



Species distribution modelling for plant communities: stacked single species or multivariate modelling approaches?

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Keywords

Nearest-neighbor imputation; Plant community composition; Random forest; Species distribution modelling; Vegetation mapping; Western Oregon

Nomenclature

USDA NRCS (2000)

Received 21 March 2013

Accepted 14 November 2013

Co-ordinating Editor: Sarah Goslee

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Abstract

Aim: Landscape management and conservation planning require maps of vegetation composition and structure over large regions. Species distribution models (SDMs) are often used for individual species, but projects mapping multiple species are rarer. We compare maps of plant community composition assembled by stacking results from many SDMs with multivariate maps constructed using nearest-neighbor imputation.

Location: Western Cascades ecoregion, Oregon and California, USA.

Methods: We mapped distributions and abundances of 28 tree species over 4,007,110 ha at 30-m resolution using three approaches: SDMs using machine learning (random forest) to yield: (1) binary (RF_Bin); (2) basal area (abundance; RF_Abund) predictions; and (3) multi-species basal area predictions using a nearest-neighbor imputation variant based on random forest (RF_NN). We evaluated accuracy of binary predictions for all models, compared area mapped with plot-based areal estimates, assessed species abundance at two spatial scales and evaluated communities for species richness, problematic compositional errors and overall community composition.

Results: RF_Bin yielded the strongest binary predictions (median True Skill Statistics; RF_Bin: 0.57, RF_NN: 0.38, RF_Abund: 0.27). Plot-scale predictions of abundance were poor for RF_Abund and RF_NN (median Agreement Coefficient (AC): -1.77 and -2.28), but strong when summarized over 50-km radius tessellated hexagons (median AC for both: 0.79). RF_Abund's strength with abundance and weakness with binary predictions stems from predicting small values instead of zeros. The number of zero value predictions from RF_NN was closest to counts of zeros in the plot data. Correspondingly, RF_NN's map-based species area estimates closely matched plot-based area estimates. RF_NN also performed best for community-level accuracy metrics.

Conclusions: RF_NN was the best technique for building a broad-scale map of diversity and composition because the modelling framework maintained inter-species relationships from the input plot data. Re-assembling communities from single variable maps often yielded unrealistic communities. Although RF_NN rarely excelled at single species predictions of presence or abundance, it was often adequate to many (but not all) applications in both dimensions. We discuss our results in the context of map utility for applications in the fields of ecology, conservation and natural resource management planning. We highlight how RF_NN is well-suited for mapping current but not future vegetation.

Introduction

Maps of current vegetation are an essential component of landscape management and conservation planning. Vegetation maps can be used to inform conservation strategies (US Geological Survey 2011), understand the distribution of invasive forest pathogens (Václavík et al. 2010) and initialize modelling efforts that explore how the future might look in the context of human management and climate change (e.g. Scheller & Mladenoff 2004; Hemstrom et al. 2007).

Species distribution modelling (SDM) techniques have been widely used for mapping geographic ranges of common trees (e.g. Schroeder et al. 2010), rare species (e.g. Engler et al. 2004) and species richness (e.g. Guisan & Rahbek 2011). Projects mapping multiple species are less common (but see Elith & Leathwick 2007; Baselga & Araújo 2009; Ohmann et al. 2011; Wilson et al. 2012). There is a strong focus in the SDM literature on mapping species presences, perhaps a legacy of the abundant studies estimating rare species' habitat from presence-only data sets (Newbold 2010). However, many applications require species-specific information on presences and abundances as well as community composition and diversity.

Stacking maps from individual species distribution models often yields problematic community-level results because errors in each model are combined (Dubuis et al. 2011; Guisan & Rahbek 2011). These errors in single species models result from a variety of factors. Species' distributions may not be in equilibrium with current climate conditions (Elith et al. 2010). Presence (or abundance) may be constrained by land-use and disturbance history (e.g. Motzkin et al. 1996) as well as interspecific interactions (e.g. Ettinger et al. 2011). SDMs often overestimate species ranges because they tend to illustrate potential rather than realized niches (Jiménez-Valverde et al. 2008), perhaps because they are often unconstrained by some of the above-mentioned factors due to lack of available data. The net effect of overestimating species presences is that stacked models yield inflated estimates of species richness (Dubuis et al. 2011; Guisan & Rahbek 2011; Pottier et al. 2012), and predicted communities may not reflect those that currently exist in nature (Baselga & Araújo 2010). Inaccurate compositional representation renders maps less fit for some uses, such as designing reserves that adequately represent biodiversity (Margules & Pressey 2000). Community composition and species richness estimates can be improved with additional layers of analysis and information (e.g. Clark et al. 2011; Guisan & Rahbek 2011). However, these extended analyses will not be easy to accomplish across broad areas, with major constraints to time, data and existing knowledge and expertise.

We use the random forest machine learning algorithm, a SDM technique that can yield strong results for mapping individual species (Evans & Cushman 2009) and that has grown in popularity in recent years (Cutler et al. 2007). The random forest algorithm performs well for species distribution modelling for several reasons: (1) it is non-parametric, and hence flexible in terms of the explanatory variables that it can handle; (2) it can represent non-linear relationships between response and explanatory variables and also hierarchical interactions of explanatory variables; and (3) it uses information on species presence and absence, a useful trait when complete data are available.

Of particular importance to our work here, the random forest algorithm has been extended in utility to inform the distance matrix used in nearest-neighbor imputation (Crookston & Finley 2008), which can yield multivariate predictions. Hence it is a useful technique for comparing single species and multivariate approaches to mapping communities.

Nearest-neighbor imputation techniques have been rising in popularity within the forestry community (Eskelson et al. 2009), as forest management planning activities often require multivariate maps describing forest structure and composition. Imputation is defined as filling in missing values within a data set with known values from that same data set. In our application, the 'missing values' are pixels within a raster data set, and the known values come from vegetation survey plot data. In our implementation, each prediction is a link to a single plot. Therefore, model predictions are constrained to communities represented in the input plot data. This means that the net effects of species interactions and site history on community structure are preserved from the original plot data. Imputation mapping can be viewed as an extension to Ferrier & Guisan's (2006) third approach for community-level mapping: rather than 'assemble and predict together', it could be called 'no assembly necessary, simply predict together'.

In this paper, we explore the trade-offs and consequences that are inherent in two approaches: mapping plant communities as cohesive units or stacking single species models. We compare results built through single species random forest models (stacked models of presence and abundance, hereafter referred to as RF_Bin and RF_Abund) with those generated by random forest-based nearest-neighbor imputation (community-level mapping, hereafter referred to as RF_NN). We test the hypothesis that nearest-neighbor imputation mapping can yield solid predictions for many dimensions of plant community composition, even though single species models may out-perform imputation in a single dimension at a time.

Methods

Study area

We built maps of forest composition across the Oregon Western Cascades ecoregion (Fig. 1). The forested area encompasses 4,007,110 ha and stretches from the Washington state border at its northern end into northern California at its southern end. The vegetation of the region varies along three primary gradients: latitude, elevation and climate. Regional climate ranges from maritime in the west to continental in the east (low seasonality to high seasonality) and interacts with the elevation gradient (colder temperatures and more snow at high elevations). Elevations modelled range from near sea level to upper tree line (ca. 1500 m). The latitudinal gradient is biogeographic, with elements of Alaskan flora in the north (e.g. *Callitropsis nootkatensis*) and species that reach their peak in California (e.g. *Pinus lambertiana*) in the south.

Data

We used 1948 US Forest Service Forest Inventory and Analysis (FIA) annual plots, located within 10 km of the study region (including plots within this 10 km buffer that decrease edge effects). We summarized the basic FIA data across whole plots, generating a matrix of basal area ($\text{m}^2 \cdot \text{ha}^{-1}$) by species and plot. These survey plots contain

information on presence and absence as well as abundance.

Our mapped explanatory variables were rasters (30-m ground pixel resolution) encompassing five thematic areas: (1) spectral reflectance (tasseled cap transformation of Landsat imagery, brightness, greenness and wetness: Crist & Ciccone 1984); (2) climate (PRISM: Daly et al. 2008; 11 variables); (3) topography (elevation from the National Elevation Dataset, and derivatives: Gesch et al. 2009; nine variables); (4) soil parent material (mosaic of SSURGO: Soil Survey Staff 2006 and the US Forest Service Soil Resources Inventory; nine variables); and (5) location (latitude and longitude). Details on each variable are available in the online appendix (Appendix S1).

For mapping and modelling exercises where the most accurate mapping of a single region is the primary goal, attention to variable selection would be merited. However, in our experience, model accuracy changes subtly with variable reductions as long as the five thematic areas mentioned above are well represented. Also, random forest is relatively robust to colinearity in explanatory variables. Because of these two factors, and because model comparison was our primary purpose, we included all variables in all models.

Modelling approach

We built maps of 28 tree species using three approaches, all based on the random forest technique: (1) binary prediction of presence/absence for each species independently (28 total models, one for each species, approach referred to as: RF_Bin); (2) continuous prediction of basal area ($\text{m}^2 \cdot \text{ha}^{-1}$) for each species independently (28 total models, approach referred to as: RF_Abund); and (3) continuous prediction of basal area for all species simultaneously using a random forest-based imputation model (one multivariate model: RF_NN). Basal area predictions from RF_NN and RF_Abund were transformed to binary for comparison with RF_Bin.

The random forest model builds on the functionality of single classification trees (or regression trees for continuous predictions) by extracting a single prediction from an ensemble of tree models (we used 1000). Each individual classification tree within a random forest is built from a random subset of observations and explanatory variables (Breiman 2001). We built RF_Bin and RF_Abund models within the R environment for statistical computing (v 3.0.1; R Foundation for Statistical Computing, Vienna, Austria), using the R-package 'randomForest' (Liaw & Wiener 2002). For our binary model, predictions range from 0 to 1 and reflect the proportion of classification trees within the random forest predicting a given species to be present rather than absent. We translate this continuous

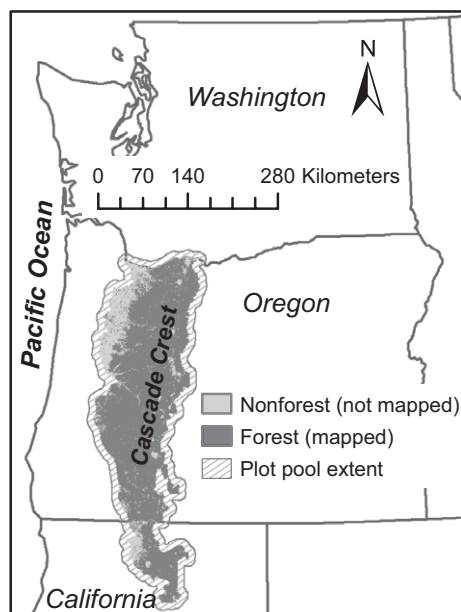


Fig. 1. Study area includes the forests of the Western Cascades ecoregion, stretching from the northern Oregon border into northern California. Plots used for modelling are drawn from the forested area within the boundaries of a 10-km buffer around the ecoregion.

output to binary by applying a cut-off threshold. This threshold was identified by the precision-recall F-measure (Parviainen et al. 2008) using the R-package 'rocr' (Sing et al. 2005) with an alpha value of 0.5 to balance the weight of false positives and negatives. Predicted values from our RF_Abund models were the average basal area predicted by the regression trees within the random forest.

We also built RF_NN maps using the R-package 'yaImpute' (Crookston & Finley 2008). The method implemented in this R-package amalgamates multiple random forest models, each tuned to a single response variable that is a summary of species compositional data (we used three: dominant species, basal area of the dominant species, total basal area). To generate predictions, RF_NN chooses neighbor plots based on a non-Euclidean distance measure built from the nodes matrix of the amalgamated random forest models. This nodes matrix holds a plot identifier for each terminus (or 'leaf') of each classification tree in the random forest models. For new locations (map pixels), the terminal nodes where the pixel falls in the random forest models are recorded. The nearest-neighbor plot for the new pixel is the most frequent plot within its set of nodes.

Mapping and accuracy assessment

Each model prediction was mapped with our in-house R-package 'SDMap' (Henderson unpubl; available upon request to first author). We calculated all accuracy assessment statistics on cross-validated predictions (ten-fold).

For each species and modelling approach, we calculated three binary accuracy assessment measures: sensitivity, specificity and the true skill statistic (TSS; Fielding & Bell 1997). We defined binary model success, for each metric, as a value of 0.3. We also assessed the area occupied by each species in the projected map surfaces. We estimated actual areas of species distributions from FIA annual plots, which are a systematic sample of the landscape. We calculated 95% confidence intervals for those area estimates based on a binomial distribution ('binom.confint' function in R-package 'binom').

We assessed the accuracy of continuous predictions through the protocol outlined in Riemann et al. (2010). The first half of that protocol uses three metrics of agreement (described in Ji & Gallo 2006): (1) an overall agreement coefficient (AC); (2) a measure of systematic agreement (AC.sys); and (3) a measure of unsystematic agreement (AC.uns). For each of these metrics, values approaching or less than zero indicate no agreement while values approaching one indicate strong agreement between observations and predictions. We also plotted empirical cumulative distribution functions (ECDF) for observations and predictions, and calculated the Kolmogorov–Smirnov statistic: the maximum distance between two

ECDF curves (K-S; Massey 1951). All of the continuous accuracy metrics were calculated at two scales: (1) the plot scale and (2) average values for plots falling within tessellated hexagons across the study area (centers spaced 50 km apart: 9128 ha, each containing 44 FIA annual plots on average).

We assessed several measures of accuracy related to community composition. For RF_Bin and RF_Abund, we developed community matrices by combining predictions from each individual species model into a single matrix, with rows for plots and columns for species. Post-modelling aggregation was unnecessary for RF_NN since predictions were generated for all species simultaneously. We compared observed and predicted species richness at the plot locations with a generalized linear model (Poisson family, with a log link function). We determined the prevalence of problematic types of compositional accuracy, calculating how frequently species that rarely co-occur within our plot sample co-occur within our predicted species matrix. We also calculated compositional distance between observed and predicted communities at each plot: Sørensen distance on binary matrices and Bray–Curtis distance on abundance matrices using the 'vegdist' function in the R-package 'vegan'. We illustrate distributions of these distances with ECDF plots.

Results

Single species predictions – binary

The RF_Bin models were strongest in differentiating species presence and absence, combining strong sensitivity with outstanding specificity to yield generally strong TSS statistics and an 86% success rate (Fig. 2a–c). RF_Abund often yielded predictions with high sensitivity, low specificity and poor TSS and a success rate of just 43% (Fig. 2b–d). The RF_NN model showed moderate sensitivity, high specificity and moderate TSS (Fig. 2d–f) as well as an intermediate success rate (64%).

These differences in sensitivity and specificity were expressed in the maps. Models with high sensitivity and low specificity (most of the RF_Abund models) drastically over-mapped species presence, while mapped estimates of species areas from RF_NN aligned well with plot-based estimates of area (Fig. 3). Because TSSs were generally reasonable for RF_NN (Fig. 2), we concluded that this area was mapped to reasonable locations as well as having the correct spatial extent.

Single species predictions – abundance

We found significant errors in abundance predictions at the plot scale for RF_Abund and RF_NN (Fig. 4a,d). At the plot level, unsystematic agreement was generally low for

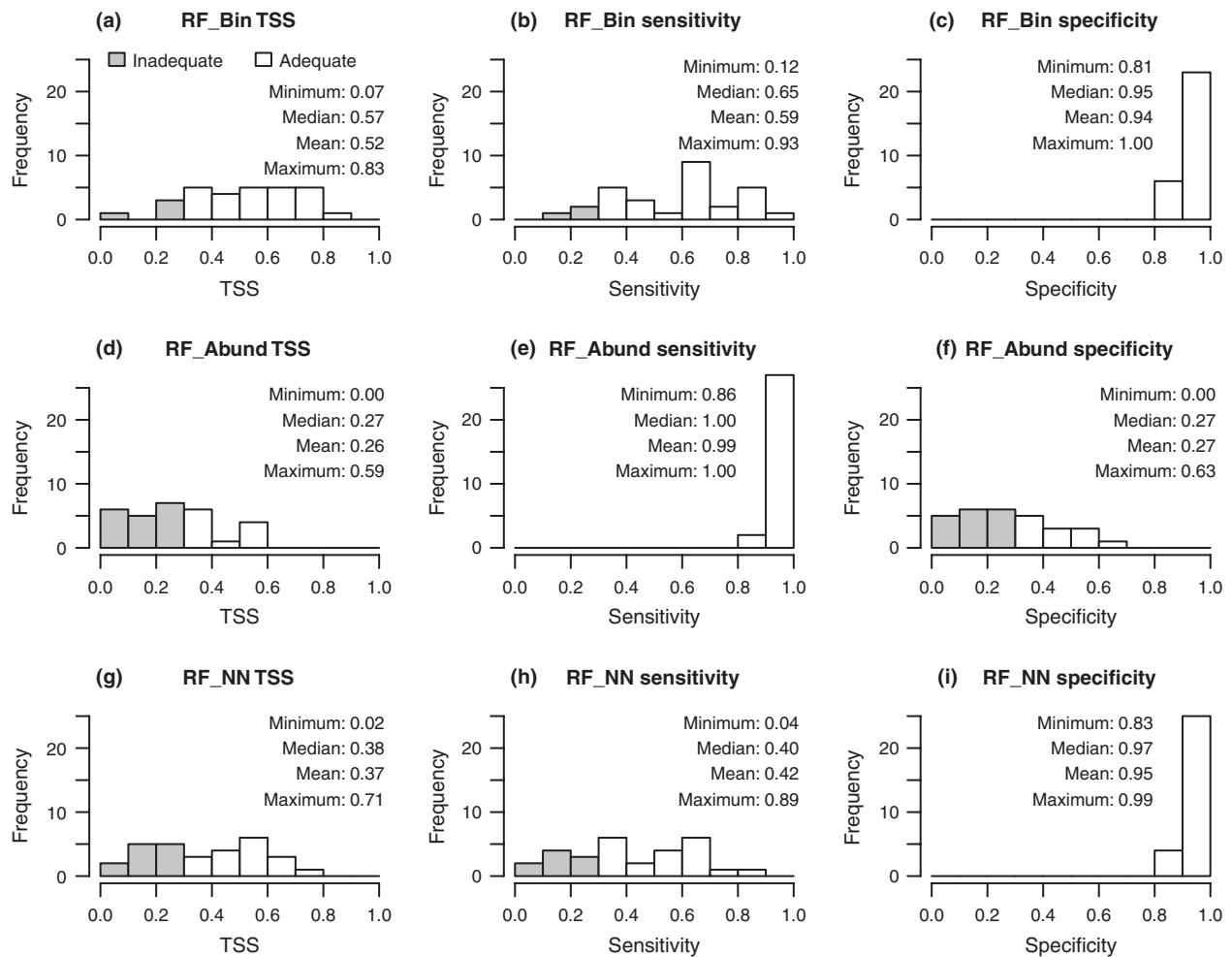


Fig. 2. Panels show histograms of accuracy statistics for single species binary predictions for all 28 tree species (TSS: True Skill Statistic, sensitivity and specificity; Fielding & Bell 1997). The numbers shown within each panel are summary statistics for the set of values illustrated by the histogram. Our three model types (two species-by-species and one multivariate model) are: (1) RF_Bin, binary random forest; RF_Abund, random forest abundance; and RF_NN, random forest, nearest-neighbor imputation. Legend in panel (a) applies to all panels.

both methods, although slightly higher for RF_Abund (Fig. 4b,e). Systematic agreement was often low for RF_Abund, but often moderate for RF_NN (Fig. 4c,f). This pattern suggests that RF_NN errors were mostly random rather than systematic while RF_Abund errors stemmed from both sources. Results were stronger for both approaches at the 50-km hexagon scale of summary (Fig. 4g–i). RF_NN showed a small failure rate for AC and AC.uns (Fig. 4j,k). At this scale, mean values for all AC statistics were lower for RF_NN, but median values were nearly identical for both approaches.

The RF_Abund models consistently under-predicted absences, especially at the plot scale (Fig. 5). Of the non-zero predictions for zero-value plots, RF_Abund predicted values of $<5 \text{ m}^2 \cdot \text{ha}^{-1}$ for the relevant species 59% of the time. Although we only show results for *Pseudotsuga menzi-*

esii and *Tsuga mertensiana*, the pattern described above was pronounced and consistent for RF_Abund for every species at the plot level. It was less pronounced but still consistent over the 50-km hexagons. High K-S values for RF_Abund also reflect this trend (Table 1). In contrast, ECDFs for RF_NN predictions consistently matched those for the observations, and K-S statistics were correspondingly low (Fig. 5, Table 1). RF_NN predicted absences more reliably, but non-zero predictions for zero-value plots were $>5 \text{ m}^2 \cdot \text{ha}^{-1}$ 90% of the time (i.e. the prevalence of species absence errors was low but the magnitude was typically high).

Community composition

Species richness was best predicted by RF_NN (Table 2). RF_Abund drastically over-predicted species richness for

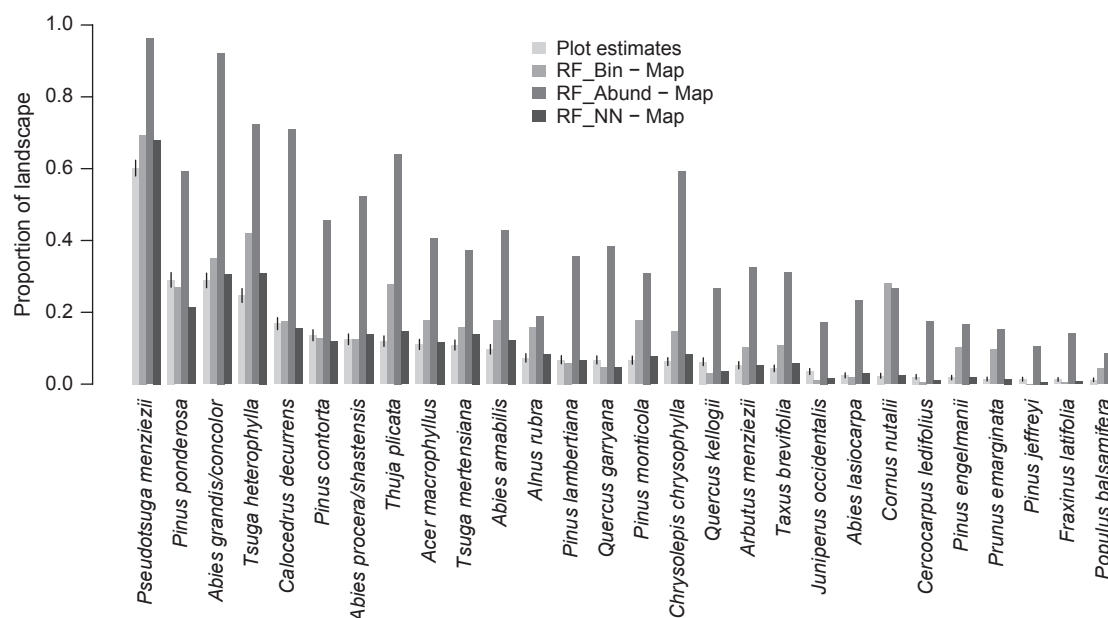


Fig. 3. Area of species presence estimated from FIA plots and from the spatial predictions of each modelling approach. Plot estimates are built directly from the FIA plot sample and represent the proportion of the forested landscape that contains each species. Error bars for plot estimates represent 95% confidence intervals for the mean binomial probability (R function 'binom.confint' in R-package 'binom', method = 'exact'). Map estimates show the proportion of the area mapped (forested land only) that contains each species. Single species models are: RF_Bin and RF_Abund. RF_NN is a multivariate model.

each plot, while RF_Bin yielded mildly inflated species richness estimates (Table 2).

The RF_NN and RF_Bin modelling approaches yielded realistic predictions of species composition. Species pairs that rarely co-occurred within our plot data were correspondingly separated within the RF_NN and RF_Bin predictions (Fig. 6). In contrast, RF_Abund often generated communities that were ecologically unrealistic. For example, RF_Abund often mapped *Pinus ponderosa* and *T. mertensiana* together (Fig. 7). These species rarely co-occur: *T. mertensiana* occupies cold, moist areas that experience deep winter snow pack, whereas *P. ponderosa* occupies warmer and drier environments, especially near the lower tree line in the eastside rain shadow (Franklin & Dyrness 1988). These two species are generally mapped to separate locations with RF_Bin and RF_NN (Fig. 7).

The RF_Bin predictions of community composition appeared slightly stronger than the corresponding RF_NN predictions (0.35 and 0.40 average Sørensen distances, respectively). For the Sørensen metric, short distances indicate similarity in species lists between observations and predictions, while for the Bray–Curtis metric short distances indicate similarities in both species identities and relative abundances between observations and predictions. RF_Abund predictions performed poorly with Sørensen distance (average distance: 0.76); however, RF_Abund

performed as well as RF_NN with Bray–Curtis distance (average distance for both: 0.45). ECDF curves of observed-to-predicted distances for all plots illustrate patterns that drive the averages (Fig. 8). There was striking similarity in the performance of RF_Bin and RF_NN with the Sørensen metric, while the RF_Abund model yielded few strong (short-distance) predictions by this metric (Fig. 8a). The Bray–Curtis curves for RF_Abund and RF_NN had distinctly different shapes, even though their mean values were almost identical (Fig. 8b). RF_NN contained more short-distance predictions and more long-distance predictions than did RF_Abund.

The high species richness of RF_Abund predictions likely drives the patterns in the distance metrics. Each community prediction from RF_Abund contained significant errors of commission (Table 2), which would lead to long Sørensen distances, but only moderate Bray–Curtis distances (because erroneous inclusions usually have low values). RF_Abund predictions also contain few errors of omission (Table 2). The net effect of these tendencies was that RF_Abund was unlikely to yield high-quality predictions due to pervasive errors of commission, and also less likely to yield poor-quality predictions because of the rarity of errors of omission. Given the pattern of RF_Abund chronically over-predicting species ranges, the rarity of omission errors is unsurprising. An extreme example

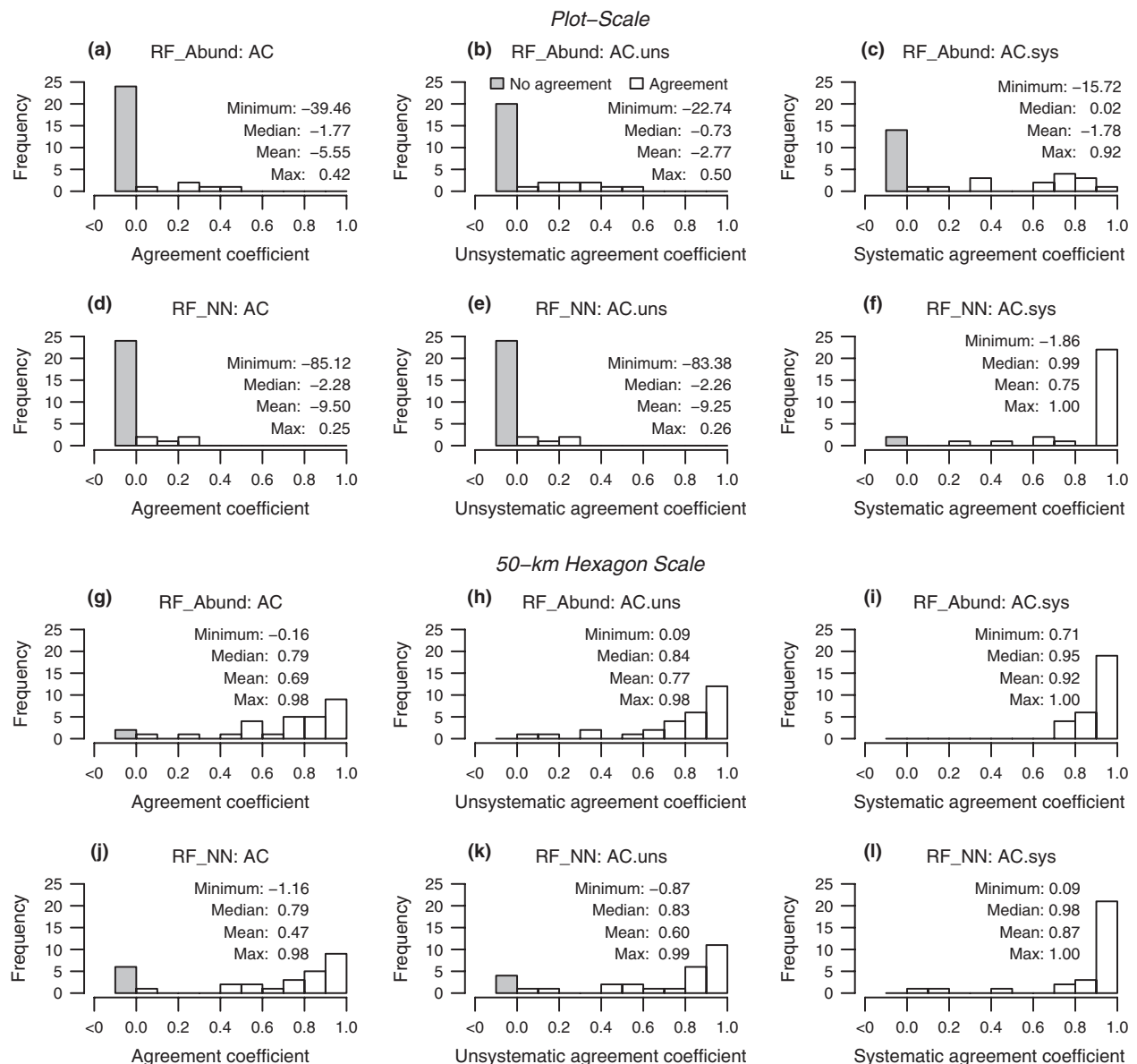


Fig. 4. Histograms of agreement coefficient statistics (Ji & Gallo 2006), calculated individually for all 28 species for continuous predictions from RF_Abund (single species) and RF_NN (multivariate) models. Statistics are shown for plot-scale (panels a–f), and broader-scale (calculated over 50-km radius hexagons, panels g–l). The left-hand column of graphs shows histograms of the overall agreement coefficient statistics for individual species predictions from RF_Abund and RF_NN models. The second column of graphs shows histograms of unsystematic agreement coefficient statistics for individual species predictions. This coefficient illustrates correlation strength between observations and predicted values (analogous to R^2 ; high values indicate low scatter around regression line). The third column of graphs shows histograms of systematic agreement coefficient statistics for the individual species predictions. This statistic measures how close the slope of an observed–predicted regression line is to 1. High values indicate that regression slope is close to 1:1. All histograms are compressed for values below zero, as these values all indicate the same thing: a lack of agreement between observations and predictions. Legend in panel 'b' applies to all panels.

illustrates why: a model that predicts all species as present everywhere will contain no errors of omission, and many errors of commission. RF_NN contained fewer errors of commission and more errors of omission than RF_Abund, and hence yielded more high- and low-quality predictions at the plot level.

Discussion

Implications for landscape analysis and conservation planning

Maps built from SDMs are used for a broad range of applications, each with different requirements in a map. Within

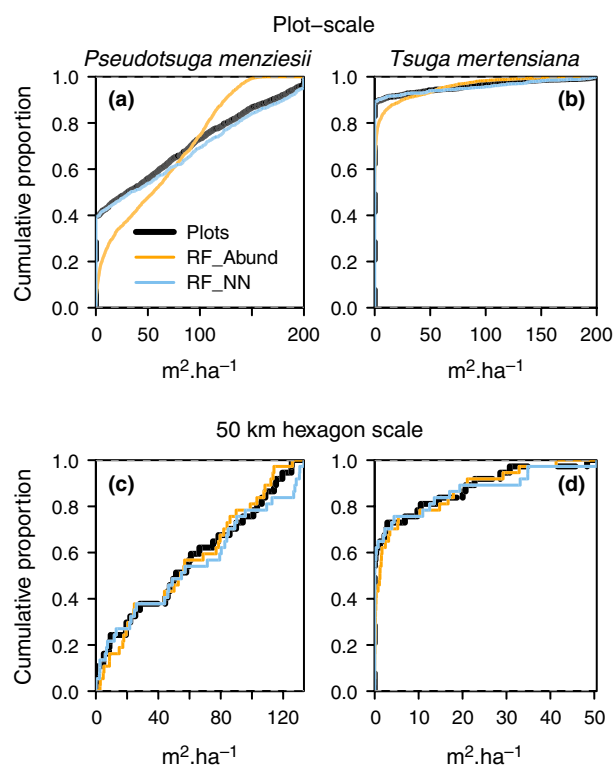


Fig. 5. Empirical cumulative distribution functions for observations (plots) and spatial predictions of *P. menziesii* and *Tsuga mertensiana* basal area, from RF_Abund (single species) and RF_NN (multivariate) models, at the plot scale (**a**, **b**), and summarized for all plots within 50-km hexagons (**c**, **d**). Legend in panel 'a' applies to all panels.

Table 1. Kolmogorov–Smirnov test statistics comparing the distribution of observed and predicted values for each species. Summaries presented here are for all species at two scales of summary: plot level and within the 50-km hexagons. RF_Abund is a single species model. RF_NN is a multivariate model.

	Min.	Mean	Max.
RF_Abund–Plot	0.37	0.64	0.88
RF_NN–Plot	0.00	0.01	0.04
RF_Abund–Hex	0.11	0.51	0.84
RF_NN–Hex	0.05	0.10	0.19

Table 2. Average plot-level species richness and types of error in plot-level species lists by model type. Values represent the average number of species per plot. For each column, letter labels indicate which values are significantly different from the others according to a generalized linear model ($\alpha < 0.01$). Within a column, cells with different letters are statistically different. Single species models are: RF_Bin and RF_Abund. RF_NN is a multivariate model.

	Species richness	Omissions	Commissions
RF_Bin	3.83 b	0.72 b	1.58 b
RF_Abund	21.51 c	0.01 a	18.54 c
RF_NN	2.89 a	1.19 c	1.10 a
Plots	2.98 a	NA	NA

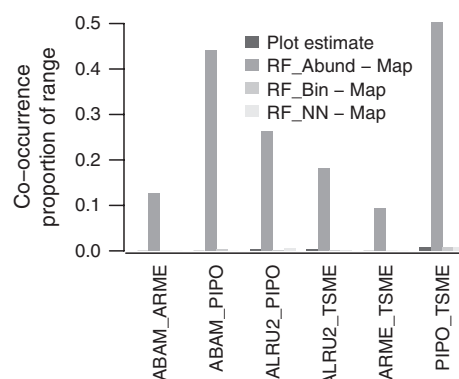
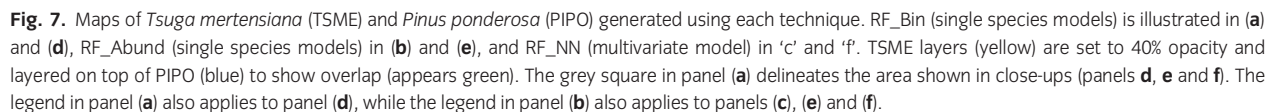


Fig. 6. Species pair co-occurrences in plot data and spatial predictions. This graph shows the range overlap of six species pairs, expressed as a proportion of the total joint range for both species (e.g. for the area occupied by either ABAM or ARME in the RF_Abund map, they co-occur over 12% of that area). These pairs were chosen from a pool of common species to represent species that rarely co-occur within the plot data. Pairs are described by USDA Plants codes for species. ABAM, *Abies amabilis*; ALRU2, *Alnus rubra*; ARME, *Arbutus menziesii*; PIPO, *Pinus ponderosa*; TSME, *Tsuga mertensiana*. Single species models are: RF_Bin and RF_Abund. RF_NN is a multivariate model.

the field of conservation planning, fine and coarse filter applications (Noss 1987) have distinctly different needs in terms of map performance. Forestry applications require unbiased multivariate information on forest composition and structure (Eskelson et al. 2009). Ecological studies of invasive pests may require information on many species simultaneously (e.g. Václavík et al. 2010). Simulation models often need input information on community composition, species abundances, as well as vegetation structure (e.g. Scheller & Mladenoff 2004; Hemstrom et al. 2007). Our maps have differing strengths and weaknesses, and none is clearly 'best' for all applications. Here, we highlight some of the trade-offs inherent in different conservation applications, and also place our work in the context of estimating future vegetation under climate change.

For fine filter conservation focused on individual species, our RF_Bin approach had clear advantages. This finding was not surprising as random forest often performs well in comparison with other techniques for building single species binary maps (e.g. Marmion et al. 2009). Although we have not modelled any threatened or endangered species, the trade-offs we highlight are relevant to that application. In particular, the balance between sensitivity and specificity has important implications for map utility (Loiselle et al. 2003). Conservation or development plans formulated from low-sensitivity maps may fail to protect missed populations, placing rare species at risk. On the other hand, low-specificity maps may trigger costly and unnecessary surveys.

For coarse filter conservation, RF_NN is well suited. Community-level information is needed to identify



RF_Abund highlight similar problems to those identified by Baselga & Araújo (2010), who found novel (i.e. currently non-existent) communities indicated by stacked maps. Our work highlights the problem at a finer spatial resolution and describes some problematic compositional errors in species co-occurrence. RF_NN is most robust for community-level predictions, not because it accounts for

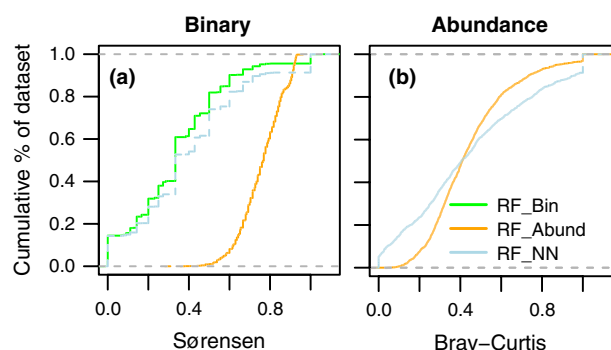


Fig. 8. Empirical cumulative distribution functions for the multivariate distance between observed and predicted communities by modelling method. Models with more short observed–predicted distances have stronger community-level predictions (greater similarity between observed and predicted communities). Values for **(a)** are calculated as the Sørensen distance, which is the binary equivalent of Bray–Curtis distance (shown in panel **b**). Sørensen distance analyses community similarities with respect to species presence/absences while Bray–Curtis distance also accounts for abundance. Single species models are: RF_Bin and RF_Abund. RF_NN is a multivariate model.

the species interactions that constrain distributions in nature, but because its predictions are constrained to assemblages that reflect the outcomes of those interactions as they are represented within the input plot data. Put another way, the RF_NN procedure does not build realistic species assemblages, but rather refrains from dis-assembling them in the first place.

The RF_NN approach to mapping is a poor choice for estimating future communities for the same reason that it is a good choice for estimating current communities. Because it can only predict species assemblages that are present within the input plot data, it cannot estimate the novel combinations that will likely emerge as species respond individually to climate change (Huntley 1991). Single species approaches still provide a better alternative (e.g. Iversen & Prasad 1998), although the problem of inflated species richness in stacked models will remain because species interactions will shape new communities that emerge as climate shifts (Walther 2010). Alternative strategies to modelling communities (e.g. Clark et al. 2011; Guisan & Rahbek 2011), or simulation modelling (e.g. Scheller & Mladenoff 2004) may be more appropriate for estimating future forest communities. For the latter, RF_NN maps are well suited to provide a starting point.

Conclusions

Single species distribution models often yielded stronger predictions for individual species for either presence or abundance, but rarely both. Imputation often yielded adequate estimates of both while also providing high-quality,

community-level information on diversity and composition. Imputed multivariate maps are therefore adequate for many purposes, from conservation reserve design to regional forest management plans, to simulation model initialization.

Acknowledgements

The ideas behind this paper stemmed from work conducted for the Nationwide Forest Imputation Study (NaFIS), a collaboration between researchers at Oregon State University, Michigan State University and the US Forest Service (the Forest Health Technology Enterprise Team, the Western Wildland Environmental Threat Assessment Center, the Remote Sensing Applications Center, Forest Inventory and Analysis, the Pacific Northwest Research Station, and the Northern Research Station).

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Supporting Information

Additional supporting information may be found in the online version of this article:

Appendix S1. Descriptions of explanatory variables for all models.