1	Assessing Climate Change Impacts on Live Fuel Moisture and Wildfire Risk
2	Using a Hydrodynamic Vegetation Model
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Abstract: Live fuel moisture content (LFMC) plays a critical role in wildfire dynamics, but little 26 27 is known about responses of LFMC to multivariate climate change, e.g., warming temperature, CO₂ fertilization and altered precipitation patterns, leading to a limited prediction ability of 28 29 future wildfire risks. Here, we use a hydrodynamic demographic vegetation model to estimate LFMC dynamics of chaparral shrubs, a dominant vegetation type in fire-prone southern 30 California. We parameterize the model based on observed shrub allometry and hydraulic traits, 31 32 and evaluate the model's accuracy through comparisons between observed and simulated LFMC of three plant functional types (PFTs) under current climate conditions. Moreover, we estimate 33 the number of days per year of LFMC below 79% (which is a critical threshold for wildfire 34 35 danger rating of southern California chaparral shrubs) from 1960 to 2099 for each PFT, and 36 compare the number of days below the threshold for medium and high greenhouse gas emission 37 scenarios (RCP4.5 and 8.5). We find that climate change could lead to more days per year (5.2-38 14.8% increase) with LFMC below 79% between the historical (1960-1999) and future (2080-39 2099) periods, implying an increase in wildfire danger for chaparral shrubs in southern 40 California. Under the high greenhouse gas emission scenario during the dry season, we find that 41 the future LFMC reductions mainly result from a warming temperature, which leads to 9.1-42 18.6% reduction in LFMC. Lower precipitation in the spring leads to a 6.3-8.1% reduction in LFMC. The combined impacts of warming and precipitation change on fire season length are 43 44 equal to the additive impacts of warming and precipitation change individually. Our results show that the CO₂ fertilization will mitigate fire risk by causing a 3.5-4.8% increase in LFMC. Our 45 results suggest that multivariate climate change could cause a significant net reduction in LFMC 46 and thus exacerbate future wildfire danger in chaparral shrub systems. 47

48 Keywords: FATES-HYDRO, chaparral shrubs, plant functional types, southern California, CO₂
49 enrichment, climate change

50 **1. Introduction**

Historical warming and changes in precipitation have already impacted wildfire at a 51 global scale (e.g. Stocks et al. 1998; Gillett et al. 2004; Westerling et al. 2003, 2006) and it is 52 expected that accelerating future warming will continue to significantly influence global wildfire 53 regimes (e.g. Flannigan et al. 2009; Liu et al. 2010; Moritz et al. 2012). So far, prior studies have 54 55 mainly focused on impacts of dead fuel moisture, fuel loads, and weather conditions on wildfire. Limited studies have applied proxies of live fuel moisture in global-fire models. For example, 56 57 dead fuel moisture is found to be related to fire ignition and fire spread potential (or potential 58 area burnt) (Aguado et al. 2007), specific weather conditions such as increased vapor pressure deficit (Williams et al. 2019) can lead to a vast increase in fire activity (Goss et al. 2020), and 59 wildfire fuel loads are projected to increase under climate change (Matthews et al. 2012; Clarke 60 et al. 2016). In global-fire models, studies have used proxies of live fuel moisture (Bistinas et al. 61 2014; Kelley et al. 2019) as well as explicit representation of live fuels (Hantson et al. 2016; 62 Rabin et al. 2017). While previous studies provide great insights into fire risks with changes in 63 climate, dead fuel moisture, fuel loads, and representation of live fuel moisture, there is still 64 limited understanding of how climate change influences live fuel moisture content (LFMC) and 65 66 the consequent wildfire risks. This is particularly true for the combined impacts of warming temperature, altered precipitation, and increasing CO₂ fertilization (Chuvieco et al. 2004; 67 Pellizzaro 2007; Caccamo et al. 2012a, b; Williams et al. 2019; Goss et al. 2020). 68

A measure of water content within living plant tissue in relation to their dry weight,
LFMC has been found to be one of the most critical factors influencing combustion, fire spread,

and fire consumption (e.g. Agee et al. 2002; Zarco-Tejada et al. 2003; Bilgili & Saglam 2003; 71 Yebra et al. 2008; Dennison et al. 2008; Anderson & Anderson 2010; Keeley et al. 2011). This is 72 73 because a low LFMC leads to increased flammability and a higher likelihood of ignition (Dimitrakopoulos & Papaioannou 2001). For instance, LFMC was found to be a significant 74 factor contributing to the occurrence of wildfires in Australia (Plucinski 2003; Nolan et al. 2016; 75 76 Yebra et al. 2018; Rossa & Fernandes 2018; Pimont et al. 2019), Spain (Chuvieco et al. 2009) and California (Santa Monica Mountains; Dennison et al. 2008; Dennison & Moritz 2009; 77 Pivovaroff et al. 2019). Dennison & Moritz (2009) found strong evidence of a LFMC threshold 78 79 (79%) for southern California chaparral shrubs, which may determine when large fires can occur in this region. 80

Vegetation moisture content is dependent on both ecophysiological characteristics of the 81 species and environmental conditions, including both climatic variables and soil water 82 83 availability (Rothermel 1972; Castro et al. 2003; Castro et al. 2003; Pellizzaro 2007; Pivovaroff 84 et al. 2019; Nolan et al. 2020). So far, little is known about the relative importance of different climate variables to future LFMC dynamics. On the one hand, warming could contribute to a 85 86 higher atmospheric demand and higher evapotranspiration (Rind et al. 1990) and thus lead to a 87 lower LFMC. On the other hand, higher CO₂ concentration will decrease stomatal conductance (Wullschleger et al. 2002) and plant water loss, and thus lead to a higher LFMC. The impacts of 88 CO₂ and warming could be complicated by local changes in precipitation patterns and humidity 89 90 (Mikkelsen et al. 2008).

91 The sensitivity of LFMC to climate change is likely to be affected by plant hydraulic
92 traits (the plant properties that regulate water transport and storage within plant tissues), which
93 affect plant water regulation (Wu et al. 2020). Variations in hydraulic traits reflect contrasting

plant drought adaptation strategies when responding to dry conditions. Two contrasting overall 94 strategies are: 1) water stress avoiders and 2) water stress tolerators (Tobin et al. 1999; Wei et al. 95 2019). The "avoiders" are generally characterized by a more conservative hydraulic strategy 96 under water stress by either closing stomata early, dropping leaves or accessing deep water to 97 avoid more negative water potentials and therefore xylem cavitation. Meanwhile, the "tolerators" 98 99 typically build xylem and leaves that are more resistant to cavitation so that they can tolerate more negative water potential and continue to conduct photosynthesis under water stress. 100 101 Therefore, compared with the tolerators, the avoiders normally have a lower sapwood density 102 and higher plant water storage capacity in their tissues to avoid cavitation (Meinzer et al. 2003, 2009; Pineda-Garcia et al. 2013). Because the avoiders rely on water storage capacity as one way 103 to avoid cavitation thereby maintaining a relatively high LFMC, and water loss from storage 104 should increase with warming, LFMC could be more sensitive to climate change in avoiders 105 relative to tolerators. 106

107 While over half of terrestrial landscapes on Earth are considered fire-prone (Krawchuk et al. 2009), Mediterranean-type climate regions are routinely impacted by fire, often on an annual 108 basis. This is partly because Mediterranean climate regions are characterized by winter rains 109 110 followed by annual dry season, when little to no rainfall occurs for several months. Multiday periods of extreme high temperatures, as well as katabatic hot, dry, and intense winds, often 111 112 punctuate the annual drought, leading to some of the worst fire weather in the world (Schroeder et al. 1964). This can result in wildfires that are large, high-intensity, and stand-replacing 113 114 (Keeley 1995; Keeley & Zedler 2009; Balch et al. 2017). Globally, Mediterranean climate regions are characterized by evergreen sclerophyllous-leaved shrublands. The Mediterranean 115 climate region in California is dominated by chaparral, which is adapted to the periodic fire 116

117	regime in California (Venturas et al. 2016). Previous studies have proposed a variety of
118	relationships between chaparral LFMC and fire danger in southern California (Dennison et al.
119	2008; Dennison & Moritz 2009), but less is known about how climate changes could alter LFMC
120	and fire danger. In chaparral, LFMC is usually high during the winter and spring (wet season)
121	and then gradually declines during the dry season (summer and fall), which leads to a typical fire
122	season approximately six months long in southern California (Pivovaroff et al. 2019). One key
123	risk is that severe drought conditions are becoming exacerbated under climate change, which
124	might lead to the occurrence of larger and higher-intensity fires in chaparral (Dennison et al.
125	2008; Dennison & Moritz 2009).
126	There has been a long history of wildfire modeling, with three types of models: 1) fine-
127	scale fire behavior models (e.g. FIRETEC by Linn et al. 2002); 2) landscape-scale fire
128	disturbance models (e.g. LANDIS-II by Sturtevant et al. 2009); and 3) global-scale fire dynamics
129	models (e.g. Hantson et al. 2016; Rabin et al. 2017; SPITFIRE by Thonicke et al. 2010). While
130	these models focus on simulation at different scales, fire measures of the simulation are mainly
131	calculated from climate and dead fuel moisture and currently lack prediction of LFMC dynamics.
132	One key limitation is that most previous models have not yet considered plant hydrodynamics
133	(Holm et al. 2012; Xu et al. 2013; Seiler et al. 2014), which is integral to LFMC prediction.
134	Recently, there have been important improvements to global dynamic and demographic
135	vegetation models by incorporating plant hydrodynamics (McDowell et al. 2013; Xu et al. 2016;
136	Fisher et al. 2018; Mencuccini et al. 2019). These models have been used to study the interaction
137	between elevated CO ₂ and drought (Duursma & Medlyn, 2012), the impact of hydraulic traits on
138	plant drought response (Christofferson et al. 2016), the role of hydraulic diversity in vegetation
139	response to drought (Xu et al. 2016) and hydroclimate change (Powell et al. 2018), and

vegetation water stress and root water uptake (Kennedy et al. 2019). While the main purpose of
the new hydraulic components is to improve the vegetation response to drought, the fact that
hydrodynamic models consider tissue water content as a prognostic variable provides an
opportunity to assess the climate impacts on LFMC.

The objective of this study is to quantify LFMC dynamics and associated changes in fire 144 145 season duration for a chaparral ecosystem in southern California under climate change using a vegetation demographic model (that resolves the size and age-since-disturbance structure of 146 plant populations) (Xu et al. 2016; Fisher et al. 2018) that incorporates plant hydraulics. We test 147 one overarching hypothesis: future climate change will decrease LFMC and consequently result 148 in a longer fire season as determined by a critical threshold of LFMC (H₀). Specifically, we test 149 the following four sub-hypotheses: 1) warming has a stronger impact on LFMC than CO₂ 150 fertilization (H₁); 2) the reductions in spring and autumn precipitation lead to a longer fire season 151 as determined by LFMC (H₂); 3) the combined impacts of warming and precipitation on fire 152 153 season length are equal to the additive impacts of warming and precipitation change individually (H₃); and 4) LFMC for plants with more conservative hydraulic strategies ("avoiders") will be 154 155 more vulnerable to warming (H₄).

156 **2. Materials and Methods**

To understand climate change impacts on LFMC for the chaparral ecosystem, we applied
the Functionally Assembled Terrestrial Simulator (FATES; Fisher et al. 2015; Massoud et al.

159 2019; Koven et al. 2020) coupled with a hydrodynamic vegetation module (FATES-HYDRO;

160 Christoffersen et al. 2016) in the Santa Monica Mountains in California. We validated the model

- using the observed LFMC for three chaparral shrub plant functional types (PFTs). Then, we
- applied FATES-HYDRO to estimate long-term dynamics of leaf water content (LWC) during

163 1960-2099 for each PFT using downscaled Earth System Model (ESM) climate scenarios. We
164 converted simulated leaf water content (LWC) to LFMC within leaves and shoots. Based on the
165 simulated LFMC, we evaluated wildfire danger based on the number of days per year of LFMC
166 below the critical value of 79% from 1960 to 2099 for each PFT under RCP 4.5 and 8.5. Finally,
167 we assessed the relative importance of changes in individual and combined climate variables
168 including CO₂, temperature, precipitation, and tested the corresponding hypotheses.

169 **2.1 Study site**

170 The study site is located at the Stunt Ranch Santa Monica Mountains Reserve, in the Santa Monica Mountains in California, USA (N 34° 05', W 118° 39'). Stunt Ranch is dominated 171 by chaparral vegetation, with an elevation of approximately 350 m, a west-facing slope, and a 172 173 Mediterranean-type climate. The study site harbors an abundance of fauna, particularly birds and reptiles. The mean annual temperature is 18.1°C. The mean annual precipitation is 478 mm, 174 175 occurring mostly during the wet season (i.e. November-March) with almost no rainfall during the dry season (i.e. April-October). Stunt Ranch last burned in year 1993. We focused on PFTs 176 177 representing 11 study species (Fig. 1), including chamise (Adenostoma fasciculatum - Af), red 178 shank (Adenostoma sparsifolium - As), big berry manzanita (Arctostaphylos glauca - Ag), buck brush (Ceanothus cuneatus - Cc), greenbark ceanothus (Ceanothus spinosus - Cs), mountain 179 mahogany (*Cercocarpus betuloides* - Cb), toyon (*Heteromeles arbutifolia* - Ha), laurel sumac 180 181 (Malosma laurina - Ml), scrub oak (Quercus berberidifolia - Qb), hollyleaf redberry (Rhamnus ilicifolia - Ri), and sugar bush (*Rhus ovata* - Ro). Detailed information about the study site and 182 183 species characterizations found at Stunt Ranch can be found in Venturas et al. (2016) and 184 Pivovaroff et al. (2019).

185 2.2 FATES-HYDRO model

FATES is a vegetation demographic model (Fisher et al. 2015), which uses a size-186 structured group of plants (cohorts) and successional trajectory-based patches based on the 187 188 ecosystem demography approach (Moorcroft et al. 2001). FATES simulates the demographic process including seed production, seed emergence, growth and mortality (Koven et al. 2020). 189 Because the main purpose is to assess LFMC, we controlled for variation in plant size structure 190 191 that could arise from plant traits or climate differences between model runs by using a reducedcomplexity configuration of the model where growth and mortality are turned off and ecosystem 192 structure is held constant. FATES has to be hosted by a land surface model to simulate the soil 193 194 hydrology, canopy temperature and transpiration. These host land models include the Exascale Energy Earth System Model (E3SM, Caldwell et al., 2019) land model (ELM) as well as the 195 Community Earth System Model (Fisher et al 2015) and the Norwegian Earth system model 196 (NorESM, Tjiputra et al 2013). In this study, we used the DOE-sponsored ELM as our host land 197 model. The time step of FATES to calculate carbon and water fluxes is 30 minutes and it can 198 199 downscale the data from 6-hourly climate drivers.

A key component of FATES, the plant hydrodynamic model (HYDRO, based on Christoffersen et al. 2016), simulates the water flow from soil through root, stem and leaf to the atmosphere. In this model, water flow is calculated based on water pressure gradients across different plant compartments (leaf, stem, transporting roots, absorbing roots and rhizosphere). Specifically, flow between compartment *i* and i + 1 (Q_i) is given by

$$Q_i = -K_i \Delta h_i,\tag{1}$$

where K_i is the total conductance (kg MPa⁻¹ s⁻¹) at the boundary of compartments *i* and *i* + 1 and Δh_i is the total water potential difference between the compartments:

$$\Delta h_i = \rho_w g(z_i - z_{i+1}) + (\psi_i - \psi_{i+1}), \tag{2}$$

where z_i is compartment distance above (+) or below (-) the soil surface (m), ρ_w is the density of 207 water (10³ kg m⁻³), g is acceleration due to gravity (9.8 m s⁻²), and ψ_i is tissue or soil matric 208 water potential (MPa). K_i is treated here as the product of a maximum boundary conductance 209 between compartments i and i + 1 ($K_{max,i}$), and the fractional maximum hydraulic conductance 210 of the adjacent compartments (FMC_i or FMC_{i+1}), which is a function of the tissue water content. 211 A key parameter that controls FMC is the critical water potential (P₅₀) that leads to 50% loss of 212 hydraulic conductivity. The tissue water potential is calculated based on pressure-volume (PV) 213 theory (Tyree & Hammel, 1972; Tyree & Yang, 1990; Bartlett et al., 2012). For leaves, it is 214 215 described by three phases: 1) capillary water phase with full turgor, 2) elastic drainage phase 216 before reaching turgor loss point; and 3) post-turgor loss phase. For other tissues, it only has phases 2 and 3. Compared to a non-hydrodynamic model, this formulation allows the simulation 217 218 of plant water transport limitation on transpiration. For the non-hydrodynamic version of 219 FATES, the water limitation factor for transpiration (Btran) is calculated based on the soil 220 moisture potential (Fisher et al. 2015). For the hydrodynamic version, B_{tran} is calculated based on the leaf water potential (ψ_l) (Christoffersen et al. 2016) as follows, 221

222
$$B_{tran} = \left[1 + \left(\frac{\psi_l}{P_{50_gs}}\right)^{a_l}\right]^{-1}$$
(3)

where P_{50_gs} is the leaf water potential that leads to 50% loss of stomatal conductance and a_l is the shape parameter. Please refer to Christoffersen, et al. 2016 for details of formulations of *FMC* for different plant tissues.

226 **2.3** Allometry and trait data for model parameterization

FATES-HYDRO has a large number of parameters (>80; see Massoud et al. 2019 for a 227 complete list except for hydraulic parameters). Based on a previous sensitivity analysis study 228 (Massoud et al. 2019), we focused our parameter estimation efforts on the most influential 229 parameters for allometry, leaf and wood traits, and hydraulic traits from observations of 11 230 chaparral shrub species (see Supplementary, Table S2), collected from Jacobsen et al. (2008) and 231 232 Venturas et al. (2016). For this study, we assumed that the allometry of a shrub is analogous to that of a small tree. However, we did make several important modifications to accommodate the 233 234 allometry of shrub as their height and crown area relationships to diameter could be different 235 from trees. First, instead of using the diameter at breast height as the basis for allometry to calculate the height, crown area and leaf biomass, we used the basal diameter as the basis for 236 shrubs. Second, in the allometry of trees, the diameter for maximum height $(d_1$: 237 Fates_allom_dbh_maxheight, Table S1) is the same as the diameter for maximum crown area (d₂: 238 Fates_allom_d2ca_max, Table S1). As our data showed that d1 and d2 are different for shrubs, 239 240 we have modified the codes so that the d_1 and d_2 can be set for different values. It is possible that different branching and path length patterns for stems of chaparral species could impact the 241 hydraulics compared to trees; however, FATES-HYDRO treats all the aboveground xylem as a 242 243 single pool and thus it should not affect our model simulation results.

Based on a hierarchical cluster analysis (Bridges 1966) of allometry and trait data, there are a clear separation among the shrub species. First, the dendrogram is built and every data point finally merges into a single cluster with the height shown on the y-axis. Then we cut the dendrogram in order to create the desired number of clusters determined by a pragmatic choice based on hydraulic traits of eleven chaparral shrub species (Fig. 1). R's rect.hclust function (https://www.rdocumentation.org/packages/stats/versions/3.6.2/topics/rect.hclust) was used to

see the clusters on the dendrogram. All parameters of allometry, leaf and wood traits, and 250 hydraulic traits were collected from observations shown in the Table S2 and S3 of the 251 supplementary. According to the principle of model parsimony, we do not want to classify the 252 species into more than 3 PFTs. Meanwhile, we also want to differentiate the fundamental plant 253 growth and water use strategies that will determine plant transpiration rate and the corresponding 254 255 LFMC. If we choose to classify the species into two PFTs (based on the solid horizontal line in Fig. 1), then we will not be able to differentiate species with aggressive and conservative 256 257 hydraulic strategies in the second group and not be able to test H4. Therefore, the chaparral shrub 258 species were classified into three PFTs (based on the dotted horizontal line in Fig. 1 and Table S3), that are able to differentiate plant growth and hydraulic strategy. The three PFTs include a 259 low productivity, aggressive drought tolerance hydraulic strategy PFT (PFT-LA) with a relative 260 low V_{c,max25} (the maximum carboxylation rate at 25 °C) and a very negative P₅₀ (the leaf water 261 potential leading to 50% loss of hydraulic conductivity); a medium productivity, conservative 262 drought tolerance hydraulic strategy PFT (PFT-MC) represented by a medium V_{c,max25} and a less 263 negative P₅₀, turgor loss point and water potential at full turgor; and a high productivity, 264 aggressive drought tolerance hydraulic strategy PFT (PFT-HA) with a relatively high V_{c,max25} 265 266 and a very negative P₅₀. The mean of species-level trait data weighted by species abundance at the site were used to parameterize FATES-HYDRO. 267

268 **2.4 Model initialization**

Our model simulation is transient in terms of soil water content, leaf water content, carbon and water fluxes. The forest structure (plant sizes and number density) is fixed and is parameterized based on a vegetation inventory from Venturas et al. (2016). The soil texture and depth information are parameterization based on a national soil survey database

(https://websoilsurvey.sc.egov.usda.gov/App/WebSoilSurvey.aspx; Table S1). The soil moisture
is initialized with 50% of the saturation and the tissue plant water content is initialized so that it
is in equilibrium with the soil water potential. We run the model for 10 years based on 19501960 climate so that the simulated soil moisture, leaf water content, carbon and water fluxes are
not depending on their initial conditions.

278 **2.5 Live Fuel Moisture Content for model validation**

In this study, we used measured LFMC to validate simulated LFMC. FATES-HYDRO
does not directly simulate the LFMC. Thus, we estimated the LFMC based on simulated LWC.
The LWC in the model is calculated as follows,

282
$$LWC = \frac{fw - dw}{dw} * 100,$$
 (4)

where, *fw* is the fresh weight and *dw* is the dry weight, which are simulated within FATESHYDRO. Then, we estimated the LFMC within leaves and shoots (< 6 mm diameter) using the
empirical equation derived from shrub LFMC and LWC data including the three regenerative
strategies [seeder (S), resprouter (R) and seeder–resprouter (SR)], in summer, autumn and winter
from Fig. 4 and 5 in Saura-Mas and Lloret's study (2007) as follows (Fig. S4),

288 LFMC = 31.091 + 0.491LWC, (5)

The climate in Saura-Mas and Lloret's study is Mediterranean (north-east Iberian Peninsula),
which is consistent with the climate of our study area. LFMC was measured on our site
approximately every three weeks, concurrently with plant water potentials in 2015 and 2016.
LFMC measurement details can be found in Pivovaroff et al. (2019). For comparison with our
model outputs, we calculated the mean LFMC within leaves and shoots for each PFT weighted

by the species abundance (Venturas et al. 2016). Species abundance was calculated by dividing
mean density of a specific species by the mean density of all species.

296 **2.6 Climate drivers**

We forced the FATES-HYDRO model with 6-hourly temperature, precipitation, 297 downward solar radiation, and wind components. Historical climate data during 2012-2019, 298 which were used for FATES-HYDRO calibration, were extracted from a local weather station 299 (https://stuntranch.ucnrs.org/weather-date/). Historical and future climate data during 1950-2099, 300 301 which were used for simulations of LFMC by FATES-HYDRO model, were downloaded from 302 the Multivariate Adaptive Constructed Analogs (MACA) datasets (Abatzoglou & Brown 2012; 303 http://maca.northwestknowledge.net). The MACA datasets (1/24-degree or approximately 4-km; Abatzoglou & Brown 2012) include 20 ESMs with historical forcings during 1950-2005 and 304 305 future Representative Concentration Pathways (RCPs) RCP 4.5 and RCP8.5 scenarios during 2006-2099 from the native resolution of the ESMs. The gridded surface meteorological dataset 306 METDATA (Abatzoglou, 2013) were used with high spatial resolution (1/24-degree) and daily 307 timescales for near-surface minimum/maximum temperature, precipitation, downward solar 308 309 radiation, and wind components. Then we downscaled the MACA daily data to 6-hourly based on the temporal anomaly of the observed mean daily data to the hourly data for each day during 310 311 2012-2019. The model is driven by yearly CO_2 data obtained from Meinshausen et al (2011).

312 **2.7 Hypothesis testing**

To test H₀ (future climate change will decrease LFMC and consequently result in a longer fire season as determined by a critical threshold of LFMC), we compared the simulated mean LFMC, derived from modeled leaf water content, under the climate projections from 20 ESMs

under RCP 4.5 and 8.5. We then tested if the LFMC during the April-October dry season in the 316 historic period of 1960-1999 is significantly higher than that in the future period of 2080-2099. 317 For the fire season duration, we estimated the number of days per year below a critical threshold 318 of LFMC (79%). Similarly, we tested if the number of days per year below the critical threshold 319 of LFMC during the historical period are significantly different from that during the future 320 321 period. We used a bootstrapped approach (Jackson 1993) to test if the mean of LFMC or fire season duration are significantly different between these two periods. Specifically, we randomly 322 323 draw 10000 samples from the simulated residuals of LFMCs or fire season durations estimated 324 by 20 ESMs for these two periods under the null hypothesis that there is no difference in the mean. We then calculated p-values by comparing the simulated mean difference to the empirical 325 distribution of difference estimated from these 10000 samples. See Supplementary section 5.2 326 within Xu et al. (2019) for the details. 327

328 To test H_1 (warming has a stronger impact on LFMC than CO₂ fertilization), we 329 compared mean simulated LFMC and fire season length for three PFTs with/without CO₂ changes (fixed CO₂ at 367 ppm vs dynamic CO₂ concentrations from RCP 4.5 or RCP 8.5) and 330 warming. To remove the future warming trend, future temperature was replaced with historical 331 332 (1986-2005) temperature data for every 20 year period. Similarly, to test H₂ (the reductions in spring and autumn precipitation lead to a longer fire season as determined by LFMC), we 333 334 compared the model outputs of LFMC and fire season length for three PFTs with/without 335 precipitation changes. To test H₃ (the combined impacts of warming and precipitation on fire 336 season length are equal to the additive impacts of warming and precipitation change individually), we compared model outputs of LFMC and fire season length for three PFTs under 337 three scenarios: 1) without warming; 2) without precipitation changes; and 3) without warming 338

339	and precipitation changes. Finally, to test H4 (LFMC for plants with more conservative hydraulic
340	strategies will be more vulnerable to warming), we compared model outputs of LFMC and fire
341	season length across the three different PFTs with different hydraulic strategies.

342 **3. Results**

343 **3.1 Comparison between simulated and measured LFMC**

344 Our results showed that FATES-HYDRO was able to capture variation in the LFMC for 345 different PFTs and soil water content in 5-cm depth (Fig. 2 and S3), also for Chamise in 2018 346 (Fig. S5) although we had limited observed LFMC data. Specifically, the model was able to capture 96%, 86%, and 80% of the variance in observed LFMC for the period of 2015-2016 for 347 348 three PFTs, respectively (Fig. 2 b, d, f). The model was also able to capture the seasonal 349 dynamics of soil water content, LFMC, and LFMC below the threshold 79% in comparison to observed data (Fig. 2 a, c, e and S3). To validate that FATES-HYDRO is able to capture the 350 interannual variability of LFMC, we compared the simulated LFMC for PFT-LA with the long-351 term observations of LFMC for the chamise species (Adenostoma fasiculatum; Fig. S5). Our 352 353 results showed that the model is able to reasonably capture the seasonal and interannual variability for the period of 2006-2019 ($R^2=0.7$), although it underestimates peaks in LFMC in 4 354 of 14 years. 355

356 **3.2** Changes in the LFMC and fire season length from historical to future periods

Using the validated model driven by climate projections from 20 ESMs under greenhouse gas emission scenarios RCP4.5 and RCP 8.5, we found that the daily mean LFMC during the future period of 2080-2099 was projected to become significantly lower than that during the historical period of 1960-1999 for all three PFTs (Fig 3, P<0.000001). Our results also showed

that the spread among models increase with time, suggesting a larger uncertainty in the
projection into the future. Specifically, the histogram of daily mean LFMC during the AprilOctober dry season showed that there was a higher probability of low LFMC under future
climate conditions (Fig. S1). The daily mean LFMC decreased from 84.7%, 101.3%, and 78.4%
during the historical period of 1960-1999 to 81.0-82.8%, 96.3-98.8%, and 74.8-76.6% during the
future period of 2080-2099 under both climate scenarios for PFT-LA, PFT-MC, PFT-HA,
respectively (Fig 3).

368 Based on the projected LFMC, there was a significant increase in the fire season length with the critical threshold of LFMC from the historical period of 1960-1999 to the future period 369 of 2080-2099 for three PFTs. With the critical threshold of 79% LFMC, the fire season length 370 371 was projected to increase by 20, 22, 19 days under RCP 8.5 (Fig. 4 and Table S4), and to increase by 9, 11, 8 days under RCP 4.5 (Fig. 4 and Table S4). Our results also showed that the 372 373 spread among models increase with time, suggesting a larger uncertainty in the projection into 374 the future. The above results for mean LFMC and fire season length support hypothesis H₀ that future climate change will decrease LFMC and consequently result in a longer fire season, as 375 determined by critical thresholds for LFMC, for all three PFTs. 376

377 **3.3 Relative effects of individual climate changes on the length of the fire season**

In order to better understand the relative contribution to fire season length of different climate variables, we ran FATES-HYDRO for three PFTs using meteorological forcings that isolated and removed changes in individual specific variables. Our results showed that the increase in fire season length mainly resulted from warming, which led to 16-23 days (9.1-18.6%) per year increase in fire season length for the critical threshold of 79% LFMC under RCP 8.5 (Fig. 5). This is because warming is pushing Vapor Pressure Deficit (VPD) higher, resulting

in increased fire season length. For RCP 4.5, the warming contributed to 5-6 days (3.8-4.3%) per
year increase in fire season length (Fig. 5). We also found that elevated CO₂ concentrations
decreased fire season length with 6-7 days (3.5-4.8%) per year decrease in fire season length
under RCP 8.5 (Fig. 5). Under RCP 4.5, CO₂ increases led to 2-3 days (1.5-2.2%) per year
decrease in fire season length (Fig. 5). Because the impact of warming on fire season length was
stronger than the mitigation from CO₂ enrichment, our results support hypothesis H₁ (warming
has a stronger impact on LFMC than CO₂ fertilization).

Even though total precipitation was projected to increase in the future, lower precipitation in the spring and autumn (Fig. S2 a, b) led to 8-10 days (6.3-8.1%) per year increase in fire season length with the critical threshold of 79% LFMC under RCP 8.5 (Fig. 5). Under RCP 4.5, the precipitation changes contributed to 1-3 days (0.8-1.6%) increase in fire season length (Fig. 5). This result supported hypothesis H₂ that the reductions in spring and autumn precipitation lead to a longer fire season as determined by LFMC.

Our results showed that the combined impacts of warming and precipitation on fire season length were equal to the additive impacts of warming and precipitation change individually. This supported hypothesis H₃. Specifically, the combined changes in temperature and precipitation caused 24-33 days per year (15.6-26.8%) increase in fire season length with the critical threshold of 79% LFMC under RCP 8.5 (Fig. 5). Under RCP 4.5, the combined changes in temperature and precipitation caused a 6-9 days per year (4.8-6.1%) change in fire season length.

404 **3.4** Comparison of changes in fire season length among three PFTs under climate change

405	Regarding three PFTs under both climate scenarios, fire season length of PFT-HA was
406	the longest (167-176 days per year), while fire season length of PFT-MC was the shortest (114-
407	124 per year) during 2080-2099 (Fig. 4). However, the response of fire season length to warming
408	was strongest for PFT-MC. Specifically, for PFT-MC, warming under RCP 8.5 led to an increase
409	of 21.6% (22 days) in fire season length (Fig. 5 b) and warming under RCP 4.5 led to an increase
410	of 10.8% (11 days) in fire season length. For PFT-LA, warming under RCP 8.5 led to an increase
411	of 14.7% (19 days) in fire season length (Fig. 5 a) while warming under RCP 4.5 led to an
412	increase of 7.4% (9 days) in fire season length. Finally, for PFT-HA, warming under RCP 8.5 led
413	to an increase of 10.2% (18 days) in fire season length (Fig. 5 c) and 5.3% (8 days) in fire season
414	length with under RCP 4.5. Because PFT-MC has a more conservative hydraulic strategy with
415	less negative P50, turgor loss point and water potential at full turgor, this result supported
416	hypothesis H ₄ that the LFMC for plants with more conservative hydraulic strategy will be more
417	vulnerable to warming.

To validate our classification scheme, we compared these PFT-level results to those obtained with single-PFT and 2-PFT simulations finding that using the three PFTs defined by our cluster analysis gives a qualitatively different view of LFMC change than a single- or 2-PFT simulation. We found significant differences in the percentage changes of LFMC and fire season length between future period (2080-2099) and historical period (1960-1999) using three distinct PFTs, but no significant differences between PFTs in 2-PFT simulations under the different climate scenarios (Fig. S6).

425 4. Discussion

Low LFMC within shrub leaves and shoots increases the flammability and likelihood of
combustion, making it vitally important to monitor temporal variations in LFMC, especially

during the dry season (Dennison et al, 2008). The strong relationships between observed and 428 simulated LFMC of all PFTs (Fig. 2) suggested that the plant hydrodynamic model, FATES-429 HYDRO, could accurately estimate LFMC seasonal dynamics as a function of modeled leaf 430 water content, and consequently be useful to predict fire risks in Mediterranean-type climate 431 regions, although only small amount of validation data were used and the underlying assumption 432 433 that a shrub was analogous to a small tree. Based on the simulated monthly mean LFMC during 2006-2019 for PFT-LA, which includes the chamise species, we found that our model can 434 435 capture the seasonal variation and interannual variability, but underestimates the highest wet 436 season peaks in LFMC in 4 of 14 years (Fig. S5). This may cause biases for future projections while it would not highly affect the long-term trend of LFMC and fire season length. During the 437 future period (2080-2099) and the historical period (1960-1999), both periods displayed lower 438 values in the dry season (April - October), which is consistent with lower LFMC during the 439 summer-fall dry season, rather than the winter-spring wet season (Chuvieco et al, 2004; 440 Pellizzaro et al, 2007; Pivovaroff et al. 2019). Extremely low daily LFMC was more likely to 441 occur during the future period, which had higher temperature than the historical period. From the 442 historical to the future period, fire season length could increase by 5.2-14.8% under climate 443 444 change for chaparral shrub ecosystems (H_0). The fire season length was not validated, rather it was defined as number of days with LFMC below 79%. 445

Quantifying influences of climatic variables on LFMC is crucial to predicting future fire
risks (Dennison & Moritz, 2009). Our results showed that future warming was the most
important driver of LFMC. This finding suggested that warming would substantially push Vapor
Pressure Deficit (VPD) higher and decrease LFMC and strongly increase the fire season length,
which may greatly increase fire risks in the future (e.g. Dennison et al, 2008; Chuvieco et al,

2009; Pimont et al, 2019). CO₂ fertilization is expected to reduce stomatal conductance (Pataki et 451 al. 2000; Tognetti et al. 2000) and thus could mitigate the impacts of warming on LFMC. Our 452 results illustrated that, even though the CO₂ impact did cause a 3.5-4.8% reduction in fire season 453 length, the impact of warming on fire season length is about 5.6-13.8% larger than the CO₂ effect 454 $(H_1, warming has a stronger impact on LFMC than CO₂ fertilization). This result suggests that$ 455 456 CO₂ fertilization cannot offset the LFMC impacts from warming. The FATES-HYDRO model assumes a consistent stomatal sensitivity to CO₂ concentration across Mediterranean shrub 457 458 species. While Mediterranean shrub functional types in arid and semi-arid systems would vary in 459 their stomatal response in the real world (Pataki et al. 2000). Therefore, our model may overestimate/underestimate the CO₂ effect on stomatal conductance and its mitigating influence 460 might be smaller in reality for some species. 461

Previous studies implied that the timing of precipitation may have a strong impact on 462 subsequent LFMC (e.g. Veblen et al. 2000; Westerling et al. 2006; Dennison & Moritz 2009). In 463 464 this study, precipitation was also a key driver of LFMC under future climate conditions. Our results showed that, even though total precipitation was projected to increase, the reduction in 465 spring and autumn precipitation (Fig. S2) was projected to cause a longer fire season length (H₂, 466 467 the reductions in spring and autumn precipitation lead to a longer fire season as determined by LFMC; Fig. 5). This result was in agreement with a prior study indicating that spring 468 precipitation, particularly in the month of March, was found to be the primary driver of timing of 469 470 LFMC changes (Dennison & Moritz 2009). We also found that the combined impacts of warming and precipitation on fire season length were equal to the linearly additive impacts of 471 warming and precipitation change individually (H₃). Our results suggested that, when evaluating 472

473 future fire risks, it is critical that we considered the seasonal changes in precipitation and its474 interaction with the warming impact.

475 Modeled vegetation responses to environmental changes is a function of variation in plant 476 functional traits (Koven et al, 2020). The three PFTs represented in this study have similar patterns in LFMC in response to climate change during 1960-2099, but we did see some critical 477 478 differences. Specifically, the plant functional type PFT-MC with more conservative hydraulic strategy had the strongest responses to climate change (Fig. 5). This could be related to the fact 479 480 that the PFT-MC is a more conservative drought tolerant PFT in terms of hydraulic strategy with less negative P50, turgor loss point, and water potential at full turgor. The PFT-MC plants had a 481 482 relatively high saturated water content based on observed data (Fig 2) and the water within plant tissues thus changes more quickly in response to the environmental condition changes (H₄, 483 LFMC for plants with more conservative hydraulic strategies will be more vulnerable to 484 485 warming). However, the three different PFTs were coexisting at the same location in model 486 simulations, coexistence and heterogeneity in LFMC might impact fire behavior and fire season length. 487

Because the moisture content of live fuels (~50–200%) are much higher than that of dead 488 fuels (~7–30%), leaf senescence induced by drought stress and subsequent mortality are 489 490 potentially vital factors to cause large wildfires (Nolan et al. 2016, 2020). Thus drought-induced 491 canopy die-back and mortality could largely increase surface fine fuel loads and vegetation flammability, which can increase the probability of wildfire (Ruthrof et al. 2016). Since growth 492 493 and mortality are turned off in model runs by using a reduced-complexity configuration, it is 494 possible that vegetation density might decrease and LFMC could be conserved under future 495 scenarios. In addition, potential vegetation transitions (e.g., shrubs to grassland and species

composition changes) might substantially affect flammability and thus fire intensity and
frequency. In this study, we used the static mode of FATES-HYDRO to simulate LWC dynamics
under climate change. If we need to assess how the leaf senescence and vegetation dynamics will
impact the fire behavior, we can use the same model with dynamic mode to assess their impacts
on fire behaviors under future drought and warming conditions.

501 Application of a hydrodynamic vegetation model to estimate LFMC dynamics could potentially benefit wildfire modeling at the fine-scale, landscape-scale, and global-scale. This is 502 503 because LFMC is one of the most critical factors influencing combustion, fire spread, and fire consumption while previous wildfire models mainly focus on impacts of dead fuel moisture, 504 weather conditions on wildfire, fuel loads, and representation of live fuel moisture (Anderson & 505 Anderson 2010; Keeley et al. 2011; Jolly & Johnson 2018). The implications of this are that fire 506 potential will vary with plant water potential and uptake from soils, photosynthetic and 507 508 respiratory activity, carbon allocation and phenology with variability across species and over 509 time (Jolly & Johnson 2018). Therefore, future work to incorporate LFMC dynamics in wildfire models could potentially play a vitally important role in the future studies of wildfire modeling 510 under climate change. 511

512 **5.** Conclusions

A hydrodynamic vegetation model, FATES-HYDRO, was used to estimate leaf water status and thus LFMC dynamics of chaparral shrub species in southern California under historical and future conditions. FATES-HYDRO model was validated using monthly mean LFMC for three PFTs. The fire season length was projected to substantially increase under both climate scenarios from 1960-1999 to 2080-2099. This could increase wildlife risk over time for chaparral shrubs in southern California. Our results showed that temperature was the most

519	important driver of LFMC among all climatic variables. The LFMC estimated by the FATES-
520	HYDRO model offered a baseline of predicting plant hydraulic dynamics subjected to climate
521	change and provided a critical foundation that reductions in LFMC from climate warming may
522	exacerbate future wildfire risk. Longer fire season might have a significant impact on overall
523	public health and quality of life in the future.
524	
525	Data availability. LFMC measurement data can be found in Pivovaroff et al. (2019). Species
526	abundance data can be found in Venturas et al. (2016). All other data are available within this
527	paper and in the Supplement.
528	
529	Supplement. The supplement related to this article is available online at
530	
531	Author contributions. WM, LZ, and CX were involved in designing the study. WM and CX
532	conducted the data analysis and ran model simulations. All authors contributed to writing,
533	reviewing, and improving the manuscript.
534	
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873 Fig 1. Hierarchical cluster analysis of allometry and hydraulic traits for eleven chaparral shrub species used to define three plant functional types at Stunt Ranch. The plant functional types 874 with a low productivity and an aggressive drought tolerance hydraulic strategy (PFT-LA) was 875 defined based on traits of red shank (Adenostoma sparsifolium - As), toyon (Heteromeles 876 arbutifolia - Ha), chamise (Adenostoma fasciculatum - Af), big berry manzanita (Arctostaphylos 877 glauca - Ag); the plant functional types with a high productivity and an aggressive drought 878 tolerance hydraulic strategy (PFT-HA) was defined based on traits of mountain mahogany 879 (Cercocarpus betuloides - Cb), greenbark ceanothus (Ceanothus spinosus - Cs), buck brush 880 (Ceanothus cuneatus - Cc), hollyleaf redberry (Rhamnus ilicifolia - Ri); the plant functional 881 types with a medium productivity and an conservative drought tolerance hydraulic strategy 882 (PFT-MC) was defined based on traits of laurel sumac (Malosma laurina - Ml), scrub oak 883 (Quercus berberidifolia - Qb), sugar bush (Rhus ovata - Ro). 884





Fig.2 Simulated and observed monthly live fuel moisture content and related R² values for three
PFTs (refer to Figure 1 for explanation of the PFTs).



Fig.3 Temporal changes in daily mean live fuel moisture content (black solid line) and 95%
confidence interval (black dash-dot line) from 1960 to 2099 for three PFTs (refer to Figure 1 for
explanation of the PFTs) under climate scenario RCP 4.5 and 8.5 with 20 Earth System Models
considering all climatic variables changes. The P values were calculated using bootstrap
sampling to test whether the daily mean live fuel moisture content across different models during
the future period (2080–2099) was significantly lower than that during the historical period
(1960-1999). The grey horizontal dotted line represents the ensemble mean for 2080–2099.



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Fig.4 Temporal changes in average number of days per year of live fuel moisture content below 898 79% (black solid line) and 95% confidence interval (black dash-dot line) from 1960 to 2099 for 899 three PFTs (refer to Figure 1 for explanation of the PFTs) under climate scenario RCP 4.5 and 900 8.5 with 20 Earth System Models considering all climatic variables changes. The P values were 901 calculated using bootstrap sampling to test whether the number of days across different models 902 during the future period (2080–2099) was significantly higher than that during the historical 903 period (1960-1999). The grey horizontal dotted line represents the ensemble mean for 2080-904 905 2099.



Fig.5 Differences on number of days per year of live fuel moisture content below 79% from
2080 to 2099 for three PFTs (refer to Figure 1 for explanation of the PFTs) under climate

- scenario RCP 4.5 and 8.5 between considering all climatic variables changes and without
- 912 considering CO₂, precipitation, temperature, and precipitation & temperature changes.