Operational strategies to achieve and maintain malaria elimination

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Summary

Present elimination strategies are based on recommendations derived during the Global Malaria Eradication Program of the 1960s. However, many countries considering elimination nowadays have high intrinsic transmission potential and, without the support of a regional campaign, have to deal with the constant threat of imported cases of the disease, emphasising the need to revisit the strategies on which contemporary elimination programmes are based. To eliminate malaria, programmes need to concentrate on identification and elimination of foci of infections through both passive and active methods of case detection. This approach needs appropriate treatment of both clinical cases and asymptomatic infections, combined with targeted vector control. Draining of infectious pools entirely will not be sufficient since they could be replenished by imported malaria. Elimination will thus additionally need identification and treatment of incoming infections before they lead to transmission, or, more realistically, embarking on regional initiatives to dry up importation at its source.

This is the third in a Series of four papers about malaria elimination
Introduction

The Roll Back Malaria strategy of Scaling Up for Impact through universal coverage with effective interventions, supported by an increase in malaria funding, has achieved low rates of malaria transmission in some areas and consequently a much reduced disease burden. Some countries, including several with historically medium-to-high transmission, are nearing a state of controlled low-endemic malaria, and policy makers have a decision to make: accept low rates of malaria transmission with a strategy of consolidation of control or redirect activities with the aim to eliminate malaria.

During the Global Malaria Eradication Program (GMEP), WHO Expert Committee reports described specific activities of an elimination programme through its defined phases, and provided advice based on years of experience from field campaigns. Since the 1970s, when WHO shifted the short-term strategic aim to control and relegated eradication to a long-term goal, there has been little scientific inquiry into or strategic thought about the theory, goals, and best approaches for national elimination. At the same time, many countries considering elimination nowadays have higher intrinsic transmission potential than do those that eliminated malaria during the GMEP and have to plan to maintain elimination despite continual importation of infections. Accordingly, the decision to move from controlled low-endemic malaria to elimination needs politicians, policy makers, and programme managers to have an informed understanding of the operational requirements for a contemporary elimination strategy so that they can set realistic goals and timelines that are relevant to malaria epidemiology nowadays.

The decision to convert a malaria programme that has successfully achieved a high level of control, such that malaria is no longer a major public health problem, into an elimination programme is complex and should take into account technical, operational, and financial feasibility. There is a broad consensus about the strategies that are needed to achieve controlled low-endemic malaria, which are based on universal coverage with prevention and treatment measures—all of which have a strong evidence base from empirical trials, observational studies, and routine monitoring and evaluation. However, elimination cannot be achieved by doing more of the same; transition from sustaining control to elimination demands additional activities. In the third paper in this Series, we review the activities needed to achieve and maintain malaria elimination in areas that have already reduced transmission to very low rates by identification of the essential operational changes that have to accompany a switch in focus from burden reduction to interruption of transmission. In doing so, several important knowledge gaps are identified that, in some cases, makes it challenging to provide evidence-based guidance about how to eliminate malaria.

Key messages

- The most important operational difference between a control and an elimination programme is the concentration of activities to identify and attack foci of clinical and asymptomatic infections that perpetuate transmission
- Detection and curing of the high proportion of infections needed to interrupt transmission requires a robust surveillance system that combines passive and active case detection methods with rapid response, with radical treatment and targeted vector control
- Most malaria-endemic countries considering elimination should aim to prevent importation of infections through proactive case detection at the border, screening of high-risk migrants, and implementation of cross-border and regional initiatives that can reduce transmission at the source of migration
- Because elimination has a known quantitative goal to end endemic transmission and reduce the number of locally acquired cases to less than a specific threshold, monitoring systems incorporating extremely sensitive laboratory techniques such as PCR, genotyping, and serology have to be put in place to track progress.

- Malaria elimination cannot be business as usual, but needs a systemic and new programmatic approach supported by political and financial commitment, ideally throughout an entire region of nations.

**Differences between control and elimination**

The programmatic focus of a country seeking to control malaria as a public health problem involves the effective treatment of clinical malaria that is detected through passive surveillance integrated into the public health infrastructure and prevention of disease through high coverage with vector control measures. The main determinant of an elimination campaign is that, by contrast with a programme designed to maintain controlled low-endemic malaria, it seeks to interrupt endemic transmission and prevent its re-establishment. Prerequisites for either state include scaling up and maintaining high rates of effective coverage of control measures such as longlasting insecticide-treated nets or indoor residual spraying, or both; rapidly detecting, diagnosing, and treating malaria cases with effective drugs; and securing sufficient funding to sustain the broader control programme. Only after low rates of malaria transmission have been achieved (a community parasite prevalence of around 1% or less for *Plasmodium falciparum*) can activities of an elimination programme substantially differ from a programme that consolidates control. The most important difference between acceptance of low-parasite prevalence and seeking to interrupt endemic transmission is the concentration of activities towards identification of residual transmission foci and intensification of efforts to eliminate the last few infections. Such an active campaign of case detection and response, coupled with directed vector control efforts, should root out not only clinical cases but also asymptomatic infections that potentially perpetuate transmission.

The strategy of the GMEP was to eradicate malaria everywhere, and guarding against imported malaria was given only little attention. Nowadays, however, as individual countries consider elimination, without accompanying reductions by their neighbours, this strategic reorientation from general scale-up of control measures to focused case detection and intervention has to be accompanied by the development of effective strategies to identify imported cases and prevent reintroduction of transmission. Imported malaria from outside the country could otherwise replenish the endemic reservoir of infections, and where a competent vector remains, imported infections can lead to indigenous (secondary) transmission and resurgence. Elimination therefore theoretically needs: (1) elimination of the mosquito vector so that transmission cannot re-emerge; (2) blockade of the flow of imported infections from endemic areas; or (3) reduction of the risks of infection at their origin. The first of these options is considered operationally unachievable and is not recommended. The second would necessitate either closing the borders of the country seeking elimination or setting up a system of border screening that could successfully identify and treat incoming infections. The third requires that all neighbouring countries from which substantial population movement into the eliminating country occurs also achieve transmission reductions to very low rates.

**Methods and strategies to interrupt local transmission**

Halting endemic transmission and draining the reservoir needs reduction of $R_0$, the basic reproduction number under control, to less than 1. To drain the reservoir within a reasonable
timeframe, mathematical models suggest that $R_c$ should be less than 0.5. Although reductions to such a rate might be possible on average throughout a country through the same scale-up of vector control activities that are necessary to achieve low parasite prevalence, foci will remain in which such reductions are not achieved. Interruption of transmission in these areas will require additional active measures, including identification of infections even if asymptomatic, effectively treating infections before onward transmission can occur, and intensification and adaption of focal vector control activities.

**Detection of infections through surveillance**

Although historically a strong emphasis was placed on surveillance, nowadays it is more often perceived as a monitoring and evaluation method. However, the GMEP, and recent mathematical modelling to assess the feasibility of malaria elimination on the islands of Zanzibar, show that surveillance is a pivotal component of any programme aiming to interrupt transmission completely. The surveillance package should always include the response, such as targeted vector control measures or radical treatment, which is triggered by case identification and thus directly contributes to the reduction of transmission. As such, surveillance becomes an essential elimination intervention in itself. However, surveillance methods and the different laboratory techniques used differ in their ability to detect all clinical and asymptomatic cases, and the pool of *Plasmodium vivax* and *Plasmodium ovale* infections cannot be directly defined because there is no known method to detect the hypnozoite stages in the liver.

Passive case detection involves a system in which data are routinely received by a central health authority based on a set of rules and laws that need a health-care provider or health facility to report some diseases or disorders on an ongoing basis and at specific intervals (weekly, monthly, yearly). During the GMEP, health systems were generally considered to have little geographical coverage and thus generally to be insufficient for surveillance for elimination. Additionally, apart from limitations related to the precision of the diagnostic methods and completeness of treatment, passive case detection has other inherent weaknesses in detection of all new infections in the population (figure 1). For Zanzibar, taking into account the factors shown in figure 1, passive case detection is estimated to identify at best 40% of all new infections.

Additional effort needs to be made to optimise passive case detection, not only by ensuring access to malaria diagnosis and treatment, which should ideally be free in both public and private sectors, but also by improving health-seeking behaviour to reporting facilities and ensuring high testing rates with highly sensitive diagnostic tests (panel 1). Many of these conditions are equally important for controlled low-endemic malaria, but passive case detection alone is unlikely to be an adequate method as a pathway to elimination. In some settings, maintenance of elimination might be adequately supported only through routine passive case detection, as exemplified in the USA and large parts of Europe. However, case investigation remains a minimum requirement as long as a competent vector exists to perpetuate transmission (table).

Pampena defined active case detection as “the search for fever cases [...] performed by house-to-house visits at regular intervals in every locality of the malarious area”, pointing out that its major advantage is “total coverage in space”. Active case detection often served as the entry point to symptomatic and asymptomatic treatment and radical cure. Strategies and methods for this detection can be broadly categorised under reactive or proactive case detection.

Reactive case detection is triggered whenever a case is identified by passive case detection. In the absence of a history of travel to an endemic area, local transmission could have
occurred, and both the index case and any other locally acquired infections have to be identified, investigated, and treated to prevent additional onward transmission. Reactive case detection will involve visiting the household of the locally acquired case, screening family members, and screening neighbours within a defined radius. Screening around the index case is based on evidence from South America, Asia, and sub-Saharan Africa that malaria cases tend to cluster spatially. For example, in a low-endemic area in the Peruvian Amazon, actively seeking cases in a 100-m zone around a passively detected case yielded an increase in prevalence four to five times greater than that estimated by passive case detection. Determination of the appropriate radius within which to screen is a challenge and, because of a lack of evidence, has often been decided arbitrarily. Further scientific investigation in different eco-epidemiological settings is merited to lend support to spatial definitions of radii, sampling strategies, and numbers of people to be screened.

Data generated from passive case detection, case investigations, and reactive screenings can be used to map cases, identify risk factors for transmission, and target malaria control interventions. In Mexico, such an integrated epidemiologically driven system is used to identify residual transmission foci (figure 2) and to measure transmission risk within those foci (figure 3), allowing for targeted and context-specific malaria control interventions. Additionally, the ratio of locally acquired to imported cases from these case identification methods can be used to assess progress made in reduction of $R_c$, with a ratio of less than one locally acquired case for each imported case suggesting that the desired goal of $R_c$ less than 0.5 has been reached.

Proactive case detection involves the screening of focal populations without the trigger of a passively identified case. This approach is based on the knowledge that transmission is more likely during some periods of the year, in specific high-risk groups, or in target geographical areas. In Morocco, for example, the populations of some regions are screened routinely in what was historically the malaria transmission season. Mass blood screening has also been used in Taiwan, southern China, and Brazil. Proactive case detection is probably best suited for a circumscribed and non-mobile population, in geographically restricted settings (eg, islands), where transmission is seasonal, where mass surveys are socioculturally acceptable, and where treatment remains safe and effective for use in asymptomatic populations. Limitations to this approach include difficult logistics, testing fatigue in the population, population movements that restrict completeness of screening, and cost. Proactive case detection of the Brazilian Yanomami, which involved monthly screenings of about 10 000 people, was estimated to cost 2-3 times more than would passive case detection per positive smear.

Once elimination is achieved, the need for continued active case detection ultimately depends on the risk of onward transmission from an imported case, as defined by the $R_c$ of a specific area. For countries with low $R_c$ such as the USA and UK, passive case detection, complemented by case investigation, has been enough to avoid resurgence (table). However, most countries that have recently eliminated or are planning elimination will most likely need to complement passive case detection with some form of rigorous active case detection (table). Further research is needed to establish the most cost-effective and context-adapted combinations and the systemic approaches that are needed for implementation.

**Killing the parasite with appropriate treatment**

Depletion of the parasite reservoir in the human host requires detection of both symptomatic and asymptomatic individuals and killing all forms of the parasite that they carry. Primaquine, which is recommended by WHO in elimination settings, and the related but unregistered 8-aminoquinoline tafenoquine, are very effective against the mature gametocytes of *P. falciparum* but have a variable effect on the hypnozoites of *P. vivax* and *P.*
Both drugs can cause haemolysis in individuals who are deficient in glucose-6-phosphate dehydrogenase (G6PD), with increasing risk when high-dose regimens or long treatment schedules are used. Even low-dose use in individuals with mild G6PD deficiency can induce transient haemolytic episodes. The need for pretreatment G6PD testing with high-quality rapid tests is evident, and large-scale trials have only recently been launched.

Artemisinin-combination therapies kill developing gametocytes of *P. falciparum*, thus reducing malaria transmissibility. The additional transmission-blocking effects of including primaquine, especially in passively detected symptomatic cases that seek prompt treatment, has not been confirmed. However, recrudescent or chronic infections detected by population screening are likely to be gametocytaemic, and there is most probably a potential benefit from addition of single-dose primaquine to kill mature gametocytes unaffected by the drugs used for therapy in such cases. More research in different epidemiological settings is needed to provide evidence of the benefit of single-dose primaquine in view of the risk of haemolytic events.

Specific targeting of gametocytes is not needed to clear the blood stages of *P. vivax*, *P. ovale*, or *Plasmodium malariae*, which are susceptible to the recommended therapeutic treatment; however, primaquine is the only drug available as a hypnozoitocidal agent that can prevent the relapse infections that cause onward transmission. The standard treatment with 15 mg of primaquine daily for 14 days has not always proved effective at elimination of relapsing episodes. Alternative therapeutic regimens with increased doses or increased periods of administration are potentially more effective, although still not completely so, and safety and adherence to these regimens are a serious cause for concern.

Primaquine, combined with chloroquine, is standard treatment policy in many countries endemic for *P. vivax*. However, in most cases there is an important gap between policy and practice, probably because of health workers’ safety concerns about primaquine use. For *P. falciparum*, a few countries in the Americas use primaquine in combination with chloroquine (which is no longer recommended), but only three countries—Oman, Sri Lanka, and the Philippines—add it to their first-line artemisinin-based combination treatment. Safety concerns related to G6PD deficiency and the short half-life of primaquine make it difficult to make clear recommendations about best use of the only drug that is available for radical treatment, and alternatives are urgently needed because this remains the main obstacle to elimination of *P. vivax*.

Treatment of cases identified by passive case detection alone is unlikely to deplete the parasite reservoir or prevent onward transmission in most settings, even if radical treatment is given and there is near perfect adherence (which is difficult to achieve when primaquine is needed for 14 days or longer). Because additional active surveillance measures to detect asymptomatic carriers are expensive, alternative strategies such as mass drug administration (MDA) and mass screening and treatment have been proposed to reduce the parasite reservoir in human beings. Treatment regimens used for these schemes vary widely but almost always include primaquine for radical treatment. WHO does not encourage MDA, but, if applied in geographically defined regions or to specific target groups, might still have a role in containing or preventing outbreaks or reducing the risk of importation, especially for *P. falciparum*. To lessen the risk of resistance developing, combinations of drugs other than those used for first-line treatment, including one partner drug that has a long half-life and, ideally, acts against sexual stages, should be used. MDA has been used for *P. vivax*—but since hypnozoites in the liver can be unaffected, strategies to deal with relapses and to accompany focal infection need to be assessed further. More research is needed to define the potential role and operational feasibility of MDA and mass screening and treatment.
including identification of which drugs should be used and how frequently, with assessments of safety, coverage, effectiveness, and acceptability.

**Elimination-specific vector control activities**

Vector control strategies targeted at specific and identifiable foci during an elimination programme might need different approaches from those routinely used in control programmes. Identification of areas of high risk, with use of geostatistical analysis of incident cases and rigorous entomological surveillance, are important to continuously reassess transmission potential and possibilities of emerging insecticide resistance, and replacement of the dominant vector species. Regular assessments of vector competence for transmission based on changes in dominant vector species and abundance will enable a decision about scaling down or changing approaches to vector control in existing or newly defined foci. In Morocco, entomological investigations of the last foci of transmission showed that larval control had reduced the vectorial capacity to such low rates that resurgence of malaria was unlikely despite the presence of gametocyte carriers in the human host population. Resistance to chemical agents that target adult and larval stages of the vectors poses a major threat to elimination, and mitigation of these risks demands intensive and regular molecular and bio-assay surveillance. In South Africa, for example, pyrethroids became largely ineffective and the pre-elimination stage was reached only after DDT was reintroduced. Similarly, in central Sudan, resistance to commonly used insecticides has prevented elimination of transmission foci in this low-endemic setting.

Control programmes generally focus interventions on the most efficient vectors, which are usually indoor biting. Achievement of the complete cessation of endemic transmission might need programmes to target all vectors including those that rest or bite outdoors. At the start of the GMEP, outdoor biting and resting vectors were considered of minor importance because they were thought to be limited to few anopheline species. Later in the programme, vector behaviour, similar to the emerging resistance to DDT, was recognised to affect transmission. These outdoor biting vectors were not specifically targeted, however, and how far they contributed to the overall receptivity of transmission foci remains unclear. Although impregnated bednets have been shown to affect transmission by outdoor biting vectors, further research is needed to establish the most effective schemes for entomological surveillance and what additional vector control measures might be necessary to completely halt transmission.

One of the potential advantages of moving towards elimination is that over time some expensive and intensive vector control measures might be scaled back or stopped altogether. GMEP publications suggest that after the attack phase most vector control measures can be scaled down, but several examples from countries that have eliminated malaria clearly show that this approach is highly context dependent and largely defined by both the intrinsic transmission risk and the number of infections imported (table). Most malaria control programmes contemplating elimination will need to considerably strengthen their entomological activities and expertise not only to identify receptive foci but also to be able to make evidence-based decisions about vector control strategies and when to scale them down safely.

**Reduction of the importation of infections**

Even for countries that have long eliminated malaria, some importation is inevitable. In the USA, for example, 1298 cases were reported in 2008. However, intrinsic transmission potential is sufficiently low that only occasional cases of local transmission result from these importations. If comprehensive health-care systems and disease-reporting mechanisms exist, passive case detection can be sufficient to avoid resurgence from imported cases if the
intrinsic transmission potential is low. In most malaria-endemic countries considering elimination the vectorial capacity is such that this rate of importation might result in substantial transmission. Elimination of malaria transmission will therefore need the stream of imported infections to be slowed through: proactive case detection and treatment in migrants and travellers before they lead to transmission; and cross-border and regional initiatives that can reduce endemicity in countries from where migrants originate. Achievement of malaria elimination will probably need a vertical approach, especially for surveillance and response, whereas the prevention of reintroduction measures can be integrated in larger communicable disease programmes, as shown in Mauritius and Singapore.

Border screening

Border screening of immigrants is a specific form of proactive case detection that tries to restrict the importation of infections. Some contexts will be more conducive to monitoring and responding to importation risk than will others; islands such as Mauritius and Vanuatu, for example, have few means of entry, with boat and air travel quantifiable and easy to screen. Regions sharing large, poorly monitored, and sometimes inaccessible land borders will have great difficulty controlling importation risk, because of both the large number of potential entry routes and the less readily available data for numbers and characteristics of people using them.

Historically, importation has only been quantified once elimination has been achieved and border screening is part of the country's strategy to avoid resurgence. The potential has been shown for alternative assessment of importation risk as an integral part of assessment of the feasibility of an elimination strategy; in Zanzibar, for example, importation risk was estimated with mobile phone data as a proxy for human population movement. Robust methods combining travel history data from health facilities and border surveys in geographical information systems will provide the necessary evidence base for understanding the magnitude of importation risk and designing appropriate strategies to control it. Additionally, border screening that is focused on specific, high-risk groups with high prevalence could be a more viable strategy than generalised population screening that is likely to prove extremely inefficient because of low prevalence. Border screening might not always be necessary once elimination has been achieved, notably in areas where receptivity is very low (table).

Cross-border and regional initiatives

Border screening is not always practical or desirable, especially in areas with artificial administrative borders and frequent border crossing by a large proportion of the population on a daily basis, sometimes even facilitated for economic reasons. Namibia and Angola, for example, have an agreement allowing free movement in an area of 60 km on either side of the border. Cross-border collaborations such as the Lubombo Spatial Development Initiative are potentially more effective in reducing importation than is border screening (webappendix). The political mechanisms, motivations, financing, and responsibilities for cross-border, regional approaches to elimination are likely to be complex. However, some have argued that this approach provides the only means of reducing importation risk between countries.

Measurement of progress

One of the differences emphasised by the WHO Expert Committee between eradication and control programmes was that the administrative standard of progress for control was measurement of accomplishments, whereas for eradication it would change to measurement.
of what remains to be accomplished. Because elimination has a known quantitative goal of ending endemic transmission and reducing the number of locally acquired cases below a specific threshold, monitoring systems are essential to track progress towards that goal. Measures of the effect of control programmes typically include population-based surveys of parasite prevalence. At very low prevalence, however, the number of samples that has to be collected to find a positive result will probably be prohibitively high. WHO therefore proposes to use incidence measures collected through the routine data collection systems (passive and active). However, present and commonly used diagnostic methods for clinical management, microscopy, and rapid diagnostic tests are not ideal for surveillance because they have limited sensitivity for infections of low-parasite density, which are common in low-transmission settings.

To improve precision of measurements of transmission, new diagnostic methods and approaches will therefore become increasingly important (panel 2). PCR methods provide a more sensitive means of testing than do microscopy or rapid diagnostic tests, but better standardised and field-applicable methods with robust quality-assurance mechanisms are needed. Genotyping the detected parasites for molecular epidemiology could potentially allow for the identification of the source of imported infections. Seroprevalence testing is an old technique that has recently been improved and standardised. It has been used in population-based surveys in Djibouti, Sudan, Swaziland, and Vanuatu, but needs to be validated as a routine surveillance method. Additionally, whereas sample collection (dry blood spots on filter paper) for these diagnostic techniques is easy, most malaria programmes are not equipped to analyse samples. A strong national or regional collaboration with research institutes will be needed not only for the analysis but also for the interpretation of results, especially in the absence of standardised quality-assured techniques.

**Conclusions**

Politicians and policy makers need to understand that malaria elimination should not take a business-as-usual approach. The most notable change will involve the evolution of a surveillance system linked to rapid response, robust epidemiological data, and sustained vigilance over a long time. Most countries aiming for elimination do not yet have the surveillance systems required for an elimination effort and will need to invest substantially to improve disease notification and analysis. Furthermore, the challenges of reaching the highly efficient operational levels that are needed to stop transmission will be great. Even if such a programme is undertaken with great precision, its success within a country is likely to be contingent on a commitment throughout an entire region of countries.

Nationally, political leaders will need to create an environment within which strategies to support elimination would operate successfully. The factors that would contribute to this enabling environment—such as well functioning health systems, community participation, sufficient skilled human resources, sustainable financing, a national and regional legal framework to facilitate elimination, and political stability—have not been discussed here but are reviewed in broad terms in other papers in this Series. Nevertheless, even for the elimination of one disease, planning needs to be approached systemically to be successful. We have not addressed the complex issues related to the national certification process of malaria-free status. We do, however, recognise that this process remains imperfect and needs more investigative modelling, new methods to measure transmission (or its absence), and empirical research to improve the specificity of the present recommendation to prove the absence of local transmission “beyond reasonable doubt”.

Systematic reviews of surveillance and response methods, and case studies from countries that have recently eliminated malaria, are essential to build a stronger evidence-base and generate practical, context-specific recommendations for future guidance. Research to improve available methods for diagnosis, treatment, and vector control is also needed. Long-term research needs have been discussed elsewhere; short-term priorities should include the development of methods to assess the overall feasibility of elimination strategies, understand the epidemiology of asymptomatic infections, quantify effect sizes of imported infection risks, and compare cost-effectiveness of different surveillance and response models.

Although the GMEP has left a legacy of technical reports and guidelines, most were focused on epidemiological contexts in the Americas and Europe. The wealth of information available from the GMEP—one of WHO’s best documented programmes—should be used to inform present and future elimination efforts, but a contemporary evidence base to support cost-effective decision making is only now beginning to be generated for the more complex transmission settings of sub-Saharan Africa and Asia. “The problem of finding an effective and economical method of [eliminating] malaria in tropical Africa has not yet been solved”, stated the WHO Expert Committee in 1957. With the subsequent shift away from elimination activities, the ensuing 50 years have not provided solutions, especially for sub-Saharan Africa where large-scale interventions have only been implemented for the first time in the past decade. In the absence of clear guidance, the decision to pursue malaria elimination might be made on a political basis without careful and rigorous assessment of the technical, operational, and financial feasibility of pursuing such a course.

Historically, it had been hoped that one approach could succeed everywhere as long as it was undertaken with “military precision”. An unfortunate consequence of this model was that it essentially replaced malariologists with technicians who were skilled in the logistics of directing insecticide spray campaigns but did not have a crucial understanding of the disease and its subtleties. Such a strategy was very successful in many settings, but the challenge of eliminating malaria in high-endemic regions will need more versatile and locally tailored, systemic approaches. Some have argued that elimination of malaria from a country can only be achieved through a global eradication effort since constant importation will otherwise make local elimination a precarious achievement. In the absence of such a global campaign, elimination in individual countries can be achieved and maintained only through robust surveillance and response combined with targeted vector control to eliminate residual or re-emerging foci. Although the GMEP needed great precision to achieve universal coverage with DDT, national elimination with the challenge of continued importation will require an equal level of precision for surveillance and response. The challenges nowadays are different from those faced by the GMEP and will need the development of new approaches, novel technologies, and sustainable financing to change the distribution of malaria prevalence progressively and to eventually eradicate the disease worldwide.

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Web Extra Material

Supplementary Material 1. Supplementary webappendix.

Acknowledgments

BM, JMC, and DLS conceived the idea for the report. BM, JMC, RWS, and GAT wrote the report. BM and JMC did the literature review. BM, CD, MHR, RM, and GAT wrote the panels. LS, RRA, and MT contributed substantial technical input to the content of the report. All authors took part in the review, preparation, and final approval of the report.

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Improving passive case detection for elimination

Improving health-seeking behaviour for fever to increase the use of and reporting by private and public facilities with adequate diagnostic capacity and treatment

People take different actions when confronted with fever, varying from a wait-and-see attitude to immediate consultation with a health-care professional. The effectiveness of passive case detection is limited by the number of patients contacting the public health system, which shows the need to ensure financial access to care, both in the public and private sector, ideally with free malaria treatment as recommended by WHO. Health education, adapted to the local practices and the epidemiological context, should be an integral part of surveillance activities, not only to increase the acceptance of screening activities for active case detection but also to increase the efficiency of the passive system. Ideally, this approach would change attitudes in the community, making seeking testing for malaria the norm.

Ensuring high testing rates in fever and history of fever cases

Although Global Malaria Eradication Program documents insisted on testing all fevers, WHO guidelines recommend to test only fevers that have no other obvious cause. Clear and easy-to-follow testing algorithms are very important to ensure high testing rates, especially for low-level health workers. Additionally, in Zanzibar, regular supervision substantially improves testing rates in primary health-care facilities. Facilities participating in the newly set-up Malaria Early Detection System initially tested only around 15% of all people attending the outpatient department services, whereas around 30% of all attendees were estimated to present with fever. After intensive supervision, testing rates doubled and more malaria cases were reported. More research is needed to define evidence-based testing criteria for elimination settings that are cost effective.

Improving sensitivity of diagnostic tests

Light microscopy and quality rapid diagnostic tests are, when done well, sufficiently sensitive to detect malaria parasites for the parasitological confirmation of patients presenting with symptoms—a requirement for case management in all endemic settings. However, at low rates of endemicity, low-parasite-density infections are not only more common, but their detection is also more important because these sometimes asymptomatic carriers will continue to cause onward transmission. Both tests have limitations in detection of low-density infections, and quality assurance for microscopy is operationally challenging and labour intensive in elimination settings. Standardised protocols for quality assurance of rapid diagnostic tests, especially to verify potentially large numbers of negative results, are not available. Positive control wells or retesting negative samples with pooled DNA PCR techniques or loop-attenuated isothermal amplification are promising, but more research is needed to provide robust recommendations. Point-of-care DNA PCR would provide the desired sensitivity but this method is unlikely to be available in the near future. For the time being, DNA PCR seems most adapted to active case detection.
Panel 2

Laboratory techniques to improve surveillance and monitoring of malaria infection and transmission

In an elimination setting, detection of all infections, whether asymptomatic, low-parasite density, or imported, and measurement of transmission intensity, both to measure progress and to enable targeting foci, are key requirements to interrupt local transmission and avoid resurgence. Laboratory techniques detecting parasite DNA, specific molecular markers, and antibodies will therefore be needed to complement conventional diagnostic methods such as rapid diagnostic tests and microscopy.

DNA-PCR

Standard surveillance systems rely on diagnosis by microscopy or use of rapid detection tests that are not sufficiently specific or sensitive to detect low-parasite-density infections. PCR-based methods are simple to do, show greatly improved sensitivity, and are able to detect mixed infections. They are particularly valuable for screening large numbers of samples because analyses of dried blood spots can be done, and a single low-grade parasitaemic sample can be detected in a pool of 50 samples, providing a resource-saving and more cost-effective procedure.

Genotyping

Elimination strategies need to know where infections originate; specifically, if they are autochthonous or imported and, if imported, from where. Malaria parasites are genetically highly variable, and genotyping has been used as a means to distinguish between indigenous and imported cases. Genetic variants can be measured from single drops of dried blood, and the assays are relatively cheap, rapid, and accurate. Molecular epidemiology can establish not only where infections originate, but also construct transmission networks in space and time to show relations between parasites in areas selected for elimination. Most studies so far have used molecular markers that are under selection (drug or immune) pressure, but there are alternative multilocus markers in Plasmodium falciparum and Plasmodium vivax that are not under selection but still show the variability that is needed.

Serology

Measurement of antimalarial antibodies in exposed populations is a valuable method because it integrates malaria exposure over time. The advent of recombinant antigen technology makes serology a robust and standardisable method. Antibodies can be detected in blood from a finger prick, and samples can be assayed quickly in large numbers. Detection of antibody responses is highly sensitive and specific, allowing an assessment of exposure to both P falciparum and P vivax (and potentially other forms of malaria) in the same sample.

For monitoring and assessment, seroprevalence rates can be used to define malaria endemicity and distinguish between areas of differential exposure when parasite rates are zero. In these areas, residual or potential foci of infection can be identified by geospatial analysis of individual or household level antibody response. Detailed examination of age-specific seroprevalence profiles can be used to monitor changes in transmission or to identify risk of exposure associated with behaviour such as travel or residence. Absence of antimalarial antibodies was used to show the success of elimination programmes in Mauritius, Greece, and on the island of Aneityum in Vanuatu.

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Figure 1.
Effect of passive case detection on transmission is limited by a cascade of factors
The percentage of infections identified through passive case detection depends on the proportion of new infections that produce clinical symptoms, the proportion of clinical cases that seek treatment in a reporting facility, the proportion of treatment-seeking cases that are tested for malaria, and the sensitivity and quality (performance) of diagnostic tests. Furthermore, the effect of passive case detection on transmission will depend on the proportion of infections identified by diagnostics that are prescribed and receive appropriate treatment, the proportion of those receiving treatment that adhere to it, and the efficacy of the drug.
Figure 2.
Residual malaria transmission foci in the states of Oaxaca and Chiapas, Mexico
Localities are depicted as dots; blue dots indicate localities where transmission has been interrupted.
Figure 3.
Residual transmission focus in southern Oaxaca state, Mexico
The annual parasite index (API) is depicted by municipality. Localities are shown as dots with risk of transmission from low to high.
## Table

Malaria control measures in WHO-certified malaria-free countries and in endemic and non-certified malaria-free countries

<table>
<thead>
<tr>
<th>WHO-certified malaria-free countries</th>
<th>Endemic and non-certified malaria-free countries</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>USA</strong></td>
<td><strong>Oman</strong></td>
</tr>
<tr>
<td><strong>UK</strong></td>
<td><strong>Sierra Leone</strong></td>
</tr>
<tr>
<td><strong>Singapore</strong></td>
<td><strong>South Africa</strong></td>
</tr>
<tr>
<td><strong>Mauritius</strong></td>
<td><strong>Mexico</strong></td>
</tr>
</tbody>
</table>

### Dominant vector(s)

- USA: Anopheles quadrimaculatus, Anopheles freeborni, Anopheles albimanus
- UK: NA
- Singapore: Anopheles suslicus, Anopheles maculata
- Mauritius: Anopheles gambiae sensu lato
- Oman: Anopheles culicifacies, Anopheles fluviatlis
- Sri Lanka: Anopheles culicifacies, Anopheles subpictus
- South Africa: Anopheles arabiensis
- Mexico: Anopheles pseudopunctipennis, A albimanus

### Total number of cases (year)

- USA: 1298 (2008), 1495 (2009)
- UK: 0 (2008), 1 (2008)
- Oman: 898 (2009)
- Mexico: 2703 (2009)

### Local cases (year)

- USA: 0 (2008), 0 (2009)
- UK: 1 (2008), 0 (2008)
- Mauritius: 0 (2008)
- Oman: 0 (2009)
- Mexico: 2698 (2009)

### Imported cases (year)

- USA: 1298 (2008), 1495 (2009)
- UK: 0 (2008), 0 (2008)
- Mauritius: 0 (2008)
- Oman: 898 (2009)
- Mexico: 2703 (2009)

### Plasmodium falciparum

- USA: 40.6%, 78.9%
- UK: 76.2%
- Singapore: 30%
- Mauritius: 18%
- Oman: 18%
- Sri Lanka: 4%
- South Africa: 95%
- Mexico: 1%

### Plasmodium vivax

- USA: 14.6%
- UK: 13.7%
- Singapore: 21.2%
- Mauritius: 70%
- Oman: 80%
- Sri Lanka: 94%
- South Africa: 99%
- Mexico: 99%

### Other (including mixed infections)

- USA: 3.6%
- UK: 7.4%
- Singapore: 2.6%
- Mauritius: 2%
- Oman: 2%
- Sri Lanka: 2%
- South Africa: 2%
- Mexico: 2%

### Species unreported

- USA: 41.2%
- UK: 41.2%
- Singapore: 41.2%
- Mauritius: 41.2%
- Oman: 41.2%
- Sri Lanka: 41.2%
- South Africa: 41.2%
- Mexico: 41.2%

### Treatment

- **P. falciparum**
  - USA: AV+PG, AL, QN+D/T/CL
  - UK: AV+PG, AL, QN+D/T/CL
  - Singapore: AN+D/CL
  - Mauritius: AL, QN, AL+PQ (if gametocytes identified)
  - Oman: AL+PQ
  - Sri Lanka: AL
  - South Africa: AL
  - Mexico: CQ+PQ

- **P. vivax**
  - USA: CQ+PQ
  - UK: CQ+PQ
  - Singapore: CQ+PQ
  - Mauritius: CQ+PQ
  - Oman: CQ+PQ
  - Sri Lanka: CQ+PQ
  - South Africa: CQ+PQ
  - Mexico: CQ+PQ

### G6PD screening

- USA: Yes
- UK: Yes
- Singapore: Yes
- Mauritius: No
- Oman: Yes
- Sri Lanka: Yes
- South Africa: No
- Mexico: No

### PCD

- USA: No
- UK: No
- Singapore: No
- Mauritius: Yes (when local transmission is suspected)
- Oman: Yes (outbreaks and migrant populations)
- Sri Lanka: Yes (high-risk groups)
- South Africa: Yes (high-risk areas)
- Mexico: Yes (starting 2010)

### ACD

- USA: Only if local transmission is suspected
- UK: All cases
- Singapore: All cases
- Mauritius: All cases
- Oman: All cases
- Sri Lanka: All cases
- South Africa: All cases
- Mexico: All cases

### Case investigation

- USA: All cases
- UK: All cases
- Singapore: All cases
- Mauritius: All cases
- Oman: All cases
- Sri Lanka: All cases
- South Africa: All cases
- Mexico: All cases

### Population screening

- USA: Yes (outbreaks and migrant populations)
- UK: Yes (high-risk groups)
- Singapore: Yes (high-risk areas)
- Mauritius: Yes (high-risk groups)
- Oman: Yes (outbreaks)
- Sri Lanka: No
- South Africa: No
- Mexico: No

### Border screening

- USA: Negative test needed for people applying for a work permit (rescreening for renewal)
- UK: No
- Singapore: No
- Mauritius: No
- Oman: No
- Sri Lanka: No
- South Africa: No
- Mexico: No

### Vector control

- USA: IRS
- UK: No
- Singapore: No
- Mauritius: No
- Oman: Yes
- Sri Lanka: No
- South Africa: No
- Mexico: No

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<tr>
<td>Targeted</td>
<td>Some counties augment blanket coverage with targeted interventions based on entomological surveillance</td>
</tr>
<tr>
<td>Outbreak response</td>
<td>Transfusion transmitted: test the involved units/donors, treat the donor, embargo remaining units; mosquito-transmitted malaria: active case detection, vector control, community education</td>
</tr>
</tbody>
</table>

NA=not applicable. AV=atovaquone. PG=proguanil. AL=artemether-lumefantrine. QN=quinine. D=doxycycline. T=tetracycline. CL=clindamycin. CQ=chloroquine. G6PD=glucose-6-phosphate dehydrogenase. PCD=passive case detection. RDT=rapid diagnostic test. ACD=active case detection. IRS=indoor residual spraying. MDA=mass drug administration.

*1031 cases are of unknown origin.