





Key points:

- Mountain pine beetle (MPB)-caused tree mortality increases water yield in most wet
 years, and a decrease in water yield mainly happens in dry years; therefore, interannual
 climate variability is an important driver of water yield response to beetle-caused tree
 mortality.
- A long-term (multi-decade) aridity index is a reliable indicator of water yield response to

 MPBs: in a dry year, decreases occur mainly in "water-limited" areas and vegetation

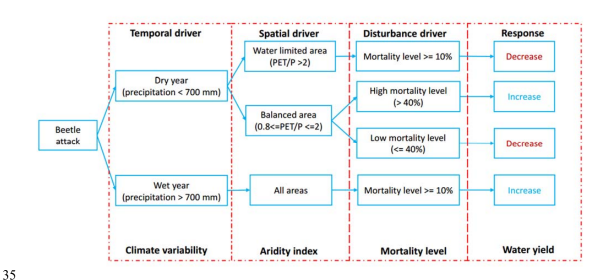
 mortality levels have only minor effects; in wetter areas, decreases only occur at low

 mortality levels.
 - Generally, in a dry year, low to medium MPB-caused vegetation mortality decreases
 water yield, and high mortality increases water yield; this response to mortality level is
 nonlinear and varies by location and year.





34 Graphical abstract



Abstract

Mountain pine beetle (MPB) outbreaks in western United States result in widespread tree mortality, transforming forest structure within watersheds. While there is evidence that these changes can alter the timing and quantity of streamflow, there is substantial variation in both the magnitude and direction of responses and the climatic and environmental mechanisms driving this variation are not well understood. Herein, we coupled an eco-hydrologic model (RHESSys) with a beetle effects model and applied it to a semiarid watershed, Trail Creek, in the Bigwood River basin in central Idaho to evaluate how varying degrees of beetle-caused tree mortality influence water yield. Simulation results show that water yield during the first 15 years after beetle outbreak is controlled by interactions among interannual climate variability, the extent of vegetation mortality, and long-term aridity. During wet years, water yield after beetle outbreak increases with greater tree mortality. During dry years, water yield decreases at low to medium mortality but increases at high mortality. The mortality threshold for the direction of change is





49 location-specific. The change in water yield also varies spatially along aridity gradients during 50 dry years. In relatively wetter areas of the Trail Creek basin, water yield switches from a 51 decrease to an increase when vegetation mortality is greater than 40 percent. In more water-52 limited areas on the other hand, water yield typically decreases after beetle outbreaks, regardless of mortality level. Results suggest that long-term aridity can be a useful indicator for the 53 54 direction of water yield changes after disturbance. 55 1 Introduction 56 In recent decades, mountain pine beetle (MPB) outbreaks in the Western U.S. and Canada have 57 killed billions of coniferous trees (Bentz et al. 2010). Coniferous forests can provide essential 58 ecosystem services, including water supply for local communities (Anderegg et al. 2013). 59 Therefore, it is essential to understand how ecosystems and watersheds respond to beetle 60 outbreaks and to identify the dominant processes that drive these responses (Bennett et al. 2018). A growing number of studies have qualitatively examined hydrologic responses to beetle 61 62 outbreaks and disturbance; however these studies have produced conflicting results (Adams et al. 63 2012; Goeking and Tarboton 2020). While some studies show increases in water yield following beetle outbreak (e.g., Bethlahmy 1974; Potts 1984; Livneh et al. 2015), many others show no 64 65 change or even decreases (e.g., Guardiola-Claramonte et al. 2011; Biederman et al. 2014; Slinski et al. 2016). To determine which mechanisms control change in water yield following beetle 66 67 outbreak, more quantitative approaches are needed. 68 Water yield is often thought to increase after vegetation is killed or removed by disturbances 69 such as fire, thinning, and harvesting (Hubbart 2007; Robles et al. 2014; Chen et al. 2014; Buma 70 and Livneh 2017; Wine et al. 2018). In the Rocky Mountain West, beetle outbreaks have 71 increased water yield through multiple mechanisms. First, defoliation/needle loss can reduce





plant transpiration, canopy evaporation, and canopy snow sublimation losses to the atmosphere 73 (Montesi et al. 2004). Increased canopy openings can also enable snow accumulation and allow 74 more radiation to reach the ground surface, resulting in earlier and larger peak snowmelt events, 75 which can in turn reduce soil moisture and therefore decrease summer evapotranspiration (ET). 76 Several studies have documented decreases in water yield following disturbances (e.g., mortality, 77 fire, beetle outbreaks; Biederman et al. 2014; Bart et al. 2016; Slinski et al. 2016; Goeking and 78 Tarboton 2020). For example, in the southwestern U.S., beetle outbreaks have decreased 79 streamflow by opening forest canopies and increasing radiation to the understory and at the 80 ground surface, which leads to increases in understory vegetation transpiration (Guardiola-81 Claramonte et al. 2011), soil evaporation, and therefore increases total ET (Bennett et al. 2018). 82 Tree mortality or removal can reduce streamflow because surviving trees and/or understory 83 vegetation compensates by using more water (Tague et al. 2019). 84 In a review of 78 studies, Goeking and Tarboton (2020) concluded that the decrease in water 85 yield after tree-mortality mainly happens in semiarid regions. Previous studies also provide rule-86 of-thumb thresholds above which water yield will increase: at least 20 percent loss of vegetation 87 cover and mean precipitation of 500 mm/year (Adams et al. 2012). However, many watersheds 88 in the western U.S. experience high interannual climate variability (Fyfe et al. 2017), and local 89 environmental gradients (e.g., long-term aridity gradients) may strongly influence vegetation and hydrologic responses to disturbances, including beetle outbreaks, making predictions difficult 90 91 (Winkler et al. 2014). Given the possibility of either increases or decreases in water yield 92 following beetle outbreaks, modeling approaches are crucial for identifying the specific 93 mechanisms that control these responses.





94 The overarching goal of this study is to identify mechanisms driving the direction of change in 95 annual water yield after beetle outbreaks in semi-arid regions (note that in the following text, 96 "water yield" refers to means annual water yield). The following specific questions address this 97 goal: 98 **Q1:** What is the role of **interannual climate variability** in water yield response? 99 • Q2: What is the role of mortality level in water yield response? 100 • O3: How does long-term aridity (defined as temporally averaged potential 101 evapotranspiration relative to precipitation for a period of 38 years) modify these 102 responses, and how do responses vary spatially within a watershed along aridity 103 gradients? 104 We hypothesize that multiple ecohydrologic processes (e.g., snow accumulation and melt, 105 evaporation, transpiration, drainage, and a range of forest structural and functional responses to 106 beetles) could interactively influence how water yield responds to beetle outbreaks—however, in 107 certain locations one or more processes may dominate. In addition, the dominant ecohydrologic 108 processes may vary over space and time due to interannual climate variability (i.e., 109 precipitation), vegetation mortality, and long-term aridity. In Sect 2, we present a conceptual 110 framework for identifying and depicting dominant hydrological processes through which forests 111 respond to beetle infestation. We use this framework to interpret the modeling results. In Sect 3, 112 we describe our mechanistic modeling approach, i.e., using the Regional Hydro-Ecological 113 Simulation System (RHESSys), which can prescribe a range of vegetation mortality levels, 114 capture the effects of landscape heterogeneity and the role of lateral soil moisture redistribution,

and project ecosystem carbon and nitrogen dynamics, including post-disturbance plant recovery.





In Sects 4 and 5, we then present modeling results that explore how multiple mechanisms 116 117 influence water yield responses. 118 2 Conceptual framework 119 2.1 Vegetation response to beetle outbreaks 120 Mountain pine beetles (MPB) introduce blue stain fungi into the xylem of attacked trees, which 121 reduces water transport in plants and eventually shuts it off (Paine et al. 1997). During outbreaks, 122 MPBs prefer to attack and kill larger host trees that have greater resources (e.g., carbon), while 123 smaller diameter host trees and non-host vegetation (including the understory) remain unaffected 124 (Edburg et al. 2012). After MBP outbreak, trees mainly go through three phases (i.e., red, gray, 125 and old) over time (Hicke et al. 2012). During the red phase, the trees' needles turn red. During 126 the gray phase, there are no needles in the canopy. During old phase, killed trees have fallen, and 127 understory vegetation and new seedlings experience rapid growth (Hicke et al. 2012; Mikkelson 128 et al. 2013). 129 2.2 Hydrologic response to beetle outbreaks 130 Figure 1 describes the main processes that alter evapotranspiration to either decrease or increase 131 water yield, depending on which processes dominate (Adams et al. 2012; Goeking and Tarboton 132 2020). During the red and gray phases, needles fall to the ground, and there is lower leaf area 133 index (LAI) and a more open canopy (Hicke et al. 2012). This can reduce plant transpiration of 134 infected trees, though remaining trees may compensate to some extent by increasing 135 transpiration in water limited environments (Adams et al. 2012, Tague et al. 2019). A more open 136 canopy intercepts less precipitation, reducing evaporation from the canopy but potentially 137 increasing it from soil and litter layers (Montesi et al. 2004; Sexstone et al. 2018). Meanwhile, an

open canopy can increase the proportion of snow falling to the ground and, therefore, increase



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snowpack accumulation. With more solar radiation reaching the ground, earlier and larger peak snowmelt can also occur (Bennett et al. 2018). Generally, earlier snowmelt increases water for spring streamflow and decreases water for summertime ET (Pomeroy et al. 2012). However, once snags fall, reductions in longwave radiation can actually lead to later snowmelt (Lundquist et al. 2013). The open canopy and less competition for resources, such as solar radiation and nutrients, can also promote understory vegetation growth, which may increase understory transpiration (Biederman et al. 2014; Tague et al. 2019). Whether water yield increases or decreases will ultimately depend on the balance of these processes that can alter transpiration and evaporation in different ways. Finally, interannual variability in climate (e.g., dry versus wet years) can affect forests' hydrological responses (Winkler et al. 2014; Goeking and Tarboton 2020). For instance, during wet years, remaining plants are not water-limited, and reductions in plant transpiration due to beetle-caused mortality dominate increases in soil evaporation or remaining plant transpiration, resulting in a higher water yield. In contrast, during dry years, plants are already under water stress and decreases in plant transpiration caused by tree mortality may be compensated by increasing soil evaporation and transpiration by remaining trees or understory vegetation, leading to declines in water yield. Moreover, these responses are also affected by land cover types (e.g., young vs old pine, different tree species, etc.), which is not currently well documented (Perry and Jones 2017; Morillas et al. 2017). 2.3 Review of modeling approaches Many models, ranging from empirical and lumped to physically-based and fully-distributed, have been used to study hydrologic responses to disturbances. Goeking and Tarboton (2020)





162 argue that only physically-based and fully-distributed models can capture how disturbances alter 163 water yield because they represent fine-scale spatial heterogeneity and physical process that vary 164 over space and time. Despite their advantages, process-based models, such as the coupled CLM-165 ParFlow model (Mikkelson et al. 2013; Penn et al. 2016), the Distributed Hydrology Soil 166 Vegetation Model (Livneh et al. 2015; Sun et al. 2018), and the Variable Infiltration Capacity 167 Model (Bennett et al. 2018) also have some limitations. For example, 1) they may assume 168 constant LAI after disturbances and static vegetation growth (e.g., VIC and DHSVM), 2) they 169 may not include lateral flow to redistribute soil moisture (VIC), and 3) in some cases, the 170 approach to represent the effects of beetle outbreaks may be too simplified (e.g., changing only 171 LAI and conductance without considering two-way beetle-vegetation interactions in post-172 disturbance biogeochemical and water cycling e.g., as in CLM-ParFlow). Thus, improving 173 current fully distributed process-based models to capture the coupled dynamics between 174 hydrology and vegetation at multiple scales is a critical step for projecting how beetle outbreaks 175 will affect water yield in semiarid systems (Goeking and Tarboton 2020). Here we use 176 RHESSys7.1, which captures these processes. 177 3 Model, data, and simulation experiment design 178 3.1 Study area 179 Our study watershed is Trail Creek, which is located in Blaine County between the Sawtooth 180 National Forest and the Salmon-Challis National Forest (43.44N, 114.19W; Fig. 2). It is a 167-181 km² sub-catchment in the south part of Big Wood River basin, and is within the wildland-urban 182 interface where residents are vulnerable to the flood and debris flows caused by forest 183 disturbances (Skinner 2013). Trail Creek has frequently experienced beetle outbreaks, notably in 184 2004 and 2009, when beetles killed 7 and 19 km² of trees, respectively (Berner et al. 2017).





185	Trail Creek has cold, wet winters and warm, dry summers; mean annual precipitation is
186	approximately 978 mm with 60% snow (Frenzel 1989). The soil is mostly permeable coarse
187	alluvium (Smith 1960). Vegetation is clustered into two major groups along the elevation which
188	ranges from 1760 to 3478 m: sagebrush, riparian species, and grasslands in lower to middle
189	elevation areas and Douglas-fir (Pseudotsuga menziesii), lodgepole pine (Pinus contorta var.
190	latifolia), subalpine fir (Abies lasiocarpa), and mixed shrub and herbaceous vegetation in
191	middle-to-higher elevations (Buhidar 2002).
192	A strong upper to lower vegetation and long-term aridity gradient exists for Trail Creek (Fig. 3).
193	The northern (higher elevation) portion of the basin is mesic and covered principally by
194	evergreen forest; the southern (lower elevation) portion is xeric and covered by shrubs, grasses,
195	and mixed herbaceous species. In total, Trail creek contains 72 sub-basins and two of them (e.g.,
196	Fig. 3, sub-basin 412 and 416) are urban areas. If we classify this basin into different zones
197	according to an aridity index, i.e., the ratio of 38-year average annual potential
198	evapotranspiration (PET) to precipitation (P) (Sect 3.4), there is a distinct gradient: the northern
199	and high elevation area is balanced (i.e., PET/P between 0.8 and 2) and evergreen tree coverage
200	is more than 50%; the southern part is water-limited (i.e., $PET/P > 2$) and evergreen tree
201	coverage is less than 30% (Figs. 2 and 3).
202	3.2 Model descriptions
203	3.2.1 Ecohydrologic model
204	The Regional Hydro-ecologic Simulation System (RHESSys) (Tague and Band 2004) is a
205	mechanistic model designed to simulate the effects of climate and land use change on ecosystem
206	carbon and nitrogen cycling and hydrology. RHESSys fully couples hydrological processes
207	(including streamflow, lateral flow, ET, and soil moisture, etc.), plant growth and vegetation



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biogeochemical cycling (including soil organic matter decomposition, mineralization, nitrification, denitrification, and leaching, etc.). It has been widely tested and applied in several mountainous watersheds in western North America, including many in the Pacific and Inland Northwest (e.g., Tague and Band 2004; Garcia and Tague 2015; Hanan et al. 2017; Hanan et al. 2018; Lin et al. 2019; Son and Tague 2019). RHESSys represents a watershed using a hierarchical set of spatial units, including patches, zones, sub-basins, and the full basin, to simulate various hydrologic and biogeochemical processes occurring in these multiple scales (Tague and Band 2004). The patch is the finest spatial scale at which vertical soil moisture and soil biogeochemistry are simulated. In every patch, there are multiple canopy strata layers to simulate the biogeochemical processes related to plant growth and nutrient uptake. Meteorological forcing inputs (e.g., temperature, precipitation, humidity, wind speed, and solar radiation) are handled at the zone level, and spatially interpolated and downscaled for each patch based on elevation, slope, and aspect. Sub-basins are closed drainage areas entering both sides of a single stream reach (the water budget is closed in sub-basins). The largest spatial unit is the basin, which aggregates the streamflow from subbasins (Tague and Band 2004; Hanan et al. 2018). In RHESSys, streamflow is the sum of overland flow and baseflow, and we consider streamflow as the water yield of each sub-basin. RHESSys models vertical and lateral hydrologic fluxes, including canopy interception, plant transpiration, canopy evaporation/sublimation, snow accumulation, snowmelt and sublimation, soil evaporation, soil infiltration, and subsurface drainage. Canopy interception is based on the water-holding capacity of vegetation, which is also a function of plant area index (PAI). Both the canopy evaporation and transpiration are modeled using the standard Penman-Monteith equation

dynamics (including photosynthesis, maintenance respiration, and mortality, etc.), and soil



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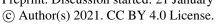
to fall evenly across each zone. Snowmelt is based on a quasi-energy budget approach accounting for radiation input, sensible and latent heat fluxes, and advection. Soil evaporation is constrained by both energy and atmospheric drivers, as well as a maximum exfiltration rate, which is controlled by soil moisture (Tague and Band 2004). Vertical drainage and lateral flow is a function of topography and soil hydraulic conductivity, which decays exponentially with depth (Tague and Band 2004; Hanan et al. 2018). Vegetation carbon and nitrogen dynamics are calculated separately for each canopy layer within each patch, while soil and litter carbon and nitrogen cycling are simulated at the patch level. Photosynthesis is calculated based on the Farquhar model considering the limitations of nitrogen, light, stomatal conductance (which is influenced by soil water availability), vapor pressure deficit, atmospheric CO₂ concentration, radiation, and air temperature (Farquhar and von Caemmerer 1982; Tague and Band 2004). Maintenance respiration is based on Ryan (1991), which computes respiration as a function of nitrogen concentration and air temperature. Growth respiration is calculated as a fixed ratio of new carbon allocation for each vegetation component (Ryan 1991; Tague and Band 2004). Net photosynthesis is allocated to leaves, stems, and roots at daily steps based on the Dickinson partitioning method, which varies with each plant development stage (Dickinson et al. 1998). LAI is estimated from leaf carbon and specific leaf area for each vegetation type. The soil and litter carbon and nitrogen cycling (heterotrophic respiration, mineralization, nitrification, and denitrification, etc.) are modified from the BIOME BGC and CENTURY-NGAS models (White and Running 1994; Parton et al. 1996; Tague and Band 2004). A detailed description of RHESSys model algorithms can be found in Tague and Band (2004).

(Monteith 1965). Snow accumulation is calculated from incoming precipitation and is assumed





254 3.2.2 Beetle effects model 255 Edburg et al. (2012) designed and developed a model of MPB effects on carbon and nitrogen 256 dynamics for integration with the Community Land Model Version 4 (CLM4) (Lawrence et al. 257 2011, Fig. 4). Here we integrated this beetle effects model into RHESSys (Fig. 4). Beetles attack 258 trees mainly during late summer, and needles will turn from green to red at the beginning of the 259 following summer. We simplify this process with prescribed tree mortality on September 1 to 260 represent a beetle outbreak of the current year. The advantage of this integration is that RHESSys 261 accounts for the lateral connectivity in water and nitrogen fluxes among patches which is not 262 represented in CLM4 (Fan et al. 2019). Differences in our approach compared to other 263 hydrological models of beetle effects (e.g., VIC, CLM-ParFlow, and DHSVM) include dynamic 264 changes in plant carbon and nitrogen cycling caused by beetle attack, plant recovery, and effects 265 on hydrological responses. Previous studies of hydrologic effects of beetle outbreaks have 266 mainly focused on consequences of changes in LAI and stomatal resistance during each phase of 267 beetle outbreak but have missed feedbacks between carbon and nitrogen dynamics, vegetation 268 recovery, and hydrology (Mikkelson et al. 2013; Livneh et al. 2015; Penn et al. 2016; Sun et al. 269 2018; Bennett et al. 2018). 270 To better represent the effects of beetle-caused tree mortality, we added a snag pool (standing 271 dead tree stems) and a dead foliage pool (representing the red needle phase) in RHESSys (Fig. 272 4). All leaf biomass (including carbon and nitrogen) become part of dead foliage pools. After one 273 year (Hicke et al. 2012; Edburg et al. 2011) the dead foliage is transferred to litter pools at an 274 exponential rate with a half-life of two years (Edburg et al. 2012). Similarly, stem carbon and 275 nitrogen are moved to the snag pool immediately after outbreak. After five years (Edburg et al. 276 2012), carbon and nitrogen in snags begin to move into the coarse woody debris (CWD) pool at







277 an exponential decay rate with a half-life of ten years (Edburg et al. 2011). After outbreak, the 278 coarse root pools that are killed move to the CWD and fine root pools move to litter pools. To 279 simplify, we assume a uniform mortality level for all evergreen patches across landscape. Due to 280 the limitation of land cover data, we cannot separate pine and fir in these evergreen patches. 281 However, this will not affect the interpretation of our results because we analyze them based on 282 mortality level and evergreen vegetation coverage rather than different species. 283 In the integrated model, the reduction of leaf carbon and nitrogen after beetle outbreak can 284 directly decrease LAI and canopy height, which consequently affects energy (i.e., longwave 285 radiation and the interception of shortwave radiation) and hydrologic (i.e., transpiration and 286 canopy interception) fluxes. We calculate two types of LAI: Live LAI (i.e., only live leaf is 287 included), and Total LAI (i.e., both live and dead leaves are included). The calculation of plant 288 transpiration is based on Live LAI, while the calculation of other canopy properties, including 289 interception and canopy evaporation, is based on Total LAI. The calculation of canopy height 290 includes the living stem and the snag pool. 291 3.3 Input data 292 We used the US Geologic Survey (USGS) National Elevation Dataset (NED) at 10 m resolution 293 to calculate the topographic properties of Trail Creek, including elevation, slope, aspect, basin 294 boundaries, sub-basins, and patches. Using NED, we delineated 16705 100-m resolution patches 295 within 72 sub-basins. We used the National Land Cover Database (NLCD) to identify five 296 vegetation and land cover types, i.e., evergreen, grass/herbaceous, shrub, deciduous, and urban 297 (Homer et al. 2015). We determined soil properties for each patch using the POLARIS database 298 (probabilistic remapping of SSURGO; Chaney et al. 2016). Parameters for soil and vegetation





299 were based on previous research and literature (White et al. 2000; Law et al. 2003; Ackerly 300 2004; Berner and Law 2016; Hanan et al. 2016). 301 Climate inputs for this study, including maximum and minimum temperatures, precipitation, 302 relative humidity, radiation, and wind speed, were acquired from gridMET for years from 1980 303 to 2018. GridMET provides daily high-resolution (1/24 degree or ~4 km) gridded meteorological 304 data (Abatzoglou 2013). It is a blended climate dataset that combines the temporal attributes of 305 gauge-based precipitation data from NLDAS-2 (Mitchell et al. 2004) with the spatial attributes of 306 gridded climate data from PRISM (Daly et al. 1994). 307 3.4 Simulation experiments 308 To quantify how water yield responds to beetle-caused mortality, we designed the following 309 simulation experiment. We prescribed a beetle outbreak in September 1989, the mortality level 310 (%) is applied to all evergreen patches for each sub-basin. After beetle outbreak, red needles stay 311 on the trees for one year before they start to fall (transferred to the litter pool) at an exponential 312 rate with a half-life of two years. The snag pools stay in the standing trees for five years and then 313 start to fall and are added to the CWD pool which decays at an exponential rate with a half-life 314 of ten years. 315 To address Q1 (i.e., the role of interannual variability), we compared water yield responses 316 during a dry water year, 1994 (i.e., five years after beetle outbreak with precipitation 611 mm), 317 to responses during a wet year, 1995 (i.e., six years after beetle outbreak with precipitation 1394 318 mm). This enabled us to estimate the role of interannual climate variability in driving changes in 319 water yield following beetle attack. The dry year are selected based on years that have precipitation below the 15th percentile across 38 years of annual precipitation data (from 1979 to 320 321 2017) (Searcy 1959, see Fig. S1). During these early period after beetle outbreak (e.g., 1994 and





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1995) the forest is experiencing large changes in vegetation canopy cover, plant transpiration, and soil moisture. We chose these two successive years because they have almost similar canopy and vegetation status in terms of fallen dead foliage and residual vegetation regrowth, which makes this comparison reasonable. However, it is possible that antecedent climate conditions may affect the following year's response. For example, soil moisture can be depleted during a drought year, affecting initial conditions the following year. Moreover, under drought conditions, less reactive nitrogen is taken up by the plants or leaching is reduced, so more nitrogen will be left for the following year. Therefore, the difference in water yield responses between 1994 and 1995 might be affected by not only climate variations but also initial conditions in the hydrology and the biogeochemistry. To consider the time lag effect (antecedent conditions affecting the current year's response), we also analyzed other dry and wet years. To address Q2 (i.e., the role of vegetation mortality), we prescribe a range of infestation-caused mortality levels (i.e., from 10% to 60% by a step of 10% in terms of carbon, uniformly applied to all evergreen patches for each sub-basins) and a control run (no mortality) to quantify the response of forests in water yield to vegetation mortality level (for each sub-basin vegetation mortality is evergreen mortality multiplied by evergreen coverage of that basin). The differences in water yield between each mortality level and the control run represent the effects of beetle kill: a positive value means that mortality increased water yield, and vice versa. We quantified the water budget for each sub-basin to examine which hydrological process contribute to the water yield responses: water yield (Q), precipitation (P), canopy evaporation (Ecanopy, canopy evaporation and snow sublimation), transpiration (T), ground evaporation (Eground, includes bare soil evaporation, pond evaporation, and litter evaporation), snow sublimation (Sublim, ground), soil storage change (${}^{dS_{soil}}/_{dt}$), litter storage change





- 345 $({}^{dS_{litter}}/_{dt})$, snowpack storage change $({}^{dS_{snowpack}}/_{dt})$ and canopy storage change
- 346 $\left(\frac{dS_{canopy}}{dt}\right)$.
- 347 The storage components include soil, litter, and canopy. According to Eq. (1), if the storage
- increases, water yield decreases.

$$Q = P - E_{canony} - E_{around} - Sublim -$$

$$T - \frac{d(S_{soil} + S_{litter} + S_{canopy} + S_{snowpack})}{dt} / dt$$
 (1)

- 351 Calculating water balance differences between different mortality scenarios and control scenario
- 352 results in Eq. (2):

$$\Delta Q = \Delta E_{canopy} + \Delta E_{ground} + \Delta Sublim + \Delta T +$$

$$\Delta \left(\frac{d(S_{soil} + S_{litter} + S_{canopy} + S_{snowpack})}{dt} \right)$$
 (2)

- To address Q3 (i.e., the role of long-term aridity), we calculated the long-term aridity index
- 356 (PET/P, Fig. 3) across the basin and analyzed the relationship between long-term aridity index
- and hydrologic response. As mentioned earlier, the long-term aridity index is defined as the ratio
- of mean annual potential ET (PET) to annual precipitation (P), averaged over 38 years (water
- year 1980-2018) of historical meteorological data. Based on the long-term aridity index, we
- 360 classified our sub-basins into three types (McVicar et al. 2012, Table 1).





362 4 Results 363 4.1 Simulated vegetation response to beetle outbreak at basin-scale 364 4.1.1 Vegetation response to beetle outbreaks 365 Figure 5 shows the basin-scale vegetation response after beetle outbreak in 1989. Live LAI 366 dropped immediately after beetle outbreak, then gradually recovered to pre-outbreak levels 367 during following years (Fig. 5a). Total LAI (i.e., including dead foliage) showed a slight increase 368 during the first ten years after beetle outbreak (1990 – 2000), which is due to the retention of 369 dead leaves in the canopy and the simultaneous growth of residual (unaffected) overstory and 370 understory vegetation (Fig. 5b). The dead foliage pool (Fig. 5c) remained in place for one year 371 and then began to fall to ground (converted to litter) exponentially with a half-life of two years, 372 and the snag pool (Fig. 5d) remained in place for five years and then began to fall to ground 373 (converted to CWD) exponentially with a half-life of ten years. These behaviors of the dead 374 foliage and snag pools are similar to Edburg et al. (2012), which demonstrates that the integrated 375 model is simulating expected vegetation dynamics following beetle outbreak. 376 4.1.2 Time series of hydrologic response to beetle outbreak 377 Figure 6 shows the changes in simulated water fluxes and soil moisture over the basin after 378 beetle outbreak with various evergreen mortality levels. During the first 15 years after beetle 379 outbreak, scenarios where the evergreen mortality level was larger than zero had higher basin-380 scale water yield than the control scenario (where the evergreen mortality level was zero). This 381 was especially true during wet years; however, there was no significant increase during dry years 382 (i.e., 1992, 1994, 2001, and 2004; Fig. 6a). The year-to-year soil storage fluxes responded 383 strongly in the first two years after beetle outbreak, then stabilized to the pre-outbreak condition 384 (Fig. 6b). Note that year-to-year soil storage change is not the same as soil water storage. After





385 beetle outbreak, the soil can hold some portion of water that not being up taken by the plants, but 386 it was confined by the soil water holding capacity. This phenomenon indicates that the soil has 387 some resilience to vegetation change. 388 Beetle outbreaks reduced transpiration during wet years but did not have significant effects in 389 dry years (Fig. 6c). This is because transpiration in dry years was water-limited and so was much 390 lower than the potential rate (more water is partitioned to evaporation; Biederman et al. 2014). 391 Thus, killing more trees had little effect on stand scale transpiration because remaining trees 392 utilized any water released by the dead trees in dry years. On the other hand, plant transpiration 393 in wet years was close to the potential rate; therefore, decreases in canopy cover reduced 394 transpiration. The simulation results did not show any apparent effect on snowmelt after beetle 395 outbreak. 396 The evaporation response was opposite in dry and wet years: evaporation increased in dry years, 397 while it decreased in wet years (Fig. 6d). This phenomenon is caused by tradeoffs and 398 interactions among multiple processes, as will be explained in more detail in the next section. 399 4.2 The role of spatial heterogeneity in water yield response 400 4.2.1 Spatial patterns of hydrologic response along long-term aridity gradient 401 4.2.1.1 Evaporation 402 Beetle outbreak had opposite effects on evaporation between a dry year and a wet year (Fig. 7). 403 In the dry year, most sub-basins experienced higher evaporation for beetle outbreak scenarios 404 than in the control scenario (Fig. 7a). This was the cumulative consequence of decreased canopy 405 evaporation and increased ground (soil, litter, pond) evaporation due to decreases in LAI (caused 406 by mortality). In the dry year, the latter effect (i.e., increased ground evaporation) dominated



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4.2.1.3 Total ET



over the former effect so that overall consequence was increased evaporation. When the vegetation mortality level (calculated as the percentage of evergreen patches in a sub-basin multiplied by the mortality level of evergreen caused by beetles) was higher than 20%, a few subbasins in the balanced (more mesic) area showed some decrease, indicating that the effects of decreasing canopy evaporation exceeded the effects of increasing ground evaporation. In the wet year, most of the sub-basins located in the balanced area showed decreases in evaporation, and the decreasing trend showed linear relationship with vegetation mortality level (where canopy evaporation decreases are dominant, Fig. 7b). However, sub-basins located in much drier regions (aridity >3.5) had relatively insignificant responses to vegetation mortality levels and some of them even had slight increases in evaporation (where ground evaporation increases are dominant due to drier long-term climate and less pine coverage resulted in lower canopy mortality). 4.2.1.2 Transpiration Beetle outbreak decreased transpiration in both dry and wet years, and with higher mortality levels the decrease became larger (Fig. 8). However, during the dry year, the water-limited area showed less change than the balanced area; some sub-basins even showed slight increases. This increase in the water-limited part of the basin occurred because after beetles kill some overstory evergreen trees, the living trees and understory plants together can exhibit higher transpiration rates in dry years (Tsamir et al. 2019). In the wet year, when most canopies reach potential transpiration rates (less competition for water), beetle outbreaks can reduce transpiration rates by decreasing Live LAI.



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Figure 9 depicts the spatial pattern of changes in total ET (i.e., evaporation and transpiration) after beetle outbreak. In a dry year, the balanced and water-limited areas showed opposite responses to mortality: the balanced area showed a decrease in ET and the water-limited area showed a slight increase. In the balanced area, larger ET decreases occurred with higher mortality levels. However, increases in ET in water-limited regions were less sensitive to vegetation mortality level, and even for high vegetation mortality levels (>40%), ET still increased (Fig. 9a). During the wet year, most sub-basins experienced decreasing ET after beetle outbreak and the magnitude was larger with higher vegetation mortality. The different responses of ET were driven by different hydrologic responses (transpiration, ground evaporation and canopy evaporation) competing with each other; this competition was influenced by climate conditions, mortality level, and spatial heterogeneity in long-term aridity. 4.2.1.4 Water yield In the dry year (1994), beetle-caused vegetation mortality affected water yield (Fig. 10), but the responses differed between the balanced and water-limited areas. For the balanced area, most sub-basins showed slight decreases in water yield after beetle outbreak and no significant differences among low vegetation mortality level (<=40%, Fig. 10a). However, with increased mortality levels, more sub-basins showed increases in water yield, particularly with vegetation mortality higher than 40% (Fig. 10a). Moreover, the vegetation mortality threshold that changed the direction of water yield response was altered by long-term aridity, e.g., it was 40% for aridity 2.0 but 20% for aridity 1.0. For the water-limited area, water yield decreased and was independent from mortality level (Fig. 10a). In the wet year (1995), the water yield in most subbasins increased after beetle outbreak, and the balanced area increased more significantly than the water-limited area. Furthermore, for the balanced area, higher mortality levels caused larger





451 increases in water yield which responded more linearly (Fig. 10b). In summary, for a wet year, 452 increases in water yield occurred for most sub-basins, driven by a decrease in ET. However, 453 during dry years, the water yield and ET responses were spatially heterogeneous, and the 454 competing changes in evaporation and transpiration changed the direction and magnitude of ET 455 and thus water yield response. The competing effect among different hydrologic fluxes for a dry 456 year is explored in more detail in the next section. 457 4.2.2 Water budgets to understand decreasing water yield in the dry year 458 We analyzed the fluxes in greater detail in a dry year (1994) to understand the response of 459 hydrologic fluxes and resulting water yield. Based on Eq. (2), we identified four hydrological 460 fluxes that can potentially affect water yield: canopy evaporation (canopy evaporation and 461 canopy snow sublimation), ground evaporation (bare soil evaporation, ground snow sublimation, 462 litter evaporation, pond evaporation), plant transpiration, and year-to-year storage change (soil, 463 canopy, litter, snowpack). These three storage terms (canopy, litter, snowpack) were considered 464 together with soil storage since their contribution was minor in comparison with other fluxes. 465 Figure 11 summarized different combinations of these four dominate processes during the dry 466 year (1994) based on their directions (increase or decrease in water yield) after beetle outbreak. 467 In total, fourteen combinations of changes in these fluxes (referred to as "response types") were 468 found. Five of them resulted in an increase in water yield, and the others resulted in a decrease. 469 Water yield responses caused by the competition of different hydrologic fluxes showed different 470 patterns across the aridity gradient (Figs. 3&10). For the balanced area (upper part of the basin), 471 with low evergreen mortality (<=30%), the major response types were D1 and D2, in which the 472 increase in ground evaporation dominated over the decrease in transpiration and canopy 473 evaporation (Fig. 11a, b, and c). However, with higher evergreen mortality (>30%), the major





474 response type became W2, where the increase in ground evaporation did not exceed the decrease 475 in canopy evaporation and transpiration (Fig. 11e, f, and g). This indicates that, in a dry year, 476 when more evergreen stands are killed, the increase in ground evaporation reaches a limit while 477 transpiration and canopy evaporation continue to decrease with decreasing LAI. The increase in 478 ground evaporation was triggered either by decreased Total LAI and open canopy, which allowed more solar radiation penetration to the ground for evaporation (Fig. S5c), or less 479 480 transpiration from plants, which left more water available to evaporate (Fig. 8a). The decrease in 481 plant transpiration and canopy evaporation was driven by a lower Live LAI and a lower Total 482 LAI, respectively (Fig. S5 a&c and Fig. 8a). 483 The decrease in water yield in the water-limited area (lower part of the basin) was driven by 484 different hydrologic flux competitions in different mortality levels. When evergreen stand 485 mortality level was low (<=30%), the response types were D5 and D7, in which the increase in 486 ground and canopy evaporation dominated over the decrease of transpiration (Fig. 11a, b, and c). 487 However, with high evergreen stand mortality (>30%), the response types became D1 and D2 488 (Fig. 11e, f, and g), in which the canopy evaporation changed from an increase to a decrease that 489 was driven by a decrease in Total LAI (Fig. S5c). When mortality was low, the increases in 490 growth from residual plants and understory outstripped the litter fall of dead foliage; thus, Total 491 LAI increased, and vice versa when mortality was high. 492 **5 Discussion** 493 5.1 Role of interannual climate variability 494 During the first 15 years after beetle attack, various hydrologic processes opposed and/or 495 reinforced one another to either increase or decrease water yield: a decrease in Live LAI can 496 reduce transpiration, while a decrease in Total LAI can enhance ground evaporation but diminish







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canopy evaporation (Montesi et al. 2004; Tsamir et al. 2019). Interannual climate variability played an important role in determining which of these competing effects dominate and, therefore, drove the direction of water yield response to beetle outbreak (Winkler et al. 2014; Goeking and Tarboton 2020). Our results show that mainly decreases in water yield occurred in dry years, while increases occurred in wet years. During a wet year, plant ET can reach its potential so that any reductions in actual plant ET will dominate over any increases in ground evaporation, resulting in a net increase in water yield. During a dry year, the relative dominance of these competing effects had greater spatial heterogeneity because the water stress status of the plants varied across the basin (as explained in Sect 4.2.2; Fig. 11). However, the responses we observed in the dry year (1994) and in the wet year (1995) were also affected by the previous year's climate (mainly precipitation) and its effects on hydrologic and biogeochemical processes, which set the initial conditions for the dry and wet year (e.g., soil moisture, nitrogen availability, etc.). Therefore, we also analyzed other water years during the first ten years after beetle outbreak to examine whether our findings for dry and wet years follow a general pattern and to what extent they are influenced by antecedent conditions. Results indicate that our findings are robust through the study time period. For example, water yield generally decreased during dry years (1992, 1994, and 2001, see Figs. S1 and S2) and always increased during wet years (1993 and from 1995 to 2000, see Fig. S1 and S2). Adams et al. (2012) provide a threshold of precipitation under which water yield increases after disturbances: at least 500 mm/year (Goeking and Tarboton 2020). The average annual precipitation over this study basin is 600-900 mm in dry years, and higher than 900 mm in wet years. Recent field work observation also find annual climate variability can affect the magnitude of evapotranspiration fluxes that change the water yield direction (Biederman et al. 2014). Our





521 which tree removal increases water yield (Figs. 10, S1 and S2). 522 5.2 Role of vegetation mortality 523 Vegetation mortality is another important factor that influences water yield response. We found 524 that during the wet year, beetle outbreak increased water yield across the basin and the 525 magnitude of these increases grew linearly with the level of vegetation mortality (Fig. 10b). In 526 the dry year, however, the response of water yield to the level of vegetation mortality was more 527 complicated because mortality influenced not only the magnitude of change but also the 528 direction (Fig. 10a). These opposing results (due to mortality level) mainly occurred in the 529 "balanced" northern part of the basin, where the competing effects of mortality (i.e., increases in 530 ground evaporation versus decreases in transpiration) are more balanced (Fig. 11). The level of 531 vegetation mortality played a less significant role in changing water yield in the southern "water-532 limited" area. Vegetation mortality level determined the magnitudes of Live LAI, Total LAI, 533 transpiration, canopy evaporation, and ground evaporation in such a way that it governed the 534 direction of change in both ET and water yield. Thus, when vegetation mortality level was higher 535 than 40%, its effect of decreasing transpiration became the dominant process and its effect of 536 increasing soil evaporation became minor (Fig. 11 f&g; Guardiola-Claramonte et al. 2011). 537 Besides the precipitation threshold of at least 500 mm/year, Adams et al. (2012) also estimate 538 that when at least 20% of vegetation cover is removed, water yield can increase. According to 539 previous analysis (Sect 4.1), for a dry year, water yield increases when more than 40% of 540 vegetation is removed (Fig. 10a). Our model simulations indicate similar mortality thresholds 541 exist for driving water yield increases during the dry year, however, we did not find evidence 542 that such a threshold exists during wet years. These differences between dry and wet years

results corroborate these earlier studies by revealing that there are precipitation thresholds above





suggest that the effects of mortality on water yield depend on climate variability. Other studies 543 544 corroborate this finding by demonstrating that the relationship between mortality level and water 545 yield response is complicated and nonlinear (Moore and Wondzell 2005). 546 5.3 Role of long-term aridity index (PET/P) 547 Long-term aridity indices can be used to predict where water yield will decrease after 548 disturbance. We found that water yield always increased in a wet year, irrespective of the 549 climatic aridity index (Fig. 10a). For dry years, long-term aridity index became important in 550 driving the direction of water yield responses to beetle outbreak. In areas that are less water-551 limited (balanced areas), the direction of water-yield responses to beetle outbreak in a dry year 552 was mixed and depended on mortality level. For water-limited areas, in a dry year, water yield 553 showed a more consistent decrease trend, and it was also less affected by mortality level. These 554 results agree with previous studies finding that water yield decreases largely happen in semiarid areas (Guardiola-Claramonte et al. 2011; Biederman et al. 2014). 555 556 The decrease in water yield for water-limited area can be driven by increases in canopy 557 evaporation or transpiration, which were different in the hydrologically-balanced area (driven by 558 increase of ground evaporation). There, the increase in canopy evaporation was due to an 559 increase in total LAI which is a combined effect of delayed decay of dead foliage and fast 560 growth of residual and understory plants (Fig. 11d type D5, D7, D8 & D9; Fig. S5). The 561 surviving and understory plants in the water-limited area can also have higher transpiration rates 562 after mortality (Fig. 11d type D6 and Fig. 8). Similarly, in field studies, Tsamir et al. (2019) 563 found an increase in photosynthesis and transpiration after thinning in a semi-arid forest. These 564 findings illustrate that in addition to top-down climate variability, the long-term aridity index





565 (which also varies with bottom-up drivers such as vegetation and local topography) can be 566 another useful indicator of how water yield will respond to disturbances. 567 5.4 Uncertainties 568 While our findings revealed how topoclimatic gradients influenced water yield responses to 569 beetle infestation, some uncertainties remain. For one, we used uniform mortality levels for all 570 patches across the watershed rather than location and vegetation-specific mortality levels. 571 However, in reality beetles usually attack older trees first (Edburg et al. 2011). Thus, 572 incorporating a more mechanistic understanding of beetle attack patterns with our beetle effects 573 model could enable us to simulate more realistic outbreak scenarios moving forward. Another 574 source of uncertainty came from the model treatment of litter pools. In the current 575 implementation, we ignored the effects of litter on ground albedo and snowmelt (Lundquist et al. 576 2013), which could have an effect on rates of AET and PET and therefore our calculated long-577 term aridity index. Also, because we focused on water yield responses during the first 15 years 578 after beetle outbreak, we may have missed some of the long-term effects (e.g., after the 579 ecosystem has begun to recover) on forest hydrology. Future research should integrate the short-580 term and long-term effects and interactions among beetle outbreak, vegetation dynamics, and 581 hydrology. Since Trail Creek is either "balanced" or "water-limited" in terms of aridity, other 582 "energy-limited" regions could also be investigated. 583 **6 Conclusion** 584 We tested a coupled ecohydrologic and beetle effects model in a semi-arid basin in southern 585 Idaho to examine how watershed hydrology responds to beetle outbreak and how interannual 586 climatic variability, vegetation mortality, and long-term aridity influence these responses. 587 Simulation results indicate that each factor can play a discrete role in driving hydrological





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processes (e.g., the direction and magnitude of changes in plant transpiration, canopy and soil evaporation, soil and litter moisture, snow sublimation, etc.). These combined effects determine the overall water budget and water yield of the basin. While interannual climate variability is the key factor driving the direction of change in water yield, vegetation mortality levels and longterm aridity modify water yield responses. In dry years, the water yield of most sub-basins slightly decreased after beetle outbreak when vegetation mortality level was lower than 40%; while during wet years in most sub-basins it increased. Our results show that long-term aridity index is a reliable indicator of the water yield decreases that occur during dry years due to the fact that there is a consistent decrease in water yield in the most water-limited portion of the basin. Generally, the effects of vegetation mortality on water yield during dry years is less uniform and depends on local, long-term aridity conditions. During wet years, on the other hand, mortality typically causes increases in water yield. This illustrates that together interannual climate variability and mortality can have a stronger effect on the direction of water yield response in water-limited regions than interannual climate variability alone. Future studies to predict water yield response to disturbance should consider the interactions of these factors and capture the fluctuations of competing water fluxes and storage change that control overall water budget and water yield. Using our novel RHESSys-beetle effects modeling framework, we demonstrate that the direction of hydrologic response is a function of multiple factors (e.g., interannual climate variability, vegetation mortality level, and long-term aridity) and that these results do not necessarily conflict with each other but are representative of different conditions. The mechanisms behind these changes compete with each other resulting in a water yield increases or decreases (Fig. 1). Contradictory findings in previous studies may result from differing mortality levels (disturbance





611 severity), or differences in aridity, and consequently, the emergent drivers that dominate water 612 yield responses differ. Disentangling these drivers is difficult or impossible using a purely 613 empirical approach where it can be challenging or cost-prohibitive to experiment under a broad 614 range of controlled conditions. Distributed process-based models on the other hand, provide a 615 useful tool for examining these dynamics. 616 Findings from this study can assist water supply stakeholders in risk management in beetle 617 outbreak locations. For example, during wet years, more attention might be focused on 618 "balanced" areas, i.e., wet regions, for flooding and erosion risks after beetle outbreaks since 619 these regions may experience large increase in runoff due to decreases in plant transpiration and 620 increases in soil moisture. During the dry years, attention might need to shift to "water-limited" 621 areas for managing wildfire risk since these regions will experience elevated ET and lower soil and litter moisture. Because multiple factors interact to influence hydrological processes after 622 623 beetle outbreak, water and forests management must respond to spatial and temporal variations 624 in climate, aridity, and vegetation mortality levels. 625 Code and data availability 626 The coupled RHESSys model code is available online at: 627 https://github.com/renjianning/RHESSys/tree/historical fire 628 The data used in this study are available at: 629 https://osf.io/tsu9z/?view_only=72bfa7b376ad40c59278312f49b03a69 630 **Author contributions** 631 JR, JA and JAH conceived of study. JR designed study with support from JA, JAH and EH. JR 632 and EH developed RHESSys code for coupling beetle effect model and parallelizing model runs





633 with help from JA, JAH, NT, ML, CK, and JTA. JR performed model simulations and developed 634 figures with help from all authors. ML and JTA generated downscaled meteorological data. JR 635 wrote manuscript with input from all authors. 636 **Competing interests** 637 The authors declare that they have no conflict of interest. 638 Acknowledgments 639 This project is supported by National Science Foundation of United States under award number 640 DMS-1520873. 641



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References

- Ackerly, David D. 2004. "Adaptation, Niche Conservatism, and Convergence: Comparative Studies of Leaf Evolution in the California Chaparral." *The American Naturalist* 163 (5): 654–71. https://doi.org/10.1086/383062.
- Adams, Henry D., Charles H. Luce, David D. Breshears, Craig D. Allen, Markus Weiler, V.
 Cody Hale, Alistair M. S. Smith, and Travis E. Huxman. 2012. "Ecohydrological
 Consequences of Drought- and Infestation- Triggered Tree Die-off: Insights and
 Hypotheses." *Ecohydrology* 5 (2): 145–59. https://doi.org/10.1002/eco.233.
- Anderegg, William R. L., Jeffrey M. Kane, and Leander D. L. Anderegg. 2013. "Consequences of Widespread Tree Mortality Triggered by Drought and Temperature Stress." *Nature Climate Change* 3 (1): 30–36. https://doi.org/10.1038/nclimate1635.
 - Baret, F., A. Olioso, J. L. Luciani, J. F. Hanocq, and J. C. Monterrot. 1989. "Estimation à partir de mesures de réflectance spectrale du rayonnement photosynthétiquement actif absorbé par une culture de blé." *Agronomie* 9 (9): 885–95. https://doi.org/10.1051/agro:19890906.
- Bart, Ryan R., Christina L. Tague, and Max A. Moritz. 2016. "Effect of Tree-to-Shrub Type
 Conversion in Lower Montane Forests of the Sierra Nevada (USA) on Streamflow."
 Edited by Julia A. Jones. PLOS ONE 11 (8): e0161805.
 https://doi.org/10.1371/journal.pone.0161805.
- Bennett, Katrina E., Theodore J. Bohn, Kurt Solander, Nathan G. McDowell, Chonggang Xu,
 Enrique Vivoni, and Richard S. Middleton. 2018. "Climate-Driven Disturbances in the
 San Juan River Sub-Basin of the Colorado River." *Hydrology and Earth System Sciences* 22 (1): 709–25. https://doi.org/10.5194/hess-22-709-2018.
- Bentz, Barbara J., Jacques Régnière, Christopher J Fettig, E. Matthew Hansen, Jane L. Hayes,
 Jeffrey A. Hicke, Rick G. Kelsey, Jose F. Negrón, and Steven J. Seybold. 2010. "Climate
 Change and Bark Beetles of the Western United States and Canada: Direct and Indirect
 Effects." BioScience 60 (8): 602–13. https://doi.org/10.1525/bio.2010.60.8.6.
 - BERNER, L. T., B. E. LAW, A. J. MEDDENS, and J. A. HICKE. 2017. "Tree Mortality from Fires and Bark Beetles at 1-Km Resolution, Western USA, 2003-2012." Collection. Tree Mortality from Fires and Bark Beetles at 1-Km Resolution, Western USA, 2003-2012. 2017. https://doi.org/10.3334/ornldaac/1512.
- Berner, Logan T., and Beverly E. Law. 2016. "Plant Traits, Productivity, Biomass and Soil
 Properties from Forest Sites in the Pacific Northwest, 1999–2014." Scientific Data 3 (1):
 1–14. https://doi.org/10.1038/sdata.2016.2.
- 675 Bethlahmy, Nedavia. 1974. "More Streamflow after a Bark Beetle Epidemic." *Journal of Hydrology* 23 (3): 185–89. https://doi.org/10.1016/0022-1694(74)90001-8.
- Biederman, J. A., A. A. Harpold, D. J. Gochis, B. E. Ewers, D. E. Reed, S. A. Papuga, and P. D.
 Brooks. 2014. "Increased Evaporation Following Widespread Tree Mortality Limits
 Streamflow Response." *Water Resources Research* 50 (7): 5395–5409.
 https://doi.org/10.1002/2013WR014994.
- Buhidar, Balthasar. 2002. "The Big Wood River Watershed Management Plan."
 https://www.deq.idaho.gov/media/450316-
- _water_data_reports_surface_water_tmdls_big_wood_river_big_wood_entire.pdf.
- Buma, Brian, and Ben Livneh. 2017. "Key Landscape and Biotic Indicators of Watersheds Sensitivity to Forest Disturbance Identified Using Remote Sensing and Historical



693

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695



- 686 Hydrography Data." *Environmental Research Letters* 12 (7): 074028. https://doi.org/10.1088/1748-9326/aa7091.
- Chaney, Nathaniel W., Eric F. Wood, Alexander B. McBratney, Jonathan W. Hempel, Travis W.
 Nauman, Colby W. Brungard, and Nathan P. Odgers. 2016. "POLARIS: A 30-Meter
 Probabilistic Soil Series Map of the Contiguous United States." *Geoderma* 274 (July):
 54–67. https://doi.org/10.1016/j.geoderma.2016.03.025.
 - Chen, Fei, Guo Zhang, Michael Barlage, Ying Zhang, Jeffrey A. Hicke, Arjan Meddens, Guangsheng Zhou, William J. Massman, and John Frank. 2014. "An Observational and Modeling Study of Impacts of Bark Beetle–Caused Tree Mortality on Surface Energy and Hydrological Cycles." *Journal of Hydrometeorology* 16 (2): 744–61. https://doi.org/10.1175/JHM-D-14-0059.1.
- Daly, Christopher, Ronald P. Neilson, and Donald L. Phillips. 1994. "A Statistical-Topographic
 Model for Mapping Climatological Precipitation over Mountainous Terrain." *Journal of Applied Meteorology* 33 (2): 140–58. https://doi.org/10.1175/1520-0450(1994)033<0140:ASTMFM>2.0.CO;2.
- Dickinson, Robert E., Muhammad Shaikh, Ross Bryant, and Lisa Graumlich. 1998. "Interactive
 Canopies for a Climate Model." *Journal of Climate* 11 (11): 2823–36.
 https://doi.org/10.1175/1520-0442(1998)011
- Edburg, Steven L., Jeffrey A. Hicke, Paul D. Brooks, Elise G. Pendall, Brent E. Ewers, Urszula
 Norton, David Gochis, Ethan D. Gutmann, and Arjan JH Meddens. 2012. "Cascading
 Impacts of Bark Beetle-Caused Tree Mortality on Coupled Biogeophysical and
 Biogeochemical Processes." Frontiers in Ecology and the Environment 10 (8): 416–24.
 https://doi.org/10.1890/110173.
- Edburg, Steven L., Jeffrey A. Hicke, David M. Lawrence, and Peter E. Thornton. 2011.
 "Simulating Coupled Carbon and Nitrogen Dynamics Following Mountain Pine Beetle
 Outbreaks in the Western United States." *Journal of Geophysical Research:* Biogeosciences 116 (G4): G04033. https://doi.org/10.1029/2011JG001786.
- Fan, Y., M. Clark, D. M. Lawrence, S. Swenson, L. E. Band, S. L. Brantley, P. D. Brooks, et al.
 2019. "Hillslope Hydrology in Global Change Research and Earth System Modeling."
 Water Resources Research 0 (0). https://doi.org/10.1029/2018WR023903.
- Farquhar, G. D., and S. von Caemmerer. 1982. "Modelling of Photosynthetic Response to
 Environmental Conditions." In *Physiological Plant Ecology II: Water Relations and* Carbon Assimilation, edited by O. L. Lange, P. S. Nobel, C. B. Osmond, and H. Ziegler,
 549–87. Encyclopedia of Plant Physiology. Berlin, Heidelberg: Springer.
 https://doi.org/10.1007/978-3-642-68150-9
 17.
- Frenzel, Steven A. 1989. "Water Resources of the Upper Big Wood River Basin, Idaho." US
 GEological Survey. https://idwr.idaho.gov/files/legal/CMR50/CMR50-1989-Water Resources-of-the-Upper-Big-Wood-River-Basin-Idaho.pdf.
- Fyfe, John C., Chris Derksen, Lawrence Mudryk, Gregory M. Flato, Benjamin D. Santer, Neil C.
 Swart, Noah P. Molotch, et al. 2017. "Large Near-Term Projected Snowpack Loss over
 the Western United States." *Nature Communications* 8 (1): 14996.
 https://doi.org/10.1038/ncomms14996.
- Garcia, E. S., and C. L. Tague. 2015. "Subsurface Storage Capacity Influences Climate—
 Evapotranspiration Interactions in Three Western United States Catchments." *Hydrology* and Earth System Sciences 19 (12): 4845–58. https://doi.org/10.5194/hess-19-4845-2015.





- Goeking, Sara A., and David G. Tarboton. 2020. "Forests and Water Yield: A Synthesis of
 Disturbance Effects on Streamflow and Snowpack in Western Coniferous Forests."
 Journal of Forestry 118 (2): 172–92. https://doi.org/10.1093/jofore/fvz069.
- Guardiola-Claramonte, M., Peter A. Troch, David D. Breshears, Travis E. Huxman, Matthew B.
 Switanek, Matej Durcik, and Neil S. Cobb. 2011. "Decreased Streamflow in Semi-Arid
 Basins Following Drought-Induced Tree Die-off: A Counter-Intuitive and Indirect
 Climate Impact on Hydrology." *Journal of Hydrology* 406 (3): 225–33.
 https://doi.org/10.1016/j.jhydrol.2011.06.017.
- Hanan, Erin J., Carla M. D'Antonio, Dar A. Roberts, and Joshua P. Schimel. 2016. "Factors
 Regulating Nitrogen Retention During the Early Stages of Recovery from Fire in Coastal
 Chaparral Ecosystems." *Ecosystems* 19 (5): 910–26. https://doi.org/10.1007/s10021-016-9975-0.
- Hanan, Erin J., Christina Tague, Janet Choate, Mingliang Liu, Crystal Kolden, and Jennifer
 Adam. 2018. "Accounting for Disturbance History in Models: Using Remote Sensing to
 Constrain Carbon and Nitrogen Pool Spin-Up." Ecological Applications: A Publication
 of the Ecological Society of America 28 (5): 1197–1214.
 https://doi.org/10.1002/eap.1718.
- Hanan, Erin J., Christina (Naomi) Tague, and Joshua P. Schimel. 2017. "Nitrogen Cycling and Export in California Chaparral: The Role of Climate in Shaping Ecosystem Responses to Fire." *Ecological Monographs* 87 (1): 76–90. https://doi.org/10.1002/ecm.1234.
- Hicke, Jeffrey A., Morris C. Johnson, Jane L. Hayes, and Haiganoush K. Preisler. 2012. "Effects of Bark Beetle-Caused Tree Mortality on Wildfire." *Forest Ecology and Management* 271 (May): 81–90. https://doi.org/10.1016/j.foreco.2012.02.005.
- Homer, Collin G., Jon Dewitz, Limin Yang, Suming Jin, Patrick Danielson, George Z. Xian,
 John Coulston, Nathaniel Herold, James Wickham, and Kevin Megown. 2015.
 "Completion of the 2011 National Land Cover Database for the Conterminous United
 States Representing a Decade of Land Cover Change Information." *Photogrammetric Engineering and Remote Sensing* 81: 345354.
- Hubbart, Jason A. 2007. "Timber Harvest Impacts on Water Yield in the Continental/Maritime
 Hydroclimatic Region of the United States," 12.
 Law, B. E., O. J. Sun, J. Campbell, S. Van Tuyl, and P. E. Thornton. 2003. "Changes in Carbor
 - Law, B. E., O. J. Sun, J. Campbell, S. Van Tuyl, and P. E. Thornton. 2003. "Changes in Carbon Storage and Fluxes in a Chronosequence of Ponderosa Pine." *Global Change Biology* 9 (4): 510–24. https://doi.org/10.1046/j.1365-2486.2003.00624.x.
- Lawrence, David M., Keith W. Oleson, Mark G. Flanner, Peter E. Thornton, Sean C. Swenson,
 Peter J. Lawrence, Xubin Zeng, et al. 2011. "Parameterization Improvements and
 Functional and Structural Advances in Version 4 of the Community Land Model."
 Journal of Advances in Modeling Earth Systems 3 (1).
 https://doi.org/10.1029/2011MS00045.
- Lin, Laurence, Lawrence E. Band, James M. Vose, Taehee Hwang, Chelcy Ford Miniat, and
 Paul V. Bolstad. 2019. "Ecosystem Processes at the Watershed Scale: Influence of
 Flowpath Patterns of Canopy Ecophysiology on Emergent Catchment Water and Carbon
 Cycling." Ecohydrology 0 (0): e2093. https://doi.org/10.1002/eco.2093.
- Livneh, Ben, Jeffrey S. Deems, Brian Buma, Joseph J. Barsugli, Dominik Schneider, Noah P.
 Molotch, K. Wolter, and Carol A. Wessman. 2015. "Catchment Response to Bark Beetle
 Outbreak and Dust-on-Snow in the Colorado Rocky Mountains." *Journal of Hydrology* 523 (April): 196–210. https://doi.org/10.1016/j.jhydrol.2015.01.039.



786 787

788

789

790

791



- Lundquist, Jessica D., Susan E. Dickerson-Lange, James A. Lutz, and Nicoleta C. Cristea. 2013.
 "Lower Forest Density Enhances Snow Retention in Regions with Warmer Winters: A
 Global Framework Developed from Plot-Scale Observations and Modeling." Water
 Resources Research 49 (10): 6356–70. https://doi.org/10.1002/wrcr.20504.
- Lundquist, Jessica D., Paul J. Neiman, Brooks Martner, Allen B. White, Daniel J. Gottas, and F.
 Martin Ralph. 2008. "Rain versus Snow in the Sierra Nevada, California: Comparing
 Doppler Profiling Radar and Surface Observations of Melting Level." *Journal of Hydrometeorology* 9 (2): 194–211. https://doi.org/10.1175/2007JHM853.1.
 - McVicar, Tim R., Michael L. Roderick, Randall J. Donohue, Ling Tao Li, Thomas G. Van Niel, Axel Thomas, Jürgen Grieser, et al. 2012. "Global Review and Synthesis of Trends in Observed Terrestrial Near-Surface Wind Speeds: Implications for Evaporation." *Journal of Hydrology* 416–417 (January): 182–205. https://doi.org/10.1016/j.jhydrol.2011.10.024.
 - Meddens, Arjan, Jeffrey A Hicke, and Charles A Ferguson. 2012. "Spatiotemporal Patterns of Observed Bark Beetle-Caused Tree Mortality in British Columbia and the Western United States." *Ecological Applications : A Publication of the Ecological Society of America* 22 (October): 1876–91. https://doi.org/10.2307/41723101.
- Mikkelson, K. M., R. M. Maxwell, I. Ferguson, J. D. Stednick, J. E. McCray, and J. O. Sharp.
 2013. "Mountain Pine Beetle Infestation Impacts: Modeling Water and Energy Budgets at the Hill-Slope Scale." *Ecohydrology* 6 (1): 64–72. https://doi.org/10.1002/eco.278.
- Mitchell, Kenneth E., Dag Lohmann, Paul R. Houser, Eric F. Wood, John C. Schaake, Alan
 Robock, Brian A. Cosgrove, et al. 2004. "The Multi-Institution North American Land
 Data Assimilation System (NLDAS): Utilizing Multiple GCIP Products and Partners in a
 Continental Distributed Hydrological Modeling System." In .
 https://doi.org/10.1029/2003JD003823.
- Monteith, J. L. 1965. "Evaporation and Environment." *Symposia of the Society for Experimental Biology* 19: 205–34.
- Montesi, James, Kelly Elder, R. A. Schmidt, and Robert E. Davis. 2004. "Sublimation of Intercepted Snow within a Subalpine Forest Canopy at Two Elevations." *Journal of Hydrometeorology* 5 (5): 763–73. https://doi.org/10.1175/1525-7541(2004)005<0763:SOISWA>2.0.CO;2.
- Moore, R Dan, and S M Wondzell. 2005. "PHYSICAL HYDROLOGY AND THE EFFECTS OF FOREST HARVESTING IN THE PACIFIC NORTHWEST: A REVIEW," 22.
- Morillas, L., R. E. Pangle, G. E. Maurer, W. T. Pockman, N. McDowell, C.-W. Huang, D. J.
 Krofcheck, et al. 2017. "Tree Mortality Decreases Water Availability and Ecosystem
 Resilience to Drought in Piñon-Juniper Woodlands in the Southwestern U.S." Journal of
 Geophysical Research: Biogeosciences 122 (12): 3343–61.
 https://doi.org/10.1002/2017JG004095.
- Mu, Qiaozhen, Faith Ann Heinsch, Maosheng Zhao, and Steven W. Running. 2007.
 "Development of a Global Evapotranspiration Algorithm Based on MODIS and Global Meteorology Data." *Remote Sensing of Environment* 111 (4): 519–36.
 https://doi.org/10.1016/j.rse.2007.04.015.
- Mu, Qiaozhen, Maosheng Zhao, and Steven W. Running. 2011. "Improvements to a MODIS Global Terrestrial Evapotranspiration Algorithm." *Remote Sensing of Environment* 115 (8): 1781–1800. https://doi.org/10.1016/j.rse.2011.02.019.



841 842



- Nash, J. E., and J. V. Sutcliffe. 1970. "River Flow Forecasting through Conceptual Models Part I
 A Discussion of Principles." *Journal of Hydrology* 10 (3): 282–90.
 https://doi.org/10.1016/0022-1694(70)90255-6.
- NRCS. n.d. "SNOTEL." https://www.wcc.nrcs.usda.gov/about/mon_automate.html.
- Paine, T. D., K. F. Raffa, and T. C. Harrington. 1997. "Interactions Among Scolytid Bark
 Beetles, Their Associated Fungi, and Live Host Conifers." *Annual Review of Entomology* 42 (1): 179–206. https://doi.org/10.1146/annurev.ento.42.1.179.
- Parton, W. J., A. R. Mosier, D. S. Ojima, D. W. Valentine, D. S. Schimel, K. Weier, and A. E.
 Kulmala. 1996. "Generalized Model for N2 and N2O Production from Nitrification and Denitrification." *Global Biogeochemical Cycles* 10 (3): 401–12.
 https://doi.org/10.1029/96GB01455.
- Penn, Colin A., Lindsay A. Bearup, Reed M. Maxwell, and David W. Clow. 2016. "Numerical Experiments to Explain Multiscale Hydrological Responses to Mountain Pine Beetle Tree Mortality in a Headwater Watershed." *Water Resources Research* 52 (4): 3143–61. https://doi.org/10.1002/2015WR018300.
- Perry, Timothy D., and Julia A. Jones. 2017. "Summer Streamflow Deficits from Regenerating Douglas-Fir Forest in the Pacific Northwest, USA: Summer Streamflow Deficits from Regenerating Douglas-Fir Forest." *Ecohydrology* 10 (2): e1790. https://doi.org/10.1002/eco.1790.
 - Pomeroy, John, Xing Fang, and Chad Ellis. 2012. "Sensitivity of Snowmelt Hydrology in Marmot Creek, Alberta, to Forest Cover Disturbance: SENSITIVITY OF SNOWMELT HYDROLOGY TO FOREST DISTURBANCE." *Hydrological Processes* 26 (12): 1891–1904. https://doi.org/10.1002/hyp.9248.
- Potts, Donald F. 1984. "Hydrologic Impacts of a Large-Scale Mountain Pine Beetle
 (Dendroctonus Ponderosae Hopkins) Epidemic1." *JAWRA Journal of the American*Water Resources Association 20 (3): 373–77. https://doi.org/10.1111/j.17521688.1984.tb04719.x.
- Robles, Marcos D., Robert M. Marshall, Frances O'Donnell, Edward B. Smith, Jeanmarie A.
 Haney, and David F. Gori. 2014. "Effects of Climate Variability and Accelerated Forest
 Thinning on Watershed-Scale Runoff in Southwestern USA Ponderosa Pine Forests."
 PLOS ONE 9 (10): e111092. https://doi.org/10.1371/journal.pone.0111092.
- Ryan, Michael G. 1991. "Effects of Climate Change on Plant Respiration." *Ecological Applications* 1 (2): 157–67. https://doi.org/10.2307/1941808.
- Searcy, James Kincheon. 1959. "Flow-Duration Curves." Report 1542A. Water Supply Paper. USGS Publications Warehouse. https://doi.org/10.3133/wsp1542A.
- Sexstone, Graham A., David W. Clow, Steven R. Fassnacht, Glen E. Liston, Christopher A.
 Hiemstra, John F. Knowles, and Colin A. Penn. 2018. "Snow Sublimation in Mountain
 Environments and Its Sensitivity to Forest Disturbance and Climate Warming." Water
 Resources Research 54 (2): 1191–1211. https://doi.org/10.1002/2017WR021172.
- Skinner, Kenneth D. 2013. "Post-Fire Debris-Flow Hazard Assessment of the Area Burned by
 the 2013 Beaver Creek Fire near Hailey, Central Idaho." USGS Numbered Series 2013–
 1273. Open-File Report. Reston, VA: U.S. Geological Survey.
 http://pubs.er.usgs.gov/publication/ofr20131273.
- Slinski, Kimberly M., Terri S. Hogue, Aaron T. Porter, and John E. McCray. 2016. "Recent Bark Beetle Outbreaks Have Little Impact on Streamflow in the Western United States."



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- 866 Environmental Research Letters 11 (7): 074010. https://doi.org/10.1088/1748-867 9326/11/7/074010.
- 868 Smith, Frederick W., D. Arthur Sampson, and James N. Long. 1991. "Comparison of Leaf Area 869 Index Estimates from Tree Allometrics and Measured Light Interception." *Forest Science* 870 37 (6): 1682–88. https://doi.org/10.1093/forestscience/37.6.1682.
 - Smith, Rex Onis. 1960. "Geohydrologic Evaluation of Streamflow Records in the Big Wood River Basin, Idaho." USGS Numbered Series 1479. Water Supply Paper. U.S. Govt. Print. Off., http://pubs.er.usgs.gov/publication/wsp1479.
- 874 Son, Kyongho, and Christina Tague. 2019. "Hydrologic Responses to Climate Warming for a Snow-Dominated Watershed and a Transient Snow Watershed in the California Sierra." 876 *Ecohydrology* 12 (1): e2053. https://doi.org/10.1002/eco.2053.
- 877 Sun, Ning, Mark Wigmosta, Tian Zhou, Jessica Lundquist, Susan Dickerson-Lange, and
 878 Nicoleta Cristea. 2018. "Evaluating the Functionality and Streamflow Impacts of
 879 Explicitly Modelling Forest–Snow Interactions and Canopy Gaps in a Distributed
 880 Hydrologic Model." *Hydrological Processes* 32 (13): 2128–40.
 881 https://doi.org/10.1002/hyp.13150.
- Tague, C. L., and L. E. Band. 2004. "RHESSys: Regional Hydro-Ecologic Simulation System—
 An Object-Oriented Approach to Spatially Distributed Modeling of Carbon, Water, and
 Nutrient Cycling." *Earth Interactions* 8 (19): 1–42. https://doi.org/10.1175/10873562(2004)8<1:RRHSSO>2.0.CO;2.
- Tague, Christina L., Max Moritz, and Erin Hanan. 2019. "The Changing Water Cycle: The Eco-Hydrologic Impacts of Forest Density Reduction in Mediterranean (Seasonally Dry) Regions." *Wiley Interdisciplinary Reviews: Water* 0 (0): e1350. https://doi.org/10.1002/wat2.1350.
 - Tsamir, Mor, Sagi Gottlieb, Yakir Preisler, Eyal Rotenberg, Fyodor Tatarinov, Dan Yakir, Christina Tague, and Tamir Klein. 2019. "Stand Density Effects on Carbon and Water Fluxes in a Semi-Arid Forest, from Leaf to Stand-Scale." *Forest Ecology and Management* 453 (December): 117573. https://doi.org/10.1016/j.foreco.2019.117573.
- White, Joseph D., and Steven W. Running. 1994. "Testing Scale Dependent Assumptions in Regional Ecosystem Simulations." *Journal of Vegetation Science* 5 (5): 687–702. https://doi.org/10.2307/3235883.
- White, Michael A., Peter E. Thornton, Steven W. Running, and Ramakrishna R. Nemani. 2000.

 "Parameterization and Sensitivity Analysis of the BIOME–BGC Terrestrial Ecosystem
 Model: Net Primary Production Controls." *Earth Interactions* 4 (3): 1–85.

 https://doi.org/10.1175/1087-3562(2000)004<0003:PASAOT>2.0.CO;2.
 - Wine, Michael L, Daniel Cadol, and Oleg Makhnin. 2018. "In Ecoregions across Western USA Streamflow Increases during Post-Wildfire Recovery." *Environmental Research Letters* 13 (1): 014010. https://doi.org/10.1088/1748-9326/aa9c5a.
- Winkler, Rita, Sarah Boon, Barbara Zimonick, and Dave Spittlehouse. 2014. "Snow
 Accumulation and Ablation Response to Changes in Forest Structure and Snow Surface
 Albedo after Attack by Mountain Pine Beetle." *Hydrological Processes* 28 (2): 197–209.
 https://doi.org/10.1002/hyp.9574.
- Zhang, Ke, John S. Kimball, Qiaozhen Mu, Lucas A. Jones, Scott J. Goetz, and Steven W.
 Running. 2009. "Satellite Based Analysis of Northern ET Trends and Associated
 Changes in the Regional Water Balance from 1983 to 2005." *Journal of Hydrology* 379
 (1): 92–110. https://doi.org/10.1016/j.jhydrol.2009.09.047.





Zhao, Maosheng, Steven W. Running, and Ramakrishna R. Nemani. 2006. "Sensitivity of
 Moderate Resolution Imaging Spectroradiometer (MODIS) Terrestrial Primary
 Production to the Accuracy of Meteorological Reanalyses." *Journal of Geophysical Research: Biogeosciences* 111 (G1). https://doi.org/10.1029/2004JG000004.

Table 1. Classification of aridity index.

Aridity Index (i.e. PET/P)	Туре
> 2	Water - limited
0.8 - 2	Balanced
< 0.8	Energy - limited





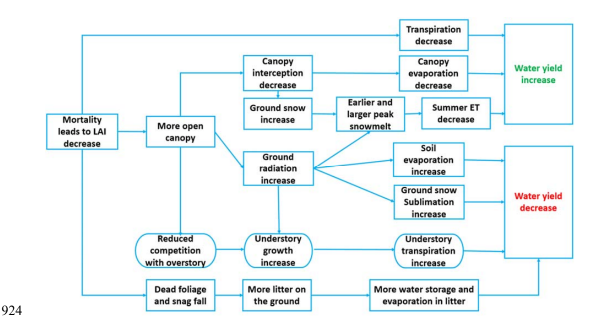
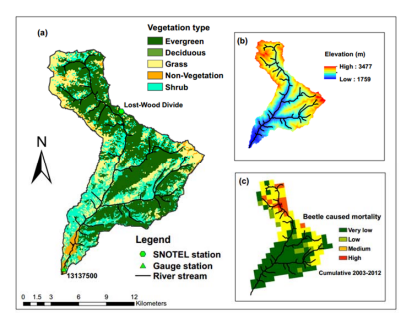


Figure 1. Mechanism of water yield responses to beetle-caused mortality during the red and gray phases (0-10 years after beetle outbreak), semicircle boxes represent understory responses and square boxes represent overstory responses.

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930 Figure 2. Land cover, el

Figure 2. Land cover, elevation, and tree mortality for Trail Creek. (a) is the land cover map with the main vegetation type, (b) is the elevation gradient, and (c) is the severity of beetle caused tree mortality (during the period 2003-2012 Meddens et al. (2012)). Note that, for our modeling experiments, we prescribe beetle outbreak uniformly across evergreen patches instead of using historical beetle outbreak data.

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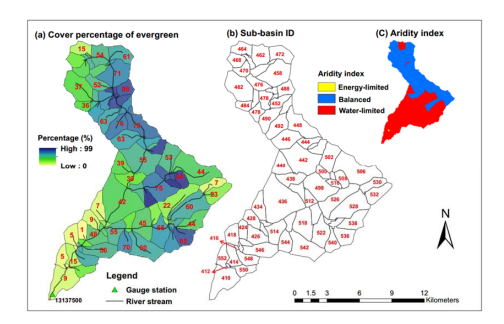


Figure 3. Trail creek evergreen forest cover percentage for each sub-basin, sub-basin ID, and long-term aridity index. Aridity index is defined as annual mean potential evapotranspiration (PET) / precipitation (P) from 38 years of data (see Sect 3.4), PET/P > 2 is water-limited, PET/P < 0.8 is energy-limited, PET/P between 0.8 and 2 is balanced. Recall that only evergreen forest trees are attacked during beetle outbreaks.





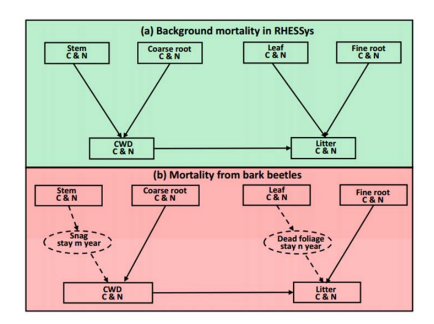


Figure 4. Conceptual framework of the beetle effect model.

(a) Normal background mortality routine in RHESSys before beetle outbreak. (b) Mortality from bark beetles. We add snag (standing dead trees) and dead foliage (needles still on dead trees) pools, shown in the dashed circle. After a beetle outbreak, carbon (C) and Nitrogen (N) move from stems to snag pools (black dashed arrow). After staying in the snag pool for m years, C and N move from snag to coarse wood debris pools (CWD) with an exponential decay rate to represent the snag fall (gray dashed arrow). It is a similar process for leaf C and N, which move from leaf to dead foliage to litter pools (black dotted arrow). Furthermore, C and N in the CWD and fine root pools move to the litter pool immediately after outbreak (solid black and gray arrows). Figure modified from Edburg et al. (2012).



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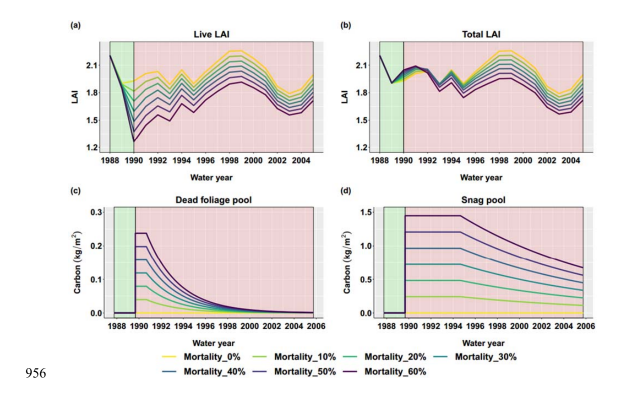


Figure 5. Basin-scale vegetation responses after beetle outbreak for different evergreen mortality level. (a) Annual live leaf area index (Live LAI), (b) Annual total LAI (LAI calculated including dead foliage pool), (c) Daily dead foliage pool, and (d) Daily snag pool after outbreak. The green background color is the period before beetle outbreak, and the red background color is after the beetle outbreak.



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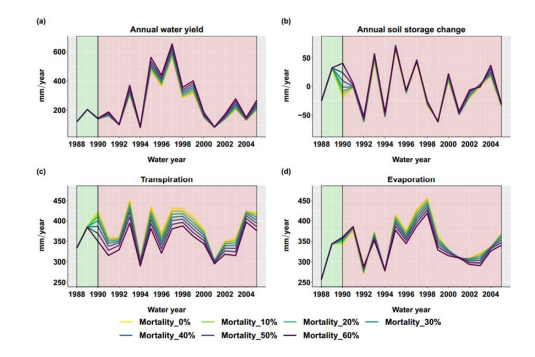
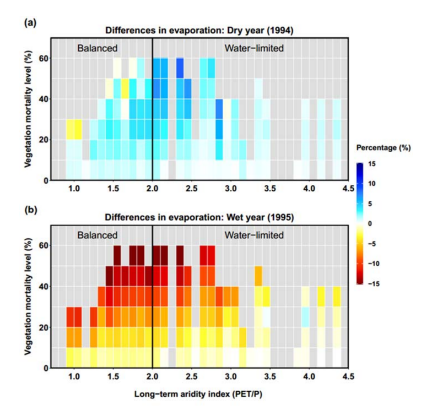


Figure 6. Basin-scale annual sum of hydrologic fluxes responses after beetle outbreak (1989) for different evergreen mortality levels. (a) Annual water yield calculated as annual sum of basin streamflow, and (b) annual soil water storage change calculated as water year soil water storage at the end of water year minus soil water storage at the beginning of water year. (c) Transpiration is the annual sum of transpiration for both overstory and understory. (d) Evaporation is calculated as the annual sum of canopy evaporation, ground evaporation, and snow sublimation.





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Figure 7. Relationship among long-term aridity, vegetation mortality level, and differences in evaporation for a dry year (1994, a) and wet year (1995, b). Differences are calculated as the normalized differences (%) of evaporation between each evergreen mortality scenario and the control run for no beetle outbreak. Vegetation mortality for each sub-basin is calculated as the percentage of evergreen patches multiplied by the mortality level of evergreen caused by beetles. Long-term aridity is defined as temporally averaged (38 years) potential evapotranspiration relative to precipitation.



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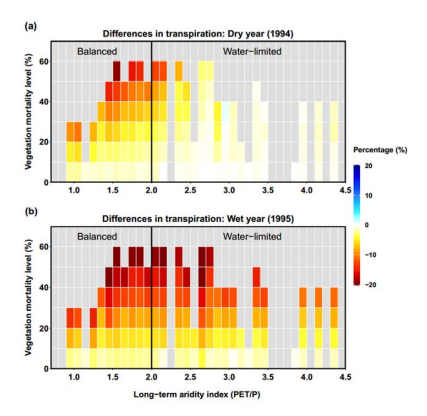


Figure 8. Relationship among long-term aridity, vegetation mortality, and differences in transpiration for a dry year (1994, a) and wet year (1995, b).



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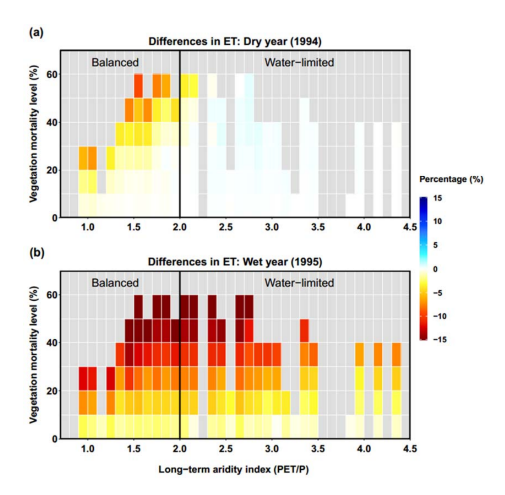


Figure 9. Relationship among long-term aridity, vegetation mortality level and differences in ET for a dry year (1994, a) and a wet year (1995, b).



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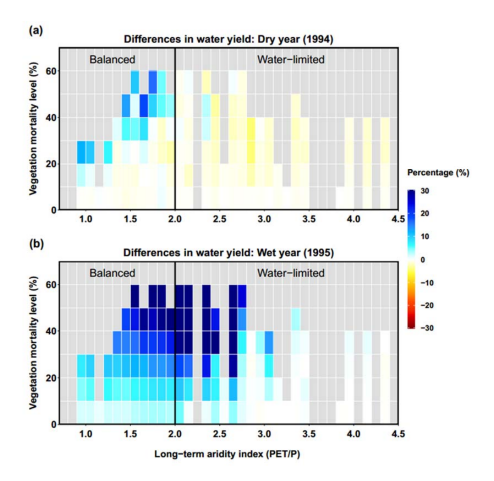


Figure 10. Relationship among long-term aridity, vegetation mortality level and Differences in water yield for a dry year (1994, a) and wet year (1995, b).





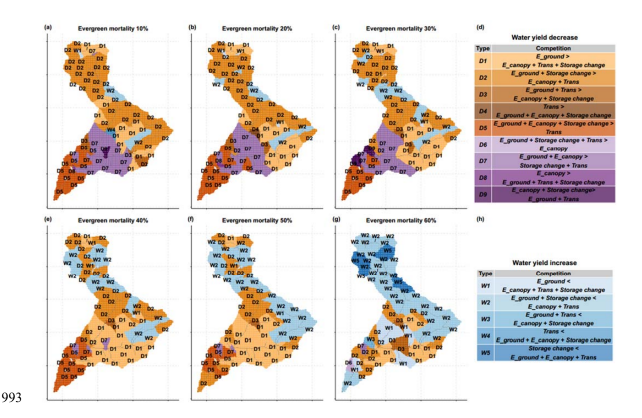


Figure 11. Water yield response types after beetle outbreak for different evergreen mortality scenarios compared with control scenario. D1 to D9 are water yield decrease types and W1 to W5 are water yield increase types. In panel D and H, the left side of each type are increasing fluxes that cause water yield decreases and the right side are decreasing fluxes that cause water yield increase. If the left side is larger than the right side, water yield increases, and vice versa. (Note: this mortality is evergreen mortality, which is different from vegetation mortality.)