



Using species distribution models in paleobiogeography: A matter of data, predictors and concepts

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ABSTRACT

The increasing interest in the effects of climate changes on species distributions has been followed by the development of Species Distribution Models (SDMs). Although these techniques are starting to be used to study the location and dynamics of past species distributions, a sound theoretical framework for their use in paleoecology is still lacking. In this paper we are reviewing the main challenges for constructing Paleo-Species Distribution Models to describe and project the past distribution of species, namely data limitations, selection of predictors and choice of a biologically-relevant modeling procedure. We also review and discuss the current state-of-the-art in Paleo-SDMs, providing a series of recommendations for their use, and proposing future research lines to improve the use of these techniques in paleobiogeography.

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1. Introduction

Research in paleobiogeography is currently moving from narrative biogeographic descriptions and interpretations of fossil data toward describing and estimating past species range shifts and extinction events through quantitative statistics and modeling techniques (Rodríguez-Sánchez and Arroyo, 2008). GIS data on past and current species distributions, climate, topography or geology and new analytical tools are now used to simulate the geographic consequences of climatic changes through time. This has allowed the relating of the environmental changes caused by Pleistocene climatic oscillations with spatial and temporal turnovers in species composition, the location of glacial refugia, the fragmentation of distributional ranges, migration and extinction events, and even speciation processes (Svenning et al., 2011).

Species distribution models (SDMs) are a heterogeneous group of techniques used to model species' geographic ranges by relating their known occurrences with the environmental – typically climatic – conditions in these locations (Guisan and Zimmermann, 2000; Guisan and Thuiller, 2005). The relationships identified with these techniques are thought to provide a description of the adequacy of each combination of environmental predictors for the maintenance of populations of the studied species, thus identifying the areas where the net reproductive rate of its populations would be positive (Soberón, 2007, 2010; Soberón and Nakamura, 2009). These descriptions can be refined by including species interactions, non-climatic predictors or spatial autocorrelation terms (Araújo and Luoto, 2007; Dormann et al., 2007; De Marco et al., 2008; Guisan and Rahbek, 2011).

SDMs are among the most widely used methods in biogeography and macroecology; although they are typically employed to map species ranges, they are also utilized to address many evolutionary and ecological questions (Lobo et al., 2010). In paleontology, SDMs are now applied to study a number of topics (see Svenning et al., 2011) including: the effects of climate changes on the temporal dynamics of species distributions and human cultures (Banks et al., 2006, 2008a; Maguire and Stigall, 2009; McDonald and Bryson, 2010; Polly and Eronen, 2011; Walls and Stigall, 2011); the nature and causes of extinction events (Nogues-Bravo et al., 2008; Varela et al., 2010); the location of glacial refugia (Jakob et al., 2007; Carnaval and Moritz, 2008; Schmickl et al., 2010); the retention of niche-related ecological traits over time (Martínez-Meyer and Peterson, 2006; McDonald and Bryson, 2010); or the fingerprints of past climatic changes on current genetic structure (Alexandrino et al., 2007). Herein we will refer to all these paleontological applications of SDMs as Paleo-Species Distribution Models, or PSDM for short. The generalization of PSDMs as research tools could be a major step forward for paleoecology and paleobiogeography (Nogues-Bravo, 2009). PSDM applications may enhance our understanding of the determinants of species distributions and their evolution, helping to address some fundamental questions such as how tight are species' responses to environmental changes?, which are the biological consequences of an extreme climatic event?, or whether glacial refugia are a key factor for explaining current biogeographic patterns.

To take full profit of the potential of PSDMs it is necessary to establish robust and scientifically-based theoretical and methodological frameworks. Although their principal strength is that they allow studying past changes in species distributions quantitatively, the adequacy of SDMs to attain certain research questions and the accuracy and reliability of their results are currently a matter of discussion (e.g. Jiménez-Valverde et al., 2008; Colwell and Rangel, 2009; Godsoe, 2010; Sinclair et al., 2010). Here we examine the main methodological and conceptual issues in the application of PSDMs. More specifically, we discuss: (i) the relevance of the biases and scarceness of the available species distribution data from fossil records; (ii) the influence of the environmental predictors used; and

(iii) which model techniques are the most appropriate to derive geographical representations of species distributions for different time scenarios, when the only reliable information are data on species' occurrences. In addition, we review the different methodologies used to project data on the current responses of species to climate into past scenarios (i.e., hindcasting; see Nogues-Bravo, 2009). Based on the above, we propose future research guidelines, suggesting key gaps of knowledge that need to be investigated before the use of PSDMs is widespread in paleontology, with the general purpose of stimulating debate and discussion on this novel and potentially fruitful field of research.

2. Methodological and conceptual issues for Paleo-Species Distribution Modeling

2.1. The paleontological data

The ideal manner to describe the past distribution of a species in a context of climate change would be to use a technique capable of identifying the causal relationships that limit species' geographic ranges. This would allow creating accurate projections of the distribution in any temporal scenario. A prerequisite for this is that the dependent variable modeled by PSDMs is either a direct measure or a proxy of the fitness of the species in any locality – in particular with regard to changes in climate at range limits – such as reproductive rate, mortality rate, abundance or physiological basal cost, among others (Kearney and Porter, 2009). However, rather than any of these measures, quite often the only dependent variable available for PSDMs – and SDMs in general – is georeferenced data on species' presences. It is well known that the information about current species' occurrences is usually not enough to represent the full spectrum of environmental conditions at which a species may have a net positive demographic rate (Hortal et al., 2008). The geographical and environmental coverage of such information is typically limited, a lack of completeness that results in a number of biases and flaws (Rocchini et al., 2011). Even if the data on species' occurrences is not biased or incomplete, it may still not be able to reflect their fundamental niche simply because some environmentally suitable localities may not have been colonized (Jiménez-Valverde et al., 2011).

The geographical information provided by fossil data is subject to similar limitations (Fernández-Jalvo, 1996; Hadly, 1999; Fernández-Jalvo et al., 2011), which perhaps are even more severe due to the spatio-temporal character of these data. Contrary to recent data, occurrences extracted from fossil data cannot be directly interpreted as being part of the realized distribution of the species because they originate from a diverse array of processes – both climatic and non climatic – that operate at different taxonomic, spatial and temporal scales (Fernández-Jalvo, 1996; Barnosky et al., 2005; Chew and Oheim, 2009; Fernández-Jalvo et al., 2011). This implies that a direct analysis based on the recorded location of the occurrences could lead to misleading conclusions about the species–climate relationships (Hadly, 1999). Here we summarize the different sources of bias that should be taken into consideration when building PSDMs from fossil data.

2.1.1. Spatial, temporal and taphonomic bias

Fossils occur in sedimentary deposits, so the geographic distribution of fossil records is biased by the distribution of sedimentary layers. In addition, the absence of fossils does not necessarily mean that the species was absent when these layers were forming, because the lack of fossil remains depends also on the high number of physical, chemical and biological factors that are behind the process of a carcass being converted into a fossil. This fundamental characteristic of fossil data adds up to other factors that make difficult the inference of absences from recent distributional data, such as lack of survey effort

or species' detectability (Mackenzie and Royle, 2005). These latter factors are discussed in detail elsewhere (e.g., Lobo et al., 2007, 2010; Hortal et al., 2008; Rocchini et al., 2011) so we will only discuss the taphonomic processes that make it difficult to separate apparent from real absences in the fossil record.

The intrinsic ecological and biological characteristics of each species can either prevent some remains from being deposited in a given sedimentary area, or increase the prevalence of others (Reed, 2005; Andrews, 2006; Davis and Pyenson, 2007). The fossilization of the organic material is determined by its composition and size in relation to the local sedimentary conditions (Arroyo-Cabrales et al., 2007), so fossil samples are usually biased toward species with large body sizes (Lucas et al., 1997; Muñoz-Durán and Van Valkenburgh, 2006). It follows that many species, and in particular the smaller ones, could be absent in a fossil deposit despite being present during the period when the deposit was formed. This characteristic of fossil data increases spatial bias, so the biotas of certain biomes are either over- or underrepresented in the available occurrence data (Nieto et al., 2003). For example, since many Spanish regions lack sedimentary basins from the Lower Pleistocene our knowledge on the Iberian faunas of this period is spatially biased, and so is the coverage of the environmental conditions that could be used to estimate the response of any species to climate during that period of time (Fig. 1). It follows that some knowledge about the taphonomic processes that may have

affected the prevalence of the studied species in the deposits is important to interpret datasets on fossil occurrences.

In addition to spatial bias, fossil data are also subject to temporal bias. Certain periods have rendered larger sedimentary areas than others, and therefore the extent of area that could host fossil records varies through time. Due to this, the variations in the abundance of fossil remains through time cannot be directly used as an indicator of changes in population size. Rather, to infer population size or compare species' abundances between periods it is necessary to consider the number of fossil remains of the species, the number of deposits from the same period and region, and the total survey effort (i.e. organized excavation campaigns or irregular prospectings). For example, sedimentary deposits are more abundant in Spain for the Lower Pleistocene than for the Upper or Middle Pleistocene (Fig. 2), so the fossil record of the Lower Pleistocene contains comparatively more rare species—such as the exceptionally rare hominid remains found in Atapuerca (Bermúdez de Castro et al., 2008). The methodologies used to date the fossil remains also generate temporal bias; the use of methods capable of dating fossil remains older than 50,000 Years Before Present (YBP) – the lower confidence limit of ¹⁴C (Magee et al., 2009) – is not widespread. Thus, while Late Pleistocene fossil records are typically dated accurately, Middle and Lower Pleistocene fossils often remain undated, which could give the false impression of higher species' abundances or occurrences in the Late Pleistocene.

To summarize, the fossil record is not evenly distributed, neither in time nor in space (Jass and George, 2010); therefore, occurrence data gathered from fossil records cannot be used as a direct indicator of the distribution and abundance of a species (Chew and Oheim, 2009). This problem can be overcome by weighting occurrence data to balance the effects of the biases discussed above (Signor, 1982; Smith et al., 1988; Crampton et al., 2003). However, doing this requires not only estimating the common biases on distributional data that are similar to those affecting data on recent species, but also the geographic and temporal bias in the location of the sedimentary basins and the eventual bias derived from dating deficiencies.

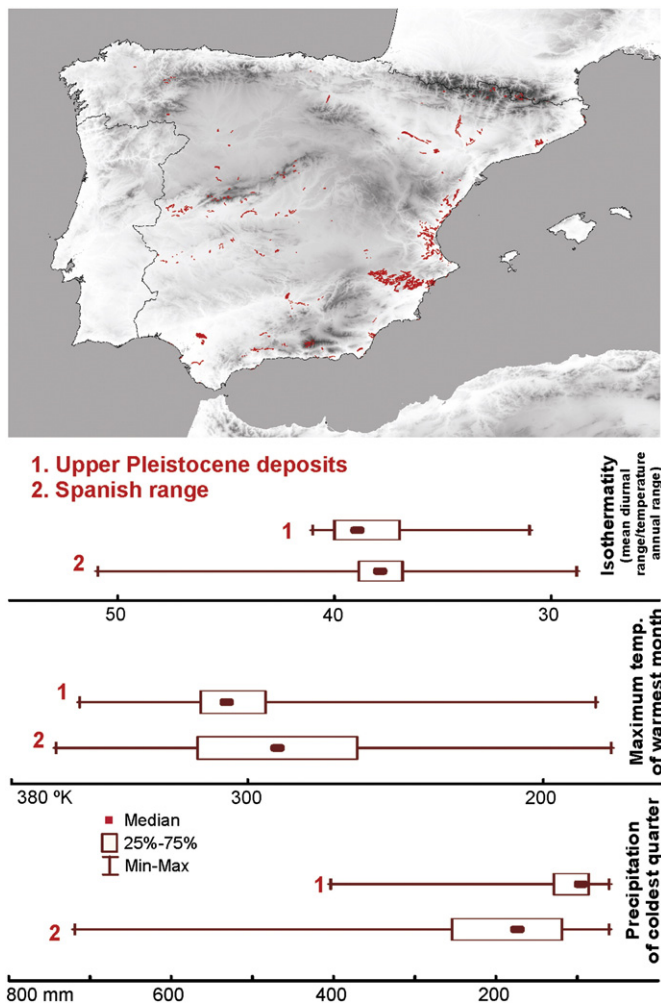


Fig. 1. Geographic distribution of Upper Pleistocene deposits in Spain. Note that the locations of these deposits do not cover the entire climatic conditions of the Iberian Peninsula, so the Iberian ranges of both temperature and precipitation are underestimated using the sample provided by the available fossil sites. Climatic variables extracted from Worldclim (<http://www.worldclim.org/>).

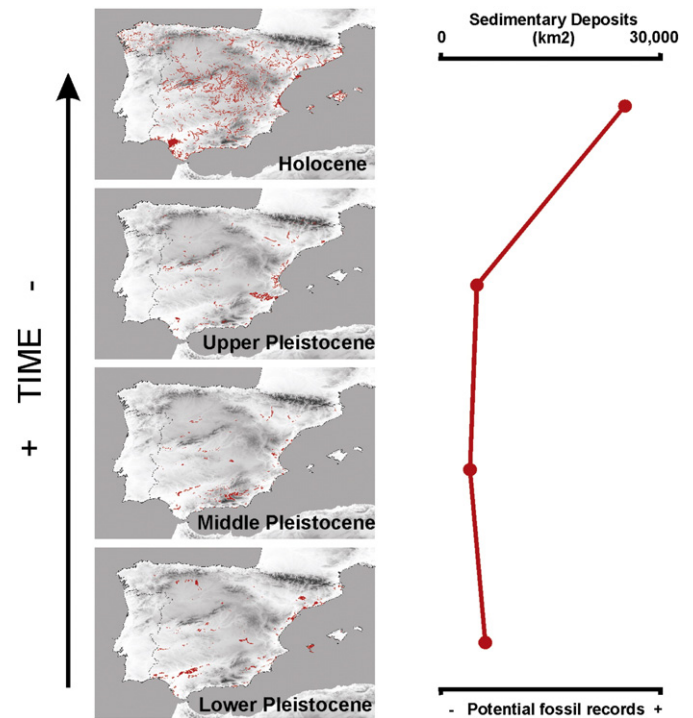


Fig. 2. Quaternary deposits in the Iberian Peninsula show different geographic biases across time. In this region, Holocene deposits are more abundant than Pleistocene deposits, and specifically, Lower Pleistocene deposits are more abundant than Middle or Upper Pleistocene deposits.

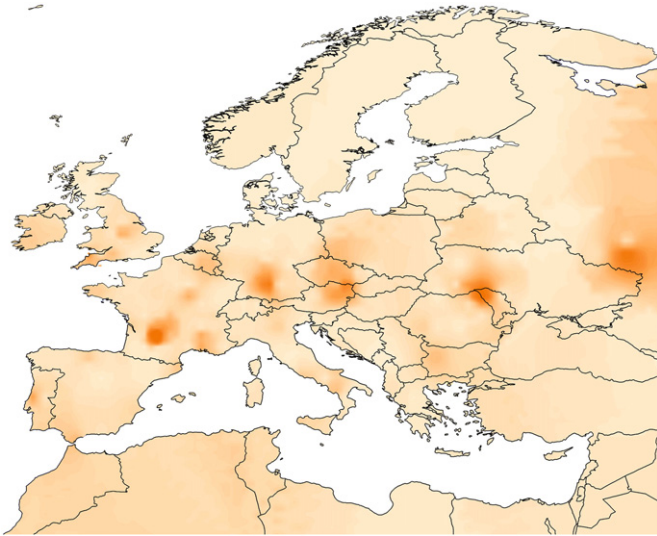


Fig. 3. Abundance of European mammal fossil sites during the OIS 3 (according to the Stage 3 Project database; <http://www.esc.cam.ac.uk/research/research-groups/oistage3/>).

2.1.2. Collectors' bias

Certain idiosyncrasies of the paleontologists themselves may also bias our knowledge on past species distributions. On the one hand, the between-country differences in paleontological tradition result in spatially uneven survey efforts. The distribution of mammal fossil sites for the European OIS3 may illustrate this point; almost one third of the fossil records for this period are located in France, the country with the longest paleontological tradition (Fig. 3). Furthermore, some temporal or geographic scenarios receive more attention – and thus survey effort – due to their interest in solving certain questions. For example, the extinction of the Neanderthals in Europe – at around 30,000 YBP – is explicitly addressed by the Stage 3 Project (<http://www.esc.cam.ac.uk/research/research-groups/oistage3/>). This project studies in detail the environmental and biotic conditions of this particular period, thus rendering data in much larger quantity and of better quality than for any other moment of the European Pleistocene. Similarly, the effort is unevenly distributed between different groups; a search in the ISI Web of knowledge (October 2010) rendered 784 papers using “Pleistocene” and “hominid” as topic keywords, 259 using “Pleistocene” and “large mammal”, and 147 using “Pleistocene” and “small mammal”. Nonetheless, the excavation methodologies can bias the fossil record, typically toward large-sized mammals. Rodents and insectivores are sometimes not recorded on a fossil site simply because the excavation method was not designed to collect small-sized species.

The taxonomic identification of the fossil remains is also highly dependent on the paleontologists' interpretation (Lister et al., 2005; Mounier et al., 2009), a taxonomic unevenness that may condition the results and conclusions obtained from the application of SDMs (see Lozier et al., 2009). Species are not homogenous entities (Quintyn, 2009), turning the decision of when to discriminate two different species into an arbitrary exercise, especially because the genotypic and phenotypic variability of the populations of related organisms are spatially and temporally structured (Holliday, 2003). This already makes difficult the discrimination of individuals of recent organisms into species (Tobias et al., 2010), but when it comes to fossil species the temporal dimension adds an extra level of complexity (Escude et al., 2008; Ogino, 2009). Temporal variations in morphology are interpreted by paleontologists like recent taxonomists interpret geographic variations in current species; depending on the criteria used individuals with similar levels of morphological differentiation

can be classified as morphotypes from the same species, different subspecies, or even distinct species or chronospecies. The spatial and temporal amplitude used to aggregate the fossil remains into discrete species plays a central role in this process, so any use of the fossil record necessarily implies taking a position on this controversial question (Hlusko, 2004). If the assumption is that all past occurrences of the studied taxon represent different individuals of the same taxonomic entity – even when separated by long periods such as 1 million yr – all fossil and current records can be used as training dataset for SDM (Varela et al., 2009). If, on the contrary, the model is intended to describe a given period and/or region of interest, the temporal or spatial dimension of analysis – or both – can be kept fixed (e.g., Alba-Sánchez et al., 2010), assuming that all species' occurrences within such spatio-temporal window behave as a separate and coherent unit in relation to their climatic requirements, and by implication as a “distinct” species. Both approaches can be correct depending on the context, but the decision to include or exclude species' occurrences from each particular analysis needs to be based in the particular question addressed, and the reasons for such decision should be explained and discussed in detail.

The combination of all these different collectors' biases leaves a footprint in the fossil record (Fig. 4). As a consequence, the relationships between the occurrence data and the predictors may offer a flawed picture of species' responses to the environment. Thus, while evaluating the quality of data on fossil occurrences, the main objective should be to determine whether these data provide an adequate sample of the conditions present during each period. Therefore, our suggestion here is to subsample the raw data on fossil occurrences to compensate for the different biases in the data. If the extent and distribution of the biases in the data are ignored or simply not taken into account, the error and/or uncertainty in the estimate of the species' climatic requirements will remain unaccounted for and/or unknown, and SDM results will fail to predict the dynamics of the species' geographic ranges through time (Elith et al., 2010). It is however possible to take profit of fossil data from an understanding of its limitations. Fossil remains may be considered extremely long temporal surveys of the geographic distribution of the species in a global scenario of changing climate. Thus, the fossil record provides the most complete knowledge on the responses to environment of many species, certainly better than the one obtained using just their current distributions (Varela et al., 2010). This makes fossil data quite helpful to understand the processes behind current biogeographical patterns; analyzing the whole “movie” of the dynamics of the realized distribution of the species through time provides much more information than what can be obtained from the mere “picture” of its current distribution (Varela et al., 2009).

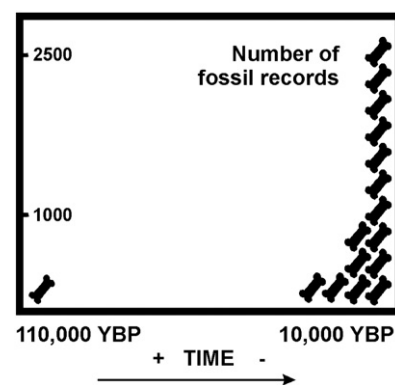


Fig. 4. Temporal distribution of the Late Pleistocene fossil records of North America (extracted from FAUNMAP database; <http://www.museum.state.il.us/research/faunmap/>). The temporal distribution of data on fossil remains is the result of the temporal changes in sampling effort bias, plus the difficulty for dating fossil records older than 50,000 yr (the lower limit of C¹⁴-based dating).

2.2. The predictors

The accuracy of SDM predictions also relies on the explanatory variables used to build the models (Stockwell and Peterson, 2002). Thus, a key step while building a PSDM is how to choose these predictors. The ideal variables should have the adequate spatial and temporal resolution to describe the response of the studied species, while at the same time be descriptors or proxies of the causal factors that truly determine the occurrence of the species (Austin, 2002) and/or have the greatest impact on the fitness of its populations (Soberón, 2010).

Unfortunately, the predictors available for use with paleontological data are in general coarse-grained outputs of the climatic scenarios provided by different General Circulation Models (GCMs; see Braconnot et al., 2007 and <http://pmip3.lscce.ipsl.fr/> for detailed information). GCMs simulate the general circulation of the atmosphere and oceans of the Earth; together with sea-ice and land-surface components, they can be used to generate climatic forecasts for a particular moment in the future, or hindcasts for a moment in the past (Foley et al., 1998). To do this, they assume certain initial conditions (i.e. CO₂ atmospheric concentration, ice-covered surface, etc.) and are run a large number of iterations, until they stabilize (Thompson and Pollard, 1995a,b). Due to the difficulty of modeling complex climatic processes at small scales, the spatial resolution of GCM-derived maps is often coarse, typically of about 2°. Only regional models that use GCMs as the boundary conditions to run more detailed climatic simulations allow finer-resolution outputs of up to 60 km in width (Barron and Pollard, 2002).

Coarse-grain variables can lead to imprecise results (Seo et al., 2009). Due to this, a diverse array of procedures is used to downscale the climatic layers provided by GCM simulations (Banks et al., 2008b; Peterson and Nyari, 2008; Galbreath et al., 2009). Unfortunately, there is not a general agreement about the best way of downscaling these simulations. Different algorithms are used to define the relationship between large- and local-scale climate variables, based on either data about current climate or proxies of past climate such as vegetation from pollen cores (Hugall et al., 2002; Bigg et al., 2008; Buckley et al., 2009). Downscaling these regional predictions presents two main challenges: avoiding the propagation of the uncertainties of the GCMs themselves into the downscaled projections (Katz, 2002), and taking the temporal variations of the local climate into consideration. Both sources of error can be accounted for using an adequate calibration of the local data. However, it is difficult to determine fine-grained climate variations from pollen data – because pollen deposits represent the flora of relatively large regions –, and methods based on current data simply extrapolate current local climatic anomalies into past scenarios. This strategy may be flawed due to the effect of the changes in air-mass circulation on local climate. The extent of the Polar icecaps influences the global atmospheric circulation—global circulation at LGM was radically different from the present due to the effect of the large extensions of ice sheets (Bowen, 1995). These temporal changes in the extension of permanent ice are likely to change the relationship between global and local circulations, preventing the extrapolation of past relationships from current ones, and thus invalidating the use of current local anomalies as proxies for past climate anomalies.

An additional problem may come from the fact that climatic variables are almost the only predictors used for PSDMs, which to some extent implies assuming that climate factors are the main drivers of species distributions. Although this strong assumption is supported by the often close correlations between current climate and species ranges (see Field et al., 2009 and references therein) it is not exempt from controversy, since other environmental factors, biotic interactions and historical and population processes may also constrain species distributions (Araújo and Pearson, 2005; Lobo et al., 2006; Araújo and Luoto, 2007; Soberón, 2007, 2010; Beale et al., 2008; Jimenez-Valverde et al., 2008; Peterson et al., 2009; Hortal et al.,

2010). However, selecting predictors to account for causal relationships from correlative analyses is, at least, complex (Shipley, 2000), and SDMs are no exception to this (Bahn and McGill, 2007). Both species distributions and climatic variables are spatially structured; this may allow describing the occurrence of a species from many climatic predictors simply because their spatial structures covary to some extent, with or without the existence of a truly causal relationship. This problem can be overcome by selecting predictors that are likely to affect the persistence of the studied species based on previous physiological or biological knowledge (Kearney and Porter, 2009), a kind of information that is quite difficult to obtain for extinct species. In the absence of such information, several statistical techniques can be used to select the most appropriate set of explanatory variables for SDMs, based on either presence (Calenge et al., 2005; Basille et al., 2008) or presence/absence data (Murtaugh, 2009). A key factor here is to select the minimum number of predictors possible, since the more variables are included in the model, the narrower its predictions will be (Beaumont et al., 2005; Bulluck et al., 2006; Soberón, 2010). No matter how thoughtful such predictor selection is, SDMs are basically correlative techniques, so the variables entering into the model may not be causally related with the distribution of the species. This implies that they are forecasting rather than predictive models (sensu Legendre and Legendre, 1998, p. 493), and therefore that although they can sometimes identify the environmental conditions accounting for the distribution of a species in a given period accurately, their predictions can be inaccurate when transferred to other temporal scenarios (Varela et al., 2009).

2.3. The modeling technique

SDM techniques include a diverse array of statistical and mathematical procedures with the only common characteristic being their general purpose: relating data on the observed species distribution to a set of spatially-defined variables – which usually are environmental factors – (Guisan and Zimmermann, 2000; Franklin et al., 2009). These techniques can be divided into: (i) those based on the statistical fitting of data to the theoretical relationships between the species' presence and the environment (e.g. GLMz, GAMs, PLS, CART, and MaxEnt; Austin et al., 1990; Yee and Mitchell, 1991; Austin, 2002; Vaz et al., 2008; Carrascal et al., 2009; Phillips et al., 2009); (ii) machine-learning techniques, that capture the complex spatial patterns in the observed data without assuming any underlying probability distribution (e.g. ANN and GARP; Stockwell and Peters, 1999; Benito Garzón et al., 2007; Olden et al., 2008); and (iii) those based on methods derived from geometrical or set theory, that only use the information coming from the observed presences (e.g. BIOCLIM, DOMAIN, BIOMAPPER, and Mahalanobis Distance; Busby, 1986; Carpenter et al., 1993; Hirzel et al., 2002; Farber and Kadmon, 2003). The first two kinds of techniques require both presence and absence data as input, and are therefore considered *discrimination* procedures, because they aim to generate a function to discriminate among the different states of a variable.

Obtaining reliable absence information for the SDMs carried out with current data is problematic (Lobo et al., 2010) because: (i) it is difficult to distinguish true absences from lack of information when an associated measure of survey effort does not exist; and (ii) the absence of species may have been caused by the effects of non-environmental and contingent factors that are hard to include in any modeling process. These drawbacks force SDM users to incorporate the so-called pseudo-absences and/or background data in order to use discrimination techniques. These absences are either selected at random from areas where the species has not been recorded or the environmental conditions are a priori unsuitable (Ferrier and Watson, 1997; Zaniwski et al., 2002; Engler et al., 2004; Lobo et al., 2006) or simply chosen at random from the entire study area (Stockwell and Peters, 1999; Elith et al., 2006). However, these pseudo-absences

should be used with caution (Lobo and Tognelli, 2011) because the way they are selected has a great influence on the geographic representation of the distribution of the species that is finally obtained (Chefaoui and Lobo, 2008; Anderson and Raza, 2010; Lobo et al., 2010).

Depending on the protocol used to select pseudo-absences (or the decision to use only presences), the geographical response described by SDM results can oscillate between the realized and the potential distribution of the species (Jimenez-Valverde et al., 2008; Colwell and Rangel, 2009; Soberón and Nakamura, 2009). In fact, background absence data selected at random may be appropriate to estimate the probability of use of a resource against its availability in order to identify environmental or habitat preferences (i.e., Resource Selection Functions; Boyce and McDonald, 1999), but not to model realized or potential distributions (McDonald et al., 2002). Choosing absence data for SDMs that are going to be projected into different temporal scenarios separated by many generations presents additional problems. The common spatial, collector and taphonomic biases of paleontological data hamper their use for the validation of SDMs calibrated with current data. Importantly, the same species will show different realized distributions in each period due to non-environmental and contingent effects that prevent it from holding populations in many suitable areas. This prevents the direct use of absence information to calibrate PSDMs; even if these factors can be included in the modeling process for a given temporal scenario, there is no guarantee that their effect would be similar in a different period.

Based on the conceptual and methodological drawbacks described above, in this article we recommend not using discrimination or correlative SDM techniques to model fossil data. This leaves presence records from fossil remains as the only reliable source of information on the past distribution of a species. These data provide valuable information on the environmental conditions for which the species may have had a positive net rate of demographic growth. The location of the regions with similar conditions to those where the species occurred offers a preliminary picture of its potential distribution—the places that could be inhabited by the species in the absence of significant dispersal limitations, local extinctions, competitive exclusion and/or survey biases (Jimenez-Valverde et al., 2008; Colwell and Rangel, 2009; Soberón and Nakamura, 2009; Lobo et al., 2010).

Establishing a robust methodological framework for the use of GCM-based projections of past climate and PSDMs in the emerging field of paleobiogeography should be rooted in careful methodological choices. Although complex methods provide powerful ways of fitting the data, they often provide responses that are too narrow for the environment (Jimenez-Valverde et al., 2008). Choices should instead be based on the capacity of SDM techniques to account for the species' responses to climatic and non-climatic factors for a given period and provide reliable projections of these responses to other moments of time. These conditions are met by the methods based on geometrical or set theory, which are safe from the errors caused by the use of absence data in the training dataset. Given the concerns above, and the problems found when using presence/absence approaches to estimate the past distribution of species (Varela et al., 2009), we argue that although presence/absence SDMs may be suitable for describing the distribution of a species in a single period, they are certainly not adequate for predicting the species' response throughout several scenarios of changing climate. Thus, we stress that discrimination techniques and similar procedures should be avoided for estimating the past distribution of species, because they are quite likely to underestimate their potential distribution (see Jiménez-Valverde et al., 2011).

The different SDM techniques available to predict species distributions from presence-only data (Tsoar et al., 2007; Calenge and Basille, 2008) are based on the estimation of both the species' tolerance range and the species' optimum conditions according to the selected environmental predictors. The potential distribution of a

species can be partially estimated from observed occurrences by means of a Multidimensional Enveloping procedure (MDE; see Busby, 1986), taking into account that: (i) the species may be able to survive outside the environmental conditions provided by the observed localities (Soberón and Peterson, 2005; Soberón and Nakamura, 2009; Varela et al., 2009); and (ii) the species will only inhabit a portion of its fundamental niche that depends on the readily available environmental space (Jackson and Overpeck, 2000). MDE can be used to generate either binary or continuous suitability maps. Binary geographical projections (i.e., suitable versus unsuitable localities) can be obtained by estimating the extreme maximum and minimum environmental values that may be inhabited by the species, to then delimit the suitable conditions in the multidimensional environmental space by means of a generalized intersection procedure, and finally transfer these conditions to the geographical space.

In contrast to MDE, continuous representations of suitability require estimating both the species' tolerances and its environmental optimum, to provide a gradient from more to less favorable conditions. Estimations of the environmental tolerance are highly dependent on the inclusion of extreme occurrences, so reliable information on presences near the physiological limits of the species is very important to estimate its potential distribution. In the specific case of paleontological data, we also recommend training PSDMs with as much past and recent information on presences as possible, thus maximizing the proportion of the full spectrum of environmental conditions inhabited by the species that is sampled by the data (Varela et al., 2009).

If the aim is to obtain information about the variability in the climatic suitability of the species, we suggest using a measure of the environmental distance from each site to the optimum, such as the scale-invariant Mahalanobis Distance (Kadmon et al., 2003; Allouche et al., 2008; Calenge et al., 2008; Etherington et al., 2009). Needless to say, the selection of the climatic optimum is a key point in this case, since it may also highly influence the obtained results. Although MDE can include the distribution of the data within the variables while modeling climatic suitability (Ruegg et al., 2006), this approach will only produce accurate results when using complete and unbiased presence data sets. In the same manner, the mean, median or any other central tendency measure can only provide good estimations of the species optimum when the data set constitutes a reliable subsample of the species' requirements (Nogues-Bravo et al., 2008). Therefore, we argue here that when studying paleontological data it is important to emphasize the role of environmental limits (Huston, 2002) while at the same time avoiding the effect of bias in current and past fossil distribution data. We thus recommend calculating the central point of the n -dimensional environmental space used by the species as the central point of its amplitude ($(\text{maximum} - \text{minimum})/2 + \text{minimum}$), to ensure that the assumed optimum environmental conditions are equidistant from the extreme values (Varela et al., 2010).

3. Former species distribution modeling applications in paleontology

During our literature search for applications of SDMs in paleontological research (ISI Web of knowledge, September 2010; Search criteria: species distribution model + fossil, species distribution model + last glacial maximum, species distribution model + Pleistocene) we found a total of forty-two papers (Tables 1 and 2; see also Nogues-Bravo, 2009; Svenning et al., 2011). These works study changes in the distribution of species from a wide range of taxa, including plants, vertebrates and insects (Jakob et al., 2009; Jezkova et al., 2009; Marske et al., 2009), as well as changes in the areas occupied by forests or biomes (Bonaccorso et al., 2006; Carstens and Knowles, 2007; Hilbert et al., 2007). The extent of analysis is also heterogeneous, varying from regional (e.g. Iberian Peninsula; Benito Garzón et al., 2007) to continental (e.g. Europe, North America; Banks et al., 2006) or even global (e.g. Yesson and Culham, 2006). In this section we review the occurrence data, predictors and SDM

Table 1

Summary of the occurrence data used in the forty-two papers using Paleo-Species Distribution Modeling analyzed in this work.

Geographic extent	Climatic variables			SDM technique	References
	GCM past	Downscaling/variables	Resolution		
Iberian Peninsula	CCSM, MIROC	Current local climate to estimate past local climate	200 m	MaxEnt	Alba-Sánchez et al. (2010)
Western Europe	PMIP2 protocol	Refined grid over Europe	60 km	GARP	Banks et al. (2008a)
Western Europe	PMIP1 protocol	Refined grid over Europe	60 km	GARP	Banks et al. (2008b)
Western Europe and North America	HadCM3	–	100–200 km	GARP	Banks et al. (2006)
Switzerland	–	Use current climate to estimate past climate	1 km	LGMz	Baumann et al. (2005)
Iberian Peninsula	ECHAM3, UGAMP	Current local climate to estimate past local climate	–	Random forest	Benito Garzón et al. (2007)
North Atlantic	Ocean GCM	Interpolation	5'	MaxEnt, "ecophysiological ranges"	Bigg et al. (2008)
Amazon Basin	HadCM3	–	0.1°	GARP	Bonaccorso et al. (2006)
New Zealand	–	Based on current climate and indirect data	100 m	MaxEnt	Buckley et al. (2009)
Iberian Peninsula	ECHAM3	Anomaly data and interpolating by thin-plane splines	1 km	Random forest	Calleja et al. (2009)
South America	ECHAM3	Bilinear interpolation	30'	MaxEnt, Bioclim	Carnaval and Moritz (2008)
Brazilian Atlantic rainforest	–	–	–	MaxEnt	Carnaval et al. (2009)
Western North America	CCSM3	Interpolation	–	MaxEnt	Carstens and Knowles (2007)
Central Europe	CCSM	Current local climate to estimate past local climate	10'	Bioclim	Depraz et al. (2008)
South Africa	–	Use current climate to estimate past climate	15'	Bioclim	Eeley et al. (1999)
Eurasia	S3P, LMDZHR	Current local climate to estimate past local climate	30'	MaxEnt, Bioclim	Flojgaard et al. (2009)
North America	CCSM3, MIROC	Current local climate to estimate past local climate	2.5'	MaxEnt	Galbreath et al. (2009)
America	ECHAM3	Current local climate to estimate past local climate	1 km	MaxEnt, Bioclim, Domain, GAM	Hijmans and Graham (2006)
North Queensland, Australia	–	Use current climate to estimate past climate	80 m	Bioclim	Hugall et al. (2002)
North Queensland, Australia	–	Use current climate to estimate past climate	1 ha	Artificial neural networks	Hilbert et al. (2007)
Eastern North America	MIROC	Interpolation	2.5'	MaxEnt	Jezkova et al. (2009)
South America	CCSM, MIROC	Interpolation	0,01°–0,04°	GARP	Jakob et al. (2009)
North America	CCSM3	Interpolation	1'	MaxEnt	Knowles et al. (2007)
North America	–	Based on fossil and sedimentary data	1°	GARP	Araújo and New (2007, 2009)
New Zealand	–	Use current climate to estimate past climate	100 m	MaxEnt	Marske et al. (2009)
North America	HadCM2	Resampled	0.1°	GARP	Martínez-Meyer and Peterson (2006)
North America	–	Use current climate to estimate past climate	–	GARP	Martínez-Meyer et al. (2004)
Australian wet tropics	–	Estimated from pollen	–	Averaged GLMz	Moussalli et al. (2009)
Eurasia	GENESIS	–	2°	Mahalanobis Distances, MaxEnt, Bioclim	Nogues-Bravo et al. (2008)
Europe	UBRIS-HadCM3	Current local climate to estimate past local climate	1 km	Boosted regression trees	Pearman et al. (2008)
North America	HadCM	Current local climate to estimate past local climate	0.1°	GARP	Peterson et al. (2004)
Central and South America	MIROC, CCSM	Current local climate to estimate past local climate	0.04°	GARP, MaxEnt	Peterson and Nyari (2008)
Europe and the Mediterranean	ECHAM3, UGAMP	Current local climate to estimate past local climate	100 km	MaxEnt	Rodríguez-Sánchez and Arroyo (2008)
America	ECHAM	Current local climate to estimate past local climate	10 km	Bioclim	Ruegg et al. (2006)
Central and South America	ECHAM	Current local climate to estimate past local climate	100 km	MaxEnt	Solomon et al. (2008)
Europe	S3P, LMDZHR	Refined grid over Europe	60 km	MaxEnt, Bioclim	Svenning et al. (2008)
Eurasia and Africa	GENESIS	–	4.5° × 7.5°	LGMz, Bioclim	Varela et al. (2009)
Eurasia and Africa	GENESIS	–	2°	Mahalanobis Distances, Bioclim	Varela et al. (2010)
Indiana and Ohio (United States of A.)	–	Interpolation-ordinary kriging based on sedimentary data	15'	GARP	Walls and Stigall (2011)
North America	CCSM, MIROC	Current local climate to estimate past local climate	2.5'	MaxEnt, GARP	Waltari et al. (2007)
North America	CCSM, MIROC	Current local climate to estimate past local climate	2.5'	MaxEnt, GARP	Waltari and Guralnick (2009)
Global and Australia	BRIDGE	–	2°	Bioclim	Yesson and Culham (2006)

Table 2
Geographic extent, General Circulation Models projected to the past, and downscaling methodologies used to construct the predictors of the Paleo-Species Distribution Models in the forty-two papers analyzed here.

Training data			References
Current data	Fossil data	Target species	
Species distribution maps	–	<i>Abies</i> spp.	Alba-Sánchez et al. (2010)
Georeferenced occurrences	Georeferenced fossil sites	<i>Rangifer tarandus</i> , <i>Cervus elaphus</i>	Banks et al. (2008a)
–	Georeferenced archeological sites	<i>Homo sapiens</i>	Banks et al. (2008b)
–	Georeferenced archeological sites	<i>Homo sapiens</i>	Banks et al. (2006)
Georeferenced occurrences	Georeferenced archeological sites	<i>Rupicapra rupicapra</i>	Baumann et al. (2005)
Species distribution maps	–	19 tree species	Benito Garzón et al. (2007)
Georeferenced occurrences	–	<i>Gadus</i> spp.	Bigg et al. (2008)
Georeferenced occurrences	–	6 trees, 11 birds	Bonaccorso et al. (2006)
Georeferenced occurrences	–	<i>Argosarchus horridus</i>	Buckley et al. (2009)
Georeferenced occurrences	–	<i>Prunus lusitanica</i>	Calleja et al. (2009)
Georeferenced occurrences	–	Forest spp.	Carnaval and Moritz (2008)
Georeferenced occurrences	–	<i>Hypsiboas</i> spp.	Carnaval et al. (2009)
Georeferenced occurrences	–	Frogs, trees and mammals	Carstens and Knowles (2007)
Georeferenced occurrences	–	<i>Trochulus villosus</i>	Depraz et al. (2008)
Species distribution maps	–	Indigenous forests	Eeley et al. (1999)
Species distribution maps	–	Rodents	Flojgaard et al. (2009)
Georeferenced occurrences	–	<i>Ochotona princeps</i>	Galbreath et al. (2009)
Species distribution maps	–	100 plant spp.	Hijmans and Graham (2006)
Georeferenced occurrences	–	<i>Gnarosophia bellendenkerensis</i>	Hugall et al. (2002)
Species distribution maps	–	Forest classes	Hilbert et al. (2007)
Georeferenced occurrences	–	<i>Chaetodipus penicillatus</i>	Jezkova et al. (2009)
Georeferenced occurrences	–	<i>Hordeum</i> species (Poaceae)	Jakob et al. (2009)
Georeferenced occurrences	–	<i>Melanoplus marshalli</i>	Knowles et al. (2007)
–	Georeferenced fossil sites	Subfamily Equinae	Maguire and Stigall (2009)
Georeferenced occurrences	–	<i>Agyrtodes labralis</i>	Marske et al. (2009)
Georeferenced occurrences	Georeferenced fossil sites	8 tree species	Martínez-Meyer and Peterson (2006)
Georeferenced occurrences	Georeferenced fossil sites	23 mammal species	Martínez-Meyer et al. (2004)
Georeferenced occurrences	–	<i>Saprosincus</i> spp.	Moussalli et al. (2009)
–	Georeferenced fossil sites	<i>Mammuthus primigenius</i>	Nogues-Bravo et al. (2008)
Species distribution maps	Georeferenced fossil sites	tree spp.	Pearman et al. (2008)
Georeferenced occurrences	–	<i>Aphelocoma jays</i>	Peterson et al. (2004)
Georeferenced occurrences	–	<i>Schiffornis</i> sp.	Peterson and Nyari (2008)
Georeferenced occurrences	Georeferenced fossil sites	<i>Laurus</i> sp.	Rodríguez-Sánchez and Arroyo (2008)
Georeferenced occurrences	–	<i>Catharus ustulatus</i>	Ruegg et al. (2006)
Georeferenced occurrences	–	<i>Atta</i> spp.	Solomon et al. (2008)
Species distribution maps	–	22 tree spp.	Svenning et al. (2008)
Georeferenced occurrences	–	<i>Crocota crocuta</i>	Varela et al. (2009)
Georeferenced occurrences	Georeferenced fossil sites	<i>Crocota crocuta</i>	Varela et al. (2010)
–	Georeferenced fossil sites	8 brachiopod species	Walls and Stigall (2011)
Georeferenced occurrences	–	20 terrestrial vertebrates	Waltari et al. (2007)
Georeferenced occurrences	–	13 mammal species	Waltari and Guralnick (2009)
Georeferenced occurrences	Georeferenced fossil sites	<i>Drossera</i> sp.	Yesson and Culham (2006)

techniques used in these papers to provide a synthesis of the basic characteristics of the Paleo-Species Distribution Modeling applications conducted so far. This complements the former reviews by [Nogues-Bravo \(2009\)](#) and [Svenning et al. \(2011\)](#) by discussing in detail the limitations of former PSDM papers based on the theoretical and practical needs of this type of studies discussed above.

3.1. The occurrence data

The training datasets include current species' occurrences in 29 of the 42 papers, and current species distribution maps in eight papers; only five papers are based just on georeferenced fossil records and another eight use both current and past occurrences (see [Table 1](#)). In spite of such variety of data types, these papers rarely comment on the geographic and temporal extent of the data, the completeness of the presence data or how absence data was selected (but see [Nogues-Bravo et al., 2008](#) and [Varela et al., 2010](#)). The geographic extent of the training data is usually the region where the model will be projected; in some papers this extent covers the total distribution area of the species but in others it does not ([Banks et al., 2008a](#)), a practice that can be a major issue for the reliability of PSDMs. Failing to include data

from the entire geographic distribution of a species can result in sampling a spuriously narrow range of environmental tolerance, which ultimately results in an underestimation of its potential distribution ([Thuiller et al., 2004](#); [Sánchez-Fernández et al., 2011](#)).

Importantly, the majority of PSDM applications have so far been trained using only a single temporal scenario ([Table 1](#)). In almost three quarters of the studies (29 cases) the SDM was calibrated using only the current distribution of the species (see [Table 1](#)), a practice that may fail to represent its past distribution ([Varela et al., 2009](#)). In fact, the only two papers that evaluate the assumption that the projections from a PSDM calibrated in a given period are able to predict the presence of the species in a different moment of time give mixed results; while [Nogues-Bravo et al. \(2008\)](#) found that the transferability between periods provided adequate results, [Varela et al. \(2009\)](#) found inconsistency between the projections for different periods. This lack of evaluation of the assumption of temporal transferability is further complicated by the use of absence data to train the models in 36 papers (see [Table 2](#)), 27 of which use either MaxEnt or GARP with background absences. As discussed above, the use of these absence data for training the models prevents them from identifying the potential distribution of the species.

3.2. Predictor variables

Twenty-four out of the 42 papers analyzed downscale the climatic predictors (see Table 2). Fifteen of these studies use the spatial patterns, amplitude and sign of current climatic anomalies at the local scale to construct fine-scale climatic layers, assuming that they are identical through time. Seven papers adopt an even simpler strategy and construct the variables describing past climate simply by adding or subtracting certain values to the current climatic layers. As discussed above, both methods are likely to include significant errors in the climatic predictors. Nevertheless, the lack of a standard for downscaling past climate scenarios includes a source of variability in the predictors between studies, that hampers the direct comparison of their results, even when they refer to the same geographic area and/or temporal scenario.

Although it is well known that the number and identity of predictors have a major influence on SDM results, few papers choose these independent variables based on an ecologically meaningful strategy. In fact, the predictors used in the analysis are generally chosen on the basis of their easy availability. As a result, the 19 WorldClim variables (available at <http://www.worldclim.org/>) are the most used predictors for current time. GCM-derived climatic layers for past scenarios are more difficult to access, and therefore a diverse array of GCMs has been used for this task. Given the difficulty of extrapolating GCMs to the past and the limitations in projecting complex aspects of climate, Annual Mean Temperature and Annual Precipitation are also the predictors most commonly included in PSDMs. There are, however, some exceptions to these opportunistic strategies, and in a few studies the selection of predictors is based on the researcher's knowledge about the specific factors that may affect the distribution of the species (e.g., Bigg et al., 2008; Walls and Stigall, 2011).

3.3. Modeling techniques

Most of the analyzed studies use discrimination techniques which incorporate some kind of pseudoabsence data to estimate the past distribution of species, including MaxEnt, GLMz, GAM, GARP, RF, BRT and ANN (see Table 2). From these, MaxEnt is the most popular, being used in nearly half of the papers (19 cases), followed by GARP (13 cases). Geometrical or set theory based techniques are used in 12 papers; all of them use Bioclim, together with Mahalanobis Distances in two occasions (Nogues-Bravo et al., 2008; Varela et al., 2009).

3.4. Problems and limitations of former PSDM approaches

A diverse array of methods has been used so far to estimate the past distribution of species. According to the conceptual and methodological issues discussed above, we suspect that most former PSDM approaches could be overestimating the role that climatic changes had on past species' range shifts and/or extinction events. In general, the choice of the distributional data used to train the models is based only on the type of data available (current, fossil, or both), rather than on their adequacy for the question at hand. These works make the implicit assumption that the available information about the species is sufficient to estimate its climatic requirements and therefore to calibrate a model that can safely be projected to the past. Unfortunately, we believe that this assumption may prove to be false in many cases. The limited quality of the data on both recent and fossil occurrences may often fail to represent the full spectrum of climatic conditions at which the species can inhabit, so given how SDM results depend on the data used to train them, current common practice in PSDM could be regularly leading to underestimate the geographic ranges of the species in different periods of time. The probability of underestimating the species' potential requirements will, however, diminish if the occurrences in the training dataset

cover the entire geographic and temporal extent in which the species has lived, as well as if the inclusion of any kind of absence data is avoided (see above, and also Jimenez-Valverde et al., 2008).

We also advocate that PSDMs should be built based on a careful selection of climatic predictors, a step that is commonly overlooked in the reviewed literature. To do this, and in the absence of previous knowledge on the actual species' requirements, we recommend using multivariate niche description techniques such as Ecological-Niche Factor Analysis (ENFA; Hirzel et al., 2002; Basille et al., 2008) to select those variables with higher probabilities of being causally related to the species' occurrences. Also, the projections of PSDMs at different moments of time should be cross-examined whenever possible, to ensure that models calibrated with data from a certain period are able to describe the occurrence of the species in other moments of time.

Finally, we want to stress that the SDM techniques used should be appropriate to the questions posed. Different SDM procedures produce wider or narrower geographic predictions and consequently they can have a profound impact in the biological interpretation of the results (Jimenez-Valverde et al., 2008). We thus believe that in the absence of any supplementary information about the species' climatic requirements, the methodological approach used for PSDMs needs to be conservative. While data for these analyses is typically composed of samples of the species' geographic range at different moments of time, these ranges are dynamic and vary as a consequence of the interactions between the potential distribution of the species and both climatic and non-climatic factors. Given the importance of non-climatic factors in preventing species from occurring in climatically suitable areas, discrimination techniques based on absence information should be discarded for the estimation of the climatic niche and the past distribution of species, because they would underestimate its potential distribution. In this context, we think that the most adequate option to predict the potential distribution of a species through time to use the more conservative multidimensional climatic envelopes, such as Bioclim. Similarly, Mahalanobis Distance would be appropriate when the aim is to estimate the climatic suitability based on a hypothetical climatic optimum.

4. Future prospects for paleobiogeography

The generalization of Paleo-Species Distribution Modeling approaches has great potential for generating new paleontological information and hypotheses in the forthcoming years. To take full profit of such potential it is however crucial: (i) to create a global database to compile all distributional information available for the Pleistocene; (ii) to develop high resolution climatic layers for different past scenarios by means of a widely-agreed on standardized downscaling protocol; and (iii) the in-depth investigation of the nature and temporal variation of species-climate relationships, as well as of which SDM methodologies provide the most adequate simulations of these changing relationships.

The analysis of the Pleistocene distribution of any species implies using data from different fossil sites. Therefore, one of the first goals for the development of modern paleobiogeography should be to establish a global open-access database, similar to the ones available for current species, such as the GBIF network (<http://www.gbif.org/>). Some initiatives are already aiming to provide databases on fossil data on the Internet, including the Paleobiology Database (<http://www.paleodb.org/cgi-bin/bridge.pl>), Faunmap (<http://www.museum.state.il.us/research/faunmap/>), Neotoma (<http://www.neotomadb.org/>), the printed compilation of data in *Evolution of Tertiary Mammals in North America* (Janis et al., 2005, 2008), Stage Three Project (<http://www.esc.cam.ac.uk/research/research-groups/oistage3/>) and NOW (Neogene Old World database; <http://www.helsinki.fi/science/now/>). However, several points need to be improved in these databases. First, the available information about fossil dating is extremely broad. As an

example, *Homo antecessor* is roughly assigned to the Pleistocene (1.810 to 0.011 Ma) (query on The Paleobiology Database) when it has been dated by paleomagnetism as being not older than 780 Ma, and assigned to the end of the Earlier Pleistocene using ESR and U-series (Falgueres et al., 1999). Second, the temporal and spatial scopes of these databases are limited. For example, the Stage 3 Project compiles European fossil sites previous to the Last Glacial Maximum, and FAUNMAP is geographically restricted to North America. Third, all of these databases have rigid taxonomic classifications. Pleistocene mammal taxonomy has been changed in the light of new evidence (Alroy, 2003) and fossil records could be reclassified in the future due to the appearance new evidence based on ancient DNA studies or any other new technology. Therefore, any durable database should allow using as many taxonomic classifications as would be necessary, providing taxonomic fields that allow eventual changes in the systematic of the group or the reassignment of specific fossil remains to different taxa in the light of future revisions. The experience of neosystematicians in building standards for biodiversity information (see the most updated information on the Taxonomic Database Working Group at <http://www.tdwg.org/>) will make the design of such a paleodistributional database relatively easy. In any case, a good knowledge on the species and/or taxonomic group studied is crucial for the taxonomic reliability of the information gathered in the database. Similar to biodiversity databases on recent species, an authoritative and comprehensive revision of all the fossil remains recorded in the database and a periodic review of the taxonomic status of all these records are necessary for the database to provide accurate information on known past species' occurrences.

Most difficulties in gathering information about the Pleistocene fossil record may be trivial. Estimating the past geographic location of the fossil remains should not be a significant problem, for the distribution of landmass in the Pleistocene is similar to the current one except for the coastline variations in relation to the glacial cycles (Peltier, 2007). Therefore, in the absence of translocations the original location of the remains can be safely assigned to the current location of the deposit. To guarantee the success of the database on paleodistributional data we suggest following the philosophy of the community of molecular biologists, where many journals require the submission of any sequence information used in the manuscript to a unified database, the GenBank (<http://www.ncbi.nlm.nih.gov/genbank/>), prior to publication. A similar strategy followed by Paleontology, Ecology and Biogeography journals would help by establishing the same philosophy in paleontological research, encouraging researchers to upload the faunal lists of their fossil sites into an open access database. The unified source of information that would be obtained with such a database will constitute a basic resource for the study of the dynamics of species ranges through time, the relationships between species distribution and climatic change, or any other spatio-temporal pattern, and hence should be the pillar of the development of modern Paleobiogeography and Paleocology.

A unified database on the distribution of the diversity in the past is not the sole requisite for the advancement of paleobiogeography. Species' relationships with climate are ultimately determining the long-term viability of local populations, and are therefore dependent on fine-scale climate patterns (Seo et al., 2009), so understanding these relationships also requires the development of high-resolution climatic layers for past periods. Given the uncertainty inherent to GCMs, the development of different temporal simulations may also help to assess the robustness of PSDM results to differences in the parameterization of the climatic models (see also Araújo and New, 2007). Also, the projection of GCMs to continuous temporal series of climatic projections rather than to key moments in time could permit the creation of temporally dynamic models, which would in turn help to analyze hypotheses on the dispersal of species and their range dynamics. Finally, we argue that more theoretical research is needed to understand the

species–climate relationships, following the line of research recently reopened by Soberón (2007, 2010) and Soberón and Nakamura (2009) (see also Kearney, 2006; Jimenez-Valverde et al., 2008 or Colwell and Rangel, 2009). Such research will allow the identification of the factors that should be taken into account to simulate the impacts that climatic changes have on species distributions.

5. Conclusions

The study of Pleistocene biogeography could provide new information about the biological consequences of climatic changes. The development of Paleo-Species Distribution Models can be a central part of such research, benefitting from the information on the past occurrence of species available from the fossil record, the development of Global Climate Models and their projection to past scenarios, and the current theoretical advances on the relationship between the fundamental niche of the species and their geographical distribution. However, three particular aspects of PSDMs must be taken into account for such development, (i) the difficulty in obtaining reliable information on species' absences; (ii) that their results must be extrapolated to transfer their projections over long time periods; and (iii) the difficulty of validating these projections. Here we argue that the fossil data used to calibrate the PSDMs should include as much information as possible, trying to sample the entire geographic and temporal extent of the species' distribution. Furthermore, in our opinion, presences are the only data that are both available and reliable enough to be used in PSDMs; by taking advantage of this information it would be possible to obtain partial estimates of the climatic niche of the species that can subsequently be used to generate hypotheses on the distribution of the species in other periods. On the other hand, absence and pseudo-absence data should be avoided, because they could add misleading information that would make the description of species–climate relationships difficult.

Although the predictors used to build PSDMs should preferably have adequate resolution to describe these relationships, we recommend not to downscale the geographical projections of GCMs using simplistic rules. Here it would be preferable to assume that the resolution of the study is limited to coarse grains, or to wait for the development of regional GCMs. The selection of the adequate predictors is also a key point in the construction of PSDMs; here we suggest using ENFA analysis to select the most biologically meaningful variables. Based on the restriction to presence-only SDMs that we suggest, the most adequate PSDM methodologies would be either (a) climatic envelopes based on the climatic tolerance range of the species when the goal is to detect geographic range shifts in relation to climatic changes; or (b) distance-based techniques such as Mahalanobis Distances when some continuous information about the climatic suitability for the species is required, although in this case the selection of the species' optimum should be discussed and justified. By following all of these recommendations we believe that we will be able to diminish the errors in the estimations of the potential distribution of the species, and therefore ensure that the future use of PSDMs will bring robust results and at the same time stimulate ideas in the fields of Paleocology and Paleobiogeography.

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