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Future malaria environmental suitability in Africa is sensitive to hydrology

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Abstract: Changes in climate shift the geographic locations that are suitable for malaria transmission because of the thermal constraints on vector *Anopheles* mosquitos and *Plasmodium spp.* malaria parasites and the lack of availability of surface water for vector breeding. Previous Africa-wide assessments have tended to solely represent surface water using precipitation, ignoring many important hydrological processes. Here, we applied a validated and weighted ensemble of global hydrological and climate models to estimate present and future areas of

hydroclimatic suitability for malaria transmission. With explicit surface water representation, we predict a net decrease in areas suitable for malaria transmission from 2025 onward, greater sensitivity to future greenhouse gas emissions, and different, more complex, malaria transmission patterns. Areas of malaria transmission that are projected to change are smaller than those estimated by precipitation-based estimates but are associated with greater changes in transmission season lengths.

Main Text:

Malaria is a climate-sensitive, vectorborne disease that caused 619,000 deaths among 247 million cases in 2021 (1). This burden is acutely focused on Africa, where 95% of global cases are reported (1). Reductions in cases in Africa have slowed or even reversed in recent years, attributed in part to a stall in investments in global responses to malaria control. To build resilient health systems, we must understand present-day malaria transmission and potential future areas of malaria risk (2). Malaria transmission is a function of many complex and interacting parameters (3), including strong environmental controls. Given these interactions and in the face of rapidly changing climate in Africa, it is important not only to model the impact of these controls on changing malaria transmission hazard per se, but also to improve the climate process representation. Doing so will facilitate future integration with estimated changes in other key sectors (e.g., agriculture) to identify intersecting impacts and potential trade-offs (4–6).

Ambient air temperature influences sporogonic and gonotrophic development rates, biting rates, and vector longevity (7). Persistent surface water is required for vector breeding sites that remains at an appropriate temperature for larval development (8–10). Thermal response models for malaria are relatively well understood (11) and readily implemented within studies examining future climate-driven shifts in disease transmission suitability (12). By contrast, the controls on surface water availability for vector breeding are either ignored or poorly represented within climate-driven geospatial models of transmission suitability.

Assessments at regional or village scales typically represent detailed hydrological processes (8, 13), and they are increasingly incorporated into malaria transmission models (14, 15). However, continental and global scale analyses lack detailed parameterization of hydrological processes and conventionally apply a rainfall threshold as a proxy for water body

availability (typically 60 to 80 mm month⁻¹ , although a wide range has been observed) (9, 16–19). Such a simple threshold bypasses many important hydrological processes. Rainfall is not a good indicator of standing water availability (12), and the resulting estimates for malaria occurrence are sensitive to the specific choice of precipitation threshold (20). Initial findings suggest that process-based representations of breeding habitat availability result in a more complex pattern of areas estimated to be suitable for malaria transmission across Africa, because infiltration, evaporation, runoff, and river discharge are directly represented (20).

Here, we address the challenge of incorporating robust estimates of vector breeding site availability into continental-scale estimates of areas of suitability for malaria transmission. The application of a single hydrological model presented in our earlier work (20) is problematic because diverse different hydrological models are available (21), each reflecting the perceptual models and focuses of their creators (22). This variability means that the outputs (e.g., surface water) of one model can be quite different from another (23), and performance (i.e., simulation compared with observation) can vary considerably (24). Recognizing these limitations, we present a multimodel, multi- scenario ensemble of global hydrological and global climate models (25) weighted based on model performance to estimate present and future distribution of malaria suitable areas across Africa. We examine the sensitivity of malaria distribution estimates to different hydrological models and evaluate multiple hydrologically informed metrics to represent breeding site availability. Critically, the inclusion of hydrological modeling identifies a different pattern of future shifts in the distribution of malaria suitable areas to a precipitation-based estimate. Finally, we quantify the human populations affected by changing lengths of malaria transmission season as future climate changes alter hydroclimatic suitability for malaria transmission.

Variability in malaria suitability estimates

We calculated the distribution of malaria suitable areas for 1986 to 2005 using hydrological outputs from seven global hydrological models (GHMs), which were each driven with climate data from four global climate models (GCMs). When a precipitation threshold is used to represent surface water availability, estimates of areas suitable for malaria transmission across the GCMs show very limited variability (Fig. 1B and fig. S1), as shown by the strong agreement among estimates. By contrast, explicit representation of hydrological processes using the GHMs shows greater variability in projections for areas of malaria transmission, reflecting different structures among hydrological models. Although total extents of hydroclimatic suitability [i.e., the length of transmission season (LTS) >1 month] were similar across the GHMs (fig. S2), estimates from the ORCHIDEE model ([https:// orchidee.ipsl.fr/](https://orchidee.ipsl.fr/)) show particularly large areas of endemic malaria (>9 months of continuous transmission suitability), whereas the LPJml (Lund-Potsdam-Jena managed land, [https:// www.pik-potsdam.de/en/institute/departments/ activities/biosphere-water-modelling/lpjml](https://www.pik-potsdam.de/en/institute/departments/activities/biosphere-water-modelling/lpjml)) model showed the least extent of endemic malaria.

Continental-scale malaria distribution patterns are broadly similar for each hydrological representation that we examined, although hydrology-based estimates are patchier than precipitation-based estimates. However, hydrological estimates are more physically realistic and identify larger areas with year-round malaria (Fig. 1). Large-scale river networks are highlighted as local foci of endemic malaria, particularly the Nile. With human population centers typically located around large waterways, these differences are more pronounced when considering the number of people residing in potentially malaria endemic areas (29 million people at risk using precipitation based estimates compared with 107 million people at risk when discharge is used).

Although malaria transmission may not be observed in all these locations today thanks to successful interventions, they nevertheless remain hydro-climatically suitable for transmission.

To represent the likelihood of temporary rain-fed puddles forming while also incorporating water fluxes within continental-scale river networks, the potential evapotranspiration to precipitation ratio was calculated and hydrological suitability computed. This ratio provides a basic surface water balance and was added to the discharge suitability layer (Q_PE). Including the local surface runoff expands the area further with larger areas suitable for >3 months per year. Conversely, the additional criteria of a minimum relative humidity threshold restricts this suitable area, notably at the northern extent, particularly Chad, Mali, and Sudan. However, active transmission in eastern Sudan has been observed in proximity to rivers even in hot, dry seasons (26).

Calculation of variance ratios showed that GHMs are the primary influence, with GCMs contributing only 2% of variance in endemic malaria areas, 7% for areas of seasonal malaria, and 16% for all areas with >1 month suitability for the Q_PE hydrological representation. For discharge alone, the values are lower (all 2% or less), but when including the additional relative humidity threshold, the values are slightly higher because of the additional climate dependency (Fig. 1G). When examining sources of variance in LTS estimates for each grid cell, Fig. 1G highlights the overall dominance of GHMs, although GCMs are locally important in the Ethiopian highlands and other high altitude areas of East Africa, in areas at the northern fringe of transmission suitability and an isolated area in Mozambique.

Ensemble weighting and model performance

Validation of estimates of hydroclimatic suitability is problematic because malaria transmission is driven by nonclimatic factors, notably successful interventions reducing the geographic spread of transmission (3). We also recognize that sub-Saharan vector species vary in their larval habitat requirements (27) and often use small, shallow, sunlit, and water filled depressions such as hoof prints, wheel ruts, and puddles for breeding sites (e.g., *Anopheles gambiae sensu stricto* and *A. arabiensis*), although even here habitat hydrology and geomorphology have been shown to control the distribution of malaria vector larva (28). Conversely, other major vectors (e.g., *A. funestus*) require semipermanent freshwater bodies such as swamps, ponds, and lake edges (27), and secondary vectors such as *A. coustani* and *A. squamosus* breeding in large, vegetated floodplains connected to rivers can be locally and regionally important (29, 30). Local variability in vector ecology presents a challenge to any geographically uniform approach to modeling malaria suitability (31), although the inclusion of both local runoff and large-scale water fluxes in the composite hydrological metric (Q_PE) is an attempt to ensure that both habitat types are represented.

We validated malaria suitability estimates for the preindustrial period 1875 to 1900 against the preintervention map of Lysenko and Semashko (32) (Fig. 2A), which approximates malaria extents at ~1900 AD. The performance of each model is used to allocate a model weight in an ensemble for future predictions of extent (see the materials and methods, figs. S3 to S5, and tables S2 to S5 for details).

Ensemble estimates are shown in Fig. 2B and figs. S6 and S7, with estimates compared against observations in Fig. 2F and fig. S8. Hydrology-based estimates (discharge with potential evapotranspiration) performed only very slightly better than estimates made with precipitation alone (ensemble score of 0.817 versus 0.816), and a relative humidity filter reduced this performance due to underestimation in the northern fringes of transmission. All models are

slightly biased toward underprediction (negative bias scores in Fig. 2C). The spatial extent of preindustrial malaria suitability estimates is broadly similar to those of the historical period presented in Fig. 1, with the more substantial increases in suitability observed at high elevations in East Africa focused on Ethiopia and Uganda (fig. S3).

5 Overall, validation shows a close match between modeled and observed suitability. All models overestimate the extent of areas suitable for mosquito breeding in South Africa, but the northern fringes of the transmission zone are correctly represented, with only minor underestimates (Fig. 2F). Similarly, areas suitable for malaria transmission in the Horn of Africa are underestimated in all models. All models underestimate the potential for malaria transmission along the North
10 African coast, although here hydrology-based estimates outperform the precipitation-based estimates. The presence of different malaria vector species in North Africa with different thermal ranges from the typical sub-Saharan species (33) may explain this discrepancy. Some overestimation is evident in the Sahara for the hydrological model output (without the relative humidity filter), where the course of the now dry Tamanrasset palaeoriver (34) is identified as
15 suitable for malaria transmission for individual months. However, the Nile corridor of malaria risk in Egypt, where malaria was endemic until the 1990s, is only identifiable in the models that describe discharge and discharge with potential evapotranspiration (35). The relative humidity threshold applied at this grid scale appears to be overly restrictive and cannot represent the fine-scale local influence of water bodies on air humidity. Overall, there is no strong evidence to
20 distinguish between the performance of precipitation and discharge with potential evapotranspiration representations (Fig. 2D). Further validation of weighted layers for the 1986 to 2005 time period against anopheline observations (36), parasite ratios (37), and modeled probabilities of occurrence of *A. gambiae* (38) (fig. S9 and table S6) confirm that both

precipitation and discharge with potential evapotranspiration representations perform equally well. Thus, future suitability estimates from both are considered in Fig. 3.

Future projections of declining malaria risk

For the representative concentration pathway (RCP) 2.6 and 6.0 model ensembles, there is agreement that a modest initial increase in areas of hydroclimatic suitability for malaria transmission (2006 to 2025; 0.5, both hydrological and precipitation based estimates agree that the overall hydroclimatic suitability for malaria transmission decreases from the baseline period (Fig. 3). The decreasing area suitable for malaria transmission becomes more pronounced through time and for higher greenhouse gas emissions scenarios (RCP 6.0 and RCP 8.5), although it should be noted that hydrological estimates show greater sensitivity to the choice of RCP than do precipitation-based malaria suitability estimates (compare Fig. 3, B and F).

Precipitation-based estimates show a larger shift in the geographical locations of malaria transmission suitability, with large areas (3 to 4 Mn km²) experiencing increasing malaria risk in all RCPs and time periods (mostly greater than twice that of the hydrology-based estimates; table S9) that are offset by growing areas of decreasing suitability for malaria transmission (>2 million km²). By contrast, hydrology-based estimates show smaller areas increasing in suitability for malaria and a greater sensitivity to the RCP, particularly for the areas diminishing in malaria suitability (Fig. 3). Precipitation-based estimated changes are mostly small (a reduction in the LTS by 1 month; Fig. 3D); conversely, hydrology-based estimates show substantial areas where suitability for malaria transmission decreases by 4 months or more (Fig. 3H and fig. S12). Thus, projected changes using hydrology-based estimates are more concentrated but also more intense when compared with precipitation based estimates. Calculation of the overall magnitude of the

change (i.e., the product of the change in LTS in months and the area) shows that the reduction in suitability is more pronounced in the hydrology-based estimates, but only for RCP 8.5, where it is more than double the precipitation-based estimates by 2076 to 2100 (fig. S10 and table S10).

The pattern of projected changes in malaria distribution is markedly different between precipitation- and hydrology-based estimates (Fig. 3, C and G, and fig. S11). Although both approaches identify future declines in malaria transmission across West Africa, the area experiencing a reduction in LTS is more extensive with hydrology-based estimates, stretching as far east as South Sudan. Areas of malaria risk in the highlands of Ethiopia are expected to expand [although to a lesser extent than modeled previously (39)], but the increase in projected malaria prevalences for Kenya, Uganda, and the north of the Democratic Republic of the Congo estimated by precipitation-based models are not seen for any RCP. Small decreases in the LTS of <1 month across Tanzania and Zambia are also not observed with any hydrology-based estimates, but for RCP 8.5, a stronger focus of decreasing risk emerges further south in Mozambique, Zimbabwe, Botswana, and Namibia. Finally, the small area of increasing malaria risk centered on Lesotho is seen in all estimates, but also stretches west along waterways [specifically the Orange River, where flood-driven malaria has been observed historically (40)] when fluvial water transfer is represented in the hydrology-based estimates.

At the country level, inclusion of hydrological processes in malaria distribution models results in larger modeled decreases in LTS compared with precipitation-based estimates in those countries where the projected changes are focused (particularly West Africa and the northern fringes of malaria transmission suitability; fig. S13). Future estimated populations at risk show considerable growth in populations residing in areas with longer estimated periods of malaria transmission suitability (Fig. 4). The difference between precipitation- and hydrology-based estimates is particularly pronounced for endemic areas with LTS >9 months, with the

populations affected being four times greater when hydrology is represented. Again, this highlights the combination of waterways as centers for human population growth and foci for malaria risk (figs. S14 to S16 and table S11).

Discussion

Changes in malaria transmission are complex and not driven by climate alone. However, the effect of climate change on malaria prevalence is pronounced, particularly in Africa (2). Many models have been used to estimate future transmission risk using simple precipitation based approaches, but positive correlations between malaria and rainfall are not consistent (41). We present a multimodel ensemble of hydrological and global climate models to estimate changes in malaria transmission risk, which reveals substantial variability in estimates from different hydrological models and the value of an ensemble approach in which models are evaluated on the basis of historical performance (42, 43). GHM ensembles are now routinely used within climate change impact assessments (e.g., 5, 23, 44–46) but have not previously been applied to represent malaria transmission.

Because river systems are directly represented in a hydrological approach, the largest differences between precipitation- and hydrology-based malaria estimates are observed near rivers and floodplains, which are also the largest population foci. Therefore, hydrology based approaches indicate that the number of people residing in potentially malaria endemic areas is almost four times that estimated with precipitation-based estimates. There is also a greater sensitivity of malaria outcomes to the choice of RCP when hydrology is simulated explicitly. There is general agreement that malaria transmission will decrease across much of West Africa, as also predicted in previous studies (e.g., 13, 18, 20, 47), although when the hydrological processes that determine viable vector habitat formation are modeled explicitly, estimates for

transmission decrease more extensively. The warming and drying trends underlying this decrease in areas suitable for malaria themselves present profound environmental and social challenges (48), including other predictions that this trend will increase dengue suitability (49).

Embedding GHM estimates of hydrological suitability for vector breeding directly into process-based malaria models such as VECTRI and LMM (14, 15) will permit a more direct evaluation of transmission intensity in the future. Differences in mosquito ecology and breeding habitat preferences across Africa are not presently included in the model. Representing the regional variability in mosquito vectors while also partitioning modeled surface water bodies by mosquito species breeding site preferences would more directly represent this important source of variability. However, this information is currently beyond the capabilities of GHM ensembles and would require a more hydrodynamic approach. Although more detailed hydrological treatments have already been tested at regional or village scales (8), the spatial resolution of available data currently limits our ability to model individual water bodies explicitly at the continental scale or to identify specific *Anopheles* vector niches.

The continental-scale evaluation of malaria climate change impacts using the GHM ensemble presented here imposes the relatively coarse half-degree grids that are presently the main limitation of this approach. Finer details of water body dynamics such as the flushing of habitat require higher-resolution models of individual floodplains. Situations in which drought and falling river levels causes flow disconnection and the formation of isolated stream-bed water bodies that function as suitable breeding habitats (9) are not well captured by our current approach. As these data sources become increasingly available, we will benefit from their explicit incorporation in projections of hydrological processes to explain physically realistic malaria transmission risk at scales that can inform national operational malaria control strategies

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Study design and methods: MWS with support from SNG

Data processing: MWS, TW and W.H.M.J.

Data analysis: MWS, EM, MJK and WHMJ

Writing initial draft: MWS

Writing review and editing: all authors.

5 **Competing interests:** Authors declare that they have no competing interests.

Data and materials availability: Global Climate Model data (40) and Global Hydrological

Model data (50) used in this study are available via the ISIMIP Repository. Population
grids (total and age-structured) are available to download from the WorldPop dataset

10 (51). UN country level population estimates and projections (total and age-structured) are
available to download from the United Nations, Department of Economic and Social
Affairs (52). Malaria suitability estimates from this paper and source data for figures are
available for download from the Research Data Leeds Repository (53).

Supplementary Materials

15 Materials and Methods

Figs. S1 – S16

Tables S1 – S11

References (54 – 78)

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Figures

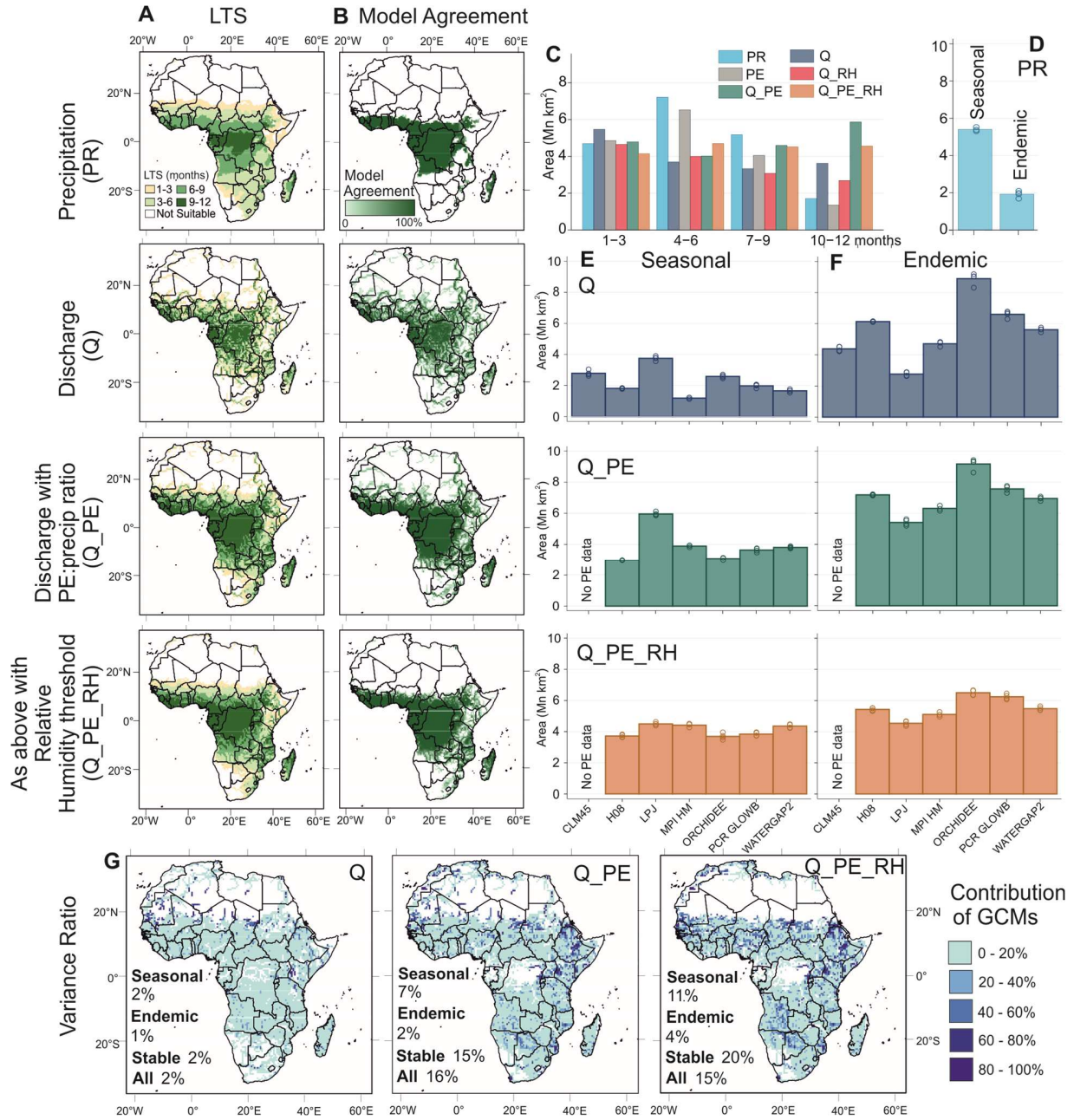


Fig. 1. Variability in malaria hydro-climatic suitability estimates (1986-2005). (A) Mean

Length of Transmission Season (LTS, in months) estimates for each hydrological representation

(mean of all GCM and GHM combinations). (B) Model agreement across each ensemble. Maps

indicate the percentage of models in the ensemble that estimate LTS > 6 months (full data presented in Fig. S1 and Table S1). (C) LTS estimates for each hydrological representation calculated within 3-month bands using the model ensemble weightings established in later validation. (D-F) Estimates of seasonal (LTS > 6 months) and endemic malaria suitability (LTS > 9 months) showing all individual model estimates across different hydrological models (bars showing mean value) and global climate models (individual points for each climate model, $n = 4$). Precipitation is plotted separately in (D). (G) Variance ratios for all hydrology-derived layers showing the percentage of variance in malaria LTS contributed by GCMs for each model cell.

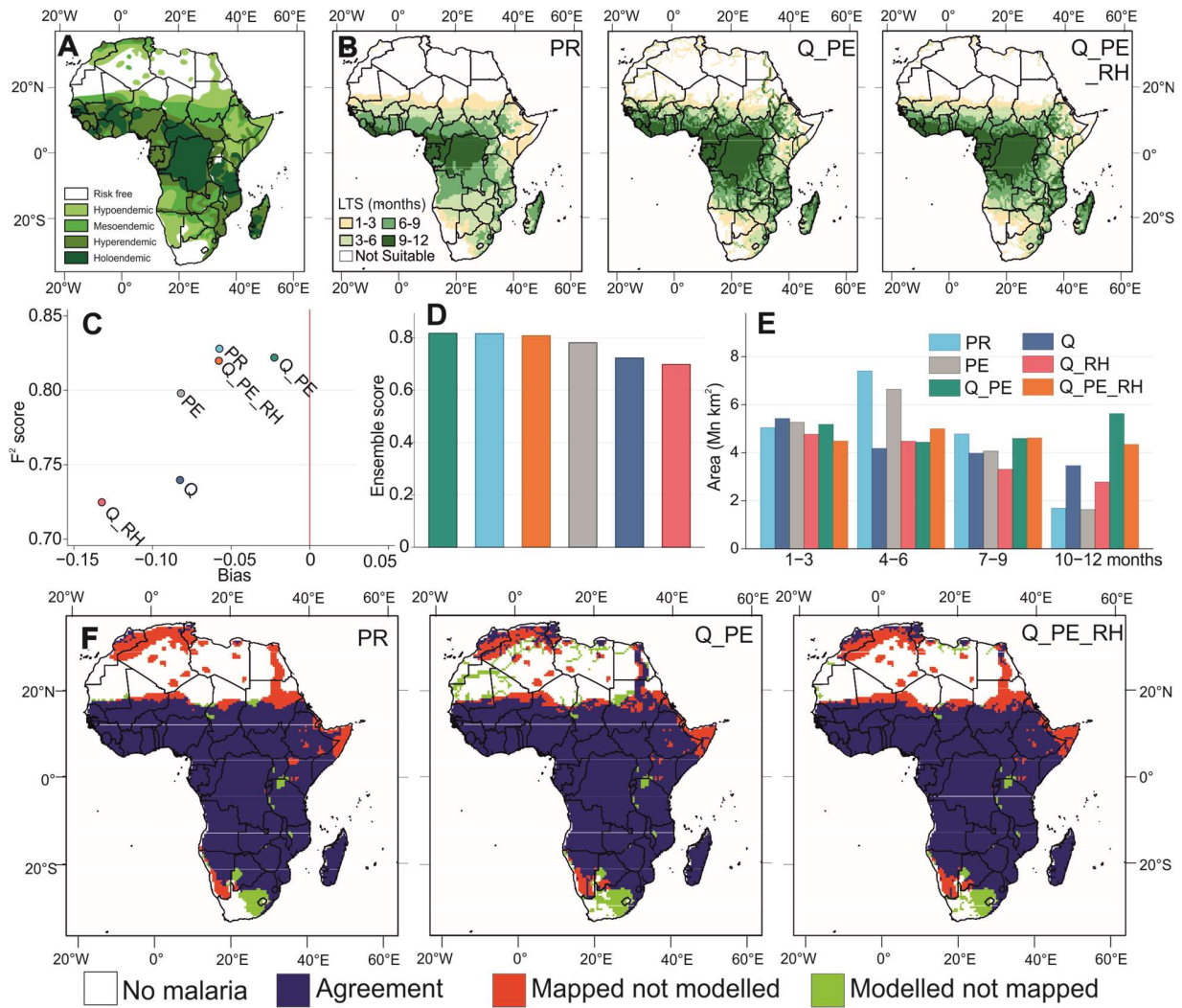


Fig. 2. Validation of model hydroclimatic malaria suitability estimates. (A) Pre-intervention malaria map of Lysenko and Semashko (32); (B) weighted ensemble model estimates of hydroclimatic malaria suitability LTS (1875-1900); (C-D) summary of ensemble validation metrics for each hydrological representation, with both metrics plotted separately and combined as an ensemble score (full data in Tables S2-S3); (E) estimates of malaria suitability by the weighted ensembles for each hydrological representation calculated within 3-month bands; (F) selected validation maps comparing observed and modelled malaria suitability (where at least one month is suitable). Figs. S6-S9 present validation maps and LTS maps for all hydrological representations.

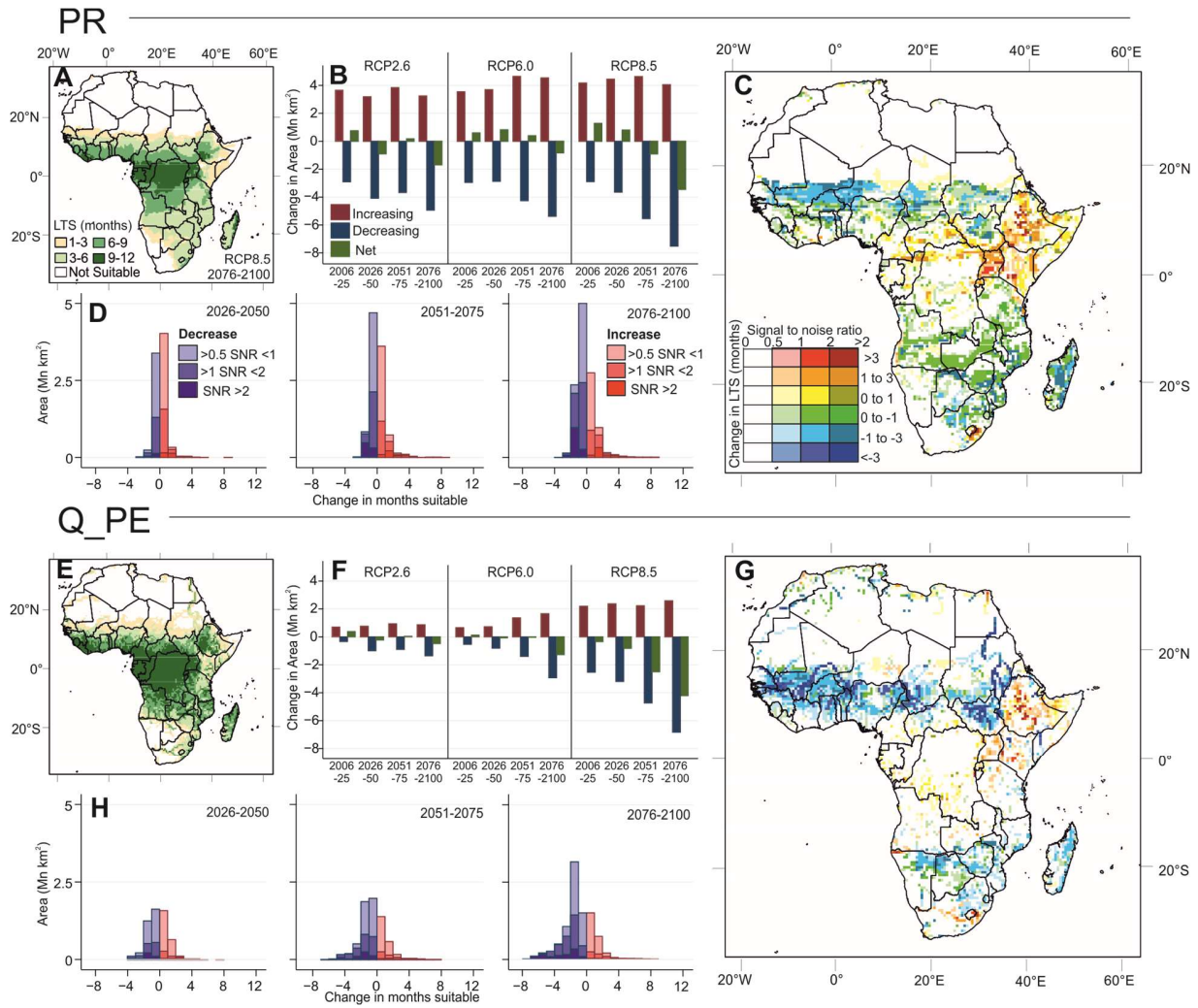


Fig. 3. Future projections of malaria hydro-climatic suitability. (A, E) Projected Length of Transmission Season in RCP 8.5 for 2076-2100. Areas showing an increase or decrease in suitability since 1986-2005 across all scenarios and time periods are plotted in (B, F) showing only areas with a signal-to-noise ratio (SNR) > 0.5. The locations of these changing areas are shown for RCP 8.5 (2076-2100) in (C, G) and for all RCP scenarios and time periods in Fig. S11. The extent of change in suitability (number of months change) is plotted by signal-to-noise ratio as stacked histograms for RCP 8.5 in (D, H) and for all scenarios in Fig. S12.

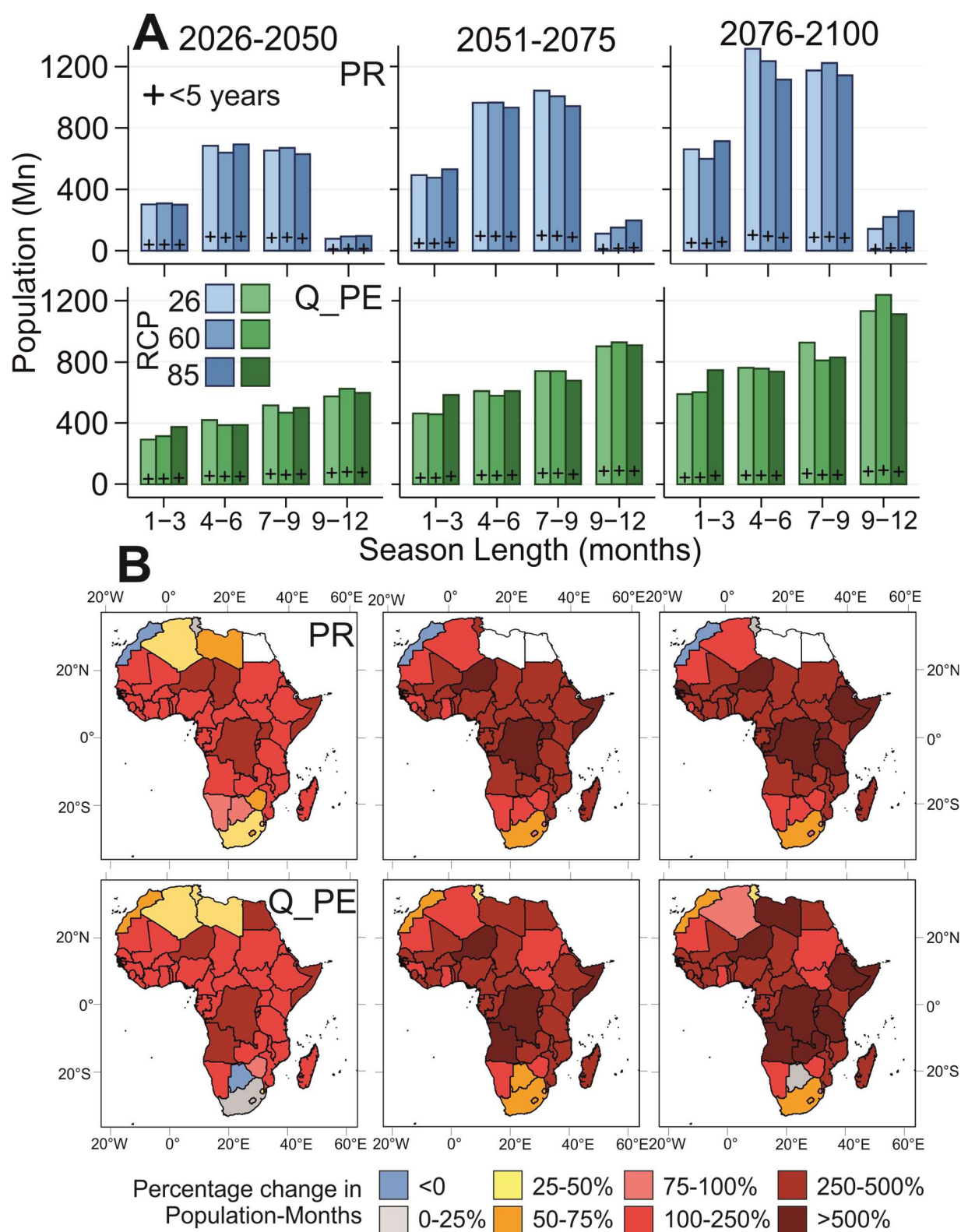


Fig. 4. Future populations in areas hydro-climatically suitable for malaria transmission.

(A) Populations within 3-month bands of LTS for both precipitation and discharge with potential evapotranspiration hydrological representations. Bars show total populations for each RCP for each weighted ensemble. Points indicate population under-5 years old. Populations calculated using UN medium variant projections for each period. (B) Predicted percentage changes in population-months (i.e., population exposed multiplied by the number of additional months of exposure) since 1986-2005 by country for RCP 8.5 only (see Fig. S13 for each RCP). White indicates no estimated malaria suitability.

Materials and Methods

Climate data

Climate data were downloaded for four Global Climate Models (GCMs) (GFDL-ESM2M, HadGem2-ES, IPSL-CM5A-LR and MIROC-ESM-CHEM) for three RCPs (RCP 2.6, RCP 6.0 and RCP 8.5) chosen to represent a range of radiative forcings and uncertainty regarding future greenhouse gas emissions. These data were retrieved from the database of the Inter-Sectoral Impact Model Intercomparison Project (ISIMIP2b) (25, 54). All climate variables were obtained on a latitude-longitude grid of 0.5 x 0.5 degrees covering the extent of Africa. Climate variables comprised of bias-adjusted (25, 55) daily mean surface temperature (in K), daily total precipitation (in $\text{kg m}^{-2} \text{s}^{-1}$), and daily relative humidity (in %). Historical daily climate data were downloaded for the period 1986-2005; future climate simulations for each RCP were split into four periods: 2006-2025 (to represent present conditions), 2026-2050, 2051-2075 and 2076-2100.

Hydrological modelling

Climate data from each combination of GCM and RCP were used to drive seven global hydrological models on a daily time step. Global hydrological models included were CLM45 (56), H08 (57), LPJml (58), MPI-HM (59), ORCHIDEE (60), PCR-GLOBW (61) and WaterGAP2-2c (62). The GHMs represent an ensemble of opportunity and while some models use similar methods to represent some hydrological processes, there are many differences in the ways that the models parameterize others (21). The differences in LTS between LPJmL and ORCHIDEE (Fig. S2) reflect these model differences. The different approaches for calculating potential evapotranspiration between GHMs often explains some of the differences between the ISIMIP2b GHM simulations of runoff (63, 64), for example, ORCHIDEE applies a simplified Penman–Monteith equation whereas LPJmL applies the Priestley–Taylor equation (21). While historical society scenarios were used to model human influences on hydrology (e.g., land use and irrigation) for the preindustrial and historical model runs (25), 2005 society models were used for all future estimates to isolate the climate change signal. Daily modelled discharge ($\text{m}^3 \text{s}^{-1}$) and monthly modelled potential evapotranspiration (both in $\text{kg m}^{-2} \text{s}^{-1}$) output from GHMs were used as variables to represent hydrological malaria suitability. These data were available via the ISIMIP2b database.

Estimation of malaria hydrological suitability

Average daily surface temperature, relative humidity, discharge, potential evapotranspiration and total precipitation were calculated for each month. For each combination of GCM and GHM, six methods of estimating hydrological suitability for malaria transmission were calculated based on monthly values of climate and hydrological parameters averaged across each of the above time periods.

(i) Precipitation (PR): the 60 mm monthly precipitation threshold of Tanser et al. (65) was implemented, including the requirement of a catalyst 80 mm month (i.e., a single month where precipitation exceeds 80 mm).

(ii) Discharge (Q): a minimum of 1 mm depth across the grid cell was used as a threshold for daily hydrological model discharge output, indicating sufficient water availability (equating to approximately $30 \text{ m}^3 \text{s}^{-1}$). Previous sensitivity analysis (20) indicated limited sensitivity of hydro-climatic malaria suitability estimates to this value, particularly when compared to the sensitivity of precipitation thresholds. The number of days this condition is met per month was

then averaged for each time period; when this exceeded the temperature-dependent development period of *Anopheles* mosquitoes (calculated using the model of Bayoh and Lindsay (10)), the month was deemed to be hydrologically suitable for malaria transmission. As per Smith et al. (20), the average daily mean surface air temperature was used to establish the development rate to maturity, though where this is below the 31 °C optimum water temperature, we apply a +2 °C offset as suggested by Paaijmans et al. (66) to account for the typical difference observed between surface air temperature and water temperature. Since shading is often available, we do not apply this offset for temperatures above the optimum as larvae are likely to seek optimum conditions where available.

(iii) Discharge with relative humidity (Q_RH): an additional criterion was imposed on the discharge estimates of a minimum relative humidity of 40% (67).

(iv) Potential evapotranspiration (PE): the ratio of total precipitation and potential evapotranspiration was calculated and following Lindsay et al. (68) a minimum value of 0.5 was considered suitable for malaria transmission.

(v) Discharge with potential evapotranspiration (Q_PE): a union of the discharge and potential evapotranspiration suitability layers was computed as the most physically realistic of representing locally-generated surface water alongside larger discharge transfers. Combining suitability layers in this way avoids any double-counting of water in the hydrological model outputs.

(vi) Discharge with potential evapotranspiration and relative humidity (Q_PE_RH): the relative humidity criterion was also applied to (v).

All data were transformed into an equal area projection (Africa Albers Equal Area Conic) appropriate for calculation of areas of malaria transmission over the African continent.

Hydro-climatic suitability and model metrics

We combine our hydrological suitability layers with the temperature curve of Mordecai et al. (8). Following (17) and given our focus on hydro-climatic suitability, the full range of viable temperatures (16-34°C) is considered suitable. The temperature suitability was then combined with each hydrological suitability layer to create a monthly mask of hydro-climatically suitable areas for malaria suitability. The Length of Transmission Season (LTS) was then determined as the maximum number of continuous months of hydro-climatic suitability. Subsequent data analysis was undertaken in Stata 12.1.

Calculation of Model Variance

To calculate whether the variance in estimated malaria suitability is introduced primarily through GCMs or GHMs, the variance ratio (R_{var}) of GCMs to GHMs was calculated (69). For each grid cell, the average variance in malaria LTS for all GCMs (σ^2_{GCM}) and the average variance in malaria LTS for all GHMs (σ^2_{GHM}) are computed and the variance ratio calculated as

$$R_{var} = \frac{\sigma^2_{GCM}}{\sigma^2_{GCM} + \sigma^2_{GHM}} . \quad (\text{Equ. 1})$$

To provide an overall metric of the source of variance, Equ. 1 was also applied to model estimates of areas of stable (>3 months), seasonal (>6 months) and endemic (>9 months) malaria transmission and all suitable areas (>1 month).

Model Validation and Ensemble Creation

Estimates of hydro-climatic suitability for malaria transmission are challenging to evaluate quantitatively, as there are numerous non-climatic controls on malaria transmission (70) including the successful reduction in transmission ranges due to interventions over the last century (3). For this reason, we focus our validation efforts on the pre-intervention map of Lysenko and Semashko (32) which is an approximation of malaria extents at ~1900AD, though the exact boundaries should be viewed with caution. ISIMIP2b climate and hydrological data were extracted for the years 1875-1900 and malaria hydro-climatic suitability calculated. Preindustrial climate and CO₂ scenarios were used where available; only 1986-2005 were available for ORCHIDEE (the differences and possible effects are demonstrated in Fig. S3). Two validation metrics are used: the most comprehensive performance score is the F^2 score (71) calculated from observed (F_o) and modelled (F_m) extents as

$$F^2 = \frac{F_m \cap F_o}{F_m \cup F_o}, \quad (\text{Equ. 2})$$

where $F_m \cap F_o$ is the intersection of modelled and observed malaria suitability extent (i.e. correct forecasts) and $F_m \cup F_o$ is the union of modelled and observed extents. Since the F^2 score is biased towards overprediction (72), we follow Bernhofen et al. (73) in also calculating the bias metric (71) which measures whether an estimate is biased towards underprediction (negative bias score) or overprediction (positive) calculated as

$$\text{Bias} = \frac{(F_m \cap F_o) + F_m}{(F_m \cap F_o) + F_o} - 1. \quad (\text{Equ. 3})$$

The performance of each individual model on each of these metrics is shown in Fig. S4. Finally, following (73) a single Ensemble Score (ES) was calculated as

$$ES = \text{Average } F^2 - |0.2 * \text{Average Bias}|, \quad (\text{Equ. 4})$$

where the bias adjustment factor of 0.2 was selected to penalise for bias while still weighting F^2 as the most important score. ES was calculated for all hydro-climatic layers at two malaria LTS levels: >1 month (i.e., all malaria suitability) or >3 months continuous transmission suitability to indicate meso-endemicity, following Chemison et al. (74) and Martens et al. (75) (Tables S2-S3).

ES was used to weight all models within the ensemble (i.e., all combinations of global climate models and hydrological models for each hydro-climatic suitability layer). Weights from unity-based normalisation (scaled 0-1) were awarded to each model separately for each malaria level. Finally, model agreement layers (i.e., minimum number of models in the ensemble agreeing that malaria transmission is suitable at a given location) were created for all agreement levels (ranging from a layer showing where at least one model identified suitability to a layer showing only where all models in the ensemble agreed was suitable for malaria). Each of these agreement layers was validated based on ES (Fig. S5). The optimum number of model agreements for each hydro-climatic suitability layer was calculated (see Table S4) and that number of the top performing models in each ensemble were then awarded a maximum weight (73). The weights for all and meso-endemic malaria levels were merged and then renormalized such that the total weights in each ensemble are 1 (Fig. S6 and Table S5). These ensemble weights were then used to make future predictions using the model ensembles, noting that some rescaling is needed for RCP 8.5 where MPI-HM and PCR-GLOWB hydrological model outputs were unavailable. This ensemble approach is common practice and allows for a representation of the range of possible

outcomes across the models used. However, the cascade of uncertainty that prevails in climate change impact assessment (76), means that even with model intercomparison projects like ISIMIP and CMIP, it is impossible to analyse the full range of uncertainty because such projects are not exhaustive in their selection of models and scenarios.

Further validation of the final weighted layers was performed for each hydrological representation using the historical time period (1986-2005). The anopheles inventory for sub-Saharan Africa (1898-2016) of Kyalo et al. (36) is independent of climate but provides only a discontinuous distribution of observation locations. Given this limitation and the more reduced range of contemporary malaria and anopheles prevalence, the percentage of positive observations that are correctly modelled was calculated as the most appropriate metric (Table S6). We also compare model predictions with both malaria parasite ratio data from the Malaria Atlas Project (considering areas where the parasite ratio $>0.01\%$ in the year 2000 (37)) and the estimate of the mean modelled relative probability of occurrence of *Anopheles gambiae* (considering areas where the probability >0.05) of Wiebe et al. (38) (Fig. S9). However, it should be noted that both these datasets rely in some part on gridded climate data for interpolation and do not therefore serve for validation analysis independent of climate.

Population data

Human population analysis was carried out using the methodology outlined in Smith et al. (20). Gridded estimates of the human population in Africa were provided by WorldPop (51) at a resolution of 30 arc seconds (~ 1 km at the equator). These were used to evaluate the number of individuals residing within areas climatically suitable for malaria transmission. Analysis was performed for individuals of all ages and for children aged under five years of age as this group is particularly susceptible to malaria (77, 78). Analysis was performed for the mid-point of each time period using the closest WorldPop grids (available annually between 2000 and 2020). For past and contemporary scenarios, grids were rescaled to match country level UN estimates (total population and under 5s) at the mid-point year of each time period (52). A similar approach was used for future scenarios, using country level UN projections (52) to maintain consistency with estimates for the historical periods. Primary analysis was based on the UN medium projection variant, whilst supplementary analysis was performed using the UN low and high projection variants for comparison and sensitivity analysis. Final results were aggregated to the continental and country level.

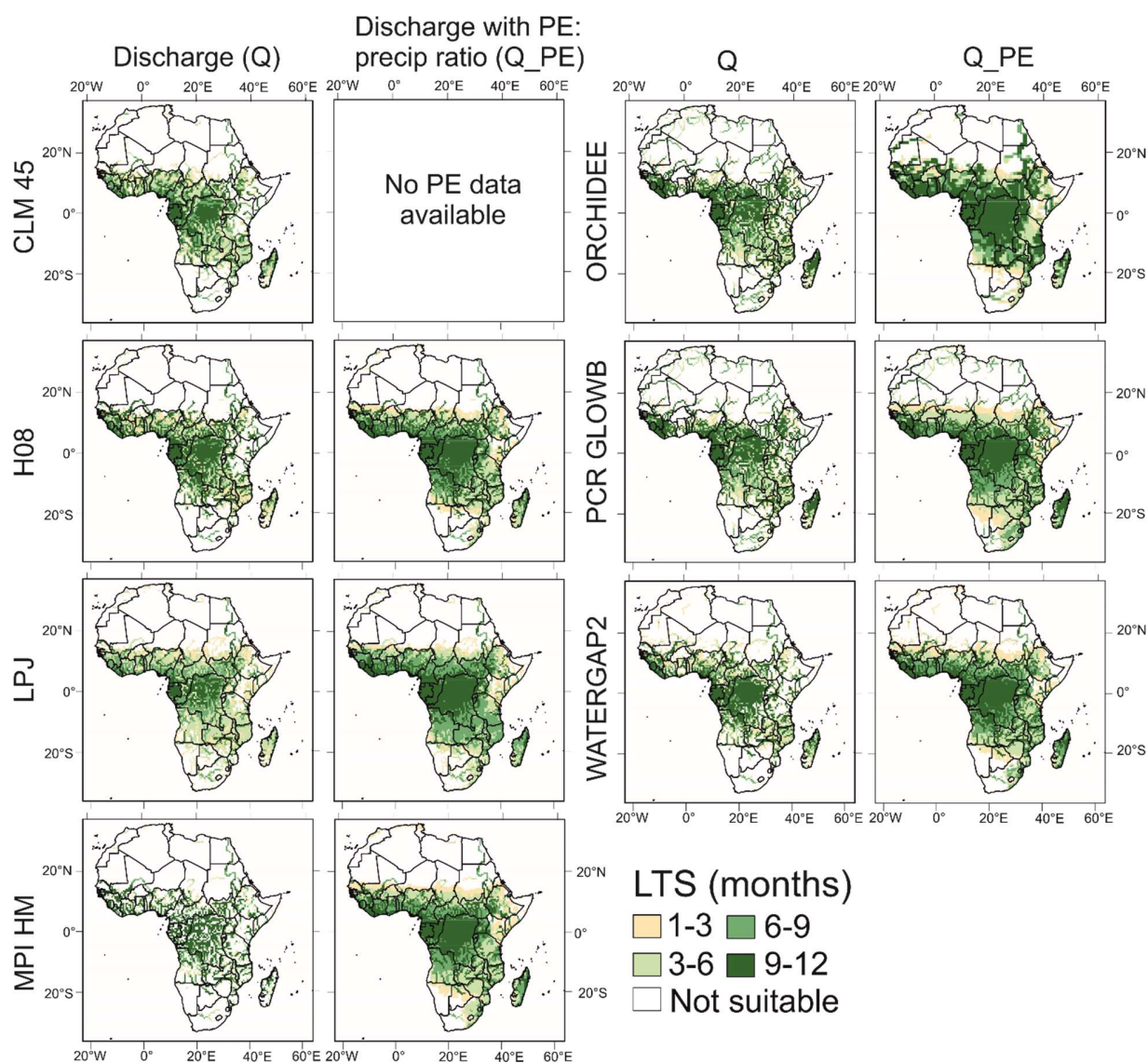


Fig. S2.

Comparison of historical (1986-2005) malaria length of transmission season (LTS) estimates for each hydrological model, for both 'discharge' and 'discharge with potential evapotranspiration' hydrological representations. LTS estimates are averages across the four global climate models.

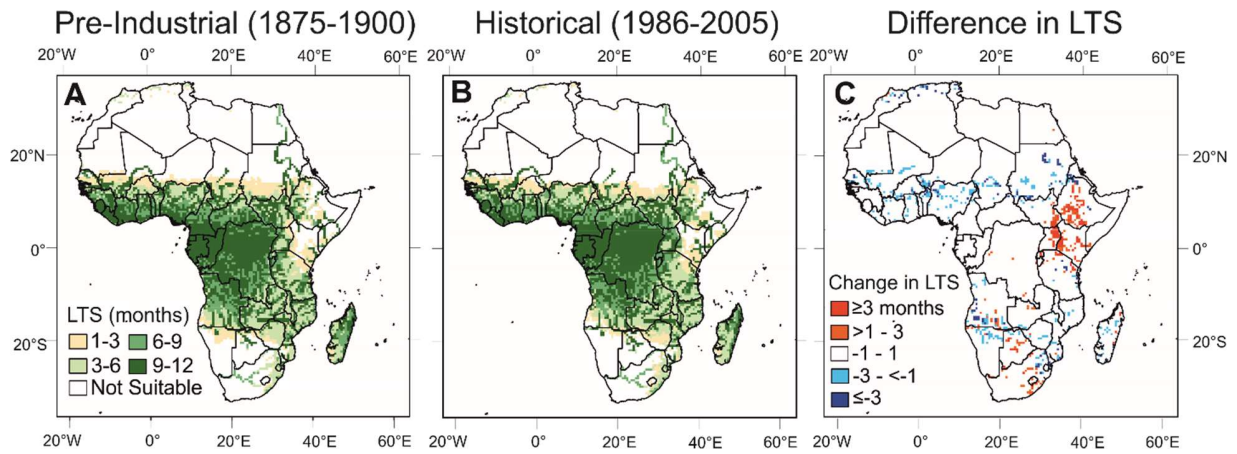


Fig. S3.

Example difference in LTS of malaria transmission hydro-climatic suitability for validation layers (Q PE hydrological representation, IPSL GCM, H08 hydrological model). Pre-Industrial society (1875-1900) (A) is used for model validation; however, due to data availability, ORCHIDEE hydrological models use 1986-2005 data (B) as an approximation. (C) Shows the changes in LTS estimates since the preindustrial period (i.e., historical – preindustrial) where red shades indicate an extension of the transmission season.

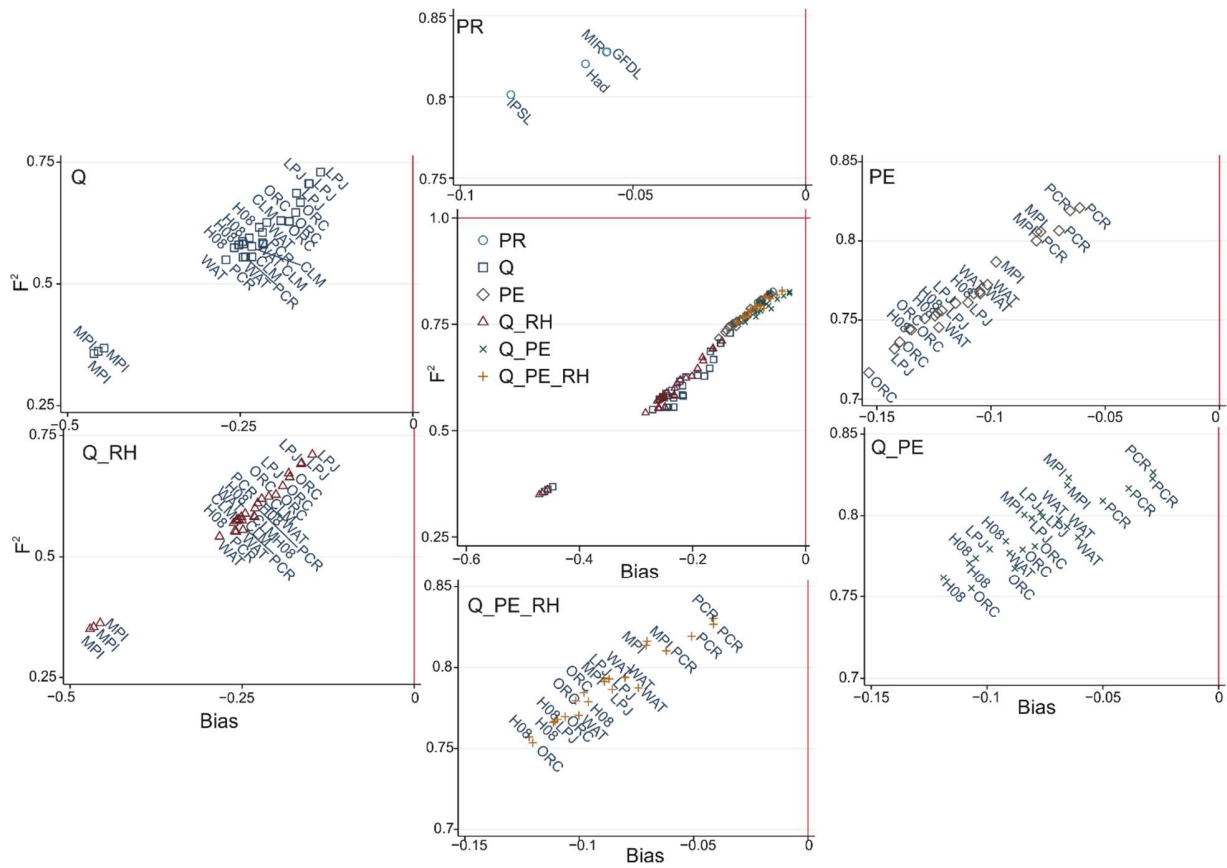


Fig. S4.

Validation metrics of model performance (F^2 and Bias) for all models in each ensemble (shown here for the >1 month malaria suitability level). The central panel shows all ensembles together, while those around the periphery separate these by hydrological representations and labels for each point indicate the hydrological model (or GCM for precipitation models).

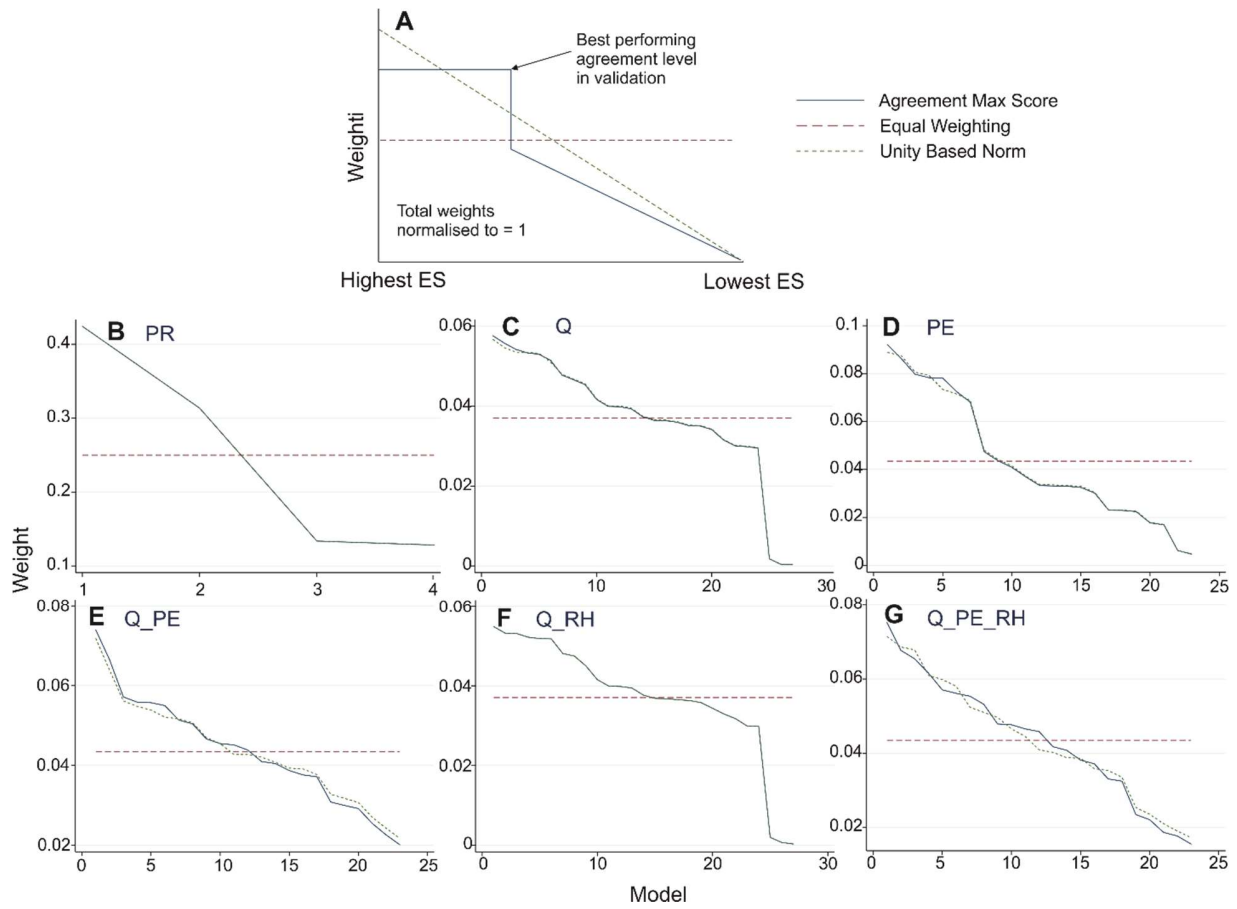


Fig. S5.

Weighting curves applied to each model ensemble based on validation performance. **(A)** Schematic of weighting curve calculation. Ensemble score (ES) is used to rank each model and a unity-based normalisation used to assign a weight. The best models up to the best-performing number of model agreements are then assigned a maximum weighting, with weights calculated separately for ‘all malaria’ and >mesoendemic season length (>3 months). These are summed and rescaled to 1. Models are ordered from best performing (left) to worst performing (right). **(B-G)** Weights are shown for ensembles for each hydrological representation.

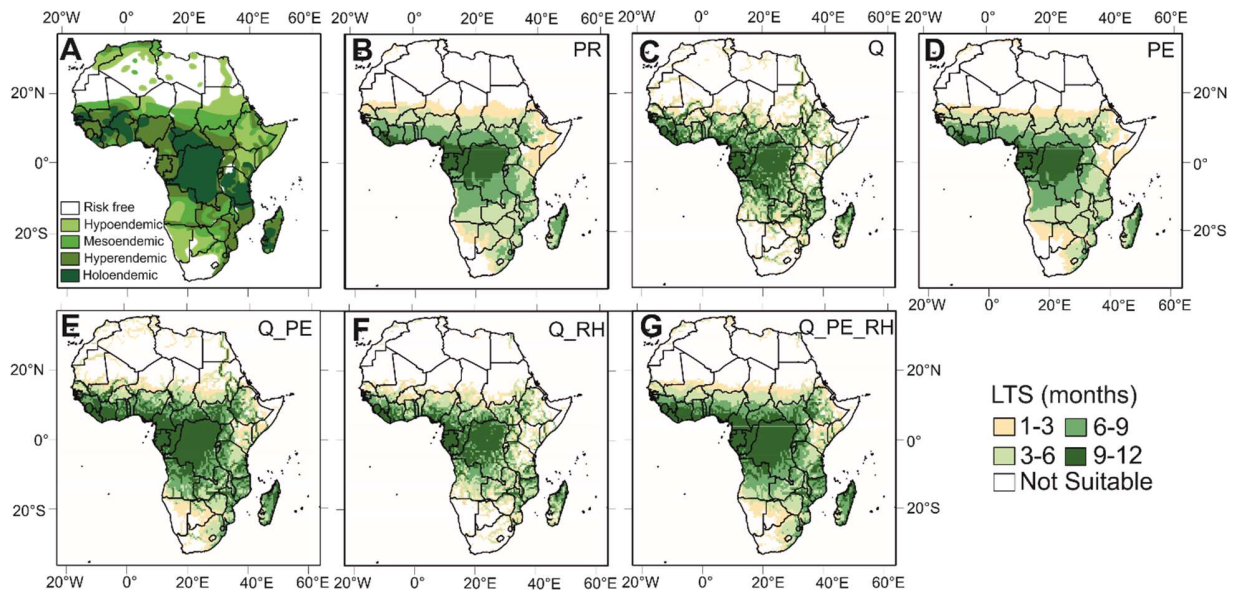


Fig. S6.

Malaria season length maps (1875-1900) for each hydrological representation model ensemble. Fig. 2E displays the areas by 3-month wide classification.

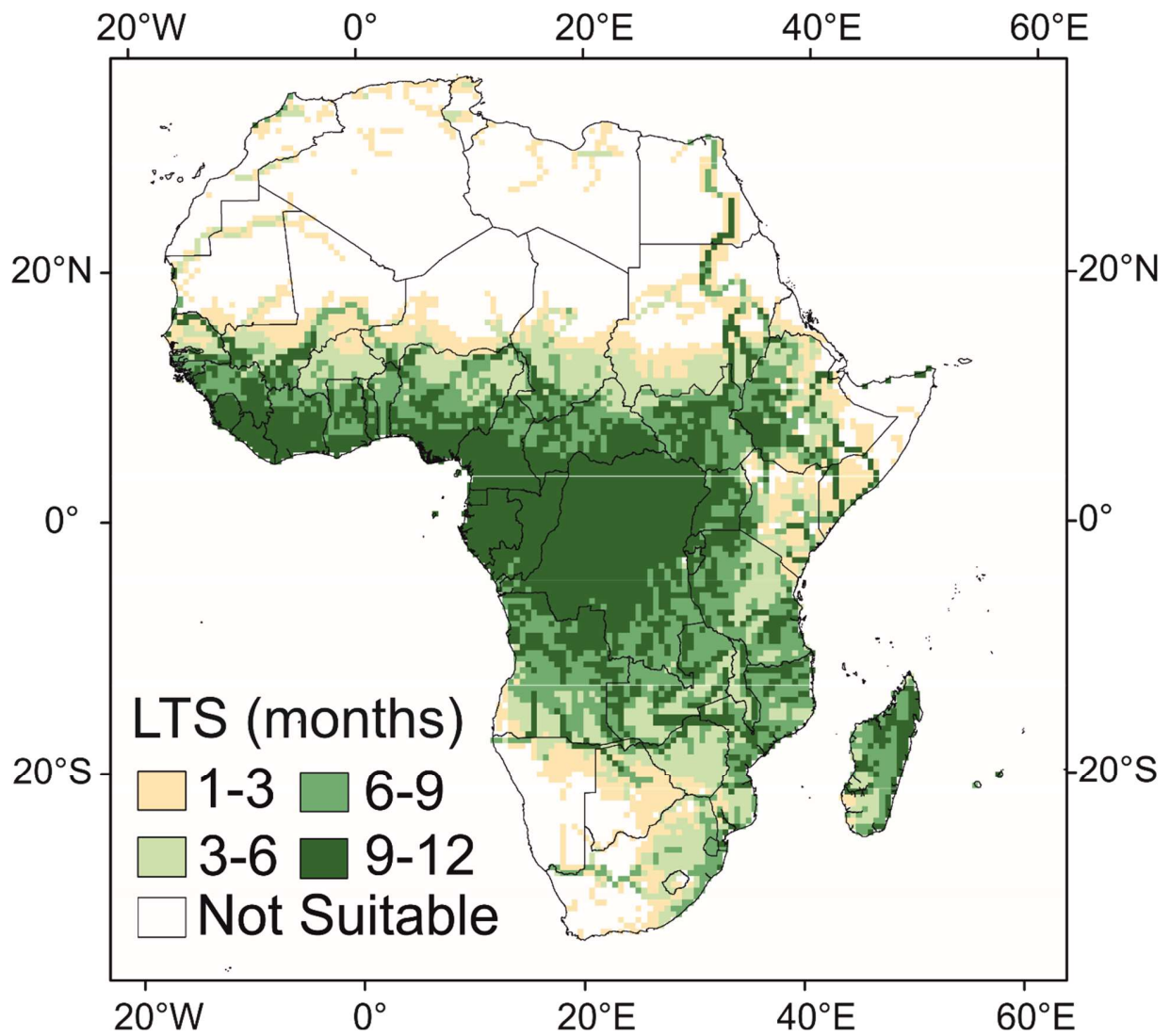


Fig. S7.

An enlarged version of the malaria season length predictions from the weighted ensemble of hydrologically-informed (Q_PE) malaria season length models for the historical period (1986-2005) for each hydrological representation model ensemble.

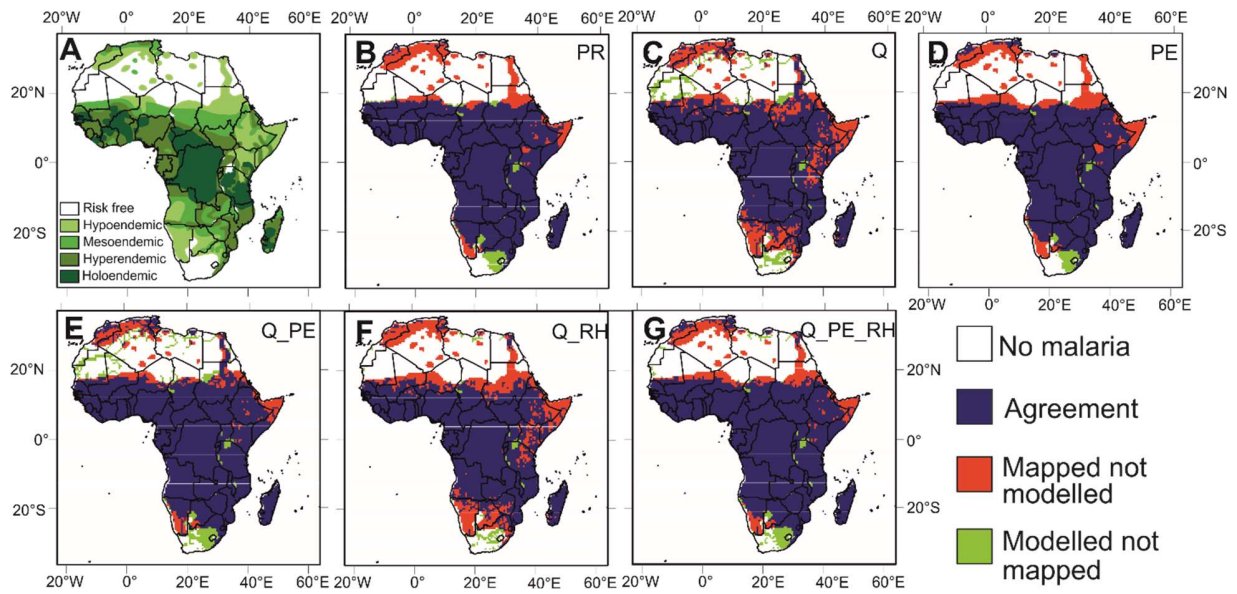


Fig. S8.

Validation layers for final weighted ensembles for each hydrological representation. The Lysenko and Semashko (32) pre-intervention malaria map is compared against model estimates for 1875-1900 with a pre-industrial society model.

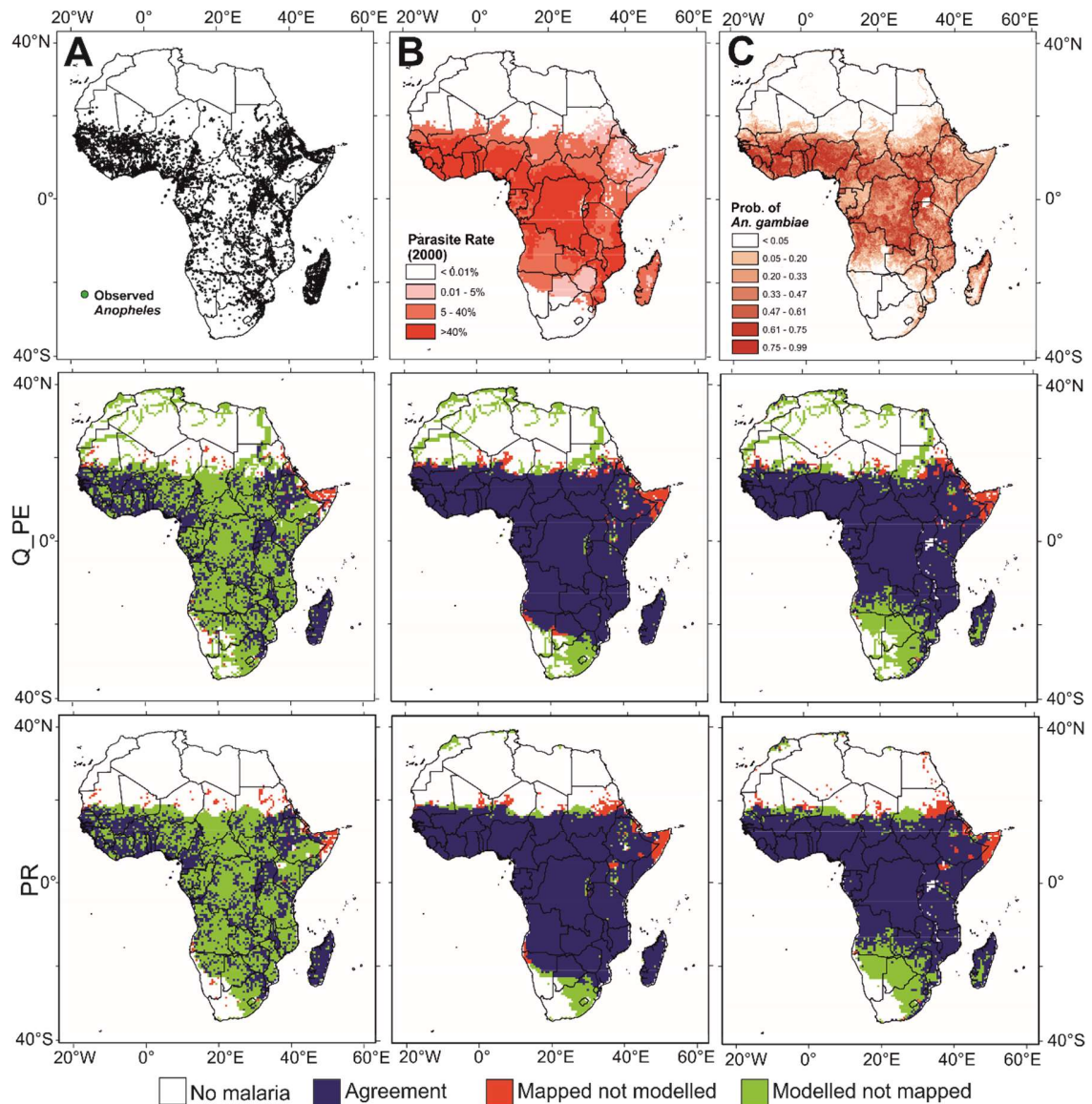


Fig. S9.

Validation of weighted malaria suitability estimates (for discharge with potential evaporation and precipitation-based estimates) against more recent data sources for ‘All malaria’ (>1 month suitability) transmission suitability calculated over the time period 1986-2005. (A) Kyalo et al. (36) anopheles inventory of sub-Saharan Africa (1898-2016) displaying all observations of anophelines. (B) Malaria Atlas Project modelled estimates of parasite rate from Bhatt et al. (37) for 2000. (C) Wiebe et al. (38) estimate of the mean modelled relative probability of occurrence of *Anopheles gambiae*.

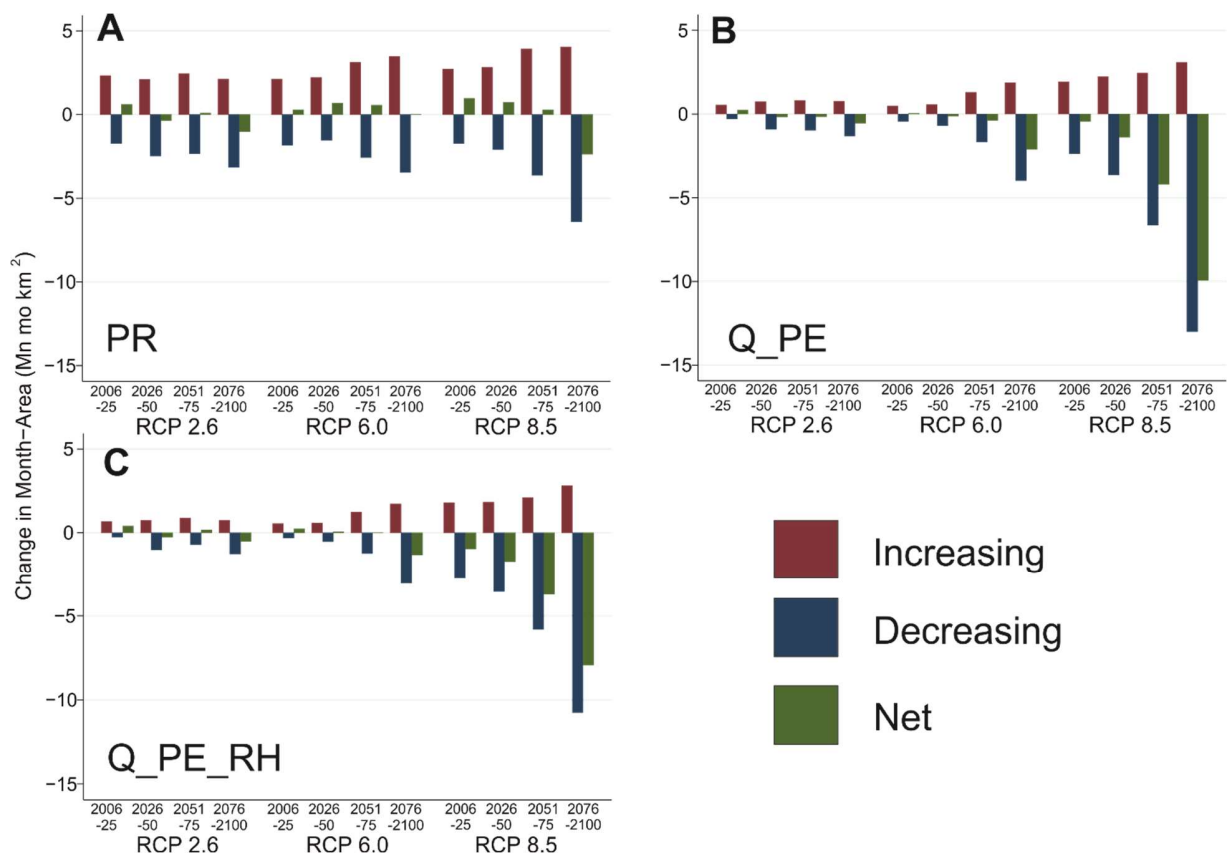
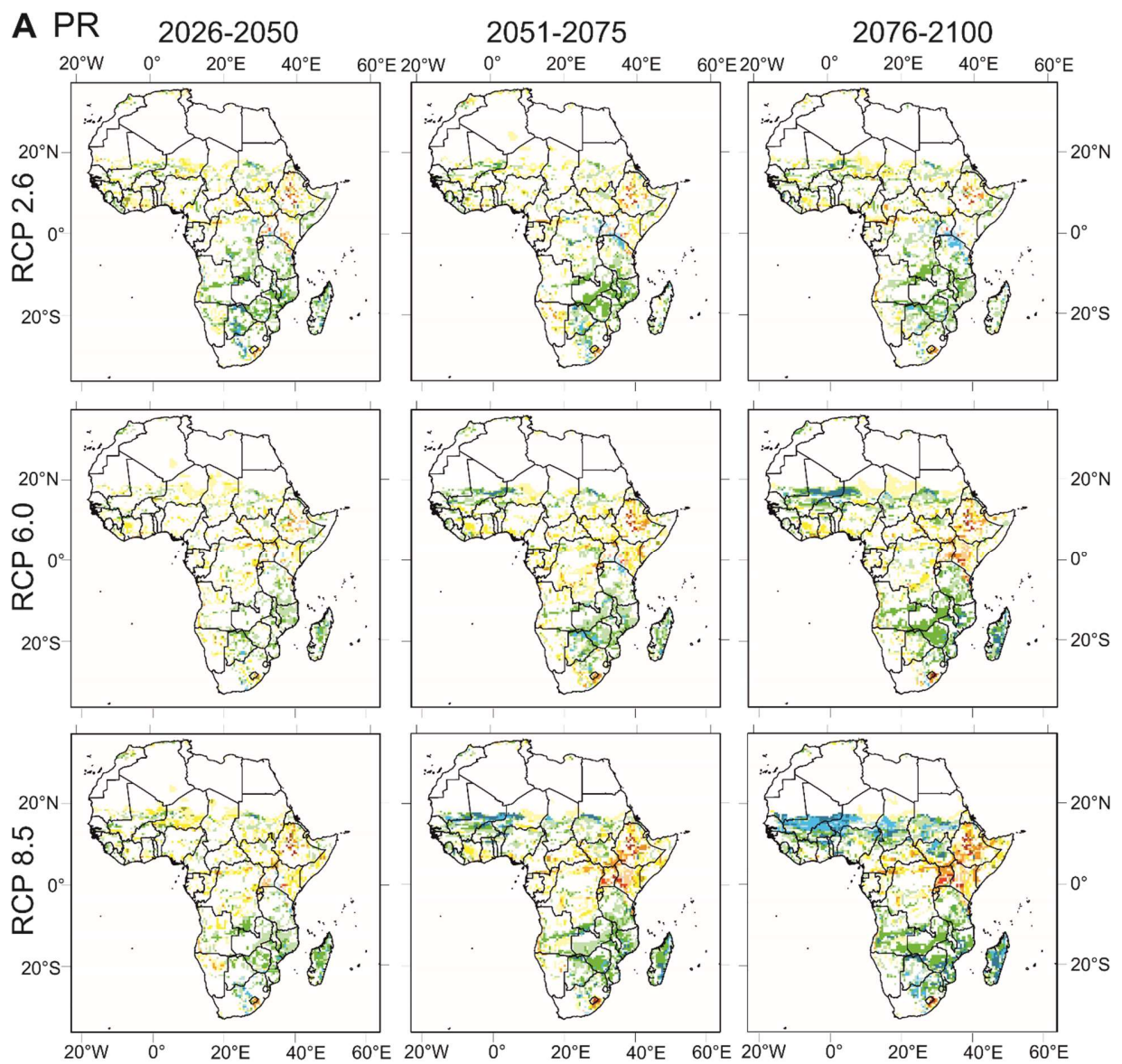


Fig. S10.

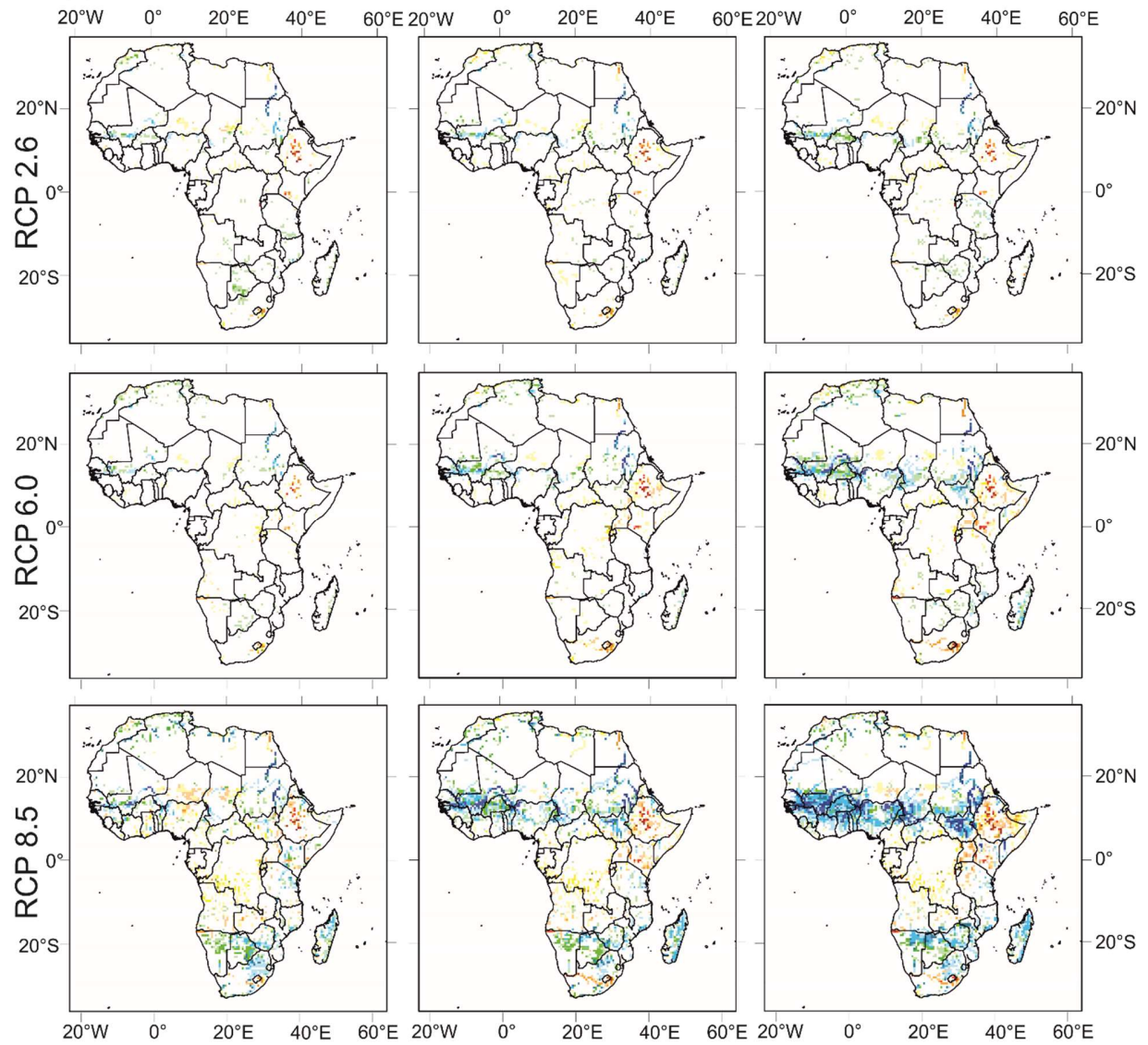
Projected changes from 1985-2005 in malaria suitability 'month-areas' suitable (in millions mo km²) in future projections where the areas are multiplied by the number of months change in malaria climatic suitability to provide a clearer indication of the magnitude of changes) for each time period, RCP and hydrological representation.



BQ_PE 2026-2050

2051-2075

2076-2100



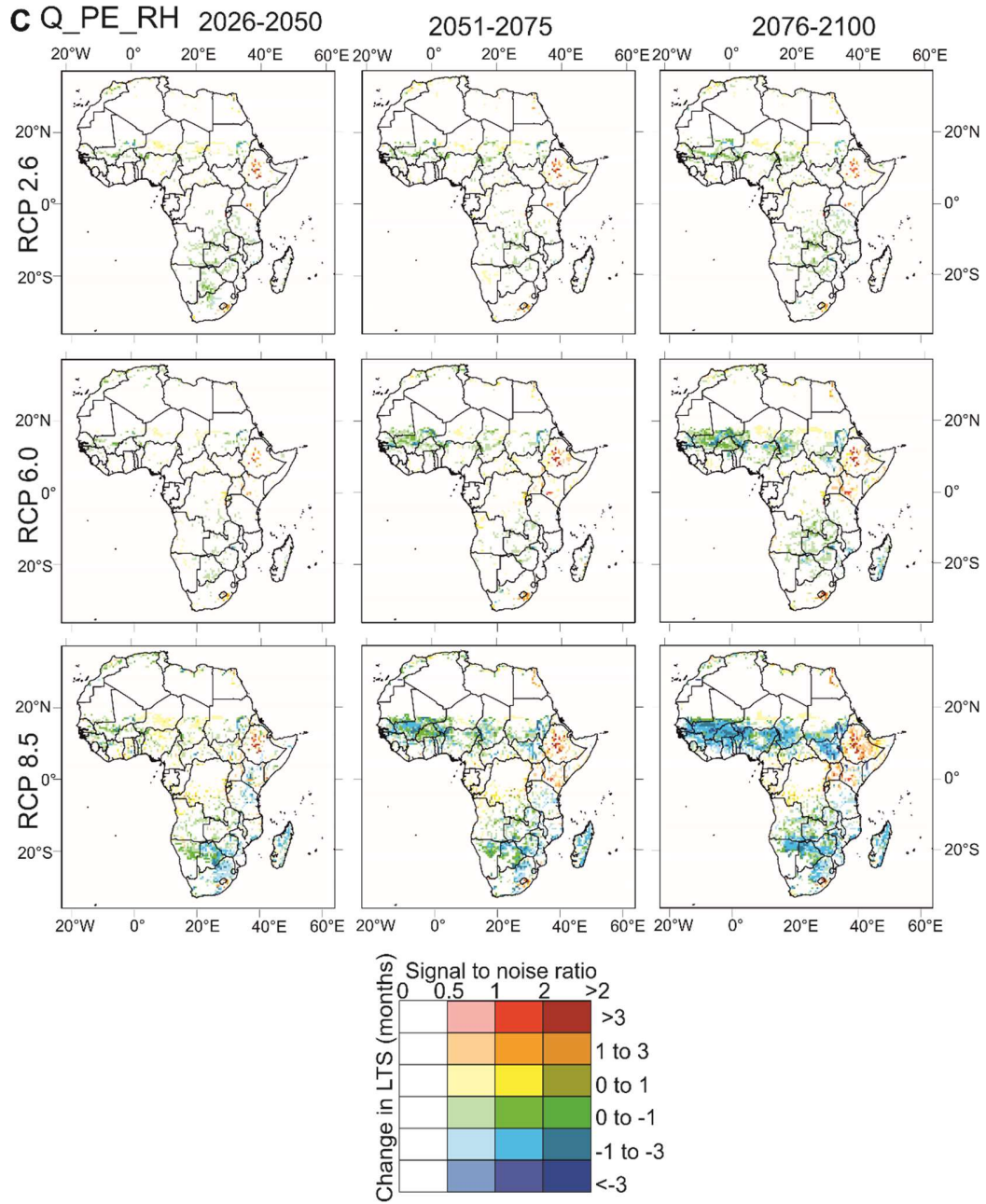


Fig. S11.

Changes in duration of malaria transmission climatic suitability between the period 1985-2005 and each future period for each RCP and hydrological representation. Different saturations indicate the signal-to-noise ratio across the model ensemble, with the noise defined as the standard deviation of estimates across the projections, as per Fig. 3.

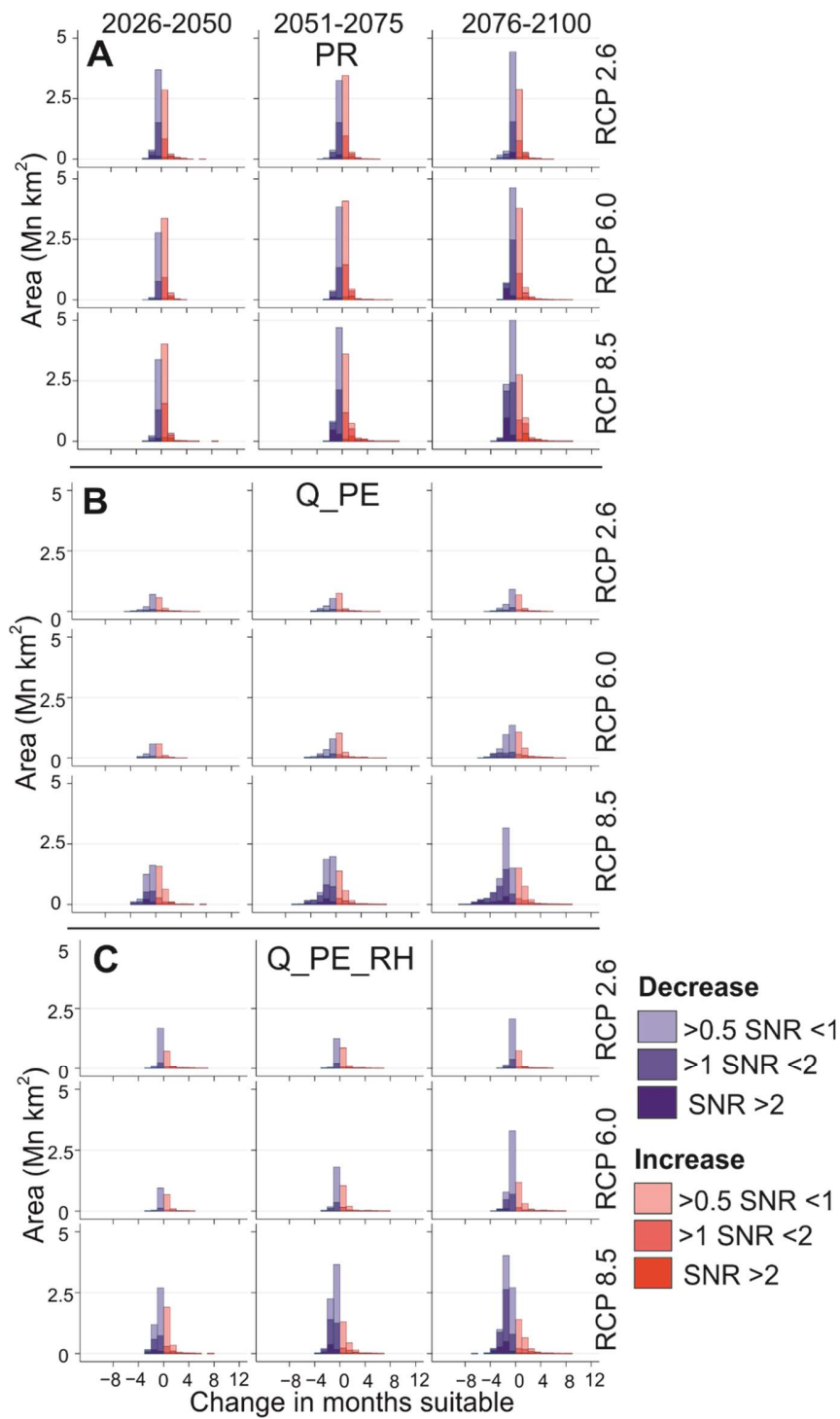


Fig. S12.

Areas with changing malaria transmission suitability. Stacked histograms showing changes in number of months climatically suitable between the period 1985-2005 and all future time periods for each RCPs and hydrological representation, split into categories of signal-to-noise ratio as per Fig. 3.

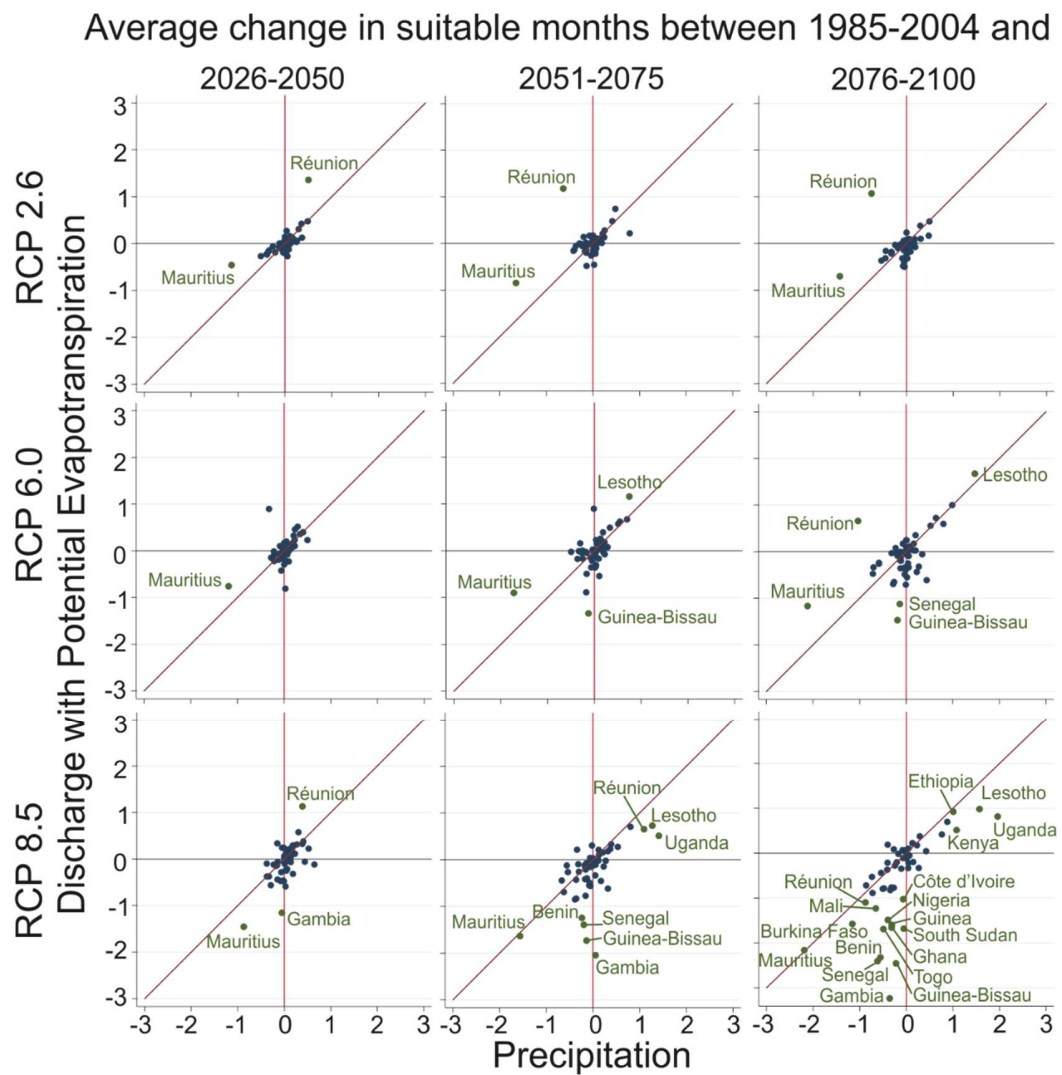


Fig. S13.

By-country comparison of average changes in malaria transmission suitability season length between the period 1985-2005 and each future period. Discharge and potential evapotranspiration and precipitation-based estimates are plotted on separate axes and the 1:1 line shown in red. Countries with an estimated average change of >1 month are named in each plot.

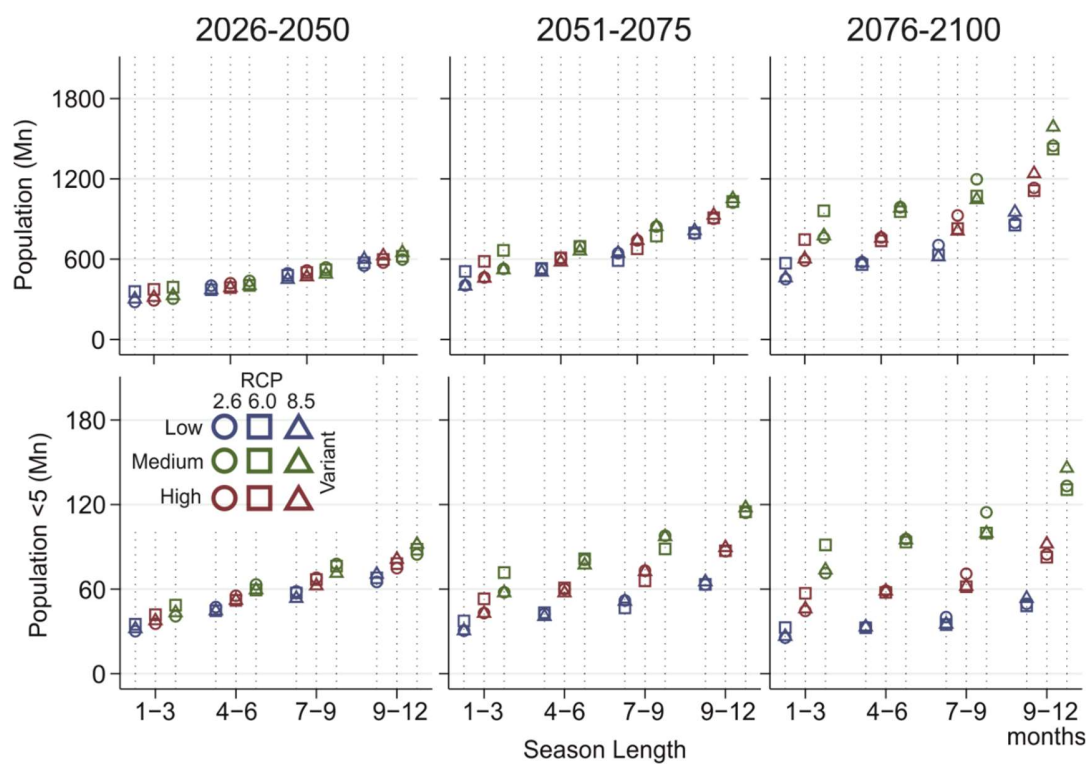


Fig. S14.

Comparison of UN population variants. The effect of UN population variant (High, Medium or Low) on future population predictions of malaria suitability in Africa for the discharge with potential evapotranspiration (Q_{PE}) hydrological representation. Results from each RCP are displayed separately.

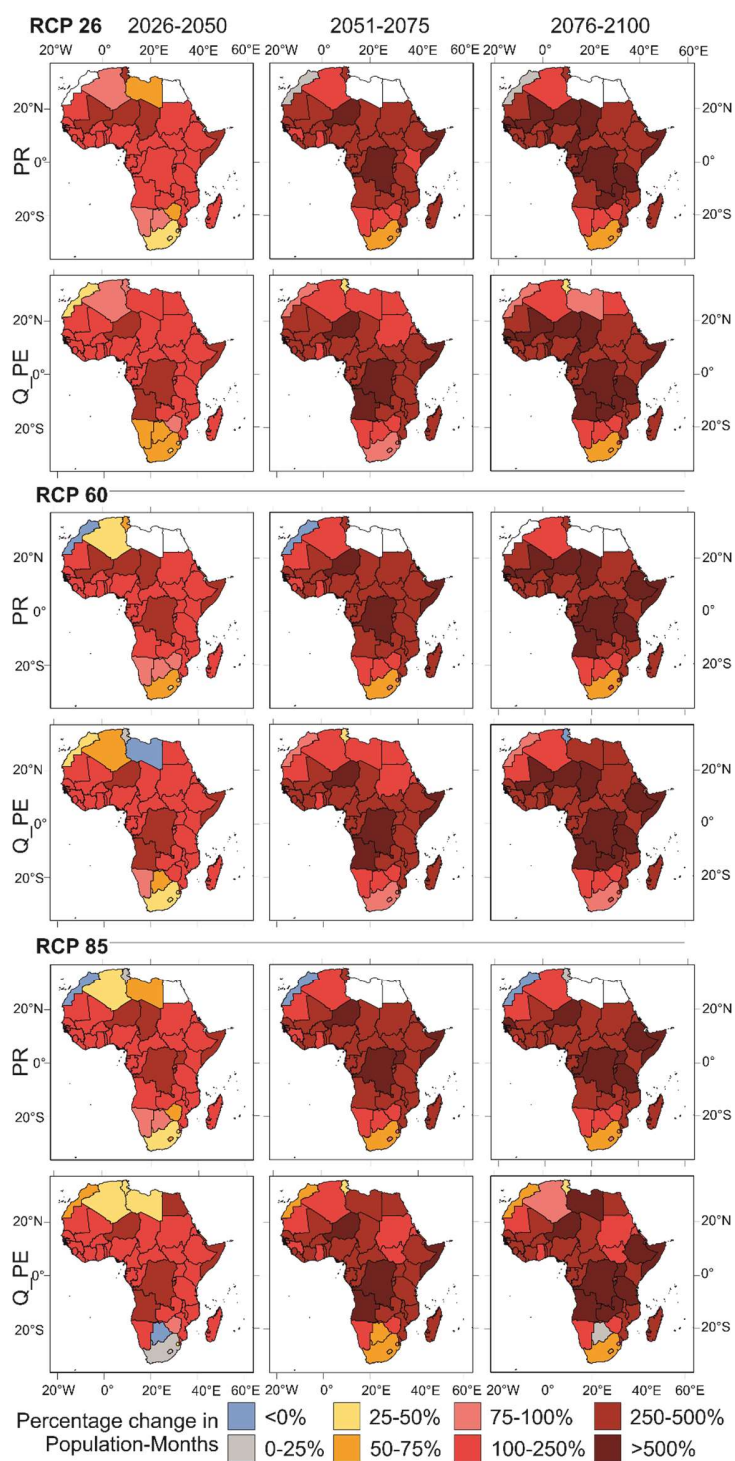


Fig. S15.

Predicted percentage changes in population-months (i.e., population exposed multiplied by the number of additional months of exposure) since 1986-2005 by country and for each RCP. White shading indicates no estimated malaria suitability.

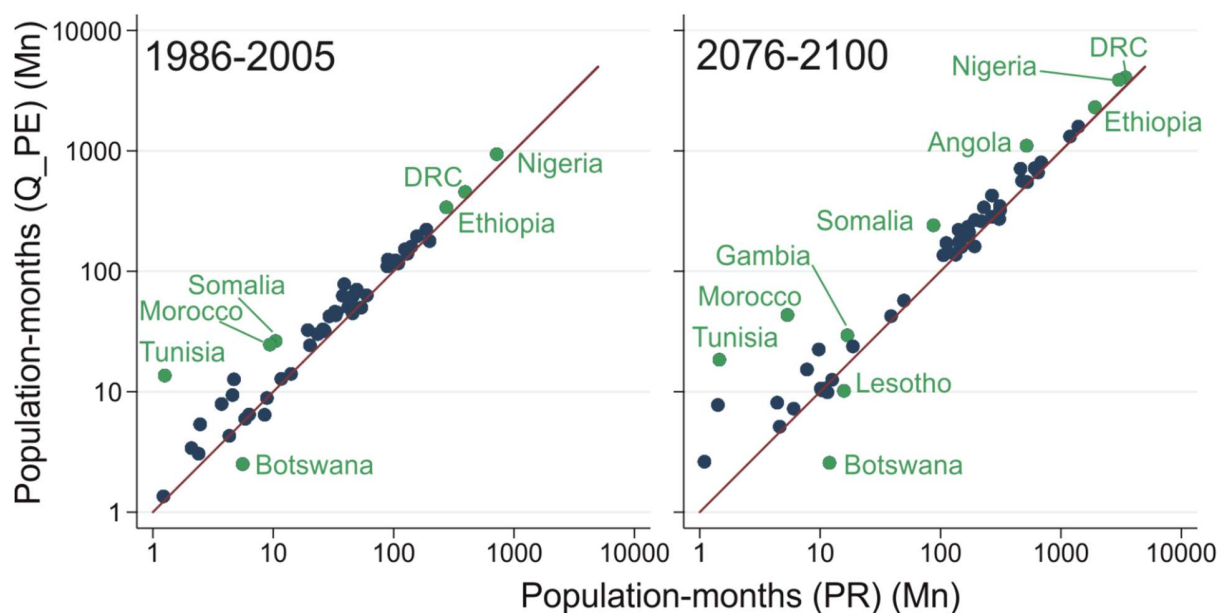


Fig. S16.

Country-by-country comparison of population-months. Comparison of population-month estimates (i.e., population exposed multiplied by the number of additional months of exposure) from the Q_PE and PR hydrological representations is presented for both the historical period 1986-2005 (using population estimates) and 2076-2100 (using RCP 8.5 and the Medium UN population estimates). The 1:1 line is shown in red. Individual countries are highlighted when a large difference in predictions is observed or for countries with high overall values.

Table S1. Areas of different model agreement levels for model ensembles for each hydrological representation (areas stated in Mn km²) and four malaria season length classifications: ‘All’ (LTS >1 month), ‘Stable’ (LTS > 3 months), ‘Seasonal’ (LTS > 6 months) and ‘Endemic’ (LTS > 9 months).

Malaria level	Hydrology	Total Models	No Models Predict	Minority Agree	Majority Agree	All Models Agree
All	PR	4	10.13	0.63	0.79	18.26
	Q	27	9.14	7.45	7.47	5.75
	PE	23	10.11	2.43	1.97	15.30
	Q_PE	23	7.28	4.23	2.48	15.82
	Q_RH	27	10.58	6.19	7.35	5.68
	Q_PE_RH	23	8.57	3.19	2.33	15.72
Stable	PR	4	14.67	0.63	0.76	13.74
	Q	27	11.92	7.25	6.16	4.48
	PE	23	14.96	2.23	1.86	10.75
	Q_PE	23	10.37	4.95	2.58	11.91
	Q_RH	27	14.47	5.28	5.87	4.19
	Q_PE_RH	23	12.62	3.05	2.39	11.74
Seasonal	PR	4	21.94	0.61	0.59	6.67
	Q	27	14.41	8.09	4.74	2.57
	PE	23	21.29	2.26	1.57	4.68
	Q_PE	23	13.38	5.74	3.53	7.16
	Q_RH	27	17.67	5.80	4.20	2.14
	Q_PE_RH	23	16.64	3.59	3.06	6.51
Endemic	PR	4	27.57	0.39	0.24	1.61
	Q	27	17.04	7.69	3.92	1.17
	PE	23	26.79	1.37	0.74	0.91
	Q_PE	23	16.91	6.20	3.49	3.21
	Q_RH	27	21.02	4.85	2.94	1.00
	Q_PE_RH	23	20.81	3.59	2.52	2.88

Table S2. Ensemble Score validation metric (Equ. 4) for each model for ‘All malaria’ (>1 month suitability) transmission suitability based on the pre-intervention (~1900) map of Lysenko and Semashko (32). The final row shows the ensemble score for the final weighted ensembles.

GCM	Hydrological Model	PR	Q	PE	Q PE	Q RH	Q PE RH
GFDL	-	0.816	-	-	-	-	-
	CLM45	-	0.533	-	-	0.526	-
	H08	-	0.529	0.730	0.753	0.522	0.746
	LPJ	-	0.676	0.732	0.782	0.661	0.774
	MPI_HM	-	0.270	0.790	0.810	0.260	0.802
	ORCHIDEE	-	0.593	0.708	0.750	0.588	0.748
	PCR_GLOWB	-	0.509	0.806	0.817	0.506	0.819
	WATERGAP2	-	0.507	0.747	0.775	0.500	0.773
HadGEM	-	0.808	-	-	-	-	-
	CLM45	-	0.584	-	-	0.575	-
	H08	-	0.572	0.745	0.766	0.565	0.760
	LPJ	-	0.703	0.739	0.784	0.681	0.769
	ORCHIDEE	-	0.635	0.717	0.762	0.628	0.759
	PCR_GLOWB	-	0.541	0.792	0.809	0.537	0.809
	WATERGAP2	-	0.562	0.746	0.784	0.554	0.778
IPSL	-	0.784	-	-	-	-	-
	CLM45	-	0.530	-	-	0.525	-
	H08	-	0.522	0.717	0.738	0.517	0.733
	LPJ	-	0.653	0.703	0.760	0.635	0.745
	MPI_HM	-	0.265	0.767	0.784	0.255	0.776
	ORCHIDEE	-	0.592	0.686	0.734	0.582	0.730
	PCR_GLOWB	-	0.506	0.784	0.799	0.501	0.798
	WATERGAP2	-	0.495	0.721	0.759	0.485	0.750
MIROC5	-	0.816	-	-	-	-	-
	CLM45	-	0.547	-	-	0.540	-
	H08	-	0.538	0.728	0.749	0.532	0.744
	LPJ	-	0.675	0.737	0.787	0.658	0.776
	MPI_HM	-	0.279	0.790	0.806	0.270	0.800
	ORCHIDEE	-	0.613	0.725	0.765	0.608	0.765
	PCR_GLOWB	-	0.538	0.809	0.821	0.535	0.822
	WATERGAP2	-	0.530	0.752	0.780	0.525	0.778
WEIGHTED ENSEMBLE		0.816	0.723	0.782	0.817	0.698	0.808

Table S3. Ensemble Score validation metric (Equ. 4) for each model for ‘Mesoendemic’ (>3 months) transmission suitability based on the pre-intervention (~1900) map of Lysenko and Semashko (32).

GCM	Hydrological Model	PR	Q	PE	Q PE	Q RH	Q PE RH
GFDL	-	0.777	-	-	-	-	-
	CLM45	-	0.596	-	-	0.598	-
	H08	-	0.560	0.730	0.751	0.562	0.757
	LPJ	-	0.653	0.700	0.762	0.648	0.770
	MPI_HM	-	0.316	0.769	0.774	0.309	0.779
	ORCHIDEE	-	0.639	0.705	0.756	0.651	0.773
	PCR_GLOWB	-	0.495	0.762	0.724	0.512	0.765
	WATERGAP2	-	0.489	0.724	0.744	0.491	0.759
HadGEM	-	0.751	-	-	-	-	-
	CLM45	-	0.631	-	-	0.629	-
	H08	-	0.569	0.732	0.756	0.569	0.761
	LPJ	-	0.670	0.707	0.764	0.666	0.766
	ORCHIDEE	-	0.685	0.698	0.756	0.692	0.776
	PCR_GLOWB	-	0.531	0.738	0.706	0.545	0.736
	WATERGAP2	-	0.525	0.708	0.740	0.522	0.747
IPSL	-	0.765	-	-	-	-	-
	CLM45	-	0.605	-	-	0.605	-
	H08	-	0.567	0.724	0.751	0.570	0.756
	LPJ	-	0.672	0.689	0.766	0.666	0.771
	MPI_HM	-	0.321	0.756	0.766	0.319	0.770
	ORCHIDEE	-	0.655	0.696	0.750	0.665	0.763
	PCR_GLOWB	-	0.517	0.750	0.735	0.532	0.765
	WATERGAP2	-	0.504	0.705	0.742	0.506	0.752
MIROC5	-	0.765	-	-	-	-	-
	CLM45	-	0.614	-	-	0.614	-
	H08	-	0.565	0.719	0.745	0.569	0.751
	LPJ	-	0.667	0.696	0.766	0.663	0.772
	MPI_HM	-	0.327	0.756	0.762	0.323	0.766
	ORCHIDEE	-	0.676	0.720	0.766	0.692	0.790
	PCR_GLOWB	-	0.535	0.747	0.722	0.554	0.762
	WATERGAP2	-	0.529	0.711	0.745	0.530	0.753

Table S4. Optimum number of model agreements used to award maximum weight to the best performing models up to that limit (following (73)) for each hydrological representation and a range metrics.

Hydrological Representation	Optimum agreement levels					
	Ensemble Score		F ²		Bias	
	All	Mesoendemic	All	Mesoendemic	All	Mesoendemic
PR	1	1	1	1	1	2
Q	1	5	1	5	1	5
PE	1	3	1	3	1	2
Q_PE	3	9	3	9	3	9
Q_RH	1	2	1	1	1	3
Q_PE_RH	2	6	2	5	1	7

5

Table S5. Weights applied to model ensemble for each hydrological representation.

GCM	Hydrological Model	Q	PE	Q_RH	Q_PE	Q_PE_RH
GFDL PR 0.4244	CLM45	0.0393		0.0394		
	H08	0.0364	0.0409	0.0365	0.0299	0.0234
	LPJ	0.0533	0.0230	0.0522	0.0550	0.0465
	MPI_HM	0.0004	0.0863	0.0003	0.0740	0.0752
	ORCHIDEE	0.0466	0.0177	0.0475	0.0308	0.0477
	PCR_GLOWB	0.0301	0.0921	0.0318	0.0514	0.0677
	WATERGAP2	0.0295	0.0436	0.0299	0.0376	0.0381
HadGEM PR 0.1337	CLM45	0.0454		0.0450		
	H08	0.0399	0.0474	0.0399	0.0387	0.0330
	LPJ	0.0576	0.0301	0.0549	0.0559	0.0417
	ORCHIDEE	0.0530	0.0169	0.0532	0.0451	0.0532
	PCR_GLOWB	0.0350	0.0680	0.0363	0.0371	0.0408
	WATERGAP2	0.0360	0.0330	0.0358	0.0405	0.0324
IPSL PR 0.1283	CLM45	0.0399		0.0398		
	H08	0.0364	0.0325	0.0368	0.0226	0.0155
	LPJ	0.0542	0.0062	0.0518	0.0438	0.0458
	MPI_HM	0.0004	0.0781	0.0007	0.0557	0.0479
	ORCHIDEE	0.0478	0.0047	0.0481	0.0201	0.0186
	PCR_GLOWB	0.0316	0.0725	0.0329	0.0455	0.0554
	WATERGAP2	0.0299	0.0224	0.0299	0.0292	0.0220
MIROC5 PR 0.3136	CLM45	0.0417		0.0415		
	H08	0.0374	0.0331	0.0377	0.0254	0.0177
	LPJ	0.0558	0.0229	0.0531	0.0572	0.0616
	MPI_HM	0.0018	0.0783	0.0020	0.0665	0.0571
	ORCHIDEE	0.0515	0.0333	0.0518	0.0466	0.0561
	PCR_GLOWB	0.0351	0.0797	0.0368	0.0504	0.0654
	WATERGAP2	0.0341	0.0370	0.0344	0.0410	0.0371

5 **Table S6.** Validation of weighted malaria suitability estimates against more recent data sources for ‘All malaria’ (>1 month suitability) transmission suitability calculated over the time period 1986-2005.

Hydrological Representation	Percentage of positive occurrences correctly identified by malaria suitability model		
	Anopheles inventory of sub-Saharan Africa (1898-2016) from Kyalo et al. (29)	Malaria Atlas Project modelled estimates of parasite rate for 2000 from Bhatt et al. (30)	estimate of the mean modelled relative probability of occurrence of <i>Anopheles gambiae</i> by Wiebe et al. (31)
PR	93.1	95.3	94.1
Q	84.7	85.2	87.5
PE	89.9	91.9	91.4
Q_PE	93.3	94.6	94.6
Q_RH	82.5	83.1	85.6
Q_PE_RH	92.1	94.2	93.8

Table S7. Areas estimated to be hydro-climatically suitable for malaria transmission by the weighted ensembles of each hydrological representation for each RCP and time period. Areas suitable for > 1 month ('All') and > 3 months (i.e., stable areas) are calculated.

	Total Area (Mn km ²)			Stable Area (Mn km ²)		
	Q PE	PR	Q PE RH	Q PE	PR	Q PE RH
1875-1900	19.825	18.907	18.457	14.646	13.863	13.968
1986-2005	19.315	18.776	18.121	14.588	14.113	13.936
RCP 2.6						
2006-2025	19.814	19.011	18.396	14.675	14.252	14.031
2026-2050	19.486	18.762	18.127	14.521	13.892	13.828
2051-2075	19.579	18.953	18.338	14.544	14.057	13.869
2076-2100	19.466	18.733	18.176	14.298	13.814	13.626
RCP 6.0						
2006-2025	19.544	18.979	18.367	14.570	14.182	13.901
2026-2050	19.312	18.773	18.089	14.515	14.147	13.855
2051-2075	19.307	18.724	18.156	14.530	13.901	13.872
2076-2100	19.205	18.362	17.976	14.281	13.773	13.620
RCP 8.5						
2006-2025	19.170	19.034	17.857	14.408	13.979	13.773
2026-2050	19.109	18.843	17.692	14.353	14.139	13.579
2051-2075	19.049	18.486	17.547	14.055	13.521	13.315
2076-2100	18.892	17.895	17.240	13.455	13.037	12.785

Table S8. Changes in total (suitable for > 1 month) and stable areas (suitable for > 3 months) between the historical period (1985-2005) and each time period. Results for each weighted ensemble are presented.

	Total Area (Mn km ²)			Stable Area (Mn km ²)		
	Q PE	PR	Q PE RH	Q PE	PR	Q PE RH
RCP 2.6						
2006-2025	0.499	0.235	0.275	0.087	0.139	0.096
2026-2050	0.171	-0.014	0.006	-0.067	-0.220	-0.107
2051-2075	0.264	0.177	0.217	-0.043	-0.055	-0.067
2076-2100	0.151	-0.043	0.055	-0.290	-0.299	-0.310
RCP 6.0						
2006-2025	0.229	0.203	0.246	-0.017	0.070	-0.035
2026-2050	-0.003	-0.003	-0.032	-0.072	0.035	-0.081
2051-2075	-0.009	-0.052	0.035	-0.058	-0.212	-0.064
2076-2100	-0.110	-0.414	-0.145	-0.307	-0.339	-0.316
RCP 8.5						
2006-2025	-0.145	0.258	-0.264	-0.180	-0.133	-0.162
2026-2050	-0.206	0.067	-0.429	-0.235	0.026	-0.357
2051-2075	-0.267	-0.290	-0.574	-0.533	-0.591	-0.620
2076-2100	-0.423	-0.881	-0.881	-1.133	-1.075	-1.151

Table S9. Areas (in Mn km²) experiencing an increase and decrease in malaria suitability in future ensemble projections since the historical period (1985-2005). Note that the net value includes areas with an average change of <1 month and only where the signal to noise ratio >0.5.

	Q PE			PR			Q PE RH		
	Increasing	Decreasing	Net	Increasing	Decreasing	Net	Increasing	Decreasing	Net
RCP 2.6									
2006-2025	0.710	0.328	0.383	3.658	2.907	0.751	0.971	0.493	0.478
2026-2050	0.771	0.991	-0.220	3.188	4.087	-0.899	0.890	1.791	-0.901
2051-2075	0.942	0.901	0.041	3.840	3.669	0.171	1.070	1.307	-0.238
2076-2100	0.878	1.356	-0.478	3.246	4.936	-1.690	0.910	2.220	-1.310
RCP 6.0									
2006-2025	0.661	0.530	0.130	3.551	2.954	0.597	0.803	0.580	0.223
2026-2050	0.733	0.806	-0.072	3.698	2.875	0.823	0.809	0.977	-0.168
2051-2075	1.368	1.403	-0.035	4.649	4.255	0.394	1.359	2.017	-0.658
2076-2100	1.672	2.939	-1.267	4.539	5.380	-0.841	1.664	4.168	-2.504
RCP 8.5									
2006-2025	2.206	2.539	-0.333	4.177	2.890	1.287	2.510	3.307	-0.797
2026-2050	2.380	3.206	-0.826	4.452	3.646	0.806	2.345	4.052	-1.707
2051-2075	2.238	4.748	-2.510	4.635	5.542	-0.907	1.980	6.139	-4.159
2076-2100	2.603	6.823	-4.220	4.058	7.516	-3.458	2.380	7.965	-5.585

Table S10. Increase and decrease in malaria suitability ‘month-areas’ (in millions mo km²) between future projections and the historical period (1985-2005). Areas used to calculate totals in Table S9 are multiplied by the number of months change in malaria climatic suitability to provide a clearer indication of the magnitude of changes.

	Q _{PE}			PR			Q _{PE} RH		
	Increasing	Decreasing	Net	Increasing	Decreasing	Net	Increasing	Decreasing	Net
RCP 2.6									
2006-2025	0.532	-0.297	0.235	2.333	-1.726	0.607	0.668	-0.264	0.404
2026-2050	0.742	-0.913	-0.171	2.101	-2.472	-0.371	0.746	-1.014	-0.268
2051-2075	0.816	-0.975	-0.159	2.439	-2.339	0.100	0.874	-0.709	0.165
2076-2100	0.770	-1.325	-0.555	2.127	-3.155	-1.027	0.748	-1.270	-0.522
RCP 6.0									
2006-2025	0.481	-0.442	0.039	2.123	-1.839	0.284	0.555	-0.310	0.245
2026-2050	0.572	-0.693	-0.120	2.226	-1.548	0.678	0.587	-0.535	0.052
2051-2075	1.296	-1.672	-0.377	3.130	-2.574	0.557	1.239	-1.249	-0.010
2076-2100	1.874	-3.974	-2.100	3.464	-3.455	0.009	1.725	-3.061	-1.336
RCP 8.5									
2006-2025	1.926	-2.374	-0.448	2.709	-1.738	0.971	1.794	-2.761	-0.967
2026-2050	2.242	-3.623	-1.381	2.829	-2.104	0.725	1.827	-3.553	-1.726
2051-2075	2.458	-6.646	-4.188	3.914	-3.632	0.283	2.094	-5.817	-3.723
2076-2100	3.078	-13.005	-9.927	4.040	-6.413	-2.372	2.816	-10.756	-7.940

Table S11. Total populations (Mn) and children under 5 years old within areas of climatic suitability for malaria transmission in Africa (>1 month) and climatic suitability for stable malaria transmission (>3 months). Population figures are calculated from the mid-point of each time period using the UN Medium Variant figures, except 2006-2025 which use Worldpop gridded estimates.

	LTS >1 month			LTS >3 months		
	Q PE	PR	Q PE RH	Q PE	PR	Q PE RH
Total Population (Mn)						
RCP 2.6						
2006-2025	1060.8	990.4	1034.4	866.4	826.9	845.5
2026-2050	1803.0	1716.2	1765.8	1510.5	1414.3	1466.7
2051-2075	2714.8	2610.7	2663.0	2251.7	2118.0	2183.8
2076-2100	3410.6	3292.9	3345.9	2821.1	2632.1	2705.0
RCP 6.0						
2006-2025	1059.6	994.2	1031.8	850.0	807.7	821.5
2026-2050	1794.9	1710.0	1747.9	1479.8	1401.8	1447.5
2051-2075	2704.9	2597.6	2658.7	2247.0	2122.1	2177.4
2076-2100	3407.6	3276.6	3337.3	2805.3	2678.1	2698.5
RCP 8.5						
2006-2025	1083.0	997.9	1039.5	850.7	814.2	832.5
2026-2050	1860.1	1717.5	1791.3	1485.4	1417.7	1434.5
2051-2075	2780.0	2601.2	2717.8	2196.2	2071.0	2119.3
2076-2100	3424.6	3229.0	3325.7	2678.4	2515.3	2548.1
Under 5s (Mn)						
RCP 2.6						
2006-2025	172.0	163.6	168.2	143.9	137.7	140.2
2026-2050	233.7	226.2	229.6	198.2	186.2	192.3
2051-2075	262.8	255.9	258.4	219.9	207.3	212.5
2076-2100	258.9	252.2	254.4	214.4	200.9	204.9
RCP 6.0						
2006-2025	171.9	164.0	167.8	140.7	134.5	135.7
2026-2050	233.0	225.6	228.0	195.5	185.4	190.4
2051-2075	262.1	255.0	258.1	219.3	206.9	212.1
2076-2100	258.6	251.0	253.4	212.6	202.7	203.7
RCP 8.5						
2006-2025	175.0	164.5	169.3	142.2	135.9	138.9
2026-2050	238.7	226.5	232.0	196.9	187.1	189.7
2051-2075	267.1	255.0	261.7	214.0	201.0	205.6
2076-2100	259.0	246.9	251.7	202.0	189.0	191.5