CLIMATE CHANGE AND AGRICULTURE RESEARCH PAPER Finding potential high-yield areas for Mexican maize under current and climate change conditions

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SUMMARY

Analyses of the geographic patterns of wild species abundance have been carried out in studies ranging from those interested in hot-spots for biodiversity and conservation to basic ecological analyses. Based on the methodological approaches used, the present study searched for areas of higher yield among native races of maize in Mexico, its centre of origin and diversification. Ecological theory suggests that population fitness, and thus abundance, is maximal at the centroid of the multi-dimensional ecological niche of a particular species, and decreases progressively as distance from it increases. In the present study, yield was used instead of abundance, assuming it to be higher under optimal environmental conditions. It was assessed whether nine Mexican maize races exhibited higher yields in areas that are ecologically closer to their niche centroid (NC), or a niche optimum (NO) that did not always coincide with the geometric centroid. Environmental and geographical clusters for each race were also created to identify additional NOs in widely distributed races. All races showed significant correlations between yield and distance, both to the NC or NOs, but in only six of them was the chosen model better than the null model. Three races and two sub-groups were selected for projection under climate change conditions: Celaya ($R^2 = 0.288$), Celaya 3 ($R^2 = 0.288$) 0.774), Vandeño ($R^2 = 0.277$) and Vandeño 2 ($R^2 = 0.466$) and Tepecintle ($R^2 = 0.537$). The Celaya race improved with environmental clustering, while Vandeño improved under geographical clustering. Finally, Tepecintle was projected at a race level. Projections under climate change suggested some potential areas for high yields in the future. It was concluded that the approach used in the present paper, considering the distance to NC/NO as a predictor of maize landrace yield, may contribute important information to agro-ecological projects.

INTRODUCTION

Maize is one of the three key crops around the globe in terms of production and as a dietary staple (Nuss & Tanumihardjo 2010). Mexico, its centre of origin and diversification (Kato *et al.* 2009), harbours around 59 native races and thousands of local varieties (created through natural and human selection over thousands of years), representing the most important genetic reservoir for this species in the world

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(Ruiz Corral *et al.* 2008) and the basis of Mexican culture and cuisine (Barros & Buenrostro 2011). Although the taxonomic term 'race' is not commonly used in agricultural plants, in the case of maize it has been very useful to assist with organizing the species' great diversity (Anderson & Cutler 1942). A well-accepted definition of race is as follows: 'a group of related individuals with enough characteristics in common to permit their recognition as a group' (Anderson & Cutler 1942). Some Mexican maize experts suggest that since there is a continuous genetic flow between maize populations, they are not

isolated enough to create sub-species and consequently it is better to classify them as races (Ortega Paczka, personal communication). It is also known that within each race a wide range of variants exists (Rocandio-Rodríguez *et al.* 2014).

In Mexico, c. 0.85 of farmers growing maize are smallholders (<5 ha producing 1.6-2 t/ha) (Cruz Delgado et al. 2012) who grow mostly native races using open-pollinated populations of traditional 'owned' seed stocks and agro-technologies, with yields depending strongly on environmental conditions (Turrent Fernández 2008). Yield depends on the biological characteristics of the race, such as cycle length (Chávez-Servia et al. 2004), but each race presents a wide range of yield variation across the race distribution, thus exhibiting plasticity depending on a range of environmental growth conditions (Acevedo et al. 2011; CONABIO 2011).

In recent years, some efforts have been focused on modelling geographic patterns of fitness using presence-only data for wild species (VanDerWal et al. 2009; Tôrres et al. 2012; Yañez-Arenas et al. 2012, 2014; Martínez-Meyer et al. 2013; Nagaraju et al. 2013; Lira-Noriega & Manthey 2014). A study carried out in India with the economically important tree Myristica malabarica showed that highly suitable sites projected with ecological niche modelling were correlated significantly with higher plant fitness. To evaluate fitness, Nagaraju et al. (2013) evaluated functional traits of the plant, such as fluctuating asymmetry, leaf weight, recruitment and genetic variability. Other studies have tested whether ecological niche modelling outcomes are related to the abundance (as an expression of fitness) of several species, resulting in marginal success in the best cases (VanDerWal et al. 2009; Tôrres et al. 2012). On the other hand, a novel method to model geographic patterns of abundance based on the ecological niche theory, named the Distance to the Niche Centroid approach, was proposed and tested by Martínez-Meyer et al. (2013), and implemented by Yañez-Arenas et al. (2012). According to Maguire (1973), population fitness should be maximal where the highest birth rate and the lowest death rate coincide in the multi-dimensional niche space; he proposed that such optimal conditions should occur at or near the centroid of the niche hypervolume for each species. Based on this body of theory, Martínez-Meyer et al. (2013) successfully tested the hypothesis that population abundances of 11 species of vertebrates follow a centre-abundant pattern in

ecological space, where maximal abundance occurs around the niche centroid (NC) and decreases progressively as populations depart from this point. An implementation of this procedure for the white-tailed deer (Odocoileus virginianus) produced robust spatially explicit models of abundance in two regions of Mexico (Yañez-Arenas et al. 2012). Other studies have already incorporated this concept to infer potential risk areas for snakebite (Yañez-Arenas et al. 2014) and to test whether there was a relationship between genetic diversity and distance to the NC (Lira-Noriega & Manthey 2014). However; it is theoretically feasible that optimum conditions for a given species are not always at or near the central values of specific critical ecological variables (i.e. NC); for example, some monkeyflower species have higher fitness at their marginal elevation range (Angert 2009). Consequently, the niche optimum (NO) may be off the centre of the niche hypervolume.

Following this line of thinking, it was hypothesized that Mexican maize races will exhibit higher yields in areas that are ecologically closer to the optimum of their ecological niches (NO), which may or may not coincide with their centroids (NC). Even when yield does not necessarily reflect higher fitness for the plant (as it has been selected to cover human needs and not to reach its own 'biological success'), both natural and human selection have operated under specific environmental conditions and one of the key traits targeted in the latter is yield (Herrera-Cabrera et al. 2004; Yamasaki et al. 2005). Today, each race is distributed across a range of environmental conditions, which may be considered equivalent to its ecological niche (Ruiz Corral et al. 2008; Ureta et al. 2012; Ruiz Corral et al. 2013) and where it should be possible to find an NC/NO. Furthermore, higher yields have already been documented for cereals (such as maize, wheat and oats) under specific optimum environmental conditions for the plant (Boyer 1982).

The present study tested whether there was a relationship between yield and the distance to the NC/ NO of nine different local Mexican maize races for which sufficient and relatively well-distributed yield data across their geographic range were found. Additionally, geographically and environmentally homogeneous clusters were identified for each race studied and the same analyses carried out to evaluate whether different NOs may exist in one single race. Finally, the yields of races Celaya, Vandeño and Tepecintle were projected under current and future climate change conditions (2050) at the race level, as well as the clusters with higher fitness for Celaya and Vandeño. The information generated in the present study may aid in identifying which factors are critical for yield in maize races. Thus, in turn, this may help with searching for new areas of production for a particular race that mimic their current optimal yield conditions, or may contribute with variables and models to estimate future yields under different conditions and in different geographical areas. If it is possible to demonstrate that this method works with domesticated plants under traditional management, then it is a step forward in conservation planning for key food and genetic resources in the face of environmental change.

METHODS

Occurrence and yield data

Occurrence records from 1980 to 2010 (Acevedo et al. 2011) and yield information from 2000 to 2010 were obtained from a database compiled by the Mexican Commission for Biodiversity (CONABIO 2011). The yield information was taken from a column named 'rendimiento uniformizada1': as indicated by the document accompanying this database, this column takes into account all information that was taken from the field and was assigned a single numerical number in kg/ha units (in cases where the information constituted intervals or some other kind of measurement, it was transformed into the corresponding units to unify quantities and make them comparable). For some localities two types of data could be found: uniformizada1 and uniformizada2, representing two different seasons (spring/summer and autumn/winter). The column uniformizada1 was the only one taken into account in the present study, since this is the season in which >0.80of Mexican maize is produced. At present, there is no official institution gathering exact estimates of native race yields, so farmers were interviewed and provided an approximate value of their annual productivity. However, the method proposed in the present study is expected to work even if approximate yield data is available. Consequently, interviews might still give valuable information at a race level that can provide an idea of the yield magnitudes and variation along their distribution ranges, under current and future conditions. Furthermore, it was decided to take out statistical outliers (mean ± 2 sD) that made no biological or agronomic sense. For example, race Bolita presents an average yield of 2.5 t/ha, but two values out of 35 were >8 t/ha. These two values are obviously statistical outliers as yields above 8 t/ha would only be expected in the high production areas of hybrid maize varieties using modern agro-technology, such as irrigation, fertilizers and pesticides (Cruz Delgado et al. 2012). Using this procedure no more than three records per race were discarded (one for Celava out of 86, one for Cónico Norteño out of 235, three for Tabloncillo out of 63, and three for Tuxpeño out of 534; for the remaining five races all records were taken into account for the analysis). The races analysed were: Bolita, Celaya, Cónico Norteño, Olotillo, Ratón, Tabloncillo, Tepecintle, Tuxpeño y Vandeño (Table 1). To characterize the bioclimatic profile of these races a 1 km² resolution climatology map generated for Mexico covering 1980-2009 was used (Cuervo-Robayo 2014).

Calculating distances to ecological centroids and optima

For all data relating to presence, the value for each of the 19 bioclimatic variables evaluated was obtained (1: annual mean temperature, 2: mean diurnal range, 3: isothermality, 4: temperature seasonality, 5: maximum temperature of the warmest month, 6: minimum temperature of the coldest month, 7: temperature annual range, 8: mean temperature of the wettest quarter, 9: mean temperature of the driest guarter, 10: mean temperature of the warmest quarter, 11: mean temperature of the coldest quarter, 12: annual precipitation, 13: precipitation of the wettest month, 14: precipitation of the driest month, 15: precipitation seasonality, 16: precipitation of the wettest quarter, 17: precipitation of the driest quarter, 18: precipitation of the warmest quarter and 19: precipitation of the coldest quarter). All these variables were included even though the races are mostly grown during the spring-summer season, because it has been shown that climatic variables at other times of the year are still significantly related to the distribution of most races (Ureta et al. 2013). The ecological niche modelling program MaxEnt (Phillips et al. 2006) was then used to identify which variables contributed most to the distribution of each race and these were used for the correlation analysis between the yield and distance to NC and NO. Only bioclimatic variables were used in the present study because the aim was to project the effect of climate change on the distribution of future yield, but it should be noted that other environmental factors (e.g. soil and slope) are also

Races	I's P-value	Best Models		AIC			R^2		$R^2 P$ value		Slope		Sample size	
		С	0	0	С	Null	С	0	С	Ο	С	0	Occur	Yield
Bolita	NS	LN	Linear	440 .583	438·421	443 · 076	0.226	0.158	0.013	0.33	Ν	n	104	26
Celaya	<0·001	Linear	LN	1443-497	1453.318	1469.329	0 ∙241	0.288	<0.001	<0.001	Ν	n	260	85
Cónico Norteño	<0.001	Linear	LN	3876.597	3872-933	3891.457	0·229	0·221	<0.001	<0.001	Р	р	512	234
Olottillo	<0.001	LN	Linear	2821.055	2819.690	2821.215	0.223	0.233	<0.001	<0.001	Р	р	474	177
Ratón	<0.001	Linear	LN	1527.343	1526.899	1525.405	0.091	0.094	0.004	0.03	Ν	n	236	91
Tabloncillo	<0.001	Linear	Linear	1052.921	1052.824	1055.450	0.153	0.138	0.002	0.002	Ν	n	276	60
Tepecintle	<0.001	Linear	LN	1062.686	1063.78	1062.97	0.531	0.537	<0.001	<0.001	Ν	n	252	68
Tuxpeño	<0.001	Linear	Linear	8971.722	8972.205	8970.274	0.150	0.151	<0.001	<0.001	Р	n	1263	530
Vandeño	<0.001	Linear	Linear	1348.617	1348-226	1350.461	0·277	0.267	<0.001	<0.001	Ν	р	194	78

Table 1. Linear models describing the relationship between yield and distance to the ecological niche centroid/optimum for nine maize races

NS, not significant.

I's *P* value: *P* value of the Moran's I test on spatial autocorrelation (H_0 : no spatial autocorrelation exists), Best model: best of two models; Linear: Yield = $\beta_0 + \beta_1 \times$ Distance, LN: Yield = $\beta_0 + \beta_1 \times \ln$ (Distance), C: centroid, O: optimum, AIC: Akaike Information Criterion of the best model, bold: highlights a model with a smaller AIC than the one of the null model (Yield = β_0), R^2 : coefficient of determination associated to the best model, $R^2 P$ value: R^2 significance *P* value, Slope: sign of the value of the β_1 estimate of the best model (n, negative; p, positive), Sample size: number of occurrence and yield data. important for characterizing the niche of a species such as maize (Ureta *et al.* 2013; Dyer *et al.* 2014).

Once the bioclimatic variables were associated with the presence data, distances were calculated to the NC (Yañez-Arenas et al. 2012; Martínez-Meyer et al. 2013) and NO. The NC was calculated as the standardized mean assuming a normal distribution of the range of each bioclimatic variable where the race was present. Although ecological theory suggests that higher suitability and consequently higher fitness should be found in the NC (Maguire 1973), it may be the case that the optimal conditions do not coincide with the centroid because a species may be adapted to extreme (or at least off-centre) conditions for a given environmental variable (Angert 2009). Therefore, the Euclidian distance was measured to a multi-dimensional point, the NO. To calculate NO, a function for building the response curves of the maize race presence records with respect to each environmental variable was used and implemented in MaxEnt, which identified the value of each environmental variable for which the presence response for that taxon was most frequent. To run MaxEnt, 0.70 of the dataset was used for training the model and the remaining 0.30 to test it. Validation of the model was performed with a partial-receiver operating characteristic (ROC) test (Peterson et al. 2008), which has the advantage of taking into account only the projected area when using the niche algorithm and does not evaluate omissions and commissions (i.e. false positive errors in Ecological Niche Modeling) equally (Peterson et al. 2011). Omissions should be punished harder. Once the NC and NO values were identified, all bioclimatic variables were z-standardized, thus the NC (multivariate mean) was zero. Then, the multidimensional Euclidian distance of every occurrence record to the NC and the NO was calculated. Thus, the distance (d) of the data point P_i to the niche (N = NC or NO) is:

$$d(P_i,N) = (S_j(P_{i,j} - N_j)^2)^{1/2}$$

where $P_{i,j}$ and N_j are the values of the *j*th bioclimatic variable of the data point P_i and NC or NO, respectively. The calculation of the Euclidian distance to NC or NO was done for every race record under current climatic conditions. However; under future climatic conditions the potential distribution area was necessary; consequently, the calculation of the Euclidian distance was done for every 10 km² (the resolution of the climate change map), to show where the race was modelled to be distributed in the future (see below).

Relationship between yield and distance to ecological centroids and optima

Once the distance to the NC and NO were obtained for every presence datum point, those occurrences for which yield data were available were taken to model the relationship between yield and distance to the NC/NO. Since spatial autocorrelation existed among the data for each race (using the Moran's I test, Table 1) (Fortin & Dale 2005), three autoregressive models were fitted:

(a) Yield = β_0 , (b) Yield = $\beta_0 + \beta_1 \times \text{Distance, and}$ (c) Yield = $\beta_0 + \beta_1 \times \ln$ (Distance).

For those cases where the Moran's I test showed existing autocorrelation, a conditional autoregressive model was used to deal with univariate models (Gelfand & Vounatsou 2003); and the spdep package in R was used to fit these models (Bivand *et al.* 2014; R Development Core Team 2014). The Akaike Information Criterion (AIC, Akaike 1974) and determination coefficient (R^2) were calculated to perform model selection and evaluate model fitting, respectively. It was also evaluated if differences could be found between models using the distance to the NC and to the NO.

Potential distribution maps under current and future climatic conditions

Most races evaluated are distributed widely throughout Mexico, encompassing different climates. Consequently, data were split into environmental and geographic clusters. These clusters were created through the 'partitioning around medoids' clustering algorithm. This algorithm has the advantage of not requiring an a priori fixed number of clusters. The environmental clustering was performed with 13 of the 19 bioclimatic variables (1, 5, 6, 8, 9, 10, 11, 12, 13, 16, 17, 18 and 19) because these showed low correlations among them and with the others. The algorithm was implemented through the fpc package in R (Hennig 2010; Pinheiro et al. 2011). The environmental clustering helped to identify sub-groups with similar ecological profiles, while geographic clustering identified sub-groups that, due to their geographic closeness, are expected to be genetically similar. It was decided to create clusters because there is evidence supporting an important ecological and genetic variation within races. If clusters improve correlations between yield and distance to the NC or NO, it indicates that there is more than one NC or NO within a race and there is an important amount of yield variation (Herrera-Cabrera *et al.* 2004).

To project occurrences of higher yields under current and future climatic conditions, three races were chosen based on their AIC and R^2 values: Celaya, Vandeño and Tepecintle. An environmental cluster for Celaya (Celaya 3) and a geographic cluster for Vandeño (Vandeño 2) were also projected. No cluster was projected for Tepecintle because the proportion of explained deviance was greater at the race level than with any cluster. For the current climate conditions (1980-2011), the same climatology as above was used, and for the future (2040-2069, hereafter called 2050), climatologies drawn from the Moscow Forestry Sciences Laboratory (http://forest.moscowfsl.wsu.edu/climate/) were used. The general circulation model (GCM) used was the Hadley Centre Global Environmental Model version 1 (HadGEM1), developed in the UK: this has been evaluated by Mexican climatologists as one of the best in representing Mexico's current climate and thus produces reasonable future climatic scenarios (Conde Álvares & Gay Garciá 2008). The worst case emission scenario A2 ('business as usual') was assessed (IPCC 2007). Although there is awareness of the new scenarios proposed by the IPCC, 'Representative Concentration Pathways' (IPCC 2014), it was decided not to use these models, because they have not been downscaled specifically for Mexico to a 1 km² resolution. Until now, Mexican government downscaling efforts (10 km²) have taken place through the Reliability Ensemble Averaging method, that takes into account all General Circulation Models giving higher weight to those performing better in specific areas of the country (Ibarra-Cardeña et al. 2013).

Current climatology was generated for Mexico using the thin-plate spline technique available in the ANUCLIM program (http://fennerschool.anu.edu.au) at 1 km² spatial resolution (Cuervo-Robayo 2014). Under current climatic conditions, the potential distribution area was modelled with the only purpose of calculating the proportion of potential distribution area that would be lost in the future. The distance to NC/NO was calculated only with present data as explained above.

To locate future possible regions of higher yields, firstly future potential distribution areas were modelled

by running ten model replicates for each emission scenario with the MaxEnt algorithm (Phillips et al. 2006). Replicate probability output was then converted into a binary map using the maximal threshold value that minimized the training and test omission rate. As a way to reduce uncertainty, binary maps were assembled and the final map was the consensus of all ten maps (Araújo & New 2007). A final map was combined with the 19 bioclimatic variables obtained for 2050 under the emission scenario A2. The bioclimatic profile for each pixel of the potential distribution area was obtained and the distance to the current NC or NO was calculated (assuming that these values will be maintained through time). In this way, the geographic areas with potentially higher yields in the future for the races Celaya, Vandeño, Tepecintle and the corresponding clusters were identified.

RESULTS

Relationship between yield and distance to ecological centroids and optima

The NO was identified via ecological niche modelling. All ecological niche models presented good performance as indicated by their high average Partial ROC: AUC ratios and *P* values that were always highly significant (P < 0.001) (see Table S1 in the Supplementary Material for details, available from http://journals.cambridge.org/AGS).

Moran's I tests performed on the data for each race showed that negative autocorrelation existed among the data and was statistically significant (P < 0.001) in eight of the nine races (Table 1). These negative autocorrelations persisted for all the ecological and geographic clusters (Tables S1 and S2, available from http://journals.cambridge.org/AGS). As expected, when geographic clustering was performed, autocorrelation within each of these clusters was lost more often than within ecological clusters.

At the race level, all showed a significant (P < 0.015) correlation between yield and the distance to NC and/ or NO. However, the best models were selected over the null model (model a) in only six of the nine races: Bolita, Celaya, Cónico Norteño, Olotillo, Tabloncillo, Tepecintle and Vandeño (Table 1). From these six races, only Cónico Norteño presented a positive correlation with NC and NO (Table 1). Bolita, Celaya, Olotillo, Tepecintle and Vandeño presented negative correlations between yield and distance to both NC and to NO. For most of these races, R^2 values were higher when evaluating the distance to the NC than to the NO (Table 1). The exceptions were Celaya and Tepecintle, which were also the races presenting the highest R^2 values (0.288 and 0.537, respectively).

When carrying out the environmental cluster analysis, sub-groups for some races presented better models than the null model and higher proportions of explained deviance. For example, Celaya increased its R^2 up to 0.774 in one of its environmental clusters (Celaya 3) (Table 1, Table S2, available from http:// journals.cambridge.org/AGS). In races that showed no significant correlations at the race level, such as Olotillo, Ratón and Tuxpeño, significant (P < 0.001) negative correlations were found with R^2 values of up to 0.298 in some of their sub-groups. With this clustering, it was possible to find five negative correlations and two positive ones. Higher yields for these environmental sub-groups were better explained by distance to the NO than by distance to the NC, except for Vandeño which presented only one subgroup with a significant (P < 0.001) positive correlation. The geographic clustering worked better in the subgroups of races Cónico Norteño, Tuxpeño and Vandeño (Table S2, available from http://journals.cambridge.org/AGS). In this geographic clustering, most sub-groups were better explained by distance to the NC than to the NO. For Bolita, Tabloncillo and Tepecintle the race level worked better than the clusters.

Potential distribution maps under current and future climatic conditions

To project the geographical location of higher yields under current and future conditions, three races that had significant (P < 0.001) negative correlations with models b or c that were better than their corresponding null models and with the highest R^2 values at the race level were used. Celaya and Vandeño increased their explained deviance with at least one clustering type. Consequently, Celaya ($R^2 = 0.288$) and Celaya 3 ($R^2 = 0.774$) were projected. This race and its environmental sub-group had a greater deviance explained by distance to the NO and the environmental clustering increased this proportion. However, Vandeño $(R^2 = 0.277)$ and Vandeño 2 $(R^2 = 0.466)$ were better explained by distance to the NC and the clustering that increased its R^2 value was the geographic one. Tepecintle presented an $R^2 = 0.537$ at the race level and this value did not increase with the cluster analysis. In other words, example is presented of a race that improved with the environmental clustering

(Celaya), another that improved with the geographic clustering (Vandeño) and finally a race that worked better at a race level (Tepecintle).

For Celaya, yield data indicate that higher yields (3500-7000 kg/ha) were found in northern Mexican states such as Chihuahua, Durango and westerncentral states such as Jalisco and Guanajuato. Projections under current conditions (using the correlation model explored in the present paper) also show that higher yields could potentially be found in Chihuahua, Durango but also in San Luis Potosí, Oaxaca and Chiapas, where yield data have not yet been collected for this race. In the future (2050), the potential distribution area is expected to decrease about 60% in comparison with its current distribution and that higher yields will shift towards the centre of its geographic distribution area. Projections suggest high yields in northern states such as Durango but not in Chihuahua. Higher yields are also projected in the south of San Luis Potosí, Jalisco, Michoacán, Guanajuato and in a small portion of Chiapas (Fig. 1a). On the other hand, sub-group Celaya 3 is currently distributed in the central and southern part of the potential distribution area of Celaya's race. Under current conditions, higher yields are projected in states such as San Luis Potosí, Michoacán, Guerrero and Oaxaca. Areas where higher yields could potentially be found are not very abundant (six out of 63 records). Under future conditions, the potential distribution area will be concentrated in only a few states (a reduction of about 70%), mainly in Jalisco where most potentially high-yield areas can be found. Higher yields are also expected in Guanajuato and Michoacán. Under future climate conditions, the potential distribution area in the North is expected to disappear. Some other highyield areas are projected to be found in Michoacán and Guerrero (Fig. 1a).

For Vandeño, yield data were mainly collected in Morelos, Jalisco and Chiapas. In these three states, potentially high-yield areas (3000–6000 kg/ha) were found (see Fig. S1 in the Supplementary Material available from http://journals.cambridge.org/AGS). Although most observations come from these three states, there are other records without yield information located in Sonora. The projected high-yield areas under current conditions are in Sonora but also in Oaxaca and Chiapas. Under future climatic conditions, the potential distribution area is reduced by 70% and high yields will be found in the western states of Michoacán and Guerrero. The sub-group

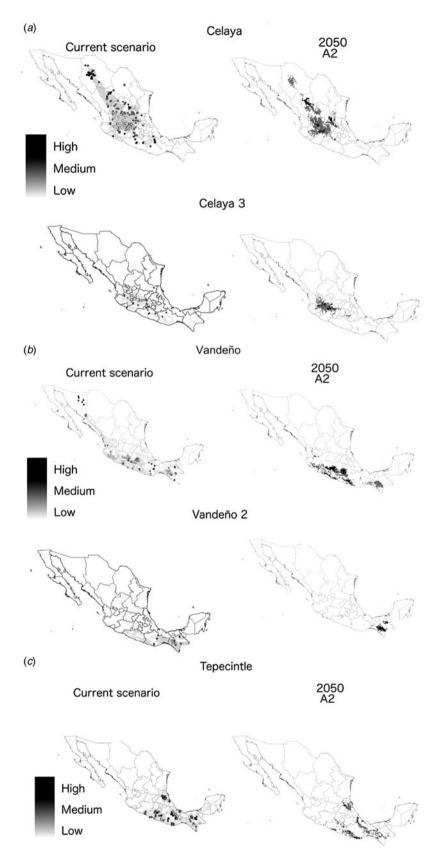


Fig. 1. Projection of areas with higher yields under current and future conditions for three maize races and their clusters.

Vandeño 2, whose distribution is in the southern part of the country, is projected to potentially have higher yields in both states where observations have taken place: Oaxaca and Chiapas. Under future climatic conditions, distribution decreases by 85% and higher yields can only be found in Chiapas (Fig. 1*b*).

Finally, for Tepecintle, observations registered high yields in the three states where yield values were obtained: Veracruz, Chiapas and Oaxaca (1601-4000 kg/ha) (Fig. S1 in the Supplementary Material available from http://journals.cambridge.org/AGS). Under current conditions the model projected potential high-yield areas in the same three states and in Guerrero, where field observations with yield values do not exist. Under climate change conditions, potentially high-yield areas are found in Veracruz, Chiapas, Oaxaca and Guerrero. The distribution of potentially high-yield areas is very similar to that presented under current conditions, but it is still possible to see a small shift closer to the coast (Fig. 1c). In terms of potential distribution area there is a reduction of c. 75% (Fig. 1c).

DISCUSSION

Relationship between yield and distance to ecological centroids and optima

Ecological theory has proposed that optimal environmental conditions for a species should be close to its ecological NC, and thus abundance should follow a 'centre-abundant' pattern in ecological space (Maguire 1973). Studies have empirically demonstrated that this relationship does exist with abundance and other characteristics, such as genetic diversity (Yañez-Arenas et al. 2012, 2014; Martínez-Meyer et al. 2013; Lira-Noriega & Manthey 2014). In the present study, it is possible to observe an inverse relationship between yield and the distance to NC/NO for most races evaluated and consequently it might be a useful approach to predict areas with potential current or future high yields, in cases that such data has not been gathered. The latter is the case for many races and sites of distribution in Mexico.

The current results showed that six races presented a model where distance explained more deviance than the null model. Most of these six races had a greater R^2 value when using distance to the NC than distance to the NO. In contrast, distance to the NO better explained yield changes for Celaya and Tepecintle. Additional factors, besides the bioclimatic variables, might be playing a more important role in the

distribution of these two races and consequently the model does not embrace the entire bioclimatic range and fails to find a more realistic environmental centroid. Still, areas with the potential of having greater yields were identified by calculating the distance to NC.

On the other hand, it is intriguing why races such as Cónico Norteño had significant positive correlations between yield and the distance to NC/NO. This result has already been found with wild species (Lira-Noriega & Manthey 2014; Yañez-Arenas et al. 2014), but in this particular case it might be related to the condition of a domesticated species, in which factors such as farming technology or farmers' preferences in growing a specific race, rather than environmental factors significantly affect yield (Bellon & Brush 1993; Brush & Perales 2007; Ureta et al. 2013). Cónico Norteño is a race with a wide distribution range and probably has more than one NC or NO; consequently when it is divided into clusters, more meaningful results are recovered (see Supplementary Material Table S2, available from http://journals.cambridge.org/AGS). This race embraces an important amount of varieties adapted to different conditions and some of them probably have strong specific local adaptations, making it very difficult to assign a single optimum condition for the entire race.

Similarly, in other widely distributed races (Olotillo, Ratón and Tuxpeño), there might be more than one NC or NO, and consequently identifying sub-groups of populations that shared ecological/genetic conditions and traits, respectively, helped to better project areas with higher yields. Hence, yields of some such widely distributed races (i.e. Tuxpeño, Vandeño and Cónico Norteño) were better explained by geographic clusters rather than by environmental ones.

In maize, geographic distance has been related to genetic distance (Vigouroux et al. 2008) and consequently, populations within the same clusters are expected to have a similar genetic composition and might be similarly impacted by environmental conditions (Mercer & Perales 2010). Regardless of geographic distance, human and natural selection facilitates local adaptation to specific environmental conditions (Mercer et al. 2008; Mercer & Perales 2010) and consequently, although groups of populations might not be geographically close, their distant localities may be environmentally similar. Both types of clustering provided information on the intra-racial ecological variability that has already been reported (Doebley et al. 1985; Mercer & Perales 2010; Ruiz Corral et al. 2013). In the present study, both clustering types increased the deviance explained for some races, so it cannot be generalized that one was better than the other. But geographic clusters generated a higher number of sub-groups, which is generally not convenient because it reduces within sub-group sample size, an essential element if we are to draw robust statistical inferences. However, in the current work splitting races into bioclimatic clusters produced more accurate models. Therefore, when looking for higher yields in other cultivated species (or abundances in wild species) with large distributions, exploring the cluster approach is recommended. Positive correlations were also found in these subgroups; still, the majority of races and sub-groups presented a negative correlation between yield and NC/ NO. In general, the distance to the NC/NO approach is promising in the field of agro-ecology.

The data generated in the present study could be the basis for recommendations to peasant communities in terms of trying to plant some landraces or varieties (clusters) in areas that are predicted to maximize yield, based on their similarities to those of optimum yield predicted here for each race and/or cluster. It would be interesting to validate the models used here, by empirically testing yield in such predicted areas of high or low yield in areas that are not planted with a particular race or variety now. Once the models and predictions are tested, the tool could be iteratively improved for providing more precise recommendations. Additional efforts to recover more complete yield data in areas where the maize races and varieties are being planted by peasants today should also be made. These data will also be a valuable means to improve the tool proposed in the current work and better predict the areas of high maize yield using native varieties and races.

On the other hand, variables correlated with yield variation as a function of NC/NO distance may be used in landrace breeding programmes such as those implemented by some Mexican native maize races (Smith *et al.* 2001; Aragón-Cuevas *et al.* 2003). Such an approach could assist in predicting which traits could be used, as correlated markers, to increase yield under contrasting environmental conditions along a landrace geographic distribution.

Potential distribution and yield under current and future climatic conditions

From the nine races evaluated, three were used because they had a significant negative correlation

with the NC/NO and high R^2 values, namely Celaya, Vandeño and Tepecintle, to create maps identifying areas with higher yields under current and future climatic conditions. The future potential distribution area of the three races decreased significantly (~58-84%). Higher yields for Celaya under the future climatic scenario were concentrated in its central area of distribution. High-yield areas in the north and south of its distribution at the race and the sub-group level can be expected to disappear based on the current paper's projections. For Vandeño, at the race level the potentially high-yield areas that are present in the northern state of Sonora were projected to disappear and become concentrated in the central and southern distribution areas. For the Vandeño 2 sub-group, potentially higher yields were only projected for Chiapas. For Tepecintle, potentially high-yield areas in the future are expected to remain in the same states where they have been projected under current conditions. There is a slight shift to areas closer to the coast.

Celaya has been classified as a native race adapted to water stress in at least one part of its life-cycle (Ruiz Corral *et al.* 2013); consequently it might be able to resist harsh environmental conditions such as those expected in the northern states of Mexico in future. Even if higher yields are not projected in Chihuahua for this race in the future, medium yields are. Under such climatic conditions (as the ones projected for 2050 A2), races such as Celaya would maintain medium and high yields in certain parts of their distribution ranges, or may be used for maize production in other colder areas, which could become warmer in the future.

On the other hand, Vandeño has been identified to be adapted to temporal humid and very humid environments; and Tepecintle to very humid ones (Ruiz Corral *et al.* 2013). Consequently, higher impacts of climate change are expected for these races. Indeed, clear reductions in their projected distribution areas under the climate change scenarios that were modelled were observed.

Native races are expected to have greater success under harsher environmental conditions than improved varieties (Smith *et al.* 2001), but it is important to know how well each race may perform in different areas of the country. Agro-biodiversity plays a major role in agriculture under changing environmental conditions, and will continue to do so, because it is required for 'evolutionary resilience' (Bellon & van Etten 2014). Nevertheless, the contraction of highyield areas under climate change conditions might have an important influence on farmers' choices of what to grow where, and also how to manage their seed stocks. The projections made in the present paper can be used to advise peasants to keep different seed stocks for each local variety and make sure to keep sufficient seed for those varieties that are more resistant to water or high-temperature stress. On the other hand, peasants who rely on races adapted to milder and more humid conditions might be advised to test breeding with varieties that are more resistant to harsh conditions. Directed seed-exchange programmes among peasants managing seed for races adapted to different environmental conditions should be planned. In general, seed exchange may on its own favour adaptability under climate change scenarios. Hence, besides the locally adapted varieties, programmes to generate varieties with more genetic variation that enable them to grow under more marginal conditions should be also developed. In any case, programmes that identify genotypes and phenotypes that enable high yield under harsher conditions, in terms of water and temperature stress, for each race or variety/ cluster, should be implemented in cooperation with local communities. In each case, local seed-banks for possible harsher conditions should be promoted and tested. The type of modelling proposed in the present study is a first step towards this aim. As stated above, this type of platform should be extended to additional races, more complete yield data gathered and incorporated, and should be improved with empirical data.

SUPPLEMENTARY MATERIAL

The supplementary material for this paper can be found at http://journals.cambridge.org/AGS

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REFERENCES

Acevedo, F., HUERTA, E., BURGEFF, C., KOLEFF, P. & SARUKHÁN, J. (2011). Is transgenic maize what Mexico really needs? *Nature Biotechnology* **29**, 23–24.

- AKAIKE, H. (1974). A new look at the statistical model identification. *IEEE Transactions on Automatic Control* **19**, 716–723.
- ANDERSON, E. & CUTLER, H. C. (1942). Races of Zea mays.
 I. Their recognition and classification. Annals of the Missouri Botanical Garden 29, 69–88.
- ANGERT, A. L. (2009). The niche, limits to species' distributions, and spatiotemporal variation in demography across the elevation ranges of two monkey flowers. *Proceedings of the National Academy of Science of the United States of America* **106** (Suppl 2), 19693–19698.
- ARAGÓN-CUEVAS, F., TABA, S., CASTRO-GARCÍA, F. H., HENRNÁNDEZ-CASILLAS, J. M., CABRERA-TOLEDO, M., ALCALÁ, L. O. & RAMÍREZ, N. D. (2005). In situ conservation and use of local maize races in Oaxaca, Mexico: a participatory and decentralized approach. In *Latin American Maize Germplasm Conservation: Regeneration, in Situ Conservation, Core Subsets, and Prebreeding; Proceedings of a Workshop held at CIMMYT, April* 7–10, 2003 (Ed. S. Taba), pp. 26–38. Mexico: CIMMYT.
- ARAÚJO, M. B. & NEW, M. (2007). Ensemble forecasting of species distributions. *Trends in Ecology and Evolution* 22, 42–47.
- BARROS, C. & BUENROSTRO, M. (2011). Pozol, popo, champurrado. *Revista Digital Universitaria* **12**, 3–9.
- BELLON, M. R. & BRUSH, S. B. (1993). Keepers of maize in Chiapas, Mexico. *Economic Botany* **48**, 196–209.
- BELLON, M. R. & VAN ETTEN, J. (2014). Climate change and onfarm conservation of crop landraces in centres of diversity.
 In *Plant Genetic Resources and Climate Change* (Eds M. Jackson, B. Ford-Lloyd & M. L. Parry), pp. 137–150.
 Wallingford, UK and New York, USA: CABI Publishing.
- BIVAND, R., ALTMAN, M., ANSELIN, L., ASSUNÇÃO, R., BERKE, Ö., BERNAT, A., BLANCHET, G., BLANKMEYER, E., CARVALHO, M., CHRISTENSEN, B., CHUN, Y., DORMANN, C., DRAY, S., GÓMEZ-RUBIO, V., HALBERSMA, R., KRAINSKI, E., LEGENDRE, P., LEWIN-KOH, N., LI, H., MA, J., MILLO, G., MUELLER, W., ONO, H., PERES-NETO, P., PIRAS, G., REDER, M., TIEFELSDORF, M. & YU, D. (2014). Package 'spdep': Spatial Dependence: Weighting Schemes, Statistics and Models. Vienna, Austria: R Foundation for Statistical Computing. R Package Version 0.5-74. Available from: http://CRAN.Rproject.org/package=spdep (verified 26 June 2015).
- BOYER, J. S. (1982). Plant productivity and environment. *Science* **218**, 443–448.
- BRUSH, S. B. & PERALES, H. R. (2007). A maize landscape: ethnicity and agro-biodiversity in Chiapas, Mexico. *Agriculture, Ecosystems & Environment* **121**, 211–221.
- CHÁVEZ-SERVIA, J. L., TUXILL, J. & JARVIS, D. I. (2004). Manejo de la Diversidad de los Cultivos en los Agroecosistemas Tradicionales. Cali, Colombia: Instituto Internacional de Recursos Fitogenéticos. Available from: http://www.bioversityinternational.org/e-library/publications/detail/manej o-de-la-diversidad-de-los-cultivos-en-los-agroecosistemas -tradicionales/ (verified 26 June 2015).
- CONABIO (2011). Base de Datos del Proyecto Golbal de Maíces. Mexico, D.F.: CONABIO. Available from: http:// www.biodiversidad.gob.mx/genes/proyectoMaices.html (verified 29 July 2015).

- CONDE ÁLVARES, A. C. & GAY GARCIÁ, C. (2008). *Guía para la Generación de Escenarios de Cambio Climático a Escala Regional*. México, D.F.: Centro de Ciencias de la Atmósfera, UNAM. Available from: http://www.atmosfera.unam.mx/cclimat/Taller_CCA_INE_dic08/Guia_escenarios.pdf (verified 26 June 2015).
- CRUZ DELGADO, M. S., GÓMEZ VALDEZ, M. M., ORTIZ PULIDO, M. E., ENTZANA TADEO, A. M., SUÁREZ HERNÁNDEZ, C. Y. & SANTILLÁN MOCTEZUMA, V. (2012). *Situación Actual y Perspectivas del Maíz en México* 1996–2012. México, D.F.: Servicio de Información Agroalimentaria y Pesquera.
- CUERVO-ROBAYO, A. (2014). Fortaleciendo los Análisis de Cambio Climático en la Aplicación de los Modelos de Distribución Potencial. Ph.D. Thesis. Universidad Autónoma del Estados de México, México.
- DOEBLEY, J. F., GOODMAN, M. M. & STUBER, C. W. (1985). Isozyme variation in the races of maize from Mexico. *American Journal of Botany* **72**, 629–639.
- DYER, G. A., LÓPEZ-FELDMAN, A., YÚNEZ-NAUDE, A. & TAYLOR, J. E. (2014). Genetic erosion in maize's centre of origin. *Proceedings of the National Academy of Sciences of the United States of America* **111**, 14094–14099.
- FORTIN, M. J. & DALE, M. R. T. (2005). *Spatial Analysis: A Guide for Ecologists*. Cambridge, UK: Cambridge University Press.
- GELFAND, A. E. & VOUNATSOU, P. (2003). Proper multivariate conditional autoregressive models for spatial data analysis. *Biostatistics* **4**, 11–25.
- HENNIG, C. (2010). *Package 'fpc': Flexible Procedures for Clustering*. Vienna, Austria: R Foundation for Statistical Computing. R Package Version 2, 0-3. Available from: http://cran.r-project.org/web/packages/fpc/index.html (verified 26 June 2015).
- HERRERA-CABRERA, B. E., CASTILLO-GONZÁLEZ, F., SÁNCHEZ-GONZÁLEZ, J. J., HERNÁNDEZ-CASILLAS, J. M., ORTEGA-PACZKA, R. A. & GOODMAN, M. M. (2004). Diversidad del maíz Chalqueño. Agrociencia 38, 191–206.
- IBARRA-CARDEÑA, C., CAVAZOS, T., SALINAS, J. A., MARTÍNEZ, B., COLORADO, G., DE GRAU, P., PRIETO GONZÁLEZ, R., CONDE ÁLVAREZ, A. C., QUINTANAR, A., SANTANA SEPÚLVEDA, J. S., ROMERO CENTENO, R., MAYA MAGAÑA, M. E., ROSARIO DE LA CRUZ, J. G., AYALA ENRÍQUEZ, M. R., CARRILLO TLAZAZANATZA, H., SANTIESTEBAN, O. & BRAVO, M. E. (2013). Actualización De Escenarios De Cambio Climático Para México Como Parte De Los Productos De La Quinta Comunicación Nacional. Informe Final del Proyecto al INECC. Mexico, D.F.: INECC.
- IPCC (2007). Summary for policymakers. In Climate Change 2007: The Physical Science Basis. Contribution of Working Group I to the Fourth Assessment Report of the Intergovernmental Panel on Climate Change (Eds S. Solomon, D. Qin, M. Manning, Z. Chen, M. Marquis, K. N. Averty, M. Tignor, M. & H. L. Miller), pp. 1–18. Cambridge, UK and New York: Cambridge University Press.
- IPCC (2014). Summary for policymakers. In *Climate Change* 2014: Impacts, Adaptation and Vulnerability. Part A: Global and Sectoral Aspects. Contribution of Working Group II to the Fifth Assessment Report of the

Intergovernmental Panel on Climate Change (Eds C. B. Field, V. R. Barros, D. J. Dokken, K. J. Mach, M. D. Mastrandrea, T. E. Bilir, M. Chatterjee, K. L. Ebi, Y. O. Estrada, R. C. Genova, B. Girma, E. S. Kissel, A. N. Levy, S. MacCracken, P. R. Mastrandrea & L. L. White), pp. 1– 34. Cambridge, UK and New York: Cambridge University Press.

- KATO, T. A., MAPES, C., MERA, L. M., SERRATOS, J. A. & BYE, R. A. (2009). Origen y Diversificación del Maíz: una Revisión Analítica. México, DF: Universidad Nacional Autónoma de México, Comisión Nacional para el Conocimiento y Uso de la Biodiversidad.
- LIRA-NORIEGA, A. & MANTHEY, J. D. (2014). Relationship of genetic diversity and niche centrality: a survey and analysis. *Evolution* **68**, 1082–1093.
- MAGUIRE, B., Jr. (1973). Niche response structure and the analytical potential of its relationships to the habitat. *The American Naturalist* **107**, 213–246.
- MARTÍNEZ-MEYER, E., DÍAZ-PORRAS, D., PETERSON, A. T. & YAÑEZ-ARENAS, C. (2013). Ecological niche structure and rangewide abundance patterns of species. *Biology Letters* **9**, 20120637, http://dx.doi.org/10.1098/rsbl.2012.0637
- MERCER, K. L. & PERALES, H. R. (2010). Evolutionary response of landraces to climate change in centers of crop diversity. *Evolutionary Applications* **3**, 480–493.
- MERCER, K., MARTÍNEZ-VÁSQUEZ, Á. & PERALES, H. R. (2008). Asymmetrical local adaptation of maize landraces along an altitudinal gradient. *Evolutionary Applications* **1**, 489–500.
- NAGARAJU, S. K., GUDASALAMANI, R., BARVE, N., GHAZOUL, J., NARAYANAGOWDA, G. K. & RAMANAN, U. S. (2013). Do ecological niche model predictions reflect the adaptive landscape of species?: a test using *Myristica malabarica* Lam., an endemic tree in the Western Ghats, India. *Plos ONE* **8**, e82066, doi:82010.81371/journal.pone.0082066.
- Nuss, E. T. & TANUMIHARDJO, S. A. (2010). Maize: a paramount staple crop in the context of global nutrition. *Comprehensive Reviews in Food Science and Food Safety* **9**, 417–436.
- PETERSON, A. T., PAPEŞ, M. & SOBERÓN, J. (2008). Rethinking receiver operating characteristic analysis applications in ecological niche modeling. *Ecological Modelling* **213**, 63–72.
- PETERSON, A. T., SOBERÓN, J., PEARSON, R. G., ANDERSON, R. P., MAZRTÍNEZ-MEYER, E., NAKAMURA, M. and ARAÚJO, M. B. (2011). *Ecological Niches and Geographic Distributions*. Princeton, NJ: Princeton University Press.
- PHILLIPS, S. J., ANDERSON, R. P. & SCHAPIRE, R. E. (2006). Maximum entropy modeling of species geographic distributions. *Ecological Modelling* **190**, 231–259.
- PINHEIRO, J., BATES, D., DEBROY, S., SARKAR, D., EISPAK Authors & R Core Development Team (2011). Package 'nlme': Linear and Nonlinear Mixed Effects Models. R Package Version 3.1-97. Vienna, Austria: R Foundation for Statistical Computing. Available from: http://cran.r-project. org/web/packages/nlme/index.html (verified 26 June 2015).
- R Development Core Team (2014). *R: A Language and Environment for Statistical Computing*. Vienna, Austria: R Foundation for Statistical Computing.
- Rocandio-Rodríguez, M., Santacruz-Varela, A., Córdova-Téllez, L., López-Sánchez, H., Castillo-González, F.,

LOBATO-ORTIZ, R. & GARCÍA-ZAVALA, J. J. (2014). Detection of genetic diversity of seven maize races from the high central valleys of Mexico using microsatellites. *Maydica* **59**, 144–151.

- RUIZ CORRAL, J. A., DURÁN PUGA, N., SÁNCHEZ GONZÁLES, J. J., RON PARRA, J., GONZÁLEZ EUIARTE, D. R., HOLLAND, J. B. & MEDINA GARCÍA, G. (2008). Climatic adaptation and ecological descriptors of 42 Mexican maize races. *Crop Science* 48, 1502–1512.
- RUIZ CORRAL, J. A., SÁNCHEZ GONZÁLEZ, J. J., HERNÁNDEZ CASILLAS, J. M., WILLCOX, M. C., RAMÍREZ OJEDA, G., RAMÍREZ DÍAZ, J. L. & GONZÁLEZ EGUIARTE, D. R. (2013). Identificación de razas mexicanas de maíz adaptadas a condiciones deficientes de humedad mediante datos biogeográficos. *Revista Mexicana de Ciencias Agrícolas* 4, 829–842.
- SMITH, M. E., CASTILLO, F. G. & GÓMEZ, F. (2001). Participatory plant breeding with maize in Mexico and Honduras. *Euphytica* **122**, 551–563.
- TORRES, N. M., DE MARCO JÚNIOR, P., SANTOS, T., SILVEIRA, L., DE ALMEIDA JÁCOMO, A. T. & DINIZ-FILHO, J. A. F. (2012). Can species distribution modelling provide estimates of population densities? A case study with jaguars in the Neotropics. *Diversity and Distributions* **18**, 615–627.
- TURRENT FERNÁNDEZ, A. (2008). Producción bajo temporal. In *El Cultivo del Maíz en México. Temas Selectos* (Eds R. Rodríguez Montessoro & C. de Léon), pp. 107–113. México: Colegio de Postgraduados y Mundi Prensa México.
- URETA, C., MARTÍNEZ-MEYER, E., PERALES, H. R. & ÁLVAREZ-BUYLLA, E. (2012). Projecting the effects of climate change on the distribution of maize races and their wild relatives in Mexico. *Global Change Biology* **18**, 1073–1082.

- URETA, C., GONZÁLEZ-SALAZAR, C., GONZÁLEZ, E.J., ÁLVAREZ-BUYLLA, E. R. & MARTÍNEZ-MEYER, E. (2013). Environmental and social factors account for Mexican maize richness and distribution: a data mining approach. *Agriculture, Ecosystems & Environment* **179**, 25–34.
- VANDERWAL, J., SHOO, L., JOHNSON, C. N. & WILLIAMS, S. E. (2009). Abundance and the environmental niche: environmental suitability estimated from niche models predicts the upper limit of local abundance. *The American Naturalis* **174**, 282–291.
- VIGOUROUX, Y., GLAUBITZ, J. C., MATSUOKA, Y., GOODMAN, M. M., SÁNCHEZ GONZÁLES, J. J. & DOEBLEY, J. (2008). Population structure and genetic diversity of New World maize races assessed by DNA microsatellites. *American Journal of Botany* **95**, 1240–1253.
- YAMASAKI, M., TENAILLON, M. I., VROH BI, I., SCHROEDER, S. G., SANCHEZ-VILLEDA, H., DOEBLEY, J. F., GAUT, B. S. & MCMULLEN, M. D. (2005). A large-scale screen for artificial selection in maize identifies candidate agronomic loci for domestication and crop improvement. *The Plant Cell* **17**, 2859–2872.
- YAÑEZ-ARENAS, C., MARTÍNEZ-MEYER, E., MANDUJANO, S. & ROJAS-SOTO, O. (2012). Modelling geographic patterns of population density of the white-tailed deer in central Mexico by implementing ecological niche theory. *Oikos* **121**, 2081–2089.
- YAÑEZ-ARENAS, C., PETERSON, A. T., MOKONDOKO, P., ROJAS-SOTO, O. & MARTÍNEZ-MEYER, E. (2014). The use of ecological niche modeling to infer potential risk areas of snakebite in the Mexican state of Veracruz. *Plos ONE* **9**, e100957, doi:100910.101371/journal.pone.0100957