Landscape factors and how they influence whitefly pests in cassava fields across East Africa

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Abstract

Context African production landscapes are diverse, with multiple cassava cultivars grown in small patches amongst a diversity of other crops. Studies on how diverse smallholder landscapes impact herbivore pest outbreak risk have not been carried out in sub-Saharan Africa.

Objectives Bemisia tabaci is a cryptic pest species complex that cause damage to cassava through feeding and vectoring plant-virus diseases and are known to reach very high densities in certain contexts. However, the factors driving this phenomenon are unclear.

Methods Bemisia density data in cassava across a large number of sites representing a geographic gradient across Uganda, Tanzania and Malawi were collected. We tested whether in-field or landscape factors associated with land-use patterns underpinned Bemisia density variability and parasitism.

Results We found the B. tabaci SSA1 species dominated our study sites, although other species were also common in some cassava fields. Factors associated with the surrounding landscape were unimportant for explaining variability in adult density, but the in-field variables of cassava age and cultivar
were very important. The density of nymphs and the parasitism of nymphs was heavily influenced by a diversity of landscape factors surrounding the field, including the size of focal cassava field, and area of cassava in the landscape. However, unlike the trend from many other studies on drivers of natural enemy populations, this pattern was not solely related to the amount of non-crop vegetation, or the diversity of crops grown in the landscape.

Conclusions

Our findings provide management options to reduce whitefly abundance, including describing the characteristics of landscapes with high parasitism. The choice of cassava cultivar by the farmer is critical to reduce whitefly outbreak risk at the landscape-scale.

Keywords

Crop diversity · Bemisia tabaci · Bemisia afer · Habitat manipulation · Parasitoids · Natural enemies · Pest density

Introduction

Small-holder production landscapes are inherently diverse both spatially and temporally. Large scale stochastic processes influence the environmental variability, and therefore the resources (such as water) available to farmers each season. This, together with planned household food consumption and external market demands influence the area committed to growing specific crops by smallholder farmers in East Africa. At a field scale, individual farmers make decisions about what crops they plant where, when they plant, apply inputs (fertilisers and pesticides) and harvest, and how they manage the non-productive parts of their land. Farmers’ management decisions directly influence biotic processes, such as the amount of suitable host-plant material available at any time for invertebrate herbivores. The change in the amount of host-plant resources across a landscape can influence population density within a field unit and population persistence from one season to the next. For widely dispersing pest species, landscape-scale effects can undermine or completely ameliorate field-scale management actions (Gurr et al. 2018). An understanding of landscape-scale effects on pest species are needed for the development of area-wide management (AWM) approaches. AWM has been advocated for reducing the risk of pest outbreaks, increasing the activity of natural enemies, and reducing the risk of insecticide resistance (Schellhorn et al. 2015). However, AWM can sometimes be difficult to achieve because it requires coordination across different landholders.

There have been many studies examining the relationship between the composition, configuration, and diversity of land-use in agricultural landscapes and pest outbreaks. Empirical studies testing these relationships come predominantly from intensively farmed landscapes (often consisting of monoculture crops), in high-income countries (Chaplin-Kramer et al. 2011; Karp et al. 2018). Some studies show a clear link between increased amounts of non-crop habitats and increased natural enemy diversity at the landscape-level, but subsequent pest suppression is not always observed (Chaplin-Kramer et al. 2011; Veres et al. 2013; Karp et al. 2018; Duarte et al. 2018). It remains unclear whether such patterns also exist in highly diverse smallholder production landscapes (Zhou et al. 2014). Using household surveys in smallholder systems in Nigeria, Zhang et al. (2018) reported crop pest severity was less where the proportions of forest and unused land at the landscape scale was higher. However, other agronomic and socio-economic factors were also important for explaining reported pest severity. In contrast, increased landscape diversity, through management actions to increase the area of grasslands, has been shown to increase the risk of Lepidoptera stemborer damage to maize and sorghum, by providing a new source habitat for these pests (Midega et al. 2014). Given the low numbers of studies we can’t yet say if the drivers associated with lower populations of pests in intensively farmed, homogenous landscapes will differ substantially from the drivers in more diverse smallholder landscapes.

Cassava is a common crop included in smallholder production landscapes in East Africa. Whilst it is grown over large areas of land, it is not managed as a monoculture crop, and has a diversity of cultivars, soil preparation techniques, and harvesting options. Cassava is a long-duration crop (remaining in the ground for periods between 8 and 18 months), with relatively low inputs in terms of fertilisers and pesticides, and therefore represents an ideal environment for population growth and activity of natural enemies. Furthermore, in some parts of East Africa, cassava is planted
in two seasonal windows per year, resulting in
temporal overlap between fields at different stages of
maturity. Farmers plant a diversity of different cassava
cultivars with different traits, including tolerance or
resistance to two major diseases; cassava mosaic
disease (CMD) caused by a multiple Begomoviruses,
and Cassava Brown Streak Disease (CBSD) caused by
two RNA Ipomoviruses (Legg and Thresh 2000;
Maruthi et al. 2002; Alicai et al. 2007). Both diseases
are vectored by whitefly pests in the
*Bemisia tabaci* complex, consisting of more than 36 different but
morphologically similar species, while CBSD is
potentially also vectored by other *Bemisia* species
(Maruthi et al. 2002; 2005; Legg et al. 2011, 2014a, b;
Ateka et al. 2017). However, these diseases can also be
spread between fields in a region via the transfer of
infected cuttings from one farmer to the next. The
realization in recent year that *B. tabaci* is a pest species
complex (Maruthi et al. 2004; Mugerwa et al. 2018;
Elfekih et al. 2018; Vyskočilová et al. 2018; Mugerwa
2019; Kanakala and Ghanim 2019; Kunz et al. 2019)
has required a re-assessment of the virus-vector
relationships, and research on the phylogenetic posi-
tion of species sampled from East Africa, using
molecular approaches (Alicai et al. 2016; Boykin
et al. 2018). The combination of disease impacts and
damage caused by high *B. tabaci* populations in
cassava production regions of East Africa is one of the
key constraints on the productivity of smallholder
farmers. Cassava breeding efforts have produced
several new cultivars that are resistant or tolerant to
either CMD or CBSD (Adriko et al. 2011; Katono
et al. 2015). However, improved cultivars are usually
planted alongside un-improved local landraces in a
farming landscape as these cultivars have other traits
that are desirable for farmers. Furthermore, a diversity
of other crops are grown in small fields and gardens
that may also act as host plants for *B. tabaci*. This
mosaic of host plant resources may potentially influ-
ence the behaviour (Kalyebi et al. 2018) and land-
scape-wide population abundance of *B. tabaci* (Parry
et al. 2020).

Until recently, research on and control of agricul-
tural pests, such as species in the *B. tabaci* complex,
was focussed at the individual field level with little
reference to surrounding land-use (although see Krist-
tensen et al. (2013) and Bianchi et al. (2015) for
models of a whitefly parasitoids at multiple scales). These field-scale studies together with discrete
behavioural experiments provide information about
an insect’s biology, by way of life-table analysis.
Although laboratory and field-scale research has
revealed valuable information about *B. tabaci*, they
do not reflect other important aspects such as the
spatial variation in the density of these pests in more
complex field environments. Controlled studies have
shown that *B. tabaci* populations respond differently
to different cassava cultivars (Ariyo et al. 2005;
Omongo et al. 2012; Katono et al. 2015; Kalyebi et al.
2018). Environmental conditions like temperature and
rainfall also impact population dynamics, either
directly by causing mortality of immature stages, or
indirectly through altering populations of natural
enemies (see Macfadyen et al. 2018 for a review).
Therefore, we can observe very different levels of *B.
tabaci* abundance across a landscape between regions
and seasons (Kriticos et al. 2020). In this study, we aim
to understand how the in-field and land-use factors in
the surrounding landscape influence the observed
variability in the density of this pest in cassava fields.
Ultimately, we can use this knowledge to develop
novel landscape management approaches that can be
implemented by smallholder farmers and adopted at a
relatively low cost. To achieve this, we collected data
on *Bemisia* abundance from smallholder cassava
farms, at a landscape-scale, at sites in three East
African countries; Uganda, Tanzania, and Malawi.
This large geographic gradient allowed us to examine
*Bemisia* whitefly populations across a diversity of
different cassava production contexts. Given that the
species in the *B. tabaci* pest complex cannot be
identified in the field using morphology alone, we
developed a high-throughput sequencing method to
identify the common species present in our focal fields
(Tay et al. 2020). Using this data set we asked:

1. What are the common species of whitefly found on
cassava and on nearby crops and non-crop host
plants?
2. Which landscape or in-field factors influence the
density of *Bemisia* adults and nymphs in cassava
fields?
3. Which landscape or in-field factors influence the
parasitism of *Bemisia* nymphs in cassava fields?
Methods

Cassava fields were selected in at least three different geographical regions in Uganda (Lira called “UG4”, Kamuli “UG1” and “UG3”, Kyotera “UG2” and “UG5”) and Tanzania (Mwanza called “TZ1” and “TZ5”, Dodoma “TZ4”, and Dar es Salaam “TZ2” and “TZ3”), and two regions in Malawi (Northern called “ML1” and Central “ML2”, Fig. 1). Kamuli, Kyotera, Dar es Salaam and Mwanza were sampled at two different periods (referred to as 12 different regions throughout for ease) (Fig. 1, Online Appendix 1, Table A1.1). These regions were chosen to represent different agro-ecological zones that grow significant amounts of cassava. The exception is TZ4, in the Dodoma region, where cassava production has only recently started. In total, we conducted three data collection trips across two years (trip 1: 1/8/2015—26/08/2015, trip 2: 5/4/2016—25/4/2016, trip 3: 29/10/2016—9/11/2016) but complete regions were sampled at each trip (i.e. all fields in a region). Each of these regions differ in their altitudes, temperature and rainfall profiles, and the agricultural systems used by farmers. For example, in Kamuli (average altitude of 1090 m above sea level, msl) and Kyotera (average altitude of 1210 msl) in Uganda, there are generally two planting windows for cassava per year, but in northern Malawi (average altitude of 547 msl), there is one (Online Appendix 1). Before sampling, regions were visited by extension officers and contact was made with local farmers to obtain their permission to sample in their fields.

In each of these 12 regions we searched for up to ten focal cassava fields to sample and each focal field was sampled once. Focal cassava fields selected needed to have: cassava of between three to seven months after planting; cassava as the dominant crop (although it could have an intercrop of another crop type if cassava was still dominant); cassava cultivars that could be identified and that were consistently planted across the field; at least 30 plants of the same cultivar to survey. Each focal field had to be at least 4 km (straight line distance) from the next nearest focal field. Prior to sampling farmers were interviewed to confirm information about the cultivars of cassava and other crops that they were growing and associated management activities, i.e. crop rotation and fungicide and pesticide use (OnZine Appendix 2). A field deemed suitable for sampling was assigned a focal field code and a full sampling protocol conducted. All data were recorded on pre-designed and trialled electronic forms that were constructed using Open Data Kit (ODK) software (Hartung et al. 2010). ODK Software was run using Android tablets, and field collection identifiers related to unique barcodes which allowed all information and samples to be referenced back to the field and individual plant. Data was collected offline and uploaded to secure cloud servers after the sampling was completed.

There were 36 different cassava cultivars recorded from the focal fields, some of which were unique to a certain region (although they may have been dominant within the region). Each cassava cultivar that was surveyed was categorized into one of four groups; susceptible to both CMD and CBSD, tolerant to CMD (CBSD could be susceptible or resistant), tolerant to both CMD and CBSD, resistant to CMD (CBSD could be susceptible or resistant), based on the knowledge of scientists involved in the project. Four cultivars had an unknown disease rating and were categorized as susceptible. Four cultivars had an unknown disease rating and were categorized as susceptible.

Mapping the landscapes

Land-use information was captured within a minimum radius of 100 m from the centroid of the focal cassava field. Given that all the mapping was completed manually, this minimum distance was chosen as it captured a large amount of the land-use diversity surrounding the focal field, and at the same time could be completed in a reasonable time-period. Field boundaries and landscape features were digitised on Android tablets using offline maps and satellite image base layers, authored in ArcGIS Collector (ESRI 2015) and resulting spatial layers checked and cleaned in ArcGIS desktop (ESRI 2010). This information was confirmed by walking field boundaries and ground truthing the details which were then digitized and used to produce maps. Different land uses and features including roads, buildings and different crop and host plant, and non-host plant categories were then added. We focussed on the dominate crop and vegetation types that could be mapped with some accuracy and were present across enough fields to enable statistical analysis. Simple landscape metrics were generated,
such as percentage cover of land-use types around the focal field, size of the focal field, amount of cassava in the landscape, diversity and number of other crops, the amount of non-crop vegetation (i.e. grassland or wooded areas, example maps of three sites are provided in Online Appendix 1).

Fig. 1 Focal cassava field sampling sites across 12 regions. At each site, a focal cassava field was sampled and the landscape around the focal field mapped. See Online Appendix 1 for examples of focal field maps.
Counting adult and nymph *Bemisia* species, and collecting parasitoids

It is not possible to identify the different species within the *B. tabaci* pest complex morphologically in the field, so we collected samples of *Bemisia* species adults and nymphs for later molecular-based identification. Throughout the methods we refer to "whitefly" to encompass *Bemisia* whitefly species later clarified using molecular identification. To sample for whitefly adults and nymphs 30 cassava plants were randomly chosen and sampled in each focal field. The top five fully expanded leaves of each cassava plant were carefully turned over, (to minimise disturbing the adults) and the numbers of adult whitefly were counted and entered as a range. During analysis each range was given a number (0 = 0, 1–9 = 1, 10–50 = 2, 51–100 = 3, 101–200 = 4, > 200 = 5), hereafter referred to as density categories. Similarly using a five point scoring (1–5 with 1 being no symptoms and five the most severe) each plant was scored for CBSD symptoms, CMD symptoms, cassava green mite (*Mononychellus tanajoa*) infestation symptoms, and sooty mould severity on the upper surfaces of the lower leaves (Sseruwagi et al. 2004). We attempted to use this data to determine whether the density of adult or nymph of *Bemisia* species influenced CMD and CBSD symptom expression in cassava fields. Previous studies have sought to link high abundance of whitefly with greater incidence or severity of cassava diseases at the field and pandemics at the regional scale (e.g., Maruthi et al. 2005; Rwegasira et al. 2011) usually with limited success. In our study we also found no clear relationship between CMD and CBSD symptoms and *B. tabaci* density in the focal fields (either for adults or nymphs). We show the full results in Online Appendix 4 but will not discuss them further in this study.

Further down each cassava plant stem, three leaves per plant were turned over and visually inspected for third to fourth instar nymphs. Cassava leaves are palmate and divided into lobes. If there were third-fourth instar nymphs anywhere on the central lobe of any of the three leaves a 10 cm diameter metal cutter was placed over the leaf and a leaf disc removed. Leaf discs (maximum three per plant) were then placed sequentially in a partitioned leaf disc holder, a modified clipboard (See images in Online Appendix 1). Each disc was labelled with a barcode, which related to the field identifiers. A digital image of each barcoded leaf disc was taken in the field. Images were examined later and the number of live third-fourth instar nymphs, and the number of empty pupal cases (where the adult had emerged) were recorded. We were unable to determine if a nymph had been parasitised from the photographs, so the nymph density count includes both parasitised and unparasitised live third to forth instar nymphs. The rearing data were used to estimate parasitism rate, once the adult parasitoids and whitefly had emerged.

For the adult whitefly data, the density categories were summed across the five leaves assessed per plant. We then calculated the mean from the 30 plants sampled in each focal field. We refer to this as adult density throughout. For nymph density, we used the count of the number of live nymphs per leaf disc. Before analysis, we examined the relationship between nymph count and actual leaf area included in the disc (as not all cassava lobes were wide enough to occupy the whole disc area). However, no obvious relationship was found, therefore we ignored leaf disc area, and used live nymph numbers. We summed the number found on each of the three discs, then took the mean across the 30 plants in the focal field.

Cassava discs cut from each of the focal field plants were combined (max three per plant) and placed into small black emergence containers (see Online Appendix 1 for images and details of the materials used to construct emergence containers). A barcode was stuck to the 5 ml clear screw-top vial on top of the emergence container. When the whitefly adults or parasitoids emerged, attracted by the light transmitted through the clear plastic vial they moved up into the larger 5 ml vial. The emergence containers with leaf discs were kept at room temperature out of direct sunlight and monitored while the whitefly adults and parasitoids emerged. After a minimum of 14 days the containers were checked for a final time. Ethanol was added to the vial to preserve the whitefly and parasitoids. Given the high numbers of parasitoids that emerged in the first sampling trips (over 80% parasitism in some fields in Uganda, and a maximum of 923 live rearings from one field), we had to reduce the number of leaf discs collected in the final trip. In two regions, only three focal fields (out of ten) were selected for rearing nymphs to obtain parasitism data (UG3 and UG5). To calculate the rate of parasitism per focal field, we used the numbers of adult whitefly and
parasitoids emerging from the emergence containers. We summed the data from all containers in each focal field. We divided the parasitoid adults by the sum of the live emergences (parasitoid adults plus whitefly adults) giving a proportion between zero to one. Eight fields with zero counts were removed before analysis: firstly, fields where no leaf discs were collected due to low/absent nymph numbers, and secondly fields with few leaf discs collected (due to low nymph density) and therefore no live emergences. Fields with leaf discs collected and live emergences recorded, but no parasitoids emerged were considered true zeros and retained in the data set.

The other crops beyond cassava and areas of non-crop vegetation were mapped and given a field code. A timed search method was used to quickly assess these fields for whitefly adults or nymphs. For 15 min plants were examined for the presence or absence of whitefly nymphs or adults. The search protocol was modified slightly in response to the different growth habits of crops. The presence and absence of whitefly adults/nymphs and the total number of plants searched were recorded. Timed searches of non-crop areas involved timed searches of any plants in the areas for whitefly adults and nymphs. When the search was completed, the name and/or description of the plant on which adults and nymphs were found was recorded. If the name of the plant could not be confidently confirmed a specimen was barcoded and pressed for identification later. If whitefly nymphs were found on the plant a sample was collected for genotyping. Genotype samples were barcoded and cross-referenced to the field code and preserved as above.

Molecular identification of whitefly nymphs on cassava

In addition to the density data, a sample of nymphs was collected for genotyping using a flat-based 5 ml plastic vial without a cap. Using the edge of the vial small ~ 7 mm diameter leaf discs complete with nymphs were cut from the leaves. The leaf discs were placed into the vial (the same vial used to cut the leaf disc) and filled with > 95% ethanol and barcoded and sealed with a screw-top lid. A maximum of three vials were collected per focal field. Adults collected using aspirators were also added to these vials. We targeted nymph specimens for genotyping because we could be certain of the host plant relationship as per Sseruwagi et al. (2006). In some cases, the field did not meet all our criteria to be considered a focal field (cassava too young or old, mixed varieties, too few plants). However, we still collected a sample of whitefly for genotyping and basic information on the characteristics of the cassava field (size of field, GPS location, etc.) (“genotype only” fields in Online Appendix 3).

DNA extraction and sequencing library preparation followed the methods described in Tay et al. (2020), based upon the Illumina 16S Metagenomic Sequencing Library Preparation (Part # 15044223 Rev. B) protocol. For the full molecular protocol see Tay et al. (2020). Briefly, nymphs were dislodged from leaves and visually sorted into parasitised and unparasitised groups under a dissecting microscope (by looking for evidence of developing hymenopteran larvae in the nymphal case). Samples of 20 or 40 unparasitised individuals per vial (representing populations from each field) were randomly selected for DNA extraction. Genomic DNA (gDNA) was extracted from each population using the Qiagen DNEasy Blood and Tissue Kit (cat. #69506), including the optional RNase A treatment. Each sample of gDNA was quantified using a Qubit Fluorometer and dsDNA HS Assay Kit and samples were standardised to 0.5 ng/μl in preparation for PCR amplification.

Two sets of primers (wfly-PCR-F1/R1 and wfly-PCR-F2/R2; Tay et al. 2020) were used to amplify the target mtCO1 region. PCR product was visualised on an agarose gel to determine success, before being cleaned and purified using Agencourt AMPure XP beads (cat. #A63882). Purified PCR product was quantified using Qubit and standardised to 0.5 ng/μl, before Index PCR to construct amplicon libraries. Indexed amplicons were then quantified and samples were combined in equal amounts, to create an F1/R1 pool and an F2/R2 pool. The pooled samples were run on a gel, the expected fragment size excised, and the amplicons cleaned and purified using the Zymoclean Gel DNA Recovery Kit (cat. # D4007). Quality and quantity of the amplicon libraries was ascertained using Qubit and Agilent Technologies TapeStation. Purified indexed amplicon pools were diluted to 4 nM and the sequencing run was performed with MiSeq Reagent Kit V3 (600 cycles).

High throughput amplicon sequence data from populations were analysed using Geneious 11.1.5 (Biomatters Ltd, Auckland, New Zealand) based on the workflow pipeline as described in Tay et al. (2020).
to determine the proportions of amplicons from each population that belonged to known and unknown *Bemisia* species. We ascertained *Bemisia* species identity based on characterised and NUMT-free/pseudogene-free partial mtCOI genes as described by Kunz et al. (2019). The species nucleotide boundary was set at 3% in the first instance (Vyskočilová et al. 2018; Kunz et al. 2019). Unmatched sequences were relaxed to 5% to allow for PCR-introduced nucleotide polymorphisms to be mapped to the reference sequences. We mapped the amplicon sequences to a total of 18 African *Bemisia* species, and included two putative novel species (Tay et al. 2020 GenBank accession numbers MN646951 and MN646952). Parasitoid, bacterial, fungal, and potential NUMTs/pseudogene sequences were assembled and identified by de novo assembly (for parameter see Tay et al. 2020).

Data analysis

All the data manipulation, graphing and statistical analysis were completed using R and associated packages (RStudio version 1.2.5033, R version 3.5.2, R core team 2018). In total, we had 101 focal fields in the three to seven months after planting age range with adult and nymph density data, and 79 fields with parasitism rate data that were included in the analysis. For all the response variables we checked for spatial autocorrelation between focal field sites by plotting spline correlograms (using spline.correlog in ncf package, Bjornstad 2019), however, we found no evidence of strong spatial autocorrelation. The explanatory variables were grouped into three categories: in-field factors, landscape factors relating to cropping components of the landscape, and landscape factors relating to the non-crop components of the landscape (Online Appendix 1). Before developing models, we tested for collinearity between pairwise combinations of the explanatory variables using Spearman correlation coefficient (magnitudes ± 0.5 were considered problematic, Online Appendix 1) and variance inflation factors (in the car package, Fox and Weisberg 2019, VIF > 10 were considered problematic). We removed the landscape suitability score (LSS, description in Online Appendix 1), area of “other crops” in the landscape (a catch-all for rare and infrequently recorded crop types) as these were highly correlated with other variables. The area of beans was removed as this land-use type was not consistently recorded across regions. We standardized the continuous explanatory variables by subtracting the mean and dividing by the standard deviation.

We used a model selection approach to identify the most parsimonious models for each response variable (denisyt of adults, nymphs, and parasitism rates in the focal fields). Each model was constructed with and without groups of explanatory variables from each of the three categories and examined the change in AICc and the weight of each model (“model.sel” function in R package MuMIn, Barton 2018). To assist in this process, we simplified the months after planting variable, by making continuous in all models (rather than categorical). In all models the variable “region” was included as a random factor. For the adult and nymph models, a generalized linear mixed model with a negative binomial distribution was used. For the final model we calculated the pseudo-R-squared value using the “r.squaredGLMM” function. We include both the marginal R-squared (R^2_m) which includes fixed effects and the conditional R-squared (R^2_c) which includes fixed and random effects, both using the delta method. We assessed the model adequacy of the final model using residual plots, QQ plots, and checking for over dispersion and zero inflation (using functions in the package DHARMa, Hartig 2019). For the parasitism rate in the final model we plotted the predictions from the model with a few of the explanatory variables. We used “predictInterval” functions from the merTools package (Knowles and Frederick 2019) to produce a fitted line and the 95% confidence intervals.

For parasitism rate we used a binomial model (glmer function in the R package lme4, Bates et al. 2015) as the response variable was a proportion (0–1) and added a weight associated with the number of live emergences per field. Given that there were less fields (N = 79) in this data set we struggled to get a full model with all explanatory variables (i.e. the ratio of observations to explanatory variables was getting too small). This model failed to converge and was removed from the analysis.

Results

Our detailed mapping data quantified how each region differs in the mean area covered by crop and non-crop
vegetation around focal cassava fields (Online Appendix 1, Fig. A1.5). From the Lira sites in the north of Uganda (UG4) having 55% of the landscape covered by crops, and 31% by un coppred forest, grass or shrubland, through to sites in southern Malawi (ML2) having 24% crops and 66% forest, grass or shrubland (Online Appendix 1, Fig. A1.5). The amount and types of crops planted exhibited significant changes across regions, with the amount of cassava on average at 20% of the landscape but varying from 1 to 84%. Sites in Uganda and the Lake Zone of Tanzania (TZ1 and TZ5) had significant amounts of maize and sweet potato fields surrounding each focal cassava field (Online Appendix 1, Figs. A1.4 and A1.5). The field sizes in East Africa are generally very small, with the cassava fields chosen as focal fields being slightly larger than a usual cassava field (as we need at least 30 plants of a known cultivar). The cassava focal field sizes were largest in Uganda at 4053 m² (± 1322), medium in Malawi at 2476 m² (± 583), and the smallest in Tanzania at 1754 m² (± 236). Change in the average number of crops grown in the landscape differed between regions, with Malawi central region having low diversity (ML2 mean = 1.2 crops, N = 5, s.e. = 0.2), and the Lira region in northern Uganda having the highest crop diversity (UG4 mean = 8.9, N = 10, s.e. = 0.6). Overall, insecticide use was very low across all the cassava regions surveyed, with about 15% of farmers reporting that they used insecticides on their crops (Online Appendix 2). There were only two farmers who confirmed insecticide use in cassava, but the pest being targeted was not always whitefly.

What are the common species of whitefly found on cassava and on nearby crops and non-crop host plants?

A large number of individual whiteflies were processed as part of the sequencing runs, with 149 samples processed in total (Table 1). Twelve of these samples consisted of adults only, and two samples had adults and nymphs combined (to increase the number of individuals per sample). The majority of the whitefly collected came from cassava host plants, but a few samples from other host plants also contained enough individuals to warrant inclusion (see details in methods). Overall most regions had on average less than one individual whitefly with unknown sequences per sample (exceptions being UG4, mean = 2.09, and TZ1 mean = 1.2 individuals). A complete list of the molecular identification results can be found in Online Appendix 3.

Of the 18 whitefly species used in our reference library, not all were detected in our samples. Notably the B. tabaci SSA3, B. tabaci MEAM1 and B. tabaci MEAM2 sensu Mugerwa et al. (2018) (c.f Tay et al. 2017) were not detected. Of the SSA species, B. tabaci SSA1 was the most commonly detected in the cassava fields we sampled across all regions (Fig. 2). B. tabaci SSA2 was present but only in a few Ugandan cassava fields (UG4 and UG1). On cassava, both B. tabaci Indian Ocean (IO) (TZ4) and B. afer (ML1, ML2) were relatively common at certain sites (Fig. 2). Some studies suggest that B. tabaci IO does not use cassava as a host plant (Misaka et al. 2020), however nymphs of this species were recorded on cassava and other plants throughout our study. B. tabaci SSA1 was also recorded on other crop and non-crop plants (e.g., wild groundnut, sweet potato), but often not in the frequency with which it was recorded on cassava (Fig. 3). Species such as B. tabaci MED and B. tabaci UGANDA1 were recorded only on non-cassava host plants (cowpea, sweet potato, and a rosella-like leafy vegetable). Sanger sequencing of individual specimens confirmed the sub-sampling of the HTS amplicon results.

Parasitoid DNA was consistently detected across our samples, demonstrating the utility of this method and associated primer pairs (Tay et al. 2020) for detecting species interactions. Despite deliberately selecting nymphs with no visual signs of parasitoid development, our detection of parasitoid partial mtCOI gene indicated that some nymphs were in the early stages of parasitism. Parasitoid DNA was absent in the samples that contained adults only, further confirming the robustness of this HTS method developed for this study. The proportion of parasitoid sequences seen in samples was highest in the Ugandan regions, and negligibly low in TZ4 and Malawi samples. Parasitoid sequences could be related to both the Eretmocerus and Encarsia genera that commonly parasitise the B. tabaci complex in East Africa (Polaszek et al. 1992; Otim et al. 2005).
Table 1 Summary of the molecular data collected on the identity of *Bemisia* whitefly species collected from cassava, other crops, and non-crop host plants

| Country | Region | Number of samples processed | Number of fields or non-crop areas | Total number of whitefly sequenced (adults and nymphs) | Total number of reads obtained |
|---------|--------|-------------------------------|-----------------------------------|-------------------------------------------------------|------------------------------|
| Uganda  | UG4    | 19                           | 18                                | 620                                                   | 167,065                      |
| Uganda  | UG1    | 16                           | 10                                | 233                                                   | 287,717                      |
| Uganda  | UG3    | 9                            | 8                                 | 220                                                   | 429,589                      |
| Uganda  | UG2    | 10                           | 10                                | 400                                                   | 406,131                      |
| Uganda  | UG5    | 10                           | 10                                | 380                                                   | 494,024                      |
| Tanzania | TZ1   | 11                           | 8                                 | 124                                                   | 187,086                      |
| Tanzania | TZ5   | 8                            | 7                                 | 240                                                   | 378,108                      |
| Tanzania | TZ4   | 12                           | 11                                | 245                                                   | 558,093                      |
| Tanzania | TZ2   | 13                           | 10                                | 193                                                   | 279,502                      |
| Tanzania | TZ3   | 17                           | 16                                | 440                                                   | 321,660                      |
| Malawi  | ML1    | 9                            | 8                                 | 301                                                   | 364,853                      |
| Malawi  | ML2    | 15                           | 10                                | 383                                                   | 740,583                      |

A total of 149 samples analysed. The regions run from north UG4 Lira in Uganda, to south ML2 southern Malawi (see map in Fig. 1).

Fig. 2 The proportion of sequences allocated to *Bemisia* species from a reference library from samples collected from cassava, other crops and non-crop host plants across East Africa. In total 149 samples were assessed across these regions (Table 1)
Which landscape or in-field factors influence the density of *Bemisia* adult and nymphs in cassava fields?

The densities of adults and nymphs in each of the focal cassava fields differed across the regions (Fig. 4). The highest adult densities were observed in cassava fields in the Kyotera region of Uganda (UG2, $n = 10$, mean = 4.78, s.e. = 0.63), the Kamuli region of Uganda (UG1, $n = 10$, mean = 3.71, s.e. = 0.54), and the Coastal region near Dar es Salaam in Tanzania (TZ3, $n = 8$, mean = 4.71, s.e. = 0.64). In contrast, the highest nymph densities were observed in cassava fields in the Kyotera region of Uganda (UG2, $n = 10$, mean = 6.79, s.e. = 0.1.6.9 and UG5, $n = 4$, mean = 5.55, s.e. = 3.66). The final model for adult density included only in-field predictor variables, with months after planting and cultivar category showing the strongest patterns (Table 2, Fig. 5a, Online Appendix 1). The conditional R-squared value for the final model was 0.65. There were significantly lower numbers of adults at 5, 6, and 7 months after planting, compared to early in the season (3 and 4 months after planting). For the cultivar categories there were lower adult density on the susceptible cultivars compared to those cultivars that were tolerant to both diseases (Fig. 5a). Landscape factors outside the field appeared unimportant for predicting adult density (Table 2).

For nymph density, landscape factors outside the field were important (Table 2). The two best performing models contained firstly only in-field factors (months after planting, cultivar category, and intercrop, conditional R-squared of 0.57); and secondly in-field factors along with landscape factors associated with crops (size of focal cassava field, and area of cassava in the landscape, conditional R-squared 0.69).
Landscape factors associated with non-crop components of the landscape appeared unimportant for nymph density in the focal fields. In the final nymph model, the factors months after planting (Fig. 5b), cultivar category (Fig. 5b), cassava area in the landscape, and area of the focal field were all significant (Table 2, Online Appendix 1). As for adults, there were significantly lower numbers of nymphs at 5, 6,
Table 2  The importance of in-field and landscape factors for predicting *Bemisia* adult and nymph density and parasitism rate

|          | ADULTS | NYMPHS | PARASITOIDS |
|----------|--------|--------|-------------|
|          | $a_2^a$ | $a_6$  | $a_5$ | $a_3$ | $a_7$ | $a_4$ | $a_1$ |
| In-field | months after planting* | *** |     |     |     |     |     |     |
|          | var.cat | *** |     |     |     |     |     |     |
|          | intercrop |     |     |     |     |     |     |     |
| Landscape, crops area |     |     |     |     |     |     |     |     |
|          | Cassava |     |     |     |     |     |     |     |
|          | no.crops |     |     |     |     |     |     |     |
|          | Sweet pot |     |     |     |     |     |     |     |
|          | Soybean |     |     |     |     |     |     |     |
|          | Pumpkin |     |     |     |     |     |     |     |
| Landscape, non-crop non-crop |     |     |     |     |     |     |     |     |
|          | Banana |     |     |     |     |     |     |     |
|          | Maize |     |     |     |     |     |     |     |
| AICc     | 375.1  | 380.2 | 385.2 | 398.6 | 405.0 | 396.8 | 391.6 |
| $R^2m$   | 0.164  |       |       |       |       |       |       |
| $R^2c$   | 0.646  |       |       |       |       |       |       |
|          | $n_2^a$ | $n_6$  | $n_5^a$ | $n_3$ | $n_7$ | $n_4$ | $n_1$ |
| In-field | months after planting* | *** | ** |     |     |     |     |     |
|          | var.cat | ** | ** |     |     |     |     |     |
|          | intercrop |     |     |     |     |     |     |     |
| Landscape, crops area |     | ** |     |     |     |     |     |     |
|          | Cassava |     | ** |     |     |     |     |     |
|          | no.crops |     | ** |     |     |     |     |     |
|          | Sweet pot |     |     |     |     |     |     |     |
|          | Soybean |     |     |     |     |     |     |     |
|          | Pumpkin |     |     |     |     |     |     |     |
| Landscape, non-crop non-crop |     |     |     |     |     |     |     |     |
|          | Banana |     |     |     |     |     |     |     |
|          | Maize |     |     |     |     |     |     |     |
| AICc     | 396.2  | 397.8 | 396.2 | 401.9 | 408.5 | 409.8 | 401.5 |
| $R^2m$   | 0.176  | 0.297 |       |       |       |       |       |
| $R^2c$   | 0.567  | 0.700 |       |       |       |       |       |
|          | $p_2$  | $p_6$  | $p_5$  | $p_3$ | $p_7^a$ | $p_4$ | $p_1$ |
| In-field | months after planting* |     |     |     |     |     | NA    |
|          | var.cat |     |     |     |     |     |     |     |
|          | intercrop |     |     |     |     |     |     |     |
| Landscape, crops area |     |     |     |     |     |     |     |     |
|          | Cassava |     |     |     |     |     | ***   |     |
|          | no.crops |     |     |     |     |     | ***   |     |
|          | Sweet pot |     |     |     |     |     | ***   |     |
|          | Soybean |     |     |     |     |     | ***   |     |
|          | Pumpkin |     |     |     |     |     | ***   |     |
| Landscape, non-crop non-crop |     |     |     |     |     |     | ***   |     |
|          | Banana |     |     |     |     |     | ***   |     |
|          | Maize |     |     |     |     |     | ***   |     |
| AICc     | 1126.7 | 1124.0 | 1044.8 | 1197.0 | 1036.0 | 1264.6 |       |
| $R^2m$   | 0.117  |       |       |       |       |       |       |
| $R^2c$   | 0.262  |       |       |       |       |       |       |
and 7 months after planting, compared to early in the season (3 and 4 months after planting), and lower nymph density on the susceptible cultivars compared to cultivars that were tolerant to diseases (Fig. 5b). Interestingly, nymph density was higher in larger fields but was significantly lower in fields surrounded by a higher proportion of cassava (Online Appendix 1). Note that the area of the focal field and the amount of cassava in the landscape are related to each other although not strongly (Spearman correlation of 0.44, Online Appendix 1).

Which landscape or in-field factors influence the parasitism of *Bemisia* nymphs in cassava fields?

The parasitism rate of nymphs varied greatly between regions with the highest average of 83% parasitism in the Kamuli region of Uganda (UG1, n = 10, s.e. = 5.24), and 76% in the coastal region of Tanzania (TZ3, n = 8, s.e. = 5.64) (Fig. 6). There was a weak asymptotic relationship between nymph numbers in the focal field and parasitism rate (Online Appendix 1) suggesting that it is not only the density of nymphs that leads to high or low parasitism rate.

There are likely to be greater than 10 species of parasitoids that attack *B. tabaci* on cassava in East Africa (Guastella et al. 2015; Macfadyen et al. 2018). Given the numbers of parasitoids reared in this study we have identified and categorised them into three groups; *Encarsia* genera, *Eretmocerus* genera, and other parasitoids not easily grouped into these two genera. When we examined the live adult parasitoids, there was a greater proportion from the *Eretmocerus* genera in the Ugandan regions (Online Appendix 1), relative to *Encarsia* genera.

Landscape factors outside the field were more important in determining the variability in parasitism rate between the focal fields (Table 2) than in-field factors. Overall, in-field factors, such as cassava cultivar had no influence on the parasitism rate in the focal fields. The most parsimonious model contained all groups of crop and non-crop landscape factors, however, the model overall did not explain a lot of the variation in parasitism rate with a conditional R-squared of 0.26 (Table 2). The amount of cassava in the landscape was significant in the final model, but not the area of the focal field. The area of other crops (sweet potato, pumpkin, banana) displayed negative coefficients (Online Appendix 1), but this was sometimes a complex relationship with parasitism rate. For example, the highest parasitism rates were observed in fields with < 20% cassava in the landscape (with a peak at about 10%), and then decreased rates with greater amounts of cassava (Fig. 7). There was a decrease in parasitism rate with an increasing cover of non-crop land-use until 40–50% of the landscape, then no impact after that point (Fig. 7). The number of crops in the landscape (crop diversity) showed a positive coefficient in the final model, however again this pattern was complex. The parasitism rate increased from zero to four crops in the landscape, peaked at five crops, and decreased with higher diversity of crops in the landscape (Fig. 7).

**Discussion**

Cassava and other crop cultivation practices in East African smallholder production landscapes are incredibly diverse across space and seasonal conditions and we have a limited understanding of how pests and natural enemies respond to this diversity. In this study,
we used a broad geographic survey approach to describe how these diverse production landscapes impact the variability in *Bemisia* density in cassava

![Graph](image)

**Fig. 5** Results from the final statistical model. *Bemisia* adults (a) and nymphs (final model 2, Online Appendix 1) in cassava fields concerning months after planting and cassava cultivar category. The cassava categories represent cultivars that are tolerant to CMD and CBSD (“tolerant”), tolerant to CMD (“tolerant CMD”), resistant to CMD (“resistant CMD”), and landraces that are susceptible to both disease (“susceptible”). Full model outputs can be found in Online Appendix 1.
identified a range of in-field factors and landscape factors that contribute to the variability in Bemisia density observed between fields. For adults the in-field factors of cassava cultivar category and age of cassava (months after planting) were important, but for the nymph counts and parasitism factors outside the field in the broader landscape (e.g., area of cassava, size of the focal field) became important. These landscape factors will interact with regional factors (that were not examined in this study) to lead to high or low populations of Bemisia species in focal cassava fields. For example, we know that long-term climate and short-term weather patterns at the regional level can dampen or facilitate populations (Macfadyen et al. 2018). However, the in-field and landscape factors identified in this study have some potential for management by farmers, whereas climate patterns are generally outside of the control of individual smallholder farmers.

What are the common species of whitefly found on cassava and on nearby crops and non-crop host plants?

Given that the B. tabaci pest complex is still undergoing significant taxonomic and nomenclatorial changes (Boykin et al. 2018; Kunz et al. 2019) based on new molecular approaches for identifying species, reciprocal-crossing studies and consideration of implications from NUMTs/pseudogenes on Bemisia cryptic species status delimitation (Tay et al. 2017; Kunz et al. 2019), we spent a significant amount of time identifying samples of whitefly nymphs collected from cassava fields. This data set represents the most comprehensive molecular identification of samples from this region. We found that the Bemisia cryptic species community in cassava is dominated by the geographically widespread B. tabaci SSA1 species, however other B. tabaci and ‘non-tabaci’ species were also relatively common in some regions. This supports
previous studies that also found *B. tabaci* SSA1 to be widespread on cassava in Tanzania (Tajebe et al. 2015), Kenya and Uganda (Mugerwa et al. 2012), although with many fewer samples. In this study, we have used the phylogenetic mtCOI grouping of SSA1 and consider it to be a single, putative species, based exclusively on the mtCOI classification. However, previous studies have identified mtCOI sub-groups and sub-clades within SSA1 using various methods, and in the future, there may be more biological diversity described within this putative species (Ally et al. 2019; Elfekin et al. 2019). Future molecular characterisation of the mtCOI barcoding gene region used for species identification and ascertainment of

Fig. 7 Results from the final model. Parasitism rate (as a proportion of parasitoid adults emerged over all live rearings, between zero to one) of *Bemisia* nymphs in cassava fields in relation to the amount of cassava in the landscape, number of crops in the landscape and amount of non-crop in the landscape. Including region as a random effect. The blue line shows the fitted values and the two red lines the upper and lower 95% confidence interval.
genetic diversity should consider the impact of NUMTs/pseudogenes especially when utilising sub-optimal PCR markers (e.g., see Tay et al. 2017; Mugerwa et al. 2018; Elfekih et al. 2018; Kunz et al. 2019). In Malawi, \textit{B. afer} was also present in our nymph samples and could represent an important vector for some of the cassava diseases (Maruthi et al. 2005). In Tanzania, the \textit{B. tabaci} IO species was also relatively common in some cassava fields, on cassava plants. Misaka et al. (2020) did not record \textit{B. tabaci} IO on cassava in a survey conducted in South Sudan, however, their study focussed on collecting adults and sequenced low numbers of individuals from cassava. In our study, \textit{B. tabaci} SSA2 was found relatively frequently but in low density on other crop plants and non-crop plants in cassava production landscapes (Fig. 3). The \textit{B. tabaci} SSA3 species was not detected in our study but its presence in cassava was previously reported further west in the Democratic Republic of Congo (Legg et al. 2014a, b).

We know that species in the \textit{B. tabaci} complex differ in the range of plant species that they utilise as host plants (i.e. those that support successful growth and reproduction, Malka et al. 2018; Vyskočilová et al. 2018, 2019). Whilst adults were detected on a large diversity of plants in every production landscape, nymphs were recorded on a smaller selection of plant species and these were the focus of our molecular identifications. Sweet potato, given its frequent planting in many regions of East Africa, maybe a potential alternate host plant for \textit{B. tabaci} SSA1 and its parasitoids. \textit{B. tabaci} SSA1 was more detected on Tanzanian sweet potato samples than samples from Uganda (Fig. 3). However, more controlled host plant choice experiments are required to confirm this.

Which landscape or in-field factors influence the density of \textit{Bemisia} adults and nymphs in cassava fields?

When examining factors that influence the density of adult \textit{Bemisia} in cassava fields we found that in-field factors were the most important. In particular, the cassava cultivar sampled and the age of the cassava, with highest densities observed on improved cassava cultivars at 3 months after planting. It was surprising to see such dominance of cassava cultivars from such a geographically broad field-survey. Differences in whitefly infestation between cassava cultivars has been shown in replicated small plot trials in multiple locations in Nigeria (Ariyo et al. 2005) and Uganda (Omongo et al. 2012; Katono et al. 2015). However, these studies do not consider the alternative host plants present surrounding these trials, which may also be attractive and accessible to adult \textit{Bemisia} species. In these plot trials the local landraces (unimproved cultivars that are assumed to be susceptible to diseases) also experienced lower adult \textit{Bemisia} whitefly pressures.

For the variation in nymph density observed in the focal fields, factors in the surrounding landscape were important (along with cassava cultivar and months after planting). The area occupied by the focal field (positive relationship) and the amount of cassava surrounding the focal field (negative relationship) were both important factors in the final model. The number of nymphs on a cassava plant will a priori be related to both the numbers of adults, the number of eggs laid, and the presence of factors that induce mortality at the egg, first and second instar stage of development. These include processes such as competition, host plant defenses, environmental events, and predatory and parasitic natural enemies. For example, high rainfall events may wash away nymphs and deter adults from ovipositing on the upper leaves (Katono et al. 2019). However, these events should be unrelated to the landscape patterns observed in our study. There was a negative relationship between nymph density in the focal field and the area of cassava in the landscape. If mortality due to predatory natural enemies (e.g. ladybeetles and ants) is the main causative agent behind this pattern, then the natural enemy species may be gaining some benefit from increased amounts of cassava in the landscape. This theory is supported by the fact that non-crop factors appeared unimportant for nymph density in the focal fields. It may be that predatory natural enemies associated with the cropped components of the landscape can reach higher densities and therefore increase the mortality of whitefly nymphs in cassava.

Which landscape or in-field factors influence the parasitism of \textit{Bemisia} nymphs in cassava fields?

Although unusual for intensive agricultural production systems, the high levels of parasitism observed in this survey were not unexpected for smallholder systems in
High parasitism rates on cassava (58–67%) have been recorded in trials in Uganda (Otim et al. 2005, 2008). Furthermore, the detection of parasitoid DNA in the *Bemisia* nymph samples also suggests that parasitoids are a common part of this community. The next step is to determine the impact these parasitoids have on pest population reduction at the field and landscape-level.

The parasitism rate of *Bemisia* in focal cassava fields was heavily influenced by factors occurring in the landscape surrounding the field. However, this pattern was not solely related to the amount of non-crop vegetation, or the diversity of crops grown in the landscape. Highest rates of parasitism were recorded in fields that were in landscapes with < 20% cassava (relatively low), < 40–50% non-crop vegetation (low-moderate) and had an intermediate level of crop diversity (~ 5 crops in the landscape) (Fig. 7). However, our final model did not explain much of the variation in parasitism rate observed, suggesting that other factors (that we did not measure here) may also be influencing parasitoid behaviour and ultimately their ability to cause mortality to whitefly. It would be impossible to disentangle potential causative mechanisms without a manipulative field study. However, it is likely that the parasitoid species are using multiple whitefly hosts in the landscape, some of which may only be found on cassava, but others may have multiple host plants. For example, Guastella et al. (2015) recorded some parasitoid species attacking both *B. tabaci* and *B. afer* hosts in cassava growing regions of Tanzania. For a host-specific parasitoid, that parasitizes a herbivore that is also specific to certain host plants, landscape factors become less important. In southeast Asia, the cassava mealybug, *Phenacoccus manihoti*, colonized cassava fields earlier in high diversity landscapes, but overall abundance was the same in high and low diversity landscapes. Furthermore, there were no landscape effects seen for the parasitism of the mealybug by the host-specific parasitoid, *Apoanagyrus lopezi* (Le et al. 2018). In contrast, if the community of parasitoids attacking *Bemisia* species also attack other closely related herbivore hosts that use a range of host plants, then factors outside the field may influence their density and behaviour more.

Whilst we sampled a large number of focal cassava fields from different regions across East Africa, there are regions that may differ in their climate, agronomic practices and whitefly community which we have missed in this survey. Furthermore, there may be interactions between climate and landscape variables we have not included in this study. The static spatial landscape descriptors used in our study do not account for the dynamic nature of these smallholder production landscapes across the season. The 100 m radius we selected around centroid of the focal field was relatively small compared to other landscapes studies (e.g. Thies et al. 2003; Janković et al. 2016), however it was large enough to capture the land-use diversity surrounding the focal fields (see maps in Online Appendix 1). The limitation here is that we cannot rule out other drivers of whitefly population dynamics that may take place at larger spatial scales. For example, the spatial scale we used would capture the regular movements of between local land-use units as they search for oviposition sites (Kalyebi et al. 2018), but would not capture large-scale movements from one region to another as the cassava season comes to an end. Importantly, we did not classify the age structure of the surrounding cassava and the suitability of growth stage of other crop types and how these resources changed across time (see basic rotation information in Online Appendix 2). We know that *Bemisia* adults move around these production landscapes easily and maybe responding to a diversity of cues concerning oviposition choice (Kalyebi et al. 2018). In some regions, and on non-cassava host plants, there was a relatively high proportion of species outside of the *B. tabaci* complex recorded in our genotype samples. This may lead to some of the whitefly-parasitoid interactions recorded here to also involve *B. afer* and/or other *Bemisia* species. Currently, our high throughput sequencing approach cannot differentiate individual whitefly host species-parasitoid interactions (as we batch-processed 20–40 individuals in each population).

How can our findings help improve farmers’ management of whitefly pests?

Given the low usage of pesticides in these production landscapes, control options for smallholder farmers are limited to cultural options and choice of cultivars. Avoiding some of the improved cultivars that appear to support higher population densities of *B. tabaci* species is very important. Fortunately, our study has
shown that cassava cultivar does not impact the top-down reduction in nymphs due to parasitoids (also seen by Otim et al. 2006). Conducting small plot trials to rank commonly used cassava cultivars in each region by their ability to support the cassava *Bemisia* species would provide a useful resource for farmers. There may be cultivars that are tolerant to disease but are relatively poor at supporting the *Bemisia* species found in each region. In the future, the development of cassava cultivars that are simultaneously resistant to *Bemisia* whitefly and diseases may be available to farmers. However, careful consideration relating to the deployment of these cultivars in complex small-holder production landscapes will nevertheless remain necessary (see modelling conducted by Parry et al. 2020). In the meantime, farmers can potentially reduce whitefly adult and nymph densities on cassava by facilitating a mosaic landscape that consists of smaller sized cassava fields, and by avoiding placing new cassava fields adjacent to fields with existing high adult whitefly populations (Kalyebi et al. 2018). Furthermore, in our study, the highest rates of adult whitefly populations (Kalyebi et al. 2018). In the future, the development of cassava cultivars that are simultaneously resistant to *Bemisia* whitefly and diseases may be available to farmers. However, careful consideration relating to the deployment of these cultivars in complex small-holder production landscapes will nevertheless remain necessary (see modelling conducted by Parry et al. 2020). In the meantime, farmers can potentially reduce whitefly adult and nymph densities on cassava by facilitating a mosaic landscape that consists of smaller sized cassava fields, and by avoiding placing new cassava fields adjacent to fields with existing high adult whitefly populations (Kalyebi et al. 2018). Furthermore, in our study, the highest rates of parasitism were seen in fields that were in landscapes with relatively low amounts of cassava (< 20% cassava), low-moderate amounts of non-crop vegetation (< 40–50%), and an intermediate level of crop diversity (~ 5 crops in the landscape). There may be several ways that farmers may enhance landscapes with these characteristics. Finally, agronomists and extension workers supporting smallholder farmers need to be aware of the larger-scale and longer-term processes that operate on *Bemisia* species populations and also impact the risk of pest outbreak. For example, we know that some parts of Uganda have gone through an extended dry period which may make these regions more suitable for *B. tabaci* SSA1 growth and development (Kriticos et al. 2020), and in some cases, the double-cropping season may also support higher population abundances of *Bemisia* species at the landscape-level (Parry et al. 2020).

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