Ministry of Medical Services &
Ministry of Public Health and Sanitation

EFFECTIVE MANAGEMENT OF
LABORATORY COMMODITIES

PARTICIPANTS MANUAL

NOVEMBER 2012
RECOMMENDED CITATION

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Enquiries regarding the Effective Management of Laboratory Commodities Participants Manual Guide should be addressed to:

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FOREWORD

Laboratory health commodities are essential to the provision of quality health services. Effective Commodity Management which includes rational selection, efficient procurement, effective logistics management systems and promoting rational use is important to ensure improved access to and use of quality diagnostics and laboratory supplies. Several gaps exist in the laboratory supply chain both at national and peripheral levels. These include lack of skills in commodity management, inadequate supply of laboratory commodities, weak inventory management, poor records and weak information flow systems.

The Effective Management of Laboratory Commodities course is designed to address some of these gaps by equipping health care providers with the necessary knowledge and skills on commodity management. The application of this curriculum will go a long way in increasing awareness of the various commodity management aspects and ultimately availability of quality laboratory commodities.

Sincere appreciation goes to all those who dedicated their efforts and resources in the development of this curriculum.

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ACKNOWLEDGEMENTS

We acknowledge the efforts and dedication of those who worked tirelessly towards the development of these training materials and countless health facility staff who shared their experiences and challenges on health commodity management thereby informing the content of the training package.

We appreciate the financial support provided by USAID Kenya and the technical guidance and leadership from MSH/ HCSM program.

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CONTENTS

ACKNOWLEDGEMENTS 3
ACRONYMS AND ABBREVIATIONS 5

1. INTRODUCTION 6
2. PURPOSE OF THE COURSE 6
3. TARGET GROUP 6
4. COURSE DURATION 6
5. CERTIFICATION 6
6. COURSE ORGANIZATION 7
7. TRAINING/FACILITATION 7
8. PERFORMANCE ASSESSMENT 7
9. CURRICULUM IMPLEMENTATION 7
10. CURRICULUM REVIEW AND CHANGE 7

Module 1: Overview of Laboratory Services 8
Module 2: Laboratory Commodity Management 16
Module 3: Good Laboratory Inventory Management 27
Module 4: Laboratory Commodity and Information flow 74
Module 5: Monitoring, Training and Planning (MTP) for laboratory commodities 97
Module 6: Monitoring and Evaluation in Commodity Management and data for decision making 106

Appendix 1: Sample letter for practicum site visit 120
Appendix 2: Action Planning 121
Appendix 3: Sample course timetable 122
Appendix 4: Inventory Management tools 125
Appendix 5: Issue and Receipt of commodities 135
Appendix 6: Disposal of Lab reagents or items (F.O.58) 140
Appendix 7: Reporting tools or LMIS tools with instructions; 142
Appendix 8: Session and Overall Course Evaluations 151

Reporting Losses of Laboratory Commodities 152
ACRONYMS AND ABBREVIATIONS

AFB  Acid-Fast Bacilli
AIDS  Acquired Immune Deficiency Syndrome
AMC  Average Monthly Consumption
AMR  Antimicrobial resistance
ART  Antiretroviral Therapy
CD4  Cluster of Differentiation Number 4
CD8  Cluster of Differentiation Number 8
DAR  Daily Activity Register
D-CDRR  District Consumption Data Report and Request form
DH  District Hospital
DMLT  District Medical Laboratory Technologist
FBC  Full Blood Count
FBO  Faith Based Organization
F-CDRR  Facility Consumption Data Report and Request
FEFO  First to expire, first out
FIF  Facility Improvement Fund
FIFO  First in, first out
HCSM  Health Commodities & Services Management program
HIV  Human Immunodeficiency Virus
KEMSA  Kenya Medical Supplies Agency
LMIS  Logistics Management Information System
LMU  Logistics Management Unit
M&E  Monitoring and Evaluation
MOMS  Ministry of Medical Services
MOPHS  Ministry of Public Health and Sanitation
MSH  Management Sciences for Health
MTP  Monitoring, Training and Planning
OJT  On Job Training
NASCOP  National AIDS & STI Control Programme
NPHLS  National Public Health Laboratory Services
PGH  Provincial General Hospital
PLWHA  People Living with HIV & AIDS
PITC  Provider initiated counseling and testing
PMLT  Provincial Medical Laboratory Technologist
PMTCT  Prevention of mother to child transmission
RDT  Rapid Diagnostics Tests (Malaria)
S11  Counter Requisition and Issue Voucher
S12  Issue and Receipt Voucher
SCMS  Supply Chain Management System
SOPs  Standard Operating Procedures
STD  Sexually Transmitted Diseases
TB  Tuberculosis
TOTs  Trainers of Trainers
USAID  United States Agency for International Development
VCT  Voluntary Counseling and Testing
1. INTRODUCTION

Currently the laboratory services are wanting in terms of service delivery and availability of laboratory commodities. Laboratory service is essential in disease management, diagnosis and surveillance. This can only be achieved when Lab commodities are available and managed appropriately to address shortages, stock outs, expiries and wastages. This is partially as a result of inadequate training on commodity management.

Most Laboratory personnel have undertaken their training in local medical training institutions that do not incorporate Laboratory commodity management in their curricula. To bridge the gap in the number of staff requiring these skills and knowledge, MOMS/MOPHS with support from various stakeholders developed this curriculum and therefore reduce the challenges noted above.

2. PURPOSE OF THE COURSE

The goal of this course is to equip participants with the necessary knowledge and skills in Laboratory commodity Management practices.

Specific objectives of the course are to enable participants to:
- Discuss the background situation of laboratory services in Kenya.
- Describe the concepts of effective management of laboratory commodities.
- Discuss good Inventory management
- Outline the national system for flow of laboratory commodities and laboratory commodity information
- discuss quality improvement of laboratory commodity management
- describe the concept of monitoring and evaluation and its importance in program activities implementation

3. TARGET GROUP

This course is designed for staff in Primary Healthcare settings who are involved in managing laboratory commodities. These include Laboratory Technologists and Technicians, Nurses & Clinical Officers, store personnel, procurement officers and health records personnel.

The course is also useful for pre-service and in-service laboratory commodity management trainers and On-the-job training (OJT) by laboratory commodity management Mentors.

4. COURSE DURATION

This is a three day residential training.

5. CERTIFICATION

The participants of the course will be awarded a certificate of attendance after attending all sessions outlined in this curriculum.
6. COURSE ORGANIZATION

This course is organized into six (6) modules which are closely related.
Module I: Overview of Laboratory Services
Module II: Laboratory Commodity Management
Module III: Laboratory Inventory Management
Module IV: Introduction to Laboratory Commodity and Information Flow and Use of data for decision-making
Module V: Planning to implement Changes in Laboratory commodity management practice
Module VI: Overview of M&E, Data Quality and Use of Data for Decision Making

7. TRAINING/FACILITATION

This course will involve various teaching methods and will emphasize methods appropriate for adult learners. These methods will include overview and illustrated lectures, small group discussions, class exercises, case studies and demonstrations.

8. PERFORMANCE ASSESSMENT

Facilitators will assess participants through question and answer sessions. Pre- test and post-test questions will be administered at the beginning and at the end of the course. This will form an important part of the assessment. In addition, evaluations will be administered to obtain feedback on the course content and delivery.

9. CURRICULUM IMPLEMENTATION

The trainers/facilitators for the course will be drawn from healthcare providers who have expertise and experience in laboratory commodity management. The time schedule for the course will be between 8 am to 5 pm with tea breaks and a lunch break in between. The course will last approximately 3 days providing 8 hours of didactic and practical sessions as outlined in this curriculum.

10. CURRICULUM REVIEW AND CHANGE

There will be a course evaluation by participants and implementers at the end of each course and appropriate changes should be made where necessary. After the first course there will be a review of the curriculum by all stakeholders. The stakeholders will evaluate the course objectives, course content and all the resources for the course. Subsequent periodic reviews will be conducted after 5 years.
MODULE 1: OVERVIEW OF LABORATORY SERVICES

Duration: 1 hour

Goal: To describe the background situation of laboratory services in Kenya.

Objectives

At the end of this module, the participants will be able to:
• Describe the types of laboratory services provided in Kenya
• Identify and describe health policies related to laboratory services
• Identify the challenges in Laboratory services at various levels of the healthcare system.

Content

Components of the health system supported by laboratory services; KEPH levels providing laboratory services; health policies related to laboratory services; challenges in providing laboratory services.

Teaching Aids:
• Kenya Medical Laboratory Policy 2012
• Power point presentation
• LCD Projector, Flip Charts, Flip Chart Stand, Markers, Masking tape

Module 1: Overview of Laboratory Services

Lesson Plan Guide:

<table>
<thead>
<tr>
<th>UNIT</th>
<th>CONTENT</th>
<th>ACTIVITY</th>
<th>TIME</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Overview of Laboratory services</td>
<td>Lectures Group discussion</td>
<td>1 hour</td>
</tr>
</tbody>
</table>

References and Recommended Readings

• Ministry of Medical Services and Ministry of Public Health & Sanitation. Kenya Medical Laboratory Policy 2012
Course Overview

Why the course?

- Currently the laboratory services are wanting in terms of service delivery and availability of laboratory commodities.

- Increased attention to the sector has provided unprecedented opportunity to improve access to quality laboratory services.

- Overriding need to strengthen laboratory commodity management systems.

Course Program

<table>
<thead>
<tr>
<th>Day</th>
<th>Activities</th>
</tr>
</thead>
<tbody>
<tr>
<td>Day 1</td>
<td>Participant registration, introduction and climate setting, pre-test, official opening, Course overview, Overview of Laboratory services, Introduction to Laboratory commodity management (LCM), Roles and responsibilities in LCM</td>
</tr>
<tr>
<td>Day 2</td>
<td>Recap, Introduction to Laboratory Inventory Management, Determining quantities to order, good storage practices</td>
</tr>
<tr>
<td>Day 3</td>
<td>Recap, Receiving and Issuing commodities, Records and Tools supporting Inventory management, Introduction to Laboratory commodity and information flow (LMIS)</td>
</tr>
<tr>
<td></td>
<td>Recap, Use of data for decision-making (monitoring and evaluation), Planning to implement changes in Laboratory commodity management practice (MTP approach), presentation of action plans, Overview of M&amp;E, Data Quality and Use of Data for Decision Making, post test and course evaluation, closing ceremony and awarding of certificates</td>
</tr>
</tbody>
</table>
**Target Audience**

- Laboratory Commodity Management Staff in Primary Healthcare settings  
  *Laboratory Technologists and Technicians, Nurses & Clinical Officers, store personnel, procurement officers and health records personnel OR those health care workers who handle Laboratory commodities*
- In-service laboratory commodity management trainers
- Training of Mentors or TOT in Laboratory commodity Management as a scale up and sustainability strategy
- On-the-job training (OJT) by laboratory commodity management Mentors or TOTs

**Purpose of this Course**

- The purpose of the course is to:  
  Provide the necessary knowledge and skills to enable staff to efficiently and effectively manage laboratory commodities in support of laboratory services

**Specific Objectives of the Course (1)**

- To introduce the principles of effective management of laboratory commodities.
- To impart skills on how to practice good inventory management.
- To describe and apply on good laboratory commodity flow and logistic management information system.
Specific Objectives of the Course (2)

• To apply the concepts of monitoring and evaluation to Lab commodity management

• To enable implementation of changes in Laboratory commodity management practice using the MTP approach

• To orient health care workers on the use of laboratory commodity management tools

Course Structure and Methodology (1)

Structure:
3-day residential training
• 3 days didactic class sessions
• Separate Half or 1-day practicum

or

Individual Module level application according to identified needs or gaps

Course Structure and Methodology (2)

Methodology:
• Pre and Post Course assessment
• Adult learning Training Methods applied:
  – Lectures, Discussions, Buzz sessions,
  Démonstrations, exercises/practice sessions,
  Case studies
• Module and Facilitator evaluations
• Certification
### Course Outline

*This course comprises of 6 broad modules:*

- Module I: Overview of Laboratory Services
- Module II: Laboratory Commodity Management
- Module III: Laboratory Inventory Management
- Module IV: Introduction to Laboratory Commodity and Information Flow and Use of data for decision-making
- Module V: Planning to implement Changes in Laboratory commodity management practice
- Module VI: Overview of M&E, Data Quality and Use of Data for Decision Making

### Course Material

- **Stationery materials for Participants**
  - Pen, Writing pad / Notebook, Name tag, Pocket folder
- **Participant Handouts:** Power-points and Session handouts
- **Sample commodity management tools**
- **Job aids**
- **Calculators**
- **Flipcharts**
Module 1

Overview of Laboratory Services

The Situation of Laboratory services in Kenya

Objectives

• The main goal of this module is to give the participant some background information on situation of laboratory services in Kenya.

• On completion of this module, the participant should be able to:
  – Describe the types of laboratory services provided in Kenya
  – Identify and describe health policies related to laboratory services
  – Identify and discuss the best practices and challenges in current Laboratory services at various levels of the healthcare system.

Background (1)

Laboratory services, as part of the national health system, support the following health service components:
  • Curative
  • Preventive
  • Palliative (monitoring)
  • Health promotion
  • Counseling
  • Surveillance
  • Research
All six levels of health care offer medical laboratory services differentiated according to the KEPH level:

- 6: Tertiary hospitals
- 5: Secondary hospitals
- 4: Primary hospitals
- 3: Health centers, maternities, nursing homes
- 2: Dispensaries/clinics
- 1: Community: Villages/Households/Families/Individuals

Medical laboratory services have existed for many years in Kenya. However the sector has been faced with tremendous challenges:

- Inadequate allocation of resources
- Inadequate infrastructure
- Staffing (HR) - inadequate numbers and skills
<table>
<thead>
<tr>
<th>Challenges (2)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>• Value of the laboratory in supporting diagnosis and management of patients is generally overlooked</td>
<td></td>
</tr>
<tr>
<td>• Limited capacity of Facility laboratories to meet the increased demand for Lab services</td>
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<tr>
<td>• Limited capacity to deal with emerging and re-emerging diseases and cross-border health concerns</td>
<td></td>
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<tr>
<td>• Inadequate capacity to address surveillance in a coordinated manner</td>
<td></td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Rationale for the Effective Management of Laboratory Commodities Training (1)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>• Increased demand for the laboratory services has prompted the need to have an uninterrupted supply of laboratory commodities</td>
<td></td>
</tr>
<tr>
<td>• Efforts to improve Lab commodity has been previously mainly limited to key MoH programs (TB, HIV and Malaria) - therefore need to enhance skills to address management of all Lab commodities</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Rationale for the Effective Management of Laboratory Commodities Training (2)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>• Due to the above, there is need to train personnel involved in laboratory commodity management on good laboratory practices and efficient handling of the commodities</td>
<td></td>
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</tbody>
</table>
Module 2: Laboratory Commodity Management

Duration: 2 Hours 15 Minutes

Goal: To discuss the concepts of effective management of laboratory commodities.

Objectives:

At the end of this module, the participants will be able to:
• Define the terms used in commodity management
• Describe the components of the Laboratory Commodity Management cycle
• Identify and discuss the best practices and challenges in current Laboratory commodity management practices at health facilities
• Describe the roles and responsibilities of healthcare workers handling laboratory commodities.

Content

Unit 1: Laboratory commodity management: Definition of key terms, components of the commodity management cycle, current Laboratory commodity management practices at health facilities, roles and responsibilities of health care workers handling laboratory commodities in service delivery points.

Module 2: Laboratory Commodity Management

Lesson Plan Guide

<table>
<thead>
<tr>
<th>UNIT</th>
<th>CONTENT</th>
<th>ACTIVITY</th>
<th>TIME</th>
</tr>
</thead>
</table>
| 1    | Laboratory commodity management | • Lecture  
|      |                     | • Group discussion / buzz sessions | 2 Hours 15 Minutes |

References and Recommended Readings


Module 2

Laboratory Commodity Management

Objectives

By the end of the session, participants will be able to:

• Define the terms used in commodity management

• Describe the components of the Laboratory Commodity Management cycle

• Identify and discuss the best practices and challenges in current Laboratory commodity management practices at health facilities

• Describe the roles and responsibilities of health care workers handling laboratory commodities in service delivery points

Buzz session

(10 minutes)

• Identify best practices and challenges in current Laboratory commodity management practices in your health facility
Examples of Challenges in Laboratory Commodity Management (LCM)

- Erratic supply of commodities
- Staff shortages
- Staff attitude
- Lack of skills and competency
- Lack of inventory management tools (e.g. bin cards, temperature logs, Expiry tracking charts)
- Lack of recording and reporting tools (e.g. DAR, F-CDDR)
- Inadequate/poor infrastructure

What is Commodity Management?

Commodity management is a set of practices and procedures for ensuring the:
- Availability
- Accessibility
- Effectiveness
- Quality
- Appropriate use of lab commodities in all service delivery points.

Importance of Commodity Management (1)

- **Ensure continuous availability of, and access to,** laboratory commodities for the patients at a health facility at all times
- **Demand:** The availability of supplies, such as HIV test kits, increases the demand for Lab services and encourages health-seeking behaviour
- **Quality** of services is dependent on availability of commodities, and improves staff motivation
Importance of Commodity Management (2)

- **Cost**: Many lab commodities are *costly* to procure. Effective management maximises utilisation of scarce resources and prevents waste.
- Facilitates *task-shifting* to non-lab staff handling lab commodities.
- Promotes *good governance*, including transparency and accountability.

---

**Selection**

- Selection process involves establishing and using a list of carefully selected Laboratory commodities.
- Selection is perhaps the single most cost-effective action to:
  - promote regular supply of quality commodities
  - prevent the wasting of scarce resources on unnecessary or ineffective commodities
- Selection helps you to decide which laboratory commodities to purchase.
- Example: when using funds available at the facility level such as facility improvement fund (FIF) or health sector services fund (HSSF) to procure commodities at the facility.
Criteria for selection of laboratory commodities

The equipment or supplies selected should include:

- Local availability of essential supplies
- Well established brand of simple sturdy design
- Local possibilities for servicing and spare parts for equipment
- Local possibilities for training staff in use, service and maintenance.
- Cost considerations

Procurement

- Procurement is the process through which items are purchased or obtained from suppliers
- The procurement management process is described in the flow diagram below
Key Components of Procurement

• Need to know how much to procure and for what periods (Forecasting & Quantification)
• Who to buy from (supplier selection)
• How to buy (e.g. pooled procurement)
• How to ensure quality of the products you buy (quality assurance)
• Systematic and sequential follow-up of the procurement activity (procurement planning and monitoring)

Benefits of Effective Procurement

• Right commodities or equipment
• Right quantities
• Competitive purchase price
• Meet recognized quality standards
• Timely delivery of commodities
• Reliable supplies
• Sufficient support in service and maintenance

Distribution

• Goal is to deliver procured commodities and supplies to the point of use
• Components of effective distribution include:-
  – Delivering in a timely manner
  – Keeping appropriate records
  – Monitor available stock:
    • Expiry dates
    • Inventory levels
    • Storage conditions, such as light and temperature
Distribution involves.....

- Port Clearing
- Receipt and Inspection
- Inventory Control
- Storage
- Requisition for supplies
- Delivery
- Receipt and Inspection
- Inventory control
- Storage
- Use of the commodities and supplies
- Reporting on use/consumption

Buzz session (5 minutes)

- What are the elements of a good distribution system?

Use (1)

- The appropriate use of laboratory commodities completes the laboratory commodity management cycle

- What are some of the important aspects of appropriate use of laboratory commodities?
Use (2)
Appropriate Use of Laboratory commodities entails:

- Correct use of commodities and supplies to perform tests effectively and efficiently
- Adhering to SOPs, technical specifications and instructions
- Taking into account the cost, i.e. selecting a cost-effective test
- Inclusion of Laboratory managers in the health facility’s Medicines & Therapeutics Committee (MTC) to improve appropriate use of lab commodities in the facility

Group discussion (15 min)
In groups, discuss factors influencing appropriate lab commodity use in relation to the following:

- Prescriber
- Lab staff
- Patients/clients and community
- Health supply system

Factors Influencing appropriate lab commodity use

- Prescriber: prescribing the wrong or unnecessary tests/ investigations
- Lab staff: Lack of competency and skills
- Patients/clients and community: preference for particular tests, prevalent conditions, personal beliefs, mis-information
- Health supply system: poor inventory management, inadequate funding, poor policies
Examples of Inappropriate Use of Lab Commodities

• Requesting for tests that are not necessary.
  – E.g. requesting for full haemogram when only the Hb levels are required
• Requesting for unnecessary investigations for monetary gains
• Prescriber preference for particular tests that may not be necessary
• Lack of knowledge and competency

Types of interventions to improve rational use of laboratory commodities

• Educational ➡️ Persuade
• Regulatory ➡️ Force
• Managerial ➡️ Guide

Management Support

• Enables each component of the Commodity management cycle to function well
• It includes:
  – Organizational structure with a clear flow of authority
  – Demonstrate lab revenue generation and advocate for involvement in the health facility’s budgeting processes
  – Maintenance of accurate, useful, and up-to-date information system
  – Capacity development for lab staff
Management Support

- Good record keeping to provide information
- Adequate financing
- Staff that is adequately trained and supervised
- A program that is well planned and involves the community

Policy, Laws & Regulations

- Affect the above other components of commodity management
- Provides the environment for effective delivery of lab services

E.g.
- Set quality standards at each level
- Define requirements for licensing, i.e. personnel
- Policies on tests, techniques and equipment
- Policies affects allocation of resources / budgets e.g. FIF, HSSF, decentralization

Buzz session
(10 minutes)

- Discuss the Roles and Responsibilities of Laboratory Staff in the Laboratory Commodity Management cycle
Summary of Roles and Responsibilities of Laboratory Staff in LCM (1)

• Selection and decision on what commodities to be ordered
• Procurement: Quantify for the commodity, supplies and equipment requirements
• Distribution: Ensuring safety and security of the commodities, inventory management, daily maintenance of equipment and cold chain management
• Use: observe the guidelines, algorithms, SOPs

Summary of Roles and Responsibilities of Laboratory Staff in LCM (2)

• Management support: awareness of lines of authority, involvement in budgeting, reporting, mentoring staff
• Policy, laws and regulation: be up-to-date and aware
MODULE 3: GOOD LABORATORY INVENTORY MANAGEMENT

Duration: 9 Hours

Goal: To discuss good Inventory management practices

Objectives:

At the end of this module, the participants will be able to:

• Define the key terms used in inventory management and give reasons for the importance of good inventory management
• Define key terms used in quantification and the importance of good quantification for laboratory commodities and supplies; and be able to use one quantification method
• Describe good storage practices in laboratories and in other facility stores and identify and articulate problems of storage at their facilities
• Describe the procedures for receiving and issuing laboratory commodities within a health facility, and the appropriate forms and records used
• Identify and demonstrate the use of records and tools supporting inventory management for lab commodities.

Content:

Session 1: Introduction to Inventory Management: Definition of the terms ‘Inventory’ and ‘Inventory management’ and other key terms; identification of the key components of the Inventory management cycle; the purpose of Inventory management; the characteristics of good inventory management for laboratory commodities; description of the challenges in inventory management.

Session 2: Determining Quantities to order for Lab Commodities: Definition of key terms used in quantification, importance of quantification of laboratory commodities and supplies, challenges / problems in quantification, appropriate choice of quantification method based on the data available, application of concepts of quantification in facility setting by quantifying the commodity needs for a facility using the Consumption method.

Session 3: Requesting Lab Commodities: Appropriate procedures used to request laboratory commodities; appropriate forms and records used when requesting laboratory commodities for a health facility.

Session 4: Good Storage Practices: Various aspects of good storage practices in laboratories and in facility stores, special storage conditions that are applicable to laboratory commodities, problems of storage at the facilities.

Session 5: Receiving and Issuing: Definition of terms (receiving, issuing); procedures for receiving and issuing laboratory commodities within a health facility; the appropriate forms and records used to receive and issue commodities for a health facility; receiving and issuing practices

Session 6: Records and Tools supporting Good Inventory Management practices: Records and tools used in support of good inventory management practices, demonstrating use of the tools and identifying the problems/challenges encountered in using the tools.
Module 3: Good Laboratory Inventory Management

Lesson Plan Guide

<table>
<thead>
<tr>
<th>UNIT</th>
<th>CONTENT</th>
<th>ACTIVITY</th>
<th>TIME</th>
</tr>
</thead>
</table>
| 1    | Introduction to Inventory Management | • Lecture  
• Group discussion / buzz sessions | 20 minutes  
10 minutes |
| 2    | Determining Quantities to order for Lab Commodities | • Lecture  
• Group Exercise  
• Exercise | 1 hour  
1 hour 15 minutes  
30 minutes |
| 3    | Requesting Lab Commodities | • Lecture  
• Group discussion / buzz sessions | 20 minutes  
10 minutes |
| 4    | Good Storage Practices | • Lecture  
• Group discussion | 1 hour  
30 minutes |
| 5    | Receiving and Issuing | • Lecture  
• Group discussion  
• Exercises | 1 hour  
15 minutes  
30 minutes |
| 6    | Records and Tools Supporting Good Inventory Management Practices | • Lecture  
• Demonstration / Exercises | 1 hour  
1 hour |

References and Recommended Readings

Introduction to Inventory Management

Module 3 Session 1

Objectives
By the end of this session, participants should be able to
• Define the terms ‘Inventory’ and ‘Inventory management’ and other key terms
• Identify the key components of the Inventory management cycle
• Describe the purpose of Inventory management
• List the characteristics of good inventory management for laboratory commodities
• Describe the challenges in inventory management

What is Inventory?
• **Inventory** is the commodities, supplies, equipment, and other materials that are available in stock in an institution (e.g. a medical store) at a given time
What is Inventory management?

• Inventory management refers to the processes of ordering, receiving, storage and distribution or issuing of items

• The goal of efficient inventory management is to ensure that commodities are available and accessible so as to provide uninterrupted services.

Buzz session
(5 minutes)

• What are the challenges you see in Inventory management in your health facility?

Challenges in Inventory Management

• Frequent stock-outs of commodities
• Wastage due to expiry
• Weak or no Standard ordering procedures (SOPs)
• Rationing of commodities
• Associated products not ordered concurrently (buffer solution, lancets, etc)
• Multiple vertical commodity sources and pipelines (for HIV lab, TB, Malaria, general Lab, donations)
• Inadequate and/or lack of supply of standard inventory management tools
Inventory Management cycle

- Determining Quantities to Order
- Requesting
- Receiving
- Storage
- Issuing

Should be guided by Standard Operating Procedures (SOPs) AND Supported by good Record keeping

Brain-storming session
(Time: 5 minutes)

Answer the following question:

“What is the purpose of Inventory management?”

Purpose of Inventory Management

- To ensure continuous supply for required items (no stock-outs)
- To determine the quantity of an item to order and the right time to place the order
- To ensure appropriate, safe and secure storage
- To minimize losses, e.g. expiries
Good Inventory management system

A good inventory management system informs the storekeeper -
• When to order or issue
• How much to order or issue
• How to maintain an appropriate stock level of all products to avoid both shortages and overstocks.

Characteristics of Good Inventory Management (1)
• Availability of an established or approved stock list for each store/facility
• Adequate security, e.g. protection against theft
• Accurate records of all stock movements
• Accurate physical stock balances
• All staff members are aware of good inventory management practices
Characteristics of Good Inventory Management (2)

- Clear guidelines stating the maximum and minimum stock levels within which stocks should stay
- Set re-order level that guides on when to order
- SOPs to handle re-distribution of excess stock, and disposal of expired or obsolete stock

Inventory Control - Record-keeping

- To achieve these aims we need inventory records which must always be:
  - legible
  - complete
  - accurate
  - up-to-date

- Inventory management records and tools include: bin / stock control cards, charts to track expiry dates, S11, Temperature monitoring log, etc
Determining Quantities to order for Lab Commodities

Module 3 Session 2

Objectives
At the end of this session, participants should be able to:

– Define key terms used in quantification
– Identify the challenges / problems in quantification
– Outline the importance of quantification of laboratory commodities and supplies
– Quantify the commodity needs for a facility using the Consumption method

What is Quantification?

• A process that involves
  – estimating quantities of a specific item needed for procurement for a specific period of time,
  – the financial requirements needed to procure those items

• Quantification is an important step in procurement and in ordering supplies
Inventory Management cycle

- Determining Quantities to Order
- Requesting
- Receiving
- Storage
- Issuing

Should be guided by Standard Operating Procedures (SOPs) AND Supported by good Record keeping

Buzz session
(10 minutes)

- What are the challenges / problems in Quantification that you have experienced in your health facility?

Terms used in Quantification (1)

- **Consumption**
  - Quantity of an item used to carry out diagnostic tests

- **Consumption review period (CP)**
  - Period over which consumption is being reviewed, usually 1 month

- **Average monthly consumption (AMC)**
  - A measure of the number of units of a commodity that was used in an average month over a specified period (e.g. in the last 1 or 3 or 6 or 12 months)
### Terms used in Quantification (2)

- **Adjusted consumption**
  This is consumption adjusted to cater for when the item was in stock and when it was out of stock, within the consumption review period

- **Maximum stock level (MSL):**
  The maximum quantity of a laboratory commodity that should be held at the facility at any given time

- **Minimum stock level (Min):**
  The minimum quantity of a laboratory commodity that the facility should hold at any given time

### Terms used in Quantification (3)

- **Maximum Months of stock**
  This is the highest stock level that the facility should hold of an item at any given time, expressed in months

- **Stock on hand (SoH) / Closing stock**
  - The quantity of a lab commodity in stock at a specified time (best when based on the Physical count)

- **Quantity to Order (QO)**
  The quantity required to be ordered for the next consumption period, e.g. 1 month

### Terms used in Quantification (4)

- **Stock-out**
  When there is no stock of a commodity, in the whole facility

- **Safety / Buffer stock**
  - This is the minimum quantity that should be kept in stock in the lab at any given time, to cushion against uncertainties in demand and supply.

- **Unit of issue**
  - The quantity of a commodity in a container or pack size (e.g. tests, pieces, mls, litres, gms, vials)
Importance of Quantification

- To avoid surpluses that may lead to over-stocking, expiries and/or wastage of commodities
- To avoid shortages/stock-outs
- To assist with the preparation and justification of a budget
- To make informed adjustments to procurement when faced with budgetary constraints
- To plan for new policies and new or expanding programs
- To estimate how much storage space may be required in the future

Some key issues to consider when Quantifying

- **Disease burden** - To treat a certain proportion of the priority disease burden in the region
- **Laboratory KEPH levels** - Caters for different levels of care
- **Available Laboratory equipment**
- **Budgetary allocation** - To ensure that a certain product accounts for no more than a certain proportion of the available budget
How do you quantify?

Quantification Methods

Two main methods:

• Consumption-based

• Morbidity-based

Consumption-based method

• It estimates the commodity needs based on records of past consumption

• Works well where:
  – All records are available, up-to-date and the information is reliable
  – There are minimal stock out periods
  – Consumption patterns are stable

• Adjustments are made for stock-outs, wastage and losses.
### Morbidity-based Method

- Estimates needs based on the frequency of health problems and their standard management
- Is used to:
  - estimate needs for a new program or in disaster situations or where consumption data is not available/reliable
  - developing and justifying budgets

### Records that provide Information for Quantification

1. **For Consumption**
   - Daily Activity Register
   - Issues Register (issues to clients/patients)
   - Consumption data report & request forms
   - Service data (e.g., number of clients/patients)
   - Lab stock or Bin cards

2. **For Morbidity**
   - District Health records
   - Hospital Health records

### Consumption-based method: Steps

**Step 1:**
**Select the Consumption period (CP)**

Determine the period over which the consumption is being reviewed in months.

Normally for supplies received from central level, the CP is 4 months
Consumption-based method: Steps

Step 2:
Determine the Consumption (C)

Determine the quantity used during the consumption period.
This information is obtained from the DAR or other consumption record, or the lab stock card.

Step 3:
Determine the Average Monthly Consumption (AMC)

\[
AMC = \frac{\text{Consumption (C)}}{\text{Consumption period (CP)}}
\]

Step 4:
Determine the Maximum Months of Stock (Max MoS)

\[
\text{Max MoS} = \text{Desired Consumption period (CP)} + \text{Buffer (in months)}
\]
**Consumption-based method: Steps**

**Step 5:**

Calculate the Maximum Stock level (MSL)

\[ MSL = AMC \times \text{Max MoS} \]

**Step 6:**

Conduct the Physical count to get the Stock on Hand (SoH)

Ensure that the SoH covers the stock at the various service points as well as the lab store, if any.

**Step 7:**

Calculate the Quantity to Order (QO)

\[ QO = MSL - SoH \]
Consumption-based method: Adjusting for stock-outs

To Adjust Consumption for stock-outs:

- Calculate the Adjusted Consumption (C2)

\[ C2 = C \times \left( CP \right) \]

Where:
- \( C \) is the consumption
- \( CP \) is the period in stock (in days)

Appropriate Quantification avoids....
Objectives

At the end of this unit, the participants will be able to:

• Describe the appropriate procedures used to request laboratory commodities

• Describe the appropriate forms and records used when requesting laboratory commodities

• Be able to prepare the appropriate intra-facility orders for a health facility

Inventory Management cycle

Determining Quantities to Order

Requesting

Receiving

Storage

Issuing

Should be guided by Standard Operating Procedures (SOPs) AND Supported by good Record keeping
Requesting Commodities

- Requesting
  The process of ordering for commodities from a point of issue (e.g. the main facility store) or from a selected supplier (e.g. KEMSA)

- Requesting for commodities occurs
  - after quantification is done
  - whenever commodities fall below Re-order level or Minimum stock levels

- Requesting should be documented on official Requisition forms

- There are standard MoH forms for requesting supplies, for example:
  - Counter Requisition & Issue voucher (S11)
  - Facility Consumption Data Report & Request (CDRR) form

Who should Request for commodities?

- Staff in charge of commodities and supplies
  - Laboratory manager (or Laboratory In-charge)
  - Laboratory technologist or technician
  - Stores officer or Procurement officer

- Staff designated by the program / facility
  - E.g. at VCT sites
When to Request?

- When stocks reach or fall below the pre-determined Minimum stock level
- At the scheduled order times for your facility
- Also depends on how long it takes to order and receive (Lead Time)
  **NB:** Lead time (period between ordering and receiving) may have an effect on the order time!

Records that provide information for Requesting commodities

**At Service delivery point / Facility level:**
- Daily activity register (MoH 642) or other Consumption record
- Facility Consumption Data Report & Request (F-CDRR, e.g. MoH 643)
- Service data, e.g. Number of tests conducted, Number of clients
- Bin cards

**At the Central warehouse (e.g. KEMSA):**
- Reports from the facilities i.e. Facility Consumption data report & request (F-CDRR)
- Central warehouse Stock cards (e.g. Bin cards)
- Procurement plans/schedules
**Good Storage Practices**

 Module 3 Session 4

**Objectives**

By the end of this session, participants should be able to:

• Describe the different aspects of good storage practices in laboratories and in other facility stores
• Describe special storage conditions that are applicable to laboratory commodities
• Identify and articulate problems of storage at their facilities
• Discuss safe disposal and waste management

**Inventory Management cycle**

- **Determining Quantities to Order**
- **Requesting**
- **Receiving**
- **Storage**
- **Issuing**

Should be guided by

Standard Operating Procedures (SOPs) AND Supported by good Record keeping
Importance of Proper Storage (1)

- **Stability**: wastage reduced and constant availability assured
- **Retrieval**: easy for issuing or conducting physical counts
- **Quality**: assures maintenance of viability of the items
- **Security**: ensures safety of held stocks, prevents loss from damage, fire, pilferage etc

Importance of Proper Storage (2)

- **Record-keeping**
  
  NB. An inventory can be either in hard or soft copies and should be updated regularly.

- **Retiring** of obsolete equipment

- **Disposal** of expired commodities

**Exercise 1**

Whole group discussion:

10 mins

What factors should be considered for optimal storage of commodities in the laboratory?
Guidelines to Good Storage Practices (1)

- Provide appropriate space and security for stored stock
- Provide safe and orderly arrangement of stock in storage
- Maintain correct storage conditions to safeguard quality
- Good stock control and rotation-(practice FEFO&FIFO)

Guidelines to Good Storage Practices (2)

- Maintain updated laboratory stock cards and the Expiry tracking chart
- Label clearly all reagents and chemicals with their name, date of preparation, date of expiry and biohazard symbol.
  
  NB: For reagents prepared/constituted in the lab, the name of the officer who prepared the reagent should be indicated on the label.
- Store chemicals and reagents in correct zones to make sure incompatible chemicals are not stored together
Guidelines to Good Storage Practices (3)

- Secure storage areas for expensive or dangerous items
- Maintain the correct temperature and other conditions for commodities
- Conduct regular physical stock counts and record the results
- Develop and implement SOPs for storage of laboratory commodities and other supplies

Guidelines for Zoning Lab Items
Exercise 2: Defining Zoning
10 min

1. What does zoning mean/imply?
2. Have you had to zone your lab commodities?

Zoning: Introduction (1)

• What is a Zone?
Is a part of a store; it can be a separate building, room, locked cupboard, a refrigerator, a freezer or a cold room

• What is Zoning?
It is a process of putting items in a particular area or in a separate area due to their special storage requirements or other characteristics.

Zoning: Introduction (2)

• Why Zone Lab Commodities in stores?
  — Safety and Security reasons
  — Reduction of accident risks or contamination
  — Allows for special requirements, e.g. Cold Chain
  — Easier Stock management (e.g. FEFO and FIFO arrangement, stock taking)
Types of zones

- Temperature zones
  - Storage at uncontrolled room temperature
  - Storage at controlled temperature and humidity
  - Cold storage

- Secure storage
  - Fuel should be stored in a locked compound for safety purposes

- Flammables
  - Keep in separate building or store, well ventilated, fireproof, fitted with explosion hatch

Buzz Session 1
(10 min)

In pairs,
match each of the following symbols presented with what it stands for:-

What do these laboratory safety symbols stand for? (1)
What do these laboratory safety symbols stand for? (2)

Zoning - Special Precaution items

Always Check and understand the hazard symbols such as:

a. Very toxic  k. Toxic
b. Irritant      l. Extremely flammable
  c. High voltage m. Explosive
d. Harmful      n. Highly flammable
e. Corrosive    o. Radioactive
f. Oxidizing    p. Dangerous for
  g. Biohazard   Environment
h. Radioactive  q. No open flames
i. Chemical hazard  r. Fire extinguisher
j. Non portable water

Meanings of symbols

Answers to Buzz session exercise

Correct answers for slide 1 are:
1. Electrical hazard
2. Combustible materials
3. No open flames
4. Corrosive materials
5. Flammable
6. Fire extinguisher
7. Toxic chemical
8. Non-potable water
Meanings of symbols
Answers to Buzz session exercise

Correct answers for slide 2 are:
1. Environmental hazard
2. Toxic materials hazard
3. Explosive materials
4. Radioactive
5. Biohazard sign
6. Chemical hazard label

Zoning:- Storage requirement

Optimal storage conditions
• If no special storage conditions are given normal storage conditions apply, i.e.
  – Dry
  – Well ventilated premises
  – Temperatures of +15°C to +25°C or up to +30°C (WHO 1990)
• Each storage zone should have at least one thermometer and temperatures should be recorded daily at the hottest time of the day.

Stock location within a Zone (1)

Within each zone, there may be several location types:-
• **Fixed**: Each stock item is always stored in the same place
• **Fluid**: Items are stored wherever there is space at time of delivery
• **Semi-fluid locations**: A combination of the previous two. Each item is assigned some fixed space for picking stock. The remainder of the store is filled on the fluid location principle.
Stock classification within a zone (2)

- Items should be clearly organized within each zone of the store
  - Alphabetical order
  - Level of use
  - Commodity code: flammable, inflammable, corrosive, radio active, toxic, carcinogenic etc

Stock storage and handling

- Items may be stored in Four basic ways:
  - Shelves
  - Floor pallets
  - Block shaped pallets
  - Pallet racks
- Choice depends on
  - Quantity of products to be stored
  - Nature of the product and its package
  - Volume of each product
  - Internal height of store
  - Local availability of mechanical handling equipment

Housekeeping tasks

These include
- Cleaning and pest control
- Regular inspection system
- Disposal of stock (expired or damaged)
- Precautions against fire
- Strict security measures
- Regular stock taking and communicating the findings to higher levels.
Group Discussion
10 min

How do you dispose off expired, damaged, or deteriorated laboratory commodities?
(Refer to the guidelines by MOH)

Summary

• Know your Laboratory store space
• Know your reagents/supplies/chemicals - keep a list
• Effect the Zoning system
• Label the various zones/storage areas appropriately

What problems do you face at your facility with regard to laboratory commodity storage practices?
Objectives

By the end of this session, participants should be able to:

• Define the terms – receiving, issuing
• Describe the procedures for receiving and issuing laboratory commodities within a health facility
• Describe and be able to complete the appropriate forms and records used to receive and issue commodities for a health facility
• Enumerate good receiving and issuing practices

Definitions

• Receiving: Process of accepting requested commodities from a supplier or point of issue and making inventory entries

• Issuing: Process of supplying commodities from a main or bulk store to another store or to a point of use on demand

Both Receiving and Issuing require making necessary inventory entries
Inventory Management: Receiving

Inventory Management cycle

- Determining Quantities to Order
  - Requesting
  - Receiving
  - Storage
  - Issuing

Should be guided by Standard Operating Procedures (SOPs) AND Supported by good Record keeping

Receiving Commodities

- A health facility may receive commodities from:
  - The central stores (e.g. NPHLS, KEMSA)
  - Direct from suppliers
  - From other health facilities
  - Donors, i.e. as donations
### How commodities should be received

- They should always be received by an authorized person at the health facility.
- There should be clear written guidelines on the procedures for receiving commodities.
- The commodities received should be inspected.
- Any discrepancies or damages should be documented immediately and communicated to the supplier within the agreed time frame (within 48 hours).

### Checklist for Receiving commodities

1. Is the Delivery Note / Invoice for your facility?
2. Are the goods delivered the same as the ones that were ordered?
3. Are the quantities delivered the same as those in the Delivery note or Invoice?
4. Is the condition of the boxes at the time of delivery acceptable?
5. Are the goods delivered in good condition? (check liquids for leakages; check also for broken containers, unsealed, unusual odors)
6. Is the Expiry date of the commodities acceptable to your facility?
7. Document any discrepancies and follow up with supplier.
Receipt and inspection

Conduct a visual inspection of the commodities assessing the following:

– Quantity
– Quality (packaging, cleanliness, integrity of container)
– Expiry date (if available)
– Breakages
– Packing instructions (to ensure appropriate transportation conditions were maintained)
– Source

Good Receiving Practices (1)

• Only authorized persons should receive laboratory commodities
• Verify that the supply is for your facility/district
• Inspect goods for quality and quantity before receiving
• Check contents against your original order and supplier’s invoice/delivery note.
• Sign accompanying delivery documents if satisfied

NB: authorized officers are - the laboratory officer, Health Facility in-charge, Procurement officer or any other designated officer

Inventory Management:
Issuing
Good Receiving Practices (2)

- Enter received items in the laboratory stock card
- Store cold-chain dependent commodities immediately in the appropriate fridge or freezer according to manufacturer’s specifications
- Ensure your facility has clear, written SOPs for ordering/receiving laboratory commodities
- Always use a separate Laboratory stock control card for each inventory item

Inventory Management: Issuing

Inventory Management cycle

Determining Quantities to Order
- Requesting
- Receiving
- Storage
- Issuing

Should be guided by Standard Operating Procedures (SOPs) AND Supported by good Record keeping
How are commodities issued?

- From Central stores (e.g. NPHLS, KEMSA) to the health facilities
- **Inter**-facility (e.g. from one health facility to another)
- **Intra**-facility (e.g. from laboratory bulk store to the lab benches)

Inter-facility issuing

- Between sites in the same organization, e.g. from a MoH district hospital to a MoH health centre or vice versa
- Between sites in different organizations, e.g. from a government health facility to a faith-based facility (FBO)

Intra-facility issuing

Examples:-
- From the bulk store to laboratory
- From lab main store to lab benches
- Any other examples?
**Guidelines for issuing commodities**

- Ensure that the sites you are issuing to are authorized by the GOK
- Ensure proper records are used and maintained, e.g. S11 or other facility based order/issue forms
- Develop SOPs for issuing commodities
- During the issuing process, ensure the following:
  - Correctly fill the referenced information
  - Correctly determine the issue quantity
  - Fill in the issue form correctly and completely

**Tools used for Issuing of commodities**

Forms used for issuing:
- S11 (Counter Requisition and Issue voucher) – for **Intra**-facility issuing
- S12 (Issue and Receipt voucher) – for **Inter**-facility issuing

Records used for reference when issuing:
- S5 (Bin card) or Lab stock card

**Procedure for Issuing**

1) At the facility, fill in the following in the S11 or S12 clearly:-
   - Indicate point to which the issue is being made and the point of use
   - Include description of commodities, unit of issue and quantities to be issued
   - Depending on available quantities, determine the quantities to be issued compared to the quantities requested
   - Sign, date and stamp the form
2) Forward the appropriate copies to the relevant officers once the transaction is complete.
Take home messages (1)

- Ordering should be done on the right form which should be completely filled with all required information
- Ensure accurate quantification and timely submission of orders
- Prompt and accurate inspection of all shipments is essential to ensure that suppliers fulfill their contract terms

Take home messages (2)

- There should be a clear procedure for receiving laboratory commodities
- Confirmation of delivery is a vital part of this procedure
- Correct and timely documentation of the receipt of lab commodities is important
- Share documents with higher level

Group Discussion
(30 min)

Identify and list current challenges and their possible solutions in your facility/district in the following areas:
  - Receiving
  - Issuing
Records and Tools supporting Good Inventory Management practices

Module 3 Session 6

Objectives

By the end of this session, participants will be able to
• Identify records and tools used in support of good inventory management practices
• Demonstrate use of the tools to support good inventory management practices in their facilities
• Identify the problems/challenges encountered in using the tools

Tools for Inventory Management

1. Lab Stock card (Bin card)
2. Stock ledger card
3. S11
4. DAR (MoH 642)
5. F-CDRR (MoH 643)
6. Expiry tracking chart
7. Top-up form
8. Temperature monitoring charts
Stock records in inventory management

- Lab stock cards / Bin cards – These cards are kept with the physical stock. This makes a visual check easy and serves as a reminder to maintain records.

These are:
- The core records in the inventory management system
- Primary source of information used when calculating orders
- Source of data used to compile reports

Note: They may be manual or computerized records

Other Commonly used Inventory records (1)

- Lab Top-up forms – These cards are physically kept on the bench. This makes it easy to monitor Bench stock levels closely.

- Expiry tracking chart – Is kept in a strategic position in the commodity store. This assists in tracking expiry dates of identified commodities.

- Temperature monitoring chart – Is attached to the door of the store. Used in monitoring the temperatures of the storage areas.

Other Commonly used Inventory records (2)

Tracking and reporting commodity usage:

- DAR – This is kept on the bench to keep track of the daily usage of commodities

- F-CDRR - Kept in the office to be filled at the end of the month or any time one has to make an order
Instructions for Laboratory commodity management forms, cards and charts

- Provide description of Lab tools
- Accompanied by instructions for completing the tool, which highlight:
  » Description of the tool
  » Resources needed to fill it
  » Location / where kept
  » Who fills it
  » Steps taken when filling it

Laboratory stock card

- Description: Serially numbered two-sided card
- Keeps track of stock movements within the facility
  - A Stock card is the basic record needed to establish accurate quantities of individual items to re-order
  - The Lab commodity usage reports summarize information from the stock cards on a regular (usually monthly) basis.
- All stock cards and Lab commodity usage forms must be maintained in a clean, legible, and up-to-date manner.

Demonstration on filling the Lab Stock cards

- Examine the Laboratory Stock card
- Look at the Instruction sheet for completing the Lab Stock card
- Identify various sections/rows/columns of the tool
Practice Exercise on Lab Stock card (20 min)

Take your blank form and complete it using the information provided below:

- **District**: Nakuru
- **Facility**: Nakuru PGH
- **Serial No.**: 0082
- **Commodity**: Determine
- **Unit of issue**: tests
- **Item code**: 0012
- **Storage requirement**: room temperature
- **Average Monthly Consumption**: 100
- **Minimum Level**: 150
- **Maximum level**: 300

**Receipts**

- **Date**: 12/07/08
- **Received from**: KEMSA
- **Doc no**: 004
- **Quantity received**: 200
- **Batch no**: 0060
- **Expiry Date**: 12/09/09
- **Name**: Jackson Onyango

Let Us Check

```
MINISTRY OF HEALTH
Laboratory Stock Card

District: __________________________ Name: ________________________

Commodity name and Description | Unit of issue | Item Code | Storage requirements
---------------------------------|--------------|----------|---------------------
                                    |              |          | Average Monthly Consumption | Minimum Level | Maximum Level |
                                    |              |          | 100 | 150 | 300 |

RECEIPTS

Date | Received from | Doc No. | Quant | Batch No. | Expiry | Loc. | Name | Doc. No. | Quant | Dest. | Name | Balance | Unit Value |
---|----------------|--------|-------|-----------|--------|------|------|----------|-------|-------|------|---------|------------|
12/07/08 | KEMSA | 004 | 200 | 0060 | Lab store 1 | Jackson | 001 | 004 | 100 | 0060 | 150 | 300 | 15,000 |

Doc. No. = Document Number, Quant. = Quantity, Loc. = Location, Dest. = Destination
```

S11

(Counter Requisition and Issue voucher)

- This assists in tracking movement of commodities
- Used for Intra-facility issuing
- Contents:
  - Point to which the issue is being made (issue point) and the point of use
  - Description of commodities, unit of issue, quantities requested, quantities issued
  - Actioning staff
Expiry tracking chart

- This assists in tracking expiry dates of identified commodities
- Alerts the laboratory in-charge when stocks should be removed for exchange, re-distribution or destruction
- Kept in a strategic position in the commodities store
- Laminated, re-usable chart that accommodates information covering three consecutive years
Practice Exercise (10 min)

Expiry tracking chart

- Record the following data on a blank chart:
  - Commodity: Determine HIV Tests
  - Batch No: 2908765
  - Expiry date: 30/07/2013

In the Year column, fill in the following:
- Mark the month indicating SIX months BEFORE the Expiry date, i.e. 30/01/2013
- Mark the month indicating THREE months BEFORE the Expiry date, i.e. 30/04/2013

Laboratory Top-Up form

- Used to order supplies for use on the bench or section from the laboratory store / laboratory in-charge
- It allows for documentation of all issues from the lab store made to the bench or section in order to replenish or top-up lab supplies
- It helps the lab in-charge to see at a glance the usage of a particular commodity on the bench or section
- Location: kept in a file within the section or next to the bench
Demonstration how to Fill the Lab Top-Up Form

• Examine the Laboratory Top-up form
• Look at the Instruction sheet for completing the Lab Top-up form
• Identify various sections/rows/columns of the tool

Practice Exercise (10 min) for Lab Top-Up form
Take your blank Top-Up form and complete it using the information provided below:-

• Serial no.: 0040
• Facility: Rift Valley PGH
• Department/section: VCT
• Date: 12/10/2011
• Commodity: Determine
• Units of issue: Tests
• Current balance: 200
• Tests done: 400 tests
• Order quantity: 600
• Issue quantity: 300
• Name (Issued): Kamau
• Name (Received): Omondi
• Remarks: No Losses
### LET'S CHECK

**MINISTRY OF HEALTH**

**NATIONAL PUBLIC HEALTH LABORATORY SERVICES**

**LABORATORY TOP-UP FORM**

Name of Facility: Rift Valley PGH

Department/Section: VCT

<table>
<thead>
<tr>
<th>Date</th>
<th>Commodity</th>
<th>Unit of Issue</th>
<th>Current Balance</th>
<th>Tests done</th>
<th>Order</th>
<th>Issue</th>
<th>Quantity</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>12/10/2011</td>
<td>Determine</td>
<td>Tests</td>
<td>200</td>
<td>400</td>
<td>600</td>
<td>300</td>
<td>Kamau KA</td>
<td>Omondi OM</td>
</tr>
</tbody>
</table>

**+2°C to +8°C**

**Temperature Monitoring chart**

- Used for regular charting of the temperature of the refrigerator storing reagents/media/samples/blood units for the purpose of monitoring the maintenance of the cold chain.
- Location: should be physically located or pinned on the door of each 4°C fridge in the laboratory or section.
- Filled TWICE a day at 9.00 am and 4.00 pm
Demonstration On how to Fill the Temperature monitoring chart

- Look at the projected +2°C to +8°C Temperature monitoring chart
- Look at the Instruction sheet for completing the Temperature monitoring chart
- Identify various sections/rows/columns of the tool

Practice Exercise (10 min)
Take your blank form and complete it using the Information provided below:
- District: Nakuru
- Facility: Nakuru PGH
- Month: July
- Year: 2008
- Refrigerator: LG
- Operation: Electric

Faults and Problems:
- Date fault noted: 14/07/08
- Fault: Breakdown
- Action taken: Changed switch
- Date action taken: 14/07/08

Lets Check

[Image of Temperature monitoring chart]

Faults and Problems:
- Date fault noted: 14/07/08
- Fault: Breakdown
- Action taken: Changed switch
- Date action taken: 14/07/08
Summary: Key points about Inventory records

• All records must be kept current / up-to-date, legible, clean, complete and accurate
• It is impossible to manage the ordering/re-ordering process if stock movements cannot be tracked
MODULE 4: LABORATORY COMMODITY AND INFORMATION FLOW

Duration: 2 hours

Goal: To outline the national system for flow of laboratory commodities and related information

Objective

At the end of this module, participants should be able to:
• Define a Logistics Management Information System, its purpose and importance
• Describe the laboratory commodity logistics and information flow pipelines and accompanying records/documentation

Content

Session 1: Introduction to Logistics Management Information System (LMIS)
Definition of terms; basic concepts of LMIS; purpose of LMIS; importance of LMIS in the laboratory Commodity management cycle

Session 2: Laboratory Commodity and Information Flow
Commodity flow and Commodity Information flow; Current laboratory commodity and information flow pipelines; Rationale of tracking and reporting lab commodities; necessary data collection and reporting tools (DAR, CDRR)

Lesson Plan Guide:

<table>
<thead>
<tr>
<th>UNIT</th>
<th>CONTENT</th>
<th>ACTIVITY</th>
<th>TIME</th>
</tr>
</thead>
</table>
| 1    | Introduction to Logistics Management Information System (LMIS) | • Lecture  
• Group activity | 1 hour |
| 2    | Laboratory Commodity and Information Flow - Tracking use and reporting on laboratory commodities | • Lecture  
• Group activity and report back | 1 hour |

References and Recommended Readings


Introduction to Logistics Management Information System (LMIS) and Laboratory Commodity and Information flow

Module 4

Introduction to Logistics Management Information System (LMIS)

Module 4 Session 1

Session Objectives

By the end of this session, participants should be able to

• Define Logistics Management Information System (LMIS)
• Describe the purpose of LMIS
• Highlight the importance of LMIS in the laboratory Commodity management cycle
Definition of LMIS

- A Manual or computerized system that comprises:
  - Recording logistics data
  - Processing the data into information
  - Information interpretation and presentation for decision-making

LMIS Design Considerations

- Data sources and reporting tools
- Reporting frequency
- Reporting deadline
- Reporting officer
- Receiving office/officer
- Number of levels in the Logistics system
- Mode of transmitting the reports
- Feedback content and flow
Importance of LMIS

- LMIS helps answer the following questions:
  - Which commodities are required?
  - What kind and in what quantities?
  - Where in the pipeline (level) are they required?
  - When are they required?
  - How will they get there?

The LMIS indicates
- The rate of consumption
- Whether more stock is required and when to place orders for re-supply
- Location of the commodities in the pipeline
- Where there are losses in the pipeline needing action
- Where there are expired and short-expiry commodities needing action
- Where there is need for stock adjustments

Essential Data elements

- **Stock on Hand (End of Month Physical count):**
  Quantities of stock available at the end of the review period
- **Consumption (Quantity used)**
  Quantity consumed in testing/analysis
- **Losses & Adjustments**
  a) **Losses** – Quantity of stock removed from the pipeline for any reason other than consumption, e.g. expiries
  b) **Adjustments** – Quantity received or given to other facilities or difference between the physical count and what is recorded in the stock control cards: positive (+) or negative (-)
LMIS: Other Data Elements collected

- **Beginning Balance**: Quantity of stock on hand at the beginning of the review period
- **Quantity Received**: Quantity received during the review period
- **Quantity expiring in less than 6 months**: Quantity of stock with short expiry
- **Days out of stock**: Number of days that the whole facility was stocked out of the item
- **Quantity Requested for Re-supply**: Quantity calculated and ordered needed for re-supply

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**Role of LMIS in Commodity Management Cycle**

![Role of LMIS in Commodity Management Cycle](image)

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**Feedback**

Important because:

- Provides information from the higher to the lower levels of the system
- Inform the lower levels about the performance of their LMIS
- LMIS informs managers at higher levels about how the lower level facilities are functioning
- May help solve problems by highlighting errors seen on facility reports and how to correct them
- It may inform facilities about how other facilities are reporting.
Brainstorm
5 min

• What will make a LMIS not function properly?

What will make a LMIS not function properly?
• No reports!
• Late reports
• Inaccurate reports
  – data collection errors; incorrect filling of the report; incorrect calculations
• Incomplete data reporting
  – not all data elements are reported, e.g. missing facility details
• Lack of required reporting tools in the pipeline
• Use of Old reporting formats / tools

Group Exercise
(10 min)

• In your respective groups, please discuss and write down the problem areas experienced with LMIS in your facilities related to:
  – Tracking commodity use
  – Reporting commodity consumption
### Challenges in Laboratory Commodities

**LMIS (1)**

- Difficult to track and measure laboratory commodities
  - Many laboratory commodities are liquids or powders that are difficult to count
  - Same commodity may be used for a variety of different tests and by a number of different people in a single laboratory, thereby making actual consumption of the commodity difficult to quantify
  - Distinguishing the use of commodities for QC from the use of commodities for testing is difficult and time-consuming.

**LMIS (2)**

- Because of the short shelf life of reconstituted reagents, they may be discarded before being completely consumed
- A certain amount of wastage should be expected in laboratory services. This wastage differs from loss caused by damage, expiry, or theft.
- Loss should be tracked in an LMIS, while it is difficult to separate wastage from consumption.

### Recommendations for LMIS (1)

- Consider two ways of capturing consumption:
  - Issues from stock as proxy consumption, or
  - Tracking actual use through Daily registers, e.g. DAR.
- Review each product, including how it is issued and used, to determine the appropriate unit for stock keeping and reporting.
- Use and maintain stock-keeping records; consider the advantages and disadvantages of having one or multiple stock cards for the same commodity.
Recommendations for LMIS (2)

- Routinely report stock levels, issues, losses, adjustments and stock outs.
- Link reporting with re-supply
- Computerize the LMIS where possible.

LMIS as a Monitoring, Evaluation & Supervision Tool

**Indicators Monitored:**

- Accuracy and completeness of reporting – **Data Quality**
- Stock-out rates at any point during a given period
- Stock levels between Maximum and Minimum
- Rate of loss of commodity by reason (Expiration, damage, pilferage, etc)
- Frequency of reporting and list of non-reporting facilities
- Rate of consumption in a given period

Conclusion – Benefits of LMIS

- Accountability for all products in the supply chain
- A reduction in supply imbalances (stock outs, overstocks) at the facilities and stores.
- Efficient, cost-effective supply chain management
  - Improved quality of care
  - Improved Impact
  - Informed decisions on supply chain management
Laboratory Commodity and Information Flow

Module 4 Session 1

Session Objectives

By the end of this session participants should be able to:

• Describe Commodity flow and Commodity Information flow
• Describe the current laboratory commodity and information flow pipelines
• Understand rationale of tracking and reporting on laboratory commodities
• Identify the necessary data collection and reporting tools and fill in the same (DAR, CDRR)

Commodity and Information flow

• A Commodity pipeline is the entire chain of facilities through which commodities move from the manufacturer / supplier to get down to the user / patient / client
• Commodity flow describes, in form of a diagram, the flow of commodities along the pipeline
• Commodity information flow describes, in form of a diagram, the flow of information concerning the commodities along the pipeline
• Documentation is needed for each transaction in order to track the item from ordering/purchase, delivery/receipt, distribution/issuing, to actual use.
Group Exercise  
(10 min)

• In your groups, draw a flowchart that depicts the flow of commodities within the health facility from the initial laboratory request to the facility store to when commodities are availed at the user points.

• For each step, indicate the documentation (records, tools) required.

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*SDPs include Provincial General Hospitals (PGHs), Regional Blood Transfusion Centres (RBTCs), the 2 Referral hospitals, District hospitals, Health centres, Dispensaries, etc*
Key notes – Commodity flow
• Central (and regional) stores e.g. KEMSA supply commodities directly to SDPs
• Copy of items supplied is provided to DMLTs
• SDPs should submit a copy of the delivery note to the DMLT once commodities are delivered.
• Within SDPs / Facilities, stock is issued from the store to the Lab user points
• Inventory management tools, e.g. Lab stock cards, are used to track inventory
• Test registers and LMIS tools, e.g. DAR, used to record tests done and consumption on daily basis

Key notes – Commodity information flow
• SDPs complete consumption reports (F-CDRR)
• Required reports are prepared on monthly basis to summarize consumption and guide requisition for resupply
• SDPs channel their F-CDRRs through the DMLT for further transmission to the central level
• Data from SDPs aggregated at central level
• Re-supply decisions made and orders sent to KEMSA
• Feedback between DMLTs, PMLTs, central level and the SDPs

Tracking and reporting use of Laboratory Commodities
• Learn how to complete Laboratory Commodities Daily Activity Register (DAR)

• Learn how to complete usage reporting tool for Laboratory Commodities, i.e. the Facility Consumption Data Report & Request (F-CDRR)
## Introduction (1)

- Data on stock, transactions and consumption should be recorded promptly in the relevant recording tool.
- Records must be carefully controlled but accessible to responsible officers through:
  - Storing them in locked and secured record storage areas.
  - Limited access areas.
  - Use of computer/computer programs with restricted access.

## Introduction (2)

- Inventory records at all levels must:
  - Be accurately kept.
  - Reflect at all times the quantities on hand and the quantities disbursed.
  - Meet the needs of external and internal audits.
- For quality control purposes, the required information is documented in the LMIS.

## Role of LMIS in tracking commodities (1)

LMIS enables personnel:
- Make routine decisions that affect commodity availability.
- Determine how much of each commodity to order or resupply.
- Forecast future demand for commodities and plan procurement and shipments.
- Identify potential supply problems and handle other issues related to commodity management.
Role of LMIS in tracking commodities (2)

- Collect the three essential data items needed to make logistics decisions: **stock on hand, consumption, losses and adjustments**
- Those data are recorded on stock-keeping records, transaction records, and consumption records.
- The data are then used at the facility and are reported to higher levels for re-supply and management purposes.

Why Report Lab commodity Consumption?

- To support Planning at the National level, e.g. through Forecasting and quantification
- To ensure continuous supply of Lab commodities and supplies
- For accountability and commodity tracking
- For monitoring and evaluation
Records / Forms / Tools used

- Stock-keeping and Transaction Tools
  - S11, S12, S13
  - Lab Top-up form
  - Lab Stock card / Bin cards
  - Expiry Tracking chart
  - Temperature Monitoring chart

- LMIS Consumption Data collection tools
  - Daily Activity Register (DAR) for Laboratory commodities (MoH 642)

- LMIS Consumption Reporting tools
  - Facility Consumption Data Report & Request Form (F-CDRR) for Laboratory commodities (MoH 643)
  - Facility Consumption Data report and request (F-CDRR) form for ART Laboratory monitoring reagents

Refer to Appendices
Laboratory LMIS Reporting tools and Deadlines

- **Reporting Tool:** Facility Consumption Report & Request (F-CDRR)
- **Who reports:** Facility Lab in-charge
- **Who do you report to:** Facility -> District -> LMU
- **By when do you report:**
  - SDPs report monthly by 5th to the DMLT. DMLT sends all SDP reports for his district by latest 10th to the LMU.
- **How do you transmit:** Electronic (aggregation tool), courier or post

**Laboratory commodity information flow**

- **NPHLS and Programs**
  - Information on Distribution, Stocks, Procurement, re-supply orders
- **Central level Data units (CDU, LMU)**
  - DMLT collects all F-CDRRs for district
  - Reporting deadline: 10th of the month – to the Central level
- **DMLT**
  - Monthly report (F-CDRR)
  - Reporting deadline: 5th of the month – to the District
- **SDP**
  - Feed back on reporting rates, data quality, etc.

**Exercise: Problem identification (10 min)**

- Identify the problems encountered in tracking consumption and reporting the use of laboratory commodities in your facility
Demonstration of Lab LMIS tools  
(15 min)

<table>
<thead>
<tr>
<th>Lab LMIS tools</th>
</tr>
</thead>
<tbody>
<tr>
<td>• DAR (MoH 642)</td>
</tr>
<tr>
<td>• F-CDRR (MoH 643)</td>
</tr>
<tr>
<td>• F-CDRR for ART Lab monitoring reagents</td>
</tr>
</tbody>
</table>

Group Exercise  
(30 min)

Using the sample data on the records and registers availed

| • complete the register with the transactions |
| • complete the summary report tool provided |

Exercise: Tracking commodity usage  
(data collection)

<table>
<thead>
<tr>
<th>Tools in use:</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Daily Activity Register (DAR) for Lab commodities (MoH 642)</td>
</tr>
</tbody>
</table>

**Exercise 1**

| • Practicum on completing the DAR form |

NB: *Avail exercise and tool*
Exercise: Reporting Commodity Consumption

Tools in use:
• F-CDRR (MoH 643)
• F-CDRR for ART Lab monitoring reagents

Exercise 2
• Practicum on how to complete the Facility CDDR form

Avail exercise and the tool
Exercises on Laboratory commodity logistics reporting tools

Exercise 1: Filling in the Daily Activity Register (DAR)

Materials needed:

For this exercise you will need:

- blank copies of the DAR for Laboratory Reagents and Consumables (MoH 642) – provided below
- a copy of the job aid for the completion of the DAR
- Pen and paper for rough-work or a calculator.

Assignment

Use the following to fill in the DAR:-

You are John Mali, working on the Serology, Microbiology and Haematology bench at Serowe District Hospital (DH). You are working on the day shift for the entire month of March 20XX, and there is no night shift.

You started the month with quantities indicated in the table below. On 1st March, using the Lab Top-up form, you requested and received the following commodities from the Lab store:-

<table>
<thead>
<tr>
<th>Commodity name</th>
<th>Batch No.</th>
<th>Expiry date</th>
<th>Beginning balance</th>
<th>Quantity received</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rapid Test kit – HIV screening</td>
<td>0060</td>
<td>Dec 20XX</td>
<td>200</td>
<td>500 tests</td>
</tr>
<tr>
<td>Carbol Fuschin solution</td>
<td>0020</td>
<td>Sept 20XX</td>
<td>0</td>
<td>500ml</td>
</tr>
<tr>
<td>Malaria Rapid diagnostic test (RDT)</td>
<td>A0254</td>
<td>Dec 20XX</td>
<td>0</td>
<td>40 tests</td>
</tr>
<tr>
<td>FACS Count CD-4CD3 reagent (Adult)</td>
<td>00125</td>
<td>June 20XX</td>
<td>0</td>
<td>50 tests</td>
</tr>
</tbody>
</table>

During the month, you conducted the following transactions:

March 2nd
- 20 tests for HIV

March 5th
- 30 tests for HIV
- 15 TB AFB microscopy smears (Note: each smear uses 3ml of Carbol fuschin solution)

March 18th
- 50 tests for HIV
- 15 malaria tests using the RDT, including 1 repeat test. Of these, 10 were for patients < 5years and 4 of these turned positive, while 4 were for patients aged 5 years and above, and all were negative.
- 15 TB AFB microscopy smears (note: each smear uses 3ml of Carbol fuschin solution)

March 29th
- 30 CD4 tests plus 2 controls
- 100 tests for HIV

During the month, you also made the following adjustments:

- On 30th March, you noticed 100 Rapid HIV tests had been damaged by rain water
- On 29th March, you gave 30 Rapid HIV tests to the MCH clinic for HIV testing under PMTCT
- Assume you had no stock-out days

Instructions

Please complete the DAR pages provided below using the information provided above.
## EFFECTIVE MANAGEMENT OF LABORATORY COMMODITIES

### PARTICIPANTS GUIDE

**Filling DAR (blank)**

<table>
<thead>
<tr>
<th>Date</th>
<th>Shift</th>
<th>Unit of Issue</th>
<th>Beginning Balance at the End of the Shift</th>
<th>Quantity Received during the Shift</th>
<th>Origin of the received stock</th>
<th>Batch Number</th>
<th>Expiry Date</th>
<th>Quantity consumed during the Shift</th>
<th>Number of Tests done during the Shift</th>
<th>Leased during the Shift</th>
<th>Adjustment during the Shift [Indicate Yes/No]</th>
<th>Ending Balance at the End of the Shift</th>
<th>Remarks [Including explanation for Leased or Adjustment]</th>
<th>Name of Officer</th>
<th>Signature</th>
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</tbody>
</table>

**Page Totals:**
- Balance carried forward: 
- Days out of stock: 

**Blank page of DAR 2:**

<table>
<thead>
<tr>
<th>Date</th>
<th>Shift</th>
<th>Unit of Issue</th>
<th>Beginning Balance at the End of the Shift</th>
<th>Quantity Received during the Shift</th>
<th>Origin of the received stock</th>
<th>Batch Number</th>
<th>Expiry Date</th>
<th>Quantity consumed during the Shift</th>
<th>Number of Tests done during the Shift</th>
<th>Leased during the Shift</th>
<th>Adjustment during the Shift [Indicate Yes/No]</th>
<th>Ending Balance at the End of the Shift</th>
<th>Remarks [Including explanation for Leased or Adjustment]</th>
<th>Name of Officer</th>
<th>Signature</th>
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</tbody>
</table>

**Page Totals:**
- Balance carried forward: 
- Days out of stock: 

**Blank page of DAR 3:**

<table>
<thead>
<tr>
<th>Date</th>
<th>Shift</th>
<th>Unit of Issue</th>
<th>Beginning Balance at the End of the Shift</th>
<th>Quantity Received during the Shift</th>
<th>Origin of the received stock</th>
<th>Batch Number</th>
<th>Expiry Date</th>
<th>Quantity consumed during the Shift</th>
<th>Number of Tests done during the Shift</th>
<th>Leased during the Shift</th>
<th>Adjustment during the Shift [Indicate Yes/No]</th>
<th>Ending Balance at the End of the Shift</th>
<th>Remarks [Including explanation for Leased or Adjustment]</th>
<th>Name of Officer</th>
<th>Signature</th>
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</tbody>
</table>

**Page Totals:**
- Balance carried forward: 
- Days out of stock: 

**Blank page of DAR 4:**

<table>
<thead>
<tr>
<th>Date</th>
<th>Shift</th>
<th>Unit of Issue</th>
<th>Beginning Balance at the End of the Shift</th>
<th>Quantity Received during the Shift</th>
<th>Origin of the received stock</th>
<th>Batch Number</th>
<th>Expiry Date</th>
<th>Quantity consumed during the Shift</th>
<th>Number of Tests done during the Shift</th>
<th>Leased during the Shift</th>
<th>Adjustment during the Shift [Indicate Yes/No]</th>
<th>Ending Balance at the End of the Shift</th>
<th>Remarks [Including explanation for Leased or Adjustment]</th>
<th>Name of Officer</th>
<th>Signature</th>
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</tbody>
</table>

**Page Totals:**
- Balance carried forward: 
- Days out of stock: 

(Note: End of DAR quantity from last row on the page.)
Exercise 2: Filling in the Facility Consumption Data Report and Request Form (FCDRR)

Materials needed:

For this exercise you will need:
- blank copy of the F-CDRR for ART Laboratory monitoring reagents
- blank copy of the F-CDRR for Laboratory commodities (MoH 643)
- filled DAR for Laboratory Reagents and Consumables (MoH 642) – from Exercise 1 above
- a copy of the job aids for the completion of the F-CDRRs
- Pen and paper for rough-work or a calculator.

On 1st April, the Lab Tech in/charge, Felix Mambo, and the hospital store person sat down together to compile the monthly report for the month of March 20XX for Serowe DH, which is located in Central district of Northern province.

They conducted a physical count and found the following closing stock balances at the District hospital store, and then compared with the Laboratory stock cards for the various commodities. They then compiled the following information:

Stock count at the beginning and end of the month of March 20XX:

<table>
<thead>
<tr>
<th>Item</th>
<th>Rapid Test kit – HIV screening</th>
<th>Carbol Fuschin solution</th>
<th>Malaria Rapid diagnostic test (RDT) kit</th>
<th>FACS Count CD4CD3 reagent (Adult)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stock Balance (as at 29th February)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>At the hospital store</td>
<td>150</td>
<td>0</td>
<td>120</td>
<td>50</td>
</tr>
<tr>
<td>Received during the month</td>
<td>600 Batch/Lot No. 0065</td>
<td>1000 ml Batch/Lot No. 0020</td>
<td>0</td>
<td>50 Batch/Lot No. 00126</td>
</tr>
<tr>
<td>Date of receipt: 1st March 20XX</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Expired during the month</td>
<td>0</td>
<td>0</td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td>Stock Balance (as at 31st March)</td>
<td>250</td>
<td>300</td>
<td>70</td>
<td>100</td>
</tr>
</tbody>
</table>

The stock receipts from KEMSA and the District store during the month are noted in the table above.

Adjustments: During the month, the Lab Tech in/charge issued to Kutus Health centre 200ml of Carbol Fuschin solution. In the same month, the DH received 50 tests of FACS Count CD4CD3 reagent (Adult) from Faidha PGH.

The PMTCT clinic, being the other user point in the DH, provided the following report for the month:-

<table>
<thead>
<tr>
<th>Name of commodity</th>
<th>Beginning balance</th>
<th>Adjustments</th>
<th>Quantity used</th>
<th>Number of Tests done</th>
<th>Balance carried forward</th>
<th>Days out of stock</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rapid Test kit – HIV screening (tests)</td>
<td>10</td>
<td>+30 (from Serology bench)</td>
<td>10</td>
<td>10</td>
<td>30</td>
<td>0</td>
</tr>
</tbody>
</table>

The PMTCT clinic had no losses and adjustments. The Lab Tech in-charge also noted that the F-CDRR for ART Laboratory monitoring reagents was nearly finished so another copy of the tool should be ordered.

Using the data provided above and the completed DAR from No. 1, you are required to complete the following reports on behalf of Felix Mambo:
- F-CDRR for ART Laboratory monitoring reagents, and
- F-CDRR for Laboratory commodities (MoH 643) for Serowe DH for the month ending 31st March 20XX.
Filling in the F-CDRR for ART Laboratory monitoring reagents (Blank)

**FACILITY CONSUMPTION DATA REPORT & REQUEST (F-CDRR) FOR ART LABORATORY MONITORING REAGENTS**

<table>
<thead>
<tr>
<th>COMMODITY CODE</th>
<th>COMMODITY NAME</th>
<th>COMMODITY GROUP</th>
<th>QUANTITY RECEIVED FROM CENTRAL WAREHOUSE (e.g. KEMSA)</th>
<th>QUANTITY REQUESTED</th>
<th>QUANTITY USED</th>
<th>QUANTITY REQUESTED</th>
<th>QUANTITY RECEIVED FROM OTHER SOURCES</th>
<th>QUANTITY REQUESTED</th>
<th>QUANTITY USED</th>
<th>QUANTITY REQUESTED</th>
<th>QUANTITY RECEIVED</th>
<th>QUANTITY REQUESTED</th>
<th>QUANTITY USED</th>
<th>QUANTITY REQUESTED</th>
</tr>
</thead>
<tbody>
<tr>
<td>FACS 001</td>
<td>Sheath fluid</td>
<td>FACS Calibur</td>
<td></td>
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<tr>
<td>FACS 002</td>
<td>Cleaning fluid</td>
<td>FACS Calibur</td>
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<td>FACS 003</td>
<td>Sheath fluid</td>
<td>FACS Calibur</td>
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**Hematology Analyser**

<table>
<thead>
<tr>
<th>COMMODITY CODE</th>
<th>COMMODITY NAME</th>
<th>COMMODITY GROUP</th>
<th>QUANTITY RECEIVED FROM CENTRAL WAREHOUSE (e.g. KEMSA)</th>
<th>QUANTITY REQUESTED</th>
<th>QUANTITY USED</th>
<th>QUANTITY REQUESTED</th>
<th>QUANTITY RECEIVED FROM OTHER SOURCES</th>
<th>QUANTITY REQUESTED</th>
<th>QUANTITY USED</th>
<th>QUANTITY REQUESTED</th>
<th>QUANTITY RECEIVED</th>
<th>QUANTITY REQUESTED</th>
<th>QUANTITY USED</th>
<th>QUANTITY REQUESTED</th>
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<tr>
<td>BAE 001</td>
<td>Sodium</td>
<td>Biochemistry</td>
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<td>Chloride</td>
<td>Biochemistry</td>
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<td>BAE 003</td>
<td>Urea</td>
<td>Biochemistry</td>
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</table>

**Comments**

Order for extra LMIS tools:

1. Daily Activity Register for Laboratory Regents and Consumables (MOH 642):
   - To be requested only when your Data collection or reporting Tools are nearly full. Indicate quantity required for each tool type.
2. F-CDRR for ART Lab Monitoring Reagents:
   - All tools required to be filled in blank.

Date Filled: ____________________________

Approved by: ____________________________

**Note:**

- Data submitted in this form will be processed in NHLSL / NASCOP MINISTRY OF HEALTH FACILITY CONSUMPTION DATA REPORT & REQUEST (F-CDRR) FOR ART LABORATORY MONITORING REAGENTS

- Filling in the F-CDRR for ART Laboratory monitoring reagents (Blank)
**Filling F-CDRR for Laboratory Commodities (MoH 643) (Blank):**

**MINISTRY OF HEALTH**

**FACILITY CONSUMPTION DATA REPORT AND REQUEST (F-CDRR) FOR LABORATORY COMMODITIES**

<table>
<thead>
<tr>
<th>Commodity Name</th>
<th>Unit of Issue (e.g. Test)</th>
<th>Beginning Balance</th>
<th>Quantity Received</th>
<th>Quantity Used</th>
<th>Number of Tests done</th>
<th>Losses</th>
<th>End of Month Physical count (e.g. Test)</th>
<th>Quantity Requested for Re-supply</th>
<th>Quantity expiring in less than 6 months</th>
<th>Days out of stock</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
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</tbody>
</table>

**HAEMATOLOGY COMMODITIES**
- Haematology reagents
- Normal control
- Abnormal high control
- Abnormal low control
- Grouping anti-sera - Anti-A
- Grouping anti-sera - Anti-B
- Grouping anti-sera - Anti-D
- Grouping anti-sera - Anti Human Globulin (AHG)
- Haemocue cuvettes
- Leishman stain

**BIOCHEMISTRY COMMODITIES**
- Albumin
- Alkaline phosphatase
- ALT (SGPT)
- AST (SGOT)
- Creatinine
- Electrolytes
- Gamma GT
- Glucose test strips
- HDL Cholesterol
- Multistix
- Pregnancy test strips
- Serum Amylase test kit
- Serum protein kit
- Total Cholesterol
- Triglycerides
- Urea

**CONSUMABLES**
- Applicator sticks
- Blue tips
- EDTA Vacutainer Stabilizer tubes
- EDTA Vacutainer tubes
- Latex Gloves
- Microscope slides
- Plain Vacutainer tubes
- Stool Polypots
- Vacutainer needles G21
- Vacutainer needles G23
- Yellow tips

**MICROBIOLOGY COMMODITIES**
- Macconkey agar
- CLED agar
- Methanol
- Glycerol
- Oil Immersion
- Universal bottles
- Culture Plates

**MALARIA-RELATED LABORATORY COMMODITIES**
- Malaria Rapid Diagnostic Test (RDT)
- Field Stain A
- Field Stain B
- Giemsa Stain

**TB-RELATED LABORATORY COMMODITIES**
- Auramine-O (*for District hospital labs)
- Carbol Fuschin (solution)
- Falcon tubes
- Hydrochloric acid (HCL)
- Lens Tissue
- Methylene Blue
- Microscope slides
- Potassium Permanganate (*for District hospital labs)
- Sputum mugs (AFB Polypots with lids)
- Sulphuric acid

**HIV-RELATED LABORATORY COMMODITIES**
- Rapid HIV 1+2 Test 1 - Screening
- Rapid HIV 1+2 Test 2 - Confirmatory
- Rapid HIV 1+2 Test 3 - Tiebreaker
- Hepatitis B (ELISA) Test (Hepanostika)
- Hepatitis C (ELISA) Test (Murex HCV)
- ELISA HIV & AIDS 1+2 Test (Vironostika)
- ELISA HIV & AIDS 1+2 Test (Murex HIV)
- Rapid Syphilis Test (RPR)

**Table of Services**
- Type of service
  - STAT
  - DENT
  - PHCU

**Adjustments**
- Indicate if (+) or (-)

**Explanation Losses & Adjustments**

**Report for Period**
- Beginning: Ending:
  - dd/mm/yyyy               dd/mm/yyyy

**Explain Losses & Adjustments:**

**Completed by:**
- Name: Tel: Designation: Sign: Date:

**Approved by:**
- Name: Tel: Designation: Sign: Date:

**Table for extra LMIS tools:**
- To be requested only when your Data collection or reporting Tools are nearly full. Indicate quantity required for each tool type.

**Order for extra LMIS tools:**

**English Notes & Adjustments:**

**MOH 643 (Blank):**

**Name of Facility:**
- Facility Code:

**District:**
- Province / County:

**Affiliation:**
- Ministry of Health
- Local Authority FBO
- NGO
- Private

**Report for Period:**
- Beginning: Ending:
  - dd/mm/yyyy               dd/mm/yyyy

**Test Category**

**FILLING F-CDRR FOR LABORATORY COMMODITIES (MOH 643) (Blank):**

**Name of Facility:**
- Facility Code:

**District:**
- Province / County:

**Affiliation:**
- Ministry of Health
- Local Authority FBO
- NGO
- Private

**Report for Period:**
- Beginning: Ending:
  - dd/mm/yyyy               dd/mm/yyyy
References and recommended readings

MODULE 5: MONITORING, TRAINING AND PLANNING (MTP) FOR LABORATORY COMMODITIES

Duration: 3 hours 15 minutes

Goal: To describe the concept of continuous performance and quality improvement for laboratory commodity management

Objectives
- Define Monitoring, Training and Planning (MTP) approach
- Differentiate between problems and challenges
- Identify priority challenges on commodity management practices at the facility
- Convert problems to challenges
- Set SMART targets for their priority challenges
- Develop a plan of action for the challenges identified throughout the course

Content:

Session 1: Introduction to MTP: Definition of terms; introduction to MTP and the MTP approach; determining key problems / challenges in laboratory commodity management at facility level; prioritizing problems; and converting problems to challenges

Session 2: Action planning: Developing Action plans for the challenges identified in laboratory commodity management at facility level; setting SMART targets for the priority challenges; identifying activities with time lines to address the challenges and discuss benchmarks for monitoring the implementation process.

Monitoring, Training and Planning (MTP) for laboratory commodities

Lesson Plan Guide:

<table>
<thead>
<tr>
<th>UNIT</th>
<th>CONTENT</th>
<th>ACTIVITY</th>
<th>TIME</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Introduction to MTP</td>
<td>• Lecture&lt;br&gt;• Group activity</td>
<td>45 minutes</td>
</tr>
<tr>
<td>2</td>
<td>Action planning</td>
<td>• Lecture&lt;br&gt;• Practice of action plans&lt;br&gt;• Presentation of action plans</td>
<td>1 hour 45 minutes 45 minutes</td>
</tr>
</tbody>
</table>

References and Recommended Readings


Applying the Monitoring Training and Planning (MTP) approach to Laboratory commodity management

Module 5

Objectives

• Define Monitoring, Training and Planning (MTP) approach
• Differentiate between problems and challenges
• Identify priority challenges on commodity management practices at the facility
• Convert problems to challenges
• Set SMART targets for the priority challenges
• Develop a plan of action for the challenges identified throughout the course

What is MTP?

Monitoring – Training – Planning
What is the MTP approach?

Description:

- **MTP (Monitoring, Training, Planning)** is a performance improvement methodology
- May be used at facility or district level to improve laboratory services

### Monitoring

**Problem Identification**

- First Visit
  - Discuss the gaps identified during the visit
  - Identify specific Problems
  - Measure the extent of the problem(s)
  - Prioritize problems

**Subsequent visits**
- Follow up on commitments from the previous visit
- Evaluate the result of the intervention
- Document and communicate outcomes

### Prioritizing Problems:

**Example of Criteria**

- **Priority A** - Those which if **not** corrected would cause a program or service to collapse (**Immediate action required**) e.g. Stock-out of lab commodities

- **Priority B** - Those which if corrected would improve service delivery (**Action required in the short term**) e.g. increased staffing levels

- **Priority C** - Those which if **not** corrected would hinder improvement of service delivery (**Action required in the long term**) e.g. infrastructural improvements
Training

• Training / Problem solving

Discuss the problem:-
1. Why is it happening? What are the underlying factors?
   - Root Cause Analysis
2. Decide how to solve the problem.
   Find a solution, i.e. the Intervention

Root cause analysis

• It is important to differentiate between the cause of a problem and the symptom of a problem
• Treating the symptoms only leads to short-term solutions
• Asking the question “Why?” several times helps to identify the “real” reason or cause of a problem
• A single problem can have more than one root cause

Root cause analysis: Example 1

Problem: No Malaria RDTs in stock
Why?

Did not request in Time
Why?

Do not have the Commodity report & request forms
Why?

Do not know where to get report & request forms
Why?
Root cause analysis: Example 2

Problem: No HIV test kits in Stock

Why?

Did not know when they ran out

Why?

Do not record how much we have on a daily basis

Why?

Did not know this was important

Why?

Root cause analysis: Example 3

Gap: Stock-out of HIV test kits

Why?

Received less than required quantity

Why?

Did not know how much to order

Why?

Lack of quantification skills

Why?

???

Why?

Examples of Problems converted into Challenges (1)

Problem

- Our district often runs out of stock for HIV rapid test kits and does not meet the needs of all the testing facilities

- The manual lab tools (DARs, Lab Stock Card, F-CDRRs, etc) have not been availed in adequate quantities in our district

Challenge

- How can we ensure that the district always has sufficient rapid test kits to issue to all the testing areas?

- How can we obtain additional copies of the manual tools (DARs, Lab Stock cards, CDRRs) in adequate quantities for all facilities in the district?
Examples of Problems converted into Challenges (2)

<table>
<thead>
<tr>
<th>Problem</th>
<th>Challenge</th>
</tr>
</thead>
<tbody>
<tr>
<td>o Our facility staff do not know how to use the new inventory management tools</td>
<td>o How can we make sure that all facility staff have the knowledge and skills required to use the new manual inventory management tools correctly?</td>
</tr>
<tr>
<td>o Our facility store has limited secure space for storage of Lab commodities and supplies</td>
<td>o How can we obtain additional storage space for the Lab commodities and supplies?</td>
</tr>
</tbody>
</table>

Group activity (30 minutes)

• In your groups, convert your identified problems into challenges

Finding Solutions

• The solution should address the root cause
• Find local solutions:- think of
  – “What can I do to solve the problem”
  – Consider solutions that do not require external assistance (e.g. authorities, stakeholders, government) as the first option
• Be creative
• If there are no local solutions, then the facility in-charge or the lab in-charge becomes the liaison to outside authorities
  – Prepare information on the extent of the problem and the difficulties in solving the problem locally
  – Take the problem to regional/national authorities
  – Assist the staff to write a letter/proposal/follow-up
Planning

- Set a target for improvement
- Set measures to achieve the target
- Assign someone the responsibility for execution of the plan
- Set the date for the next progress review

Action Plans (1)

- Once the solution is agreed on, then a plan to implement the solution is needed

- An Action Plan answers
  - What will be done
  - Who will do what
  - When will they do it
  - Where will they do it
  - How will they do it

- Action Plans should also include Targets and means of monitoring progress towards targets

Action Plans (2): Class Discussion

- Review the action plan below and think about what might be problematic about it

<table>
<thead>
<tr>
<th>No.</th>
<th>Problem</th>
<th>Solution</th>
<th>By Whom</th>
<th>By When</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Frequent Stock outs</td>
<td>Order from KEMSA</td>
<td>In-Charge</td>
<td>Immediately</td>
</tr>
<tr>
<td>2</td>
<td>Inadequate space for storage</td>
<td>Request for Donor Funding</td>
<td>In-Charge</td>
<td>Two weeks</td>
</tr>
<tr>
<td>3</td>
<td>Inadequate staff</td>
<td>Hire more Staff</td>
<td>Government</td>
<td>ASAP</td>
</tr>
</tbody>
</table>
Benefits of applying the MTP concept

- Staff are oriented on **quality improvement** by using the MTP approach
- MTP assists staff in
  - Identifying **root causes**
  - Identifying **creative solutions**
  - Developing **practical Action plans**
  - **Reviewing progress** towards targets
- MTP allows for **follow-up support**

Summary of MTP
**Monitoring – Training – Planning**

Activity: Action plan session (25 min)

<table>
<thead>
<tr>
<th>Objectives</th>
<th>Activities</th>
<th>Responsible person</th>
<th>Resources</th>
<th>Time / By when</th>
<th>Indicators</th>
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### Activity: Action plan session (25 min)

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<th>Responsible person</th>
<th>Resources</th>
<th>Time / By when</th>
<th>Indicators</th>
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### Plenary session / Discussion (20 min)

- Feedback and peer review of the Action plans

---

**Objectives:**

**Activities:**

**Responsible person:**

**Resources:**

**Time / By when:**

**Indicators:**

---

**Activity:**

**Action plan session**

(25 min)

- Feedback and peer review of the Action plans
MODULE 6: MONITORING AND EVALUATION IN COMMODITY MANAGEMENT AND DATA FOR DECISION MAKING

Duration: 2 hours

Goal: To describe the concept of monitoring and evaluation and its importance in program activities implementation

Objectives:

At the end of this module, the participants should be able to:
• Identify the basic purposes and scope of M&E
• Differentiate between monitoring functions and evaluation functions
• Identify criteria for the selection of indicators
• Discuss M&E indicators and what should be monitored for commodity management
• Describe data quality dimensions and how to detect data quality errors
• Describe the process of using Data for Decision-making

Content:
Introduction to monitoring and evaluation
Definition of key terms; Importance of monitoring and evaluation; what programs should monitor for commodity management

Session 1: Overview of Monitoring & Evaluation (M&E) for commodity management
Definition of terms; Basic concepts of M&E; M&E indicators and what should be monitored for commodity management

Session 2: Improving Data Quality
Definition of terms; Data quality dimensions; how to detect types of errors that impact on data quality; identify data quality challenges in facilities

Session 3: Use of Data for Decision Making
Concepts of decision-making; components of decision-making cycle; Steps of the framework for using data for decision-making

Lesson Plan Guide:

<table>
<thead>
<tr>
<th>UNIT</th>
<th>CONTENT</th>
<th>ACTIVITY</th>
<th>TIME</th>
</tr>
</thead>
</table>
| 1    | • Introduction to monitoring and evaluation  
      • Overview of Monitoring & Evaluation (M&E) for commodity management | • Lecture  
      • Group activity | 45 minutes |
| 2    | • Improving data quality  
      • Use of Data for Decision-making | • Lecture | 45 minutes |
| 3    | • Post Test  
      • Course evaluation  
      • Closing ceremony | | 30 minutes |

References and Recommended Readings


FHI and Division of Reproductive Health under MOH. Training Module on Data for Decision Making. May 2007
Overview of M&E, Data Quality and Use of Data for Decision Making

Module 6

Module Objectives

By the end of the session, the participants should be able to:

- Discuss basic concepts of M&E
- Discuss M&E indicators and what should be monitored for commodity management
- Describe data quality dimensions and how to detect data quality errors
- Describe the process of using Data for Decision-making

Overview of Monitoring & Evaluation (M&E) for commodity management

Module 6 Session 1
Definition of Terms

• Monitoring:
  – The continuous review of the degree to which program activities are completed and targets are met
    *Trying to know if we are doing things in the right way*

• Evaluation:
  – To assess progress toward meeting established objectives and goals, while providing feedback on the performance of the program and direction for future plans
    *Trying to know if we did the right things*

How do you monitor and evaluate?

• M/E is done using **Indicators**

• An Indicator is:-
  – A variable that measures one aspect of a program or project
  – A unit of information measured over time that documents changes in a specific condition.

• An indicator is easy to use, useful, measurable, reliable and valid.

Importance of M&E

M&E helps to:

• Track project/program implementation

• Determine whether the project/program is relevant, efficient and having the intended effects

• Provide an insight into what is working and what is not working

• Identify any operational problems while they can still be corrected, and thus ongoing performance improvement
What programs should monitor for commodity management

- Accurate information on consumption at the Service delivery point (SDP) or facility
- Stock at hand at every facility store
  - Does physical inventory fall within Max-Min levels, over- or under-stock?
  - Is there short expiry stock that can be re-distributed or exchanged?
- Losses (due to expiry, damage, etc)
- Timeliness, accuracy, and completeness of Commodity usage Reporting
- Performance in Procurement and Distribution

Examples of indicators used in Lab commodity management

<table>
<thead>
<tr>
<th>What to monitor</th>
<th>Indicator used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stock status</td>
<td>Percentage of facilities providing the service that experienced stock-out of a tracer commodity* on the day of the visit</td>
</tr>
<tr>
<td></td>
<td>Percentage of facilities providing the service that experienced stock-out of a tracer commodity during a specified period of time</td>
</tr>
<tr>
<td></td>
<td>Proportion of health facilities having expiries of at least one lab commodity from the Tracer Laboratory commodities list</td>
</tr>
<tr>
<td>Inventory management</td>
<td>Proportion of facilities with updated Lab Stock / Bin cards</td>
</tr>
<tr>
<td></td>
<td>Proportion of health facilities where physical stock and record counts are in agreement for a set of tracer laboratory commodities</td>
</tr>
<tr>
<td>Reporting</td>
<td>Percentage of facilities submitting (timely, complete, accurate) lab commodity consumption reports to the central level</td>
</tr>
</tbody>
</table>

*A tracer commodity should be a product whose stock-out would result in cessation of services

Improving Data Quality

Module 6 Session 2
What is Data quality?

- Data quality is an assessment of the ability of the data to serve its purpose in a given context, e.g. planning, decision-making, etc
- High quality data serves its intended purpose
- Data quality also describes:
  The processes and technologies involved in ensuring that data conforms to its organizational requirements

Data quality and Data Quality assurance (DQA)

- Data quality is affected by the way data is entered, stored and managed
- Data quality assurance (DQA) is the process of verifying the reliability and effectiveness of data

Dimensions of Data quality (1)

- Validity
- Reliability
- Precision
- Integrity
- Timeliness
Dimensions of Data quality (2)

- **Validity:**
  Validity refers to the extent to which a measure actually represents what we intend to measure

- **Reliability:**
  Data should reflect stable and consistent data collection processes and analysis methods over time

- **Precision:**
  Precise data have a sufficient level of detail to present a fair picture of performance and enable decision-making

Dimensions of Data quality (3)

- **Integrity:**
  Integrity focuses on whether there is improper manipulation of data

- **Timeliness:**
  Data should be available and up-to-date enough to meet management needs

- **Relevance:**
  Data should be relevant to the program. Data collection tools should include relevant data

Detecting Data quality errors

- What data is missing or unusable?
  - E.g. data for a whole reporting period missing
- Which data values are in conflict?
  - E.g. Contradictions between variables
  - 10 paediatric patients (5 girls and 7 boys)
  - A pregnant male
- Which items or records are duplicated?
- What linkages are missing?
- Calculation and Typing errors
  - E.g. 5000 + 4 = 50004
- Guessing / cooked data
  - Guessing a likely looking number.
- Intentional errors
  - Increase count of clients served to show improved workload
Ways of Improving Data quality

- **Use of Visual scanning**
  Look across each line and then down each column. Able to show errors, e.g. missing data, fluctuations,

- **Trend analysis**
  Shows fluctuations in data over time, inconsistency

- **Calculations**
  Shows up mathematical errors

- **Documented data review procedures** to be performed at all levels

- **Feedback** to reporting facilities on data quality problems picked up from reports

---

**Group Discussion 1**

10 min

- What data quality problems are you experiencing currently at your facility / district related to commodity management?

- How can they be resolved?

---

**Group Discussion 2**

5 min

<table>
<thead>
<tr>
<th>Commodity consumption</th>
<th>Year</th>
<th>2011</th>
<th>2011</th>
<th>2011</th>
<th>2011</th>
<th>2011</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Period</td>
<td>Feb</td>
<td>Mar</td>
<td>Apr</td>
<td>May</td>
<td>Jun</td>
</tr>
<tr>
<td>Name of item</td>
<td>Qty used (Units)</td>
<td>Qty used (Units)</td>
<td>Qty used (Units)</td>
<td>Qty used (Units)</td>
<td>Qty used (Units)</td>
<td></td>
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<tr>
<td>HIV Rapid Test kit</td>
<td>8,023</td>
<td>9,225</td>
<td>7,846</td>
<td>38,225</td>
<td>6,746</td>
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<td>(Determine)</td>
<td>(Units)</td>
<td>(Units)</td>
<td>(Units)</td>
<td>(Units)</td>
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<td></td>
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<tr>
<td>Malaria Rapid</td>
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<td>31,511</td>
<td>35,662</td>
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<td>(Units)</td>
<td>(Units)</td>
<td>(Units)</td>
<td>(Units)</td>
<td></td>
</tr>
</tbody>
</table>

What data quality issues do you see in the above data?
Group Discussion 3  
15 min

Discussion using Samples of some of the Facility reports received at the LMU (central level)

Use of Data for Decision-making

Module 6 Session 3

Why Data for decision making?

- **Scenario 1**: A manager is in the process of hiring someone. Interviews have been conducted, and two candidates emerged tops but they both have very similar qualifications. The manager made a decision on who to hire based on his intuition and opinion.
- **Scenario 2**: When determining the cost of a lab test, e.g. HIV rapid test, a Lab manager may believe that KSh 100 is a reasonable price per test. He/she dictated that the price always be set at KSh 100.

1. For each scenario, do you think the manager made the right decision? Why or why not?
2. What other factors might the manager have considered before making his/her decision?
3. Who else might the manager consult about his/her decision?
The concept of decision-making (1)

• What is a decision?
A Decision is a choice arrived at on the basis of information received that may lead into a course of action.

• What is decision-making?
This is the process involved in identifying alternative courses of action and selecting one or more to undertake with the aim of maximizing attainment of the desired objective. Every decision making process produces a final choice.

The concept of decision-making (2)

• Who makes decisions?
All staff in the organization do

• At what level are decisions made?
Every level of the organization, from executive level to operational level.

The concept of decision-making (3)

• How does the decision-making process happen?
  i. The decision-maker looks at the benefits and disadvantages of the decision situation
  ii. Selects criteria to define the alternatives
  iii. Identifies the alternatives
  iv. Makes a choice from among the alternatives
Components of the Data for Decision-making cycle

The data for decision-making cycle has the following components:

i. Data collection
ii. Data analysis
iii. Decision-making
iv. Action
v. Feedback

Notes:
• The steps flow logically
• Each step influences the next just as all members in an organization influence each other.

Steps of the framework for using data for decision-making

• Step 1: Collect data in a timely way
Two healthcare providers are compiling their monthly commodity consumption report, but they cannot find the relevant registers. Besides they never received feedback on the last two monthly reports they sent in, nobody gave them feedback on. Their supervisor in the district calls to say that the report is already late by two days.

• What problems can you identify here?
• What do you think happened next?
Steps of the framework for using data for decision-making

- **Step 1: Collect data in a timely way**
  - Decide which data should be collected
  - Decide at what level the data will be collected
  - It is important to
    - set deadlines for data collection and
    - ensuring that deadlines are kept

Steps of the framework for using data for decision-making

- **Step 2: Put data into perspective**
  - Facility B reports that they used 400 test kits during the month. Is there a problem with this report?
  To answer that question, a data manager needs to several questions:
  - Does the number reflect an acceptable or unacceptable usage rate?
  - Is it higher or lower than the previous month?
  - Is it what was expected or unexpected?
  - How does it compare to other projects (e.g. service data)?
  - Does it indicate that you are likely to meet your district-wide target to testing?

Steps of the framework for using data for decision-making

- **Step 2: Put data into perspective**
  - Determine its validity and accuracy
  - This enables us to compare it with other data to verify it.
    - Data is said to be valid if it is true in all interpretations
    - Data is said to be accurate if it is found to be close to its true value
Steps of the framework for using data for decision-making

• Step 3

*Find the story – what does the data tell us?*

— Raw data on its own is not useful, it needs to be transformed into information through a process of analysis and interpretation

— One has to
  • Describe the event
  • Find the causes of the event
  • Explain how to control the phenomena

---

Steps of the framework for using data for decision-making

Step 3

*Find the story – what does the data tell us?*

Ask the following questions:

• Have there been any significant differences over time?

• Has the context changed over the period of time during which the data were being collected?

• What were the expectations versus actual outputs/targets according to the work plan and/or other project

• Was there new, unexpected information?

• What lessons were learned?

---

Steps of the framework for using data for decision-making

• Step 4

*Develop explanations and determine actions to take. Take action.*

— Interpret the data so that you can tell the story

— Always ask why something happened

— Examine trends to note, changes over time

— Compare actual outputs to targets

— Test if your conclusions are true
Steps of the framework for using data for decision-making

- Step 5
  
  **Report on analysis and actions**
  
  - State lessons learnt
  - Prepare a summary brief or complete report in simple language for ease of understanding

Steps of the framework for using data for decision-making

- After reporting, remember to
  
  **obtain feedback**
  
  - Enables adjustment actions to be undertaken
  - Different types of feedback are suitable for different audiences

Steps of the framework for using data for decision-making

1. Collect data in a timely fashion
2. Put data into perspective
3. Find the story
4. Develop explanations and take action
5. Report on analysis and actions
Example of commodity logistics impact

“A poor manager uses statistics as a drunken man uses lamp-posts - for support rather than for illumination”
Andrew Lang
APPENDIX 1: SAMPLE LETTER FOR PRACTICUM SITE VISIT

Insert Letter Head

Date:……………..

Address: …………..

RE: USE OF XXXXX FACILITY LABORATORY AS PRACTICUM SITE FOR TRAINING COURSE ON EFFECTIVE MANAGEMENT OF LABORATORY COMMODITIES

This letter is to kindly request your authorization to use the ______________________________ Hospital Laboratory as a practicum site for the above mentioned training workshop on ______________________________ from 9.30 am to 12 noon.

The training is organized by ______________________________ and shall take place from _____ to _____ at the _______________________ in Nairobi. The goal of this workshop is to provide the necessary knowledge, skills and attitudes to Laboratory staff to enable them function efficiently and effectively in the provision of Laboratory commodity management services. The course strives to provide participants the opportunity to visit a site that is already providing efficient laboratory services.

Thank you for your continued support.

Yours Sincerely

Dr.___________________________

Cc: Chief Laboratory Technologist, XXXX Hospital
<table>
<thead>
<tr>
<th>MTP Action Plan Province</th>
<th>County</th>
<th>District</th>
<th>Objectives</th>
<th>Activities</th>
<th>Responsible person</th>
<th>Resources</th>
<th>Time / By when</th>
<th>Indicators</th>
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<tr>
<td>TIME</td>
<td>MODULE/SESSION</td>
<td>FACILITATOR</td>
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<td>DAY 1</td>
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</tr>
<tr>
<td>8.00 – 8.30 am</td>
<td>Arrival and Registration</td>
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<td></td>
</tr>
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</table>
| 8.30 – 9.30 am| • Welcome and Introductions  
• Remarks from MOH representative  
• Expectations  
• Group Norms and Schedule announcements  
• Course Overview |             |
<p>| 9.30 – 10.30 am | Module 1: Overview of Laboratory Services                                    |             |
| 10.30 – 10.45 am | TEA BREAK                                                                     |             |
| 10.45 – 1.00 pm | Module 2: Laboratory Commodity Management                                     |             |
| 1.00 – 2.00 pm  | LUNCH BREAK                                                                   |             |
| 2.00 – 2.30 pm  | Module 3: Good Laboratory Inventory Management                                |             |
|                | Session 1: Introduction to Inventory Management                              |             |
| 2.30– 3.30 pm  | Module 3: Session 2: Determining Quantities to order and Requesting for Lab Commodities |             |
| 3.30 - 3.45 pm | TEA BREAK                                                                     |             |
| 3.45 – 5.00 pm  | Group Exercise: Quantification of Lab Commodities                             |             |</p>
<table>
<thead>
<tr>
<th>Time</th>
<th>Session</th>
<th>Facilitator</th>
</tr>
</thead>
<tbody>
<tr>
<td>8.30 - 8.45am</td>
<td>Recap of Day 1</td>
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<tr>
<td>8.45 - 9.15am</td>
<td>Plenary Discussion: Quantification Exercise</td>
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<tr>
<td>9.15 – 9.45 am</td>
<td>Module 3 Session 3: Requesting for Lab Commodities</td>
<td></td>
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<tr>
<td>9.45 – 10.45 am</td>
<td>Module 3 Session 4: Good Storage Practices</td>
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<tr>
<td>10.45 – 11.15am</td>
<td>TEA BREAK</td>
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<tr>
<td>11.15 – 11.45am</td>
<td>Group discussion on Storage</td>
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<tr>
<td>11.45 – 1.00 pm</td>
<td>Module 3 Session 5: Receiving and Issuing</td>
<td></td>
</tr>
<tr>
<td>1.00 – 2.00 pm</td>
<td>LUNCH BREAK</td>
<td></td>
</tr>
<tr>
<td>2.00 – 2.30 pm</td>
<td>Exercise: Receiving and Issuing</td>
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<tr>
<td>2.30 – 3.30pm</td>
<td>Module 3 Session 6: Records and Tools supporting Inventory Management Practices</td>
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<td></td>
<td>Demonstration and exercises on use of various tools (1)</td>
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<tr>
<td>3.30 - 4.00pm</td>
<td>TEA BREAK</td>
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<tr>
<td>4.00 – 5.00pm</td>
<td>Demonstration and exercises on use of various tools (2)</td>
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<td></td>
<td>• S11, Lab Stock Card, Laboratory Top-up form, Lab Expiry tracking chart, Lab Temp monitoring chart</td>
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</tr>
<tr>
<td>TIME</td>
<td>SESSION</td>
<td>FACILITATOR</td>
</tr>
<tr>
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</tr>
<tr>
<td>DAY 3</td>
<td></td>
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<tr>
<td>8.15 – 8.30 am</td>
<td>Recap of previous day</td>
<td></td>
</tr>
<tr>
<td>8.30 - 9.30 am</td>
<td>Module 4: Laboratory Commodity and Information flow  Session 1: Introduction to Logistics Management Information System (LMIS)</td>
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<tr>
<td>9.30 – 10.30am</td>
<td>Demonstration and Exercise: Lab LMIS</td>
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<tr>
<td>10.30 – 10.45 am</td>
<td>TEA BREAK</td>
<td></td>
</tr>
<tr>
<td>10.45- 11.30pm</td>
<td>Module 5: Applying the Monitoring Training and Planning (MTP) approach to Laboratory commodity management Session 1: Introduction to MTP and the MTP approach</td>
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<td>11.30-12.30pm</td>
<td>Session 2: Action Planning for facility level Laboratory Commodity Management practice</td>
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<tr>
<td>12:30- 1.15pm</td>
<td>Presentation of action plans (1)</td>
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</tr>
<tr>
<td>1.15 – 2.15 pm</td>
<td>LUNCH BREAK</td>
<td></td>
</tr>
<tr>
<td>2.15 – 3.00pm</td>
<td>Presentation of action plans (2)</td>
<td></td>
</tr>
</tbody>
</table>
| 3.00- 3.45pm     | Module: 6. Overview of M&E, Data Quality and Use of Data for Decision Making: Session 1: Overview of Monitoring & Evaluation Demonstration and exercises on use of various tools (2)  
S11, Lab Stock Card, Laboratory Top-up form, Lab Expiry tracking chart, Lab Temp monitoring chart |             |
| 3.45-4.30pm      | Session 2: Improving Data Quality  
Session 3: Use of Data for Decision-making |             |
| 4.30 – 5.00 pm   | • Post Test  
• Course Evaluation  
• Closing Ceremony |             |
| 5.00 pm          | TEA BREAK                                                               |             |
APPENDIX 4: INVENTORY MANAGEMENT TOOLS

Bin card/Stock card:

1. Description of the stock card
   • Is a serially numbered, single-copy, two-sided card
   • Must exist for each laboratory stock preparation
   • Is used for inventory control of laboratory commodities (including reagents, consumables, chemicals etc) in the facility or district laboratory store
   • Is used for checking physical stock against records in the laboratory store
   • A supply of Laboratory Stock cards is to be kept as a stock item by each laboratory in-charge of a PGH and DMLT

2. Other forms or resources needed to complete the stock card:
   • Invoice or delivery note number accompanying the commodity from KEMSA or supplier
   • Counter Requisition and Issue Voucher (S11) serial number accompanying goods from the general stores of the hospital/facility or district

3. Location
   Must be located next to each item if room available, or in a file or box in close proximity to the commodities—preferably within the laboratory store

4. Who fills the card
   Laboratory in charge/laboratory manager (or his/her designated proxy) is responsible for documentation and recording of all laboratory commodity receipts and issues from the laboratory stores (including storage refrigerators, cool rooms etc).

5. When is the card filled
   Every new receipt and issue must be entered at the time that stock is received or removed, without exception, for purposes of tracking movement of laboratory stock in storage

6. Steps for completing the stock card:
   6.1 Open/complete a separate stock record card for differing strengths/concentrations and units of issue

   6.2 Record for a new stock card:
      • District: laboratory in charge enters the name for the district where facility is located followed by name of facility
      • Commodity name and description: enters the generic name (trade name in brackets) and strength/concentration of the item as appropriate
      • Unit of Issue: enters pack size (e.g. number of tests per kit, or number of pieces per box etc)
      • Item code: enters the code number allocated by the supplier/requisitioning office and normally reflected against the item name on the delivery note from KEMSA/supplier, or S11, S12, S13 or S3
      • Add storage requirements: Enters special storage conditions (e.g. refrigeration at +2 to 8°C)
      • Average monthly consumption: enters a figure which he/she has calculated from
      • Minimum level of stock for that commodity—as calculated by the manager
• Maximum levels of stock—calculated by the manager (Note—since these values change over time, the calculations are best done using the electronic lab commodity tool)

6.3 Receipts: In the receipts column the Lab in charge will enter information as follows: -
  • Date: enters the date the transaction occurred as dd/mm/yyyy
  • Received from: enters the supply source of the stock item - for example KEMSA, NTLP, a name of Supplier, Hospital stores, NPHLS etc
  • Document Number: Writes the number of the invoice or delivery note accompanying the commodity, or the serial number of the S11 or S12 that prompted the order being received.
  • Quantity: enters the amount received (e.g. 1000 tests, 500 pairs of gloves)
  • Batch Number: indicates this as is written on the packaging
  • Expiry date: indicates as written on the item container
  • Location: indicates where goods are received and stored e.g. cold room, or Lab store, or refrigerator
  • Name: enters name of the officer receiving the commodity

6.4 Disbursements /Issues: In the disbursements/ issues column the Laboratory in charge will enter information as follows: -
  • Document number: enters the serial number of S11 or Top-Up Form, or any other order/ request form accompanying the goods that are disbursed/ issued
  • Quantity: inserts the amount/ quantity of goods disbursed (e.g. 10 liters of ethanol, 2 boxes of slides)
  • Destination: specifies the unit or bench or ward to which the goods are issue For the district and PGH Laboratory store, destination could be a lower level/satellite facility such as a health centre or sub-district hospital, or police clinic
  • Name: enters the name of person who physically receives and takes the commodity goods away

6.5 Stock: In the Stock column the Laboratory in charge will enter the following information: -
  • Balance: inserts quantity of commodity in stock after each transaction i.e. after each receipt or issue
  • Unit value: enters value in Kenya shillings for each unit of issue if known
  • Total value: calculates and indicates the total value of commodity left in stock, calculated as balance multiplied by unit value

6.6 At bottom of each page, once the card is full indicate the balance carried forward and to which card number the balance is moved.
### Laboratory Stock Card

<table>
<thead>
<tr>
<th>Commodity name and Description</th>
<th>Unit of issue</th>
<th>Item Code</th>
<th>Storage requirements</th>
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</thead>
<tbody>
<tr>
<td>Average Monthly Consumption</td>
<td>Minimum Level</td>
<td>Maximum Level</td>
<td></td>
</tr>
</tbody>
</table>

### RECEIPTS

| Date | Received From | Doc. No. | Quant. | Batch No. | Expiry | Loc. | Name | Doc. No. | Quant. | Dest. | Name | Balance | Unit Value | Total Value |
|------|---------------|----------|--------|-----------|--------|------|------|----------|--------|-------|------|---------|-----------|------------|-------------|

### DISBURSEMENTS/ISSUES

<table>
<thead>
<tr>
<th>Doc. No.</th>
<th>Quant.</th>
<th>Dest.</th>
<th>Name</th>
<th>Balance</th>
<th>Unit Value</th>
<th>Total Value</th>
</tr>
</thead>
</table>

### STOCK

<table>
<thead>
<tr>
<th>Balance</th>
<th>Unit Value</th>
<th>Total Value</th>
</tr>
</thead>
</table>

Doc. No. = Document Number, Quant. = Quantity, Loc. = Location, Dest. = Destination

Balance C/F ………………. to Card No. …………..
TOP UP FORM

Instructions for completing the laboratory Top-Up form

1. Title:
Laboratory Top-Up Form

2. Objective of instructions:
To describe the correct use of the supply Top-Up Form

3. Description and location of the form
a. Is a serialized two-sided card used to order supplies for use on the bench or section from the laboratory store / laboratory in-charge.
b. It allows for documentation of all issues from the lab store made to the bench or section in order to replenish or top-up lab supplies
c. It helps the lab in-charge to see at a glance the usage of a particular commodity on the bench or section.

4. Location:
The card under current use is kept in a file within the section or next to the bench.

5. Who fills the Top-Up Form?
a. Both the staff who is requesting /ordering items for the bench or section and the one issuing from Laboratory store(s) will fill the form
b. The staff of the bench/section/unit will fill in this card every time he/she is requesting initial or additional commodities from the laboratory store(s).
c. The laboratory in-charge /or the designated laboratory staff will fill relevant columns of this card to the bench or section.

6. Steps for filling the top up form
6.1 The staff or section head who initiates the order fills in the following information:
• Name of Facility: enter name of the laboratory, e.g. “Coast PGH” or “Hola D/H”
• Department/Section: indicate the bench or section, e.g. serology, parasitology, etc or the unit/area ordering this item, e.g. “VCT” or “MCH”, etc
• Date: write down the date of the transaction
• Commodity: indicate the name of the commodity as it appears on the laboratory stock card in the store(s)
• Unit of issue: enter as it appears on the laboratory stock card
• Current balance: enter the amount/quantity of units that are remaining for use on the bench/section
• Tests done: enter the number of tests done between your last order from the store and this order
• Order quantity: indicate the number of units you are ordering through this request

Present the top-up form with this order to the laboratory in-charge or the staff member designated to man the laboratory store(s).

6.2 The laboratory in-charge or his/her designated proxy fills the relevant column as follows:
• Issue quantity: enter the number of units the laboratory in-charge is issuing to the bench
• Issued by: enter name of the issuing officer who also appends his/ her signature
• Received by: enter name of the staff officer collecting/receiving the issued stock for the bench/section
• Remarks: enter any relevant remarks.
<table>
<thead>
<tr>
<th>Date</th>
<th>Commodity</th>
<th>Unit of Issue</th>
<th>Current balance</th>
<th>Order Quantity</th>
<th>Tests done</th>
<th>Issue Quantity</th>
<th>Received by</th>
<th>Name</th>
<th>Sign</th>
<th>Remarks</th>
</tr>
</thead>
</table>

**Name of Facility**: 
**Department/Section**: 
**Serial No.**
Temperature Control Log

Steps for completing Temperature Control Log:

- Facility: Enter the facility name
- Location: Enter the location where the temperature is being checked. i.e. commodities, commodities fridge and Lab fridge.
- A.M./P.M. Time: Enter the time the temperature is checked
- Recorded temp (°C): write down the temperature
- Indicate if the temperature recorded is within the acceptable range by writing Yes or No
- Enter the initials of the person who recorded the temperature

Distribution:

- The logs in use are pinned up in the relevant area.
- Completed logs are filed and stored in the LAB store or Lab working area as may be relevant name of the staff officer collecting/receiving the issued stock for the bench/section
TEMPERATURE CONTROL LOG

<table>
<thead>
<tr>
<th>Date</th>
<th>A.M. Time</th>
<th>Recorded Temp (°C)</th>
<th>Acceptable Yes/No</th>
<th>Initials</th>
<th>P.M. Time</th>
<th>Recorded Temp (°C)</th>
<th>Acceptable Yes/No</th>
<th>Initials</th>
</tr>
</thead>
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# Republic of Kenya
## Ministry of Health
### +2° to +8° C Temperature Monitoring Chart

<table>
<thead>
<tr>
<th>District</th>
<th>Name of Health Facility</th>
<th>Month</th>
<th>Year</th>
</tr>
</thead>
</table>

**Name and Type of Refrigerator:**

- GAS
- ELEC
- SOLAR
- KEROSENE

<table>
<thead>
<tr>
<th>OPERATING ON</th>
<th>GAS</th>
<th>ELEC</th>
<th>SOLAR</th>
<th>KEROSENE</th>
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<table>
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<tr>
<th>Acceptable Temperature -16°C or Colder</th>
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<tr>
<td>12</td>
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<tr>
<td>AM PM</td>
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<table>
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<tr>
<th>Faults and Problems</th>
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</thead>
<tbody>
<tr>
<td>Date</td>
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**DOT**
- The temperature morning and afternoon in degrees centigrade

**Acceptable Temperature:** -16°C or COLDER

**Republic of Kenya**
- Ministry of Health

**District**
- Name of Health Facility
- Month
- Year

**Sign**
- Initials
- Faults and Problems

**From The American People**
- USAID

**USAID**
- Health Commodities and Services Management
Tracking Expiry dates of Laboratory Commodities

Date Approved: October 2012
Approved by: Ministries of Health

1. Objective
To describe the procedure for tracking the expiry dates of Laboratory commodities in order to avert expiries.

2. Responsible Persons
2.1. The facility staff member in charge of managing Laboratory commodities or his/her designated proxy.

3. Tools Needed (Refer to appendices)
3.1. Expiry Tracking Chart for Laboratory reagents and consumables
3.2. Erasable Marker Pen or stickers

Note:
• The Expiry Tracking Chart for Laboratory reagents and consumables is designed to be reusable.
• Use an erasable marker pen or stickers. DO NOT USE A PERMANENT MARKER. If you use a permanent marker, you will need to use spirit or other ink removers to erase the entries.
• The chart has three columns for three years. The first column should be used for the current year and the other two for the following two consecutive years. Fill in the corresponding years.
• At the beginning of each year, delete the old year. Thereafter shift the originally second year to become the first year. Finally, add the subsequent two years in the remaining two columns.

4. Procedure
The staff in charge of the Lab commodities or his/her designated proxy shall
4.1. Write the name of the product in the column marked “Commodity”.

4.2. Indicate the batch number of the product in the second column. This column has space to list up to three different batches or lot numbers.
   If you have more than three batches/lots, record the three that have the shortest expiry.
4.3. Indicate the expiry date for each of the 3 batches listed.
4.4. Mark the month and year when the commodities expire (in the correct year/month column) using a bold red X or sticker
4.5. Mark the month and year when the commodities will reach 3 months prior to expiry date (in the correct year/month column) using a bold green X or sticker
4.6. Take action 6 months prior to the expiry date by issuing, re-distributing or exchanging the short-dated commodities.
4.7. Erase the entry from the chart once the short-dated product has been removed from stock and indicate the next batch to expire.

Note:
• If a commodity is to expire after the three years covered on the chart, do not enter it on the chart. Wait until it is within three years expiry then enter the details of the reagents accordingly.
• If a commodity has a shorter shelf-life than 6 months (e.g. hematology controls), adjust the timelines accordingly.
• You may need to use more than one chart, depending on the number of commodities and Laboratory commodity storage areas.

5. Storage of Tools
5.1. The Expiry Tracking Chart for Laboratory reagents and consumables should be hung on a wall in the Laboratory commodity storage area and should be visible at all times.
**Expiry Tracking Chart for Laboratory Reagents and Consumables**

<table>
<thead>
<tr>
<th>Commodity</th>
<th>Batch No</th>
<th>Expiry Date (dd/mm/yyyy)</th>
<th>Year:</th>
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APPENDIX 5: ISSUE AND RECEIPT OF COMMODITIES

Issue and Receipt Voucher (S12):

- Is serially numbered, bound in a book. Each voucher has six pages, one original and five copies.
- Is used to transfer LAB stocks from one facility to another external site or other peripheral facility.

Steps for completing the Issue and Receipt Voucher (S12):

1. The Lab staff in charge of the LAB commodities or designated proxy records: Under Supplier/Issuing Office
   - Name of Facility: requesting reagents
   - Supplier/Issuing Office: e.g. CPGH
   - Min./Dept: Health/Medical
   - Address: of CPGH
   - Issuing officer: Lab staff’s name
   - Designation and Lab stamp
   - Merchant name/Address: is left blank
   - Date: date stock issued
   - Issue approved by: name of designated authorizing officer signing letter of request
   - Date: date of signed letter
   - Stores packed by: issuing laboratory staff’s name
   - Stores recorded by: issuing laboratory staff’s name
   - Mode of transportation: record how Reagents were sent, or if collected, by whom
   - Designation and Lab stamp
   - L.P.O. No.: is left blank
   - Delivery note: is left blank
   - Invoice: is left blank

2. The requisitioning/receiving facility records upon receipt of Reagents: Under Requisitioning/Receiving Office:
   - Min./Dept.: e.g., Health /Port Reitz District Hospital
   - Indenting unit: Lab
   - Address: of indenting unit
   - Receiving officer: name
   - Designation and stamp:
   - Reasons for demand: Write A
   - Indent approved by: the name of the designated authorizing officer signing letter of request
   - Date: date of letter of request
   - Address for delivery: address of receiving facility
   - Receipt recorded by: name of person recording the receipt of Reagents
   - Chargeable to: is left blank
   - Vote/Head: is left blank
   - S/Head/Item No.: is left blank
   - Quantity Issued Received: number of packs received
   - Certified that the above items/s has/have been Received/Uploaded and recorded on Ledger/Inventory: signed by receiving Lab personnel.
   - Signature of Receiving Officer and Date

Distribution:
- The S12 document has six pages.
  - The original is retained by issuing facility Lab.
  - The receiving facility receives the second and third pages.
  - The fourth page is sent to the receiving facility’s accounts department.
  - The fifth page is sent to the issuing site’s accounts department.
  - The sixth page remains in the book.
- The S12 will be filed together with the letter of request and accompanying needs list from the peripheral facility in the issuing site
  - S-11 has three pages
  - Retain the original copy and file it chronologically for easy retrieval.
  - The duplicate is retained by the receiving facility.
  - The triplicate is left in the S11 book.

Where S12 is not available facilities may use the Counter Requisition and Issue Voucher (S11), and S13
<table>
<thead>
<tr>
<th>Item</th>
<th>Item code</th>
<th>Description</th>
<th>Unit</th>
<th>Quantity Required/Ordered</th>
<th>Quantity Issued/Received</th>
<th>Quantity to follow</th>
<th>Rate Sh.</th>
<th>Total value Cts.</th>
<th>Stock Balance</th>
<th>Ledger Folio. No.</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
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</table>

Total

Certified that the above items/s has/have been Received/Issued and recorded on Ledger/Inventory

Signature of Issuing Officer and Date

Signature of Receiving Officer and Date
# COUNTER RECEIPT VOUCHER

<table>
<thead>
<tr>
<th>Code No.</th>
<th>Item Description</th>
<th>Unit</th>
<th>Quantity</th>
<th>Value</th>
<th>Remarks</th>
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Order No. | Invoice No. | OR

Issuing Officer: ___________________________ Signature: ___________________________ Designation: ___________________________

Account No: ___________________________ Date: ___________________________

GPK 090-1980bKS—5/2002
COUNTER REQUISITION AND ISSUE VOUCHER (S11):

Counter Requisition and Issue Voucher (S11):
- Is used for issue, delivery, and receipt of stock.
- Is a serially numbered triplicate form.
- Is used by the outpatient Lab to order from the Lab commodities, and by wards and other departments to order from the outpatient Lab.
- Is kept in each department requiring its use.

Steps for completing the Counter Requisition and Issue Voucher (S11):
1. Designated person, authorized to requisition the stock, records:
   - Ministry (of Health) Depart./Branch (Medical) and Unit (e.g., CPGH/ART)
   - To (Issue point): e.g., Lab store
   - Issue to (Point of use): receiving location
   - Item Description:
   - Unit of Issue: the quantity in the container or pack size
   - Quantity required: the number of units of issue required
   - Account No.: designated number if appropriate
   - Date: of requisition
   - Requisitioning Officer: name, designation, and signature

2. Designated person, authorized to issue the stock, records:
   - Quantity Issued: the number of units of issue which are actually issued
   - Value: of total amount of stock issued (calculated using average unit price)
   - Remarks/Purpose: expiry date of stock issued
   - Issued by: issuing officer’s name and signature
   - Date: issued

3. Designated person authorized to receive the stock checks identity and quantity of supplies issued and records:
   - Received by: name, designation, and signature

Distribution:
The Counter Requisition and Issue Voucher (S11) is to be completed in triplicate:
- The original is kept by the Lab personnel.
- Duplicate is retained by the initiating department. Triplicate remains in the S11 book.
## COUNTER REQUISITION AND ISSUE VOUCHER

**Republic of Kenya**

<table>
<thead>
<tr>
<th>Code No.</th>
<th>Item Description</th>
<th>Unit of Issue</th>
<th>Quantity Required</th>
<th>Quantity Issued</th>
<th>Value</th>
<th>Remarks/Purpose</th>
</tr>
</thead>
<tbody>
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</table>

**Account No:** ....................................................  **Designation:** (Laboratory in charge) ....................................................  **Date:** ....................................................

**Requisitioning Officer:** ....................................................  **Signature:** ....................................................  **Sign:** ....................................................

**Issued by:** ....................................................  **Designation:** ....................................................  **Date:** ....................................................

**Received by:** ....................................................  **Signature:** ....................................................  **Sign:** ....................................................

---

**Ministry ....................................................**  **Dept/Branch....................................................**  **Unit....................................................**

**To (Issue point)....................................................**

Please issue the stores listed below to (Point of use)....................................................

---

**FORM S11**  **Serial No....................................................**
### APPENDIX 6: DISPOSAL OF LAB REAGENTS OR ITEMS (F.O.58)

**REPUBLIC OF KENYA**

**REPORT OF THE BOARD OF SURVEY ON STORES (UNSERVICEABLE AND SURPLUS TO REQUIREMENTS)**

Ministry of ..........................................................Department ..................................................

N.B.-Column Nos. 1–6 to be completed by the Department prior to the assembly of the Board.

<table>
<thead>
<tr>
<th>Item No.</th>
<th>Article</th>
<th>Quantity</th>
<th>Date Of Purchase</th>
<th>Original Value</th>
<th>State Whether Unserviceable Or Surplus</th>
<th>Board's Report On Condition</th>
<th>Recommendation Of Board For Disposal</th>
<th>Estimated Local Saleable Value If Sale Is Recommended</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
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</table>

**DECISION OF:** -

ACCOUNTING OFFICER

TREASURY

For Minister for Finance

**Signature and Official Designation of Officer-In-Charge of Stores**

Date..................................................

Station..................................................
Steps for completing Report of the Board of Survey on Stores (Unserviceable and Surplus to Requirements) (F.O.58) – applicable to GOK sites only

1. Record:

   • Ministry of: Health; Department: Medical; Station: Facility name

   • Complete the table:
     • Column 1: Item No.
     • Column 2: Article: generic name; strength/concentration; dosage form
     • Column 3: Quantity: write in quantity to be destroyed
     • Column 4: Date of Purchase: write date the product was delivered to facility
     • Column 5: Original value: insert value if known
     • Column 6: State whether Unserviceable or Surplus: Unserviceable

2. Signature and Official Designation of Officer-in-Charge of Stores: Sign; state position, and date the form.

Distribution:

ART Programme: Notification of Reagents Set Aside for Disposal

   • The original is forwarded to the ACCU or KEMSA/LMU

   • One copy of the completed and signed ART Programme: Notification of Reagents Set Aside for Disposal is retained by the Laboratory in charge of the ART Programme.

   • One copy of the completed and signed ART Programme: Notification of Reagents Set Aside for Disposal is retained by the Laboratory in charge.
## APPENDIX 7: REPORTING TOOLS OR LMIS TOOLS WITH INSTRUCTIONS;
### DAILY ACTIVITY REGISTER (DAR-MOH 642)

<table>
<thead>
<tr>
<th>Date</th>
<th>Staff</th>
<th>Unit of Issue of the Commodity</th>
<th>Beginning balance at the start of the Shift</th>
<th>Quantity Received during the Shift</th>
<th>Origin of the received stock</th>
<th>Rent Number</th>
<th>Expiry Date</th>
<th>Quantity used during the Shift</th>
<th>Number of Tests done during the Shift</th>
<th>Losses during the Shift</th>
<th>Adjustments during the Shift</th>
<th>Ending Balance at the end of the Shift</th>
<th>Remarks (including explanations for losses or adjustments)</th>
<th>Name of officer</th>
<th>Signature</th>
</tr>
</thead>
<tbody>
<tr>
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</tbody>
</table>

Positive  Negative

Page Tools:

Date:   Balance carried forward:   Days out of stock:  

(Use the ending balance quantity from last row on the page)
How to fill the daily activity register (DAR).

1. **Title of Register**
   Daily Activity Register for Laboratory Reagents and Consumables (MoH 642)

2. **Objective**
   To describe the procedure for completing the register accurately

3. **Description**
   This tool is to be used for recording and tracking reagents and consumables issued and used daily / in each shift in the laboratory or the SDP.
   It also records and tracks the number of tests done, quantity used and ending balance by end of each day/shift.

4. **Location of the register**
   Must be physically located or placed on every bench, lab section or user point e.g. service delivery point (SDP) such as VCT.

5. **Who will fill it?**
   The laboratory technician/technologist or primary user of the item during the particular shift duty / day at the particular bench/section/SDP.

6. **When to fill the register?**
   For each shift/day, (i) Immediately you receive new stock of a commodity, complete the first eight (8) columns of the register (ii) Each time you undertake a test.

7. **How to use the register for different commodity items**
   Before using this register, the Lab In-Charge must allocate each commodity a number of pages according to frequency with which the commodity is used/ consumed. Each different unit of issue of a commodity should be on a separate page.

8. **Steps for filling the register**
   On the first page of the DAR book, fill in the following:-
   Facility name: Write the name of your health facility.
   Facility code: Write your Master facility list code.
   Province/County: Write the province or county where the health facility is located.
   District: Write the district where the health facility is located.

   During each shift, the Laboratory staff or In-Charge must fill in the details as follows:

   8.1 Enter the bench or section each time you open a new page
   8.2 Enter the name of the commodity being tracked in the space provided
   8.3 Fill in the date in the format dd/mm/yyyy in the first column
   8.4 Enter the shift duty as appropriate (e.g. 8 to 5; 8 to 2; 2 to 7; etc)
   8.5 Enter the unit of issue e.g. 1 kit of 50 tests each (1x50 tests)
   8.6 Enter the Beginning balance of the commodity at the bench or section. This is the amount of the particular commodity left over from the previous shift or filled page
8.7 Fill in the quantity received by the bench during your shift

8.8 Origin of the received stock: State the source or where the item originated from e.g. Lab i/c, Lab Store, DTLC, main hospital store

8.9 Indicate the batch number and expiry date of the received items as indicated on the package/container

8.10 Quantity used during the shift: Enter the total quantity of the commodity used during the shift/day

Note: This quantity should include the sum of the quantity of commodity actually used for testing as well as quantity used for repeats and controls.

8.11 Number of Tests done during the shift: At the end of the shift/day, count the total number of tests done using information from the relevant Test register (e.g. Haematology register), and enter the total in the relevant row and column

8.12 Losses during the shift: In this column, enter the total amount/volume of commodity that was lost if any (e.g. from spills, damage, pilferage, etc)

8.13 In the Adjustments column, enter the number of tests either received from another facility or excess quantities counted when stock-taking (Positive adjustments) or number of tests given to another site or quantities of stock found to be missing when stock-taking or those quantities used for training purposes (Negative adjustments).

8.14 Ending balance at the end of the shift: Enter the quantity of the specific commodity that you have on hand at the end of the shift/day

8.15 In the “Remarks” column, record any special comments, including an explanation for the losses or adjustments

8.16 Enter the name and signature of the Laboratory technician/technologist or primary user who is filling the register, in the appropriate columns

8.17 Every time a page is full, add up the figures in the following columns “Quantity received during the shift”, “Quantity used during the shift”, “Number of tests done during the shift”, “Losses during the shift”, “Positive adjustments”, “Negative adjustments”. Fill the totals in the Row marked Page totals.

8.18 On the full page, also enter the Balance carried forward from the last row’s Ending balance, the date and complete the number of Days out of stock, if any, for that commodity.

The totals of each individual commodity from the various pages should be added up at the end of the month and used to fill the Facility CDRR.
### Facility Consumption Data Report and Request (F-CDRR) for ART Laboratory monitoring reagents

#### Table: Facility Consumption Data Report & Request (F-CDRR) for ART Laboratory Monitoring Reagents

<table>
<thead>
<tr>
<th>COMMODITY CODE</th>
<th>COMMODITY NAME</th>
<th>UNIT</th>
<th>ORDERED</th>
<th>ORDERED BASED ON</th>
<th>UNIT PRICE</th>
<th>ORDERED QUANTITY</th>
<th>頂頭消費</th>
<th>OVERDUE QUANTITY</th>
<th>OVERDUE</th>
<th>ORDERED QUANTITY</th>
<th>頂頭消費</th>
<th>OVERDUE QUANTITY</th>
<th>OVERDUE</th>
</tr>
</thead>
<tbody>
<tr>
<td>D002</td>
<td>Test Tubes</td>
<td>Each</td>
<td>100</td>
<td>ART Laboratory</td>
<td>$1.50</td>
<td>200</td>
<td>200</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
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<td>0</td>
</tr>
<tr>
<td>D003</td>
<td>Paper</td>
<td>Sheet</td>
<td>100</td>
<td>ART Laboratory</td>
<td>$0.50</td>
<td>200</td>
<td>200</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>D004</td>
<td>Test Strips</td>
<td>Each</td>
<td>100</td>
<td>ART Laboratory</td>
<td>$2.00</td>
<td>200</td>
<td>200</td>
<td>0</td>
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</tbody>
</table>

**Note:**
- The table above lists the consumables used for ART laboratory monitoring reagents.
- The columns include: Commodity Code, Commodity Name, Unit, Ordered, Ordered Based On, Unit Price, Ordered Quantity, and Overdue Quantity.
- The data is organized for easy tracking and management of laboratory supplies.

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**Effective Management of Laboratory Commodities Participants' Guide**

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**Username:**

---

**Date:**

---

**Signed:**

---

**Approved:**

---

**Remarks:**

---

**Reference:**

---

**Page:** 145
Facility Consumption Data Report and Request (F-CDRR) for ART Laboratory monitoring reagents

How to complete the Facility Consumption Data report and request (F-CDRR) form for ART Laboratory monitoring reagents

Use the following instructions to complete the F-CDRR for ART Laboratory monitoring reagents:

Use of the F-CDRR:
- Used to report data on the following commodities: Laboratory reagents for ART treatment monitoring (i.e. reagents for CD4 count, Haematology, Biochemistry) and related consumables at facility level
- It is used to summarize and report on consumption of individual commodities used by a facility each month, and also to request stock for the facility from KEMSA or other supply chains.
- Source of Data: It draws its information from the Daily Activity Register for Laboratory Regents and Consumables (MOH 642).

Who will fill the F-CDRR and When?
- The Facility Laboratory in-charge or his/her designated deputy will complete the form.
- The F-CDRR is completed by the facility at the end of each month and sent to the Logistics Management Unit (LMU) before the 10th of the following month.

To be filled on the second page of the F-CDRR book:

Name of Facility: Write the name of your health facility/laboratory.
Facility code: Write your Master Facility List (MFL) code.
Province/County: Write the province or county where the health facility/laboratory is located.
District: Write the district where the health facility is located.

To complete a report each month:

Facility name: Write the name of your health facility/laboratory
Facility code: Write your Master Facility List (MFL) code.
District: Write the district where the health facility/laboratory is located.
Province/County: Write the province/county where the health facility/laboratory is located.
Affiliation: Indicate the affiliation for your health facility/laboratory by ticking (√) in the appropriate box.
(Options: Ministry of Health; Local authority; FBO (Faith-based organization); NGO (Non-governmental organization; Private)
Report for Period: Beginning (date): Write the date of the first day of the period covered by the report. Enter as day-month-year (in format dd-mm-yyyy), e.g. 01/09/2011.
Report for Period: Ending (date): Write the date of the last day of the period covered by the report. Enter as day-month-year (in format dd-mm-yyyy), e.g. 30/09/2011.
Number of CD4 Tests conducted: In this section, write the total number of CD4 tests conducted at the facility during the month, i.e. the actual number of tests done without repeats or failed runs.
Commodity code, Commodity name, Unit of Issue: The commodities are pre-printed on the form along with a code and unit of issue. For each commodity, complete the columns in the main part of the reporting form as follows:
Beginning Balance: Enter the quantity, in units, of each usable ART laboratory reagent or consumable available for use in the facility at the beginning of the reporting month.

Note: Please include all commodities irrespective of the supplier from whom they were sourced.

NB. The Beginning balance for this month should be equal to the Physical count (Ending balance) at the end of the previous month. If it is not, indicate the loss or adjustment in the respective columns of this F-CDRR and explain in the Comments section.

Quantity Received from the Central Warehouse (e.g. KEMSA): Enter the total Quantity, in units, of each ART laboratory reagent or consumable received by your facility from the Central Warehouses, i.e. KEMSA or SCMS during the reporting month under consideration. If no stock was received at the facility during the month, enter a zero (“0”) in this column.

The Quantity received is obtained from the Quantity Received or Receipts section of the Bin card / Stock card.

Quantity Received from Other Source(s): Enter the total Quantity, in units, of each ART laboratory reagent or consumable received by your facility from any other sources (e.g. partner, donor) during the reporting month under consideration. If no stock was received at the facility during the month, enter a zero (“0”) in this column. The Quantity received is obtained from the Quantity Received or Receipts section of the Bin card / Stock card.

Quantity Used: Enter the total Quantity, in units, of each ART laboratory reagent or consumable used in the facility during the reporting month. If no reagent or consumable was used during the month, enter a zero (“0”).
Note: This quantity should include the sum of the quantity of commodity actually used for testing as well as quantity used for repeats and controls.

The Quantity used is obtained from the Daily Activity Register for Laboratory Regents and Consumables (MOH 642) or equivalent records of consumption in the facility. If several pages of the DAR have been used over the month, be sure to aggregate the figures across all the pages used that month for each laboratory reagent or consumable.

**Losses / Wastage:** Enter the total Quantity, in units, of each ART laboratory reagent or consumable lost or wasted at the facility during the month, e.g. defective, damaged or expired stock. The reason for the loss/wastage should be written in the “Comments” section.

**Adjustments [Indicate if (+) or (-)]:** Enter the total positive or negative adjustments for the month, for each ART laboratory reagent or consumable.

An **adjustment** is a change in stock balance for any reason other than reagents used in the laboratory or quantities received from suppliers.

Positive adjustment: Enter the quantity of each commodity that was added to the facility stock for any reason except receipt from the official supplier during the month. Examples: transfer of stock from another health facility to your facility, excess quantities counted when stock-taking.

Negative adjustment: Enter the quantity of each commodity that was removed from the facility stock for any reason except use in laboratory testing or loss/wastage. Examples: transfer of stock from your facility to another health facility, quantities of stock found to be missing when stock-

Adjustments should be recorded in the Bin card when they occur. The reason for the adjustment should be written in the “Comments” section.

** Ending balance (PHYSICAL COUNT at end of Month):** At the end of each month, conduct a physical count of each ART laboratory reagent or consumable, irrespective of their source. Enter the total Quantity, in units, of each ART laboratory reagent or consumable counted in the facility at the end of the reporting month. If there is no stock at the facility for a commodity, enter a zero (“0”).

Report any differences between the Physical count and the expected Stock Balance in the Bin card as Adjustments or Losses. The reason for the adjustments or losses should be written in the “Comments” section.

**Quantity requested:** Calculate the quantity, in units, for each ART laboratory reagent or consumable that your health facility should order, guided by the following formula:-

- Multiply the reported Quantity used by 4
- Subtract this value from it the Ending balance (Physical count)
- This gives the Quantity requested in units.

**Tick Type of Machine at the Facility as appropriate [Haematology Analyser / Biochemistry Analyser]:** For Haematology and Biochemistry sections, tick, in the boxes provided, the relevant machine that your health facility is using.

**Comments:** Use this space to explain losses and adjustments that have been reported as explained above. Additional remarks related to the commodities reported may also be made, e.g. need for stock re-distribution to prevent expiry.

**Order for extra LMIS tools:** Use this section to request for additional DAR or F-CDRR only when the tools in the facility are nearly full. Indicate quantity required for each type of tool.

**Filled by:** As the person who has filled in this form, write your name, designation, telephone contact, and date that the report was prepared, and sign the form.

**Approved by:** Take the report to the Facility in-charge (or other authorized person) for review. Once reviewed, this person should write their name, designation, telephone contact, and date that the report was approved, and sign the form.

The signature of the programme manager or other authorized person confirms that the report has been reviewed and is valid.
Facility Consumption Data Report and Request (F-CDRR) form for Laboratory commodities (MOH 643)

<table>
<thead>
<tr>
<th>Facility Consumption Report and Request (F-CDRR) form for Laboratory Commodities</th>
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<tbody>
<tr>
<td><strong>Facility Name</strong>:</td>
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<tr>
<td><strong>Local Authority</strong>:</td>
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<td><strong>Date of Request</strong>:</td>
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<tr>
<td><strong>Date of Consumption Report</strong>:</td>
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<td><strong>Type of Report</strong>:</td>
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<td><strong>No. of Items Requested</strong>:</td>
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<td><strong>No. of Items Consumed</strong>:</td>
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</tbody>
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<thead>
<tr>
<th>Commodity Name</th>
<th>Unit of Issue</th>
<th>Beginning Balance</th>
<th>Quantity Delivered</th>
<th>Quantity Issued</th>
<th>Opening Balance</th>
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<td><strong>Commodity 1</strong></td>
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<td><strong>Commodity 3</strong></td>
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**Adjustments**

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<th>Department</th>
<th>Physical Issue</th>
<th>Physical Return</th>
<th>Physical Loss</th>
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<td><strong>Department 3</strong></td>
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**Commodity Balance**

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<tr>
<th>Commodity Name</th>
<th>Opening Balance</th>
<th>Physical Issue</th>
<th>Physical Return</th>
<th>Physical Loss</th>
<th>Closing Balance</th>
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**Order for FCDRR**

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<th>Facility Name</th>
<th>Local Authority</th>
<th>Date of Request</th>
<th>Date of Consumption Report</th>
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<td><strong>Facility 3</strong></td>
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**Approval Date**

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<td><strong>Date 3</strong></td>
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**Effective Management of Laboratory Commodities**

**Participants Guide**

Facility Consumption Data Report and Request (F-CDRR) form for Laboratory commodities (MOH 643)
Instructions for completing the Facility Consumption Data Report and Request (F-CDRR) form for Laboratory commodities (MOH 643)

1. Use of the F-CDRR:
   - Used to report data on the Laboratory (Lab) commodities, including reagents, consumables and other commodities in the Lab tracer item list at the health facility level, e.g. a hospital or other type of service delivery point (SDP). [Use the tracer item list identified by NPHLS as a primary priority list for regular tracking; however additional items may be tracked as per individual district or provincial/county tracking requirements.]
   - Each month, it is used to summarize and report on consumption of individual commodities used by all lab sections / areas within a facility.
   - Also used to request stock for the facility from KEMSA or other supply chains, or any other appropriate source.
   - Source of Data: It draws its information from the Daily Activity Register for Laboratory Reagents and Consumables (MOH 642), the Lab Stock cards and / or Top-Up forms in use by the main Lab, Lab store and the various user points within the facility, as applicable.

Who will fill the F-CDRR and When?
   - The Facility Laboratory in-charge or his/her designated deputy will complete the form.
   - Do not leave any blank cells. In case there is no value, put a zero (0).
   - The F-CDRR is completed by the facility at the end of each month and sent to the Logistics Management Unit (LMU) before the 10th of the following month.

To be filled on the second page of the F-CDRR book:

Name of Facility: Write the name of your health facility/laboratory.
Facility code: Write the Master Facility List (MFL) code of your health facility/laboratory.
Province/County: Write the province or county where the health facility/laboratory is located.
District: Write the district where the health facility/laboratory is located.

To complete a report each month:

Name of Facility: Write the name of your health facility/laboratory
Facility code: Write the Master Facility List (MFL) code of your health facility/laboratory.
District: Write the district where the health facility/laboratory is located.
Province/County: Write the province/county where the health facility/laboratory is located.
Affiliation: Indicate the affiliation for your health facility/laboratory by ticking (√) or putting a cross (X) in the appropriate box. Select from the allocated options: - Ministry of Health; Local authority; FBO (Faith-based organization); NGO (Non-governmental organization; Private.

Report for Period: Beginning (date): Write the date of the first day of the month covered by the report. Enter as day-month-year (in format dd-mm-yyyy), e.g. 01/09/2011.

Report for Period: Ending (date): Write the date of the last day of the month covered by the report. Enter as day-month-year (in format dd-mm-yyyy), e.g. 30/09/2011.

Type of Service; No. of Tests done: In this section, write the total number of tests conducted at the facility during the month, for the following services:
   (i) HIV Testing: disaggregate the number of tests done by Voluntary Counseling & Testing (VCT), Provider-initiated Counseling & Testing (PITC), Prevention of Mother to Child transmission (PMTCT), Blood screening and any other.
   (ii) Malaria testing: This applies to Malaria tests done with either Rapid Detection Test (RDTs) or with Microscopy in category of above or below 5 years of age. Disaggregate the number of tests done by RDT or Microscopy for each age category given, as well as the number of positive tests obtained.

Commodity name: The various Lab commodities are pre-printed on the form categorized into the various laboratory sections, e.g. Haematology.
Use the additional empty rows to record any item not pre-printed – use the commodity name as it appears on the stock card.
For each commodity, complete the columns in the main part of the reporting form as follows:

Unit of Issue: Enter the unit pack size for each lab commodity available for use in your health facility, e.g. Test, piece, strip.

Beginning Balance: Enter the quantity, in units, of each usable laboratory commodity available for use in the facility at the beginning of the reporting month.

NB: The Beginning balance for this month should be equal to the Physical count (Ending balance) at the end of the previous month. If it is not, indicate the loss or adjustment in the respective columns of this F-CDRR and explain in the section labelled “Explain Losses & Adjustments”.

Quantity Received: Enter the total quantity, in units, of each usable laboratory commodity received by your facility from the central warehouses (e.g. KEMSA, SCMS, NPHLS) or any other sources (e.g. the District store, partner; donor, stock purchases), during the
EFFECTIVE MANAGEMENT OF LABORATORY COMMODITIES

PARTICIPANTS GUIDE

The signature of the Facility in-charge or other authorized person confirms that the report has been reviewed and is valid.

Approved by:

Take the report to the Facility in-charge (or other authorized person) for review. Once reviewed, this person should write their name, designation, telephone contact, and date that the report was approved, and sign the form.

The signature of the Facility in-charge or other authorized person confirms that the report has been reviewed and is valid.

150

EFFECTIVE MANAGEMENT OF LABORATORY COMMODITIES

PARTICIPANTS GUIDE
APPENDIX 8: SESSION AND OVERALL COURSE EVALUATIONS
Effective Management of Lab Commodities by Health Workers in Kenya

Module _________________________ Session _____________________________

1. Did this session meet its stated objectives? If not, explain in what way:

2. What were the most useful features of this session?

3. What were the least useful features of this session?

4. Please comment on the speaker’s presentation style:

Any other comment?
**Report of Losses of Laboratory Commodities**

**GOVERNMENT OF KENYA**

**LOSS REPORT/AUTHORITY TO WRITE-OFF**

**Ministry:**

**Department:**

**Unit:**

The items/plant/equipment/vehicles listed below are the subject of loss to the Government due to the circumstances shown. All the circumstances concerned have been examined and are as reported below and overleaf. Remedial action has been carried out. Submitted for decision.

**Officer-in-Charge:**

**Designation:**

**Date:**

<table>
<thead>
<tr>
<th>Item No.</th>
<th>Code No.</th>
<th>Full Description</th>
<th>Unit</th>
<th>Value of Stock Charge</th>
<th>Estimated Value at the time of loss</th>
<th>Difference in value</th>
<th>Circumstances and Nature of the loss</th>
<th>Remarks</th>
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**Case History**

1. Loss was from (full address) .......................................................................................... and discovered by .................................................. on ..........................................
2. Time and date it occurred .................................................................................................. and discovered by .................................................. on ..........................................
3. Date and time discovered and in what circumstances ........................................................
4. Reason for the delay and who was responsible ..................................................................
5. Brief details of the circumstances of the loss ..................................................................
6. Name of officer responsible for custody ............................................................................ How long was he responsible for custody ............................................................
7. (a) Describe the arrangements for safeguarding the stores ................................................ (b) What periodical checking of the stores is carried out and by whom? ..............................
(e) Were the checks properly carried out? Give the date of the last check and the name of the officer who carried it out.

(d) If proper checking was not performed who is responsible for the omission?

(e) Was the loss, damage, deterioration attributable to any of the following:—(Delete which is not applicable)

1. Unavoidable accident
2. Negligence
3. Incompetence
4. Misconduct?

(f) If through (e) 2, 3 or 4, the name of the person concerned and his involvement in the loss.

8. (a) In the case of theft or fraud when were the police informed?

(b) By whom

(c) When was their investigation performed?

(d) The result of their enquiries

(e) Whether prosecution will or has resulted? (A copy of the report should follow)
   A copy of the judgement should be forwarded.

9. Proposals to prevent a recurrence

10. (a) If the responsibility for the loss in part or whole is attributable solely to the officer referred to at 7 should he be required to make restitution and to answer for his dereliction of duty?

(b) If not, why not?

11. If 10(a) applies, give details of the officer’s emoluments, other income, property and family commitments.

   To the best of my knowledge the above replies cover all the circumstances pertaining to the loss to the Government in respect of the stores referred to.

   Sgd.

   Officer-in-Charge

   Date

REPORT OF THE ACCOUNTING OFFICER OR OTHER COMPETENT AUTHORITY

To: Ministry

Department

Unit

The circumstances concerning the loss valued at…………………………….have been investigated and authority is hereby given to write off the items shown on this report subject to the following:—

1. Disciplinary/Recovery action is to be taken against…………………………………………………………

2. Action to improve the security/safety is to be taken…………………………………………………………

3. Disposal of the items is to be arranged as follows…………………………………………………………

Date…………………………………….

Sgd……………………………………

Permanent Secretary