




ORIGINAL RESEARCH

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# Patterns of woodboring beetle activity following fires and bark beetle outbreaks in montane forests of California, USA

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## Abstract

**Background:** Increasingly frequent and severe drought in the western United States has contributed to more frequent and severe wildfires, longer fire seasons, and more frequent bark beetle outbreaks that kill large numbers of trees. Climate change is expected to perpetuate these trends, especially in montane ecosystems, calling for improved strategies for managing Western forests and conserving the wildlife that they support. Woodboring beetles (e.g., Buprestidae and Cerambycidae) colonize dead and weakened trees and speed succession of habitats altered by fire or bark beetles, while serving as prey for some early-seral habitat specialists, including several woodpecker species. To understand how these ecologically important beetles respond to different sources of tree mortality, we sampled woodborers in 16 sites affected by wildfire or bark beetle outbreak in the previous one to eight years. Study sites were located in the Sierra Nevada, Modoc Plateau, Warner Mountains, and southern Cascades of California, USA. We used generalized linear mixed models to evaluate hypotheses concerning the response of woodboring beetles to disturbance type, severity, and timing; forest stand composition and structure; and tree characteristics.

**Results:** Woodborer activity was often similar in burned and bark beetle outbreak sites, tempered by localized responses to bark beetle activity, burn severity, tree characteristics, and apparent response to ignition date. Larval woodborer activity was inversely related to bark beetle sign within a sampling quadrat, was higher on pines, and—in burned sites—was higher on the south-facing sides of smaller trees. Adults—especially buprestids—were more abundant where burn severity was higher. Fires with intermediate ignition dates during the study period supported higher rates of larval woodborer activity and higher numbers of adult buprestids as well as cerambycids in the genus *Monochamus* Dejean 1821.

**Conclusions:** Woodboring beetle abundance was related to bark beetle activity, burn severity, tree characteristics, and ignition date. Considering these patterns when managing disturbed forest stands could yield improved outcomes for wildlife, including species that prey on woodboring beetles. We also reported preliminary evidence that the current trend toward more frequent wildfires might not stimulate larger woodboring beetle populations if those fires increasingly occur outside the historical fire season.

**Keywords:** Buprestidae, burn severity, Cerambycidae, fire season, forest management, Scolytinae, snags, succession, wildfire, woodborers

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## Resumen

**Antecedentes:** El aumento frecuente y severo de sequías en el oeste de los EEUU ha contribuido a incrementar la frecuencia y severidad de los incendios forestales, el alargamiento de la temporada de estos y el incremento de brotes de escarabajos descortezadores que matan un gran número de árboles. Se espera que el cambio climático perpetúe estas tendencias, especialmente en ecosistemas montañosos, lo que llama a mejorar las estrategias para el manejo de los bosques del oeste de los EEUU y conservar la vida silvestre que estos sostienen. Los escarabajos barrenadores (*i.e.*, Buprestidae y Cerambycidae) colonizan árboles muertos y débiles y aceleran la sucesión secundaria en hábitats alterados por fuego o por escarabajos descortezadores; al mismo tiempo, sirven de presa para algunos especialistas que actúan durante las etapas tempranas de la sucesión, incluyendo diferentes especies de pájaros carpinteros. Para entender ecológicamente cómo los escarabajos barrenadores responden a las diferentes causas de mortalidad de árboles, estos fueron muestreados en 16 sitios afectados por incendios o por brotes de escarabajos descortezadores ocurridos de uno a ocho años antes. Los sitios de estudio se ubicaron en la Sierra Nevada, en el Plateau Modoc, en las montañas Warner y las Cascadas del Sur de California, EEUU. Se usaron modelos mixtos lineales generalizados para evaluar las hipótesis concernientes a la Respuesta de los escarabajos barrenadores al tipo de disturbio, severidad, tiempo de ocurrencia, la composición del rodal y su estructura, y las características de los árboles.

**Resultados:** La actividad de los barrenadores fue frecuentemente similar en áreas quemadas y en sitios que presentaron brotes de escarabajos descortezadores, y atemperada por respuestas localizadas por la actividad de escarabajos descortezadores, la severidad de los incendios, las características de los árboles, y la aparente respuesta a la fecha de ignición. La actividad de las larvas de barrenadores presentó una relación inversa a la presencia de escarabajos descortezadores dentro de un cuadrante de muestreo; esta relación fue más alta en pinos y en los lados con orientación sur de árboles pequeños en los sitios quemados. Los adultos de la familia Buprestidae fueron más abundantes donde la severidad del incendio fue mayor. Los incendios con fechas de ignición intermedias durante el período de estudio mantuvieron mayores tasas de actividad larval de barrenadores y mayores números de buprestidos adultos como así también de cerambícidos del género *Monochamus* Dejean 1821.

**Conclusiones:** La abundancia de los escarabajos barrenadores estuvo relacionada con la actividad de escarabajos descortezadores, la severidad del incendio, las características de los árboles, y la fecha de ignición. Considerar estos patrones cuando se manejan rodales perturbados puede beneficiar a la vida silvestre; incluyendo a aquellas especies que presan sobre escarabajos barrenadores. Así mismo, reportamos evidencia preliminar que la tendencia actual hacia incendios forestales más frecuentes podría no estimular mayores poblaciones de escarabajos barrenadores si estos incendios forestales ocurren de manera incremental fuera del lapso histórico de la temporada de estos.

## Abbreviations

AIC: Akaike's information criterion  
 BA: basal area  
 BBO: bark beetle outbreak  
 BUPR: buprestids  
 CERA: cerambycids  
 CWHR: California Wildlife Habitat Relationships  
 DBH: diameter at breast height  
 DOY: day of year  
 EPN: Eastside Pine (forest)  
 MECO: *Melanophila consputa*  
 MONO: *Monochamus* species  
 RdNBR: Relative differenced Normalized Burn Ratio  
 SE: standard error  
 SMC: Sierran Mixed Conifer (forest)  
 QAIC: quasi-AIC (see AIC)

VRP: variable radius plot

## Background

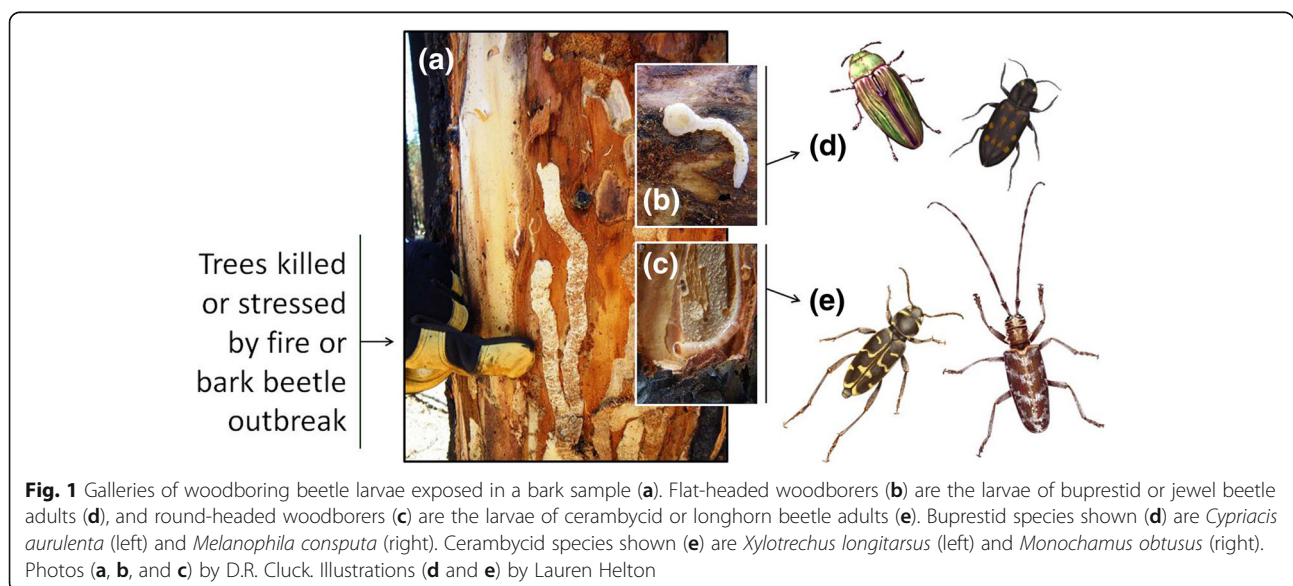
Forests in the western United States have entered a period of intensifying disturbance. Drought years have become increasingly common and severe, and projected changes in precipitation are expected to exacerbate these trends (Seager et al. 2007; Diffenbaugh et al. 2015). Severe drought and increasing temperatures leave forests drier and more susceptible to bark beetle (Coleoptera: Scolytinae) outbreaks triggered by the lower resistance of drought-stressed trees (Bentz et al. 2010; Williams et al. 2013; Kolb et al. 2016; Marini et al. 2017). The frequency and area of large fires (Littell et al. 2009; Dennison et al. 2014), as well as fire-season length (Westerling et al. 2006; Westerling 2016), have increased in recent decades across the western US, where forest fires are

expected to become more common in response to predicted changes in climate, especially in montane forests (Abatzoglou and Williams 2016). Larger and more severe wildfires have been facilitated by increased forest density, fuel loading, and the loss of forest heterogeneity resulting from fire exclusion (Miller et al. 2009a; Taylor et al. 2014; Boisramé et al. 2016). Over the next two decades, as many as 11 states in the western US might see the average annual area burned increase by 500 percent (Kitzberger et al. 2017), and temperature-driven models indicate that there is potential for extensive bark beetle outbreaks in forests of the western US throughout this century (Raffa et al. 2008; Bentz et al. 2010). The increase in tree mortality resulting from these large and frequent disturbances is expected to provide abundant habitat for woodboring beetles that typically colonize trees killed or weakened by other agents (Cranshaw et al. 2000). Many woodboring beetles aid decomposition and nutrient recycling in disturbed forest stands (Jacobsen et al. 2015; Kahl et al. 2017). Understanding how woodborers respond to different types of forest disturbance and stand characteristics will be important for predicting the response of forest communities to changes in the disturbance regime.

Woodboring beetle families Cerambycidae and Buprestidae contain hundreds of species, many of which are common and widespread throughout coniferous forests of the western US. As reviewed in Furniss and Carolin (1977), most woodboring beetles are considered secondary forest pests because they generally colonize trees already killed or weakened by disturbance, including wildfire, blowdown, timber harvest activities, and outbreaks of primary forest pests such as “primary” bark beetles that can kill healthy trees. In trees colonized by bark beetles, the spatial distribution of woodborer and bark beetle larvae is sometimes

disjunct, and there is some evidence of competition for phloem between species within trees (Coulson et al. 1976; Foelker et al. 2018), as well as some evidence that woodboring larvae consume bark beetle larvae (Dodds et al. 2001). Larval woodborers feed within the cambium layer during early development (Fig. 1) and later tunnel into sapwood and heartwood. Their tunneling introduces fungi that reduce the economic value of harvested timber but also facilitate the decomposition of dead wood, recycling nutrients and reducing fuel loads (Harrington 1993; Lowell and Cahill 1996). The life cycle of both cerambycid and buprestid beetles is dominated by the larval stage, which typically lasts one year but can extend to several years (Kariyanna et al. 2017). Adults emerge and fly during the warmer months, seeking out recently dead or weakened trees. Most woodborers are unable to attack healthy trees and are more often attracted to trees that release chemical signals of stress (Ikeda et al. 1980; Kelsey and Joseph 2003; Miller 2006). Some woodborers locate burned trees by sensing heat or smoke (Schmitz et al. 1997; Schütz et al. 1999; Álvarez et al. 2015). Females deposit eggs in bark crevices, under bark scales, or in small niches cut into the bark, and have been observed to avoid oviposition where bark beetle activity is high (Gardiner 1957).

Research on woodboring beetles has concentrated primarily on native species that cause damage to wood products (Álvarez et al. 2015) or exotic and invasive species that disrupt entire ecosystems (Aukema et al. 2010; Aukema et al. 2011). Much less is known about how native woodboring beetles colonize and use trees damaged by fire and primary bark beetles, and how woodborers respond to the timing and severity of disturbance, forest composition and structure, host tree attributes, and interspecific competition (Costello et al. 2011; Costello



2013; Brin and Bouget 2018). Given the observed potential for ecological and economic impacts of many woodborer species, characterizing these responses could inform management of disturbed forests. Management of post-disturbance forests often involves multiple objectives, such as recovering the economic value of dead trees (Eklund et al. 2009), mitigating hazards associated with disturbance, facilitating reforestation (Collins and Roller 2013; Bohlman et al. 2016), reducing fuel levels and the risk of type conversion (Coppoletta et al. 2016), and providing a diversity of wildlife habitats (White et al. 2015). Achieving these objectives might be facilitated by manipulating woodborer impacts on woody fuel decomposition, nutrient cycling, and succession, as well as increasing the availability of woodboring larvae as prey for early-seral habitat specialists. For example, the black-backed woodpecker (*Picoides arcticus* Swainson, 1832) is a species of management interest (Saracco et al. 2011; Siegel et al. 2018) that feeds primarily on the larvae of woodboring beetles (Villard and Beninger 1993; Murphy and Lehnhausen 1998). Relative to bark beetles, woodborer larvae are larger and provide a more concentrated prey resource (Powell 2000).

To inform forest management strategies designed to maintain processes dependent on woodborers, we characterized woodborer activity in 16 sites representing 11 wildfires and five bark beetle outbreaks. We tested several hypotheses (Table 1) regarding important controls on woodborer activity, based on our own experiences in the field as well as previous research on these species (e.g., Costello et al. 2011; Costello 2013). In general, our hypotheses address processes that we expected to affect

the activity (abundance) of both larval and adult woodborers, under the simplifying assumption that processes affecting one life stage should result in similar effects on the subsequent stage or generation. However, we conducted separate analyses of larvae and several adult taxa to differentiate responses by stage and taxon. Our results represent the first characterization of woodborer response to disturbance type and stand conditions across multiple forests in California.

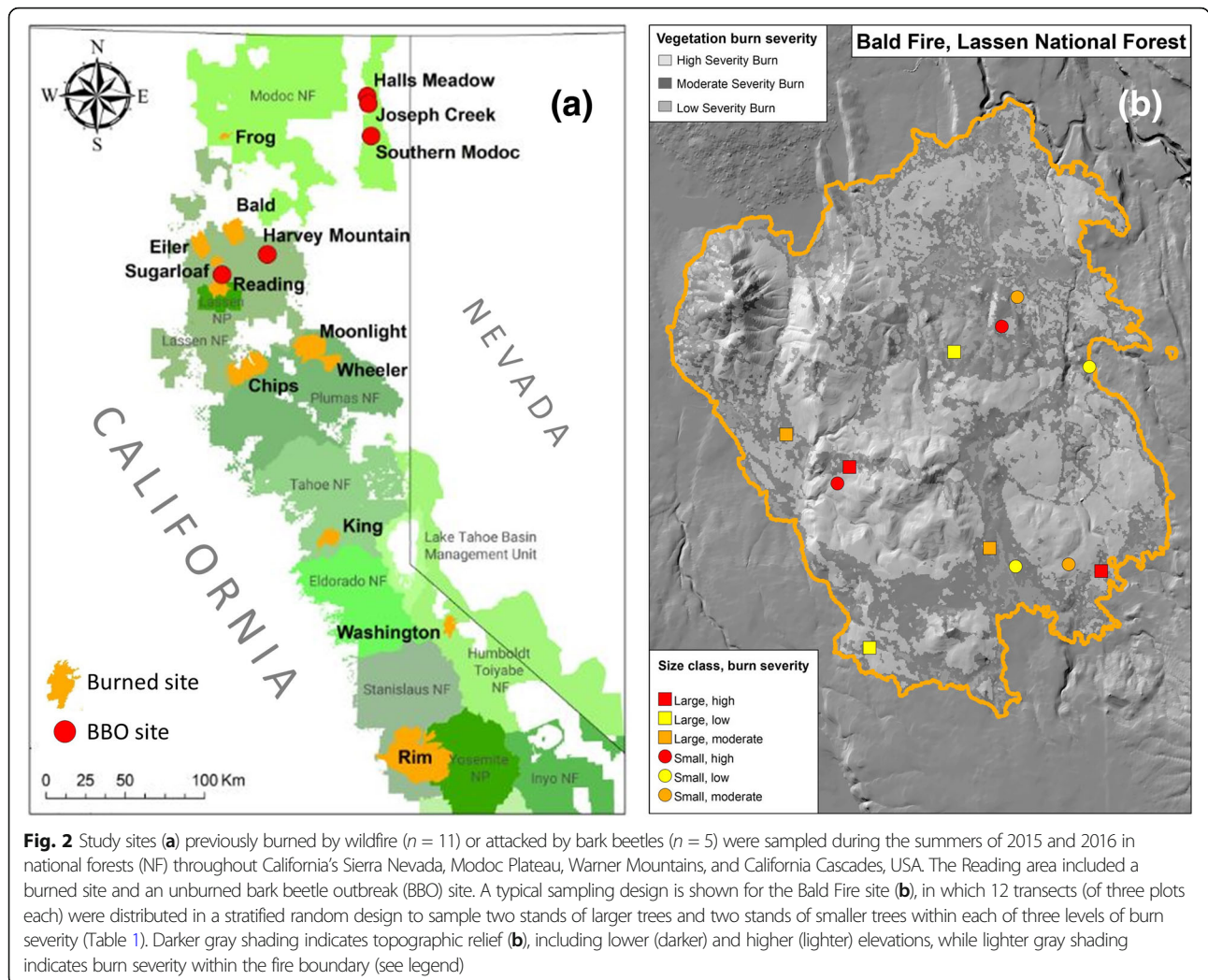
## Methods

### Site selection

During the summers of 2015 and 2016, we sampled 11 burned sites and five bark beetle outbreak (BBO) sites in the greater Sierra Nevada region of California, USA (Fig. 2), including the physiographic Sierra Nevada, Modoc Plateau, Warner Mountains, and southern Cascades. Sierran Mixed Conifer (SMC) and Eastside Pine (EPN) (California Department of Fish and Game 2005) forests were selected for sampling due to their common occurrence in the study region, and variety of constituent tree species. SMC forests typically consist of white fir (*Abies concolor* [Gordon and Glendinning] Lindley ex Hildebrand), Douglas-fir (*Pseudotsuga menziesii* [Mirb] Franco), ponderosa pine (*Pinus ponderosa* Douglas ex C.Lawson), sugar pine (*Pinus lambertiana* Douglas), incense-cedar (*Calocedras decurrens* [Torrey] Florin), and California black oak (*Quercus kelloggii* Newb). EPN forests consist of ponderosa pine with less representation by Jeffrey pine (*Pinus jeffreyi* A.Murray bis), lodgepole pine (*Pinus contorta* Douglas ex Loudon), white fir,

**Table 1** Non-exclusive, alternative hypotheses ( $H_A$ ) regarding processes expected to affect the activity (abundance) of larval or adult woodboring beetles that colonize trees killed or weakened by wildfire or bark beetle outbreak (BBO). These hypotheses were addressed with respect to five BBO sites and 11 burned sites that were sampled in the greater Sierra Nevada region of California, USA, during the summers of 2015 and 2016

$H_A$	Process
T	<b>Type of disturbance:</b> Woodborer activity is higher in burned than BBO stands because heat and smoke attract some woodborer taxa and because fire can create a pulse of resources (dead and weakened trees) not previously exploited by bark beetles.
M	<b>Magnitude of disturbance:</b> Woodborer activity is higher where dead trees are more abundant, due to the attractiveness of a large resource base.
S	<b>Severity of disturbance:</b> Woodborer activity is higher in stands burned at higher severity because more severe fires produce more smoke and heat as well as more dead and dying trees. Outbreak severity also affects woodborer activity, through positive effects on resource production and negative effects of increased interspecific competition.
A	<b>Age of disturbance:</b> Woodborer activity eventually declines with time after a disturbance, as the resource is depleted. Conversely, the activity of larval woodborers increases within the first year or two after a disturbance, as eggs continue to be deposited and larvae grow.
P	<b>Phenology:</b> Woodborer activity is lower in stands that ignite early or late in the fire season, if most adults disperse during the middle of the fire season when ignitions are most frequent.
H	<b>Host quality:</b> Woodborer activity varies by tree taxon due to taxon-specific differences in the quality of trees as a host resource.
G	<b>Growth rate:</b> Woodborer activity is higher on the southern sides of smaller snags (standing dead trees) in locations where diurnal warming can facilitate larval growth.
C	<b>Competition:</b> At scales appropriate to each life stage, woodborer activity varies inversely with bark beetle activity, due to adult host preferences and larval competition for phloem.



incense-cedar, Douglas-fir, California black oak, and western juniper (*Juniperus occidentalis* Hooker).

We selected sampling sites non-randomly, based on specific criteria and logistical concerns. All sites were selected prior to 2016, before large bark beetle outbreaks became evident in the southern Sierra Nevada. Burned and BBO sites were situated mainly in Plumas, Lassen, and Modoc national forests for easy access by our primary field crew, augmented by additional burned sites within the King, Washington, and Rim fires that were accessible by a secondary field crew (Fig. 2a). Burned sites (Table 2) were selected from fires that occurred throughout the prior decade and that burned enough forest on national forest land to accommodate 12 sampling transects (Fig. 2b). BBO sites (Table 3) were smaller and accommodated fewer transects.

To guide transect placement, we used several spatial data products (Additional file 1, a through g) to exclude areas (a) on private property, (b) with slopes exceeding 30%, (c) more than 250 m from roads, or d) where

post-disturbance logging had occurred; and to include only areas (e) within targeted forest types (SMC or EPN), and within either (f) mapped wildfire boundaries, or (g) areas with a high density of trees ( $>50 \text{ ha}^{-1}$ ) killed by bark beetles in the year prior to BBO site surveys. The remaining area was stratified by the dominant (pre-disturbance) tree size class (Additional file 1, e). Using California Wildlife Habitat Relationships (CWHR) classifications (California Department of Fish and Game 2005), we focused our study on trees in two size classes: class 3 trees (15 to 28 cm in diameter at breast height [DBH]), and classes 4 and 5 trees ( $>28 \text{ cm DBH}$ ).

In burned sites, burn severity was classified as low, moderate, or high (Additional file 1, f) based on a Relative differenced Normalized Burn Ratio (RdNBR) calculated from pre- and post-fire imagery and calibrated using the Composite Burn Index (Miller et al. 2009b; Miller and Quayle 2015). Within each fire's perimeter, transects were positioned randomly among the available levels of our two focal tree size classes and three burn

**Table 2** Characteristics of 11 burned sites sampled in the greater Sierra Nevada region of California, USA, during the summers of 2015 and 2016, including the date on which each wildfire ignited (Ignition date)

Burned site (fire name)	National forest	Area <sup>a</sup> (ha)	Ignition date	Survey		
				Year	Transects (n)	Elevation (m)
Wheeler	Plumas	9 484	5 Jul 2007	2015	12	1588–1726
Moonlight	Plumas	26 509	3 Sep 2007	2015	12	1565–2016
Sugarloaf	Lassen	3 897	1 Aug 2009	2015	11	1225–1590
Reading <sup>b</sup>	Lassen	11 478	23 Jul 2012	2015	12	1659–2003
				2016	12	1669–2162
Chips	Plumas and Lassen	31 210	28 Jul 2012	2016	12	1221–1834
Rim	Stanislaus	104 460	17 Aug 2013	2015	12	1160–2073
Bald	Lassen	16 258	30 Jul 2014	2016	12	1269–1599
Eiler	Lassen	13 624	31 Jul 2014	2015	10	1154–1715
King	Eldorado	39 843	13 Sep 2014	2015	12	1362–2206
Washington	Humboldt-Toiyabe	7 444	19 Jul 2015	2016	6	1866–2050
Frog	Modoc	2 036	30 Jul 2015	2016	6	1340–1358

<sup>a</sup>Total area within fire boundary (Additional file 1, g), regardless of jurisdiction  
<sup>b</sup>n = 24 separate transects sampled during 2015 and 2016

severities; in most cases, 12 transects were positioned in order to sample two stands of smaller trees and two stands of larger trees within each available level of burn severity. In some cases, some combinations of size class and burn severity were limited or unavailable within a fire boundary, resulting in fewer transects at those sites (Table 2).

BBO sites (Table 3) were dominated by white fir killed by fir engraver beetle (*Scolytus ventralis* LeConte), but also featured ponderosa pine killed by western pine beetle (*Dendroctonus brevicomis* LeConte) or Jeffrey pine killed by Jeffrey pine beetle (*Dendroctonus jeffreyi* Hopkins). Random transect positions in BBO sites were stratified by tree size class but not by severity of disturbance because the density of trees killed by bark beetles continued to increase unevenly across areas suitable for transect placement before our sampling could begin.

**Table 3** Characteristics of five bark beetle outbreak sites sampled in the greater Sierra Nevada region of California, USA, during the summer of 2016

BBO site	National forest	Area <sup>a</sup> (ha)	Survey		
			Year	Transects (n)	Elevations (m)
Harvey Mountain	Lassen	1162	2016	4	1967–2046
Reading <sup>b</sup>	Lassen	795	2016	9	1418–1619
Halls Meadow	Modoc	542	2016	6	1797–1908
Joseph Creek	Modoc	403	2016	8	1698–1907
Southern Modoc	Modoc	245	2016	3	1710–2087

<sup>a</sup>Total area of 2015 Aerial Detection Survey polygons per site (see Additional file 1, g)

<sup>b</sup>Adjacent to the Reading Fire but unburned

### Data collection

We established 159 transects (Tables 2 and 3), each 100 m long and aligned with a cardinal or inter-cardinal azimuth selected to fall within the defined tree size class and (if applicable) burn severity polygon. Each transect consisted of three plots (n = 477 total). Plot centers were established at 0, 50, and 100 m along each transect.

Larval activity and adult activity (Table 4) were selected as response variables for statistical modeling, and were measured at scales appropriate to each life stage. Adult woodborer activity was sampled at the plot scale using unbaited traps to avoid influencing the beetles' natural distribution. Each plot contained one panel trap (Alpha Scents, Inc., West Linn, Oregon, USA), mounted on a metal pole near the plot center and positioned at least 2 m from any tree. Traps were lubricated with Fluon (AGC Chemicals, Exton, Pennsylvania, USA), and each contained a pest strip in its collection cup. Traps were emptied every two weeks (mean = 13.98 ± 0.12 days), and remained in place for nearly eight weeks (52.65 ± 0.39 days) during June through August, the period of peak woodborer flight in the greater Sierra Nevada (D.R. Cluck, USDA Forest Service, Forest Health Protection, Susanville, California, USA, unpublished data).

Larval woodborer activity (Table 4) was sampled in bark quadrats on the north and south sides of up to 6 (5.19 ± 0.09) of the closest snags within 25 m of each trap or plot center. Bark quadrats were 15 cm × 15 cm and were positioned at breast height (1.37 m) in snags with at least one side suitable for larval sampling (bark attached or sapwood surface intact) (Powell et al. 2002). We placed quadrats in trees of a representative size by targeting snags 15 to 40 cm DBH where trees were in

**Table 4** Response variable definitions for the two life stages of woodboring beetles analyzed in this study of 11 burned sites and five bark beetle outbreak sites sampled in the greater Sierra Nevada region of California, USA, during the summers of 2015 and 2016

Response	Definition
Larval activity	Abundance of larval woodborer sign within a bark sampling quadrat, scored as 0 to 4 based on presence or absence of sign in each of four sub-quadrats
Adult activity	Number of adult woodborers (by taxon) captured per trap-visit within a plot

size class 3, and snags >40 cm DBH where trees were in size classes 4 and 5. Each quarter of each quadrat was scored separately for presence (1) or absence (0) of larval woodborer activity, including living or dead specimens and characteristic frass or boring dust, exit or transit holes, or galleries on any part of the quarter-quadrat, including the sapwood surface, bark interior, or (except frass or boring dust) bark exterior (Furniss and Carolin 1977).

Larvae collected from quadrats were stored in vials and identified to family, while adults collected from traps were sealed in plastic bags and identified to species, genus, or family using reference collections (USDA Forest Service, Forest Health Protection, Reference Insect Collection, Susanville, California, USA), online resources (<http://www.mtent.org/projects/wood-boring/index.html>), identification keys (Furniss and Carolin 1977, Arnett Jr. et al. 2002, Evans and Hogue 2006, Schiff et al. 2006, LaBonte et al. 2013a, LaBonte et al. 2013b), and consultation with Oregon Department of Agriculture for identification of *Melanophila* Eschscholtz 1829 species. Non-target insects captured in traps were generally identified to family.

Bark quadrats used for sampling larval woodborers were also used to quantify several tree characteristics that were targeted as potential predictors of woodborer activity (detailed in Table 5). Each full quadrat was scored for bark condition, char depth (Ryan 1982), and bark beetle activity (characteristic pitch tubes and other taxon-specific signs as listed above for woodboring larvae). Snags sampled via quadrat were also identified to species; measured for DBH, height, char height, and percent needles retained; and examined for broken top.

To quantify predictor variables related to stand characteristics (Table 5), we established a variable-radius plot (VRP) centered on each trap. We used a slope-compensating angle gauge to count trees  $\geq 10$  cm DBH, and estimated stand basal area ( $\text{m}^2 \text{ha}^{-1}$ ) from stem counts using a basal area factor of 10 or 20. For each stem counted within the VRP, we also recorded species and status (live or dead).

Predictors related to geographic gradients (Table 5) were calculated using a digital elevation model (Additional file 1). Elevation was adjusted by 130.1 m per degree of latitude, to account for the declining elevation of tree line with latitude in California (Brock and Inman 2006; Tingley and Beissinger 2013).

## Data analyses

We tested our focal hypotheses (Table 1) using an information-theoretic approach, fitting a set of alternative models for each response (Table 4) based on one to four fixed effects and selected interactions drawn from our list of potential predictor variables (Table 5). Response variables measured in burned sites were modeled separately from those in BBO sites, to address hypotheses regarding the effects of ignition date and burn severity not applicable in unburned sites. Mixed-effects multiple regression was used to account for the random effects in this system (Table 5), including spatial dependence among samples and repeated measures at each trap. Larval activity was modeled as a binomial process, and support for each larval model was ranked with respect to a null (random effects) model using AICc, an information criterion corrected for sample size (Burnham and Anderson 2002). Adult activity was modeled as a Poisson process adjusted for over-dispersion, and each adult model was ranked using QAICc (Burnham and Anderson 2002). For each numeric predictor variable  $x$ ,  $x_i$  ( $i$ th value of  $x$ ) was standardized by subtracting the mean of  $x$  and dividing by the standard deviation of  $x$ . Multicollinearity among fixed effects was avoided by screening predictor variables for high pairwise correlations (Kendall's tau > 0.5), or by replacing a predictor variable with the residual of its linear regression on a correlated predictor (following Graham 2003).

We reported the relative support for models using both (a)  $\Delta(\text{Q})\text{AICc}$  = focal model (Q)AICc – minimum (Q)AICc across each candidate model set, and (b)  $\Delta_{\text{Null}}$  = null model (Q)AICc – focal model (Q)AICc. We assumed that  $\Delta(\text{Q})\text{AICc} > 2$  indicated models with lower support than the top model, while (by extension)  $\Delta_{\text{Null}} > 2$  indicated models with higher support than the null model. We also summarized key data and results using means  $\pm$  standard errors (SE), and adopted  $\alpha = 0.05$  as our threshold for determining the significance of  $P$ -values. All analyses were conducted using the R environment for statistical computing (R Core Team 2016), and mixed-effect models were fit using package `lme4` (Bates et al. 2015).

## Results

### Patterns in potential covariates of woodborer activity

We measured 5683 trees (43% of which were snags) across our variable-radius plots, including 1683 white fir, 1268 ponderosa pine, 1265 Jeffrey pine, 568

**Table 5** Predictor variable definitions and associated hypotheses ( $H_A$  from Table 1) regarding woodborer response to disturbance severity (S), magnitude (M), age (A), and phenology (P), as well as host quality (H) and factors affecting or indicating the potential for larval growth (G) and interspecific competition (C). Values for each predictor variable in this study were drawn from 11 burned or five bark beetle outbreak sites sampled in the greater Sierra Nevada region during the summers of 2015 and 2016. RdNBR = Relative differenced Normalized Burn Ratio, BA = basal area, VRP = variable-radius plot, DOY = day of year, DBH = diameter at breast height, CWHR = California Wildlife Habitat Relationships

Predictor	Definition and unit of measurement	$H_A$
Bark beetle activity	Bark beetle sign (present = 1, absent = 0) within a bark sampling quadrat <sup>a</sup>	C
Burn severity <sup>b</sup>	RdNBR class (low, moderate, high) at the transect level	S
Char depth	Bark char depth class (deep = 3, moderate = 2, light = 1, unburned = 0, bark missing = NA) within a bark sampling quadrat <sup>a</sup>	S
Char height	Mean of two measures of maximum height (m) of char on a tree <sup>a</sup>	S
Snag BA ratio	Snag BA : total BA (unitless) within a VRP	S
Snag BA	Dead tree basal area ( $m^2 ha^{-1}$ ) within a VRP	M, S
Live tree BA	Live tree basal area ( $m^2 ha^{-1}$ ) within a VRP	M, S
Needles retained	Percent of needles (estimated in 5% increments) retained on a tree <sup>a</sup>	A
Fire age	Time (yr) since ignition of a burned site	A
Bark condition	General condition (tight, loose, missing) of bark within a quadrat	A
Broken-top tree	Top of tree is broken or missing (true, false) <sup>a</sup>	A
Ignition date	DOY of site-level ignition and its nonlinear effect ( $DOY \times DOY$ )	P
Host tree taxon	Taxon (yellow pine, true fir, or incense-cedar) of a tree <sup>a</sup>	H
Tree size <sup>b</sup>	Small (13–28 cm DBH = CWHR size class 3) or large (>28 cm DBH = CWHR size classes 4 and 5)	G
DBH	Diameter (cm) at breast height of a tree <sup>a</sup>	G
Height	Maximum vertical height (m) of a tree <sup>a</sup>	G
Bark aspect <sup>b</sup>	Aspect (northern or southern exposure) of a bark-sampling quadrat	G
Elevation	Latitude-adjusted plot elevation (m) and its nonlinear effect ( $m^2$ )	G
Latitude, longitude	Coordinates of plot position (UTM WGS 84)	G
Year <sup>p</sup>	Sampling year for the site	
Random effects <sup>b</sup>	Site, transect, plot, and tree	

<sup>a</sup>For models of adult woodborer activity, sub-plot data were averaged at the plot level, except bark condition and aspect (not used in adult models) and host tree taxon (replaced by dominant host tree taxon in each plot).

<sup>b</sup>Study design variables

incense-cedar, and several less common taxa. Yellow pines dominated in burned sites and white fir dominated in BBO sites (Table 6). Basal area (BA) of all trees (live and dead) calculated at the plot level and averaged at the transect level was significantly higher at BBO sites, while snag BA was significantly higher at burned sites. Tree size class and BA were correlated at the transect level (Kendall's tau = 0.13,  $P = 0.041$ ), but the measured size of trees averaged at the transect level did not differ between site types (Table 6).

In the subset of snags sampled for larval activity and additional covariates, mean DBH was slightly lower in burned sites (Table 6), which contained very few snags in the small size class. A similar proportion of trees were snags, regardless of size class: snag proportion was  $0.50 \pm 0.05$  in stands of larger trees and  $0.49 \pm 0.05$  in stands of smaller trees. Bark beetle sign was found in

nearly one third and over two thirds, respectively, of quadrats in burned and BBO sites (Table 6). Covariance among transect-level metrics of tree size (mean height, BA, and CWHR class) tended to be low but significant ( $0.07 < \text{Kendall's tau} < 0.26$ ,  $P < 0.015$ ).

Among our potential metrics of burn severity (Table 5, Additional file 2), we found high and significant correlations between RdNBR, char depth, and snag ratios (Kendall's tau  $\geq 0.516$ ,  $P < 0.001$ ). However, percent needles retained and bark condition were less correlated with RdNBR, and were instead significantly correlated with fire age (Kendall's tau  $\geq 0.152$ ,  $P \leq 0.027$ ). Snag ratios were most highly correlated with RdNBR (Additional file 3).

#### Larval woodborer activity

We rated larval woodborer activity on 2297 dead trees in 443 of 477 plots, including 355 plots within burned



**Table 6** Comparisons of raw data from 11 burned sites and five bark beetle outbreak sites sampled in the greater Sierra Nevada region of California, USA, during the summers of 2015 and 2016, summarized by transect as count ratios or mean  $\pm$  standard error. BBO = bark beetle outbreak, BA = basal area, DBH = diameter at breast height

Feature <sup>a</sup>	Burned sites	BBO sites
Dominant tree taxon (count ratio)	Yellow pine (2.71:1)	White fir (1.48:1)
BA (m <sup>2</sup> ha <sup>-1</sup> )**	31.69 $\pm$ 1.27	39.28 $\pm$ 2.59
Snag BA (m <sup>2</sup> ha <sup>-1</sup> )**	14.87 $\pm$ 1.31	11.20 $\pm$ 1.20
Mean DBH (cm) <sup>ns</sup>	46.17 $\pm$ 1.22	45.42 $\pm$ 1.52
Snag mean DBH (cm) <sup>ns</sup>	42.38 $\pm$ 1.26	44.45 $\pm$ 2.27
Larval sampling snag DBH (cm)**	36.55 $\pm$ 1.23	40.54 $\pm$ 1.50
Bark beetle presence per quadrat***	0.61 $\pm$ 0.04	1.46 $\pm$ 0.05
Larval woodborer activity per quadrat <sup>ns</sup>	1.99 $\pm$ 0.08	1.74 $\pm$ 0.12
Adult buprestids captured (n) <sup>ns</sup>	1.00 $\pm$ 0.12	0.88 $\pm$ 0.11
Adult cerambycids captured (n) <sup>ns</sup>	0.72 $\pm$ 0.07	0.90 $\pm$ 0.11

<sup>a</sup>ns ( $P > 0.05$ ), \* ( $P < 0.05$ ), \*\* ( $P < 0.01$ ), \*\*\* ( $P < 0.001$ )

sites and 88 plots within BBO sites; in the remaining 34 plots, no dead trees were available for sampling. Larvae collected were identified as buprestids ( $n = 158$ ), cerambycids ( $n = 221$ ), bark beetles ( $n = 223$ ), and a variety of less frequent taxa ( $n = 162$ ). Larval woodborer activity (Table 4) was evident mainly from galleries, frass or boring dust, and exit or transit holes, rather than direct observations of larvae. Although these indirect signs were sufficient for distinguishing larval woodborers from other taxa, they were often insufficient for differentiating woodborers by family due to intermingled galleries of unrelated taxa and deterioration of the inner bark and sapwood interface on older snags. Based on these data, the activity per quadrat of all larval woodborers combined was similar between burned and BBO sites for data summarized at the transect level (Table 6).

Our null model of larval activity for burned sites included nested random effects of site, transect, and tree. We omitted plot effects because they explained <5% of the random variance and because larval activity was sampled without repeated measures. A fixed effect of survey year (2015 or 2016) was not supported ( $\Delta_{Null} = -1.84$ ), nor were fixed or random effects of burn year ( $\Delta_{Null} \leq -5.43$ ). For BBO sites, our null model of larval activity included a fixed effect of site ( $n = 5$  sites), as well as nested random effects of transect, plot, and tree.

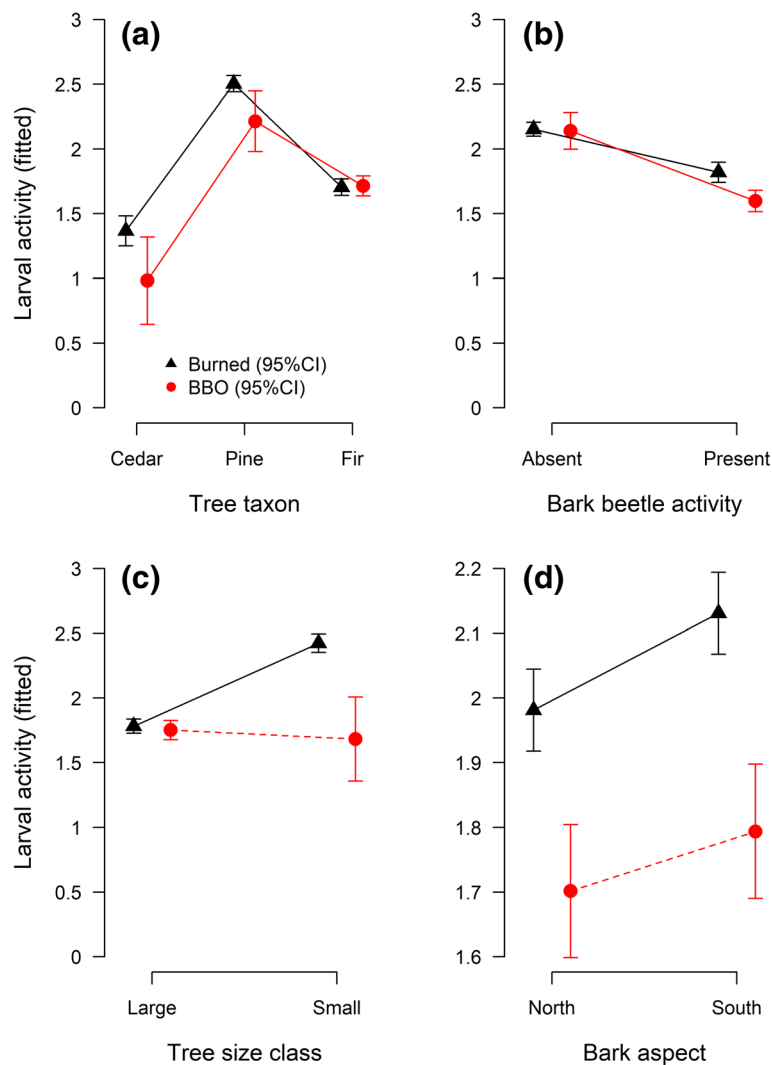
Adding our predictor variables (Table 5) one at a time to the null model for each site disturbance type revealed some similarities in the significant predictors of larval activity between site types. Apparent effects of tree taxon and bark beetle presence were similar in burned and BBO sites (Fig. 3). Regardless of disturbance type, larval woodborer activity was highest in yellow pine (hereafter, pine), followed by true fir (fir) and incense-cedar (cedar) (Fig. 3a). Lower activity was recorded in quadrats where bark beetle activity was

present (Fig. 3b). Larval activity was significantly higher on smaller trees in burned sites (where small and large trees were sampled equally) but not in BBO sites (where there were few small trees available for sampling) (Fig. 3c). Bark aspect had a significant but minor effect on larval activity after fire, and no significant effect in BBO sites (Fig. 3d).

Model rankings confirmed the support for negative effects of bark beetle activity on larval woodborer activity in both burned and BBO sites. In burned sites (Table 7), the top four models of larval activity confirmed support for effects of bark beetle activity, host tree taxon, tree size, and bark aspect, as in the often simpler models used to generate Fig. 3. In the fifth-ranked model, there was also support for linear and quadratic effects of fire ignition date, with negative effects suggesting lower larval activity in fires ignited at earlier and (especially) later dates during our study period. Support was much higher, however, for effects that we associated with hypotheses related to larval development, resource quality, and competitive interactions within the host resource.

In BBO sites, site was confounded with tree taxon and bark beetle activity (Additional file 4), reducing our ability to confirm these effects. Nonetheless, the BBO site richest in pine and with lowest bark beetle activity showed much higher activity of woodborer larvae ( $12.11 \pm 0.01$  per plot) than other BBO sites ( $8.05 \pm 0.01$ ), and negative effects of bark beetle activity were included in models ranked 1 and 5 (Table 8).

Aside from competition, data from BBO sites did not support the same hypotheses as data from burned sites (Table 7); instead, top models from BBO sites were based on potential metrics of disturbance age. Signs of larval activity were more abundant on snags with fewer needles retained, with broken tops, and with loose (as opposed to tight or missing) bark. The lower activity associated with



**Fig. 3** Larval woodborer activity in burned (black) and bark beetle outbreak (BBO; red) sites in relation to host tree taxon (a), bark beetle activity (b), tree size class at the transect level (c), and aspect of the bark sampling quadrat (d), in the greater Sierra Nevada region of California, USA, sampled during the summers of 2015 and 2016. Point estimates and error bars represent mean responses and 95% confidence intervals (CIs) from fits to models exemplified in Tables 7 and 8. Dashed lines connect means that do not differ significantly. Response variables are defined in Table 4 and predictor variables are defined in Table 5

missing bark was expected, because signs of activity were sometimes recorded from bark alone. Trees with broken tops might have suffered damage prior to bark beetle outbreaks, allowing more time for woodborer colonization. Needle loss appeared to represent outbreak age, rather than outbreak severity, as bark beetle activity and needles retained were uncorrelated at the transect level within BBO sites (Kendall's tau = 0.085).

#### Adult woodborer activity

We identified 10 412 adults belonging to six families: 1718 Buprestidae, 1277 Cerambycidae, 5765 Elateridae,

285 Siricidae, 396 Melandryidae, and 971 Cupedidae. The most frequently captured buprestids were *Cypriacis aurulenta* Linneaus 1767 ( $n = 466$ ), *Melanophila consputa* LeConte 1857 ( $n = 200$ ), and *Chalcophora angulicollis* LeConte 1857 ( $n = 193$ ), while *Xylotrechus longitarsus* Casey 1912 ( $n = 190$ ), *Monochamus obtusus* Casey 1891 ( $n = 89$ ), and *Ortholeptura valida* LeConte 1857 ( $n = 88$ ) were most frequent among cerambycids (Additional file 5). Annual differences in buprestid and cerambycid captures could be explained by site (Additional file 6). Raw capture numbers were similar in burned and BBO sites for both buprestids and cerambycids (Table 6).

**Table 7** Fixed effects and associated hypotheses ( $H_A$  from Table 5) in top models of larval woodborer activity across 11 burned sites sampled in the greater Sierra Nevada region of California, USA, during the summers of 2015 and 2016. AICc = Akaike’s information criterion, corrected for small sample size; DOY = day of year; Neg = negative; Pos = positive

Model rank	Predictor	Effect <sup>a</sup>	$\Delta AICc$	$\Delta_{Null}$	$H_A$
1	Bark beetle activity	Neg <sup>***</sup>	0.00	115.86	C
	Host tree taxon = pine	Pos <sup>***</sup>			H
2	Host tree taxon = pine	Pos <sup>***</sup>	14.93	100.93	H
	Host tree taxon = fir	Pos <sup>ns</sup>			H
	Host tree taxon = cedar	Pos <sup>ns</sup>			H
3	Host tree size-class = small	Pos <sup>***</sup>	51.69	64.17	G
	Bark aspect = south	Pos <sup>***</sup>			G
	Fire age	Pos <sup>ns</sup>			A
	Bark aspect × Fire age	Neg <sup>***</sup>			G, A
4	Bark aspect = south	Pos <sup>***</sup>	71.91	43.95	G
	Longitude	Pos <sup>***</sup>			G
	Elevation	Pos <sup>ns</sup>			G
5	Ignition date (DOY)	Neg <sup>*</sup>	107.41	8.45	P
	DOY × DOY	Neg <sup>**</sup>			P
	Fire age	Pos <sup>ns</sup>			A
	DOY × DOY × Fire age	Pos <sup>ns</sup>			P, A

<sup>a</sup>ns ( $P > 0.05$ ), \* ( $P < 0.05$ ), \*\* ( $P < 0.01$ ), \*\*\* ( $P < 0.001$ )

Mixed models of adult activity included nested random effects of site and transect for burned sites, or a fixed effect of site and random effect of transect for BBO sites. Model rankings confirmed predictive effects of bark beetle activity, tree size, and burn severity (Fig. 4), as well as ignition date (Fig. 5), among others (Tables 9, 10, 11).

Except for negative effects of ignition date and fire age, top cerambycid models differed from top buprestid models either in hypotheses supported or direction of effects. For example, *Xylotrechus longitarsus* activity in

**Table 8** Fixed effects and associated hypotheses ( $H_A$  from Table 5) in top models of larval woodborer activity across five bark beetle outbreak sites sampled in the greater Sierra Nevada region of California, USA, during the summers of 2015 and 2016. AICc = Akaike’s information criterion, corrected; Neg = negative; Pos = positive

Model rank	Predictor	Effect <sup>a</sup>	$\Delta AICc$	$\Delta_{Null}$	$H_A$
1	Bark beetle activity	Neg <sup>**</sup>	0.00	20.22	C
	Needles retained	Neg <sup>***</sup>			A
2	Needles retained	Neg <sup>***</sup>	5.37	14.85	A
	Bark condition = tight	Neg <sup>***</sup>			A
3	Bark condition = missing	Neg <sup>*</sup>	7.73	12.49	A
	Broken-top tree = true	Pos <sup>**</sup>			A
4	Bark beetle activity	Neg <sup>**</sup>	12.73	7.49	A
5	Bark beetle activity	Neg <sup>**</sup>	13.90	6.32	C

<sup>a</sup>ns ( $P > 0.05$ ), \* ( $P < 0.05$ ), \*\* ( $P < 0.01$ ), \*\*\* ( $P < 0.001$ )

burned stands supported strong positive effects of latitude-adjusted elevation (Table 10) that were not supported by buprestids (Table 9), and *Monochamus* Dejean 1821 species supported strong effects of burn severity but were more common in stands burned at low to moderate severity (Table 10), in contrast with buprestid *Melanophila consputa* (Table 9).

Across multiple buprestid and cerambycid taxa, we found support for negative linear and quadratic effects of ignition date as well a negative effect of fire age and a positive interaction between fire age and the quadratic effect of ignition date. Buprestid data also repeatedly supported the burn severity hypothesis.

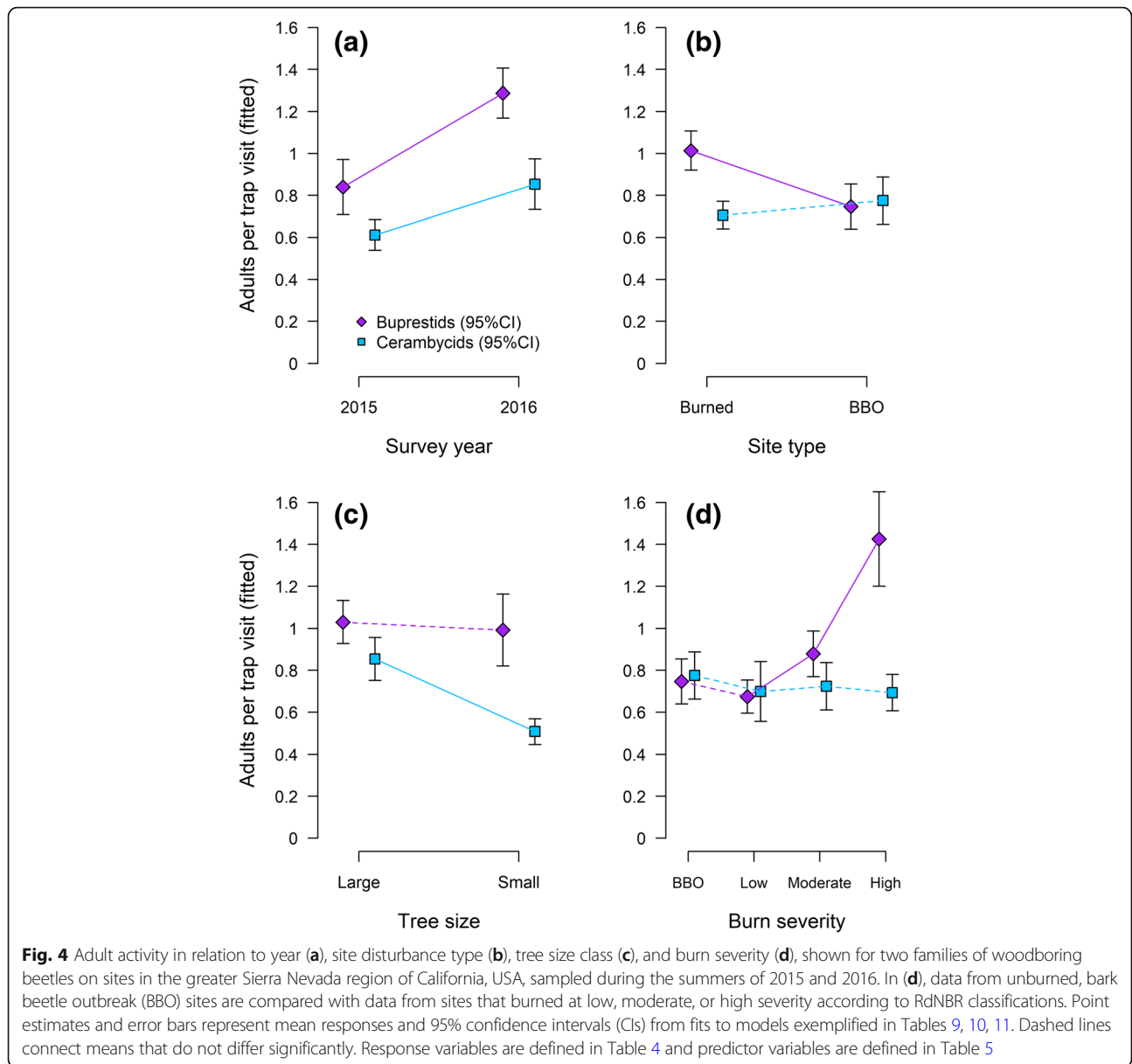
For buprestid captures in BBO sites (Table 11), the top model included a negative effect of live tree BA, suggesting support for disturbance magnitude and severity hypotheses. All other models of buprestid and cerambycid adults in BBO sites had much lower relative support and suggested that processes related to habitat quality, larval growth, and competition explained adult activity.

**Comparison of larval and adult activity within fires**

We compared fire site effects on larval activity and adult captures by adding a fixed effect of site to null models of larval activity or adult captures. In no case did this effect of site differ in direction between models of larval activity and adult captures. Both larval activity and adult captures were significantly lower than average in samples from the King, Moonlight, and Rim fires (Table 12), which all ignited relatively late in the season (Table 1). At the Washington Fire (1 yr old), larval activity was significantly lower than average but adult captures were moderate. At the Wheeler fire (8 yr old), larval activity was moderate but adult captures were significantly lower than average.

**Discussion**

In these montane forests of California, the range of woodborer response to forest conditions suggests that managing for a diversity of disturbance types and severity will support a diverse woodborer community. The different taxa and life stages that we sampled appeared to respond in characteristic ways. Buprestid adults were much more frequent in stands that burned at high severity, a pattern especially evident in *Melanophila consputa*, which is known for its attraction to heat and smoke (Schmitz et al. 1997; Schütz et al. 1999). In contrast, cerambycids that sense heat and smoke (*Monochamus* spp.; Álvarez et al. 2015) were more frequent in transects burned at low and moderate severity. Larval activity responded more often to features that might represent the capacity for larval growth and development. Disturbance type, however, appeared to have a relatively minor influence on the activity of most woodborers in our

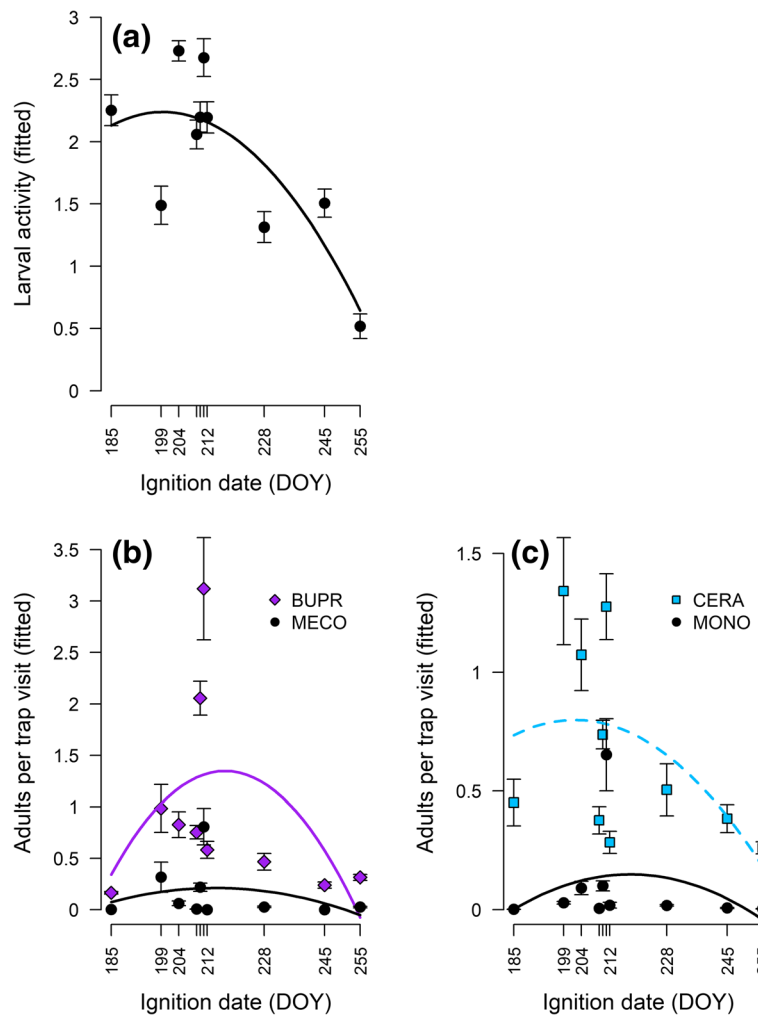


study. Except where burn severity was high, cerambycids and buprestids of both stages appeared to occur with similar frequency in burned and BBO sites.

The scale and timing of each analysis should be considered when interpreting an appropriate scope of inference. Burned sites were distributed throughout the Sierra Nevada, while BBO sites were clustered in the northern half of this region, and disturbance type also covaried with disturbance age, because large bark beetle outbreaks had just begun to develop as we initiated the study. We also sampled during a period of extended drought in our study region, when primary and secondary forest pests were likely to be responding in complex ways to stress on the forests in this system (Kolb et al. 2016). Regardless of the unique features of this study,

strong patterns were revealed in our data that are likely relevant to many dry coniferous forests that occur throughout the western US. Evidence for these relationships should be investigated for generality in other regions, and across a diversity of fire ages and outbreak sites, to facilitate inferences currently constrained by the naturally high covariance and spatial clustering in values of key predictor variables.

The strong patterns we found, however, can help inform where—and at what scale—management actions might be most effectively applied. Within a disturbed patch of forest, the deliberate retention of pockets or islands of snags with specific characteristics might limit the negative ecological impacts of a reduction in woodborer activity. For example, larval woodborers were



**Fig. 5** Activity of larvae (a) and adults (b, c), as defined in Table 4, in relation to fire ignition date (DOY = day of year) at sites in the greater Sierra Nevada region of California, USA, sampled during the summers of 2015 and 2016. Curves represent relationships between squared DOY and fitted point estimates of woodborer response ( $\pm 95\%$  confidence interval) obtained from linear mixed effects models that included fixed effects of DOY and an interaction between squared DOY and fire age (Tables 7, 9, and 10). BUPR = all buprestids, MECO = *Melanophila consputa*, CERA = all cerambycids, MONO = *Monochamus* species. Effects of squared DOY were not supported for CERA. Data from 12 surveys (Table 2) are represented here by only 10 point estimates in each panel because some fires shared the same ignition date (DOY = 210 for Bald and Frog fires, and DOY = 204 for 2015 and 2016 samples from the Reading Fire)

more common in particular quadrats where bark beetles were absent, suggesting that the local retention of woodborers might be enhanced by prioritizing retention of trees less heavily infested by bark beetles. Our observations that woodborer larvae were more abundant in smaller pine snags with more southerly exposure, that *Xylotrechus* Chevrolat 1860 adults were more common where the dominant host tree taxon was true fir, and that *Monochamus* species were more common where burn severity was low to moderate, can further inform design of snag retention patches that might effectively promote successional processes and provide prey for wildlife species like the black-backed woodpecker (Siegel et al. 2018; Tarbill et al. 2018; Stillman et al. 2019).

The design of post-fire forest management strategies often involves balancing multiple economic and ecological objectives with clear links to woodborer dynamics. Taken together, our results suggest a dynamic process controlling local initiation and accumulation of woodborer activity that likely leads to temporal trends in the faunal communities supported by woodborer larvae and adults as prey and agents of tree decomposition. Beginning with the environment inhabited by larvae, our finding of higher larval activity on smaller trees and southern bark aspects suggests that there are microclimatic effects on larval development. Smaller-diameter trees have thinner bark and lower thermal mass of sap and heartwood, as well as lower sapwood volume (water holding capacity), leading to more

**Table 9** Fixed effects and associated hypotheses ( $H_A$  from Table 5) in top models of adult buprestid activity across 11 burned sites sampled in the greater Sierra Nevada region of California, USA, during the summers of 2015 and 2016. QAICc = quasi-Akaike's information criterion, corrected for small sample size; DOY = day of year; Neg = negative; Pos = positive; BA = basal area

Response	Model rank	Predictor	Effect <sup>a</sup>	$\Delta$ QAICc	$\Delta$ Null	$H_A$
Adult buprestid activity	1	Ignition date (DOY)	Neg <sup>ns</sup>	0.00	11.96	P
		DOY × DOY	Neg <sup>***</sup>			P
		Fire age	Neg <sup>***</sup>			A
		DOY × DOY × Fire age	Pos <sup>***</sup>			P, A
	2	Snag BA	Pos <sup>**</sup>	6.17	5.79	M, S
		Fire age	Neg <sup>**</sup>			A
		Snag BA × Fire age	Pos <sup>ns</sup>			M, S, A
	3	Snag BA	Pos <sup>***</sup>	6.32	5.64	M, S
	4	Char height	Pos <sup>**</sup>	7.57	4.39	S
	5	Char height	Pos <sup>**</sup>	7.66	4.30	S
Fire age		Neg <sup>**</sup>	A			
Char height × Fire age		Neg <sup>ns</sup>	S, A			
Adult <i>Cypriacis aurulenta</i> activity <sup>b</sup>	1	Host tree taxon = pine	Pos <sup>**</sup>	0.00	0.39	H
Adult <i>Melanophila consputa</i> activity	1	Fire age	Neg <sup>***</sup>	0.00	20.19	A
		Char height	Pos <sup>ns</sup>			S
		Live tree BA	Neg <sup>**</sup>			M, S
		Ignition date	Neg <sup>***</sup>			P
	2	Fire age	Neg <sup>***</sup>	2.33	17.86	A
		Burn severity = low	Neg <sup>***</sup>			S
		Burn severity = moderate	Neg <sup>***</sup>			S
		Ignition date (DOY)	Neg <sup>*</sup>			P
		DOY × DOY	Neg <sup>*</sup>			P
	3	Fire age	Neg <sup>***</sup>	3.84	16.35	A
		Char height	Pos <sup>***</sup>			S
		Ignition date (DOY)	Neg <sup>*</sup>			P
		DOY × DOY	Neg <sup>*</sup>			P
	4	Fire age	Neg <sup>***</sup>	4.46	15.73	A
		Burn severity = low	Neg <sup>***</sup>			S
Burn severity = moderate		Neg <sup>**</sup>	S			
5	Live tree BA	Neg <sup>***</sup>	7.55	12.64	M, S	

<sup>a</sup>ns ( $P > 0.05$ ), \* ( $P < 0.05$ ), \*\* ( $P < 0.01$ ), \*\*\* ( $P < 0.001$ )

<sup>b</sup>Only one model of adult *Cypriacis aurulenta* activity was supported relative to the null model

rapid desiccation of phloem tissue (Six et al. 2002; Hayes et al. 2008). Southern bole aspect would intensify these microclimatic conditions, likely creating a more favorable environment for rapid larval development. An inverse relationship between tree size and larval woodborer activity was also noted by Powell (2000) in the northern Rocky Mountains. Hanks (1999) suggested that phloem-boring larvae might be especially vulnerable to moisture conditions because they are embedded in the tissues of their host plant, and sensitivity to bark moisture could account for the association between drought stress and host susceptibility that is typical for many phloem-boring insects,

especially cerambycid beetles. Thatcher (1961) reported that drought conditions can enable a larger number of cerambycid larvae to survive in systems in which adults attempt to attack living trees. Subcortical and sapwood temperatures in trees and logs exposed to sunlight can be 5 to 30 °C greater than shaded portions, and these warmer temperatures can decrease the larval development time of phloem-feeding woodborers by 1 to 2 yr (Graham 1925, Savely 1939, Hosking 1977; reviewed in Kariyanna et al. 2017). Alternatively, these patterns could be related to adult female selection of oviposition locations, although we found no evidence of such selectivity in the literature.

**Table 10** Fixed effects and associated hypotheses ( $H_A$  from Table 5) in top models of adult cerambycid activity across 11 burned sites sampled in the greater Sierra Nevada region of California, USA, during the summers of 2015 and 2016. QAICc = quasi-Akaike's information criterion, corrected for small sample size; DOY = day of year; Neg = negative; Pos = positive; BA = basal area

Response	Model rank	Predictor	Effect <sup>a</sup>	$\Delta QAIC_c$	$\Delta_{Null}$	$H_A$	
Adult cerambycid activity	1	Null		0.00	0.00		
Adult <i>Xylotrechus longitarsus</i> activity	1	Fire age	Neg <sup>**</sup>	0.00	4.66	A	
		Elevation	Pos <sup>***</sup>			G	
		Ignition date	Neg <sup>ns</sup>			P	
	2	Elevation	Pos <sup>***</sup>	0.02	4.64	G	
		3	Host tree taxon = fir	Pos <sup>**</sup>	0.76	3.90	H
			4	Elevation	Pos <sup>***</sup>	2.04	2.62
Adult <i>Monochamus</i> species activity	5	Elevation <sup>2</sup>	Pos <sup>ns</sup>			G	
		Snag BA	Pos <sup>***</sup>	2.68	1.98	M, S	
	1	Fire age	Neg <sup>*</sup>	0.00	9.06	A	
		Burn severity = low	Pos <sup>**</sup>			S	
		Burn severity = moderate	Pos <sup>***</sup>			S	
	2	Burn severity = low	Pos <sup>**</sup>	1.35	7.71	S	
		Burn severity = moderate	Pos <sup>***</sup>			S	
	3	Ignition date (DOY)	Neg <sup>ns</sup>	1.56	7.50	P	
		DOY × DOY	Neg <sup>ns</sup>			P	
		Fire age	Neg <sup>***</sup>			A	
DOY × DOY × Fire age		Pos <sup>***</sup>			P, A		
4	Needles retained	Pos <sup>***</sup>	2.64	6.42	A		
5	Fire age	Neg <sup>*</sup>	3.16	5.90	A		
	Burn severity = low	Pos <sup>**</sup>			S		
	Burn severity = moderate	Pos <sup>***</sup>			S		
	Ignition date (DOY)	Neg <sup>ns</sup>			P		
		DOY × DOY	Neg <sup>ns</sup>			P	

<sup>a</sup>ns ( $P > 0.05$ ), \* ( $P < 0.05$ ), \*\* ( $P < 0.01$ ), \*\*\* ( $P < 0.001$ )

**Table 11** Fixed effects and associated hypotheses ( $H_A$  from Table 5) in top models of adult buprestid or cerambycid activity across five bark beetle outbreak sites sampled in the greater Sierra Nevada region of California, USA, during the summer of 2016. QAICc = quasi-Akaike's information criterion, corrected for small sample size; BA = basal area; Neg = negative; Pos = positive; DBH = diameter at breast height

Response	Model rank	Predictor (fixed effects)	Effect <sup>a</sup>	$\Delta QAIC_c$	$\Delta_{Null}$	$H_A$
Adult buprestid activity	1	Live tree BA	Neg <sup>***</sup>	0.00	25.85	M, S
	2	Bark beetle activity	Pos <sup>**</sup>	20.97	4.88	C
	3	Elevation	Neg <sup>**</sup>	21.20	4.65	G
	4	Longitude	Neg <sup>**</sup>	21.97	3.88	G
	5	Needles retained	Pos <sup>**</sup>	22.36	3.49	A
Adult cerambycid activity	1	Host tree taxon = cedar	Neg <sup>**</sup>	0.00	4.02	H
	2	Broken-top tree = true	Neg <sup>**</sup>	1.00	3.02	A
	3	DBH	Neg <sup>**</sup>	1.38	2.64	G
	4	Host tree taxon = fir	Pos <sup>*</sup>	3.46	0.56	H
	5	Null		4.02	0.00	

<sup>a</sup>ns ( $P > 0.05$ ), \* ( $P < 0.05$ ), \*\* ( $P < 0.01$ ), \*\*\* ( $P < 0.001$ )

**Table 12** Fixed effects of burned site on larval and adult woodborer activity across 11 burned sites sampled in the greater Sierra Nevada region of California, USA, during the summers of 2015 and 2016. Neg = negative; Pos = positive

Burned site (fire name)	Fire age (yr)	Fixed effect on activity <sup>a</sup>	
		Larval model	Adult model
Eiler	1	ns	Pos <sup>**</sup>
Frog	1	ns	ns
King	1	Neg <sup>***</sup>	Neg <sup>**</sup>
Washington	1	Neg <sup>*</sup>	ns
Bald	2	ns	Pos <sup>*</sup>
Rim	2	Neg <sup>***</sup>	Neg <sup>*</sup>
Reading (2015)	3	Pos <sup>**</sup>	ns
Reading (2016)	4	ns	ns
Chips	4	ns	ns
Sugarloaf	6	ns	ns
Moonlight	8	Neg <sup>*</sup>	Neg <sup>*</sup>
Wheeler	8	ns	Neg <sup>***</sup>

<sup>a</sup>ns ( $P > 0.05$ ), \* ( $P < 0.05$ ), \*\* ( $P < 0.01$ ), \*\*\* ( $P < 0.001$ )

The inverse relationship between bark beetle and larval woodborer activity that we observed was likely due to competition for phloem resources or other factors affecting adult or larval microhabitat partitioning among beetle species. Because woodboring beetles typically colonize dead and dying trees, such as those killed or weakened by bark beetles, the nutrient-rich phloem resource can be greatly reduced by bark beetle activity prior to colonization by woodboring beetles, limiting its availability for larval woodborer development (Coulson et al. 1979; Powell 2000). Thus, variation in the availability of phloem for woodboring beetle larvae could depend on the timing of bark beetle attack versus woodborer colonization. For example, the western pine beetle (*Dendroctonus brevicomis* Erichson 1836) produces more than one generation per year (Miller and Keen 1960), which might allow these bark beetles to attack trees outside the peak flight period for woodboring beetles. If bark beetles attack trees in the weeks or months before woodborer dispersal, the resource might be depleted before woodborer eggs hatch. If bark beetles attack trees after woodborer dispersal, changes in the resource during the ensuing year might combine with resource depletion to make trees unattractive or unsuitable for some woodboring beetles. Alternatively, tree size can play a role in competition with bark beetles such as the fir engraver (*Scolytus ventralis* LeConte 1868), which tends to attack higher on the bole of larger-diameter trees, leaving the lower portions of the bole available for woodboring beetles (Ferrell and Hall 1975). In addition, because primary bark beetles avoid fire-killed trees,

especially those with foliage completely scorched or consumed, the phloem resource in fire-killed trees might often be more available for woodboring beetle colonization (Parker et al. 2006). In contrast, competition with secondary bark beetles, such as *Ips* De Geer 1775 spp., in fire-killed trees may occur in the upper boles and limbs of larger trees and all portions of smaller trees. This is especially true for secondary bark beetles attacking thicker-barked tree species such as those sampled in this study. However, Costello (2013) found that no bark beetles emerged from fire-killed tree bolts, even though outbreaks of the mountain pine beetle (*Dendroctonus ponderosae* Hopkins 1902) were abundant near the fires in that study.

Costello (2013) also found significantly more woodborers emerging from fire-killed trees than from trees killed by bark beetles, in contrast with our finding of similar woodborer activity in burned and BBO stands. This contrast could reflect a difference in the timing of post-fire monitoring between their study of fires 1 to 2 yr old and our study of fires 1 to 8 yr old. Furthermore, bark beetle activity was common in our burned sites, occurring in over 30% of bark samples. Some snags in older burned sites might have been killed by bark beetles attacking fire-stressed trees in the years following a burn (Davis et al. 2012), reducing the distinction between burned and BBO sites as a food resource for woodboring beetles. Potential similarities in post-disturbance ecology of burned and BBO sites warrant further investigation.

Fire age also helped explain the pattern of adult captures. For all but one taxon modeled in burned sites, adult capture numbers declined with fire age, as expected if older fires were less attractive to dispersing adults or less productive as source habitats for these species. Data from buprestids (especially *M. consputa*) and cerambycids (*Monochamus* species) also supported a positive interaction between fire age and squared ignition date. Although these species were more common at burned sites with intermediate ignition dates, those fires did not appear to act as long-term sources of these species in the landscape. This dynamic process should lead to temporal trends in the faunal communities woodborers support. Indeed, black-backed woodpecker abundance in burned forests in California generally peaks within five years after fire, followed by sharp declines (Tingley et al. 2018).

Our results also suggest that there are competing influences of disturbance age. Adult woodborers might cue on current signs of a snag-producing disturbance, but signs of larval activity accumulate over time, so we might expect stronger or opposing effects of disturbance age on adult captures when compared with larval activity. In fact, we found a positive effect of fire age on the combined activity of larval woodborer species, but negative effects of fire age



on captures of adult *M. consputa*, *X. longitarsus*, and *Monochamus* species, as well as all adult buprestids combined. Negative effects of fire age were most evident for *M. consputa*, as expected for members of a genus notably adapted for detecting fires in progress. Emerging adults of species that are attracted to fire might often emigrate, seeking fires in progress and reducing local recruitment that would otherwise tend to obscure a negative relationship between fire age and adult captures. We found negative effects of fire age on both buprestids (especially *M. consputa*) and cerambycids (*Monochamus* species) despite allowing up to eight years of post-fire recruitment, suggesting that there is low incidence of local recruitment in these species. Other woodborers less attracted to heat and smoke might be more likely to colonize trees that succumb to post-fire processes (Angers et al. 2011), such as bark beetle attack.

Finally, the effects of fire ignition date deserve special consideration in further analyses of woodboring beetle ecology. We found surprisingly consistent support across life stages and taxa for negative linear and quadratic relationships between ignition date and woodborer activity, suggesting that there is a temporal peak and decline in the efficacy of woodborer colonization following forest disturbances. The relatively late-season King, Moonlight, and Rim fires provided important support for this apparent effect that should be investigated for generality. Throughout the western US, fire season has lengthened substantially in recent decades (Abatzoglou and Williams 2016; Westerling 2016), due to changes in climate as well as increasing anthropogenic ignitions, which often occur well outside the range of dates associated with lightning ignitions (Balch et al. 2017). This lengthening of the fire season is expected to continue and perhaps even accelerate in the coming decades (Yue et al. 2013, Westerling 2016), yielding more fires that ignite outside the optimal period for driving woodborer population growth and the ecosystem processes that depend on woodborers.

However, we caution that our analysis is based on only 11 fires with a relatively modest spread in ignition dates (5 July to 13 September). Only two of these fires ignited earlier than the interquartile range of lightning ignition dates reported by Balch et al. (2017) for an area that includes the Sierra Nevada, and only two others ignited later than the interquartile range. Furthermore, ignition date is at best a rough index of burn date for the specific areas that we surveyed with our transects. We therefore cannot pinpoint thresholds in ignition date that are likely to yield robust colonization by woodborers, nor can we state with certainty that the apparent effect of ignition date in this study does not stem from another process associated with site. Rather, we view the

pervasive effect of ignition date in our models as a starting point for further research on this topic.

## Conclusions

Woodboring beetle activity at our study sites was often similar between burned stands and unburned stands damaged by bark beetles, but varied dramatically with stand composition, burn severity, and the intensity of bark beetle activity associated with each type of disturbance. In burned sites, the relationship between woodborer activity and ignition date suggested that increased prevalence of wildfire might not increase woodboring beetle populations if fires increasingly occur outside the historical fire season. Effects of these patterns on ecological processes like snag longevity and woodpecker activity should be further explored to inform decisions about management of disturbed forest stands. The patterns observed in this study should be investigated for generality in other regions, and across a diversity of fires and bark beetle outbreaks, to facilitate inference in this system, which naturally exhibits high covariance and spatial clustering in the values of key predictor variables.

## Additional files

**Additional file 1:** Spatial data sources used to guide transect placement in burned (a–f) and bark beetle outbreak (a–e, g) sites in the greater Sierra Nevada region of California, USA, during the summers of 2015 and 2016. (PDF 107 kb)

**Additional file 2:** Correlations (Kendall's tau) among potential metrics of burn severity and fire age, 1 to 8 years post burn, from 11 California wildfires in the greater Sierra Nevada region of California, USA, that burned during the summers of 2015 and 2016. BA = basal area, RdNBR = Relative differenced Normalized Burn Ratio. (PDF 119 kb)

**Additional file 3:** Comparison of the Relative differenced Normalized Burn Ratio (RdNBR) with metrics of burn severity measured in the field (panels a–c), and correlations between burn severity metrics based on snag ratios (d), from 11 burned sites and five bark beetle outbreak (BBO) sites in the greater Sierra Nevada region of California, USA, sampled during the summers of 2015 and 2016. Points represent transect-level averages. BA = basal area. (PDF 591 kb)

**Additional file 4:** Tree species composition (a) and indices of bark beetle activity (b) in bark beetle outbreak (BBO) sites in the greater Sierra Nevada region of California, USA, sampled during the summers of 2015 and 2016. The Reading BBO site differed markedly in tree composition and bark beetle activity, and was also lowest in elevation (see main text, Table 3) and lowest in latitude-adjusted elevation (mean  $\pm$  SE = 1901.10  $\pm$  0.07 m for Reading BBO, 2354.13  $\pm$  0.03 m for other BBO sites). BA = basal area. (PDF 796 kb)

**Additional file 5:** Buprestids (a) and cerambycids (b) identified from unbaited beetle traps operated during June to August of 2015 and 2016 in burned and bark beetle outbreak sites within mixed-conifer montane forests in the greater Sierra Nevada region of California, USA. (PDF 862 kb)

**Additional file 6:** Adult buprestid and cerambycid captures per trap visit averaged by day of year from 11 burned sites and five bark beetle outbreak sites in the greater Sierra Nevada region of California, USA, sampled during the summers of 2015 and 2016. Trends by day of year were not significant after accounting for effects of year. Differences in raw captures per trap visit between years for buprestids (0.82  $\pm$  0.17 in

2015 and  $1.14 \pm 0.11$  in 2016) and cerambycids ( $0.63 \pm 0.08$  in 2015 and  $0.88 \pm 0.09$  in 2016) were not significant after accounting for effects of site. (PDF 1113 kb)

### Acknowledgements

We thank IBP field crews (A. Marquez, D. Mauer, M. Lerow, J. Rivera, C. Seay, M. Seeley, and B. Wiesner) and USFS staff (B. Bualon, M. Easton, A. Eglitis, M. Johnson, R. Mahnke, A. Nelson, K. Ripley, S. Smith, and B. Woodruff) for help with this project, which was funded by USFS Region 5 and the Evaluation Monitoring component of the USFS Forest Health Monitoring program. Manuscript development benefited greatly from two anonymous reviews. This is Contribution No. 624 of The Institute for Bird Populations.

### Funding

Funding for this research was provided by the USDA Forest Service through the Forest Health Monitoring Program (Evaluation Monitoring), Pacific Southwest Region, and Forest Health Protection Program.

### Availability of data and materials

Data and materials used in this study are available from the corresponding author upon reasonable request.

### Authors' contributions

DRC, RLW, RBS, AMW, and CAH conceived and designed the study. DRC, RLW, RBS, AMW, and GLT collected the data. CR analyzed the data and drafted the manuscript with input from DRC, RLW, RBS, AMW, GLT, SCS, and CAH. All authors approved the final manuscript.

### Ethics approval and consent to participate

Not applicable.

### Consent for publication

Not applicable.

### Competing interests

The authors declare that they have no competing interests.

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Received: 21 December 2018 Accepted: 8 May 2019

Published online: 05 July 2019

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