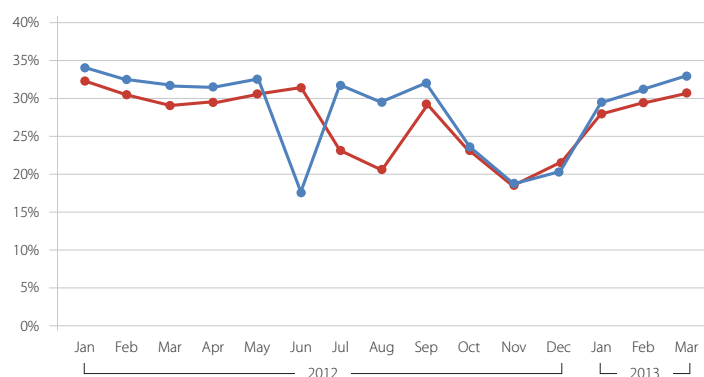
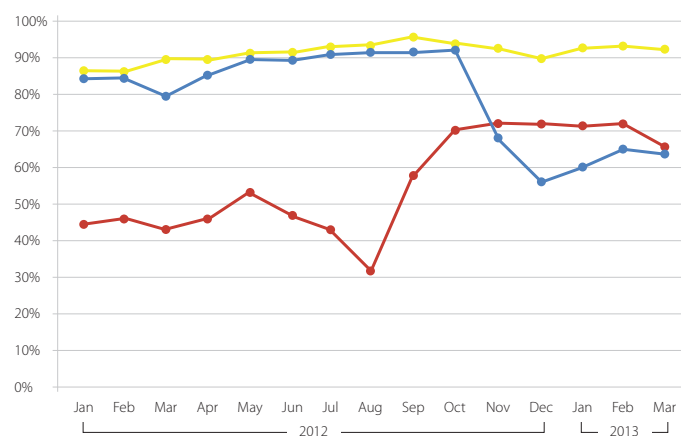
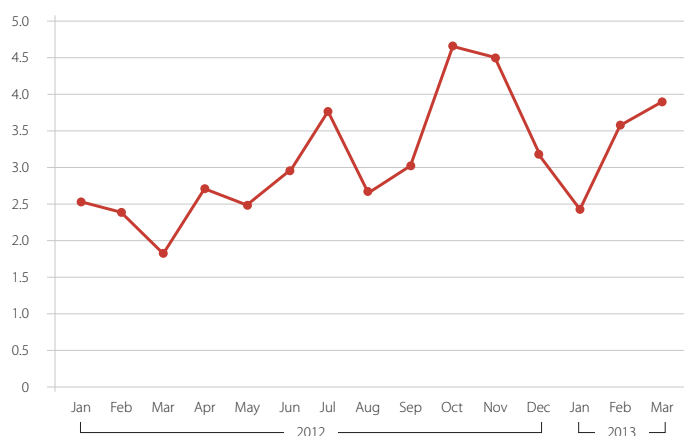




MINISTRY OF HEALTH

MALARIA SURVEILLANCE AND RESPONSE: A COMPREHENSIVE CURRICULUM AND IMPLEMENTATION GUIDE



Participant's Manual

MEASURE Evaluation is funded by the U.S. Agency for International Development (USAID) under terms of Cooperative Agreement GHA-A-00-08-00003-00 which is implemented by the Carolina Population Center, University of North Carolina at Chapel Hill in partnership with Futures Group, ICF International, John Snow, Inc., Management Sciences for Health, and Tulane University. The views expressed in this publication do not necessarily reflect the views of USAID or the United States government. (ms-13-77c)

List of Contributors

Dr. Ahmeddin H. Omar	Division of Malaria Control
Ms. Beatrice Machini	Division of Malaria Control
Ms. Jacinta O. Omariba	Division of Malaria Control
Ms. Jacinta Opondo	Division of Malaria Control
Mr. James Kiare	Division of Malaria Control
Mr. John O. Nyamuni	Division of Malaria Control
Dr. Kiambo Njagi	Division of Malaria Control
Mr. Maurice K'Omollo	Division of Malaria Control
Dr. Rebecca Kiptui	Division of Malaria Control
Mr. Urbanus Kioko	Division of Malaria Control
Ms. Caroline Maina	Division of Disease Surveillance and Response
Dr. Samuel Muiruri	Division of Vector-Borne and Neglected Tropical Diseases
Mr. Patrick Warutere	Division of Health Information System
Dr. Evan Mathenge	Kenya Medical Research Institute
Prof. Simon Kang'ethe	Kenya Methodist University
Dr. Geoffrey Lairumbi	MEASURE Evaluation
Mr. Peter Nasokho	MEASURE Evaluation
Dr. Ann Buff	CDC/PMI
Mr. Paul Malusi	National Public Health Laboratory Services
Mr. Charles Njuguna	World Health Organization

List of Reviewers

Dr. David Soti	DOMC
Dr. Nabie Bayoh	KEMRI/CDC—Kisumu
Dr. Abdisalan Noor	KEMRI Wellcome Trust—Nairobi
Dr. Charles Mbogo	KEMRI Wellcome Trust—Kilifi
Dr. Abdinasir Amin	MEASURE Evaluation
Dr. Daniel Wacira	USAID/PMI
Dr. Dunstan Mukoko	DVBNTD
Dr. Ayub Many	HMIS
Dr. Elizabeth Juma	KEMRI
Dr. Akpaka Kalu	WHO
Dr. Gausi Khoti	WHO
Dr. Daniel Langat	DDSR
Steve Yoon	CDC/PMI
Christine Hershy	CDC/PMI
Dr. Villegas Leopoldo	ICF International
Dr. Yazoume Ye	ICF International
Raphael Pudo	Abt Associates

List of Editors

MEASURE Evaluation
Division of Malaria Control

Foreword

Strengthening of malaria surveillance, monitoring and evaluation systems with the aim of routinely monitoring and evaluating key malaria indicators at all levels of health service delivery is a key objective of the National Malaria Strategy (NMS) 2009–2017, which aims to achieve our ultimate vision of a malaria-free Kenya.

Data from a variety of surveys and operational research show declines in malaria parasite prevalence, malaria trends, vector densities and other entomological indices over the last ten years and the Division of Malaria Control (DOMC) plans to ensure that further reductions are achieved by strengthening surveillance and response. It is anticipated that the generation of focused, timely, scientifically sound information through robust surveillance systems will provide evidence to counties/sub counties and the DOMC to make decisions on interventions for sustaining control of and eventually eliminating malaria.

This curriculum on *Malaria Surveillance and Response—A Comprehensive Curriculum and Implementation Guide* will facilitate attainment of a key malaria control objective of reducing malaria incidence and mortality by equipping health care workers with the knowledge, skills and attitude to effectively undertake and implement a robust malaria surveillance system. This curriculum will be used in the roll out of national malaria surveillance systems to the whole country.

This curriculum will be reviewed periodically in response to expressed need to improve the surveillance systems in the country. I am confident that this curriculum and implementation guide will be found extremely useful.



Dr. S. K. Sharif MBS MBChB, MMED, DLSHTM, MSc
Director Public Health

Acknowledgments

The development of the *Malaria Surveillance and Response—A Comprehensive Curriculum and Implementation Guide* involved an elaborate consultative process involving several key stakeholders in malaria control. The Department of Disease Prevention and Control would like to thank the Director Public Health Dr. S. K. Sharif for providing policy guidance and technical directions to the development of this curriculum.

The commitment, technical support and overall stewardship from the members of the Malaria Inter-agency Coordinating Committee and the United States President's Malaria Initiative (USAID/CDC) through MEASURE Evaluation is highly appreciated. I acknowledge the contribution and technical support from the World Health Organization (WHO–Kenya country office), Division of Disease Surveillance and Response (DDSR), Division of Health Information System (DHIS), the National Public Health Laboratory Services (NPHLS) and Division of Vector Borne & Neglected Tropical Diseases (DVBNTD) to the finalization of this malaria surveillance curriculum.

My sincere gratitude to the United States President's Malaria Initiative (USAID/CDC) through MEASURE Evaluation for financing the development of the malaria surveillance curriculum.

I also like to acknowledge and appreciate both the internal and external reviewers for their valuable contributions and critical review without which this curriculum would not have been realized.

I would like to thank the staff of the Division of Malaria Control for coordinating the development of the malaria surveillance curriculum.

A handwritten signature in black ink, appearing to read 'Willis S. Akhwale', with a horizontal line drawn through the middle of the signature.

Dr. Willis S. Akhwale MBS

Head of Department of Disease Prevention and Control

Acronyms

ACSM	Advocacy, Communication and Social Mobilization
ACT	Artemisinin-based Combination Therapy
AL	Artemether-Lumefantrine
ANC	Ante-Natal Clinic
CDC	Centers for Disease Control and Prevention
CFR	Case Fatality Rate
CHW	Community Health Worker
CM	Case Management
DDSC	District Disease Surveillance Coordinator
DDSR	Division of Disease Surveillance and Response
DHIS	Division of Health Information Systems
DMCC	District Malaria Control Coordinator
DOMC	Division of Malaria Control
DOMT	Disease Outbreak Management Teams
DVBNTD	Division of Vector-Borne and Neglected Tropical Diseases
DPH	Dihydro-artemesinin Piperaquine
eIDSR	Electronic Integrated Disease Surveillance and Response
ELISA	Enzyme Linked Immunosorbent Assay
EPR	Epidemic Preparedness and Response
EWS	Early Warning Systems
GIS	Geographic Information System
GoK	Government of Kenya
HMIS	Health Management and Information Systems
IDSR	Integrated Disease Surveillance and Response
IEC	Information, Education and Communication
IP	In-Patient
IPTp	Intermittent Preventive Treatment in Pregnancy
IRS	Indoor Residual Spraying
ITN	Insecticide Treated Nets
IV	Intravenous
LLIN	Long Lasting Insecticidal Nets
M&E	Monitoring and Evaluation
MIS	Malaria Indicator Survey
MoH	Ministry of Health
NMS	National Malaria Strategy
OJT	On-Job Training
OP	Out-Patient
OPD	Out-Patient Department
PC	Personal Computer
PCR	Polymerase Chain Reaction
PSI	Population Services International
PSCM	Procurement and Supply Chain Management
QA	Quality Assurance
QBC	Qualitative Buffy Coat
QC	Quality Control

Acronyms

RBC	Red Blood Cell
RDT	Rapid Diagnostic Test
SD	Standard Deviation
SOP	Standard Operation Procedure
SP	Sulfadoxine Pyrimethamine
TPR	Test Positivity Rate
WBC	White Blood Cells
WHO	World Health Organization

Part A: How to Use This Curriculum and Implementation Guide

This Curriculum is designed in a simple, easy to use format. It is divided into two parts, A and B.

PART A Presents the foundation of The Curriculum and Implementation Guide showing detailed front matter, the module titles, objectives, and content.

PART B Presents power-points used for all modules of the course.

The facilitators will need to use teaching methods that are appropriate for adult learners, including brainstorming, group discussions, overview lectures and participant presentations.

Malaria Surveillance Course Objectives

The main broad objectives or outcomes of the Malaria Surveillance and Response course are that, at the end of the training, the health care workers will be able to effectively:

1. Explain and articulate the importance of DISEASE SURVEILLANCE
2. Participate in MALARIA IDENTIFICATION, CONFIRMATION AND REPORTING
3. Carry out tasks specified under MALARIA SURVEILLANCE DATA MANAGEMENT
4. Generate and explain CORE MALARIA SURVEILLANCE GRAPHS
5. Illustrate the significance of MALARIA ENTOMOLOGICAL SURVEILLANCE
6. Undertake MALARIA EPIDEMIC PREPAREDNESS AND RESPONSE activities.
7. Participate in and undertake activities pertaining to malaria surveillance SUPERVISION AND FEEDBACK.

Content

1. Introduction
2. Purpose of Course
3. Target Group
4. Course Duration
5. Certification
6. Course Organization

Module 1 Introduction and Overview of Disease Surveillance

Unit 1: Introduction and Overview to Disease Surveillance
Unit 2: Basic malaria epidemiology
Unit 3: Overview of the National Malaria strategy
Unit 4: Malaria control interventions

Module 2 Malaria Identification, Confirmation, and Reporting

Unit 1: Identification of malaria cases
Unit 2: Case confirmation
Unit 3: Reporting

Module 3 Malaria Surveillance Data Management

Unit 1: Data collection, processing and flow
Unit 2: Data quality
Unit 3: Data analysis, presentation and interpretation
Unit 4: Data demand and use for policy and program management

Module 4 Core Malaria Surveillance Graphs

Unit 1: Malaria surveillance indicators, targets and data sources
Unit 2: Introduction to WHO core malaria surveillance graphs
Unit 3: Malaria surveillance graphs and interpretations
Unit 4: Malaria surveillance summary tool

Module 5 Malaria Entomological Surveillance

Unit 1: Introduction to malaria entomology
Unit 2: Surveillance of malaria vectors
Unit 3: Mapping of malaria vectors
Unit 4: Insecticide susceptibility and cone bioassay tests

Module 6 Malaria Epidemic Preparedness and Response

Unit 1: Introduction to malaria epidemics
Unit 2: Malaria epidemics thresholds setting in Kenya
Unit 3: Methods of malaria epidemic prevention
Unit 4: EPR Planning, and response to malaria epidemics
Unit 5: Post epidemic assessment

Module 7 Supervision and Feedback

Unit 1: Introduction to malaria supervision

Unit 2: Planning for malaria supervision

Unit 3: Conducting the malaria support supervision

Unit 4: Report writing and feedback

7. Training and Facilitation

8. Performance Assessment

9. Curriculum Implementation

10. Curriculum Review and Change

11. References and Recommended Readings

1. Introduction

Division of Malaria Control in Kenya has been in the process of operationalizing the WHO Manual for Surveillance in endemic settings. The Division of Malaria Control has so far realized the adoption of common indicators and dashboards for malaria program monitoring. With the aim of determining whether the data required for malaria surveillance indicators exists at the national, sub-national (district and health facility) level a series of international and national consultative workshops were held, a gap analysis of the existing systems carried out and a pilot of malaria surveillance data collection tool conducted in selected districts.

In this regard, the DOMC has developed the curriculum to train health workers on how to carry out an effective malaria surveillance at all service levels in the awareness that surveillance systems consists of tools procedures, people and structures which are required to generate information for planning, monitoring and evaluating malaria programs.

2. Purpose of the Course

The purpose of this course is to equip health care workers across the health care delivery system with the necessary knowledge, skills and attitudes that will enable them to effectively carry out malaria surveillance activities.

3. Target Group

The course is designed for all health care workers at all service levels who in the course of their duty participate in carrying out the malaria surveillance activities. The target group includes, but not limited to the following; disease surveillance teams, malaria control coordinators, medical practitioners, clinical officers, nurses, laboratory technologists, public health officers, health records information officers and pharmaceutical technologists.

4. Course Duration

The course is designed in a modular format which allows for very flexible implementation. It can be implemented in a period of 5 days as an intensive course.

However for busy working health professionals several modules can be covered at a time with subsequent coverage of the remaining modules as planned by organizers.

5. Certification

Upon successfully attending all the modules of the course as outlined in this curriculum, participants will be awarded a certificate.

6. Course Organization

Course organization is the comprehensive description of all the modules of the course, and is as indicated below (See EXAMPLE MODULE 1)

- Module 1 Introduction and Overview of Disease Surveillance**
Unit 1: Introduction and Overview to Disease Surveillance
Unit 2: Basic malaria epidemiology
Unit 3: Overview of the National Malaria strategy
Unit 4: Malaria control interventions
- Module 2 Malaria Identification, Confirmation, and Reporting**
Unit 1: Identification of malaria cases
Unit 2: Case confirmation
Unit 3: Reporting
- Module 3 Malaria Surveillance Data Management**
Unit 1: Data collection, processing and flow
Unit 2: Data quality
Unit 3: Data analysis, presentation and interpretation
Unit 4: Data demand and use for policy and program management
- Module 4 Core Malaria Surveillance Graphs**
Unit 1: Malaria surveillance indicators, targets and data sources
Unit 2: Introduction to WHO core malaria surveillance graphs
Unit 3: Malaria surveillance graphs and interpretations
Unit 4: Malaria surveillance summary tool
- Module 5 Malaria Entomological Surveillance**
Unit 1: Introduction to malaria entomology
Unit 2: Surveillance of malaria vectors
Unit 3: Mapping of malaria vectors
Unit 4: Insecticide susceptibility and cone bioassay tests
- Module 6 Malaria Epidemic Preparedness and Response**
Unit 1: Introduction to malaria epidemics
Unit 2: Malaria epidemics thresholds setting in Kenya
Unit 3: Methods of malaria epidemic prevention
Unit 4: EPR Planning, and response to malaria epidemics
Unit 5: Post epidemic assessment
- Module 7 Supervision and Feedback**
Unit 1: Introduction to malaria supervision
Unit 2: Planning for malaria supervision
Unit 3: Conducting the malaria support supervision
Unit 4: Report writing and feedback

7. Training and Facilitation

Trainers and facilitators for the course will be drawn from among various experts in the areas of malaria case management, laboratory, entomology, epidemiology and monitoring and evaluation.

8. Performance Assessment

The learners will be assessed through pre-tests and post-tests. Continuous assessments will also be used through question and answer sessions, practicum and attendance for all the modules will be mandatory. Assignments and group activities will also be assessed and feedback given.

9. Implementation

This is a 5 day course for health care workers. Ideally, the course ought to begin at 8.00 am on a Monday and stretch through to 5.00 pm every day. This implies that participants travelling from far-out districts will have to arrive at the workshop venue by Sunday preceding the week of training to be in time for the starting of the course on Monday morning.

Various teaching/learning methods, appropriate for adult learners will be applied including, overview lectures, brainstorming, demonstrations, small group discussions, case studies, role plays, assignments, practicum, and attendance at all sessions. This course will emphasize innovative methods, appropriate for adult learners.

10. Curriculum Review and Change

Each course will be evaluated by the participants and the facilitators, and the observations recorded. A workshop to review the curriculum will be held after the first five trainings are implemented to incorporate changes and recommendations made, and there after every 2 years.

11. Reference and Recommended Readings

These are appended at the back of each module.

Module 1: Introduction and Overview of Disease Surveillance

OBJECTIVES

By the end of this module participants will be able to:

1. Describe basic disease surveillance concepts
2. Explain basic concepts of malaria epidemiology
3. Explain the objectives and pillars of the National Malaria Strategy (NMS) (2009–2017)
4. Describe main malaria control interventions

CONTENT

- Definition of surveillance, methods/types of surveillance, functions and systems of surveillance (IDSR and HMIS)
- Describe malaria, parasite and vector, prevalence and endemicity in Kenya
- NMS goal, vision, mission, objectives and pillars
- Case management including malaria in pregnancy, vector control, epidemic preparedness and response, surveillance monitoring and evaluation , advocacy communication and social mobilization

LESSON PLAN GUIDE: MODULE 1 (2 ½ hours)

Unit	Content	Activity	Time
Unit 1	Definition of surveillance, methods/types of surveillance, functions and systems of surveillance (IDSR and HMIS)	Lecture and discussion	45 min
Unit 2	Describe malaria, parasite and vector, prevalence and endemicity in Kenya	Lecture and discussion	30 min
Unit 3	NMS goal, vision, mission, objectives and pillars	Lecture and discussion	30 min
Unit 4	Case management including malaria in pregnancy, vector control, epidemic preparedness and response, surveillance monitoring and evaluation , advocacy communication and social mobilization	Lecture and discussion	45 min

REFERENCES AND RECOMMENDED READINGS

1. Ministry of Public Health & Sanitation, Kenya. *Integrated Disease Surveillance and Response in Kenya*. Technical guidelines 2011.
2. WHO 2012. *Disease surveillance for malaria control, operational manual*.
3. WHO 2012. *World Malaria Report*
4. Division of Malaria Control 2009. *National Malaria Strategy 2009–2017*. Ministry of Public Health & Sanitation, Kenya.
5. Division of Malaria Control 2010. *National Malaria Policy*. Ministry of Public Health & Sanitation, Kenya.
6. Division of Malaria Control 2010. *National Malaria Indicator Survey 2010*. Ministry of Public Health & Sanitation, Kenya.
7. Ministry of Public Health & Sanitation, Kenya 2011. *Integrated Vector Management policy guideline*.
8. Noor et al. *The risks of malaria infection in Kenya , BMC Infectious disease 2009*

Module 2: Malaria Identification, Confirmation, and Reporting

OBJECTIVES

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

1. Identify/detect cases of malaria using the standard case definition
2. Describe malaria parasitological diagnostic methods
3. Demonstrate malaria recording and reporting format using appropriate tools

CONTENT

- Clinical presentation of malaria, standard case definition, differential diagnosis
- Test procedures of performing malaria microscopy and rapid diagnostic testing
- Case recording, reporting tools, reporting requirements,

LESSON PLAN GUIDE: MODULE 2 (2 hrs 45 mins)

Unit	Content	Activity	Time
Unit 1	Clinical presentation of malaria, standard case definition, differential diagnosis	Lecture	30 min
Unit 2	Test procedures of performing malaria microscopy and rapid diagnostic testing	Lecture and demonstration	1hr 30 min
Unit 3	Case recording, reporting tools, reporting requirements	Demonstration and group work	45 min

REFERENCES AND RECOMMENDED READINGS

1. Ministry of Public Health & Sanitation, Kenya. *Integrated Disease Surveillance and Response in Kenya*. Technical guidelines 2011.
2. Ministry of Public Health & Sanitation, Kenya. *Quality manual for laboratory diagnosis in Kenya 2013*.
3. Division of Malaria Control 2010. *National Malaria Policy*. Ministry of Public Health & Sanitation, Kenya.
4. Ministry of Public Health & Sanitation, Kenya. *Health information systems manual 2003*.
5. WHO 2012. *Disease surveillance for malaria control, operational manual*.
6. WHO 2011. *Universal access to malaria diagnostic testing, Operational Manual*

Module 3: Malaria Surveillance Data Management

OBJECTIVES

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

1. Identify different types of data sources, and describe the process involved in the malaria surveillance data collection, processing and flow using the existing MOH tools
2. Perform data quality checks to review the reports.
3. Perform simple data analysis tasks, present, interpret and share the results
4. Promote data demand and use for policy and program management

CONTENT

- Types of data sources, the process of data collection, processing, storage and data flow.
- Elements of data quality (accuracy, completeness timeliness, precision, validity, reliability and integrity)
- Definition of statistical measures (mean, median, mode, variance, ratio, proportion, percentage, rate).
- Methods of data analysis and presentation.
- The role of data in decision making, challenges faced in data demand and use

LESSON PLAN GUIDE: MODULE 3 (3 hrs)

Unit	Content	Activity	Time
Unit 1	Types of data sources, the process of data collection, processing, storage and data flow	Overview lecture	45 min
Unit 2	Data quality improvement	Overview lecture	30 min
Unit 3	Data analysis and interpretation, routine and non-routine data	Overview lecture and exercise	1hr 15 min
Unit 4	Data demand and use	Overview lecture	30 min

REFERENCES AND RECOMMENDED READINGS

1. Laurie Liskin. "Dissemination and Data Use Tools". MEASURE DHS. PowerPoint Presentation. 17 June 2009.
2. MEASURE DHS. "Module 7: Disseminating and Using Data for Change". PowerPoint Presentation. Kenya, June 2010.
3. Statistical Service Centre. (1998, March). Retrieved February 2013, from www.reading.ac.uk/ssc.
4. MoH 2010. *HIS training manual for health workers*.
5. MoH 2010. *DHIS training manual*.

Module 4: Core Malaria Surveillance Graphs

OBJECTIVES

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

1. Define the malaria surveillance indicators, data sources and targets
2. Identify the Core Malaria Surveillance Graphs adapted from WHO
3. Explain malaria surveillance graphs/dashboards
4. Demonstrate how the malaria core surveillance graphs are generated and update the summary tools

CONTENT

- Malaria surveillance indicator data sources and targets
- Introduction to WHO core malaria surveillance graphs
- Malaria surveillance graphs and interpretation
- Use of malaria surveillance summary tool in excel

LESSON PLAN GUIDE: MODULE 4 (3hrs)

Unit	Content	Activity	Time
Unit 1	Malaria surveillance indicators and targets	Overview lecture	30 min
Unit 2	Introduction to WHO core malaria surveillance graphs	Overview lecture	30 min
Unit 3	Malaria surveillance graphs and interpretation	Overview lecture	60 min
Unit 4	Malaria surveillance summary tool	Overview lecture, demonstration, and exercise	60 min

REFERENCES AND RECOMMENDED READINGS

1. Division of Malaria Control. (2009b) *National Malaria Strategy 2009–2017*. Ministry of Public Health & Sanitation, Republic of Kenya, November.
2. Division of Malaria Control. (2009c). *Kenya Monitoring & Evaluation Plan 2009–2017*. Ministry of Public Health & Sanitation, Nairobi, June.
3. MEASURE and EVALUATION (2012). *Operationalizing WHO Malaria Surveillance Guidelines in Kenya*.
4. World Health Organization. (2009). *Programme management: Guidelines for countries with moderate to high transmission of malaria*.
5. WHO 2012. *Disease Surveillance for Malaria control: An Operation manual*.

Module 5: Malaria Entomological Surveillance

OBJECTIVES

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

1. Describe the role of mosquitoes in malaria transmission
2. Describe different types of mosquito surveys and their roles in malaria vector surveillance
3. To stratify the distribution, density, behavior of vectors in relation to malaria transmission and control options
4. Describe how to conduct insecticide susceptibility and cone bioassay tests

CONTENT

- Life-cycle of the Anopheles mosquito; main bio-ecological traits of medical importance; vector incrimination and differentiation between other non-vector mosquitoes; interactions between mosquito, parasite and man
- Importance of mosquito sampling; types of mosquito surveys; methods of mosquito sampling;
- Importance of vector maps, key vector parameters in maps, generation of entomological profile maps and their use in selection of vector control options.
- Reasons for determining susceptibility of vectors and residual efficacy of insecticides on sprayed surfaces and insecticide treated materials; WHO tests: susceptibility of adult and larval mosquitoes to insecticides; cone bioassay tests; data interpretation and use

LESSON PLAN GUIDE: MODULE 5 (6hrs 30 min)

Unit	Content	Activity	Time
Unit 1	The role of mosquitoes in malaria transmission	Overview Lecture	1 hr 30 min
Unit 2	Different types of mosquito surveys and their roles in malaria vector surveillance	Lecture, discussions, and demonstrations	1 hr 30 min
Unit 3	Stratifying the distribution, density, behavior of vectors in relation to malaria transmission and control options	Lecture, discussions, and demonstrations	1 hr 30 min
Unit 4	Bioassays for determining the insecticide susceptibility of mosquito populations and residual efficacy of insecticides on sprayed surfaces and insecticide treated materials	Discussions, demonstrations	2hr

REFERENCES AND RECOMMENDED READINGS

1. WHO (2003). *Malaria Entomology and Vector Control: Learners and Facilitators Guide*.
2. RTI International (2012). *Training Manual on Malaria Entomology*.
3. Bruce Chawatt (2000). *Essential Malariology*.
4. Mbogo, C; et al (2012). *Entomological Manual for use by the Technical Teams within the Context of Integrated Disease Surveillance and Integrated Vector Management at the District Level*. Kenya Medical Research Institute, Centre for Geographic Medicine Research Coast, P.O. Box 428, 80108 Kilifi, Kenya.
5. WHO (2005). *Guidelines on Testing Residual Efficacy of Insecticide on Sprayed Surfaces and Insecticide Treated Materials*.
6. WHO 2012. *Global Plan for Insecticide Resistance Management in Malaria Vectors*.

Module 6: Malaria Epidemic Preparedness and Response

OBJECTIVES

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

1. Describe malaria epidemics
2. Demonstrate malaria threshold setting
3. Describe methods of malaria epidemic prevention
4. Develop malaria epidemic preparedness and response plans
5. Describe post malaria epidemic evaluation

CONTENT

- Definition of epidemics, types of epidemics, contributing/predisposing/triggering factors and consequences of epidemics
- Definition of threshold, scientific methods of setting malaria threshold (constant count, third quartile, Cullen method and cumulative sum methods), thresholds proposed for Kenya
- Strategies of malaria epidemics prevention (surveillance—early detection system, vector control—LLINs, IPTp, ACSM), epidemic cycle
- County/district/facility EPR plans (personnel, referral services, diagnostics, commodity supplies, resource mobilization, ACSM, surveillance), rapid assessments
- Assessments (what went wrong, lessons learnt and what can be done better) and preparedness

LESSON PLAN GUIDE: MODULE 6 (5 hrs)

Unit	Content	Activity	Time
Unit 1	Definition of epidemics, types of epidemics, contributing/predisposing/triggering factors and consequences of epidemics	Lecture and discussion	40 min
Unit 2	Definition of threshold, scientific methods of setting malaria threshold (constant count, third quartile, Cullen method and cumulative sum methods), thresholds proposed for Kenya	Lecture and group work	2 hr
Unit 3	Strategies of malaria epidemics prevention (surveillance—early detection system, vector control—LLINs, IPTp, ACSM), epidemic cycle	Lecture and discussion	30 min
Unit 4	County/district/facility EPR plans (personnel, referral services, diagnostics, commodity supplies, resource mobilization, ACSM, surveillance), rapid assessments	Lecture and group work	1 hr 30 min
Unit 5	Assessments (what went wrong, lessons learned and what can be done better) and preparedness	Lecture and group work	20 min

REFERENCES AND RECOMMENDED READINGS

1. Division of Malaria Control 2011. *Epidemic preparedness and response guidelines*. Ministry of Public Health & Sanitation, Kenya.
2. Ministry of Public Health & Sanitation, Kenya. *Integrated Disease Surveillance and Response in Kenya*. Technical guidelines 2011.
3. Division of Malaria Control 2009. *National Malaria Strategy 2009–2017*. Ministry of Public Health & Sanitation, Kenya.
4. WHO 2003. *Prevention and control of malaria epidemics*.
5. WHO 2006. *Systems for early detection of malaria epidemics in Africa*.
6. WHO 2012. *Disease surveillance for malaria control, operational manual*.

Module 7: Supervision and Feedback

OBJECTIVES

At the end of the module the health care workers will be able to:

1. Describe malaria support supervision
2. Develop a plan for Malaria supervision and use the planning tools
3. Perform malaria supervision using the supervisory checklists
4. Write a supervision report and give feedback using the reporting and feedback template

CONTENT

- Define supervision, characteristics of support supervisors, roles of a supervisor, roles of a supervisee, frequency of supervisory visits, supervision approaches.
- Developing a contact list, advance scheduling of the visit, supervisory team, role of the malaria coordinators and disease surveillance coordinators(including introduction to planning tools
- Conducting supervision (including introduction to supervision checklists), Tracking supervision visits
- Analyzing the supervision visits results, report writing, reporting templates, submission of reports, feedback, incentives and other follow up actions (including introduction to reporting and feedback templates)

LESSON PLAN GUIDE MODULE 7 (4 hrs)

Unit	Content	Activity	Time
Unit 1	Introduction to malaria support supervision	Overview lecture	45 min
Unit 2	Planning for malaria supervision	Lecture, practicals on filling the planning tools based on a case study(small group discussion)	45 min
Unit 3	Conducting the malaria support supervision	Lecture, role play ,practicals on filling the supervisory checklist based on a case study(small group discussion)	1hr
Unit 4	Report writing and feedback	Lecture, role play, practical on calculating scores and report writing based on a case study(small group discussion)	1hr 30 min

REFERENCES AND RECOMMENDED READINGS

1. MOPHS (2012) Manual for malaria supervision Nairobi Kenya
2. MOPHS (2012) Integrated disease surveillance technical guidelines

Malaria Surveillance System Training Course Schedule

Venue:

Dates:


Time	Monday	Tuesday	Wednesday	Thursday	Friday
8:30–9:30 am	Climate Setting Introductions Group Norms Expectations	Recap of Day 1 (15 minutes) Module 2: Malaria Identification, Confirmation and Reporting	Recap of Day 2 (15 minutes) Module 4: Malaria Surveillance Graphs	Recap of Day 3 (15 minutes) Module 5: Malaria Entomological Surveillance	Recap of Day 4 (15 minutes) Module 6: Epidemic Preparedness and Response (EPR)
9:30–10:30 am	Malaria Surveillance course objectives by: Opening Remarks Pre-test	Module 2: Malaria Identification, Confirmation and Reporting	Module 4: Malaria Surveillance Graphs	Module 5: Malaria Entomological Surveillance	Module 7: Supervision and Feedback
10:30–11:00 am	Module 1: Introduction and Overview of Disease Surveillance	Module 3: Malaria Surveillance Data Management	Module 5: Malaria Entomological Surveillance	Module 5: Malaria Entomological Surveillance	Module 7: Supervision and Feedback
11:00–11:30 am	TEA & COFFEE BREAK				
11:30–12:00 pm	Module 1: Introduction and Overview of Disease Surveillance	Module 3: Malaria Surveillance Data Management	Module 5: Malaria Entomological Surveillance	Module 6: Epidemic Preparedness and Response (EPR)	Module 7: Supervision and Feedback
12:00–1:00 pm	Module 1: Introduction and Overview of Disease Surveillance	Module 3: Malaria Surveillance Data Management	Module 5: Malaria Entomological Surveillance	Module 6: Epidemic Preparedness and Response (EPR)	Module 7: Supervision and Feedback
1:00–2:00 pm	LUNCH BREAK				
2:00–3:00 pm	Module 1: Introduction and Overview of Disease Surveillance	Module 3: Malaria Surveillance Data Management	Module 5: Malaria Entomological Surveillance	Module 6: Epidemic Preparedness and Response (EPR)	Module 7: Supervision and Feedback
3:00–4:00 pm	Module 2: Malaria Identification, Confirmation and Reporting	Module 4: Malaria Surveillance Graphs	Module 5: Malaria Entomological Surveillance	Module 6: Epidemic Preparedness and Response (EPR)	Post test Course Evaluation Certification Vote of Thanks Final Remarks & Closure
4:00–4:30 pm	TEA & COFFEE BREAK				
4:30–5:00 pm	Module 2: Malaria Identification, Confirmation and Reporting	Module 4: Malaria Surveillance Graphs	Module 5: Malaria Entomological Surveillance	Module 6: Epidemic Preparedness and Response (EPR)	Departure


Part C: Power Point Presentations Slides


Slide 1


Module 1


INTRODUCTION AND OVERVIEW OF
DISEASE SURVEILLANCE


Republic of Kenya
Ministry of Health


Division of Malaria
Control


World Health Organization


MEASURE
Evaluation


USAID
President's Malaria Initiative

Slide 2

OBJECTIVES

- By the end of this module participants will be able to:
 - Describe basic disease surveillance concepts
 - Explain basic concepts of malaria epidemiology
 - Explain the objectives and pillars of the National Malaria Strategy (NMS) (2009 – 2017)
 - Describe malaria control interventions

Slide 3

Unit 1

Introduction to Disease Surveillance

Slide 4

Brainstorming (5 min)

What is Disease surveillance?

Slide 5

Disease Surveillance

- Ongoing, systematic collection, analysis, and interpretation of health-related data essential to the planning, implementation, and evaluation of public health practice, closely integrated with the timely dissemination of these data to those responsible for prevention and control
- **WHO Definition**
 - Regardless of the type of surveillance, remember that surveillance is data that is used for action!

Slide 6

Brainstorming (5 min)

Why do disease surveillance?

Slide 7

Functions of Disease Surveillance

1. Monitor trends, patterns and estimate magnitude of health problem
2. Detect sudden changes in disease occurrence and distribution (Epidemics/outbreaks)
3. Portray the natural history of a disease
4. Monitor changes in infectious agents
5. Detect changes in health practices
6. Evaluate control measures
7. Generate hypotheses, stimulate research
8. Facilitate planning

Slide 8

Surveillance link to action

- Disease control
 - Interruption of transmission
 - Vaccination / prophylaxis
 - Elimination of cause
- Outbreak investigation
- Development and targeting of programs (education, risk reduction, etc.)
- Development of policies, regulations

Slide 9

Components of Surveillance System

- Surveillance systems consists of tools, procedures, people and structures required to generate information for planning, monitoring and evaluating malaria programmes.
- Tools: report forms, tally sheets, registers, patient records
 - Procedures: case definitions, reporting frequency, information flow, data analysis, dissemination
 - People: health workers, community, decision makers
 - Structures: health systems

Slide 10

Level of Surveillance in Health Systems	
1. Community	
2. Health facility (include Laboratory)	
3. District (sub county)	
4. County	
5. National Level	

Slide 11

Types of Surveillance	
• Community-based surveillance	
• Health facility-based surveillance	
• Sentinel surveillance	
• Laboratory based surveillance	

Slide 12

Approaches to Surveillance	
▪ Active vs. Passive (active case search vs routine reporting)	
▪ Categorical / Integrated (One disease or Many)	
▪ Syndromic /Laboratory-based (Case definition or laboratory confirmation)	

Slide 13

Brainstorming (5 min)

What are the systems & tools used for malaria surveillance in Kenya?

Slide 14

Malaria Surveillance in Kenya

1. Health Management and information systems (HMIS)

- Routine malaria surveillance in all epidemiological zones (monthly facility reporting-DHIS2)

2. Integrated Disease Surveillance and Response (IDSR)

- Weekly reporting for priority diseases (e-idsr) for early detection

3. Sentinel Surveillance

- Weekly threshold data from 45 epidemic prone sub-counties (districts) of western Kenya highlands

Slide 15

Malaria Surveillance in Kenya Cont'd

HMIS (monthly)

- OPD clinical & confirmed malaria cases
- Laboratory tested and positive cases
- Inpatient (malaria admissions) & Deaths

IDSR (weekly)

- OPD clinical malaria cases
- Laboratory tested and positive cases
- Malaria related Deaths

Sentinel Surveillance

- Weekly threshold data from 45 epidemic prone districts in western highlands

Slide 16

What are the basic ingredients of a
good surveillance system?

Slide 17

Questions?

Slide 18

Unit 2

Slide 19

Basic Malaria Epidemiology

Slide 20

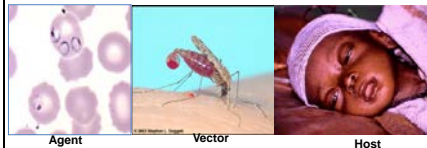
Brainstorm (5 min)

What is Malaria?

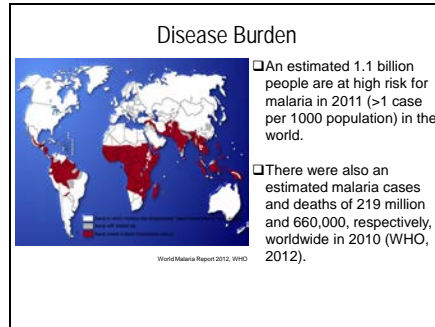
Slide 21

Background

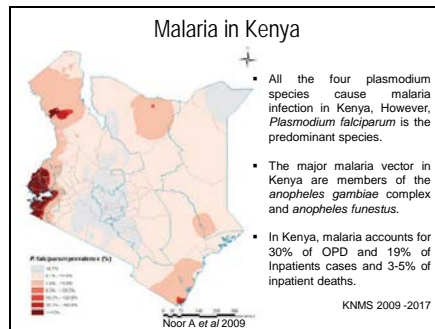
- ❑ Malaria is an acute febrile infection caused by protozoan parasites of the genus *Plasmodium*.
 - *Plasmodium falciparum*, *P. vivax*, *P. ovale*, *P. malariae* and (*P. knowlesi*).
- ❑ Vector: Female Anopheles
- ❑ **Susceptible persons:** Children < 5 years, Pregnant women & non-immune individuals



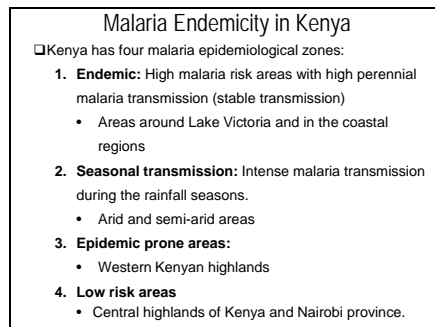
Slide 22



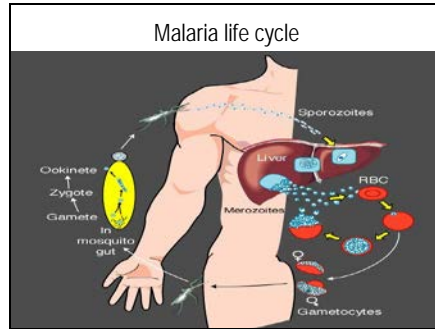
Slide 23



Slide 24



Slide 25

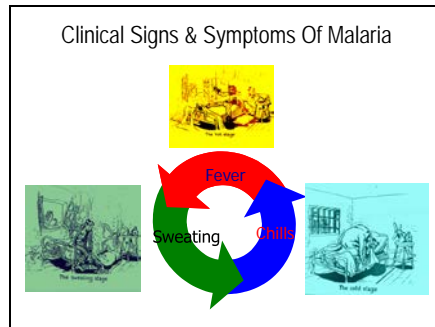


Slide 26

Incubation period

- The time between the infective bite and the appearance of clinical symptoms is approximately
 - 10-14 days for *P. falciparum*,
 - 10-17 days for *P. Vivax* & *P. Ovale*
 - 18-40 days for *P. malariae*
 - 10-14 days for *P. Knowlesi*
 - Some strains of *P. Vivax* & *P. Ovale* mostly from temperate areas may have an incubation period of 8-10 months and even longer.

Slide 27





Slide 28

WHO recommendation on malaria diagnosis

❖ **Parasitological confirmation before treatment**

1. Microscopy
2. Rapid diagnostic tests



Slide 29

Treatment of Uncomplicated Malaria

☐ **First line treatment**

- Artemether-Lumefantrine (AL)
- 6 doses given over 3 days

☐ **Second Line Treatment**

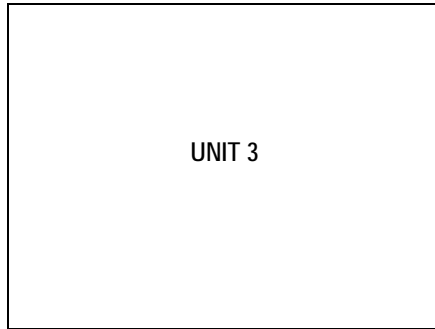
- Dihydroartemisinin-Piperaquine (DHP)

☐ In absence of DHA-PPQ oral quinine should be used

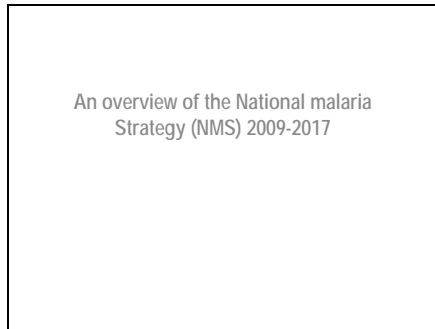
Slide 30

Questions?

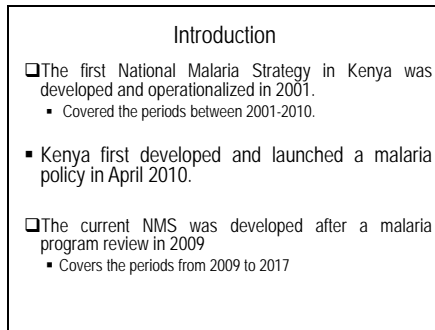
Slide 31



Slide 32



Slide 33



Slide 34

NMS 2009 - 2017

- **Vision:** Malaria free Kenya
- **Mission:** To direct and coordinate efforts towards a malaria free Kenya through effective partnerships
- **Goal:** By 2017, to have reduced morbidity and mortality caused by malaria in the various epidemiological zones by 2/3 of the 2007/2008 levels

Slide 35

Brainstorm (5 min)

What are the Objectives of NMS 2009-2017?

Slide 36

Objective 1

- To have at least 80% of people living in malaria risk areas using appropriate malaria preventive interventions by 2013 through:
 1. Universal LLIN coverage for populations at risk
 2. Indoor Residual Spraying in targeted areas
 3. Prevention of malaria in Pregnancy

Slide 37

Objective 2

- To have 80% of all self-managed fever cases receive prompt and effective treatment and 100% of all fever cases who present to health facilities receive parasitological diagnosis and effective treatment by 2013 by:
 1. Strengthening capacity for malaria diagnosis & treatment
 2. Increase access to affordable malaria medicines
 3. Strengthening home management of malaria

Slide 38

Objective 3

- To ensure that all malaria epidemic prone districts have the capacity to detect and the preparedness to respond to malaria epidemics annually by 2010 through:
 1. Capacity strengthening for epidemic preparedness and response
 2. Strengthen disease surveillance at district level
 - ✓ Surveillance sites
 - ✓ Analysis and interpretation of data
 - ✓ Planning for activities

Slide 39

Objective 4

- To strengthen surveillance, monitoring and evaluation systems so that key malaria indicators are routinely monitored and evaluated in all malarious districts by 2011 through:
 1. Malaria surveillance in all districts
 2. Health facility and school based sentinel surveillance
 3. Malaria data management
 4. Community surveys
 5. Monitoring
 6. Operations Research and Translation
 7. Capacity building

Slide 40

Objective 5

- To strengthen advocacy, communication and social mobilization capacities for malaria control to ensure that at least 80% of people in malarious areas have knowledge on prevention and treatment of malaria by 2014
 1. Capacity strengthening
 - Guidelines
 - Training
 - Monitoring and evaluation
 2. Support for implementing partners
 3. Support for various malaria control interventions

Slide 41

Objective 6

- By 2013, to strengthen capacity in program management in order to achieve malaria programmatic objectives at all levels of the health care system
 1. Planning and partnerships coordination
 2. Program management at provincial and district level
 - Infrastructure strengthening
 3. Resource mobilization
 4. Activity and performance monitoring
 5. Human resource strengthening
 6. Strengthen coordination of PSM for malaria commodities

Slide 42

Pillars of NMS 2009 - 2017

- To achieve the 6 main objectives of NMS, several cross-cutting supportive steps need to be taken.
- These can be referred to as Pillars or Strategic orientations.

Slide 43

Pillars of NMS 2009 - 2017

1. Adopting multi-sector approach to malaria control
2. Decentralizing malaria control operations
3. Basing malaria control interventions on prevailing epidemiology
4. Strengthening the malaria control performance monitoring systems

Slide 44

Unit 4

Malaria Control Interventions

Slide 45

Epidemiological Triad

The diagram illustrates the Epidemiological Triad, a model used in epidemiology to understand the transmission of infectious diseases. It consists of three components arranged in a triangle:

- Agent:** Represented by a microscopic image of red blood cells, indicating the presence of a pathogen.
- Vector:** Represented by a close-up image of a mosquito, which is the organism that transmits the pathogen.
- Host:** Represented by a photograph of a young child, who is the individual that the pathogen infects.

A central triangle connects these three elements, signifying their interrelationship in the disease cycle.

CDC Stock photos

Slide 46

Brainstorming (5 min)

What are the main malaria control interventions?

Slide 47

Malaria Control Interventions


•Seven primary malaria control interventions


1. Case management (CM)
2. Intermittent preventive treatment in pregnancy (IPTp)
3. Long-lasting insecticidal Nets (LLIN)
4. Indoor residual spraying of insecticide (IRS)
5. Monitoring and Evaluation
6. Epidemic preparedness and response (EPR)
7. Advocacy, communication and social mobilization (ACSM)


Slide 48


Activity (3 min)

What malaria control intervention is shown in each photo?

#1

#2





#3

#4

48
CDC Stock Photos CDC Stock Photos

Slide 49

Answers

<p>#1 Case management (ACTs)</p>  <p>#3 Indoor residual spraying (IRS)</p>  <p>CDC Stock Photos</p>	<p>#2 Long-lasting insecticide-treated bed nets (LLINs)</p>  <p>PSI</p> <p>#4 Intermittent preventive treatment in pregnancy (IPTp)</p>  <p>CDC Stock Photos</p>
---	--

49

Slide 50

Malaria Case Management (1)

- Early recognition of malaria
- Diagnostic testing
- Use of effective antimalarial medication
- Prompt treatment of uncomplicated illness
- Recognition and treatment of severe / complicated illness
- Appropriate in all epidemiological zones

Slide 51

Malaria Case Management (2)

Consists of two primary components

1. All suspected malaria cases should be tested
 - Microscopy or
 - Rapid diagnostic test (RDT)
2. All confirmed malaria cases should be treated with artemisinin-based combination therapy (ACT)
 - Artemether-lumefantrine (AL) – 1st line
 - Dihydroartemisinin-piperaquine – 2nd line




❖ Except women in 1st trimester of pregnancy

- Quinine – recommended

Slide 52

Malaria Case Management (3)

- A full 3-day course with an ACT is required
 - Ensures >90% reduction in parasitemia
- Decreases the "pool" of persons with parasites who can transmit to mosquitos
- Therefore, case management prevents secondary cases of malaria



CDC Stock photos

Slide 53

Intermittent Preventive Treatment in Pregnancy (IPTp)

- Appropriate only in endemic areas
- All pregnant women should receive sulfadoxine-pyrimethamine (SP)
 - At each antenatal care visit after quickening (doses at least 4 weeks apart)
 - Prevents maternal anemia, placental malaria
 - Prevents infant low-birth weight, premature delivery and deaths

Slide 54


Long-lasting Insecticidal Nets (LLINs)

- In endemic and epidemic-prone areas
- Initially, all pregnant women & children <5 years
- Now, all persons in household
- Universal coverage = 1 net per 2 persons in household via mass net distributions
- Protects persons sleeping under the LLIN
 - Decreases number of persons infected with malaria parasites
- Kills mosquitos and thus reduces transmission intensity

Slide 55

Indoor Residual Spraying with Insecticide (IRS)

- In endemic and epidemic-prone areas
- Optimal IRS application is before the rainy season
- Augments LLIN usage
- Prevents malaria infections in persons in sprayed households
- Kills mosquitos and thus reduces transmission intensity




Abt Associates

Slide 56

Surveillance

- Appropriate for all epidemiological zones
- Accurate diagnosis and confirmation via testing will improve malaria surveillance data
- Malaria case reporting via health information systems (IDSR and DHIS2)
- Detects changes in malaria cases over time
- Provides data to evaluate malaria control interventions



Maggie Hallahan

Slide 57

Epidemic Preparedness and Response (EPR)

- Appropriate in epidemic-prone and seasonal epidemiological zones
- Requires accurate and timely surveillance data
- Allows prompt implementation of control measures
- Prevents or minimizes malaria morbidity and mortality during epidemics

Slide 58

Advocacy, Communication and Social Mobilization (ACSM)

- Appropriate in all epidemiological zones
- Community awareness of malaria prevention and treatment
 - Decreases testing and treatment delays
 - Increases community utilization of malaria control interventions (CM, LLINs, IRS)
- Prevents or minimizes malaria morbidity and mortality in communities

Slide 59

Summary of Malaria Control Interventions

Epidemiological Zone	CM	IPTp	LLINs	IRS	Surveillance	EPR	ACSM
Endemic - Lake - Coast	X	X	X	X	X		X
Epidemic-prone - Highland	X		X	X	X	X	X
Seasonal, low transmission - Semi-arid - Arid	X				X	X	X
Low risk	X				X		X

Slide 60

Activity: Name at least four malaria control interventions appropriate for each area

1. Endemic areas

- High transmission
- Affects children, pregnant women
- Many asymptomatic carriers

2. Epidemic-prone areas

- Low transmission
- All age groups
- Few asymptomatic carriers

Slide 61

Answers

1. Endemic area	2. Epidemic-prone area
– Case management with RDTs and ACTs	– Case management with RDTs and ACTs
– IPTp	– LLINs for everyone
– LLINs for everyone	– IRS
– IRS	– Surveillance
– Surveillance	– EPR
– ACSM	– ACSM

Slide 62

Questions?






Slide 63

THANK YOU

Slide 1

MODULE 2

MALARIA CASE IDENTIFICATION,
CONFIRMATION AND REPORTING



Slide 2

Objectives

1. Identify/ detect cases of malaria using the standard case definition
2. Describe malaria parasitological diagnostic methods
3. Demonstrate malaria data recording and reporting format using appropriate tools

Slide 3

Unit 1

Identification of Malaria cases

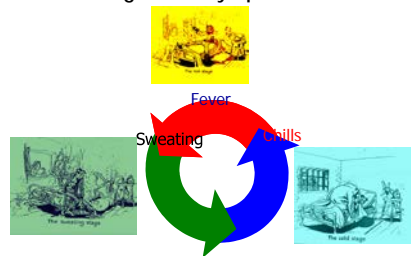
Slide 4

Brain storm (5 mins)

What is the clinical presentation of malaria?

Slide 5

Clinical Signs and Symptoms of Malaria



Slide 6

Common Signs and Symptoms of Uncomplicated Malaria

- Fever
- Chills
- Profuse sweating
- Muscle pains
- Joint pains
- Abdominal pain
- Diarrhoea
- Nausea
- Vomiting
- Irritability
- Refusal to feed
- (Sometimes the symptoms may be non-specific)

Slide 7

Standard Case Definition

- Standard description of a disease
- Or standard set of criteria used to describe if a person has a particular disease
- Standard case definitions are used for reporting by all health workers
- Importance:
 - Easier to follow trends in diseases and recognize outbreaks
 - Data can be compared more accurately from one area to the other
 - Increase the specificity of reporting

Slide 8

Types of Case Definitions

- Surveillance case definition for health staff (Standard Case Definition)
- Case definition for Community Health Workers (Lay Case Definition)

Slide 9

How to use the standard case definition

- Patient comes to consulting room
- Ask about symptoms and duration
- Conduct physical examination and record findings on OPD card
- Make diagnosis based on signs and symptoms

Slide 10

How to use the standard case definition

- Match signs and symptoms with that of case definition
- Record Information about suspected cases in the health facility register and patients card
- Report case based information for immediate notifiable diseases using the IDSR reporting tools
- N/B use the local Lab capacity to diagnose suspected cases

Slide 11

Malaria standard case definition

- **Uncomplicated malaria:** Any person living in area at risk of malaria with fever or history of fever within 24 hours; without signs of severe disease (vital organ dysfunction) is diagnosed clinically as malaria.
- **Confirmed uncomplicated malaria:** Any person with fever or history of fever within 24 hours; and with laboratory confirmation of diagnosis by malaria blood film or other diagnostic test for malaria parasites.

Slide 12

Malaria standard case definition Cont'd

- **Unconfirmed severe malaria:** Any patient living in area at risk of malaria hospitalized with severe febrile disease with accompanying vital organ dysfunction diagnosed clinically
- **Confirmed Severe malaria:** Any patient hospitalized with *P. falciparum* asexual parasitaemia as confirmed by laboratory tests with accompanying symptoms and signs of severe disease (vital organ dysfunction) diagnosed through laboratory.

Slide 13

Brainstorming (5 min)

What are the differential diagnosis of malaria?

Slide 14

Differential diagnosis

- Influenza
- Dengue fever
- Enteric fever
- Gastroenteritis
- Brucellosis
- Hepatitis
- Acute Schistosomiasis (Katayama Fever)
- HIV seroconversion

Slide 15

Unit 2

Case confirmation



Slide 16

Brain storm (5 mins)

1. Rationale of malaria parasitological diagnosis
2. Challenges of confirmatory diagnosis and how to address them.
3. Do clinicians always use lab results to make clinical decision

Slide 17

Rationale for malaria parasitological diagnosis

1. To differentiate malaria cases from other diseases with similar presentations
2. To monitor response to malaria treatment
3. To confirm/ or predict out breaks

Slide 18

Methods of malaria parasitological diagnosis

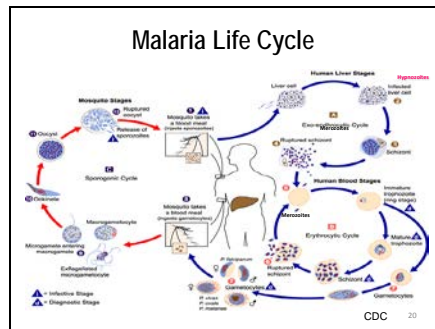
- Visualization
 - Microscopy
 - Qualitative Buffy Coat (QBC)
- Rapid Diagnostic Tests (RDTs)
- Detection of parasite products
 - Enzyme linked immunosorbent assay (ELISA)
 - Polymerase Chain Reaction (PCR)

Slide 19

Microscopic Diagnosis of Malaria

- It is the 'Gold standard' for detection of malaria parasitaemia
- Has sensitivity >90% if performed well
- Used to confirm diagnosis, monitor treatment outcome, confirm epidemics and in clinical trials of drugs and vaccines

Slide 20



Slide 21

Procedure

- i. Specimen collection
- ii. Specimen processing
- iii. Blood slide examination
- iv. Blood slide reporting
- v. Results interpretation

Slide 22

Specimen Collection

- Label the patient identity and date on slide
- Disinfect the puncture site
- Prick the finger firmly with a sterile lancet
- Wipe the first drop of blood
- Collect a drop of blood on a glass slide
- Make a thin and thick smear

Slide 23

Specimen Collection Cont'd

Thick Smear

- Pre-cleaned/Washed grease free slides
- Proper labeling
- Correct amount of blood (5-15ul)
- Right diameter (10-15mm)
- Right thickness (0.05-0.09mm)

Slide 24

Specimen Collection Cont'd

Thin Smears

- Correct amount of blood (2-4ul)
- Smooth spreader
- Correct angle (45°)
- Right length (25-30mm)

Slide 25

Specimen Processing

- i. Fix thin film with methanol
- ii. Allow to air dry
- iii. stain appropriately
- iv. Wash, let dry and examine

Slide 26

Examination and Reporting

- **Examine using the x100, oil immersion objective**
 - **Thick film:** if parasites present, count trophozoites against WBCs until 200 WBCs are counted
 - If no parasites are seen, examine 100 high power fields
 - **Thin film:** Species identification

Slide 27

Reporting/ Interpretation

- *Report on parasite seen, developmental stage and species*
- Parasite density (parasites/200 WBC or per microlitre of blood)
- No of parasites counted $\times 8000$ / WBC counted = parasites / μ l
 - e.g. $35/200 \times 8000$ per μ l gives you 1400 parasites per microlitre of blood

Slide 28

Quality Assurance for Microscopy

Quality Assurance (QA)

is a broad spectrum of plans, policies and procedures which together ensure that a system conforms to established technical requirements

Quality Control (QC)

deals with the techniques and procedures that monitor performance

Slide 29

Malaria Rapid Diagnostic Tests (mRDT)

• Test Principle

- The test contains a strip with antibodies against malaria parasites
- If malaria parasite antigens are present two bands are formed: a control band and a positive band
- In the absence of malaria parasite antigens, only the control band is formed

Slide 30

Kit Format

- Dipsticks
- Cassettes
- Card

Slide 31

Materials required to Perform RDTs

- RDT kit. (Test cassette, Buffer, Blood collecting device)
- Sterile Lancet
- Alcohol Swab
- Pencil/ Pen for Labeling
- Gloves
- Sharps Container
- Waste Disposal container
- Timer/ Clock
- Instruction Manual for the specific RDT/SOP
- Dry cotton wool.

Slide 32



Slide 33

Preparing to Perform the Tests

1. Gather the necessary materials in the testing area.
2. Check the expiry date at the back of the test package. If the test kit has expired use another test.
3. Ensure the RDT packaging is not damaged by squeezing gently and feel/listen for air leakage.
NOTE: If the foil packaging is damaged, use another test kit.
4. Explain to the patient what the test is for and procedure

Slide 34

Preparing to Perform the Tests Cont'd

5. Open the package tearing along the nick and look for the following : - a.) Desiccant b). Cassette and c). Dropper
- 6 Remove the cassette from the foil packaging and label it with patient particulars and reading time
7. Wear Gloves
8. Disinfect the puncture site with an alcohol swab or appropriate disinfectant.

Slide 35

Finger Prick

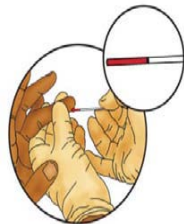
- Make a gentle prick with a sterile lancet at the disinfected site.
- By applying gentle pressure to the finger express the first drop of blood and wipe it away with a dry piece of cotton wool. Make sure no strands of cotton remain on the finger to contaminate blood.
- Apply gentle pressure to the finger until a new blood drop appears.



Slide 36

RDT Test Procedure


1. Using the blood collection device (Pipette or Capillary tube) provided in the RDT kit, gently immerse the open end in the blood drop. Collect the required volume of blood as per manufacturer's instructions.



Slide 37

RDT Test Procedure

2. Transfer the collected blood to the sample well (as indicated on the RDT cassette).




Slide 38

RDT Test Procedure

3. Place dry cotton wool over the puncture site to stop the bleeding.

4. Holding the buffer bottle vertically, add the recommended number of drops of buffer into the buffer well.



Slide 39

Results of the RDT

Time the test as recommended by the manufacturer.

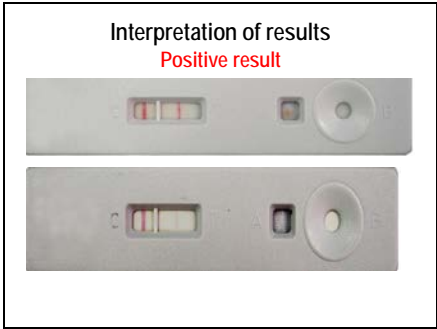
NOTE: Do not read the results before or after the set time.

- One band (control) – negative
- Two bands – positive
- No control line – invalid test

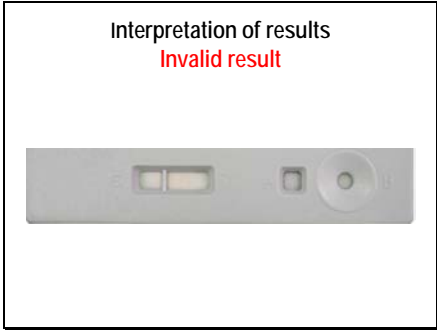
Slide 40



Slide 41



Slide 42



Slide 43

Reporting

- Report the results as "RDT Negative" or "RDT Positive" or "RDT Invalid" (in which case the RDT should be repeated).

Clinic/OPD Reporting

- If the RDT is performed in the clinic, outpatient department or in the wards, the result, even if it is negative, should be reported on
 - The appropriate patient card/form
 - As well as in the OPD register, RDT Daily activity register and any other register.

Slide 44

Advantages of RDTs

- Simple and fast
- Can be performed anywhere
- Portable
- Kit components easily packed

Slide 45

Discussion (5 min)

What are the strengths of each parasitological method as we scale up Testing? Microscopy vs. RDTs

Slide 46

Microscopy vs. mRDT	
Microscopy	RDT
Technical	Simple and fast
Needs longer training	Shorter training
Needs Equipment	No equipment
Used in Epidemic confirmation	Used in epidemic confirmation
Used in treatment treatment monitoring	Can not be used for treatment monitoring

Slide 47

Quality Assurance & Sources of Common Error
<ul style="list-style-type: none">• Read the manufacturer's instructions prior to performing the test.• Follow the test procedure, precautions and interpretation of results for this test. (Use of SOPs and Job aids)• Use the correct amount of blood and buffer. (Incorrect amount of Buffer and blood may lead to inaccurate results)• Read the test at the recommended time.

Slide 48

Quality Assurance & Sources of Common Error
<ul style="list-style-type: none">• Check expiry date of the test kit before use.• Only open the foil packaging and remove the RDT immediately before performing the test. If preparation is delayed after opening the packaging, the RDT may be damaged by humidity and results may not be accurate.• Label correctly the patient details on the test cassette to avoid mix ups.• Proper storage conditions as per manufacturer's instructions

Slide 49

Biohazard, Safety and Waste Management

- Protect yourself and others
 - Laboratory coat
 - Gloves
 - Wash hands
 - Disinfect working bench

Slide 50

Biohazard, Safety and Waste Management Cont'd

- Segregate waste material as follows
 - Sharps
 - Collect in puncture-proof container
 - Pathological hazardous waste
 - Collect in hazardous waste bags (Red bag)
 - Non-pathological waste (Black)
 - Pour in sink, latrine, or waste pit
- All bio-hazardous waste should be incinerated

Slide 51

Practicum (30 min)

Practical session by carrying out RDT test performance

Slide 52

UNIT 3

Reporting

Slide 53

Background on Reporting

- Every level of the health system has a role in carrying out ongoing surveillance for priority diseases, conditions and events.
- If a disease is identified at a local level, for example, but the information is not reported to the next level, an opportunity for timely response is lost.

Slide 54

Background on Reporting

- What is reported to each level and how often is usually guided by national policy .It can be immediately, weekly, monthly, or quarterly.
- How the information is reported depends on the capacity in your area. For example, reporting may be done by electronic methods such as email or other electronic transmission, or cell phone SMS reporting.

Slide 55

Brainstorm (5 mins)

1. Which tools are used for recording malaria cases?
2. How often are malaria cases reported?
3. Which tools are used in malaria reporting?

Slide 56

Case recording

- Tools for recording
 - OPD cards
 - Registers (MOH 705A, MOH 705B, Lab registers)
 - Tally sheets
- In many health facilities, more than one person is responsible for recording information about patients seen in the facility.
- Example
 - The clinician records the patient's name and diagnosis in a clinic register.
 - Later in the day, a nurse tallies the number of cases and deaths seen in an outpatient service.
 - A ward nurse tallies the number of hospitalized cases.
- Then: Each week and month
 - A data clerk will calculate summaries for all the diseases and records the totals in a standard form.

Slide 57

Reporting tools

- Health Facility Line listing Form (MOH 503)
- Monthly Surveillance Report Form (MOH 504)
- Epidemic Monitoring Form (MOH 505)
- Outpatient monthly summary for <5 years (MOH 705A)
- Outpatient monthly summary for >5 years (MOH 705B)
- Lab test data summary report form (MOH 706)

Slide 58

Reporting requirements for malaria

- Weekly (IDSR)
 - Epidemiological week starts on Monday and ends on Sunday
 - The total number of cases both clinical and confirmed and deaths seen in a particular week are reported
 - The cases are summarized in the facility and sent to district by Monday which are then collated and entered on the e-idsr system by Wednesday

Slide 59

Reporting requirements for malaria

- Monthly reporting (HMIS)
 - the total number of cases and deaths seen in a particular month are reported through the DHIS 2 system

Slide 60

Group Work (30 min)


- Demonstration of how to fill and upload malaria data on the e-IDSR & DHIS2 systems





Slide 1


Module 3


MALARIA SURVEILLANCE DATA
MANAGEMENT


Republic of Kenya
Ministry of Health


Division of Malaria
Control


World Health Organization


MEASURE
Evaluation


USAID
President's Malaria Initiative

Slide 2

Objectives

1. Identify different types of data sources, describe the process involved in the Malaria surveillance data collection, processing and flow using the existing MOH tools.
2. Perform data quality checks to review the reports.
3. Perform simple data analysis tasks, present, interpret and share the results.
4. Promote data demand and use for policy and program management

Slide 3

Unit 1

Data collection, processing and flow

Slide 4

Definitions

- What is:
 - Data collection?
 - Data Source?
 - Data processing?
 - Data flow?

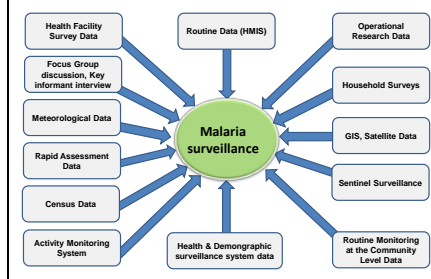
Slide 5

Types of data

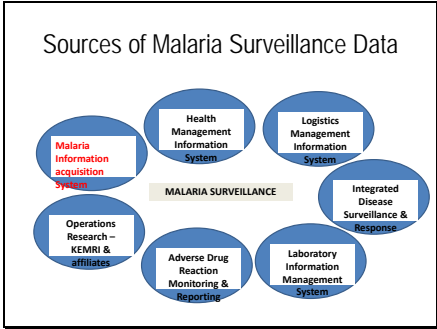
- Survey data
- Surveillance data
- Service data
- Routine data
- Primary data
- Secondary data

Slide 6

Potential Data Sources for Malaria



Slide 7



Slide 8

Group Activity:

- Exercise on Identifying MOH Data Management Tools

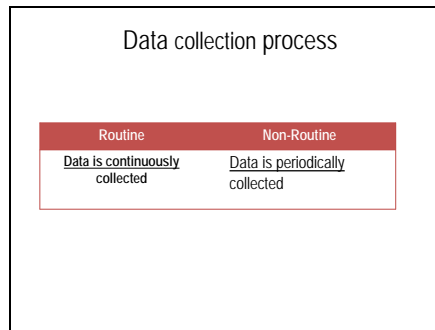
Module 5.1: Data Collection, Collation, & Aggregation 8

Slide 9

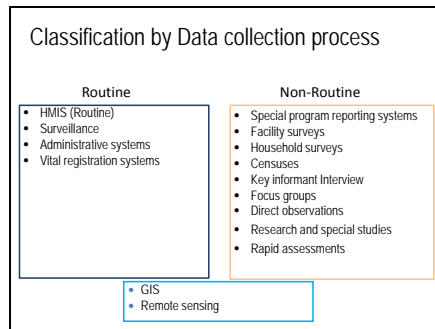
Health Facility Data Sources

- Base Registers
 - MOH 204A Outpatient < 5 yrs Register
 - MOH 204B Outpatient >= 5 yrs Register
 - MOH 240 Lab Register
 - MOH 405 ANC Register
 - MOH 511 CWC
 - MOH 301 in-patient register
- Summaries and Frequencies
 - MOH 705A-OP Summary Sheet Under 5yrs (Daily)
 - MOH 705B-OP Summary Sheet Over 5yrs (Daily)
 - MOH 711A-Facility Integrated (Monthly)
 - MOH 715-Health Facility template (Monthly)
 - MOH 105-Facility Service Delivery template (Monthly)
 - MOH 711A-Facility Integrated (Monthly)

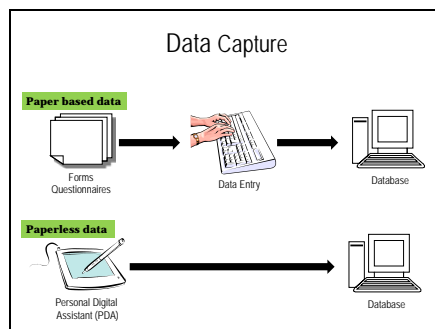
Slide 10



Slide 11



Slide 12



[illegible][illegible]

HMIS DATA FLOW STRUCTURE

```

graph TD
    MOH-DHIS2[MOH- DHIS2] --> DOMC[DOMC]
    DOMC --> MME[MME Manager & MAE Program Officers]
    MOH-DHIS2 --> PHRIO[PHRIO's office]
    PHRIO --> District[District level  
(Service Delivery -MOH 105)]
    District --> DHISD[DHISD]
    District --> Health[Health facility level  
(MOH 705 A/B)]
    Health --> CHS[CHS Office/Chargé de Bureau & Data Clerks]
    Health --> OPD[OPD Registrar -MOH 204A/B]
    Health --> ANC[ANC Registrar -MOH 605]
    Health --> PNC[PNC Registrar -MOH 301]
    OPD --> Health
    ANC --> Health
    PNC --> Health
    
```

The flowchart illustrates the HMIS Data Flow Structure. At the top is **MOH- DHIS2**, which connects to **DOMC** (with responsibility for MME Manager & MAE Program Officers) and **PHRIO's office**. **PHRIO's office** reports to the **District level (Service Delivery -MOH 105)**. The **District level** reports to **DHISD** and the **Health facility level (MOH 705 A/B)**. The **Health facility level** reports to **CHS Office/Chargé de Bureau & Data Clerks** and three registrars: **OPD Registrar -MOH 204A/B**, **ANC Registrar -MOH 605**, and **PNC Registrar -MOH 301**. Feedback loops exist from each registrar back to the Health facility level. Reporting timelines are specified: 15th of the following month for PHRIO to District, and 5th of the following month for District to Health facility level.

The flowchart illustrates the Kenya IDSR Data flow. At the top, three boxes labeled 'Health Facilities' are connected by dashed lines to a central box labeled 'District (IDSR) Collate data and report'. To the right of this central box, a box labeled 'Reporting predominantly via phone (SMS)' has a dashed arrow pointing to the central box. Below the central box is a box labeled 'Provincial (IDSR) Receive and review reports', connected by a solid double-headed arrow. To the right of this box, another box labeled 'Reporting predominantly via phone (SMS) or email' has a dashed arrow pointing to the provincial box. Below the provincial box is a box labeled 'National level (IDSR - Provincial focal person) Receive data, update database, analyse and provide feedbacks', connected by a solid double-headed arrow. At the bottom is a box labeled 'WHO and other partners', connected by a solid double-headed arrow to the national level box. Dashed lines also connect the 'Health Facilities' boxes to the 'National level' box.

Slide 16

Purpose of Understanding Data Flow

- Helps us better understand our role in the health information system and the importance of collecting data
- Identify opportunities for improving data collection and analysis, increasing availability, and ensuring data use

Slide 17

Unit 2

Data quality

Slide 18

What is Data Quality?

Slide 19

Data quality is defined as “the totality of features and characteristics of data set that bear on its ability to meet the needs that result from the intended use of the data.”

Slide 20

Elements of data Quality

- Timeliness
- Completeness
- Validity
- Accuracy
- Precision
- Reliability
- Integrity

Slide 21

How do you improve data quality?

Slide 22

Improving data quality

- Check completeness of the data
- Check consistency- compare variables
- Check plausibility (value with acceptable range)
- Check for duplicates
- Check for outlier (run basic freq, mean)

Slide 23

Unit 3

Data analysis, presentation and interpretation

Slide 24

What is Data Analysis?

Slide 25

Data Analysis

- The process of **understanding** and **explaining** what findings actually mean. Turning raw data into useful information
- Provide answers to questions being asked at a program site or research questions being studied
- The greatest amount and best quality data mean nothing if not properly analyzed, or, if not analyzed at all

Slide 26

What is Data Analysis? *Cont'd*

Analysis is looking at the data in light of the questions you need to answer



How would you analyze data to determine, "Is my program meeting its objectives?"

Slide 27

Data Analysis Tools

- Examples of data analysis tools include:
 - Pen, pencil
 - Paper
 - Calculators
 - Spreadsheet (e.g., Excel®)
 - Database (e.g., Access®)
 - Epidemiological information
 - Statistical software (e.g., SPSS, STATA)

Slide 28

What is the importance of Data Analysis and interpretation?

- Know the size of the health problem
- Monitor trends and take prompt action
- Identify the cause of the problem
- Monitor progress of public health programs

28

Slide 29

Statistical Measures

- Measure of central tendency
 - Mean
 - Median
 - Mode
- Measure of variation
 - Range
 - Variance and standard deviation
 - Inter-quartile range
 - Proportion, Percentage

Slide 30

Mean

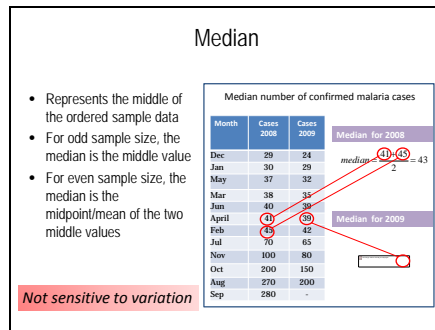
Sum of the values divided by the number of cases

Also called *average*

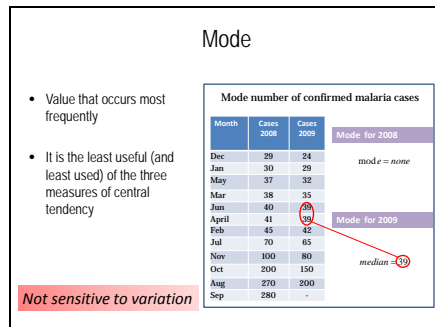
Very sensitive to variation

Average number of confirmed malaria cases per month		
Month	Cases 2008	Total number of cases
Jan	30	1,180
Feb	45	
Mar	38	
Apr	41	
May	37	Number of observations
Jun	40	12
Jul	70	
Aug	270	
Sep	280	
Oct	200	Mean number of cases
Nov	100	$\frac{1,180}{12} = 98.2$
Dec	29	

Slide 31



Slide 32



Slide 33

Practice Calculations

- What is the mode, mean and median Hb for the following set of observations?
11.5, 10.8, 12.5, 14.1, 8.3, 9.2, 7.9, 10.6
- Answers:
 - Mean = 10.61
 - Median = 10.7
- Would you use Mean or Median?
- Answer: Median
- Use Median when you have a large variation between high and low numbers
- Use Mean when there is not a huge variation between the values

Slide 34

Proportion

- A ratio in which all individuals in the numerator are also in the denominator
- Example: If a clinic has 12 female clients and 8 males clients, then the proportion of male clients is 8/20 or 2/5

MMMM
MMMM

FFFF
FFFF
FFFF

Slide 35

Percentage

- A way to express a proportion
- Proportion multiplied by 100
- Example: Males comprise 2/5 of the clients or, 40% of the clients are male (0.40 x 100)

Important to know: What is the whole? An orange?
An apple? All clients? All clients on with a fever?

Helps us standardize so that we are able to compare data across facilities, regions, countries

Slide 36

Rate

- A quantity measured with respect to another measured quantity
- Number of cases that occur over a given time period divided population at risk in the same time period

(Under five mortality rate)
Probability of Dying Under Age Five
per 1,000 Live Births

Nation	Under five mortality rate per 1,000 live births in 2008
France	4
Ghana	76
Sierra Leone	194
Afghanistan	257

Source: UNICEF: Statistics and Monitoring by Country

Slide 37

Data presentation

Slide 38

Effective presentation

- Clear
- Concise
- Practical
- Actionable
- Attractive

Slide 39





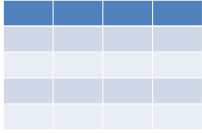
Effective presentation

- For all communication formats it is important to ensure that there is:
 - Consistency
 - Font, Colors, Punctuation, Terminology, Line/ Paragraph Spacing
 - An appropriate amount of information
 - Less is more

Slide 40

Data Presentation

- Tables
 - Rows
 - Columns
- Figures



40

Slide 41

Summarizing data

- Tables
 - Simplest way to summarize data
 - Data is presented as absolute numbers or percentages
- Charts, maps and graphs
 - Visual representation of data
 - Usually data is presented using percentages

Slide 42

Tables and graphs

- Have titles and axis labels
- Tables and graphs are used to:
 - Convey a message
 - Stimulate thinking
 - Portray trends, relationships and comparisons
- The most informative graphs are simple and self-explanatory

Slide 43

Choosing a Title

- A title should express
 - Who
 - What
 - When
 - Where

Slide 44

Tables: Frequency distribution

Year	Number of cases
2005	4 216 531
2006	3 262 931
2007	3 319 339
2008	5 338 008
2009	7 545 541
2010	9 181 224
2011	8 926 058
2012	9 610 691

Slide 45

Tables: Relative frequency

Percent contribution of reported malaria cases by year between 2005 and 2012

Year	Number of malaria cases (n)	Relative frequency (%)
2005	4 216 531	8
2006	3 262 931	6
2007	3 319 339	7
2008	5 338 008	10
2009	7 545 541	15
2010	9 181 224	18
2011	8 926 058	17
2012	9 610 691	19
Total	51 400 323	100.0

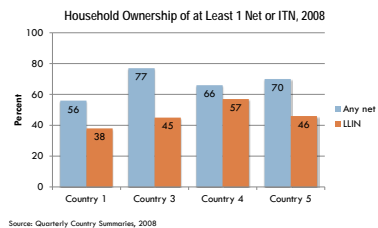
Slide 46

Use the right type of graphic

- Charts and graphs
 - **Bar chart:** comparisons, categories of data
 - **Histogram:** represents relative frequency of continuous data
 - **Line graph:** display trends over time, continuous data (ex. cases per month)
 - **Pie chart:** show percentages or proportional share

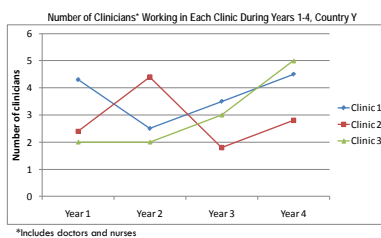
Slide 47

Bar Chart

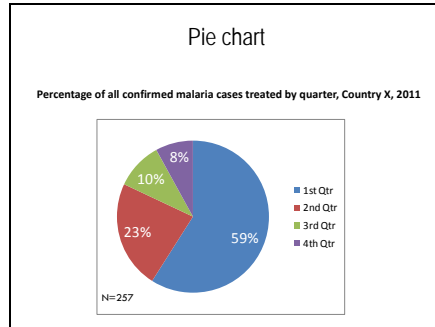


Slide 48

Line graph



Slide 49



Slide 50

Exercise:
How should you present...

1. Prevalence of malaria in 3 countries over a 30 year period?
2. Data comparing prevalence of malaria in 10 different countries?
3. Data on reasons why individuals not using ITNs (out of all individuals surveyed who own an ITN and are not using it)?
4. Distribution of patients tested for malaria by parasite density

Slide 51

Data Interpretation

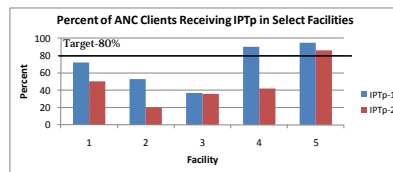
Slide 52

Analysis vs. Interpretation

- **Analysis:** describing data with tables, graphs, or narrative; transforming data into information
- **Interpretation:** adding meaning to information by making connections and comparisons and by exploring causes and consequences

Slide 53

Are facilities reaching coverage targets?



Slide 54

Interpreting Data

- Does the indicator meet the target?
- What is the programmatic relevance of the finding?
- What are the potential reasons for the finding?
- What other data should be reviewed to understand the finding (triangulation)?
- How does it compare? (trends, group differences)
- Conduct further analysis

Slide 55

Additional Questions

- Which facility is performing better/worse than expected?
- What is the trend over time for these facilities?
- How would you assess each facility's performance based on the data?
- What other data or information should you consider in providing recommendations or guidance to the facilities?

Slide 56

Unit 4

Data demand and use

Slide 57

Definitions

- Data Demand
- Data Use
- Decisions

Slide 58

Why Data Demand and Use

- Increased financial investments for service delivery
- Increased accountability requirements
- Improved national HMIS
- Increased demand for evaluation and other research

Slide 59

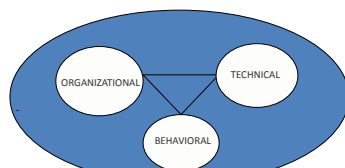
Data Demand & Use



Source: MEASURE Evaluation

Slide 60

What Determines Data Demand & Use?



* Based on PRISM analytical framework (Lafont, Fields et al. (2005). The PRISM: An analytical framework for understanding performance of health information systems in developing countries. MEASURE Evaluation).

Slide 61

Group Participation

What barriers have you faced to using or getting others to use data and information?

Slide 62

Barriers to Data Demand and Use

Technical constraints

- Technical skills
- Availability of computers
- Data system design
- Definition of indicators
- Lack of data quality assurance protocols

Slide 63

Barriers to Data Demand and Use Cont'd

Organizational constraints

- Structural – roads, telecommunications
- Organizational – clarity of roles, support, flow of information
- Political interference

Slide 64

Barriers to Data Demand and Use Cont'd

Individual constraints

- Decision-maker attitudes
- Staff motivation
- Lack of "data use culture"

Slide 65

Group Participation

What challenges have you faced trying to use data and information?

Slide 66

Challenges

- Integrated HMIS still not fully functioning
- Little or no communication between data producers and data users
- Low capacity to collect, analyze, & interpret data
- Limited or no culture of data use
- Data collection and use not a priority

Slide 67

Importance of Feedback

- Information needs to be shared:
 - At timely and regular intervals
 - Within, between, up, and down
- Paves path between data collectors and users at all levels of the health system

Slide 68

Importance of Feedback Cont'd

- Leads to greater appreciation of data:
 - Improved data quality
 - Influences collection of appropriate data
- Important element of management and supervision:
 - Creates opportunity to monitor & improve program services
 - Incentive for staff

Slide 69

Types of Feedback

Type of Feedback	Example	Audience
Written	Tables of monthly reports	Staff, Managers
	Short program reports	Staff, Managers
	Comparison tables by facility	Staff, Managers
	Graphs	Staff
	Quarterly, biannual, and annual reports	Staff, Managers, Community, NGOs
	Standard reports	Staff, Managers
	Special reports	Policymakers, NGOs
Oral	Staff assessments	Staff
	Staff appraisals	Staff

Slide 70

Examples of Feedback

- Sharing information within a facility or organization
- Sharing aggregated service provision data from facilities within a district or between provinces
- Meetings between facility and supervising agency to review and discuss information
- Meetings between donor and NGO to review information and discuss challenges and opportunities

Slide 71

Beware of information overload!








Slide 72

THANK YOU!

Slide 1

Module 4

Core Malaria Surveillance Graphs



Slide 2

Objectives

1. Define the malaria surveillance indicators, data sources and targets
2. List the Core Malaria Surveillance Graphs based on WHO requirements
3. Describe malaria surveillance graphs/dashboards
4. Demonstrate how the malaria core surveillance graphs are generated and update the summary tools

Slide 3

Unit 1

Malaria Surveillance Indicators, Targets and Data sources

Slide 4

Review of surveillance indicators

- Indicators measured monthly
 - OP confirmed malaria cases
 - Clinical Malaria cases
 - OP malaria TPR
 - IP malaria cases
 - IP malaria deaths
 - Diagnostics: %OP suspected tested
 - ITN routine distribution
 - IPTp
 - Stockouts
 - Completeness of reports

Slide 5

Review of surveillance indicators

- Indicators measured annually
 - IRS coverage (population)
 - IRS coverage (households)

Slide 6

Surveillance Indicators

Indicator	Numerator, Denominator	Targets	Comments
Indicators measured monthly			
1. Outpatient confirmed malaria cases	Numerator – Number of outpatient confirmed malaria cases (by microscopy or RDT) reported by health facilities per year Denominator for rate – Resident population by age >5 years, all ages per 1000 people resident in areas at risk of malaria	Case rate trend: → 50% reduction by 2010 → 75% reduction by 2015 Rate: → <1 confirmed case per 1000 people indicates excellent control	Rate of <1 confirmed case per 1000 people indicates readiness for elimination phase
2. Outpatient malaria TPR	Numerator – Number of outpatient laboratory-confirmed malaria cases Denominator – Total number of outpatient suspected malaria cases tested = 100	TPR trend: → 50% reduction by 2010 → 75% reduction by 2015 Annual TPR: → 10–20% = intermediate control → 5–10% = good control → <5% = excellent control	Annual rate should be used, not just the rate during the peak season → <5% in peak season indicates readiness for elimination phase
3. Inpatient malaria cases	Numerator – Cases (confirmed and unconfirmed) with a primary diagnosis of malaria at discharge (and not admission) Denominator for rate – Resident population by age >5, all ages per 1000 people resident in areas at risk of malaria	Trend: → 50% reduction by 2010 → 75% reduction by 2015	

Slide 7

Surveillance Indicators Cont'd			
4. Inpatient malaria deaths	Numerator = Deaths with a primary diagnosis of malaria at discharge Denominator for rate = Mid-year resident population by age <5 all ages (per 1000 people resident in areas at risk of malaria)	Target +150% reduction by 2010 +75% reduction by 2015 (Reduction of malaria deaths by 2015)	
5. Diagnosis rate = percentage of outpatient/suspected malaria cases that undergo laboratory diagnosis	Numerator = Number of outpatient/suspected malaria cases that received laboratory examination for malaria (microscopy or RDT) Denominator = Number of outpatient/suspected malaria cases × 100	90%	
6. Treatment (ACT) = percentage of outpatient malaria cases that received appropriate antimalarial treatment according to national policy	Numerator = Number of malaria cases receiving appropriate antimalarial treatment at health facility Denominator = Number of outpatient malaria cases expected to be treated at health facility with appropriate antimalarial medicines (all those with a diagnosis of malaria) × 100	100%	

Slide 8

Surveillance Indicators Cont'd			
Indicator	Numerator, denominator	Targets	Comments
Indicators measured monthly			
7. ITN = routine ITN distribution to populations at high risk (pregnant women)	Numerator = Number of ITNs distributed or delivered to target population = pregnant women attending ANC Denominator = Total number of pregnant women attending an ANC for the first time	≥95%	
8. IPT = IPT in pregnant women	Numerator = Number of pregnant women receiving second dose of IPT Denominator = Number of pregnant women with at least one ANC visit	≥80%	
9. Stock-outs = percentage of health facilities without stock-outs of first-line antimalarial medicines, microscopy nets and diagnostics, by month?	Numerator = Number of health facilities in areas at risk of malaria, without stock-outs of first-line antimalarial medicines (according to national policy), ITNs and RDT in a month Denominator = Number of reporting health facilities in the same areas at risk of malaria × 100	100%	

Slide 9

Surveillance Indicators Cont'd			
10. Completeness of monthly health-facility reports on surveillance and logistics?	Numerator = Number of health facility monthly reports received on surveillance and logistics, by month Denominator = Number of health facility reports expected each month	100%	

Slide 10

Malaria Surveillance Data Sources		
#	Indicator Numerator	Data Source Register(s)
1	Total suspected malaria cases	MoH 204 A/B
2	Number of Malaria cases tested (Microscopy)	MoH 240 Lab Register
3	Number of outpatient confirmed malaria cases (Microscopy)	MoH 240 Lab Register
4	Number of Malaria cases tested (RDT)	RDT Facility Registers

Slide 11

Malaria Surveillance Data Sources Cont'd		
#	Indicator Numerator	Data Source Register(s)
5	Number of outpatient confirmed Malaria cases (RDT)	RDT Facility Registers
6	Total Number of outpatient confirmed Malaria cases (Microscopy + RDT)	MoH 240 and RDT Facility Registers
7	Total number of confirmed Malaria cases treated with ACTs	AL registers
8	Total suspected malaria cases treated with ACTs	AL registers

Slide 12

Malaria Surveillance Data Sources Cont'd		
#	Indicator Numerator	Data Source Register(s)
9	No of Nets distributed to under 1 yrs	MOH 511 - CWC
10	Nets distributed to pregnant women	MoH 405 ANC Register
11	Inpatient Malaria cases (confirmed with primary diagnosis of malaria at discharge)	MoH 301 MoH 268 (Dist. Hosp.)
12	Inpatient malaria cases (confirmed & unconfirmed with primary diagnosis of malaria at discharge)	MoH 301 MoH 268 (Dist. Hosp.)
13	Total inpatient malaria deaths (with primary diagnosis as malaria)	MoH 301 MoH 268 (Dist. Hosp.)
14	IPT 1 & IPT 2	MoH 405

Slide 13

Unit 2

Introduction to WHO core Malaria Surveillance graphs

Slide 14

WHO Core Malaria Surveillance Graphs

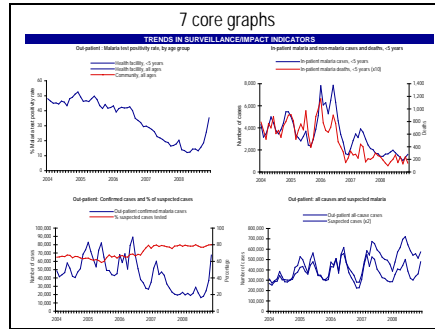
- The core graphs are grouped into two categories – surveillance (four graphs) and logistics (three graphs), as follows:
- **Surveillance graphs:**
 - outpatient malaria TPR in children under 5 years of age and all ages
 - inpatient malaria cases and deaths in children under 5 years of age (double-axis graph);
 - outpatient confirmed malaria cases and percentage of suspected malaria cases tested with parasite-based test (double-axis graph); and
 - outpatient all-cause cases and suspected malaria cases, all ages (double-axis graph).

Slide 15

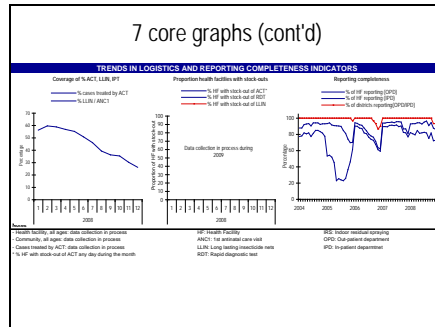
WHO Core Malaria Surveillance Graphs
Cont'd

- **Logistics and completeness of reporting graphs:**
 - percentage coverage with patients treated with ACT (of number expected to be
 - treated according to national policy), and of ANC clients receiving ITN or IPT2 (i.e. second dose of IPT)
 - percentage of health facilities without stock-outs of ACT, RDT and LLIN; and
 - percentage of health facilities and districts that reported

Slide 16



Slide 17

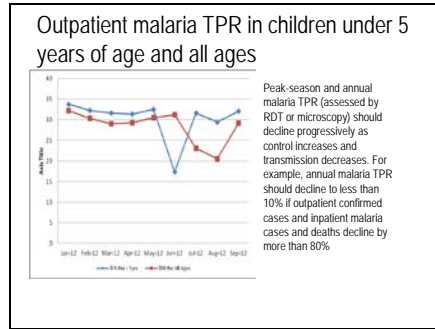


Slide 18

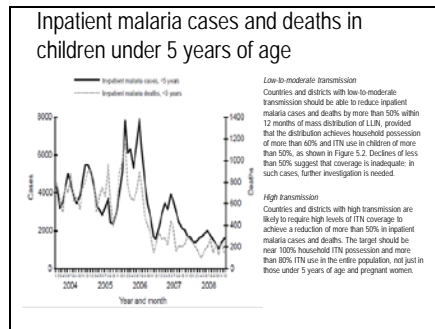
Unit 3

Malaria Surveillance Graphs and Interpretation

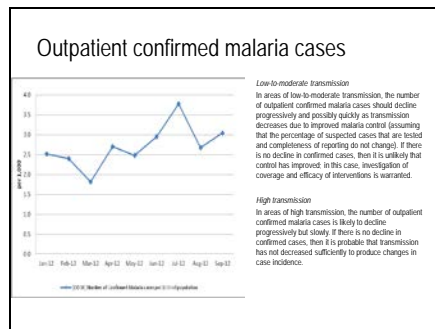
Slide 19



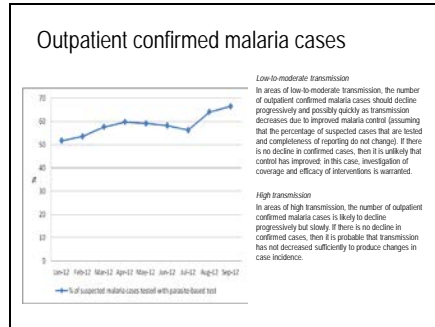
Slide 20



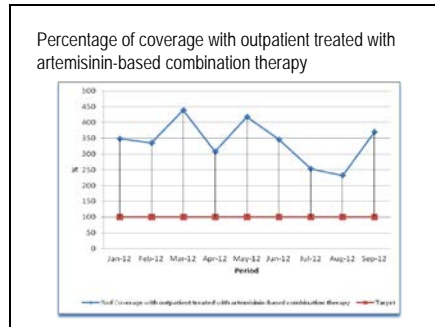
Slide 21



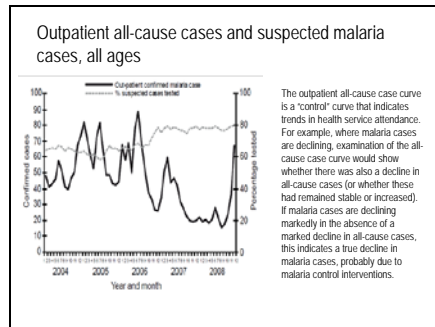
Slide 22



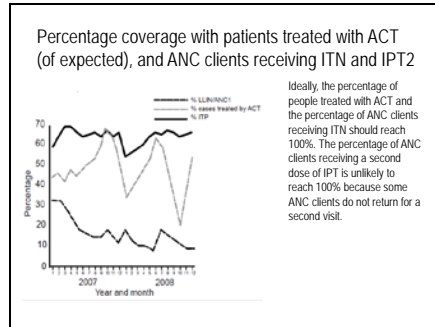
Slide 23



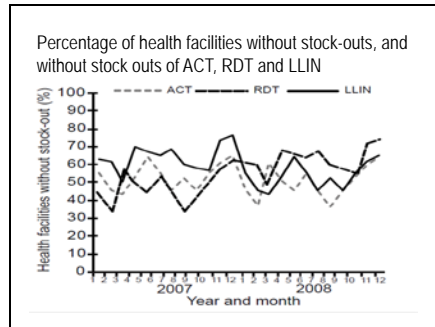
Slide 24



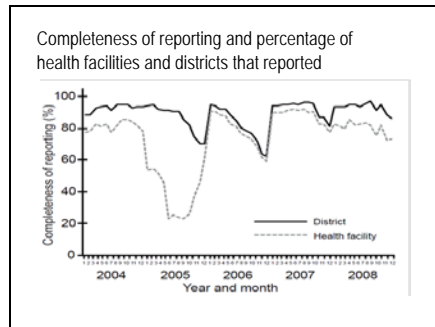
Slide 25



Slide 26



Slide 27



Slide 28

Unit 4

Malaria Surveillance Summary Tool

Slide 29

Filling the Electronic Tool

The Electronic DMCC Tool is an excel workbook with 14 worksheets containing:

- Facility List
- Jan-Dec worksheets with the indicators for each facility
- District Indicator Summary worksheet

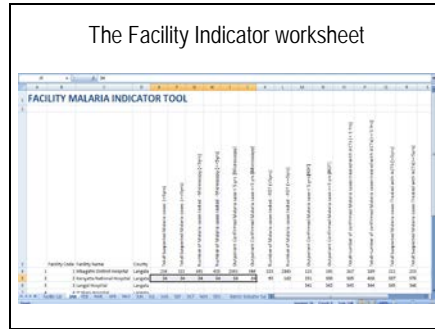
Some parts of the worksheet are protected to avoid accidental deletions

Slide 30

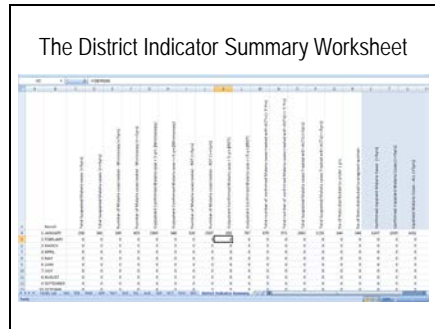
The Facility List worksheet

	Facility Code	Facility Name	County
1	1	1 Mbagathi District Hospital	Langata
2	2	2 Kenyatta National Hospital	Langata
3	3	3 Langat Hospital	Langata
4	4	4 St Mary Hospital	Langata
5	5		
6	6		
7	7		
8	8		

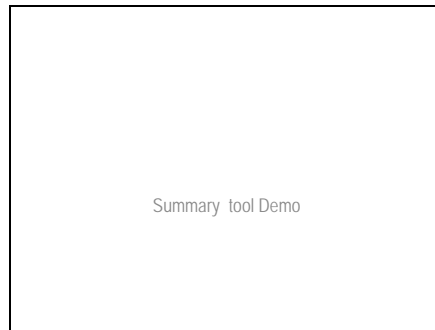
Slide 31



Slide 32



Slide 33








Slide 1


Module 5


MALARIA ENTOMOLOGICAL
SURVEILLANCE


Republic of Kenya
Ministry of Health


Division of Malaria
Control


World Health Organization


MEASURE
Evaluation


USAID
President's Malaria Initiative

Slide 2

Objectives

- Describe the role of mosquitoes in malaria transmission
- Describe different types of mosquito surveys and their roles in malaria vector surveillance
- To stratify the distribution, density, behavior of vectors in relation to malaria transmission & control options.
- Describe how to conduct insecticide susceptibility & cone bioassay tests

Slide 3

Unit 1

Introduction to Malaria Entomology

Slide 4

Activity (10 mins)

Question and Answer Session

- What is malaria entomology?
- How is malaria transmitted?
- Do all mosquitoes transmit malaria?

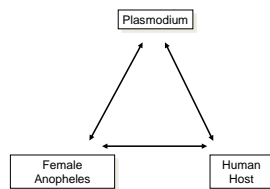
Slide 5

Definition

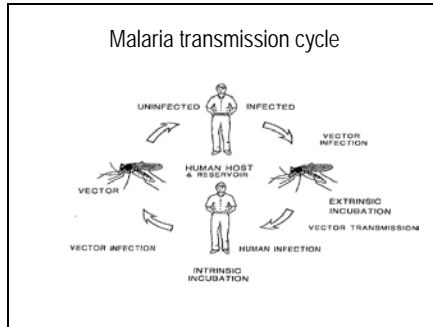
- Malaria is a parasitic disease caused by a protozoan parasite of the genus *Plasmodium* transmitted by an infective female *Anopheles* mosquito
 - Only female mosquitoes feed on blood as a requirement for their eggs maturation
 - In Kenya only *Anopheles gambiae* and *funestus* are known malaria vectors

Slide 6

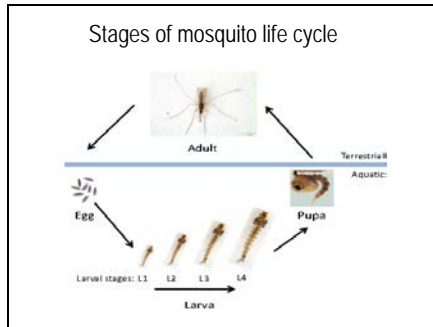
Malaria Transmission Triangle



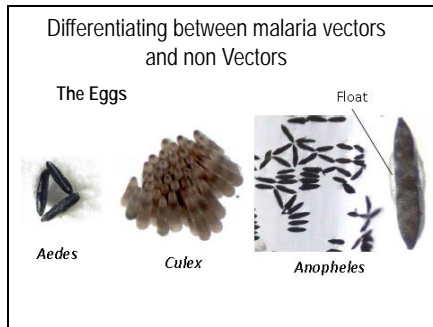
Slide 7



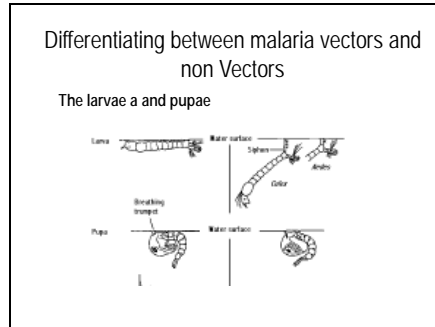
Slide 8



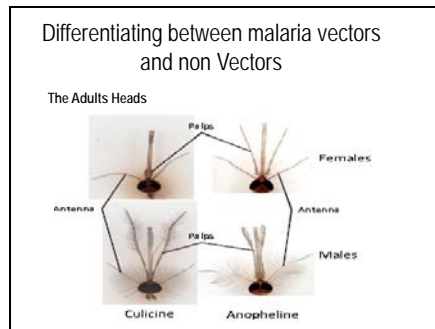
Slide 9



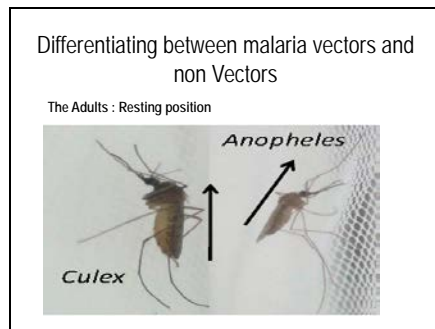
Slide 10



Slide 11




Slide 12



Slide 13

Bio-ecological traits of malaria vectors



- A female mosquito enters a house in search of a blood meal – sits on the wall to orientate
stop mosquitoes from entering into houses (Target = Adults)
- After biting, the mosquitoes usually rest on the wall to digest the blood meal
stop mosquitoes from biting people to get a blood meal (Target = Adults)
- Becomes gravid, searches for suitable water body to lay eggs on
stop gravid mosquitoes from laying eggs on water (Target = Adults)

Slide 14

Bio-ecological traits of malaria vectors

- Feeding preferences (Host choice): Man or other animals?
- Time of feeding: Early evening or late at night?
- Place of feeding (Indoors or outdoor)
- Resting behavior (Indoor or outdoors)
- Effects of bio-ecological traits on choice of vector control methods and their effectiveness

Slide 15

Unit 2

Surveillance of Malaria Vectors

Slide 16

Brainstorming (15Minutes)

- What is vector surveillance?
- Why do vector surveillance?
- What is the use of vector surveillance data?
- How do you collect vector surveillance data?

Slide 17

Definition

- Vector surveillance is a regular and systematic collection, analysis and interpretation of entomological data.

Slide 18

Why vector surveillance

- To know the type and density of mosquitos
- To determine the entomological inoculation rates (EIR)
- To know the feeding and resting behaviour of mosquitos
- To evaluate interventions and resistance studies

Slide 19

Usefulness of vector surveillance data

- Planning, implementation and evaluation of vector control interventions
- Early detection, prediction and prevention of vector borne disease outbreaks through a systematic data collection, analysis and evidence based decision making procedures.
- Early resistance detection and management
- Timely dissemination of the data to those responsible for vector control interventions.
- Development of malaria entomological profile

Slide 20

Types of entomological surveys

There are four main types of mosquito surveys:

- **Preliminary surveys:**
 - original, basic and short-term.
 - used to gather baseline data usually for the purpose of planning a vector control intervention.
 - Emphasis on vector species, density, resting & feeding behavior, larval habitats, longevity, infection rates & insecticide susceptibility.
- **Regular or trend observations:**
 - routine or long-term observations (longitudinal or operational surveys of monitoring).
 - carried out regularly (e.g. weekly, monthly) in order to evaluate the impact of control measures.

Slide 21

Types of entomological surveys cont'd

- **Spot checks:**
 - carried out in randomly chosen localities other than the fixed monitoring stations
 - provide supplementary information from areas otherwise not represented in routine monitoring.
- **Foci investigations:**
 - carried out in areas of new or persistent malaria transmission to investigate reasons for disease transmission, or why implemented interventions are ineffective in reducing disease burden.

Slide 22

Methods of mosquito sampling

- The choice of method for mosquito sampling depends on the purpose and the desired outcome.
- The following are the main methods of sampling
 - Pyrethrum Spray Catches (PSC);
 - Hand collections
 - Light traps
 - Human Landing Catches
 - Window (entry/exit) trap
 - Larval collection

Slide 23

Pyrethrum Spray Catches (PSC)

- Method used to collect indoor resting mosquitoes to establish densities, species composition, physiological status, human blood index and infection rates
- Resting mosquitoes are knocked down using aerosols and collected on white calico sheets as shown in the photo.

Slide 24

Pyrethrum spray collection



Slide 25

Hand collections

- Method used to collect resting mosquitoes to establish densities, species composition, physiological status, resting behavior, human blood index and infection rates or rearing for insecticide resistance monitoring
- Resting mosquitoes are picked using a sucking tube (aspirator) & placed in mosquito cages or paper cups.
- This method picks live mosquitoes

Slide 26

Hand collections and main materials used



1. mouth aspirator, 2. mechanical aspirator, 3. flashlight, 4. spare batteries, 5. adhesive tape, 6. rubber bands, 7. paper-cups with netting, 8. cotton wool.

Slide 27

Light traps

- Method used to collect mosquitoes to establish densities, species composition, resting behaviour, physiological status, human blood index and infection rates
- Light trap is a battery powered device fitted with a motor, a fan and light source.
- Mosquitoes are collected either dead or alive

Slide 28

Light trap



Slide 29

Window (exit/entry) trap

- Method used to collect mosquitoes to establish densities, species composition, physiological status, human blood index, infection rates, resting and exit behaviors
- Exit trap is device fitted to a window such that all exiting/entering mosquitoes are trapped within it
- Mosquitoes are collected either dead or alive

Slide 30

Window (exit/entry) trap



Slide 31

Human landing Catches

- Method used to collect mosquitoes to establish densities, species composition, association between man and mosquito, biting patterns, infection rates, feeding behavior and transmission intensities.
- This method of collection act as a bait and any mosquitoes landing are picked before they bite
- Mosquitoes are collected alive

Slide 32

Human Landing Catch

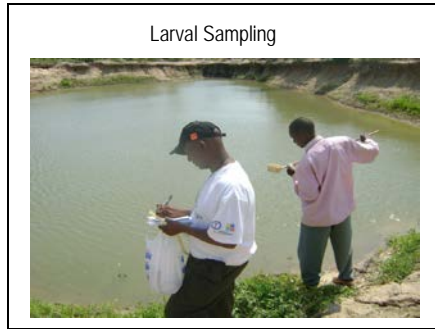


Slide 33

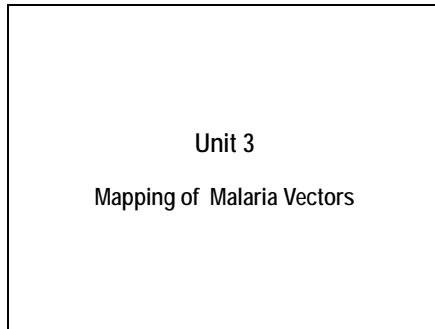
Larval sampling

- Larval sampling is important for
 - Determination of the vector species present in the study area.
 - Identification of preferred active breeding sites for each species.
 - Determination of the geographical distribution of vectors.
 - Evaluation of anti-larval measures on larval density.
 - Collecting samples for rearing to adults in the for insecticide susceptibility studies

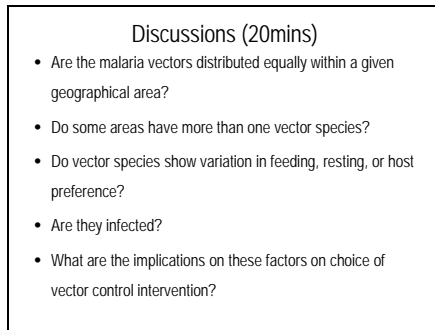
Slide 34



Slide 35



Slide 36



Slide 37

Mapping

- A map is a visual *representation* of an area – a symbolic depiction highlighting relationships between elements of that space
- Vector maps depict the distribution, species, relative abundance, vectorial capacity and other parameters related to malaria transmission in space and time.
- A detailed knowledge of the distribution, behaviour, & malaria transmission potentials of the main *Anopheles* malaria vectors guide the choice & targeting of vector control interventions.

Slide 38

Mapping of malaria vectors

- Malaria transmission in endemic countries is not uniform thus resulting in differences in its epidemiology
- Climatic conditions such as temperature, humidity & precipitation have a direct bearing on vector breeding, density, distribution, longevity, feeding frequency, resting behavior and the rate at which the parasites develop in the vector

Slide 39

Mapping of malaria vectors Cont'd

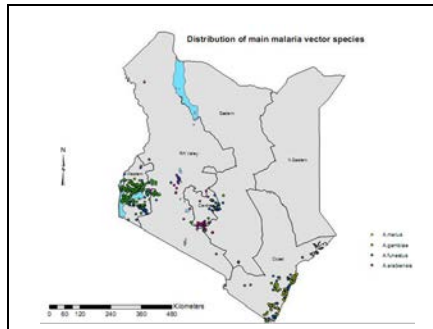
- These vector traits have direct impact on malaria transmission potentials and the choice of vector control interventions
- Thus it is important to map the spatial and temporal variation of the key vector parameters and reviewing them from time to time as vector control interventions are scaled up

Slide 40

Development of vector maps

- Data for development of vector maps can be obtained by actively conducting vector surveillance or retrospective records
- These data are geo-referenced using GIS and other techniques (Google maps)
- The key parameters are keyed in into a data base
- Maps with specific area generated by using a mapping software (e.g. Arcview GIS)

Slide 41



Slide 42

Unit 4

Insecticide Susceptibility and Cone Bioassay Tests

Slide 43

Why determine the susceptibility of malaria vectors to insecticides?

- If a vector is susceptible to an insecticide, then it means that the vector will be killed when it comes into contact with the insecticide used for the particular intervention (indoor residual spray, insecticide-treated bed net or larvicide).
- Decreasing susceptibility means that the vector becomes increasingly tolerant to the insecticide, up to a point where it becomes resistant.

Slide 44

Why determine the susceptibility of malaria vectors to insecticides? Cont'd

- If a vector develops resistance to an insecticide, it means it can withstand the dose that normally would have killed it and this may undermine the effectiveness of the intervention.
- It is therefore important to know the susceptibility level of the local vector to the insecticides to be used in the intervention.

Slide 45

Preparation of test vectors for susceptibility and cone bioassay evaluations

Two general methods are used to prepare/obtain test vectors for bioassays:

- Larvae may be collected from a range of local breeding sites and reared to adults
- Alternatively, blood fed & gravid local mosquito species are hand collected using adult sampling techniques and kept to lay eggs. The eggs are then reared to adults

Slide 46

Determining the susceptibility of adult mosquitoes

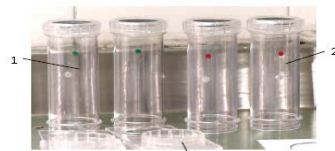
- There are standardized methods for determining vector susceptibility to insecticides in adult mosquitoes.

WHO Tube Assay:

- The standardized methodology is provided by the World Health Organization (WHO) for assessing the susceptibility of female *Anopheles*
- Mosquito vectors of a known species are exposed in special test tubes containing filter papers, impregnated with a lethal concentration (discriminating dose) of a given insecticide dissolved in oil.

Slide 47

WHO test tubes for susceptibility testing



1. Control/holding tubes (green dot), 2. exposure tubes (red dot), 3. slide units

Slide 48

The WHO kit

- The WHO tube test kit is made up of two plastic tubes
- One of the tubes is marked with a red dot & is used as "exposure tube" as it is lined with insecticide impregnated filter paper
- The other tube, with a green dot, serves as a "holding tube", which has its inner walls lined with plain paper.
- Another exposure tube (also marked with a green dot) is lined with a filter paper impregnated only with the oil used to dissolve the insecticide and serves as a control

Slide 49

The bioassay

- 15-20 mosquitoes are exposed to each insecticide for one hour & an equal number kept as control
- Observation for dead mosquitoes is done at 15min intervals
- After the exposure period the mosquitoes are transferred to the holding tubes and mortality recorded after 24 hrs

Slide 50

Interpretation of the results

WHO classification for insecticide resistance as follows:-

- 98-100% - Susceptible
- 90-97% - Resistance suspect
- < 90% - Resistance

Slide 51

Residual Efficacy of Insecticide on Sprayed Surfaces and insecticide treated materials

- The residual efficacy of an insecticide on a sprayed surface is determined by cone bioassay tests
- Done by checking mortality of the target mosquito vector species exposed to the sprayed surface at intervals of weeks or months after the spraying.
- This technique can be also used to evaluate the quality of a residual spraying operation
- Also used to determine residual efficacy of an insecticide on bed nets.

Slide 52

The WHO Cone Bioassay kit

- The WHO cone bioassay kit includes:
 - plastic cones,
 - adhesive sponge tape,
 - bent aspirator or sucking tube,
 - normal aspirators or sucking tubes,
 - cardboard paper, s
 - mall nails,
 - hammer,
 - cotton
 - wool,
 - paper cups with cover nets,
 - rubber bands, markers,
 - mosquito cage, wooden box with large holes, towels

Slide 53

The bioassay

- The cones are fixed on the test surface (wall or net)
- Untreated surfaces or materials are used as control
- 10 mosquitoes from a fully susceptible *Anopheles* strain from an insectary are introduced into each cone & a piece of cotton wool inserted in the opening of the cone
- After the exposure period (usually 30 mins) the mosquitoes are removed and dead ones counted
- Mosquitoes are transferred to holding paper cups and mortality recorded after 24 hours

Slide 54

WHO cone bioassay on a wall



Slide 55

WHO cone bioassay on an insecticide treated net



Slide 56

Demonstrations

1 Hr






Slide 57

Thank You

Slide 1

Module 6

MALARIA EPIDEMIC
PREPAREDNESS AND RESPONSE



Slide 2

Objectives

1. Describe malaria epidemics
2. Demonstrate malaria epidemic threshold setting
3. Describe methods of malaria epidemic prevention
4. Develop malaria epidemic preparedness and response plans
5. Describe post malaria epidemics evaluation

2

Slide 3

Unit 1

Introduction to Malaria Epidemics

Slide 4

Brainstorming (5 Min)

What is an Epidemic?

Slide 5

Definition of an Epidemic

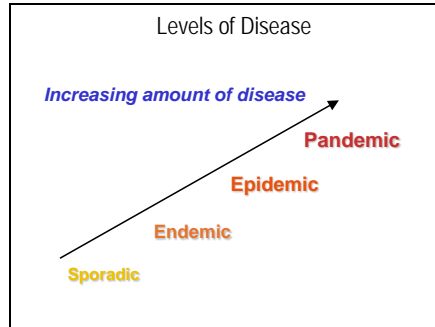
- Occurrence of more cases of disease than expected in a given area among a specific group of people over a particular period of time
- Synonym-Outbreak

Slide 6

What is an Epidemic?

- A public health emergency
- A political emergency
- An economic emergency
- An unusual event
- An event requiring rapid action
- Surveillance failure
- Control failure
- An opportunity

Slide 7



Slide 8

Malaria epidemic

- An increase in morbidity (cases) beyond what is normal for the area
- Excessive case-fatality rates in *P. falciparum* malaria (>1% for all cases and >20% for severe cases)
- Malaria caseload exceeding the capacity of the existing health care facilities to handle
- A disturbance of a previously existing epidemiological equilibrium

8

Slide 9

Causes of Malaria epidemics

Human related Factors

- Relative immunity
- Population movement, displacement, resettlement
- Land use practices
- Vulnerability due to other factors – malnutrition, HIV etc

Vector related Factors

- Increased breeding possibilities of vectors due to abnormal heavy rains or flooding downstream
- Changing agricultural practices especially irrigation
- New and more efficient vectors
- Breakdown of vector control program
- Insecticide resistance

Parasite related Factors

- Resistance to anti-malaria drugs

9

Slide 10

Factors Triggering Malaria epidemics	
Determinants (Direct)	Influencing Factors (Indirect)
Vector density	Rainfall, drought, incorrect maintenance of irrigation systems changed in vector breeding habitats
Human biting	Housing, behaviour, disaster, socio-economic factors
Rate of gametocyte carriers	Importation of malaria parasite
Length of sporogony	Temperature
Daily survival rate of vectors	Temperature, humidity

Slide 11

Types of Malaria epidemics

a) **True epidemics**—infrequent/cyclical outbreaks in relatively non-immune populations related to climatic anomalies (mainly arid and semi-arid zones). E.g. Eastern Kenya

b) **Strongly seasonal transmission**—variable but relatively predictable transmission influenced by variations in normal climatology. Population living in western Kenya highlands

c) **Neglect/breakdown of control**—where malaria has re-emerged due to neglected control activities

d) **Complex emergencies**—malaria transmission exacerbated by population movements and country political instability.

11

Slide 12

Brainstorming (5 Min)

What are the consequences of Epidemics?

Slide 13

Consequences of malaria epidemics

- Considerable morbidity and mortality in affected population
- Vulnerable groups more susceptible to other diseases
- Disrupt health care services
- Long-term consequences for the health of unborn children
- Additional costs at family, community & health sector level for both prevention and cure
- Economic losses through decline in agriculture output
- School and work absenteeism


13

Slide 14

Malaria Epidemics

Thresholds Setting In Kenya

Unit 2

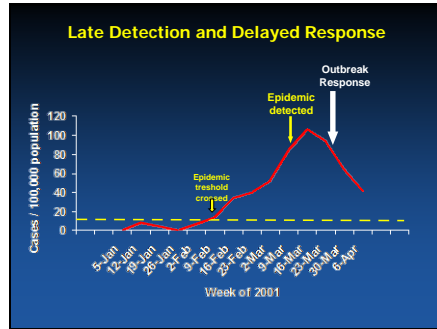


Slide 15

Brainstorming (5 min)

What is a threshold?

Slide 16



Slide 17

Malaria epidemics thresholds

Definition of threshold:

- Threshold is a science base indicator used to determine when a situation has developed into another situation.
- A malaria epidemic alert threshold is reached when there is an increase above the expected cases seen over a period of time in weekly or monthly summary reporting.
- It helps surveillance and programme managers to decide when to take action and what that action will be.

Slide 18

Malaria epidemics thresholds Cont'd

It can be summarized as follows:

- A malaria epidemic management tool based on weekly case-based reliable data
- Provides an early warning and very short lead time for increasing preparedness and response
- A situation analysis describing **who** is at risk for the disease, **what** are the risks, **when** is action needed to prevent a wider outbreak and **where** do the epidemics usually occur
- An evidence based tool for declaration of an epidemic

Slide 19

Determination of epidemics thresholds

Thresholds can be divided in ALERT and ACTION thresholds.

- An ALERT threshold suggests to health staff further investigations are needed. A malaria alert threshold is reached when there is an increase above the expected cases seen over a period of time in weekly summary reporting.

Slide 20

Response to an alert threshold

- Reviewing past data and reporting malaria incidence increase to the next level
- Suspect all cases of fever reported to health facility as malaria especially during high season malaria
- Use Laboratory confirmation methods for all suspected cases that fit the standard case definition
- Being more alert to new data and actively follow up trends in malaria incidence
- Alert the epidemic response team to a potential epidemic or outbreak

Slide 21

What can account for an apparent increase in cases?

- Change in reporting procedures / change in surveillance system
- Change in case definition
- Improvements in diagnostic procedures
- Increased awareness
- Increased access to health care
- New clinician- may see more referred cases, may make diagnosis more often, or report more consistently
- Laboratory or diagnostic error
- Batch reporting
- Change in denominator
- True increase in incidence

Slide 22

Action Threshold

- A malaria ACTION threshold is reached when there is a steady increase above the expected cases seen over a period of time in weekly summary reporting.
- This increase is proved real after investigations triggered by alert threshold.

Slide 23

Response to an Action threshold

This can be

- Net distribution
- Enhancing public awareness
- Improve case detection and management
- Ensuring adequate stocks of reagents, drugs and non pharmaceuticals
- Indoor residual spraying (IRS)
- On the Job training (OJT)
- Enhanced surveillance

Slide 24

Types of epidemics thresholds

- Constant case count:
- Third quartile:
- Cullen method:
- C-SUM (Cumulative sum) Method:

Slide 25

Types of epidemics thresholds

Constant case count:

- This is used where there is little or no malaria, or not much variation by season.
- It sets a threshold that is constant all year. (Botswana).
- An epidemic occurs if the number is above threshold.
- 400 cases/week in one district indicates an ALERT which should be acted upon
- 800 cases/week in one district indicates the national authorities should be informed
- 1200 cases/week indicates a national emergency

Slide 26

Types of epidemics thresholds

Third quartile:

- It calculates the thresholds as the third or upper quartile value of the number of cases per week for at least the last 5 years.
- This mean that $\frac{3}{4}$ (75%) of the time, we expected the number of cases to be below the threshold.
- Epidemics years are included in the calculation of this type of threshold.

Slide 27

Types of epidemics thresholds (cont)

Cullen method:

- It sets a threshold from the mean+2SD of the 5 years or more previous years number of cases for the week or month.
- This mean roughly 97.5% of the time, the number of cases will be below the threshold.
- Epidemic years must not be factored into the calculation of threshold.

Slide 28

Types of epidemics thresholds Cont'd

C-SUM (Cumulative sum) Method:

- This uses a running total of cases for each year rather than the weekly or monthly average.
- The threshold for each week or month is based on a moving average of that week or month plus the preceding and following weeks or months, to account for yearly variation in onset of the malaria "season".
- This method can give a good picture of whether the number of cases is rising faster than usual in a certain year.
- ❖ WHO recommends the 3Q method, especially at health facility level. District level aggregates can use mean+2SD threshold.

Slide 29

Thresholds Proposed for Kenya

1. **Health Facility Level (level 2-3):**
 - Third quartile as ALERT threshold and communicate with district for early investigation
2. **District aggregates and District Hospitals with large catchment areas:**
 - Third quartile as ALERT threshold and Mean + 1.5 SD Mean threshold as ACTION threshold
3. **Provincial/County Aggregates:**
 - Long term mean versus Current incidence to follow on trends. It is not a threshold for epidemic detection

Slide 30

Calculation of ALERT thresholds

By Hand:

1. Write down the OPD data by week for each health facility from the last 5 years.
2. For each week sort the numbers by ascending order from the lowest to the highest and write them in the table. Week one to week 52
3. The middle number in each group is the median. Take the median for each week and plot the points with a line. This is the median number of cases expected per week.
4. The 4th highest number in each series is the 3Q. Take the 3Q number for each week and plot the on the graph of cases by week and join the points. This is the ALERT threshold level.

Slide 31

Calculation of ALERT thresholds Cont'd

Using an Excel spreadsheet:

1. Open a file Malaria epidemic threshold and save with the name of district or health facility
2. Save with the name of Health facility or district
3. Click on the sheet and name it "weekly data"
4. Enter weekly or data year 1 to year 5. For 3Q include all years for Cullen we should exclude epidemic years.

Slide 32

Steps in setting up Malaria weekly threshold using quartiles

1. Collect weekly Malaria data for 5 or 7 years and the current year.
2. Make a trend graph on Malaria data collected
3. Rank the data in ascending order across the period i.e. week 1 for all the years---week 52 for all the years.
4. Get the median of the distribution. This becomes 2Q (second quartile of the distribution).
5. Identify the median of the distribution below the median (2Q) and this becomes the first quartile (1Q).

Slide 33

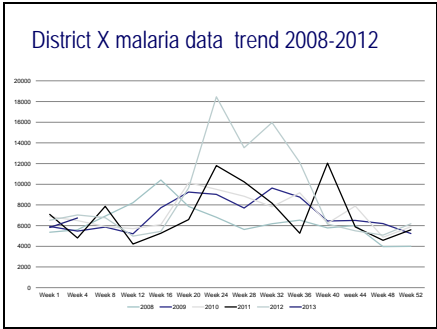
Steps Continued....

6. Identify the median of the distribution above the median (2Q) and this becomes the third quartile (3Q).
 7. Plot a graph using figures in columns 1Q, 2Q, 3Q, and the current years data.
 8. Name the zones as follows
 - I. Success zone - The area below the 1Q
 - II. Security zone - The area between lines 1Q and 2Q
 - III. Alarm zone - The area between lines 2Q and 3Q
 - IV. epidemic zone - The area above line 3Q (replaced by mean + 1.5 SD)
- NB: Health facilities should use data in 3Q to monitor malaria trends.

Slide 34

District X Malaria Data, 2008-2013						
	2008	2009	2010	2011	2012	2013
Week 1	5357	5913	6877	7085	6514	5824
Week 4	5589	5464	6493	4807	7028	6732
Week 8	6922	5854	5914	7866	6765	
Week 12	8218	5206	5714	4220	4988	
Week 16	10411	7719	6095	5265	5448	
Week 20	7845	9250	10143	6584	9648	
Week 24	6811	9025	9513	11798	18454	
Week 28	5629	7704	8835	10232	13533	
Week 32	6186	9635	7828	8170	15969	
Week 36	6530	8751	9172	5266	12114	
Week 40	5772	6449	6229	12038	6159	
week 44	6012	6503	7890	5880	5500	
Week 48	3972	6201	4865	4583	5077	
Week 52	4001	5231	5885	5589	6177	
Fill data for all weeks (1-52)						

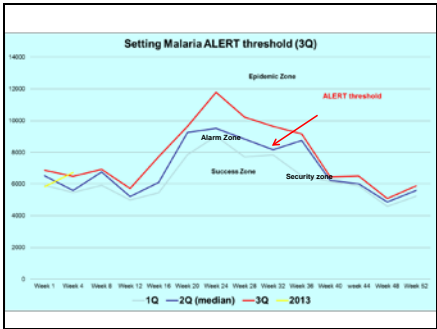
Slide 35



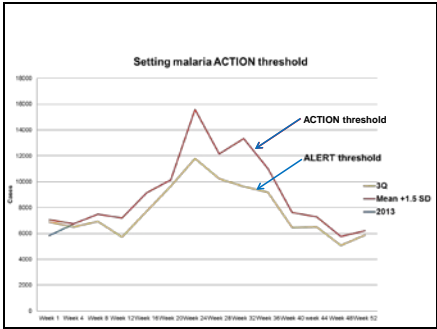
Slide 36

Malaria data in ascending order						
	1Q	2Q (median)	3Q	2013		
Week 1	5357	5913	6514	6877	7085	5824
Week 4	4807	5464	5589	6493	7028	6732
Week 8	5854	5914	6765	6922	7866	
Week 12	4220	4988	5206	5714	8218	
Week 16	5265	5448	6095	7719	10411	
Week 20	6584	7845	9250	9648	10143	
Week 24	6811	9025	9513	11798	18454	
Week 28	5629	7704	8835	10232	13533	
Week 32	6186	7828	8170	9635	15969	
Week 36	5266	6530	8751	9172	12114	
Week 40	5772	6449	6229	12038	6159	
week 44	5500	5880	6012	6503	7890	
Week 48	3972	4583	4865	5077	6201	
Week 52	4001	5231	5589	5885	6177	
Fill data for all weeks (1-52)						

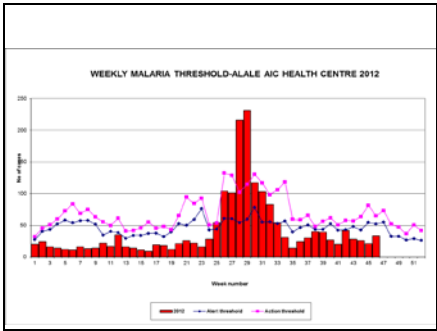
Slide 37



Slide 38



Slide 39



Slide 40

Investigating & Declaration of Epidemics

- **Malaria Prevention interventions:**
 - IRS coverage and Timeliness
 - ITN distribution and Re-treatment
 - Insecticide Resistance
 - EPR planning and implementation
 - Cross border movements
- **Malaria Case management:**
 - Drug availability & consumption
 - Blood Transfusions
 - CFR
 - Slide & RDT positivity rates
 - Drug resistance
- ❖ **Provincial/county Authorities declare epidemics**

Slide 41

Constant Case Count	Mean + 2DS or Mean + 1.6 SD	3 rd Quartile
<ul style="list-style-type: none"> • High sensitivity in season (detects most epidemics) • Ease of calculation (time & process) • Results in high False positives • Based upon Weekly Data • Little early warning 	<ul style="list-style-type: none"> • Based upon weekly data • Appears to give a valid 'epidemic threshold' • Varies throughout the season • Must exclude epidemic year • Difficult to calculate • Requires a PC 	<ul style="list-style-type: none"> • Based upon weekly data • Relatively easy to calculate • Varies throughout the season • Good Early Warning Indicator • Don't need to exclude epidemic year • Perceived as difficult to use • Time consuming

Slide 42

Group work (1 hour)

1. Use dummy data to set thresholds
2. Demonstrate using the excel tool to set the threshold.
3. Use data from your health facility or district to set malaria ALERT and ACTION thresholds

Slide 43

Unit 3

Methods of Malaria Epidemic Prevention

Slide 44

Malaria epidemic prevention strategies

What are the main malaria epidemic prevention strategies?

44

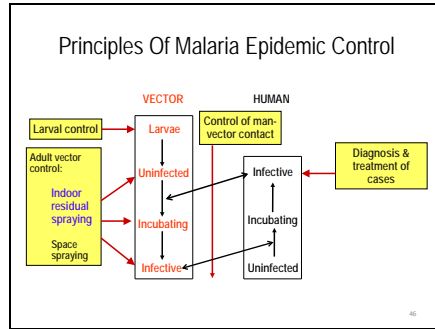
Slide 45

Malaria Epidemic Prevention Strategies

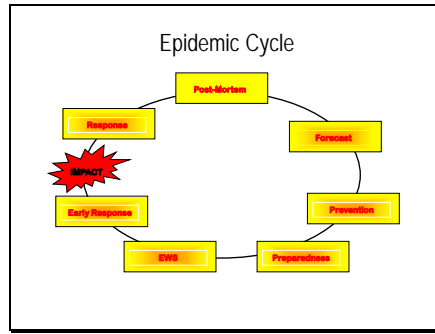
- Vector control
 - LLINs:
 - Environmental management – drainage of stagnant water
- Surveillance
 - Early detection of all cases
- IPTp
 - IPTp for pregnant women residing in malaria endemic regions
- ACSM
 - Awareness creation and reinforcement of preventive strategies

45

Slide 46



Slide 47



Slide 48

Malaria Epidemic Prevention: - Vector Control

- What are the known malaria vector/s?
- Do they rest or feed indoor/outdoor?
- Is there an ongoing malaria vector control program?
- Do people use LLINs if yes what is the current coverage of households/high risk groups?
- Is there reason to suspect insecticide resistance?

48

Slide 49

Unit 4

Epidemic Preparedness and Response plans

Slide 50

Brainstorming session

What are the key components of an epidemic preparedness and response plan?

Slide 51

OUTLINE OF AN EPIDEMIC PREPAREDNESS AND RESPONSE PLAN

- Introduction
- Problems
- Objectives
- Strategies
- Targets/Priorities
- Activities
- Resources
- Implementers
- Time Lines
- Monitoring indicators
- Evaluation indicators

Slide 52

EPR planning levels

- Facility level
- Sub county level
- County level
- National level

Slide 53

Brainstorming (5 min)

What do you take into consideration when making EPR plans?

Slide 54

Considerations for EPR Plans

- Vector control
 - Establish efficacy of existing IRS
 - Train teams for IRS
 - Make insecticide and pumps and logistics available for IRS
 - LLIN coverage and use
- Case management
 - Diagnosis and treatment
- Communication
 - Information campaigns, health education
- Monitoring and evaluation
- Partner mobilisation

54

Slide 55

Logical Framework for Malaria EPR Plan						
Problem statement	Strategy/Intervention on	Activities	Resources	Responsible person/unit	Timeline	Progress Indicators
Strengthen analysis to inform	Program Management	Training EPR Managers	GOK	National level	Specify timelines	No of teams trained
	Vector Control	Training Purchasing of commodities Spraying	GOK	county	Specify timelines	300 spray personnel trained
	Case Management	Training	GOK	County and sub county	Specify timelines	No of personnel trained on case management
	IEC	Training CHW	GOK	County and sub county	Specify timelines	150 CHW Key issues identified for IEC. Materials printed and distributed
	Surveillance and M&E	Strengthen MIS Dev Database for M&E	GOK	County and sub county	Specify timelines	MIS updated with critical key indicators Computer system updated.

Slide 56

Practicum (1 hr)

- Teams to assemble and come up with epidemic preparedness and response plans

Slide 57

Malaria Epidemic Response

- Introduction
 - Rapid assessment
 - Epidemic notification
 - Resource mobilisation
 - Response activities

Slide 58

Malaria Epidemic Response

- Rapid assessment:
 - Determine extent of the problem
 - Define type and size of intervention/s and priority activities
 - Plan the implementation of the activities
 - Pass information to stakeholders, international organizations to mobilize additional resources

58

Slide 59

Epidemic Notification

- Upon confirmation, disease outbreak management teams (DOMT) should notify health facilities and in the sub county / county, DDSR and DOMC.
- A team should be sent to confirm the epidemic before notifying WHO and other partners.

Slide 60

Resource Mobilisation

- Personnel
- Equipment (vehicles and fuel)
- Commodities (anti malarial drugs, IV fluids, syringes etc)
- Lab supplies
- Insecticides
- Emergency response funds

Slide 61

Response activities

- Interventions to be selected according to eco epidemiological zones (Refer to eco epidemiological zonal table)
- Key activities include:
 - Strengthening treatment services & vector control services
 - Strengthen disease surveillance
 - Community mobilisation and health education
 - Coordination and response activities

Slide 62

Unit 5

Post epidemic assessment

Slide 63

Introduction

- This is the final activity aimed at documenting the preparedness and effectiveness of epidemic response
- It provides experiences and lessons learnt to guide future EPR actions
- All indicators specific to EPR should be assessed. (Refer to national malaria M&E plan)

Slide 64

Brainstorming (5 min)

What indicators are used for assessing epidemic preparedness and response?

Slide 65

EPR indicators

- Malaria death rate among target population
- Proportion of out patient and inpatient malaria cases
- Percentage of health facilities reporting no stock of anti malarial for more than one week in the last three months.
- Percentage of IRS coverage (where implemented)

Slide 66

Assessment activities and levels

Levels	Assessment Activities			
	Preparedness	Prevention	Early detection	Response
Community	- Availability of community systems, treatment practices	- Availability of bed nets, IRS coverage, timing between IRS & occurrence of epidemics	- Reports of acute deaths in the community	- Involvement of community in dissemination of information
Health facility	- Whether adequate surveillance data was collected		- Whether facilities used surveillance to draw charts the thresholds to detect outbreak	

Slide 67

Assessment activities and levels Cont'd				
	Preparedness	Prevention	Early detection	Response
Sub county	- Whether health facility teams were trained on EPR - Whether the district has adequate EPR committees - No of EPR meetings held at the district	- LLIN coverage - No. of people protected by indoor residual spraying	- Proportion of health facilities with - Proportion of surveillance graphs - No. of health facilities - Notification and response	- Whether there were sufficient committees for rapid response - Whether there were enough personnel to handle the epidemic - Any stock outs - No. of cases confirmed and treated
County	- Proportion of districts with functional EPR teams and plans - Proportion of districts with adequate committees - Frequency of support supervision	- Whether CHRT's support risk and monitor district preventive activities	- Whether the affected districts were timely epidemic response - Whether the county has an updated risk map	- Whether the CHRT's conducted support supervision for epidemic response
National	- All health resources were allocated for epidemic response - Whether there were adequate buffer stock for EPR - Whether EPR planning meetings were held	- Whether adequate resources were allocated for epidemic prevention in high risk areas	- Whether the national level prepared materials risk maps - Proportion of epidemic response - Whether adequate budget was allocated for epidemic response	- Timely communication of epidemic risk data - National level effectiveness of national level in setting epidemic - Whether adequate budget was allocated for epidemic response

Slide 68

Remember

Failing to plan, Means planning to fail!

THANK YOU








68

Slide 1

Module 7

Supervision and feedback



Slide 2

Objectives

- Describe malaria support supervision
- Develop a plan for Malaria supervision and use the planning tools
- Perform malaria supervision and use the supervisory checklist
- Write a supervision report and give feedback using the reporting and feedback template

Slide 3

Unit 1

Introduction to Malaria Supervision

Slide 4

Brainstorming (5mins)

What is supervision?

Slide 5

Definition of supervision

- This is an activity carried out to by supervisors to oversee the productivity and progress of employees who report directly to them
- Supportive supervision is a processes of guiding, supporting and assisting service providers to assigned tasks so as to achieve organizational goals

Slide 6

Brainstorming (5 mins)

- What are the characteristics of a support supervisor

Slide 7

Characteristic of support supervisors

- Supports the staff in a way that helps them develop problem-solving skills.
- Helps workers to think critically, prioritize tasks and to communicate effectively.
- Observes, provides feedback, discusses technical issues with staff, updates staff on policies.

Slide 8

Characteristic of Support Supervisors

- Trains on -job and works with staff to jointly identify problems and develop action plans.
- Ensures that after each encounter, decisions are documented and appropriate follow up is done.

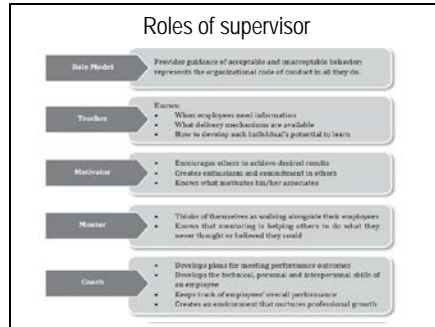
Slide 9

In addition

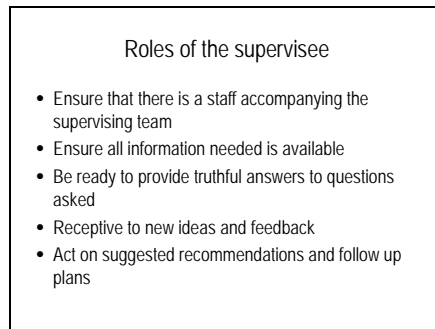
A support supervisor ensures that:

- Adequate resources are allocated and provided for carrying out the required task
- Facilities have adequate infrastructure and are adequately equipped
- Appropriate written procedures and guidelines are available and understood by staff
- Clients' rights are respected at all times.

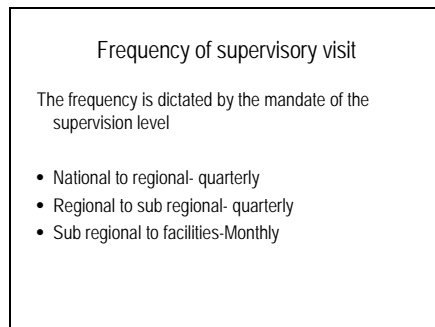
Slide 10



Slide 11



Slide 12



Slide 13

Integrated supervision

- For logistics reasons it is reasonable that supervision at sub-national levels be part of integrated supervision activities undertaken at these levels.
- Integration also allows for cost effective utilization of limited resources, reduces overburdening of health workers at the lower levels, and also minimizes interference without patient service provision.

Slide 14

Brainstorm (3)

What supervisory approaches do you know?

Slide 15

Supervision approaches

- Assessment
- Immediate feedback(onsite discussions)
- On job training

Slide 16

Unit 2

Planning for Malaria supervision

Slide 17

Brainstorming (5 mins)

How do you usually plan for your supervisory visit?

Slide 18

Introduction to planning

- Effective supportive supervision requires proper planning and coordination. The following steps should assist a supervision team while planning for and undertaking malaria supervisory visits.
- Creation of a contact list
- Advance scheduling of the visit
- Selection of team members

Slide 19

Contact list

- Contains the list of the person(s) at the facilities or districts the teams will communicate with during the supervision
- Should be updated regularly
- Allows the teams quick access to the relevant staff
- Will facilitate organization of the supervision visit logistics.

Slide 20

Advance Scheduling of Visits (1)

To avoid disruption to normal service delivery, the following tasks should be done in scheduling for a supervision visit :

- Plan for the supervision visit in advance, harmonizing the supportive supervision timetable with other programmatic schedules
- Consult with the proposed supervisory team members including the sub county / facility teams to ensure their availability
- Let the supervisory teams jointly select supervision visit dates and facilities/ districts to be visited.

Slide 21

Advance Scheduling of Visits (2)

- Communicate the agreed upon dates to the staff to be supervised well in advance so that they can be prepared for the visits.
- Review the previous reports and schedule the action points as necessary.
- Arrange for the necessary logistics for the visit e.g. transport and accommodation.
- Arrange to take along any supplies that will need to be replenished during the supervision e.g. reporting tools, guidelines, stationary etc.

Slide 22

Selection of supervisory team (1)

To enrich the support supervision experience, the following considerations should be made in composing a supervisory team:

- Allocate team members in a manner that ensures mix of skills, competencies and experience.
- Actively work to maintain team cohesion since no one member is competent in all areas of health care provision.
- Allocate each team member specific tasks beforehand, preferably according to their expertise and training.

Slide 23

Selection of supervisory team(2)

If the visiting team does not usually directly supervise the staff, the team needs to include a team member who is an immediate supervisor because:

- the staff will feel more comfortable to discuss their challenges, problems and needs with their immediate supervisors.
- the immediate supervisor has a better understanding of the staff and would therefore be in a position to give practical recommendations and assist the staff to achieve them.

Slide 24

Role of Malaria Control Coordinators (1)

The following are the supervisory responsibilities of malaria control coordinators:

- to ensure monthly site support supervision of health facilities under their jurisdiction.
- To ensure quality control procedures for data capture and transmission system

Slide 25

Brainstorming (5mins)

What is your role during supervision?

Slide 26

Role of Malaria Control Coordinators (2)

- Coordinate the supervisory visits
- Play a key role in planning the logistics for the visit
- Liaise with all persons to be involved to ensure availability and full participation.

Slide 27

Role of Disease surveillance Coordinator

- To assist Malaria control coordinator in surveillance supervision by:
 1. Conducting record search
 2. Use the health facility surveillance checklist

NB. The epidemic preparedness section must be applied in epidemic prone district and seasonal transmission areas

Slide 28

Introduction to planning tools

Slide 29

Practicals in filling the planning tools (30 mins)

Mavuno county has 5 districts with 5 facilities in each district. The CHMT of Mavuno county is planning to conduct supervisory activities to all their sub counties. How will they ensure that that the supervisory activity is well planned.

Use the planning tools available.

Slide 30

Unit 3

Conducting the Malaria support supervision

Slide 31

Conducting supervision visits

The following tasks should be undertaken during the supervisory visit:

- Meet with the facility/district in-charge and introduce yourselves and explain the purpose of the visit.
- Review the previous supervision report together and discuss the findings/challenges identified during that visit.
- Agree on how to carry out the supervision tasks during this visit and the debriefing afterwards.
- Assign specific supervisory tasks to different members of the team.
- Proceed to carry out tasks using the structured tools.

Slide 32

Debriefing after supervisory Visit (1)

- a. Thank the staff for participation and cooperation during visit
- b. Give feedback on the supervision findings covering:
 - i. what they have done well
 - ii. weak areas
 - iii. problems identified
- c. Congratulate the staff for positive findings
- d. Brainstorm for possible solutions on the identified problem areas e.g.
 - i. better planning
 - ii. better coordination
 - iii. extra training for the staff
 - iv. redeployment of staff

Slide 33

Debriefing after supervisory Visit(2)

- e. Give recommendations
- f. Agree on the way forward:
 - i. action points for the staff
 - ii. action points for the supervisors
 - iii. identify resources required
 - iv. define timelines for the action points
 - v. establish monitoring and evaluation mechanism for the agreed action points
- g. Update the staff on new knowledge, procedures and policies
- h. Thank the staff once again for the positive findings and participation in the supervision.

Slide 34

Tracking supervision visits

Each supervision visit to a health facility should be documented in the supervision logbook, which remains at the health facility. The following details should be included when signing the logbook:

- i. date(s) of the supervision visit
- ii. objective(s) of the visit
- iii. summary of the findings
- iv. recommendations and agreed action points
- v. signatures(s) of at least two members of the supervision team.

This logbook summary should be reviewed during the subsequent visits, before actual supervision is carried out.

Slide 35

Introduction to health facility surveillance checklists

- Health facility surveillance checklist
- Facility supervision checklist
- District supervision checklist
- County supervision checklist

Slide 36

Role play (45mins)

Divide yourself in groups of 5 appoint 4 members of the CHMT and one facility staff. The CHMT of Mavuno county should conduct a facility supervision and , administer the health facility supervision checklists including the health facility surveillance checklist

Slide 37

Unit 4

Report Writing and feedback

Slide 38

Brainstorming (5mins)

- Do you usually write supervision reports?
- How do you do them?
- Do you analyze your findings?

Slide 39

Analyzing the Supervision Visit Results

- The broad aspects looked at during the supervision and whose results should be analyzed include:
 - delivery of malaria services and best practices
 - human resources capacity and training status
 - availability of malaria supplies e.g. anti-malaria medicines
 - data management and reporting
 - availability of relevant malaria documents e.g. guidelines, job aids, etc.
 - Any problems and their priorities

Slide 40

Demonstration on how to score using the supervision checklist

Slide 41

Interpretation of supervisory scores

The performance of the supervisee under each of these categories should be calculated and graded as follows:

1. Excellent (80%-100%)

Interpretation:

- a. Performance frequently exceeded standards for the job
- b. Supervisee understood all matters and consistently provided high quality service
- c. Minimum problems were identified

Slide 42

Interpretation of supervisory scores

2. Good (50%-79%)

• *Interpretation:*

- a. Performance met the requirements of the job
- b. Supervisee performed these in a competent and satisfactory manner
- c. Supervisee is familiar with all the aspects of malaria control

3. Poor (<50%)

• *Interpretation:*

- a. Performance falls below average standard
- b. Severe constraints were identified
- c. Supervisee requires urgent intervention to improve service delivery.

Slide 43

Report writing

- The supervision team should compile detailed report soon after the visit (within 1week)
- The supervision report should be sent to the next supervision/management level, and a feedback report sent to the facility/sub county/county concerned.

Supervision summary forms should be used to give a quick overview of the results of the supervision visit

- These should be filled immediately after the supervision visits are over after several facilities/ sub county/counties are visited. The purpose of these forms is to summarize the findings of the visit before the teams submit the detailed reports.
- A copy will be sent back to the facility/county visited

Slide 44

Reporting Templates

Standardized reporting templates will be used to allow for objective supervision visits.

- These allows comparison of supervision results between Counties/facilities and between different visits.
- The teams should also include an addendum of issues that need to be reported but are not provided for in the template.
- [Detailed Supervision Report Format.docx](#)

Slide 45

Submission of the Reports

Supervision reports should be written and sent to the next management level within 1 week of completion of the activity

- The malaria focal is responsible for compiling, completing and ensuring that the report is on time
- The report should be countersigned by the chief health officer

Slide 46

Submission of the Reports (2)

The report should describe in details the following:

- i. how the supervision was conducted
- ii. the findings after the visit and their implications
- iii. immediate actions taken including updates given and on the job training conducted
- iv. action plans agreed on, their timelines and the responsible parties.

Slide 47

Brainstorming (5mins)

- How do you motivate service providers that have shown exemplary performance

Slide 48

Incentives and Other Follow up Actions

The supervision team should decide how to recognize staffs/facilities/districts/provinces that show exemplary performance. Some of the forms of recognition and/or incentives could include:

- i. letters of recommendation.
- ii. involving the staff in a mentorship program.
- iii. positively mentioning good performers during important meetings and gatherings.
- iv. certificates of recognition.

Slide 49

Practical on calculating scores and report writing (30 mins)

- The Mavuno CHMT has completed its Supervisory visits to 1 sub-county team and 2 facilities (the filled out supervision checklists have been given to you).
- Fill in the appropriate summary score sheets and summary reports

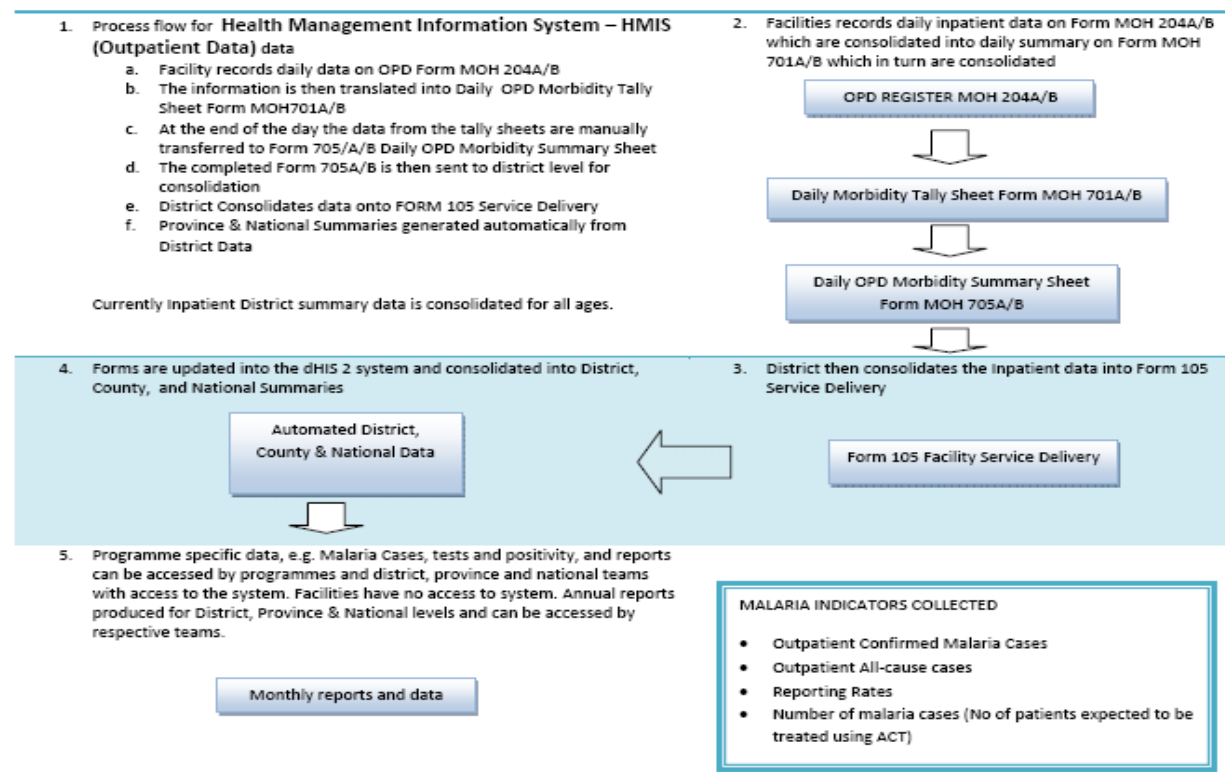
Slide 50

Thank you

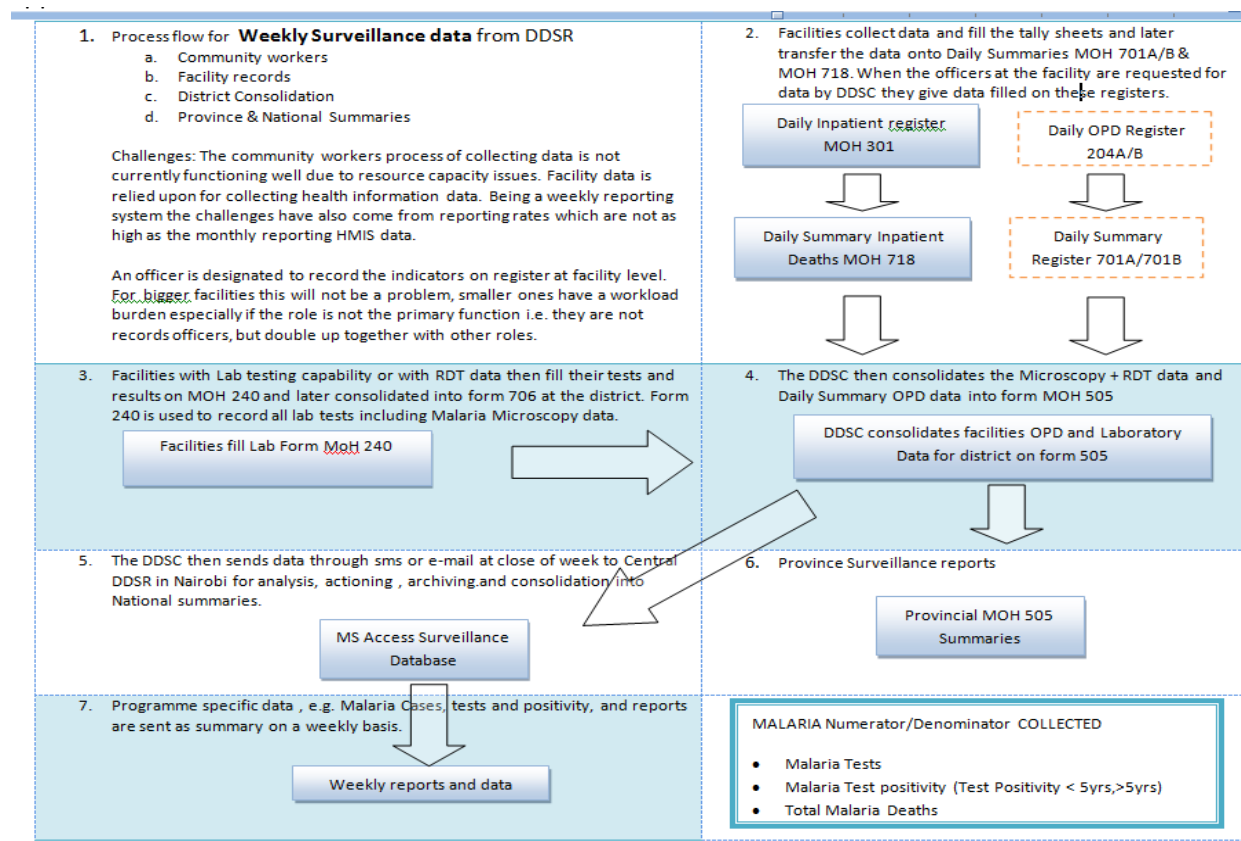
Appendices

1. HMIS Data Flow
2. DDSR Data Flow
3. Malaria Surveillance Indicators and Targets
4. Core Malaria Surveillance Graphs and Interpretations
5. Checklist for Supervising Surveillance and Response Activities at the Health Facility
6. Field Data Sheet
7. Malaria Facility Supervision Checklist
8. Malaria District Supervision Checklist
9. Facility Contact List
10. DHMT Contact List
11. District Supervision Activity Schedule
12. Facility Score Sheets
13. District Score Sheet
14. Supervision Summary Report
15. Detailed Supervision Report Format
16. IDSR Weekly Summary Reporting Form

Appendix 1: HMIS Data Flow



Appendix 2: DDSR Data Flow



Appendix 3: Malaria Surveillance Indicators and Targets

Most of the targets given here are the same as those published in the World Malaria Report 2008. However, a new indicator for malaria mortality reduction has been included—“near zero preventable deaths in 2015”. This indicator comes from the Roll Back Malaria (RBM) Global Malaria Action Plan for 2008–2015, which was published in September 2008.

These guidelines are the first to list preliminary targets for malaria test positivity rate (TPR). The targets are based on observations from five African countries and three recently published studies. These TPR targets may need to be revised once more experience is available.

INDICATORS AND TARGETS FOR MONITORING AND EVALUATING MALARIA PROGRAMMES

Indicator (measured monthly)	Numerator, denominator	Targets	Comments
1. Outpatient confirmed malaria cases ¹	<p><u>Numerator:</u> Number of outpatient confirmed malaria cases (by microscopy or RDT) reported by health facilities per year</p> <p><u>Denominator for rate:</u> Resident population by age (<5 years, all ages) per 1000 people resident in areas at risk of malaria²</p>	<p><u>Case/rate trend:</u></p> <ul style="list-style-type: none"> • >50% reduction by 2010 • >75% reduction by 2015 <p><u>Rate:</u></p> <ul style="list-style-type: none"> • <1 confirmed case per 1000 people indicates excellent control 	Rate of <1 confirmed case per 1000 people indicates readiness for elimination phase
2. Outpatient malaria TPR	<p><u>Numerator:</u> Number of outpatient laboratory—confirmed malaria cases</p> <p><u>Denominator:</u> Total number of outpatient suspected malaria cases tested × 100</p>	<p><u>TPR trend:</u></p> <ul style="list-style-type: none"> • >50% reduction by 2010 • >75% reduction by 2015 <p><u>Annual TPR:</u></p> <ul style="list-style-type: none"> • 10–20%—intermediate control • 5–9%—good control • <5%—excellent control 	Annual rate should be used, not just the rate during the peak season. • <5% in peak season indicates readiness for elimination phase
3. Inpatient malaria cases	<p><u>Numerator:</u> Cases (confirmed and unconfirmed) with a primary diagnosis of malaria at discharge (and not admission)</p> <p><u>Denominator for rate:</u> Resident population by age (<5, all ages) per 1000 people resident in areas at risk of malaria</p>	<p><u>Trend:</u></p> <ul style="list-style-type: none"> • >50% reduction by 2010 • >75% reduction by 2015 	
4. Inpatient malaria deaths	<p>Numerator – Deaths with a primary diagnosis of malaria at discharge</p> <p>Denominator for rate – Mid-year resident population by age (<5, all ages) per 1000 people resident in areas at risk of malaria</p>	<p><u>Trend:</u></p> <ul style="list-style-type: none"> • >50% reduction by 2010 • >75% reduction by 2015 <p>Elimination of malaria deaths by 2015</p>	

Indicator (measured monthly)	Numerator, denominator	Targets	Comments
5. Diagnostics: percentage of outpatient suspected malaria cases that undergo laboratory diagnosis	<p><u>Numerator:</u> Number of outpatient suspected malaria cases that received laboratory examination for malaria (microscopy or RDT)</p> <p><u>Denominator:</u> Number of outpatient suspected malaria cases $\times 100$</p>	$\geq 90\%$	
6. Treatment (ACT): percentage of outpatient malaria cases that received appropriate antimalarial treatment according to national policy	<p><u>Numerator:</u> Number of malaria cases receiving appropriate antimalarial treatment at health facility</p> <p><u>Denominator:</u> Number of outpatient malaria cases expected to be treated at health facility with appropriate antimalarial medicine (all those with a diagnosis of malaria) $\times 100^3$</p>	100%	
7. ITN: routine ITN distribution to populations at high risk (pregnant women)	<p><u>Numerator:</u> Number of ITNs distributed or delivered to target population—pregnant women attending ANCs</p> <p><u>Denominator:</u> Total number of pregnant women attending an ANC for the first time</p>	$\geq 80\%$	
8. IPT: IPT in pregnant women	<p><u>Numerator:</u> Number of pregnant women receiving second dose of IPT</p> <p><u>Denominator:</u> Number of pregnant women with at least one ANC visit</p>	$\geq 80\%$	
9. Stock-outs: percentage of health facilities without stock-outs of first-line antimalarial medicines, mosquito nets and diagnostics, by month ¹	<p><u>Numerator:</u> Number of health facilities, in areas at risk of malaria, without stock-outs of first-line antimalarial medicine (according to national policy), ITN and RDT in a month</p> <p><u>Denominator:</u> Number of reporting health facilities in the same areas at risk of malaria $\times 100$</p>	100%	

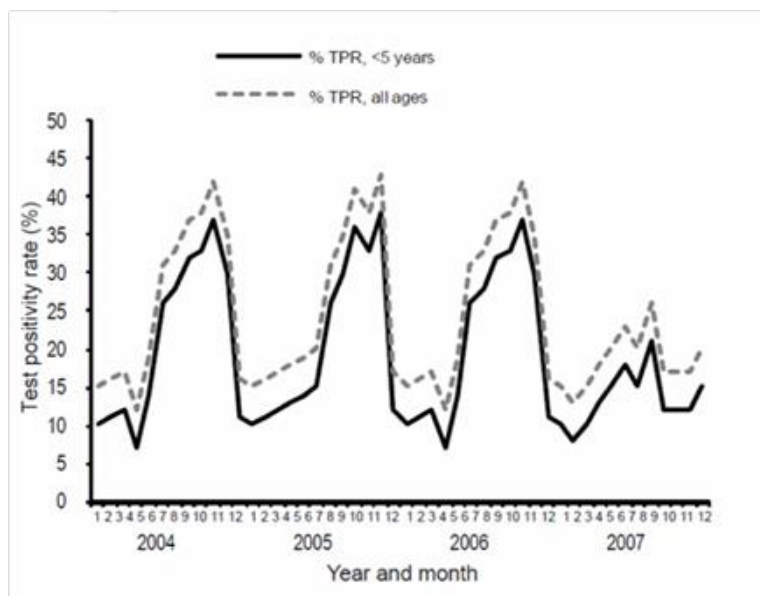
Indicator (measured monthly)	Numerator, denominator	Targets	Comments
10. Completeness of monthly health-facility reports on surveillance and logistics	<u>Numerator:</u> Number of health facility monthly reports received on surveillance and logistics, by month <u>Denominator:</u> Number of health facility reports expected each month	100%	

(Footnotes)

1. Epidemiological trends can be followed for any time interval—weekly, monthly or yearly. Trends of numbers of cases and deaths, without calculating rates, are the easiest to understand. Rates are useful if the trend period is long or comparisons are made, for example, with other countries, districts or provinces.
2. Several indicators use the term “population at risk.” In most high-burden African countries, almost everyone is at risk of malaria, except for those living at >2000–2500 m and those near the centre of some large cities.
3. Denominator is composed of those <5 years old and those ≥5 years old. Treatment and policy for those age groups may be different—for example, testing of all suspected malaria cases in those ≥5 years of age and treatment of only confirmed cases, but presumptive treatment of all suspected malaria cases (without regard for testing) for those <5 years old. Denominator comes from surveillance data— either suspected or confirmed malaria cases by age group, depending on treatment policy.

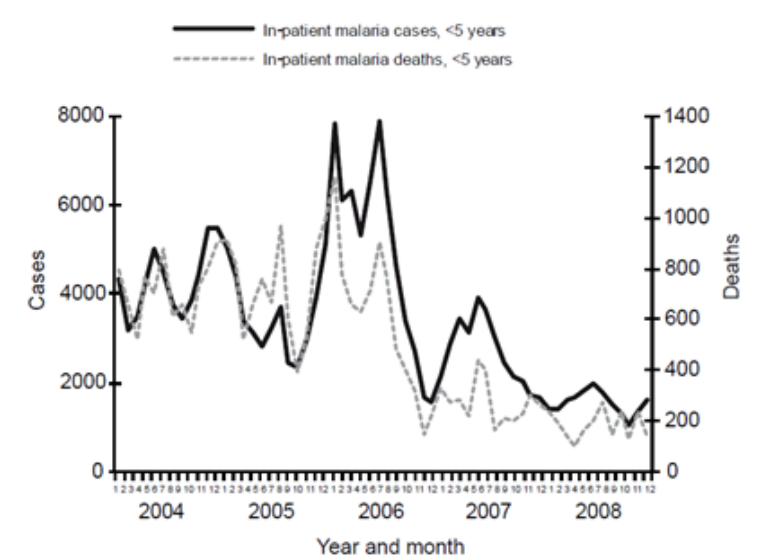
Appendix 4: Core Malaria Surveillance Graphs and Interpretations

Outpatient malaria TPR in children under 5 years of age and all ages



Peak-season and annual malaria TPR (assessed by RDT or microscopy) should decline progressively as control increases and transmission decreases. For example, annual malaria TPR should decline to less than 10% if outpatient confirmed cases and inpatient malaria cases and deaths decline by more than 80%

Inpatient malaria cases and deaths in children under 5 years of age



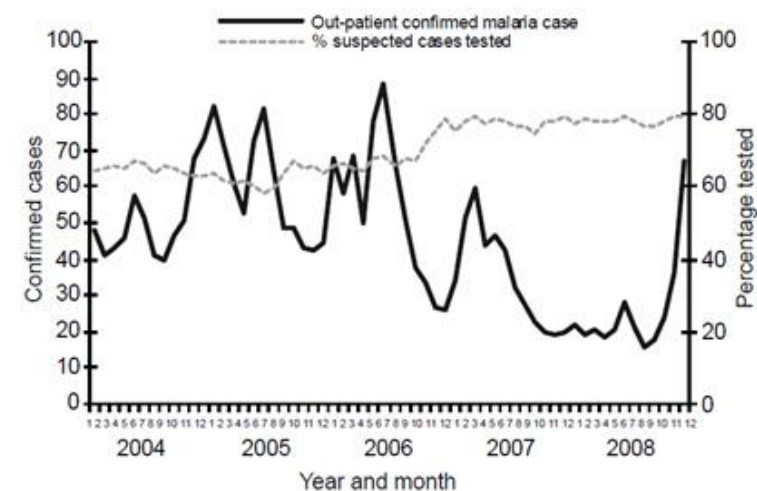
Low-to-moderate transmission

Countries and districts with low-to-moderate transmission should be able to reduce inpatient malaria cases and deaths by more than 50% within 12 months of mass distribution of LLIN, provided that the distribution achieves household possession of more than 60% and ITN use in children of more than 50%, as shown in Figure 5.2. Declines of less than 50% suggest that coverage is inadequate; in such cases, further investigation is needed.

High transmission

Countries and districts with high transmission are likely to require high levels of ITN coverage to achieve a reduction of more than 50% in inpatient malaria cases and deaths. The target should be near 100% household ITN possession and more than 80% ITN use in the entire population, not just in those under 5 years of age and pregnant women.

Outpatient confirmed malaria cases and percentage of suspected malaria cases tested with parasite-based test



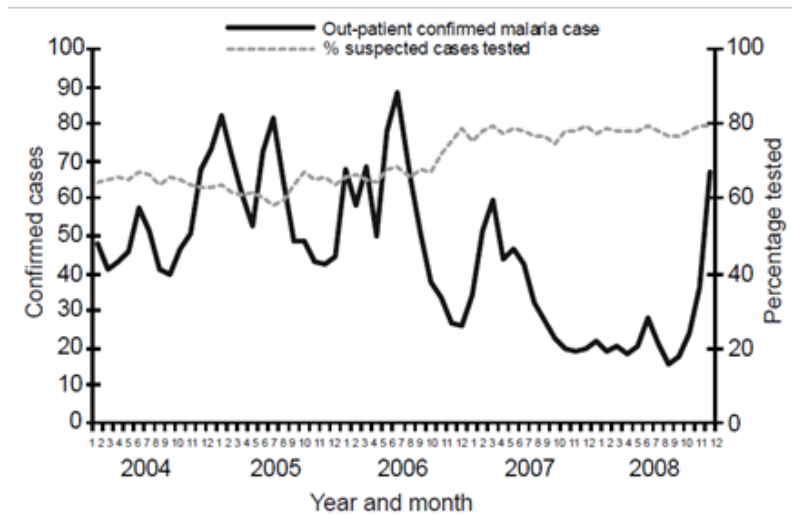
Low-to-moderate transmission

In areas of low-to-moderate transmission, the number of outpatient confirmed malaria cases should decline progressively and possibly quickly as transmission decreases due to improved malaria control (assuming that the percentage of suspected cases that are tested and completeness of reporting do not change). If there is no decline in confirmed cases, then it is unlikely that control has improved; in this case, investigation of coverage and efficacy of interventions is warranted.

High transmission

In areas of high transmission, the number of outpatient confirmed malaria cases is likely to decline progressively but slowly. If there is no decline in confirmed cases, then it is probable that transmission has not decreased sufficiently to produce changes in case incidence.

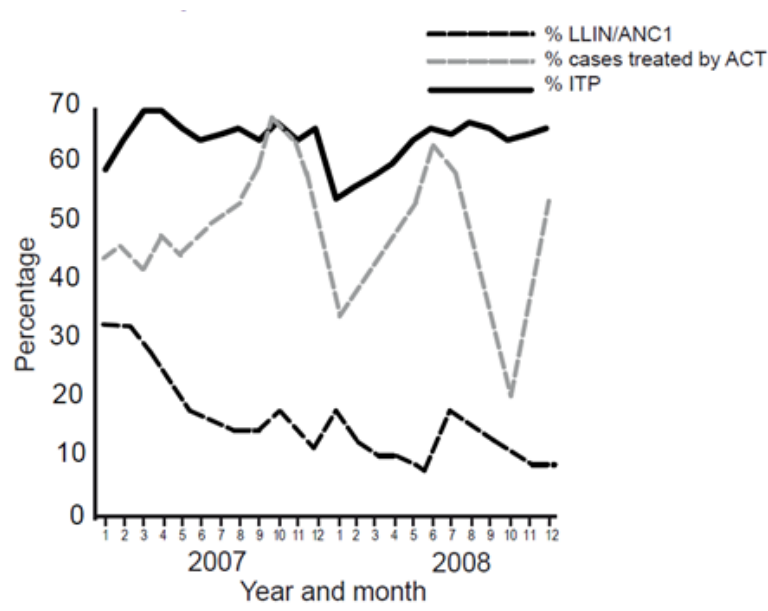
Outpatient all-cause cases and suspected malaria cases, all ages



The outpatient all-cause case curve is a “control” curve that indicates trends in health service attendance. For example, where malaria cases are declining, examination of the all-cause case curve would show whether there was also a decline in all-cause cases (or whether these had remained stable or increased). If malaria cases are declining markedly in the absence of a marked decline in all-cause cases, this indicates a true decline in malaria cases, probably due to malaria control interventions.

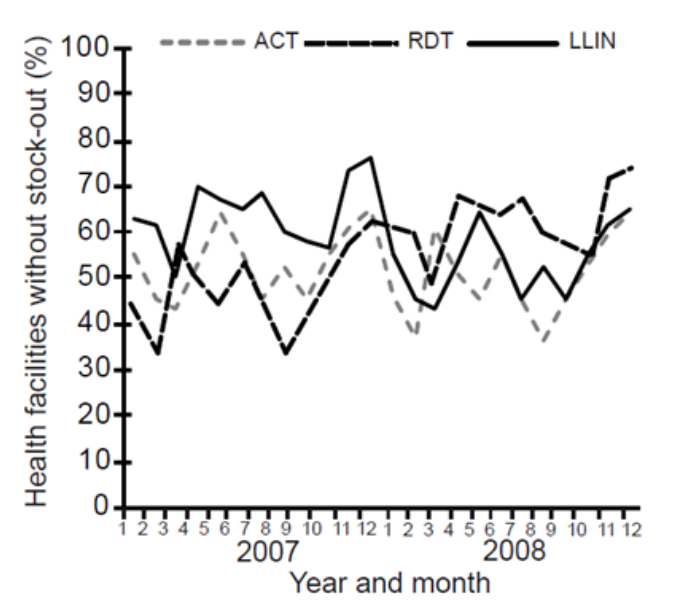
Interpretation of logistics and completeness-of-reporting graphs

Percentage coverage with patients treated with ACT (of expected), and ANC clients receiving ITN and IPT2



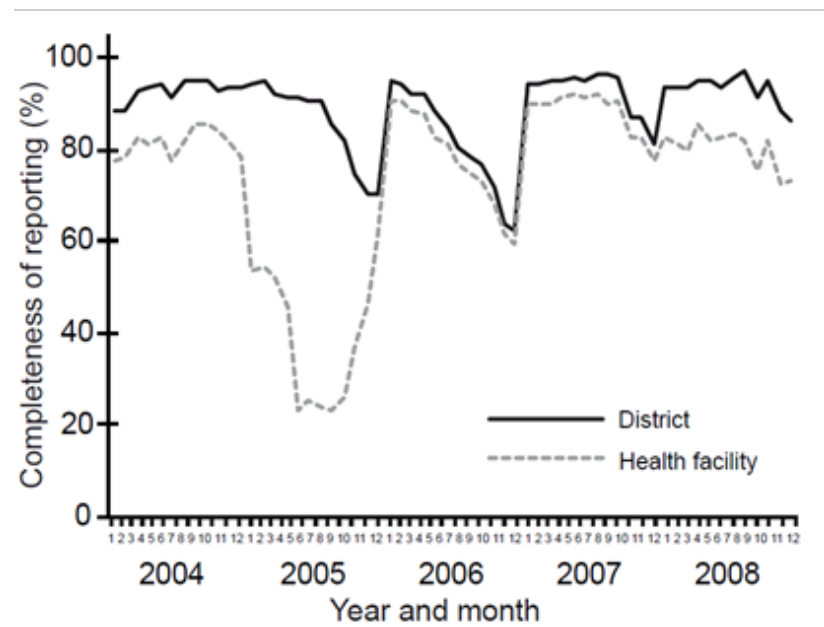
Ideally, the percentage of people treated with ACT and the percentage of ANC clients receiving ITN should reach 100%. The percentage of ANC clients receiving a second dose of IPT is unlikely to reach 100% because some ANC clients do not return for a second visit.

Percentage of health facilities without stock-outs, and without stock outs of ACT, RDT and LLIN



This indicator refers to the percentage of health facilities without stock-outs, rather than the percentage with stock-outs. Ideally, the percentage of health facilities without stock-outs should reach 100% each month.

Completeness of reporting and percentage of health facilities and districts that reported



The percentage of districts with complete reporting should reach 100%. The percentage of health facilities with complete reporting should reach more than 95%. Supervision should be targeted at health facilities and districts with incomplete reporting.

Appendix 5: Checklist for Supervising Surveillance and Response Activities at the Health Facility

District: _____ Health Facility: _____ Date of Supervisory Visit: _____

ACTIVITY	SUPERVISORY QUESTION	ANSWER	COMMENT (What Caused Problem)
Data collection to identify Suspected Cases within health facilities	How often do you collect information from the community about reports of suspected cases or deaths due to a priority disease or condition?		
Register cases	Are diagnoses of cases of priority diseases recorded in the clinic register according to the standard case definition?	Yes No	
Report	Do health staff use a standard case definition to report the suspected cases and outbreaks? Do you record information about immediately notifiable diseases on a case form or line list?	Yes No Yes No	
Analyze and Interpret	Do you plot the numbers of cases and deaths for each priority disease on a graph? Do you plot the distribution of cases on a map?	Yes No Yes No	
Investigate and Confirm Reported Cases and Outbreaks	If an epidemic-prone disease was suspected, was it reported immediately to the district office? For the cases of priority diseases needing laboratory tests seen since the last supervisory visit, how many had laboratory results? Are appropriate supplies available or set aside for collecting laboratory specimens during an urgent situation and show me the supply?	Yes No Number of results obtained: _____ Number of expected cases seen: _____ Yes No	
Respond	Are appropriate supplies available for responding to a confirmed case or outbreak (for example, immunization supplies and vaccine, ORS, antibiotics, and so on)? Please show me the supplies for carrying out a recommended response. Who is the outbreak coordinator for this facility? How often do you provide information and training in outbreak response to the staff of this facility?	Yes No Yes No Name: _____ Designation: _____ Training is done _____	
Provide Feedback	How often do you report information to the community? Do you receive the latest bulletin from the (central, subnational) level?	Report it _____ Yes No	

ACTIVITY	SUPERVISORY QUESTION	ANSWER	COMMENT (What Caused Problem)
Evaluate and Improve the System	Were the last 3 routine monthly reports sent to the district office? Were the last 3 routine monthly reports sent on time?	Yes No Yes No	
Epidemic Preparedness	What precautions do health staff (including laboratory staff) take routinely with all patients regardless of the patients' infection status? How do you estimate the number of supplies to set aside for use during an emergency situation?	Minimum level of standard precautions: _____ How supplies are estimated: _____	

FIELD DATA SHEET - ADULT COLLECTION

Other Anophelines N/A

John J. Jones

Appendix 7: Malaria Facility Supervision Checklist

Facility Supervision Checklist *(To be completed in duplicate and copy left at the facility)*

[Indicate N/A where the question is not applicable]

[NOTE: Even when asking YES/NO questions, in addition, kindly observe the practice to confirm the answers provided]

A. General section

1. Name of facility_____ Level of facility_____

2. Facility in charge_____ Contact: Tel. No._____

Email_____

3. Ownership (GoK, Private, NGO, FBO) _____

4. District_____ Province _____ Date of Supervision_____

5. Supervision Team Members:

	Name	Organization/Division	Designation
1			
2			
3			
4			
5			

6. Respondents:

	Name	Designation
1		
2		
3		

7. Does the facility have inpatient facility ☐ Yes ☐ No

B: Human Resource Capacity [Maximum YES score Available = 9]

8. Technical HR Capacity (where applicable)

CADRE	Available? Y/N	Number in health facility
Medical Officer		
Pharmacist		
Clinical Officer		
Pharmaceutical Technologist		
Nurses		
Lab. Technicians/Technologists		
Health Records Officer		
Public Health Technician/Public Health Officer		
Others (please specify)		

9. Training Details (Where applicable).

CADRE	Number in health facility	Number trained in malaria case management	Number trained in the last 1 year
Medical Officer			
Pharmacist			
Clinical Officer			
Pharmaceutical Technologist			
Nurses			
Lab. Technicians / Technologists			
Proportion trained		=No. trained /No. in health facility	=No. trained in last one year /No. in health facility

C: Delivery of Malaria Services and Best Practices [Maximum YES score available = 33]

The following questions should be asked to the clinicians engaged in malaria clinical management. Observations should be used to confirm the answers.

10. Istesting of ALL suspected malaria cases undertaken at you facility? If **No** skip Q11

☐ Yes ☐ No

11. Which test do you carry out to confirm malaria diagnosis? (Tick all that apply)

Microscopy ☐ Yes ☐ No

RDT ☐ Yes ☐ No

If the facility has RDTs, check the following

12. During the visit observed a health worker performing an RDT for malaria? If none, skip to 14

If observed, has the following been done correctly (As per the RDT Job aid)

- a. Blood collection ☐ Yes ☐ No
- b. Blood and buffer placed in the correct wells ☐ Yes ☐ No
- c. Enough time allowed before reading test results. ☐ Yes ☐ No
- d. Were the readings read correctly ☐ Yes ☐ No

13. RDTs storage at facility?

- a. Are RDTs stored in a cool, dry place away from the floor ☐ Yes ☐ No

14. What recommended 1stline anti-malaria medicine is used at your facility for the treatment of uncomplicated Malaria? [Tick YES if answer is given is Artemether- Lumefantrine (AL)]

☐ Yes ☐ No

15. What medicine is used at your facility for treatment of malaria in 1st trimester of pregnancy?

Quinine tablets ☐; AL ☐; SP ☐; Other (Specify).....

(Tick Yes if answer given is **Quinine tablets**) ☐ Yes ☐ No

16. What AL dosing schedule is used for a 20kg patient visiting your facility?

(Tick YES if answer is given is **"6 doses given over 3 days and 1st dose Directly Observed"**)

☐ Yes ☐ No

17. What is the 2nd line anti-malaria medicine used at your facility for treatment of uncomplicated malaria? (Tick Yes if DHAP) ☐ Yes ☐ No

18. Please mention 3 signs of severe malaria that a patient may present with?

(Correct responses include: Prostration; Altered level of consciousness; Multiple convulsions; Respiratory distress; Circulatory collapse; Pulmonary oedema; Jaundice; Haemoglobinuria; Abnormal bleeding)

(Tick YES if at least 3 correct signs are named) ☐ Yes ☐ No

19. What anti-malaria medicine is used to treat severe malaria in your facility?

(Tick YES if answer is given is **IV Quinine**) ☐ Yes ☐ No

20. In addition to giving the anti-malaria medicine, what other steps do you take in the management of severe malaria in this facility?

(Tick YES if **any** of the answers below are provided) ☐ Yes ☐ No

- Organize for referral (apply only to facilities without inpatient facilities)
- Manage complications

21. **Observe** the malaria drug dispensing procedure and state whether the health worker

(i) Directly observed the first dose ☐ Yes ☐ No

(ii) Gave adequate dispensing instructions to the patient which includes

a. Dosage ☐ Yes ☐ No

b. Timing ☐ Yes ☐ No

c. Advice on side effects profile ☐ Yes ☐ No

d. Advice on follow-up ☐ Yes ☐ No

22. Does the facility provide pregnant women with ITNs / LLINs? (Please check the ANC register to confirm) ☐ Yes ☐ No ☐ N/A ☐

If not, why not? (specify)

.....

LAB Questions

23. If the facility has a lab, **examine the Lab Register** to determine how reporting for parasitemia is done. Tick Yes if facility records any of the following:

a. + + + + ☐ Yes ☐ No

b. Parasites/200WBC ☐ Yes ☐ No

c. Parasites/microlitre of blood) ☐ Yes ☐ No

24. Does the lab report malaria parasite species? ☐ Yes ☐ No

25. Are adequate microscopic fields examined before a negative smear is reported? (Tick YES if 100 microscopic fields are examined) ☐ Yes ☐ No

ANC QUESTIONS (To be asked in Nyanza, Western and Coast provinces only)

26. Are pregnant women given SP as IPTp when they come for ANC your health facility?

☐ Yes ☐ No (Please check the ANC register to confirm)

27. What procedure is used in giving IPTp at your facility? *[Tick Yes if answer given is DOT and (if possible) you actually observe the IPTp being administered correctly (DOT)]*

☐ Yes ☐ No

Please comment below if incorrect procedure was observed:

28. At what times/intervals is IPT administered at your facility? *[Tick Yes if the following two answers are given: (i) 'every four weeks after quickening' or (ii) whenever the mother presents herself if interval between her visits is greater than 4 weeks]*

☐ Yes ☐ No

29. If a woman comes to the clinic when her pregnancy is later than 36 weeks, would you still administer IPTp? ☐ Yes ☐ No

30. Is IPTp given to HIV-positive pregnant women who are on daily cotrimoxazole? *(Please check the ANC register to confirm)[The correct answer is **NO**.*

Tick YES if this correct answer is given ☐ Yes ☐ No

31. **Observe** for availability of following in the ANC room

SP ☐ Yes ☐ No

Drinking water ☐ Yes ☐ No

Drinking cups ☐ Yes ☐ No

D: Availability of Malaria Commodities / Medicines [Maximum YES score Available = 7]

32. Have you had stockouts of any anti-malaria medicines over the last three months?

(Use the response to this question to complete the table below)

Malaria Commodity / Medicines	NO stock out was recorded in the last 3 months (Yes/No)	Duration of Stock out, if any
Sulfadoxine-Pyrimethamine(SP)		
Quinine tablets		
Quinine injection		
Artemether- Lumefantrine ⁴		
DHAP		
RDTs		
ANC / CWC Nets		

⁴ Stockout for AL implies total stockout of all bands of this medicine

E: Data Management and Reporting [Maximum YES score Available = 25]

33. **Review** the following documents and comment on their status

Document	Correctly filled and up to date? (Y/N)	Other Status**
Artemether- Lumefantrine- dispenser register		
Health Facility Monthly summary form for Malaria Medicines		
Bin card/ stock control card		
Receipt/ issue vouchers		
Laboratory register		
HMIS Inpatient register		
HMIS Outpatient Under 5 register		
HMIS Outpatient over 5 register		
ANC Register (check IPTp1 and IPT2 columns)		
CWC Register		
Facility Supervision Log Book		
ADR reporting Form (Yellow form) ⁵		
Poor quality medicine reporting form (pink form) ⁶		

**Document status key: a. *Correctly filled but not up to date* b. *Incorrectly filled* c. *Not available*

34. **Verify facility data for the previous month** (check the relevant Daily Activity Registers and compare actual figures with those reported to the district)

	Please indicate the previous month's tally obtained below		Are these the same values contained in the district report?
Out -patient malaria indicators	<5yrs	>5yrs	Y/N
Total number of outpatient malaria cases			
Number of malaria cases tested (Microscopy)			
Number of outpatient confirmed malaria cases (Microscopy)			
Number of malaria cases tested (RDT)			
Number of outpatient confirmed Malaria cases (RDT)			
Total number of outpatient confirmed Malaria cases (Microscopy + RDT)			
No of nets distributed to under 1 yrs			
Nets distributed to pregnant women			
In- patient Malaria indicators(<5 and>5)	<5yrs	>5yrs	Y/N
Inpatient malaria cases (confirmed with primary diagnosis of malaria at discharge)			
Inpatient malaria cases (confirmed & unconfirmed with primary diagnosis of malaria at discharge)			

⁵ The health worker to give a scenario where the form may be used

⁶ The health worker to give a scenario where the form may be used.

Total inpatient malaria deaths (with primary diagnosis as malaria)			
---	--	--	--

35. When did you last send your malaria medicines consumption summary report to the District?
(Ask to see copy at the facility - Tick YES if the last month's report was sent to district by the
5th day of the subsequent month) ☐ Yes ☐ No

F: Availability of Relevant Malaria Documents

[Maximum YES score Available = 15]

36. Check for the availability of the following documents.

	Document	Document Available (Y/N)	Comments
i.	Abridged NMS 2009 - 2017		
ii.	The National Guidelines for Diagnosis, Treatment and Prevention of Malaria in Kenya 3rd Edition		
iii.	2010 Diagnostics, Treatment and Drug Management set of Job Aids		
iv.	MIP orientation package (in MCH/FP clinic)		
v.	MIP Job Aids (in MCH/FP clinic)		
vi.	Pharmacovigilance guidelines		
vii.	Bench aides for microscopy (in the Laboratory)		
viii.	SOP or Job aid for performing RDT test procedure		
ix.	Laboratory diagnosis of malaria user's guide (in the Laboratory)		
x.	Inventory of ACSM materials		

37. Has the facility displayed health promotion materials covering the following areas:
- a. Need to seek prompt treatment for fevers ☐ Yes ☐ No
 - b. Recognition of symptoms and signs of severe malaria ☐ Yes ☐ No
 - c. Adherence to malaria treatment plan ☐ Yes ☐ No
 - d. Use of appropriate malaria prevention measures:
 - IPTp poster/brochures ☐ Yes ☐ No
 - LLINs posters/brochures ☐ Yes ☐ No

38. Overall achievements and challenges

.....

G: List at Most Three Gaps Identified and Actions Needed

No	Problems/ gaps	Action needed	Person to take action	By when action to be taken
1				
2				
3				

Name of Facility in charge: Signature:

Date: Rubber stamp:

Name of Leader of Supervision team:.....

Signature..... Date:.....

Appendix 8: Malaria District Supervision Checklist

District Supervision Checklist *(To be completed in duplicate and copy left at the District)*
[Indicate N/A where the question is not applicable]

A. General section

1. District _____ Province/County _____

1. District in charge _____ Contact: Tel _____
Email _____

2. Date of Supervision _____

4. Supervision Team Members:

	Name	Organization/Division	Designation
1			
2			
3			
4			
5			

5. Respondents:

	Name	Designation
1		
2		
3		

6. How many facilities does the district have?

- a. GoK _____
- b. Private _____
- c. NGO _____
- d. Faith-Based _____
- e. Municipal _____
- f. Others _____

B. Planning and Management [Maximum YES score Available = 4]

7. Does the district have a dedicated malarial focal person? ☐ Yes ☐ No

8. Are malaria control activities included in the district annual operational plan (AOP)? (*Asks for a copy of AOP*) ☐ Yes ☐ No

9. Does the district hold review meetings during which malaria control activities are discussed?

☐ Yes ☐ No

If yes, what is the frequency of holding such meetings? ☐ Monthly ☐ Quarterly

☐ Biannual ☐ Other (specify) _____

10. Has the district updated the partners' database detailing their contribution to malaria control in the district? (**Obtain the updated copy**) ☐ Yes ☐ No

C. Data Reporting and Analysis [Maximum YES score Available = 16]

11. Does the district have a Health Records Information Officer? ☐ Yes ☐ No

12. Has at least 1 district staff been trained on malaria medicines data management?

☐ Yes ☐ No

13. Review the following data reporting documents and comment on their status

Document	Correctly filled and up to date? (Y/N)	Other Status**
District Monthly Aggregation forms for malaria medicines		
District Monthly Summary Tool for malaria medicines		
HMIS Summary Reports		
DDSR Weekly Reports		
Malaria Partners' Database		

****Document status key:** a. Correctly filled but not up to date b. Incorrectly filled c. Not available

14. Review last quarter's reporting pattern for malaria medicines and calculate the average reporting rate.

(a) Is the overall reporting rate $\geq 70\%$? (i.e. number of facilities reporting out of the total facilities in the district) ? ☐ Yes ☐ No

(b) If $< 70\%$, what are the reasons for the low reporting rate?

15. Has the district been sending its malaria medicines consumption summary report to the national level in a timely manner? (**Ask to see copy at the district- Tick YES if the last month's report was sent to national level by the 20th day of the subsequent month**)

☐ Yes ☐ No

16. Does the district analyze its malaria data? ☐ Yes ☐ No

17. Are the following data elements calculated and up to date?

Analysis	Analysis done (Y/N)	Status
Annual trends of outpatient malaria cases (over 5yrs & under 5yrs) over the last 5 years		
Annual trends of confirmed malaria cases over the last 5 years		
Annual trends of confirmed malaria admissions over the last 5 years		
Annual trends of inpatient malaria deaths over the last 5 years		
Trends of the average facility monthly reporting rate for malaria medicines (for all facilities in the district)		
Trends for IPTp 1 and IPT 2 provision		

Status Key 1-Complete and up to date. 2-Availalable but not up to date 3-Not available

D. Training and Supervision [Maximum YES score Available = 12]

18. What malaria-related trainings have been undertaken in your district over the last 12 months? [Please insert details in the table below]

Course Name	Course Provider/Training Organization	Month of training

19. Is there an updated Facilities Contact List for all facilities in the district?

Yes ☐ No ☐

20. Does the district have a documented facilities supervision schedule? ☐ Yes ☐ No

21. How often is the integrated supervision conducted?

☐ Monthly ☐ Quarterly ☐ Not regular

• Tick Yes, if supervision done at least once every quarter. ☐ Yes ☐ No

22. (a). How many facilities has the district supervised in the last 3 months? Is the percentage of facilities supervised $\geq 70\%$? (compared with total facilities in district)

☐ Yes ☐ No

(b). If $<70\%$, what are the reasons for the low supervision coverage?

23. Does the district document supervision visits? ☐ Yes ☐ No (*ask to see the documentation*)
24. (a). Does the district give written feedback to the facilities after supportive supervision?
☐ Yes ☐ No
- (b). If yes, what is the date of the last report (*ask to see a copy*) _____
- (c). If no, what is the reason? _____
25. Did the district send a timely supervision report to the province and national level after completion of last supervision visits? (i.e. within 2 week of completing the supervision)
☐ Yes ☐ No
26. Has any team from the province level come for integrated supportive supervision in the last 6 months? ☐ Yes ☐ No
27. IF YES, did the supervisors perform any of the following activities?
- (a) Record Reviews ☐ Yes ☐ No
- (b) Review of the malaria data analysis done by district ☐ Yes ☐ No
- (c) Discuss problems associated with supervision and other malaria control activities in the district and provided recommendations? ☐ Yes ☐ No
28. Has the district received any written feedback from the supervisor after a supervisory visit in the last 6 months? (*ask to see report or documentation*) ☐ Yes ☐ No

E. Availability of Relevant Malaria Documents
[Maximum YES score Available = 13]

29. Indicate availability of the following malaria documents.

Document	Available? (Y/N)	Comments
NHSSP II		
National Malaria Policy 2010		
National Malaria Strategy 2009-2017		
(i). Complete version		
(ii). Abridged version		
Malaria Monitoring and Evaluation Plan 2009- 2017		
Inventory of ACSM Material		
Global Fund Operations Manual		
The National Guidelines for Diagnosis, Treatment and Prevention of Malaria in Kenya (3rd edition)		
IRS training manual (where applicable)		

Support Supervision Manual and Tools for supervision of Malaria Control Activities		
IVM guidelines		
EPR guidelines(whenever applicable)		
Malaria Communication strategy		
Others specify		

F. Advocacy, Communication and Social Mobilization (ACSM)

[Maximum YES score Available = 6]

30. Does the district hold stakeholders forums? ☐ Yes ☐ No
- How often are such forums held? _____
31. Are malaria issues discussed during these forums? ☐ Yes ☐ No
32. What are the channels that the district uses for health promotion/social mobilization?
- Barazas ☐ Yes ☐ No
- Religious groups ☐ Yes ☐ No
- Road shows/theatre groups ☐ Yes ☐ No
- Print media ☐ Yes ☐ No
- Others *(Please note them down)*

G. Emergency Preparedness [Maximum YES score Available = 5]

33. Check the district's DDSR reporting rate for the last four weeks – is the average rate $\geq 70\%$?
☐ Yes ☐ No
34. Is this weekly information shared by the following week with the following:
- i. DOMC ☐ Yes ☐ No
- ii. DDSR ☐ Yes ☐ No

(Questions 35 – 37 are for epidemic prone districts only)

35. Does the district have a written plan of epidemic preparedness and response?
☐ Yes ☐ No
36. Has the district had adequate emergency stocks of malaria drugs and supplies (that would last at least 4 weeks) at all times in the past 3 months? ☐ Yes ☐ No
37. How many malaria sentinel surveillance sites exist in the district?

38 Overall achievements and challenges

.....

.....

.....

.....

.....

.....

.....

I: List at Most Three Gaps Identified and Actions Needed

No	Problems/ gaps	Action needed	Person to take action	By when action to be taken
1				
2				
3				

Name of District Medical Officer of Health:Signature:

Date: Rubber stamp:

Name of Leader of Supervision team:

Signature..... Date:.....

Appendix 9: Facility Contact List

District: _____ Province: _____ Year: _____

Facility Name	Address	Office Phone No.	Contact Person	Designation	Mobile Phone No.	Email address
1						
2						
3						
4						
5						
6						
7						
8						
9						
10						
11						
12						
13						
14						
15						
16						
17						
18						
19						
20						
21						

Date last updated: _____

Appendix 10: DHMT Contact List

Province: _____ Year: _____

	District Name	Address	Office Phone No.	Contact Persons	Designation	Mobile Phone No.	Email address
1							
2							
3							
4							
5							
6							
7							
8							
9							
10							
11							
12							
13							
14							
15							

Appendix 11: District Supervision Activity Schedule

District: _____ Province: _____ Schedule for Period from: _____ to _____

Health Facility	Contact Person	Phone Number	SUPERVISION SCHEDULE											
			Jan	Feb	Mar	Apr	May	Jun	Jul	Aug	Sep	Oct	Nov	Dec
1														
2														
3														
4														
5														
6														
7														
8														
9														
10														

Appendix 12: Facility Score Sheets

Supervision Aspect	Maximum YES score Available	Total YES Recorded	Total N/A Recorded	Calculated % SCORE	COMMENTS
HR Capacity and Training Status				0.0%	
Delivery of Malaria Services and Best Practices				0.0%	
Availability of Malaria Commodities / Medicines				0.0%	
Data Management and Reporting				0.0%	
Availability of Relevant Malaria Documents				0.0%	
OVERALL SCORE	0	0	0	0.0%	

NOTE:

% Score Obtained = Total "YES" Recorded x 100 / (Max. "YES" Score - Total "N/A" recorded)

Obtain the score for each supervision aspect before calculating the overall score for the supervisee.

Appendix 13: District Score Sheet

Supervision Aspect	Maximum YES score Available	Total YES Recorded	Total N/A Recorded	Calculated % SCORE	COMMENTS
Planning and Management				0.0%	
Data Reporting and Analysis				0.0%	
Supervision				0.0%	
Availability of Relevant Malaria Documents				0.0%	
Advocacy, Communication and Social Mobilization (ACSM)				0.0%	
Emergency Preparedness (for districts)				0.0%	
OVERALL SCORE	0	0	0	0.0%	

NOTE:

$$\% \text{ Score Obtained} = \frac{\text{Total "YES" Recorded} \times 100}{(\text{Max. "YES" Score} - \text{Total "N/A" recorded})}$$

Obtain the score for each supervision aspect before calculating the overall score for the supervisee.

Appendix 14: Supervision Summary Report

District: _____ Province: _____ Report for
Period: From: _____ To: _____

	Facility Name	Date of supervision	Main findings	Actions taken	Recommendations	Responsible	Required support	Date Actions Due
1								
2								
3								
4								
5								
6								
7								
8								
9								
10								
11								

Report approved by:

Name: _____

Signature: _____

Date: _____

Appendix 15: Detailed Supervision Report Format

[For use by All Levels]

Within two weeks of completing the supervision, the supervision team should compile the detailed report using the report format below:

1. Introduction

- a. Report Background, e.g., a brief introduction of the district/province.
- b. Objective(s) of the supervisory visit
- c. Dates of the visit
- d. List of names/designations of members of the supervision team

2. Methods

- a. How the supervision was organized and carried out
 - i. Courtesy calls
 - ii. Review of previous reports
 - iii. Allocation of tasks to team members
- b. Tools used
- c. Approaches used
 - i. Direct observation
 - ii. Interviews
 - iii. Review of data
- d. Documents reviewed
- e. Other sources of information

3. Finding

- a. Detailed description of the findings
- b. Relating the current findings to the previous reports
 - i. Have action plans and recommendations been accomplished
 - ii. Are there recurrent issues

4. Actions taken

- a. Details of immediate actions taken during supervision
 - i. Corrective actions taken
 - ii. On the job training
 - iii. Facilitation

5. Recommendations and action plans

- a. Recommendations given and to whom.
- b. Action plans and the agreed timelines.

6. Annexes

- a. Supervision summary
- b. Facility/District/Provincial Score sheet
- c. List of partners (Partners' Database)
- d. Financial statement

Appendix 16: IDSR Weekly Summary Reporting Form

MOH 505

County _____ District _____ Health Facility _____ Epi Week _____ Week ending _____ Month _____ Year _____

[illegible]

Diseases, Conditions or Events	< 5 years		≥ 5 years		Total	
	Cases	Deaths	Cases	Deaths	Cases	Deaths
AEFI*						
Acute Jaundice						
Acute Malnutrition						
AFP (Poliomyelitis)**						
Anthrax						
Cholera						
Dengue						
Dysentery (Bacillary)						
Guinea Worm Disease (Dracunculiasis)						
Malaria						
Maternal deaths						
Measles						
Laboratory	< 5 years		≥ 5 years		Total	
	Tested	+ve	Tested	+ve	Tested	+ve
Malaria						
Shigella Dysentery						
Tuberculosis (MDR/XDR)						
Typhoid						

*****Any public health disease, condition or event of national or international concern (Infectious, zoonotic, food borne, chemical, radio nuclear, or due to unknown condition

Reported by: _____

Designation _____

Sign _____

Date _____

Reporting Instructions

Health Facility Level: Send a copy to DMOH or DDSC every Monday and file a copy

District Level: Reviews all health facility reports for correctness then enters the data in the electronic IDSR system. Files the health facility copies

Surveillance week: A week starts on Monday and ends on Sunday

DIVISION OF MALARIA CONTROL
Ministry of Public Health and Sanitation
P.O Box 19982 – 00202 KNH
Nairobi, Kenya
head.domc@domckkenya.or.ke

