

Predicting climate-induced range shifts: model differences and model reliability

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Abstract

Predicted changes in the global climate are likely to cause large shifts in the geographic ranges of many plant and animal species. To date, predictions of future range shifts have relied on a variety of modeling approaches with different levels of model accuracy. Using a common data set, we investigated the potential implications of alternative modeling approaches for conclusions about future range shifts and extinctions. Our common data set entailed the current ranges of 100 randomly selected mammal species found in the western hemisphere. Using these range maps, we compared six methods for modeling predicted future ranges. Predicted future distributions differed markedly across the alternative modeling approaches, which in turn resulted in estimates of extinction rates that ranged between 0% and 7%, depending on which model was used. Random forest predictors, a model-averaging approach, consistently outperformed the other techniques (correctly predicting >99% of current absences and 86% of current presences). We conclude that the types of models used in a study can have dramatic effects on predicted range shifts and extinction rates; and that model-averaging approaches appear to have the greatest potential for predicting range shifts in the face of climate change.

Keywords: climate change, climate-envelope models, extinction, geographic range, model averaging, model prediction, random forest predictors

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Introduction

Global temperatures have risen an average 0.6 °C over the past century (Houghton *et al.*, 2001). Recent studies suggest that this climate change has caused shifts in the geographic ranges of both plants and animals (Parmesan & Yohe, 2003; Root *et al.*, 2003). Given that average global temperatures are predicted to rise between 1.4 and 5.8 °C over the next century (Houghton *et al.*, 2001), it is likely that many species will undergo dramatic range shifts in the future. To anticipate the effects of climate change, and to identify conservation strategies that might mitigate the undesirable consequences of climate change, it is essential that we develop models that link the distributions of species to alternative scenarios of climate change.

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Several studies have attempted to predict future range shifts, often with the goal of estimating climate-induced extinction rates (Williams *et al.*, 2003; Thomas *et al.*, 2004). Most predictions of future species distributions rely on what are commonly called climate-envelope or niche models. Collectively, these models can be referred to as bioclimatic models because they relate biotic distributions to climate. At large spatial scales, the distributions of plant and animal species are, in part, determined by climatic factors (Lomolino *et al.*, 2005). Bioclimatic models attempt to relate species current geographic distributions to a set of current climatic factors. Relatively simple climate variables are used to define the abiotic conditions, or 'climate envelope' in which a species exists. Predicted future climate variables, usually derived from a general circulation model (GCM), are used as input for these models to predict future distributions.

These predictive models are generally either based on statistical techniques (e.g. Thuiller *et al.*, 2004c) or machine learning approaches (e.g. Peterson *et al.*,

2002). There are several limitations to this correlative approach. In general, these models do not account for biotic interactions, evolutionary change, or dispersal (Pearson & Dawson, 2003). Furthermore, because they are correlative in nature, there is no guarantee that the current relationships between a species' distribution and the current climate will adequately predict the future distribution of a species. Despite these limitations, these approaches currently provide the best methods for predicting climate-induced range shifts for large numbers of species. Consequently, they have been used extensively in a wide range of studies (Pearson *et al.*, 2002; Huntley *et al.*, 2004; Thomas *et al.*, 2004; Thuiller *et al.*, 2005a, b).

Unfortunately, researchers have reported large uncertainties and error rates in these bioclimatic model predictions, and we have little understanding of which, if any of the various modeling approaches is most reliable (Thuiller, 2003; Segurado & Araújo, 2004). In this paper, we report on a systematic comparison of all the major approaches to predicting range shifts with a common data set and common metrics for estimating error rates. Our goal was to quantify the types of errors associated with bioclimatic models and to determine whether any approach clearly outperforms the alternatives. The approaches we examined were: generalized linear models (McCullagh & Nelder, 1989), classification trees (Breiman *et al.*, 1984), generalized additive models (GAM, Hastie & Tibshirani, 1990), random forest predictors (Breiman, 2001), artificial neural networks (Ripley, 1996), and genetic algorithms for rule-set prediction (GARP, Peterson *et al.*, 2002).

Methods

We compared the model accuracy and the future predictions of the six different modeling approaches described below by applying each approach to 100 randomly selected mammal species in the western hemisphere. All analyses were conducted on a 50 km × 50 km resolution grid consisting of 15 323 cells. Current species distributions were based on digital range maps (Patterson *et al.*, 2003). We selected the 100 species at random from 1022 mammals with ranges occupying at least 50 grid cells. We chose this threshold to eliminate many species for which it was impossible to build predictive models while still including species with a wide range of geographic range sizes.

Current climate data were derived from average monthly precipitation and temperature values from 1961 to 1990 for the land surface of the globe at 0.5° resolution (Leemans & Cramer, 1991). For that 30-year period, we calculated mean annual temperature, average temperature of the hottest and coldest months, and

degree-days over 5 °C. We also calculated average yearly precipitation as well as precipitation in the hottest, coldest, wettest, and driest months.

In addition to climate data, we used land-cover data to predict current and future species distributions. Most models used to predict climate-induced range shifts have used only climate data, making the assumption that climate will act as a surrogate for land cover for species that respond to vegetation patterns. Although climate might act as a suitable surrogate for vegetation in a static environment, climate-induced shifts in vegetation will depend in part on responses to changing CO₂ levels, as well as the distribution of soil types. Therefore, for many animal species, we should be able to make more accurate future projections if models include vegetation.

Current land cover was derived from both predicted current potential vegetation and measured land cover derived from Advanced Very High Resolution Radiometer (AVHRR) satellite data (Loveland *et al.*, 1999). Predicted current vegetation types were produced using the Mapped Atmospheric-Plant-Soil System (MAPSS) model (Neilson, 1995). Although measured vegetation provides a more accurate representation of current vegetation, we chose to use the predicted current vegetation to best correspond with the classification of predicted future vegetation for the years 2061–2090. MAPSS predictions of current potential vegetation have been shown to closely approximate other potential vegetation classifications (Bachelet *et al.*, 2001). We overlaid the 44 land-cover classes of predicted potential current vegetation from the MAPSS model with five agriculture classes and one urban and suburban land-cover class from the AVHRR-derived land-cover data to produce the new 50-class land-cover data set used for building the models.

Predicted future climate data were produced using the Hadley Climate Centre's HADCM2SUL model (Johns *et al.*, 1997) using Intergovernmental Panel on Climate Change predicted future greenhouse gas contributions (IS92a) for the years 2061–2090 (Kattenberg *et al.*, 1996). This model and greenhouse gas contribution scenario together generally predict larger increases in precipitation and smaller increases in temperature (particularly for North America) than do more recent models. Although a wide array of more recent GCM predictions based on alternative emissions scenarios exist, the purpose of our study was not to draw conclusions about the future, but to compare the differences in predictions resulting from using different bioclimatic modeling approaches. Using the future climatic predictions, we calculated the same set of nine climate variables for all 0.5° grid cells. Predicted future land cover was produced with the MAPSS model using

the predicted climate data for input. For the purposes of these analyses, we assumed no change in the distribution of agriculture and urban–suburban areas. We overlaid the predicted future potential vegetation data with the current agriculture and urban–suburban data to produce predicted future land cover. All data compiled at 0.5° resolution were projected to the 50 km resolution grid. For comparison, a 0.5° cell is approximately 3025 km² at the equator, 2139 km² at 45° latitude, and 514 km² at 80° latitude.

Modeling approaches

For all six modeling approaches, we used the presence and absence of a species as the response variable and the set of nine continuous variables representing current climate and one categorical variable representing the 50 land-cover classes as predictors. All models except the GARP models were built using the R software package (version 1.9.1). For all 100 species, we selected a training- and a test-data set. For the training set, we randomly selected 80% of all species presences and 80% of all species absences. For each species, we then used the remaining 20% of the data for testing the models and determining their errors in terms of absences falsely predicted as presences (commission error), and presences falsely predicted to be absences (omission error).

Generalized linear models. Generalized linear models offer a slightly more flexible modeling framework than basic linear regression models as they allow for the modeling of alternative distributions in the response variable and nonconstant variance functions (Guisan *et al.*, 2002). We built logistic regression models (generalized linear models with an assumed binomial error distribution) using a combined backward- and forward-stepwise selection process. Variable inclusion was based on Akaike's information criterion (Chambers & Hastie, 1991). We modeled all linear and second-order polynomials of the climatic predictor variables. Because the test-data sets for 21 species contained land-cover classes that were not found in the training sets of those species, we chose to drop the land-cover variable from the models for these species.

Classification tree models. Classification trees, and regression trees, their counterpart for analyzing continuous response variables, are nonparametric modeling approaches (Breiman *et al.*, 1984; Venables & Ripley, 2002). Both techniques involve the recursive binary partitioning of data. Each split of the data is made using the predictor variable and the point along that variable's distribution that divides the data into the

two most homogeneous groups with respect to the response variable. The result is a tree-like structure with one root node and a number of terminal nodes. In a classification tree, the proportional class membership of the observations in a terminal node form the basis for predicted probabilities. De'ath & Fabricius (2000) provide excellent examples of the use of tree-based models for ecological analyses. We fit classification trees using the RPART package in R originally designed for S-Plus (Therneau & Atkinson, 1997). Because most trees tend to over-fit the data, we selected the optimal tree size using the modal size suggested by fifty 10-fold cross-validations applying a 1-SE rule (De'ath & Fabricius, 2000).

GAM. GAMs are similar to generalized linear models, but they are more flexible because they do not require a specific response curve to be fit to the predictor variables (Hastie & Tibshirani, 1990). Smoothing functions allow data-driven response curves to be fit for each predictor variable. We fit GAMs using penalized regression splines (Wood & Augustin, 2002). This approach takes advantage of generalized spline smoothing (Wahba, 1990) but can be equally or less computationally expensive than backfit GAMs. To increase the speed of the modeling process, we prescreened each variable by fitting a GAM model for that variable alone. We dropped all variables for which the fitting algorithm was unable to converge. Variable selection for those variables included in the modeling process was based on smoothness penalties in conjunction with a shrinkage parameter. Variables were effectively dropped from a model based on the fit smoothing parameter. We used the MGCV package in R to fit all GAM models (Wood & Augustin, 2002). As for the generalized linear models, we did not include the categorical land-cover variable in the models built for the 21 species for which the test-data set contained land-cover classes not found in the training-data set.

Random forest predictors. Random forest predictors are a model-averaging approach based on regression or classification trees (Breiman, 2001). Instead of building one tree model, the random forest algorithm builds multiple trees using randomly selected subsets of the observations and random subsets of the predictor variables. The predictions from the trees are then averaged (in the case of regression trees) or tallied using a voting system (for classification trees). We used the R package RandomForest to build random forest predictors. As part of the random forest procedure, 500 classification trees were built for each species. To build each tree, 12 258 observations were selected at random, with replacement, from the training

set. For each split in these trees, three predictor variables were selected at random from the full set of 10 predictor variables as candidates for that particular split.

Artificial neural networks. Artificial neural networks are a machine-learning approach based on real neural networks (Ripley, 1996). The networks are composed of a series of interconnected nodes (neurons) which receive and process input signals and potentially generate output signals. A network is trained on a data set to recognize the patterns in the data. We built artificial neural networks using the NNET package in R which was based on the S-Plus package NNETW (Venables & Ripley, 2002). These feed-forward networks had one hidden layer with eight nodes. To train the network, we used 5000 presence and 5000 absence observations selected at random, with replacement, from the training-data set. Trial and error determined that these 10 000 observation data sets were most effective and efficient for training the networks. To produce more robust predictions, we built 10 networks for each species and averaged the model predictions (Thuiller, 2003; Segurado & Araújo, 2004).

GARP. GARP is a machine learning-based approach that uses a genetic algorithm (a stochastic optimization technique) to assemble a set of rules to define a species' range (Stockwell & Noble, 1992). The approach was developed expressly for predicting species distributions. The rules used by the GARP algorithm include logistic relationships, climate envelopes (Nix, 1986), and simple Boolean rules. We used the Unix version of GARP to build 500 models for each species. All models were selected from all rule types. GARP limits model training sets to 2500 observations. For each of the 500 models, we selected 1250 presences and 1250 absences, with replacement, from the training-data set for the given species. For each species, we used Cohen's κ statistic (Monserud & Leemans, 1992), calculated using the training-data set, to select the 10 best performing models from the set of 500 models. We combined the binary predictions of these 10 models to produce a predicted probability of presence.

Model comparisons

Using the reserved test-data set, we computed four different metrics to compare the performance of the six different modeling approaches. The first three of these approaches included the percentage of the presences correctly classified, the percentage of the absences correctly classified, and Cohen's κ . Because all six modeling approaches produced predicted prob-

abilities, calculating these three metrics required selecting a threshold with which to classify predicted presences and absences. We used receiver-operating characteristic (ROC) curves to select the optimal threshold, assuming that predicting presences correctly was twice as important as predicting absences correctly (Fielding & Bell, 1997). This is a conservative approach and should generally reduce the chances of overestimating future range contractions. In addition to the three metrics listed above, we used the area under the ROC curve (AUC) to provide an assessment of model performance that was independent of a specific classification threshold (Fielding & Bell, 1997).

There are advantages and disadvantages to using each of these different measures of model accuracy. The percentage of correctly predicted presences and absences are the simplest and most straightforward measures. The main drawback to using these measures is that both are required to assess the accuracy of a model. The large extent of our study is also likely to inflate the percentage of correctly predicted absences. This inflation will be more pronounced for species with small ranges. Both κ and AUC are commonly used statistics for assessing overall model accuracy taking both omission and commission error into account. The κ statistic makes an adjustment for chance agreement and that adjustment can produce different accuracy estimates that depend on the structure of the data set in question (Stehman, 1997). Because AUC assesses accuracy independent of a given classification threshold, it likely produces an overly optimistic estimate of model accuracy when applied to test-set data. Given the various advantages and disadvantages to using these different measures, we chose not to use any one single measure to assess model accuracy in our analyses.

For all four measures of accuracy, we compared model performance across model types using Wilcoxon's signed-ranks tests with a Holm correction for conducting multiple tests (Holm, 1979).

Future predictions

We used the models to predict future geographic ranges under two alternative dispersal scenarios. First, we assumed that a species would be able to completely disperse into any new geographic range. For the second scenario, we assumed that a species would be unable to disperse from its current range. These two extreme assumptions have been made in several recent studies with which we wish to draw comparisons (Peterson *et al.*, 2002; Thomas *et al.*, 2004). Realistic future range shifts are likely to fall somewhere between these two extremes.

Table 1 Accuracy of six different modeling approaches used to model the current geographic ranges of 100 mammal species in the western hemisphere

Model	% presences correct	% absences correct	κ	AUC
GLM	77.7 (17.3), a	98.9 (1.4), a	0.68 (0.13), a	0.970 (0.017), a
TREE	55.5 (19.0), b	99.6 (0.5), b	0.63 (0.13), b, c	0.838 (0.072), b
GAM	68.9 (19.3), a	99.1 (1.4), a	0.62 (0.15), a, b	0.966 (0.022), c, d
RF	86.0 (12.1), c	99.6 (0.3), c	0.86 (0.09), d	0.995 (0.003), e
ANN	75.6 (12.5), a	98.2 (2.1), d	0.58 (0.13), c, e	0.968 (0.017), a, c
GARP	85.0 (6.2), c	95.9 (2.7), e	0.53 (0.17), e	0.962 (0.023), d

Accuracy was assessed using a reserved test-data set composed of a randomly selected 20% of the presences and 20% of the absences for each species. Values reported are the medians and one half of the interquartile range of the accuracy of the model predictions for 100 species. Values with the same letters were not significantly different ($P > 0.05$).

GLM, generalized linear model; TREE, classification tree; GAM, generalized additive model; RF, random forest; ANN, artificial neural network; GARP, genetic algorithm for rule-set prediction; AUC, area under the receiver-operating characteristic curve.

Results

How did alternative modeling approaches affect the types of error and uncertainty in our analyses? The amounts and types of error were markedly influenced by which approach was used to predict range shifts (Table 1). The most significant consistencies in model performance were the over-prediction of current presences (commission error) by the neural networks and GARP models, the under-prediction of current presences (omission error) by the classification tree models, and the small number of errors predicted by the random forest models. For example, classification trees often incorrectly predicted current presences (median of 56% correct). This is a higher rate of omission error than produced by the other five approaches (medians of 69–86% correct presences). GARP models tended to have higher commission error rates than the other approaches, correctly predicting 96% of test-set absences compared with correct prediction rates of between 98% and 100% of absences for the other types of models. The spatial patterns of both commission and omission errors also differed across the six modeling approaches (e.g. Fig. 1). Whereas the commission errors of the GARP models and artificial neural networks tended to be relatively widely distributed, the few errors that the random forest models produced were generally clustered tightly around the area occupied by the species (Fig. 1).

We also found that different modeling approaches produced dramatically different predictions of future range shifts for many species. Not surprisingly, these differences were heavily influenced by assumptions regarding dispersal. On average, if one assumes no dispersal, so that species cannot move to occupy newly predicted portions of their range, only 19% of the cumulative future range of a species was similarly

predicted by all six models. The percent agreement was even lower (11%) when full dispersal (species can fully exploit new range space that arises in the future) was assumed. For example, for the black tufted-ear marmoset (*Callithrix penicillata*), assuming unlimited dispersal, the generalized linear model and classification tree predicted contractions of 70% and 58% of the current range, respectively, whereas the artificial neural network and the GARP model, respectively, predicted expansions of 180% and 53% of the range (Fig. 2).

These differences in model prediction translated into different estimates of overall range contractions and expansions as predicted by the alternative modeling approaches (Fig. 3). When we assumed unlimited dispersal, classification trees predicted range contractions of over 50% for 36% of the species in the study compared with neural networks and GARP models, which respectively predicted similar range contractions for 16% and 17% of all species. Because these models are often used to predict extinction rates, it is worth noting that depending on the modeling approach used, extinction rates ranged from 0% to 7% assuming unlimited dispersal and from 6% to 14% assuming no dispersal. In general, GARP models predicted the most drastic range expansions including at least a tripling in range size for 19% of all species compared with classification tree models that predicted at least a tripling in range size for only 7% of the species.

All of the differences among models would be daunting were it not for the finding that one modeling approach clearly performed better than all of the alternatives. In particular, random forest models had the highest median performance scores across all four measures of model accuracy (Table 1), and were consistently ranked the best performing of the six model types (Fig. 4). Random forests were the best performing models with respect to AUC and κ for 88% of species.

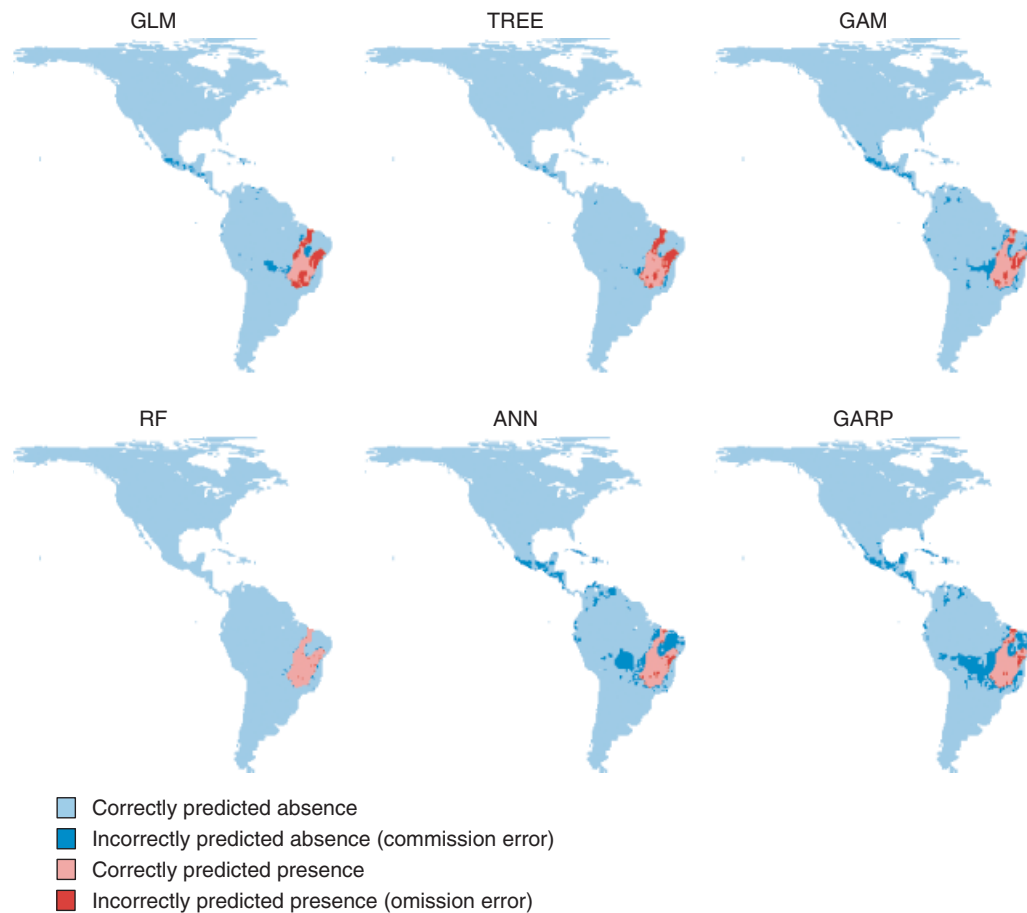


Fig. 1 Maps of the current range of the black tufted-ear marmoset (*Callithrix penicillata*) as predicted by six alternative modeling approaches. See Table 1 for an explanation of model abbreviations.

The superiority of the random forest models as measured by AUC, κ , and the percentage of correctly predicted presences was independent of species range size. Range size did, however, affect the accuracy of the models and the ranking of some of the approaches. For all approaches, the percentage of correctly predicted presences increased with initial range size (Fig. 5), whereas the percentage of correctly predicted absences decreased with initial range size (Fig. 6). Range size had little effect on the ranking of the modeling approaches with respect to the percentage of correctly predicted absences (Table 2) but more substantially affected the ranking of the approaches with respect to correctly predicted presences (Table 3). In particular, GARP models were the best at predicting presences for the species with the smallest ranges. This reduced omission error came at a cost, however, because GARP models had the highest commission error rates.

We noted some distinct differences in the models built for the 21 species for which land-cover data were not used in the modeling process. For all but the generalized linear models and random forest models,

the predictions for these species had higher commission error rates and lower omission error rates than the other 79 species for which land-cover data were used.

Discussion

Differences in bioclimatic modeling approaches

There are several different approaches to predicting changes in species distributions as a result of climate change (Iverson & Prasad, 1998; Shafer *et al.*, 2001; Pearson *et al.*, 2002; Araújo *et al.*, 2004; Meynecke, 2004; Thomas *et al.*, 2004). With few exceptions, previous studies have found very little consistency in the performance of these alternative approaches (Moisen & Frescino, 2002; Robertson *et al.*, 2003; Thuiller, 2003; Segurado & Araújo, 2004). We have found similar inconsistency among models. Others have demonstrated that certain modeling approaches work differently for groups of species that demonstrate qualitatively different relationships with their environments (Segurado & Araújo, 2004). The six modeling techniques that we applied in

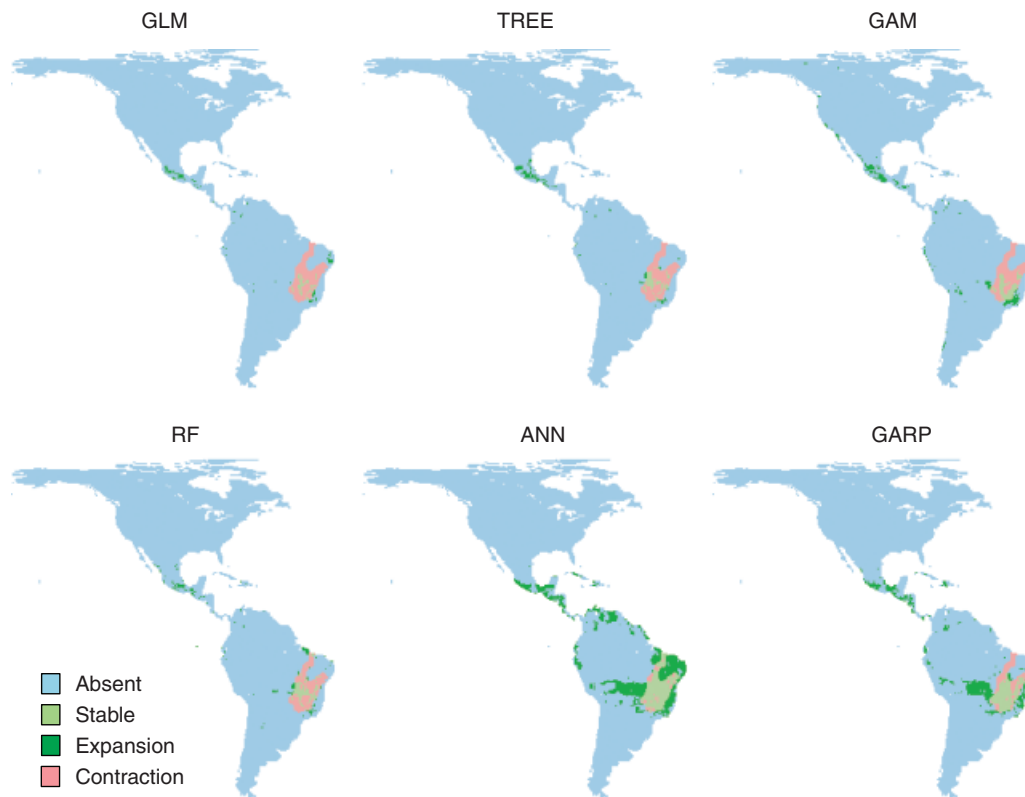


Fig. 2 Maps of the predicted future range of the black tufted-ear marmoset (*Callithrix penicillata*) as predicted by six alternative modeling approaches. See Table 1 for an explanation of model abbreviations.

this study make different assumptions about the relationships between species and their environments (Guisan & Zimmermann, 2000). For example, generalized linear models assume a given response curve that defines the relationship between the probability of presence and various environmental gradients. These models will generally work well for species with relatively simple relationships to environmental gradients. The other five techniques that we tested are more flexible with respect to the complexity of the relationships that they can model. For example, GAMs allow for complex relationships with individual variables to be modeled. They are not, however, as adept at modeling complex interactions between variables as are classification tree models or random forests. Artificial neural networks and GARP models, the two machine-learning-based approaches tested here, are in part an attempt to model both complex relationships with individual variables and complex interactions among those variables.

Inconsistencies in bioclimatic model predictions

The inconsistency among bioclimatic models has led some to suggest innovative methods for addressing

model uncertainty that involve finding consensus among different models and then selecting the model that best represents these commonalities (Thuiller, 2003; Thuiller *et al.*, 2004b). Another approach to reducing uncertainty is to ask whether some models might simply perform better than others, and hence we need not consider all of their predictions. Pursuing that strategy, our study compares essentially the full suite of correlative bioclimatic modeling approaches with a common data set, several metrics of model performance, and alternative assumptions about dispersal. The lessons are clear. First, random forest predictors, which averaged the predictions of hundreds of models, were consistently the best performers, and for the data we examined, performed remarkably well. They achieved error rates of less than 15% for presences and less than 1% for absences. We are aware of only one other study that has compared the performance of random forest predictors to other models for use as climate-envelope models. Prasad *et al.* (2006) found that random forest models and bagging (another tree-based model-averaging approach) consistently produced better predictions than multivariate adaptive regression splines and regression trees for predicting the distribu-

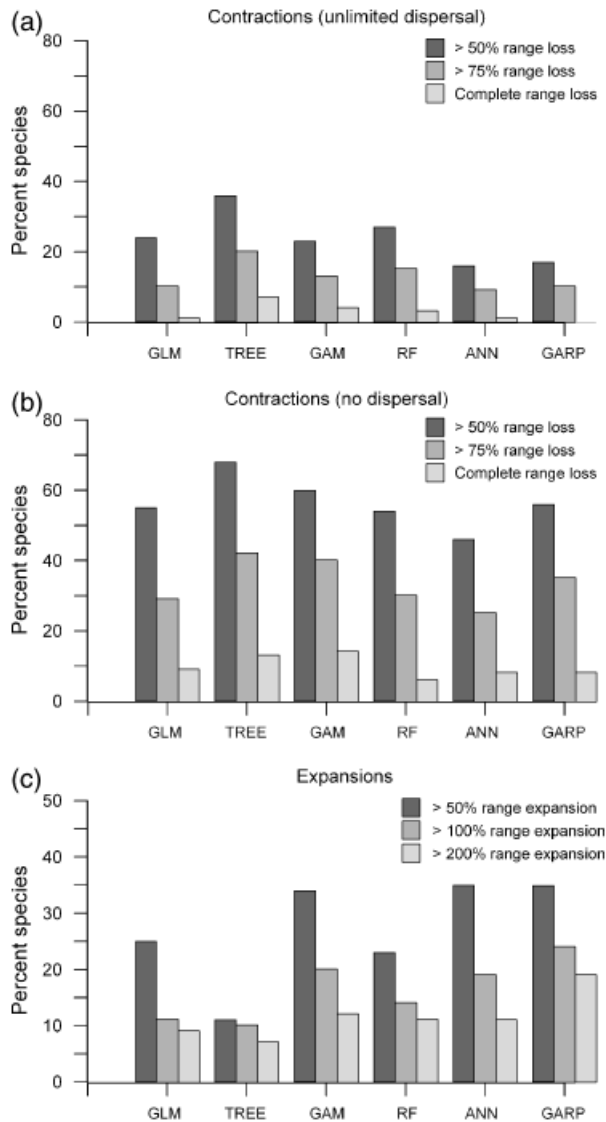


Fig. 3 Climate-induced range contractions and expansions for 100 species as predicted by six different modeling approaches. We report the percentage of species predicted to experience each of three levels of range contraction when (a) individuals are assumed to be able to disperse completely into their future range and (b) when individuals cannot disperse out of their current range. We also report the percentage of species predicted to experience three levels of range expansion (c).

tions of four tree species. The performance of each of the other five modeling approaches tested here, but not by Prasad *et al.*, is generally comparable with the performance of models of the same type tested elsewhere (Thuiller *et al.*, 2003; Pearson *et al.*, 2004; Segurado & Araújo, 2004).

Our results raise the obvious question of why random forest models work so remarkably well. The strength of this approach likely lies in the power derived from

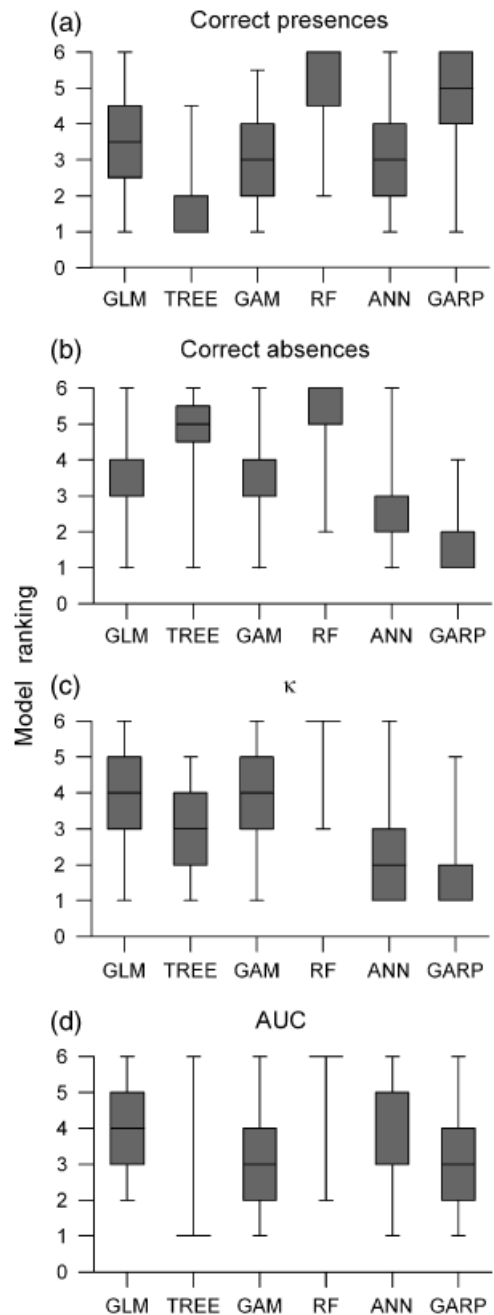


Fig. 4 Ranking of the performance of six different modeling approaches for predicting the current distribution of 100 mammal species. Performance was assessed as (a) the percentage of correctly predicted presences, (b) the percentage of correctly predicted absences, (c) the κ statistic, and (d) the area under the receiver-operating characteristic curve (AUC). Each set of box and whiskers represents the median, first and third quartiles, and the maximum and minimum values. See Table 1 for an explanation of model abbreviations.

averaging hundreds of different models (Breiman, 2001). The individual models are built with randomly selected subsets of the data and randomly selected

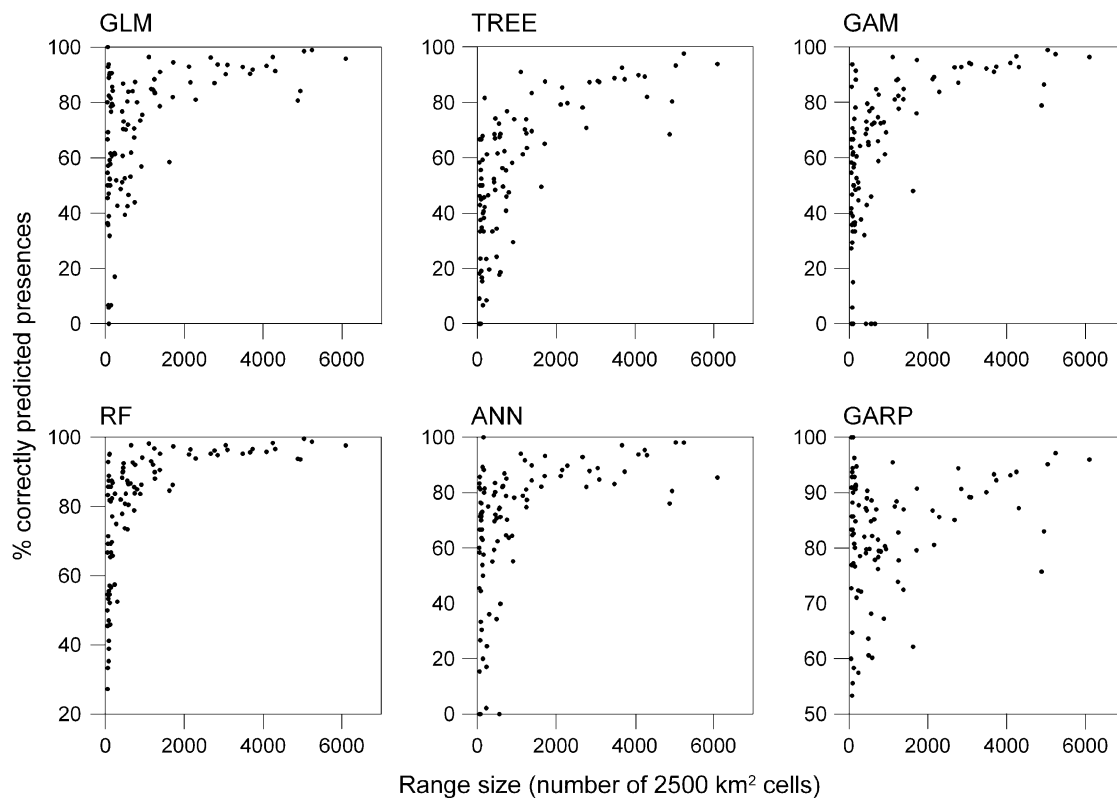


Fig. 5 Relationships between model accuracy (as measured by the percentage of correctly predicted presences) and species range size for 100 mammal species using six different modeling approaches. See Table 1 for an explanation of model abbreviations.

subsets of the predictor variables. Although we averaged 10 artificial neural networks and 10 GARP models to produce predictions for each species, the model averaging accomplished by random forest predictors is much more comprehensive. Although it is possible that model averaging applied similarly to techniques other than the classification trees on which random forests are based would produce models of comparable accuracy, the tree-based models themselves provide added advantages over other modeling approaches. In addition to providing a method for modeling complex interactions without having to specify them *a priori*, tree-based models allow the relationships between the response and the predictors to vary over the domain of the study. This is particularly advantageous for modeling data that cover large and diverse geographic areas.

The second lesson to be taken from our study is that the different modeling approaches tend to be relatively consistent in the types of errors they make. For example, classification trees produced the most omission errors whereas GARP models had the highest commission error rates. These errors, in turn, lead to different predicted range shifts, extinction rates, and changes in species composition at specific sites. The large number

of commission errors produced by the GARP models may, in part, reflect a difference in philosophy inherent in the design and execution of GARP. The model is generally used with presence-only data (e.g. Peterson *et al.*, 2002). Without true absences, it is impossible to fully assess model accuracy; one cannot determine whether predicted presences that do not coincide with the presence data represent commission error or unsampled presences. Indeed, when GARP is applied, many of the predicted presences that do not correspond with presence data points are generally assumed to either represent unrecorded presences or the unrealized portions of a species' fundamental niche (Anderson *et al.*, 2003). While this assumption may be true, our results indicate that it may lead to an overly optimistic view of model performance. In our study, because we used both presence and absence data, we were able to identify commission error and thus fully test the GARP models.

It is important to recognize that no correlative modeling approach can accurately model the fundamental niche of a species. Whether using presence only data or data on presences and absences, the best one can do with a correlative approach is to approximate a species'

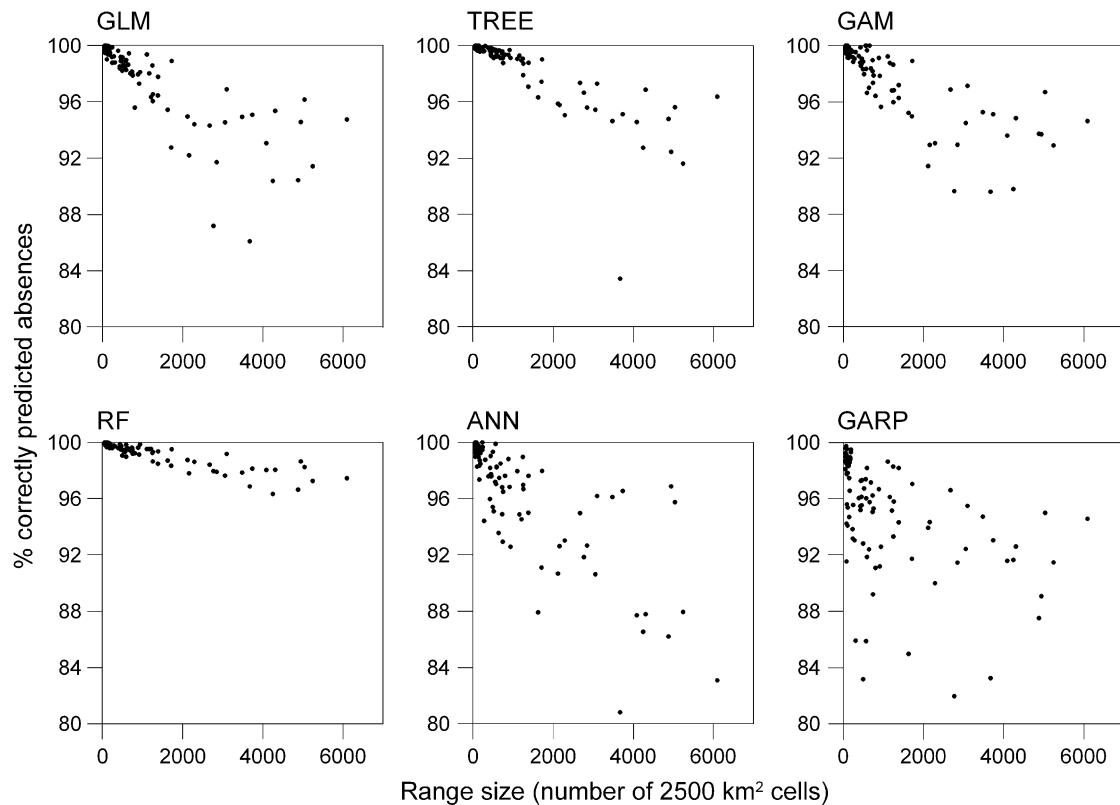


Fig. 6 Relationships between model accuracy (as measured by the percentage of correctly predicted absences) and species range size for 100 mammal species using six different modeling approaches. See Table 1 for an explanation.

Table 2 Median rankings of the accuracy of six different modeling approaches for predicting current absences of 100 mammal species

Median model rankings (for correctly predicted absences)		
Species with ranges of 50–200 cells	Species with ranges of 201–1000 cells	Species with ranges >1000 cells
RF	RF	RF
TREE	TREE	TREE
GLM	GLM	GAM
GAM	GAM	GLM
ANN	ANN	ANN
GARP	GARP	GARP

Species have been divided into three groups based on current range size. The three columns in the table represent model rankings for (from left to right) 39 species with ranges consisting of 50–200 grid cells, 31 species with ranges of 201–1000 grid cells, and 30 species with ranges of >1000 grid cells. The highest ranked models are at the tops of the columns and models tied in rank are linked with a vertical line. See Table 1 for an explanation of model abbreviations.

current realized niche and hope that the modeled relationships hold in the future. Although there is no assurance that the model that most accurately predicts the current distribution of a species will always produce the most accurate future predictions, it is likely that minimizing known errors in the current predictions will reduce the total amount of error in projections of future or past ranges.

In addition to being prone to committing specific types of errors, different modeling approaches may also be more or less sensitive to various attributes of the data used in the modeling process. For example, some modeling approaches may be more robust to changes in spatial resolution (Thuiller *et al.*, 2003) and some may be more robust to the changes in spatial extent (Thuiller *et al.*, 2004c). Some modeling approaches may be more sensitive to the ratio of presences to absences in the data set (Fielding & Haworth, 1995). Finally, some modeling approaches may be more or less sensitive to the type of predictor variables used in the modeling process (Thuiller *et al.*, 2004a). Determining the degree to which these attributes of data sets differentially affect modeling approaches will require a concerted research effort in the future.

Table 3 Median rankings of the accuracy of six different modeling approaches for predicting current presences of 100 mammal species

Median model rankings (for correctly predicted presences)		
Species with ranges of 50–200 cells	Species with ranges of 201–1000 cells	Species with ranges > 1000 cells
GARP	RF	RF
RF	GARP	GLM
GLM	GLM	GAM
ANN	GAM	ANN
GAM	ANN	GARP
TREE	TREE	TREE

Species have been divided into three groups based on current range size. The three columns in the table represent model rankings for (from left to right) 39 species with ranges consisting of 50–200 grid cells, 31 species with ranges of 201–1000 grid cells, and 30 species with ranges of > 1000 grid cells. The highest ranked models are at the tops of the columns and models tied in rank are linked with a vertical line. See Table 1 for an explanation of model abbreviations.

The last lesson we can take from our study is that the models differed greatly in the extent to which they predicted shrinking ranges vs. expanding ranges in the face of climate change. For example, when we assumed unlimited dispersal, classification tree models predicted extinctions for 7% of the species compared with GARP models, which predicted no extinctions. Similarly, Thuiller *et al.* (2004b) demonstrated potential differences in predicted extinction rates across modeling approaches ranging from less than 1% to roughly 5% over a 50-year period.

Limitations and advances in bioclimatic modeling

Although bioclimatic models are a useful tool for investigating the effects of climate change on biodiversity at large spatial scales, they are not without their limitations. Our analyses address one aspect of the uncertainty associated with current bioclimatic models and highlight a tool for reducing this uncertainty. There are, however, several other points at which uncertainty enters the bioclimatic-modeling process. The limitations of bioclimatic models have been thoroughly reviewed by Pearson & Dawson (2003). Here, we discuss four of these limitations.

First, most correlative approaches do not directly model biotic interactions. These interactions can have strong influences on species' responses to climate change (Davis *et al.*, 1998). As a first step to addressing biotic interactions, we included vegetation in our mod-

els as a proxy for animal–habitat interactions. Although simple vegetation associations cannot capture all biotic interactions, they likely represent some of the most basic, resource use, predator–prey, and competitor interactions. Models that assume climate variables will serve as a proxy for vegetation, will fail to capture the effects of changes in atmospheric CO₂ concentrations on animal habitat. Including vegetation in bioclimatic models for animals is only a first step to addressing biotic interactions. Explicitly modeling interspecific interactions will involve linking bioclimatic models for multiple species or further integrating mechanistic and correlative models.

The second limitation of correlative models is that they do not address dispersal. Assuming that organisms can fully disperse into their projected future range or that they will be limited to that portion of their projected future range that overlaps their current range is overly simplistic. One solution is to link bioclimatic model projections with simulated dispersal patterns (e.g. Peterson *et al.*, 2002). Such integrated modeling approaches will provide more accurate predictions of future distributions.

The third limitation of bioclimatic models is that they cannot account for evolutionary change. For species with rapid adaptation rates, evolutionary changes may influence the impacts of climate change on species distributions (Hoffmann & Parsons, 1991; Thomas *et al.*, 2001). However, for many species, evolutionary change will likely lag far behind climate change (Peters & Darling, 1985; Etersson & Shaw, 2001). With respect to evolutionary change, bioclimatic approaches will most accurately model species with poor dispersal capabilities and long generation times (Pearson & Dawson, 2003).

Finally, the fourth limitation of bioclimatic approaches is that the models are exceedingly difficult to validate. Ideally, models are validated with data that are completely independent of the data used to build them. However, many models are evaluated with the same data used in model building (e.g. Huntley *et al.*, 2004). In these cases, there is a complete lack of independence of the data sets, which prevents any assessment of whether or not the models over-fit the data. Another common approach is the one taken in this and many other studies (Iverson & Prasad, 1998; Pearson *et al.*, 2002; Thuiller, 2003) in which data are split into two sets, one of which is used to build the models and the other of which is reserved for model validation. Although this approach provides some independence of the model building and validating data sets, the reserved data are not completely independent because of spatial autocorrelation (Koenig, 1999). To obtain a completely independent data set, one must find data

from a geographically distinct region (Fielding & Haworth, 1995) or from a historical period (Araújo *et al.*, 2005) – although the latter may still be both spatially and temporally autocorrelated.

For continental analyses, truly independent data seldom exist. Many species that occur on multiple continents do so because they are invasive exotics and, thus, may not be at equilibrium with their new environments. Historic data for most species do not exist. When they do, they often provide few data points for model validation. Fortunately, there is evidence that bioclimatic model validation estimates based on semi-independent reserved validation data sets may approximate estimates based on more independent data sets. Araújo *et al.* (2005) found that model performance estimates based on historic bird ranges were similar to performance estimates based on a reserved data set.

Despite these limitations, we should not underestimate the role of bioclimatic models in assessing the potential effects of climate change. Bioclimatic model predictions should be seen as a first approximation of the potential effects of climate change on biota at large spatial scales and not as accurate predictions of future distributions of individual species (Pearson & Dawson, 2003). Although dynamic global vegetation models (DGVMs) currently provide a process-based alternative for projecting climate-induced shifts in vegetation types or biomes (Bachelet *et al.*, 2001; Sitch *et al.*, 2003), building purely mechanistic models for large numbers of individual species would be a massive undertaking because of the lack of knowledge of species' life histories and physiologies and the amount of work each individual model would require. Indeed, mechanistic models come with their own uncertainties as DGVM-comparison studies demonstrate (Cramer *et al.*, 2001; Bachelet *et al.*, 2003). Our results highlight one specific modeling approach that will reduce the uncertainty in bioclimatic-model predictions. Reducing the uncertainty associated with biotic interactions, dispersal, and evolutionary change will involve even more creative approaches that combine mechanistic and correlative models.

Conclusions

The uncertainties in future range predictions that can be attributed to the errors in the bioclimatic models currently in use are likely to be greater than the uncertainties of actually predicting the underlying climate change (i.e. the differences among climate models and emissions scenarios) (Thuiller, 2004). This means that unless we can produce more accurate bioclimatic models, they cannot really be used to compare the consequences of different greenhouse gas emissions scenarios. Looking

forward, it appears that random forest models or other model-averaging approaches may yield robust predictions of range shifts in the face of climate change. It will still be difficult to translate these predictions into expected extinctions and species turn-over rates because actual range shifts will depend on dispersal, evolutionary flexibility, and species interactions. Nonetheless, for the sake of adaptive management and conservation planning, random forest models provide a useful and reliable tool. By minimizing the uncertainty in bioclimatic models, studies of climate-induced range shifts can concentrate on elucidating the effects of the more important uncertainties in climate-change predictions.

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Appendix

Table A1

Table A1 Future predicted range sizes of 100 mammal species in the western hemisphere as a proportion of current range

Scientific name	English name	Current range (km ²)	Predicted future range as a proportion of current range											
			Unlimited dispersal					No dispersal						
			GLM	TREE	GAM	RF	ANN	GARP	GLM	TREE	GAM	RF	ANN	GARP
<i>Akodon albiventer</i>	White-bellied grass mouse	437500	0.51	0.55	1.03	1.58	0.92	1.71	0.34	0.31	0.54	0.68	0.46	0.68
<i>Akodon cursor</i>	Cursor grass mouse	1430000	0.64	0.21	0	0.53	1.06	0.38	0.39	0.17	0	0.29	0.31	0.17
<i>Alouatta caraya</i>	Black howling monkey	3127500	1.08	0.34	0.79	0.31	0.97	0.62	0.41	0.19	0.23	0.17	0.58	0.25
<i>Alouatta pigra</i>	Black howling monkey	275000	5.71	0.45	0.75	4.48	1.51	3.61	0.54	0.05	0.03	0.04	0.06	0.21
<i>Alouatta sara</i>	Bolivian red howling monkey	410000	7.03	0.15	3.85	1.27	10.09	1.07	0	0	0	0.12	0.71	0.37
<i>Amorhynchillus schmablii</i>	Smoky bat	370000	0.88	0.09	0.92	1.01	1.01	2.91	0.37	0.07	0.43	0.36	0.34	0.68
<i>Anoura latidens</i>	Broad-toothed tailless bat	1417500	2.75	3.29	7.13	7.39	0.33	5.35	0.64	0.57	0.81	0.82	0.12	0.63
<i>Aotus vociferans</i>	Tropical night monkey	1157500	1.35	0.05	0.19	0.28	1.67	0.91	0.73	0.01	0	0.08	0.83	0.55
<i>Artibeus fraterculus</i>	Fraternal fruit-eating bat	297500	0.84	0.29	0.87	0.33	1.28	1.81	0.19	0.11	0.29	0.18	0.44	0.3
<i>Auliscomys pictus</i>	Painted big-eared mouse	320000	0.8	0.18	1.16	0.73	0.82	0.96	0.39	0.13	0.38	0.45	0.39	0.45
<i>Blastocercus dichotomus</i>	Marsh deer	2005000	1.56	0.5	1.49	0.77	0.9	1.01	0.39	0.22	0.26	0.21	0.44	0.15
<i>Bradylops torquatus</i>	Maned three-toed sloth	162500	0.14	0.38	0.02	0.14	0.46	0.51	0	0	0	0	0	0.02
<i>Cabreramops aequatorianus</i>	Equatorial dog-faced bat	127500	2.37	0.39	0.16	0.08	1.24	2.53	0.29	0.12	0	0.06	0.14	0.25
<i>Callicebus moloch</i>	Titi monkey	975000	0.02	0.01	3.64	0.03	1.05	0.02	0	0	0	0	0.31	0
<i>Callicebus nigrifrons</i>	Black-fronted titi	472500	0.49	0.22	2.58	0.25	1.15	0.04	0.32	0.04	0.15	0.05	0.28	0
<i>Callicebus personatus</i>	Northern masked titi	175000	0.11	0.11	0.09	0.06	0.37	0.73	0.01	0.03	0	0.01	0	0.01
<i>Callicebus regulus</i>	Titi monkey	230000	0	0.03	0	0.04	0.08	0.01	0	0	0	0.04	0.04	0
<i>Callicebus torquatus</i>	Collared titi	265000	0.17	0.03	0.01	0	8.93	0.03	0.01	0	0	0	0.93	0
<i>Callithrix penicillata</i>	Black tufted-ear marmoset	1602500	0.3	0.42	0.71	0.54	2.8	1.53	0.14	0.2	0.28	0.29	0.85	0.56
<i>Cavia apera</i>	Brazilian guinea pig	5725000	0.57	0.77	0.62	0.53	0.78	1.04	0.4	0.49	0.4	0.43	0.58	0.65
<i>Cavia magna</i>	Greater guinea pig	172500	0.29	0	0.38	0	0.19	0.87	0.12	0	0.07	0	0.06	0.23
<i>Centronycteris centralis</i>	Bat	1242500	1.42	4.55	7.32	6.34	0.31	3.2	0.3	0.41	0.71	0.77	0.16	0.44
<i>Chinchilla lanigera</i>	Chinchilla	205000	1.06	0.32	0.7	0.44	0.6	2.05	0.63	0.26	0.59	0.43	0.46	0.73
<i>Chinchillula sahamae</i>	Altiplano chinchilla mouse	267500	1.08	0.29	1.65	0.59	1.19	1.8	0.55	0.15	0.63	0.36	0.58	0.73
<i>Chiroderma trinitatum</i>	Little big-eyed bat	9347500	1.15	1.07	1.07	1.12	0.99	0.7	0.95	0.86	0.87	0.86	0.87	0.66
<i>Chiropotes albinasus</i>	White-nosed bearded saki	145000	3.16	1.1	0.07	0.48	10.74	1.12	0.78	0.17	0.07	0.28	0.76	0
<i>Chiropotes albinasus</i>	White-nosed bearded saki	1107500	0.31	0.63	0.26	0.32	0.89	0.02	0.14	0.17	0.11	0.06	0.24	0
<i>Chiropterus auritus</i>	Big-eared woolly bat	9182500	1.33	1.49	1.63	1.47	1.71	0.97	0.88	0.92	0.98	0.94	0.98	0.6
<i>Coendou bicolor</i>	Bicolor-spined porcupine	1237500	0.42	0.73	0.67	3.95	1.72	1.02	0.13	0.22	0.22	0.67	0.58	0.21
<i>Cryptotis mayensis</i>	Maya small-eared shrew	147500	0.78	0.78	1.34	0.29	10.59	0.17	0	0	0	0	0.66	0
<i>Ctenomys torquatus</i>	Collared Tucú-tucú	475000	1.1	0.64	0.26	0.59	0.86	0.74	0.6	0.39	0.06	0.39	0.24	0.28
<i>Cyclopes didactylus</i>	Silky anteater	8700000	1.39	1.41	1.32	1.45	1.17	0.9	0.98	0.97	0.98	0.98	0.89	0.82
<i>Cynomys parvus</i>	Dog-faced bat	1855000	1.53	3.81	3.23	5.58	3.42	3.68	0.57	0.73	0.88	0.92	0.89	0.64
<i>Dasyprocta azarae</i>	Azara's Agouti	1637500	0.39	0.5	0	0.64	0.85	0.12	0.29	0.26	0	0.34	0.47	0.02
<i>Dasyprocta sabanicola</i>	Llanos long-nosed armadillo	695000	3.25	1.18	0.4	1.4	7.44	3.37	0.46	0.23	0.05	0.29	0.76	0.39
<i>Delomys dorsalis</i>	Striped Atlantic forest rat	142500	0.18	0.14	0.33	0.05	0.05	0.26	0.12	0.07	0.09	0.05	0	0.02

Table A1. (Contd.)

Scientific name	English name	Current range (km ²)	Predicted future range as a proportion of current range											
			Unlimited dispersal						No dispersal					
			GLM	TREE	GAM	RF	ANN	GARP	GLM	TREE	GAM	RF	ANN	GARP
<i>Delomys sublineatus</i>	Pallid Atlantic forest rat	212500	0.27	0	1.72	0.18	1.05	0.39	0.08	0	0.08	0.02	0.26	0.01
<i>Eptesicus diminutus</i>	Diminutive Serotine	3152500	1.04	0.45	0.83	0.79	0.56	0.63	0.51	0.34	0.49	0.54	0.4	0.32
<i>Eptesicus fuscus</i>	Big brown bat	12362500	1.32	1.13	1.55	1.5	1.18	1.34	0.9	0.83	0.92	0.94	0.86	0.87
<i>Heteromys anomalus</i>	Trinidad spiny pocket mouse	592500	4.04	15.59	15.08	17.14	2.83	7.92	0.49	0.68	0.78	0.77	0.38	0.46
<i>Histiotes macrotus</i>	Big-eared brown bat	2215000	0.97	0.8	1.07	1.07	0.57	1.07	0.55	0.54	0.61	0.75	0.42	0.58
<i>Hydrochaeris hydrochaeris</i>	Capybara	13105000	1.12	1.22	1.12	1.05	1.24	1.06	0.94	0.99	0.94	0.94	0.98	0.93
<i>Leopardus pardalis</i>	Ocelot	15232500	1.12	1.11	1.17	1.12	1.31	0.98	0.99	0.98	0.99	0.99	0.92	0.94
<i>Lepus alleni</i>	Antelope jackrabbit	250000	3.2	5.75	1.03	3.79	1.47	3.59	0.85	0.55	0.05	0.6	0.54	0.72
<i>Lepus townsendii</i>	White-tailed jackrabbit	3465000	1.52	0.6	1.72	1.14	0.94	1.24	0.73	0.54	0.71	0.72	0.59	0.89
<i>Lichonycteris obscura</i>	Dark long-tongued bat	7737500	1.17	1.26	1.36	1.36	1.06	0.86	0.97	0.94	0.97	0.96	0.89	0.78
<i>Lonchophylla hesperia</i>	Western nectar bat	227500	0.18	0	2.93	0.02	0.16	3.49	0	0	0.09	0.02	0.03	0.21
<i>Lonchorhina orinocensis</i>	Orinoco sword-nosed bat	262500	9.93	0.98	17.76	0.62	1.97	1.79	0.43	0.28	0.56	0.07	0.43	0.38
<i>Lyncodon patagonicus</i>	Patagonian weasel	1070000	0.93	0.56	0.94	0.72	1.77	1.69	0.44	0.41	0.5	0.56	0.64	0.32
<i>Marmosa robinsoni</i>	Robinson's mouse opossum	765000	6.85	2.82	12.44	11.6	4.67	10.93	0.75	0.43	0.79	0.8	0.58	0.64
<i>Marmosops noctivagus</i>	White-bellied slender mouse opossum	1825000	0.47	0.61	0.2	0.75	1.79	0.26	0.3	0.37	0.1	0.43	0.64	0.13
<i>Marmosops parvidens</i>	Delicate slender mouse opossum	2282500	1.17	0.69	1.33	0.98	1.83	0.1	0.23	0.26	0.35	0.25	0.63	0.02
<i>Martes pennanti</i>	Fisher	3455000	1.43	1.27	1.51	1.25	0.91	0.83	0.82	0.75	0.83	0.71	0.51	0.47
<i>Megascorex gigas</i>	Mexican shrew	155000	0.6	0	3.82	2.31	1.42	6.58	0.1	0	0.02	0.34	0.42	0.16
<i>Mephitis macroura</i>	Hooded skunk	1832500	1.18	0.53	0.81	1.19	2.21	1.82	0.62	0.33	0.46	0.64	0.74	0.67
<i>Micromycteris hirsuta</i>	Hairy big-eared bat	4282500	1.84	2.03	2.2	1.91	1.69	0.79	0.95	0.87	0.95	0.88	0.92	0.46
<i>Microtus oeconomicus</i>	Tundra Vole	2352500	0.81	1.1	1.52	1.18	1.52	1.77	0.44	0.46	0.42	0.55	0.47	0.66
<i>Microtus pinetorum</i>	Woodland vole	2767500	1.27	1.18	1.42	1.23	1.49	0.89	0.93	0.92	0.97	0.92	0.95	0.83
<i>Mustela frenata</i>	Long-tailed weasel	12200000	1.11	0.8	1.09	2.03	1.09	1.06	0.8	0.65	0.76	0.94	0.69	0.76
<i>Mioprocta acouchy</i>	Red acouchy	1300000	1.91	1.08	1.67	1.27	4.29	0.57	0.77	0.37	0.28	0.25	0.93	0.02
<i>Myotis evotis</i>	Long-eared myotis	3115000	1.12	0.79	1.04	1.01	0.8	1.32	0.75	0.69	0.75	0.85	0.68	0.77
<i>Myotis levis</i>	Yellowish myotis	2900000	0.86	0.52	1.25	0.72	0.99	1.06	0.39	0.38	0.41	0.55	0.5	0.59
<i>Myotis ruber</i>	Red myotis	1725000	0.61	0.33	0.59	0.4	0.36	0.1	0.46	0.29	0.45	0.31	0.23	0.04
<i>Nasua olivacea</i>	Mountain coati	210000	0.74	32.48	21.83	35.04	0	17.68	0.25	0.3	0.43	0.29	0	0.36
<i>Necomys lasiurus</i>	Hairy-tailed bolo mouse	5395000	0.87	1.28	0.89	0.61	1.62	1.25	0.56	0.76	0.52	0.4	0.86	0.67
<i>Neotoma goldmani</i>	Goldman's woodrat	175000	0.3	0.64	0	0.67	5.07	0.61	0.17	0.23	0	0.43	0.83	0.14
<i>Nyctinomys femorosaccus</i>	Pocketed free-tailed bat	1132500	1.7	0.6	1.97	1.29	2.21	3.11	0.84	0.41	0.7	0.62	0.78	0.89
<i>Oecomys speciosus</i>	Arboreal rice rat	185000	0.35	1.43	0.38	0.46	10.62	6.86	0.12	0.05	0.11	0.04	0.85	0.15
<i>Oryzomys angouya</i>	Rice rat	1142500	1.62	0.28	3.17	0.96	1.09	0.88	0.38	0.2	0.56	0.38	0.21	0.23
<i>Oryzomys rostratus</i>	Long-nosed rice rat	370000	3.26	0.83	0.89	2.36	3.05	3.06	0.62	0.11	0.03	0.3	0.08	0.34
<i>Oryzomys yunganus</i>	Yungas rice rat	5302500	1.56	0.96	1.61	0.87	1.78	0.51	0.9	0.67	0.9	0.54	0.96	0.37
<i>Oxymycteris rufus</i>	Red Hoctiudo	1395000	0.96	0.3	0.04	0.43	0.62	0.44	0.28	0.23	0.01	0.34	0.3	0.24
<i>Peromyscus leviceps</i>	Nimble-footed mouse	582500	1.28	0.53	3.73	6.76	0.01	4.36	0.5	0.26	0.28	0.7	0	0.42

Table A1. (Contd.)

Scientific name	English name	Current range (km ²)	Predicted future range as a proportion of current range											
			Unlimited dispersal						No dispersal					
			GLM	TREE	GAM	RF	ANN	GARP	GLM	TREE	GAM	RF	ANN	GARP
<i>Peromyscus polionotus</i>	Oldfield mouse	425000	0.33	0.39	0.25	0.49	3.23	0.05	0.21	0.12	0.08	0.18	0.79	0.01
<i>Platyrrhinus helleri</i>	Heller's broad-nosed bat	10765000	1.06	0.99	0.96	1.23	1.26	0.83	0.93	0.84	0.84	0.97	0.92	0.77
<i>Proechimys cajannensis</i>	Cayenne spiny rat	3030000	0.7	1.39	0.44	1.58	2.15	0.88	0.24	0.68	0.15	0.78	0.9	0.18
<i>Proechimys oris</i>	Para spiny rat	225000	0.12	0	0.04	0	1.54	0.5	0	0	0	0	0	0
<i>Proechimys roberti</i>	Spiny rat	1077500	0.3	0.82	2.79	0.86	1.13	1.03	0.17	0.5	0	0.5	0.56	0.02
<i>Pteronotus dacyi</i>	Davya's naked-backed bat	4062500	1.3	2.42	2.82	3.32	2.57	1.58	0.61	0.73	0.83	0.94	0.86	0.57
<i>Rhogeessa tumida</i>	Black-winged little yellow bat	7637500	1.48	1.38	1.58	1.76	1.48	1.1	0.97	0.94	0.96	0.99	0.94	0.85
<i>Saccopteryx gymnura</i>	Amazonian sac-winged bat	615000	0.09	0.7	0.38	0.96	1.58	1.26	0	0.11	0	0.18	0.23	0.11
<i>Scapanus latimanus</i>	Broad-footed mole	337500	1.39	0.67	2.78	1.1	1.01	2.93	0.46	0.44	0.49	0.61	0.4	0.96
<i>Sciurus aestivus</i>	Guianan squirrel	6922500	0.5	0.62	0.82	0.69	1.29	1.01	0.32	0.39	0.4	0.45	0.81	0.44
<i>Sciurus niger</i>	Eastern fox squirrel	4307500	1.56	1.15	1.45	1.52	1.52	0.91	0.99	0.86	0.96	0.96	0.96	0.75
<i>Scotinomys teguina</i>	Alstona's brown mouse	207500	0.33	—	0.43	0.24	0.19	3.87	0.01	—	0.01	0.04	0	0.17
<i>Signodon arizonae</i>	Arizona cotton rat	347500	1.68	1.89	1.78	1.37	2.4	1.76	0.73	0.49	0.36	0.58	0.57	0.73
<i>Sorex dispar</i>	Long-tailed shrew	327500	1.09	0.89	2.45	1.56	0.61	0.4	0.41	0.29	0.59	0.5	0.1	0.14
<i>Sorex trowbridgii</i>	Trowbridge's shrew	402500	1.1	0.4	0.63	0.84	2.45	1.8	0.58	0.27	0.42	0.55	0.88	0.75
<i>Speothos venaticus</i>	Bush dog	10617500	1.05	1.11	1.19	1.23	1.32	0.78	0.83	0.89	0.91	0.93	0.94	0.67
<i>Spermophilus elegans</i>	Wyoming ground squirrel	272500	0.6	0.23	0.66	0.17	0.39	3.41	0.29	0.1	0.18	0.06	0.15	0.69
<i>Sturnira magna</i>	Greater yellow-shouldered bat	1470000	0.27	0.53	0.73	1.04	0.6	0.84	0.08	0.22	0.35	0.49	0.34	0.32
<i>Sturnira nana</i>	Lesser yellow-shouldered bat	137500	1.51	0.27	0.73	0.75	1.25	1.36	0.38	0.04	0.09	0.13	0.22	0.16
<i>Tamias rufus</i>	Hopi chipmunk	140000	0.09	0	1.43	0.02	0.2	6.09	0	0	0.07	0	0	0.54
<i>Tamias umbrinus</i>	Uinta chipmunk	287500	1.57	0.1	2.09	0.17	0.1	4.63	0.23	0.05	0.23	0.05	0.02	0.61
<i>Tamiasciurus hudsonicus</i>	Red squirrel	10217500	1.01	1.05	1.02	1.06	1.2	1.19	0.85	0.89	0.87	0.89	0.9	0.94
<i>Thomomys bottae</i>	Botta's pocket gopher	1885000	1.81	1.11	1.75	1.66	1.56	2.13	0.84	0.64	0.87	0.84	0.67	0.85
<i>Tonatia saurophila</i>	Stripe-headed round-eared bat	7115000	1.68	1.46	1.65	1.52	1.35	1.18	0.98	0.93	0.98	0.96	0.89	0.87
<i>Uroderma bilobatum</i>	Tent-making bat	12600000	1.02	1.12	0.99	1.14	1.08	0.88	0.95	0.96	0.94	0.99	0.98	0.83
<i>Vampyressa bidens</i>	Bidentate yellow-eared bat	6682500	1.27	0.99	0.64	0.77	1.01	0.51	0.97	0.81	0.57	0.5	0.84	0.46
<i>Zaedyus pichiy</i>	Pichi	14775000	0.76	0.61	0.78	0.64	0.9	1.09	0.52	0.43	0.56	0.56	0.53	0.6

Future ranges were predicted using six different modeling approaches given two different dispersal scenarios. The models included generalized linear models (GLM), classification trees (TREE), generalized additive models (GAM), random forest predictors (RF), artificial neural networks (ANN), and genetic algorithms for rule-set prediction (GARP). The dispersal scenarios assumed that individuals could disperse completely into the predicted new range (unlimited dispersal) or conversely, that they were restricted to areas in which the current and future predicted ranges overlapped (no dispersal).