

# Predictive mapping of mosquito distribution based on environmental and anthropogenic factors in Taita Hills, Kenya

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## ABSTRACT

Mosquitoes are vectors for numerous pathogens, which are collectively responsible for millions of human deaths each year. As such, it is vital to be able to accurately predict their distributions, particularly in areas where species composition is unknown. Species distribution modeling was used to determine the relationship between environmental, anthropogenic and distance factors on the occurrence of two mosquito genera, *Culex* Linnaeus and *Stegomyia* Theobald (syn. *Aedes*), in the Taita Hills, southeastern Kenya. This study aims to test whether any of the statistical prediction models produced by the Biomod2 package in R can reliably estimate the distributions of mosquitoes in these genera in the Taita Hills; and to examine which factors best explain their presence. Mosquito collections were acquired from 122 locations between January–March 2016 along transects throughout the Taita Hills. Environmental-, anthropogenic- and distance-based geospatial data were acquired from the Taita Hills geo-database, satellite- and aerial imagery and processed in GIS software. The Biomod2 package in R, intended for ensemble forecasting of species distributions, was used to generate predictive models. Slope, human population density, normalized difference vegetation index, distance to roads and elevation best estimated *Culex* distributions by a generalized additive model with an area under the curve (AUC) value of 0.791. Mean radiation, human population density, normalized difference vegetation index, distance to roads and mean temperature resulted in the highest AUC (0.708) value in a random forest model for *Stegomyia* distributions. We conclude that in the process towards more detailed species-level maps, with our study results, general assumptions can be made about the distribution areas of *Culex* and *Stegomyia* mosquitoes in the Taita Hills and the factors which influence their distribution.

## 1. Introduction

Environmental and anthropogenic disturbances such as climate change, urbanization and deforestation are crucial factors in the distribution of pathogen vectors and the emergence of diseases that they transmit (Crowl et al., 2008). This is already evident as the global rise in human population densities, and correlated mean temperatures leading to land-use changes, has created new suitable breeding sites for mosquitoes (Roiz et al., 2011; Campbell et al., 2015). Geographic

information system (GIS) and species distribution modeling (SDM) approaches have been used to understand these connections and their implications for the spread of invasive species, particularly with regard to northern regions (Neteler et al., 2011). Other important areas, however, remain unexplored, including biodiversity rich but increasingly fragmented parts of Africa, where infectious diseases can be highly prevalent but access to health care is limited (Bhutta et al., 2014). As a result, tools such as GIS and SDM that can estimate and predict infectious disease risks have the potential to optimize the use of

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limited resources and improve public health outcomes.

Due to their ability to act as vectors for a suite of pathogens, mosquitoes (Diptera, Culicidae) are among the most economically and socially important taxa on the planet (The World Health Organization (WHO, 2017)). Two notable genera are *Culex* Linnaeus and *Stegomyia* Theobald (following the classification of Reinert et al., (2009)). *Culex* is a large genus of mosquitoes, with 769 species in 26 subgenera (Mosquito taxonomic inventory (MTI, 2017)), accounting for 21.6% of all mosquito species worldwide. Species of *Culex* have an almost worldwide distribution from the tropics to cool temperate regions, but do not extend into extreme northern latitudes (Mosquito taxonomic inventory (MTI, 2017)). While species from other subgenera are known to vector pathogens, subgenus *Culex* (*Culex*) contains many of the significant human vector species, including those demonstrated to transmit West Nile fever virus, Rift Valley fever virus and Japanese Encephalitis virus (Mosquito taxonomic inventory (MTI, 2017; The World Health Organization (WHO, 2017)). *Stegomyia* is a moderately sized mosquito genus comprising 128 species, and distributed in the Afrotropical, Oriental and Australasian regions (Mosquito taxonomic inventory (MTI, 2017)). Following human dispersal, at least two species, *St. aegypti* and *St. albopicta*, are also present in Neotropical, Nearctic and Palearctic regions (Mosquito taxonomic inventory (MTI, 2017; Paupy et al., 2009)). Member species are competent vectors of yellow fever virus (Huang, 1986), dengue virus serotypes 1–4, Zika virus, and Chikungunya virus, among others (Huang, 1990; Mosquito taxonomic inventory (MTI, 2017)).

SDM helps to understand the relationship between wildlife and their environments, to estimate their distributions where empirical data is missing, and to forecast range expansions in the face of environmental change. SDM is used in decision-making for a range of global challenges, such as informing protected areas for wildlife conservation (Guisan et al., 2013), the distribution potential for invasive species (Václavík & Meentemeyer, 2009), and investigating the effects of climate change on species that carry pathogens of public health concern such as mosquitoes (Dukes et al., 2009). Species distribution modeling, therefore, provides a means to understand and predict vector responses to changing climate patterns in dynamic and fragmented environments.

The purpose of this study is to test whether any of the statistical prediction models produced by the Biomod2 package in R, can reliably estimate the distributions of mosquitoes in genera *Culex* and *Stegomyia* in the Taita Hills; to examine which factors best explain their presence; and to create maps of their predicted occurrence. The target area, Taita Hills in rural south-eastern Kenya, is an ecologically diverse area with little previous mosquito research, strong variability in rainfall and a rapidly growing human population. While machine-learning techniques such as maximum entropy modeling is often used to understand mosquito distributions (Mughini-Gras et al., 2014; Sallam et al., 2016), the employed R package, the Biomod2 (Thuiller et al., 2016), has important benefits for estimating mosquito distributions. This is due to its ability to provide improved simulations across initial conditions, test multiple model classes and parameterizations, and include unlimited number of boundary conditions (Thuiller et al., 2009).

## 2. Methods

### 2.1. Study area and mosquito collections

The Taita Hills (03°20'S, 38°15'E) in Taita-Taveta county, south-eastern Kenya (Fig. 1.) is a range of peaks varying in altitude from 600 to 2200 m above sea level (a.s.l.) (Erdogan et al., 2011). The study area of 286 km<sup>2</sup> ranges from the Taita Hills at 1800 m a.s.l. to the surrounding lowlands at 900 m a.s.l., as outlined in Fig. 1.

Kenya has two annual wet and dry seasons, with the total rainfall varying across the country (Kaplan et al., 1976; Fig. 2.). The Taita Hills region receives on average 1330–1910 mm/annually, which contributes to the formation of many suitable mosquito breeding habitats

(Erdogan et al., 2011).

Mosquito collections were acquired at over 122 locations across the region between January and March 2016. This figure includes 'sub-locations', which fell within 500 m of the primary target location, and repeat collections from the same locations across the study time (Fig. 1.). Of these 122 attempts, mosquitoes were present in 107 collections and absent from 15. Mosquito collections started after the rainy season on January 25<sup>th</sup>, 2016 and continued until March 16<sup>th</sup>, 2016. Collections did not follow transects up to February 12<sup>th</sup>, since the goal at this time was to focus on human-biting species around human dwellings. From February 13<sup>th</sup> until the end of the collection period a stratified sampling scheme was applied based on the road network. Each main road from the lowlands to the highest reachable locations was used to collect mosquitoes on 100 m elevation intervals. This sampling method enabled reliable spatial extrapolation and interpolation, as systematic samples minimize the distance from any point in the study area to a sample point (Franklin and Miller, 2010).

Collection locations and sub-locations were mainly around human dwellings close to roads, but collections were also made in forest fragments and in croplands. Adults were collected using commercially available Prokopacks (The John W. Hock Company, Gainesville, USA) or by using CDC Miniature Light Traps (The John W. Hock Company, Gainesville, USA). Immature life stages (larvae, pupae and eggs) were collected from stagnant water using a 1 L plastic dipper, a fine meshed aquarium net or a turkey baster. All water sources were considered, including septic tanks, discarded tires, tree cavities, leaf axils and other items filled with rain water, such as discarded artificial containers or water tanks. At each collection location, the GPS point was recorded using a Garmin Map 64S handset. Collections were initially sorted in a field laboratory and stored in ethanol or RNA-later to preserve the RNA and DNA for future studies. Genus-level identifications were made in Finland using suitable identification keys (Service, 1991; Huang, 2001, 2004).

### 2.2. Explanatory environmental, anthropogenic and distance variables

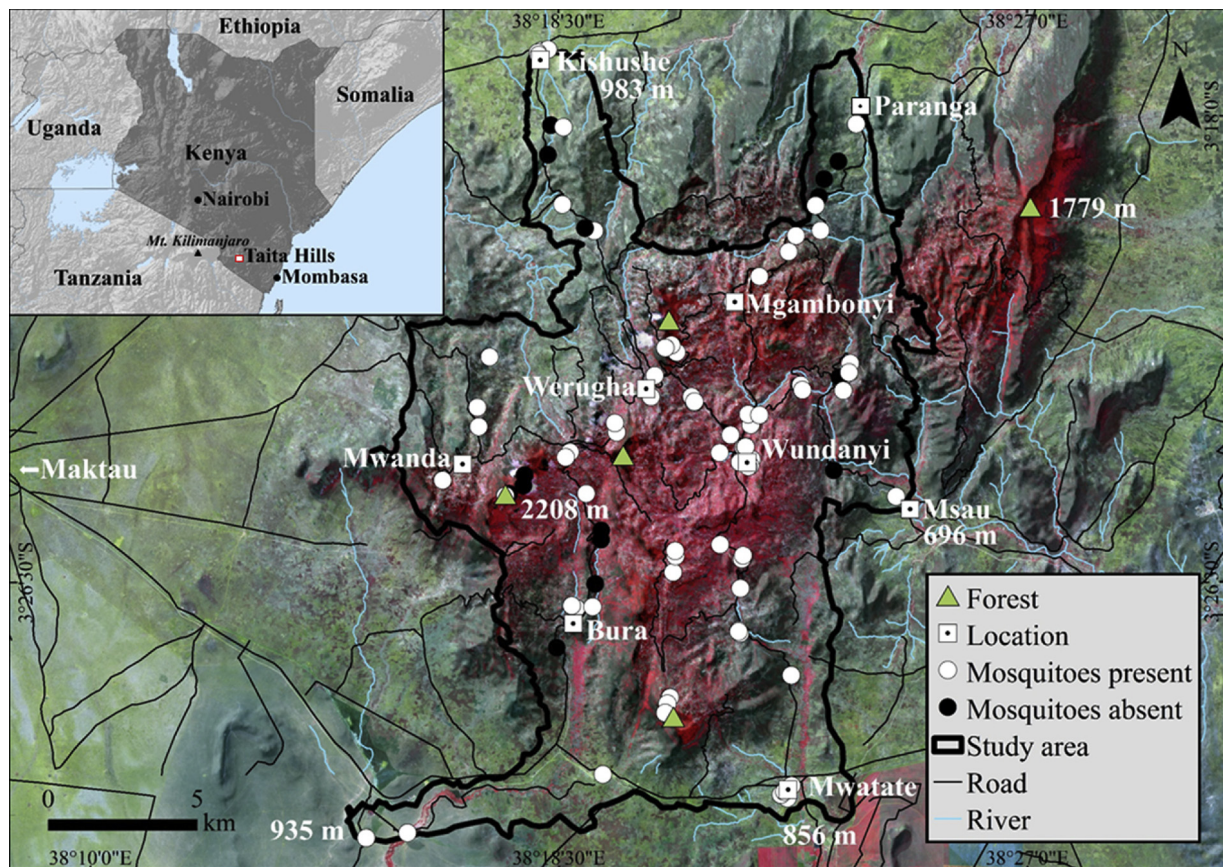
The selection of explanatory variables to predict mosquito distributions was based on those used in existing literature. Environmental and anthropogenic data for the Taita Hills region were obtained directly from satellite imagery and airborne digital data, or by deriving them from the satellite imagery in ArcGIS (Esri, Redlands, CA, USA, v.10.3.1) (Table 1). Distance variables were created from an existing Taita Hills geo-database. For the distance to buildings calculations, existing airborne laser scanning (ALS) data was used. In this data set, buildings were classified from point cloud data that were derived from 2014 to 2015 Taita Hills ALS flights (Adhikari et al., 2017). LAsTools software (rapidlasso GmbH) was used to classify ground, building and vegetation returns.

Normalized difference vegetation index (NDVI) is a measure of plant greenness, ranging between -1 to 1 (Tucker, 1979), and was used to approximate land cover type. Mean precipitation, mean temperature and mean relative humidity from January - March was calculated for each collection location in ArcGIS. Mean monthly precipitation was derived from long-term precipitation grids between 1987 and 2005 (Hutchinson, 1991; Erdogan et al., 2011). A digital elevation model (DEM) was used to calculate the elevation, slope and mean monthly solar radiation in ArcGIS, and the values were extracted to each mosquito collection location.

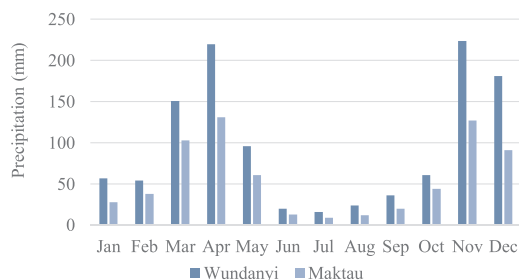
### 2.3. Data analysis and modeling

All the GIS datasets including environmental and other attributes were set to the same spatial extent, geographic coordinate system (WGS 1984 UTM Zone 37S) and resolutions (20 m x 20 m). The compiled data consisted of presence-absence data and explanatory variables for each sampling location. We predicted the distribution of *Culex* and *Stegomyia*





**Fig. 1.** The Taita Hills and sampling locations in southeastern Kenya. Sampling was conducted along the main roads, from the surrounding lowlands to mountain peaks.



**Fig. 2.** Monthly average rainfall characteristics for Wundanyi town in the Taita Hills and Maktau village in the lowland plains. Weather data was collected between 1982 and 2012, with a resolution of 30 arc seconds (Climate-Data.org, 2018).

mosquitoes using the Biomod2–platform (version 3.3–7) in R software (version 3.0.3; R Development Core Team, 2013). Prior to prediction modelling, multicollinearity of the variables was tested using Pearson correlation coefficients in R, which resulted in the exclusion of four variables. Models were processed mainly using the default settings of the Biomod2. The following eight predictive modeling techniques were used: generalized linear models (GLM), generalized additive models (GAM), classification tree analysis (CTA), artificial neural networks (ANNs), multivariate adaptive regression splines (MARS), general boosting method (GBM), random forest (RF) and maximum entropy (Maxent). Flexible discriminant analysis (FDA) and surface range envelope (SRE) were excluded due to their methodological weaknesses (Hastie et al., 1994).

The original data set of 122 collection locations was randomly divided into model training (70%,  $n = 85$ ) and model evaluation sets (30%,  $n = 37$ ) (split-sample approach: Guisan and Zimmermann,

2000). Models were built using the training set, and the models were validated using the model evaluation set. The area under the curve (AUC) of a receiver operating characteristic (ROC) plot was produced based on each model to estimate the predictive power of the model (Fielding and Bell, 1997), to assess the agreement between the presence-absence records and the predictions. The model types with highest AUC values ( $0.7 < \text{AUC} < 1.0$ ) and statistically significant variables ( $p \leq 0.05$ ) were used to conduct the predictive modeling (Drew et al., 2011). In order to receive the highest possible prediction accuracy values, we compared the different sets and orders of predictor variables. Depending on their strength of relationship to the response variable, we selected the best combination of predictors to include in a model. We compared the variable contributions in the models to define the most powerful variables, and their relative magnitude. Spatial autocorrelation (SAC) of the predictor variables was measured using Moran's Index (Moran, 1950).

### 3. Results

A total of 3130 mosquitoes were collected from 107 locations across the Taita Hills region. The majority (~2600 mosquitoes from 73 locations), belonged to genus *Culex* and included at least three subgenera: *Culex* (*Culex*), *Cx.* (*Culiciomyia*) and *Cx.* (*Eumelanomyia*). *Culex* mosquitoes were common across the whole study area, including around human settlements and in forests. *Stegomyia* mosquitoes were the second most abundant genus in the collections, (~180 individuals from 28 locations). Larvae of both genera were found in water tanks and small ponds, and adults were collected from houses along the roadsides and in villages. *Stegomyia aegypti* larvae were especially common in water tanks in the villages of Paranga and Kishushe in the Taita Hills north, and also in car tires in Mwatate village. Both genera were present

**Table 1**  
Value ranges and sources for explanatory variables used in species distribution modeling.

Environmental, anthropogenic or distance factor	Min.	Max.	Avg.	Data source
Distance to houses (m)	0	1270	52	Distance to houses was determined from the building data computed from airborne laser scanning (ALS) data (Adhikari et al., 2017), modified by digitizing more houses in the study area using QGIS, and calculating Mean Euclidean distance to houses.
Elevation (m)	694	2079	1330	Mean elevation was derived from digital elevation model (DEM). DEM was obtained from scanned Survey of Kenya 1:50 000 scale topographic map from 1991, in which a 20 m planimetric resolution DEM was interpolated from 50 feet interval contours (Clark and Pellikka, 2005).
Distance to roads (m)	0	927	127	Road data was digitized in 2004 from Survey of Kenya 1:50 000 scale topographic map from 1991. (Broberg et al., 2004). Modified by calculating Mean Euclidean distance to roads using ArcGIS.
Mean precipitation (mm)	20	113	47	Mean precipitation was obtained from long-term mean precipitation grids, which were interpolated on to a 20 m resolution grid from monthly available meteorological data and surrounding areas between 1987 and 2005, obtained from Kenya Meteorological Department using ANUSPLIN (Australian National University Splines) software (Hutchinson, 1991; Erdogan et al., 2011).
Mean radiation (kWh/m <sup>2</sup> )	176	228	216	Mean radiation was received on a given surface area in a given time (kWh/m <sup>2</sup> ), was calculated from the DEM using ArcGIS Area Solar Radiation tool (Esri, Redlands, CA, USA).
Mean relative humidity (%)	71	94	77	Mean relative humidity data was created based on data logger observations between April 2013 and May 2014 (Virtanen, 2015).
Mean temperature (°C)	15	25	21	Mean temperature data was created based on data logger observations between April 2013 and May 2014 (Virtanen, 2015).
Slope (°)	0	43	11	Slope degree was derived from DEM using Slope function in ArcGIS.
Human population density (persons/km <sup>2</sup> )	0	9090	1260	Human population density was estimated from ALS buildings data and from a random non-stratified household survey (n = 100) carried out in October 2006, which found an average of 6 persons per dwelling in the area (Siljander et al., 2011). The data was modified by digitizing more houses in the study area using QGIS.
Normalized Difference Vegetation Index (NDVI)	−0.4	0.2	−0.2	NDVI was derived from a Sentinel-2 A MSI Level-1C satellite image from 8 October 2016, downloaded from the Sentinel's Scientific DataHub (ESA, 2015).

in lowland and upland areas, including elevations up to 1900 m.

The environmental predictors, elevation, mean precipitation, mean temperature and mean relative humidity, were positively or negatively correlated with each other ( $r \geq 0.9$  or  $r \leq -0.9$ ), as were slope and mean radiation ( $r \geq -0.75$ ); therefore, only one of each variable group was retained in the final model. Normalized difference vegetation index, distance to roads, distance to houses, and human population density did not result in high correlations, and were retained in the model process. The combination of non-correlating variables, which resulted in highest evaluation values, was run to obtain reliable estimations for *Culex* and *Stegomyia* (Appendix A). Mean temperature, mean radiation, elevation, slope and NDVI were influential factors in the models (Appendix B). Human population density had the greatest effect on the distribution of both *Culex* and *Stegomyia* in all models, but its importance in each model varied (Appendix B). Moderately explaining variables - NDVI, slope, distance to roads and elevation - were not ranked consistently in the models. The model that best explained *Culex* mosquito distribution included the variables slope, population density, NDVI, distance to roads, and elevation. Mean radiation, NDVI, human population density, distance to roads, and mean temperature were most influential in explaining the distribution of *Stegomyia* mosquitoes.

Spatial autocorrelation (SAC) of the predictor variables was measured using Moran's Index (Moran, 1950) (Appendix C). For *Culex*, human population density and elevation were highly spatially autocorrelated (Moran's  $I \geq 0.8$ ) for short distances ( $p < 0.05$ ) but not at longer distances. Human population density and mean temperature were highly spatially autocorrelated at short distances, but not at longer distances for *Stegomyia*. The models with highest AUC value and significant p-values for *Culex*, were GAM (AUC = 0.791) and MARS (AUC = 0.809) (Table 2). Altogether, six of the eight models provided reliable estimates for *Culex* mosquito distribution (AUC > 0.7). For *Stegomyia*, only two of the eight models resulted in evaluation values considered reliable (AUC > 0.7); GBM (AUC = 0.708) and RF models (AUC = 0.708).

For the predictive models (described below), we focus on the GAM model for *Culex* and the RF model for *Stegomyia* due to their ability to explain the data reliably and produce the best explaining variable contributions for this study. According to the GAM model, the probability of *Culex* presence was high ( $\geq 80\%$ ) in locations with NDVI-values ranging from −0.4 to 0.2 (Fig. 3.). The likelihood of *Culex* presence remained high when roads were within 500 m, and at elevations between 800 and 2000 m, in locations with moderate slope angles ( $0^\circ$ – $35^\circ$ ), and with population densities of 500–6000 people/km<sup>2</sup>.

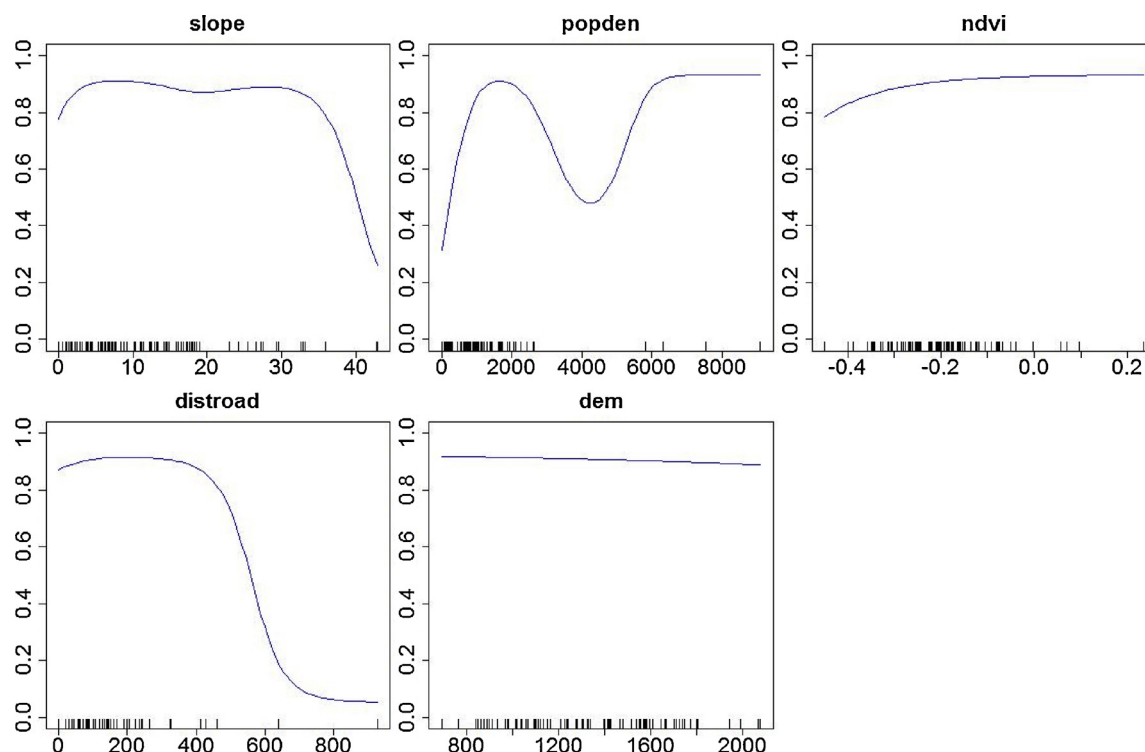
According to the RF model, the presence of *Stegomyia* was highest in locations with solar radiation levels  $\geq 230$  kWh/m<sup>2</sup> (Fig. 4.). They favored temperatures between 15 °C and 20 °C, and over 23 °C, and locations with poor (NDVI  $\leq -0.2$ ) or moderate vegetation (NDVI  $\geq 0.2$ ). All distances between 0 and 600 m from roads resulted in moderate probabilities of *Stegomyia* presence.

The prediction model risk maps (Figs. 5A and B) show that village areas and forests, including elevations above 1500 m in the Taita Hills,

**Table 2**

Area under the curve (AUC) values of all models used to estimate the presence of *Culex* and *Stegomyia* mosquitoes.

<i>Culex</i>		<i>Stegomyia</i>	
Model	AUC	Model	AUC
GLM	0.730	GLM	–
GAM	0.791	GAM	0.643
GBM	0.750	GBM	0.708
CTA	0.620	CTA	0.616
ANN	0.764	ANN	0.612
MARS	0.806	MARS	–
RF	0.729	RF	0.708
Maxent	0.585	Maxent	0.690



**Fig. 3.** Response curves for *Culex* mosquito estimations based on the GAM model. The Y-axis represents the probability of presence, and the black marks along the x-axis represent empirical observations.

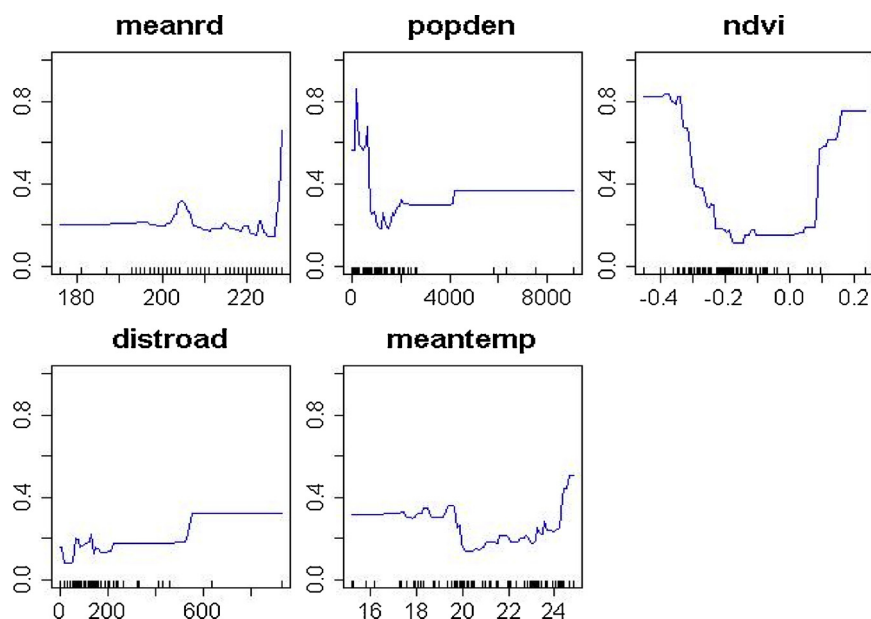
were most suitable for *Culex* mosquitoes (80–100%), whereas the likelihood of their presence was reduced in lowland plateaus, apart from Mwatate village in the South (0–20%). Conversely, the likelihood of *Stegomyia* mosquitoes was high in villages on the plateaus and close to roads (Fig. 5.B.). Areas with high *Stegomyia* mosquito probability were sporadically dispersed.

#### 4. Discussion

We show that the Biomod2 ensemble platform in R, was able to

predict the distribution of *Culex* and *Stegomyia* mosquitoes in the Taita Hills in Kenya. These two genera were found to overlap in geographical range, but also to be common in different areas based on environmental and anthropogenic factors. *Culex* and *Stegomyia* mosquitoes contain member species that are well-known vectors of significant human pathogens (Lounibos, 2002). This approach can be applied to better understand how these vectors, and the infections they carry, vary across space and time, and later, this may be extrapolated to species-level studies and to other understudied areas in rural Africa.

The majority of the models utilized herein produced high AUC



**Fig. 4.** The response curves of predictors for *Stegomyia* mosquito estimations in RF model. The Y-axis represent the probability of presence, and the black marks along the x-axis represent empirical observations.



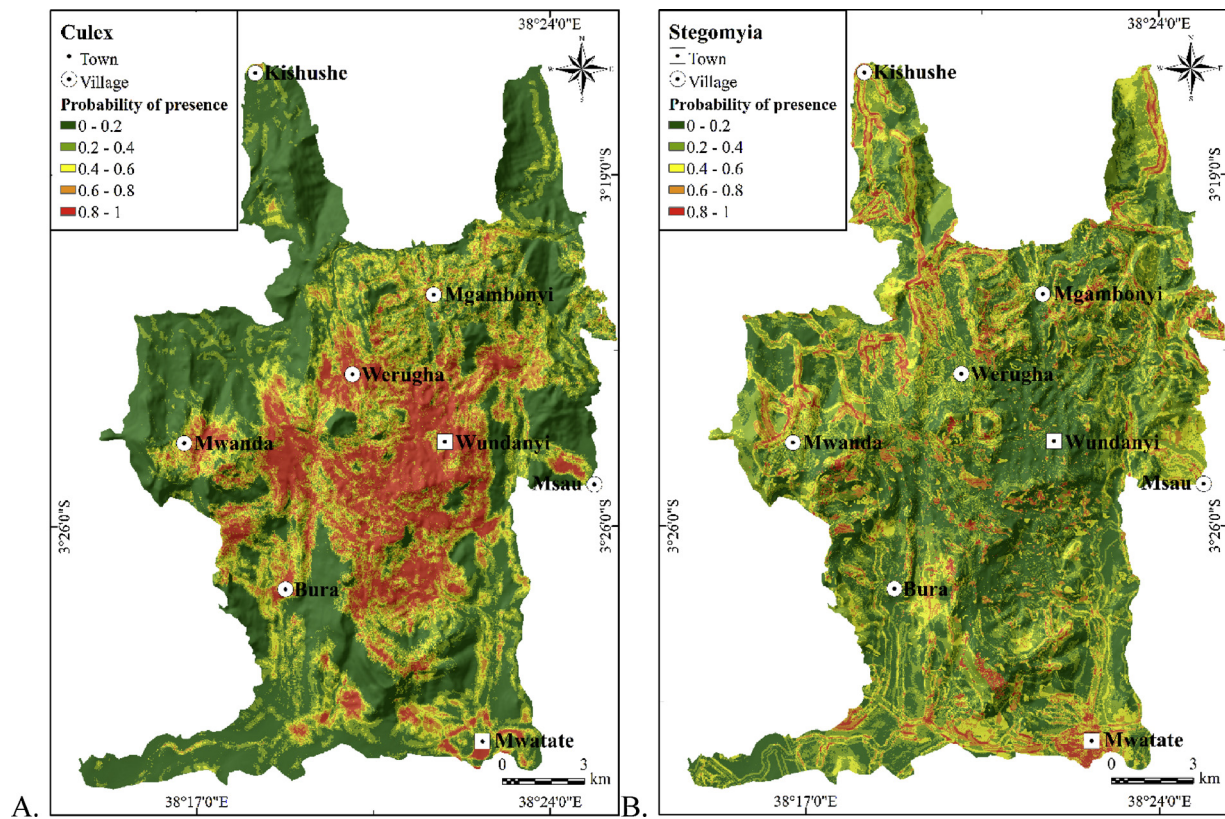


Fig. 5. Predictive risk maps for *Culex* and *Stegomyia* mosquito presence over the study area in the Taita Hills. A. The probability of *Culex* presence was highest in the central part based on the GAM model. B. The probability of *Stegomyia* presence was highest in plateau villages and fragmented areas based on a RF model.

values for *Culex* mosquitoes, indicating reliable estimations. We concentrated on the GAM model because it resulted in high AUC values with relevant variable contributions having an automated approach to identify non-linear relationships, and it may result in more accurate predictions than other model types (Yee and Mitchell, 1991). For *Stegomyia* estimations, we focused on the RF model because of its advantages over other models, including high classification accuracy and the ability to model interactions among predictors (Cutler et al., 2007). However, some uncertainties exist concerning *Stegomyia* estimations due to a higher number of absences (94 collections) than presences (28 collections) in the sampling data. Environmental, anthropogenic and distance variables were spatially autocorrelated to some extent, which may, among other things, affect the precision of coefficients (Diniz-Filho et al., 2003).

In our study, the spatial resolution of environmental data was 20 m. This resolution may have produced some bias to the study results because mosquitoes are also breeding e.g. in tree holes and water tanks in different microclimates. The lack of very high-resolution environmental data (such as the resolutions of 2 m and 5 m) is a general problem and not limited just to the Taita Hills. In addition, increasing the resolution of environmental data by deriving them from a high-resolution environmental data does not always improve the species distribution models (Pradervand et al., 2014). Instead, very high-resolution environmental predictors should be produced by taking into account more local field measurements such as fine environmental mapping or in-situ measurements (Pradervand et al., 2014; Lembrechts et al., 2018). The potential solution for the lack of microclimate data could be the use of unmanned aerial vehicles (UAVs) in order to map the sampling areas in real time, and to produce higher resolution environmental data of even 3 cm (Fornace et al., 2014; Anderson and Gaston, 2013). Furthermore, there certainly exist other explanatory factors affecting the mosquito distributions which were not included in the modelling process including landscape fragmentation indicators. Fragmentation variables,

such as distance from forest patch, patch size, distance from patch edge and the landscape metric of PPU (patches per unit) have resulted significant statistical relations with mosquito distributions (Richman et al., 2018; Reiter and LaPointe, 2007). For example, the use of FRAGSTATS—a spatial pattern analysis program may add value for our future mosquito habitat studies (McGarigal et al., 2002).

Consistent with previous research from other global locations (Sallam et al., 2017; Ding et al., 2017), environmental and anthropogenic variables were important determinants of *Culex* and *Stegomyia* mosquito distribution in the Taita Hills. For example the response curve for slope (Fig. 3.) shows high probability for *Culex* presence for slopes less than 35°. In the Taita Hills, dwelling units are mainly absent on slopes greater than 35° (Siljander et al., 2011). High probability for *Culex* presence were found with high NDVI values caused by strongly reflecting orchard trees and croplands adjacent to dwelling units in the Taita Hills. Specifically, we found that locations with high population densities, short distances to roads, and gentle slope angles positively influenced *Culex* presence. Residential and urban areas are often recognized as important for *Culex* distributions (Reiter and LaPointe, 2007; Conley et al., 2014). Rich and poor vegetation and all elevations were suitable for *Culex* mosquito presence, strengthening previous findings of its widespread distribution (Mosquito taxonomic inventory (MTI, 2017). However, we cannot rule out some effects of potential sample biases (e.g. time of the day and traps involved) which may have affected the mosquito spectrum captured. Sampling bias may also have been the reason for an unexpected response curve for population density (*popden*) at values from ca. 4000 to 5000 (Fig. 3).

Our study suggests, that *Stegomyia* mosquitoes prefer locations with lower human population densities, higher temperatures and solar radiation, and poor or moderate vegetation. This finding somewhat contradicts the notion that *St. aegypti* distribution is linked to growing human population (Fatima et al., 2016), but is consistent with historic records of the variety of *Stegomyia* species suggesting that most

members of the *Stegomyia* genus are forest dwelling, with only some adapted to breeding close to human habitation (Powell and Tabachnick, 2013). Since our collections included several different *Stegomyia* species - with *St. aegypti* being the only species collected from around human habitation, and other species being restricted to the forest - the model findings are consistent with these historical reports (Powell and Tabachnick, 2013). *Stegomyia* favored locations with intermediate and high temperatures, confirming the argument that *Stegomyia* mosquitoes have temperature-based limits to survival (Brady et al., 2013). Overall, anthropogenic and distance factors appeared to be more important than environmental drivers for the distribution of both *Culex* and *Stegomyia* mosquitoes. We note that other variables, not considered in this study, may also affect the distribution of *Culex* and *Stegomyia* mosquitoes and that the genus-level pooling of the species will mask species distribution determinants, which will be targeted in future work with larger and more detailed datasets.

## 5. Conclusions

Our results affirm the utility and reliability of the Biomod2 package in R as a valid modeling method for species distributions, resulting in insights into vector-ecological interactions. Such work will be further refined with species-level identification and screening for arboviruses. With our study results, general assumptions can be made about the

distribution of mosquitoes belonging to *Culex* and *Stegomyia* genera in the Taita Hills and the factors that influence their distribution. High population densities, short distance to roads and gentle slope angles were associated with the occurrence of *Culex* mosquitoes. For *Stegomyia* mosquitoes, on the other hand, low human population densities, high temperatures and solar radiations as well as poor or moderate vegetation were suitable factors. Together, these results have implications for applying the approach for identifying risk areas and optimizing the use of limited resources for mitigation strategies.

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**Appendix A. Multicollinearity of the variables included in the models. All statistical significance values (p-values), are marked as follows: statistically highly significant \*\*\* = < 0.001, statistically significant \*\* = < 0.01, statistically significant \* = < 0.05 and ns = not statistically significant**

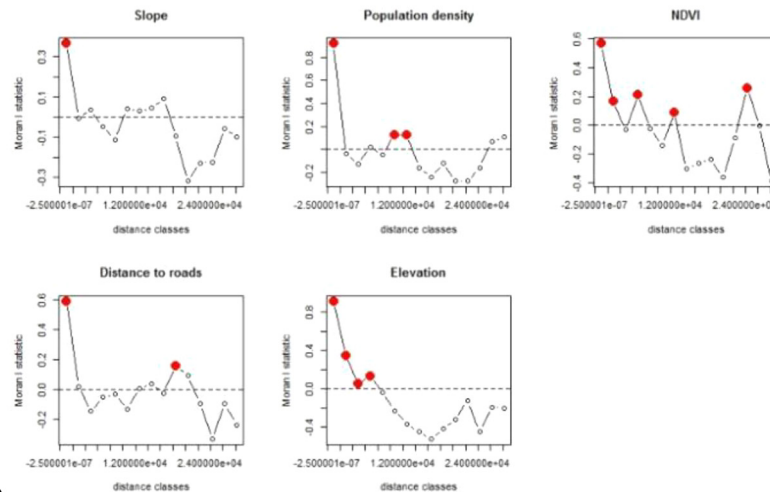
Culex					
Variable	Slope	NDVI	Population density	Distance to roads	Elevation
Slope		0.115 ns	-0.294**	0.172ns	0.279**
NDVI	0.115 ns		-0.226*	0.391***	0.557***
Population density	-0.294**	-0.226*		-0.26**	-0.313***
Distance to roads	0.172ns	0.391***	-0.26**		0.143ns
Elevation	0.279**	0.557***	-0.313***	0.143ns	
Stegomyia					
Variable	Mean radiation	NDVI	Population density	Distance to roads	Mean temperature
Mean radiation		-0.04ns	0.312***	-0.12 ns	0.10ns
NDVI	-0.04ns		-0.226*	0.391***	-0.543***
Population density	0.313***	-0.226*		-0.26**	0.286**
Distance to roads	-0.12 ns	0.391***	-0.26**		-0.12 ns
Mean temperature	0.10ns	-0.543***	0.286**	-0.12 ns	

## Appendix B. Variable importance presented in each model for *Culex* and *Stegomyia* estimates

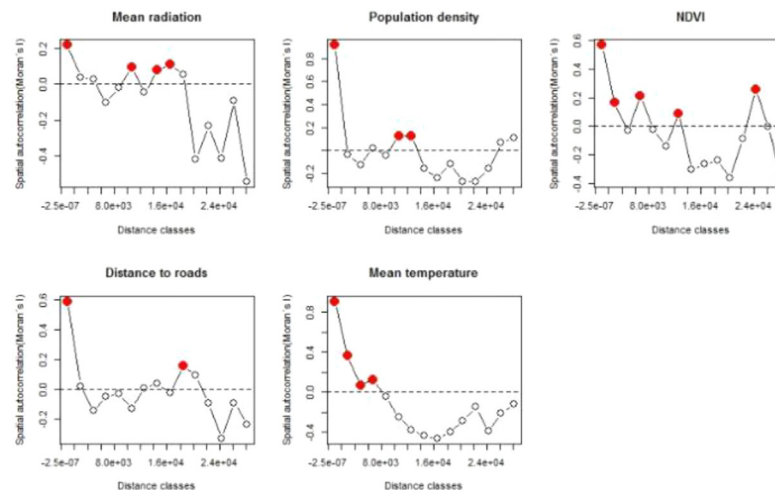
Culex								
Variable	GLM	GAM	GBM	CTA	ANN	MARS	RF	Maxent
Slope	0	0.147	0.078	0	0.147	0.385	0.200	0.432
Population density	0.658	0.764	0.678	1	0.773	0.702	0.429	0.513
NDVI	0.261	0.165	0.088	0	0	0	0.114	0.382
Distance to roads	0	0.179	0.039	0	0.092	0	0.064	0.453
Elevation	0	0.028	0.006	0	0.097	0	0.030	0.436
Stegomyia								
Variable	GLM	GAM	GBM	CTA	ANN	MARS	RF	Maxent
Mean radiation	-	0.041	0	0	0.289	-	0.059	0.176
Population density	-	0.575	0.680	0.966	1	-	0.206	0.357
NDVI	-	0.388	0	0	0	-	0.210	0.291
Distance to roads	-	0.337	0	0	0.215	-	0.087	0.028
Mean temperature	-	0	0.358	0	0.081	-	0.089	0.026

**Appendix C. A. SAC of Culex predictors. B. SAC of Stegomyia predictors. Red circles indicated a significant p-value ( $p < 0.05$ ) and indicate distances where the variable was autocorrelated**

A.



B.



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