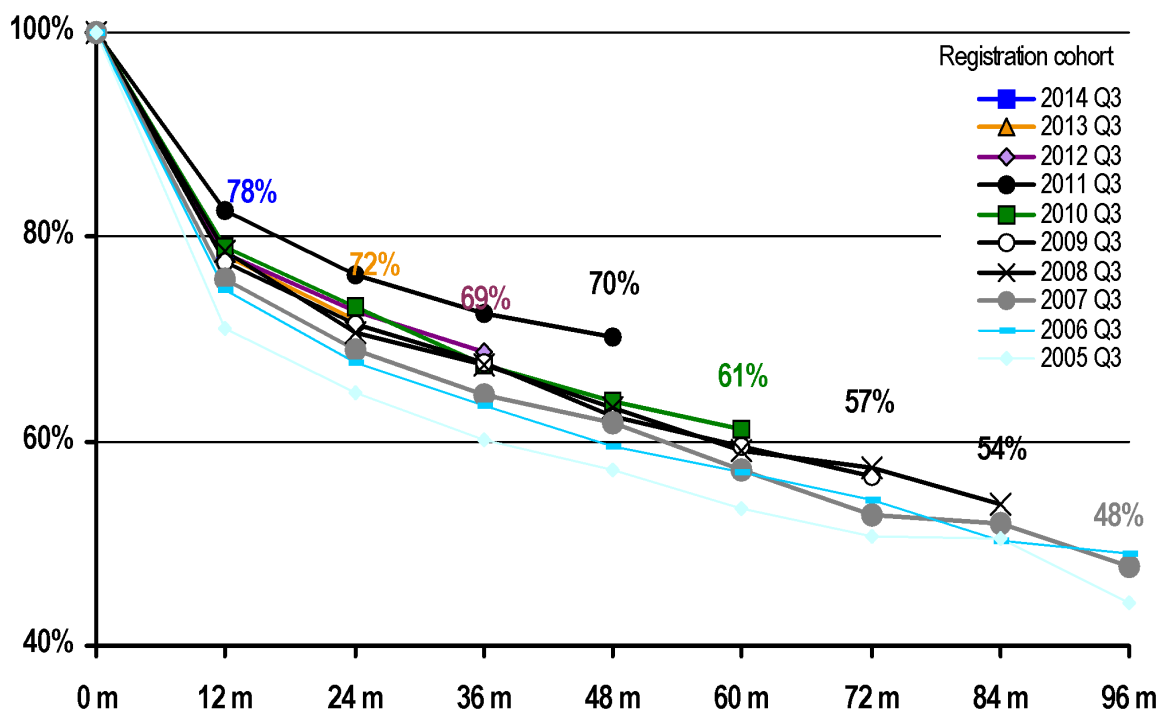
15.4 ART Cohort Survival Analysis

A 12, 24, 36, 48, 60, 72, 84, 96 and 108-month **'cohort outcome survival analysis'** was conducted for patients registered in Q3 of 2006, 2007, 2008, 2009, 2010, 2011, 2012, 2013 and 2014, respectively. A separate 12-month cohort outcome analysis was conducted for children who were under 15 years at the time of ART initiation and who registered for ART in Q3 2014. For the 13th time, a further subgroup analysis was done for women who started ART under **Option B+** during Q3 2012, Q3 2013, Q3 2014 and Q1 2015. **78% of adults** and **79% of children** were retained alive on ART after 12 months on treatment. This is similar to the previous quarter and remains below the WHO target of 85%. The majority of patients classified as lost to follow-up are likely to have stopped/ interrupted ART, but others will have transferred to another facility without notifying the previous site. Actual retention rates are thought to be about **10%** higher due to this misclassification of 'silent transfers' as 'defaulters' in clinic-based survival/retention analysis. A population-based study in Karonga district with individual linkage showed that **92%** of patients started in 2011-2012

were retained after 12 months on ART while routine monitoring data showed **79%** retention rates for the same period.¹³

Figure 7 shows the continuous improvement of long-term treatment outcomes over time. **61%** and **54%** of patients registered 5 and 7 years ago had been retained alive on ART.

Figure 7: Group cohort survival analysis: Proportion of patients retained alive on ART 12, 24, 36, 48, 60, 72, 84 and 96 months after ART initiation



6-month group cohort survival outcomes were known for **7,928** women registered as having started ART under *Option B+* in Q1 2015.¹⁴ This number is 122 (2%) higher than the number of women that registered as having started ART of 7,806. This discrepancy is likely due to data abstraction inaccuracies. The 7,928 represents 502 (6%) women who transferred out and are therefore double counted and **7,426 (94%)** patients not transferred. **5,781 (78%)** of these were retained at 6 months after registration. **1,560 (95%)** of those not retained were lost to follow-up, **31 (2%)** were known to have stopped ART and **54 (3%)** were known to have died.

12-month group cohort survival outcomes were known for **9,082** women registered as having started ART under *Option B+* in Q3 2014.¹⁴ This number is 300 (3%) higher than the number of women that registered as having started ART of 8,782. This discrepancy is likely due to data abstraction inaccuracies. The 9,082 represents **752 (8%)** women who transferred out and are therefore double counted and **8,330 (92%)** patients not transferred. **5,909 (71%)** of these were retained at 12 months after registration. **2,352 (97%)** of those not retained were lost to follow-up, **25 (1%)** were known to have stopped ART and **44 (2%)** were known to have died.

24-month group cohort survival outcomes were known for **8,633** women registered as having started ART under *Option B+* in Q3 2013.¹⁴ This number is 368 (4%) higher than the number of women that registered as having started ART of 8,265. Similar to the 6 and 12 months cohorts, the discrepancy is likely due to data abstraction inaccuracies. The 8,633 number represents **956 (11%)** women who transferred out and are

¹³ Koole, O., Houben, R. M. G. J., Mzembe, T., Van Boeckel, T. P., Kayange, M., Jahn, A., Crampin, A. C. (2014). Improved retention of patients starting antiretroviral treatment in Karonga District, northern Malawi, 2005–2012. *Journal of Acquired Immune Deficiency Syndromes* (2014), 67(1), e27–33. doi:10.1097/QAI.0000000000000252

¹⁴ Group cohort survival analyses were not available from some sites with electronic data systems. ‘Reason for starting’ may be reclassified for some patients, leading to minor inconsistencies in patients included in group cohort survival analyses.

therefore double counted and **7,677 (89%)** patients not transferred. **5,225 (67%)** of these were retained at 24 months after registration. **2,282 (91%)** of those not retained were lost to follow-up, **50 (>1%)** were known to have stopped ART and **166 (7%)** were known to have died.

1,834 (21%) of the women in the 24-month Option B+ survival cohort had initiated ART in the breastfeeding period and **2,103 (24%)** started in the third trimester / in labour; considering the 24 month median breastfeeding period in Malawi (2010 MDHS), more than half of the women in this cohort can be assumed to have stopped breastfeeding. The **67% and 66% retention rate at 24 and 36 months** after ART initiation confirms that a high proportion of women started under Option B+ **remain on ART beyond the cessation of breastfeeding.**

The 6-month retention rate is slightly higher than the previous quarter. These are satisfactory results. Most of the women lost to follow-up failed to return after their first visit and many of these may have never actually started ART due to inadequate counselling and preparation in the initial phase of implementation. The program is examining the different modes of delivery closely to further improve uptake and retention on *Option B+*.

6 month survival OptionB+

Survival and retention in ART program

*

ART cohort registration group outcomes

Total ART clinic registrations	7,928	100%
Transfers out (double counted)	502	6%
Total not transferred out (patients in cohort)	7,426	94%
Total alive on ART	5,781	78%
Total not retained	1,645	22%
Defaulted	1,560	95%
Stopped ART	31	2%
Died	54	3%

12 month survival OptionB+

Survival and retention in ART program

*

ART cohort registration group outcomes

Total ART clinic registrations	9,082	100%
Transfers out (double counted)	752	8%
Total not transferred out (patients in cohort)	8,330	92%
Total alive on ART	5,909	71%
Total not retained	2,421	29%
Defaulted	2,352	97%
Stopped ART	25	1%
Died	44	2%

24 month survival OptionB+

Survival and retention in ART program

*

ART cohort registration group outcomes

Total ART clinic registrations	8,633	100%
Transfers out (double counted)	956	11%
Total not transferred out (patients in cohort)	7,677	89%
Total alive on ART	5,179	67%
Total not retained	2,498	33%
Defaulted	2,282	91%
Stopped ART	50	2%
Died	166	7%

36 month survival OptionB+

Survival and retention in ART program

*

ART cohort registration group outcomes

Total ART clinic registrations	10,261	100%
Transfers out (double counted)	1,275	12%
Total not transferred out (patients in cohort)	8,986	88%
Total alive on ART	5,967	66%
Total not retained	3,019	34%
Defaulted	2,717	90%
Stopped ART	48	2%
Died	254	8%

15.4.1 Secondary outcomes of patients retained on ART

Secondary outcomes are known for the **582,384** patients alive on ART who remained at their sites at end of the quarter.

ART Regimens

574,610 (99%) of patients were on first line and **7,334 (1%)** were on second line regimens; **440 (<1%)** were on non-standard regimens. Non-standard regimens are not necessarily substandard regimens and include patients continuing an ART regimen that was started outside Malawi, patients in research programmes and patients in specialist care.

Among patients on first line regimens, **26,063 (5%)** were on paediatric formulations and **25,001 (96%)** of these were on the new standard first line for children (regimen 2P: AZT/3TC/NVP). By the end of September 2015, **510,798 (93%)** of patients on adult first line were receiving regimen **5A** (tenofovir / lamivudine / efavirenz). **27,334 (5%)** were on regimen 2A (zidovudine / lamivudine / nevirapine), which was the main alternative regimen for patients with stavudine side-effects before transition to regimen 5A and **1,098 (<1%)** were on regimen 1A (stavudine / lamivudine / nevirapine).

Adherence to ART

Pill counts and the number of missed doses were documented for **570,105 (98%)** out of all patients retained on ART and **520,401 (91%)** of these were classified as >95% adherent in Q3 2015. Manual estimation of adherence from pill counts is practically difficult and classification can be misleading. The ART program has switched to a direct evaluation of doses missed in 2010 to improve on accuracy of adherence assessment and plausible adherence levels are recorded with this method. However, there have also been persistent challenges with the analysis of adherence levels at sites with Electronic Data Systems (EDS) and adherence data from several of these sites could not be included in this report.

ART Side Effects

481,934 (83%) patients on ART had information on drug side effects documented at their last clinic visit before end of September 2015. This is an increase from the previous quarter (73%). This was due to a software update for the national EMR that interfered with the side effect recording. **8,457 (2%)** of patients with information had documented side-effects. The prevalence of side effects seems to have stabilized at very low levels following the full transition to regimen 5A (tenofovir / lamivudine / efavirenz) that started in July 2013.

15.5 Viral Load (VL) Monitoring

The National Treatment Program has started rolling out routine VL monitoring for patients on ART to facilitate early detection of treatment failure and timely switching to second line ART. Routine VL monitoring is scheduled at 6 months after ART initiation, at 2 years and every 24 months thereafter. Additional targeted VL testing may be carried out for patients with clinically suspected treatment failure. During Q3 2015, **9** laboratories in the national program provided VL testing for patients enrolled at the respective facilities and associated sites. All labs used the MOH information management system (**LIMS**) for registration of samples and storage of results. The following results are based on an analysis of exported LIMS data.

A total of **27,859** VL results were produced between July and September 2015. **8,065 (29%)** of samples processed were plasma and **19,783 (71%)** were DBS. For 11 results, the specimen type was not specified.

Lab	Samples Processed				Turn-around Time (Days) [§]
	Plasma	DBS	Oth/unk	Total	
DREAM Blantyre	1,708	1,603	0	3,311	41
DREAM Balaka	1,572	216	5	1,793	34
Kamuzu CH	2,336	612	1	2,949	28
Mzimba DH	0	104	0	104	8
Mzuzu CH	0	4,468	0	4,468	35
Partners in Hope	1,430	5,004	1	6,435	30
QUECH	11	971	0	982	91
Thyolo DH	1,008	3,317	3	4,328	14
Zomba CH	0	3,488	1	3,489	38
Total	3,031	10,173	20	13,224	30

§ Median days between sample collection and printing of results in the lab

Partners in Hope lab (Lilongwe) achieved the highest outputs, contributing 23% of all results this quarter. The median interval between sample collection and printing of results was **30 days** at the national level, ranging from **8 days** at Mzimba DH to **91 days** at Queen Elizabeth CH. The most significant delays occurred between sample receipt and processing in the lab (median 15 days), while on average only 7 days elapsed between sample collection and receipt in the lab.

Reason	0-999		1000-4999		5000+		Total
Routine	23,311	86%	1,119	4%	2,732	10%	27,162
Targeted	215	62%	24	7%	109	31%	348
Other/unk	240	69%	14	4%	95	27%	349
Total	23,766	85%	1,157	4%	2,936	11%	27,859

27,162 (97%) of all VL samples were classified as *routine scheduled*. This is equivalent to **39%** of the estimated 70,000 ART patients passing a VL monitoring milestone this quarter. **348 (1%)** of samples were classified as *targeted (suspected treatment failure / repeat)* and for **349 (1%)** the reason for the sample was 'other' or not specified. **23,766 (85%)** of all results were below 1,000 copies/ml. The proportion of results with 5,000+ copies was higher among the *targeted* samples (31%) and those with *unspecified* reason (27%), compared with 10% among *routine* samples.

The time on ART was entered for only **6,766 (25%)** of 27,162 routine samples registered on the LIMS and only **2,105 (31%)** of these were drawn on schedule (from 1 month before to 3 months after a VL milestone). The proportion of patients with VL < 1000 was **87%, 85%, 85%, 90%, 84%** and **88%** at 6, 24, 48, 72, 96 and 120 months on ART respectively. Viral suppression rates of samples drawn on schedule were similar to those of 'catch-up' (extra-schedular) samples (**82%**) or those with unknown timing (**85%**).

Patient age was recorded for all routine monitoring samples. Among these, 5%, 6%, 11%, 31% and 48% were from the age groups 0-9, 10-19, 20-29, 30-39 and 40+ years. Viral suppression rates (VL<1000/ml) were significantly lower among children (0-9 yrs: **61%**) and adolescents (10-19 yrs: **60%**) compared with adults (**81%, 85%** and **88%** for the age groups 20-29, 30-39, 40+ years, respectively).

VL monitoring outputs are expected to increase further over the next quarters.

Given the relatively low access to VL monitoring (estimated 39% of all ART patients due for VL monitoring this quarter), the measured **86% viral suppression rate** may not be representative for the entire national ART cohort. With generally limited access to testing, the VL samples analyzed this quarter may over-represent patients with poor adherence and/or treatment failure. Conservatively, the national viral suppression rate can be estimated as **503,668 (86%)** of 585,660 patients on ART, which is equivalent to **50%** of the total 1 million HIV infected population.

16 TB / HIV Management

Approximately **89%** of HIV infected TB patients were receiving ART in Q3 2015. This estimate is based on the following triangulation of TB and ART program data:

TB Program Data: A total of **4,346** TB patients were registered during Q3 2015. Assuming an average HIV prevalence of 60% among TB patients, **2,608** TB patients were HIV positive and therefore in need of ART. Given that **1,573** TB patients registered were already on ART at the time of starting TB treatment, $2,608 - 1,573 = 1,035$ TB patients needed to initiate ART.

ART Program Data: An estimated **649** patients¹⁵ started ART with a current or recent episode of TB in Q3 2015. This is **63%** (649 of 1035) of the TB patients who needed to start ART. This means that a total of $1,573 + 649 = 2,222$ (**85%**) of the estimated 2,608 HIV infected TB patients were receiving ART in Q3 2015.

TB program report

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TB clinic registrations

Total TB patients registered	4,346	100%
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HIV status ascertainment

HIV status not ascertained	373	9%
HIV status ascertained	3,973	91%
HIV negative	1,743	44%
HIV positive	2,230	56%
Already on ART	1,573	71%
Not on ART when starting TB treatment	657	29%

TB / ART program triangulation

*

HIV-burden among TB patients (estimated)

HIV negative (est. 40%)	1,738	40%
HIV positive (est. 60%) in need of ART	2,608	60%
Not on ART	280	11%
Total on ART (coverage)	2,328	89%
Already on ART (TB prog)	1,573	68%
Started ART within 24m of TB diagnosis (ART prog)	755	32%
ART initiations with current TB (ART prog)	499	66%
ART initiations after recent TB (ART prog)	256	34%

17 STI Treatment

STI reports were actively collected during the Integrated HIV Program Supervision exercise for the 10th time this quarter. This decision was taken due to the persistent challenges with 'passive' reporting based on data aggregated by the district STI coordinators. This quarter, supervision teams collected STI data from 656 out of 928 facilities offering STI management according to the *2013-14 Service Provision Assessment*¹⁶ in Malawi. The site-level reports included here may therefore only represent 71% of all STI services in Malawi. The supervision teams re-emphasized the importance of complete and accurate documentation at the sites and the data quality is expected to improve with resumption of regular site supervision for the STI program. The complete set of STI program data collected is included in the Appendix.

¹⁵ 21% of the 1,012 ART patients who were registered this quarter with a recent or current episode of TB at the time of ART initiation were assumed to be transfers and were subtracted to adjust for double-counting.

¹⁶ Ministry of Health, & ICF International. (2015). Malawi Service Provision Assessment (SPA) 2013-14. Lilongwe, Malawi and Rockville, Maryland, USA. Retrieved from <http://dhsprogram.com/pubs/pdf/SPA20/SPA20.pdf>

17.1 Access to STI treatment and coverage

Based on the data collected at the facilities, a total of **61,188** STI cases were treated in Q3 2015. Considering the 71% site-level completeness of reporting, this number is estimated to represent a total of **86,180** STI cases treated. This is equivalent to **87% STI treatment coverage** of the expected 98,600 STI cases in the population.

Out of **61,188** documented clients treated, **24,598** (40%) were male and **36,590** (60%) were female. **4,451** (12%) of female STI clients were pregnant. **41,595** clients (68%) were 25 years and above, **14,503** (24%) were 20-24 years and **5,090** (8%) were under 20 years old.

17.2 Client Type and STI History

54,201 (89%) of clients were symptomatic and **6,987** (11%) were asymptomatic (treated as partners). Among symptomatic clients, **49,166** (91%) of were index cases and **5,035** (9%) were partners. A total of **16,660** partner notification slips were issued, equivalent to an average of 0.34 slips per index case. Considering the 16,660 partner notification slips issued, **72%** (12,022) of those notified presented to the clinic. **45,638** (75%) of clients presented with their first lifetime episode of STI, **10,858** (18%) clients reported to have had an STI more than 3 months ago and **4,692** (8%) of clients reported having had an STI within the last three months. Re-occurrence of an STI after a recent episode may be due to re-infection or treatment failure.

17.3 HIV Status

HIV status was ascertained for **35,406** (58%) clients and **8,044** (23%) of these were HIV positive. **2,077** (26%) of positives were identified through a new test initiated at the STI clinic, while **5,967** (74%) presented with a documented previous positive HIV test result. **4,764** (80%) of clients with a previous positive HIV test result were on ART.

The rate of HIV status ascertainment at STI clinics remained low. This is likely due to poor implementation of provider initiated testing and counselling, combined with weak back-referral systems which may lead to incomplete documentation of new HIV test results at the STI clinics. It is worth noting that a substantial proportion of clients who are aware of their HIV infection present with a new episode of an STI. This may suggest poor translation of positive living strategies promoted during counselling, but could also be due to the increased risk of recurrence of HSV-2 and balanitis among HIV-infected clients.

17.4 STI Syndromes and Referrals

The most common syndrome was abnormal vaginal discharge (AVD) with **19,309** (29%) cases, followed by urethral discharge (UD, **15,172** cases), genital ulcers (GUD, **11,947** cases) and lower abdominal pain (LAP, **10,290** cases). Similar to previous reports, balanitis, bubo, warts and neonatal conjunctivitis each accounted for 1 – 2% of cases.

Given the high risk of recent HIV infection among STI clients, all clients with unknown status and those with a new negative test result should be referred for (repeat) HIV testing and counselling. **20,693 (39%)** of the 53,144 STI clients with unknown or new negative test result were referred for repeat HTC. **1,288 (62%)** of 2,077 clients who were newly tested HIV positive were referred for ART eligibility assessment.

18 Supply of HIV Program Commodities

18.1 Quantification, procurement planning and distribution

The program finalized the quantification, procurement plan and budget review for all HIV commodities as a pre-requisite to grant approval of the New Funding model (NFM). This covers the period ending December 2017 plus 9 months' buffer.

During Q3 2015, ARVs and medicines for opportunistic infections worth \$13.6 million were received by the Bollore Africa Logistics managed warehouses dedicated for Department of HIV and National Malaria Control Program commodities. This included Tenofovir/Lamivudine/Efavirenz 300/300/600mg (Regimen 5A; 90% of the value of adult ARVs) and other ARV formulations (80% of the value for all medicines received during the period). To maintain adequate stocks in the pipeline and ensure uninterrupted supply for subsequent orders, the Ministry has continued processing HIV commodity orders for ARVs, OI, RDTs and other related commodities through Partnership for Supply Chain Management (ARVs and RDTs) and IDA Foundation (laboratory commodities and medicines for opportunistic infections).

Two scheduled rounds of the bimonthly distribution of HIV & Malaria commodities (Distribution Round 24 & 25) took place between August and October 2015. A total of 86 different commodities (anti-malarials, ARVs, OI medicines, STI medicines and laboratory commodities) were distributed to 726 health facilities. These were the fifth and six successful consolidated distributions for HIV and malaria commodities.

The DHA Logistics Team developed distribution lists for all HIV commodities for 726 sites for both rounds, covering the consumption period for the period ending December 2015. During Q3 2015, the logistics team at the Department of HIV and AIDS also coordinated a total of over 990 individual commodity transactions between ART sites to avert stock outs and/or expiries. The transactions are all managed using the HIV Department Supply Chain Hot Line, a toll free facility that was set up to facilitate communication between the health facilities and the central level. Health workers are able to communicate supply chain and other HIV commodities related issues that need to be resolved by the technical team at the department in a timely manner.

18.2 Logistics support during integrated site supervision

District and central level Supply Chain and Logistics Officers provided stock management support at 200 sites during the Q3 2015 integrated ART/PMTCT site supervision. This included a physical inventory at all sites and ad-hoc mentoring in stock management at health facilities with poor performance. There was an overall improvement in the logistics management of ARVs and medicines for OI medicines. Health care providers have continued to use RDT daily activity registers and relocation books for registration of redistributed commodities to health facilities. Authorization codes must be obtained for all such transactions as a measure for tracking, audit and to foster accountability at all sites.

18.3 Stock Status of HIV Commodities

Physical stock counts for ARVs and other medicines for HIV-related diseases were performed at all sites during the supervision visits in October 2015. **Table 6** shows the total medicine stocks found at the sites and the estimated consumption patterns.

Minimum stocks of TDF/3TC 300/300mg and AZT/3TC 300/150mg are maintained at all sites for post-exposure prophylaxis (PEP) and the total stocks at the sites therefore far exceeds the actual consumption from patients using this regimen in alternative ART regimens or as PEP. There are some residual quantities of stavudine- and efavirenz-containing regimens at the warehouse and site level following the transition to a tenofovir-based regimens. The program continues to monitor the trend of patients on such regimens to inform future procurements for alternative regimens such as Abacavir/Lamivudine formulations given that stavudine containing regimens will be phased out completely in 2016.

513,549 patients were on regimen 5A (510,798 at their last registration site + 2,751 patients in transit), which was **14,390 (2.7%)** less than projected in the previous forecast for the end of this quarter (**527,939**). The national ART program forecast and quantification was updated in August 2015 to inform procurement planning and budgeting for HIV commodities for the period ending December 2017.

18.4 Availability of standard first line ARVs

510,798 of all ART patients retained at their last site of registration were on the standard first line regimen (5A; tenofovir / lamivudine / efavirenz). This is equivalent to 87% of patients overall or 93% of patients on first line adult regimens. As of October 2015, the total stock of this regimen was equivalent to 5.0 and 6.3 months of consumption at the warehouse and site-level, respectively. Total national stocks may conceal stock imbalances at the facility level and a key supply management indicators is therefore the availability of this regimen at each site. The physical stock count carried out during supportive supervision in October 2015 confirmed that 698 (99.1%) of all 704 ART sites with patients on this regimen had available stocks. This translates into a 'stock-out' rate of only 0.9% of sites. Such stock-out events typically affect small peripheral sites and are usually short due to the bi-monthly scheduled distribution cycle and the ad-hoc stock relocation facility coordinated through the toll-free supply hotline. This healthy supply chain has enabled the program to consistently implement three monthly drug dispensations for patients.

Table 6: Total stocks of HIV program commodities at all sites visited during the 2015 Q3 supportive site supervision. Stock positions are from the date of the visit (between 1-4 weeks after the end of the quarter). Warehouse stock positions are from 26/11/2015

Inventory unit	Item	Sites with any Stock	Total Physical Stock		Consumption/ Month	Months of Stock *	
			At Sites	In Warehouse		At Sites	Wareh.
tins	ABC / 3TC 60 / 30mg tins (60 tabs)	179	22,488	85,388	4,692	4.8	18.2
	ABC / 3TC 600 / 300mg tins (30 tabs)	36	875	16,198	711	1.2	22.8
	ATV / r 300 / 100mg tins (30 tabs)	195	6,715	33,765	6,342	1.1	5.3
	AZT / 3TC / NVP 300 / 150 / 200mg tins (60 tabs)	629	123,075	290,675	27,334	4.5	10.6
	AZT / 3TC / NVP 60 / 30 / 50mg tins (60 tabs)	638	441,083	496,243	62,503	7.1	7.9
	AZT / 3TC 300 / 150mg tins (60 tabs)	363	14,012	15,830	3,045	4.6	5.2
	AZT / 3TC 60 / 30mg tins (60 tabs)	583	28,946	35,589	2,359	12.3	15.1
	d4T / 3TC / NVP 30 / 150 / 200mg tins (60 tabs)	161	25,135	856	1,098	22.9	0.8
	d4T / 3TC 30 / 150mg tins (60 tabs)	207	12,272		86	142.7	
	EFV 200mg tins (90 tabs)	163	1,983	6,456	327	6.1	19.8
	EFV 600mg tins (30 tabs)	231	15,551	227	815	19.1	0.3
	LPV / r 100 / 25mg tins (60 tabs)	85	8,001	13,953	2,976	2.7	4.7
	LPV / r 200 / 50mg tins (120 tabs)	72	1,442	1,113	456	3.2	2.4
	NVP 200mg tins (60 tabs)	475	34,663	54,700	8,231	4.2	6.6
	NVP 50mg tins (60 tabs)	160	10,276	12,171	1,430	7.2	8.5
	TDF / 3TC / EFV 300 / 300 / 600mg tins (30 tabs)	704	2,566,373	3,198,696	510,798	5.0	6.3
TDF / 3TC 300 / 300mg tins (30 tabs)	639	46,278	55,934	13,319	3.5	4.2	
bottles	Fluconazole (Diflucan) 50mg / 5ml bottles (35 ml)	13	1,125		74	15.3	
	NVP 10mg/ml bottles (25 ml)	579	131,446	55	17,201	7.6	0.0
vials	Benzathine Penicillin 1.44g vials (50 each)	540	337,130		40,505	8.3	
	Bleomycine 15,000IU vials (1 each)	21	5,718	3,970			
	Ceftriaxone 1g vials (50 each)	596	73,956		109,330	0.7	
	Depo-Provera 150mg/1ml vials (25 each)	589	821,734	567,175	278,291	3.0	2.0
	Gentamicin 80mg / 2ml vials (50 each)	546	164,090		102,884	1.6	
	Streptomycin 1 gm vials (50 each)	51	21,256				
Vincristine 1mg / 1ml vials (1 each)	82	31,597	730	3,876	8.2	0.2	
tabs	Acidovir 200mg blister packs (500 tabs)	599	3,847,044		659,037	5.8	
	Azithromycin 500mg blister packs (3 tabs)	406	168,194	25,845	10,876	15.5	2.4
	Ciprofloxacin 500mg blister packs (100 tabs)	327	876,781	1,589,900	311,734	2.8	5.1
	Clotrimazole 500mg boxes (1 each)	464	72,472	16,490	40,070	1.8	0.4
	Codeine 30mg tins (100 tabs)	51	304,734	710,500	51,594	5.9	13.8
	Cotrimoxazole 100 / 20mg blister packs (1000 tabs)	580	30,836,385	27,037,000	7,458,256	4.1	3.6
	Cotrimoxazole 400 / 80mg tins (1000 tabs)	542	28,180,544	41,460,000	17,283,119	1.6	2.4
	Cotrimoxazole 960mg blister packs (1000 tabs)	484	18,980,826	26,668,000	18,392,669	1.0	1.4
	Doxycycline 100mg tins (1000 tabs)	551	6,315,338		4,618,940	1.4	
	E thambutol (E) 100 mg blister packs (100 tabs)	46	104,843				
	E thambutol (E) 400 mg blister packs (672 tabs)	6	16,712				
	Erythromycin 250mg tins (1000 tabs)	305	2,152,326	5,237,000	4,132,109	0.5	1.3
	Fluconazole (Diflucan) 200mg tins (28 tabs)	208	619,441	116,592	42,170	14.7	2.8
	Ibuprofen 200mg tins (100 tabs)	138	1,128,562		883,454	1.3	
	Isoniazid (H) 100mg blister packs (100 tabs)	147	213,350		157,656	1.4	
	Isoniazid (H) 300mg blister packs (672 tabs)	68	509,611		1,152,857	0.4	
	Isoniazid (H) 300mg tins (1000 tabs)	621	15,734,844	9,963,000	1,153,894	13.6	8.6
	Metronidazole 200mg tins (1000 tabs)	575	13,444,704	9,293,000	5,017,667	2.7	1.9
	Morphine 10mg blister packs (60 tabs)	84	329,911		225,136	1.5	
Pyridoxine 50mg tins (1000 tabs)	510	12,434,719	6,361,000	1,231,685	10.1	5.2	
sheets	ART pat. card adult (yellow) bundles (100 sheets)	636	225,320		10,289	21.9	
	ART pat. card paed. (blue) bundles (100 sheets)	587	81,802		869	94.1	
	Exposed child card (pink) bundles (50 sheets)	600	58,152	53,500	3,786	15.4	14.1
	Family HTC Referral Slip bundles (100 sheets)	232	27,433				
	Polythene sleeve bundles (100 sheets)	561	129,429		17,253	7.5	
	Pre-ART pat. card (green) bundles (100 sheets)	581	138,252		2,310	59.9	
	STI Partner Referral Slip bundles (100 sheets)	319	32,362				
tests	DBS kit (filter paper, lancet, etc.) boxes (50 each)	498	63,255	144,000	33,662	1.9	4.3
	Determine HIV1/2 boxes (100 each)	637	703,275	1,661,400	203,875	3.4	8.1
	Determine syphilis boxes (100 each)	489	139,819	282,100	52,231	2.7	5.4
	Uni-Gold HIV1/2 boxes (20 each)	573	55,669	229,340	21,887	2.5	10.5
pieces	Condoms female boxes (1000 each)	358	557,401		193,467	2.9	
	Condoms male boxes (144 each)	423	7,138,533	15,258,816	5,609,140	1.3	2.7

* 'Consumption per month' and 'Months of stock' for ARVs, CPT, INH and HIV test kits are based on the respective patient-regimen groups in the standard service reports. Estimates are based on the number of patients on the respective regimen at the end of the quarter evaluated and do not account for potential (positive or negative) growth. Facility stock positions for OI and STI drugs include HIV Program and other supply sources. Total national consumption and MoS estimates are used for these commodity groups. 'Months of stock' is calculated from the day of the physical stock count, which is on average 1 month after the end of the quarter.

19 Training and Mentoring

19.1 ART/PMTCT trainings

158 providers were newly trained in the 2014 national guidelines in the quarter. 87 were nurses, 42 clinicians, 14 statistical clerks, 11 health surveillance assistants and 4 pharmacy technicians.

19.2 HIV Testing Services

107 were trained in initial HIV testing services this quarter.

19.3 VMMC training

26 clinical officers and **14** nurses were trained in VMMC clinical skills. This VMMC initial training covers forceps guided & dorsal slit methods of circumcision, VMMC group education, HIV counselling and testing, circumcision clients screening and postoperative care services.

20 Participants in Q3 2015 Supervision (Site visits 5 to 16 October 2015)

Absalom Kaunda (CO, MOH, Mzimba DHO)	Felix Magwira (Clinical Coordinator, indep NGO)	Mike Kalulu (CO, MOH)
Afred Kamoto (Logistics Fellow, MOH)	Frazer Mkwawa (Nurse, MOH)	Mike Nyirenda (CO, Lighthouse)
Agnes Kalitsiro (Nurse, Mlambe Mission Hospital)	Geoffrey Makhalaria (, NTP)	Miriam Thindwa (Clinician, Limbe H/C)
Alefa Fikira (CMT, MOH)	Gerald Zomba (Program Officer, Dept for HIV and AIDS)	Monica Simfukwe (Nurse, MOH, Chinthethe RH)
Alexander Malunguza (, NTP)	Grant Gondwe (, NTP)	Mphatso Magwaya (, JHPIEGO)
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Andraida Mtoseni (Nurse, MOH)	Hannock Matupi (ARV clinician, MOH, Rumphidh)	Noel Mphasa (TB Zonal Supervisor, NTP)
Andrew Dimba (, NTP)	Harrison Tembo (CO, MOH)	Noxy Mphaka (, MSH)
Andrew Gompho (Clinician, MOH)	Harry Tsapa (CO, MOH)	Nyembezi Chibonga (, NTP)
Andrew Mganga (M&E Fellow, Dept for HIV and AIDS)	Henry Kanyerere (TB/HIV Program Officer, MOH)	Offrey Mnduwira (CO, Police)
Annie Biza (Nurse, MDF)	Isaac Makhonya (, PIH)	Oscar Kasiyamphanje (Nurse, CHAM)
Austin Nkute (CLINICAL COORDINATOR, indep NGO)	Isaiah Dambe (, NTP)	Overtone Ndhlovu (CO, MOH)
Austins Namondwe (CO, CHAM)	Issa Sulemani (, MOH)	Patrick Ngwira (, NTP)
Batoni Upindi (TB Zonal Supervisor, MOH)	Janet Chikonda (Nurse, MOH)	Patrick Paul J M Chirwa (TB Zonal Supervisor, NTP)
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Benardetta Chunda (Nurse, Lighthouse)	Jesse Lobeni (Nurse, MOH)	Paul Nyasulu (CO, Dept of HIV & AIDS)
Catherine Kassam (, MOH)	John Kabichi (CO, MOH)	Pax Mkupani (Logistics Fellow, MOH)
Cecelia Tenesi (Nurse, MOH)	Johnbosco Mwafilaso (Clerk, MOH)	Peter Chalusa (, EGPAF)
Cecilia Manyawa (Nurse, MOH)	Juliana Soko (ARV nurse, MOH, Livingstonia MH)	Peter Donda (CO, Dedza DH)
Cecilia Sambakunsi (Logistics Fellow, HIV Dept)	Julie Kazima (Nurse, MSH)	Peter Mzumara (ART clinician, MOH)
Chancy Kamba (, NTP)	Juliet Nyirenda (Nurse, MOH)	Phillip Chitowe (Nurse, MOH)
Chifundo Makuluni (Nurse, MOH)	Justice Kaphiri (, NTP)	Precious Mtegha (CO, MOH)
Chikayiko Majamanda (Nurse, MOH)	Kelvin Makina (Logistics, Kasungu)	Priscilla Milongo (Nurse, Lighthouse)
Chikumbutso Pendame (MA, MOH)	Kingsley Makwale (MA, MOH)	Relia Nkhata Mandindi (Logistics, HIV Dept)
Chimwemwe Francis Mkandawire (IT Fellow, Dept for HIV and AIDS)	Kingsley Mbewa (CO, MOH)	Rhoda Banda (, NTP)
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Chisomo Thondolo (Nurse, EGPAF)	Kondwani Chikoti (CO, MOH)	Rodrick Kaulere (CO, CHAM (Sister Tereza))
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Christopher Mkwazalamba (CO, MOH)	Lameck Mlauzi (, NTP (MOH))	Rumours Lumala (CO, MOH)
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We thank all facility staff for their sincere welcome and co-operation with the HIV Department and its partners during these supportive visits. We congratulate all staff for their excellent work.

29 January 2016

21 Appendix (Full National HIV Program Data)

HTC site report

Malawi (national)

2015 Q3 (1st month of quarter, 2nd month of quarter, 3rd month of quarter)

Clients at health facility (static)

HTC client details

*

Total HTC clients served

Total HIV tested	586,926	100%
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Sex

Males tested	215,480	37%
Females tested	371,446	63%
Females non-pregnant	203,837	55%
Females pregnant	167,609	45%

Age

Children 0-14 yrs	61,772	11%
Children below 12 mths (Age group A)	3,112	5%
Children 12 mths - 14 yrs (Age group B)	58,660	95%
Adults 15+ years	525,154	89%
Young adults 15-24 years (Age group C)	227,418	43%
Older adults 25+ yrs (Age group D)	297,736	57%

HTC access type

PITC	345,143	59%
Family Referral Slip (FRS)	2,878	0%
Other (VCT, etc.) HTC access	238,905	41%

HTC first time / repeat

Never tested before	177,613	30%
Previously accessed HTC	409,313	70%
Last negative	393,730	96%
Last positive	13,033	3%
Last exposed infant	1,979	0%
Last inconclusive	571	0%

Counseling session type / Partner present

Counseled with partner / partner present	140,089	24%
Counseled alone / Partner not present	446,837	76%

Outcome summary (HIV test)

Single test negative	539,079	92%
Single test positive	206	0%
Test 1&2 negative	820	0%
Test 1&2 positive	44,730	8%
Test 1&2 discordant	2,091	0%

Final result given to client

Results among clients never tested / last negative	574,155	98%
New negative	538,713	94%
New positive	32,541	6%
New exposed infants	1,109	0%
New inconclusive	1,792	0%
Confirmatory results (previous positive clients)	12,771	2%
Confirmatory positive	12,201	96%
Confirmatory inconclusive	570	4%

HTC site report

Malawi (national)

2015 Q3 (1st month of quarter, 2nd month of quarter, 3rd month of quarter)

HTC client details

*

Partner / Family HTC referral slips

Sum of slips given	15,002	100%
Total clients presenting with referral slip	2,878	19%
Total failed referrals (slips not returned)	12,124	81%

Clients tested in the community

HTC client details

*

Total HTC clients served

Total HIV tested	24,314	100%
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Sex

Males tested	11,098	46%
Females tested	13,216	54%
Females non-pregnant	10,824	82%
Females pregnant	2,392	18%

Age

Children 0-14 yrs	2,796	11%
Children below 12 mths (Age group A)	111	4%
Children 12 mths - 14 yrs (Age group B)	2,685	96%
Adults 15+ years	21,518	89%
Young adults 15-24 years (Age group C)	10,037	47%
Older adults 25+ yrs (Age group D)	11,481	53%

HTC access type

PITC	7,012	29%
Family Referral Slip (FRS)	15	0%
Other (VCT, etc.) HTC access	17,287	71%

HTC first time / repeat

Never tested before	8,249	34%
Previously accessed HTC	16,065	66%
Last negative	15,606	97%
Last positive	450	3%
Last exposed infant	3	0%
Last inconclusive	6	0%

Counseling session type / Partner present

Counseled with partner / partner present	1,438	6%
Counseled alone / Partner not present	22,876	94%

Outcome summary (HIV test)

Single test negative	22,432	92%
Single test positive	22	0%
Test 1&2 negative	55	0%
Test 1&2 positive	1,720	7%
Test 1&2 discordant	85	0%

HTC site report

Malawi (national)

2015 Q3 (1st month of quarter, 2nd month of quarter, 3rd month of quarter)

HTC client details

*

Final result given to client

Results among clients never tested / last negative	24,139	99%
New negative	22,463	93%
New positive	1,361	6%
New exposed infants	10	0%
New inconclusive	305	1%
Confirmatory results (previous positive clients)	175	1%
Confirmatory positive	169	97%
Confirmatory inconclusive	6	3%

Partner / Family HTC referral slips

Sum of slips given	918	100%
Total clients presenting with referral slip	15	2%
Total failed referrals (slips not returned)	903	98%

Clients at stand-alone HTC sites

HTC client details

*

Total HTC clients served

Total HIV tested	14,563	100%
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Sex

Males tested	7,535	52%
Females tested	7,028	48%
Females non-pregnant	6,062	86%
Females pregnant	966	14%

Age

Children 0-14 yrs	321	2%
Children below 12 mths (Age group A)	5	2%
Children 12 mths - 14 yrs (Age group B)	316	98%
Adults 15+ years	14,242	98%
Young adults 15-24 years (Age group C)	4,841	34%
Older adults 25+ yrs (Age group D)	9,401	66%

HTC access type

PITC	9,471	65%
Family Referral Slip (FRS)	12	0%
Other (VCT, etc.) HTC access	5,080	35%

HTC first time / repeat

Never tested before	4,189	29%
Previously accessed HTC	10,374	71%
Last negative	9,948	96%
Last positive	421	4%
Last exposed infant	0	0%
Last inconclusive	5	0%

Counseling session type / Partner present

Counseled with partner / partner present	1,727	12%
Counseled alone / Partner not present	12,836	88%

HTC site report

Malawi (national)

2015 Q3 (1st month of quarter, 2nd month of quarter, 3rd month of quarter)

HTC client details

*

Outcome summary (HIV test)

Single test negative	13,029	89%
Single test positive	2	0%
Test 1&2 negative	6	0%
Test 1&2 positive	1,444	10%
Test 1&2 discordant	82	1%

Final result given to client

Results among clients never tested / last negative	14,416	99%
New negative	13,054	91%
New positive	1,064	7%
New exposed infants	3	0%
New inconclusive	295	2%
Confirmatory results (previous positive clients)	147	1%
Confirmatory positive	143	97%
Confirmatory inconclusive	4	3%

Partner / Family HTC referral slips

Sum of slips given	1,014	100%
Total clients presenting with referral slip	12	1%
Total failed referrals (slips not returned)	1,002	99%

Blood safety

Malawi (national)

2015 Q3 (1st month of quarter, 2nd month of quarter, 3rd month of quarter)

Infect. disease screening among potential donors

*

HIV screening

HIV testing not done	2,114	20%
Tested for HIV	8,571	80%
HIV negative	8,090	94%
HIV positive	481	6%

Hepatitis B screening

HepB testing not done	2,182	20%
Tested for Hepatitis B	8,503	80%
HepB Negative	8,073	95%
HepB Positive	430	5%

Hepatitis C screening

HepC testing not done	5,602	52%
Tested for Hepatitis C	5,083	48%
HepC Negative	4,992	98%
HepC Positive	91	2%

Syphilis screening

Syphilis testing not done	2,105	20%
Tested for Syphilis	8,580	80%
Syphilis Negative	8,357	97%
Syphilis Positive	223	3%

Malaria screening

Malaria testing not done	3,541	33%
Tested for malaria	7,144	67%
Malaria Negative	6,685	94%
Malaria Positive	459	6%

Summary screening outcome

Not donated	3,366	32%
Donated	7,319	68%
Screened for at least HIV, HepB and syphilis	6,191	85%
Screened for HIV, HepB, HepC, Syphilis, Malaria	3,368	54%
Screened for HIV, HepB, Syphilis	2,823	46%
Screened for HIV, HepB	17	0%
Screened for HIV only	2	0%
Screened with any other combination of tests	1,109	15%

Cross-matching report

*

Blood group typing (for units and patients)

Total blood group typing done	21,138	100%
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Blood units cross-matched (by source)

Total blood units cross-matched	11,534	100%
Total units from MBTS (estimated)	4,215	37%
Total units from replacement donors	7,319	63%

Blood units cross-matched by patient group

Units cross-matched for maternity	2,371	21%
Units cross-matched for paediatrics	3,925	34%
Units cross-matched for other ward	5,238	45%

Blood safety

Malawi (national)

2015 Q3 (1st month of quarter, 2nd month of quarter, 3rd month of quarter)

Cross-matching report

*

Transfusion reactions

Units transfused without adverse events	11,486	100%
Units with suspected transfusion reactions	41	0%
Units with confirmed transfusion reactions	7	0%

HIV exposed child follow-up

Malawi (national)

2015 Q3 (1st month of quarter, 2nd month of quarter, 3rd month of quarter)

Age 2 months

Age cohort outcomes

*

Total children in birth cohort

Total children registered	7,800	100%
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CPT status

On CPT	6,954	89%
Not on CPT	846	11%

HIV status

Current HIV infection status unknown	5,438	70%
HIV infection not confirmed, not ART eligible	5,418	100%
HIV infection not confirmed, ART eligible (PSHD)	20	0%
Current HIV infection status known	2,362	30%
Confirmed not infected	2,319	98%
Confirmed infected (ART eligible)	43	2%

ART eligibility summary

Not eligible for ART	7,737	99%
ART eligible	63	1%
ART not initiated	30	48%
Initiated ART	33	52%

Primary follow-up outcome

Discharged uninfected	34	0%
Continue follow-up	7,020	92%
Started ART	33	0%
Defaulted	526	7%
Died	19	0%

Transfers between sites

Total not transferred out	7,632	98%
Transferred out	168	2%

Age 12 months

Age cohort outcomes

*

Total children in birth cohort

Total children registered	8,434	100%
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CPT status

On CPT	6,216	74%
Not on CPT	2,218	26%

HIV status

Current HIV infection status unknown	4,378	52%
HIV infection not confirmed, not ART eligible	4,358	100%
HIV infection not confirmed, ART eligible (PSHD)	20	0%
Current HIV infection status known	4,056	48%
Confirmed not infected	3,886	96%
Confirmed infected (ART eligible)	170	4%

HIV exposed child follow-up

Malawi (national)

2015 Q3 (1st month of quarter, 2nd month of quarter, 3rd month of quarter)

Age cohort outcomes

*

ART eligibility summary

Not eligible for ART	8,244	98%
ART eligible	190	2%
ART not initiated	15	8%
Initiated ART	175	92%

Primary follow-up outcome

Discharged uninfected	73	1%
Continue follow-up	6,148	75%
Started ART	175	2%
Defaulted	1,747	21%
Died	74	1%

Transfers between sites

Total not transferred out	8,217	97%
Transferred out	217	3%

Age 24 months

Age cohort outcomes

*

Total children in birth cohort

Total children registered	8,057	100%
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CPT status

On CPT	1,311	16%
Not on CPT	6,746	84%

HIV status

Current HIV infection status unknown	4,127	51%
HIV infection not confirmed, not ART eligible	4,117	100%
HIV infection not confirmed, ART eligible (PSHD)	10	0%
Current HIV infection status known	3,930	49%
Confirmed not infected	3,705	94%
Confirmed infected (ART eligible)	225	6%

ART eligibility summary

Not eligible for ART	7,822	97%
ART eligible	235	3%
ART not initiated	23	10%
Initiated ART	212	90%

Primary follow-up outcome

Discharged uninfected	3,540	45%
Continue follow-up	842	11%
Started ART	212	3%
Defaulted	3,132	40%
Died	103	1%

Transfers between sites

Total not transferred out	7,829	97%
Transferred out	228	3%

2015 Q3 (Quarter)

Registration details

*

HCC clinic registrations

Total HCC registrations	18,286	100%
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Registration type

Patients enrolled first time	17,250	94%
Patients re-enrolled	51	0%
Patients transferred in	985	5%

Sex

Males (all ages)	8,511	47%
Females (all ages)	9,775	53%
Non-pregnant	9,751	100%
Pregnant	24	0%

Age at registration

Adults 15+ yrs	6,486	35%
Children 0-14 yrs	11,800	65%
Children 24 months - 14 years	587	5%
Children below 24 months (exposed children)	11,213	95%
Children 2 - below 24 months	3,655	33%
Infants below 2 months	7,558	67%

Reason for HCC registration

Exposed infants	11,357	62%
Confirmed infected patients (pre-ART)	6,929	38%

2015 Q3 (Cumulative)

Registration details

*

HCC clinic registrations

Total HCC registrations	374,404	100%
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Registration type

Patients enrolled first time	360,839	96%
Patients re-enrolled	1,168	0%
Patients transferred in	12,397	3%

Sex

Males (all ages)	162,829	43%
Females (all ages)	211,575	57%
Non-pregnant	210,619	100%
Pregnant	956	0%

Age at registration

Adults 15+ yrs	185,233	49%
Children 0-14 yrs	189,171	51%
Children 24 months - 14 years	16,853	9%
Children below 24 months (exposed children)	172,318	91%
Children 2 - below 24 months	80,106	46%
Infants below 2 months	92,212	54%

Reason for HCC registration

Exposed infants	172,600	46%
Confirmed infected patients (pre-ART)	201,804	54%

Pre-ART follow-up outcome

*

Primary follow-up outcomes

Total retained in pre-ART	43,217	22%
Started ART	102,297	53%
Defaulted	46,939	24%
Died	1,917	1%

Transfers between sites

Total not transferred out	194,870	97%
Transferred out	6,934	3%

Antenatal Care

Malawi (national)

2015 Q3 (1st month of quarter, 2nd month of quarter, 3rd month of quarter)

New ANC registrations in reporting period

*

Women with first visit in reporting period

New women registered	150,456	100%
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ANC cohort analysis

*

Trimester of first visit

Started ANC 0-12 wks	13,629	9%
Started ANC 13+ wks	136,827	91%

HIV status ascertainment

HIV status not ascertained	15,045	10%
HIV status ascertained	135,411	90%
Valid previous test result	12,636	9%
Previous negative	7,268	58%
Previous positive	5,368	42%
New test at ANC	122,775	91%
New negative	118,081	96%
New positive	4,694	4%

HIV status summary

Total women HIV negative	125,349	93%
Total women HIV positive	10,062	7%

PMTCT regimen mother

No ARVs	1,019	10%
Any ARVs	9,043	90%
ART (by time of initiation)	9,043	100%
Already on ART when starting ANC	4,849	54%
Started ART at 0-27 weeks of pregnancy	3,469	38%
Started ART at 28+ weeks of preg.	725	8%

ANC women after 6 months

ANC cohort analysis

*

Total women completing ANC in the reporting period

Total women in booking cohort	156,851	100%
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Visits per woman

Women with 1 visit	32,900	21%
Women with 2 visits	39,494	25%
Women with 3 visits	46,134	29%
Women with 4 visits	30,761	20%
Women with 5+ visits	7,562	5%

Pre-eclampsia

No pre-eclampsia	153,042	98%
Pre-eclampsia	3,809	2%

TTV doses

0-1 TTV doses	77,784	50%
2+ TTV doses	79,067	50%

SP tablets

0 SP doses	17,632	11%
1 SP dose (1 x 3 tabs)	36,618	23%
6+ SP tablets (2 x 3 tabs)	102,601	65%

Antenatal Care

Malawi (national)

2015 Q3 (1st month of quarter, 2nd month of quarter, 3rd month of quarter)

ANC cohort analysis

*

FeFo tablets

0-119 FeFo tablets	116,023	74%
120+ FeFo tablets	40,828	26%

Albendazole (Deworming)

0 Albend. doses	28,188	18%
1 Albend. dose	128,535	82%

ITN (bednets)

No ITN	24,425	16%
ITN received	131,133	84%

Syphilis status

Not tested for syphilis	126,777	81%
Tested for syphilis	30,074	19%
Syphilis negative	29,462	98%
Syphilis positive	612	2%

HIV status ascertainment

HIV status not ascertained	14,150	9%
HIV status ascertained	142,701	91%
Valid previous test result	12,197	9%
Previous negative	6,416	53%
Previous positive	5,781	47%
New test at ANC	130,504	91%
New negative	125,172	96%
New positive	5,332	4%

HIV status summary

Total women HIV negative	131,588	92%
Total women HIV positive	11,113	8%

CPT status (among HIV pos)

Not on CPT	796	7%
On CPT	10,317	93%

PMTCT regimen mother

No ARVs	1,014	9%
Any ARVs	10,099	91%
ART (by time of initiation)	10,099	100%
Already on ART when starting ANC	5,045	50%
Started ART at 0-27 weeks of pregnancy	4,047	40%
Started ART at 28+ weeks of preg.	1,007	10%

Baby's ARVs dispensed

No ARVs dispensed for infant	1,680	15%
ARVs dispensed for infant	9,433	85%

Maternity

Malawi (national)

2015 Q3 (1st month of quarter, 2nd month of quarter, 3rd month of quarter)

Maternal details

*

Admissions in the reporting period

Total admissions (referrals double-counted)	132,255	100%
Not referred to other site (total women)	124,983	95%
Referred out before delivery (multiple admissions)	7,272	5%

HIV status ascertainment

HIV status not ascertained	4,044	3%
HIV status ascertained	129,887	97%
Valid previous test result	126,231	97%
Previous negative	116,717	92%
Previous positive	9,514	8%
New test at maternity	3,656	3%
New negative	3,302	90%
New positive	354	10%

HIV status summary

Total women HIV negative	120,019	92%
Total women HIV positive	9,868	8%

ARVs during pregnancy (among HIV pos)

No ARV in pregnancy	209	2%
Any ARVs	9,659	98%
ART (by time of initiation)	9,659	100%
ART initiated before pregnancy	7,401	77%
ART initiated in 1st / 2nd trimester	1,207	12%
ART initiated in 3rd trimester	890	9%
ART initiated during labour	161	2%

Obstetric complications

No obstetric complications	117,621	88%
Any obstetric complications	16,310	12%
Haemorrhage	2,668	16%
Haemorrhage ante-partum	857	32%
Haemorrhage post-partum	1,811	68%
Obstr / prol labour	5,637	35%
(pre-) Eclampsia	1,367	8%
Maternal sepsis	126	1%
Ruptured uterus	128	1%
Other obstetric complications	6,384	39%

Emergency obstetric care

Oxytocin	122,493	94%
Anticonvulsive	1,025	1%
Antibiotics	6,470	5%
Blood transfusion	316	0%
Manual removal of placenta	382	0%

Vitamin A

Vit A not given	35,631	27%
Vit A given	98,300	73%

Maternity

Malawi (national)

2015 Q3 (1st month of quarter, 2nd month of quarter, 3rd month of quarter)

Maternal details

*

Staff conducting delivery

Category A: MO, CO, nurse/midwife, MA	121,450	96%
Category B: PA, WA, HSA	604	0%
Category C: Other	4,605	4%

Mother survival

Mother alive	126,556	100%
Mother died	103	0%

Infant details

*

Single babies / multiple deliveries

Total babies delivered	128,967	100%
Single babies	124,549	97%
Twin / multiple babies	4,418	3%

Delivery place

Total deliveries at a health facility	124,100	96%
This facility	123,837	100%
Other facility	263	0%
Total deliveries before reaching the facility	4,867	4%
In transit	3,282	67%
Home / TBA	1,585	33%

Delivery mode

Spontaneous vaginal	116,177	90%
Vacuum extraction	1,776	1%
Breech	2,199	2%
Caesarean section	8,815	7%

Infant complications

No infant complications	112,805	87%
Total infants with complications	16,162	13%
Prematurity	3,501	22%
Weight less 2500g	4,918	30%
Asphyxia	5,328	33%
Sepsis	609	4%
Other newborn complication	1,806	11%

Infant survival

Total live births	126,776	98%
Discharged alive	125,653	99%
Neonatal deaths	1,123	1%
Stillbirths	2,191	2%
Stillbirth, fresh	1,164	53%
Stillbirth, macerated	1,027	47%

Maternity

Malawi (national)

2015 Q3 (1st month of quarter, 2nd month of quarter, 3rd month of quarter)

Infant details

*

HIV exposure / ARV proph. (among discharged alive)

Infants with unknown HIV exposure status	3,106	2%
Infants with known HIV exposure status	122,547	98%
Not HIV exposed	113,528	93%
HIV exposed	9,019	7%
Received no ARVs	605	7%
Received ARVs	8,414	93%
Nevirapine	8,414	100%

Breastfeeding initiated

BF not started within 60min	9,886	8%
BF started within 60min	119,081	92%

Tetracycline eye ointment given

TO not given	14,072	11%
TO given	114,874	89%

ART cohort analysis

Malawi (national)

2015 Q3 (Cumulative)

Registration details

*

ART clinic registrations

Total ART clinic registrations	1,060,580	100%
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Registration type

First time ART initiations (total patients)	849,368	80%
ART re-initiations	11,866	1%
ART transfers in	199,346	19%

Sex

Males	382,186	36%
Females	678,394	64%
Non-pregnant	551,091	81%
Pregnant	127,303	19%

Age at ART initiation

Adults 15+ yrs	968,577	91%
Children 0-14 yrs	92,003	9%
Children 2-14 yrs	70,806	77%
Children below 24 mths	21,197	23%

Reason for starting ART

Presumed severe HIV Disease	3,620	0%
Confirmed HIV infection	1,056,960	100%
WHO stage 1 or 2	469,368	44%
Total lymphocytes <threshold	267	0%
CD4 below threshold	311,347	66%
CD4 unknown or >threshold	157,754	34%
PCR infants	2,919	2%
Children 12-59 mths	7,511	5%
Pregnant women	108,157	69%
Breastfeeding mothers	39,167	25%
WHO stage 3	478,064	45%
WHO stage 4	102,109	10%
Unknown / reason outside of guidelines	7,419	1%

TB at ART initiation

Never TB / TB > 24 months ago	988,576	93%
TB within the last 24 months	36,580	3%
Current episode of TB	35,424	3%

Kaposi's sarcoma at ART initiation

No KS	1,040,824	98%
Patients with KS	19,756	2%

ART cohort analysis

Malawi (national)

2015 Q3 (Cumulative)

ART outcomes

*

Primary follow-up outcomes

Total alive on ART	582,523	68%
Alive on ART at site of last registration	582,384	100%
ART patients in transit between sites	139	0%
Defaulted	195,982	23%
Stopped ART	3,448	0%
Total died	79,280	9%
Died month 1	19,212	24%
Died month 2	12,258	15%
Died month 3	7,294	9%
Died month 4+	40,516	51%

Transfers between sites

Total not transferred out	861,095	81%
Transferred out	199,485	19%

ART regimens

First line regimens	574,610	99%
Adult formulation	548,547	95%
Regimen 0A	271	0%
Regimen 1A	1,098	0%
Regimen 2A	27,334	5%
Regimen 3A	86	0%
Regimen 4A	729	0%
Regimen 5A	510,798	93%
Regimen 6A	8,231	2%
Paed. formulation	26,063	5%
Regimen 0P	471	2%
Regimen 1P	101	0%
Regimen 2P	25,001	96%
Regimen 3P	49	0%
Regimen 4P	441	2%
Second line regimens	7,334	1%
Adult formulation	6,342	86%
Regimen 7A	4,557	72%
Regimen 8A	1,785	28%
Paed. Formulation	992	14%
Regimen 9P	992	100%
Other regimen (adult / paed)	440	0%

Adherence

Adherence unknown (not recorded)	12,279	2%
Adherence recorded	570,105	98%
0-3 doses missed	520,401	91%
4+ doses missed	49,704	9%

ART side effects

Side effects unknown (not recorded)	100,450	17%
Side effects recorded	481,934	83%
No side effects	473,477	98%
Any side effects	8,457	2%

ART cohort analysis

Malawi (national)

2015 Q3 (Cumulative)

ART outcomes

*

Current TB status among ART patients (ICF)

ICF not done (Current TB status unknown/ not circ)	4,943	1%
ICF done	577,441	99%
TB not suspected	573,369	99%
TB suspected	3,270	1%
TB confirmed	802	0%
TB confirmed, not on treatment	73	9%
TB confirmed, on TB treatment	729	91%

ART cohort analysis

Malawi (national)

2015 Q3 (Quarter)

Registration details

*

ART clinic registrations

Total ART clinic registrations	33,474	100%
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Registration type

First time ART initiations (total patients)	25,737	77%
ART re-initiations	409	1%
ART transfers in	7,328	22%

Sex

Males	12,325	37%
Females	21,149	63%
Non-pregnant	14,664	69%
Pregnant	6,485	31%

Age at ART initiation

Adults 15+ yrs	30,867	92%
Children 0-14 yrs	2,607	8%
Children 2-14 yrs	1,935	74%
Children below 24 mths	672	26%

Reason for starting ART

Presumed severe HIV Disease	104	0%
Confirmed HIV infection	33,370	100%
WHO stage 1 or 2	21,603	65%
Total lymphocytes <threshold	4	0%
CD4 below threshold	12,723	59%
CD4 unknown or >threshold	8,876	41%
PCR infants	103	1%
Children 12-59 mths	608	7%
Pregnant women	6,475	73%
Breastfeeding mothers	1,690	19%
WHO stage 3	10,008	30%
WHO stage 4	1,401	4%
Unknown / reason outside of guidelines	358	1%

TB at ART initiation

Never TB / TB > 24 months ago	32,492	97%
TB within the last 24 months	333	1%
Current episode of TB	649	2%

Kaposi's sarcoma at ART initiation

No KS	33,151	99%
Patients with KS	323	1%

2015 Q3 (Quarter)

12 month survival children**Survival and retention in ART program**

*

ART cohort registration group outcomes

Total ART clinic registrations	2,767	100%
Transfers out (double counted)	277	10%
Total not transferred out (patients in cohort)	2,490	90%
Total alive on ART	1,970	79%
Total not retained	520	21%
Defaulted	440	85%
Stopped ART	7	1%
Died	73	14%

12 month survival all ages**Survival and retention in ART program**

*

ART cohort registration group outcomes

Total ART clinic registrations	35,515	100%
Transfers out (double counted)	3,401	10%
Total not transferred out (patients in cohort)	32,114	90%
Total alive on ART	25,107	78%
Total not retained	7,007	22%
Defaulted	6,052	86%
Stopped ART	48	1%
Died	907	13%

24 month survival all ages**Survival and retention in ART program**

*

ART cohort registration group outcomes

Total ART clinic registrations	33,262	100%
Transfers out (double counted)	4,499	14%
Total not transferred out (patients in cohort)	28,763	86%
Total alive on ART	20,660	72%
Total not retained	8,103	28%
Defaulted	6,627	82%
Stopped ART	76	1%
Died	1,400	17%

36 month survival all ages**Survival and retention in ART program**

*

ART cohort registration group outcomes

Total ART clinic registrations	38,483	100%
Transfers out (double counted)	5,917	15%
Total not transferred out (patients in cohort)	32,566	85%
Total alive on ART	22,397	69%
Total not retained	10,169	31%
Defaulted	8,032	79%
Stopped ART	130	1%
Died	2,007	20%

2015 Q3 (Quarter)

48 month survival all ages**Survival and retention in ART program**

*

ART cohort registration group outcomes

Total ART clinic registrations	32,695	100%
Transfers out (double counted)	6,701	20%
Total not transferred out (patients in cohort)	25,994	80%
Total alive on ART	18,239	70%
Total not retained	7,755	30%
Defaulted	5,528	71%
Stopped ART	137	2%
Died	2,090	27%

60 month survival all ages**Survival and retention in ART program**

*

ART cohort registration group outcomes

Total ART clinic registrations	22,203	100%
Transfers out (double counted)	5,893	27%
Total not transferred out (patients in cohort)	16,310	73%
Total alive on ART	9,981	61%
Total not retained	6,329	39%
Defaulted	4,297	68%
Stopped ART	85	1%
Died	1,947	31%

72 month survival all ages**Survival and retention in ART program**

*

ART cohort registration group outcomes

Total ART clinic registrations	19,503	100%
Transfers out (double counted)	5,832	30%
Total not transferred out (patients in cohort)	13,671	70%
Total alive on ART	7,734	57%
Total not retained	5,937	43%
Defaulted	3,979	67%
Stopped ART	92	2%
Died	1,866	31%

84 month survival all ages**Survival and retention in ART program**

*

ART cohort registration group outcomes

Total ART clinic registrations	18,599	100%
Transfers out (double counted)	5,594	30%
Total not transferred out (patients in cohort)	13,005	70%
Total alive on ART	7,005	54%
Total not retained	6,000	46%
Defaulted	3,748	62%
Stopped ART	82	1%
Died	2,170	36%

2015 Q3 (Quarter)

96 month survival all ages**Survival and retention in ART program**

*

ART cohort registration group outcomes

Total ART clinic registrations	13,721	100%
Transfers out (double counted)	4,518	33%
Total not transferred out (patients in cohort)	9,203	67%
Total alive on ART	4,405	48%
Total not retained	4,798	52%
Defaulted	2,992	62%
Stopped ART	47	1%
Died	1,759	37%

108 month survival all ages**Survival and retention in ART program**

*

ART cohort registration group outcomes

Total ART clinic registrations	11,107	100%
Transfers out (double counted)	3,509	32%
Total not transferred out (patients in cohort)	7,598	68%
Total alive on ART	3,533	46%
Total not retained	4,065	54%
Defaulted	2,103	52%
Stopped ART	48	1%
Died	1,914	47%

120 month survival all ages**Survival and retention in ART program**

*

ART cohort registration group outcomes

Total ART clinic registrations	6,844	100%
Transfers out (double counted)	2,089	31%
Total not transferred out (patients in cohort)	4,755	69%
Total alive on ART	1,991	42%
Total not retained	2,764	58%
Defaulted	1,140	41%
Stopped ART	45	2%
Died	1,579	57%

6 month survival OptionB+**Survival and retention in ART program**

*

ART cohort registration group outcomes

Total ART clinic registrations	7,928	100%
Transfers out (double counted)	502	6%
Total not transferred out (patients in cohort)	7,426	94%
Total alive on ART	5,781	78%
Total not retained	1,645	22%
Defaulted	1,560	95%
Stopped ART	31	2%
Died	54	3%

2015 Q3 (Quarter)

12 month survival OptionB+**Survival and retention in ART program**

*

ART cohort registration group outcomes

Total ART clinic registrations	9,082	100%
Transfers out (double counted)	752	8%
Total not transferred out (patients in cohort)	8,330	92%
Total alive on ART	5,909	71%
Total not retained	2,421	29%
Defaulted	2,352	97%
Stopped ART	25	1%
Died	44	2%

24 month survival OptionB+**Survival and retention in ART program**

*

ART cohort registration group outcomes

Total ART clinic registrations	8,633	100%
Transfers out (double counted)	956	11%
Total not transferred out (patients in cohort)	7,677	89%
Total alive on ART	5,179	67%
Total not retained	2,498	33%
Defaulted	2,282	91%
Stopped ART	50	2%
Died	166	7%

36 month survival OptionB+**Survival and retention in ART program**

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ART cohort registration group outcomes

Total ART clinic registrations	10,261	100%
Transfers out (double counted)	1,275	12%
Total not transferred out (patients in cohort)	8,986	88%
Total alive on ART	5,967	66%
Total not retained	3,019	34%
Defaulted	2,717	90%
Stopped ART	48	2%
Died	254	8%

STI site report

Malawi (national)

2015 Q3 (1st month of quarter, 2nd month of quarter, 3rd month of quarter)

STI clients treated in the reporting period

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Total STI clients

Total STI clients treated	61,188	100%
Index patients treated (symptomatic)	49,166	80%
Partners treated	12,022	20%

Sex

Males	24,598	40%
Females	36,590	60%
Non-pregnant	32,139	88%
Pregnant	4,451	12%

Age group

Age group A (0-19 years)	5,090	8%
Age group B (20-24 years)	14,503	24%
Age group C (25+ years)	41,595	68%

Client type

Symptomatic cases	54,201	89%
Index cases	49,166	91%
Partners symptomatic	5,035	9%
Partners asymptomatic	6,987	11%

STI treatment history

Never treated for STI	45,638	75%
Previously treated for STI	15,550	25%
Old >3 months ago	10,858	70%
Recent ≤3 months ago	4,692	30%

STI syndromic diagnosis

GUD	11,947	18%
UD	15,172	23%
AVD	19,309	29%
Low risk	7,552	39%
High risk	11,757	61%
LAP	10,290	16%
SS	973	1%
BU	682	1%
BA	1,167	2%
NC	331	1%
Genital Warts	584	1%
Syphilis RPR VDRL	1,821	3%
Other STI	3,851	6%

STI partner notification

Total partner notification slips issued	16,660	100%
Total partners returned	12,022	72%
Total partners not seen	4,638	28%

STI site report

Malawi (national)

2015 Q3 (1st month of quarter, 2nd month of quarter, 3rd month of quarter)

STI clients treated in the reporting period

*

HIV test / ART status

HIV status not ascertained	25,782	42%
HIV status ascertained	35,406	58%
HIV negative (new test)	27,362	77%
HIV positive	8,044	23%
New positive	2,077	26%
Previous positive	5,967	74%
Not on ART	1,203	20%
On ART	4,764	80%

STI clients referred for services

Lab	590	2%
Gynae review	578	2%
Surgical review	626	2%
Repeat HTC	20,693	82%
ART (for assessment)	1,288	5%
PMTCT	176	1%
Other (service referrals)	1,419	6%