

# USE OF FOREST INVENTORY AND ANALYSIS INFORMATION IN WILDLIFE HABITAT MODELING: A PROCESS FOR LINKING MULTIPLE SCALES

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**ABSTRACT.**—We describe our collective efforts to develop and apply methods for using FIA data to model forest resources and wildlife habitat. Our work demonstrates how flexible regression techniques, such as generalized additive models, can be linked with spatially explicit environmental information for the mapping of forest type and structure. We illustrate how these maps of forest structure can be used to model wildlife habitat, focusing on the prediction of suitable habitat for cavity-nesting birds in forest systems in the Intermountain West.

Landscape data often have scale-specific resolutions and extents as well as thematic content due to methods of observation, making it difficult to scale measured responses of ecological systems either upwards or downwards. For example, use of satellite-derived data such as the National Oceanic and Atmospheric Administration's 1.1-km resolution Advanced Very High Resolution Radiometer (AVHRR) for mapping animal habitat automatically limits the scale of animal study to a 1.1-km resolution. Any gains in the ability to systematically map habitat over large spatial extents are offset by a loss of resolution relating back to the animal(s) of interest. Similarly, the kinds of ecological characteristics that plants often are associated with (e.g., microclimates, forest structure attributes) are frequently of such fine resolution that they cannot be systematically mapped or modeled over large spatial extents. As before, gains in the understanding of the ecological processes that may determine plant species distributions are offset by an inability to map these distributions over large spatial extents.

This limitation places serious constraints on the application of landscape theory to a wide range of forestry issues ranging from the development of spatially explicit, predictive maps of

forest resources for use in management to wildlife habitat modeling. The full exploration of landscape relationships requires spatially explicit depictions of habitat and other variables at fine resolutions over large spatial extents. Such depictions would allow for simultaneous exploration of relationships of variables at small spatial extents (e.g., canopy closure within forest stands) and over large landscapes (e.g., pattern of canopy closure within an ecoregion). Although it is possible to model structural attributes of habitats and vegetation on small regions using satellite imagery, the regional-scale focus of many cover-mapping efforts makes it difficult to build vegetation structure into cover maps. Current efforts provide good maps of broad cover classes at landscape levels (Homer and others 1997) but typically provide no information on the structure of the cover type or the spatial distribution of structure within the cover type. Recently, emphasis has been placed on linking forest data with satellite-based information not only to improve the efficiency of estimates of forest population totals, but also to produce regional maps of forest class and structure and to explore ecological relationships (Frescino and others 2001, Moisen 2000, Moisen and Edwards 1999, Moisen and Frescino in press). The accuracy of these types of map products is reasonably high (Edwards and others 1998, Frescino and others 2001).

Here we describe our collective efforts to develop and apply methods for using FIA data to model forest systems, focusing on the application of these FIA-based models for wildlife management. Our process requires two steps. The first step focuses on methods for modeling habitat that provide fine-grained estimations of forest habitat type and structure over

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large spatial extents. The second step is to use these representations of landscapes for modeling habitat use by terrestrial vertebrates at multiple scales. We illustrate how flexible regression techniques, like generalized additive models (GAM), can be linked with spatially explicit environmental information to map forest habitat structure. We next illustrate how the spatially explicit maps of forest structure can be used to model wildlife habitat, focusing on the prediction of suitable habitat for cavity-nesting birds in forest systems at landscape scales.

## EXAMPLE APPLICATIONS

### Study Area

The two example studies outlined here were conducted in the Uinta Mountains of northern Utah. The Uintas are characterized by an east-west orientation, and they have an approximate length of 241 km and a width of 48 to 64 km. Elevation ranges from ~1,700 to ~4,000 m. The area contains conspicuously deep, V-shaped canyons on the south side of the range and less pronounced canyons on the north side of the range. The distribution of vegetation in the Uinta Mountains is highly influenced by topographic position and geographic location. Lodgepole pine (*Pinus contorta*) is the dominant vegetation type, ranging from 1,700 to 3,000 m elevation. At elevations between 2,400 and 3,000 m, lodgepole is mixed with aspen (*Populus tremuloides*), with a few homogeneous aspen stands at lower elevations. As elevation increases, lodgepole forests are gradually replaced by spruce-fir (*Picea engelmannii*-*Abies lasiocarpa*) forest types and are frequently interspersed with large patches of wet and dry meadows. Other forest types include pinyon-juniper (*Pinus edulis*-*Juniperus osteosperma*) at lower elevations on the northeastern slope, Douglas-fir (*Pseudotsuga menziesii*) on steep, protected slopes, and ponderosa pine (*Pinus ponderosa*) forests on exposed slopes on the south side of the range (Cronquist and others 1972).

### Modeling Forest Habitat Pattern and Structure

If a major objective of landscape modeling is to enhance understanding of relationships at multiple scales as a precursor for forest management, then methods for modeling scale-related ecological parameters are paramount. From a vegetation perspective, the principal question is how to accurately and efficiently model vegetation structure and

patterns at multiple scales. Recent advances in statistical modeling techniques (Hastie and Tibshirani 1990, Hastie and others 2001, McCullagh and Nelder 1989) and geographical tools, such as remote sensing and Geographical Information Systems (GIS), have increased the opportunities for the delineation and analysis of vegetation structure and pattern.

Readers are referred to Frescino (1998), Frescino and others (2001), Moisen (2000), Moisen and Edwards (1999), and Moisen and Frescino (in press) for details regarding the complexities of generating spatially explicit models of forest structure using FIA data. The process is necessarily complex, and only a short overview of work in the Uinta Mountains is presented below. In this study area, five response variables collected on FIA plots were modeled as functions of a wide variety of digitally available explanatory variables through GAMs (Frescino 1998, Frescino and others 2001). Response variables included binary forest and lodgepole presence, as well as continuous basal area, percent shrub cover, and snag density (table 1). A variety of explanatory variables were considered, including those related to topography, precipitation, geology, spatial position, as well as several from TM and AVHRR platforms (table 2). As noted above, the GAMs used for modeling are non-parametric extensions of the more commonly used generalized linear models (GLM). The GAM, like the GLM, uses a link function to establish a relationship between the mean of the response variable(s) and a smoothed function of the explanatory variable(s). The main attraction of GAMs for vegetation modeling is their ability to handle non-normal features in the data such as bimodality or asymmetry. GAMs are best described as data-driven rather than model-driven, such that the data determine the shape of the response curves rather than fitting a known function to the data. The major weakness of GAMs is the danger of overfitting the data (Austin and Meyers 1996).

For forest and lodgepole presence, a logit link was used to transform the mean of the response to a binomial scale. For the continuous variables (basal area, percent shrubs, snag density), a Poisson link was used to transform the data to the scale of the response. A loess smoothing function (see Venables and Ripley 1997 for description) was chosen to summarize the relationship between the predictors and the response. One limitation of smoothed functions obtained from GAMs is their inability to extrapolate outside the range of the data used to build the model. To handle this problem, values of the prediction and validation data sets that were

Table 1.—Summary of response variables for modeling forest attributes in the Uinta Mountains, Utah. See Frescino and others (2001) for additional details. *P* = proportion of model-building points defined as forest or lodgepole pine.

Forest attribute	Type	Description	Distribution
Forest presence	Binomial	>10% tree cover	<i>P</i> = 0.77
Lodgepole pine presence	Binomial	Majority of forest cover	<i>P</i> = 0.31
Basal area (m <sup>2</sup> /ha)	Continuous	Area of trees at 1.37 m basal ht. (Trees > 2.5 cm d.b.h.)	Range: 0 to 70 Median: 16
Shrubs (%)	Continuous	Sum of total cover from upper, middle, and lower layers	Range: 0 to 92 Median: 15
Snag density	Continuous	Total salvable and non-salvable (Snags > 10.2 cm d.b.h.)	Range: 0 to 248 Median: 5

Table 2.—Summary of explanatory variables used to model forest attributes in the Uinta Mountains, Utah. See Frescino and others (2001) for additional details.

Variable	Type	Resolution	Source
Elevation (m)	Continuous	90 m	DMA
Aspect (°)	Continuous	90 m	Derived from DMA
	Discrete	90 m	Relative annual solar radiation (Swift 1976)
	Continuous	90 m	Radiation/wetness index (Roberts and Cooper 1989)
Slope (%)	Continuous	90 m	Derived from DMA
Precipitation	Continuous	90 m	Downscaled from PRISM; yearly precipitation climate maps
Geology	Discrete	1:500,000	Hintze (1980) Timeframe (1-Precambrian, 2-Mississippian to Eocene, 3-Alluvium)
	Discrete	1:500,000	Nutrients (1-sandstone and limestone, 2-sedimentary, 3-alluvial)
	Discrete	1:500,000	Rock Type (1-sedimentary, 2-alluvial)
Easting	Continuous	-	UTM Easting coordinates
Northing	Continuous	-	UTM Northing coordinates
District	Discrete	-	7 National Forest Ranger Districts
TM-classified	Discrete	90 m	GAP Analysis (Homer and others 1997)
AVHRR	Continuous	1,000 m	NOAA (June 1990)
TM	Continuous	30 m	Landsat TM (June 1990/August 1991) TM Band 3 (Red)
	Continuous	30 m	TM Band 4 (Near-infrared)
	Continuous	30 m	TM Band 5 (Mid-infrared)

outside the range of the model-building data set were assigned the maximum/minimum value of the respective variable in the data set.

The functional relationships between each explanatory variable and the respective response variables were analyzed for potential parametric fits following guidelines in Hastie and Tibshirani (1990) and Yee and Mitchell (1991). If a potential parametric fit existed, piecewise and second- and third-order polynomial functions were fit to the data and assessed based on the relative degree of change to the residual deviance (Cressie 1991). All explanatory variables, including all potential parametric fits, were run through a stepwise procedure to determine the best-fit model for prediction (see Chambers and Hastie 1992) using Akaike's Information Criterion (AIC). A percent deviance reduction ( $D^2$ ) was also calculated for each model, representing the percent of deviance explained by the respective model (Yee and Mitchell 1991). Once the model fits were derived (see Frescino and others 2001, tables 3 and 4), the model was applied to all the explanatory digital layers (table 2) and predictive map surfaces generated. The result was a series of predictive maps of forest attributes having fine resolution (~0.8 ha) and covering large spatial extents (>1 million ha) (fig. 1).

Accuracy of the models predicting forest and lodgepole presence was high, ranging from 86 to 80 percent,

respectively. Sixty-seven percent of the basal area validation points fell within  $\pm 15$  percent (11.5 m<sup>2</sup>/ha) of the true value, 75 percent of the shrub density validation points fell within  $\pm 15$  percent of the true cover, but only 54 percent of the points fell within  $\pm 15$  percent of the true snag count.

### Modeling Cavity Bird Nesting Habitat in Forested Systems

Once the maps of forest attributes are generated, the next step is to generate models of bird presence based partly on the spatially explicit forest maps. As an example in the Uinta Mountains, habitat associations based on landscape patterns were modeled for four species of cavity-nesting birds nesting in aspen (Lawler 1999; Lawler and Edwards, in press). These species include red-naped sapsuckers (*Sphyrapicus nuchalis*), northern flickers (*Colaptes auratus*), tree swallows (*Tachycineta bicolor*), and mountain chickadees (*Parus gambeli*).

Habitat models were built for each of the four species using classification trees (Breiman and others 1984, Venables and Ripley 1997). Classification trees are a flexible and simple tool for modeling complex ecological relationships (De'ath and Fabricius 2000). Classification trees work by recursive partitioning of the data into smaller and more homogeneous groups with respect to the response variable. Each split is

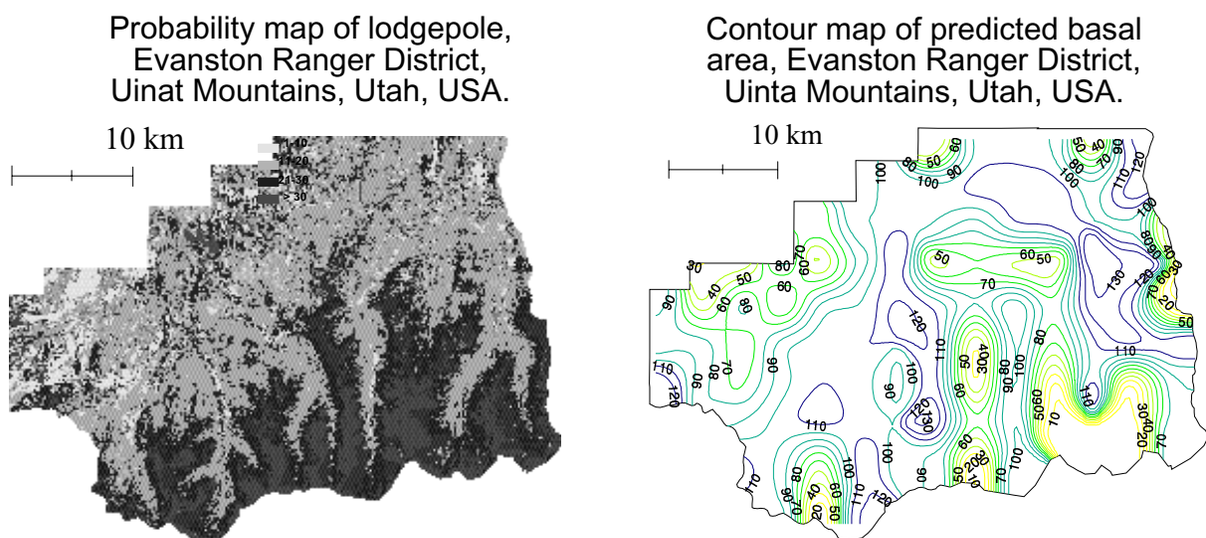


Figure 1.—Example maps of nominal (lodgepole presence) and continuous (basal area) responses generated for a ~100,000-ha region of the Uinta Mountains, Utah (from Frescino 1998).

made by the explanatory variable and the point along the distribution of that variable that best divides the data. See De'ath and Fabricius (2000) and Lawler and Edwards (in press) for a more thorough discussion of the use of classification trees in ecological modeling.

The four species models included a number of variables pertaining to the amount and configuration of forest and open area (fig. 2). Spatially explicit predictions for each of the four species were produced from these models (fig. 3). The spatial configuration of forest that was predicted as suitable nesting habitat differed among the four species, and the models varied in their ability to correctly predict nests at the new sites (Lawler and Edwards, in press). The northern flicker model was the most accurate (84 percent of nests correctly classified). The red-naped sapsucker and tree swallow models were also relatively accurate (80 percent and 75 percent of the nests correctly classified, respectively). The mountain chickadee model was far less accurate, correctly predicting only 50 percent of the nests at the test sites. These estimates are within ranges of accuracy reported elsewhere (Edwards and others 1996).

## CART model Red-naped sapsucker

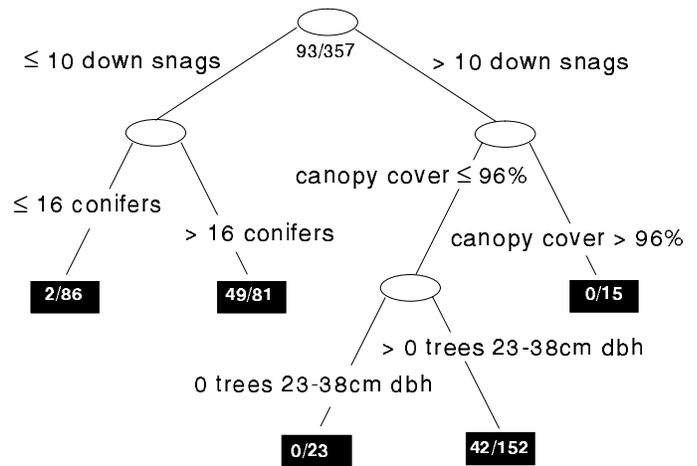


Figure 2.—Classification and regression tree model predicting nesting habitat for red-naped sapsuckers. Models for the other species were similar in structure, varying only in the predictor variables and tree complexity (see Lawler and Edwards, in press).

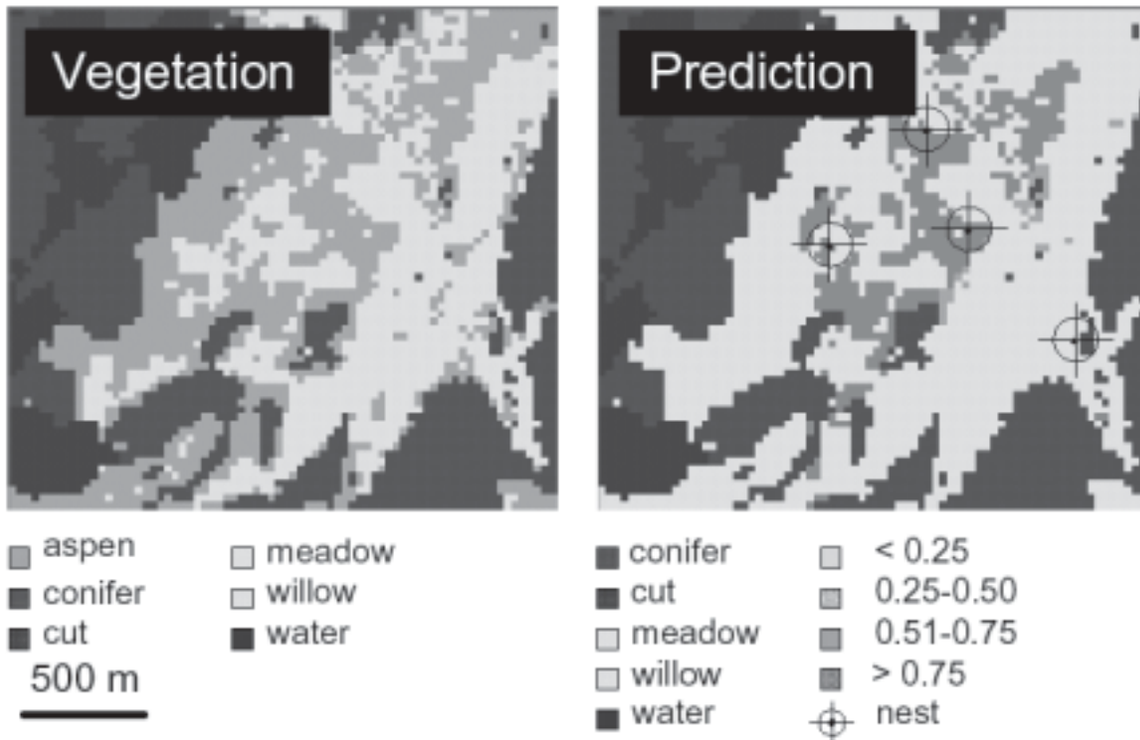


Figure 3.—Vegetation and spatially explicit prediction maps for northern flicker nesting habitat. Medium gray in the vegetation map represents suitable nesting habitat and is based on classical WHR approaches (see text). Note how the amount and distribution of gray are reduced under the refined vegetation models, which then are incorporated in the wildlife models as described in the text. Nests are represented as circles with crosshairs.

## DISCUSSION

The ability to create spatially explicit depictions of vegetation type and structure is dependent, in part, on the flexibility and capability of the models used to predict vegetation characteristics. GAMs, in contrast to some analytical procedures (e.g., ordination and linear regression models), do not make *a priori* assumptions about underlying relationships, thus allowing the data to drive the fit of the model instead of the model driving the data. The graphical nature of GAMs also allows a visualization of the additive contribution of each variable to the respective response using smoothed functions. One limitation of GAMs is the uncertainty associated with extrapolation of the smoothed functions, particularly at the tails of the distribution. As suggested by Hastie and Tibshirani (1990) and Yee and Mitchell (1991), parametric functions were to the model whenever statistically allowable, thus constraining the behavior of the functions in the extreme ranges of the data. See Moisen and Frescino (in press) for a comparison of alternative modeling techniques for predictive mapping applications.

Once the vegetation type and structure are modeled, the resultant maps can be linked with wildlife models and used to create predictive maps. Although predictive models based on landscape patterns may prove to be a promising technique in light of their ease of use and relative accuracy, like all models they have distinct shortcomings. The ability to build such models depends on having access to remotely sensed data. Fortunately, remotely sensed data are becoming not only more diverse, but also more widely available.

Because the field of landscape ecology is relatively young, associations between given species and landscape patterns are not as prevalent in the literature (Karl and others 1999) as are associations with the composition and structure of vegetation at relatively fine spatial scales (e.g., Cody 1985). Thus, many of the basic habitat associations related to landscape patterns will need to be determined in the field for the first time.

Selecting the scales at which to measure landscape patterns is difficult when modeling several different species. Different species are likely to respond to their environment at different spatial scales (Wiens 1989). Models built solely at coarse spatial scales and using only vegetation type are likely to be less accurate when fine-scale associations with structural attributes are strong. Our approach, which employs

techniques capable of modeling fine-scale attributes (e.g., canopy closure, stem density) at fine resolutions, overcomes this issue and generally increases model predictive capabilities. The use of new, more flexible modeling techniques such as classification trees (De'ath and Fabricius 2000) may further improve the predictive capability of models of forest resources, and the wildlife dependent on these resources, as well as the ease of model building and interpretation. Although our results indicate that our approach may not work equally well for all species, we found that when tested and refined, models that rely on landscape patterns derived from FIA data may provide a reliable alternative to traditional wildlife models that require the collection of habitat data in the field and have no spatial resolution.

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