



Article

<https://doi.org/10.1038/s41591-023-02551-w>

Modeling policy interventions for slowing the spread of artemisinin-resistant *pfkelch* R561H mutations in Rwanda

Received: 20 December 2022

Accepted: 18 August 2023

Published online: 21 September 2023

Robert J. Zupko ¹, Tran Dang Nguyen¹, J. Claude S. Ngabonziza^{2,3}, Michee Kabera⁴, Haojun Li^{1,5}, Thu Nguyen-Anh Tran ¹, Kien Trung Tran¹, Aline Uwimana^{4,6} & Maciej F. Boni ^{1,7}

Check for updates

Artemisinin combination therapies (ACTs) are highly effective at treating uncomplicated *Plasmodium falciparum* malaria, but the emergence of the new *pfkelch13* R561H mutation in Rwanda, associated with delayed parasite clearance, suggests that interventions are needed to slow its spread.

Using a Rwanda-specific spatial calibration of an individual-based malaria model, we evaluate 26 strategies aimed at minimizing treatment failures and delaying the spread of R561H after 3, 5 and 10 years. Lengthening ACT courses and deploying multiple first-line therapies (MFTs) reduced treatment failures after 5 years when compared to the current approach of a 3-d course of artemether–lumefantrine. The best among these options (an MFT policy) resulted in median treatment failure counts that were 49% lower and a median R561H allele frequency that was 0.15 lower than under baseline. New approaches to resistance management, such as triple ACTs or sequential courses of two different ACTs, were projected to have a larger impact than longer ACT courses or MFT; these were associated with median treatment failure counts in 5 years that were 81–92% lower than the current approach. A policy response to currently circulating artemisinin-resistant genotypes in Africa is urgently needed to prevent a population-wide rise in treatment failures.

The introduction of artemisinin combination therapies (ACTs) has been instrumental in reducing the burden of *Plasmodium falciparum* malaria, but the continued evolution of drug resistance by malaria parasites has the potential to undermine these advances. Since the first appearance of artemisinin resistance in Cambodia in the 2000s^{1,2}, the spread of molecular markers associated with artemisinin resistance has largely been concentrated in Southeast Asia^{3,4}. However, the de novo appearance of confirmed markers for artemisinin resistance in Rwanda^{5–10}

and Uganda^{11,12} signals the need for interventions to be considered in the African context¹³ to minimize the expected health, mortality and economic costs should artemisinin resistance become widespread¹⁴.

The ACT artemether–lumefantrine (AL) was adopted by Rwanda as the first-line therapy for uncomplicated falciparum malaria in 2006 as part of a comprehensive national strategic plan for malaria control¹⁵. Since adoption of AL, the *P. falciparum* kelch protein 13 (*pfkelch13*, PF3D7_1343700) R561H mutation has emerged and been validated as a

¹Center for Infectious Disease Dynamics, Department of Biology, Pennsylvania State University, University Park, PA, USA. ²Research, Innovation and Data Science Division, Rwanda Biomedical Center (RBC), Kigali, Rwanda. ³Department of Clinical Biology, University of Rwanda, Kigali, Rwanda. ⁴Malaria and Other Parasitic Diseases Division, Rwanda Biomedical Centre (RBC), Kigali, Rwanda. ⁵Department of Computer Science, Columbia University, New York City, NY, USA. ⁶Louvain Drug Research Institute, Université Catholique de Louvain, Ottignies-Louvain-la-Neuve, Belgium. ⁷Centre for Tropical Medicine and Global Health, Nuffield Department of Medicine, University of Oxford, Oxford, UK. e-mail: rbz5100@psu.edu

marker for partial artemisinin resistance in samples collected as part of clinical drug efficacy studies between 2012 and 2015 (ref. 3). In contrast to the wild-type clearance rate of 2.7 h, the 561H mutant is associated with a delayed clearance half-life of 7.2 h¹⁶. This is similar to the clearance half-life of the 580Y mutation that emerged in Cambodia^{6,8,16}. Following the original identification of 561H in the Gasabo district of Rwanda⁵, additional studies have found 561H in more districts, with recent allele frequency measurements ranging from 0.045 to 0.219 (refs. 6,7,9,10). These findings indicate that drug-policy interventions are now needed to delay the spread of 561H within the local *P. falciparum* population and to reduce the impact of treatment failures due to artemisinin-resistant parasites.

As of March 2023, the World Health Organization (WHO) recommends the following six ACTs for the treatment of uncomplicated *P. falciparum* malaria: AL, artesunate–amodiaquine (ASAQ), artesunate–mefloquine (ASMQ), dihydroartemisinin–piperaquine (DHA–PPQ), artesunate–sulfadoxine–pyrimethamine (AS + SP) and artesunate–pyronaridine (AS–Pyr)¹⁷. Within sub-Saharan Africa, the predominant therapies deployed are AL and ASAQ¹⁸, while the non-ACT formulation of sulfadoxine–pyrimethamine + amodiaquine (SPAQ) is commonly used for seasonal malaria chemoprophylaxis¹⁹. As such, sub-Saharan Africa faces a constrained drug landscape that requires national drug-policy interventions to be balanced between delaying drug-resistance evolution by the parasite—which leads to increased drug failures over the long term—and ensuring that therapies currently given are highly efficacious.

Studies detecting the mutant 561H allele indicate both increasing allele frequency and geographic spread from 2014 to 2019; however, there may still be a window of opportunity to delay or prevent the fixation of 561H in Rwanda and avert high numbers of treatment failures. Accordingly, using a mathematical model of malaria transmission, we examined 26 possible drug-policy interventions (the majority of which use existing therapies) and their ability to slow down 561H evolution and reduce long-term treatment failures. These include the replacement of the existing first-line therapy, introduction of multiple first-line therapies (MFTs) and lengthening the dosing schedule for AL from a 3-d course of treatment to up to 5 d of AL treatment in accordance with previous clinical trials^{20–22}. Additionally, a more logically complicated strategy of drug rotation is evaluated along with sequential therapy approaches (for example, 3-d AL course followed by a 3-d DHA–PPQ course) and the deployment of triple ACTs. Table 1 summarizes our findings and policy implications.

Results

Status quo

To provide a common point of comparison for policy interventions, a baseline (or status quo) scenario was run ($n = 100$ replicates) in which no interventions were implemented. Our spatially calibrated model initially uses the 2017 Malaria Atlas Project (MAP) projections for Rwanda's malaria prevalence (that is, the *P. falciparum* prevalence in ages 2–10 years ($PfPR_{2-10}$)) through the end of 2020, after which prevalence is scaled down in agreement with current incidence estimates for Rwanda²³. Following this transmission reduction to the 2021 malaria incidence, the model presumes that incidence will remain stable over time. The allele frequency of 561H is calibrated (Fig. 1) to its measured distribution and frequency in Rwanda from 2014 to 2019 (Extended Data Fig. 1 and Supplementary Table 1). Using this calibration, the simulation forecasts that at the end of 2023, the national 561H allele frequency will be 0.36 (interquartile range (IQR): 0.29–0.45) and will reach a frequency of 0.98 (IQR: 0.97–0.99) by 2033. This projects that 561H will be effectively fixed as the dominant allele if current status quo conditions continue, a pattern typical for most but not all past patterns of antimalarial drug-resistance evolution.

The increase in 561H allele frequency is projected to be associated with an increase in treatment failures. Based on previously calibrated

Table 1 | Policy summary

Background	The identification of artemisinin-resistant <i>P. falciparum</i> parasites in Rwanda carrying the <i>pfc kelch13 R561H</i> allele requires a change to antimalarial strategy that explicitly aims to (1) slow down the spread of artemisinin-resistant genotypes and (2) minimize drug-resistance-associated treatment failures over the next decade.
Main findings and limitations	Using a stochastic individual-based <i>P. falciparum</i> transmission model calibrated to Rwanda's malaria epidemiology and known distribution of 561H mutations between 2014 and 2019, we project that by January 2024 the 561H allele frequency in Rwanda will be between 0.29 and 0.45 (IQR). The population-level treatment failure rate is projected to surpass 10% sometime between 2024 and 2026. Lengthening the course of the currently recommended AL treatment from 3 to 5 d, or switching away from AL altogether, is projected over a 5-year span to slow down the spread of 561H and to reduce the treatment failure rate. The model projects that the most effective single policy switch is to an MFT approach with ASAQ used for 75% of treatments and DHA–PPQ used for 25% of treatments. Under this MFT strategy, the 5-year projected treatment failure rate is between 9% and 11%, whereas the status quo of using 3-d AL is projected to result in 16–20% treatment failure after 5 years.
Policy implications	The major uncertainty in these evaluations is that the future evolutionary path of piperaquine resistance in Africa is unknowable. If a piperaquine-resistant phenotype emerges in Africa that has similar characteristics to the Southeast Asian phenotype, DHA–PPQ use will likely need to be restricted. If low-grade piperaquine resistance is observed in Africa, DHA–PPQ usage can be expanded if paired with real-time molecular surveillance.
	Certain next-generation approaches—such as the usage of triple artemisinin combination therapies or 6-d sequential courses of approved ACTs—have the potential to keep treatment failures low for 5 years and in some cases 10 years. Both triple therapies evaluated here and all 6-d mixed regimens of two different ACTs were projected to keep the median treatment failure percentage below 4% after 5 years.

AL efficacy on commonly circulating genotypes in East and Central Africa^{24,25}, our model estimates that in 2024, presuming no interventions, 11.15% (IQR: 9.9–12.11%) of treated *P. falciparum* malaria cases will fail treatment with the present first-line therapy of AL, or about 154,000 individuals for the calendar year (median monthly average of 12,800 (IQR: 11,200–14,000)). This monthly average is forecast to increase to 39,400 (IQR: 36,200–42,400) per month by 2033 (Supplementary Table 2). Under the current status quo conditions, the 10% treatment failure threshold recommended by the WHO for a change in first-line therapy is likely to be met or exceeded between 2024 and 2026 (12.7% median estimate for 2026, IQR: 11.2–12.9%; Supplementary Table 2).

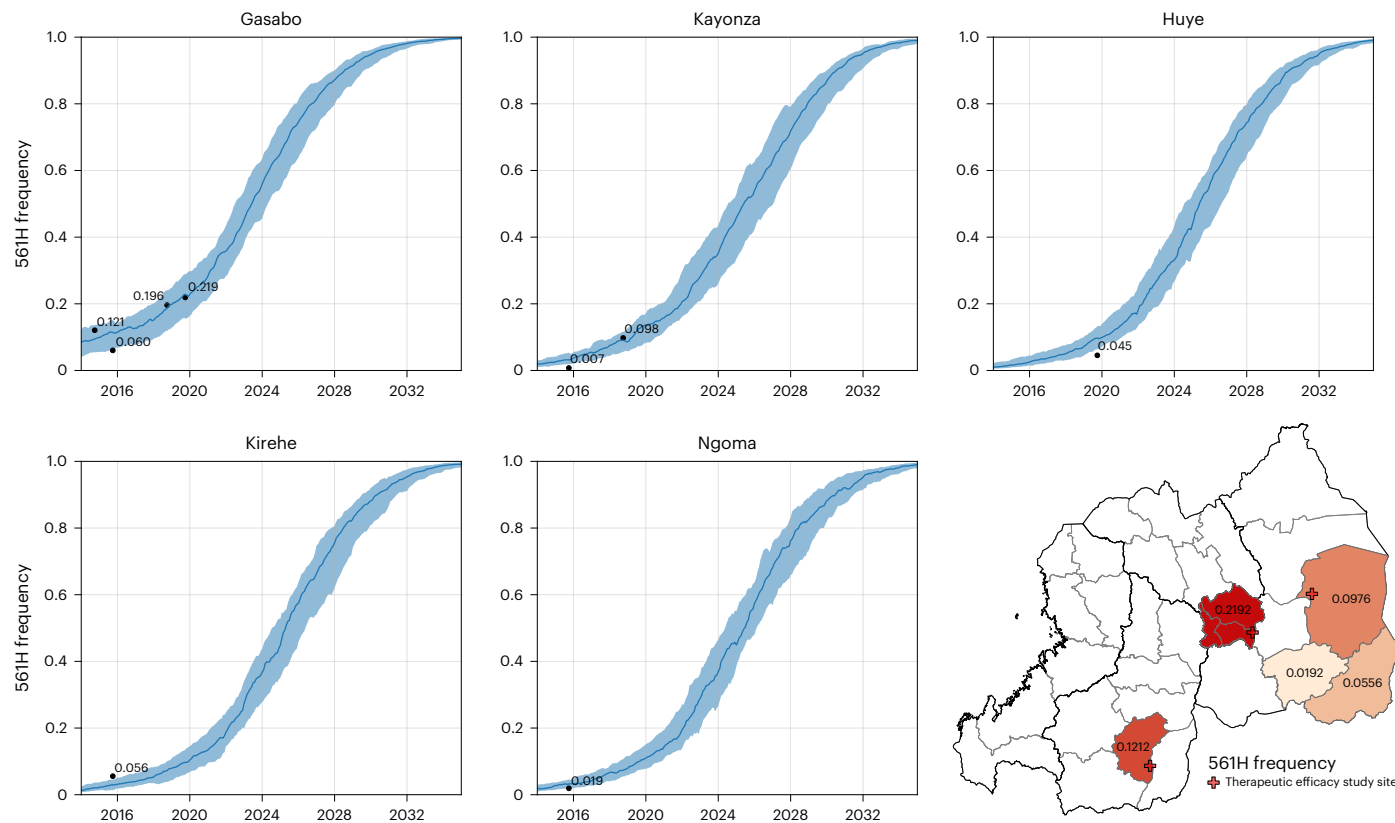


Fig. 1 | Calibration of simulated 561H allele frequency versus known frequency. The 561H mutations are artificially introduced into the simulation (10 years before detection) in the Gasabo district and allowed to evolve and increase in frequency, as shown here through model completion in 2035. During model execution, the 561H alleles spread across the simulated landscape via human migration and are selected via local drug pressure. The simulated allele

frequencies in five districts (median and IQR shown with blue line and shaded area, $n=100$) are compared to known allele-frequency data^{5–7,9} (black dots). Gasabo is a district of Kigali City; Huye district is in southern Rwanda; Kirehe, Kayonza and Ngoma districts are in eastern Rwanda. The calibrated model is largely in agreement with known 561H spatial evolutionary patterns.

To evaluate the impact of potential drug-policy interventions, we examined 26 national-scale drug-resistance-response strategies (Table 2), with a presumed implementation date of 1 January 2024, and calculated the relevant metrics using 3-, 5-, and 10-year endpoints, with the objectives of (1) reducing the near- and long-term numbers of treatment failures and (2) minimizing the increase in 561H allele frequency. The interventions can broadly be placed into the following five categories: (1) a change in first-line recommendation to a readily available and deployable therapy, (2) a change in strategy to deployment of MFT, (3) a change to a more intensive management approach, where multiple approaches are used sequentially with different goals at different times (for example, lowering prevalence and delaying resistance), (4) sequential dosing of two ACTs and (5) a switch to high-efficacy triple ACTs, assuming triple ACTs are approved and immediately available for emergency use.

Currently approved interventions

Among alternate first-line therapies, extending the course of AL from 3 to 4 or 5 d is the most immediately available option due to stocks of AL already being present and available. Continued use of a 3-d AL course for 5 years is projected to lead to a median 561H allele frequency of 0.82 (IQR: 0.73–0.86), whereas 4-d AL results in a 561H allele frequency of 0.76 (IQR: 0.67–0.85, $P=0.0044$; Wilcoxon rank-sum test) and 5-d AL results in a 561H allele frequency of 0.69 (IQR: 0.57–0.78, $P<10^{-4}$; Fig. 2). After 5 years, the national average monthly treatment failures are expected to be 23,500 (IQR: 20,900–26,500). These figures drop to 17,100 (IQR: 15,200–19,300, $P<10^{-4}$) under 4-d AL and 13,100

(IQR: 10,500–14,700, $P<10^{-4}$) under 5-d AL. Treatment failure numbers are improved under a longer course of AL because of the combined effect of lower 561H frequency and higher treatment efficacy of the longer course. AL efficacy generally remained high in the model's parameterization with the majority of AL efficacies, across all genotypes in the simulation, above 85%.

Replacing AL with an alternative first-line therapy such as ASAQ or DHA-PPQ is the next most practical intervention to implement. A switch to ASAQ gives results similar to a 4-d or 5-d course of AL with a 5-year 561H allele frequency of 0.70 (IQR: 0.58–0.77, $P<10^{-4}$ compared to 3-d AL) and a median of 14,100 (IQR: 12,500–14,700, $P<10^{-4}$) treatment failures per month. However, a switch to DHA-PPQ results in an acceleration of the fixation of 561H with the allele frequency reaching 0.90 (IQR: 0.87–0.93, $P<10^{-4}$) within 5 years and fixation within 10 years (Fig. 2). The projected number of treatment failures is also high with a monthly average of 60,800 (IQR: 57,100–64,000, $P<10^{-4}$) after 5 years and 83,300 (IQR: 82,900–83,400, $P<10^{-4}$) after 10 years. When conducting sensitivity analysis for scenarios where DHA-PPQ efficacy remains relatively high, switching to DHA-PPQ is still projected to reach the 10% treatment failure threshold within 5 years of deployment, although the long-term treatment failure rates are lower (Supplementary Table 3). The rapid fixation of the 561H allele when switching to DHA-PPQ as the first-line therapy is due to the presence of artemisinin resistance coupled with the model's projected rapid evolution of PPQ resistance leading to partner-drug failure, resulting in an environment favorable for rapid selection for artemisinin resistance.

Table 2 | Summary of the primary drug therapy interventions examined using the simulation

Intervention	Therapy
AL extension	AL (4-d course)
	AL (5-d course)
	AL (3-d course on days 0, 1 and 2) followed up with a second course on days 7, 8 and 9; labeled 'AL789'
AL replacement	ASAQ
	DHA-PPQ
	ASAQ (75%)+DHA-PPQ (25%)
	ASAQ (50%)+DHA-PPQ (50%)
	ASAQ (25%)+DHA-PPQ (75%)
	AL (75%)+ASAQ (25%)
	AL (50%)+ASAQ (50%)
	AL (25%)+ASAQ (75%)
	AL (75%)+DHA-PPQ (25%)
MFTs	AL (50%)+DHA-PPQ (50%)
	AL (25%)+DHA-PPQ (75%)
	AL on days 0, 1 and 2, followed by ASAQ on days 3, 4 and 5; labeled 'AL, then ASAQ345'
	AL on days 0, 1 and 2, followed by DHA-PPQ on days 3, 4 and 5
	ASAQ on days 0, 1 and 2, followed by AL on days 3, 4 and 5
	DHA-PPQ on days 0, 1 and 2, followed by AL on days 3, 4 and 5
	AL on days 0, 1 and 2, followed by ASAQ on days 7, 8 and 9; labeled 'AL, then ASAQ789'
	AL on days 0, 1 and 2, followed by DHA-PPQ on days 7, 8 and 9
	ASAQ on days 0, 1 and 2, followed by AL on days 7, 8 and 9
Sequential courses of 3-d ACT	DHA-PPQ on days 0, 1 and 2, followed by AL on days 7, 8 and 9
	DHA-PPQ (3 years), then AL (50%)+ASAQ (50%)
	DHA-PPQ (3 years), then 5-d course of AL (50%)+ASAQ (50%)
Switch to DHA-PPQ, followed by switch to MFT	ALAQ
	ASMQ-PPQ

In contrast to extending the duration of AL treatment or replacing the first-line therapy, the introduction of MFT will require additional logistical and operational effort, but this is likely to be offset by the effectiveness of MFT in slowing the spread of drug-resistant genotypes (due to the more complex evolutionary environment that parasites face under MFT^{26–28}). Nine combinations of AL, ASAQ and DHA-PPQ with distribution ratios of 25/75, 50/50 and 75/25 were considered (Table 2) with drug choice at the time of treatment based upon a random draw in the simulation. Except for MFTs with a high (that is, 75%) proportion of DHA-PPQ treatments, MFT strategies outperformed the status quo with regard to 561H allele frequency (Fig. 3) and treatment failures (Fig. 4 and Supplementary Table 2). However, only an MFT consisting of 75% ASAQ and 25% DHA-PPQ is projected to be under the 10% treatment failure threshold after 5 years at 9.9% (IQR: 8.7–10.6%, $P < 10^{-4}$ compared to 3-d AL). Within 10 years, all MFT approaches are projected to exceed 10% treatment failure, although combinations of 50% AL + 50% ASAQ and 25% AL + 75% ASAQ have comparably acceptable outcomes with 15.5% (IQR: 14.8–16.0%, $P < 10^{-4}$) and 14.5% (IQR: 14.0–14.8%, $P < 10^{-4}$) treatment failure rates, respectively.

The high percentage of treatment failures in MFTs incorporating DHA-PPQ at high levels (Fig. 4) is once again due to the loss of DHA-PPQ efficacy as PPQ-resistance evolution accelerates. After 5 years, the optimal MFT policy (75% ASAQ and 25% DHA-PPQ) is projected to generate 11,900 (IQR: 10,300–12,800, $P < 10^{-4}$) monthly treatment failures (Extended Data Fig. 2).

While DHA-PPQ deployment, compared to all other strategies, is associated with a higher frequency of 561H over the 5- and 10-year time frame—and an associated higher level of treatment failure—over a 3-year time frame DHA-PPQ is projected to be more efficacious, suggesting that a more complex rotation strategy is worth considering. To evaluate this, we introduced the use of DHA-PPQ as the first-line therapy for 3 years, followed by a switch to an MFT using either 50% AL and 50% ASAQ or 50% 5-d AL and 50% ASAQ. These two strategies were comparable to or marginally better than pure MFT approaches. Only the strategy involving a switch to an MFT with 50% receiving a 5-d course of AL and 50% receiving ASAQ remained under the 10% threshold at the end of 5 years (9.5% (IQR: 8.8–10.1%, $P < 10^{-4}$ compared to 3-d AL); Supplementary Table 2) with projected monthly treatment failures of 11,800 (IQR: 11,000–12,600, $P < 10^{-4}$). The procurement and distribution required to deploy a drug rotation coupled with MFT suggest that compliance and operations would have a considerable influence on the success of this approach. In general, MFT strategies promote the emergence of a large number of distinct genotypes, but selection pressure is weak on each genotype, limiting their ability to reach high allele frequency (Extended Data Fig. 3).

Next-generation interventions

We evaluated an intervention using two ACT courses sequentially for a single case of uncomplicated *P.falciparum* malaria. This approach has the advantage of an increased total dose of artemisinin (6 d), minimization of safety risks by changing partner drugs and potential exploitation of partner drugs selecting for opposite alleles. We explored this approach in the following two ways: following the protocol discussed in ref. 29 with six consecutive days of treatment from day 0 to day 5 inclusive, and a modified protocol in which the second course is taken on days 7, 8 and 9 (labeled as '789' in Extended Data Fig. 4). This modified protocol was chosen as one that may sustain better adherence in settings where village health workers assist patients in completing their malaria treatment courses, as it allows the second course to be taken on the same days of the week as the first course. At 5 years after introduction, all sequential courses have a median 561H frequency between 0.50 and 0.63, with ASAQ followed by AL having the lowest of 0.50 (IQR: 0.38–0.59, $P < 10^{-4}$ compared to 3-d AL). These are lower than the median 561H frequencies projected for all therapy switches, MFT approaches and rotations considered thus far. ASAQ followed by AL is also projected to have the lowest treatment failure rate 5 years after introduction at 1.8% (IQR: 1.5–2.0%, $P < 10^{-4}$). All sequential courses are projected to have median treatment failure rates below 4% after 5 years and below 12% after 10 years. As in other scenarios, 10 years of DHA-PPQ use as part of a strategy of sequential ACT courses still results in strong selection pressure for PPQ resistance with long-term treatment failures increasing correspondingly (Supplementary Table 2).

While the previous 24 national-scale response strategies make use of currently available therapies, the results of past and ongoing clinical trials of the triple ACTs artemether–lumefantrine–amodiaquine (ALAQ) and artesunate–mefloquine–piperaquine (ASMQ-PPQ)³⁰ suggest that they are likely to be highly efficacious and should be considered as an emergency intervention in response to rising treatment failure rates. For these two scenarios, we presume that ALAQ or ASMQ-PPQ are deployed as first-line therapy, replacing AL. As expected based upon previous modeling studies^{31,32}, in our analysis, triple ACTs outperformed most other drug-policy interventions with ALAQ resulting in average monthly treatment failures of 2,100 (IQR: 1,900–2,500, $P < 10^{-4}$ compared to 3-d AL) and a 561H allele

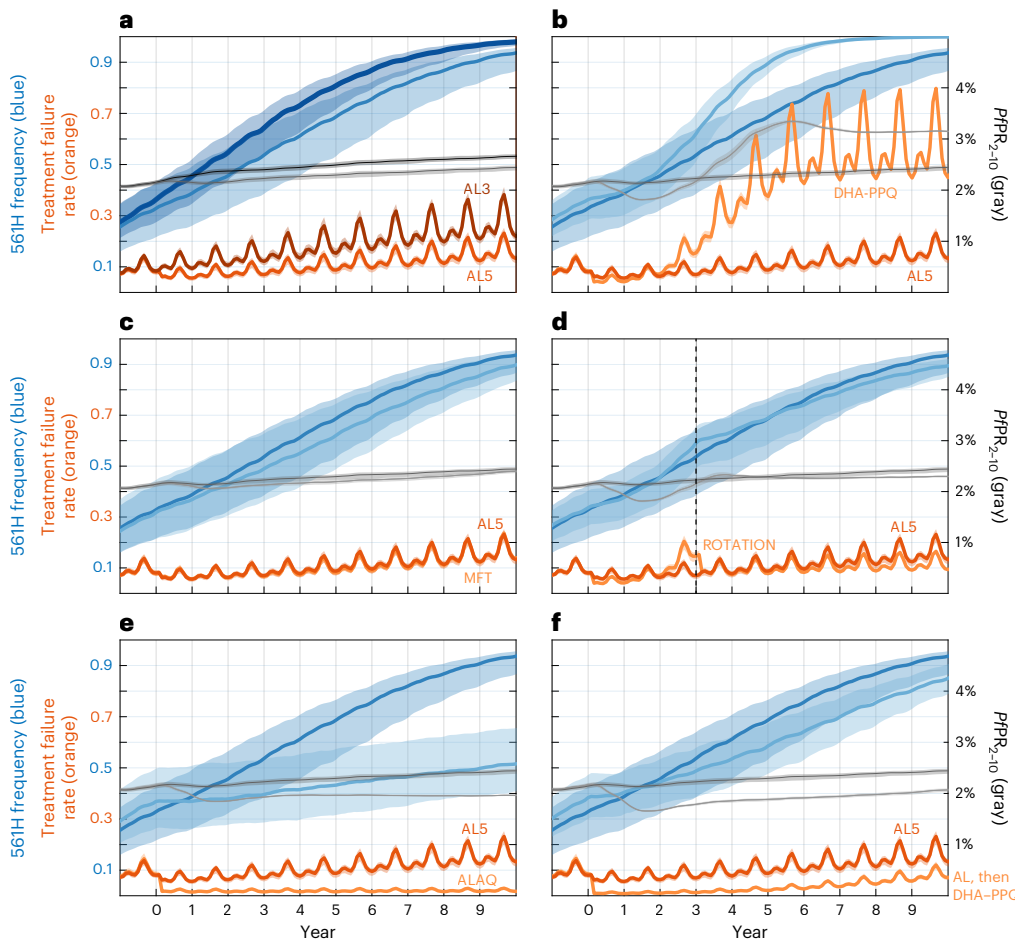


Fig. 2 | Projected 561H allele frequencies and treatment failure percentages under potential intervention scenarios. The right-hand axis shows 12-month-smoothed malaria prevalence ($PfPR_{2-10}$). All strategies are evaluated over 10 years with year 0 corresponding to the year the strategy was first implemented. Shaded bands show IQR. **a**, Comparison between the status quo treatment of 3 d of AL therapy (AL3, darker lines) and 5 d of AL therapy (AL5). **b**, Comparison between AL5 and switching to DHA-PPQ (lighter lines) as first-line therapy. Note here that under DHA-PPQ prevalence drops early (to <2%) and later rises to nearly 3.5%. This occurs because piperaquine resistance evolves quickly and reaches

>0.50 genotype frequencies after 3 years, resulting in high prevalence and high levels of treatment failure. **c**, Comparison between AL5 (darker colors) and an MFT policy using ASAQ (75% of treatments) and DHA-PPQ (25%).

d, Comparison between AL5 (darker colors) and a rotation strategy where DHA-PPQ is used for 3 years and then replaced with an MFT policy using AL5 (50%) and ASAQ (50%). **e**, Comparison between AL5 (darker colors) and the triple therapy ALAQ. **f**, Comparison between AL5 (darker colors) and sequential courses of AL on days 0, 1 and 2 and DHA-PPQ on days 7, 8 and 9.

frequency of 0.43 (IQR: 0.34–0.57, $P < 10^{-4}$) after 5 years, while ASMQ-PPQ resulted in 1,900 (IQR: 1,200–2,600, $P < 10^{-4}$) monthly treatment failures and a 561H allele frequency of 0.47 (IQR: 0.35–0.58, $P < 10^{-4}$). Median treatment failure rates are projected to be 1.9% and 1.8%, respectively, after 5 years. The usage of ASMQ-PPQ comes with an increased risk of PPQ failure leading to projected treatment failures of 18.8% (IQR: 15.6–21.0%, $P < 10^{-4}$) at 10 years after deployment. In contrast, treatment failures are likely to still be low if ALAQ is deployed reaching only 2.1% (IQR: 1.8–2.3%, $P < 10^{-4}$) 10 years after deployment (Fig. 2).

Patient adherence to treatment regimens

The scenarios modeled here assume that all courses of treatment will be completed in full; in practice, 100% compliance is unlikely and adherence rates may be complicated by under-dosing, over-dosing and formulation design (that is, fixed-dose combination versus copackaged blister packs), resulting in real-world compliance rates between 65% and 90% for a 3-d course of treatment^{33,34}. To evaluate the possible impacts that compliance with treatment regimens would have, additional scenarios were evaluated in which ASAQ, DHA-PPQ and 3-, 4- or 5-d courses of AL were administered using low (25–70%), moderate

(50–80%) and high (70–90%) compliance rates for complete courses (Supplementary Table 4). As expected, failure to comply with the prescribed course of treatment results in an increase in treatment failures; however, extended courses of AL (4 or 5 d) and ASAQ, even under low compliance, still outperform perfect compliance with a 3-d course of AL (Supplementary Table 5). No differences were seen in policy evaluation or prioritization when evaluating scenarios with imperfect compliance.

Discussion

The projected national frequency of the 561H allele in Rwanda under a continuation of status quo treatment with a 3-d course of AL suggests that treatment failures will increase over the next 5 years and that drug-policy interventions are required to mitigate this risk as much as possible. The current spread of the 561H allele to other districts^{6–10}, from its initial identification⁵, together with confirmation of 561H at frequencies similar to those projected in our simulation, suggests that the 561H allele is likely present throughout Rwanda. As recommended by the WHO, the Rwandan National Malaria Control Program (NMCP) should consider several strategies for mitigating the spread of *pfkkelch13* alleles associated with artemisinin resistance¹³.

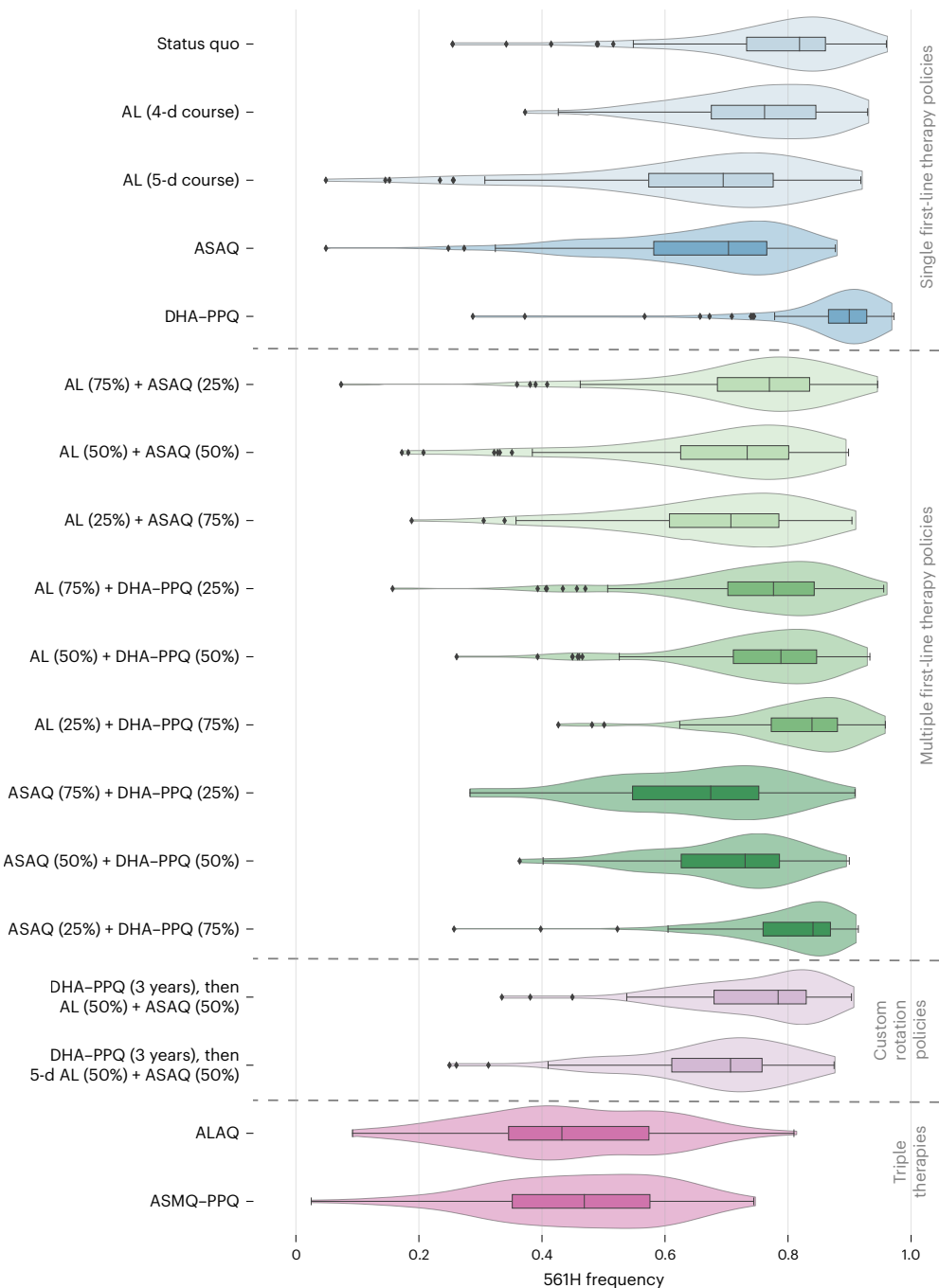


Fig. 3 | Projected 561H allele frequency after 5 years. Box plots ($n = 100$ model replicates per policy, median line with IQR) and violin plots (full data range) show the projected 561H frequency, after 5 years, under 17 policy options and the status quo of continued usage of 3-d AL. Box-plot whiskers show $1.5 \times$ IQR,

and all outliers (outside $1.5 \times$ IQR) are plotted individually as diamonds. The first section includes the status quo and changes to the first-line therapies, followed by the second section containing MFT approaches, then custom drug rotation strategies and finally triple artemisinin combination therapies.

Our findings suggest that over a 5-year time frame extending the use of AL from 3 to 5 d may hold treatment failure rates at or near the 10% threshold and switching to an MFT strategy is also worth considering with the optimal MFT approach—a 75% ASAQ and 25% DHA-PPQ deployment—projected to hold treatment failure rates to 9.9% after 5 years. Extending the course of treatment with AL has minimal logistical considerations beyond ensuring sufficient quality of doses being distributed and appears to be beneficial compared to the status quo scenario. Although there are concerns regarding the cardiotoxicity of antimalarial drugs^{35,36}, the incidence of adverse cardiac events recorded during clinical trials has been low^{37,38}. Switching to one of several MFT

options where future treatment failure rates can be kept close to 10% will ensure there are no concerns with extended artemisinin dosing, and the success of these MFT deployments will depend on operational capability around the supply and distribution of ACTs. Based on the structure of the distribution network for antimalarials in Rwanda and the current availability of different ACTs, an MFT deployment is more likely to be feasible than a custom rotation approach, although it lacks the logistical ease of a single recommended first-line therapy.

The major uncertainty in these strategy comparisons is that the future course of PPQ resistance in Africa cannot be predicted through any modeling approaches, *in vitro* studies or clinical trials.



Fig. 4 | Comparison of projected treatment failures after 5 years. Box plots ($n = 100$ model replicates per policy, median line with IQR) and violin plots (full data range) show the projected population-level treatment failure rate, after 5 years, under 17 policy options and the status quo of continued usage of 3-d artemether-lumefantrine. Box-plot whiskers show $1.5 \times$ IQR, and all outliers

(outside $1.5 \times$ IQR) are plotted individually as diamonds. The first section includes the status quo and changes to the first-line therapies, followed by the second section containing MFT approaches, then custom drug rotation strategies and finally triple artemisinin combination therapies.

The DHA-PPQ-resistant lineage that emerged in Southeast Asia led to 58% treatment failure under 3 d of DHA-PPQ treatment³⁹, and a number of the key mutations associated with these phenotypes are now known⁴⁰. However, there is no guarantee that the same PPQ-resistance mutations will emerge in Africa. Any DHA-PPQ-centered strategy in Africa should be paired with routine rapid-turnaround molecular surveillance for known markers of PPQ resistance. Similarly, for lumefantrine and amodiaquine, projected scenarios of resistance evolution come with uncertainty, but the effects of currently circulating parasite mutations on the efficacies of AL and ASAQ can be estimated.

Certain new therapeutic approaches to resistance management will need specific national approval or WHO prequalification. Triple ACTs and sequential ACT courses have the advantage of higher total parasite killing than a 3-d ACT and thus an associated lower treatment failure rate. All sequential and triple ACT approaches are projected to keep treatment failure rates below 4% over a 5-year period, with no noticeable advantage seen for sequential approaches despite their higher total dose of artemisinin. A major limitation of our modeling approach is that each therapy's pharmacodynamics and pharmacokinetics are simplified to daily killing and drug elimination rates; a more detailed analysis will be required to understand whether we should

expect true differences in failure rates for these therapies. However, for both triple ACTs and sequential ACT courses, the combination of three drugs used over a short period lowers the probability of multidrug resistance emerging during a treatment course and slows down the spread of artemisinin resistance over the long term. Triple ACTs will likely enter the WHO approval process in the next year, while approval for sequential courses may have to be sought on a case-by-case basis.

To further contextualize this study within the limitations inherent in modeling studies, due to the introduction of AL in 2006, other genomic mutations, such as those associated with reduced lumefantrine susceptibility, are present in Rwanda. The introduction of mutations in the model year 2014 represents a ‘model fitting compromise’ made necessary due to the stochastic nature of the rare mutation emergence process^{24,41} from 2006 to 2014 that has not been systematically captured in any known data collections. Furthermore, while the data and model calibration indicate that 561H initially appeared in the Gasabo district, the design of the simulation renders it incapable of fitting the highly stochastic process that would have driven the initial appearance and spread of 561H. Additionally, although it is possible that the dominant strain of *P.falciparum* circulating in Rwanda may be fitness neutral⁴², the model takes a more traditional assumption and includes a fitness penalty for the 561H mutation.

Pyronaridine–artesunate deployment was not considered in our simulations as pyronaridine-resistant falciparum phenotypes have not yet been described. Assuming similar rates of emergence and similar drops in ACT efficacy due to future pyronaridine resistance, pyronaridine–artesunate would likely make a positive contribution as an addition to any of the MFT strategies shown in Table 2. The relative rankings of the different strategies are expected to stay the same; however, this would need to be confirmed with a new set of simulations. It is unknown how early we should expect to see pyronaridine-resistant parasites and whether there will be cross-resistance with other partner drugs. As with any other deployment of a new antimalarial, regular monitoring of pyronaridine–artesunate efficacy and planned in vitro resistance studies will be crucial to the early identification of resistant phenotypes.

Another major limitation for our model calibration—and any malaria modeling exercise evaluating the last 5 years of epidemiological changes—is that the effects of the COVID-19 pandemic on malaria control are not easily quantified. A mixed-methods study of three high-endemic districts in Rwanda (Gasabo, Kayonza and Rwanamaga) suggests that while the distribution of malaria testing shifted, the overall decline in uncomplicated malaria cases continued through 2020 (ref. 43). The most recent annual case estimates for the fiscal period of July 2021–June 2022 place *P.falciparum* case numbers at around 1 million annually, with 1.16 million reported for calendar year 2021 to the WHO⁴⁴. With these recent gains, resistance management strategies should seek to keep absolute treatment failure counts below 10,000 monthly for the medium term.

Rwanda’s NMCP’s current activities to expand indoor residual spraying activities (from 2–8 districts last decade to 12–15 districts this decade) and to introduce synergistic insecticide-impregnated nets (piloted in 2019–2020 with distribution planned for 2023 (ref. 45)) are likely to reduce biting rates and number of cases in the coming years. A recent pilot study in a rural part of the Gasabo district with high malaria risk showed that larviciding activity had a moderate effect on reducing incidence; the NMCP also expects to expand larviciding as part of a new set of malaria control activities. These interventions may prove beneficial in the fight against drug resistance as they may eliminate pockets of transmission, including drug-resistant alleles, and are likely to slow the overall geographic spread of all genotypes. These activities and their effects are not included in the present modeling analysis.

For the majority of national drug-policy intervention scenarios considered here that are based upon existing therapies or protocols,

treatment failures are projected to exceed 10% within 5 years and may reach as high as 40% if, under a worst-case scenario, DHA-PPQ-resistant phenotypes similar to the known Southeast Asian lineages were to emerge or be imported into Africa (Supplementary Tables 6 and 7). As such, emphasis should be placed on the development of next-generation strategies along with continued evaluation of pyronaridine–artesunate, triple ACTs and sequential ACT courses. According to model projections, the most successful among these choices in slowing down the spread of 561H and reducing treatment failures is likely to be the adoption of triple ACTs following the completion of clinical trials and regulatory approval. Early introduction of triple ACTs, before crossing the WHO first-line therapy treatment failure threshold of 10%, would need to be accompanied by appropriate public health communication and an assessment of acceptability in affected communities⁴⁶. Model projections also show that sequential ACT courses may have comparable treatment failure benefits to triple ACTs, but the prolonged course may result in lower patient adherence.

Without additional interventions targeting drug resistance or general malaria transmission, drug resistance can spread rapidly once it is established, and this could leave NMCPs with limited forewarning to respond to a rise in malaria cases and treatment failures. The identification of 561H in Uganda¹² suggests that the spread of the 561H mutant is currently underway and that this spread is likely to co-occur with that of other markers for artemisinin resistance within sub-Saharan Africa^{11,12,47}. Urgency and speed in setting a new antimalarial policy—a policy that is specifically aimed at containing the spread of artemisinin-resistant *pfkelch13* mutants—are most likely to determine the success of our response to artemisinin resistance in Africa.

Online content

Any methods, additional references, Nature Portfolio reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at <https://doi.org/10.1038/s41591-023-02551-w>.

References

1. Ariey, F. et al. A molecular marker of artemisinin-resistant *Plasmodium falciparum* malaria. *Nature* **505**, 50–55 (2014).
2. Ashley, E. A. et al. Spread of artemisinin resistance in *Plasmodium falciparum* malaria. *N. Engl. J. Med.* **371**, 411–423 (2014).
3. Report on Antimalarial Drug Efficacy, Resistance and Response: 10 Years of Surveillance (2010–2019) (World Health Organization, 2020); <https://apps.who.int/iris/bitstream/handle/10665/336692/9789240012813-eng.pdf>
4. World Malaria Report 2021 (World Health Organization, 2021); <https://www.who.int/teams/global-malaria-programme/reports/world-malaria-report-2021>
5. Uwimana, A. et al. Emergence and clonal expansion of in vitro artemisinin-resistant *Plasmodium falciparum kelch13 R561H* mutant parasites in Rwanda. *Nat. Med.* **26**, 1602–1608 (2020).
6. Uwimana, A. et al. Association of *Plasmodium falciparum kelch13 R561H* genotypes with delayed parasite clearance in Rwanda: an open-label, single-arm, multicentre, therapeutic efficacy study. *Lancet Infect. Dis.* **21**, 1120–1128 (2021).
7. Bergmann, C. et al. Increase in Kelch 13 polymorphisms in *Plasmodium falciparum*, Southern Rwanda. *Emerg. Infect. Dis.* **27**, 294 (2021).
8. van Loon, W. et al. In vitro confirmation of artemisinin resistance in *Plasmodium falciparum* from patient isolates, Southern Rwanda, 2019. *Emerg. Infect. Dis.* **28**, 852–855 (2022).
9. Straimer, J., Gandhi, P., Renner, K. C. & Schmitt, E. K. High prevalence of *Plasmodium falciparum* K13 mutations in Rwanda is associated with slow parasite clearance after treatment with artemether-lumefantrine. *J. Infect. Dis.* **225**, 1411–1414 (2022).

10. Kirby, R. et al. Examining the early distribution of the artemisinin-resistant *Plasmodium falciparum* kelch13 R561H mutation in areas of higher transmission in Rwanda. *Open Forum Infect. Dis.* **10**, ofad149 (2023).
11. Asua, V. et al. Changing molecular markers of antimalarial drug sensitivity across Uganda. *Antimicrob. Agents Chemother.* **63**, e01818-18 (2019).
12. Asua, V. et al. Changing prevalence of potential mediators of aminoquinoline, antifolate, and artemisinin resistance across Uganda. *J. Infect. Dis.* **223**, 985–994 (2021).
13. *Strategy to Respond to Antimalarial Drug Resistance in Africa* (World Health Organization, 2022); <https://www.who.int/publications/item/9789240060265>
14. Lubell, Y. et al. Artemisinin resistance—modelling the potential human and economic costs. *Malar. J.* **13**, 452 (2014).
15. Karem, C. et al. Trends in malaria cases, hospital admissions and deaths following scale-up of anti-malarial interventions, 2000–2010, Rwanda. *Malar. J.* **11**, 236 (2012).
16. Amaratunga, C. et al. Association of mutations in the *Plasmodium falciparum* Kelch13 gene (Pf3D7_1343700) with parasite clearance rates after artemisinin-based treatments—a WWARN individual patient data meta-analysis. *BMC Med.* **17**, 1 (2019).
17. WHO Guidelines for Malaria—14 March 2023 (WHO Global Malaria Programme, 2023); <https://app.magicapp.org/#/guideline/7089>
18. Whegang Youdom, S., Tahar, R. & Basco, L. K. Comparison of anti-malarial drug efficacy in the treatment of uncomplicated malaria in African children and adults using network meta-analysis. *Malar. J.* **16**, 311 (2017).
19. *Seasonal Malaria Chemoprevention with Sulfadoxine-Pyrimethamine Plus Amodiaquine in Children: A Field Guide* (World Health Organization, 2013); https://apps.who.int/iris/bitstream/handle/10665/85726/9789241504737_eng.pdf
20. van Vugt, M. et al. Artemether-lumefantrine for the treatment of multidrug-resistant falciparum malaria. *Trans. R. Soc. Trop. Med. Hyg.* **94**, 545–548 (2000).
21. Tun, K. M. et al. Effectiveness and safety of 3 and 5 day courses of artemether-lumefantrine for the treatment of uncomplicated falciparum malaria in an area of emerging artemisinin resistance in Myanmar. *Malar. J.* **17**, 258 (2018).
22. Onyamboko, M. A. et al. A randomized controlled trial of three- versus five-day artemether-lumefantrine regimens for treatment of uncomplicated *Plasmodium falciparum* malaria in pregnancy in Africa. *Antimicrob. Agents Chemother.* **64**, e01140-19 (2020).
23. U.S. President's Malaria Initiative Rwanda: Malaria Operational Plan FY 2022 (U.S. President's Malaria Initiative, 2021); <https://d1u4sg1s9ptc4z.cloudfront.net/uploads/2022/01/FY-2022-Rwanda-MOP.pdf>
24. Watson, O. J. et al. Pre-existing partner-drug resistance to artemisinin combination therapies facilitates the emergence and spread of artemisinin resistance: a consensus modelling study. *Lancet Microbe* **3**, e701–e710 (2022).
25. Nguyen, T. D., Tran, T. N.-A., Parker, D. M., White, N. J. & Boni, M. F. Antimalarial mass drug administration in large populations and the evolution of drug resistance. *PLoS Glob. Public Health* **3**, e0002200 (2023).
26. Boni, M. F., Smith, D. L. & Laxminarayan, R. Benefits of using multiple first-line therapies against malaria. *Proc. Natl Acad. Sci. USA* **105**, 14216–14221 (2008).
27. Boni, M. F., White, N. J. & Baird, J. K. The community as the patient in malaria-endemic areas: preempting drug resistance with multiple first-line therapies. *PLoS Med.* **13**, e1001984 (2016).
28. Boni, M. F. Breaking the cycle of malaria treatment failure. *Front. Epidemiol.* <https://doi.org/10.3389/fepid.2022.1041896> (2022).
29. Schallig, H. D. et al. Randomised controlled trial of two sequential artemisinin-based combination therapy regimens to treat uncomplicated falciparum malaria in African children: a protocol to investigate safety, efficacy and adherence. *BMJ Glob. Health* **2**, e000371 (2017).
30. van der Pluijm, R. W. et al. Triple artemisinin-based combination therapies versus artemisinin-based combination therapies for uncomplicated *Plasmodium falciparum* malaria: a multicentre, open-label, randomised clinical trial. *Lancet* **395**, 1345–1360 (2020).
31. Kunkel, A., White, M. & Piola, P. Novel anti-malarial drug strategies to prevent artemisinin partner drug resistance: a model-based analysis. *PLoS Comput. Biol.* **17**, e1008850 (2021).
32. Nguyen, T. D. et al. Preventing antimalarial drug resistance with triple artemisinin-based combination therapies. *Nat. Commun.* **14**, 4568 (2023).
33. Cohen, J. L., Yavuz, E., Morris, A., Arkedis, J. & Sabot, O. Do patients adhere to over-the-counter artemisinin combination therapy for malaria? evidence from an intervention study in Uganda. *Malar. J.* **11**, 83 (2012).
34. Banek, K., Webb, E. L., Smith, S. J., Chandramohan, D. & Staedke, S. G. Adherence to treatment with artemether-lumefantrine or amodiaquine-artesunate for uncomplicated malaria in children in Sierra Leone: a randomized trial. *Malar. J.* **17**, 222 (2018).
35. White, N. J. Cardiotoxicity of antimalarial drugs. *Lancet Infect. Dis.* **7**, 549–558 (2007).
36. Chan, X. H. S. et al. The cardiovascular effects of amodiaquine and structurally related antimalarials: an individual patient data meta-analysis. *PLoS Med.* **18**, e1003766 (2021).
37. Haesler, I. L., Chan, X. H. S., Guérin, P. J. & White, N. J. The arrhythmogenic cardiotoxicity of the quinoline and structurally related antimalarial drugs: a systematic review. *BMC Med.* **16**, 200 (2018).
38. Whalen, M. E. et al. The impact of extended treatment with artemether-lumefantrine on antimalarial exposure and reinfection risks in Ugandan children with uncomplicated malaria: a randomized controlled trial. *Clin. Infect. Dis.* **76**, 443–452 (2023).
39. Witkowski, B. et al. A surrogate marker of piperaquine-resistant *Plasmodium falciparum* malaria: a phenotype–genotype association study. *Lancet Infect. Dis.* **17**, 174–183 (2017).
40. Amato, R. et al. Origins of the current outbreak of multidrug-resistant malaria in southeast Asia: a retrospective genetic study. *Lancet Infect. Dis.* **18**, 337–345 (2018).
41. Pongtavornpinyo, W. et al. Probability of emergence of antimalarial resistance in different stages of the parasite life cycle. *Evol. Appl.* **2**, 52–61 (2009).
42. Stokes, B. H. et al. *Plasmodium falciparum* K13 mutations in Africa and Asia impact artemisinin resistance and parasite fitness. *eLife* **10**, e66277 (2021).
43. Hakizimana, D. et al. The impact of COVID-19 on malaria services in three high endemic districts in Rwanda: a mixed-method study. *Malar. J.* **21**, 48 (2022).
44. World Malaria Report 2022 (World Health Organization, 2022); <https://www.who.int/teams/global-malaria-programme/reports/world-malaria-report-2022>
45. Gansané, A. et al. Design and methods for a quasi-experimental pilot study to evaluate the impact of dual active ingredient insecticide-treated nets on malaria burden in five regions in sub-Saharan Africa. *Malar. J.* **21**, 19 (2022).
46. Tindana, P. et al. Deploying triple artemisinin-based combination therapy (TACT) for malaria treatment in Africa: ethical and practical considerations. *Malar. J.* **20**, 119 (2021).
47. Balikagala, B. et al. Evidence of artemisinin-resistant malaria in Africa. *N. Engl. J. Med.* **385**, 1163–1171 (2021).

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this

article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit <http://creativecommons.org/licenses/by/4.0/>.

© The Author(s) 2023

Methods

Model description

A previously validated spatial, stochastic, individual-based model was used as the basis for the study^{48,49}, and new model calibration and validation was performed to match the malaria prevalence of the 30 administrative districts of Rwanda. The MAP mean $PfPR_{2-10}$ projections for 2017 (ref. 50) were used as the basis to calibrate local transmission parameters on a 5-by-5 km (25 km²) scale in Rwanda (Extended Data Fig. 1), followed by a switch to the malaria incidence in 2021 using aggregate data reported by the Ministry of Health²³. The seasonal variation in malaria transmission was coupled to seasonal rainfall (Extended Data Fig. 5a,b), consistent with the general coupling of seasonal transmission and increased rainfall. Treatment seeking and treatment coverage data were obtained from the 2019–2020 Demographic and Health Survey⁵¹, with treatment coverage ranging from 53.3% to 71.8% across provinces. Under the baseline model, or status quo conditions, all treated individuals in the model receive a 3-d course of AL and fully comply with the course of treatment. No private market drugs are incorporated, consistent with survey results⁵¹. During model execution, individuals move around the simulated landscape in a manner consistent with previous travel studies for sub-Saharan Africa^{52,53} and carry any *P.falciparum* clones to these destinations. The carried clones may then enter into circulation within a new region if the traveling individual is bitten by a mosquito, which then proceeds to infect another individual.

With these prevalence and treatment calibrations, the model produces a symptomatic malaria incidence ranging from 9.92 per 1,000 (Burera district) to 465.62 per 1,000 (Gisagara district) in 2020 under the MAP $PfPR_{2-10}$ projections, and 5.49 per 1,000 (Burera district) to 410.26 per 1,000 (Nyamasheke district) in 2022 following the switch to the incidence-based prevalence calibration. This results in an incidence of 139.13 per 1,000 for symptomatic cases and an incidence of 87.76 per 1,000 for treated cases at the national level in 2022, consistent with reporting that the incidence rate has been declining from a high of 403 per 1,000 in 2016 (ref. 23; Extended Data Fig. 5c,d). All policy interventions were introduced on 1 January 2024, of the simulation using version 4.1.4 of the simulation.

After calibration of prevalence, incidence and treatment coverage, the simulation's genotype frequency and trajectory were matched to known observations of 561H allele frequencies. At the time of calibration, nine data points of the *pfkelch* 561H mutation were available at the district level—four close to the capital Kigali^{5,6,9}, two from the eastern district of Kayonza^{5,6} and one each from the districts of Huye⁷, Kirehe and Ngoma¹⁰. To calibrate to these values, the simulation was seeded with a single mutation of 561H genotypes in Gasabo district (Kigali province) to generate a slowly growing exponential curve with district frequencies that are consistent with measured values (Fig. 1). This results in a mean national frequency ranging from 0.01 in 2014 to 0.12 in 2020. In the event that the seeded 561H genotypes went extinct before 2014, the replicate was discarded from analysis.

Because the mutation rate to 561H alleles is not known, these artificially introduced 561H genotypes are the only means by which 561H can be introduced into the simulation. The spatial spread of 561H is driven by human movement and migration within the simulation, which is based on the gravity model described in ref. 53 for sub-Saharan Africa and presumes that major cities (that is, Kigali) will have an oversized effect on human movement dynamics. To account for the ability of ACT partner-drug resistance to accelerate the fixation of artemisinin resistance²⁴, mutations affecting other alleles (for example, *pfcrt*, *pfmdr1*, etc.) are enabled in the simulation on 1 January 2014, following the completion of model burn-in using a previously calibrated mutation rate²⁴. This produces a slight model delay in *pfcrt* and *pfmdr1* mutations that are associated with the use of lumefantrine given the adoption of AL by Rwanda in 2006, likely resulting in a slight model bias toward low frequencies of alleles associated with lower lumefantrine susceptibility.

Model calibration and validation. The following three metrics were used for model calibration and validation: the population-weighted, district-level, annual mean $PfPR_{2-10}$ as projected by MAP⁵⁰ versus the simulated $PfPR_{2-10}$ before 2021; the district-level clinical cases projected by the simulation and the projected 561H frequency. To calibrate the $PfPR_{2-10}$, the local transmission intensity, or β , was determined using a constrained parameter space search. This was performed by first binning the population in each 25 km² cell using Jenks natural breaks optimization followed by scanning the domain of possible β values using a fixed population (that is, the bin size) along with the relevant population and climatic variables. Upon determining the possible β values, they were assigned to cells in the model by matching the population in each cell to the appropriate bin and assigning the calculated transmission intensity. Once this process was complete, the model was run and assessed against a target deviation of MAP $PfPR_{2-10}$ values to within ±10% (Extended Data Fig. 5c), although for low-prevalence districts, matching the MAP projections was a challenge when accompanied by a low population due to the increase in stochasticity present in the model. We found the overall $PfPR_{2-10}$ calibration to be acceptable, with low-prevalence districts having a simulated $PfPR_{2-10}$ that skewed slightly higher than the reference, whereas the higher prevalence districts skewed slightly lower.

The next model validation metric is a comparison of the district-level projections for clinical cases versus the true incidence for 2017 (ref. 54). Starting with the total clinical cases per 1,000, it is clear that the spatial distribution of model-generated clinical cases is distributed in a manner that is consistent with 2017 true incidence⁵⁰, although the counts of both all clinical cases (Extended Data Fig. 6) and treated cases (Extended Data Fig. 7) are lower in the model than in the reported figures²³. However, these lower model projections are consistent with the overall decline in malaria cases in Rwanda⁵⁵ and also support the simulation transition from using a calibration based on MAP projections to one based on incidence projections during model execution at the start of 2021. Overall, the good agreement between the projected $PfPR_{2-10}$ and reference values supports model calibration as being within acceptable bounds.

The final point of model calibration and validation is the 561H frequency. Presently, 561H frequency data are only available for Huye, Kayonza, Kirehe and Ngoma districts, along with the province of Kigali, consisting of Gasabo, Kicukiro and Nyarugenge districts (Supplementary Table 1). As a result, the model was calibrated to use a single introduction event on 30 September 2004, in which 6% of infected individuals in any of the cells of Gasabo district had their parasites switched from R561 to 561H. This date and quantity of mutations were selected on the basis of a parameter space search, which is the only introduction of 561H in the simulation. Although this introduction results in a generally consistent spread of the mutation, outliers are still possible (Extended Data Fig. 8). To ensure that replicates used for analysis are in good agreement with observed data, only replicates in which the 561H frequency in Gasabo is greater than 0.01 in September 2014 of the simulation are retained. The model's spread of mutations is consistent with measured allele frequencies between 2014 and 2019 (refs. 5–7,9), supporting the model as having a 561H introduction that is properly calibrated. As a note of caution, although this calibration suggests a mechanism through which 561H may have spread in Rwanda, the model was not designed, configured or calibrated to explore the nature of the introduction event or original mutation event.

Model scenarios. As a common point of comparison, a baseline scenario was run in which the calibrated model was simulated for a 10-year window past the proposed point of intervention (that is, 1 January 2024 to 1 January 2034). This baseline scenario also controls for any deviations in the model versus real-world data for Rwanda by allowing all policy interventions to be compared to the same projected outcome and presumes that the calibrated malaria incidence remains stable over

time. A total of 26 drug-policy intervention scenarios were evaluated within the simulation (Table 2), and all interventions were introduced at the same time with no delay between the introduction and the change in therapies received by individuals. The 5-year endpoints are presented in the main text, with 3- and 10-year endpoints referred to as needed; comprehensive results for all endpoints can be found in Supplementary Tables 2 and 5.

The policy scenarios run may be summarized as follows. Five evaluated interventions involved a change in first-line therapy to a 4-d course of AL, 5-d course of AL, ASAQ and DHA–PPQ. Nine of the interventions involved MFT approaches with drug distribution proportions ranging from 25/75 to 50/50 to 75/25 (Table 2), with selection of the drug given to the individual based upon a random draw upon first treatment. Functionally this can be imagined working the same as a provider randomly assigning a therapy to a patient via coin flip, randomization schedule or other means. Other MFT implementations with different approaches to drug distributions are possible²⁸ but were not evaluated here. Two interventions considered drug rotation with short-term DHA–PPQ use first (for 3 years) followed by replacement of DHA–PPQ with one of two MFT strategies using AL and ASAQ. Four interventions replicated the therapeutic arms of the sequential ACT therapy regimen proposed in ref. 29 with AL followed by either ASAQ or DHA–PPQ, or AL preceded by ASAQ or DHA–PPQ. However, the model scenario deviates from the protocol proposed in ref. 29 by applying sequential therapy to all treatment-seeking individuals as opposed to just children aged 6–120 months. The second of the sequential therapies was given either on treatment days 3, 4 and 5 or on treatment days 7, 8 and 9; results for both timings of treatment courses are presented. Finally, two interventions considered the replacement of AL with one of two triple therapies (ALAQ and ASMQ–PPQ), which are yet to be approved^{30,46}.

Sensitivity analysis. Five forms of sensitivity analysis were performed using the same base calibration previously described, with the relevant parameters adjusted as needed. These studies included assessing the sensitivity to individual movement, the fitness cost associated with drug-resistance mutations and the impact of individual compliance with drug treatments. All studies used at least 100 replicates per permutation in assessing the results.

First, the model sensitivity to individual movement was assessed by adjusting the movement to be 0.3, 0.5, 2 or 3 times more than the calibrated movement. Under these conditions, it was found that a movement rate less than the calibrated value resulted in projected 561H frequencies that were higher than observed in Gasabo while lower in other districts (Supplementary Figs. 1 and 2). However, higher movement rates (that is, 2× and 3×) have a lower variance and plausibly reproduced the observed 561H frequencies for Gasabo, Kayzona, Kicukiro and Nyarugenge districts, although the projected frequencies were substantially higher for Huye (Supplementary Figs. 3 and 4). These results show that the calibrated individual movement rate is consistent with the observed geographical frequency of 561H, although it is possible that the true individual movement rate may be slightly faster.

Next, the model sensitivity to fitness cost was assessed by adjusting all fitness costs to be 10, 25 or 50 times greater than previously calibrated values. Given that there is evidence that there is a 561H mutant strain of *P.falciparum* circulating in Rwanda that does not incur a fitness penalty⁴², lower values for the fitness cost were not assessed. Under the 10× scenario, it was found that the initial mutation event needed to be increased to 20% of the infected individuals in the Gasabo district, but the 561H frequency was similar to the observed data and the projection was similar to the model calibration (Supplementary Fig. 5). This suggested that the calibrated fitness penalty used in the model, based upon previous modeling exercises²⁴, is reasonable and the model projections would still remain valid when the actual fitness cost incurred by the 561H mutant is up to 10× the calibrated value. However, when increasing the fitness penalty to 25×, it is necessary to

increase the initial mutation event to be 50% of the infected individuals in the Gasabo district (this is unrealistic) and the allele frequency trajectories only weakly tracked the observed data points (Extended Data Fig. 9 and Supplementary Fig. 6). Under the 50× scenario, the initial mutation event was increased to 75% of the infected individuals in the Gasabo district and the projected frequencies suggest that 561H would only exist in Rwanda at a low frequency, or would soon face extinction (Supplementary Fig. 7). It is highly unlikely that higher fitness costs of 25× or 50× are plausible.

The model sensitivity to individual drug treatment compliance was assessed by administering ASAQ, DHA–PPQ and 3-, 4- or 5-d courses of AL to individuals using low (25–70%), moderate (50–80%) and high (70–90%) compliance rates for complete courses (Supplementary Table 4). Each scenario takes effect on 1 January 2024, and presumes that the individual will always complete the first day of treatment, and compliance rates then proceed to drop after that point. As expected, across all treatment options, failure to comply with the prescribed course of treatment results in higher treatment failures when compared to the full compliance outcomes. Under the 3-, 4- and 5-d AL treatment options, along with ASAQ, the number of treatment failures increases with the rate of noncompliance.

Because PPQ resistance may evolve differently in Rwanda than what was observed in Southeast Asia, the model's sensitivity to PPQ resistance was evaluated by first evaluating the sensitivity to the gene duplication rate for *plasmepsin 2/3* genes. Using the calibrated value based upon the emergence of PPQ resistance in Southeast Asia as the baseline²⁵, the probability of gene duplication was adjusted to be 0.25, 0.5, 1.25 and 1.5 times the calibrated value. These represent optimistic scenarios in which African strains of *P.falciparum* are less likely to develop PPQ resistance (0.25× and 0.5× scenarios) and pessimistic scenarios in which African strains are more likely to develop PPQ resistance (1.25× and 1.5× scenarios). For all four scenarios, the annual percentage of treatment failures exceeds the 10% treatment failure threshold within 5 years after switching from AL to DHA–PPQ (Supplementary Table 3). Ten years after the introduction of DHA–PPQ, the annual percentage of treatment failures was approximately 55% with 561H fixed for all scenarios, indicating that PPQ efficacy is a critical value to monitor.

Finally, the sensitivity of the model to changes in PPQ efficacy on PPQ-resistant parasites was evaluated by altering the configured EC₅₀ value (concentration at which killing is 50% of maximum). The details of the pharmacokinetic/pharmacodynamic model are described in ref. 48, and the model uses calibrated values of 0.58 for PPQ-sensitive and 1.4 for PPQ-resistant parasites, where 1.0 is set as the standard therapeutic dose given in the simulation. The simulation was then run with values ranging from 1.0 to 1.6 for the EC₅₀ of PPQ. As expected, an EC₅₀ value less than the configured value of 1.4 had fewer treatment failures at 5 years while 1.5 and 1.6 produced more treatment failures (Supplementary Table 3). This pattern was similar for the 561H frequency, although the difference was only statistically significant for EC₅₀ ≤ 1.2 with $P < 10^{-4}$. For all scenarios, the model projects that the 10% treatment failure threshold would still be reached within 5 years of DHA–PPQ introduction and 561H would be fixed within 10 years.

Statistical analysis. Differences between groups of simulation results (that is, comparing 100 simulations each from two scenarios) are tested with nonparametric Wilcoxon rank-sum tests, and all P values lower than 10^{-4} are shown as 10^{-4} .

Model calibration data

Spatial data. A 5-by-5 km (25 km²) cell is the primary spatial unit in the simulation, the same size as the resolution of the MAP *PfPR*₂₋₁₀ projections used as part of the model calibration⁵⁰. This cellular resolution results in a modeled space of 979 cells covering 24,475 km² or about 93% of the total area of Rwanda (Extended Data Fig. 1a). Cells that are

predominately water are not simulated, and some discrepancy in area is due to clipping along national borders. During model initialization, spatial data in the form of national districts, population, access to treatment and the beta (that is, transmission parameter) for each cell are loaded. Administrative and geographic boundaries from the World Bank^{56,57}, the Rwanda National Institute of Statistics⁵⁸ and the World Database on Protected Areas⁵⁹ were used in the preparation of model inputs (Fig. 1 and Extended Data Figs. 1, 6 and 7).

Demographics, mortality and treatment seeking. The estimated population of Rwanda was 12,663,116 in 2020 with a crude birth rate of 28.8 per 1,000 (ref. 60), and the population skews younger with about 54.3% being under the age of 20 years⁶¹ (Supplementary Table 1). Deaths due to malaria continue to have a substantial impact on the mortality rates within Rwanda⁶², and the mortality rate applied to the population was adjusted to remove the deaths that were attributed to malaria using the UN population projections⁶³ (Supplementary Tables 2 and 3). In the event an individual reaches the age of 100 years, they are removed from the simulation. When individuals are infected with the parasite, upon exhibiting clinical symptoms, they seek treatment based upon the surveyed treatment-seeking behavior for the province that contains the cell they are currently in with no distinction between under-5 and over-5 treatment-seeking rates⁵¹ (Supplementary Table 4).

Seasonal transmission. To account for the seasonal variation in transmission intensity, a seasonality calibration was conducted, and the results were applied to all cells during model execution. This seasonality calibration is based upon the correlation between rainfall and the *Anopheles* mosquito prevalence and presumes that transmission will begin to increase as favorable conditions increase. Thus, the transmission intensity is adjusted up, or down, by applying the seasonal adjustment value (Extended Data Fig. 5b). This adjustment was calculated by first calculating the 10-year daily average rainfall for Rwanda, using data from ERA5 global climate and weather projections⁶⁴. The rainfall data were then clipped to the borders of Rwanda using Google Earth Engine for January 2009 to December 2019, inclusive, and then smoothed and shifted by 10 d using a MATLAB script. The 10-d offset was selected on the basis of the *Anopheles gambiae* lifecycle. Finally, a lower bound of 0.4 was used to ensure that some malaria transmission is always present at a rate consistent with seasonal patterns. The calculated adjustment is then imported during model initialization and applied during execution.

Drug efficacy. As part of the model calibration and validation, the drug efficacies were used as previously calculated²⁵ following verification given the Rwandan population distribution. Within the current simulation, a population of 50,000 individuals, with a population distribution consistent with that of Rwanda, was infected with *P. falciparum* infections. Following transition of the infection from the liver stage to the blood stage (that is, clinical symptoms), the relevant therapy was given to the individual and the efficacy was assessed using the parasite density at 28 d after the first treatment. Individuals with a parasite density less than 10 ml⁻¹ of blood were counted as cleared (failed otherwise). Drug efficacies of longer courses—for example, 5-d AL or two sequential consecutive courses of ACT—are calculated via a one-compartment pharmacokinetic model and traditional Hill-function pharmacodynamic model, using a 1-d time step⁴⁸.

The lowest efficacy in the simulation is that of DHA–PPQ on the 561H genotype, which also carries PPQ resistance (43.3% efficacy); this is also the reason that PPQ resistance spreads so quickly in the model simulations. The efficacy of 3-d AL on wild-type *P. falciparum* is 95.5%, while the efficacy of 5-d AL is 97.5%. The complete drug efficacies are included in Supplementary Tables 5 and 6 and were within $\pm 2.5\%$ of the original efficacies calibrated in ref. 25.

Ethics and inclusion in global research. This study included researchers from the Rwanda Biomedical Research Center (who were colocated and in close collaboration with Rwanda's NMCP). Under the leadership of A. Uwimana and C. Ngabonziza, they planned the major study-design aspects for this paper and assisted the mathematical modeling team in identifying drug-resistance-management strategies that could be feasibly implemented in Rwanda. Research roles and responsibilities were discussed throughout 2022, including during in-person meetings in August and October 2022. Capacity co-enhancement collaborations have already begun by planning (1) a local modeling instance to be run in RBC Kigali with M. Kabera as the primary analyst and (2) a series of tutorials for Penn State scientists to understand the operational aspects of drug delivery and distribution in Rwanda.

Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

Data availability

Intermediate data files produced by the simulation can be found on GitHub at <https://github.com/bonilab/malariaibm-spatial-Rwanda-561H/tree/main/Data>. The configuration files used for the study described in this paper can be found at <https://github.com/bonilab/malariaibm-spatial-Rwanda-561H/tree/main/Studies>. The prevalence data for 2017, released by the MAP (version 2019), has been archived under <https://github.com/bonilab/malariaibm-spatial-Rwanda-561H/tree/main/Data/GIS/MAP>. The spatial distribution of population is derived from the WorldPop 2015 spatial distribution of population in Rwanda and can be found at <https://hub.worldpop.org/doi/10.5258/SOTON/WP00674>.

Code availability

The archived version of the simulation code base, complied binaries for Linux and scripts used for analysis and production of figures presented in this paper can be found on GitHub at <https://github.com/bonilab/malariaibm-spatial-Rwanda-561H>.

References

48. Nguyen, T. D. et al. Optimum population-level use of artemisinin combination therapies: a modelling study. *Lancet Glob. Health* **3**, e758–e766 (2015).
49. Zupko, R. J. et al. Long-term effects of increased adoption of artemisinin combination therapies in Burkina Faso. *PLoS Glob. Public Health* **2**, e0000111 (2022).
50. Weiss, D. J. et al. Mapping the global prevalence, incidence, and mortality of *Plasmodium falciparum*, 2000–17: a spatial and temporal modelling study. *Lancet* **394**, 322–331 (2019).
51. Demographic and Health Survey 2019/2020—Key Indicators (National Institute of Statistics of Rwanda, 2020); <https://www.statistics.gov.rw/publication/demographic-and-health-survey-20192020-key-indicators>
52. Wesolowski, A. et al. Quantifying travel behavior for infectious disease research: a comparison of data from surveys and mobile phones. *Sci. Rep.* **4**, 5678 (2014).
53. Wesolowski, A., O'Meara, W. P., Eagle, N., Tatem, A. J. & Buckee, C. O. Evaluating spatial interaction models for regional mobility in Sub-Saharan Africa. *PLoS Comput. Biol.* **11**, e1004267 (2015).
54. Rwanda Malaria Operational Plan FY 2019 (U.S. President's Malaria Initiative, 2019); <https://www.pmi.gov/where-we-work/rwanda/>
55. Rwanda Malaria Operational Plan FY 2020 (U.S. President's Malaria Initiative, 2020); <https://www.pmi.gov/where-we-work/rwanda/>
56. Rwanda Districts (World Bank Group, 2018); <https://datacatalog.worldbank.org/search/dataset/0041453>

57. World Boundaries GeoDatabase (World Bank Group, 2020); <https://datacatalog.worldbank.org/search/dataset/0038272/World-Bank-Official-Boundaries>
58. Rwanda—Water Bodies (Lakes) (OCHA, 2018); <https://data.humdata.org/dataset/rwanda-water-bodies?>
59. Protected Planet: The World Database on Other Effective Area-Based Conservation Measures (WD-OECM) (UNEP-WCMC and IUCN, 2023); <https://resources.unep-wcmc.org/products/4c1733823f2a451e8d5ecbaef3f1a06>
60. Rwanda Vital Statistics Report—2020 (National Institute of Statistics of Rwanda, 2021); <http://www.statistics.gov.rw/publication/1705>
61. Rwanda Malaria Indicator Survey (RMIS), Final Report 2017 (Malaria and Other Parasitic Diseases Division of the Rwanda Biomedical Center Ministry of Health, 2018); <https://dhsprogram.com/publications/publication-MIS30-MIS-Final-Reports.cfm>
62. Rwanda Health Sector Performance Report 2017–2019 (Republic of Rwanda, Ministry of Health, 2020); https://www.moh.gov.rw/fileadmin/user_upload/Moh/Publications/Reports/FINAL_Annual_Report_2017-2019_02062020.pdf
63. World Population Prospects 2019: Volume II: Demographic Profiles (United Nations, 2019); https://population.un.org/wpp/Publications/Files/WPP2019_Volume-II-Demographic-Profiles.pdf
64. Hersbach, H. et al. Complete ERA5 from 1940: fifth generation of ECMWF atmospheric reanalyses of the global climate. *Copernicus* <https://doi.org/10.24381/cds.143582cf> (2023).

Acknowledgements

This work was supported by the National Institutes of Health under grant NIAID R01AI153355 (to M.F.B., R.J.Z., T.D.N., T.N.-A.T. and K.T.T.) and Bill and Melinda Gates Foundation under grant INV-005517 (to M.F.B., R.J.Z., T.D.N., T.N.-A.T., K.T.T. and H.L.). The funders had no role in study design, data collection and analysis, decision to publish or preparation of the manuscript. Computations for this research

were performed on the Pennsylvania State University's Institute for Computational and Data Sciences' Roar supercomputer.

Author contributions

A.U., M.F.B. and R.J.Z. designed the study. R.J.Z. designed and implemented the spatial epidemiological model, ran all scenario evaluations for controlling the R561H mutation, processed the simulation output and drafted the manuscript. R.J.Z., T.D.N. and K.T.T. integrated the spatial malaria model into an existing validated individual-based malaria model. R.J.Z. and H.L. calibrated the spatial model to malaria prevalence and incidence data from Rwanda. M.F.B., R.J.Z., A.U., J.C.S.N. and M.K. designed scenarios for model evaluation. T.N.-A.T. provided genotype-specific efficacy estimates. All authors edited and approved the final manuscript.

Competing interests

The authors declare no competing interests.

Additional information

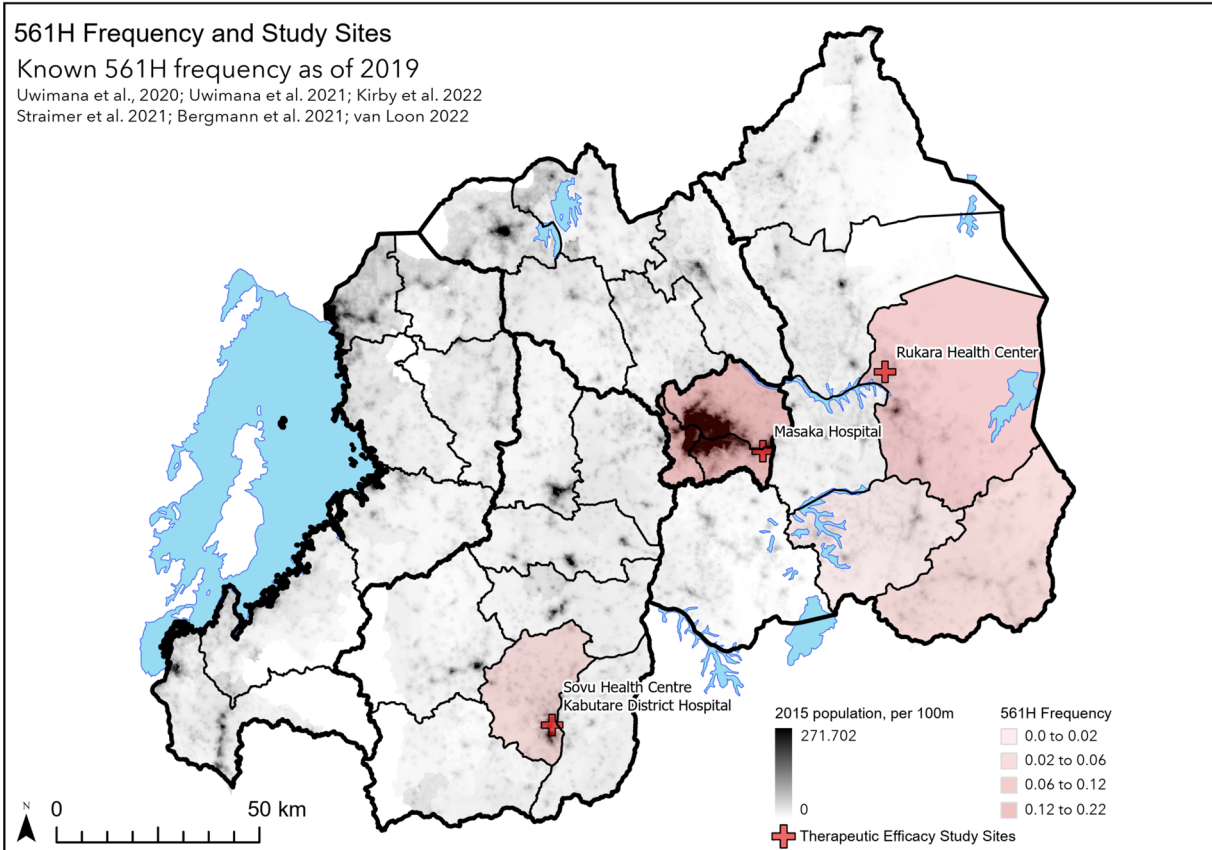
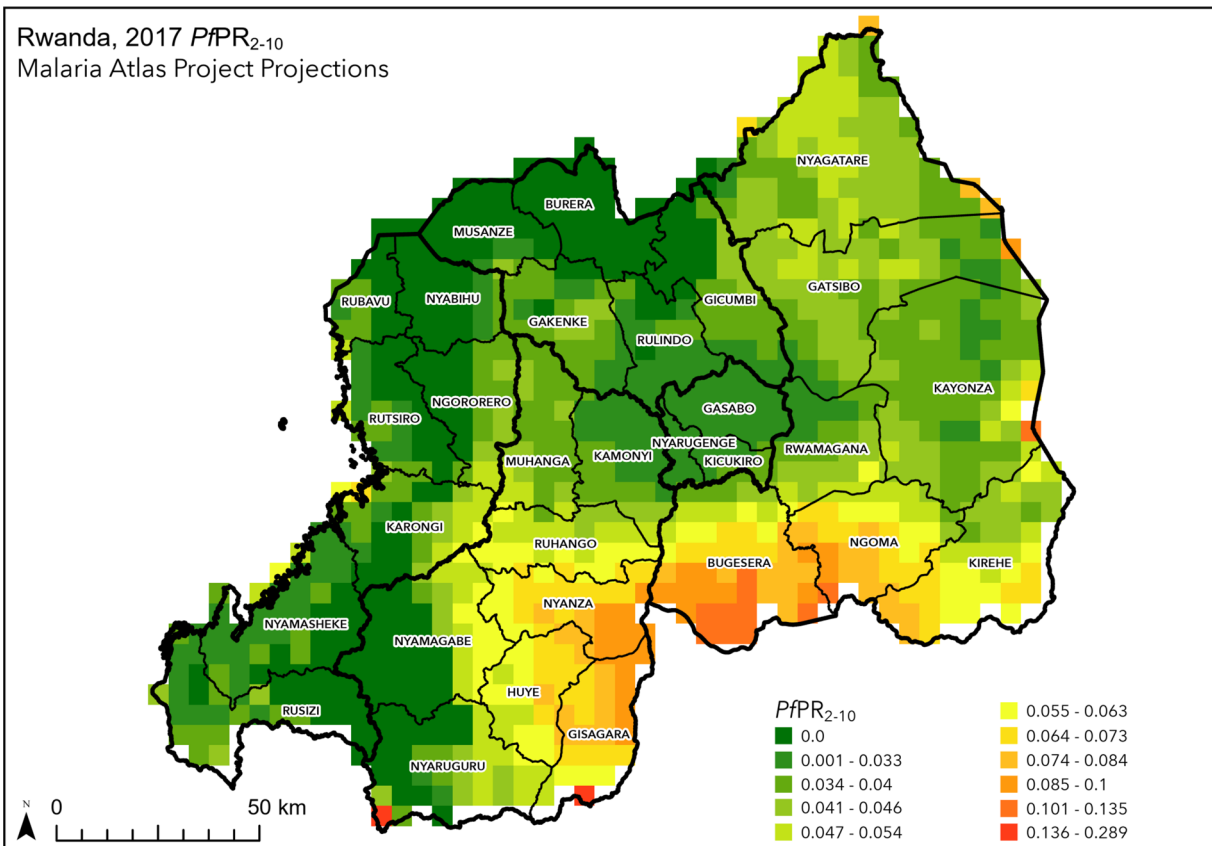
Extended data is available for this paper at <https://doi.org/10.1038/s41591-023-02551-w>.

Supplementary information The online version contains supplementary material available at <https://doi.org/10.1038/s41591-023-02551-w>.

Correspondence and requests for materials should be addressed to Robert J. Zupko.

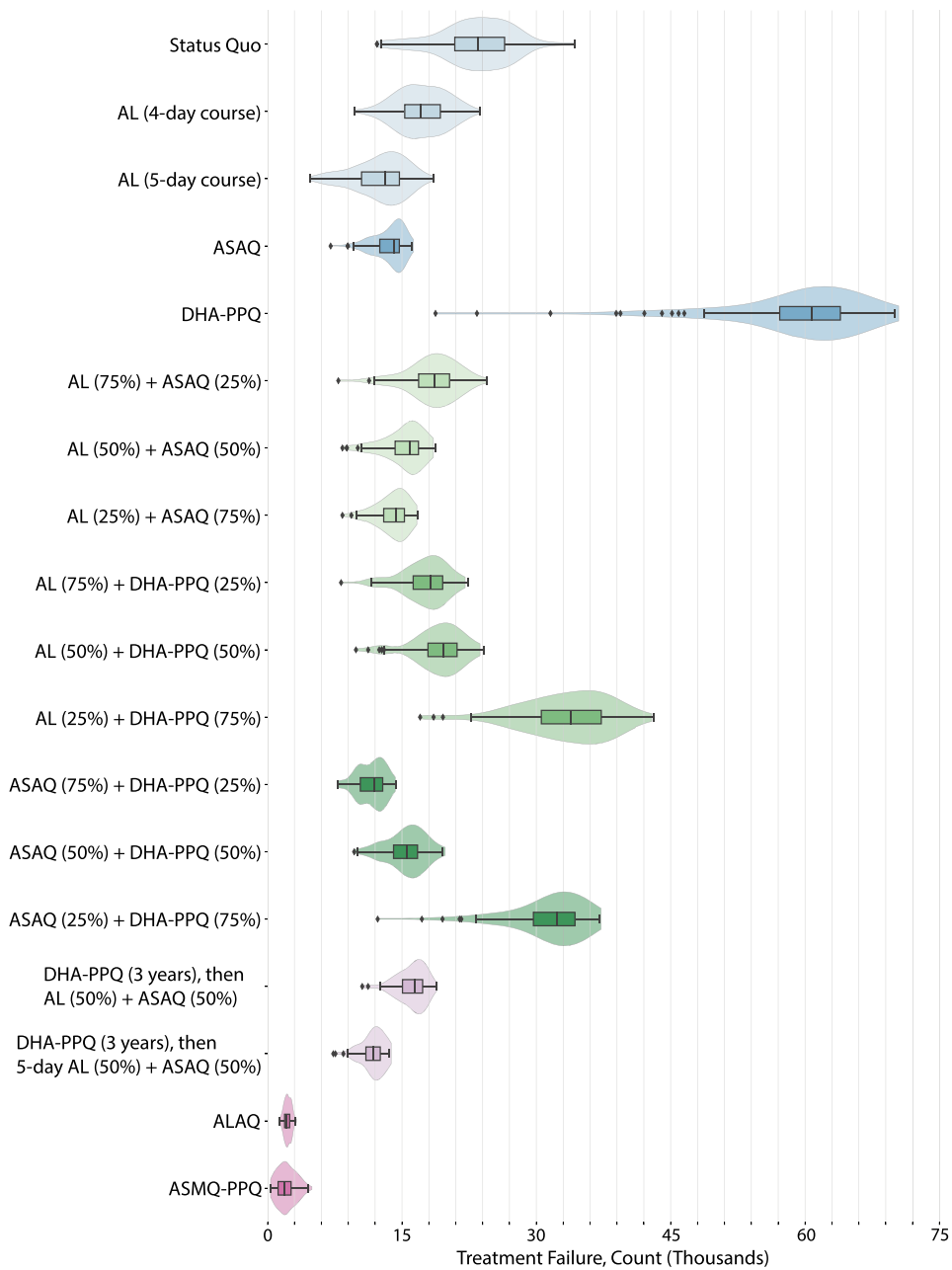
Peer review information *Nature Medicine* thanks Philip Rosenthal, Rashid Ansumana and the other, anonymous, reviewer(s) for their contribution to the peer review of this work. Primary Handling Editor: Ming Yang, in collaboration with the *Nature Medicine* team.

Reprints and permissions information is available at www.nature.com/reprints.



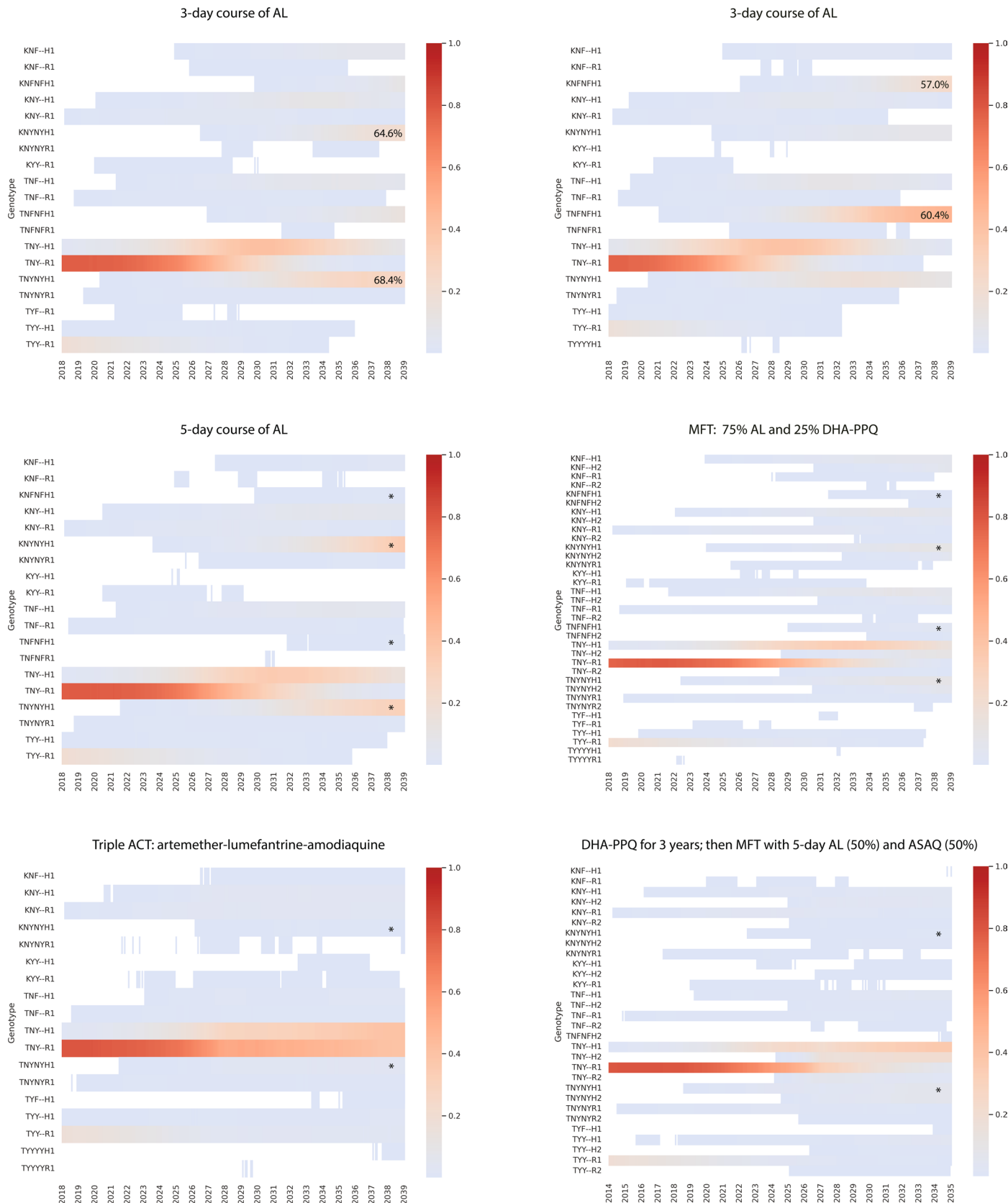
Extended Data Fig. 1 | Plasmodium falciparum prevalence and allele frequency of R561H in Rwanda. (Top) Estimates of *P. falciparum* prevalence in 2-to-10 year-olds ($PfPR_{2-10}$) from Malaria Atlas Project⁵⁰ where each cell is 5 km × 5 km (25 km²). Highest $PfPR_{2-10}$ levels above 10% are seen in the south-central

part of the country. (Bottom) Map showing five districts in Rwanda where 561H alleles have been detected, as part of therapeutic efficacy studies (TES) or cross-sectional molecular surveys. Red crosses show sites of TESs, population is per 100 m grid cell.



Extended Data Fig. 2 | Projected monthly treatment failure counts in 2028 under various treatment strategies. Box plots ($n = 100$ model replicates per policy, median line with interquartile range (IQR) and violin plots of treatment

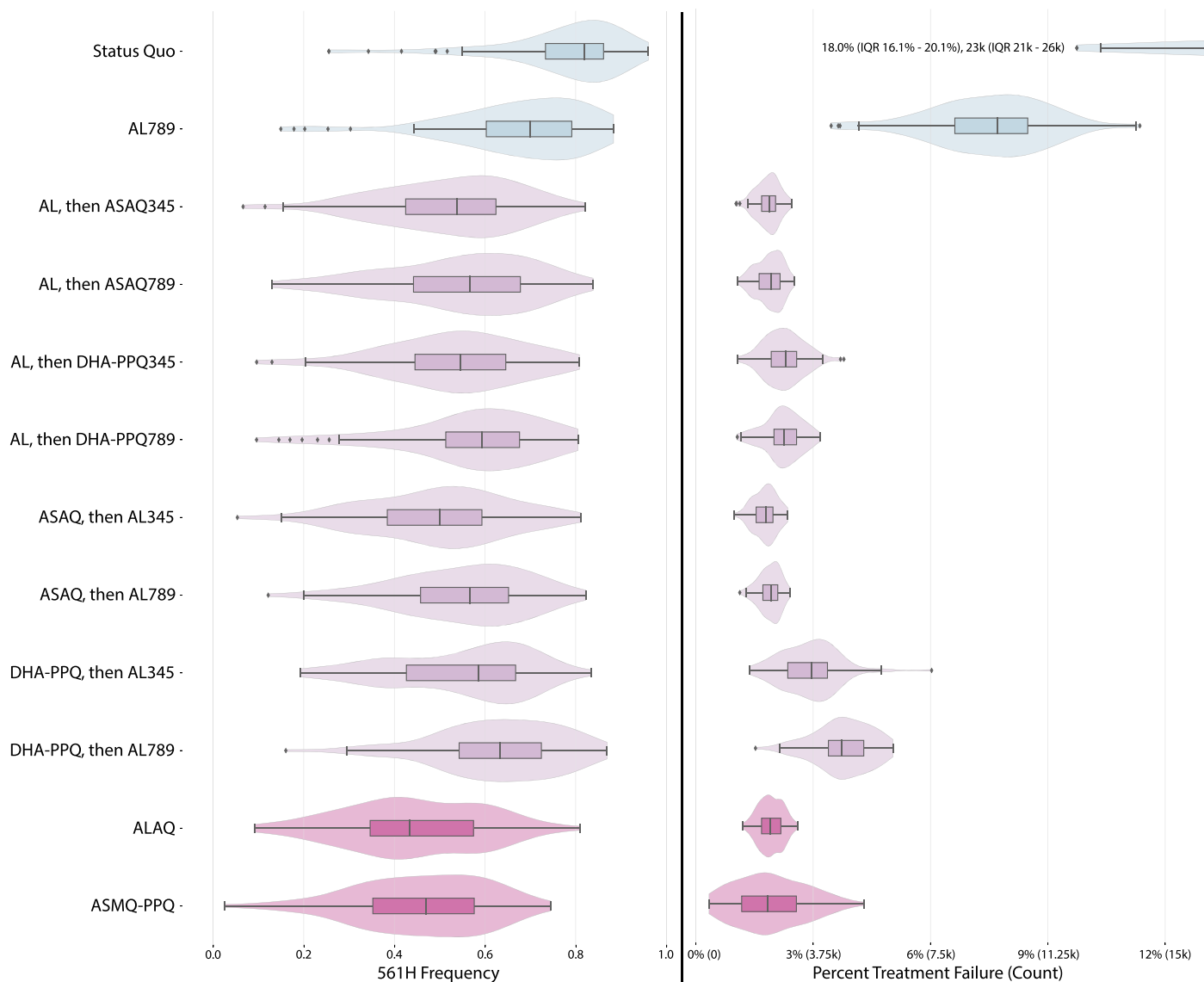
failure count for the same strategies as shown in Figs. 3 and 4 of main text. Whiskers show 1.5 times IQR while all outliers (outside $1.5 \times$ IQR) are plotted individually as diamonds.

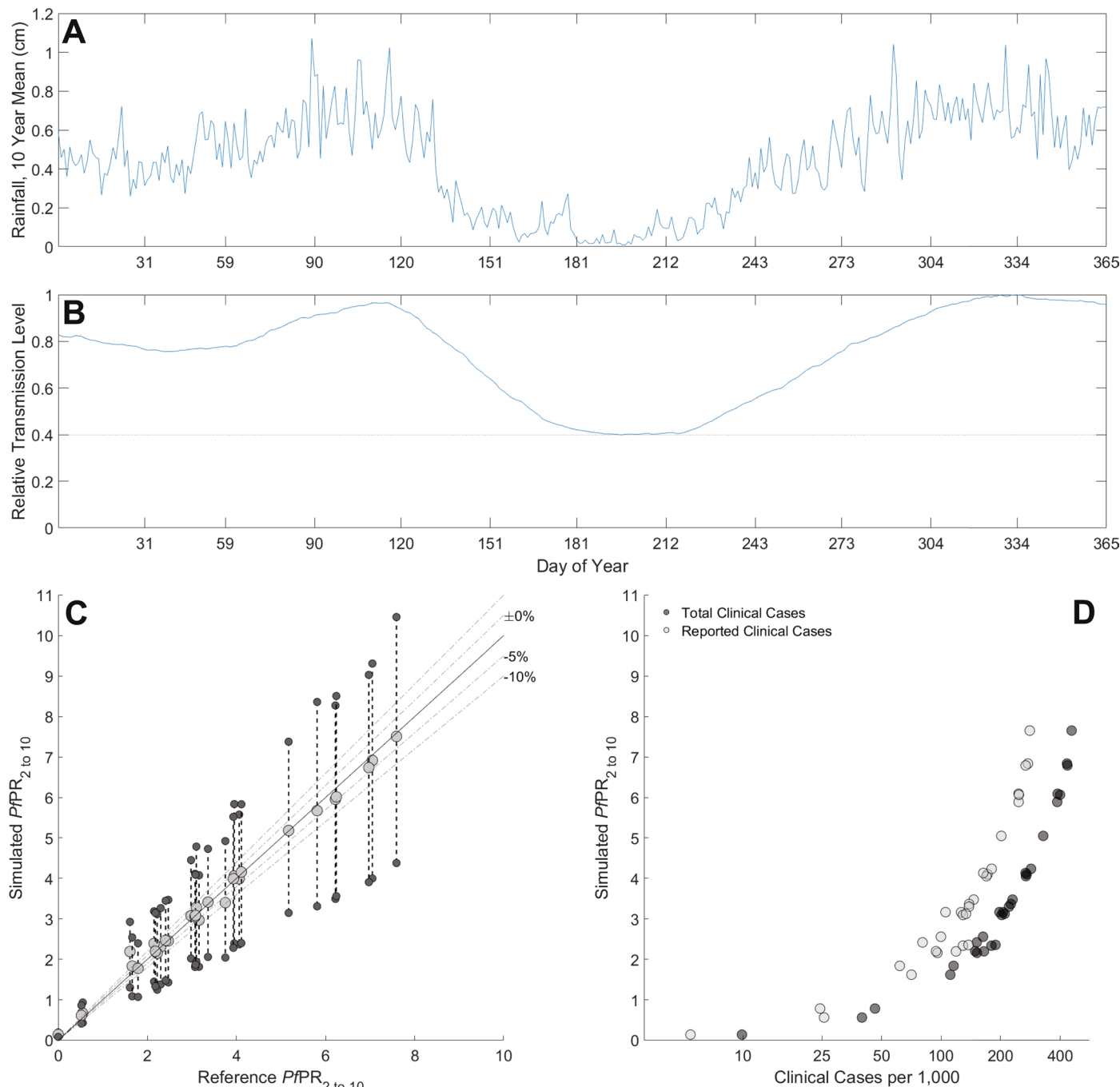


Extended Data Fig. 3 | See next page for caption.

Extended Data Fig. 3 | Individual genotype plots. Six heat maps showing the evolution of distinct genotypes for individual model runs. Each heat map shows genotype frequency, from 2018 to 2039, for all genotypes that rose above a 0.001 frequency at any point during this period. Frequencies below 0.001 are shown in white, whereas genotype frequencies above 0.001 are shown according to the colorbar on the right. The seven-letter genotype codes refer to, from left to right: the K76T locus, the N86Y locus, the Y184F locus, a second copy of N86Y, a second copy of Y184F, the R561H locus, and the presence ('2') or absence ('1') of a combination of piperazine-resistant alleles. The top two heat maps show two different model runs for the status quo – differing only in that the model's

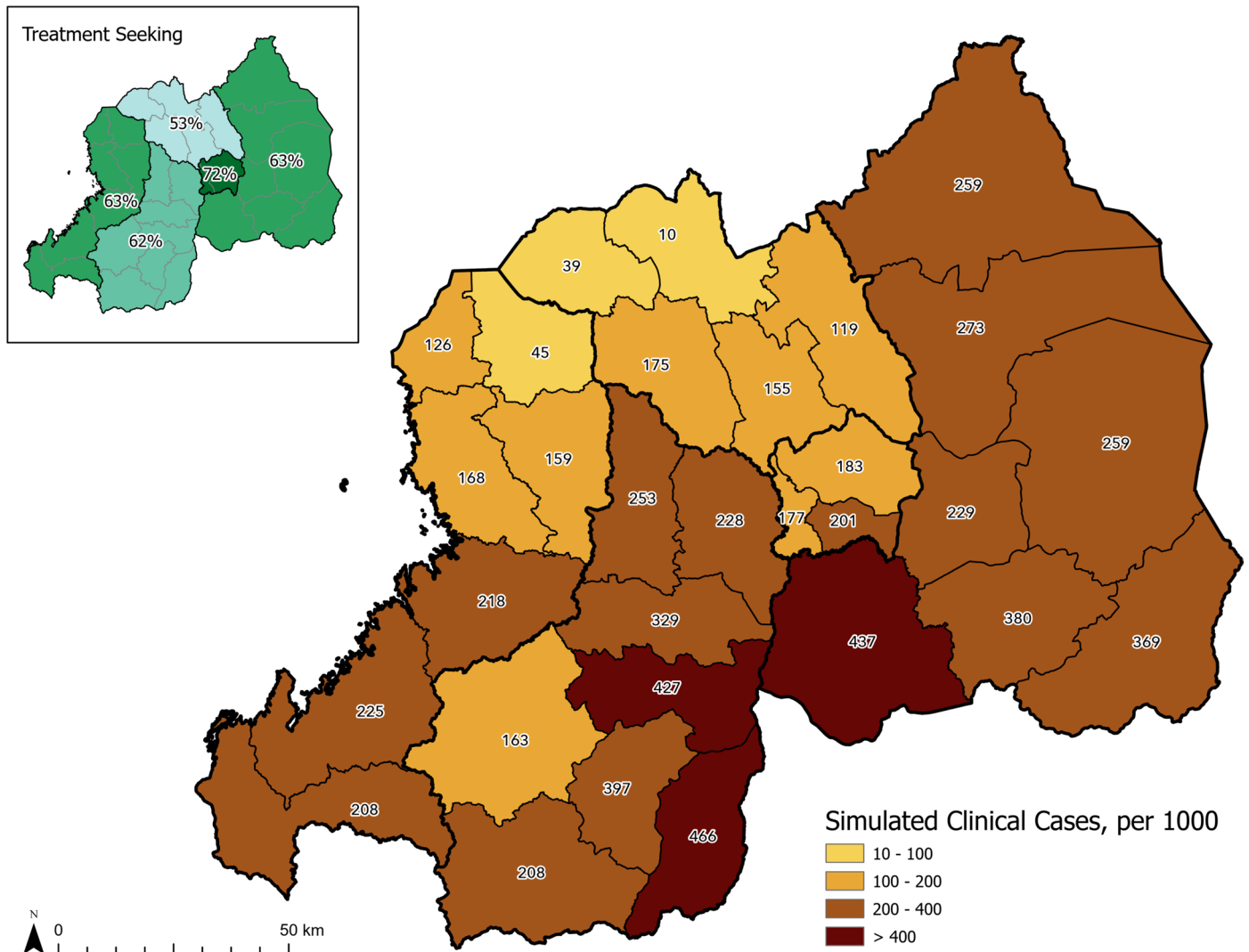
behavior is stochastic – where 3-day artemether-lumefantrine is maintained as first-line therapy. The outcomes of these runs differ, with different double-resistant (to AL) genotypes emerging and spreading. The four major AL-resistant genotypes that spread in these model runs are labeled with their efficacies in the heatmap (under a 3-day course of AL). These four double-resistant genotypes are labeled with asterisks in the four heatmaps at bottom, showing that some of the management strategies evaluated in this modeling exercise are successful at delaying the onset of AL double-resistance. For the four strategies shown at bottom, a typical model run was chosen to display. Note that under MFT, more genotypes emerge but selection pressure is weaker on each individual genotype.



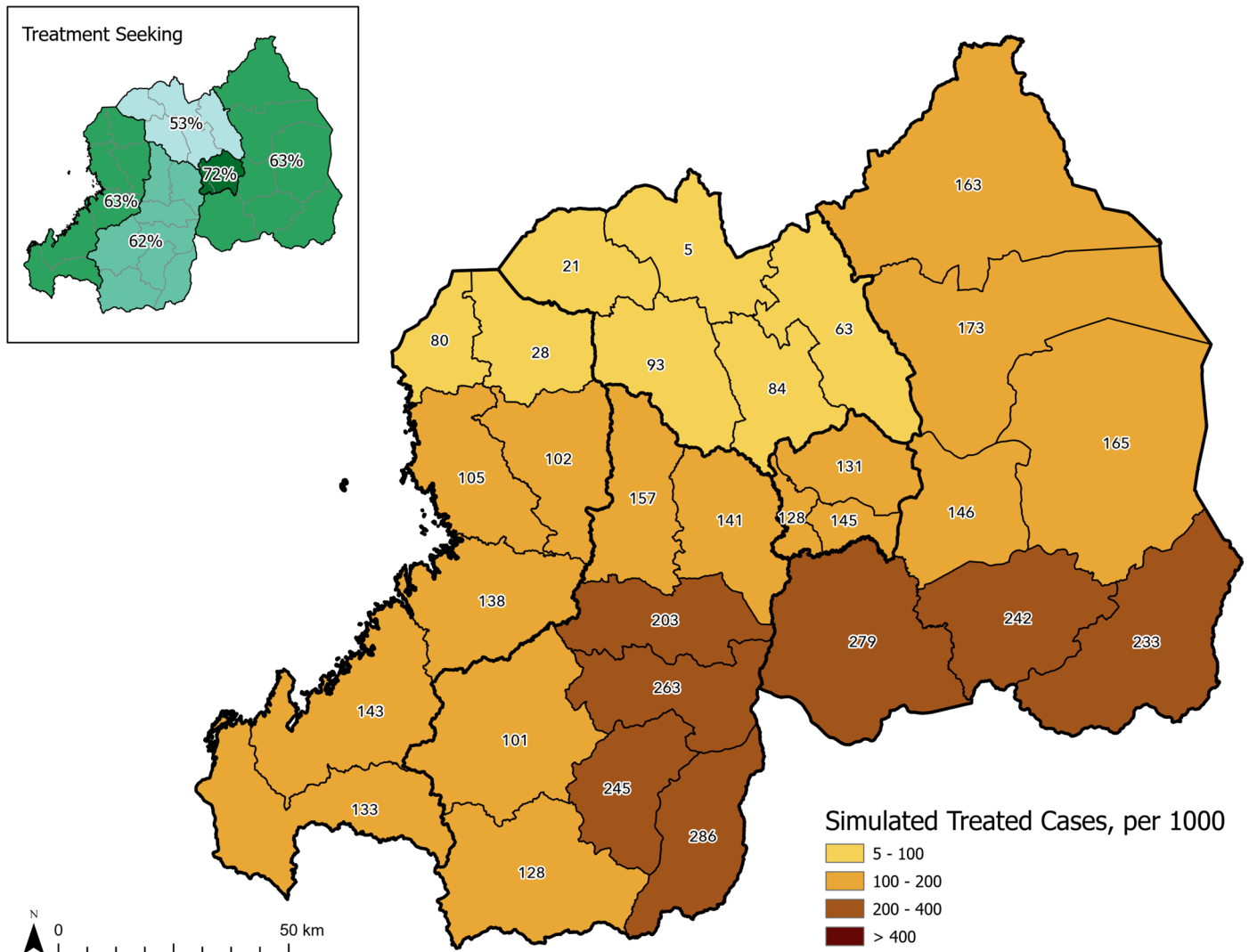


Extended Data Fig. 5 | Calibration of incidence and seasonality. (a) Mean daily rainfall over the period 2009–2019 reported from ERA5 global climate and weather projections⁶⁴. (b) Model’s relative transmission parameter (β) based on 54-day smoothing and normalization of rainfall data, note the lower bound of 0.4 during the low transmission season ensures some transmission is always present.

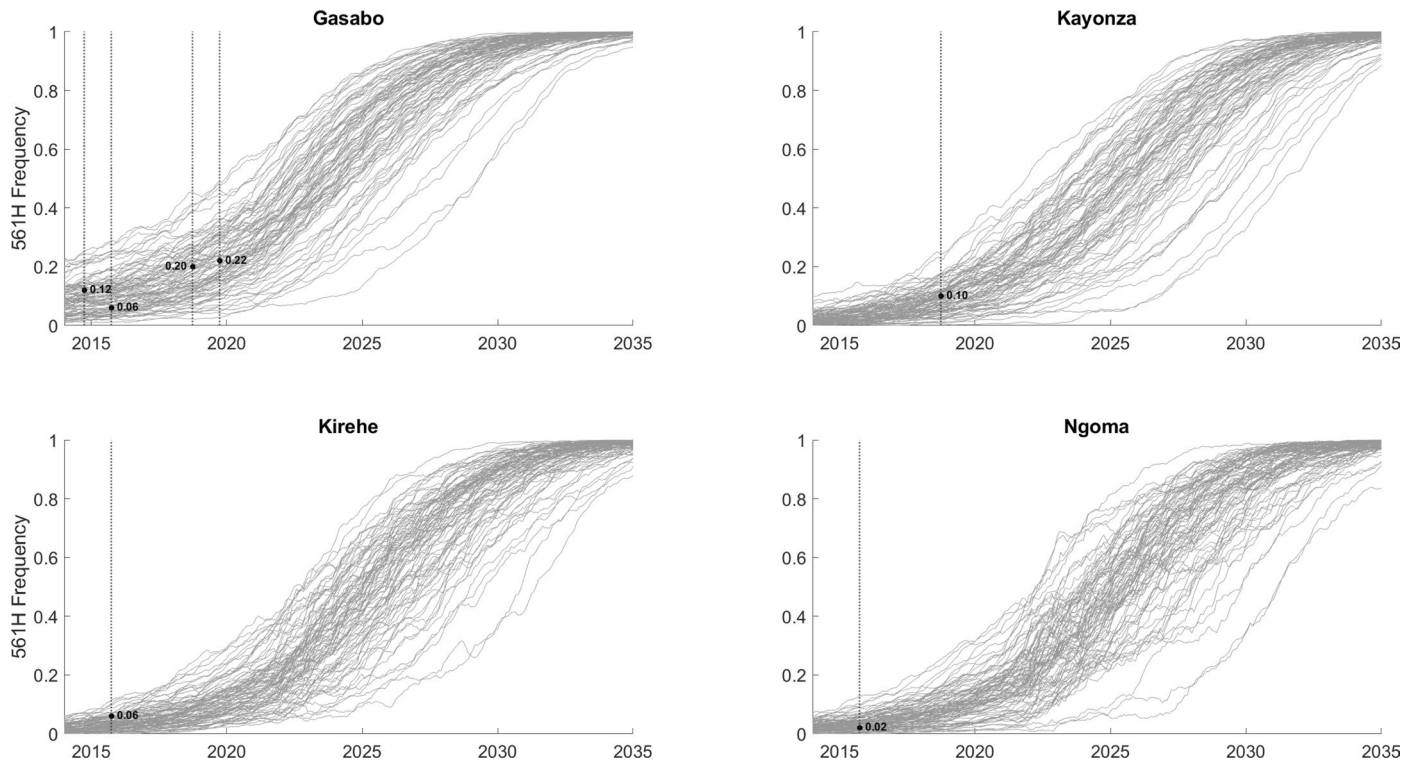
(c) Comparison of simulated $PfPR_{2 \text{ to } 10}$ to reference $PfPR_{2 \text{ to } 10}$ from the Malaria Atlas Project⁵⁰, with large gray circles showing the 12-month mean $PfPR_{2 \text{ to } 10}$ and smaller dark gray circles showing peak and trough $PfPR_{2 \text{ to } 10}$ during a single year. (d) Relationship between model $PfPR_{2 \text{ to } 10}$ and model’s malaria case counts (total and reported). Reported cases are total cases multiplied by the drug coverage.



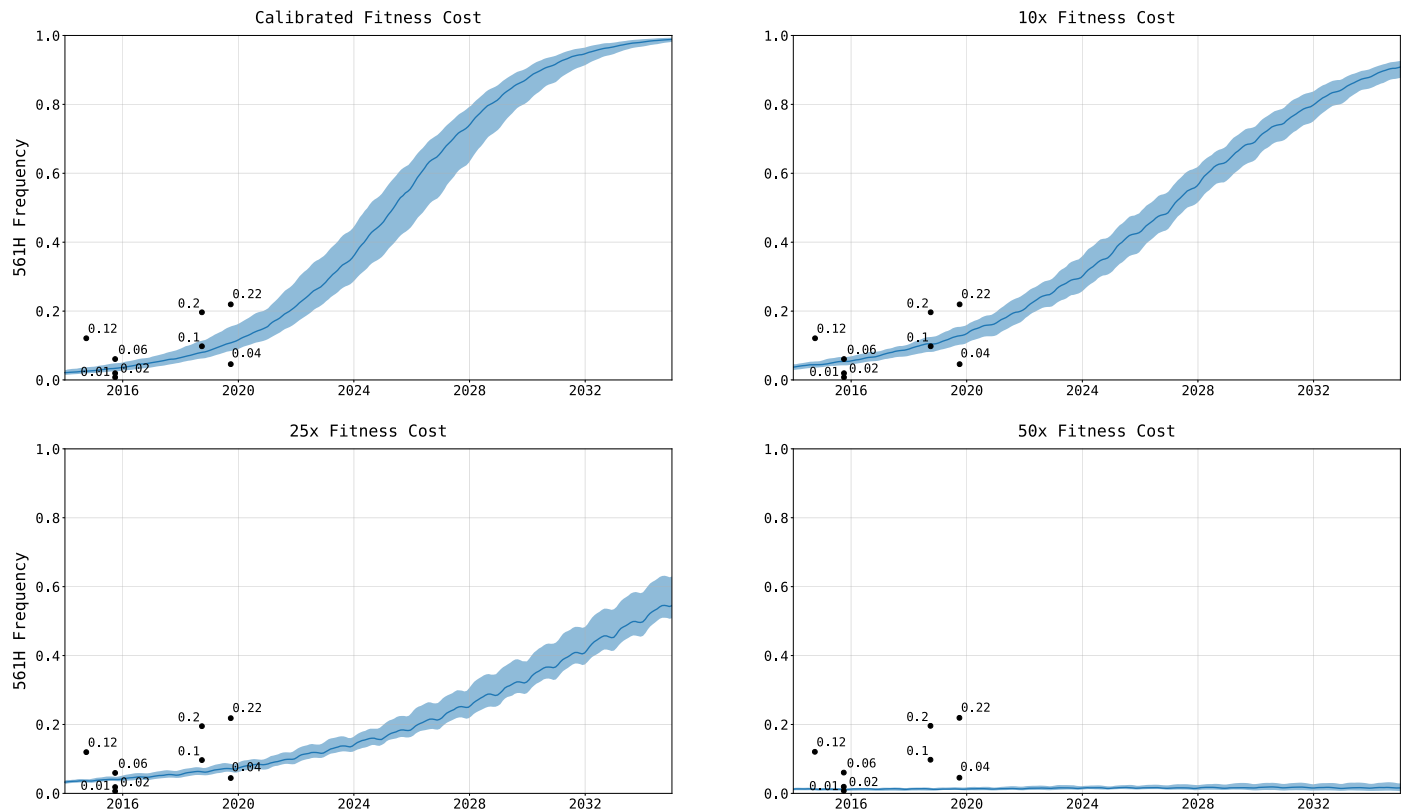
Extended Data Fig. 6 | Treatment seeking by province and model incidence by district. Map shows model incidence of *P.falciparum* cases in 2020, by district. Inset shows treatment coverage, from DHS data⁵¹, by province.



Extended Data Fig. 7 | Treatment seeking by province and treated clinical cases by district. Map shows model incidence of treated *P. falciparum* cases in 2020, by district. Inset shows treatment coverage, from DHS data⁵¹, by province.



Extended Data Fig. 8 | Individual trajectories of R561H evolution. Each panel shows model's frequency of 561H allele ($n = 100$ replicates) in a particular district or province of Rwanda: Gasabo province (Top Left), Kayonza district (Top Right), Ngoma district (Bottom Right), Kirehe district (Bottom Left).



Extended Data Fig. 9 | National 561H allele frequency under different assumptions on fitness cost of resistance. Model trajectories show median, national 561H allele frequency (shaded area is 95% range) under previously accepted fitness cost assumptions^{24,32,48,49} which result in a 17% within-host fitness cost per year. Sensitivity analysis increasing the daily fitness cost 10-fold (**Top**

Right), 25-fold (Bottom Left), and 50-fold (**Bottom Right)**** show that the fitness cost used in our simulations is consistent with allele frequency measurements (black dots) taken between 2014 and 2019. A 10-fold larger fitness cost is also compatible with the data post-2014, but it is unknown which of these values is compatible with the unseen emergence pattern from prior to 2014.**

Extended Data Table 1 | Effects of treatment compliance on treatment failure counts and percentages in 5 years

Therapy	Compliance Level (% of patients that complete treatment)	Median (IQR) Monthly Treatment Failure Count, in 5 years	Median (IQR) Treatment Failure Percentage, in 5 years
AL (3 days)	Full (100%)	23,400 (20,900 – 26,500)	18.0% (16.1% – 20.1%)
AL (3 days)	High (75%)	24,600 (21,700 – 27,300)	18.9% (16.8% – 20.9%)
	Moderate (65%)	25,000 (22,500 – 27,400)	19.2% (17.3% – 21.1%)
	Low (55%)	25,900 (23,600 – 29,900)	19.8% (18.1% – 22.6%)
AL (4 days)	High (70%)	19,500 (17,200 – 21,100)	15.2% (13.5% – 16.4%)
	Moderate (50%)	21,600 (20,600 – 23,900)	16.6% (16.0% – 18.2%)
	Low (25%)	22,800 (19,800 – 27,200)	17.6% (15.3% – 20.5%)
AL (5 days)	High (70%)	15,900 (14,300 – 17,420)	12.6% (11.4% – 13.6%)
	Moderate (50%)	17,400 (15,200 – 19,700)	13.6% (11.9% – 15.3%)
	Low (25%)	20,800 (16,800 – 23,000)	16.2% (13.3% – 17.6%)
ASAQ (3 days)	High (90%)	14,100 (12,600 – 15,300)	11.4% (10.2% – 12.3%)
	Moderate (80%)	14,600 (12,900 – 15,800)	11.7% (10.5% – 12.6%)
	Low (70%)	15,100 (13,700 – 16,200)	12.2% (11.1% – 13.0%)
DHA-PPQ (3 days)	High (90%)	65,200 (54,700 – 67,900)	43.5% (37.5% – 45.3%)
	Moderate (80%)	66,300 (62,100 – 69,900)	44.2% (41.8% – 46.1%)
	Low (70%)	64,600 (60,400 – 70,000)	43.3% (40.7% – 46.3%)

Reporting Summary

Nature Portfolio wishes to improve the reproducibility of the work that we publish. This form provides structure for consistency and transparency in reporting. For further information on Nature Portfolio policies, see our [Editorial Policies](#) and the [Editorial Policy Checklist](#).

Statistics

For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.

n/a Confirmed

- The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement
- A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
- The statistical test(s) used AND whether they are one- or two-sided
Only common tests should be described solely by name; describe more complex techniques in the Methods section.
- A description of all covariates tested
- A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons
- A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals)
- For null hypothesis testing, the test statistic (e.g. F , t , r) with confidence intervals, effect sizes, degrees of freedom and P value noted
Give P values as exact values whenever suitable.
- For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
- For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
- Estimates of effect sizes (e.g. Cohen's d , Pearson's r), indicating how they were calculated

Our web collection on [statistics for biologists](#) contains articles on many of the points above.

Software and code

Policy information about [availability of computer code](#)

Data collection

The simulation described (version 4.1.4) produced all data analyzed, which was aggregated in a Postgre (version 11.20) database, and downloaded into aggregated intermediate files using Python 3 (version 3.6.9). The intermediate files can be downloaded at <https://github.com/bonilab/malariaibm-spatial-Rwanda-561H/raw/main/Data/rwa-intermediate.zip>

Data analysis

The archived version the simulation code base (version 4.1.4), complied binaries for Linux, and scripts used for analysis and production of figures presented in this manuscript can be found on GitHub at <https://github.com/bonilab/malariaibm-spatial-Rwanda-561H>. ArcGIS Pro (version 3.1.2), MATLAB (R2021b, version 9.11.0.1809720 Update 1), and Python 3 (version 3.6.9) were used for data analysis and to prepare figures along with Adobe Illustrator 2023 (version 27.7) to combine and label figures.

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Portfolio [guidelines for submitting code & software](#) for further information.

Data

Policy information about [availability of data](#)

All manuscripts must include a [data availability statement](#). This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A description of any restrictions on data availability
- For clinical datasets or third party data, please ensure that the statement adheres to our [policy](#)

Intermediate data files produced by the simulation can be found on GitHub under <https://github.com/bonilab/malariaibm-spatial-Rwanda-561H/raw/main/Data/rwa-intermediate.zip> while the configuration files used for the study described in this manuscript can be found under <https://github.com/bonilab/malariaibm-spatial-Rwanda-561H/tree/main/Studies>. Global PfPR for children 2 to 10 years of age was used in calibrating the model, and the dataset was archived at <https://github.com/bonilab/malariaibm-spatial-Rwanda-561H/tree/main/Data/GIS/MAP> while the most recent data can be retrieved from https://data.malariaatlas.org/maps?layers=Malaria:202206_Global_Pf_Parasite_Rate. The spatial distribution of population is derived from the WorldPop 2015 spatial distribution of population in Rwanda, <https://hub.worldpop.org/doi/10.5258/SOTON/WP00674>

Human research participants

Policy information about [studies involving human research participants](#) and [Sex and Gender in Research](#).

Reporting on sex and gender

Sex and gender are not included as part of the simulation and model calibration for treatment seeking and movement is based upon the aggregated population data that has been reported in the literature.

Population characteristics

N/A

Recruitment

N/A

Ethics oversight

N/A

Note that full information on the approval of the study protocol must also be provided in the manuscript.

Field-specific reporting

Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

Life sciences Behavioural & social sciences Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see nature.com/documents/nr-reporting-summary-flat.pdf

Ecological, evolutionary & environmental sciences study design

All studies must disclose on these points even when the disclosure is negative.

Study description

A mathematical modeling analysis -- using a previously calibrated, stochastic, individual-based model -- to evaluate 26 possible options for Rwanda's National Malaria Control Program to respond to the recent emergence and frequency increase of the artemisinin-resistant R561H allele in Plasmodium falciparum.

Research sample

No independent data were collected for this study. The study presents future projections of R561H evolution under 26 different treatment scenarios over the next 10 years.

Sampling strategy

N/A

Data collection

N/A

Timing and spatial scale

Simulation/project approaches start by matching Rwanda's (1) prevalence data, (2) incidence data, and (3) R561H frequency data from 2014 to 2022. This is done on a 5km-by-5km spatial scale in Rwanda.

Data exclusions

N/A

Reproducibility

All calibrations and simulation results are fully reproducible, with source code and output files stored in a GitHub repository.

Randomization

N/A

Blinding

N/A

Did the study involve field work? Yes No

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems		Methods	
n/a	Involved in the study	n/a	Involved in the study
<input checked="" type="checkbox"/>	<input type="checkbox"/> Antibodies	<input checked="" type="checkbox"/>	<input type="checkbox"/> ChIP-seq
<input checked="" type="checkbox"/>	<input type="checkbox"/> Eukaryotic cell lines	<input checked="" type="checkbox"/>	<input type="checkbox"/> Flow cytometry
<input checked="" type="checkbox"/>	<input type="checkbox"/> Palaeontology and archaeology	<input checked="" type="checkbox"/>	<input type="checkbox"/> MRI-based neuroimaging
<input checked="" type="checkbox"/>	<input type="checkbox"/> Animals and other organisms		
<input checked="" type="checkbox"/>	<input type="checkbox"/> Clinical data		
<input checked="" type="checkbox"/>	<input type="checkbox"/> Dual use research of concern		