

Research, part of a Special Feature on [The Many Facets of Forest Resilience in the Lake Tahoe Basin](#)

Simulating wildlife habitat dynamics over the next century to help inform best management strategies for biodiversity in the Lake Tahoe Basin, California

[Angela M. White](#)¹, [Tim G. Holland](#)², [Eric S. Abelson](#)^{1,3}, [Alec Kretchun](#)⁴, [Charles J. Maxwell](#)⁴ and [Robert M. Scheller](#)⁴

ABSTRACT. Many forests of the western United States have undergone over one hundred years of anthropogenic impacts that have led to increased tree density, homogenization in forest structure, and accumulation of woody material, which combined with a changing climate pose threats to valued social and ecological features. In California, recent waves of tree mortality and unprecedented large and destructive fires have led to rising concerns about the impact of these disturbances on biodiversity and how forest management actions can mitigate negative impacts. To better understand the degree to which different management scenarios could mitigate the negative impacts of these disturbances on biodiversity, we used a spatially explicit modeling platform to model forest management impacts on habitat for terrestrial vertebrate species in the Lake Tahoe Basin of California and Nevada. Specifically, we modeled how 5 different management scenarios that differed in the type of fuel reduction treatment (e.g., fire and mechanical removal of vegetation) and extent of area treated influenced the amount, value, and distribution of reproductive habitat for the 159 species present in the study area. Our model results suggested that within the study area forest growth was predicted to out-pace disturbance leading to a higher percentage of late seral conditions; however, choice of management strategy impacted the composition and structure of the forested landscape leading to different trajectories for wildlife. In general, scenarios that allowed for more extensive use of fire led to a more equitable distribution of habitat types, whereas extensive thinning by hand and mechanical methods resulted in future forest structure that provided better outcomes in terms of reproductive habitat for wildlife. Our modeling results also suggested that low to moderate management strategies were not likely to change the current trajectory to more dense forests dominated by fewer species.

Key Words: *biodiversity; climate change; dynamic modeling; fire; forest management; resilience; wildlife habitat*

INTRODUCTION

In the seasonally dry forests of the western United States, fire is a natural disturbance that shapes habitat composition, structure, and distribution (Larson and Churchill 2012, Knapp et al. 2013, Lutz et al. 2018). In the mid-elevation forests of the Sierra Nevada Mountains in California, fire historically maintained a species-rich mixture of tree densities and size classes (Barbour et al. 2002, Beaty and Taylor 2007, North et al. 2009), but due to past practices such as logging, grazing, and highly effective fire suppression, mixed-conifer forests have become denser and simplified in structure and composition (Knapp et al. 2013, Stephens et al. 2013). Coupled with rapid changes in climate, these changes in forest structure have contributed to large-scale susceptibility of trees to bark beetle attacks during drought (Young et al. 2017) and massive, high-severity fires (Lydersen et al. 2014, Young et al. 2017). In California, these large-scale disturbances threaten the State's economy because these forests not only provide woody material, clean water, and abundant recreational opportunities, but also serve as habitat for a diverse community of plants and animals (Mayer and Laudenslayer 1988). The rapid change in these forests underscores the need to increase the pace and scale of forest treatments designed to reduce woody material and increase forest resiliency to future, large-scale disturbances (North et al. 2012, Stephens et al. 2013).

Forest treatments designed to reduce the risk of high-severity fire commonly focus on reducing the amount and connectivity of woody material on the forest floor (e.g., branches, logs, and shrubs), between the forest floor and the canopy (e.g., shrubs and small trees), and within the canopy (e.g., smaller, subdominant canopy trees; Agee and Skinner 2005). To implement large-scale

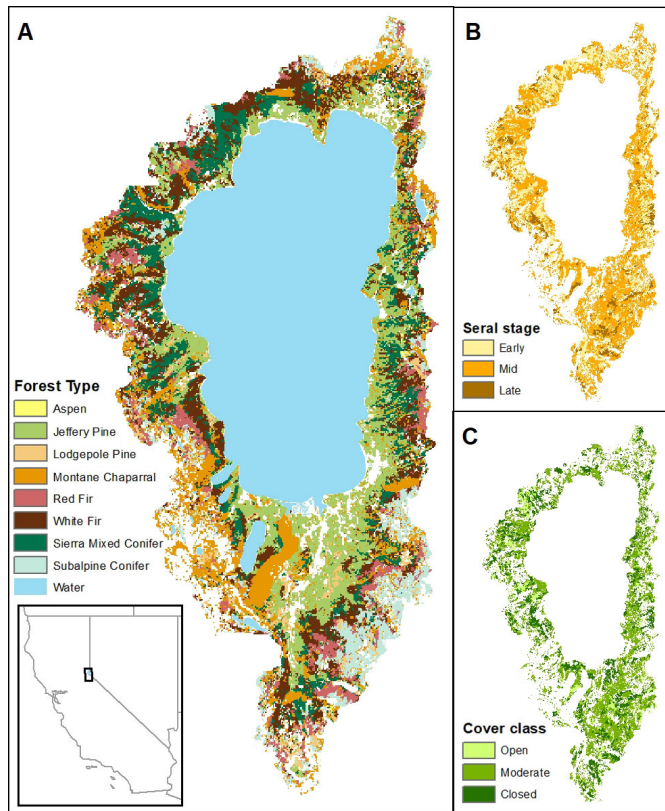
forest restoration projects will require not only mechanical thinning of trees and shrubs, but due to the relatively small percentage of forest that can be mechanically treated because of logistical and fiscal constraints (North et al. 2015), will likely require the use of prescribed burning and managed wildfire (North et al. 2012). Although data collected from forested areas receiving both mechanical/hand removal of vegetation and prescribed burning treatments have been shown to have neutral to positive effects on many wildlife species (Fontaine and Kennedy 2012, Stephens et al. 2014), many species thrive in forests burning with moderate to high-severity fire effects (Smucker et al. 2005, Tingley et al. 2016, White et al. 2016), higher cover of shrubs (White et al. 2013), or complex, early seral habitat conditions (Swanson et al. 2011). Collectively, these studies confirm that wildlife response to fuel reduction treatments will depend on the type of disturbance or restoration action and on the resulting structure.

Forest structure, and its composition, provide the environmental conditions needed to support essential food, resting, and breeding requirements for wildlife. Changes to forest structure inherently impact the degree to which environmental conditions provide suitable habitat, i.e., improving habitat conditions for some species, while reducing it for others (Fontaine and Kennedy 2012, White et al. 2013, Stephens et al. 2014). The California Wildlife Habitat Relationships (CWHR) database (<https://www.wildlife.ca.gov/Data/CWHR>) classifies the suitability of habitat for vertebrate species based on predominate vegetation (tree, shrub, herbaceous, etc.), species composition, and stage of vegetation development (Fig. 1). For each combination of these conditions, CWHR provides a habitat suitability classification (high,

¹US Forest Service, Pacific Southwest Research Station, Davis, California, ²Tetra Tech, ARD, Burlington, Vermont, ³University of Texas, Austin, Integrative Biology, Austin, Texas, ⁴Department of Forestry and Environmental Resources, NCSU, Raleigh, North Carolina

moderate, low, unsuitable) for species based on expert-solicited information (Fig. 2A). These broad habitat classifications make it possible to predict habitat suitability for a rich diversity of wildlife species over large areas and timescales where data exist on vegetation type, structure, and seral stage.

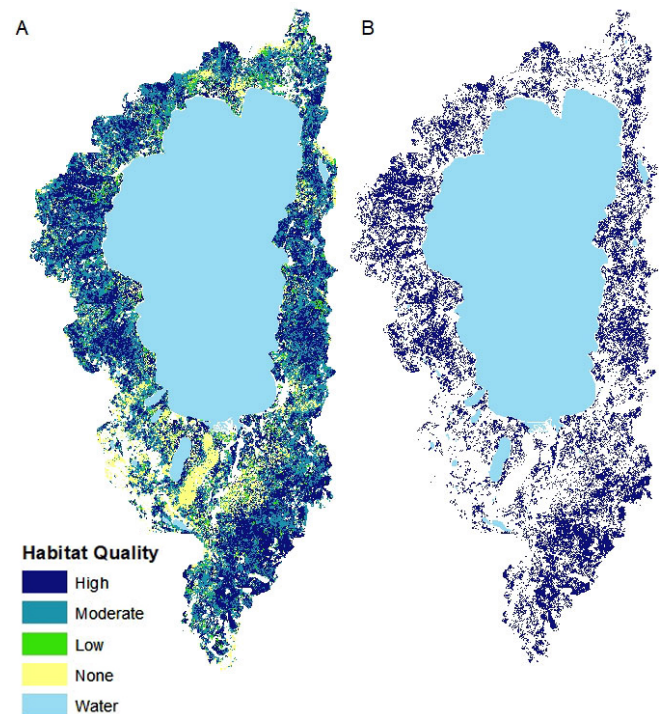
Fig. 1. Location of the Lake Tahoe Basin depicting (A) the distribution of modeled habitat types, (B) seral stage, and (C) canopy cover class at the start of LANDIS-II modeling simulations in 2010.



We build on previous efforts to better understand how the type and intensity of management treatments are likely to influence wildlife habitat (e.g., Fontaine and Kennedy 2012, White et al. 2013, Stephens et al. 2014) over large spatial and temporal scales. Although providing for habitat connectivity and a mosaic of forest conditions provides general principles for landscape scale wildlife management, specific targets are rarely quantified and will differ with species and existing forest conditions (North and Manley 2012). Using a modeling framework, we assess how different types of forest restoration treatments (i.e., fire suppression, mechanical and hand thinning, prescribed burning, and managed wildfire) and placement, i.e., wildland-urban interface (WUI) or general forest, impact the quality and distribution of reproductive habitat for terrestrial vertebrate species in the Lake Tahoe Basin of California over the next century. We predicted that management scenarios incorporating managed wildfire would lead to greater structure complexity in forest types and subsequently provide greater benefits to wildlife than management scenarios focusing on mechanical treatments in the wildland-urban interface. This effort is part of a larger initiative led by the Lake Tahoe West Restoration Partnership

to restore the social-ecological resilience of forests, watersheds, and communities on 59,000 acres of Lake Tahoe's west shore (Abelson et al 2022).

Fig. 2. An example of the outputs used for assessing wildlife metrics: (A) total value of reproductive habitat on the landscape, and (B) patches of high-quality reproductive habitat. This figure is of replicate 1 of the Northern flying squirrel (*Glaucomys sabrinus*) under the Thin scenario in 2070.



METHODS

The Lake Tahoe Basin, located on the eastern crest of the Sierra Nevada Mountains straddling the states of California and Nevada, supports high levels of biodiversity (Fig. 1). The location of the basin between two major biogeographic provinces, the Sierra Nevada and the Great Basin, creates a juxtaposition of habitats, each province of which supports a unique array of species. In addition, the basin's steep elevational gradients (ranging from 1900 m at Lake Tahoe's surface to 3400 m at the highest mountain peak) span lower montane, upper montane, subalpine, and alpine habitats. Within these habitats, differences in forest structure and composition further influence the provisioning of resources used by wildlife for reproduction, foraging, and cover. Our assessment focused on the habitat of 159 vertebrate species that have suitable habitat for breeding in the Lake Tahoe Basin.

Dynamic modeling of habitat

To model habitat change over time, we used the landscape disturbance and succession model LANDIS-II (<https://www.landis-ii.org/>) to simulate forest change as a function of growth and succession (Mladenoff 2004, Scheller et al. 2007). Within

LANDIS-II, trees and shrubs are modeled as species-age cohorts; each species has its own unique life history attributes and as such, each responds uniquely to disturbance. Every cell (1-ha resolution) across the landscape has a unique combination of cohorts, live biomass, dead material, and soils as determined by its simulated history of succession and disturbance type and intensity. In addition to fire and management, several species of bark beetles, including the Jeffrey pine beetle (*Dendroctonus jeffreyi*), Mountain pine beetle (*D. ponderosae*), and fir engraver (*Scolytus ventralis*) are a frequent disturbance in the study area that was modeled. Due to previous efforts, LANDIS-II has been calibrated to conditions in the Lake Tahoe Basin (Loudermilk et al. 2013, 2014, Scheller et al. 2019).

Using the CWHR classification scheme, we developed a rule set for LANDIS-II biomass outputs that would approximate CWHR habitats (Appendix 1). For this study, we modeled change in tree-dominated and shrub-dominated CWHR habitats (Fig. 1A). Areas that would be classified as aquatic, developed, or barren following CWHR classification rules were excluded from the landscape analysis because these areas were not projected to change over time. In addition, LANDIS-II does not model the herbaceous community, and we classified herbaceous-dominated habitat structure according to the woody vegetation component (i.e., tree or shrub-dominated) or as NULL following recent disturbances that had removed all woody vegetation. The shrub component of different habitat types can vary, and we classified an area as shrub dominated when $\geq 75\%$ of the total biomass was attributed to shrubs (Appendix 1). This biomass threshold was chosen based on LANDIS-II biomass estimates of the current landscape compared to current classification of CWHR shrub-dominated habitats. Species composition in tree-dominated habitats followed the CWHR classification scheme but relied on species-specific biomass estimates instead of estimates of canopy cover. Species-specific biomass estimates produced by LANDIS-II to classify composition included: white fir (*Abies concolor*), red fir (*A. magnifica*), lodgepole pine (*Pinus contorta*), Jeffrey pine (*P. jeffreyi*), incense-cedar (*Calocedrus decurrens*), sugar pine (*Pinus lambertiana*), mountain hemlock (*Tsuga mertensiana*), western white pine (*P. monticola*), and whitebark pine (*P. albicaulis*). Aspen (*Populus tremuloides*) is the only hardwood species that occurs regularly within the study area and was the only tree-dominated habitat modeled that was not classified as conifer. Based on this rule set, we identified eight CWHR habitat types in our study area: White Fir (WFR), Red Fir (RFR), Jeffrey Pine (JPN), Lodgepole Pine (LPN), Sierra Mixed Conifer (SMC), Subalpine Conifer (SCN), Aspen (ASP), and Montane Chaparral (MCP).

For conifer dominated habitat types, we assigned each habitat type a seral stage based on CWHR size class descriptions (Fig. 1B). There are five CWHR size classes that influence habitat suitability for wildlife based on the mean diameter at breast height (DBH) of predominant trees. Seedling trees are the first size class and are defined by trees with a mean diameter < 2.5 cm, followed by sapling trees (size class 2: 2.5-15 cm), pole trees (size class 3: 15-28 cm), small trees (size class 4: 28-61 cm), and med/large trees (size class 5: > 61 cm). We estimated these size classes by combining estimates from LANDIS-II for the biomass of different species age cohorts with data from Forest Inventory and Analysis (FIA; Burrill et al. 2021) on the distribution of stem sizes

(for equivalent species and age) in trees that have been measured in the northern Sierra Nevada ecoregion where Lake Tahoe is situated. We fit the Weibull distribution to FIA tree data for species-age groups that were equivalent to each LANDIS-II cohort. We assumed that the LANDIS-II cohorts would follow a similar size distribution to what was observed in the FIA data and distributed the LANDIS-II biomass estimates among two-inch DBH height bins (i.e., into a 0-5 cm bin, a 5-10 cm bin, etc.). To assign a pixel to a CWHR size class, we used the DBH of the 95th percentile tree in our estimated distribution and compared that to the CWHR size classes as described above (classes 1 through 5 being < 2.5 cm; 2.5-15 cm; 15-28 cm; 28-61 cm; and > 61 cm). We assessed different approaches, specifically, we also completed the classification using the size of the 90th percentile tree and the 99th percentile tree, and we found that using the 95th percentile yielded a CWHR in year 1 (2010) of the LANDIS-II model run that most closely reflected our baseline habitat distribution in the Lake Tahoe Basin. We considered this method reliable in producing three distinct tree-dominated seral stages: early (tree size approximating size classes 1-3), mid (tree size approximating size class 4), and late (tree size approximating size class 5).

Canopy cover in early seral habitat did not have a predictable relationship with canopy cover estimates, but for mid and late conifer-dominated habitats, we additionally assigned each pixel a canopy cover estimate (open, moderate, closed) based on CWHR canopy classes (Fig. 1C). Using the biomass bins described above, we again used the FIA data for average stem to biomass relationships at different size classes to translate biomass values into stem counts for each size bin (Evans et al. 2022). This process gave us an estimated size distribution of trees on each pixel as well as the estimated number of trees per pixel, which we translated into an estimate of trees per acre (TPA) to estimate canopy cover. We based our canopy thresholds on estimates of TPA whereby mid and late seral habitats with < 50 TPA were categorized as open, 51-500 TPA as moderate, and > 500 TPA as closed. Using these methods, we assigned each 1 ha pixel a CWHR habitat (composition, size class, canopy class). Although the CWHR classification scheme separates open from sparsely covered habitats, we combined these two into one category creating three canopy classes that are ecologically important and are used to establish guidelines for management. Descriptions of CWHR habitats modeled and the number of hectares of each habitat combination produced are presented in Appendix 2.

Simulated management scenarios

Management scenarios simulated in this study were selected by the Lake Tahoe Basin West Restoration Partnership to better understand the societal benefits of increasing the pace and scale of forest restoration (Table 1). Under the suppression-only (Suppress) scenario no forest management activities were modeled other than active fire suppression (i.e., if a fire started, it was suppressed). Under the wildland-urban interface (WUI) scenario, in addition to fire suppression, forest thinning (both mechanical and hand treatment) was modeled in the vegetated area adjoining structures and evacuation routes (which accounts for $\sim 58\%$ of the landscape). Similar to the WUI scenario, the thin scenario (Thin) modeled high levels of forest thinning in the WUI, as well as some treatments in the general forest and surrounding wilderness areas. The WUI and thin scenarios thus differed in the

Table 1. Names and descriptions of the five management scenarios modeled.

Scenario name	Modeled management actions	Scenario intent
Suppress	Management actions were limited to suppressing all fire activity.	Understand the societal benefits of fire suppression.
Wildland-urban interface (WUI)	In addition to fire suppression, approximately 1.8% of the vegetated area was treated annually in the WUI and defense zone.	Understand the societal benefits of current management actions focused on reducing fuels around human-built structures and evacuation zones.
Thin	In addition to fire suppression and WUI-focused fuel reduction treatments, this scenario treated approximately 6.7% of the vegetated area annually including the forest beyond the WUI.	Understand the societal benefits of expanding the use of mechanical and hand-thinning into the general forest.
Fire	Management actions were focused on reducing fuels via prescribed fire and managed wildfire with approximately 4% of the vegetated area treated each year.	Understand the societal benefits of using fire-driven fuel reduction treatments in addition to mechanical and hand-thinning mechanisms.
Fire ⁺	Management actions were focused on reducing fuels via prescribed fire and managed wildfire with approximately 7.2% of the vegetated area treated each year.	Increase the amount of vegetated area treated each year via fire-driven mechanisms (75% of all fuel reductions) to better differentiate between the societal benefits of fuel reduction mechanisms and the area treated.

amount and extent of treatment. The two remaining scenarios were modeled with the use of fire as a management tool; both modeled moderate levels of thinning in the WUI while allowing for some prescribed burns and managed wildfire ignitions outside of the WUI. The fire scenario (Fire) however, simulated moderate levels of prescribed and managed fire, whereas the Fire⁺ scenario simulated high use of prescribed and managed fire. Information on the parameterization of the model is available here: <https://github.com/LANDIS-II-Foundation/Project-Lake-Tahoe-2017>

For each of the five scenarios, annual change in forest conditions were simulated from 2010–2110 using climate data from the CanESM2 global change model based on the relative concentration pathway 4.5. Each scenario run was simulated 10 times (i.e., 10 replicates) to quantify the mean and variation in change in habitat. At each 10-year timestep, we assigned CWHR habitat types to each spatial output (5 scenarios X 10 replicates X 10 decadal time steps) resulting in 500 spatially explicit habitat maps at 1-ha resolution.

Simulating wildlife-habitat outcomes

We extracted data from CWHR (<https://www.wildlife.ca.gov/Data/CWHR>) on wildlife-habitat relationships for each vertebrate species known to occur in the Lake Tahoe Basin (Schlesinger and Romsos 2000). We excluded any species from the analysis that relied primarily on aquatic habitats (e.g., amphibians and waterfowl). This resulted in a list of 135 birds, 51 mammals, and 5 reptiles. Of these 191 species, we were unable to simulate change in reproductive habitat for 23 birds and 2 mammals because they are not known to breed in the study area, or only breed in urban environments. Additionally, we excluded species in which their reproductive habitat was rare (< 5% of the landscape) including the Loggerhead Shrike (*Lanius ludovicianus*), Bullock's Oriole (*Icterus bullockii*), White-crowned Sparrow (*Zonotrichia leucophrys*), Tree Swallow (*Tachycineta bicolor*), Lark Sparrow (*Chondestes grammacus*), and Gray-crowned Rosy Finch (*Leucosticte tephrocotis*). This resulted in 159 species in which a change in the value of habitat for reproductive purposes was modeled (Appendix 3).

For each modeled species and every habitat map (N = 500), we assigned a habitat-suitability ranking of high (1.00), moderate

(0.66), low (0.33), or unsuitable (0.00) for reproduction based on the CWHR database for each 1-ha pixel (Fig. 2A). Because our translation from LANDIS II outputs (biomass by species by age cohort) to CWHR habitat resulted in the decision to combine certain CWHR habitats (i.e., sparse and open canopy classes were combined), we assigned the maximum value of reproductive habitat (hereafter habitat) for the combined categories (Appendix 3). From each map, we generated three metrics: (1) the total sum of habitat suitability values on the landscape for each species, (2) the proportion of the landscape classified as high-value habitat (i.e., suitability = 1; Fig. 2B), and (3) the average size of habitat patches of high reproductive value using ClassStat function in package SDMTools (McGarigal et al. 2002) in Program R version 3.6.1 (R Core Team 2018). For each species*scenario*replicate*timestep, we compared the computed metric to baseline starting conditions in 2010. For each metric (overall habitat quality, proportion of landscape considered high-quality habitat, average patch size of high-quality habitat) at each scenario*replicate*timestep, we summed the number of species in which each metric was predicted to increase and decrease. Because these metrics always fluctuate due to the dynamic nature of the model, we chose an arbitrary threshold of > 30% change to quantify more significant impacts on species and quantified these species separately.

RESULTS

Changes in habitat type and composition

At the start of our simulations, the modeled landscape was dominated by Jeffrey Pine (22%), White Fir (20%), Sierra Mixed Conifer (15%), and Montane Chaparral (21%) habitat types, with Aspen, Lodgepole Pine, Red Fir, and Subalpine Conifer Habitat types each comprising less than 10% of the landscape (Fig. 3). Under all scenarios, except the Fire⁺ scenario, the amounts of White Fir and Sierra Mixed Conifer Habitat types were predicted to increase over time, becoming the dominant habitat types, whereas all other habitat types tended to decrease. In the Fire⁺ scenario, increased use of fire was predicted to have a stabilizing effect on all habitat types over time, with Montane Chaparral quickly becoming the most common habitat type. Compared to the other scenarios, the Fire⁺ scenario maintained a larger

Table 2. Comparisons of the median and range of patch sizes in 2040 and 2070 for different ecological groups of taxa under the five management scenarios relative to baseline conditions.

Group	Baseline (ha)	Scenario	2040 (ha)	2070 (ha)
Small-bodied	10.8 (6.26-17.1)	Suppress	8.9 (2.9-10.9)	7.8 (2.1-9.8)
		WUI	8.6 (3-10.7)	7 (2.2-9.9)
		Thin	9.2 (3.2-11.7)	7.1 (2.5-12.2)
		Fire	8.5 (3-10.5)	7.2 (2.2-10.3)
		Fire ⁺	6.9 (2.4-11.2)	6.8 (2.5-8.8)
Medium-bodied	9.23 (6.42-17.2)	Suppress	5.1 (4.8-11)	5.3 (3.1-9.7)
		WUI	5.7 (4.8-11.5)	5.8 (3.2-13.1)
		Thin	6.8 (5-13.1)	6.9 (3.7-13.7)
		Fire	5.4 (4.7-11.2)	6 (3.1-13.4)
		Fire ⁺	6.9 (3.1-12)	6.4 (2.8-9.4)
High mobility	17.3 (8.96-36.8)	Suppress	10.8 (4.9-44.6)	8 (5.8-76.4)
		WUI	11.2 (6.2-38.6)	7.5 (6.6-65)
		Thin	12.9 (6.9-36.8)	8.4 (7.1-53.7)
		Fire	10.9 (5.9-38.4)	7.8 (6.3-65.2)
		Fire ⁺	12.5 (6.4-28.5)	9.4 (5.2-25.1)

percentage of Red Fir and Jeffrey Pine habitat types across the landscape.

Similar to the predicted outcomes for habitat type composition, management scenarios had little influence on the proportion of the landscape comprised of early, mid, and late seral stages over time (Fig. 3). Of the conifer dominated habitat types, 63% of the cells were classified initially as mid-seral, 26% as late seral, and 11% as early seral (the latter of which primarily occurred in White Fir and Red Fir habitat types). Under all management scenarios, the amount of early and mid-seral forest decreased whereas late seral forest increased. In the Fire⁺ scenario, conversion of more areas to Montane Chaparral resulted in a lower proportions of conifer forest, with late seral habitat conditions plateauing at ~22,500 ha and the smallest percentage of early seral conditions.

Compared to habitat type composition and seral stage, forest cover was more strongly influenced by management actions that increased the rate and extent of fuel reduction activities (Fig. 3). Compared to the management scenario simulating fire suppression (Suppress), fuel reduction treatments concentrated in the WUI (WUI) and with only moderate fire use (Fire) had a stabilizing effect on the proportions of the landscape in which forest cover was classified as moderate and closed. Increasing the pace and scale of fuel reduction treatments (Thin) was projected to reduce the amount of closed habitat conditions relative to the Suppress scenario but resulted in the largest proportion of the landscape being comprised of moderate forest cover. With the simulated increase in use of fire in the Fire⁺ scenario, forest cover across the landscape was projected to become sparser, with the most limited area of closed forest conditions.

Changes in suitability of reproductive habitat for wildlife

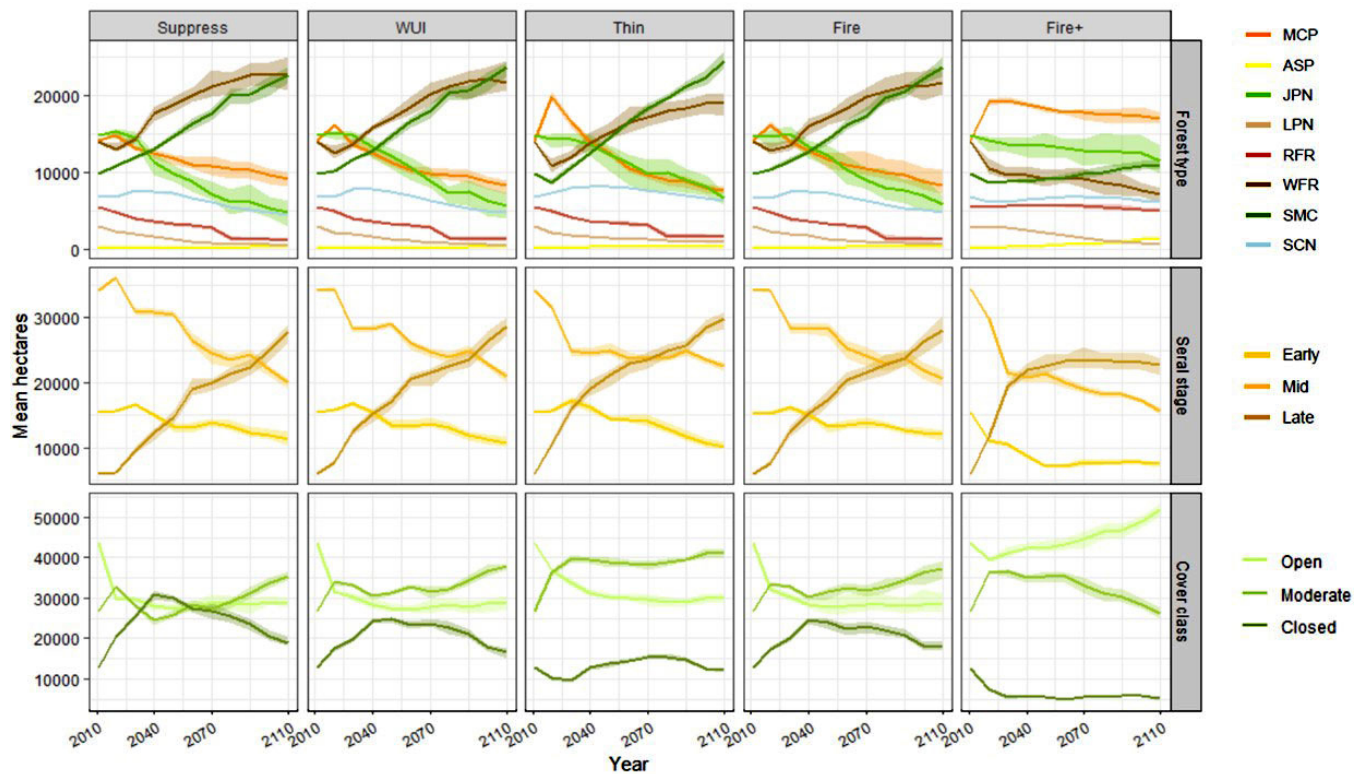
Of the 159 modeled species, 7 species were expected to experience a > 30% increase in the value of their habitat regardless of scenario: American Kestrel (*Falco sparverius*), Great Horned Owl (*Bubo virginianus*), Lewis's Woodpecker (*Melanerpes lewis*), Williamson's Sapsucker (*Sphyrapicus thyroideus*), Purple Martin

(*Progne subis*), Western Bluebird (*Sialia mexicana*), and Evening Grosbeak (*Coccothraustes vespertinus*). Four species were predicted to lose over 30% of their reproductive habitat regardless of scenario including the Mountain Bluebird (*Sialia currucoides*), Western Meadowlark (*Sturnella neglecta*), House Finch (*Carpodacus mexicanus*), and mountain beaver (*Aplodontia rufa*).

For the remaining species the value of the landscape for reproductive purposes, both in the amount and configuration, differed by scenario. In terms of the overall value of reproductive habitat, all scenarios except the Fire⁺ scenario, led to outcomes that resulted in slightly more species experiencing increases in this metric than decreases (Fig. 4A, B). Under the Fire⁺ scenario, more species were predicted to experience declines in the overall amount of habitat for reproduction, although under this scenario the number of species experiencing significant change (i.e., > 30%) was minimized relative to the other scenarios (Fig. 4B).

In terms of high-quality reproductive habitat, all scenarios tended to have relatively neutral impacts on the proportion of their habitat categorized as high value. In the first half of the century, the Fire⁺ scenario resulted in increases in this metric for the largest number of species, whereas the Suppress scenario resulted in the least number of species (Fig. 4C). In the second half of the century, differences in this metric between scenarios were minimal and largely neutral. Similar to the results for the impact of scenarios on the overall habitat quantity of individual species, the Fire⁺ scenario tended to reduce the number of species experiencing more significant fluctuation in this metric during the first half of the century (Fig. 4D). For those species with > 30% fluctuation in the proportion of their high-quality reproductive habitat, more species were predicted to see increases in this metric relative to decreases, for all scenarios (Fig. 4D). Although the overall proportion of high-quality habitat was predicted to increase for as many species as it decreased, the average patch size of individual patches of this habitat was predicted to decrease for most of the species modeled, most notably under the Fire⁺ scenario (Fig. 4E, F). Differences between all other scenarios was

Fig. 3. Changes in forest type, seral stage, and cover class over the period of modeling under each management scenario. Note forest types: White Fir (WFR), Red Fir (RFR), Jeffrey Pine (JPN), Lodgepole Pine (LPN), Sierra Mixed Conifer (SMC), Subalpine Conifer (SCN), Aspen (ASP), and Montane Chaparral (MCP).



minor and the variance in replicates overlapped, but the Thin scenario was predicted to lead to the most favorable biodiversity outcomes in the size of high-quality patches (Fig. 4C).

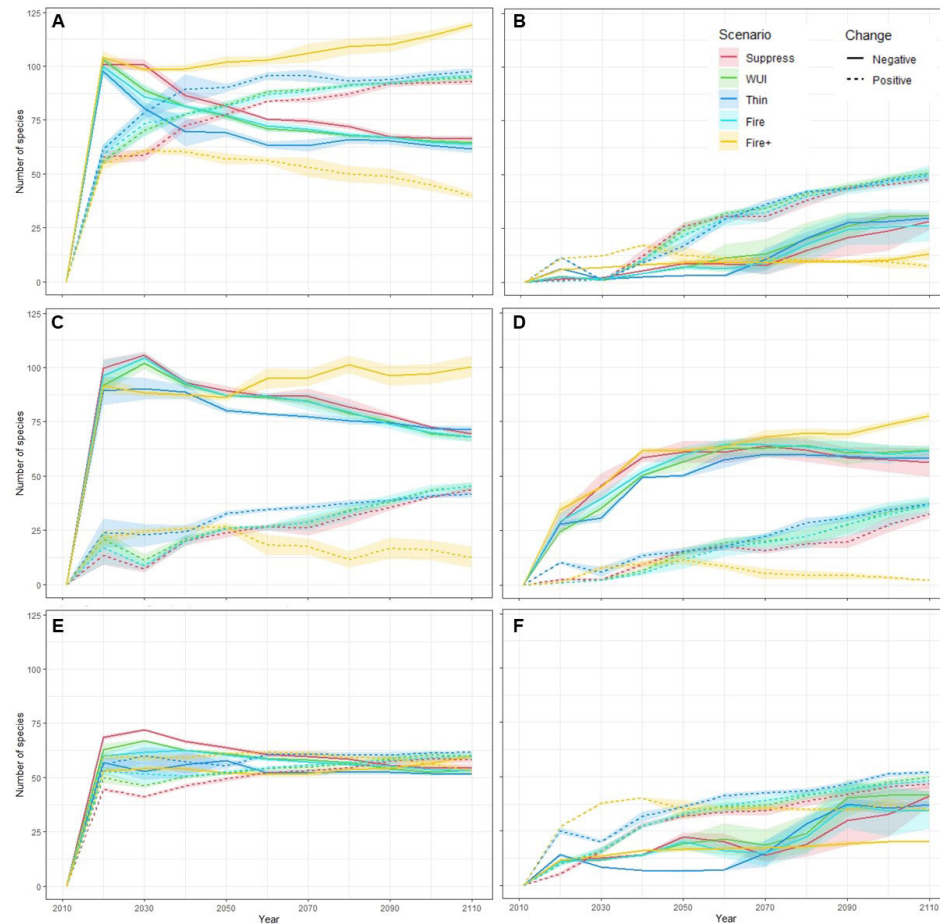
As patches of high-quality reproductive habitat were predicted to decrease over time regardless of scenario, we grouped modeled species based on their size and mobility and compared how each scenario was projected to impact the median size of high-quality reproductive patches for each grouping in 2040 (early century) and in 2070 the mid-century (Table 2, Appendix 3). For species with low mobility that tended to occupy small to moderate sized home ranges (≤ 40 ha; Manley and Schlesinger 2000), the median size of habitat patches decreased over the century regardless of scenario. These decreases were most dramatic under the Fire⁺ scenario both early and mid-century. Relative to the other scenarios, the Thin scenario was predicted to result in larger median patch sizes in the near term. In comparison, for larger-bodied species, the median patch size was predicted to be the smallest relative to other species groups and more similarly across time periods, with the Thin and Fire⁺ scenario leading to higher median patch sizes. For highly mobile species that typically occupy home ranges > 40 ha, patch size decreased over time regardless of scenario with more favorable outcomes again under the Thin and Fire⁺ scenario, both scenarios of which treated larger proportions of the landscape.

DISCUSSION

Recent extreme fire years in forested ecosystems across the globe have raised concerns regarding the potential impact of these climate-driven fires on biodiversity (Barlow et al. 2020, Kelly et al. 2020, Ward et al. 2020) and actions that can be taken to increase resilience of forests to future fires (Wintle et al. 2020, Stephens et al. 2021). In the seasonally dry forests of the western United States, forest restoration strategies that reduce tree densities and fuel loading have been shown to alter fire behavior and reduce fire severity (Stephens and Moghaddas 2005, Tubbesing et al. 2019). Our model results suggest that management regimes that incorporated more extensive use of fuel reduction treatments and fire can alter forest trajectories and habitat for biodiversity.

With the exception of a management regime incorporating more prescribed and managed wildfire (Fire⁺), our models indicated that forests surrounding Lake Tahoe will experience an increase in White Fir and Sierra Mixed Conifer and a decrease in stands dominated by Jeffrey pine and montane chaparral. This prediction is supported by previous research indicating that Jeffrey pine is adapted to a frequent-fire disturbance regime whereby frequent removal of small diameter trees and fire-intolerant tree species by low-intensity fire is needed to maintain this species on the landscape (Fulé et al. 2012). Our modeling results suggest that higher use of fire is needed to maintain Jeffrey Pine habitat in our study area. Previous research has also shown that red fir seedling establishment and growth are strongly related

Fig. 4. The number of species projected to experience positive and negative changes in (A) the value of reproductive habitat, (B) the number of species in which change in A is > 30%, (C) proportion of the landscape considered high quality, (D), the number of species in which change in C is > 30%, (E) the average size of patches of high-quality habitat, and (F) the number of species in which change in E is > 30% under each management scenario.



to periodic disturbance (Chappell and Agee 1996) and this may explain why the Fire⁺ scenario was also the only scenario predicted to maintain Red Fir habitat. Climate change is predicted to have a negative impact on red fir (Meng et al. 2015) as establishment of red fir seedlings is influenced by soil moisture and post-fire temperatures and maintaining Red Fir habitat will be a future management challenge. Although maintaining a more even mix of habitat types, increased use of fire under the Fire⁺ scenario led to the highest proportion of montane chaparral. Many non-conifer vegetation communities have been negatively impacted by fire suppression including montane chaparral and riparian communities, including aspen (White and Long 2019), and our modeling results indicate that higher use of fire may be one way to ensure persistence of these habitat types in the Lake Tahoe Basin.

Whereas frequent use of fire appeared to be important in maintaining a diversity of vegetative communities, the area being treated led to larger differences among scenarios in the density of forest canopy. Increasing levels of disturbance resulted in

landscapes with a higher percentage of the landscape comprised of moderate tree densities and associated canopy (Thin scenario) or low tree densities and open canopy (Fire⁺ scenario). Over the century, present day areas with open canopy were predicted to transition to areas with moderate canopy cover consistent with forest growth and succession with limited disturbance and competition for resources. This trajectory is similar to that observed over that last century with increasing stand densities of smaller-sized trees and less structural variability (Taylor 2004). Landscape restoration strategies incorporating more extensive fuel reduction resulted in forests with more moderate tree densities that are less prone to stand-replacing fires (Agee and Skinner 2005) and more resilient to a changing climate (North et al. 2019), but that also may be less conducive to species preferring closed canopy conditions.

One of our main objectives was to understand the impact of different landscape management scenarios on wildlife in the Lake Tahoe Basin based on wildlife-habitat associations. Our modeling results suggest that over the next century changes in wildlife

habitat will occur in quantifiable ways, but predictions for most species indicate that they will experience habitat fluctuations that are less than 30% of their current habitat regardless of management approach, suggesting that over the long-term the trajectory of forest change will have an overall neutral effect on biodiversity. Although the Fire⁺ scenario was predicted to promote greater representation of different habitat types, it was predicted to result in a reduction in the overall value of the habitat for many species despite maintaining more consistent amounts of high-quality reproductive habitat than the other scenarios (Fig. 4). This suggests that in our study area, structural variables of forest habitat were more influential in maintaining habitat for biodiversity than forest composition.

The reduced frequency and extent of fire have caused an increase in forest density, a shift to more fire-sensitive species and large-scale homogenization of forest conditions (Taylor 2001, Safford and Stevens 2017). Over the next century, our simulations project that disturbances altering forest structure will break up the landscape regardless of scenario, reducing the current size of large contiguous forest patches. The distribution of habitat across landscape will impact species differently depending on the distribution of needed resources and their ability to transverse landscapes (Kindlmann and Burel 2008). Based on the size of high-valued reproductive habitat patches, we attempted to predict how different management approaches were likely to impact different groupings of wildlife based on their typical home-range size and mobility (Manley and Schlesinger 2000). We acknowledge that the habitat matrix (i.e., inter-patch habitat) can be as important as the size of habitat patches (Tischendorf and Fahrig 2000), but this is beyond the scope of this study. In general, we found that patches of contiguous high-quality reproductive habitat would decrease for the majority of species, most notably under the Fire⁺ scenario (Fig. 4D, E). Management scenarios that were simulated at greater extents (i.e., the Thin and Fire⁺ scenarios) tended to result in larger patch sizes for the more mobile species and those with larger home ranges. This result in combination with the metric of overall habitat quality for wildlife indicates that management strategies are likely to have idiosyncratic effects of wildlife species, but biodiversity in this disturbance-prone system is fairly resilient to change (see also Sollmann et al. 2015, Kelt et al. 2017). However, in our model we assume that habitat suitability is strongly linked to forest composition and structure, and we do not account for other abiotic and biotic factors that determine the capacity of a given habitat to support the needs of a species.

CONCLUSIONS

Change is predicted, and management can have an impact on the trajectory and speed of change for habitat types and associated species. Choice of best management approach is a societal decision (Abelson et al. 2022) and will depend on prioritization of resources and outcomes, of which biodiversity is just one. Our results are consistent with other studies (Fontaine and Kennedy 2012, Stephens et al. 2014, Stevens et al. 2016) suggesting that different management actions will alter habitat suitability for biodiversity, but that the effects will be species-dependent with compensatory effects on species richness. However, our simulations are based on a representative concentration pathway (rcp 4.5) that does not match our current global emissions (Schwalm et al. 2020). It is likely that use of a higher greenhouse

gas emissions scenario would increase the amount of fire on the landscape, leading to more frequent disturbance and landscape conditions most similar to the Fire⁺ scenario. We have only considered one metric for measuring biodiversity without considering the functional role of individual species (Caro 2010, Slauson et al. 2022), which may be as important in sustaining ecosystem function as maintaining current species richness. Ensuring the sustainability of forested ecosystems into the future will continue to be a challenge for managers and society, and integrated frameworks are needed to assess the value of future landscapes (Stevens et al. 2016, Abelson et al. 2022).

Responses to this article can be read online at:

<https://www.ecologyandsociety.org/issues/responses.php/13301>

Author Contributions:

RS, AK, and CM formulated the LANDIS model and computed the outputs necessary to implement the wildlife habitat model. AW, TH, AK, and AB developed the framework for habitat compilations. AW analyzed the data and wrote the manuscript with input from all authors. AW conceived the study and was in charge of overall direction and planning.

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Data Availability:

Parameters used in the LANDIS modeling are available here: <https://github.com/LANDIS-II-Foundation/Project-Lake-Tahoe-2017>, which has been archived here: <https://doi.org/10.5281/zenodo.4644579>

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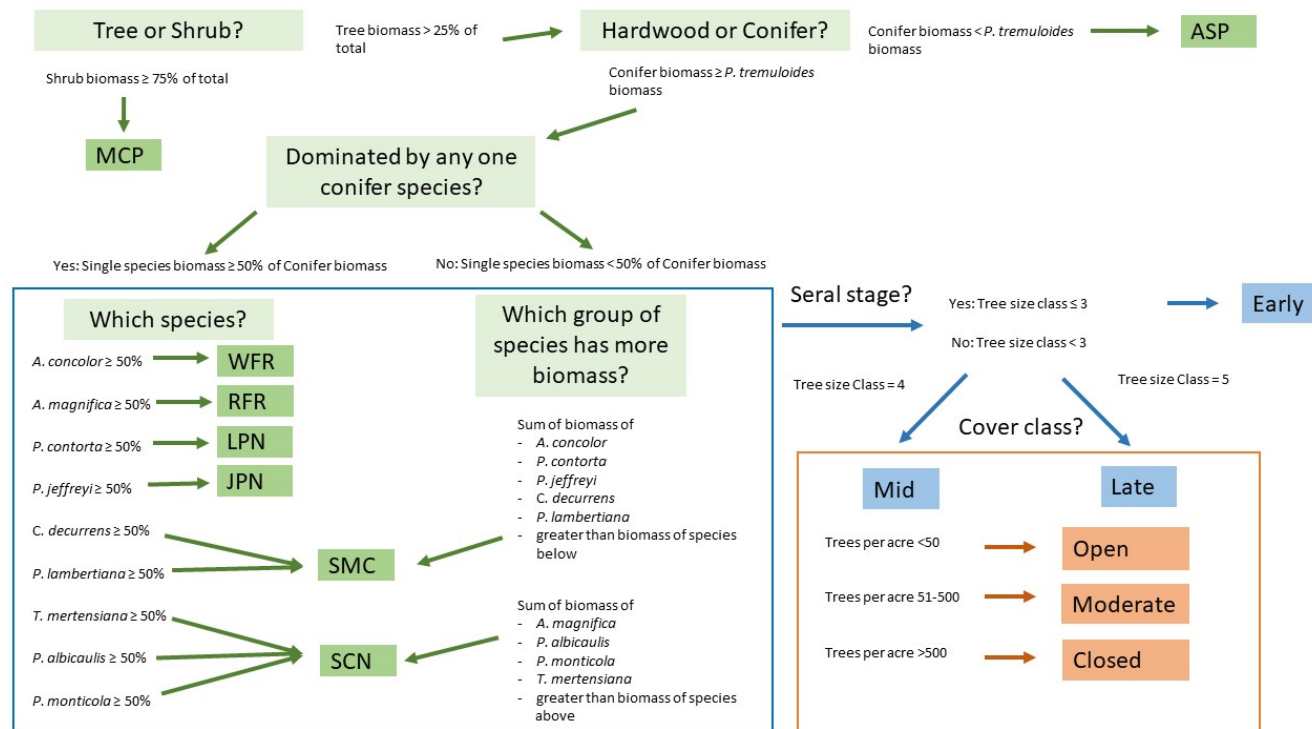
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APPENDIX 1. Schematic of the crosswalk to California Wildlife Habitats using species biomass outputs produced by LANDIS-II. The biomass composition of each pixel in the LANDIS-II output is evaluated to approximated predefined habitat types including montane chaparral (MCP), aspen (ASP), white fir (WFR), red fir (RFR), lodgepole pine (LPN), Jeffrey pine (JPN), Sierra mixed conifer (SMC), and subalpine conifer (SCN). After habitat type is determined, conifer-dominated pixels are classified as early, mid or late seral conditions according to average tree size estimates. Finally, canopy cover class is estimated in mid to late seral forests using estimates of tree density.



APPENDIX 1. Median and Range of Hectares of Each Modeled Habitat on 2010, 2040 and 2070 Under Each Management Scenario

Habitat Code	Vegetation type	2010					2040					2070				
		initial	Suppression	WUI	Thin	Fire	Fire+	Suppression	WUI	Thin	Fire	Fire+	Suppression	WUI	Thin	Fire
MCP	Montane Chaparral	14217	12549.8 (11836-13064)	12603 (12031-13534)	14120.3 (13533-14518)	12849 (12306-13370)	18888.1 (18501-19259)	10963.2 (9722-12230)	9930.6 (9323-10655)	9776.6 (9194-10187)	10581.6 (9710-12753)	17812.7 (16904-18456)				
ASP	Aspen	234	275.1 (248-329)	281.4 (257-298)	308.4 (287-323)	289.5 (271-342)	434.3 (410-445)	283.9 (251-386)	277.5 (231-323)	374.4 (323-430)	326.8 (284-420)	794.9 (725-858)				
JPN1	Jeffrey Pine (early seral)	0	259.1 (234-330)	339.4 (261-406)	354.4 (296-421)	309.3 (256-354)	116 (97-134)	70.2 (47-86)	107.1 (86-137)	105 (89-118)	111.6 (86-125)	154 (133-172)				
JPN2S	Jeffrey Pine (mid seral, open)	1366	212.7 (161-272)	395.7 (329-511)	824.5 (705-1021)	357.6 (259-487)	1620.7 (1536-1748)	180.2 (129-242)	254.4 (205-318)	709 (577-865)	242.7 (192-332)	1750.9 (1533-1914)				
JPN2M	Jeffrey Pine (mid seral, moderate)	8952	2373.3 (2119-2653)	3299.3 (3145-3524)	3815.1 (3543-4027)	3241.5 (3040-3454)	3478.9 (3274-3571)	1194.2 (1091-1500)	2012.5 (1777-2384)	2538 (2292-2856)	2048.3 (1676-2264)	4185.9 (3972-4491)				
JPN2D	Jeffrey Pine (mid seral, closed)	3883	4723.3 (4427-5161)	3467.7 (3152-3665)	1654.4 (1439-1872)	3781.6 (3439-4108)	445.7 (400-516)	2811.8 (2425-3287)	2389.9 (2131-2721)	1263.9 (1144-1421)	2558.2 (2340-2728)	964.8 (869-1061)				
JPN3S	Jeffrey Pine (late seral, open)	484	146.3 (102-174)	131.8 (92-177)	446.2 (258-613)	86.4 (34-161)	269.4 (189-335)	204 (83-367)	131.7 (92-209)	299.7 (175-442)	121.8 (57-182)	240.8 (132-321)				
JPN3M	Jeffrey Pine (late seral, moderate)	206	3697.7 (2575-4277)	5775.4 (4575-6368)	6601.8 (5265-7824)	5606.5 (4448-6310)	7593.9 (6187-8665)	2676.2 (2098-3722)	4044.7 (3139-4930)	5029.9 (2891-6234)	3892.7 (3237-5449)	5532.2 (3969-6996)				
JPN3D	Jeffrey Pine (late seral, closed)	0	0.9 (0-3)	0.5 (0-3)	0.3 (0-2)	0.5 (0-2)	0 (0-0)	4.3 (1-9)	3.8 (1-8)	2.5 (1-5)	3.2 (1-8)	0 (0-0)				
LPN1	Lodgepole Pine (early seral)	0	79.4 (59-100)	80.3 (67-94)	78 (59-92)	69.4 (65-76)	39.3 (29-48)	7.6 (2-12)	7.2 (3-12)	5.4 (4-10)	7.4 (5-10)	18.2 (13-22)				
LPN2S	Lodgepole Pine (mid seral, open)	386	6.7 (0-21)	17.2 (8-26)	65.5 (38-95)	14.4 (2-29)	140.4 (127-158)	47.1 (14-96)	55.4 (36-136)	103.6 (89-118)	56.5 (20-106)	91.4 (70-110)				
LPN2M	Lodgepole Pine (mid seral, moderate)	1807.8	496.2 (406-581)	621.6 (562-739)	967.6 (904-1050)	639.2 (570-719)	1611.2 (1563-1682)	364.4 (312-398)	417.2 (378-477)	704.3 (622-768)	498 (433-553)	992.4 (947-1061)				
LPN2D	Lodgepole Pine (mid seral, closed)	817.2	1149.3 (1110-1192)	1014.8 (971-1056)	539.1 (511-558)	1024.9 (969-1074)	616.4 (581-664)	371.3 (349-420)	341.2 (291-363)	259.9 (232-294)	355.9 (327-400)	207.1 (177-232)				
LPN3S	Lodgepole Pine (late seral, open)	1	6.5 (3-12)	6.7 (4-10)	9.2 (5-13)	10 (7-12)	40.7 (30-54)	59.1 (13-139)	36.7 (15-67)	88 (51-158)	47.3 (12-114)	92.5 (69-111)				
LPN3M	Lodgepole Pine (late seral, moderate)	0	58.1 (49-69)	64.8 (51-77)	89.8 (72-107)	61.1 (53-69)	75.1 (56-98)	88.9 (59-122)	98.2 (81-116)	147 (116-168)	119.6 (80-153)	149.5 (134-170)				
RFR1	Red Fir (early seral)	11	39.3 (33-47)	43 (38-51)	26.1 (20-39)	38.9 (28-49)	354.7 (321-382)	79.7 (62-93)	78.6 (44-91)	44.1 (35-53)	76.8 (62-93)	509.1 (457-537)				
RFR2S	Red Fir (mid seral, open)	0	10.6 (0-20)	12.3 (4-26)	11.6 (5-19)	10.7 (2-22)	20.4 (5-33)	0.9 (0-3)	3.3 (0-7)	13.2 (9-18)	3 (0-8)	12.3 (4-24)				
RFR2M	Red Fir (mid seral, moderate)	716	275.9 (259-291)	273.4 (249-303)	275.5 (225-351)	246.7 (236-278)	330.3 (276-383)	111.6 (101-124)	127.5 (108-140)	235.8 (204-264)	118 (104-146)	232.6 (198-261)				
RFR2D	Red Fir (mid seral, closed)	1393	118.6 (109-128)	130.1 (100-146)	77 (52-119)	124 (102-145)	8 (3-12)	69.7 (61-81)	64.7 (57-75)	206.9 (54-106)	69.3 (51-88)	66.2 (57-76)				
RFR3S	Red Fir (late seral, open)	802	277.3 (238-308)	277.3 (254-301)	510.9 (458-581)	312.4 (282-335)	497.8 (466-532)	991.1 (726-1211)	957.2 (774-1100)	1171.8 (918-1628)	1064.4 (733-1264)	1478.9 (1312-1650)				
RFR3M	Red Fir (late seral, moderate)	94	2959.3 (2707-3101)	2938.1 (2774-3031)	2876 (2672-3017)	3024.9 (2860-3152)	4548.3 (4387-4685)	1657.7 (1397-1797)	1727.9 (1345-1973)	1615.2 (1254-1816)	3349.8 (1341-1841)	3349.8 (3040-3778)				
RFR3D	Red Fir (late seral, closed)	2478	0 (0-0)	0 (0-0)	0 (0-0)	0 (0-0)	0 (0-0)	0 (0-0)	0 (0-0)	0 (0-0)	0 (0-0)	0 (0-0)				
WFR1	White Fir (early seral)	8881.2	5537.3 (5376-5737)	5479.8 (5326-5585)	4983.1 (4643-5234)	5335 (4987-5629)	2763.8 (2514-2974)	4704 (4480-5028)	4498.2 (4181-4868)	4110.6 (3866-4395)	4433.3 (4200-4763)	1816.2 (1620-2087)				
WFR2S	White Fir (mid seral, open)	3	25.3 (17-32)	147.8 (96-217)	552.7 (398-640)	112.5 (72-158)	900.8 (719-1129)	5.2 (0-10)	84.6 (49-113)	350.5 (270-467)	79.8 (49-117)	217.1 (154-283)				
WFR2M	White Fir (mid seral, moderate)	2304.8	2969.2 (2653-3244)	3186.3 (2896-3497)	3168.3 (2752-3787)	3186 (2776-3805)	2207.4 (1947-2601)	1599.4 (1471-1748)	1906.7 (1839-1985)	2064.9 (1739-3523)	1990.3 (1799-2231)	1740.6 (1543-2030)				
WFR2D	White Fir (mid seral, closed)	2205	6687.6 (6016-7069)	4047.9 (3870-4274)	1386.9 (1204-1633)	441.7 (3837-4737)	257.8 (210-356)	7023.1 (6561-7908)	5942.7 (5389-6357)	2860.8 (2484-3139)	5233.9 (4690-5707)	360.5 (289-445)				
WFR3S	White Fir (late seral, open)	219	98.7 (69-137)	124.5 (82-153)	223.7 (163-275)	106.4 (79-139)	441.7 (363-571)	219.5 (167-291)	415.6 (302-609)	691.8 (428-977)	345.7 (231-558)	1551.4 (1167-1958)				
WFR3M	White Fir (late seral, moderate)	288	2473 (2233-2609)	2831.5 (2616-2988)	3630.2 (3380-3898)	3001.4 (2690-3212)	3121.7 (2768-3568)	7590.1 (6835-8560)	7300.6 (6185-8279)	6288.9 (5777-6656)	7727.6 (6719-8737)	3499.5 (2947-4496)				
WFR3D	White Fir (late seral, closed)	228	0 (0-0)	0 (0-0)	0 (0-0)	0 (0-0)	0.1 (0-1)	0 (0-0)	0 (0-0)	0 (0-0)	0 (0-0)	0 (0-0)				
SMC1	Sierran Mixed Conifer (early seral)	5242	6380 (6073-6626)	6336.7 (5902-6511)	6056.3 (5791-6386)	6185.9 (5932-6426)	2486 (2323-2680)	6775.9 (6440-7050)	6707.6 (6365-7283)	6893.8 (6527-7348)	6683.1 (6327-7145)	2097 (1975-2197)				
SMC2S	Sierran Mixed Conifer (mid seral, open)	110.2	35.8 (17-62)	155.1 (114-180)	282.2 (236-341)	121.5 (86-153)	596.1 (561-648)	74.8 (33-107)	214.5 (165-288)	490.7 (419-531)	181.8 (152-260)	386.6 (327-448)				
SMC2M	Sierran Mixed Conifer (mid seral, moderate)	2312.8	1952.2 (1877-2026)	2046.4 (1989-2223)	2356.6 (2195-2496)	1947.7 (1798-2096)	1848 (1747-2004)	1775.3 (1600-1874)	2425.2 (2156-2587)	3713.2 (3494-3935)	2402.4 (2327-2475)	2421.8 (2171-2662)				
SMC2D	Sierran Mixed Conifer (mid seral, closed)	1459	2884.3 (2751-3022)	2178.2 (2048-2314)	1081.1 (974-1210)	2299.9 (2171-2453)	380.3 (337-425)	6466 (5721-6862)	5790.3 (5585-6186)	3834 (3470-4190)	5340.2 (4793-5805)	560.4 (483-678)				
SMC3S	Sierran Mixed Conifer (late seral, open)	715.8	172.8 (99-247)	175.8 (125-219)	232.3 (203-299)	149.3 (124-178)	874.4 (778-1025)	393.1 (218-494)	421.9 (303-565)	592.8 (499-671)	413.9 (295-509)	1364.4 (1154-1584)				
SMC3M	Sierran Mixed Conifer (late seral, moderate)	114.2	1641.7 (1554-1751)	2023.1 (1951-2125)	2540.4 (2393-2670)	1938.8 (1836-2051)	2768.7 (2686-2851)	2191.3 (2089-2333)	2447.5 (2312-2582)	2877.4 (2768-2996)	2443 (2306-2560)	3027.5 (2916-3125)				
SCN1	Subalpine Conifer (early seral)	0	0.6 (0-3)	0.5 (0-2)	0.2 (0-2)	0.4 (0-1)	52.8 (42-63)	0 (0-0)	0.5 (0-1)	0 (0-0)	0.1 (0-1)	52 (42-65)				
SCN2S	Subalpine Conifer (mid seral, open)	422	3.4 (1-9)	3.2 (1-6)	4.7 (1-8)	2.9 (0-5)	164.6 (138-193)	1.3 (0-4)	1.1 (0-2)	0.9 (0-4)	1 (0-3)	78.3 (64-91)				
SCN2M	Subalpine Conifer (mid seral, moderate)	6136	463.5 (429-504)	650.3 (594-705)	1947 (1762-2144)	900 (842-965)	2181 (2074-2267)	550.5 (495-617)	591.3 (552-635)	417.8 (330-457)	714.6 (610-779)	1891.3 (1822-1999)				
SCN2D	Subalpine Conifer (mid seral, closed)	35	6321.5 (5963-6535)	6337.1 (6197-6426)	4398.9 (4138-4648)	5692.6 (5410-5850)	2956.6 (2817-3155)	1776.2 (1722-1843)	1863.8 (1800-1946)	2579.2 (2472-2650)	2006.3 (1921-2123)	2098.9 (2013-2160)				
SCN3S	Subalpine Conifer (late seral, open)	65	20.1 (12-40)	15.5 (11-20)	14.2 (8-20)	17.8 (11-31)	583.1 (562-605)	4.7 (0-7)	1.5 (0-7)	0.9 (0-2)	2.1 (0-7)	541.1 (522-570)				
SCN3M	Subalpine Conifer (late seral, moderate)	213	805.9 (738-956)	816.1 (782-866)	1907.7 (1691-2137)	926.3 (883-1020)	629.5 (583-692)	3861.1 (3639-3961)	3959.9 (3893-4033)	4714.3 (4584-4831)	3690.3 (3581-3849)	2231.7 (2202-2296)				
SCN3D	Subalpine Conifer (late seral, closed)	0	0 (0-0)	0.1 (0-1)	0.3 (0-2)	0 (0-0)	0 (0-0)	0.7 (0-2)	1.5 (0-4)	22.4 (14-31)	1 (0-3)	0 (0-0)				
Null values		200334	200951.1 (200691-201330)	200806.3 (200621-201283)	200717.5 (200619-200898)	200808 (200641-201075)	202792 (202554-202352)	201886.7 (201192-202848)	201493.7 (201104-201922)	201147 (200893-201335)	201526.9 (201087-202431)	204563.5 (203890-205718)				

APPENDIX 3. Habitat Suitability Values for Each Modeled Species for Each Habitat Type

Code	Common Name	Scientific name	Range Size	Mobility
B108	Turkey Vulture	<i>Cathartes aura</i>	>40ha	high
B115	Sharp-Shinned Hawk	<i>Accipiter striatus</i>	<40ha	high
B116	Cooper's Hawk*	<i>Accipiter cooperii</i>	<40ha	high
B117	Northern Goshawk	<i>Accipiter gentilis</i>	>40ha	high
B123	Red-Tailed Hawk	<i>Buteo jamaicensis</i>	>40ha	high
B126	Golden Eagle	<i>Aquila chrysaetos</i>	>40ha	high
B127	American Kestrel	<i>Falco sparverius</i>	>40ha	high
B131	Prairie Falcon	<i>Falco mexicanus</i>	>40ha	high
B134	Blue Grouse	<i>Dendragapus obscurus</i>	<40ha	high
B138	Wild Turkey*	<i>Meleagris gallopavo</i>	>40ha	high
B140	California Quail	<i>Callipepla californica</i>	<40ha	high
B141	Mountain Quail	<i>Oreortyx pictus</i>	>40ha	high
B251	Band-Tailed Pigeon	<i>Columba fasciata</i>	<40ha	high
B255	Mourning Dove*	<i>Zenaida macroura</i>	>40ha	high
B263	Flammulated Owl	<i>Otus flammeolus</i>	<40ha	high
B264	Western Screech-Owl*	<i>Otus kennicottii</i>	<40ha	high
B265	Great Horned Owl	<i>Bubo virginianus</i>	>40ha	high
B267	Northern Pygmy-Owl	<i>Glaucidium gnoma</i>	<40ha	high
B270	Spotted Owl	<i>Strix occidentalis</i>	>40ha	high
B272	Long-Eared Owl*	<i>Asio otus</i>	<40ha	high
B274	Northern Saw-Whet Owl	<i>Aegolius acadicus</i>	<40ha	high
B276	Common Nighthawk	<i>Chordeiles minor</i>	>40ha	high
B277	Common Poorwill*	<i>Phalaenoptilus nuttallii</i>	<40ha	high
B279	Black Swift	<i>Cypseloides niger</i>	>40ha	high
B281	Vaux's Swift*	<i>Chaetura vauxi</i>	>40ha	high
B282	White-Throated Swift	<i>Aeronautes saxatalis</i>	>40ha	high
B287	Anna's Hummingbird*	<i>Calypte anna</i>	<40ha	high
B289	Calliope Hummingbird	<i>Stellula calliope</i>	<40ha	high
B290	Broad-Tailed Hummingbird	<i>Selasphorus platycercus</i>	<40ha	high
B294	Lewis's Woodpecker	<i>Melanerpes lewis</i>	<40ha	high
B299	Red-Breasted Sapsucker	<i>Sphyrapicus ruber</i>	<40ha	high
B300	Williamson's Sapsucker	<i>Sphyrapicus thyroideus</i>	<40ha	high
B304	Hairy Woodpecker	<i>Picoides villosus</i>	<40ha	high
B305	White-Headed Woodpecker	<i>Picoides albolarvatus</i>	<40ha	high
B306	Black-Backed Woodpecker	<i>Picoides arcticus</i>	<40ha	high
B307	Northern Flicker	<i>Colaptes auratus</i>	<40ha	high
B308	Pileated Woodpecker	<i>Dryocopus pileatus</i>	>40ha	high
B309	Olive-Sided Flycatcher	<i>Contopus cooperi</i>	<40ha	high
B311	Western Wood-Pewee	<i>Contopus sordidulus</i>	<40ha	high
B317	Hammond's Flycatcher	<i>Empidonax hammondii</i>	<40ha	high
B318	Dusky Flycatcher	<i>Empidonax oberholseri</i>	<40ha	high
B320	Pacific-Slope Flycatcher	<i>Empidonax difficilis</i>	<40ha	high
B326	Ash-Throated Flycatcher*	<i>Myiarchus cinerascens</i>	<40ha	high
B338	Purple Martin	<i>Progne subis</i>	<40ha	high

B340	Violet-Green Swallow	<i>Tachycineta thalassina</i>	<40ha	high
B341	Northern Rough-Winged Swallow	<i>Stelgidopteryx serripennis</i>	<40ha	high
B343	Cliff Swallow*	<i>Petrochelidon pyrrhonota</i>	>40ha	high
B344	Barn Swallow	<i>Hirundo rustica</i>	<40ha	high
B346	Steller's Jay	<i>Cyanocitta stelleri</i>	<40ha	high
B348	Western Scrub Jay*	<i>Aphelocoma californica</i>	<40ha	high
B349	Pinyon Jay*	<i>Gymnorhinus cyanocephalus</i>	>40ha	high
B350	Clark's Nutcracker	<i>Nucifraga columbiana</i>	>40ha	high
B354	Common Raven	<i>Corvus corax</i>	>40ha	high
B356	Mountain Chickadee	<i>Poecile gambeli</i>	<40ha	high
B360	Bushtit	<i>Psaltiriparus minimus</i>	<40ha	high
B361	Red-Breasted Nuthatch	<i>Sitta canadensis</i>	<40ha	high
B362	White-Breasted Nuthatch	<i>Sitta carolinensis</i>	<40ha	high
B363	Pygmy Nuthatch	<i>Sitta pygmaea</i>	<40ha	high
B364	Brown Creeper	<i>Certhia americana</i>	<40ha	high
B366	Rock Wren	<i>Salpinctes obsoletus</i>	<40ha	high
B368	Bewick'S Wren	<i>Thryomanes bewickii</i>	<40ha	high
B369	House Wren	<i>Troglodytes aedon</i>	<40ha	high
B370	Winter Wren	<i>Troglodytes troglodytes</i>	<40ha	high
B375	Golden-Crowned Kinglet	<i>Regulus satrapa</i>	<40ha	high
B376	Ruby-Crowned Kinglet	<i>Regulus calendula</i>	<40ha	high
B377	Blue-Gray Gnatcatcher*	<i>Polioptila caerulea</i>	<40ha	high
B380	Western Bluebird*	<i>Sialia mexicana</i>	<40ha	high
B381	Mountain Bluebird	<i>Sialia currucoides</i>	<40ha	high
B382	Townsend's Solitaire	<i>Myadestes townsendi</i>	<40ha	high
B385	Swainson's Thrush*	<i>Catharus ustulatus</i>	<40ha	high
B386	Hermit Thrush	<i>Catharus guttatus</i>	<40ha	high
B389	American Robin	<i>Turdus migratorius</i>	<40ha	high
B411	European Starling*	<i>Sturnus vulgaris</i>	>40ha	high
B415	Plumbeous Vireo	<i>Vireo plumbeous</i>	<40ha	high
B418	Warbling Vireo	<i>Vireo gilvus</i>	<40ha	high
B426	Nashville Warbler	<i>Vermivora ruficapilla</i>	<40ha	high
B430	Yellow Warbler	<i>Dendroica petechia</i>	<40ha	high
B435	Yellow-Rumped Warbler	<i>Dendroica coronata</i>	<40ha	high
B436	Black-Throated Gray Warbler*	<i>Dendroica nigrescens</i>	<40ha	high
B438	Hermit Warbler	<i>Dendroica occidentalis</i>	<40ha	high
B460	Macgillivray's Warbler*	<i>Oporornis tolmiei</i>	<40ha	high
B471	Western Tanager	<i>Piranga ludoviciana</i>	<40ha	high
B475	Black-Headed Grosbeak*	<i>Pheucticus melanocephalus</i>	<40ha	high
B477	Lazuli Bunting*	<i>Passerina amoena</i>	<40ha	high
B482	Green-Tailed Towhee	<i>Pipilo chlorurus</i>	<40ha	high
B483	Spotted Towhee	<i>Pipilo maculatus</i>	<40ha	high
B484	California Towhee*	<i>Pipilo crissalis</i>	<40ha	high
B489	Chipping Sparrow	<i>Spizella passerina</i>	<40ha	high
B491	Brewer'S Sparrow	<i>Spizella breweri</i>	<40ha	high
B494	Vesper Sparrow*	<i>Pooecetes gramineus</i>	<40ha	high
B504	Fox Sparrow	<i>Passerella iliaca</i>	<40ha	high

B505	Song Sparrow*	<i>Melospiza melodia</i>	<40ha	high
B506	Lincoln's Sparrow*	<i>Melospiza lincolnii</i>	<40ha	high
B512	Dark-Eyed Junco	<i>Junco hyemalis</i>	<40ha	high
B521	Western Meadowlark*	<i>Sturnella neglecta</i>	<40ha	high
B524	Brewer's Blackbird*	<i>Euphagus cyanocephalus</i>	>40ha	high
B528	Brown-Headed Cowbird*	<i>Molothrus ater</i>	<40ha	high
B535	Pine Grosbeak	<i>Pinicola enucleator</i>	<40ha	high
B536	Purple Finch	<i>Carpodacus purpureus</i>	<40ha	high
B537	Cassin's Finch	<i>Carpodacus cassinii</i>	<40ha	high
B538	House Finch*	<i>Carpodacus mexicanus</i>	<40ha	high
B539	Red Crossbill	<i>Loxia curvirostra</i>	>40ha	high
B542	Pine Siskin	<i>Carduelis pinus</i>	<40ha	high
B543	Lesser Goldfinch*	<i>Carduelis psaltria</i>	<40ha	high
B546	Evening Grosbeak	<i>Coccothraustes vespertinus</i>	<40ha	high
M003	Vagrant Shrew	<i>Sorex vagrans</i>	<40ha	low
M004	Dusky Shrew*	<i>Sorex monticolus</i>	<40ha	low
M012	Trowbridge's Shrew	<i>Sorex trowbridgii</i>	<40ha	low
M018	Broad-Footed Mole	<i>Scapanus latimanus</i>	<40ha	low
M021	Little Brown Myotis*	<i>Myotis lucifugus</i>	>40ha	high
M023	Yuma Myotis*	<i>Myotis yumanensis</i>	>40ha	high
M025	Long-Eared Myotis	<i>Myotis evotis</i>	>40ha	high
M026	Fringed Myotis*	<i>Myotis thysanodes</i>	>40ha	high
M028	California Myotis*	<i>Myotis californicus</i>	>40ha	high
M030	Silver-Haired Bat	<i>Lasionycteris noctivagans</i>	<40ha	high
M032	Big Brown Bat	<i>Eptesicus fuscus</i>	>40ha	high
M038	Pallid Bat*	<i>Antrozous pallidus</i>	>40ha	high
M039	Brazilian Free-Tailed Bat*	<i>Tadarida brasiliensis</i>	>40ha	high
M049	Snowshoe Hare*	<i>Lepus americanus</i>	<40ha	low
M051	Black-Tailed Hare*	<i>Lepus californicus</i>	<40ha	low
M052	Mountain Beaver	<i>Aplodontia rufa</i>	<40ha	low
M055	Yellow-Pine Chipmunk	<i>Tamias amoenus</i>	<40ha	low
M057	Allen's Chipmunk	<i>Tamias senex</i>	<40ha	low
M062	Long-Eared Chipmunk	<i>Tamias quadrimaculatus</i>	<40ha	low
M063	Lodgepole Chipmunk	<i>Tamias speciosus</i>	<40ha	low
M070	Belding's Ground Squirrel*	<i>Spermophilus beldingi</i>	<40ha	low
M072	California Ground Squirrel	<i>Spermophilus beecheyi</i>	<40ha	low
M075	Golden-Mantled Ground Squirrel	<i>Spermophilus lateralis</i>	<40ha	low
M077	Western Gray Squirrel*	<i>Sciurus griseus</i>	<40ha	low
M079	Douglas' Squirrel	<i>Tamiasciurus douglasii</i>	<40ha	low
M080	Northern Flying Squirrel	<i>Glaucomys sabrinus</i>	<40ha	low
M085	Mountain Pocket Gopher*	<i>Thomomys monticola</i>	<40ha	low
M117	Deer Mouse	<i>Peromyscus maniculatus</i>	<40ha	low
M119	Brush Mouse	<i>Peromyscus boylii</i>	<40ha	low
M120	Pinyon Mouse	<i>Peromyscus truei</i>	<40ha	low
M126	Desert Woodrat*	<i>Neotoma lepida</i>	<40ha	low
M127	Dusky-Footed Woodrat	<i>Neotoma fuscipes</i>	<40ha	low
M128	Bushy-Tailed Woodrat	<i>Neotoma cinerea</i>	<40ha	low

M133	Montane Vole*	<i>Microtus montanus</i>	<40ha	low
M136	Long-Tailed Vole	<i>Microtus longicaudus</i>	<40ha	low
M143	Western Jumping Mouse	<i>Zapus princeps</i>	<40ha	low
M146	Coyote	<i>Canis latrans</i>	>40ha	high
M151	Black Bear	<i>Ursus americanus</i>	>40ha	high
M153	Raccoon	<i>Procyon lotor</i>	>40ha	high
M154	Marten	<i>Martes americana</i>	>40ha	high
M155	Fisher	<i>Martes pennanti</i>	>40ha	high
M156	Ermine	<i>Mustela erminea</i>	<40ha	high
M157	Long-Tailed Weasel	<i>Mustela frenata</i>	<40ha	high
M160	Badger	<i>Taxidea taxus</i>	>40ha	high
M161	Western Spotted Skunk	<i>Spilogale gracilis</i>	>40ha	high
M162	Striped Skunk	<i>Mephitis mephitis</i>	>40ha	high
M165	Mountain Lion	<i>Felis concolor</i>	>40ha	high
M166	Bobcat	<i>Felis rufus</i>	>40ha	high
M181	Mule Deer	<i>Odocoileus hemionus</i>	>40ha	high
R022	Western Fence Lizard	<i>Sceloporus occidentalis</i>	<40ha	low
R040	Southern Alligator Lizard	<i>Elgaria multicarinata</i>	<40ha	low
R042	Northern Alligator Lizard	<i>Elgaria coerulea</i>	<40ha	low
R061	Common Garter Snake*	<i>Thamnophis sirtalis</i>	<40ha	low
R063	Western Aquatic Garter Snake*	<i>Thamnophis couchii</i>	<40ha	low

[illegible]

0	0	0	0.33	0.33	0.33	0.33	0.33	0.33	0
0	1	0	0	0	0	0	0	0	0
0	0.67	0	0	0	0	0	0	0	0
0	1	0	0	0	0	0	0	0	0
0.67	0	1	0.33	1	1	0.33	1	1	0.67
0	0.33	0	0	0	0	0	0	0	0
0	0	0	0.33	0.33	0	0.33	0.33	0	0
0	0	0.33	0.33	0.33	0.33	0.67	0.67	0.67	0.33
1	1	1	1	1	1	1	1	1	1
0.67	0	0.33	1	1	0.67	1	1	0.67	0.33
0	1	0	0	0	0	0	0	0	0
0	0	0	0.67	0.67	0.67	1	1	1	0
0	0	0.33	0.67	1	1	1	1	1	0.33
0	0	0	0.67	0.67	0.67	1	1	1	0
0	0	0.33	0	0.67	0.67	0	1	1	0.33
0	1	0	0	0	0	0	0	0	0
0	1	0	0	0	0	0	0	0	0
1	0.33	0	0	0	0	0	0	0	0
0	0	0.33	0	0.33	0.33	0	0.33	0.33	0
0	0	0.33	0.33	0.67	0.67	0.33	0.67	0.67	0.67
0	0	0.67	0.67	0.67	0.33	0.67	0.67	0.33	1
0	0.67	0	0	0	0	0	0	0	0
0	0	0	0.67	0.33	0	0.67	0.33	0	0
1	0	0.33	0.33	0	0	0	0	0	1
0	0	1	1	0.67	0.33	1	0.67	0.33	0
0	0	0	0	0	0	0	0	0	0
0.33	0	0.67	0	0.33	0.67	0	0.33	0.67	1
0.67	0	1	1	0.67	0.33	1	0.67	0.33	1
0.33	0	0.33	0.33	0	0	0.33	0	0	0
0.67	0	1	1	0.33	0.67	1	0.33	0.67	0.33
1	0	0.33	0.33	0.33	0	0.33	0.33	0	0.67
0	1	0	0	0	0	0	0	0	0
0	0.67	0	0	0	0	0	0	0	0
0.67	0	1	1	1	0.67	1	1	0.67	1
0	0.33	0	0	0	0	0	0	0	0
0	0	0.67	1	1	1	1	1	1	0
0	0.33	0	0	0	0	0	0	0	0
0	0	0.67	1	1	0.67	1	1	0.67	0
0	0	0.33	0.33	0.33	0	0.33	0.33	0	0
0	0.33	0.33	0.33	0	0	0.33	0	0	0
0	1	0.33	0.33	0	0	0.33	0	0	0.33
0	1	0.33	0	0	0	0	0	0	0
0	0.67	0	0	0	0	0	0	0	0
0.33	0	0.67	0.67	0.33	0	0.67	0	0	0.67
0	1	0	0	0	0	0	0	0	0
0	0.67	0	0	0	0	0	0	0	0
0.67	1	1	1	0.67	0	1	0.67	0	0.67

0.67	0	0.33	0.33	0	0	0.33	0	0	0
0	0	0	0	0	0	0	0	0	0.67
1	0.67	1	1	0.67	0.33	1	0.67	0.33	1
0	0	0.33	0.33	0	0	0.33	0	0	0
0	0	0.33	0.33	0	0	0.33	0	0	0.33
0.33	0	0.67	0.67	0.33	0.33	0.67	0.33	0.33	0.33
0	0	0	0	0	0	0	0	0	0
0	0	0	0.33	0.67	0.33	0.33	0.67	0.33	0
0.33	0	0	0.67	0.33	0	0.67	0.33	0	0.67
0.33	0	0.33	0.33	0	0	0.33	0	0	0
0	0	0	0.33	0.33	0.33	0.33	0.33	0.33	0.67
0	0	0	0.33	0.33	0	0.33	0.33	0	0.67
0.33	0.33	0.33	0.33	0	0	0.33	0	0	0
0.33	0	0	0	0	0	0	0	0	0
1	0.67	0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33
0	0	0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33
0	0.33	0	0	0	0	0	0	0	0
1	0	0.67	0.33	0	0	0.33	0	0	0.67
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0.67	0	1	1	1	1	1	1	1	0.33
0	0	0	0	0	0	0	0	0	0
0.33	0.67	0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33
1	0	0.67	1	0.67	0.67	1	0.67	0	0.67
0.67	1	0.67	0.67	0.67	0.67	0.67	0.67	0.67	0.67
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0.67	0	0.33	0	0	0	0	0	0	0.33
0.33	0.67	0.67	0.33	0	0	0.33	0	0	0.33
0.67	0	0.67	0.33	0.33	0	0.33	0	0	0.67
0.67	1	1	1	1	0.33	1	1	0.33	1
0.67	0.67	1	1	0.67	0.33	1	0.67	0.33	0.33
0	0	0	0	0	0	0	0	0	0
0	0.33	0.33	0.33	0	0	0.33	0	0	1
0.33	0.33	0.33	0	0	0	0	0	0	0.67
0.67	1	1	0.67	0.33	0.33	0.67	0.33	0.33	0.67
0.67	1	1	1	0.67	0.33	1	0.67	0.33	1
0.33	0	0.33	0.33	0.33	0.67	0.33	0.67	0.67	0
0.33	0	0.33	0	0.67	1	0.67	1	1	0.33
1	0	0.67	0.33	0.67	0.67	0.67	1	1	0.67
0	0.33	0.67	0.33	0	0	0.33	0	0	0.67
0.33	1	1	0.67	0.67	0.33	0.67	0.67	0.33	1
0	1	1	0.67	0.67	0	0.67	0.67	0	0
0	1	0	0	0	0	0	0	0	0
0	0.67	0	0	0	0	0	0	0	0
0	1	0	0	0	0	0	0	0	0
0	0.67	1	1	0.67	0.67	1	0.67	0.33	0.67

0.33	0.33	0.67	0.67	0.33	0.33	0.67	0.33	0.33	0.67
1	0.33	1	0.67	0.67	0.33	0.67	0.33	0	1
1	0	0.67	0.67	0.67	0.33	0.67	0.67	0.33	0.33
1	1	1	0.67	0.67	0.33	0.67	0.33	0.33	1
1	0.33	0.67	0.67	0.67	0.67	0.67	1	0.67	0.67
1	0.67	0.33	0.67	1	0.67	0.67	1	1	0.33
0.67	0	0.33	0.33	0.67	0.67	0.33	0.67	0.67	0.33
0.67	0	0	0	0.33	1	0	0.67	1	0
1	0.33	0.67	0.67	0.67	1	0.67	0.67	1	0.33
1	0.67	1	0.33	0.67	0.67	0.33	0.67	0.67	1
0.33	1	1	0.67	0	0	0.33	0	0	1
1	1	1	0.33	0.33	0.33	0.67	0.33	0	0.33
1	1	1	0.67	0.67	0.33	0.33	0.33	0.33	0.33
1	1	1	0.67	0.67	0.67	0.67	0.67	0.67	1
1	1	1	1	0.67	0.33	0.67	0.33	0.33	1
1	1	1	0.67	0.67	0.33	0.67	0.33	0.33	1
0	1	0.67	0.67	0.67	0.33	0.67	0.67	0	0.67
0	1	0	0	0	0	0	0	0	0
0	1	0.67	0.67	0.67	0.33	0.33	0.33	0	0.67
0	0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33	0
0.67	0.67	0	0	0	0	0	0	0	0

[illegible]

0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0.33	0.67	0.67	0.33	0.67	0.67	0.67	0.33	0.67	0.67
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0.67	0.67	0.67	1	1	1	0.33	0.33	0.33	0.33
1	1	1	1	1	1	1	1	1	1
1	1	0.67	1	1	0.67	0.33	1	1	0.67
0	0	0	0	0	0	0	0	0	0
0.33	0.33	0.33	0.67	0.67	0.67	0	0.67	0.67	0.67
0.67	0.67	0.67	1	1	1	0	0.33	0.33	0.33
0	0	0	0.33	0.33	0.33	0	0	0	0
0	0.67	0.67	0	0.67	0.67	0.33	0.33	0.67	0.67
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0.33	0.33	0.67	0.67
0.33	0.67	1	0.67	1	1	0.67	0.67	1	1
1	1	0.67	1	1	0.67	0.33	0.33	0.33	0.33
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0.33	0.33	0.33
1	0	0	1	0	0	0.33	0.33	0	0
0.67	0.33	0	0.67	0.33	0	0.67	0.67	0.67	0.67
0	0	0	0	0	0	0	0	0	0
0	0.67	1	0	0.67	0.67	1	0.33	0.67	1
1	0.67	0.33	1	0.67	0.33	0.67	0.67	0.67	0.67
0	0	0	0	0	0	0	0	0	0
0.33	0.33	0.33	0.33	0.33	0.33	0.67	0.67	0.33	0.33
0.67	0.67	0	0.67	0.67	0	0.67	0.67	0.67	0
0	0	0	0	0	0	0	0.33	0.33	0.33
0	0	0	0	0	0	0	0	0	0
1	1	0.67	1	1	0.67	1	1	1	0.67
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0.67	1	1	1
0	0	0	0	0	0	0.33	0	0.67	0.67
0.33	0.33	0	0.33	0.33	0	0.67	1	1	1
0	0	0	0	0	0	0.33	0.33	0.33	0.33
0	0	0	0	0	0	0	0	0	0
0.33	0	0	0.33	0	0	0.33	0.33	0.33	0.33
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0.67	0.33	0	0.67	0	0	0.67	0.67	0.33	0.33
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0.67	0.33	0	0.67	0.33	0	0.67	1	0.33	0.33

0	0	0	0	0	0	0	0	0	0
0.67	0	0	0.67	0	0	0.67	0.67	0.33	0.33
1	0.67	0.33	1	0.67	0.33	1	1	0.67	0.33
0	0	0	0	0	0	0	0	0	0
0.33	0	0	0.33	0	0	0	0	0	0
0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33
1	0.67	0.33	1	0.67	0.33	0	1	0.67	0.67
0	0	0	0	0	0	0	0.33	0.33	0
1	0.67	0.33	1	0.67	0.33	0.33	1	0.67	0.67
0	0	0	0	0	0	0	0	0	0
1	1	0.67	1	1	0.67	0	0.67	0.67	0.33
1	0.67	0.33	1	0.67	0.33	0.67	0.67	0.33	0.33
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0.67	0.67	0.67
0.33	0.33	0.33	0.33	0.33	0.33	0.67	0.33	0.33	0.33
0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33
0	0	0	0	0	0	0.33	0.33	0.33	0.33
0.33	0	0	0.33	0	0	0.67	0.33	0	0
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33
0	0	0	0	0	0	0	0	0	0
0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33
1	0.67	0.67	1	0.67	0.67	0.67	1	0.67	0.67
0.67	0.67	0.67	0.67	0.67	0.67	0.67	0.67	0.67	0.67
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0.67	0	0	0
0.33	0	0	0.33	0	0	0.33	0.33	0	0
0.67	0.33	0	0.33	0	0	0.67	0.33	0.33	0
0.67	1	0.33	0.67	1	0.33	1	1	1	0.33
0.33	0.33	0.33	1	1	0.67	1	1	0.67	0.33
0	0	0	0	0	0	0	0	0	0
1	1	0.67	1	0.67	0.33	0.33	0.33	0	0
0.33	0	0	0.33	0	0	0.33	0	0	0
0.33	0.33	0	0.33	0.33	0	0.67	0.33	0.33	0
1	0.67	0.33	1	0.67	0.33	1	1	0.67	0.33
0	0	0	0	0	0	0	0	0	0
0	0.67	1	0.67	1	1	0.33	0	0.67	1
0.33	0.33	0.67	0.67	1	1	0.67	0.33	0.67	0.67
0.33	0.67	0.33	0.33	0.67	0.33	0.67	0.33	0	0
1	1	0.67	1	0.67	0.67	1	0.67	0.67	0.33
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0.67	0.67	0.67	0.67	0.67	0.67	1	1	0.67	0.67

[illegible]

RFR3S	RFR3M	RFR3D	WFR1	WFR2S	WFR2M	WFR2D	WFR3S	WFR3M	WFR3D
1	1	1	1	1	1	1	1	1	1
0	0.33	0.33	1	0	1	1	0	1	1
0	0.33	0.33	0.67	0	0.67	0.67	0	0.67	0.67
0	1	1	0.33	0	1	1	0	1	1
1	1	1	0.67	1	1	1	1	1	1
1	0	0	1	1	1	1	1	1	1
0.67	0.33	0.33	0	0.67	0.67	0.67	1	1	1
1	1	1	1	1	1	1	1	1	1
1	0.67	0.67	0.33	0.67	0.67	0.67	1	1	1
0	0	0	0	0	0	0	0	0	0
0	0	0	0.67	0.67	0.67	0.33	0.67	0.67	0.33
0.67	0.33	0.33	1	1	0.67	0.67	1	0.67	0.67
0.33	0.33	0.33	0.33	0.67	1	1	1	1	1
0	0	0	0.67	0.67	0.67	0.33	0.67	0.67	0.33
0.67	0.67	0.33	0	1	1	0.67	1	1	0.67
0.33	0.33	0.33	0	0.33	0.33	0.33	0.33	0.33	0.33
0.67	0.67	0.67	0	1	1	0.67	1	1	0.67
0	0	0	0.33	1	1	0.67	1	1	0.67
0.33	1	1	0	0.33	0.33	0.33	0.33	1	1
0	0	0	0.67	0.33	0.67	0.67	0.33	0.67	0.67
0.33	0.67	0.67	0	0.67	1	1	0.67	1	1
0.33	0	0	1	1	0	0	1	0	0
0.33	0	0	0.33	0.33	0	0	0.33	0	0
1	1	1	1	1	1	1	1	1	1
0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33
0	0	0	1	1	1	1	1	1	1
0	0	0	0.67	0.67	0.33	0	0.67	0.33	0
0.33	0.33	0.33	1	1	0.33	0	1	0.33	0
0	0	0	0.67	0.67	0.33	0	0.67	0.33	0
0	0	0	0	1	0.67	0.33	1	0.67	0.33
0.67	0.67	0.67	0.33	1	0.67	0.33	1	0.67	0.33
1	0.67	0.33	0	0.33	0.33	0.33	0.33	0.33	0.33
1	0.67	0.67	0.33	1	1	0.67	1	1	0.67
1	0.67	0.67	0.33	1	1	0.67	1	1	0.67
0.67	0.67	1	0	0	0	0	0	0	0
0.67	0.67	0.67	0.33	0.67	0.67	0.67	1	1	0.67
0.33	0.67	0.67	0.33	0.33	0.67	0.67	0.67	1	1
1	0.67	0.67	0.33	0.33	0.67	0.67	1	1	1
1	1	1	0.67	1	1	1	1	1	1
1	1	1	0	0.33	0.33	0.67	0.33	0.67	1
0.67	0.33	0.33	1	0.67	0.67	0.33	0.67	0.67	0.33
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0	0	0	0.33	0.33	0	0	0.33	0.33	0

0	0	0	0.33	1	1	0.67	1	1	0.67
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0	0	0	1	1	0.67	0.33	1	0.67	0.33
0.67	1	1	1	0.33	1	1	0.33	1	1
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0.67	0.67	0.67	0	0	0	0	0	0	0
1	1	1	1	1	1	1	1	1	1
1	1	0.67	0.33	1	1	0.67	1	1	0.67
0	0	0	0	0	0	0	0	0	0
1	1	1	0	0.67	0.67	0.67	1	1	1
0.33	0.33	0.33	0.33	0.67	1	1	1	1	1
0	0	0	0	0.67	0.67	0.67	1	1	1
0.67	1	1	0.33	0	1	1	0	1	1
0	0	0	0	0	0	0	0	0	0
0	0	0	0.33	0.33	0	0	0.33	0	0
0	0	0	0	0	0	0	0	0	0
0.33	0.67	0.67	0.67	0.33	0.67	0.67	0.33	0.67	1
0.67	1	1	0.67	0.67	1	1	0.67	1	1
0.33	0.33	0.33	1	1	1	0.67	1	1	0.67
0	0	0	0	0	0	0	0	0	0
0.33	0.33	0.33	0	0.67	0.33	0	0.67	0.33	0
0.33	0	0	1	1	0	0	0	0	0
0.67	0.67	0.67	1	1	0.67	0.33	1	0.67	0.33
0	0	0	0.33	0	0.33	0.33	0	0.33	0.33
0.33	1	1	1	0.67	1	1	0.67	1	1
0.67	0.67	0.67	1	1	0.67	0.33	1	0.67	0.33
0	0	0	0.33	0.33	0.33	0	0.33	0.33	0
0.33	0.33	0.33	0.67	0.67	0.67	0.33	0.67	0.67	0.33
0.67	0.67	0.33	0.33	0.33	0.33	0	0.33	0.33	0
0.33	0.33	0.33	1	1	0.67	0.33	0	0	0
0	0	0	1	1	1	0.33	0	0	0
1	1	1	1	1	1	0.67	1	1	0.67
0	0	0	0.33	0.33	0.33	0.33	0.33	0.33	0.33
0.33	0.33	0.33	0.67	1	1	1	1	1	1
0	0.67	0.67	0	0	0	0	0	0	0
1	1	1	0.67	1	1	1	1	1	1
0.33	0.33	0.33	0.67	0.67	0.67	0.33	0.67	0.67	0.33
0	0	0	0.33	0.33	0	0	0.33	0	0
0.33	0.33	0.33	1	1	0	0	0.67	0	0
0	0	0	0.67	0.33	0.33	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0.67	0.33	0.33	1	1	0.67	0	1	0	0
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
1	0.33	0.33	1	1	0.67	0	1	0.33	0

0	0	0	0.33	0.33	0	0	0.33	0	0
0.67	0.33	0.33	0.33	0.33	0	0	0.33	0	0
1	1	1	1	1	0.67	0.33	1	0.67	0.33
0	0	0	0.33	0.33	0	0	0.33	0	0
0	0	0	0.67	0.33	0.67	0.67	0.33	0.67	0.67
0.33	0.33	0.33	0.67	0.67	0.33	0.33	0.67	0.33	0.33
1	0.67	0.67	0	0	0	0	0	0	0
0.33	0.33	0	0	0.67	1	0.67	0.67	1	0.67
1	0.67	0.67	0.33	0.67	0.33	0	1	0.33	0
0	0	0	0.67	0.67	0	0	0.67	0	0
0.67	0.33	0.33	0	0.33	0.33	0.33	0.67	0.67	0.33
0.67	0.67	0.33	0.33	0.67	0.33	0	0.67	0.33	0
0	0	0	0.33	0.33	0	0	0.33	0	0
0.67	0.67	0.67	0	0.67	1	1	0.67	1	1
0.33	0.33	0.33	0.67	0.33	0	0	0.33	0	0
0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33
0.33	0.33	0.33	0.67	0.67	1	1	0.67	1	1
0.33	0	0	0.67	0.33	0	0	0.33	0	0
0	0	0	0.33	0.33	0.33	0.33	0.33	0.33	0.33
0	0	0	0.67	0.67	0.67	0.67	0.67	0.67	0.67
0.33	0.33	0.33	0.67	0.67	1	1	1	1	1
0	0	0	0.33	0.33	0.33	0.33	0.33	0.33	0.33
0.33	0.33	0.33	0.67	0.67	0.67	0.67	0.67	0.67	0.67
1	0.67	0.67	0.67	1	0.67	0.67	1	0.67	0.67
0.67	0.67	0.67	1	1	1	1	1	1	1
0	0	0	0.33	0.33	0.33	0.33	0.33	0.33	0.33
0	0	0	0.33	0.33	0.33	0.33	0.33	0.33	0.33
0	0	0	0.33	0	0	0	0	0	0
0.33	0	0	0.67	0.33	0	0	0.33	0	0
0.33	0	0	1	1	0.67	0	0.67	0	0
1	1	0.33	1	1	1	0.33	1	1	0.33
1	0.67	0.33	1	1	0.67	0.33	1	1	0.33
0	0	0	1	1	0.67	0.33	1	0.67	0.33
0.33	0	0	0.67	0.67	0.33	0.33	0.67	0.67	0.33
0	0	0	0.33	0	0	0	0	0	0
0.33	0.33	0	1	0.67	0.33	0.33	0.67	0.33	0.33
1	0.67	0.33	1	1	0.67	0.33	1	0.67	0.33
0	0	0	0.33	0.33	0.67	0.67	0.67	0.67	0.33
0.67	1	1	0.33	0	0.67	1	0.67	1	1
0.67	1	1	0.67	0.33	0.67	1	0.67	1	1
0.33	0	0	0.67	0.33	0	0	0.33	0	0
0.67	0.67	0.33	1	0.67	0.67	0.67	0.67	0.67	0.67
0	0	0	1	0.67	0	0	0.67	0	0
0	0	0	0.67	0.67	0.67	0.33	0.67	0.67	0.33
0	0	0	0	0	0	0	0	0	0
0	0	0	1	1	0.33	0.33	0.67	0.67	0.33
1	0.67	0.33	0.67	0.67	0.33	0.33	0.67	0.67	0.67

[illegible]

SMC1	SMC2S	SMC2M	SMC2D	SMC3S	SMC3M	SMC3D	SCN1	SCN2S	SCN2M
1	1	1	1	1	1	1	0	0	0
1	0	1	1	0	1	1	0	0	0
0.67	0	0.67	0.67	0	0.67	0.67	0	0	0
0.33	0	1	1	0	1	1	0.67	0.33	1
0.67	1	1	1	1	1	1	0	0	0
1	1	1	1	1	1	1	1	1	0
0	0.67	0.67	0.67	1	1	1	0	0	0
1	1	1	1	1	1	1	0	0	0
0.33	0.67	0.67	0.67	1	1	1	0.33	0.67	0.67
0.67	0.67	0.33	0.33	0.67	0.33	0.33	0	0	0
0.67	0.67	0.67	0.33	0.67	0.67	0.33	0	0	0
1	1	0.67	0.67	1	0.67	0.67	1	1	0.67
0.33	0.67	1	1	1	1	1	0	0	0
0.67	0.67	0.67	0.33	0.67	0.67	0.33	0	0	0
0	1	1	0.67	1	1	0.67	0	0	0
0	0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.33
0	1	1	0.67	1	1	0.67	0	0	0
0.33	1	1	0.67	1	1	0.67	0	0	0
0	0.33	0.33	0.33	0.33	1	1	0	0	0
0.67	0.33	0.67	0.67	0.33	0.67	0.67	0	0	0
0	0.67	1	1	0.67	1	1	0	0	0
1	1	0	0	1	0	0	0	0	0
0.33	0.33	0	0	0.33	0	0	0	0	0
1	1	1	1	1	1	1	0	0	0
0.33	0.33	0.33	0.33	0.33	0.33	0.33	0	0	0
1	1	1	1	1	1	1	0	0	0
0.67	0.67	0.33	0	0.67	0.33	0	0	0	0
1	1	0.33	0	1	0.33	0	0.33	0.67	0.33
0	0	0	0	0	0	0	0	0	0
0	1	0.67	0.33	1	0.67	0.33	0	0	0
0.33	1	0.67	0.33	1	0.67	0.33	0	0	0
0	0.33	0.33	0.33	0.67	0.67	0.33	0	0	0
0.33	1	1	0.67	1	1	0.67	0.67	0.67	0.67
0.33	1	1	0.67	1	1	0.67	0	0.33	0.33
0	0.33	0.33	0.33	0.33	0.33	0.33	0	0.67	0.67
0.33	0.67	0.67	0.67	1	1	0.67	0	0.33	0.33
0.33	0.33	0.67	0.67	0.67	1	1	0	0	0
0.67	0.67	0.67	0.67	1	1	1	0	0	0
0.67	1	1	1	1	1	1	0.33	0.33	0.33
0	0	0.33	0.67	0.33	1	1	0	0	0
1	0.67	0.67	0.33	0.67	0.67	0.33	1	1	0.67
1	0.33	1	1	0.33	1	1	0	0	0
0	0	0	0	0	0	0	0	0	0
0.33	0.67	0	0	1	1	0	0	0	0

0.33	1	1	0.67	1	1	0.67	0	0	0
0	0	0	0	0	0	0	0	0	0
0.33	0.33	0.33	0.33	0.33	0.33	0.33	0	0	0
1	1	0.67	0.33	1	0.67	0.33	0	0	0
1	0.33	1	1	0.33	1	1	0.33	0	0.33
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0.33	0.67	0.67
1	1	1	1	1	1	1	1	0.67	0.67
1	1	1	0.67	1	1	0.67	0.33	0.67	0.67
0.33	0.33	0.33	0.33	0.33	0.33	0.33	0	0	0
0.67	0.67	0.67	0.67	1	1	1	0	0	0
0.33	0.33	0.67	0.67	0.67	0.67	0.67	0	0.33	0.33
0	0.67	0.67	0.33	1	1	0.33	0	0	0
0.33	0.33	1	1	0.33	1	1	0	0	0
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0.67	0.33	0.67	0.67	0.33	0.67	1	0	0	0
0.67	0.67	1	1	0.67	1	1	0.33	0.33	0.33
0.67	0.67	0.67	0.33	0.67	0.67	0.33	1	1	1
0	0	0	0	0	0	0	0	0	0
0	0.67	0.33	0	0.67	0.33	0	0	0	0
0.33	0.33	0	0	0.33	0	0	0.67	0.67	0
1	1	0.67	0.33	1	0.67	0.33	0	0	0
0.33	0	0.33	0.33	0	0.33	0.33	0	0	0
1	0.67	1	1	0.67	1	1	1	0	0.67
1	1	0.67	0.33	1	0.67	0.33	0.67	0.67	0.33
0.33	0.33	0.33	0	0.33	0.33	0	0	0	0
0.67	0.67	0.67	0.33	0.67	0.67	0.33	0	0	0
1	1	1	0.33	1	1	0.33	0	0	0
1	1	0.67	0.33	0.67	0.67	0.33	0	0	0
0.67	0.67	0.67	0.33	0	0	0	0	0	0
1	1	1	0.67	1	1	0.67	0.67	0.67	0.67
0.33	0.33	0.33	0.33	0.33	0.33	0.33	0	0	0
0.67	1	1	1	1	1	1	0	0	0
0.33	0	0.67	0.67	0	0.67	0.67	0	0	0
1	1	1	1	1	1	1	0	0	0
0.67	0.67	0.67	0.33	0.67	0.67	0.33	0	0	0
0.33	0.33	0	0	0.33	0	0	0	0	0
0.67	0.67	0.33	0	0.67	0.33	0	0	0	0
0.67	0.33	0.33	0.33	0.33	0.33	0.33	0	0	0
0	0	0	0	0	0	0	0	0	0
1	1	0.67	0.33	1	0.67	0.33	0.33	0.33	0
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
1	1	0.67	0.33	1	0.67	0.33	0	0	0

0.67	0.67	0	0	0.67	0	0	0	0	0
0.67	0.67	0.33	0	0.67	0.33	0	0	0	0
1	1	0.67	0.33	1	0.67	0.33	1	1	0.67
0	0	0	0	0	0	0	0	0	0
0.67	0.33	0.67	0.67	0.33	0.67	0.67	0	0	0
0.67	0.67	0.67	0.33	0.67	0.67	0.33	0	0	0
0	0	0	0	0	0	0	0	0.67	0.33
0	0.67	1	0.67	0.67	1	0.67	0	0	0
0.67	0.67	0.33	0.33	0.67	0.33	0.33	0.33	1	0.67
0	0	0	0	0	0	0	0	0	0
0	0.33	0.33	0.33	0.67	0.67	0.33	0	0.33	0.33
0.33	0.67	0.67	0.33	0.67	0.67	0.33	0.33	0.67	0.33
0.33	0.33	0.33	0.33	0.33	0.33	0.33	0	0	0
0	0.67	1	1	0.67	1	1	0	0	0
0.67	0.33	0	0	0.33	0	0	0.33	0.33	0.33
0.33	0.33	0.33	0.33	0.33	0.33	0.33	0.67	0.67	0.67
0.67	0.67	1	1	0.67	1	1	0	0	0
0.67	0.33	0	0	0.33	0	0	0	0	0
0.33	0.33	0.33	0.33	0.33	0.33	0.33	0	0	0
0.67	0.67	0.67	0.67	0.67	0.67	0.67	0	0	0
0.67	0.67	1	1	1	1	1	0.33	0.33	0.33
0.33	0.33	0.33	0.33	0.33	0.33	0.33	0	0	0
0.67	0.67	0.67	0.67	0.67	0.67	0.67	0	0	0
0.67	1	0.67	0.67	1	0.67	0.67	0	0	0
1	1	1	1	1	1	1	0.33	0.33	0.33
0.33	0.33	0.33	0.33	0.33	0.33	0.33	0	0	0
0.33	0.33	0.33	0.33	0.33	0.33	0.33	0	0	0
0.33	0	0	0	0	0	0	0.33	0	0
0.67	0.33	0	0	0.33	0	0	0.33	0.33	0
1	1	0.67	0	0.67	0	0	1	1	0.67
1	1	1	0.33	1	1	0.33	1	1	1
1	1	0.67	0.33	1	1	0.33	0.33	0.33	0.33
1	1	0.67	0.33	1	0.67	0.33	0	0	0
0.67	0.67	0.33	0.33	0.67	0.67	0.33	0.33	0.33	0.33
0.33	0	0	0	0	0	0	0.67	0.33	0
1	0.67	0.33	0.33	0.67	0.33	0.33	0.67	0.33	0.33
1	1	0.67	0.33	1	0.67	0.33	1	1	0.67
0.33	0.33	0.67	0.67	0.67	0.67	0.33	0	0	0
0.33	0	0.67	1	0.67	1	1	0.33	0	0.67
0.67	0.33	0.67	1	0.67	1	1	0.67	0.33	0.33
0.67	0.33	0	0	0.33	0	0	0.67	0.33	0
1	0.67	0.67	0.67	0.67	0.67	0.67	1	0.67	0.67
1	0.67	0	0	0.67	0	0	0.67	0.67	0.33
0.67	0.67	0.67	0.33	0.67	0.67	0.33	0	0	0
0	0	0	0	0	0	0	0	0	0
1	1	0.33	0.33	0.67	0.67	0.33	0	0	0
0.67	0.67	0.33	0.33	0.67	0.67	0.67	1	0.67	0.67

[illegible]

SCN2D	SCN3S	SCN3M	SCN3D
0	0	0	0
0	0	0	0
0	0	0	0
1	0.33	1	1
0	0	0	0
0	1	0	0
0	0.33	0.33	0.33
0	0	0	0
0.67	0.67	0.67	0.67
0	0	0	0
0	0	0	0
0.67	1	0.67	0.67
0	0	0	0
0	0	0	0
0	0.33	0.33	0.33
0	0.33	0.33	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0.67	0.33	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0.67	1	1	0.67
0	0.33	0.33	0
0.33	0.67	0.67	0.33
0	0.33	0.33	0
0	0	0	0
0	0	0	0
0.33	0.33	0.33	0.33
0	0	0	0
0.33	1	0.33	0
0	0	0	0
0	0	0	0
0	0	0	0

0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0.33	0	0.33	0.33
0	0	0	0
0	0	0	0
0.67	1	1	1
0.67	0.67	0.67	0.67
0.33	0.67	0.67	0.33
0	0	0	0
0	0	0	0
0.33	0.67	0.67	0.67
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0.67	0.33	0.67	0.67
0.67	1	1	0.67
0	0	0	0
0	0	0	0
0	0.67	0	0
0	0	0	0
0	0	0	0
1	0	0.67	0.67
0.33	0.67	0.33	0.33
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0.33	0.67	0.67	0.33
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0	0.33	0	0
0	0	0	0
0	0	0	0
0	0	0	0

0	0	0	0
0	0	0	0
0.33	1	0.67	0.33
0	0	0	0
0	0	0	0
0	0	0	0
0.33	0.67	0.33	0.33
0	0	0	0
0.33	1	0.67	0.33
0	0	0	0
0.33	0.33	0.33	0.33
0.33	0.67	0.33	0.33
0	0	0	0
0	0	0	0
0.33	0.33	0.33	0.33
0.67	0.67	0.67	0.67
0	0	0	0
0	0	0	0
0	0	0	0
0	0	0	0
0.33	0.33	0.33	0.33
0	0	0	0
0	0	0	0
0	0	0	0
0.33	0.33	0.33	0.33
0	0	0	0
0	0	0	0
0	0	0	0
0	0.33	0	0
0.33	0.67	0	0
0.33	1	1	0.33
0.33	1	1	0.67
0	0	0	0
0.33	0.33	0	0
0	0.33	0	0
0	0.33	0.33	0
0.33	1	0.67	0.33
0	0	0	0
1	0.67	1	1
0.67	0.67	1	1
0	0.33	0	0
0.33	0.67	0.67	0.33
0	0.67	0.33	0
0	0	0	0
0	0	0	0
0	0	0	0
0.67	0.67	0.67	0.33

0.33	0.67	0.33	0.33
0.33	0.67	0.33	0
0.33	0.67	0.67	0.33
0.33	0.67	0.33	0.33
0.67	0.67	0.67	0.33
0.67	0.67	1	1
1	0.67	1	1
1	0	0.67	1
1	0.67	1	1
0.33	0.67	0.33	0.33
0	0	0	0
0	0	0	0
0	0	0	0
0.67	0.33	0.67	0.33
0.33	0.33	0.33	0.33
0.33	0.67	0.33	0.33
0	0	0	0
0	0	0	0
0	0.33	0.33	0
0	0	0	0
0	0	0	0