Cost analysis of implementing HIV drug resistance testing in Kenya: a case study of a service delivery site at a tertiary level hospital in Kenya [version 1; peer review: 2 approved]

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Abstract

Background: HIV drug resistance (HIVDR) threatens progress achieved in response to the HIV epidemic. Understanding the costs of implementing HIVDR testing programs for patient management and surveillance in resource-limited settings is critical in optimizing resource allocation. Here, we estimate the unit cost of HIVDR testing and identify major cost drivers while documenting challenges and lessons learnt in implementation of HIVDR testing at a tertiary level hospital in Kenya.

Methods: We employed a mixed costing approach to estimate the costs associated with performing a HIVDR test from the provider’s perspective. Data collection involved a time and motion study of laboratory procedures and interviewing laboratory personnel and the management personnel. Cost analysis was based on estimated 1000 HIVDR tests per year. Data entry and analysis were done using Microsoft Excel and costs converted to US dollars (2019).

Results: The estimated unit cost for a HIVDR test was $271.78 per test. The main cost drivers included capital ($102.42, 37.68%) and reagents (101.50, 37.35%). Other costs included: personnel ($46.81, 17.22%), utilities ($14.69, 5.41%), equipment maintenance costs ($2.37, 0.87%) and quality assurance program ($4, 1.47%). Costs in relation to specific laboratory processes were as follows: sample collection ($2.41, 0.89%), RNA extraction ($22.79, 8.38%), amplification ($56.14, 20.66%), gel electrophoresis ($10.34, 3.80%), sequencing ($160.94, 59.22%), and sequence analysis ($19.16, 7.05%). A user-initiated modification of halving reagent volumes for some laboratory processes (amplification and sequencing) reduced the unit cost for a HIVDR test to $233.81 (13.97%) reduction.

Conclusions: Capital expenditure and reagents remain the most expensive components of HIVDR testing. This cost is bound to change as the sequencing platform is utilized towards maximum capacity or leveraged for use with other tests. Cost saving in offering HIVDR...
testing services is also possible through reagent volume reduction without compromising on the quality of test results.

**Keywords**
HIV, HIV drug resistance testing, implementation science, cost analysis, health systems strengthening

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Background
Unprecedented increased access to antiretroviral therapy (ART) is one of the greatest milestones in the fight against the HIV epidemic, resulting in reduced mortality from AIDS-related causes and a global decline in HIV incidence (UNAIDS, 2019). However, this success is threatened by emergence of HIV drug resistance (HIVDR). The World Health Organization (WHO) reports greater than 10% pretreatment drug resistance to non-nucleoside reverse transcriptase inhibitors (NNRTIs) among adult patients starting on a first-line ART regimen. This rate is higher in children below 18 months, with over half of newly diagnosed infants harboring resistance to NNRTIs. The prevalence of acquired HIVDR among patients on ART ranges from 3% to 29% (WHO, 2019). Moreover, recent studies in Kenya have shown an upward trend in both transmitted and acquired HIVDR (Hassan et al., 2019; Kantor et al., 2018; Milne et al., 2019).

ART is delivered through the public health approach in most low- and middle-income countries, where standardized drug regimens are administered with simplified laboratory monitoring using tests such as HIV viral load and CD4 count assays (Lessells et al., 2013; De Luca et al., 2013; Phillips et al., 2018). Access to HIVDR testing is limited for patients in resource-limited settings such as Kenya due to high costs involved and inadequate laboratory capacity (Kennedy et al., 2016; Nkengasong et al., 2018; Petti et al., 2006). On the other hand, HIVDR testing is offered routinely in resource-rich setting to inform clinical management of people living with HIV (Dunn et al., 2011; Günthard et al., 2019). However, considerable effort has been made to monitor population level of HIVDR in low- and middle-income countries by implementing HIVDR surveys according to WHO guidelines. These surveys have been crucial in informing national ART guidelines; for example, data on the high prevalence of pretreatment drug resistance to NNRTIs has been critical in the transitioning from an NNRTI-based first-line regimen to a regimen that consists of dolutegravir (DTG) in sub-Saharan countries (WHO, 2019).

In Kenya, there are 10 laboratories across the country that support HIV viral load and EID testing through a PEPFAR and Global Fund funded specimen referral network, and only four of these have capacity to perform HIVDR testing. However, there is a paucity of costing data from these laboratories and no detailed cost analysis has been done (Inzaule et al., 2013). Some of the essential costs when performing a costing analysis include equipment costs, personnel costs, and utilities costs. Previous studies have included reagents cost and consumables cost in their estimations, excluding major cost categories (Acharya et al., 2014; Inzaule et al., 2013; Novitsky et al., 2015; Zhou et al., 2011). A detailed cost analysis provides a deeper understanding of the costs of HIVDR to the health systems. Moreover, cost information is important in development of business plans, projections, planning, budgeting, pricing and resources allocation. Finally, it gives a good guide on the affordability of HIVDR testing inclusion in the standard package of HIV testing in Kenya to alleviate the rising number of cases of HIVDR.

This study we estimate the unit cost of HIVDR testing, identify the cost drivers for the HIVDR test, explore opportunities for cost saving and document challenges and lessons learnt in implementation of HIVDR testing.

Methods
Ethical statement
Ethical approval was obtained from University of Nairobi/Kenyatta National Ethics Review Committee (KNU-UNIV-ERC-P 562/01/2019). Verbal informed consent was obtained from all participants for the interviews and the ethics review board waived the need for a signed consent form. The study followed the guidelines for verbal consent, including explaining to the study participants the pertinent issues about the study including the purpose, benefits, risks and procedures. The participants were also given enough time to decide whether to participate or not as well as an opportunity to ask questions.

Study site
This costing study was conducted at the Molecular and Infectious Diseases Research Laboratory (MIDRL) located within the Kenyatta National Hospital and University of Nairobi School of Medicine. The MIDRL supports the implementation of high quality, sustainable and comprehensive HIV prevention, care and treatment in Nairobi through provision of laboratory services including HIV viral load, HIV early infant diagnosis and HIVDR testing. Data collection was performed during the initial implementation stages, that is within the first year of commencing HIVDR testing.

Costing methodology
The study utilized both micro and gross costing in quantification and valuation of the cost categories, which was done from the provider’s perspective.

Data collection and analysis
Data were collected and compiled from 1st January to 30th June 2019. Costs were assessed for various processes in the
HIVDR testing workflow including laboratory administration, sample collection and preparation, viral RNA extraction, nucleic acid amplification, gel electrophoresis, Sanger sequencing, data analysis and reporting. At the time of data collection, the laboratory used a commercial HIVDR assay supplied by Thermofisher (Cat. no. 12183018A, Waltham, USA). Cost data were collected on annualized depreciation for capital items including laboratory equipment and furniture, long term training and information technology equipment at a depreciation rate of 10%, reagents and consumables, personnel, utilities, laboratory and office space, quality assurance program, and maintenance costs. Cost details were obtained from quotations, invoices and delivery notes.

Data was collected by RG, who at the time of the study was working at MDRL as a clinical laboratory assistant and a master’s student (Health Economics and Policy) at the University of Nairobi. Data collection involved a time and motion study of the laboratory procedures for HIVDR testing and interviewing of laboratory and management personnel. The time and motion study was carried out in 12 sessions lasting between 1–3 hours based on the length of the laboratory procedure. A structured questionnaire depicting all the HIVDR testing steps and data collection tables were used to document quantity of reagents and consumables used as well as duration of each HIVDR process (Gachogo et al., 2020b). Two laboratory technical staff and one member of management were interviewed in three sessions (1 hour) each. Individuals with in-depth knowledge on HIVDR testing implementation were purposively selected to participate in the interviews. Interviews took place within their work environment. An interview guide was used to conduct the interview process and the data was recorded in form of field notes (Gachogo et al., 2020b). Technical staff were interviewed about HIVDR testing processes and experiences of implementing testing, while the member of management was interviewed about cost data, which was recorded in the data collection tables. None of the interviewees declined to participate. Administrative records such as invoices, requests for quotations and delivery notes obtained from the program archives were reviewed to obtain purchase costs. In addition, the laboratory personnel were asked to narrate their experience in setting up a HIVDR testing laboratory, with particular interest on problems encountered and solutions to these set-backs.

Data analysis

We developed a Microsoft Excel version 2010 based model to aid in estimation of the unit cost and cost for the various categories and laboratory processes. All costs were converted to US dollars (13th April 2019, $1 USD = 101.2 Kshs). The MIDRL projected to perform 1000 tests in 2019 based on the number of HIVDR tests performed in the first and second quarter of the 2019 financial year. Responses on challenges experienced during the implementation of HIVDR testing were manually scanned through by RG and FO for developing themes and coded accordingly.

Sensitivity analysis

One-way sensitivity analysis for 20% variations to cost categories was performed to establish the level of uncertainty linked with costs variation of inputs to HIVDR test. This involved varying capital, personnel, reagents, maintenance, and quality assurance program costs by ±20% and evaluating how each of them influence the HIVDR unit cost relative to the estimated cost.

Results

HIVDR testing process

HIVDR testing is carried out in five major processes; namely, sample collection and preparation, nucleic acid extraction, nucleic acid amplification, sequencing and sequence analysis. Specimen collection and preparation involves collection of whole blood from the patient into blood collection tubes that contain ethylenediaminetetraacetic acid (EDTA) anticoagulant. Once the blood is collected into the blood collection tubes, the specimen is prepared for storage by spinning, pipetting and aliquoting into storage vials. The second step in HIV resistance testing is nucleic acid extraction from the plasma. In this step, HIV ribonucleic acid (RNA) is isolated from plasma. Once extracted and purified, the nucleic acid is converted to complementary deoxy-ribonucleic acid (cDNA) and amplified by polymerase chain reaction (PCR) and sequenced. Sequencing involves amplicon purification, cycle sequencing, amplicon purification, sequence detection and visualization. The last step in HIVDR testing is sequence analysis, which involves sequence data validation, sequence assembly, interpretation and quality analysis.

HIVDR unit cost

Activity-based costing for HIVDR testing was performed at MIDR Laboratory by collecting cost data for each step in the drug resistance testing. The cost for performing HIVDR testing was US$ 271.78 per test, where capital costs took the biggest share at $102.42, followed by reagents and consumables at $101.5 (Table 1 and Figure 1). Other costs included personnel ($46.81), utilities ($14.69), maintenance cost of equipment ($2.37) and quality assurance program ($4.00) (Gachogo et al., 2020a).

Cost per laboratory process

The sequencing step had the largest cost of $160.94 per test, while DNA/RNA amplification had the second largest cost of $153.74. Figure 1 shows the cost per laboratory process.

| Item                        | Cost per test | %  |
|-----------------------------|---------------|----|
| Capital cost*               | 102.42        | 37.68|
| Reagents + consumables      | 101.48        | 37.35|
| Personnel                   | 46.81         | 17.22|
| Utilities                   | 14.69         | 5.41 |
| Maintenance cost of equipment | 2.37         | 0.87 |
| Quality assurance program   | 4.00          | 1.47 |
| Total cost                  | 271.78        | 100.00|

* The most costly component of HIV resistance testing.

Table 1. Cost breakdown for each category. Costs in USD.
analysis and sample collection had a cost of $22.79, $10.34, $19.16 and $2.41, respectively (Table 2 and Figure 2).

Cost of the modified HIVDR assay
The laboratory validated a low-cost assay, whereby reagents volumes used during the amplification and sequencing steps were half the recommended volumes by the manufacturer. The test performance was in agreement with the original assay as previously reported (Magomere et al., 2019). The unit cost as a result of halving reagents volumes at the amplification and sequencing step was $233.81, a reduction from $271.78 of the original assay. There was a notable reduction for the amplification and sequencing costs to $38.38 and $140.70 from $56.14 and $160.94, respectively. There was no change in costs for the other steps in HIVDR testing (Table 3 and Figure 3).

Cost of HIVDR test using US Food and Drug Administration (FDA)-approved assay
The cost for the HIVDR test using Viroseq HIV genotyping reagents and consumables (Abbott Molecular, Abbott Park, IL) was estimated at $379.46. This is one of the FDA-approved HIVDR tests available on the market and is used as an alternative to in-house reagents and consumables manufactured by Thermofisher. Reagents and consumables accounted for 55% ($209.18) of the unit cost of the HIVDR testing (Table 4).

Challenges and lessons learnt
As a startup laboratory, the challenges and lessons learnt during the processes of establishing such a capital-intensive undertaking in a resource-limited setting were documented. Table 5 shows some of the challenges and lessons learnt.

Sensitivity analysis
The costs presented assume that the laboratory runs 1000 HIVDR tests per year with no machine breakdown or waste of supplies. Considering that variation in input costs would have an impact on the input costs, a one-way sensitivity analysis for 20% variations to cost categories was performed. Variations to capital, reagents, and personnel inputs had a major impact on the unit cost, whereas variations to utilities, maintenance and quality assurance results had no significant impact on the unit cost. A 20% variation to capital and reagents results in changes of up to 7.5% in unit cost; approximately a $20 difference (Figure 4).

Discussion
The aim of this study was to establish a detailed cost profile for HIVDR testing from a provider’s perspective and identify cost drivers. We also report the challenges encountered and lessons learnt during the implementation of HIVDR testing at the MIDRL. The cost of performing HIVDR testing was estimated to be $271.78 per test. The cost estimate represents all the inputs required for performing HIVDR testing including; capital, personnel, reagents, consumables, quality assurance program and service contracts for the laboratory equipment for performing 1000 tests per year. Previous studies did not include all cost categories, making it difficult to compare costs for offering drug resistant tests across many laboratories (Acharya et al., 2014; Alemán et al., 2015; Inzaule et al., 2013; Novitsky et al., 2015). In this study, we offer a framework for performing laboratory cost analyses that makes it easy to compare cost categories between different laboratories. A previous study from KEMRI/CDC, Kenya, performed a cost analysis of their in-house assay and established the unit cost to be approximately $113.33, with $109.31 as the cost of reagents and consumables (Inzaule et al., 2013). The analysis included the costs of reagents and consumables and the cost of maintaining the equipment but did not account for capital, personnel and external quality assurance program costs. Considering the reagent and consumable costs, there was a correlation in the cost results for these items, as our study estimated the cost to be $101.50. The slight difference could be attributed to time difference in performing the cost analysis. In addition, the previous study estimated the equipment maintenance cost to be $4.02, while the present study estimates
Table 2. Cost breakdown by laboratory processes. Costs in USD.

| Item                        | DNA/RNA extraction | DNA/RNA amplification | Gel electrophoresis | Sequencing* | Sequence analysis | Sample collection | Total cost |
|-----------------------------|--------------------|------------------------|----------------------|-------------|------------------|-------------------|------------|
| Capital cost                | 7.66               | 4.48                   | 6.55                 | 82.57       | 1.21             | 0.10              | 102.56     |
| Reagents + consumables      | 5.42               | 37.30                  | 1.47                 | 55.63       | 0                | 1.67              | 101.48     |
| Personnel                   | 4.68               | 9.36                   | 1.87                 | 15.44       | 15.44            | 0.47              | 47.27      |
| Utilities                   | 4.39               | 3.96                   | 0.20                 | 5.20        | 0.41             | 0.11              | 14.28      |
| Maintenance cost of equipment | 0.24              | 0.24                   | 0.09                 | 0.78        | 0.78             | 0.02              | 2.15       |
| Quality assurance program   | 0.40               | 0.80                   | 0.16                 | 1.32        | 1.32             | 0.04              | 4.03       |
| Total cost in USD           | 22.79              | 56.14                  | 10.34                | 160.94      | 19.16            | 2.41              | 271.78     |
| % Total cost               | 8.38               | 20.66                  | 3.8                  | 59.22       | 7.05             | 0.89              | 100        |

* The most expensive step in HIV drug resistance testing.

Figure 2. Distribution of costs of HIV drug resistance testing laboratory processes, including DNA/RNA extraction, DNA/RNA amplification, gel electrophoresis, sequencing, sequence analysis, and sample collection. Unit cost is $271.78.

A cost of $2.37 for this category. Some of the studies performed in other parts of the world found considerably lower reagents and consumables costs than those estimated by this study. For instance, the costs reported in India and Cuba were $85.00 and $87.80, respectively (Acharya et al., 2014; Alemán et al., 2015). Conversely, one study reported higher reagent costs than those found in the present study; the estimated cost was $139.75 per test (Novitsky et al., 2015).

To answer the question of the cost drivers for HIVDR testing, the costs were categorized according to the processes involved in HIVDR testing, including sample collection, RNA extraction and amplification, gel electrophoresis, sequencing, and sequencing analysis. In terms of cost categories, capital cost took the biggest share of pie at $102.42 (37.68%), followed by reagents plus consumables at $101.50 (37.35%). High capital costs could be attributed to sub-optimal utilization of the sequencing platform. It should be noted that the equipment required for HIVDR testing can be leveraged to perform more patient tests that support HIV care and treatment, therefore bringing down the cost of equipment attributed to HIVDR testing. Our cost analysis was based on an estimated projection of 1000 HIVDR tests per year, which is a gross underestimation of the laboratory’s capacity. If the laboratory operated optimally, offering approximately ~6720 tests per year, the capital cost would reduce ~6.7 fold.

The reagent costs were considerably high as a result of acquiring
Table 3. Cost breakdown of the modified assay.

| Item                          | DNA/RNA extraction | DNA/RNA amplification | Gel-electrophoresis | Sequencing analysis | Sample collection | Total cost |
|-------------------------------|--------------------|------------------------|---------------------|--------------------|-------------------|------------|
| Capital cost                  | 7.66               | 4.48                   | 6.55                | 2.57               | 0.1               | 102.60     |
| Reagents + consumables        | 5.42               | 19.3                   | 1.47                | 35.38              | 0                 | 63.24      |
| Personnel                     | 4.68               | 9.36                   | 1.87                | 15.34              | 0.47              | 47.27      |
| Utilities                     | 4.39               | 3.96                   | 0.2                 | 5.2                | 0.11              | 14.28      |
| Maintenance cost of equipment | 0.24               | 0.47                   | 0.09                | 0.78               | 0.02              | 2.39       |
| Quality assurance program     | 0.40               | 0.8                    | 0.16                | 1.32               | 0.04              | 4.03       |
| Total cost in USD             | 22.79              | 38.38                  | 10.34               | 140.70             | 19.16             | 233.81     |
| % Total cost                  | 9.74               | 16.41                  | 4.42                | 60.18              | 8.21              | 100        |

1 Halving reagent volumes at DNA/RNA amplification step reduces the cost of this step from $56.14 to $38.38.
2 Halving reagent volumes at sequencing step reduces the cost of this step from $160.94 to 140.70.
3 Halving reagent volumes at DNA/RNA amplification and sequencing steps reduce the HIV drug resistance test cost to $233.81 from $271.78.

![Figure 3. Cost distribution of the modified assay among HIV drug resistance processes including capital, reagents + consumables, personnel, utilities, maintenance cost of equipment and quality assurance program. Total unit cost is $233.81.](image)

Figure 3. Cost distribution of the modified assay among HIV drug resistance processes including capital, reagents + consumables, personnel, utilities, maintenance cost of equipment and quality assurance program. Total unit cost is $233.81.

the sequencing machine at no upfront cost. This bound the laboratory to only procure reagents and consumables from the machine provider. This commitment denies the laboratory an opportunity to practice strategic purchasing, which would be a key factor in lowering the cost of reagents. This is not unique to HIVDR testing; a study done in Kenya to estimate cost of HIV viral load and EID reported high reagent costs as a result of machine acquisition on a placement basis (Cintron et al., 2017).

A comparison with other studies was impossible as most of the previous cost analysis included reagents and consumables costs only, omitting other categories such as capital, personnel, utilities and quality control program costs (Alemán et al., 2015; Inzaule et al., 2013; Novitsky et al., 2015). In terms of cost per process, the sequencing step, which involves purification of PCR products, cycle sequencing, purification of sequencing products and sequence detection, was the most costly step in HIVDR testing at $160.94 (59.22%). This is in keeping with
Table 4. Costs for HIV drug resistance test using US Food and Drug Administration approved reagents (Viroseq HIV genotyping). Costs in USD.

| Item                                | Cost per test | %  |
|-------------------------------------|---------------|----|
| Capital cost                        | 102.42        | 26.99 |
| Reagents and consumables*           | 209.18        | 55.13 |
| Personnel                           | 46.81         | 12.34 |
| Utilities                           | 14.69         | 3.87 |
| Maintenance cost of equipment       | 2.37          | 0.62 |
| Quality assurance program           | 4             | 1.05 |
| **Total Cost**                      | **379.46**    | **100** |

* Reagents and consumables are the most expensive inputs to HIV drug resistance testing in Viroseq HIV genotyping.

Table 5. Challenges and lessons learnt.

| Challenges: Challenges experienced in the initial implementation of HIV drug resistance testing | Lessons learnt: Some of the solutions applied to overcome initial challenges |
|------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------|
| High staff turnover.                                                                         | Building capacity through training grant application.                       |
| Insufficiencies in the supply chain management.                                              | Strategic memorandum of understanding with the suppliers.                   |
| Lack of functional laboratory network for sample flow, hence sub-optimal utilization of the facility | Instruments suboptimal utilized can be leveraged to be used for other services. |
| Financial sustainability due to decreased funding.                                           | Engagement of key stakeholders.                                              |
| Frequent electricity disconnection                                                          | Need to build relationship with other laboratories.                          |
| Premixed PCR reagents kit, limiting flexibility for use with other tests                    |                                                                               |

Comparing the cost of the HIVDR test to a HIV viral load test used in monitoring and management of people living with HIV, the HIVDR testing cost is higher. A study in Kenya estimates HIV viral load test at $24.63 for non-point-of-care viral load testing and $29.74 for point-of-care HIV viral load testing (Cintron et al., 2017). This is attributed to additional processes in HIVDR test, that is, nested PCR and cycle sequencing.

Other studies evaluating the cost of HIVDR testing. Other studies report $59.88 (52.92%) and $50 (58.82%) as the cost for the sequencing step (Acharya et al., 2014; Inzaule et al., 2013). The one-way sensitivity analysis performed illustrates a cost saving opportunity, for example through negotiating lower reagent prices and maximizing utilization of the sequencing platform, therefore ensuring sustainable use of health financing resources.

Figure 4. Tornado graph for one-way sensitivity analysis, costs in USD. Grey represents 20% reduction in the input costs, while dark grey represents 20% increase in the input costs. Unit cost is $271.78.
processes. These increase the amount of cost inputs used in HIVDR testing, especially staff hands-on time.

The study evaluated the effects of reducing the reagents volume on the cost and performance characteristics of the HIVDR testing in view of reagents being one of the cost drivers for HIVDR testing. On the cost of the HIVDR testing, there was a significant reduction in the cost from $271.78 to $247.30; a ~13.97% reduction in the cost per test. This assay modification led to a ~37.68% reduction in reagent costs. Of note is the concordance of the two assays in their performance characteristics, which increases the confidence in adoption of this cost saving undertaking by the laboratories that would like to increase their efficiency in offering the HIVDR testing service (Magomere et al., 2019). The new assay performance characteristics met the WHO HIVDR validation criteria (WHO, 2018). Cost computation using Viroseq reagents, which are FDA approved and an alternative to in-house Thermofisher (Illinois, US) reagents, gave a cost of $379.46 per test, with reagents taking the biggest share of the cost at 55.13% ($209.18). This illustrates a lower cost of HIVDR testing using Thermofisher reagents by $107.68. These findings are in keeping with other studies where the cost of HIVDR per test was lower when using the in-house system compared to the Viroseq system (Acharya et al., 2014; Inzaule et al., 2013; Zhou et al., 2011). For instance, one study reported a $132.86 difference in the two systems, while another reported a $165.01 difference (Inzaule et al., 2013).

One of the challenges encountered during the implementation of HIVDR testing was high staff turnover. This is attributed to advanced molecular skills required for sample analysis in HIVDR testing. There are a few laboratory specialists equipped with these skills, making them highly sought after in the job market. This is a challenge in a low resource set up where skill shortages often limit productivity. One solution to this challenge is to train laboratory staff to the highest level possible and provide machine service when due. Maintaining a good working relationship with other laboratories performing the test helps in the exchange of new ideas and also facilitates an inter-laboratory proficiency testing program.

Unlike HIV viral load and EID, HIVDR testing is not included in the Global Access Program, which has helped in the scaling up of HIV viral load and EID testing in Kenya at a relatively low cost (WHO, 2014). This raises sustainability concerns owing to recent reduced donor funding for HIV programs. However, HIVDR testing services can leverage on already established sample referral networks, human resources, laboratory equipment and database for HIV viral load. The multiple possible applications of the sequencing platform provides opportunities to deploy it for other tests and services, therefore reducing the overall running costs. Sensitization of key stakeholders involved in management of people living with HIV through regular stakeholders meetings has been instrumental in uptake of HIVDR test.

Other challenges experienced during the implementation of drug resistance testing included frequent electricity disconnections, which was solved by installing a backup generator to ensure a constant supply of power. This corroborates other previous studies that highlighted similar findings in resource limited settings (Kennedy et al., 2016; Nkengasong et al., 2018). Furthermore, supply chain insufficiency, which delayed timely delivery of reagents, consumables, and laboratory equipment, was a major setback in implementing HIVDR testing. Finally, the premixed PCR master-mixes limited the flexibility of their use for other tests.

**Strengths of the study**

This report presents findings from a complete cost analysis performed in the early stages of implementation of HIVDR testing, hence giving a good picture of the costs involved in the process. This report will further serve as a useful resource for planning and budgeting information for better resource management for similar projects in future. The inclusion of the cost-saving assay evaluation makes the study one of a kind, as it provides an evidence of cost reduction and comparable performance characteristics for both assays.

**Study limitations**

The study estimated costs from the provider’s perspective, thus limiting the inclusion of cost incurred by patients. The study design also excluded transport costs incurred for the transport of samples from peripheral health facilities to the testing laboratory in Nairobi. Furthermore, the cost analysis was carried out in only one facility, hence hindering the comparison across facilities offering HIVDR testing. This study presents a partial economic evaluation; a complete economic evaluation would give a clearer picture on the cost-effectiveness of HIVDR testing versus the status quo. Finally, at the time of the study the laboratory was not operating at full capacity, which increases the unit cost of HIVDR testing. It is conceivable that once uptake for the HIVDR test increases, the additional volumes would translate to reduced costs.

**Conclusion**

The MIDRL has implemented HIVDR testing capacity for patients failing ART at a cost of $271.78 per test. The most important cost driver is expenditure on capital cost, which is likely to reduce when utilization of the equipment increases. It has also been demonstrated that there are opportunities for cost saving through assay modifications such as selective reagent volume reduction.

**Data availability**

**Underlying data**

Figshare: Cost analysis of implementing HIV drug resistance testing in Kenya: a case study of a service delivery site at a tertiary level hospital in Kenya: https://doi.org/10.6084/m9.figshare.12561980.v3 (Gachogo et al., 2020a).
This project contains the following underlying data:

- HIVDR_Consumables_cost.xlsx
- HIVDR_BuildingCost.xlsx
- HIVDR_ElectricityCost.xlsx
- HIVDR_Equipment_cost.xlsx
- HIVDR_Indirectcost.xlsx
- HIVDR_Personnel_cost.xlsx
- HIVDR_Reagents_cost.xlsx
- HIDVR_Viroseq_consumbles.xlsx
- HIVDR_viroseq_reagents.xlsx

Extended data

Figshare: Cost analysis of implementing HIV drug resistance testing in Kenya: a case study of a service delivery site at a tertiary level hospital in Kenya. https://doi.org/10.6084/m9.figshare.12628031.v1 (Gachogo et al., 2020b).

- HIVDR_Questionnaire.pdf
- Interview Guide.pdf

Data are available under the terms of the Creative Commons zero “No rights reserved” data waiver (CC0 1.0 Public domain dedication).

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Joses Muthuri Kirigia
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Recommendation to the Editor:
This is an important and pertinent manuscript in the ongoing fight against HIV/AIDS. The estimated unit cost is important for planning and budgeting purposes. Therefore, I recommend acceptance after some minor suggested revisions.

Research Article
- Is the work clearly and accurately presented and does it cite the current literature?
  Yes.

- Is the study design appropriate and does the work have academic merit?
  Yes. It is a cost analysis of HIV drug resistance tests study in Kenya.

- Are sufficient details of methods and analysis provided to allow replication by others?
  Yes. However, I suggest that the authors explain more about the method used to estimate the annual cost of capital inputs. Please see the detailed suggestion to the authors below.

- If applicable, is the statistical analysis and its interpretation appropriate?
  The statistical analysis is not applicable for this kind of study.

- Are all the source data underlying the results available to ensure full reproducibility?
  Yes, the study references the repository (figshare) where data has been deposited.

- Are the conclusions drawn adequately supported by the results?
  Yes, the conclusions are adequately supported by the results.

Other discretionary suggestions for the authors

Abstract
- Under Results section (page 1), amend the last sentence slightly to read as:
  “A user-initiated modification of halving reagent volumes for some laboratory processes
(amplification and sequencing) reduced the unit cost for a HIVDR test to $233.81, i.e. a 13.97% reduction.

MAIN TEXT

BACKGROUND
○ No amendment suggested.

METHODS
Data collection and analysis
○ On page 3, first paragraph, the third sentence reads “Cost data were collected on annualized depreciation for capital items including laboratory equipment and furniture, long term training and information technology equipment at a depreciation rate of 10%, ...”.

○ Suggestions (A): I suggest that the authors explain more about the method used to estimate the annual cost of capital inputs. As Drummond et al. explain, capital inputs are assets that are used over periods of more than one year; and depreciate (wear out) with time. Capital costs have two components. (1) Opportunity cost: lost opportunity to invest money used to purchase a capital input to yield positive benefit, e.g. the money could have been put in a fixed deposit to earn interest income. (2). Capital input depreciates over time – thus, its value decreases the longer the life span. Thus, costing of capital inputs should capture both components. As Drummond et al explain, there are two best methods costing capital inputs: (1) To calculate the equivalent annual cost, i.e. annuitize the initial cost of purchase of equipment (or building) over its useful life (years). In calculating the equivalent annual cost, one should take into account both the capital input useful life (in years) and a discount rate. (2) Where competitive market exists to use the rent (per square metre) for building space, and lease price for equipment. The rental and lease price would capture both the depreciation and the opportunity cost. The accounting approach of dividing the price of a capital input by the length of its useful life does not capture fully the two cost components mentioned above. Reference: Drummond MF, Sculpher MJ, Torrance GW, O’Brien BJ, Stoddard GL. Methods for the economic evaluation of health care programmes (3rd Edition). Oxford: Oxford University Press; 2005.¹ In Drummond et al., refer to pages 64 and 65 for explanation and pages 72-75 for the formula for estimating the equivalent annual cost of a capital item.

○ Suggestions (B): Authors should justify the choice discount rate of 10%. The authors cite a published article that has used the same discount rate, and that would suffice.

○ Suggestions (C): On page 3, first paragraph, the fourth sentence reads “Cost details were obtained from quotations, invoices and delivery notes.” Please replace the word “Cost” at the beginning of the sentence to “Price”.

Data analysis
○ Page 4: Please consider modifying the first sentence to read as follows: “We developed a costing model in Excel Software (Microsoft, New York) to aid in the estimation of the unit cost and cost for the various categories and laboratory processes.”
Sensitivity analysis

○ Page 4: The first sentence in this subsection reads “One-way sensitivity analysis for 20% variations to cost categories was performed...”. Authors could cite published studies to justify the choice of 20% variation.

RESULTS

○ **Suggestion (A):** Page 4: I suggest that the authors consider moving subsection titled “HIVDR testing process” and the first sentence of subsection titled “HIVDR unit cost” to the methods section.

○ **Suggestion (B):** Slightly amend the Title of Table 1 to read as: “Table 1. Cost breakdown for each category (US$)” Please reflect the minor amendment in Tables 2, 3, and 4.

○ **Suggestion (C):** The Figures 1, 2, 3, and 4 portray results that are already contained in Tables 1, 2, 3, and 4. Therefore, I suggest for authors to choose between presenting results either Tables or Figures.

Cost of the modified HIVDR assay

○ **Suggestion (A):** Page 5: Please move the first sentence to methods section.

○ **Suggestion (B):** Page 5: Insert the percentage change of “13.97%” into the second sentence. Such that it reads as follows: “The unit cost as a result of halving reagents volumes at the amplification and sequencing step was $233.81, a 13.97% reduction from $271.78 of the original assay.”

Challenges and lessons learnt

○ In Table 5, rephrase the fifth challenge to read as “Frequent electric power outage”. Rephrase the second lesson learnt to read as “Strategic memorandum of understanding between the laboratories and the suppliers”. Rephrase the third lesson learnt to read as “Underutilised equipment can be used for other programmes”.

Sensitivity analysis

○ **Suggestion:** Page 5: In the sensitivity analysis subsection, please delete the first two sentences because they were already stated in Methods.

DISCUSSION

○ Page 5: The ninth sentence reads as “Considering the reagent and consumable costs, there was a correlation in the cost results for these items, as our study estimated the cost to be $101.50.”

○ **Suggestion (A):** Please reword the sentence to remove the word “correlation”. I guess you want to say that your finding is very similar to those from cited studies. Correlation is a statistical term.

○ **Suggestion (B):** Page 6: Please move the first sentence of the second paragraph to the Methods section. The sentence I am referring to starts as “To answer the question of the cost drivers for HIVDR testing...”. Page 9: In the second paragraph, the second and third sentences read as “On the cost of
the HIVDR testing, there was a significant reduction in the cost from $271.78 to $247.30; a ~13.97% reduction in the cost per test. This assay modification led to a ~37.68% reduction in reagent costs.”.

- **Suggestion (C):** Page 9: In the Results section, I could not find the result of $247.30. The percentage change from $271.78 to $247.30 is not ~13.97% but 9.1%. Also, I wonder whether the use of the symbol “~” is appropriate. Please check.

- **Suggestion (D):** Page 9: Should the third sentence read as follows “Unlike HIV viral load and EID, HIVDR testing is not included in the Global Access Program, which could have helped in the scaling up of HIV viral load and EID testing in Kenya at a relatively low cost (WHO, 2014).” Incase, the sentence is correct as it was, please ignore this suggestion.

**CONCLUSION**

- **Page 9:** Please consider modifying the first and second sentences as follows: “The MIDRL has implemented HIVDR testing capacity for patients with resistance to ART at a cost of $271.78 per test. The most important cost driver is expenditure on capital inputs, which is likely to reduce when utilization of the equipment increases.”

**References**

1. Drummond ME, Sculpher MJ, Torrance GW: Methods for the economic evaluation of health care programmes, 3rd ed. Oxford University Press. 2005.

**Is the work clearly and accurately presented and does it cite the current literature?**

Yes

**Is the study design appropriate and is the work technically sound?**

Yes

**Are sufficient details of methods and analysis provided to allow replication by others?**

Partly

**If applicable, is the statistical analysis and its interpretation appropriate?**

Not applicable

**Are all the source data underlying the results available to ensure full reproducibility?**

Yes

**Are the conclusions drawn adequately supported by the results?**

Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Economic evaluation; cost-effectiveness analysis; cost-utility analysis; cost-benefit analysis; hospital and programme costing; health financing; health economics; health
systems performance assessment; national health research systems

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Reviewer Report 14 September 2020

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○ The study analysed the costs involved for HIVDR testing in Nairobi, Kenya. Such studies are vital to maximize the use of available resources for laboratory assays, and to help optimize the success of HIV antiretroviral therapy outcomes.

○ AIDs – S should be capital letter - AIDS.

○ It should be noted that adherence is also a threat to the success of ART, along with the development of resistance.

○ CD4 cell counts are not recommended anymore – test-and-treat campaigns.

○ Say that DTG is an Integrase Inhibitor.

○ The study gives an overall adequate outline of the major costs involved for HIVDR. As newer technologies, such as Next-generation sequencing, become more freely available (and cheaper), cost analyses studies can be re-evaluated.

Is the work clearly and accurately presented and does it cite the current literature? Yes

Is the study design appropriate and is the work technically sound? Yes

Are sufficient details of methods and analysis provided to allow replication by others? Yes

If applicable, is the statistical analysis and its interpretation appropriate? Yes
Are all the source data underlying the results available to ensure full reproducibility? Yes

Are the conclusions drawn adequately supported by the results? Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Virology; HIV; Infectious Diseases

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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