MODELING CURRENT AND HISTORIC HABITAT FOR CANADA LYNX (LYNX CANADENSIS) IN THE UPPER PENINSULA OF MICHIGAN

By

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ABSTRACT

MODELING CURRENT AND HISTORIC HABITAT FOR CANADA LYNX (LYNX CANADENSIS) IN THE UPPER PENINSULA OF MICHIGAN

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In the ruling to list Canada lynx (Lynx canadensis) as a federally threatened species, the US Fish and Wildlife Service identified the Great Lakes region as an area of concern. While there is no current evidence of a resident population in the Upper Peninsula (UP) of Michigan, trapping and track records over the past century suggest the area was periodically invaded by lynx during population eruptions in Canada. My objectives were to quantify past and present forest conditions in the UP for lynx habitat potential, and estimate changes in habitat connectivity between Canada and the UP. I used a spatially explicit, landscape-level habitat model that required multiple layers of spatial data compiled in a GIS to describe lynx habitat components. Forest inventory plots were used to provide detailed stand attribute information. The historical range of variability in presettlement forest conditions was simulated with the landscape age-class demographics simulator (LADS). Outputs from the model indicated that potential habitat has become more widespread under current conditions than that which existed during the presettlement era. Foraging quality has increased in many areas across the UP, but remains low throughout. Non-habitat interspersion is limiting under current conditions and has increased dramatically from that of presettlement in the eastern UP. This increase has resulted in lower habitat quality and decreased connectivity of habitat in the eastern UP. The establishment of a resident lynx population in the UP is questionable.
ACKNOWLEDGEMENTS

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The US Fish and Wildlife Service (USFWS) made their final ruling on 24 March 2000 to list Canada lynx (*Lynx canadensis*) as a threatened species in accordance with the Endangered Species Act of 1973, following an investigation of the status of lynx populations in the contiguous United States (USFWS 2000). They determined that some current land management practices had the potential to negatively affect lynx and/or lynx habitat in the absence of adequate protection for the species. The ruling distinguished four regions (the Northeast, Great Lakes, Northern Rockies & Cascades, and Southern Rockies) collectively as one “Distinct Population Segment” (USFWS 2000) based on the criteria that lynx populations in those regions were discrete and significant from populations in northern Canada. The separation by international political boundaries made them discrete, mostly through disparities in management policies between the US and Canadian governments, while differences in vegetation types, climate, and ecology made them significant. In light of the final ruling, government agencies, such as the USDA Forest Service, are faced with developing and implementing management strategies that facilitate lynx populations on public lands. Attaining this management goal has been problematic because an understanding of current and historical lynx distributions and habitat use patterns in the southern regions of their range, such as the Great Lakes, is largely incomplete.

Lynx are commonly associated with the boreal forest region of northern Canada, also known as the taiga (McCord and Cordoza 1982, McKelvey et al. 2000a). Their distribution in the contiguous United States is limited to southern boreal forests, consisting of subalpine conifers in the West, and mixed coniferous-deciduous forests in...
the East (Aubry et al. 2000). These southern boreal forests have been increasingly fragmented by human disturbance, and lynx habitat patches tend to be smaller than those in the north. Some habitat patches in the southern boreal forests are able to support resident lynx populations (McKelvey et al. 2000b), but it is uncertain how source-sink dynamics among subpopulations have been affected by human-induced fragmentation, or how this impacts lynx viability in these areas. In other parts of the southern boreal forest, at the limits of the species’ range, lynx presence is attributed “solely” to the dispersal of individuals from core populations in Canada (USFWS 2000). Habitat quality in these regions is, therefore, assumed to be poor; the temporal and spatial extents to which these regions have acted as sinks to the core populations in Canada are undefined. This complicates the formulation of management policies, considering that the overall importance of these regions (e.g. the Great Lakes) to the species is questionable.

The scant amount of information concerning lynx in the Great Lakes region consists of occurrence data, as documented by trapping and/or track records from the past 150+ years (Aubry et al. 2000, Beyer et al. 2001). These records appear to illustrate the historical rarity of lynx in Michigan, though they do not provide any indication of individual fitness or population viability. Also, abundance (i.e., number of records per geographic area) can be a misleading measure of habitat quality in the absence of ancillary ecological information (Van Horne 1983), so the occurrence data alone are not useful for determining whether there were viable populations of lynx historically in the state. Wood and Dice (1924) warned that some of the historical lynx records from Michigan were unreliable given the potential misidentification of bobcat (*Lynx rufus*) for lynx, but several verified records dated between 1842-1912 do exist for the Upper and
Lower Peninsulas. The lynx was considered “extremely rare, if not already extinct” in
the ninth Biennial Report (1937-38) of the Michigan Department of Conservation, and
only 6 records of lynx occurred between 1904-1939 for the Upper Peninsula (UP), with a
lone record in 1917 for the Lower Peninsula (McKelvey et al. 2000a, Beyer et al. 2001).
Burt (1946) illustrated a map showing “questionable” records of lynx across most of
Michigan and as far south as Monroe County, but reiterated the belief that the species
was most likely extirpated from the state, except for a possible remnant on Isle Royale.
Harger (1965:152) believed that lynx were “definitely making a comeback” in the UP
due to the influx of trappings in 1962, but McKelvey et al. (2000a) pointed to an irruption
of lynx populations in Canada as the cause for this increase. McKelvey et al. (2000a)
compared lynx occurrence data from Michigan, Wisconsin and Minnesota with trapping
records from Ontario, Manitoba, and Saskatchewan spanning 50+ years, and found a
strong correlation when the Canadian data were lagged 3 years. They also examined the
potential correlation between lynx occurrence/harvest data and snowshoe hare (*Lepus
americanus*) harvest data in Minnesota and found no relationship, thus supporting the
hypothesis that migration was the primary force influencing lynx populations in the Great
Lakes region. Extensive track surveys by the Michigan Department of Natural Resources
(DNR) in the 1990s found no sign of lynx in the UP, prompting Beyer et al. (2001) to
conclude there was no evidence of a resident population. Whether or not a resident
population exists in Michigan, it is possible that individuals dispersing from Canada
during a population increase may enter the UP in search of resources (McKelvey et al.
2000a). Thus, the identification of these resources and their location within the landscape
becomes an important task for state and federal agencies in Michigan.
The objectives of this research project were to quantify current and historic (i.e., prior to European settlement) habitat potential for Canada lynx in the UP of Michigan, analyze temporal trends in the amount and distribution of habitat in the region, and to examine the dynamics of potential migrations from Canada into Michigan. The process involved formulating spatial data in a Geographic Information System (GIS) that could describe the landscape, from both time periods, in terms of the forest attributes that define the current understanding of lynx habitat requirements. The project also involved simulating the dispersal routes of individuals from Canada into Michigan, given the obstacles that may be perceived in the current landscape. Major shifts in forest dynamics across the UP due to anthropogenic influences (e.g., human development, harvesting, fire suppression) have most likely affected the amount and distribution of lynx habitat since presettlement, and the increased fragmentation due to roads and development in the Great Lakes region on a whole has affected the ability of lynx to disperse into and throughout the area. The results of these analyses may guide the formulation of management practices and policies for lynx conservation in the UP, and the methods used can be applied other regions in the contiguous United States.
CHAPTER 1:
TEMPORAL CHANGES IN HABITAT POTENTIAL FOR CANADA
LYNX IN THE UPPER PENINSULA OF MICHIGAN

INTRODUCTION

The listing of Canada lynx (*Lynx canadensis*) as a threatened species in the contiguous United States by the US Fish and Wildlife Service (USFWS) in March of 2000 necessitated the implementation of management policies that facilitate the persistence of lynx on public lands (USFWS 2000). This is complicated by an incomplete understanding of lynx ecology in the southern boreal forests of the United States, which represent the southern periphery of the species’ geographic range (McCord and Cordoza 1982). In the Great Lakes, the only information regarding the species comes from historical trapping and occurrence data (Aubry et al. 2000), and the scarcity of verified records suggests the species was traditionally rare throughout Michigan, with nearly all occurrences in the Upper Peninsula (UP). Currently, there is no evidence of a resident population in the UP (Beyer et al. 2001), though transient lynx from neighboring states and Canada may disperse into the area in search of resources (McKelvey et al. 2000a). Accordingly, the USFWS called for “accurate mapping of lynx habitat in the Great Lakes Region” (USFWS 2000:16057) to provide the location and distribution of resources necessary for the species’ persistence and identify potential areas for conservation and management.

One resource that is critical for determining lynx survival is its primary prey, the snowshoe hare (*Lepus americanus*). The close relationship between fluctuations of lynx and hare populations in the boreal forest ecosystem is well documented, and patterns of
habitat use exhibited by lynx are assumed to be highly correlated with those of hare (Keith 1963, Nellis et al. 1972, Brand et al. 1976, McCord and Cordoza 1982, Koehler and Aubry 1994). Lynx are known to prey upon squirrels and other small mammals, as well as ungulates and grouse during the summer and periods of low hare density (see diet summary in Koehler and Aubry 1994), but the availability of hare seems to limit the persistence of lynx, regardless of alternative prey abundance (Ward and Krebs 1985, Mowat et al. 2000). There is some debate among biologists as to whether or not southern populations of snowshoe hare exhibit the same predator-prey cycle found in the north (see review in Hodges 2000). In the southern boreal forests, habitat conditions are thought to be suboptimal in comparison to the northern boreal forest, and may not allow hare populations to reach the high densities found in the north (Koehler 1990). Marginal forest conditions and the presence of other predators (e.g., coyote \([\text{Canis lupus}]\), bobcat \([\text{Lynx rufus}]\) not abundant in the north could be keeping hare populations at a stable low density (Wolff 1981, Koehler and Aubry 1994), resulting in a cycle with reduced amplitude or no cycle at all (Hodges 2000).

Wolff (1980, 1981) hypothesized that hare populations in the south were unable to successfully disperse among habitat patches due to the higher densities of predators, resulting in the absence of a cycle. In the northern boreal forest, when the pressures of predation and competition on hares are both low, individuals are able to disperse from optimal habitat and persist in suboptimal areas as populations increase. The stability of predator populations in the southern boreal forest, along with increased herbivore competition, preclude hare populations from reaching the high densities needed to initiate the predator-prey cycle (Wolff 1980, 1981). This “refugium” model (Wolff 1980, 1981)
has been tested in other studies, which have found that habitat patches in some southern regions are too small (i.e., <10 ha) to provide significant refugia for hare, resulting in similarly high predation rates between optimal and suboptimal habitat conditions (Keith et al. 1993, Wirsing et al. 2002). Thus, the fragmentation of habitat patches decreases the ability of hares to successfully disperse, as well as increasing the vulnerability of philopatric individuals to predation. Patches of sufficient size (e.g., ~50 ha [Litvaitis et al. 1985]) are necessary for hare densities in southern boreal forests to rival those in the north (Wirsing et al. 2002).

One distinct difference between northern and southern boreal forests is the relatively high frequency and extent of fire disturbance in the north, which can create widespread areas of early successional vegetation types important to hares, interspersed with a mosaic of mature forest patches (Fox 1978, Keith et al. 1993, Agee 2000). Snowshoe hares select habitat based on the density of understory vegetation providing security cover and winter browse (Brocke 1975, Wolff 1980, Litvaitis et al. 1985, Parker 1986, Hodges 2000), which is generally most abundant in forests during early seral stages. The combination of fire suppression and exclusion in the contiguous United States, along with naturally longer fire return intervals across boreal regions of the Northeast and Intermountain West, results in less frequent disturbances of a lower intensity than those found in the north. Koehler (1990:849) explained that the early successional patches he found in Washington were “small, isolated islands” created by wind-throw and lightning, typically <1 ha, and unlikely to support sufficient numbers of hares. Fire disturbance is among the mechanisms believed to drive the lynx-hare cycle in northern Canada because of its high frequency and widespread impact, resulting in the
constant re-initiation of secondary succession (and consequently, abundant cover) across large tracts of forest (Fox 1978). In northern Michigan, the current lengths of fire rotations across the landscape are estimated to have increased tenfold over those that existed prior to European settlement (Cleland et al. 2004); this has undoubtedly altered the dynamic interactions between forest disturbances and hare populations in the region.

While it is apparent that hare populations require an adequate amount of early successional vegetation types to persist (Hodges 2000), lynx have been shown to benefit from an interspersion of mature forest (O’Donoghue et al. 1998, Mowat et al. 2000). Lynx are unable to effectively hunt in vegetation that is too dense for them to traverse, making early seral stands a refuge for snowshoe hare (Wolff 1980); an interspersion of mature forest would create edges that are navigable for lynx to hunt. Mature forest stands may also provide necessary denning conditions for lynx, as the most common characteristic found to be an indicator of denning habitat is the amount of downed woody debris (Koehler 1990, Koehler and Brittell 1990, Slough 1999, see review in Mowat et al. 2000), which is frequently scattered throughout older mesic forests. An old forest dominated by conifers with a sparse canopy late in succession could potentially contain a dense understory and an adequate array of woody debris, thus providing both foraging and denning opportunities for lynx (Buskirk et al. 2000).

The absence of quantitative studies on lynx-habitat relationships in the UP would require the use of theoretical habitat models to evaluate the landscape for lynx. A framework proposed by Roloff and Haufler (1997) integrates the use of habitat suitability index (HSI) modeling (USFWS 1981) with a Geographic Information System (GIS) to allow a landscape-level analysis of habitat at multiple spatial scales that could address the
viability of an individual organism, sub-population, or whole population in a given area (Roloff and Haufler 1997). This framework was similar in concept to habitat evaluation procedures (USFWS 1980), and was demonstrated using Canada lynx as the focal species (Roloff and Haufler 1997). The process involved quantifying habitat suitability at the home-range scale across a grid map of the landscape using HSI models, which assume that a species will use a certain area more frequently when it contains the necessary life requisites (USFWS 1981). The resulting grid map is used to simulate the spatial distribution of hypothetical lynx home ranges, based on quality and quantity thresholds of habitat suitability described by previous studies, which then provides a measure of habitat potential across the landscape (Roloff and Haufler 1997). An important step in the framework is the validation of model assumptions with observational data. Nylen-Nemetchek (1999) used this procedure with a lynx HSI model (Roloff 2001) to assess lynx habitat potential within the Riding Mountain National Park in Manitoba and found a significant positive correlation between model outputs and surveyed lynx presence.

Our study objectives were to determine the amount and distribution of potential Canada lynx habitat in the UP of Michigan, and examine differences between the current and historic (i.e., prior to European settlement) ranges of variability in forest conditions and subsequent habitat potential. Habitat potential for each time period was quantified using the modeling framework proposed by Roloff and Haufler (1997) with the lynx HSI model developed by Roloff (2001). Results of this analysis could help guide natural resource managers with planning objectives for lynx conservation and management in the Great Lakes region, as well as providing a context for inferences about historic lynx distributions in the UP.
STUDY AREA

The UP of Michigan contains approximately 42,610 km², located in the northern portion of the Great Lakes region. It is bounded by Lake Superior to the north, Lakes Michigan and Huron to the south, and the state of Wisconsin to the west (Figure 1). Over 80% of the land area is forested (Leatherberry and Spencer 1996), and the region falls within the Laurentian Mixed Forest Province, representing a transition from broadleaf deciduous to boreal coniferous forests (Albert 1995). The climate is humid continental, with significant influence from Lake Superior on the north side. Winters are moderately long and cold, with mean temperatures of -9°C along the shorelines and -13°C inland; summers are short and cool, with mean temperatures of 19°C along the shorelines and 17°C inland (Eichenlaub et al. 1990). The length of growing season ranges from 180 days along the shorelines to 130 days inland (Eichenlaub et al. 1990). Annual precipitation averages 76-96 cm depending on location and is evenly distributed throughout the year. Snowfall occurs most regularly between November and April, and average seasonal totals range from 150 cm across the southern UP to >500 cm along the Keweenaw Peninsula and northern portions of the UP receiving lake effect snow off Lake Superior (Eichenlaub et al. 1990).

The physiography of the UP is defined by numerous glacial landforms (e.g., outwash and lacustrine plains, ground and end moraines) that were created during the last retreat of glaciers >10,000 years before present (Dickman and Leefers 2003). A boundary of Paleozoic and Precambrian bedrock divides the UP roughly in half: the eastern section is characterized by a lower elevation (155-195 m) and relatively flat topography with sandy lake plains that are poorly drained, while the western section
Figure 1. Location of the Upper Peninsula (black outline) and the extent of forest cover (green) within the Great Lakes region. Images in this thesis are presented in color.
(at elevations ranging 184-604 m) contains varying depths of glacial drift forming ground and end moraines over igneous and metamorphic bedrock, with exposed bedrock ridges along the coast of Lake Superior (Albert 1995). Each section encompasses a variety of smaller ecoregions defined by similar landforms, soils and climate regimes; these factors interact to influence natural disturbance and successional patterns among forest communities in the UP (Albert 1995, Frelich 2002).

Human influences after European settlement have altered the structure and composition of forests in the UP. Extensive logging and catastrophic fires during the late 19th and early 20th centuries have resulted in second growth forest throughout most of the region, including an increased presence of aspen (*Populus* spp.) (Dickman and Leefers 2003). Species such as eastern hemlock (*Tsuga canadensis*) and eastern white pine (*Pinus strobus*) have a minor presence in the landscape compared to that of the presettlement era due to heavy selective logging (Dickman and Leefers 2003), and fire suppression and exclusion over the past century have favored an increased dominance of shade tolerant hardwoods (e.g., maple [*Acer* spp.]) (Zhang et al. 2000). Despite these changes, the species compositions of forests in the UP remain heavily dependent on physiographic conditions.

Hydric lowlands are populated by conifers such as (in order of predominance) northern white-cedar (*Thuja occidentalis*), balsam fir (*Abies balsamea*), black spruce (*Picea mariana*), tamarack (*Larix laricina*) and hemlock, with hardwoods including black ash (*Fraxinus nigra*), red maple (*Acer rubrum*) and balsam poplar (*Populus balsamifera*). Mesic uplands are populated by hardwoods such as sugar maple (*Acer saccharum*), aspen (*Populus tremuloides*, *Populus grandidentata*), red maple, yellow birch (*Betula*
alleghaniensis), paper birch (Betula papyrifera), beech (Fagus grandifolia), basswood (Tilia americana), and white ash (Fraxinus americana) with conifers including fir, cedar, hemlock, and white spruce (Picea glauca). Xeric uplands are populated by conifers such as jack pine (Pinus banksiana), red pine (Pinus resinosa), fir, white pine and spruce, with hardwoods such as red maple, aspen, and northern red oak (Quercus rubra).

Land use in the UP is primarily recreation and timber harvest. Out of the 33,900 km$^2$ of forested land, 40% belongs to state and federal governments, including two national forests (the Ottawa and Hiawatha), and 18% is owned by timber industry (Leatherberry and Spencer 1996). There are approximately 328,000 residents; over 75% of the land area has a population density <4 persons/km$^2$, and 95% has a population density <15 persons/km$^2$ (U.S. Census Bureau 2000). Wildlife species that are relevant to lynx ecology in the UP include coyote, bobcat, gray wolf (Canis lupus), and fisher (Martes pennanti), as well as snowshoe hare, white-tailed deer (Odocoileus virginianus) and moose (Alces alces).

METHODS

LYNX HABITAT MODEL

Roloff’s (2001) lynx habitat model uses a limiting factor approach to characterize three components (life history traits – foraging, denning, non-habitat interspersion) demonstrated as necessary for lynx survival. The hypothesized requirements of foraging, denning, and interspersion were related to those suggested by Koehler and Aubry (1994). The input for the model consists of an ecological land classification, in the form of a grid-based GIS, that can adequately stratify the variability in forest vegetation attributes relating to each habitat component (Roloff 2001). These habitat components are assessed
within the allometric home range for lynx (250 ha), a theoretical scale at which habitat selection is hypothesized to be strongest (Roloff and Haufler 1997).

Several calibrations were necessary to account for differences in biogeoclimatic conditions between the Great Lakes and the Intermountain West, for which the model was originally developed (Roloff 2001). A similar approach was taken by Nylen-Nemetchek (1999) in Manitoba. Roloff (2003) (Appendix A) incorporated these changes into a new model specifically designed for the Great Lakes region, based on information obtained from the literature and recommendations from local forest and wildlife biologists. The changes involved reductions in the magnitude of variables related to snow accumulation, topographic influence and forest stature (Roloff 2003). The indices of habitat quality and quantity that were developed for each component in the original validated model were unchanged in the new model. All references to the lynx model, hereafter, will be to this new model developed in 2003. A schematic of the modeling process is illustrated in Appendix B.

Foraging Component

The availability of prey in the winter has been shown to affect the size of a lynx home range, and the degree to which home ranges overlap (McCord and Cardoza 1982, Ward and Krebs 1985); therefore, the foraging component of the model focuses on ecological conditions during the winter season. The foraging component consists of a snowshoe hare sub-model that assesses hare habitat quality within vegetation types based on the availability of palatable browse and winter security cover in the understory (Roloff 2003). For this study, I assumed that horizontal understory cover was the most limiting factor, based on evidence from the literature regarding the demonstrated importance of
horizontal cover for hare (Brocke 1975, Wolff 1980, Litvaitis et al. 1985, Parker 1986, Hodges 2000) and the high variability in definitions of palatable browse (see review in Hodges 2000). Therefore, hare habitat quality was indexed by 2 measures: (1) horizontal cover, defined by the percentage of vegetation cover measured perpendicular from the ground within two vertical height strata, 0-1 m and 1-2 m (accounting for variable snow depths throughout winter) at a standing distance of 10 m; and (2) understory dominance, defined by the percentage of coniferous species among all trees with a height to crown <3 m. Habitat quality for each variable (portrayed by HSI score on a scale of 0-100) is determined by a production function (Figure 2); quality increases logistically with the percentage of horizontal cover (Figure 2a), and linearly with the percentage of conifers (Figure 2b). These production functions were based on relationships demonstrated in the literature concerning perceived thresholds of horizontal cover, below which habitat is not provided for hare (Brocke 1975, Wolfe et al. 1982, Parker 1986, Ferron and Ouellet 1992), and the additional importance of a conifer component in the understory for providing thermal cover during winter (Buehler and Keith 1982, Orr and Dodds 1982, Monthey 1986). A total HSI score for horizontal cover is calculated by combining the HSI scores for each height stratum with an arithmetic mean, which allows the horizontal cover in a particular height stratum to contribute to habitat quality regardless of the cover availability in the other height stratum. The overall hare habitat quality is calculated as the geometric mean of HSI scores for horizontal cover and understory dominance. This model structure insures that a vegetation type will not qualify as suitable hare habitat if horizontal cover is inadequate.
Figure 2. Production function describing the relationship between hare habitat suitability index and stand attributes for: (a) horizontal cover \( y = \frac{103}{1+1400e^{-0.11x}} \); and (b) understory dominance (for \( x < 60 \), \( y = 0.8333x + 50 \)).
The snowshoe hare sub-model produces a grid map of hare habitat quality with values ranging from 0-100, describing non-suitable to optimal conditions, respectively. Habitat units (= quality x quantity) are preferentially aggregated by a raster-based GIS function (Plume 2005) into hare home ranges of differing size. The function places seeds randomly on the landscape, and habitat units with the highest value are selected within the search window until a habitat unit goal is achieved (Plume 2005). The habitat unit goal is based on the allometric home range for hare (4.5 ha) (Boutin et al. 1986), which describes the minimum area required by a mammal according to its mass. A home range is created for each seed as long as the habitat unit goal can be achieved before the home range reaches a maximum area (20 ha for hares). As the quality of hare habitat decreases, the number of pixels required to meet the habitat unit goal, and thus, the size of the simulated hare home range, increases. For example, one hypothetical home range would have a total area equal to the minimum of 4.5 ha if the average quality of the contributing habitat units was 100, while another would be 6.0 ha if the average quality was 75.

Home ranges are considered viable (i.e., contributing to population fitness) at an average quality ≥60 and marginal (i.e., not contributing to population fitness) at an average quality ≥25 and <60. Home ranges with an average quality <25 are deemed non-viable and precluded from further analysis, under the assumption that these areas do not provide habitat for hares of a high enough quality to survive and reproduce. The thresholds are based on the inverse relationship between reproductive output and home range size of hares observed in previous studies (Behrend 1962, Dolbeer and Clark 1975, Cary and Keith 1979, Sievert and Keith 1985) (Figure 3), and the premise of these thresholds relates to forage availability for lynx (Roloff 2003). Areas with viable hare home ranges
Figure 3. Viability relationship for snowshoe hares developed from previous studies (from Roloff 2003). The viability threshold occurs at 12 young/year (HSI = 60), and the marginal threshold occurs at 5 young/year (HSI = 25)
would potentially support higher reproduction of hares, and thus, contribute to lynx
foraging more than areas with marginal hare home ranges. Areas with non-viable hare
home ranges are presumably void of snowshoe hares that can survive and reproduce.

A production function describes the relationship between lynx foraging quality and the
number of hypothetical hare home ranges within an allometric lynx home range
(Figure 4a). The function is based on the estimate that lynx need to consume 1 hare
every 2 days during the winter (Brand et al. 1976, as cited in Roloff 2003), and the winter
is assumed to last approximately 180 days in Michigan (Eichenlaub et al. 1990).

Therefore, lynx foraging quality is highest at ≥90 hare home ranges and lowest at 0 hare
home ranges, with a linear trend between. The grid map is generated by counting the
hare home ranges within a 250 ha moving window around each pixel. Hare home ranges
that are viable contribute twice as much as those that are marginal, to account for the
increased reproduction and survival of hares in higher quality habitats. The resulting grid
map describes lynx foraging availability at the scale of a lynx home range.

**Denning Component**

The denning component defines quality based on thresholds of forest structure
and the degree of juxtaposition with summer foraging habitat for lynx (Roloff 2003). In
the Great Lakes region, vegetation types are assumed to provide potential denning sites
when they contain ≥11.49 m² ha⁻¹ of trees with an average DBH of 20 cm, ≥50% canopy
closure, and mesic soil conditions (Roloff 2003). These values are intended to represent
mature, productive forest conditions, which are hypothesized to provide the necessary
requirements for lynx denning habitat (Koehler 1990, Koehler and Aubry 1994, Mowat et
al. 2000). The patches of these vegetation types are required to be at least 2.0 ha in size,
Figure 4. Production functions relating lynx habitat quality to each of the three model components within a 250 ha home range: (a) foraging (for $x < 90$, $y = 1.1x$); (b) denning (for $x > 400$ and $< 1750$, $y = -0.0741x + 129.64$); and (c) interspersion (for $x > 300$ and $< 1500$, $y = -0.0833x - 25$).
adjoined by other lynx habitat for $\geq 50\%$ of their perimeter, and within 0.8 km of suitable summer foraging habitat (defined as vegetation types providing $\geq 20\%$ understory cover) (Roloff 2003). A mosaic of lynx summer foraging and denning habitat would provide female lynx the ability to move kittens between denning sites to avoid predation (Koehler and Aubry 1994). Denning quality is, therefore, determined by a function of the average distance to lynx denning habitat from a 100 x 100 m grid of points within a lynx allometric home range (Figure 4b). Average distances to denning habitat $< 400$ m provide the most suitable conditions and those $> 1750$ m provide non-suitable conditions. The resulting grid map describes lynx denning potential at the scale of a lynx home range.

**Interspersion Component**

The interspersion component addresses travel requirements of lynx, and is a measure of the average distance to “non-lynx” habitat within a lynx allometric home range (Figure 4c). Throughout North America, daily movements of lynx have averaged 5-10 km depending on hare densities and season (Nellis and Keith 1968, Brand et al. 1976, Parker et al. 1983, Koehler 1990), and in Washington, they have demonstrated a reluctance to cross openings (e.g., fields, clear-cuts, developments) with little cover that are $> 91$ m across (Koehler 1990, Koehler and Brittell 1990). These potential “barriers” could disrupt movements between foraging and denning sites (Koehler and Aubry 1994), so this habitat model component relates the amount of “non-lynx” habitat interspersion throughout the landscape to habitat quality. An average distance of $> 1500$ m to the nearest non-lynx habitat within a lynx allometric home range provides the most suitable conditions, while an average distance $< 300$ m provides non-suitable conditions. Pixels are assigned non-lynx habitat if they are identified as having no suitability for denning or
foraging and do not provide adequate cover for travel. Inadequate cover includes human developments, water bodies, permanent openings and vegetation types having vegetation <2 m tall that are >91 m from suitable habitat (Roloff 2003). Forest patches with <440 trees ha\(^{-1}\) and <50% horizontal cover that are >91 m in length also qualify as non-lynx habitat. The hypothesized effect of habitat interspersion on lynx should be interpreted carefully, since this relationship is unknown (Roloff 2003). The resulting grid map of habitat interspersion describes the general contiguity of lynx habitat at the scale of a lynx home range.

**Final Habitat Potential**

The three components previously described are combined using a geometric mean to provide an index to the overall habitat suitability for lynx. The use of a geometric mean results in zero habitat suitability for areas that are unsuitable in any one or more of the habitat components, since each component is determined to be important for lynx survival (Koehler and Aubry 1994, Roloff 2003). The grid map of lynx habitat suitability is used to create hypothetical home ranges for lynx with the Plume (2001) GIS function, in the same way home ranges are created for hare. A habitat unit goal based on the allometric home range for lynx (250 ha) supports the simulation of viable, marginal, and non-viable lynx home ranges across the landscape (Roloff and Haufler 1997). Viable lynx home ranges meet a minimum quality threshold of 70, which is based on a relationship between lynx home range sizes and habitat quality, as indexed by fitness indicators (e.g., survival, litter size, pregnancy rate) (see Figure 3: Roloff and Haufler 1997) demonstrated in previous field studies. The threshold represents the point at which lynx home range size no longer increases rapidly with decreased habitat quality, and
fitness indicators shift from good to bad (Roloff and Haufler 1997). Marginal lynx home ranges have an average quality $\geq 25$ and $< 70$, and represent areas where lynx may persist and reproduce, but would be vulnerable to temporal fluctuations in resource availability (Roloff and Haufler 1997). The number and spatial configuration of these hypothetical home ranges would provide insight into the habitat potential for lynx in the region.

Prior to evaluating habitat potential with the lynx model, the composition and structure of vegetation types needed to be quantified and mapped across the landscape for the assessment of habitat suitability. Roloff (2003) suggested the use of an ecological land classification that described biotic and abiotic factors of the landscape to properly stratify the variation in vegetation attributes defining habitat quality in the HSI models.

**ECOLOGICAL LAND CLASSIFICATION FOR THE UPPER PENINSULA**

The ecological land classification used in this study was based upon the “Habitat Classification System” constructed by Coffman et al. (1984), describing “habitat types” (Daubenmire 1966) of the UP that were defined by associations of tree species occurring within similar ecological conditions that had the potential to support similar successional trajectories and climax vegetation communities. The ecological conditions were characterized by a specific range of variation in abiotic factors (e.g., climate, landform, soils) that affect the composition and structure of vegetation within the various seral stages that are initiated by disturbance or developed through succession (Kotar and Burger 2000). The habitat type concept has been used in the Great Lakes region (Coffman et al. 1984, Kotar et al. 1988, Kotar and Burger 2000, Felix et al. 2004) to guide forest and wildlife management strategies by improving land-cover and forest-type classifications with metrics of site potential and an understanding of community
dynamics. Using the Coffman et al. (1984) habitat types as a basis for the ecological land classification facilitated the stratification of vegetation attributes necessary for the lynx model into ecologically meaningful and homogenous classes that were developed specifically for the forest communities of the UP.

Coffman et al. (1984) described 21 habitat types for the UP, covering a gradient of soil texture and drainage classes that can influence the composition and dominance of tree species throughout seral stages and into late succession. Habitat types were named by combining the genera of the dominant climax species with that of an understory indicator species. Coffman et al. (1984) provided a detailed description for each habitat type, including the landform and soil characteristics on which it occurs, the effects of silvicultural practices on species composition at the start of secondary succession, site indices for the primary species within each seral stage, and one or more specific trajectories outlining the canopy replacement of seral species by shade-tolerant species. Coffman et al. (1984) aggregated habitat types into more general categories denoted as “series” that were defined and labeled by the genera of the dominant climax species; these categories were further aggregated (though not mutually exclusively) into series groups based on soil moisture, as determined by texture and drainage properties. In summary, each of the 21 original habitat types belonged to 1 of 8 series, and each series belonged to 1 or more of 3 series groups, depending on the range of soil conditions that could be dominated by a particular series. Group I incorporated 3 series associated with sandy soils retaining little moisture and experiencing periodic drought: the Pinus series, Acer-Quercus series, and Tsuga series; Group II incorporated 4 series associated with soils with adequate moisture and fine textures ranging from loamy sand to clay loam: the
Acer-Quercus series, Tsuga series, Acer-Tsuga series, and Acer series; Group III incorporated 5 series associated with wet mineral or organic soils with impeded or variable drainage: the Tsuga series, Acer-Tsuga series, Tsuga-Thuja series, Fraxinus series, and Picea series (Coffman et al. 1984). Each of the 3 categorizations (habitat types, series, series groups) was meant to simplify a continuum of ecological conditions.

I condensed the 21 original habitat types into 8 classes (hereafter, “habtypes”), based on combinations of similar series and series groups (Table 1). Each of the 8 habtypes contained several early successional stages with a variety of potential overstory compositions (Table 2). The PiVa habtype consisted of habitat types identified by Coffman et al. (1984) that were dominated by jack pine and red pine on sandy outwash soils with excessive drainage. The QuAc habtype consisted of red oak/red maple dominated types that had potential early stages of aspen/birch, pine, or spruce/fir occurring on sandy soils with good drainage. The TsMa habtype consisted of hemlock/maple dominated types with early stages of aspen/birch, pine, or spruce/fir, occurring on well-drained sands and sandy loams. The Acer habtype consisted of sugar maple dominated types with early stages of aspen and northern hardwoods (i.e., beech, basswood) occurring on rich loamy soils with good drainage. The TsTh-dry habtype consisted of hemlock/cedar dominated types with early stages of aspen/birch, spruce/fir and maples occurring on clays and loams with moderate to somewhat poor drainage. The Frax habtype consisted of black ash dominated types with early stages of seral hardwoods occurring on clays and loams with poor drainage. The TsTh-wet habtype consisted of cedar/hemlock dominated types with early stages of seral hardwoods and spruce/fir occurring on a variety of poorly drained soils. Finally, the Picea habtype consisted of
Table 1. Habitat types originally described by Coffman et al. (1980) for the Upper Peninsula of Michigan with associated soil properties and “habtype” groupings.

<table>
<thead>
<tr>
<th>Habitat Type (^a)</th>
<th>Soil (drainage b/texture)</th>
<th>Habtype</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pinus-Vaccinium-Deschampsia</td>
<td>E sand</td>
<td>PiVa</td>
</tr>
<tr>
<td>Pinus-Vaccinium-Carex</td>
<td>E sand</td>
<td></td>
</tr>
<tr>
<td>Quercus-Acer-Epigaea</td>
<td>W sand</td>
<td>QuAc</td>
</tr>
<tr>
<td>Acer-Quercus-Vaccinium</td>
<td>W sand</td>
<td></td>
</tr>
<tr>
<td>Tsuga-Maianthemum-Vaccinium</td>
<td>W loamy-sand</td>
<td>TsMa</td>
</tr>
<tr>
<td>Acer-Tsuga-Dryopteris</td>
<td>W loam</td>
<td>Acer</td>
</tr>
<tr>
<td>Acer-Viola-Osmorhiza</td>
<td>W loam</td>
<td></td>
</tr>
<tr>
<td>Acer-Osmorhiza-Caulophyllum</td>
<td>W silt loam</td>
<td></td>
</tr>
<tr>
<td>Tsuga-Acer-Mitchella</td>
<td>MW clay</td>
<td></td>
</tr>
<tr>
<td>Tsuga-Thuja-Lonicera</td>
<td>MW clay</td>
<td></td>
</tr>
<tr>
<td>Tsuga-Thuja-Petasites</td>
<td>SP clay</td>
<td></td>
</tr>
<tr>
<td>Tsuga-Maianthemum-Coptis</td>
<td>SP loam</td>
<td></td>
</tr>
<tr>
<td>Fraxinus-Impatiens</td>
<td>P loam/clay</td>
<td>Frax</td>
</tr>
<tr>
<td>Fraxinus-Mentha-Carex</td>
<td>P loam</td>
<td></td>
</tr>
<tr>
<td>Fraxinus-Eupatorium</td>
<td>P clay</td>
<td></td>
</tr>
<tr>
<td>Tsuga-Thuja-Mitella</td>
<td>VP sand/loam</td>
<td>TsTh-wet</td>
</tr>
<tr>
<td>Tsuga-Thuja-Sphagnum</td>
<td>VP organic</td>
<td></td>
</tr>
<tr>
<td>Picea-Osmunda</td>
<td>VP deep organic</td>
<td>Picea</td>
</tr>
<tr>
<td>Picea-Chamadaphne-Sphagnum</td>
<td>VP deep organic</td>
<td></td>
</tr>
</tbody>
</table>

\(^a\) The *Acer-Quercus-Viburnum* habitat type was not included, due to its minor extent in the Upper Peninsula.

\(^b\) Soil drainage codes: E = excessively drained; W = well-drained; MW = moderately well-drained; P = poorly drained; VP = very poorly drained.
Table 2. Dominant tree species assemblages within seral stages of each habtype for the Upper Peninsula of Michigan.

<table>
<thead>
<tr>
<th>Habtype</th>
<th>Early-seral</th>
<th>Mid-seral</th>
<th>Late</th>
</tr>
</thead>
<tbody>
<tr>
<td>PiVa</td>
<td>jack pine</td>
<td>jack pine</td>
<td>jack pine, red pine</td>
</tr>
<tr>
<td>QuAc</td>
<td>(1) aspen, white birch, (2) jack, red pine</td>
<td>(1) white pine, (2) white spruce, fir</td>
<td>red oak, red maple</td>
</tr>
<tr>
<td>TsMa</td>
<td>(1) aspen, white birch, (2) jack, red pine</td>
<td>(1) white pine, (2) white spruce, fir</td>
<td>hemlock, maple, beech, yellow birch</td>
</tr>
<tr>
<td>Acer</td>
<td>aspen</td>
<td>sugar maple, basswood, white ash, beech</td>
<td>sugar maple, hemlock</td>
</tr>
<tr>
<td>TsTh-dry</td>
<td>aspen, white birch, balsam poplar</td>
<td>(1) white spruce, fir, (2) maple</td>
<td>hemlock, cedar, fir, maple</td>
</tr>
<tr>
<td>Frax</td>
<td>aspen, white birch, balsam poplar, red maple</td>
<td>black ash, red maple</td>
<td>black ash, red maple</td>
</tr>
<tr>
<td>TsTh-wet</td>
<td>white birch, balsam poplar, red maple</td>
<td>cedar, fir, black spruce, tamarack</td>
<td>cedar, hemlock, fir</td>
</tr>
<tr>
<td>Picea</td>
<td>lowland brush, white birch, red maple</td>
<td>black spruce, tamarack, cedar, fir, jack pine</td>
<td>black spruce, tamarack</td>
</tr>
</tbody>
</table>

a Seral stages are not equally distributed through time across habtypes. Numbered species groups within seral stages represent different potential trajectories of succession.
black spruce/tamarack dominated types with early stages of cedar and fir occurring on very poorly drained organic soils and peatlands. The combination of a habtype and seral stage (as defined by overstory species) resulted in an ecological land unit (ELU), which provided the basis of the ecological land classification necessary for the lynx model.

The classification system developed by Coffman et al. (1984) was intended as a field guide for in situ classification of forest stands that could improve the predictions of forest management prescriptions; it did not involve mapping habitat types across the landscape. Mapping the spatial configuration and extent of the ELUs across the UP required the integration of multiple spatial data layers within a GIS. Each data layer represented components of the ecological conditions that could identify the occurrence of a habtype, and within each habtype, a seral stage. My objectives included the assessment of current and presettlement habitat conditions for lynx, to examine how forest composition and structure effected lynx habitat potential within each time period. Theoretically, habtypes should be similar between time periods where certain ecological conditions (e.g., soils) had not been measurably altered; however, at any given point in time the distribution of seral stages within habtypes will be dictated by forest disturbances (Kotar and Burger 2000). The current distribution of seral stages results from a disturbance history with considerable human influence (e.g., fire suppression and exclusion, development, harvesting), and is very different from the “shifting mosaic” of seral stages that historically occurred in the Great Lakes region due to natural disturbances (Zhang et al. 1999, Frelich 2002). Therefore, it was necessary to incorporate different data layers and develop separate methods to map the location and quantify the vegetation of ELUs for current and presettlement landscapes.
Current Ecological Conditions

A combination of digital spatial data and region-wide forest inventory surveys was used to delineate ELUs and quantify the current species composition and structure of forest types in the UP. Thematic spatial data included: (1) current vegetation, to separate forested from non-forested areas and describe the species composition of the overstory; (2) presettlement vegetation, which indicated potential natural communities in the absence of human disturbance; (3) soils, to provide a context for identifying habitat types; and (4) ecoregions, which defined broad ecological units having similar climate and physiography. The intersection of these thematic GIS layers created the framework for identifying ELUs on the landscape. Survey data provided plot-level measurements of stand composition and structure within each ELU to quantify specific vegetation attributes required by the lynx model. A final data layer consisted of unclassified satellite imagery that was used with the survey data to model forest structure as a continuous variable across the landscape and account for variation within ELUs. Spatial data were compiled and manipulated using GIS software which included ARC/INFO 8.1 and ArcView 3.2 (Environmental Systems Research Institute, Redlands, Calif.) and ERDAS Imagine 8.7 (ERDAS, Inc). All thematic spatial data were projected in the Michigan GeoRef Coordinate System. The ecological land classification was created in a raster-based format to facilitate the calculation of habitat units within a GIS; habitat quality for hare was assessed with a grid resolution of 30 m, while that for lynx was assessed with a grid resolution of 90 m. The multiple data sources that were collected and the methods that were employed to utilize them as input for the lynx model included the following:
Current vegetation – I obtained a land-cover dataset publicly available from the Michigan Department of Natural Resources (MDNR) which had been created for their Integrated Forest Monitoring, Assessment and Prescription (IFMAP) project in cooperation with the GAP analysis program (MDNR 2001, Donovan 2005). The IFMAP land cover was developed using supervised and unsupervised classification of 3 seasons (spring, summer, fall) of Landsat satellite imagery collected at a 30 m resolution between 1999-2001. Forested pixels were defined to have ≥25% canopy cover; forest composition was defined according to the dominant species (≥60% cover) in the canopy, with “mixed” categories for stands without dominant species. Land cover was classified to Anderson Level II with an estimated accuracy of 87%, and for some classes Level III, with an estimated accuracy between 37-87% (Donovan 2005).

Subbedi (2005) found large inaccuracies in the IFMAP prediction of some Anderson Level III categories, especially with aspen and oak, across public lands in the UP. My own accuracy assessment of the map revealed similar results (Appendix C); therefore, most Anderson Level III categories were condensed to Level II for my analysis (Table 3). Because mixed forests in the UP generally succeed to hardwoods in the uplands and conifers in the lowlands, each of the mixed forest cover types was grouped into the appropriate condensed category. The decision to group the mixed forest cover types in this manner was supported by findings from the accuracy assessment.

Although the loss of detail made it difficult to distinguish canopy species, the IFMAP layer represented the most recent land-cover data for the UP, and the Anderson Level II categories that provided broad classes of overstory composition were accurate. The accuracy and resolution of this dataset made it the most reliable spatial layer for
Table 3. Areas of land-cover classes identified by the 2001 IFMAP in the Upper Peninsula of Michigan.

<table>
<thead>
<tr>
<th>IFMAP land cover</th>
<th>Condensed</th>
<th>Area (ha)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low/High Intensity Urban</td>
<td>Developed</td>
<td>76,603</td>
</tr>
<tr>
<td>Airports</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Road/Parking Lot</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-Vegetated Farmland</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Row Crops</td>
<td>Agriculture</td>
<td>126,628</td>
</tr>
<tr>
<td>Forage Crops/Non-Tilled</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Orchards/Vineyards/Nursery</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Parks/Golf Courses</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sand, Soil, Exposed Rock</td>
<td>Non-vegetated</td>
<td>29,271</td>
</tr>
<tr>
<td>Mud Flats</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Other Bare/Sparsely Vegetated</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Herbaceous Openland</td>
<td>Openland</td>
<td>160,842</td>
</tr>
<tr>
<td>Upland Shrub/Low Density Trees</td>
<td>Upland shrub</td>
<td>84,067</td>
</tr>
<tr>
<td>Northern Hardwood Association</td>
<td>Upland deciduous</td>
<td>1,929,651</td>
</tr>
<tr>
<td>Oak Association</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aspen Association</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mixed Upland Deciduous</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Upland Mixed Forest</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pines</td>
<td>Upland coniferous</td>
<td>509,131</td>
</tr>
<tr>
<td>Other Upland Conifers</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mixed Upland Conifers</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Water</td>
<td>Water</td>
<td>239,319</td>
</tr>
<tr>
<td>Lowland Deciduous Forest</td>
<td>Lowland deciduous</td>
<td>1,562</td>
</tr>
<tr>
<td>Lowland Coniferous Forest</td>
<td>Lowland coniferous</td>
<td>708,289</td>
</tr>
<tr>
<td>Lowland Mixed Forest</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lowland Shrub</td>
<td>Lowland shrub</td>
<td>250,428</td>
</tr>
<tr>
<td>Floating Aquatic</td>
<td>Wetland</td>
<td>206,608</td>
</tr>
<tr>
<td>Emergent Wetland</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mixed Non-Forest Wetland</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
identifying landscape attributes; for this reason it was used to describe current vegetation and correct any inconsistencies in the descriptions of ecological conditions resulting from the intersection of other spatial layers.

*Presettlement vegetation* – I obtained a spatial layer of presettlement vegetation that was constructed by Comer et al. (1995) using GLO surveyor notes recorded between 1816 and 1856, before the extensive logging that took place during the late 1800s. This data layer provided information on forest types that occurred prior to European settlement, and thus, described potential species assemblages determined by natural disturbance and physiography, in the absence of significant human influence (Table 4). The map was created by interpretation of recorded landscape descriptions for each 2.56 km$^2$ section and occurrences of tree species tallied at and between 1.6 km section corners (and 0.8 km quarter corners) (Comer et al. 1995). Comer et al. (1995) combined this information with ancillary GIS data (e.g., landforms, wetlands, topography) to delineate forest cover types.

The exact spatial accuracy of the map is unknown and interpolated boundaries between section corners and quarter corners were expected to contain some error where ancillary data were unable to distinguish actual borders between cover types. Also, the potential bias of surveyors for long-lived species (e.g., beech) (Manies et al. 2001), combined with the scale of survey data which can be expected to miss small (< 20 ha) wetlands and wind-throw disturbances, has most likely resulted in the under-representation of seral communities (Comer et al. 1995). Regardless of these potential inaccuracies, the cover classes delineated by the presettlement layer identified the dominant species within many of the forest types (Table 4), and provided more detailed
Table 4. Area of cover types in the presettlement vegetation layer from Comer et al. (1995) located in the Upper Peninsula of Michigan.

<table>
<thead>
<tr>
<th>Cover Type</th>
<th>Area (ha)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Upland Non-forested</strong></td>
<td></td>
</tr>
<tr>
<td>Exposed bedrock</td>
<td>3,650</td>
</tr>
<tr>
<td>Sand dune</td>
<td>1,266</td>
</tr>
<tr>
<td>Grassland</td>
<td>37</td>
</tr>
<tr>
<td>Oak/Pine Barrens</td>
<td>387</td>
</tr>
<tr>
<td>Pine Barrens</td>
<td>25,556</td>
</tr>
<tr>
<td><strong>Upland Forested</strong></td>
<td></td>
</tr>
<tr>
<td>Aspen/Birch</td>
<td>72,085</td>
</tr>
<tr>
<td>Beech/Sugar Maple/Hemlock</td>
<td>533,844</td>
</tr>
<tr>
<td>Sugar Maple/Hemlock</td>
<td>939,430</td>
</tr>
<tr>
<td>Sugar Maple/Basswood</td>
<td>86,216</td>
</tr>
<tr>
<td>Sugar Maple/Yellow Birch</td>
<td>383,908</td>
</tr>
<tr>
<td>Hemlock/Yellow Birch</td>
<td>118,957</td>
</tr>
<tr>
<td>Hemlock/White Pine</td>
<td>215,463</td>
</tr>
<tr>
<td>White Pine/Mixed Hardwood</td>
<td>16,430</td>
</tr>
<tr>
<td>Mixed Pine/Oak</td>
<td>6</td>
</tr>
<tr>
<td>White Pine/Red Pine</td>
<td>129,552</td>
</tr>
<tr>
<td>Jack Pine/Red Pine</td>
<td>81,117</td>
</tr>
<tr>
<td>Spruce/Fir/Cedar</td>
<td>360,584</td>
</tr>
<tr>
<td><strong>Lowland Non-forested</strong></td>
<td></td>
</tr>
<tr>
<td>Wet Prairie</td>
<td>3,909</td>
</tr>
<tr>
<td>Shrub swamp/Emergent marsh</td>
<td>9,290</td>
</tr>
<tr>
<td>Muskeg/Bog</td>
<td>127,946</td>
</tr>
<tr>
<td>Lake/River</td>
<td>985,918</td>
</tr>
<tr>
<td><strong>Lowland Forested</strong></td>
<td></td>
</tr>
<tr>
<td>Black Ash</td>
<td>243</td>
</tr>
<tr>
<td>Mixed Hardwood</td>
<td>55,598</td>
</tr>
<tr>
<td>Cedar</td>
<td>104,552</td>
</tr>
<tr>
<td>Mixed Conifer</td>
<td>94,182</td>
</tr>
</tbody>
</table>
information concerning potential species assemblages than that of the IFMAP layer. The GLO data have proven useful throughout the Great Lakes region as a means to reconstruct natural disturbance regimes and to compare current forest conditions with those of the presettlement era (Frelich 1995, Zhang et al. 1999, Schulte and Mladenoff 2001, Cleland et al. 2004, Friedman and Reich 2005).

The presettlement layer was combined with the IFMAP layer after being converted to a raster format, and 2 iterations of reclassification were used to provide the most informative description of the vegetation. The first iteration involved reclassifying pixels that described inaccurate ecological conditions (e.g., upland versus lowland), using a nearest-neighbor approach. I assumed the IFMAP layer provided greater accuracy because it was constructed from remotely sensed imagery; therefore, it was given precedence in determining the final attributes for a pixel. For example, if a pixel described current vegetation as upland deciduous and presettlement vegetation as cedar swamp, its presettlement description was reassigned the nearest presettlement cover type that was an acceptable match for upland deciduous. Acceptable matches were based on the distinction between upland and lowland for each layer, with the exception of 2 presettlement cover types: spruce-fir-cedar and aspen-birch. These 2 cover types contained tree species that could tolerate a range of ecological conditions; therefore, pixels belonging to those presettlement cover types that were located on a lowland cover type for the IFMAP layer were not adjusted, in spite of the fact that their presettlement classification was originally upland (Table 4) (Comer et al. 1995). Non-forested presettlement pixels were assigned the nearest forested pixel value if the IFMAP cover type was forested.
Visual inspection of grid overlays indicated that pixels located between upland and lowland cover type boundaries were responsible for the majority of inconsistencies, and 35% of misclassified pixels, representing 6% of all forested areas, involved a mismatch between upland hardwoods and lowland conifers. This was expected due to the interpolation technique used to create the presettlement layer (Comer et al. 1995) and the heterogeneous landscape of the UP, especially in the central and western portions where a mosaic of drumlins and lowland depressions occur. The second iteration of reclassification involved reassigning pixels that were described as having a presettlement cover type of aspen-birch. This presettlement cover type would not provide useful information for identifying an ELU because aspen and/or birch forest types could occur as a seral stage in nearly every habtype (Table 2). Therefore, a nearest-neighbor approach was again utilized to assign a new presettlement cover type based on acceptable upland/lowland matches within close proximity. The 2 iterations of reclassifying pixels resulted in the most informative combination of data layers, given that I assumed: (1) any contradictions in vegetation were reflecting spatial inaccuracies in the map layers and not actual conditions, and (2) the aspen-birch cover types represented early seral stages and would have been replaced through succession by surrounding forest types.

Soils – The spatial data that were used to describe soils were obtained from the State Soil Geographic (STATSGO) database for Michigan, developed by the Natural Resources Conservation Service (USDA, NRCS 1994). The boundaries of map units in the STATSGO layer are broad groupings of soil associations, which contain the most common soil series that occur together on the landscape. Soil properties were the key factors in the determination of habitat types by Coffman et al. (1984), and aside from
regional differences in climate, would account for most variation in vegetation composition and structure within otherwise similar forest types, given the generally small differences in local topography throughout the Great Lakes region (Kotar and Burger 2000). The information provided by the STATSGO database that was used to identify habtypes for each soil series (component) within a map unit included percent coverage within the map unit, surface texture, drainage, depth to water table, depth to bedrock, common woodland species (with site indices), and understory indicator species (both woody and non-woody). Additional characteristics of each soil series were obtained from the NRCS Official Soil Series Descriptions (OSD), which provided a description of common landforms, typical pedons, and forest vegetation.

Unfortunately, the STATSGO layer was mapped at a coarse scale of 1:250,000, and the NRCS has suggested that its use be limited to interpretation at broad geographic regions (USDA, NRCS 1994). The more detailed data provided by the Soil Survey Geographic (SSURGO) database are currently unavailable for greater than half of the UP (USDA, NRCS 2006), thus, the STATSGO database was the only available source of soil information with associated spatial data that could be integrated into a GIS for my extensive study area. The limitations in the STATSGO soils data were mediated with several approaches.

Each STATSGO map unit contained multiple soil components within its boundary, and each soil component represented a soil series. There were 84 unique map units for the UP with 181 unique soil series; the extents of 15 soil series covered ~50% of the UP, while that of 35 soil series covered ~75%. Map unit polygons ranged in size from 200 ha to 370,000 ha, and consequently, large map units encompassed a variety of
soil series (including both upland and lowland) that were not spatially delineated. The
STATSGO layer was converted to raster format and combined with a reclassified IFMAP
layer that delineated upland from lowland cover types. The resulting grid delineated
STATSGO map units into upland and lowland soils, and a relational database was used to
assign properties from hydric soil components to the lowland soil pixels, and that of
mesic/xeric soil components to the upland soil pixels. The soil information from the map
unit components was interpreted alongside the vegetation described by the IFMAP and
presettlement layers to identify the most probable habtype. Similar to the adjustment of
presettlement cover types, this process resulted in a logical assignment of soil
information to the grid pixels, and avoided the inconsistent descriptions of ecological
conditions that would have occurred if map units were assigned information from only
the most dominant soil component (e.g., upland deciduous on very poorly drained
organic muck).

Ecoregions – A spatial layer of ecoregions was used to separate otherwise similar
habtypes that existed within different climatic and geophysical regions of the UP.
Several ecoregion delineations exist for the Great Lakes region, including those of
Omernik (1987), Albert (1995), Bailey (1995), and Cleland et al. (2005b), and some of
these delineations are continually being refined with increases in the amount and
resolution of available spatial data (Cleland et al. 2005b). I used the ecoregions layer
developed by Cleland et al. (2005b), which depicted the 2 units of the subregion planning
scale (sections and subsections) as outlined by the Forest Service national hierarchical
framework of ecological units (Cleland et al. 1997). The UP contained 21 subsections
that were contained within 7 sections, ranging in size from 20-670 km² and 580-
1,538 km², respectively. Ecoregions were grouped into 2 “subregions” based on their broad-scale location within the UP; sections in the eastern subregion included 212R and 212T, while those in the western subregion included 212J, 212L, 212S, 212X, and 212Y. This ecoregions map was chosen over the others because it was being actively used by the Forest Service (Cleland et al. 2005b) and, consequently, coordinated with other data collected for my study (see following). Also, the locations of ecoregion boundaries were mostly consistent with previously developed maps.

**ELU grids** – The intersection of soils and vegetation data resulted in 4,781 unique combinations of forest cover categories. The high number of unique combinations was a result of the soil layer, which was not consolidated prior to the grid intersection in order to preserve all available information from the STATSGO database. Only one seral stage and ELU could be described for most habtypes due to a lack of decisive information from the vegetation layers, and one ELU (TsTh-Picea) was created to describe areas that could not be split between the two lowland coniferous habtypes. The base ELU grid contained 14 ELUs, with 2 non-forest classes and 12 forest classes (Table 5). The ecoregions layer was used to stratify the forested ELUs into smaller units and create an ecoregion-ELU grid with 220 ecoregion-ELUs. Each grid value contained the ELU and subsection identification (e.g., PiVa-212Rb). The forested classes in the ecoregion-ELU grid ranged in total area from <1 ha to 158,715 ha, with a median area of 4,004 ha (Figure 5).

**Forest plot surveys** – Vegetation attributes necessary for the lynx model were obtained from plot surveys collected by the Forest Inventory and Analysis (FIA) program of the Forest Service (USFS 2006). The FIA program provides detailed measurements of the nation’s forests on 5-10 year cycles (5 years in Michigan), using a double sampling
Table 5. Classes within spatial layers that delineated habtypes and ELU categories in the Upper Peninsula of Michigan.

<table>
<thead>
<tr>
<th>ELU</th>
<th>Habtype</th>
<th>Current Vegetation</th>
<th>Presettlement Vegetation&lt;sup&gt;a&lt;/sup&gt;</th>
<th>STATSGO Mapunit ID&lt;sup&gt;b&lt;/sup&gt; (total count)</th>
</tr>
</thead>
<tbody>
<tr>
<td>N/A</td>
<td>N/A</td>
<td>developed, agriculture, non-vegetated, open-land, water, wetland</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>QuAc2</td>
<td>QuAc</td>
<td>upland deciduous</td>
<td>(see QuAc1)</td>
<td>(see QuAc1)</td>
</tr>
<tr>
<td>TsMa2</td>
<td>TsMa</td>
<td>upland deciduous</td>
<td>(see TsMa1)</td>
<td>(see TsMa1)</td>
</tr>
<tr>
<td>TsTh-dry2</td>
<td>TsTh-dry</td>
<td>upland deciduous</td>
<td>(see TsTh-dry1)</td>
<td>(see TsTh-dry1)</td>
</tr>
</tbody>
</table>
Table 5 (cont’d).

<table>
<thead>
<tr>
<th>ELU</th>
<th>Habtype</th>
<th>Current Vegetation</th>
<th>Presettlement Vegetation&lt;sup&gt;a&lt;/sup&gt;</th>
<th>STATSGO Mapunit ID&lt;sup&gt;b&lt;/sup&gt; (total count)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frax</td>
<td>Frax</td>
<td>lowland deciduous</td>
<td>AS-PB, BA swamp, BS-BF-WC, conifer swamp, hardwood swamp, WC</td>
<td>104, 117, 124, 127, 128, 131, 133-146, 148-210 (84)</td>
</tr>
<tr>
<td>TsTh-wet</td>
<td>TsTh-wet,</td>
<td>lowland coniferous</td>
<td>BA swamp, BS-BF-WC, conifer swamp, hardwood swamp, WC</td>
<td>104, 117, 124, 127, 131, 133-135, 137-152, 154-182, 184, 185, 187-209 (79)</td>
</tr>
<tr>
<td></td>
<td>Picea</td>
<td></td>
<td></td>
<td>143, 165 (2)</td>
</tr>
<tr>
<td>Picea</td>
<td>Picea</td>
<td>lowland coniferous</td>
<td>BA swamp, BS-BF-WC, conifer swamp, hardwood swamp</td>
<td>104, 124, 127, 128, 131, 133, 134-142, 144, 145, 147-164, 166-210 (81)</td>
</tr>
</tbody>
</table>


<sup>b</sup> Mapunit IDs are consistent with those found in the Michigan STATSGO database.
Figure 5. Spatial distribution of ecological land units (ELUs) in the Upper Peninsula of Michigan with subsection boundaries from the Cleland et al. (2005b) map of ecoregions.
for stratification approach (Smith 2002). Remote sensing imagery is used to identify forested land across a state, which is then systematically field sampled within a hexagonal grid of 5 interlocking panels, with one plot survey for every 2,428 ha (6,000 ac). The state of Michigan provided funding to increase the intensity of field sampling and essentially triple the number of plot surveys conducted, reducing the area represented by each plot survey to 809 ha (2,000 ac). The plot surveys consist of a 4-subplot cluster, each subplot having a fixed radius of 7.32 m, within which trees having a diameter at breast height (dbh) $\geq 12.7$ cm (5.0 in) are individually measured. All trees with a dbh $< 12.7$ cm are counted within a smaller fixed radius (2.07 m) microplot, and only those with a dbh $\geq 2.54$ cm (1 in) are individually measured. Data that are recorded for individually measured trees include species, dbh, height, crown ratio, and crown class (e.g., dominant, co-dominant), while seedlings (dbh $< 2.54$ cm) are simply tallied by species. Landscape attributes that define one or more “conditions” (e.g., land use, forest type, physiographic class, stand density, stand size) are also recorded for each plot.

The plot data that were available for Michigan at the beginning of my study included information for 80% of the FIA plots in the 6th cycle (4/5 sub-cycles), collected between 2000-2003. Data from the final sub-cycle (2004) became available in December 2005, resulting in a total of 4,741 plots with recorded tree data for the UP. Plot data were downloaded from the FIA website (http://fia.fs.fed.us/) in two different formats: (1) spreadsheets that could be imported to a relational database; and (2) text files that were formatted for use with the Forest Vegetation Simulator (FVS). The FVS is a framework used by the Forest Service to standardize forest growth and yield modeling, with variant growth models calibrated for specific regions (Dixon 2002). I used FVS to calculate the
per hectare stand attributes required for the HSI models (e.g., basal area, average dbh, stem density, canopy closure) from the FIA data.

The FIA data were filtered to remove plots that could not be used with my mapping methodology. Plots with <25% canopy closure were removed to ensure that vegetation attributes from the FIA data corresponded to the IFMAP definition of forested cover. Plots with >1 landscape condition were removed to ensure that survey data were describing homogenous samples that could be linked to single ELUs. This filtering process reduced the number of available plots by ~30%, leaving 3,256 plots for my analysis. The frequency distributions of stand attributes necessary for the model were compared between the original and filtered datasets to verify that no bias was introduced through the filtering process.

The stand attributes required for the snowshoe hare sub-model included horizontal cover and understory dominance. Understory dominance was calculated by the percentage of conifer stems among stem densities of trees with a height to crown <3 m. Horizontal cover was not measured in the FIA surveys, so a new method was developed to extract an estimate from the data that were gathered (Appendix D). A post-processing function in FVS was used to create tree-lists for the Stand Visualization System (SVS) (McGaughey 1997), a program that creates 3-dimensional diagrams of 0.40 ha forest plots based on survey data, with each tree rendered according to its attributes (e.g., dbh, height, species). Plant form definitions are used to define appearances for different species and can be controlled by the user; I altered plant form definitions in SVS by removing leaves from deciduous species to simulate winter conditions. For each plot, the profile-view diagram in SVS was saved and converted to a 1-bit (black and white)
bitmap. The proportion of black to white pixels within each height strata determined the horizontal cover for a given plot (Appendix D).

The method of calculating horizontal cover with SVS was complicated by the absence of size measurements for seedlings, which would have a significant influence on estimates of horizontal cover. The Forest Service assigns a common dbh value of 0.25 cm (0.1 inch) to all seedlings in the FVS-ready files, for the purposes of growth modeling. I felt that a common dbh of 0.25 cm did not accurately reflect the true variation in seedling sizes, given that seedlings were defined as any tree with a dbh < 2.54 cm. Large seedlings (e.g., dbh ~ 2.0 cm) had the potential to provide more horizontal cover than small ones (e.g., dbh ~ 0.25 cm), depending on species, and the absence of this distinction could result in conservative estimates of horizontal cover. Therefore, I assigned each plot a frequency distribution of seedling dbh values that fit a negative exponential trend and ranged 0.25-2.29 cm (0.1-0.9 inches) in increments of 0.25 cm (0.1 inches). This procedure resulted in a high ratio of small to large seedlings, which I hypothesized was a more adequate representation of the vegetation. Original plot data were retained to examine the effect of manipulating the seedling size distribution on measurements of horizontal cover. Seedling heights were determined by FVS using dbh-height models that were specific to each species.

The uncertainty in my estimates of horizontal cover warranted an additional measure of hare habitat quality from the plot data. Litvaitis et al. (1985) described a strong relationship between estimated stand densities of snowshoe hares (indexed by live-trapping and pellet counts) and “stem cover units” (SCUs), calculated as the count of trees and shrubs < 7.5 cm dbh and > 0.5 m tall, with coniferous stems weighted by a factor
of 3. The weighting factor for conifers was based on the estimated difference in visual obstruction provided by coniferous stems and deciduous stems (Litvaitis et al. 1985). I used Litvaitis et al.’s (1985) linear regression model of hare density as predicted by SCUs to create a production function for hare habitat quality (Figure 6). Hare habitat quality was highest at $\geq 55,652$ SCUs/ha and lowest at $<23,043$ SCUs/ha, corresponding to the maximum and minimum densities of hares in the regression model (Litvaitis et al. 1985). This alternative index defined hare habitat quality in a similar manner to the indices in the original hare sub-model (Roloff 2003); horizontal cover and SCUs were essentially measurements of the same stand attribute, and the presence of conifers increased quality in both indices. In summary, hare habitat quality was measured in 3 separate ways: an HSI similar to that of Roloff (2003) which combined understory dominance with SVS estimates of horizontal cover using manipulated seedling distributions (HSI-1) and original seedling distributions (HSI-2); and an HSI using the regression equation from Litvaitis et al. (1985) measuring stem cover units (HSI-3).

*Linking plots to ELU grid* – The actual coordinates of FIA plot locations were mandated confidential by Congress in 2000, in accordance with amendments to the Food Security Act of 1985, to ensure that private landowners are not identified with plots and that long-term integrity of plot data is maintained. The use of FIA plot locations by outside researchers is facilitated through the Spatial Data Services (SDS) for each FIA regional unit. I cooperated with Geoff Holden at the North Central Research Station SDS (USFS, St. Paul, Minnesota) to have the FIA plot locations overlaid with the spatial data layers. Confidentiality concerns over the amount of detail and the multiple iterations of the ELU grid restricted the amount of information that could be disclosed. Also, the final
Figure 6. Production function of hare habitat quality as predicted by stem cover units. Hare density was scaled to index habitat quality, and the regression model from Litvaitis et al. (1985) was used to determine the slope of the relationship with stem cover units (for $y > 23.04$ and $< 55.65$, $y = 3.067x - 70.667$).
20% of the FIA survey data was released after SDS had completed the analysis. For these reasons, the overlay of FIA plot locations by SDS was used only to examine the accuracy of the land cover map (Appendix C), and in modeling forest structure using satellite imagery (see following section).

The link between FIA plots and the ELU grid was established by matching the ecological conditions described by each data source. Determinant variables for the plot data included physiographic class and species composition of the canopy and sub-canopy, which were used to identify the most probable of the 12 ELUs (Table 5) for each plot. Species composition was defined by FVS calculations of species crown-cover within 3 canopy strata that I hypothesized would provide the best indication of ELU: (1) open grown, dominant, and co-dominant trees; (2) intermediate and overtopped trees; and (3) seedlings. Plot estimates of cover were analyzed with TWINSPAN (Hill 1979, Hill and Šmilauer 2005), a program that conducts 2-way indicator species analysis using ordination and provides a hierarchical classification of sample plots according to species abundance and site fidelity. Species compositions were summarized by TWINSPAN group, and each plot was assigned an ELU based on an interpretation of its TWINSPAN group and physiographic class.

Ecoregion subsection (identified in the FIA database) was used to further stratify the plots to match the ecoregion-ELU grid, and subsection ELUs that did not contain ≥10 FIA plots were aggregated by section or subregion, depending on the number of available FIA plots. For example, the PiVa ELU had >10 FIA plots within section 212R, but <10 FIA plots within subsection 212Rb. Therefore, all grid values belonging to PiVa-212Rb were aggregated at the section level to become PiVa-212R. The PiVa ELU also had <10
FIA plots within section 212S, resulting in those grid values being aggregated at the subregion level to become PiVa-212west. In this way, ecoregion-ELUs were required to contain ≥10 samples for adequate measurements of stand attributes used in the lynx model. The final grid of condensed ecoregion-ELUs contained 2 non-forested classes (non-forest and shrub) and 134 forested classes (Table 6); 88% of forested land was described by ELUs aggregated at the subsection level, while 75% and 50% was described by subsection ELUs with >20 and >60 FIA plots, respectively.

The transfer of stand attributes from the plot data to the grid classes involved the calculation and assignment of mean values, and the random simulation of a value’s spatial distribution within ecoregion-ELUs, to account for the local variation that would otherwise have been masked by the assignment of a mean. The attributes that were spatially simulated included each of the HSI measurements for hare habitat quality, since the scale of habitat selection by hares (4.5 ha) would be most affected by local heterogeneity. The local variation of other stand attributes necessary for the lynx model, including the denning and interspersion variables (e.g., basal area, average dbh, canopy cover, stem density), were assumed to have little effect on lynx habitat selection due to the larger scale of analysis (250 ha). Consequently, these attributes were averaged for each ecoregion-ELU.

I simulated the spatial variation of hare habitat quality for each of the 3 HSI estimates by randomly assigning pixels within each ecoregion-ELU into 1 of 8 quantile classes. Two of the quantile classes represented the upper and lower 5% of the distribution of HSI values from the FIA plots for an ecoregion-ELU, and 6 represented 15% intervals between. The median values were calculated for each quantile class and
Table 6. Forested classes from condensed ecoregion-ELU grid for the Upper Peninsula.

<table>
<thead>
<tr>
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<th>Subregion</th>
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Subsection codes are combined with section codes to be consistent with those found in the ecoregions map constructed by Cleland et al. (2005b).
assigned to the corresponding pixels to create the map of hare habitat quality. In this way, grid pixels would have a similar proportion of HSI values on the landscape to that as measured by the plot data, within each ecoregion-ELU. I examined the effects of the random pixel assignment on lynx model output with a sensitivity analysis. Hare habitat quality (indexed by SCUs) was randomly assigned to pixels within 20 separate grids of the UP. Mean quality within a 4.5-ha moving window was calculated for each, and a grid of the standard deviation for each pixel was created to identify a sample area (1,200 km²) with high variability. This sample area was input to the lynx model for each of the 20 grids, and 3 grids were chosen to be assessed by the lynx model across the entire study area. These 3 grids represented the range in outputs calculated for the smaller sample area, and covered the variability in hare habitat quality and subsequent lynx foraging quality that resulted from the random pixel assignment.

Non-forested ELUs were not represented by FIA plot data and were assumed to provide no habitat to hare and lynx, with the exception of upland and lowland shrub cover types, which were assigned a mid-range HSI score of 50 for hare. These areas could provide refugia for hare, especially in lowland thickets of alder and willow, but they would not qualify as habitat for lynx, based on the lack of overhead cover. Therefore, the effect of the arbitrary assignment of quality for shrub cover types was assumed to have little effect on the estimation of lynx habitat potential.

*Satellite imagery* – I obtained spectral imagery taken by the Enhanced Thematic Mapper Plus (ETM+) of the Landsat 7 satellite, with the intentions of predicting a continuous surface of stand attributes to improve upon the mapping of local variation provided by the random simulations (Appendix E). Eight images were required to cover
the entire UP, with dates of acquisition ranging between late summer and early fall of 2000-2002. Images were rectified and georeferenced to zone 16 of the UTM system, and mosaics were created where possible. The FIA data that was available at the time of analysis included surveys conducted between 2000-2003 for the 6th inventory of Michigan, resulting in 3,809 forested plots. A portion of the modeling was conducted by Geoff Holden at the North Central SDS due to the confidentiality of FIA data.

The FIA plot locations were overlaid with the spectral imagery and a $k$-nearest neighbor (KNN) analysis was used to impute plot attributes continuously across the landscape. The KNN method assigns attribute values to non-sampled pixels from those that are sampled (i.e., located underneath a ground plot) based on the distance between the pixels in a multi-dimensional feature space defined by the combination of spectral values from each wavelength band. The analysis was performed using functions within ERDAS Imagine; a 3x3 mean filter was applied to the spectral imagery, of which bands 1-5 and 7 were used with $k = 5$. The prediction accuracy of the KNN model was estimated by holding out 40% of the FIA plots from the training set, and comparing the observed attribute values with those predicted by the model.

The estimated accuracy of the KNN model was poor throughout the UP (Appendix E); as a result, I did not use the stand attribute maps for modeling lynx habitat. The proportion of explained variation ($R^2$) in the predictions ranged from 0.01 to 0.22, with most of the UP having an $R^2$ below 0.10 (Appendix E). The poor accuracy could have been caused by several factors, including limitations in the availability of cloud-free Landsat 7 imagery for the entire region, the use of one season of imagery as opposed to multiple seasons, the sampling intensity of FIA plots, and the lack of a relationship
between spectral values and stand attributes. Several alternative approaches to the KNN method, including the use of generalized additive models (Frescino et al. 2001), may have yielded better predictions of forest structure. Constraints on the access to FIA plot locations and the amount of time available prevented any further analyses.

**Presettlement Ecological Conditions**

The range of conditions that existed in the UP during the centuries prior to European settlement was estimated using spatial data that described ecological characteristics of the landscape and a simulation model that reproduced the effects of major disturbance on stand conditions. The presettlement forests exhibited a “shifting mosaic” of seral stages that was driven by disturbance and succession (Frelich 2002); therefore, it was my objective to capture a range of forest conditions, and resultant lynx habitat potential, provided by the landscape under a natural disturbance regime.

Habtypes were mapped across the UP by combining data from the STATSGO soil layer and the original presettlement vegetation map from Comer et al. (1995), using similar methods to those applied for current conditions (Table 5), aside from the exclusion of the IFMAP layer. Seral stages were simulated across the UP with the landscape age-class demographics simulator (LADS), a stochastic simulation program that mimics the temporal and spatial patterns of fire disturbance and subsequent forest age classes across a landscape through time (Wimberly et al. 2000). Historical fire regimes in the UP were estimated with a map developed by the Forest Service, similar to one that was created for northern lower Michigan (Cleland et al. 2004). The simulation output provided maps of age classes within habtypes, and formed the basis of ELUs for the presettlement era. Habitat quality of the ELUs was estimated and the lynx model was
used to predict the historical variability of lynx habitat potential prior to European settlement. The methods used to generate input for the LADS program and estimate habitat quality for ELUs included the following:

*Simulating forest age classes* – A detailed description of the LADS program can be found in Wimberly et al. (2000), and I obtained the program directly from Michael Wimberly (Associate Professor, Department of Geography, South Dakota State University). The program simulates the spatial variability in fire regimes across large-scale regions of macroclimate and small-scale changes in topography and vegetation (Wimberly et al. 2000). The required input involves 4 raster layers of landscape characteristics and a set of parameters for determining the properties of the simulation and the behavior of fire within and across scales. The raster layers include: (1) climate zones, which delineate regions with specific fire rotations; (2) topographic zones, which portray local differences in topography that affect susceptibility to fire; (3) summary zones, which identify optional subsets of the study area for which summaries are desired; and (4) a buffer zone, which identifies areas from which fires can propagate but landscape summaries are not calculated. The input parameters include the total length of the simulation, length of burn-in period, length of summary steps, fire shape calibration, wind direction and intensity, fire susceptibility within topographic zones, age-class definitions, length of fire rotation, and the mean and standard deviation of fire size. Wimberly et al. (2000) designed the program for analysis of large landscapes (>1,000,000 ha) with a coarse pixel resolution (>4 ha); the raster layers were aggregated to a resolution of 180 m as a compromise. Also, islands were masked out of the maps
due to the cellular automata approach of fire spread in the program, which did not allow for disjunct pieces of land.

The most critical input map was that of the climate zones, which determined the fire regime parameters for a given location. The input map of climate zones for the UP was based on a spatial layer obtained from the Forest Service (Dave Cleland, USFS, Rhinelander, Wisconsin), which delineated the landscape into “biophysical units” (BPUs) based on ecological characteristics that affected fire susceptibility and spread (i.e., landform, lake density, soil texture and drainage) (Cleland et al. 2005a). Cleland et al. (2005a) assigned each BPU to 1 of 6 stand-replacing fire rotation categories, and used GLO survey notes to quantify the natural frequency of stand-replacing fires within each BPU. This work was an expansion on the analysis of northern lower Michigan (Cleland et al. 2004), and BPUs were mapped across the entire Laurentian Mixed Forest Province, with natural fire rotations summarized by BPU and ecoregion section (Cleland et al. 2005a). The accuracy of fire rotation estimates decreases with the size of the landscape unit (Cleland et al. 2005a); therefore, I condensed BPU-section groups that had similar fire frequencies, and recalculated fire rotations based on the fire frequency and area of each new BPU (Figure 7). The mean and standard deviation of fire sizes in the UP were estimated to be 342 ha and 1,347 ha, respectively (Cleland et al. 2005a). The LADS program used rotation length and fire size parameters to compute the mean number of fires per decade within a climate zone, which was simulated for each decade as a Poisson random variable. Individual fires were randomly sized according to a lognormal probability distribution, calculated from the input fire size parameters.
Figure 7. Spatial distribution of climate zones created from groupings of similar biophysical units (Cleland et al. 2005a) within and between ecoregion sections, along with associated estimates of natural fire rotations (NFR) in years.
I created the topographic zones with a 30-m U.S. Geological Survey digital elevation model (DEM) by calculating grids of flow accumulation and aspect within ARC/INFO 8.1. These functions identified local ridges (having a flow accumulation = 0) and areas that were flat or had southern aspects, which were assumed to have a higher susceptibility to fire (Zhang et al. 1999). The combination of the 2 grids resulted in the following 4 topographic zones (in order of increasing susceptibility): local valleys with slopes not facing south, local ridges with slopes not facing south, flat or south-facing local valleys, and flat or south-facing local ridges. The 30 m grid was aggregated to 180 m resolution using a majority filter to match the other input maps.

The summary zone was set to all forested areas within the contiguous boundary of the UP (excluding islands). The buffer zone included a 20 km strip along the border of Wisconsin to accommodate the propagation and spread of fires from outside the UP. Age classes were defined by the amount of time since the last stand-replacing fire; I defined 11 intervals with the following endpoints: 10, 20, 30, 40, 60, 80, 100, 150, 200, 250, and >250 years. Fire susceptibility was assumed to be uniform across age classes, and fire severity was set at a constant high level to simulate only stand-replacing fires. Wind direction was set at 135° (northwesterly) with a variability of 90° and a moderately low intensity (3 on a scale of 1-10). Simulation length was 10,000 yrs, with a 1000-yr burn-in period, a 10-yr run interval and a 50-yr summary interval. The output from the LADS program included maps illustrating the distribution of age classes across the UP for each of the 200 summary intervals, as well as a table of age-class proportions for each.

Estimating habitat quality within age classes – The vegetation attributes necessary to quantify habitat quality in the lynx model could not be directly measured for
the presettlement conditions. Therefore, estimates of quality for age classes within habtypes were developed using information from the literature and simplified assumptions about the structure and composition of vegetation.

Hare habitat quality was influenced by age class and habtype by assigning high scores to early age classes (<40 yrs) and habtypes with a conifer component, both characteristics that have been consistently described as important for snowshoe hare (see review in Hodges 2000). For the 0-10 year age class, all habtypes other than PiVa were assigned a hare HSI score of 50, to account for variability in regeneration of aspen-birch within the first 10 years of a severe fire. The PiVa habtype was assigned a score of 100 for the 0-10 year age class, assuming that jack pine regeneration would be dense after several years (Frelich 2002). The 10-20 and 20-30 year age classes were assigned a score of 100, under the assumption that these age classes would provide the greatest amount of understory cover for hare. The 30-40 year age class was assumed to be the last suitable stage before stem exclusion would result in a sparse understory; it was assigned a score of 75.

For the QuAc, TsMa, and TsTh-dry habtypes, the age classes between 40-150 yrs were assigned a score of 25; these habtypes contained a potential spruce-fir and/or cedar component that might have provided cover for hare. The PiVa was no longer suitable after 40 yrs due to the assumption that poor soil quality would result in a relatively sparse mature forest. The Acer and Frax habtypes were also unsuitable after 40 yrs, since dominance by hardwoods was expected to provide inadequate understory cover during the winter, even in areas where regeneration of shade tolerant species was high (i.e., sugar maple in the Acer habtype). The QuAc, TsMa, and TsTh-dry habtypes became
unsuitable after 150 yrs, when shade tolerant hardwoods would likely attain dominance in
the absence of fire (Frelich 2002). The TsTh-wet and Picea habtypes were assigned a
score of 50 for all age classes >40 yrs, since they were dominated by spruce-fir and
cedar; it was assumed that multi-aged mature stands within these habtypes would provide
pockets of suitable cover for hare. Non-forested areas provided no suitable hare habitat,
aside from shrub types, which were assigned an HSI score of 50.

The assignment of HSI scores for the denning and interspersion components of
the lynx model were even more simplified than that for hare quality. Habtypes that
provided potential denning sites based on mesic soil conditions included QuAc, TsMa,
Acer, and TsTh-dry; within those habtypes, age classes >60 yrs qualified as denning sites.
Travel cover was assumed to be provided by age classes >10 yrs within all habtypes. All
non-forested areas were considered unsuitable for lynx.

Based on my methods of assigning HSI scores, habitat suitability for lynx during
the presettlement era would be most influenced by fluctuations in the amount of early age
classes providing highly suitable hare habitat. Therefore, the total proportion of age
classes between 10-40 yrs on the landscape was calculated for each of the 200 simulated
maps (Figure 8), and maps representing the highest and lowest proportions were selected
for analysis with the lynx model. This allowed us to examine the range in habitat
suitability that might have occurred under the presettlement disturbance regime. The
resolution of the grid maps was increased to 30 m after assessing hare habitat quality so
that habitat units could be sufficiently accumulated into hare home ranges using Plume’s
(2005) GIS function, though I recognize that this procedure did not increase the spatial
accuracy of habitat components.
Figure 8. Fluctuation in the proportion of early successional forest (10-40 yrs old) over a 10,000-yr simulation of severe fires in the Upper Peninsula of Michigan, with arrows identifying highest and lowest years. Fire was simulated with the Landscape Age-class Dynamics Simulator under the natural disturbance regime that occurred prior to European settlement.
ANALYSIS OF MODEL OUTPUTS

The lynx model generated 5 maps of habitat quality for each version of the ecological land classification, representing indices for snowshoe hare habitat, the 3 life history components for lynx (foraging, denning, interspersion), and overall lynx habitat. Each map described habitat quality on a scale of 0-100, ranging from unsuitable to optimal conditions. There were 3 versions of the ecological land classification: 1 for current conditions and 2 for presettlement conditions.

The assessment of current conditions involved several variations for describing snowshoe hare habitat on the landscape, though each utilized the same ecoregion-ELU grid. Three versions of hare habitat quality were created from the separate HSI models (HSI-1, HSI-2, HSI-3), and another 3 were created for testing the sensitivity of random pixel assignment. The outputs of lynx foraging quality were compared within each set and one representative map was chosen to calculate overall lynx habitat and illustrate current conditions. Presettlement conditions were represented by 2 simulated landscapes of forest structure resulting from a natural disturbance regime. Each version had a separate ELU grid describing the arrangement of age-classes resulting from fire within each habtype, and subsequently, separate inputs and outputs for each of the habitat quality indices.

Pairwise comparisons were made within and between the outputs from current and presettlement conditions using several statistics. The pixel by pixel differences between paired maps was quantified by the Volume of Intersection Index (VOI) (Seidel 1992, Millspaugh et al. 2000). The VOI is a measure of the overlap between utilization distributions, with values ranging from 0 (no overlap) to 1 (direct overlap). This index
provided a measure of the similarity in the location of suitable habitat between a pair of maps. Each habitat quality map was converted into a utilization distribution, with HSI values weighted so that the total volume of the utilization distribution was equal to 1.

This procedure removes the magnitude of difference in habitat quality between paired maps, so the mean and standard deviation of the differences were also calculated. In this way, if a pair of maps had a similar spatial arrangement of suitable habitat, but one suggested a much higher level of suitability, the VOI score might be close to 1, while the mean difference would be relatively high. Likewise, if a pair of maps described suitable habitat at vastly different locations, but with a similar frequency of occurrence, the VOI score would be close to 0 and the mean difference would be low. Comparisons were also made among the outputs of simulated lynx home ranges created by Plume’s (2005) GIS function, in regards to the number and location of viable home ranges across the study area resulting from each habitat quality map. Lynx home ranges were simulated 5 times for each map; the mean number of home ranges was calculated for comparison, and an average map of home ranges was chosen for illustration.

Prior to the comparative analyses, each output map of habitat quality was aggregated to a 90-m grid resolution and clipped to the same extent. All processing was completed within ARC/INFO 8.1, aside from the simulation of lynx home ranges, which used a custom program written by Plume (2001).

**RESULTS**

**CURRENT ECOLOGICAL CONDITIONS**

The assessment of hare habitat quality, and subsequent lynx foraging quality, was affected by the choice of HSI model for current conditions (Figure 9). Horizontal cover
Figure 9. Frequency distributions of quality scores for hare habitat quality and lynx foraging quality across the study area, as assessed by HSI models measuring horizontal cover with manipulated, HSI-1 (a, d) and original, HSI-2 (b, e) seedling distributions, and the HSI model measuring stem cover units, HSI-3 (c, f).
within a stand was likely overestimated when the seedling size distributions were
manipulated, resulting in an overly optimistic portrayal of habitat quality by the first
model (HSI-1). The description of habitat quality was more similar between the HSI
model measuring horizontal cover with original size distributions (HSI-2) and the one
measuring stem cover units (HSI-3) (Figure 9). Volume of Intersection (VOI) indices for
lynx foraging quality ranged from 0.79 to 0.83, indicating that the HSI models were
locating suitable habitat in similar areas. Mean differences in lynx foraging quality were
high when comparing output from HSI-1 to both HSI-2 (mean = 43.84, SD = 18.07) and
HSI-3 (mean = 39.77, SD = 18.3), and low when comparing HSI-2 to HSI-3 (mean = -
4.07, SD = 12.42). I chose to retain the output from HSI-3 for further analysis, given the
high uncertainty in the measurements of horizontal cover, but also, the relative agreement
between output from HSI-3 and HSI-2.

The random pixel assignment had little effect on the assessment of lynx foraging
quality throughout the study area. The VOI index was 0.94 for each comparison, and
mean differences in foraging quality among the 3 grids ranged from -0.60 to 0.40, with
standard deviations <4.35, suggesting a high degree of similarity. Thus, local differences
in hare habitat quality among the simulated grids did not translate into significant
differences in lynx foraging quality. The grid with a mean quality most typical of the 3
was chosen to represent lynx foraging quality for the current conditions.

The amount and distribution of suitable habitat differed greatly between each
output of the lynx habitat components. Hare habitat quality and thus, lynx foraging
quality, was mostly poor throughout the study area (Figures 10, 11), with the highest
suitability (HSI = 40-60) in lynx foraging occurring in the eastern portion of the UP. The
Figure 10. Hare habitat quality in the Upper Peninsula of Michigan between 2000-2003. The inset illustrates a histogram of the proportion of quality scores across the study area.
Figure 11. Lynx foraging quality in the Upper Peninsula of Michigan between 2000-2003. The inset illustrates a histogram of the proportion of quality scores across the study area.
poor suitability in hare habitat was evident from the FIA data, as 74% of surveyed plots had an HSI score of 0 for hare, while 93% had an HSI score <60. The only habtype with >10% of its FIA plots scoring >60 for hare suitability throughout the UP was the TsTh-wet habtype (23%). Within the eastern ecoregion subsections 212Ra and 212Rb, 42% of FIA plots representing the TsTh-wet habtype scored >60 for hare suitability; the location of these subsections corresponds with the areas of the eastern UP that contained the highest suitability in lynx foraging across the study area.

Lynx denning quality was highly suitable throughout most of the UP (Figure 12), as only a few areas (<10% total area) were at an unsuitable distance (>400 m) to the vegetation types that provided denning areas. The high suitability for the denning component was due to the fact that over 53% of FIA plots met the requirements for den sites according to the lynx model, and combined with the high heterogeneity in habitat types across the UP, den sites were always within a suitable distance from a given point on the landscape.

The interspersion of non-lynx habitat was highly variable in the study area (Figure 13), with 20% of the total area having an HSI score <10 and 36% having an HSI score = 100. The extensive areas of non-forested wetlands, agricultural land, and human developments resulted in low levels of interspersion suitability across the eastern UP, while the contiguous forests of the western UP resulted in a high suitability with pockets of low suitability scattered evenly throughout.

Overall lynx habitat quality was mostly mid-range across the UP, with 49% of the total area having an HSI between 40 and 70 (Figure 14). Quality was limited by the high amount of non-suitable (HSI = 0) habitat according to the interspersion component, and
Figure 12. Lynx denning quality in the Upper Peninsula of Michigan between 2000-2003. The inset illustrates a histogram of the proportion of quality scores across the study area.
Figure 13. Lynx interspersion quality in the Upper Peninsula of Michigan between 2000-2003. The inset illustrates a histogram of the proportion of quality scores across the study area.
Figure 14. Overall lynx habitat quality in the Upper Peninsula of Michigan between 2000-2003. The inset illustrates a histogram of the proportion of quality scores across the study area.
the generally low suitability of foraging habitat. A mean of 603.8 viable lynx home ranges (SD = 26.3) were simulated for the current conditions, with locations throughout most portions of the UP (excluding the islands) (Figure 15). Viable home ranges were concentrated in 6 of the 21 ecoregion subsections (212Jb, 212Ra/b/e, 212Sn/q) covering nearly half of the UP; only 5% of viable home ranges were located in other subsections.

**PRESETTLEMENT ECOLOGICAL CONDITIONS**

The small size and relatively infrequent occurrence of fire throughout most of the UP resulted in a mature forest with little early successional vegetation on average. Over 97% of the study area had an average fire frequency >100 yrs, and 75% had an average frequency >500 yrs. The greatest amount of early successional vegetation (10-40 yrs old) in a simulation year was still only a small percentage (8.1%) of the total study area (Figure 8), but nearly 3 times the magnitude of the lowest amount. The main difference between the 2 simulation years (high, low) that were chosen to capture the range of presettlement forest conditions was a large conflagration that occurred in the eastern portion of the UP, which created >150,000 ha of early successional vegetation. As a consequence, hare habitat quality and lynx foraging quality fluctuated significantly between the high and low simulation years (Figures 16, 17).

The large differences in lynx foraging quality were not exhibited by the other lynx habitat components. Denning quality was only slightly decreased by the large expanse of early successional vegetation created by the conflagration in the high simulation year (Figure 18). The conflagration did not result in one contiguous patch of early successional vegetation; it contained many islands of mature forest that were saved from the burn, and thus, did not present a complete loss of areas for denning. Interspersion
Figure 15. Viable lynx home ranges simulated for current conditions in the Upper Peninsula of Michigan (islands excluded) using the Plume (2005) GIS function for aggregating habitat units. Viable home ranges were defined as having a quality =70.
Figure 16. Hare habitat quality during the 2 simulation years of presettlement fire disturbance with the highest (top) and lowest (bottom) amount of early successional vegetation. In the top map, a conflagration responsible for creating a large area of early successional vegetation can be seen in the eastern UP.
Figure 17. Lynx foraging quality during the 2 simulation years of presettlement fire disturbance with the highest (top) and lowest (bottom) amount of early successional vegetation.
Figure 18. Lynx denning quality during the 2 simulation years of presettlement fire disturbance with the highest (top) and lowest (bottom) amount of early successional vegetation.
quality was nearly the same between simulation years (Figure 19), as only non-forested vegetation types, water bodies, and the relatively few recent burns (<10 yrs) contributed obstacles to lynx in the presettlement landscapes. Over 90% of the study area had an HSI score >80 for denning and interspersion quality in each of the 2 simulation years.

Similar to the current conditions, overall lynx habitat quality during the presettlement era was limited by foraging quality. The majority of the study area had an HSI score between 40 and 70 for the high (61%) and low (64%) simulation years (Figure 20); HSI scores >70 occurred across 7% of the study area for the high year, and 3% for the low year. In each simulation year, the highest amount of suitable habitat occurred in the eastern portion of the UP, where average fire frequencies where 3 times the magnitude of those in the west. A mean of 677.8 viable lynx home ranges (SD = 8.6) were simulated for the high year, while 264.8 (SD = 7.1) were simulated for the low year (Figure 21). Home ranges were mostly located in the eastern portion of the UP, and the major difference between simulation years occurred at the conflagration, where a high concentration of home ranges appeared during the high year.

DIFFERENCES BETWEEN CURRENT AND PRESETTLEMENT CONDITIONS

The differences between current and presettlement conditions were unique to each lynx habitat component. These differences reflected the spatial properties of vegetation types across the landscape, as well as the structural attributes within them.

The VOI for lynx foraging quality between the current and presettlement maps ranged from 0.56-0.59, with a mean pixel difference of 2.87 (SD = 23.01) between the current and high presettlement year, and 6.98 (SD = 18.16) between the current and low presettlement year. Although the average foraging quality was slightly higher for the
Figure 19. Lynx interspersion quality during the 2 simulation years of presettlement fire disturbance with the highest (top) and lowest (bottom) amount of early successional vegetation.
Figure 20. Overall lynx habitat quality during the 2 simulation years of presettlement fire disturbance with the highest (top) and lowest (bottom) amount of early successional vegetation.
Figure 21. Locations of viable lynx home ranges according to habitat quality for the 2 simulation years of presettlement fire disturbance with the highest (top) and lowest (bottom) amount of early successional vegetation. Habitat units were aggregated into simulated home ranges using Plume’s (2005) GIS function.
current conditions than those during presettlement, the VOI scores indicate a large
difference in the location of suitable habitat for each time period. The VOI between each
simulation year for presettlement was 0.72, with a mean difference of 4.1 (SD = 19.88),
illustrating the changes caused by the conflagration.

The VOI for lynx denning quality between the current and presettlement maps
ranged from 0.94-0.96, with mean differences of 4.63 (SD = 20.99) and 2.74 (SD = 19.09) between the current map and the high and low maps, respectively. The VOI for
denning quality between the presettlement simulation years was 0.97, with a mean
difference of -1.89 (SD = 15.79). Lynx interspersion quality was the same between
presettlement simulation years, but very different when compared to current conditions.
The VOI for interspersion was 0.70, and the mean difference between current and
presettlement maps was -35.55 (SD = 39.13), indicating a large overall decrease in
interspersion quality from the presettlement era to the present.

The VOI for overall lynx habitat quality ranged from 0.68-0.69 for current and
presettlement conditions, with mean differences of 0.14 (SD = 33.25) and 1.51 (SD =
31.66) between the current map and the high and low maps, respectively. The number of
simulated lynx home ranges for the current conditions were within the range exhibited by
presettlement conditions (Figure 22), though the locations of the home ranges were very
different (Figures 15, 21). Under present conditions, home ranges were grouped in
several locations throughout the UP, including the far western region near the border of
Wisconsin. This region in the west is void of lynx home ranges under presettlement
conditions, due the lack of foraging habitat.
Figure 22. Number of viable home ranges simulated in each of the 3 lynx habitat quality grids, using the Plume (2001) GIS function to aggregate habitat units. Viable home ranges were defined as having an average quality >70.
DISCUSSION

Roloff’s (2003) model quantified habitat suitability for Canada lynx from the scale of an allometric home range (250 ha), at which habitat selection for a species is hypothesized to be strongest (Roloff and Haufler 1997). The model consisted of 3 habitat components representing life requisites that influenced the ability of a given landscape to support an individual. I integrated multiple forms of data within a GIS to quantify the spatial arrangement and structural attributes of vegetation types for assessing lynx habitat suitability under current and presettlement forest conditions in the UP.

For the analysis, the large size of the UP made it a logistical obstacle to collect data that could adequately describe the vegetation attributes necessary for the lynx model. I used multiple spatial layers portraying biotic and abiotic characteristics of the landscape to delineate the study area into ecological land units (ELUs), representing vegetation types that had similar composition and structure. The ELUs were based on a previous field study which illustrated relationships between tree species assemblages through succession and soil/landform properties in the UP (Coffman et al. 1983). The use of “habitat type” mapping increased my ability to stratify forest conditions beyond what could be provided by simple land-cover maps. The structural attributes for ELUs were obtained in 2 different ways: for current conditions, a region-wide inventory of forest stands conducted by the FIA program provided stand-level measurements of vegetation; for presettlement conditions, a simulation program developed for modeling the effects of fire disturbance on the spatial distribution of forest age-classes was used to estimate the range in forest conditions that might have occurred prior to European settlement. The synthesis of these data provided the best available estimates of both present and past
stand conditions in the UP, and allowed us to determine the nature of lynx habitat during both time periods.

The current distribution of lynx foraging habitat according to the model is very different from that which existed prior to European settlement. The output from both time periods indicated mostly poor quality foraging throughout the UP, due to a lack of early successional (10-40 yrs) vegetation types providing the understory cover required for a high abundance of snowshoe hares. According to the 2000-2003 FIA data, the ages of forest stands throughout the UP were normally distributed with a mean of ~60 yrs; approximately 15% of forest stands were at an age between 10-40 yrs. Using a 10,000-yr simulation of natural fire under a presettlement disturbance regime in the UP, the greatest amount of early successional vegetation in a given year was estimated at 8%, and forest stands had a mean fire return interval of 1,250 yrs. While the current landscape has been drastically altered by human actions, it would appear that logging has actually improved conditions for snowshoe hare and lynx in the UP, given that presettlement disturbances may have been too infrequent to provide adequate amounts of high quality habitat for hare. Both timber harvesting and severe fires have the ability to create adequate early successional vegetation types for hare, and each has been shown to benefit lynx in the long term (Fox 1978, Litvaitis et al. 1985, Koehler 1990). It would appear that current timber harvesting strategies do not occur at a large enough scale and high enough frequency to support high densities of hare and a viable lynx population in the UP, and neither did the presettlement fire regime. One example of a noticeable difference between current and presettlement foraging quality occurred in subsection 212Jb of the western UP, where quality was much higher under current conditions, especially for the
Ottawa National Forest. In this ecoregion subsection, sugar maple forest types had twice the proportion of FIA plots with a hare HSI >60 than found elsewhere in the UP. Reasons for the increased suitability are unclear, though nearly half of the highly suitable FIA plots had a record of silvicultural treatment in this ecoregion. The forests of the presettlement era within this ecoregion were dominated by late successional sugar maple, which was assumed to be of poor suitability for hare.

Denning quality was shown to be adequate throughout the UP, both during the presettlement era and in the present. The denning requirements for lynx were once thought to be strictly associated with late successional forest (Koehler and Aubry 1994), but Slough (1999) found a wider range of conditions, including those found in regenerating forests, provided suitable denning sites for female lynx. The definition of suitable habitat in the denning component of the model reflected the need for older forests, but not necessarily late successional, and therefore nearly half the forest stands according to the FIA data met the model requirements. Forest conditions according to the presettlement simulation were expected to support an abundance of denning opportunities, given the low frequency of disturbance and subsequently, the relatively high amount of mature forest across the landscape.

Interspersion quality was the habitat component that changed most significantly between time periods. Given that the presettlement landscape would only contain non-habitat where natural features such as water bodies and non-forested wetlands occurred, as well as recently burned forest stands, it was reasonable to assume that conditions for traveling would not be limiting during that time period. The amount of non-habitat has increased considerably since presettlement due to the conversion of forest into
agricultural land throughout large portions of the eastern and southern UP, as well as the development in and around cities such as Sault St. Marie and Marquette. Another region of non-habitat for lynx in the eastern UP is the Seney National Wildlife Refuge, an area once dominated by forested swamps that was mostly converted into a managed wetland after extensive logging in the late 19th century. The combination of these factors has resulted in an overall decrease in the contiguity of forested vegetation types that are important for lynx traveling needs, most notably in the eastern UP.

A number of assumptions were required for modeling current and historic forest conditions in the UP that may have effected the results. For current conditions, the FIA data represented a point sampling of forest structure, and therefore, the mapped ELUs did not have attribute information that could be considered accurate at the pixel level, especially for the fine scale structural measurements. This was evident from my method of simulating a random pixel distribution, which inferred that variation in structure was occurring at a distance of a 30 m (pixel size) with no regard to spatial autocorrelation. Given the systematic sampling which results in FIA plots being separated by potentially long distances, and my inability to obtain accurate plot locations due to confidentiality, the scale at which forest structure varied could not be validated. I caution that the prediction of stand variables not be used to assess forest conditions at scales less than that of townships (~92 km²). The filtering of FIA plots based on condition and canopy cover requirements reduced the dataset by 30%; an examination of the distributions of stand variables did not find significant differences. Even so, by classifying ELUs into discrete mapping units I eliminated the potential importance of ecotones, which can effect the assessment of forest conditions in a heterogeneous landscape. This might have had the
most impact in sparsely forested spruce bogs, where pixels of forest (>25% canopy cover) are scattered throughout lowland shrub vegetation, and any FIA plots located within would most likely have had <25% canopy cover and been filtered out. A more intensive analytic approach to classifying the landscape would be necessary to account for ecotones.

The varying accuracy of the digital maps, especially that of soil types, may also have produced errors in assessing habitat types across the landscape. The use of STATSGO soils data should be limited to large scale analyses, and given the near completion of the SSURGO dataset for Michigan (USDA, NRCS 2006), its utility may become obsolete in the future. The accuracy of the land cover dataset made it nearly impossible to distinguish canopy species with any confidence (Appendix C). New approaches to classifying land cover using combinations of high and low resolution imagery collected across multiple seasons would improve the ability to predict more detailed vegetation attributes.

For the presettlement conditions, many assumptions were made when modeling natural fire disturbance in the UP. Fire rotations were estimated using 10+ years of data collected during the GLO surveys of the 1800s; this small sample may not have been adequate to accurately document fire disturbances in the region. Also, I assumed that fire disturbance regimes did not change over time, an assumption that could be violated with broad-scale changes in climate (Frelich 2002, Cleland et al. 2005a). The interaction of many factors results in the propagation and persistence of tree species assemblages in a given area, not all of which can be captured by modeling the physiography of a region (Frelich 2002). I did not model disturbances other than those created by severe fires, but
while small surface fires, wind, and insects have all contributed to changes in forest composition and structure in the past, none have been of the same magnitude as that of severe fires (Cleland et al. 2005a). Finally, I assumed that stand age could be correlated with habitat quality for snowshoe hare, which is not always true (Hodges 2000), as understory structure can be influenced by factors other than stand age. Despite these caveats, I feel that my methods provided an adequate way to estimate habitat quality for lynx over a large study area and across 2 time periods.

The current absence of lynx in the UP is contrary to the output, which suggested there were several areas throughout the UP and close to the eastern and western borders that were capable of supporting multiple viable home ranges for lynx (Figure 15). The model assessed habitat quality in terms of the vegetation attributes on the landscape that would affect the availability of resources necessary for survival; it did not incorporate other factors such as competition, which will invariably affect habitat quality for a given species. The presence of coyote (*Canis latrans*) and bobcat (*Felis rufus*) in the UP can reduce the availability of snowshoe hare for lynx, even in areas where hare habitat quality is high (Koehler and Aubry 1994). These predators are not found in high abundance across the northern boreal forests of Canada, and their presence, combined with the generally low densities of hare across southern boreal forests (Koehler 1990, Koehler and Aubry 1994, Hodges 2000), may ultimately create unsuitable conditions for lynx in the United States. It is difficult to quantify the effects of these competitors on lynx foraging quality in the UP. Hoving et al. (2005) found that lynx occurrence in the Northeast was best predicted by annual snowfall, and suggested a critical threshold of 270 cm/year, below which the competitive advantage of lynx over other predators in the snow is lost.
Annual snowfall throughout more than half of the northern UP is >270 cm/year, especially within the snow belts (Eichenlaub et al. 1990), and 75% of the viable lynx home ranges that were simulated fell north of the snowfall threshold. Bobcat harvest records from the past 20 yrs indicate a close agreement with the snowfall threshold, as most harvests have been in areas of the southern UP with <270 cm/year (Figure 23). The trend of a warming climate may affect lynx habitat potential in the UP and beyond if snowfall decreases and the northward expansion of species such as bobcats and coyotes is facilitated (Hoving et al. 2005).

Another factor influencing the absence of lynx in the UP may relate to their ability to disperse from Canada. Lynx are known to be capable of long-distance dispersals (>1,000 km) (Nellis et al. 1972, Mech 1977, Ward and Krebs 1985), and historical records indicate that lynx were once able to enter the UP during population eruptions in Canada (McKelvey et al. 2000a). In the early winter of 2003, a lynx was incidentally trapped in Hiawatha National Forest near the city of St. Ignace, which marked the first sighting/trapping of an individual in more than 20 years for the UP (Stephen Sjogren, USFS biologist, personal communication); the origins of the trapped lynx were undetermined. In the eastern UP, habitat quality was poor throughout the northern end and annual snowfall was below the critical threshold in the southern end (Figure 23), suggesting this area could be a potential barrier to lynx dispersal from Ontario. If the lynx trapped in 2003 was a disperser from Ontario, it is obvious that migration across this area of the UP is not impossible. An examination of the connectivity of habitat between Canada and the areas of suitable habitat within the UP may provide insight into the current status of lynx in the region.
Figure 23. Relative frequency of bobcat harvests in the Upper Peninsula of Michigan between 1985-2005. The dotted line represents the boundary below which annual snowfall is <270 cm/year.
CHAPTER 2: 
TEMPORAL CHANGES IN HABITAT CONNECTIVITY FOR CANADA LYNX IN THE UPPER PENINSULA OF MICHIGAN 

INTRODUCTION 

In listing the Canada lynx (*Lynx canadensis*) as a threatened species in the contiguous United States, the US Fish and Wildlife Service (USFWS) called for “accurate mapping of lynx habitat in the Great Lakes Region” (USFWS 2000:16057) to provide the location and distribution of resources necessary for the species’ persistence and identify potential areas for conservation and management. The habitat modeling framework proposed by Roloff and Haufler (1997) was used with a habitat suitability index (HSI) model (Roloff 2003) to quantify the Upper Peninsula (UP) of Michigan in terms of its ability to provide the necessary life requisites for lynx (Chapter 1). The analysis identified areas of the UP that were potentially important for lynx persistence in the region, though no evidence of a resident lynx population is currently present (Beyer et al. 2000), and provided an historical context by assessing forest conditions and lynx habitat suitability in a presettlement-era landscape. The results of the analysis indicated that lynx habitat potential was high in several areas of the UP, both in the past and present, and it was hypothesized that factors other than the condition of forest communities were preventing a resident population from being established. One of the competing hypotheses was that habitat connectivity between the core lynx population in Canada and suitable habitat in the UP may be poor enough to preclude the successful migrations necessary for lynx establishment in the region.
Lynx are known to travel great distances in search of habitat that can support adequate foraging opportunities, especially during times when the local abundance of their primary prey, snowshoe hare (*Lepus americanus*), is extremely low (Nellis et al. 1972, Mech 1977, Ward and Krebs 1985). Mech (1977) reported a dispersal distance of 483 km in Minnesota, and long-distance movements of lynx in Canada have been estimated to exceed 1,000 km (Slough and Mowat 1996, Poole 1997). It is clear that distance alone would not prevent lynx from migrating out of the core population in Canada to areas of the Great Lakes such as the UP. Historical accounts of lynx presence in the UP include an influx of occurrences during the 1960s, when an irruption of populations in Canada caused a large migration of individuals into the Great Lakes region (McKelvey et al. 2000a). The obstacle presented by the St. Mary’s River in the eastern UP cannot be considered insurmountable to lynx, given that migrations have been documented in the past, as well as the fact that winter ice cover on the river has averaged >90% over the past 30 years (Assel 2003). The ability, or apparent inability, of lynx to migrate into the UP may be more related to habitat quality than physical obstacles.

The objective of this analysis was to examine the connectivity of lynx habitat across the UP during the past and present with a Geographic Information System (GIS). The outputs from a prior assessment of lynx habitat suitability (Chapter 1) were used to provide input for quantifying the “cost” associated with traveling from outside the UP to areas within that could potentially support lynx. Cost would be defined by the amount of energy expended on a dispersal, and/or the risk of mortality associated with conditions on the landscape (e.g., starvation, incidental trapping). Comparing the nature of least-cost travel corridors between current and presettlement landscapes might contribute insight
regarding the current absence of lynx in the UP, and identify areas that would be important for lynx conservation and management.

STUDY AREA

(see Chapter 1)

METHODS

I assumed that dispersal by lynx would be associated with areas of good habitat, and that the least-cost corridor between 2 points on the landscape would indicate the travel route providing the highest probability of survival by an individual. Determining the least-cost corridor required a map of habitat quality that could be converted into a cost surface, over which an accumulated cost could be calculated between grid pixels to identify corridors with the least resistance. Multiple cost surfaces were compared to quantify differences in the number and location of least-cost corridors, as well as the magnitude of cost calculated for each corridor.

Maps of lynx habitat quality were created using the modeling framework proposed by Roloff and Haufler (1997) with an HSI model (Roloff 2003) constructed for lynx. A detailed description of the data and processes used to model lynx habitat can be found in Chapter 1. Lynx habitat quality was quantified for 2 time periods in the UP, with 1 map for current conditions and 2 maps for presettlement conditions. The 2 maps for presettlement conditions represented simulation years with the highest and lowest amounts of early successional forest, capturing the range in lynx habitat quality exhibited under a natural disturbance regime prior to European settlement. The maps of habitat quality ranged in HSI score from 0-100, representing unsuitable to optimal habitat conditions, respectively. These maps were converted into cost surfaces by assigning a
relative cost value based on the HSI score. I assigned a cost of 20 to HSI scores of 0 to ensure that non-habitat was heavily weighted against in the cost surface. The remaining HSI scores were grouped into 4 classes at intervals of 25 (1-25, 26-50, 51-75, 76-100), and each class was assigned a cost based on its rank, with the most suitable class (e.g., 76-100) receiving a cost of 1. In this way, poorer quality habitat had a higher cost of traversing, with non-habitat having a very high cost. Connectivity was assessed between 3 points in the UP (Figure 25), with source locations near the border of Canada on the eastern side and the border of Wisconsin near Lake Superior on the western side, and a centralized destination located in the middle of the UP. This destination represented an area where viable home ranges had been simulated under current conditions.

I used 2 functions within the GRID extension of ARC/INFO 8.1 to create the least-cost corridors for each map. The costdistance function calculated the least-accumulative-cost distance for each pixel on the cost surface grid to pixels from each of the 3 points of interest. The costdistance grids were paired using the corridor function to calculate the sum of accumulative costs between each of the 2 source locations and the destination; pixels with the lowest accumulative costs identified the least-cost corridors between points. The least-cost corridors were compared by year and point pair to quantify differences in their locations and magnitudes. I calculated the average habitat quality and total amount of non-suitable habitat crossed by each path.

RESULTS

The least-cost corridors illustrated some large differences in lynx habitat connectivity between current and presettlement forest conditions, and the direction of difference depended on the point pair. The change from presettlement conditions to
Figure 24. Location of sources (black dots) and destination (white star) for calculating the least-accumulative-cost corridor for lynx in the Upper Peninsula of Michigan.
current conditions was relatively minor in the west and more pronounced in the east (Figure 25). These differences reflect the changes in land use across each side of the UP.

The lowest accumulated cost for the eastern corridor was 36% and 12% higher for the current map than that for the high and low presettlement maps, respectively (Figure 25). Only the current corridor required the crossing of non-suitable habitat (1.08 km in length), which occurred directly next to the source point in the east. The mean HSI score for the current eastern corridor (63.5) was between those of the presettlement (high = 76.0, low = 62.5). In mapping the corridors under a similar scale of accumulative cost, the two presettlement maps provided the same route, though the high presettlement had less relative cost (Figure 26). The current corridor provided a different route from that of the presettlement maps and the relative cost was higher (Figure 26).

The lowest accumulated cost for the western corridor was 4.6% and 2.8% lower for the current map than for the high and low presettlement maps, respectively (Figure 25). None of the corridors required crossing non-suitable habitat. Similar to the eastern corridors, the mean HSI score for the current western corridor (60.0) was bounded by that of the high (61.3) and low (58.2) presettlement corridors. The current corridor provided a different route than that of each presettlement map (Figure 27), but the difference was minor. The small differences in accumulative cost between all 3 maps are reflected in the similarity between the size of each corridor (Figure 27).

**DISCUSSION**

The least-cost corridor currently available to lynx in the western UP was similar to those provided under presettlement forest conditions, while in the eastern UP, the current corridor changed significantly in a negative way. Contiguous forest patches in
Figure 25. Least-accumulative-cost values for the eastern and western corridors created for each cost surface in the Upper Peninsula of Michigan.
Figure 26. Relative travel costs for eastern corridors simulated in each cost-surface map of the Upper Peninsula of Michigan, for the high (A) and low (B) presettlement conditions and present conditions (C).
Figure 27. Relative travel costs for western corridors simulated in each cost-surface map of the Upper Peninsula of Michigan, for the high (A) and low (B) presettlement conditions and present conditions (C).
Habitat quality was lower in the west under presettlement conditions, but this did not significantly alter the corridor. Habitat quality in the east was higher under presettlement conditions; large patches of unsuitable habitat caused the corridor under current conditions to be located along a different route and accrue a higher cost. The existence of unsuitable habitat is a reflection of the widespread agricultural land and human developments in the eastern UP that were not present during the presettlement era. The only entry point for lynx from Ontario is located directly within an area of unsuitable habitat for lynx near the city of Sault St. Marie. Whether a lynx would survive a migration through this area is unknown, but it represents the only major obstacle to dispersal across the eastern UP. The western UP offers a number of travel corridors, and its proximity to lynx populations in Minnesota suggests that successful dispersals would be more probable from that region.

Relating habitat quality to cost is a difficult task, since the relative cost of traveling through an area for a species cannot be readily quantified. Little information exists in the literature for determining how lynx might view a landscape in terms of travel cost. Intuitively, areas with low habitat quality would be traversed at greater speeds than those of higher quality; how this would affect the probability of a successful dispersal is unknown. If cost is defined by the time spent or distance traveled while dispersing, good quality habitat could theoretically have a higher “cost” than that of poor quality habitat, if the good quality habitat is small in area and does not provide enough resources for home range establishment. Individuals that spend time hunting in small patches of good quality
habitat could have an increased risk of mortality compared to those that travel quickly to large contiguous patches. The time and distance over which a lynx could travel without foraging might indicate the importance of habitat quality during a dispersal event. More information on the nature of lynx dispersals in the Great Lakes region is necessary to better understand the probability of a population being established in the UP.
MANAGEMENT IMPLICATIONS

Our results indicated that lynx habitat potential in the UP has changed from that which existed prior to European settlement. Human alterations on the landscape have both increased and decreased habitat quality. The development of human-made structures and agricultural practices in the eastern UP has significantly decreased habitat potential in that region, while forestry practices throughout the UP have resulted in a wider distribution of potential habitat. Conversion of forests to other land-uses is usually not reversible, and therefore, the low habitat quality in many regions of the UP, especially in the east, cannot be changed. While it appears that timber harvesting has increased the quality of lynx habitat, this increase is realistically small. Snowshoe hares require widespread areas of early successional vegetation, the size of which can only be created by relatively large disturbances such as severe fire, or clearcutting. The negative social connotation that follows clearcut harvesting can make the practice undesirable and/or impossible across large areas, especially where public opinion seeks to prevent it. Without a consistent cycle of forest disturbance that could allow hare populations to thrive, the chances of a lynx population being established in the UP are low.

A further investigation of the areas I qualified as suitable habitat is necessary to validate the model outputs, but also to determine whether other factors (i.e., interspecific competition, snowfall) that were not modeled might have an effect on habitat quality. Given that high snowfall has been hypothesized to separate lynx from other competitors, the current trend of a warming climate will presumably increase the northern expansion of these species may further decrease habitat potential for lynx in southern boreal forests, including those of the UP.


Hill, M. O. 1979. TWINSPAN - a FORTRAN program for arranging multivariate data in an ordered two-way table by classification of the individuals and attributes. Cornell University, Ithaca, New York, USA.


APPENDIX A:

HABITAT SUITABILITY MODEL FOR THE NORTH AMERICAN

LYNX: LAKES STATES VERSION

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This habitat model was originally written for the intermountain west region of the United States. In an effort to calibrate the model for use in the Lake States, modifications to the mathematical equations were implemented in a manner consistent with how the model was applied in a verification study in Manitoba, Canada (Nylen-Nemetchek 1999). These modifications were based on the fact that most portions of the Lake States tend to have less accumulated snow, less topographic influences, and an overall smaller forest stature than the intermountain west forest communities. Although outputs from the calibrated lynx model in Manitoba corresponded to the occurrence of lynx as expected (Nylen-Nemetchek 1999), this Lake States version should be viewed as a hypothesis on lynx habitat relationships in the Region.

Throughout this document, the citations in support of model variables remained unchanged from the original intermountain west documentation.

Introduction

In 1995, concerns over lynx (Lynx canadensis) population viability in the conterminous United States prompted several resource groups to petition the species for listing under the Endangered Species Act. Since that time, lynx conservation has received considerable attention (e.g., Paquet and Hackman 1995, Washington Department of Natural Resources 1996, Ruggiero et al. 2000). Lynx were listed as federally threatened in 2000 throughout the conterminous United States range (U.S. Fish and Wildlife Service 2000). Lynx are also listed on Appendix II of the Convention on International Trade of Endangered Species (CITES) and are currently identified as a sensitive species by several U.S. Forest Service regions (Macfarlane 1994).

Concerns surrounding the effects of land management activities on lynx populations in the conterminous United States necessitated development of a model that quantitatively assessed these effects on habitat suitability. Although few data exist on lynx in the United States from which to build habitat models, considerable research has been conducted in Canada and Alaska (see review in Ruggiero et al. 2000). Lynx data from Canadian and Alaskan studies must be applied cautiously to the lower United States because of the unique features associated with the
southern range of lynx distribution. These unique features of southern lynx range, with an emphasis on western studies, include:

- The inherently peninsular and disjunct distribution of suitable habitats (Koehler and Aubry 1994).


- More consistently low hare densities, comparable to hare population lows in Canada and Alaska (Koehler and Aubry 1994).

- Consistently low lynx densities making the effects of fur-harvests on populations in some areas additive rather than compensatory (Koehler and Aubry 1994).

- Higher human densities coupled with low lynx densities potentially causing both direct (e.g., fur harvest) and indirect (e.g., land development causing displacement) anthropogenic effects on population persistence to be of greater magnitude.

- The potential importance of immigration from the north for short-term population persistence (Koehler and Aubry 1994).

- The range overlaps of lynx, bobcat (*Felis rufus*), and coyote (*Canis latrans*) (Koehler and Aubry 1994), and the propagation of bobcat and coyote range extensions that typically accompany anthropogenic development.

The habitat suitability index (HSI) modeling concept (U.S. Fish and Wildlife Service 1981) was used to develop a lynx habitat model for the southern Rocky Mountains that was field verified (Nylen-Nemetchek 1999) and as such, formed the basis for a Lake States model. In this model, habitat potential for a lynx home range was divided into foraging, denning, and travel requisites. The lynx model uses a limiting factor approach (a concept founded in ecological theory) in that the most limiting resource(s) is assumed to have the greatest impact(s) on the population. A premise of HSI models is that limiting factors can be portrayed using mathematical relationships between vegetation structures, spatial metrics, and indices of habitat quality (U.S. Fish and Wildlife Service 1981). Theoretically, these limiting factors can be expressed as an index to animal fitness.

Overview

Critical considerations prior to running the lynx model are the resolution, accuracy, and precision of the land classification system and associated vegetation attribute information for the
planning landscape. The land classification must be capable of characterizing vegetation structures and spatial arrangements at a resolution compatible with lynx and snowshoe hare habitat use. The ideal stratification is a stand-based (minimum mapping unit around 2 ha) ecological classification system that integrates existing vegetation conditions and site potentials (e.g., geology, soils). An ecological classification system is recommended to reduce the variability in quantifying understory vegetation attributes since these attributes are primary components of the lynx model. Deviations from these baseline recommendations for land stratification will reduce the robustness and utility of the model output.

The lynx model is divided into three components: 1) foraging, 2) denning, and 3) interspersion. The lynx model was specifically developed for the mountainous habitats of Washington, Idaho, and Montana, corresponding to the southern extension of lynx range in the Rocky Mountains, however, the model framework may be applied to other regions, in this case the Lake States region. In applying the model to the Lake States, each input variable was calibrated to the biogeoclimatic conditions characteristic of the region. For example, model variables that index winter browse availability for snowshoe hares will differ across regions depending on average snow depths. These types of differences must be accounted for when applying this framework to other regions.

Lynx habitat in the southern Rocky Mountains consists primarily of 2 structurally different forest types occurring at opposite ends of the forest seral gradient: 1) early successional forest structures that contain high numbers of prey (especially snowshoe hare), and 2) late-successional forest structures for denning (Koehler and Aubry 1994). Second-growth forests with dense understories also may support abundant hare populations (John Weaver, Northern Rockies Conservation Cooperative, Missoula, MT, pers. comm.). Intermediate seral stages with sparse understories serve as travel cover, functioning to provide connectivity between foraging and denning patches (Koehler and Aubry 1994). Recent data from Montana and Maine suggest that late-successional forests may not be as important as originally thought for denning, but may serve as snowshoe hare refugia thereby contributing to foraging habitat.

Literature reviews and consultation with experts on lynx and snowshoe hare ecology were used to develop this lynx model. The model is not stand-based, but rather, it is designed to evaluate habitat quality in an area that corresponds to the allometric home range of lynx (250 ha; Roloff and Haufler 1997). Within a 250 ha area, habitat quality is expressed on a scale of 0-100, denoting "poor to good" habitat, respectively. Subsequently, habitat units from each allometric home range are aggregated into viable, marginal, or non-viable areas, the size of which depends on habitat quality (Roloff and Haufler 1997).
**Lynx Foraging Component**

Forage availability during the winter months appears to be the most important criterion in the determination of lynx home range size and degree of home range overlap (McCord and Cardoza 1982, Ward and Krebs 1985). Lynx populations covary with snowshoe hare numbers (Brand et al. 1976, Brand and Keith 1979, Parker 1981, Bailey et al. 1986), and lynx tend to choose habitats where hares are most numerous (Murray et al. 1994). Although prey switching has been documented in the southern Rocky Mountains, the underlying determinant of lynx fitness appears to be related to winter snowshoe hare abundance. Thus, the foraging component of the lynx model is based on winter snowshoe hare habitat quality. Snowshoe hare habitat is assessed using an HSI model, and the results of the hare model are incorporated into a lynx foraging assessment.

**HABITAT SUITABILITY MODEL FOR SNOWSHOE HARE**

**Overview**

Important components of hare habitat have been reported for different vegetation types and include dense woody vegetation (Adams 1959, Monthey 1986, Koehler 1990, Keith et al. 1993), stem diameter of browse (Keith 1974), continuity of coniferous cover (Brocke 1975), habitat interspersion (Keith et al. 1993), the distance to lowland forest cover (Conroy et al. 1979), and patch size (Thomas et al. 1997). The snowshoe hare model is divided into two primary components: 1) foraging, and 2) security cover. These components are mathematically combined into an overall index of winter hare habitat quality at the map-polygon and home range levels.

Winter foraging and security cover conditions are assumed to be limiting to hares (Hart et al. 1965, Dolbeer 1972, Keith and Windberg 1978, Pease et al. 1979, Keith et al. 1984, Boutin et al. 1986, Keith et al. 1993). In this model, summer habitats are not considered a limiting factor. To index the quality of snowshoe hare habitat, it is assumed that measures of understory cover and species composition in different height strata can be used (summarized by Ferron and Ouellet 1992). In support of this assumption, Thomas et al. (1997) demonstrated significant relationships between hare population indices and the horizontal and vertical cover of understory vegetation. Since few snowshoe hare studies have been conducted in the Pacific Northwest and Lake States, the vegetation-hare relationships depicted in this model were inferred from Thomas et al. (1997). Studies conducted across North America were used to supplement Thomas et al.’s (1997) work.

**Snowshoe Hare Winter Food Component**

Winter availability of palatable browse is believed to be a limiting factor of snowshoe hare populations (e.g., Windberg and Keith 1976, Pease et al. 1979, Vaughan and Keith 1981,
Sinclair et al. 1988, Sullivan and Sullivan 1988, O’Donoghue and Krebs 1992). In this model, the amount of winter browse for snowshoe hares is assessed using two different measures: 1) the amount of live horizontal (or lateral) cover, and 2) the amount of live vertical cover. Both measures are used to represent the “thickness” of forage for hares. Horizontal and vertical cover correlate with understory stem density (Gysel and Lyon 1980, Litvaitis et al. 1985), although this relationship may be weak (Thomas et al. 1997). In the southern Rocky Mountains, forage for hares is quantified in three height strata; 0-1, 1-2, and 2-3 m to account for variations in availability as a result of changing snow depths and the ability of hares to "clip" down vegetation from unreachable heights (Keith et al. 1984, Sullivan and Moses 1986). In the Lake States region, however, snow depths rarely exceed 2 m in height, particularly for areas inland from the Great Lakes. Thus, this Lake States lynx model is based on 2 height strata, 0-1 m and 1-2 m.

Horizontal cover is measured along the geometric plane that corresponds to the ground (i.e., the thickness if one stands and tries to look through a vegetation type) whereas vertical cover is measured along a geometric plane perpendicular to the ground (i.e., the thickness if one looks up). The woody browse component in this model includes all live plants since hares have been documented to feed on a variety of species (Table 1). However, it is recognized that some browse is unpalatable or of higher quality to hares and if possible, this model component should include only those species used by hare for foraging.

Thomas et al. (1997) found that highest browse use occurred in vegetation types with 30 to 40% horizontal cover of live vegetation. Use of vegetation types for foraging declined as woody cover approached <20% (Ferron and Ouellet 1992, Thomas et al. 1997). These findings roughly correspond to other studies that found highest hare use during winter in vegetation types with ≥50% horizontal cover (Carreker 1985, Parker 1986). Thus, optimal foraging habitat for snowshoe hares is provided by vegetation types with ≥35% horizontal cover of live vegetation (Fig. 1). Hare winter foraging habitat quality declines as horizontal cover decreases, and habitat is unsuitable when ≤10% horizontal cover of live vegetation is provided (Fig. 1). Horizontal cover for foraging habitat is measured for the 0-1 and 1-2 m height strata.

Thomas et al. (1997) also associated vertical cover with the intensity of snowshoe hare browsing. Highest browse levels corresponded to about 80% vertical cover. Browse use approached zero as vertical cover declined to about 20%. In this model, vertical cover of live vegetation is optimum at ≥80% and provides no foraging habitat at ≤20% (Fig. 2). Similar to horizontal cover, vertical cover is measured for two height strata.

For both horizontal and vertical cover relative to snowshoe hare browsing potential, overall habitat quality is assessed independently for each strata (i.e., an increase in browse in one
stratum cannot offset a decrease in another stratum). The rationale behind this logic is that snow
levels dictate the heights at which hares can access browse, thus, the different strata cannot
compensate for each other (i.e., if the 0-1 m strata is unavailable, the quality does not matter
because hares cannot access it). Two HSI scores are calculated from figure 1: 1) horizontal cover
0-1 m tall ($HSI_{hare,wint,food,hcov,0-1}$), and 2) horizontal cover 1-2 m tall
($HSI_{hare,wint,food,hcov,1-2}$). Similarly, two HSI values are derived from figure 2: 1) vertical
cover 0-1 m tall ($HSI_{hare,wint,food,vcov,0-1}$) and 2) vertical cover 1-2 m tall
($HSI_{hare,wint,food,vcov,1-2}$). The equation for calculating hare foraging HSIs for the Lake States
from horizontal and vertical cover is presented in equations 1 and 2, respectively.

Equation 1:

$$\left(\frac{HSI_{hare,wint,food,hcov,0-1} + HSI_{hare,wint,food,hcov,1-2}}{2}\right) = HSI_{hare,wint,food,hcov}$$

Equation 2:

$$\left(\frac{HSI_{hare,wint,food,vcov,0-1} + HSI_{hare,wint,food,vcov,1-2}}{2}\right) = HSI_{hare,wint,food,vcov}$$

The foraging habitat quality for snowshoe hare is based on the arithmetic mean of
$HSI_{hare,wint,food,hcov}$ and $HSI_{hare,wint,food,vcov}$ (Equation 3). An arithmetic mean was
selected because some foraging habitat can be provided (i.e., the foraging HSI > 0) if only
horizontal or vertical foraging cover is present. For example, densely-stocked woody species
with single-stem growth forms that do not have spreading crowns [e.g., aspen (Populus
tremuloides)] will tend to exhibit suitable horizontal cover during winter months whereas the
vertical cover provided by this vegetation community may be marginal. Using the arithmetic
relationship in Equation 3, horizontal or vertical foraging cover can equal 0 and the foraging HSI
can still be >0. Both horizontal and vertical foraging cover is weighted equally in the winter food
component (Equation 3).
Equation 3:

\[
\left(\frac{\text{HSI}_{\text{hare}, \text{wint}, \text{food}, \text{vcov}} + \text{HSI}_{\text{hare}, \text{wint}, \text{food}, \text{cov}}}{2}\right) = \text{HSI}_{\text{hare}, \text{wint}, \text{food}}
\]

Snowshoe Hare Winter Security Cover Component

The presence of adequate winter security cover has been recognized as the primary determinant of hare habitat quality (Buehler and Keith 1982, Wolfe et al. 1982, Sievert and Keith 1985, Rogowitz 1988). In this model, cover is defined as any structure (live or dead) that provides security for snowshoe hares. Winter security cover for hares is assessed using three measures of structure and composition: 1) understory vegetation composition (Severaid 1942, deVos 1962, Bookhout 1965a, b, Buehler and Keith 1982, Orr and Dodds 1982), 2) horizontal cover in three height strata (Brocke 1975, Wolfe et al. 1982), and 3) vertical cover in three height strata (Wolff 1980).

Understory Vegetation Composition

Snowshoe hares appear to select habitats based on vegetation structure as opposed to species composition (Litvaitis et al. 1985, Ferron and Ouellet 1992) and will use most forest cover types if adequate understory vegetation exists. However, some researchers have demonstrated that vegetated stands <3 m tall dominated by conifer species provide better habitat as opposed to stands dominated by deciduous species (deVos 1962, Buehler and Keith 1982, Orr and Dodds 1982, Monthey 1986). A subjective evaluation of the dominant understory vegetation type ≤3 m tall is used to index winter cover composition (HSI_{hare, wint, cov, dom}). The following criteria were developed to calculate HSI_{hare, wint, cov, dom}:

<table>
<thead>
<tr>
<th>Understory Cover Dominance Class</th>
<th>HSI=50</th>
<th>HSI=75</th>
<th>HSI=100</th>
<th>HSI=0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deciduous</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mixed</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Coniferous</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>None</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

a Based on a subjective evaluation of understory cover ≤3 m tall. "Deciduous" = >60% understory in deciduous species; "Mixed" = 40% ≤ Deciduous/Coniferous Cover ≤60%; "Coniferous" = >60% understory in coniferous species. If no understory cover exists, the HSI defaults to 0.
Horizontal Security Cover

Brocke (1975) suggested that horizontal cover is the single most important stimulus in selecting cover to avoid predation. Parker (1986) found that snowshoe hare population indices were related to horizontal cover in the 1-2 and 2-3 m height strata. Winter horizontal cover \( (HSI_{hare,wint,cov,hcov}) \) includes both live and dead vegetation and inanimate objects (e.g., rocks, root wads). Optimum horizontal cover is provided at \( \geq 90\% \), and horizontal cover \( \leq 40\% \) is deemed unsuitable winter habitat (Carreker 1985, Ferron and Ouellet 1992) (Fig. 3). Separate horizontal cover HSI s are calculated for height strata 0-1 and 1-2 m and these are subsequently combined using an arithmetic mean to produce \( HSI_{hare,wint,cov,hcov} \) (Equation 4).

Equation 4:

\[
\left( \frac{HSI_{hare,wint,cov,hcov,0-1} + HSI_{hare,wint,cov,hcov,1-2}}{2} \right) = HSI_{hare,wint,cov,hcov}
\]

Vertical Security Cover

Vertical vegetation cover is also considered an important component of hare habitat quality (Wolff 1980). Vertical cover is defined as the percent cover of live or dead material. Again, multiple strata are used to account for variations in cover availability as a result of changing snow depths. Optimal vertical cover occurs at \( \geq 90\% \), and vertical cover \( \leq 40\% \) provides unsuitable habitat \( (HSI_{hare,wint,cov,vcov}) \) (Fig. 4). Separate vertical cover indices are calculated for height strata 0-1 \( (HSI_{hare,wint,cov,vcov,0-1}) \) and 1-2 \( (HSI_{hare,wint,cov,vcov,1-2}) \) m and are subsequently combined using an arithmetic mean to produce \( HSI_{hare,wint,cov,vcov} \) (Equation 5). An arithmetic mean was selected because if vertical cover is provided in one stratum, the vegetation type provides functional security cover for at least a portion of the winter (i.e., until snow covers it). Both vertical cover strata are weighted equally in the winter vertical cover component (Equation 5).
Equation 5:
\[
\frac{H_{\text{hare, wint, cov}, 0 - 1} + HSI_{\text{hare, wint, cov}, \text{vcov}}}{2} = HSI_{\text{hare, wint, cov, vcov}}
\]

Winter security cover for snowshoe hares \((H_{\text{hare, wint, cov}})\) is computed by first establishing whether suitable security cover exists (as the arithmetic mean of \(H_{\text{hare, wint, cov, hcov}}\) and \(H_{\text{hare, wint, cov, vcov}}\)) (Equation 6). Subsequently, the arithmetic mean from the cover calculation is geometrically combined with the understory composition component \((H_{\text{hare, wint, dom}})\) (Equation 6). This mathematical relationship will cause \(H_{\text{hare, wint, cov}}\) to score as unsuitable if appropriate cover conditions are not provided.

Equation 6:
\[
\left[\frac{(H_{\text{hare, wint, hcov}} + H_{\text{hare, wint, vcov}})}{2}\right] \times HSI_{\text{hare, wint, dom}}^{0.5} = HSI_{\text{hare, wint, cov}}
\]

Calculating the Snowshoe Hare Winter HSI

Winter habitat conditions are assumed to limit snowshoe hare populations, and thus, winter HSI values drive the final HSI calculation. Winter habitat components (forage and cover) are integrated into one habitat value using a geometric mean. If the winter HSI for one habitat component equals 0, the final HSI equals 0 (i.e., suitable forage and cover must be present to provide hare habitat). Equation 7 is used to calculate the snowshoe hare winter HSI \((H_{\text{hare, wint}})\).

Equation 7:
\[
(HSI_{\text{hare, wint, food}} \times HSI_{\text{hare, wint, cov}})^{0.5} = HSI_{\text{hare, wint}}
\]
Calculating the Lynx Forage Component

The index $HSI_{hare,wint}$ provides a polygon-level assessment of snowshoe hare habitat quality. The next step in the modeling process for lynx is to relate the polygon-level depiction of hare habitat quality to the allometric home range of lynx (250 ha). It is assumed that for each allometric home area to support lynx, some minimum level of foraging habitat (i.e., snowshoe hare habitat) is required. These habitats must themselves be of sufficient quality to support consistent and abundant numbers of snowshoe hares. Applying the methodology of Roloff and Haufler (1997), home range functionality thresholds were established for snowshoe hares based on an evaluation of hare studies.

Similar to relationships demonstrated for other wildlife species, the home range of Lagomorphs appears negatively associated with habitat quality (Boutin 1984, Hulbert et al. 1996). It is assumed that hares will exhibit smallest home ranges when habitat conditions are optimum and that hares have largest home ranges or become nomadic in unsuitable habitats (see Roloff and Haufler 1997). Although few home range studies have quantified habitat quality and estimates of annual home range sizes for hares are uncommon, existing literature and allometric theory were used to estimate home range functionality thresholds for snowshoe hares (Roloff and Haufler 1997).

The smallest documented home range for snowshoe hares is 1.4 ha for females (mid-summer to fall) (Ferron and Ouellet 1992). Ferron and Ouellet’s (1992) estimate is smaller than the allometric home range for snowshoe hares [4.5 ha, assuming an average mass of 1,400 g (Boutin et al. 1986) and using the equation for primary consumers from Harestad and Bunnell 1979], but note that Ferron and Ouellet (1992) did not estimate an annual range. Studies conducted over longer time periods have demonstrated larger home ranges. For example, Dolbeer and Clark (1975) estimated a home range of 8.1 ha using mark-recapture techniques from mid-April to early September in Colorado. Similarly, Sievert and Keith (1985) documented annual home ranges >10 ha in Wisconsin. Neither of these studies occurred in what would be considered optimum habitat conditions (Dolbeer and Clark 1975, Sievert and Keith 1985), thus, for an annual estimate of snowshoe hare home range in optimal habitat, the allometric scale (4.5 ha) seems to be a reasonable minimum area threshold (Fig. 5).

The maximum documented home range (excluding nomadic individuals) is 16 ha (Behrend 1962). Behrend’s (1962) study occurred at the southern edge of snowshoe hare range in presumably fragmented and thus sub-optimal habitat (Sievert and Keith 1985). Sievert and Keith (1985), working in fragmented habitats in Wisconsin, also documented home ranges >10 ha in size. Based on assumptions between home range size and expected productivity (see Fig. 5), this
model assumes that hares exhibiting home ranges of 16 ha or larger are not contributing to the viability of the population (see Roloff and Haufler 1997).

Habitat quality thresholds (Roloff and Haufler 1997) were inferred by comparing home range size to hare productivity under the assumption that larger home ranges correspond to lower quality habitats and thus lower productivity (Fig. 5). The maximum annual productivity of snowshoe hares (18 young/female) has been recorded from the center of their geographic range (i.e., central Alberta) in what many assume to be an area of high quality habitats (Cary and Keith 1979). This model assumes that maximum reproductive output corresponds to optimum habitat conditions, that habitat quality scores scale linearly with reproductive output, and that the maximum documented home range corresponds to habitat quality in which a female only replaces herself annually (i.e., 2 offspring per year assuming a 50-50 sex ratio)(Fig. 5). Using documented productivity rates and estimates of home range area, a viability relationship was established for snowshoe hare (Fig. 5).

A 4.5 ha (corresponding to the allometric home range of hares) area under optimal habitat conditions (i.e., HSI score = 100) provides 4.5 habitat units for hares (consistent with the U.S. Fish and Wildlife Service’s process for calculating habitat units). The accumulation of habitat units to the 4.5 target occurs in a grid- environment and is based on the snowshoe hare HSI score (see Roloff and Haufler 1997). The higher the quality of the habitat, the fewer number of pixels required to achieve the habitat unit target. As habitat quality declines, more pixels are required and home range size increases (Roloff and Haufler 1997). Using the viability relationship developed for snowshoe hares (Fig. 5) and the output from the snowshoe hare HSI model, the forage potential of each lynx home range is scored according to the number of viable, marginal, and non-viable hare ranges it encompasses (Fig. 6). Hare home ranges above the viability threshold (60 HSI, Fig. 5) count double towards the home range tally whereas marginal home ranges (between 25 and 60 HSI, Fig. 5) count one each. Non-viable home ranges do not contribute towards the lynx forage score. It is estimated that lynx require about 600 g of food/day (or a hare every 2 days) to subsist during winter (Brand et al. 1976). Assuming that the winter season starts in November and extends through April (about 180 days), this would imply that 90 hares would support a lynx through winter. Thus, 90 hare home ranges in a lynx home range was considered optimum (Fig. 6). The resulting HSI score from tallying hare home ranges and applying the sum to figure 6 is the foraging score for the 250 ha lynx home range.
Lynx Denning Component

Delineation of potential lynx denning habitat is a 3-phase process: I) identify vegetation types that provide vegetative structure and size deemed suitable for denning, II) identify vegetation types that are properly arranged within a home range area, and III) identify vegetation types that provide suitable denning micro-sites. Components of suitable lynx denning habitat include: 1) vegetation cover type, 2) mesic site conditions, 3) canopy closure, 4) the area of the vegetation type, 4) juxtaposition and interspersion, and 5) the amount and arrangement of downed woody debris. These stand- and site-based components are integrated into a single estimate of denning habitat quality ($H_{SI_{lynx,den}}$) for the home range area. Management for denning habitat should also emphasize minimizing human disturbance.

PHASE I: (Vegetation type, site potential, canopy closure, and the area of the vegetation type)

Potential denning sites are initially delineated by vegetation cover type and site conditions. Vegetation types classified as forested with an average diameter of 20 cm providing $\geq 11.49 \text{ m}^2/\text{ha}$ (50 ft$^2$/ac) of basal area on mesic sites are assumed to satisfy denning requirements in the Lake States region. These areas must also have $>50\%$ canopy closure (where "canopy" is defined as trees $>5 \text{ m}$ in height) and be a minimum of 2 ha in size. Also, rock crevices, caves, and overhanging banks may be used for denning sites (Hoover and Willis 1987).

PHASE II: Juxtaposition and Interspersion

Denning sites (the 2 ha patch, rocks, crevices, or banks identified in Phase I) must be in close proximity to forage cover (Koehler and Brittell 1990). At least 50% of the perimeter of 2 ha patches identified as potential denning sites in “Phase I” must be adjacent to some form of "suitable lynx habitat" (i.e., habitat identified as denning, foraging, or travel). Also, 30% of the land within 0.8 km of the potential denning sites must be in suitable summer foraging habitat. Suitable summer foraging habitat is based on the habitat potential score calculated for the 0-1 m vertical cover measurement in the snowshoe hare model.

Snowshoe hares forage on a variety of herbaceous vegetation during the spring and summer months (Wolff 1978), and thus, hare forage is not limiting. It is assumed, however, that snowshoe hares are more vulnerable to predation in open areas (Adams 1959, Dolbeer and Clark 1975, Sievert and Keith 1985), and thus, vegetation cover for security is the limiting factor during spring and summer. Snowshoe hare summer security cover can be estimated based on the amount of cover (both live and dead) 0-1 m tall. The HSI for summer cover ($H_{SI_{hare,summer,cov}}$) is derived
from a measure of vertical cover provided by all objects within the 0-1 m height strata. Optimum summer cover for hares exists in stands providing $\geq 60\%$ vertical cover, and summer cover habitat quality is 0 when $\leq 20\%$ vertical cover exists (Fig. 7).

Map polygons with a summer forage HSI value $>10$ satisfy the lynx forage requirements. A map polygon may provide both suitable forage and denning, in which case the denning area is counted towards the 30% foraging. If these criteria are satisfied, the map polygon is a potential denning site and an assessment of downed woody debris is performed. Map polygons not identified as potential denning habitat through the first 2-phases are assigned a $HSI_{lynx,den,stand}$ value of 0. These sites are not evaluated for Phase III attributes. Also, denning structures can be constructed in association with sites that satisfy the Phase I and II criteria.

**PHASE III: Dead and Downed Woody Debris**

Dead and downed woody debris include logs, stumps, and upturned root masses (Koehler 1990, Koehler and Brittell 1990). Potential lynx dens generally consist of inter-tangled woody material with interstitial spaces large enough to provide lynx cover. Lynx dens have been described as having a high density (40 pieces per 50 m) of downed woody debris that were vertically structured 0.3-1.2 m above the ground (Brittell et al. 1989, Koehler 1990). These types of structures are often dependent on micro-site characteristics (e.g., areas susceptible to wind throw; drainages) and are often uncommon across entire forest stands, thus, it was deemed impractical to systematically sample this attribute within a stand and base the estimate on a mean value. Additionally, recent data from Montana and Maine have demonstrated that lynx exhibit greater variability in den site selection than previously thought. These studies have documented lynx denning in young forest plantations in slash piles and jackstrawed tree stems. The common theme among all lynx denning data is that some minimal amount of cover is required and that these areas have access to abundant hare populations. The lynx model assumes that habitat patches identified in Phase I and Phase II, with some occurrence of downed wood or slash, provide suitable denning sites for lynx.

Within lynx home ranges, multiple denning sites are important. Females may move kittens to better foraging areas or to avoid disturbance (Koehler and Brittell 1990). In low quality habitat, the inability of females to move kittens may increase kitten mortality (Koehler and Aubry 1994). Assuming that the majority of denning stands contain suitable micro-sites (verified by walk-through inventories), the quantity and spatial extent of denning stands is used to index denning habitat quality. An optimal home range is assumed to contain a mosaic of vegetation
types that include foraging and denning habitat (Koehler and Aubry 1994). In the lynx habitat model, the denning score in a home range is based on the average distance in a home range to denning sites. The variable $H_{SI_{lynx,den}}$ is calculated on the premise that multiple, interspersed denning sites in a home range is of better habitat quality than a home range containing few, blocked sites. To assess each home range area, a 100x100 m grid of points is overlaid and the average nearest distance to a suitable denning site from all points is calculated. Optimum denning habitat is provided when the average distance to denning sites is $\leq 400$ m and denning habitat is deemed unsuitable if average distance is $\geq 1,750$ m (Fig. 8). Under these parameters, optimum conditions roughly correspond to a denning site every 16 ha.

Lynx Interspersion Component

The interspersion component is designed to address the "travel corridor" needs of lynx (Washington Department of Wildlife 1993). Lynx travel through and on a variety of vegetation cover types and landscape features including thinned and un-thinned forested stands, regeneration, open meadows ($\leq 100$ m in width), ridges just above timberline, roads, and forest trails (Taylor and Shaw 1927, Parker 1981, Brittell et al. 1989, Koehler 1990). This model assumes lynx will traverse most cover types except open or sparsely-stocked stands $>100$ m in width. The interspersion component of this model uses a 2-step process: 1) identify areas of "non-lynx" cover, and 2) index the amount and spatial distribution of "non-lynx" habitat in the home range. "Non-lynx" habitat is defined as map polygons (or portions of map polygons) with a winter foraging or denning HSI of 0 that are:

a) permanent "openings" $>91$ m in width (e.g., meadows),  
b) map polygons with perennial vegetation $<2$ m tall and $>91$ m in width, and  
c) map polygons with $<440$ trees/ha (178 trees/ac) having a 2-3 m understory providing $<50\%$ visual obstruction.

It is important to note that some map polygons may be split during this process, (i.e., a portion of the polygon is $>91$ m in width and the other portion is $<91$ m in width). These portions need to be segregated during the analysis to reduce assessment error. Suitable travel cover is subsequently delineated as map polygons not identified as forage, denning, or "non-lynx" habitat.

The interspersion index ($H_{SI_{lynx,inter}}$) is based on the average nearest distance within a home range to "non-lynx" polygons. A systematic grid (100 x 100 m) is used to estimate the average distance to “non-lynx” polygons (Fig. 9). The $H_{SI_{lynx,inter}}$ is based on the premise that a lower average distance to "non-lynx" polygons equates to a more interspersed configuration of
habitats, and thus, to a greater probability of lynx encountering travel barriers (Fig. 10). The 100 x 100 m grid corresponds to the maximum hypothetical distance lynx will traverse without sufficient cover. Model simulations conducted on ≈5,000 ha in potential lynx habitat in northeast Washington demonstrated that the size of the sample grid had negligible impact on the average nearest distance to “non-lynx” habitats (Table 2). Of more importance is the relationship depicted in figure 10. Lynx will traverse long distances to fulfill their life requisites. For example, Brand et al. (1976) and Nellis and Keith (1968) found that lynx traveled 8.8 km hunting during hare population lows and 4.7 km when hares were plentiful. Parker et al. (1983) calculated daily cruising distances of 6.5-8.8 km in winter and 7.3-10.1 km during summer in Nova Scotia. Koehler (1990) documented females foraging 6-7 km from their den sites. The habitat model for lynx penalizes landscapes that restrict these movements. Figure 10 attempts to quantify the effects of barriers to movement on habitat quality (i.e., how often can lynx encounter movement barriers without detracting from habitat quality?). A low average nearest distance to “non-lynx” habitat in a home range (i.e., the chances of encountering a “non-lynx” polygon are high) equates to a poor habitat quality rating (Fig. 10). As with all of the relationships depicted in this model, the distances in figure 10 are believed to be conservative approximations and should be refined with empirical data.

**Computation of Overall Lynx HSI**

The 3 primary components of the lynx habitat model; foraging (HSI_{lynx,food}), denning (HSI_{lynx,den}), and interspersion (HSI_{lynx,inter}) are combined into one index value (HSI_{lynx}) depicting overall habitat suitability for lynx in the 250 ha area. All components of the lynx model are weighted equally and deemed critical for a functional home range therefore a geometric mean was used to represent the final HSI (Equation 8). The geometric mean causes the final HSI to equal 0 if any of the components equal 0. These 250 ha areas can be subsequently aggregated into home ranges of differing functionality and used for resource planning and modeling (Roloff and Haufler 1997).

**Equation 8:**

\[(HSI_{lynx, \text{food}} \times HSI_{lynx, \text{den}} \times HSI_{lynx, \text{inter}})^{0.33} = HSI_{lynx}\]
Literature Review


<table>
<thead>
<tr>
<th>Tree Species</th>
<th>Un-browsed or used &lt; expected</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scotch pine (<em>Pinus sylvestris</em>) (NY)</td>
<td></td>
</tr>
<tr>
<td>striped maple (<em>Acer pensylvanicum</em>) (PA)</td>
<td></td>
</tr>
<tr>
<td>yellow birch (<em>Betula alleghaniensis</em>) (PA, Nova Scotia)</td>
<td></td>
</tr>
<tr>
<td>grey willow (<em>Salix glauca</em>) (Yukon)</td>
<td></td>
</tr>
<tr>
<td>bog birch (<em>Betula glandulosa</em>) (Yukon)</td>
<td></td>
</tr>
<tr>
<td>lodgepole pine (<em>Pinus contorta</em>) (B.C., WA)</td>
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<td>red pine (<em>Pinus resinosa</em>) (NY)</td>
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<td>paper birch (<em>Betula papyrifera</em>) (NY, Nova Scotia)</td>
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<td>willow (<em>Salix</em> spp.) (MN, UT)</td>
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<td>Douglas-fir (<em>Pseudotsuga menziesii</em>) (UT, WA, MT)</td>
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<td>red alder (<em>Alnus rubra</em>) (WA)</td>
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<tr>
<td>ponderosa pine (<em>Pinus ponderosa</em>) (MT)</td>
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<td>western serviceberry (<em>Amelanchier alnifolia</em>) (MT, WA)</td>
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<td>white cedar (<em>Thuja occidentalis</em>) (Nova Scotia)</td>
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<tr>
<td>balsam fir (<em>Abies balsamifera</em>) (Nova Scotia)</td>
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<tr>
<td>Sugar maple (<em>Acer saccharum</em>) (NY, PA)</td>
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<td>beech (<em>Fagus grandifolia</em>) (NY, PA)</td>
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<td>pin cherry (<em>Prunus pensylvanica</em>) (PA)</td>
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<tr>
<td>alder (<em>Alnus</em> spp.) (WA)</td>
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</tr>
</tbody>
</table>

Shrub Species:
- ocean spray
- ninebark (*Physocarpus* spp.) (WA)
- squaw berry
- rose (*Rosa* spp.) (WA)
- bearberry (*Arctostaphysus uva-ursi*) (WA)
- Oregon grape (*Berbis repens*) (WA)
- boxwood (*Pachystima myrsinites*) (WA)

Table 2. Effect of grid cell size on the computation of mean nearest distance to non-lynx habitat.

<table>
<thead>
<tr>
<th>Grid Cell Size (m)</th>
<th>No. Points Generated</th>
<th>Mean Nearest Distance (m)</th>
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<tbody>
<tr>
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</tr>
<tr>
<td>100x100</td>
<td>4,977</td>
<td>1,384</td>
</tr>
<tr>
<td>500x500</td>
<td>197</td>
<td>1,356</td>
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<tr>
<td>1000x1000</td>
<td>47</td>
<td>1,303</td>
</tr>
</tbody>
</table>
Figure 1. Relationship between horizontal cover of browse and HSI score for 0-1 and 1-2 m height strata. Line equation between 10 and 35% is $y = 4x-40$.

Figure 2. Relationship between vertical cover of browse and HSI score for 0-1 and 1-2 m height strata. Line equation between 20 and 80% is $y = 1.666x-33.33$. 

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Figure 3. Relationship between horizontal cover and HSI score for 0-1 and 1-2 m height strata. Line equation between 40 and 90% is $y = 2x - 80$.

Figure 4. Relationship between vertical cover and HSI score for 0-1 and 1-2 m height strata. Line equation between 40 and 90% is $y = 2x - 80$. 
Figure 5. Viability relationship for snowshoe hare developed for the lynx habitat model. Viability threshold was established at 12 young/year (60 score). The marginal threshold was established at 5 young/year (25 score).

- **a** Annual productivity assumed to be 2 in that females are only replacing themselves.
- **b** Home range presented as >10 ha in literature. Here assumed as 11 ha.
- **c** Inferred home range size based on allometric theory.
Figure 6. Relationship between the number of snowshoe hare home ranges in a lynx allometric home range and the lynx forage HSI score. Line equation between 0 and 90 home ranges is \( y = 1.1x \).

Figure 7. Relationship between summer security cover and HSI score for the 0-1 m height strata. Line equation between 20 and 60% is \( y = 0.25x-50 \).
Figure 8. Relationship between the average distance to suitable denning sites in a 250 ha lynx home range and HSI score. Line equation between 400 and 1,750 m is $y = -0.0741x + 129.64$.

Figure 9. Calculating the average distance to non-lynx habitat using a 100x100 m sample grid. Distance from each grid intersection to the nearest non-lynx habitat is measured.
Figure 10. Relationship between the average distance to “non-lynx” habitat in a 250 ha lynx home range and HSI score. Line equation between 300 and 1500 m is $y = 0.08333x - 25$. 

![Graph showing the relationship between average distance to non-lynx habitat and HSI score.](image-url)
APPENDIX B:

SCHEMATIC OF PROCESSES USED FOR THE LYNX MODEL

Combine spatial layers describing attributes of vegetation and physiography

- current land cover
- historic land cover
- soils
- ecoregions

Quantify habitat quality across ecological land classes for each lynx model component

FORAGING

Hare Habitat

Hare Home Ranges

Denning Quality

Interspersion Quality

Combine grids to compute final grid of lynx habitat quality

Lynx Habitat Quality

Lynx Home Ranges
APPENDIX C:

ACCURACY ASSESSMENT OF IFMAP LANDCOVER

A land-cover dataset for Michigan was created through the 2001 IFMAP project of the Michigan DNR, and used in the Michigan GAP Analysis Project (Donovan 2005). The map is widely used as the source of land-cover information for the state, with an estimated accuracy of 87% at Anderson Level II and 37-87% at level III, and can be obtained from the Geographic Data Library of the Michigan DNR (MiGDL) on their website (<http://www.mcgi.state.mi.us/mgdl/>). The version that is released by the MiGDL contains Anderson Level III cover classes for deciduous (e.g., northern hardwoods, oak, aspen, other upland deciduous, mixed upland deciduous, lowland deciduous) and coniferous (e.g., pines, other upland conifers, mixed upland conifers, lowland coniferous, lowland mixed) forests. A problem with using these cover classes for habitat assessment is the variability in their accuracy, which could lead to erroneous conclusions about the distribution of forest resources on the landscape.

Subdedi (2005) examined the accuracy of the IFMAP forest classifications on public lands using Forest Inventory and Analysis (FIA) plots collected by the Forest Service as ground references. He found overall accuracies of 63.6% on state forests and 64.8% on national forests, when using the Anderson Level III categories provided by the IFMAP layer. User’s accuracies ranged from 17.6-60.0% for aspen and 23.5-45.7% for oak on public lands. These results suggest that the map predictions of certain forest types may not be very useful at the Level III category.

I conducted a similar analysis to that of Subdedi (2005), using FIA plots on all ownerships as ground references for the IFMAP forest classes in the entire UP. At the
time of the analysis, the plots that were available included 80% of the 6th cycle for Michigan. Plots were filtered to include only those that were forested (>25% tree cover) and containing a single condition, to ensure that canopy information was consistent throughout the 4 subplots. This resulted in 2,328 plots spread across the UP. The tree information for each plot was used to calculate the percent cover of canopy trees for each species, and plots were labeled according to the classification rules used by the IFMAP project (MDNR 2001). The FIA plots were overlaid with a grid of the land-cover dataset by the North Central Spatial Data Services (SDS) of the Forest Service, and the cover type for each plot was determined by the majority of pixels within a 3x3 window around each plot location. An error matrix was constructed to compare the IFMAP classifications for each plot, as determined by the map and tree list data (Table C.1).

The overall accuracy of the IFMAP forest classification was estimated at 45.9%, with relatively poor user’s accuracies for most forest classes other than northern hardwoods (79.8%) (Table C.1). After grouping IFMAP classes according broader categories of forest type (Table 3), overall accuracy increased to 74.3%.

Our results further illustrate the potential problems with using the IFMAP land-cover dataset for mapping forest types in the UP. The heterogeneity of forest types across the landscape of the UP might explain the difficulty in accurately predicting their occurrence using satellite imagery. Aspen forest types that are being succeeded by conifers (e.g., balsam fir) may also present problems during interpretation. My analysis utilized a filtered set of FIA plots, which may not have provided the best representation of reference points. Using all FIA plots may have yielded different results, though plots with multiple conditions would likely have decreased the accuracy of IFMAP predictions.
Table C.1. Error matrix of forest classifications between IFMAP and FIA surveys in the Upper Peninsula.

<table>
<thead>
<tr>
<th>code</th>
<th>IFMAP cover</th>
<th>14</th>
<th>15</th>
<th>16</th>
<th>18</th>
<th>19</th>
<th>20</th>
<th>21</th>
<th>22</th>
<th>24</th>
<th>25</th>
<th>26</th>
<th>total</th>
<th>UAa (%)</th>
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<td>681</td>
<td>1</td>
<td>50</td>
<td>81</td>
<td>1</td>
<td>4</td>
<td>1</td>
<td>23</td>
<td>1</td>
<td>1</td>
<td>853</td>
<td>79.8</td>
<td></td>
</tr>
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<td>0</td>
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<td>4</td>
<td>0</td>
<td>0</td>
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<tr>
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<td>aspen</td>
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<td>4</td>
<td>6</td>
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<td>26</td>
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<td>4</td>
<td>304</td>
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<td>1</td>
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<td>14</td>
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<td>3</td>
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<td>1</td>
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<td>119</td>
<td>299</td>
<td>110</td>
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</table>

PA\(^b\) (%)

\(^a\)UA = user’s accuracy

\(^b\)PA = producer’s accuracy
APPENDIX D:

CALCULATING HORIZONTAL COVER WITH THE
STAND VISUALIZATION SYSTEM

Horizontal cover has been demonstrated as an important habitat component for snowshoe hare (Brocke 1975, Wolff 1980, Litvaitis et al. 1985, Parker 1986, Hodges 2000), and many other wildlife, as it can provide a range of life history requirements, including thermal and escape cover, and available browse. Common measurements of horizontal cover involve counting the proportion of covered grid squares on a profile board held perpendicular to the ground and viewed from a specific distance. This type of sampling is subject to large measurement error on the part of the observer (Collins and Becker 2001), as the decision to count a grid square as “covered” is subjective at best. Forest inventories do not typically measure horizontal cover as a stand attribute, which limits their use in assessing habitat for species that require it.

The Stand Visualization System (SVS) (McGaughey 1997) is a software program that creates 3-dimensional renderings of 0.40 ha stands (Figure D.1), using forest inventory data for input. SVS has the ability to interpret a number of common sampling

Figure D.1  SVS drawings of a mature red pine stand with hardwood understory.
designs that are used for forest inventory, and when incorporated with the Forest Vegetation Simulator (FVS), stands can be grown and subjected to harvest scenarios over time. SVS contains a number of tree form files that outline parameters used to create a realistic diagram for each tree species. Parameters can be manipulated by the user and include the shape of the crown, the number, positioning, and color of branches and leaves, and the manner in which these parameters change with height and diameter. For example, the leaves on deciduous stems can be removed to simulate the appearance of leaf-off season. The profile view illustrates all trees within a user-specified strip width, which can simulate the horizontal cover provided at the specified distance.

I developed a method to estimate horizontal cover (within 0-1 m and 1-2 m from the ground) using the profile view of an image in order to assess the quality of forest stands for winter hare habitat. SVS images were created for a series of FIA plots using the post-processing function of FVS, and tree forms for each deciduous species were altered to remove leaf cover. The view azimuth was changed to 0° for the perspective view, which placed the range poles at either end of the profile image. Images were saved as 1280 x 960 Windows bitmaps in SVS. A single image was examined to determine the pixel coordinates of a box encompassing each height stratum (Figure D.2). The 3 m increments on the range poles measured 28 pixels, which meant the height of each 1 m stratum would approximate to 9 pixels, with a length between range poles of 642 pixels. These dimensions should be consistent for all SVS images that are saved as 1280 x 960 bitmaps. Each image was converted to a monochrome portable bitmap and read into R as a text file. The file contained a matrix of binary values which represented the black (1)
and white (0) pixels from the bitmap. Dividing the sum of values within the 9 rows and 642 columns for each stratum by the total pixels (5778), resulted in the estimate of cover.

The following R code was used for the calculations:

```r
svs <- read.table("stand1.pbm", skip=32189, nrow=943)
stratum1 <- sum(sum(svs[641,24:36]), sum(svs[642:658,1:36]), sum(svs[659,1:17]),
    sum(svs[748,12:36]), sum(svs[749:765,1:36]), sum(svs[766,1:5]),
    sum(svs[783,32:36]), sum(svs[784:800,1:36]), sum(svs[801,1:25]),
    sum(svs[819,16:36]), sum(svs[820:836,1:36]), sum(svs[837,1:9]),
    sum(svs[854,36]), sum(svs[855:871,1:36]), sum(svs[872,1:29]),
    sum(svs[890,20:36]), sum(svs[891:907,1:36]), sum(svs[908,1:13]),
    sum(svs[926,4:36]), sum(svs[927:942,1:36]), sum(svs[943,1:33])) / 5778
stratum2 <- sum(sum(svs[1,24:36]), sum(svs[2:18,1:36]), sum(svs[19,1:17]),
    sum(svs[37,8:36]), sum(svs[38:54,1:36]), sum(svs[55,1]),
    sum(svs[72,28:36]), sum(svs[73:89,1:36]), sum(svs[90,1:21]),
    sum(svs[108,12:36]), sum(svs[109:125,1:36]), sum(svs[126,1:5]),
    sum(svs[143,32:36]), sum(svs[144:160,1:36]), sum(svs[161,1:25]),
    sum(svs[179,16:36]), sum(svs[180:196,1:36]), sum(svs[197,1:9]),
    sum(svs[214,36]), sum(svs[215:231,1:36]), sum(svs[232,1:29]),
    sum(svs[250,20:36]), sum(svs[251:267,1:36]), sum(svs[268,1:13]),
    sum(svs[286,4:36]), sum(svs[287:302,1:36]), sum(svs[303,1:33])) / 5778

stratum1 = 0.5896504 (59%)  stratum2 = 0.5705434 (57%)
```
Figure D.2. Profile view of stand with a red box around each height stratum.
APPENDIX E:

LANDSCAPE-SCALE MODELING OF FOREST STRUCTURE USING LANDSAT TM IMAGERY AND FIA DATA

The mapping of forested land using remotely-sensed imagery has become a standard procedure in the management of natural resources at the landscape scale. These classified maps of land cover contain varying levels of detail and accuracy, depending on the extent of the mapped landscape and the resolution of the imagery, but the amount of information they can relay is limited. Evaluations of wildlife habitat often require fine-scale measurements of vegetation structure, which are not provided by land-cover datasets. Obtaining this type of information for forest inventories can be costly for large landscapes, involving a combination of high resolution aerial photography and field-based measurements at the stand level. Even when this information becomes available, the temporal resolution of data acquisition may be too coarse to capture alterations in vegetation structure due to growth and disturbance through time.

Our objective was to map the local variation in forest structure across the UP, which required the integration of remotely-sensed imagery with field surveys to utilize relationships between vegetation structure and spectral reflectance from the ground. The $k$-nearest neighbor (KNN) method assigns attribute values to non-sampled pixels from those that are sampled (i.e., have an associated ground plot) based on the distance between the pixels in a multi-dimensional feature space, as defined by the combination of spectral values in each wavelength band of a remotely-sensed image. This method has been used to predict forest attributes across large landscapes (Tomppo 1991, Fazakas and Nilsson 1996, Tokola et al. 1996, Franco-Lopez et al. 2001), and in the Great Lakes
region with Forest Inventory and Analysis (FIA) plots serving as ground references (McRoberts et al. 2002, Haapanen et al. 2004). I acquired Enhanced Thematic Mapper Plus (ETM+) imagery from the Landsat 7 satellite to model forest structure using the KNN method, with ground references provided by the FIA plots collected between 2000-2003 for the 6th inventory of Michigan.

The UP required 8 scenes of Landsat imagery for complete land coverage, including row 28 of paths 21-22, rows 28-29 of path 23, and rows 27-28 of paths 24-25 (Figure E.1). Images were acquired from a range of dates between 2000-2002 due to the limited availability of cloud-free imagery; images 5 and 6 were from late August (8/20),

![Figure E.1](image-url)
images 3 and 4 from early September (9/08), images 1, 2 and 8 from mid to late September (9/17-9/29), and image 7 from early October (10/11). Each image was rectified and geo-referenced to zone 16 of the UTM system, using the GRS 1980 spheroid and NAD83 datum. A vector layer of roads from the Michigan Center for Geographic Information (MCGI) was used for geo-referencing. Small clouds or areas of haze were masked out and each image was clipped to the boundaries of the UP. Three pairs of images were successfully mosaicked together, and the final 5 images were named as follows: EAST = images 1 + 2, MIDEAST = images 3 + 4, MIDWEST = images 5 + 6, WEST1 = image 7, WEST2 = image 8.

The images were transferred to Geoff Holden (USFS) at the Spatial Data Services (SDS) of the North Central Research Station, where FIA plot locations were overlaid with the spectral imagery and values of plot attributes were imputed across the landscape using the KNN method. Spectral values from bands 1-5 and 7 were used to define the feature space, and a 3 x 3 mean filter was applied to each image to match the 4-subplot sampling design of the FIA plots. The number of nearest neighbors \((k)\) was set to 5 and the distance metric was Euclidean. The prediction accuracy of the model was estimated by holding out 40% of the FIA plots from the training set and comparing the observed with predicted pixel values using error matrices and linear regression.

In the first iteration, the plot attribute was categorical, describing the structural stage of each plot as defined by the Forest Vegetation Simulator (FVS). The stages were based on the size of trees and number of height strata, with dbh thresholds for sapling/pole and pole/large set at 12.7 cm (5 in) and 30.48 cm (12 in), respectively, and percent cover of valid height stratum set to 20 (Crookston and Stage 1999). Several
combinations of structural stages were used to vary the number of attribute categories (i.e., 4 stages versus 6 stages), but the KNN procedure was unable to produce an overall classification accuracy >40%.

The second iteration involved plot attributes that were continuous, including horizontal cover (as measured by FVS) and basal area. Horizontal cover was corrected for normality using the arcsine transformation, after converting it to a proportion, while basal area was determined to be normally distributed. Initial examination of basal area predictions on a single image were comparable to that of horizontal cover; therefore, basal area was not assessed any further. Predictions of horizontal cover were developed for each of the 5 images; all had poor relationships between predicted and observed values, with most having an adjusted R² <0.10 (Figure E.2).

Our inability to accurately predict horizontal cover using the Landsat imagery could be attributed to a number of different reasons. The data provided by the 6 ETM+ bands were most likely insufficient for capturing any differences in reflectance caused by vegetation structure, especially that of the understory. The analysis was limited to one date for each region due to the availability of satellite imagery (determined by acquisition cost and cloud cover), and the range of acquisition dates (20 August – 11 October) could arguably have spanned multiple seasons. Using 3 or more dates of imagery that spanned the growing season of vegetation, as well as ancillary GIS data that stratified the landscape based on multiple ecological characteristics (i.e., topographical position, ecoregion), may have improved the predictions (Franco-Lopez et al. 2001). The KNN method assumes homogeneity in forest composition across the analysis area, which was not the case for most of the UP. In spite of this, the greatest prediction accuracy came
Figure E.2. Relationship between predicted and observed values of arcsine transformed horizontal cover for each of the 5 spectral images in the Upper Peninsula.
from the EAST image, which had a far greater contrast in forest types than the areas captured by the western images. A stratification of the landscape, other than that created by the Landsat scenes, was not practical due to the low sampling intensity of the FIA plots. A potential source of error in the methodology was in matching FIA plots to pixels within the satellite image, as plots cannot be expected to lie perfectly centered within the 3 x 3 pixel windows, which may effect the assignment of spectral values. Finally, the measure of horizontal cover used as the dependent variable was an estimated value with an unknown accuracy; any hypothesized relationship between horizontal cover and the spectral reflectance of the vegetation would rely on an accurate estimate.

Earlier examinations of the KNN method for classifying forest structure have had some success (Franco-Lopez 2001), and the method has proven useful in predicting compositional attributes across large landscapes (McRoberts et al. 2002). The application of this method to predictions of understory structure using Landsat imagery is dependent on a relationship between understory vegetation and the reflectance of the canopy. The use of high resolution imagery and/or light detection and ranging (lidar) data in conjunction with Landsat imagery may provide a more accurate means of predicting forest structure across large landscapes (Lefsky et al. 1999).
## APPENDIX F:

### METADATA FOR SPATIAL LAYERS

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<th>Type</th>
<th>Data Layer</th>
<th>Description</th>
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<td>IFMAP/GAP Land Cover</td>
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<td>Michigan’s Presettlement Vegetation</td>
<td>Vegetation type boundaries interpreted from GLO Surveys 1816-1856 for Michigan</td>
<td>David T. Cleland&lt;br&gt;USDA Forest Service&lt;br&gt;Southern Research Station&lt;br&gt;5985 County Road K&lt;br&gt;Rhinelander, WI 54501&lt;br&gt;email: <a href="mailto:dcleland@fs.fed.us">dcleland@fs.fed.us</a>&lt;br&gt;<a href="http://www.ncrs.fs.fed.us/gla/">website</a></td>
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<td>Daniel W. Linden&lt;br&gt;Dept. of Fisheries and Wildlife&lt;br&gt;13 Natural Resources Bldg.&lt;br&gt;Michigan State University&lt;br&gt;East Lansing, MI 48824&lt;br&gt;email: <a href="mailto:lindend1@msu.edu">lindend1@msu.edu</a></td>
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