NOISY DATA AND DISTRIBUTION MAPS: THE EXAMPLE OF PHYLAN SEMICOSTATUS MULSANT AND REY, 1854 (COLEOPTERA, TENEBRIONIDAE) FROM SERRA DE TRAMUNTANA (MALLORCA, WESTERN MEDITERRANEAN)

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ABSTRACT

Distribution maps are key tools for environmental management and biogeographic analyses. However, success in predicting spatial distribution is limited when using noisy presence/absence data sets. Both false absences and presences can be related with local departures from equilibrium (for example, temporary extinctions or unsuccessful colonisations). Moreover, false absences can arise from limited sampling effort. Here we explore an analytical strategy to get additional information on the presence/absence pattern of one target species from the presence/absence of all other species in the community. The logic is simple: the target species should display higher probability of presence at a site if a sample from this site is faunistically very close to the samples from other sites where the species occurs. Therefore, we first model presence/absence of the target species as a function of between-sample faunistic similarity. Second, the observed data for the target species are readjusted as a function of the expected probability of presence: current presences at sites with extreme low probability of presence are interpreted as unstable presences, and are recoded as absences. Seemingly, absences at sites with high probability of presence are interpreted as false absences, and are recoded as presences. In the experimental case presented herein, the recoding procedure is based on the presence/absence of 174 species, covering a broad taxonomic scope (snails, beetles, spiders and isopods). 1 km² distribution maps of presence/absence of the endemic beetle Phylan semicostatus were modelled from these recoded data. Mapping is done using GARP based on four environmental explanatory variables. These maps seem to be more stable and less prone to fail in predicting presence than those derived directly from the observed data.

Keywords: Distribution maps, Occurrence patterns, Genetic algorithms, GARP modelling, Invertebrates, Biogeography.

RESUMEN

Datos imprecisos y mapas de distribución: el ejemplo de Phylan semicostatus Mulsant y Rey, 1854 (Coleoptera, Tenebrionidae) en la Serra de Tramuntana (Mallorca, Mediterráneo occidental)

Los mapas de distribución son herramientas clave para la gestión medioambiental y los análisis biogeográficos. Pero el éxito en las predicciones de distribución espacial es limitado cuando se dispone de datos imprecisos de la presencia/ausencia. Tanto falsas ausencias como falsas presencias pueden estar relacionadas con desviaciones locales del equilibrio (por ejemplo, extinciones temporales o colonizaciones no exitosas). Además, las falsas ausencias pueden surgir de un esfuerzo de muestreo limitado. Aquí se explora una estrategia analítica para obtener información adicional sobre el patrón de presencia/ausencia de una especie diana a partir de la presencia/ausencia de otras especies en la comunidad. La logica es simple: la especie diana debería tener una mayor probabilidad de presencia en un punto si
Introduction

There is a general agreement on the low budget available for faunistic and floristic surveys; unfortunately, the near future will not bring notable increases in most of the European countries. At the same time, the current level of knowledge on the spatial distribution of most species remains still inadequate for to be considered useful tools for environmental management. The case of the Balearic Islands is an example: no reliable data exists even for the mere number of invertebrate species. This stage of knowledge is evident, for example, in a contribution by Dr. Martín-Piera, who increased by eight species the checklist of the Scarabaeidae (Coleoptera) from the Balearic Islands after only two short field surveys (Martín-Piera & Lobo, 1992).

This justifies an increasing effort toward predicting species distributions (or community descriptors as diversity) from a small sample of prospected sites (Austin, 2002). The inferential logic behind this is quite simple: a species is predicted to occur in a non-surveyed site if it displays similar environmental features to other sites where the species currently occurs. The biological basis for these inferences is founded in a number of assumptions, two of which are especially relevant here. The first one is that species are in equilibrium (or at least, some kind of quasi-equilibrium) with the environment (Austin, 2002). It is thus assumed that abundance (or probability of presence) of each species is potentially deducible from environmental data, in spite of the debate on the shape of species-response curves or on the rules controlling species turnover and species packing. However, success in prediction is limited by the effects of history or disturbance (i.e., non-equilibrium). For example, in a presence/absence survey in the context of metapopulation dynamics, it is easily imaginable that one species can be detected in a site that does not display suitable (or even unsuitable) environmental conditions. This local population is prone to becoming extinct, and its presence should be considered as temporally unstable. Another source of uncertainty is that a species currently present in a site has not been effectively observed (Zaniewski et al., 2002; Dennis & Shreeve, 2003). Such false positives and negatives (Fielding & Bell, 2003) can be considered as noise over the signal (species-environment relationships), and certainly they will limit the performance of prediction.

The second assumption involves the continuum concept that, widely interpreted, points to species composition changing along a continuous environmental gradient (Fig. 1, inspired, for example, in Putman & Wratten, 1984 and ter Braak & Smilauer, 1998). Figure 1 involves the additional assumption that the species response curves to environment are unimodal and symmetric. However, independently from the current shape of the species-response curve, it is clear that some information on the presence-absence of an individual species should be potentially deducible from the presence or the absence of all other species in the community. Here we explore an analytical strategy to obtain additional information on the presence/absence of one target species from the presence/absence of all other species in the community. We first model presence/absence of the target species as a function of between-sample faunistic similarity. Second, the observed data-set for the target species is readjusted as a function of the expected probability of presence: current presences at sites with extreme low probability of presence are interpreted as unstable presences, and are recoded as absences. Similarly, absences at sites with high probability of presence are interpreted as false absences, and are recoded as presences.
We first analyse a simulated system looking for the accurate description of the procedure using a data set corresponding to a community with a simplified and known structure. Subsequently, a macro-invertebrate community is studied. The adequacy of arthropods and other macroinvertebrate relies on their large number and their high species turnover along environmental gradients (Lawton et al., 1998), while most vertebrates are insensitive to fine-scale habitat heterogeneity (Mattoni et al., 2000). The case study is based on the presence/absence of 174 species (covering snails, beetles, spiders and isopods) on 48 sites of 1 km² each along a mountain range. The overall data set are used to adjust the observed presence-absence data-set of a target species (the endemic beetle Phylan semicostatus). The spatial distribution of this species along the whole mountain range is then modelled from the adjusted occurrence records and four environmental variables. The modelling strategy adopted is based in an heuristic search using a genetic algorithm (the Genetic Algorithm for Rule-Set Prediction, GARP; Payne & Stockwell, 2001).

**Methods**

**SIMULATED GRADIENT**

In order to analyse the potential capabilities of the proposed analytical strategy, we designed a system composed of 50 sites and 254 species. For simplicity, species responses to the environment are all considered unimodal and symmetric (i.e., Gaussian, ter Braak & Smilauer, 1998; Oksanen et al., 2001), and the environmental gradient highly correlated with a single variable. Gaussian response curves are defined by three parameters. Namely optimum, tolerance, and abundance at the optimum. Species optima are randomly located along the gradient. Species tolerance is randomly allowed to move between zero and the half-width of the gradient. Finally, abundance at the optimum is randomly shifted between zero and the half-width of the gradient. Stochastic variability is simulated by simple addition (or subtraction) of a random proportion of the maximum species abundance. Figure 1 shows the species response of a random subsample (10) of the 254 species considered.

The averaged number of individuals per site is $2.1 \times 10^6$. A small number of individuals (1/50000) are sampled with replacement from each site, from which a presence/absence data set is built. The averaged number of species and individuals per sample are respectively 16.1 and 42.8.

Between-samples differences are analysed using correspondence analyses (CA, ter Braak & Smilauer, 1998) of the abundance and presence-absence data. The scores of the first CA axis are estimates of between-sample faunistic differences and were plotted against the (single) environmental variable for comparing the pattern extracted by abundance data against that extracted by presence-absence data.

One species was chosen at random, and its presence/absence pattern is modelled with the scores on the first CA axis. Instead of the usual Gaussian model (Oksanen et al., 2001) we used the full HOF model (Huisman et al., 1993, Oksanen & Minchin, 2002). The main advantage of this model is that skewed (asymmetric) unimodal species response can be managed also. Two *a priori* selected thresholds allowed us to detect false absences and unstable presences from the data. A new data set (i.e., the adjusted data) is

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Fig. 1.— Theoretical species turnover along an ecological gradient. Species response curves are assumed to be unimodal, symmetric and defined by three parameters (optimum, tolerance and abundance at the optimum).

Fig. 1.— Reemplazamiento teórico de especies a lo largo de un gradiente ecológico. Se asume que la curva respuesta de las especies es unimodal, simétrica y definida por tres parámetros (óptimo, tolerancia y abundancia en el óptimo).
then built after recoding the putative false positives and negatives. Finally, the modelled species presence using the recoded data-set is compared with the original data set. This comparison is made by plotting the proportion of correctly predicted occurrences against an increasing threshold level (Fielding & Bell, 2003).

**THE CASE OF *PHYLAN SEMICOSTATUS***

The study area (Serra de Tramuntana) is a mountain range located to the North-West of Mallorca, rising from the seashore to 1445 m a.s.l. Forty-eight sampling sites have been selected along its main axis (100 km long; Fig. 2); Their positions GPS-determined. Five pitfall traps (10 cm wide; 2 m left between traps; detergent and salt used as preservative) were set at each site. The traps remained in the field for two moths (June to July). The sampling schedule has been complemented with one hour of direct search (e.g., under stones) at each site on a 1000 m² plot around the pitfall traps. We analyse presence-absence data because between-species differences in detectability and catchability were noticeable. Data from the five pitfalls and from direct search are pooled as a single sample per site, and it is assumed to represent the fauna of the corresponding 1 km² cell.

We focus on four groups of invertebrates, namely Coleoptera, Arachnida, Isopoda and Gastropoda. As regard Gastropoda, a species is considered to be present also when empty shells only are found. Some of the species not readily determinable are identified as morphospecies (Oliver & Beattie, 1996).

Presence/absence data are analysed in a similar way to the analysis of the simulated data set. The major differences are: (1) the ecological gradient is considered to be bivariate instead of univariate (presence/absence of *Phylan semicostatus* was modelled using the scores on the first two CA axes), and (2) the spatial distribution of *Phylan semicostatus* is deducted using a more formal modelling strategy (GARP). The occurrence pattern along 584 cells of 1 km² (Fig. 2) is predicted from the adjusted occurrence pattern and four environmental variables: altitude, aspect, distance to the nearest house and NVDI (non-linear combination of reflectance at specific lambdas extracted from a satellite image). The methods for estimating the environmental values corresponding to each 1 km² cell are fully detailed in (Palmer et al., 2002).

The Genetic Algorithm for Rule-Set Prediction (GARP) is an expert system machine-learning approach to predictive modelling (Stockwell & Peters,
Genetic algorithms are inspired in evolutionary models. They present an heuristic solution after scanning broadly across the solution space (i.e., all possible solutions), and refining solutions that show high values for the optimisation criterion. GARP has not been extensively used yet, but has proved to be a useful approach (see Anderson et al., 2003 for a recent review). The ordinary strategy involves running the program a number of times to obtain several tentative models. Model selection have been carried out using the criteria suggested by Anderson et al., 2003). Briefly, the models selected display both low intrinsic omission error and intrinsic commission error near the average (see Fielding & Bell, 2003 and Anderson et al., 2003 for more details on error theory and model evaluation for occurrence data).

GARP (Stockwell & Peters, 1999) is downloadable at http://beta.lifemapper.org/desktopgarp. Correspondence analyses have been carried out using CANOCO 4 (ter Braak & Smilauer, 1998), and HOF models were fitted using R (http://cran.r-project.org; nlm functions were modified from those supplied by http://cc.oulu.fi/~jarioksa/ and Oksanen & Minchin, 2002).

Results

SIMULATED GRADIENT

As expected, the scores on the first axis from a Correspondence Analysis (CA) of the original data are clearly correlated with the environmental variable (Fig. 3). Moreover, and more interestingly, the corresponding scores derived from a presence/absence data set that simulate a low sampling effort (1 of each 50000 individuals) show a very close trend. The importance of this result relies on the fact that CA scores on the first few axes obtained using real-world presence/absence data probably reflect the main betwe-
en-sample similarity trends, even in cases of noisy data sets.

The results of modelling presence/absence data of a target species using the scores on the first CA axis are shown in Figure 4. The observed presences/absences are derived from the noise-added data. The crux of the strategy is the following recording procedure: The species was not detected at the sites labelled as open points because of the low relative abundance of the species at these sites. However, some of these sites (#45, 47, 51, 58, 59 and 61) showed CA scores that are very close to all the other sites where the species was present because they share a similar faunistic composition (except for the case of the target species). After modelling probability of presence as a function of the scores on the first two CA axes, the probability values of presence for the sites listed above is always higher than a prefixed threshold (here, 0.5); consequently, the absences of the target species on the sites are considered to be false absences and are recoded as presences.

Similarly, the species is detected at the sites labelled as closed points, but two of them (#34 and 63) display a low probability of presence (below 0.3). Consequently, these two occurrences are considered to be unstable and are recoded as absences.

The latter threshold is selected to include 90% of the observed occurrences (i.e., sites 34 and 63 are those with more deviant CCA scores).

The consequences of the recoding procedure on the prediction of presence/absence from the (single) environmental variable are shown in Figure 5. The overall performance (total misclassifications) of the two data sets (observed versus recoded) in predicting the occurrence profile of the original (noise-free, Gaussian) data is similar. However, occurrences are better predicted by the recoded data, which is exactly the final objective of the recoding procedure.

THE CASE OF *PHYLAN SEMICOSTATUS*

The target species occurs along the length of the Serra de Tramuntana mountain range, but occurrences seem to be more frequent at the central area (Fig. 6). This pattern coincides with field experience pointing to this species being characteristic of places at high-moderate altitude (in relation to the observed maximum altitude, 1445 m). This endemic beetle is easily found at sites with low vegetation cover and mountain-type shrubs. However, it is noteworthy that the species is also frequent in a number of small islets around Mallorca, demonstrating that altitude itself has no biological relevance (i.e., indirect effects; Palmer, 1994; Palmer, 1997). The current research does not cover littoral habitats.

Presence/absence data of the target species is modelled using the scores on the two first axes of the ordination.
a CA (174 species and 48 sites) as predictors. The fitted model is a full HOF model (four parameters for each of the two predictors involved; more details are provided by Oksanen & Minchin, 2002). The probability of presence predicted by the model is shown in Figure 7. The threshold for detecting a false absence is 0.5 (i.e., all sites with expected P-values higher than 0.5 are assumed to be occupied by the species), while unstable presence threshold is defined as the probability level that includes 90% of the occurrences (i.e., the occurrences that display the more extreme values on CA1 and CA2 axes are considered to be unstable; in the case of Phylan semicostatus this value was 0.2).

Twenty optimal (Anderson et al., 2003) GARP-predicted spatial distribution maps are obtained with both recoded and observed data. The median values for each of the 584 pixels are compared in Fig. 8. The maps corresponding to the observed and recoded data consistently show similar intrinsic omission rates (close to 0%). However, the between-models variability is higher among the maps built with the observed data (Fig. 9). The high rate of pixels for which the model does not offer clear-cut prediction is also noteworthy (Fig. 8).

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Fig. 7.— Standard plot of the sample scores on the first two axes of a Correspondence Analysis (CA). Solid and open points denote respectively samples with and without Phylan semicostatus. Probability of presence of Phylan semicostatus (HOF model) is indicated by a grey scale (darkness indicate higher probability values). The wider isoline indicates the threshold for considering presence as unstable. The dashed isoline indicates the threshold for considering absence as false absence.

Fig. 7.— Representación estándar de la posición de las muestras con respecto a los dos primeros ejes de un análisis canónico de correspondencias. Los puntos negros y blancos indican respectivamente las muestras con y sin Phylan semicostatus. La probabilidad de presencia de Phylan semicostatus es indicada mediante una escala de grises (tanto mas oscura al aumentar la probabilidad de presencia). La línea ancha continua indica el umbral para considerar una presencia como inestable. La línea ancha discontinua indica el umbral para considerar una ausencia como falsa ausencia.
Fig. 8.— Maps of the modelled spatial distribution of *Phylan semicostatus* in the study area. These maps show for each pixel (1 km$^2$) the median value of twenty optimal maps. The map corresponding to the recoded data seems to be more stable, and show fewer points with undefined prediction.

Fig. 8.— Mapas de la distribución espacial modelada para *Phylan semicostatus* en el área de estudio. Estos mapas muestran para cada pixel (1 km$^2$) el valor de la mediana de 20 mapas originales (i.e., modelados independientes). El mapa correspondiente a los valores recodificados parece ser más estable, y muestra menos puntos sin una predicción definida.

Fig. 9.— Maps of the modelled spatial distribution of *Phylan semicostatus* in the study area. These maps show for each pixel (1 km$^2$) the number of models predicting presence. Between-model differences in the range of potential distribution of the target species are higher in the case of the observed data.

Fig. 9.— Mapas de la distribución espacial modelada para *Phylan semicostatus* en el área de estudio. Estos mapas muestran para cada pixel (1 km$^2$) el numero presencias en 20 modelados independientes. Para la especie diana, las diferencias entre modelados para las predicciones son mas grandes en el caso de los datos observados que en el caso de los datos recodificados.

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Discussion

The results produced by the analysis using the simulated community data-set show that faunistic differences between sites are well described with presence/absence data even in cases of small sampling effort and noisy data. Canonical correspondence analyses of the original abundance data and presence/absence data would render similar patterns if the number of species considered is high enough. Simulated data illustrate also the type of noise that potentially affects the species-specific response curves to ecological gradients, and reveals ways for filtering this noise. Recoded data are less prone to fail in predicting presence (Fig. 5). The counter-part of recoding is that these data show increased risk of failing to predict absence. However, the latter is an advantage when sampling is not very exhaustive, because the high prevalence of false absences.

The results derived from the experimental data suggest that the solutions obtained using the recoded data are more stable in the sense that they are more similar to each other. However, two potential limitations should be noticed. First, the method is not applicable to species from environments very different to a single and general environmental gradient. The reason is that the first axis from correspondence analyses (or any other multivariate approach) account for the faunistic differences between these marginal sites and the rest of sites, instead of being correlated with the target environmental gradient. An extreme example of this would be to include some samples from ponds or other aquatic environments: these samples are easily detectable by plotting the CA scores, and it has no sense to include them in any further analysis that focuses on non-aquatic fauna. Another limitation of the method is it implicitly assumes similar sampling effort. This is the case of the data analysed here, but most of the data sources used for mapping do not meet this assumption (e.g., museum data) and provides biased images of species distribution (Dennis, 1999; Dennis, 2000).

In summary, the proposed analytical strategy (i.e., using the information from other species in order to improve the observed pattern of occurrence of a target species) seems to work reasonably well. This improvement, far from being considered as data manipulation, should be better defined as noise filtering. Obviously, further research should be done regarding a number of points. For example, enlarging the number of target species (and covering a wide range of prevalence), and improving the selection procedure of the thresholds to detect false positives and negatives, which are reasonably but subjectively defined. However, this or similar analytical procedures opens the possibility of making full use of all the information obtained with extensive invertebrate trapping surveys. The unbiased image of species distribution can be the input for a variety of additional analyses. One interesting approach is to integrate presence-absence data with knowledge on interspecific associations and multivariate ordination, using distance-dependent variance-covariance matrices (Wagner, 2003). The same idea is extrapolable from spatially to temporally autocorrelated data, opening new ways to estimate the relative importance of historical versus environmental forces in determining species composition.

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