


When bigger isn't better—Implications of large high-severity wildfire patches for avian diversity and community composition

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Abstract

Aim: Wildfires increasingly create large high-severity patches with interior areas far from less disturbed habitats. We evaluated how these trends impact bird communities by investigating the effect of internal distance from lower-severity areas, high-severity patch size, and years since fire on avian alpha and beta diversity.

Location: Sierra Nevada, California, USA.

Methods: Bird occurrence data were collected during 2009–2017 within high-severity patches of 27 wildfires representing 1–30 years since disturbance. A two-step multispecies occupancy method was used to account for imperfect detection of 94 species and estimate effects of patch characteristics on community richness and dissimilarity.

Results: Community richness decreased with distance from patch edge and with patch size. Richness increased with years since fire, but this pattern was dependent on distance from edge with higher peak richness (23 species) near edges than interiors (18 species). Community dissimilarity was not associated with distance, indicating that large high-severity patch interiors contain subsets of, rather than complements to, edge communities. Dissimilarity peaked later with increasing patch size. Guild richness of tree and primary cavity nesters was negatively associated with distance and patch size. Richness of ground and shrub nesters was insensitive to distance, while shrub nester richness increased somewhat with patch size. Due to declines among other species, ground and shrub nesters made up a greater percentage of the avian community within the interiors of large high-severity patches.

Main conclusions: As fire activity increases due to accumulating forest fuels and accelerating climate change, high-severity patches and their resulting early-seral habitats are becoming more extensive with less edge and more interior area. Such changes are likely to decrease avian diversity locally and shift community composition away from forest-associated species. Management actions that promote the full range of fire effects but limit high-severity patch size may best conserve bird diversity within fire-adapted ecosystems.

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KEYWORDS

avian diversity, fire ecology, fire suppression, heterogeneity, landscape ecology, multispecies occupancy model, patch size, pyrodiversity, wildfire

1 | INTRODUCTION

Wildfire is a fundamental ecological process that influences the distribution of ecosystems (Bond & Keeley, 2005; McLauchlan et al., 2020; Whittaker, 1975), landscape pattern (Turner, 2010) and vegetation structure (Agee, 1996). Wildfire patterns are shifting globally (Bowman et al., 2011; Flannigan et al., 2009) and especially in fire-prone forests of California where wildfires are creating ever larger patches of high-severity effects in which forests are converted to early-seral habitats (Abatzoglou & Williams, 2016; Miller et al., 2012; Westerling, 2016). Further, increasingly large extents of these high-severity patches are located far from lower-severity areas where fire-caused vegetation mortality and habitat structure is more variable and forests remain largely intact (Steel et al., 2018). Such shifts influence landscape heterogeneity and habitat availability, potentially driving patterns of species diversity and community composition. A greater understanding of how changes in the spatial pattern of high-severity fire are impacting bird communities is necessary to effectively manage fire-adapted ecosystems and conserve biodiversity.

Bird species vary widely in their response to wildfire and in their associations with post-fire habitats (Fontaine & Kennedy, 2012). High-severity fire that removes the majority of pre-fire vegetation (i.e., >75% canopy cover mortality) leaves fire-killed trees (snags) in the short term, with early-successional habitats such as grasslands and shrublands developing in the years immediately following the fire (White et al., 2016). If reestablishment of conifer or hardwood trees is successful and in the absence of repeated severe disturbance, the habitat will transition back to a forested state in the medium to long term (Agee, 1996; van Wagtenonk et al., 2018). In contrast, low- to moderate-severity effects reduce tree and understory density but maintain mature canopy cover (Miller et al., 2009) and habitat structure more similar to unburned areas (Roberts et al., 2021). Many bird species in the Sierra Nevada, California are adapted to particular combinations of burn severity and time since fire, including early-successional habitats created by high-severity fire (Fontaine et al., 2009; Taillie et al., 2018; White et al., 2016). Indeed, peak abundance of at least a quarter of the bird community occurs in recently burned forests (≤ 15 years) across the severity gradient with an equivalent number most abundant in unburned forests (Taillie et al., 2018). While bird communities in contrasting severities have been relatively well studied, our understanding of how spatial patterns of high-severity patches influence bird diversity is limited.

The spatial pattern of high-severity fire may influence bird diversity at both the patch and the local scales. Patch size influences the amount and connectivity of early-seral habitat created by a given fire, as well as the amount of habitat isolated from lower-severity areas. Within a given high-severity patch, distance from its edge can

drive the level of landscape heterogeneity a bird perceives. High-severity areas near surviving forests (i.e., low- to moderate-severity areas) potentially provide complementary resources not available to birds occupying the high-severity patch interior (Stillman et al., 2019, 2021). In these fire-adapted landscapes, bird diversity may maximize in areas characterized by heterogeneous fire histories (aka pyrodiversity) where variation in burn severity and other fire regime characteristics is high (Beale et al., 2018; Martin & Sapsis, 1992; Steel et al., In Press; Tingley et al., 2016), suggesting that high-severity patch edges may contain greater bird diversity than interiors. Wildfire spatial pattern is also an important determinant of vegetation succession (Turner, 2010) with likely consequences for avian compositional shifts in the decades following disturbance. In forests composed of tree species that are unable to resprout or regenerate through serotiny, the distance to surviving trees determines the rate of post-fire forest recovery (Greene & Johnson, 1995; Shive et al., 2018; Welch et al., 2016) and bird habitat quality. While individual bird species respond to fire patterns idiosyncratically, generalized responses may emerge among functional groups with species reliant on residual trees (e.g., tree nesters) showing opposing effects as those reliant on early-seral resources (e.g., shrub nesters).

To assess how the spatial pattern of high-severity patches influences avian community diversity and composition, we used a two-step multispecies occupancy model analysis along with 2044 bird community surveys conducted within high-severity patches of 27 wildfires. Specifically, we addressed the following questions: (a) How does location within a high-severity patch (i.e., distance from lower-severity areas) and patch size influence occupancy of individual species, as well as avian alpha and beta diversity? (b) How does diversity change 1–30 years after fire and are these changes dependent on distance to lower-severity areas or high-severity patch size? (c) Do these factors influence species richness of functional groups (i.e., nesting guilds) and their relative prevalence in the bird community?

2 | METHODS

2.1 | Study system

Through a combination of lightning ignitions and indigenous management, semi-arid conifer forests of California historically experienced frequent fire dominated by low- to moderate-severity effects (Safford & Stevens, 2017). However, legacies of 19th and 20th century fire and forest management as well as accelerating climate change are causing shifts in fire patterns across the region (Mallek et al., 2013; Safford & Water, 2014). Following Euro-American colonization and genocide of native peoples in the 1800s, traditional fire management ceased and naturally ignited wildfires were suppressed

(Anderson & Keeley, 2018; Stephens et al., 2007; van Wagtenonk et al., 2018). Forest structure was further altered in many areas by 19th and 20th century timber harvest and silvicultural practices (McKelvey & Johnston, 1992; Safford & Stevens, 2017). These changes led to a large increase in tree density, surface fuels and an elevated risk of high-severity fire effects (Lydersen & Collins, 2018; Steel et al., 2015; Stephens et al., 2015). Apart from vegetation pattern and structure, climate change is lengthening fire seasons and increasing instances of extreme fire weather (Westerling, 2016; Williams et al., 2019). Contemporary wildfires that escape suppression efforts are consequently burning with a greater proportion of high-severity effects, larger high-severity patches, and more high-severity area far from patch edges (Miller et al., 2012; Parks & Abatzoglou, 2020; Steel et al., 2018).

We surveyed birds within previously burned conifer forests of the Sierra Nevada and Southern Cascade ranges in California

(Figure 1). Sample locations span an elevation gradient of 1050–2200 m and a latitudinal gradient of 35.8–41.4 degrees, which encompass the lower and upper montane zones of the range. Along this gradient, dominant tree species transition from *Pinus ponderosa*, *P. lambertiana*, *Abies concolor*, *Pseudotsuga menziesii*, *Calocedrus decurrens* and *Quercus* spp in the lower montane to *A. magnifica*, *Pinus jeffreyi* and *Pinus monticola* in the upper montane. Shrubs, including *Arctostaphylos* spp., *Ceanothus* spp., *Prunus* spp., *Q. vaccinifolia*, *Chrysolepis sempervirens* and *Ribes* spp., are also found in conifer stands or as the dominant vegetation in areas characterized by xeric conditions, poor soils or past disturbance (van Wagtenonk et al., 2018). Historically, the lower montane yellow pine and mixed conifer forests were characterized by a 10- to 20-year mean fire return interval with 5%–8% of burned area experiencing high-severity effects on average. At higher elevations, red fir forests burned slightly less frequently (40-year mean fire return interval) and with

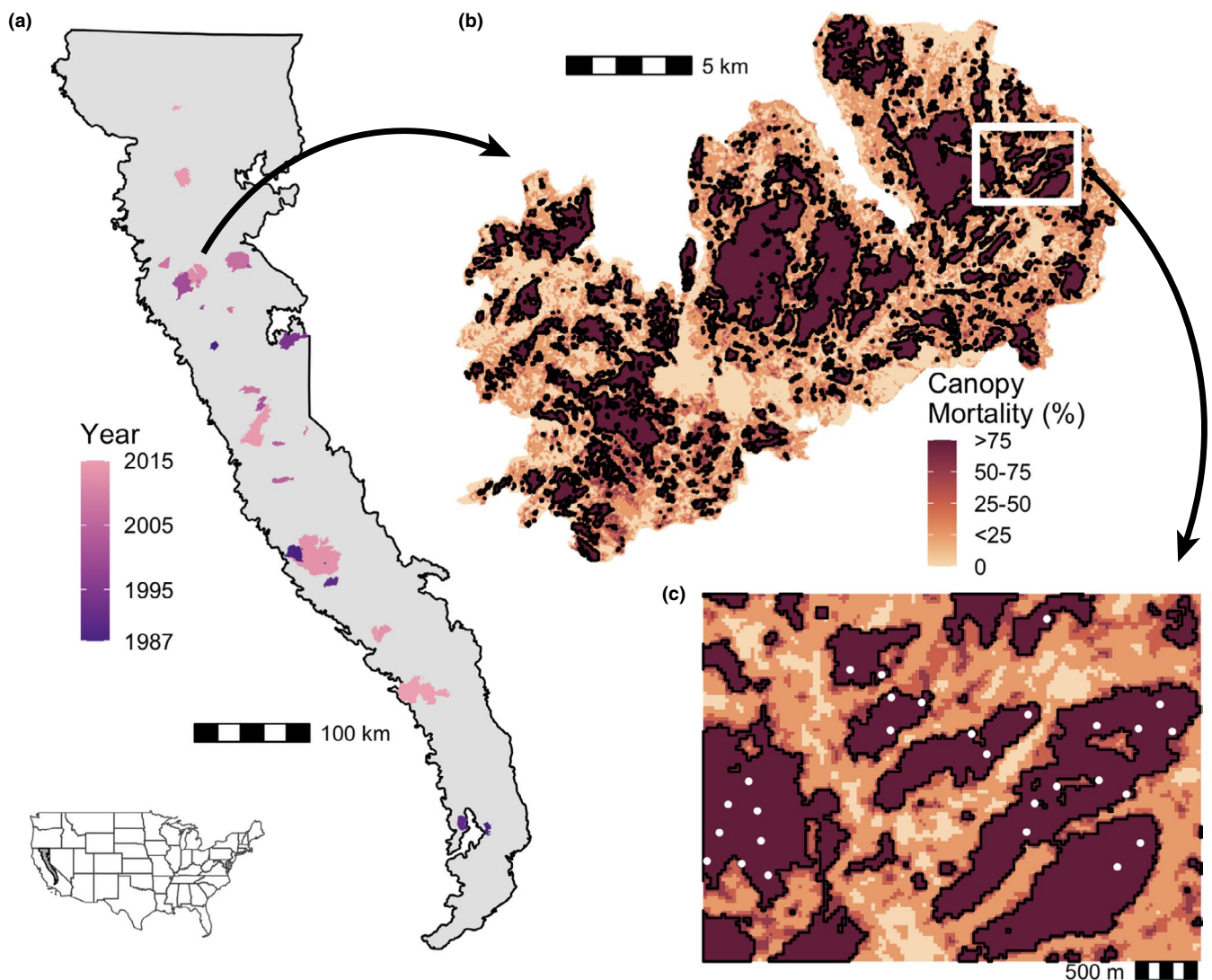


FIGURE 1 Study region map and illustration of sampling design. (a) Footprints of 27 sampled fires that burned across a range of years within California's Southern Cascades and Sierra Nevada bioregions. (b) An example severity map (the 2012 Chips Fire) with high-severity patches of various sizes outlined in black. (c) An example burned area containing bird survey locations (white dots) at various distances from patch edge

greater percentage of high severity (10%–20%; Mallek et al., 2013; Safford & Water, 2014). The historic spatial patterns of fire severity is less certain, but high-severity patches were likely relatively small (mostly <10 ha) in the yellow pine and mixed conifer forests (Safford & Stevens, 2017), and contemporary reference sites show 86% of high-severity patches in red fir and subalpine forests are less than 15 ha in size (Meyer & North, 2019).

2.2 | Environmental and bird data

Fire severity data were obtained from a Forest Service-maintained database, which includes fires that occurred at least partially on Forest Service or National Park Service land in California since 1984 (available at <http://www.fs.usda.gov/detail/r5/landmanagement/gis/>). The database was derived from LANDSAT-TM satellite multi-spectral imagery. The relativized differenced normalized burn ratio was calculated as a measure of vegetation change resulting from each fire and was subsequently used to quantify canopy cover mortality (Miller et al., 2009; Miller & Thode, 2007). Extensive calibration and validation of this database has shown high classification accuracy of high-severity areas with greater classification uncertainty within the matrix of moderate severity (25%–75% mortality; Miller et al., 2009; Miller & Quayle, 2015; Lydersen et al., 2016). We focus on high-severity areas where a minimum of 75% of the pre-fire canopy cover was killed and define high-severity patches as contiguous areas connected by at least one 900 m² pixel (not including diagonals). This mortality threshold is often employed to designate high-severity fire effects and has been used effectively to describe differences in vegetation structure, avian diversity and occurrence of sensitive species (e.g., Jones et al., 2020; Tingley et al., 2016; Welch et al., 2016). However, other thresholds have also been used in the literature to test various ecological questions (e.g., Collins et al., 2017; Lydersen et al., 2016). Patch size and the distance to survey locations within a high-severity patch from areas with <75% pre-fire mortality (i.e., patch edge) were calculated using R (R Core Team, 2020).

We combined avian point count survey data from multiple monitoring programmes in the Sierra Nevada and Southern Cascade Mountains (Figure 1). The majority of surveys (62%) were conducted as part of a post-fire monitoring programme in eight fires of the central and northern part of the study region. To extend the temporal and geographic scope of our analysis, we also drew from a region-wide monitoring programme where survey locations were coincident with areas that previously experienced high-severity fire. The post-fire monitoring programme included sampling stratified across the burn severity gradient, while the region-wide programme used a generalized random tessellation stratified sampling design with a habitat type strata (Burnett & Roberts, 2015; Roberts et al., 2011; Taillie et al., 2018). Only points surveyed following high-severity fire were considered here (Figure 1). Survey sites were spaced a minimum of 250 m apart and were located on US Forest Service lands. A standard 5-min point count survey protocol was used where each individual bird was recorded along with its initial distance from point

centre (Ralph et al., 1993). All observers were experienced avian field technicians that completed an intensive 2-week training and passed a double observer test with experts in bird identification and point count methods prior to conducting surveys. Surveys were conducted during the peak breeding season (May–June), with repeat visits conducted by different observers to reduce the likelihood that variability in observer skill affected occupancy estimates. Sites used in analysis were limited to locations that had not experienced a previous high-severity event since at least 1984 and were classified as conifer forests prior to the most recent fire, and where no major management actions (e.g., salvage logging, herbicide, or reforestation treatments) had occurred since the fire either at the location of the survey or between survey points and the nearest high-severity patch edge. The Forest Service Activity Tracking System database (available at <https://www.fs.usda.gov/main/r5/landmanagement/gis/>) was used to identify post-fire management and was verified with historical Google Earth aerial imagery. In total, 746 point count sites were surveyed in multiple years during the 2009–2017 seasons. Sites were located within one of 130 high-severity patches and 27 fires. Sampled wildfires burned between 1987 and 2015, with surveys conducted 1–30 years post-fire (median = 3 years; mean = 6 years). Sample distributions of distance from edge and patch size were log-normal with distance from lower-severity areas ranging between 1 m and 1,090 m (median = 81 m; mean = 125 m) and high-severity patch size ranging between 0.1 and 21,000 ha (median = 42 ha; mean = 680 ha). Stand structure data were also collected at point count locations during some years using a modified relevé protocol (Ralph et al., 1993) and used for modelling variation in bird detectability. For years where birds were surveyed but not vegetation, the most proximate (before or after) vegetation data were used.

2.3 | Statistical analysis

Of the 122 species observed during at least one survey occasion, we restricted our analysis to 94 species that are effectively surveyed using the point count protocol and are known to breed within the study area (see Appendix S1 in Supporting Information). Species were classified into nesting guilds according to known life histories as breeding bird territories and occurrence are tightly linked to nesting sites (Beedy & Pandolfino, 2013; Steel et al., 2012). Specifically, each species was classified as predominantly a tree, primary cavity, secondary cavity, shrub or ground nester (Appendix S1). We consider each survey season and site combination as a primary survey occasion (*i*) and within season visits as temporal replicates (*j*). The community is assumed to be closed during each season, allowing for formal estimation of rates of detection and occupancy. On average, each site was surveyed for 2.7 years, for a total of 2044 primary survey occasions; most (90%) composed of two visits per year. Observation data are binary with $y_{s,ij} = 1$, when species *s* is detected during survey *i* and visit *j* within 100 m of the observer, or $y_{s,ij} = 0$, when a species was not detected. Because detection is imperfect, we used an occupancy modelling framework $y_{s,ij} = \text{Bernoulli}(p_{s,ij} * z_{s,i})$

where $p_{s,i,j}$ is the probability of detection given a site is occupied (i.e., $z_{s,i} = 1$). A species' true occurrence is modelled as $z_{s,i} = \text{Bernoulli}(\psi_{s,i})$ where $\psi_{s,i}$ is the probability of occurrence and $z_{s,i}$ is a binary latent variable of a species' true occurrence state.

We assumed detection and occurrence probabilities vary by species and survey and are functions of habitat and survey characteristics. The detection process is modelled using the following logit-linear function:

$$\begin{aligned} \text{logit}(p_{s,i,j}) = & \alpha 0_s \\ & + \alpha_{\text{day},s} * X1_{i,j} + \alpha_{\text{day}^2,s} * X2_{i,j} \\ & + \alpha_{\text{snag},s} * X3_i + \alpha_{\text{shrub},s} * X4_i \end{aligned} \quad (1)$$

where the Julian day (α_{day} and α_{day^2}) of each visit, snag basal area (α_{snag}), and percentage shrub cover (α_{shrub}) are predictors of detection probability. The occurrence process is modelled using the following logit-linear function:

$$\begin{aligned} \text{logit}(\psi_{s,i}) = & \beta 0_s + \beta_{\text{site}}[i]_s + \beta_{\text{patch}}[i]_s + \beta_{\text{fire}}[i]_s \\ & + \beta_{\text{YSF},s} * X5_i + \beta_{\text{YSF}^2,s} * X6_i \\ & + \beta_{\text{dist},s} * X7_i + \beta_{\text{dist*YSF},s} * X7_i * X5_i \\ & + \beta_{\text{area},s} * X8_i + \beta_{\text{area*YSF},s} * X8_i * X5_i \\ & + \beta_{\text{lat},s} * X9_i + \beta_{\text{elev},s} * X10_i + \beta_{\text{elev}^2,s} * X11_i \end{aligned} \quad (2)$$

where β_{site} , β_{patch} , and β_{fire} are varying intercepts for site, patch and fire ID for species s . For each species, occupancy probability is a function of number of years since fire (β_{YSF} and β_{YSF^2}), distance (log m; β_{dist}) from high-severity patch edge, the interaction of distance and years since fire ($\beta_{\text{dist*YSF}}$), patch area (log ha; β_{area}), the interaction of patch area and years since fire ($\beta_{\text{area*YSF}}$), latitude (degrees; β_{lat}), and elevation (m; β_{elev} and β_{elev^2}). Quadratic terms for distance and patch area were also considered, but early models showed little justification for their inclusion. Variable correlations were low ($< |0.6|$) among the detection or occupancy model.

The primary predictor variables of interest are distance from patch edge, years since fire, and high-severity patch size. The additional predictors in Equations 1 and 2 were chosen based on previous work assessing species occurrence in the Sierra Nevada Mountains (e.g., Burnett & Roberts, 2015; Siegel et al., 2011; Tingley et al., 2016; White et al., 2016). We utilize a multispecies occupancy model where each species-specific parameter α_s or β_s is modelled as a random sample from a community-level hyperparameter. This allows for more precise estimates for rare species, which can be challenging to model using single-species approaches, and improves estimates of community metrics with appropriate propagation of model uncertainty (Zipkin et al., 2010). We allow intercepts of ψ_s to vary by fire and patch ID to help account for spatial clustering of points and for cluster-specific environmental variation not explicitly included in the model (e.g., fire size). Similarly, site ID is modelled as a varying intercept to account for variation in unmodelled habitat characteristics and other local influences unique to each point. This approach may be considered a correlated multispecies model as it makes explicit the assumption that occupancy at a site is not

fully independent among years. This model differs from a dynamic multispecies model (Dorazio et al., 2010) in that correlations are bi-directional (e.g., information in survey year two informs estimates of year one and vice versa) and does not estimate rates of local extinction and colonization, which are not of interest here. All continuous predictors were standardized with a mean of zero and standard deviation of one, and varying intercepts are estimated with a group population mean of zero. The multispecies occupancy model was estimated using Hamiltonian Monte Carlo sampling in Stan via the rstan package (Stan Development Team., 2020). We specified weakly regularizing priors to prevent model over-fitting (McElreath, 2016).

To estimate alpha and beta diversity at the community level, we used a two-step (aka Bayesian meta-analysis) modelling process (Kery & Royle, 2015; Tingley & Beissinger, 2013) where the detection-corrected estimates of occupancy ($z_{s,i}$) from the multispecies occupancy model were used to calculate diversity metrics for each primary survey occasion. Richness (alpha diversity) is calculated as the sum of $z_{s,i}$ for the full community or for a subset of species when conducting nesting guild analysis. Beta diversity is calculated as the compositional dissimilarity from the group centroid (multidimensional median), where groups are defined as each of the 27 sampled fire areas. We limit our assessment of alpha diversity to species richness because metrics such as evenness and Shannon diversity require estimates of species abundance which cannot be produced by occupancy models and the presence/absence data used. We use the Raup–Crick Index (Raup & Crick, 1979) as our measure of beta diversity because it adjusts for differences in species richness by evaluating dissimilarity between each survey and randomly generated communities based on the group's species pool (Chase et al., 2011). Alpha- and beta-diversity calculations were conducted for each draw of the model posterior distribution creating derived distributions for each metric and each primary survey occasion. By using the full multispecies model posterior distribution, model uncertainty is properly propagated to the community models (Kery & Royle, 2015; Tingley & Beissinger, 2013). Community models were fit using the BRMS and rstan packages (Bürkner, 2017; Stan Development Team., 2020) with the same set of predictors as equation 2 as well as a gaussian error structure and log-link. All models were run with 4 chains, each for 1,000 samples with a warmup of 500, resulting in 2,000 post-warmup samples. Proper mixing of sampling chains was checked using traceplots, and all parameters received R-hat values of less than 1.1. Model code can be found in Appendix S2 with additional supporting code found at https://github.com/zacksteel/High_Severity_Birds.

To quantify the interaction of distance to patch edge and years since fire, we calculated marginal effects of years since fire at three distance levels (10, 50, and 250 m). We limit projected effects to 250 m because surveys conducted at greater distances from patch edge were limited to the first two decades since fire. The marginal effect of patch size and its interaction with years since fire was similarly quantified using projected effects for 1, 50, and 2,500 ha patches. The use of a linear and quadratic effect of years since fire assumes a parabolic form with an average peak or nadir at some

vertex year following disturbance. We solved for the location of the vertex of each richness model by calculating the expected year since fire when the derivative of the parabola is equal to zero (i.e., when the rate of change is zero; $-\beta_{YSF}/2*\beta_{YSF}^2$). To evaluate the degree to which distance or patch size influences changes in species richness over time, we calculated the expected difference between year one and the vertex year with respect to the three distance and patch size levels. Probabilistic statements as well as derived model estimates and credible intervals were calculated using model posterior distributions. For example, the probability an effect was positive (or negative) was calculated as the proportion of the parameter posterior distribution above (or below) zero.

3 | RESULTS

3.1 | Species-level effects

Individual bird species differed greatly in how occupancy was affected by the environmental variables tested (Figure 2 & Appendix S3). Among the primary variables of interest, years since fire showed the widest range of standardized effects, with 26% of species showing clear positive (probability [Pr.] >95%) and 21% showing clear negative associations with occupancy. 15%

of species showed a concave (positive quadratic effect) and 27% showed a convex (negative quadratic effect) functional relationship between occupancy and years since fire. Occupancy of 13% of species showed clear positive associations with internal distance from patch edge, while 27% showed a clear negative relationship. The species with the strongest positive effect of distance from edge included European Starling (*Sturnus vulgaris*), Bullock's Oriole (*Icterus bullockii*), Sooty Grouse (*Dendragapus fuliginosus*), and Lewis' Woodpecker (*Melanerpes lewis*), while those with the strongest negative effect included Red-breasted Nuthatch (*Sitta canadensis*), Hermit Warbler (*Setophaga occidentalis*), Mountain Chickadee (*Parus gambeli*), and Brown Creeper (*Certhia americana*). Occupancy of 7% of species showed either a positive or a negative interaction between years since fire and distance, while 3% of species showed a clear interaction between year since fire and patch size, indicating that the rate of change following fire varies by distance from edge more often than patch size. Occupancy of only 1% of species (i.e., Lazuli Bunting; *Passerina amoena*) showed a positive association with high-severity patch size, while 9% of species occurred more often in smaller patches (Figure 2). Those species with the strongest negative effect of patch size included Red-breasted Nuthatch, Western Wood-pewee (*Contopus sordidulus*), Hutton's Vireo (*Vireo huttoni*), and Mountain Chickadee. Estimates of all multispecies model parameters can be found in Appendix S3.

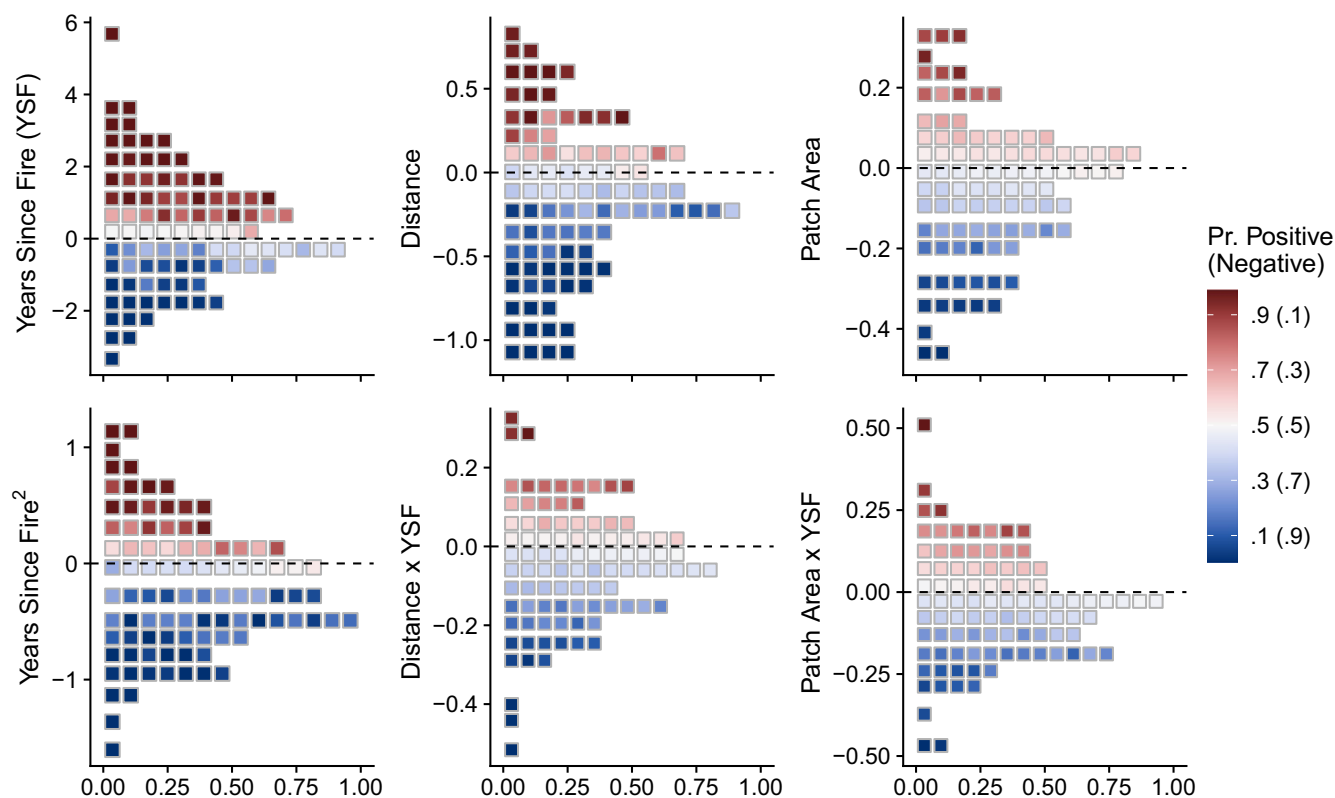


FIGURE 2 Coefficient estimates for avian multispecies occupancy model parameters of interest. Each square represents a single species with its location on the y-axis indicating the mean parameter estimate and colour indicating probability of a positive or conversely negative (1 – Pr. Positive) effect. The x-axis indicates the proportion of species in a given effect size bin relative to the bin with the maximum count. Estimates are for bird species of the Sierra Nevada, California

3.2 | Community richness and dissimilarity

The community richness model estimated very clear ($Pr. \geq 99\%$) effects of all parameters describing internal distance from patch edge, years since fire, and patch area (Figure 3a). Richness was positively associated with years since fire with a negative quadratic term indicating a convex (humped-shaped) relationship with time. Richness was negatively associated with internal distance from high-severity patch edge and to a lesser degree patch size. The interaction of years since fire and distance from patch edge was negative indicating that the positive effect of years since fire diminished with distance and conversely that the negative effect of distance grew stronger with time. The interaction of patch size and years since fire was similarly negative but with greater model uncertainty ($Pr. 90\%$). Richness declined with latitude and likely increased with elevation but with greater uncertainty ($Pr. = 94\%$; Figure 3a). The combined effects of years since fire, distance from patch edge, and patch size resulted in the most speciose communities occurring in the later years of our sample period, near patch edges, and within relatively small patches (Figure 4). After accounting for differences in species richness, community dissimilarity was negatively associated with the linear elevation term and positively with the quadratic elevation term ($Pr. >99\%$), indicating a concave (U-shaped) relationship with points located at low and high elevations most different from the average community (Figure 3b). Community dissimilarity was also negatively associated with the quadratic term of years since fire and

likely positively related to linear term of years since fire ($Pr. = 93\%$; Figures 3b and 4). Dissimilarity was positively associated with the interaction of patch area and years since fire, resulting in a delayed peak in beta diversity within large patches (Figure 4). The direction of all other effects in the community dissimilarity model was less certain ($Pr. <90\%$; Figure 3b).

3.3 | Nesting guilds

The direction and magnitude of model effects varied considerably among nesting guilds (Figure 3c). Tree and primary cavity nester richness was very clearly negatively associated with years since fire and positively associated with years since fire quadratic term resulting in concave (U-shaped) relationships over time. Conversely, secondary cavity, shrub, and ground nester richness was positively associated with years since fire and negatively associated with the years since fire quadratic term resulting in a convex (hump-shaped) relationship over time (Figures 3c, 5 and 6). Distance from patch edge had a negative effect on richness for all guilds with the exception of shrub nesters; the magnitude of the effect was strongest for tree nesters, followed by primary cavity nesters. The interaction of distance and years since fire had a negative effect on richness of for tree, primary cavity, and secondary cavity nesters, while positive for ground nesters (Figures 3c and 5). High-severity patch area was negatively associated with richness of tree and primary cavity nesters, and the

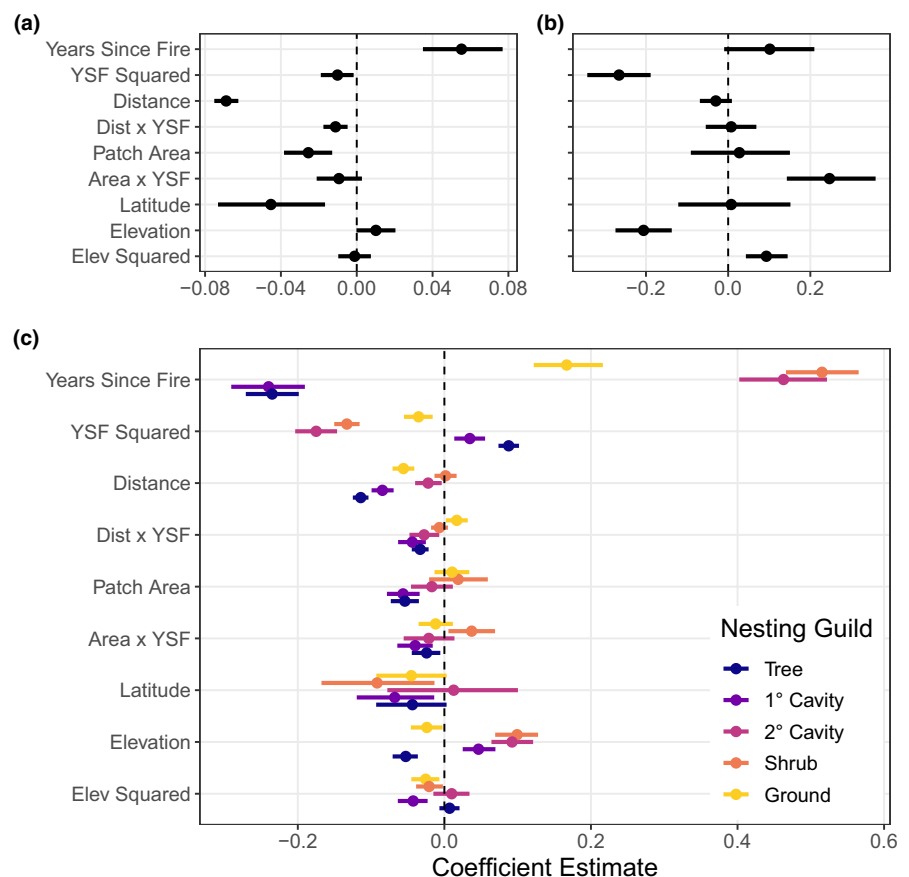


FIGURE 3 Coefficient dotplots for avian (a) community richness, (b) community dissimilarity, and (c) guild richness in the Sierra Nevada, CA. Points represent median parameter estimates and bars represent 90% credible intervals. Tabulated parameter estimates can be found in Appendix S4

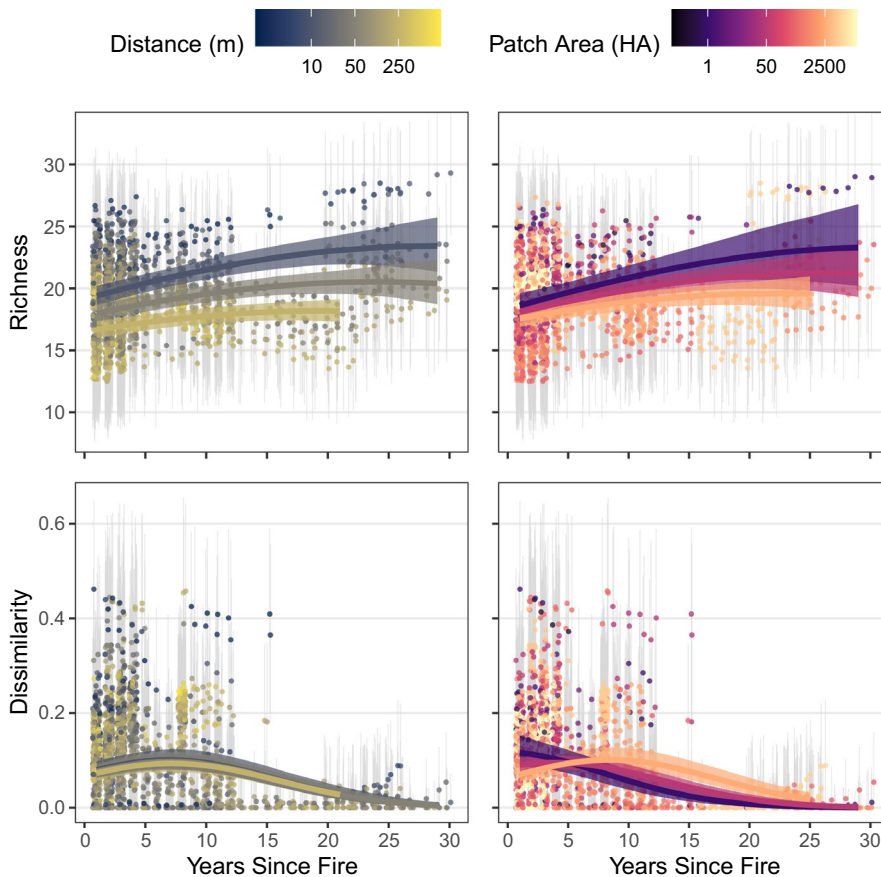


FIGURE 4 Marginal effects of years since fire, distance from high-severity patch edge, and patch size on bird community alpha and beta diversity in the Sierra Nevada, CA. Alpha diversity is quantified as species richness. Beta diversity is quantified using the Raup–Crick Index, where groups (species pools) are defined by each individual fire area. Model fits are shown as median lines and 90% credible interval bands for 10 (blue), 50 (grey), and 250 m (yellow) from patch edge or 1 (purple), 50 (pink), and 2,500 ha (orange) patches with other predictor variables held at their mean values. After accounting for imperfect detection, estimated medians and 90% credible intervals of sampled communities are shown as jittered points and vertical bars

interaction between patch area and years since fire was positive for shrub nester richness (Figures 3c and 6). Latitude was negatively associated with richness of primary cavity and shrub nesters and likely negatively with ground and tree nester richness (Pr. 94%). Elevation was positively associated with richness of primary cavity, secondary cavity, and shrub nesters and negatively with tree and ground nesters. The quadratic term of elevation was negative for primary cavity, ground, and shrub nesters (Figure 3c). While guild-level patterns were often apparent, there were within-guild exceptions. For example, in contrast to the guild overall Hairy (*Dryobates villosus*), Lewis', and Acorn (*M. formicivorus*) woodpeckers occurred more often with increasing distance from patch edge, while the Black-backed woodpecker (*Picoides arcticus*; an important species for post-fire management) showed a neutral response (Appendix S3).

3.4 | Distance- and patch size-dependent avian succession

Distance from patch edge, and to a lesser degree patch size, influenced the rate at which species richness changed following wildfire (Figure 3c). In addition to greater mean richness, patch edges also experienced greater growth in community richness with time since fire as compared to patch interiors (Figure 5f). Conversely, the declines in richness experienced by the tree and primary cavity nesting guilds

were more pronounced in patch interiors than edges. Similar to the overall community, between year one and the year of peak richness, secondary cavity nester richness increased more along edges than interiors. Temporal changes in ground and shrub guild richness were largely unaffected by distance from edge (Figure 5f). Larger patches also resulted in more rapid declines in tree and primary cavity nester richness. However, where temporal changes in shrub nester richness were unaffected by distance from edge, the guild peaked at higher levels within large high-severity patches (Figure 6f).

The strength of the interactions of distance or patch size and years since fire ultimately influenced the relative composition of communities along edges and interiors. When guilds reached their richness peak or nadir approximately two decades following a fire, tree and primary cavity nesters represented greater percentages of the full community along edges than interiors, while the ground and shrub guilds saw the opposite pattern. Specifically, the relative number of tree nesters decreased from a median of 28% of the community at 10 m from edge to 24% at 250 m from edge. Similarly, the relative number of primary cavity nesters decreased from 12% to 9%. While absolute ground and shrub nester richness was insensitive to distance, the loss in forest-dependent species resulted in relative increases for these groups. The median percentage of the community composed of ground nesters increased from 18% at 10 m from edge to 22% at 250 m from edge, while the relative number of shrub nesters increased from 22% to 28%. Secondary cavity

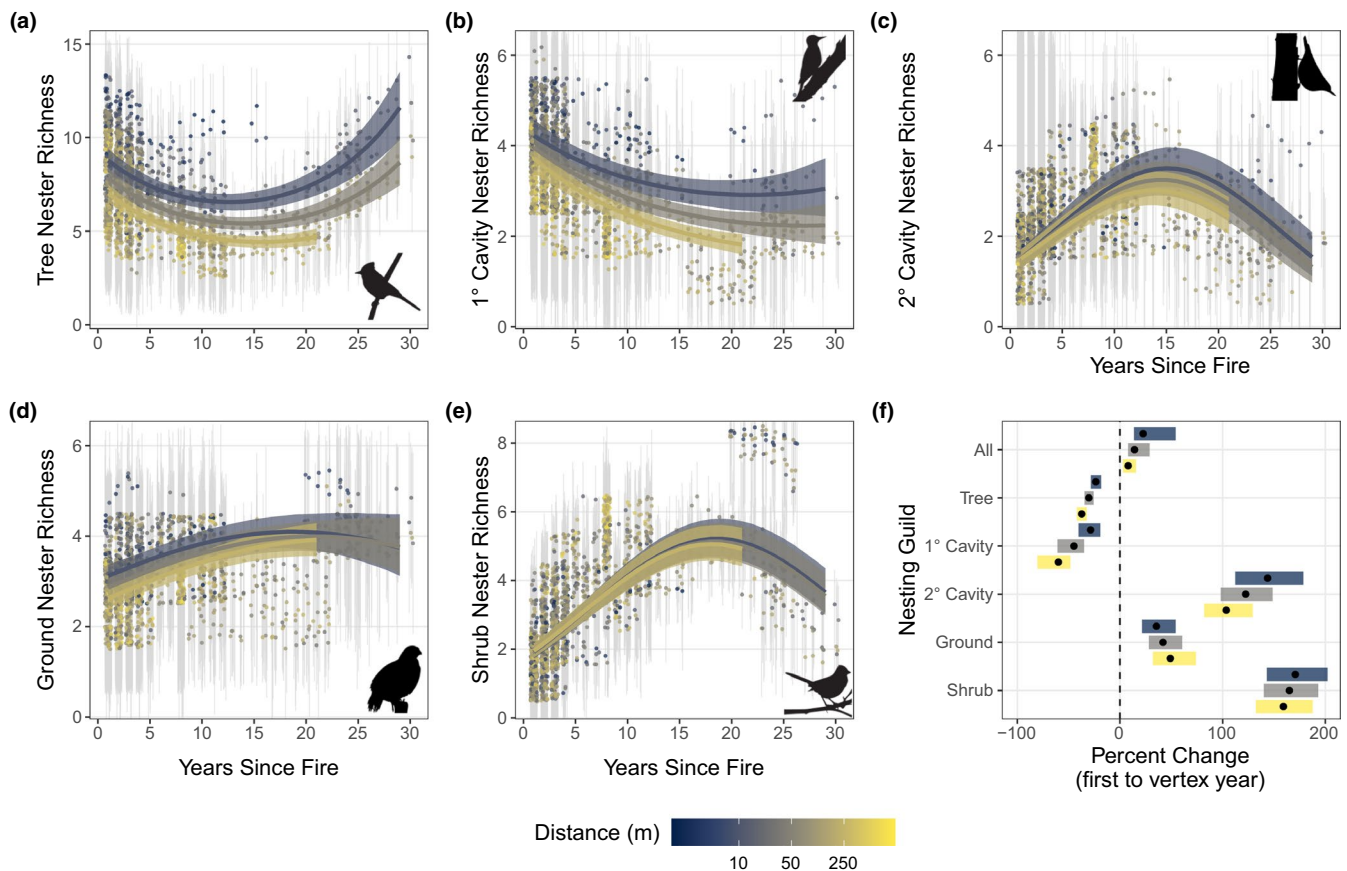


FIGURE 5 Marginal effects of years since fire and internal distance from high-severity patch edge on avian guild richness in the Sierra Nevada, CA. Panels represent (a) tree, (b) primary cavity, (c) secondary cavity, (d) ground, and (e) shrub nester richness. (f) shows the predicted per cent change in richness from 1 year since fire to the median vertex (peak or nadir) year for an average point characterized by a given distance from patch edge. For example, shrub nester richness at an average point 250 m from patch edge is expected to maximize at a mean of 5.1 species in year 18, a 159% increase from year one. Model fits are shown as median lines and 90% credible interval bands for 10 (blue), 50 (grey), and 250 m (yellow) from patch edge with other predictor variables held at their mean values. After accounting for imperfect detection, estimated medians and 90% credible intervals of sampled communities are shown as jittered points and vertical bars. Bird silhouettes from www.allaboutbirds.org, © Cornell Lab of Ornithology

nesters maintained approximately the same percentage of the community regardless of distance from edge.

Patch size had a similar effect on community composition with tree and primary cavity nesters making up a smaller component of the community on average in large patches as compared to small patches and ground and shrub species making up a relatively greater component. Due to smaller effect sizes (Figure 3c), the realized compositional shifts were weaker for most guilds, except for the shrub nesters. At its peak richness, the shrub nester group made up approximately 18% of the community at an average point in small patches (1 ha) and 27% of the community in large patches (2,500 ha). These ultimate effects of distance from patch edge and patch size on community composition were weaker in the years immediately following a fire and likely continued or strengthened into the third decade since fire. However, our sampling of patch interiors (e.g., >250 m) and very large patches (e.g., >2,500 ha) beyond 20 years since fire was limited, making projections into the third decade less certain (Figures 5 and 6).

4 | DISCUSSION

As fire activity increases across the western United States and large patches of high-severity effects become more common, it is important to understand how bird communities will respond in the decades following disturbance. Here, we show that as compared to small high-severity patches or patch edges, interiors of large patches support fewer species and a lower rate of increase in species richness within the first three decades following wildfire. Further, this lower richness is not compensated by an increase in compositional dissimilarity, indicating that high-severity patch interiors contain a subset of those species found at patch edges, rather than species unique to patch interiors. These differences within and among high-severity patches are largely attributable to tree and primary cavity nesting species (e.g., Hermit Warbler and Red-breasted Nuthatch), which were more common in small patches and along patch edges. Neither ground nor shrub nesting guilds had higher species richness in patch interiors despite the greater cover of early-seral habitat in the local

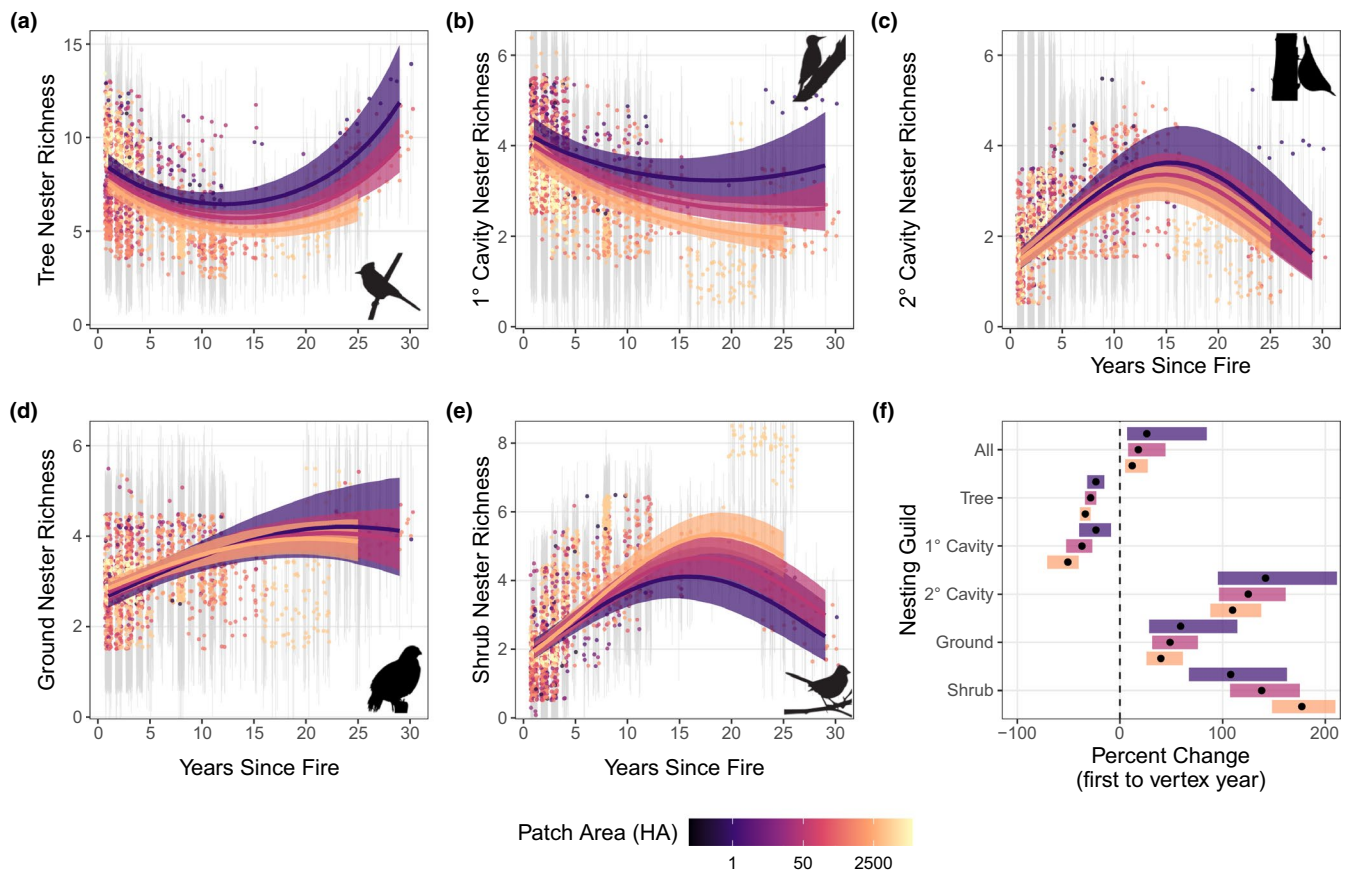


FIGURE 6 Marginal effects of years since fire and high-severity patch size on avian guild richness in the Sierra Nevada, CA. Panels represent (a) tree, (b) primary cavity, (c) secondary cavity, (d) ground, and (e) shrub nester richness. (f) shows the predicted per cent change in richness from one year since fire to the median vertex (peak or nadir) year for an average point within a given patch size. For example, shrub nester richness at an average point within a 2,500 ha patch is expected to maximize at a mean of 5.3 species in year 19, a 177% increase from year one. Model fits are shown as median lines and 90% credible interval bands for 1 (purple), 50 (pink), and 2,500 ha (orange) patches with other predictor variables held at their mean values. After accounting for imperfect detection, estimated medians and 90% credible intervals of sampled communities are shown as jittered points and vertical bars. Bird silhouettes from www.allaboutbirds.org, © Cornell Lab of Ornithology

landscape. However, differential responses among functional groups to high-severity spatial pattern resulted in relative shifts in community composition where ground and especially shrub nesting species made up an increasing percentage of interior communities within large patches with greater time since fire, while tree and primary cavity nesters experienced declines. While distance from edge generally showed stronger effects on bird diversity than patch size, it is important to highlight that these features are functionally related to each other in that for a given patch geometry (e.g., a circle) larger patches have more area far from edges and residual forests. Taken together, these results suggest that small high-severity patches or large patches with high edge-area ratios support higher levels of bird diversity than large high-severity patches containing extensive interior habitat.

These findings further our understanding of how montane bird communities respond to the spatial pattern of high-severity wildfire but also highlight areas of future research. Species occupancy is generally correlated with abundance (Kery & Royle, 2015), but the latter along with fecundity and survival data can provide a more

nuanced picture of how individual species respond to their environment (e.g., Stillman et al., 2021). Further, our data were limited to 30 years since fire with relatively few surveys in the oldest fires. While this remains an exceptionally wide range of time since fire relative to existing research, more extensive sampling of older fires would allow for a more complete understanding of the relationship between fire patterns, vegetation succession, and bird diversity. We also limit our assessment of patch dynamics to one local-level metric (distance from edge) and one patch-level metric (size). Assessments of other landscape metrics and the broader landscape context (e.g., how diversity is affected by distance from the burn perimeter) may complement this work and our understanding of post-fire habitats. Further investigation into the mechanisms of elevated avian richness at high-severity edges could also better inform conservation efforts. Due to the limitations of the remotely sensed fire severity data, it was not currently possible to confidently classify soft (gradual) versus hard (abrupt) patch edges, which may influence the nature of patch dynamic effects on the avian community. More broadly, this work can provide greater context for a growing literature assessing

the influence of disturbance on metacommunity dynamics, including how the relative importance of dispersal and competition in structuring metacommunities may vary with disturbance severity, successional stage, and patch size (Holyoak et al., 2020; Vanschoenwinkel et al., 2013). As burn severity data accumulate (availability starting in 1984) and recent large wildfires age, future studies can further elucidate the relationship between high-severity patch dynamics and long-term avian succession.

4.1 | High-severity patches and pyrodiversity

Pyrodiversity, or the variation in a landscape's history of fire severity, patch sizes, frequency, and seasonality, likely plays an important role in maintaining biodiversity in fire-adapted ecosystems (Jones & Tingley, In Press; Martin & Sapsis, 1992; Steel et al., In Press; Tingley et al., 2016; In Press). Local landscapes surrounding edges of high-severity patches contain greater variation in fire severity than patch interiors due to their proximity to lower-severity areas. Thus, our finding that edges of high-severity patches support greater species richness is consistent with past research showing many western bird species are tolerant or even benefit from creation of habitat edges (McGarigal & McComb, 1995; Sisk & Battin, 2002) and that variation in burn severity is positively associated with bird diversity (Tingley et al., 2016). Diversity of other taxa in the Sierra Nevada including bats (Steel et al., 2019) and plant-pollinator communities (Ponisio et al., 2016) also increases with variation in burn severity suggesting that pyrodiversity likely promotes biodiversity generally. Although a forested landscape composed of a variety of patch sizes of post-fire habitats consisting of multiple severity classes may promote biodiversity, many of the large patches assessed in this study far exceeded the estimated natural range of variation, where the majority of high-severity patches were likely less than 10 ha in size (Safford & Stevens, 2017). Thus, these large patches may not be optimal for promoting biodiversity and instead may perpetuate landscape homogeneity. Our results suggest that maintaining avian diversity in these burned landscapes can be achieved without the largest high-severity patches, but rather with smaller high-severity patches interspersed with a matrix of low- to moderate-severity fire. The availability of habitat features such as snags and montane chaparral necessary for early-seral birds may be sufficient in much smaller high-severity patches than the largest patches studied here. The relatively strong influence of distance from patch edge on community richness suggests that shape complexity (i.e., the amount of edge per unit area) may also be an important consideration as large patches may accommodate high bird diversity if most of its area is near less disturbed habitats. Additionally, edges of high-severity patches appear to accommodate individual bird species of conservation concern with otherwise contrasting habitat needs. While we found little effect of distance from patch edge on occupancy of breeding Black-backed Woodpeckers, other studies have found juveniles of the snag specialist preferred patch edges to interiors (Stillman et al., 2019, 2021) and nest site suitability was lower towards the centre of large

patches (Campos et al., 2020). On the other hand, the mature forest specialist Spotted Owl (*Strix occidentalis*) selects relatively small high-severity patches (i.e., <115 ha) and uses patch edges but avoids large patches and interior areas beyond approximately 100 m (Jones et al., 2020; Kramer et al., 2020).

The temporal components of pyrodiversity have been less explicitly studied but our findings that individual species and guild richness peak at different years reinforce previous work showing a variety of fire ages within a landscape promote overall bird diversity (Smucker et al., 2005; Taillie et al., 2018). Both absolute richness and the post-fire rate of change were dependent on distance from patch edge and patch size, perhaps reflecting that proximity to residual live trees and greater natural regeneration of conifers near patch edges (Greene & Johnson, 1995; Welch et al., 2016) might be important for the bird community in this region. We suspect that the relatively high richness of tree nesters soon after fire is indicative of high site fidelity for some species as well as delayed tree mortality providing temporary residual habitat (Hood et al., 2007). Primary cavity nesters also showed greater richness soon after fire and a decline with years since fire, likely reflecting woodpecker food resources associated with recently killed trees and subsequent losses of snag habitat as fire-killed trees decay and fall (Raphael et al., 1987; Ritchie et al., 2013; Smucker et al., 2005; White et al., 2016). This decline was less pronounced along edges where woodpeckers can switch to using residual live trees (Dudley & Saab, 2007). This reduced richness, especially in primary cavity nesting species, may have cascading effects on biodiversity in high-severity patches (Tarbill et al., 2015). Richness peaked at later times since fire for the other three nesting guilds. Slightly more secondary cavity nesters were found near patch edges and mean guild richness along edges is expected to reach a higher peak as compared to patch interiors. This response to distance from edge is similar to primary cavity nesters, but secondary cavity nesters showed a later peak and subsequent decline in richness reflecting their dependence on woodpeckers to create nesting habitat and less of a dependence on snag-associated prey species (Saab et al., 2004). Distance from patch edge had little influence on the succession of either ground or shrub guilds, but shrub nesters reached greater and later peak richness within large patches. Overall, high-severity patches of a range of ages contributed to montane bird diversity with edges producing greater species richness across the gradient of years since fire.

4.2 | Changing fire regimes and outlook for the future

Montane bird communities face opposing but interconnected changes to habitats shaped by fire or its absence. A century of fire exclusion in western forests has led to over-densification and in some areas a loss or degradation of early-seral habitats (Betts et al., 2010). Due to these changes and factors associated with climate change (Parks & Abatzoglou, 2020; Westerling, 2016), contemporary fires are becoming more severe with larger patches of

high-severity effects. As fires continue to grow larger and more severe, we are likely to see directional shifts from conifer forests to montane chaparral or forests dominated by resprouting oaks (Coop et al., 2020; Coppoletta et al., 2016; Safford & Vallejo, 2019; Tepley et al., 2017). Such shifts would have mixed consequences for the montane bird community. Increases in chaparral habitat will benefit shrub-associated bird species, but when conversions are primarily concentrated in large patches with proportionally less edge, overall diversity is likely to decline at least within the first 30 years after fire. On the other hand, yellow pine and mixed conifer forests of the Sierra Nevada have experienced a massive reduction in acres burned since the onset of modern fire suppression with a large deficit in low- to moderate-severity fire (Mallek et al., 2013; Stephens et al., 2007). While modern wildfires contain increasing proportions of high-severity effects (Steel et al., 2018), they often also include extensive areas of low- to moderate-severity which can promote forest resilience (Collins et al., 2018; Kane et al., 2019; Steel et al., In Review). Thus, increasing burn activity promises both positive and negative impacts with interiors of large high-severity patches representing degraded ecosystem resilience and biodiversity, while areas of more moderate and heterogeneous fire effects supporting partially restored ecological communities.

4.3 | Management implications

High-severity patches, which represent the greatest ecosystem change within a wildfire footprint, are also frequently the targets of post-fire management. To reforest a burned area quickly, practices include removing snags, controlling shrubs mechanically or chemically, and planting desired tree species (Lindenmayer et al., 2012; North et al., 2019). These management actions themselves impact the local bird community, particularly species that rely on snags, shrubs and herbaceous vegetation as habitat (Lindenmayer et al., 2012; Saab et al., 2009). For example, salvage logging can negatively affect primary cavity nesters like the Black-backed woodpecker while shrub abatement can have similarly negative consequences for shrub-associated species (Cahall & Hayes, 2009; Easton & Martin, 2002; Saab et al., 2009). In the longer term, reforestation may eventually benefit late-seral species by accelerating forest succession, whereas high-severity wildfire without intensive post-fire management may benefit early-seral species for decades. When attempting to balance multiple objectives with limited funds, reforestation on public lands may be conducted in some areas with natural succession left to proceed in others (North et al., 2019). These trade-offs are accentuated in modern wildfires, which are progressively more likely to be large and severe with increasingly large and simply shaped (low edge:area) high-severity patches (Miller et al., 2012; Steel et al., 2018; Westerling, 2016). If reforestation efforts occur, managers concerned with avian diversity can prioritize interiors of large high-severity patches where average bird diversity is lowest and where natural conifer reforestation is least likely to succeed (Shive et al., 2018; Welch et al., 2016). In contrast, avoiding intensive

management along patch edges and within high-severity patches of sizes that fall within the natural range of variation can retain pyrodiversity and diversity of the broader avian community. Further, prescribed fire and managed wildfire that increase pyrodiversity in both previously burned and fire suppressed forests can help restore fire-adapted habitats and promote bird diversity as the avian community responds to accelerating global change.

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BIOSKETCH

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

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