

Can we eliminate malaria with current intervention tools?

Lessons from mathematical models for malaria control & elimination

María-Gloria Basáñez

(with slides from Azra Ghani & Emma Dawes)

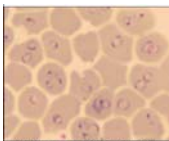
Department of Infectious Disease Epidemiology
MRC Centre for Outbreak Analysis & Modelling

m.basanez@imperial.ac.uk

a.ghani@imperial.ac.uk

Human malaria *Plasmodium* species

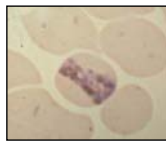
Plasmodium falciparum



Africa
SE Asia
Latin America

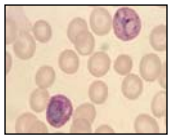
(original image provided by Steve Aitky)

Plasmodium malariae



Africa
SE Asia
Latin America

Plasmodium vivax



Middle East
Asia
Western Pacific
Latin America
Africa

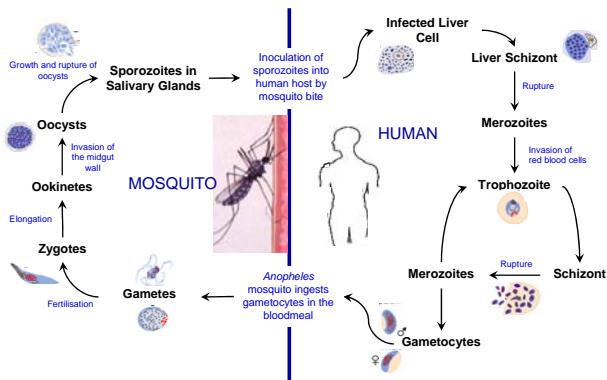
(original image by Mark Lontie)

Plasmodium ovale




Tropical Africa
West Pacific

Life-cycle of *Plasmodium falciparum*




Imperial College London **Human malaria is transmitted by *Anopheles* species mosquitoes** MRC Medical Research Council

Anopheles gambiae s.l.




Main vector in Africa

Anopheles funestus




Africa

Anopheles albimanus



Central America


Anopheles stephensi



India

A, B, C = CDC Image Library
D = The Wellcome Trust

Imperial College London **Distribution of *Anopheles* vectors** MRC Medical Research Council

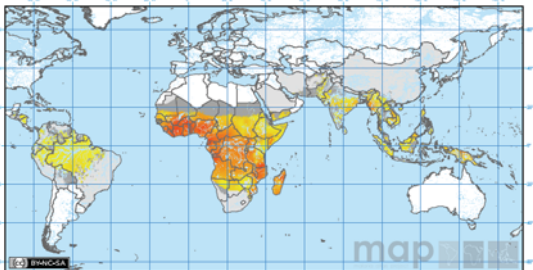


Anopheles

- No vector
- albimanus
- ambusius
- anthropophilus
- andersoni
- andersoni and funestus
- aquasalis
- amplicornis
- barbivittatus
- subulifacies
- barlingi
- grisea
- leucis
- flavipes
- funestus and andersoni
- funestus and gambiae s.s.
- gambiae s.s.
- gambiae s.s. and funestus
- maculipes
- manipurensis
- mesasiaticus
- mesasiaticus
- multicolor
- harveyi
- harveyi
- punctulata group
- phosphoreus
- psittacula
- superpictus
- sublimatus
- sectator
- argyritarsis
- argyritarsis
- albipennis
- indicus
- phosphoreus
- subulifacies

Imperial College London **Prevalence of *P. falciparum* in 2007** MRC Medical Research Council

The spatial distribution of *Plasmodium falciparum* malaria endemicity in the World



map

0 2,500 5,000 10,000 15,000 Kilometers

Water
Malaria free
PfPR < 0.1%

Parasite rate (in units of PfPR₂₋₁₀ 0-100%)
0 100

Source: www.map.ox.ac.uk

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Burden of Disease 2007

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- Estimated 450 million (95% Credible Intervals 349-552 million) cases of malaria
- Majority of cases in population-dense areas e.g. India, Nigeria

Hay et al. (2010) PLoS Med 7(6)

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The Triangle of Transmission

MRC Medical Research Council

Plasmodium

Oocysts in mosquito abdomen

Plasmodium sporozoites

Infective stages to humans

Anopheles

Human

Gametocytes in blood

Infective stages to vectors

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Interrupting the Triangle of Transmission

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Transmission-blocking vaccines (e.g. preventing oocyst formation). Refractory, GM mosquitoes

Vaccines against pre-erythrocytic stages, RTS,S

Gametocytocidal treatment (e.g. ACT, Primaquine)

Plasmodium

Oocysts in mosquito abdomen

Plasmodium sporozoites

Infective stages to humans

Anopheles

ITNs, LLINs, IRS

Human

Gametocytes in blood

Infective stages to vectors

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History of Malaria Control

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GMEP: Global Malaria Eradication Programme
 RBM: Roll Back Malaria (to halve malaria burden by 2010)
 MalERA: Malaria Eradication Research Agenda

Wernsdorfer, Hay & Shanks (2009)

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The Components of Transmission

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The probability of vectors surviving the n days of sporogony and beyond, $p^n(-\ln(p))$

Plasmodium

Oocysts in mosquito abdomen

Infective stages to humans

The probability of infection establishing in humans, b

Anopheles

The Vector to Human ratio, $m = V/H$

The biting rate per vector on humans, a

Human

The duration of infectiousness, $1/r$

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The Basic Reproduction Ratio, R_0 , for Malaria

MRC Medical Research Council

How long does a person remain infectious? $1/r$

How many times a day is a person bitten by potential vectors? ma

What fraction of bites on infectious humans infect a mosquito? c

m – ratio of mosquitoes to humans
 p – probability a mosquito survives one day
 n – number of days required for sporogony
 a – number of human bites, per mosquito, per day

What fraction of mosquitoes survive sporogony? p^n

How many human blood meals does a vector take over its lifetime? $a/-\ln p$

What fraction of infectious bites infect a human? b

$$R_0 = \frac{ma^2bc}{r(-\ln p)} p^n$$

From Smith, Smith & Hay (2009)

Imperial College London **The Basic Reproduction ratio, R_0 , for Malaria** MRC Medical Research Council

$R_0 > 1$ (~2)

$$R_0 = \frac{ma^2 b P^n}{r (-\log_e P)}$$

$R_0 < 1$ (~1/2)

Average number of secondary cases originated during infectiousness by a primary case introduced in a wholly susceptible population

Each case generates on average 2 cases, there will be an epidemic Each case generates less than 1 case on average, malaria will die down

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- Delay in the parasite development / multiplication within the mosquito (sporogony) is the "weakest link" in the transmission cycle because sporogony takes an appreciable portion of the vector life-span
- Interventions that reduce adult mosquito life-span (e.g. IRS with DDT) have the greatest potential to reduce R_0 via reducing the probability of daily survival, p (p^n is strongly non-linear)
- DDT subsequently main intervention in GMEP

n Within-vector latency period (sporogony)
 p Probability of vector daily survival
 $\mu_v = -\ln(p)$ Per capita vector mortality rate
 p^n Probability of vector surviving sporogony

The expectation of infective life

Imperial College London **Elimination Strategies** MRC Medical Research Council

- Which interventions, alone and/or in combination, have the potential to achieve local elimination and how best to combine such strategies to achieve elimination?
- When should interventions be initiated, what effort is needed & how long will it take?

Low, unstable transmission

SPR < 5% in lever cases* < 1 case/1000 population at risk* 0 locally acquired certification cases WHO

3 years ↓

High, stable transmission

Mendis et al. (2009) *Trop Med Int Hlth* 14: 802-809

Box 1 Definitions (WHO 2006a)

Malaria control is reducing the disease burden to a level at which it is no longer a public health problem.

Malaria elimination is interrupting local mosquito-borne malaria transmission in a defined geographical area, i.e. 0 incidence of locally contracted cases.

Malaria eradication is the permanent reduction to 0 of the worldwide incidence of malaria infection caused by a specific agent; i.e. applies to a particular malaria parasite species.

Mendis et al. (2009) *Trop Med Int Hlth* 14: 802-809

- *Where is elimination achievable? Where should initial efforts be focused?*

- Shrink the map?
- Focus resources on high transmission foci?



Malaria Atlas Project (www.map.ox.ac.uk)

➢ **Vector Targets**

- Long-lasting insecticide-treated nets (LLIN)
- Indoor Residual Spraying (IRS)
- Spatial repellents (House screening)
- Larval control (Source reduction)



➢ **Parasite within Human Targets**

- Switch to ACT regimens as first-line therapy
- Mass Drug Administration (MDA) or Mass Screening & Treating (MSAT)
- Pre-erythrocytic vaccine (RTS'S)



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Intervention Impact

- Currently observing wide-spread declines in malaria infection prevalence and disease
- Much of the evidence for intervention impact is observational, e.g. Malaria in KwaZulu-Natal, South Africa

Insecticide 1: DDT for IRS in KwaZulu Natal

Insecticide 2: IRS in Mozambique

ACT: Artemether-Lumefantrine for uncomplicated malaria

Barnes et al. (2005) *PLoS Med* 2(11):e330

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Questions for Mathematical Models

- In which settings are current tool sufficient to reduce prevalence to low levels (PfPR < 1%) ?
- What tools are needed to prevent re-introduction?
- How can we use models to inform strategic planning at a local, national, regional and continental scale?

- **Need models which include multiple interventions across different transmission settings**
- **Transmission measured by the Entomological Inoculation Rate (EIR): The average number of infectious bites received per person per year (if the person is maximally exposed to mosquitoes during the night) = ibppy**

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Transmission Intensity: EIR and Parasite Prevalence

- Marked variation in the average number of infectious bites individuals are exposed to (Entomological Inoculation Rate – EIR)
- Determines the reproduction number (R_0) in any setting as well as endemic prevalence

Griffin et al. (2010) *PLoS Med* 7(8)

Smith et al. (2007) *PLoS Biol* 5(3):e42

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- Increasingly apparent that sub-patent infection (not detectable under microscopy) may play a key role in sustaining transmission

Microscopy prevalence	Median PCR prevalence (IQR)
75.0 - 100	92 (84.0-96.5)
50.0 - 74.9	79 (68.4-83.6)
30.0 - 49.9	61.7 (52.4-71.1)
20.0 - 29.9	49.9 (28.0-61.9)
10.0 - 19.9	35.4 (24.3-47.3)
5.0 - 9.9	34.5 (10.0-24.2)
0.0 - 4.9	10.4 (3.4-17.6)

Okell et al. (2009) J Infect Dis 200(10):1509-17

Imperial College London **Human Infectious Reservoir** MRC Medical Research Council

- "Infectious reservoir":
 - Age-specific biting rate x prevalence x infectiousness x population size
- Defines where interventions need to be targeted to reduce transmission rather than control disease
- Depends on:
 - Parasite prevalence by age
 - Gametocytaemia
 - Onward infectiousness
 - Treatment

Griffin et al. (2010) PLoS Med 7(8)

Imperial College London **The Importance of Vector Species** MRC Medical Research Council

- Key aspects of mosquito behaviour:
 - Endophagy: propensity to bite indoors
 - Endophily: propensity to rest in the house after feeding
 - Human Blood Index (HBI): propensity to bite humans versus e.g. cattle
- Three key vector species in Africa:
 - An. gambiae s.s.* – dominant vector species, high endophagy & endophily, high HBI
 - An. arabiensis* – more common in less humid times of the year, low endophagy & low HBI
 - An. funestus* – breeds in swamp areas, dependent on landscape, high HBI

Jan Apr Jul Oct

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Transmission Settings

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- Consider different settings characterising across Africa:
 - Transmission intensity (EIR)
 - Seasonality Index (proportion of EIR occurring within the peak 3 months of transmission)
 - Vector species combinations

Location	Population	Reported (fitted) annual EIR (ibppy)	Type of transmission	Anopheles species composition
Kjenjojo, Uganda	Rural	7 (3)	Low, perennial	L 65% <i>An. gambiae</i> s.s., 35% <i>An. funestus</i>
Maputo, Mozambique	Rural	28 (46)	Moderate, perennial	M 46% <i>An. funestus</i> , 42% <i>An. arabiensis</i>
Kinkole, DRC	Rural	48 (43)	Moderate, perennial	M Nearly 100% <i>An. gambiae</i> s.s.
Nkoteng, Cameroon	Rural	94 (81)	Moderate, perennial	M 72% <i>An. funestus</i> , 28% <i>An. gambiae</i> s.s.
KND, Ghana	Rural	630 (586)	High, seasonal	H 60% <i>An. gambiae</i> s.s., 40% <i>An. funestus</i>
Matimbwa, Tanzania	Rural	703 (675)	High, seasonal	H 85% <i>An. gambiae</i> s.s., 10% <i>An. funestus</i> , 5% <i>An. arabiensis</i>

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Scenarios for Intervention Packages

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- Prior to 2000:**
 - assume the only intervention available was treatment with Sulphadoxine-Pyrimethamine (SP)
- From 2000 to 2010:**
 - increase LLIN use from 0% to 20% (*Noor et al. 2009 BMC Public Health 9:369*)
 - switch to ACT as first-line therapy
- From 2010:**
 - introduce range of intervention packages
- Range of endpoints:**
 - change in parasite prevalence
 - change in EIR
 - time to reaching parasite prevalence of <1%

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Realistic Coverage

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- RBM goals are 80% coverage of ITNs
- Perfect continued usage is unrealistic
- With decaying efficacy, relatively low *effective* coverage

Slow scale-up ———
 Rapid scale-up ———
 Rapid, no drop-out ———
 Rapid, no drop-out, 95% ———

Imperial College London **Impact of scale-up of LLINs, Low Transmission** MRC Medical Research Council

➤ Increasing coverage to RBM target of 80% can reduce transmission to <1% prevalence in low transmission areas only

Chart C: Parasite prevalence

Region	Transmission Level	Baseline	Rapid	Rapid, no drop-out
Kampala Uganda	L	~0.05	~0.05	~0.05
Kinshasa DRC	M	~0.15	~0.15	~0.15
Maputo Mozambique	M	~0.15	~0.15	~0.15
Nkolong Cameroon	M	~0.25	~0.25	~0.25
KND Ghana	H	~0.55	~0.55	~0.55
Matimba Tanzania	H	~0.55	~0.55	~0.55

Chart D: Absolute reduction in parasite prevalence

Region	Transmission Level	Baseline	Rapid	Rapid, no drop-out
Kampala Uganda	L	~0.02	~0.02	~0.02
Kinshasa DRC	M	~0.05	~0.05	~0.05
Maputo Mozambique	M	~0.05	~0.05	~0.05
Nkolong Cameroon	M	~0.08	~0.08	~0.08
KND Ghana	H	~0.12	~0.12	~0.12
Matimba Tanzania	H	~0.12	~0.12	~0.12

Imperial College London **Addition of IRS and MSAT, Moderate Transmission** MRC Medical Research Council

➤ Frequent rounds of IRS could reduce prevalence to low levels in areas with moderate transmission

➤ With additional mass screening and treatment (MSAT) further reductions can be achieved

Chart G: Parasite prevalence

Region	Transmission Level	LLINs only	MSAT	IRS	MSAT and IRS
Kampala Uganda	L	~0.05	~0.05	~0.05	~0.05
Kinshasa DRC	M	~0.15	~0.15	~0.15	~0.15
Maputo Mozambique	M	~0.15	~0.15	~0.15	~0.15
Nkolong Cameroon	M	~0.25	~0.25	~0.25	~0.25
KND Ghana	H	~0.55	~0.55	~0.55	~0.55
Matimba Tanzania	H	~0.55	~0.55	~0.55	~0.55

Chart H: Absolute reduction in parasite prevalence

Region	Transmission Level	LLINs only	MSAT	IRS	MSAT and IRS
Kampala Uganda	L	~0.02	~0.02	~0.02	~0.02
Kinshasa DRC	M	~0.05	~0.05	~0.05	~0.05
Maputo Mozambique	M	~0.05	~0.05	~0.05	~0.05
Nkolong Cameroon	M	~0.08	~0.08	~0.08	~0.08
KND Ghana	H	~0.12	~0.12	~0.12	~0.12
Matimba Tanzania	H	~0.12	~0.12	~0.12	~0.12

Imperial College London **IRS and Vector behaviour, Moderate Transmission** MRC Medical Research Council

➤ Interventions will have different impact in settings with similar EIR (e.g. moderate transmission) but different vector species

➤ IRS and ITNs unlikely to have sufficient impact if outdoor-resting mosquitoes are common (Maputo)

Chart C: Maputo, Mozambique

EIR (fitted) = 46
46% *An. funestus*, 42% *An. arabiensis*

Chart D: Nkolong, Cameroon

EIR (fitted) = 81
72% *An. funestus*, 28% *An. gambiae* s.s.

Legend:

- LLINs only
- MSAT
- IRS
- MSAT and IRS
- Twice yearly MSAT
- Twice yearly IRS
- Twice yearly MSAT and IRS
- MSAT and IRS, no drop-out from LLINs
- Twice yearly MSAT and IRS, no drop-out from LLINs

Imperial College London **High Transmission Settings** MRC Medical Research Council

- Current tools are unlikely to be sufficient to reach the pre-elimination threshold of 1% parasite prevalence in areas of high transmission
- However, substantial declines in prevalence can be achieved
- Interventions will greatly reduce incidence of disease / clinical burden

E KND, Ghana
EIR (fitted) = 586
60% *An. gambiae*,
40% *An. funestus*

F Matimba, Tanzania
EIR (fitted) = 675
85% *An. gambiae*,
10% *An. funestus*
5% *An. arabiensis*

LLINs only MSAT and IRS
MSAT Twice yearly MSAT
IRS Twice yearly IRS
Twice yearly MSAT and IRS
MSAT and IRS, no drop-out from LLINs
Twice yearly MSAT and IRS, no drop-out from LLINs

Imperial College London **Vaccine Impact, Low transmission settings** MRC Medical Research Council

- RTS,S vaccine in Phase III trials prevents infection (pre-erythrocytic vaccine – PEV)
- Efficacy ~50% from Phase II studies
- Likely to be delivered via Expanded Programme of Immunisation (EPI)
- Additional impact on transmission greatest in low transmission settings

A Kjeriyojo, Uganda

B Kjeriyojo, Uganda

LLINs only PEV at EPI MSAT MSAT and PEV at EPI MSAT and IRS Twice yearly MSAT and IRS
Mass PEV MSAT, IRS and mass PEV Twice yearly MSAT and IRS, and mass PEV

Imperial College London **Take Home Messages. I** MRC Medical Research Council

- The Basic Reproduction Ratio (R_0) of malaria depends on:
 - entomological components (vector density, biting rate on humans, probability of daily survival)
 - components of the vector-parasite interface (probability of successful establishment in the vector, duration of sporogony)
 - components of the human-parasite interface (probability of successful establishment in the human, duration of infectiousness)
- Elimination programmes aim at reducing R_0 below 1 by implementing interventions that target the above
- Mathematical models provide useful tools to summarise and update current knowledge on the biology and epidemiology of malaria and its transmission in a quantitative framework, so that impact of interventions can be measured / anticipated

Take Home Messages. II

Mathematical models are important in all stages of malaria elimination programs:

- **Planning:** Determining what is achievable, with what tools
 - **Reducing transmission:** Identifying optimal combinations and strategies
 - **Monitoring:** Helping to design appropriate surveillance strategies
 - **Holding the line:** Advising on tools needed to prevent re-introduction
- Can also aid in defining properties of new tools needed in areas where current tools are insufficient
 - Importance of local vector species composition (feeding / resting behaviour) as well as overall transmission intensity
 - Currently available tools insufficient to eliminate malaria in high transmission settings (but can help reduce disease / mortality burden)
 - So far model assumes no development of insecticide or drug resistance
 - **Need to combine epidemiological with evolutionary models**

Malaria Research Group



Neil Ferguson



Maria-Gloria Basañez



Tom Churcher



Bhargavi Rao

With additional input from



Bob Sinden



Danail Stoyanov



Déirdre Hollingsworth



John Marshall



Lucy Okell



Michael White



Teun Doustema (LSHTM)



Chris Drakeley (LSHTM)



Jamie Griffin



Azra Ghani



Wes Hinsley



Tini Garske