Modelling the potential distribution of *Bemisia tabaci* in Europe in light of the climate change scenario

Gianni Gilioli,a* Sara Pasquali,b Simone Parisi,c and Stephan Winter,d

Abstract

BACKGROUND: *Bemisia tabaci* is a serious pest of agricultural and horticultural crops in greenhouses and fields around the world. This paper deals with the distribution of the pest under field conditions. In Europe, the insect is currently found in coastal regions of Mediterranean countries where it is subject to quarantine regulations. To assess the risk presented by *B. tabaci* to Europe, the area of potential establishment of this insect, in light of the climate change scenario, was assessed by a temperature-dependent physiologically based demographic model (PBDM).

RESULTS: The simulated potential distribution under current climate conditions has been successfully validated with the available field records of *B. tabaci* in Europe. Considering climate change scenarios of +1 and +2 °C, range expansion by *B. tabaci* is predicted, particularly in Spain, France, Italy, Greece and along the Adriatic coast of the Balkans. Nonetheless, even under the scenario of +2 °C, northern European countries are not likely to be at risk of *B. tabaci* establishment because of climatic limitations.

CONCLUSION: Model validation with field observations and evaluation of uncertainties associated with model parameter variability support the reliability of model results. The PBDM developed here can be applied to other organisms and offers significant advantages for assessing the potential distribution of invasive species.

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Keywords: *Bemisia tabaci* MEAM1; population dynamics; invasive species; potential establishment; climate variability; biological variability

1 INTRODUCTION

*Bemisia tabaci* is a serious pest of agricultural and horticultural crops cultivated in tropical and subtropical climates around the world. A genetic analysis of world populations of *B. tabaci* revealed that it is a cryptic species complex composed of 11 well-defined genetic groups comprising more than 28 distinct species.1–5 The global status of *B. tabaci* as a pest refers mainly to two species: MEAM1 (formerly named biotype B) and Med (formerly named biotype Q), which are highly polyphagous, highly invasive and distributed worldwide, including the coastal regions of Mediterranean countries of Europe. Notwithstanding the major significance of MEAM1 and Med, other *B. tabaci* species can also be important agricultural pests and be particularly damaging by transmitting viruses causing serious crop diseases.5–8

*B. tabaci* occurrence in open fields in Europe is currently restricted to coastal regions of Mediterranean countries. Because of its worldwide occurrence as a major pest and the intensive global movement of agricultural and horticultural commodities, there is an inherent threat of *B. tabaci* and its associated viruses arriving from outside Europe. The pest and the viruses it transmits are currently subject to quarantine regulations within Europe, aimed at preventing introductions of *B. tabaci* and associated viruses and the further spread of existing populations. Consequently, a pest risk analysis (PRA) of *B. tabaci* to plant health in Europe was conducted by the Panel on Plant Health of the European Food Safety Authority (EFSA) at the request of the European Commission.9 The present study was initiated to help the EFSA PRA by providing a map of the potential distribution of *B. tabaci* under extant weather and its range expansion in the face of climate change. Temperature is the major environmental variable that influences the development, survival and reproduction10 and dynamics of *B. tabaci* populations, and its potential geographic range. Expansion of the pest northwards is thought to be limited by low temperatures, explaining why countries in the north of Europe are considered potentially pest-free areas, but this could change under the effect of climate variability.

Evaluation of the impact of climate variability on potential range expansion of invasive alien species requires the use of appropriate methodologies. Common correlativemethodsusedtopredict the geographic distribution of invasive alien species are classed as

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ecological niche models (ENMs) which attempt to characterise the ecological niche of a species on the basis of weather and other factors in the species’ native range. With restrictions, ENMs can also be used to estimate the potential invasiveness of the species in new areas. These methods include statistical models (e.g. GARP), physiological indexes (e.g. CLIMEX), artificial intelligence (e.g. MaxEnt) and other methods reviewed by Elith and Leathwick13 and Gutierrez et al.15 Empirical or statistical modelling approaches used to estimate the potential establishment of invasive alien species provide useful insight when the relationships within the systems are insufficiently known. However, simplified statistical and correlative models may lead to incorrect interpretation of the effects of environmental forcing variables on population dynamics. Integrative processes that give rise to counterintuitive population system responses may occur, and consequently the population dynamical patterns arising may not be simply interpretable in terms of linear responses to change in the environmental forcing variables.13

Mechanistic approaches are preferable when the underlying functional mechanisms of the processes are known in sufficient detail. Mechanistic models are crucial to evaluate the effects of biological and ecological processes on population dynamics, especially in complex systems characterised by non-linearity and acting over a wide range of space and time, as is the case when evaluating the effects of climate variation.14–16 Mechanistic approaches based on physiologically based population dynamics models (PBDMs)17 allow fine-scale predictions of the potential distribution using ecologically meaningful state variables directly related to the pest (population number or biomass). PBDMs have significant implications for the projection of population distribution and dynamics under climate change scenarios. These models take into account non-linear relationships between environmental forcing variables (e.g. temperature) and biological processes, so that population dynamics can be realistically described in sufficient detail, allowing the exploration of complex scenarios in which no simple (i.e. linear) conclusions can be drawn.

This paper reports on the development of an age-structured population model describing the local temporal dynamics of *B. tabaci* MEAM1 under current climatic conditions. The model simulates the population dynamics at the nodes (i.e. local dynamics) of a gridded map of Europe. Simulation data on whitefly population dynamics driven by extant temperature are used to define prospectively the potential distribution of the whitefly in Europe. The potential distribution is validated by comparison with data on the current field distribution of *B. tabaci*. Change in the average local population abundance and in the area of potential distribution are then investigated, considering variability in population dynamics owing to variability observed in the data used to develop the model. Finally, systematic perturbations of temperature of +1 and +2 °C are used to simulate the effects of climate change on the potential abundance and distribution of *B. tabaci*.

In the model, population dynamics is the outcome of the three fundamental demographic processes: development, mortality and fecundity regulated by rate functions (also known as biodemographic functions) that depend on environmental variables. In this first approximation, temperature was considered as the most important environmental forcing variable because of its main role in regulating the rate of biophysical processes in heterotherm species such as *B. tabaci*. Further relevant variables, including the host plant, could be easily introduced into the model.15

The daily rate of development as a function of temperature defines the change in physiological age within immature stage and adult stage (ageing process). The mortality rate function is the daily rate of individuals leaving the system owing to temperature-dependent intrinsic mortality.18 The fecundity rate function is the daily number of eggs produced by a reproductive female and describes the flux from the adult stage into the immature stage. Functional forms of development, mortality and fecundity rate functions are detailed in Appendix 2. Parameters for these functions were estimated for each stage of the population by fitting literature data for *B. tabaci* MEAM 1 using a non-linear least-squares regression method. This method also allowed estimation of the confidence bands for the biodemographic functions (Appendix 2).

Population abundance $N_j(t)$ at each stage $i$ in cell $j$ at time $t$ was obtained by considering a discretisation of the von Foerster Equations.20,21 The model simulates an entire year of *B. tabaci* population dynamics with a time step of 1 h. This small time interval guarantees convergence of the discretised von Foerster equations. The initial condition for whitefly is 0.1 adults plant$^{-1}$ on 1 March. From $N_j(t)$, an index was derived for population pressure related to average population abundance defined as the average number of adults per plant and time unit calculated for the period of the year during which the abundance of adults was above a threshold of 1 adult plant$^{-1}$. The index values were then mapped to visualise the area of potential establishment under the current climatic conditions and under two climate change scenarios.

### 2 Data and parameter estimation

Biodemographic parameters for *B. tabaci* MEAM1 were obtained from available literature on life history studies.22–52 Most references considered a dependence of life history traits on temperature as well as on host plant. The present authors only considered the dependence of biodemographic functions on temperature, and host plant as well as other factors (e.g. the origin of the experimental population) are regarded as components contributing to experimental heterogeneity. For precision, only data from reports in which the taxonomic identity of the experimental population was clearly determined were considered.

Meteorological variables are the fundamental driving forces for *B. tabaci* population dynamics. The climatological data input of the model was retrieved from the WORLDCLIM database (University of California at Berkeley, http://www.worldclim.org/) containing the monthly mean of minimum and maximum temperatures for the period 1950–2000. The WORLDCLIM database consists of gridded data assembled from a large number of sources, including: (i) the Global Historical Climate Network Dataset (GHCN), v.2. (http://www.ncdc.noaa.gov/pub/data/ghcnv2); (ii) the climatological normals (CLINO) for 1961–1990 (WMO, 1996); (iii) FAOCLIM 2.0, the global climate database (FAO, 2001). For simulation of population dynamics in Europe, the original world grid was restricted to the area $-11–41$°E, $30–73$°N. The climate data calculated for such a long reference period express a robust average
temperature input for the model. From the WORLDCLIM database, maximum and minimum monthly temperatures were obtained for each node of the grid. Subsequently, an interpolation model considering geographic position and astronomic data was used to obtain the hourly temperatures at each node (see Appendix 3).

Model validation was done with all records of *B. tabaci* presence in Europe and studies on the geographical distribution of the insects taken into account. The model performance was evaluated by superimposing the locations of *B. tabaci* findings in Europe, as reported in the literature, on the map of model prediction of *B. tabaci* occurrence.

2.3 Scenario analysis

2.3.1 Exploring biological variability

Literature data on development, survival and fecundity temperature-dependent rates of *B. tabaci* are affected by the heterogeneity inherent in the different experimental datasets, including biological variability. To take into account variations in biodemographic functions, the 95% confidence bands of the estimated development, mortality and fecundity rate functions were estimated (see Appendix 2). A commonly used approach to introducing variability into a model is to apply a Monte Carlo method performing various simulations of population dynamics by varying development, survival and fecundity rate functions in the confidence band regions, and to use these estimates to compute the mean of the simulated abundances. The high computational complexity of the present system, however, made this method inapplicable, and consequently only two extreme situations were considered here: (i) the lower bound and (ii) the upper bound of the confidence band for development, survival and fecundity rate functions. The distribution area simulated using the Monte Carlo approach was expected to fall between these two extreme cases.

2.3.2 Exploring climate change

Climate scenarios generally consider changes in temperature for time horizons far beyond that of risk assessment analyses. Projections on temperature changes, based on global climate models (GCMs), are subject to uncertainty. This uncertainty is low on a global scale, but with decreasing scale, i.e. on a continental or regional scale, errors increase. Thus, the reliability of scenarios on future spatial and temporal patterns of temperature is strongly affected, and this also disturbs an analysis of *B. tabaci* population dynamics following temperature change. Sensitivity analysis in PBDMs driven by meteorological variables can be carried out, imposing perturbation to time series of weather data by means of stochastic methods or deterministic methods modifying the dataset with predefined changing factors. The latter approach has been adopted here, and the forcing variable (temperature) was systematically perturbed by adding 1 and 2 °C to the hourly temperatures derived from the current climate average at each node for the selected temporal horizon. Further explanation and justification are given in Appendix 4. To assess the impact of climate...
change on whitefly population dynamics, a time horizon of 20 years was considered.

3 RESULTS
3.1 Model validation and delineating the potential distribution of B. tabaci in Europe

The model output obtained under current climate conditions is validated by superposing the available information on the distribution of open field records of B. tabaci in Europe on the map of the prospective spatial distribution of the B. tabaci population pressure index (Fig. 1). The simulated distribution of B. tabaci satisfactorily corresponds to the observed distribution, as all recorded findings fall within the predicted area of the potential establishment forecast by the model. Some discrepancies may result from the 25 × 25 km spatial resolution in the simulation based on temperature estimated at the node of a cell.

Simulation results presented in Fig. 1 depict the area of potential establishment for B. tabaci in Europe, and indicate areas with a population pressure greater than zero. Prospectively, B. tabaci solid establishment is expected in the Mediterranean coastal areas and in the interior regions of Spain (red). Limited presence, indicated by very low population abundance (blue), is expected in southern France, in some areas of northern Italy and in some regions of the Balkans, suggesting low favourability. Note that intermediate values are very rare. Further, it is not expected that B. tabaci will expand its distribution to colonise regions outside this coloured region because of the lower temperatures prevailing in central northern regions of Europe, particularly during winter. Thus, countries in the north of Europe (white area in Fig. 1) can be considered to be beyond the climatic limits of the pest.

A further characteristic associated with the potential establishment is the fraction of the year in which adult whitefly abundance is above a given threshold. Considering a threshold level of 1 adult plant$^{-1}$, the fraction of the year in which the populations are potentially active is computed at each node of the grid, and results are mapped in Fig. 2. In most of the area of potential establishment, the whitefly is expected to be active only 3–5 months of the year. Only in the warmer areas of southern Europe is the whitefly active for 7 months, in contrast to 1–3 months in the northern limits of its distribution.

Uncertainty associated with experimental heterogeneity was included in the model by considering the upper and lower limits of the 95% confidence bands of the biodemographic functions estimated by literature data (Figs 3 and 4). When the lower bounds for development, survival and fecundity were considered, the potential distribution of B. tabaci was greatly reduced. The whitefly distribution remained limited to the European Mediterranean coastal areas and partly to central Spain, with only lower values of population pressure (Fig. 3). In this case the simulated area of potential establishment was far smaller than the area of current distribution (see the circles in Fig. 1).

Using the worst-case scenario, where the upper limits for development, survival and fecundity are considered, the area of potential establishment increased and, with it, the putative population density and level of infestation. This is shown by a sharp increase

![Figure 2. Fraction of the year in which the simulated abundance of B. tabaci adults is above the threshold of 1 adult plant$^{-1}$.](image-url)
in the whitefly population pressure index (red) in areas where the insect is established (Fig. 4), and also by an expansion of its potential range generally (e.g. the Balkans). Nonetheless, countries in northern Europe (white) are not in the expanded range and hence not at risk that B. tabaci might establish.

3.2 Exploring the consequences of climate change scenarios
Simulating temperature increases of 1 and 2 °C, expansion of the areas currently invaded by B. tabaci is predicted (Figs 5 and 6). Further spread and an increase in abundance are expected, particularly in Spain, France, Italy, Greece and along the Adriatic coast of the Balkans. Apart from small areas in southern Germany and patches in Poland that are prospectively suitable for insect development (Fig. 6), B. tabaci is not expected to establish in such isolated areas. Even under a climatic change scenario of +2 °C, northern European countries are not likely to be suitable for B. tabaci establishment because of adverse conditions, in particular low winter temperatures.

4 DISCUSSION AND CONCLUSIONS
Several population models that can be helpful to predict the potential establishment of B. tabaci have been developed in the past years (for review, see Naranjo et al.10). The models integrate biological and ecological knowledge and are temperature-dependent 78,79 or spatiotemporal models,24,80 or aim at predicting population abundance81–84 to support pest management tactics and strategies or to explain the role of different parasitoids in the biological control of whitefly.

With respect to the models on B. tabaci previously developed, the model proposed here is novel in that it considers biological uncertainties in the projection of the population dynamics and distribution. The model accounts for the biological variation in life history traits of the organism related to heterogeneity characterising different experimental datasets reported in the literature. The model has also been used to explore the effects of climate change on the potential establishment of B. tabaci.

For model development, sufficient information on the biological characteristics of B. tabaci, from life history data to vector transmission capacity, is available. Because of their significance, most studies have focused on the former B. tabaci biotypes B (MEAM1) and Q (Med). In the initial phase of model development, the estimated biodemographic functions for MEAM1 and Med were compared, and it was found that only the number of eggs laid per female was generally higher in MEAM1 than for Med species, while development and mortality rates were very similar for the two species. Because significantly more literature references were available for MEAM1, only data for this species were used for model parameter estimation. Based on its biodemographic performances, the present authors consider that MEAM1 has prospectively a wider area of potential establishment and reaches higher average densities. The fact that the Med species...
Figure 4. Spatial distribution of the simulated *B. tabaci* population pressure in Europe, starting from an initial condition of 0.1 individuals plant$^{-1}$ on 1 March, and considering upper limits of 95% confidence bands for development, survival and fecundity rate functions. Population pressure is the average number of adults per plant and time unit calculated for the period of the year in which the adult abundance is over 1 individual plant$^{-1}$.

Figure 5. Spatial distribution of the simulated *B. tabaci* population pressure in Europe, starting from an initial condition of 0.1 individuals plant$^{-1}$ on 1 March, under a climate scenario with an increase of +1 °C from the current temperatures. Population pressure is the average number of adults per plant and time unit calculated for the period of the year in which the adult abundance is over 1 individual plant$^{-1}$.
could take advantage of its resistance to pesticides would not affect the projection of its potential prospective range as a function of climate.

For the purpose of the PRA, the goal of which was to assess the establishment and spread of \textit{B. tabaci} in Europe, the model predictions for the area suitable for establishment (Fig. 1) matched well with those locations where \textit{B. tabaci} has been reported. The regions in Fig. 1 that are highlighted (blue-red) in the graphical presentation of the model correspond to the regions in Europe where \textit{B. tabaci} is known to occur. This was taken as reasonable proof of the model’s validity. The blue areas in Fig. 1 represent the outermost limits of the \textit{B. tabaci} possible range and thus demarcates the borders of the area of establishment. Northward, beyond that boundary, \textit{B. tabaci} cannot establish because low temperatures are limiting. Insects introduced into these areas, i.e. with plant materials, may survive under conducive conditions (in summer or in greenhouses), even lay eggs and develop locally for a few generations, but adverse autumn/winter conditions will prevent establishment.

It can be assumed that \textit{B. tabaci} species around the world have temperature requirements similar to those of MEAM1 or Med, and hence it is unlikely that \textit{B. tabaci} species other than Med and MEAM1 could invade and establish in locations outside areas in which \textit{B. tabaci} MEAM1 and Med have established. Therefore, based on biological data for \textit{B. tabaci} MEAM1, the model probably captures what is probably the worst-case scenario in terms of the potential establishment and spread.

The results of model validation and the exploration of model parameters made it possible to consider the uncertainties associated with the definition of the area of potential establishment of \textit{B. tabaci}. Choosing model parameters less favourable to \textit{B. tabaci} population dynamics results in a significant reduction in the area of potential establishment. However, this projected distribution was far less than the observed distribution in Europe, making this scenario very unlikely. When highest survival, development and fecundity for \textit{B. tabaci} were used in the model, the geographic range of the pest expanded significantly into northern Europe. The area predicted in the worst-case scenario of parameter exploration corresponded well to a +2 °C climate change increase in temperature, which makes the worst-case scenario unlikely. Considering that (i) the simulated area suitable for establishment that was obtained with the estimated parameters matched well with those locations where \textit{B. tabaci} has been reported, and (ii) the probability associated with the scenario less favourable to \textit{B. tabaci} is very low and the probability associated with the scenario more favourable to \textit{B. tabaci} is low, the authors conclude that the estimated biodemographic curves are reliable and the model is only slightly affected by biological parameter uncertainty.

Exploration of the effect of the climate change scenario revealed that, even when a systematic perturbation of +1 and +2 °C is added to the current climate, northern European countries are unlikely areas for \textit{B. tabaci} establishment because of limiting thermal conditions. In northern European countries with extensive

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure6}
\caption{Spatial distribution of the simulated \textit{B. tabaci} population pressure in Europe, starting from an initial condition of 0.1 individuals plant$^{-1}$ on 1 March, under a climate scenario with an increase of +2 °C from the current temperatures. Population pressure is the average number of adults per plant and time unit calculated for the period of the year in which the adult abundance is over 1 individual plant$^{-1}$.}
\end{figure}
greenhouse agriculture, spread from greenhouse areas to the field will be only seasonal.

PBDMs such as the one developed here for the study of the potential establishment of B. tabaci can be applied to other organisms and offer significant advantages, especially when the potential distribution of invasive alien species is to be assessed. Because PBDMs are mechanistic, demographic processes of invasion can be explained from life history strategies described at the individual level.\textsuperscript{17,19} PBDMs are transparent, and the biologically relevant features used to project the population dynamics are mathematically described in detail. PBDMs are based on well-recognised and accepted models in mathematical population ecology, and, because they are realistic and flexible, they can be applied to a variety of species and systematic groups.\textsuperscript{12} Because PBDMs are biophysical models, they can consider how the physiology related to life history strategies is conditioned by biotic and abiotic environmental variables. A further great advantage of using PBDMs is that they permit assessment and quantification of sources of variability and uncertainty.\textsuperscript{20} PBDMs produce quantitative outputs (numbers or biomass), and the information on the spatiotemporal population abundance is biologically meaningful because it can be directly related to impact or used to define ecological properties of populations such as resistance and resilience, or to predict the effect of perturbations (e.g. environmental and biological stochasticity). Assessing population development/abundance is also fundamental to the design of pest management strategy and tactics.

ACKNOWLEDGEMENTS

The authors thank the members of the EFSA Panel on Plant Health and specifically the working group on Bemisia tabaci for helpful contributions and critical discussions.

APPENDIX 1

Model equations

In the model presented here, the population structure of B. tabaci is composed of two stages: immature and adult. Denote by $x$ the physiological age of an individual, i.e. the percentage of development in a stage, and by $\phi_i(t,x)$ dx the number of individuals at stage $i$ at time $t$ with physiological age in $(x,x+dx)$, $i=1,2$ ($i=1$ for immature and $i=2$ for adult).

The population dynamics is described by the system of von Foerster equations

$$\frac{\partial \phi_i(t,x)}{\partial t} + \sigma_i(t) \frac{\partial \phi_i(t,x)}{\partial x} + \mu_i(t) \phi_i(t,x) = 0, \ x \in (0,1], \ i=1,2$$

with the boundary conditions

$$\sigma_i(t) \phi_i(t,0) = \phi^i(t), \quad i=1,2$$

and the initial conditions

$$\phi_i(0,x) = \tilde{\phi}_i(x), \quad i=1,2$$

where $\sigma_i(t)$ and $\mu_i(t)$ are development and mortality rate, respectively, depending on time through temperature, and $\tilde{\phi}_i(x)$ is the initial distribution for stage $i$.

The egg production flux, $F^1(t)$, is given by

$$F^1(t) = \int_0^1 \eta(t) fec(t) \phi^2(t,x) \ dx$$

where $fec(t)$ is the number of eggs produced by a female in the time unit and depends on time through temperature, and $\eta(t)$ is a function dependent on adult development rate.

The flux of individuals from stage 1 to stage 2, $F^2(t)$, is given by

$$F^2(t) = \sigma^1(t) \phi^1(t,1)$$

For more details on von Foerster equations in population dynamics, see Hoppensteadt,\textsuperscript{86} Curry and Feldman,\textsuperscript{18} Metz and Diekmann\textsuperscript{19} and Di Cola et al.\textsuperscript{20} For a discretisation of von Foerster equations to solve the problem numerically, see Di Cola et al.\textsuperscript{20} and Buffoni and Pasquali.\textsuperscript{21}

APPENDIX 2

Biodemographic functions and parameter estimation

Development rate function

Different development rate functions have been reported for B. tabaci in the literature: Bonato et al.\textsuperscript{23} used a Logan function, while in Muñiz and Nombela\textsuperscript{38} the development rate function of B. tabaci was described using a Lactin function. For the functional form of the Logan and Lactin functions, see Kontodimas et al.\textsuperscript{87} Here, the development rate function is modelled using the function proposed by Brière et al.\textsuperscript{88} because of the advantages it offers in terms of interpolation of the experimental data:

$$\sigma(T) = aT (T-T_{inf}) \sqrt{T_{sup}-T}$$

Data collected in the literature are fitted with the function $\sigma(T)$ using the nlinfit MATLAB function based on a non-linear least-squares regression method. The estimated parameters $a$, $T_{inf}$ and $T_{sup}$ for the immature and adult are reported in Table A1. For the adult stage, development has to be interpreted in terms of ageing, and the development period defines the maximum lifespan of the adult.

The MATLAB tool nlintool for non-linear fitting and prediction makes it possible to obtain the 95% confidence bands for the estimated curves on the basis of published data on developed rate (development period\textsuperscript{-1}) at different temperatures. Confidence bands are useful when considering the effect of experimental heterogeneity in the model. The 95% confidence bands are reported in Fig. A1 for immature and adult development rate functions.

Mortality rate function

A finite survival rate function of the polynomial type

$$sur(T) = p_1T^2 + p_2T + p_3$$

Table A1. Parameters of the immature and adult development rate function

<table>
<thead>
<tr>
<th></th>
<th>$a$</th>
<th>$T_{inf}$ (°C)</th>
<th>$T_{sup}$ (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Immature</td>
<td>0.00003502</td>
<td>9.67</td>
<td>37.05</td>
</tr>
<tr>
<td>Adult</td>
<td>0.00005112</td>
<td>12.07</td>
<td>36.26</td>
</tr>
</tbody>
</table>

where $F^1(t)$ is the number of eggs produced by a female in the time unit and depends on time through temperature, and $\eta(t)$ is a function dependent on adult development rate.

The flux of individuals from stage 1 to stage 2, $F^2(t)$, is given by

$$F^2(t) = \sigma^1(t) \phi^1(t,1)$$

For more details on von Foerster equations in population dynamics, see Hoppensteadt,\textsuperscript{86} Curry and Feldman,\textsuperscript{18} Metz and Diekmann\textsuperscript{19} and Di Cola et al.\textsuperscript{20} For a discretisation of von Foerster equations to solve the problem numerically, see Di Cola et al.\textsuperscript{20} and Buffoni and Pasquali.\textsuperscript{21}
is considered, where \( p_1, p_2 \) and \( p_3 \) are constants that have to be estimated. The temperature-dependent mortality rate function is given by

\[
\mu(T) = \begin{cases} 
-\sigma(T) \cdot \log [\text{sur}(T)] & \text{for } T \leq T_{\text{inf}}, T_{\text{sup}} \\
-\sigma(T_{\text{inf}}) \cdot \log [\text{sur}(T_{\text{inf}})] (T_{\text{inf}} - T_{\text{inf}} + 1) & \text{for } T < T_{\text{inf}} \\
-\sigma(T_{\text{sup}}) \cdot \log [\text{sur}(T_{\text{sup}})] (T_{\text{sup}} - T_{\text{sup}} + 1) & \text{for } T > T_{\text{sup}}
\end{cases}
\]

where \( \sigma(T) \) is the development rate function described above.

The mortality rate function is defined piecewise. In the interval between the two development thresholds \( [T_{\text{inf}}, T_{\text{sup}}] \), the definition depends on the development rate function. Outside this interval, the mortality rate is a function that increases as the temperature increases \((T > T_{\text{sup}})\) or decreases \((T < T_{\text{inf}})\).

Owing to the density dependence regulation, the mortality has a further term depending on the number of individuals when the population abundance is greater than the threshold of 100 individuals. The temperature- and density-dependent mortality rate function \( m(T) \) is defined as follows:

\[
m(T) = 0.03 \left( \mu(T) \right) \begin{cases} 
\mu(T) & \text{if } N(t) \leq 100 \\
0.6 \log [N(t) - 100 + 1] + 1 & \text{if } N(t) > 100
\end{cases}
\]

where \( N(t) \) is the total population at time \( t \) per spatial unit (single plant).

The survival rate functions for immature and adult were estimated by fitting all the data reported in the literature using the nlinfit MATLAB function. The best fit of immature and adult survival was obtained using the values of parameters of survival function indicated in Table A2. For immature \( T_{\text{inf}} = 10, T_{\text{sup}} = 37.049 \), for adults \( T_{\text{inf}} = 12.4, T_{\text{sup}} = 36.259 \).

### Appendix 3

#### Hourly temperature reconstruction

Starting from the WORLDCLIM database of monthly maximum and minimum temperature averages, hourly temperatures have been reconstructed.

The algorithm used for reconstruction is based on the equation published in de Wit. By a first step, the daily maximum and minimum temperatures have been obtained, fitting the monthly maximum and minimum temperatures by a third-order polynomial.

Denote by \( T_{\text{min}} \) and \( T_{\text{max}} \) the minimum and the maximum temperatures of the current day, by \( T_{\text{max}+1} \) the maximum temperature of the previous day, and by \( T_{\text{min}+1} \) the minimum temperature of the next day. Let \( h_{\text{lev}} \) be the time of sunrise. The temperature at hour \( h \) is given by

\[
T(h) = \begin{cases} 
T_{\text{max}+1} + T_{\text{min}} & \text{if } 0 \leq h < h_{\text{lev}} \\
T_{\text{max}+1} - T_{\text{min}} & \text{if } h_{\text{lev}} \leq h < 14 \\
T_{\text{max}+1} + T_{\text{min}} - 2 & \text{if } 14 \leq h < 24
\end{cases}
\]
Figure A2. Estimated survival rate curves (continuous lines) as a function of temperature (in °C) with 95% confidence bands (dotted lines) for immature (left) and adult (right).

Table A3. Parameters of the fecundity rate functions

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Temperature</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\alpha$</td>
<td>143.53</td>
</tr>
<tr>
<td>$T_{\text{min}}$ (°C)</td>
<td>13.42</td>
</tr>
<tr>
<td>$T_{\text{mid}}$ (°C)</td>
<td>11.11</td>
</tr>
</tbody>
</table>

Figure A3. Estimated fecundity rate curve (continuous lines, in eggs female$^{-1}$ day$^{-1}$) as a function of temperature (in °C) with 95% confidence bands (dotted lines).

It is assumed that the maximum temperature occurs at 2:00 p.m., and the minimum temperature at sunrise.

APPENDIX 4

Exploration of climatic scenarios

Climate scenarios are frequently used for the analysis of biological and ecological system responses to change in environmental forcing variables. Nonetheless, projections carried out by GCMs are subject to inaccuracies of different kinds. Maslin and Austin$^{80}$ signalled a paradox arising from continuous increasing climate model complexity (more equations and parameters that describe processes). This feature brings increasing uncertainties in model projections. Regional downscaling emphasises uncertainties on the local scale, and model outputs sometimes give very different indications.

Hence, to avoid the uncertainties resulting from these inaccuracies, the present authors adopted a simpler approach based on a systematic perturbation of time series of temperature to analyse the responses of the $B. tabaci$ population dynamics model to temperature change.

The climate models outputs (used to generate climate scenarios) have been verified by the IPCC (IPCC, 2007: http://www.ipcc.ch/publications_and_data/ar4/wg1/en/ch8s8-1-2-1.html) and by researchers from other research institutions and organisations. Results of these verifications show that climate model outputs do not provide a deterministic forecast, but rather only a probabilistic range of values of meteorological driving variables (IPCC 2007, IPCC-TGICA, 2007: http://www.ipcc-data.org/guidelines/TGICA_guidance_sdciaa_v2_final.pdf).

The verification of climate models for the last 150 years, performed by Anagnostopoulos et al.,$^{91}$ found uncertainties in mean values and standard deviation of yearly mean temperatures. In particular, the interannual fluctuations of temperature are strongly smoothed in climate models with respect to the observations. The uncertainties on temperature (mean and standard deviation) are low on the global scale, but errors increase at the continental or lower scale.$^{75}$

The highlighted uncertainties in climate model outputs strongly affect the possibility of obtaining reliable scenarios on future spatial and temporal patterns of temperature for the purpose of the analysis of $B. tabaci$ population dynamics in response to temperature change. The sensitivity analysis of $B. tabaci$ population dynamics to temperature performed here is based on a perturbation approach in which the forcing variable (temperature) is perturbed by adding 1 and 2 °C to calculated hourly temperature. The thermal perturbation is considered for a 20 year timescale.

The worst-case scenario, a temperature perturbation of +2 °C, is the outcome of an analysis of the last 50 years of European climate. In particular, from second half of the 1980s, the Euro-Mediterranean area has been affected by an abrupt climatic change due to a trajectory reconfiguration of the westerlies. On a
macro scale, this phenomenon has been highlighted by the strong positive values of the NAO index (North Atlantic Oscillation), beginning in the second half of the 1980s.92–94 Moreover, the historical recurrence of phases with persistent positive NAO values in the last millennium has also been highlighted by means of suitable paleoclimatic proxies.95 This change in circulation caused an abrupt increase of +1°C in yearly mean temperature (1988–2010) relative to the 1973–1987 period.96 On a lower spatial scale (regional) and during the spring period, the thermal anomaly rose to 2°C. On the basis of this evidence, it was assumed that a similar kind of circulation change might happen in the next 20 years.

Another important point supporting the choice of +2°C as the temperature perturbation is the consistency of this value with climate projections. The major part of these projections is based on increased concentrations of greenhouse gases on a long timescale. An increase in temperature in the range 1.4–5.8°C by the year 2100 has been obtained for a stable CO2 increase (IPCC, 2007), or 1.5–4.5°C for a doubling of atmospheric CO2 concentration (otherwise known as the ‘equilibrium climate sensitivity’, see IPCC-TGICA, 2007). These projections are referred to a very long timespan, and hence the results cannot be interpolated for a period of 20 years.

A valid support to +2°C as the temperature perturbation is obtained from short-term climate projections referred to Europe. For example, the chosen perturbation is consistent with a projection provided by the PESETA Project (http://peseta.jrc.ec.europa.eu/docs/ClimateModel.html). This project considers change in thermal and rainfall patterns from 2010 to 2040. Projections have been obtained from the model RCA3/ECHAMS of the Danish Meteorological Institute [project PRUDENCE: Prediction of Regional scenarios and Uncertainties for Defining European Climate change Risks and Effects (http://prudence.dmi.dk/)]. The PESETA Project estimates a rise in temperature of +3°C in 30 years for summer temperatures in most of the southern European regions, and an increase of +2°C for most of the northern European regions.

According to the above historical evidence, and relating to the European climate scenario, the decision of +2°C perturbation seems to be a representative description of possible extreme thermal increase in Europe over the next 20 years. For the purpose of the analysis of B. tabaci population dynamics responses to temperature change, this can be considered to be the climate worst-case scenario.

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