Towards transboundary networks of climate-smart marine reserves in the Southern California Bight

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102 Abstract

103

Climate-smart conservation addresses the vulnerability of biodiversity to climate change 104 impacts but may require transboundary considerations. Here, we adapt and refine 16 105 biophysical guidelines for climate-smart marine reserves for the transboundary California 106 Bight ecoregion. We link several climate-adaptation strategies (e.g., maintaining connectivity, 107 representing climate refugia, and forecasting effectiveness of protection) by focusing on kelp 108 forests and associated species. We quantify transboundary larval connectivity along ~800 km 109 110 of coast and find that the number of connections and the average density of larvae dispersing through the network under future climate scenarios could decrease by ~50%, highlighting the 111 need to protect critical steppingstone nodes. We also find that although focal species will 112 generally recover with 30% protection, marine heatwaves could hinder subsequent recovery in 113 the following 50 years, suggesting that protecting climate refugia and expanding the coverage 114 of marine reserves is a priority. Together, these findings provide a first comprehensive 115 framework for integrating climate resilience for networks of marine reserves and highlight the 116 117 need for a coordinated approach in the California Bight ecoregion.

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119 Introduction

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Marine reserves can rebuild the biomass of overfished species¹, conserve biodiversity², and 121 enhance the resilience and adaptive capacity of ecosystems to climate impacts³⁻⁷. However, 122 delivering large-scale benefits requires networks of marine reserves that are functionally 123 interconnected, and are large enough to protect the underlying biophysical processes that 124 maintain species distribution and composition⁸. A rich literature exists on biophysical 125 guidelines for designing networks of marine reserves for fisheries, conservation, and climate-126 adaptation objectives⁹⁻¹², but most are limited to analysis within country or state boundaries. 127 By contrast, ecoregion-scale planning efforts may span thousands of kilometres and, in many 128 cases, cross multiple national or international jurisdictions^{13,14}. Consequently, before designing 129 networks of transboundary marine reserves, planners need to develop shared biophysical 130 guidelines and comprehensive spatial analyses across borders¹⁵. 131

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Climate change is one of the main threats to marine ecosystems¹⁶ that could be partially
addressed by large-scale, coordinated, climate-smart networks of marine reserves^{11,17,18}.
Although marine reserves cannot directly abate climate-change threats, they can indirectly

mitigate climate-change impacts by promoting ecological resilience^{3-5,19}. Climate-smart 136 conservation is a multiple-step approach that addresses the vulnerability of species and 137 ecosystems to changes in climate and ocean chemistry, and supports resilience of populations 138 and ecosystems¹⁷. Climate-adaptation strategies include protecting areas that act as climate 139 refugia^{11,18,20}, maintain ecological connectivity to ensure metapopulation persistence²¹, 140 facilitate species' range shifts, and recover important species for ecosystem functioning (e.g., 141 predators)²². Complementary strategies include supporting ecosystem resilience by addressing 142 threats not directly abated by marine reserves¹¹ (e.g., pollution and other impacts) and 143 protecting and restoring habitats that could mitigate some of the effects of climate change (e.g., 144 through carbon sequestration). Because global conservation targets aim to protect 30% of 145 marine habitats by $2030^{23,24}$ while adapting to climate change, there is a need to integrate 146 multiple adaptation strategies for designing networks of climate-smart reserves. 147

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Given the highly dynamic nature of the oceans, biophysical modelling of larval dispersal is an 149 essential tool to inform transboundary conservation^{25,26}. However, these models should 150 consider the implications of climate change on larval dynamics, such as changes in dispersal 151 distances and the availability of suitable habitats for settlement^{21,27}. These considerations are 152 153 essential because transboundary dispersal may be critical for metapopulation persistence. Notably, certain areas may be less impacted by climate change and act as climate refugia²⁸, 154 155 providing food, shelter, and habitat, despite future changes. Identifying climate refugia at large spatial scales can be challenging, requiring the use of ecosystem attributes (resistance, 156 resilience, persistence²⁹) or environmental proxies (e.g., micro climates)³⁰. If we map these 157 areas, we can prioritize their protection and assess changes in larval connectivity for future 158 159 scenarios. Climate change may undermine the effectiveness of transboundary networks of marine reserves to facilitate recovery of exploited species. Thus, it is a priority to assess 160 whether proposed protection targets^{23,24} will facilitate recovery of overexploited species in the 161 future and whether increased protection or alternative strategies will be necessary to ensure 162 163 their recovery.

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165 The Southern California Bight ecoregion (henceforth "California Bight") in the northwest 166 Pacific Ocean — shared between the state of California, USA, and the Peninsula of Baja 167 California, Mexico — has a long history of research cooperation and binational environmental 168 agreements^{31,32}. It is considered a marine climate-change "hotspot" – rapidly warming ocean 169 regions that are natural laboratories for evaluating climate adaptation options³³. Recent marine

heatwaves³⁴⁻³⁷ and prolonged hypoxic events³ exemplify the impacts of climate variability and
environmental extremes on species, ecosystems, and coastal economies of this region.
Documentation of these changes includes mass mortality events and range shifts of
economically or ecologically important species^{3,34,37-39}. This threatens the conservation and
economic sustainability outcomes that both countries seek to deliver through marine zoning.

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In 2012, California implemented a network of marine protected areas covering 16% of state waters, with more than half being fully protected marine reserves. This network was based on stakeholder input and biophysical guidelines to ensure ecological connectivity, habitat representation and replication, while balancing users' needs⁴⁰. However, the establishment of marine protected areas did not include climate-adaptation objectives⁴¹. Moreover, the network did not consider the transboundary nature of the region, where many species move across the USA-Mexico border through larval dispersal and adult movement.

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By contrast, although Baja California has extensive coastal areas protected, there is a lack of 184 an integrated network of marine reserves, with less than 1% of the coastal waters fully 185 protected^{20,42}. In this region, coastal resource management is based on territorial user rights 186 187 granted to fishing cooperatives and independent permit holders, with successful local cases of community-based marine reserves sparking other cooperatives' interest⁴². Additional 188 189 community-led marine reserves are awaiting government approval, while environmental NGOs, scientists, fishing cooperatives, and governmental agencies are promoting a region-190 wide marine spatial planning process⁴². 191

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193 Here, we first adapt and refine 16 biophysical guidelines for climate-smart, transboundary marine reserve design for the California Bight ecoregion - spanning over eight degrees of 194 195 latitude. We used kelp forest ecosystems (dominated by giant kelp, *Macrocystis pyrifera*) and focal species of fishes and invertebrates of commercial and ecological importance to identify, 196 analyze, and map areas that integrate and meet the proposed climate-smart guidelines. Then 197 we explore the magnitude and importance of transboundary connectivity in the region and ask 198 199 whether binational connections will be lost under future climate scenarios. We also ask whether sea surface temperature variability can be used as a proxy for climate refugia and whether 200 marine heatwaves in the following 50 years will undermine the effectiveness of marine reserves 201 202 for facilitating recovery of vulnerable species in the California Bight. Our research can inform

the delivery of networks of climate-smart marine reserves by 2030 in the California Bight andother regions.

- 205
- 206 Methods
- 207
- 208 Study area
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The California Bight ecoregion is located in the southern California Current System in the 210 211 northeast Pacific Ocean and spans the USA-Mexico international border, from Point Conception, California, USA, in the north to Punta Abreojos, Baja California Sur, Mexico, in 212 the south⁴³. This highly productive ecoregion is in a transitional zone between the southward-213 flowing, cold, nutrient-rich California Current and the northward-flowing, warm, nutrient-214 depleted Davidson Current⁴⁴. This transboundary ecoregion is characterized by strong 215 latitudinal gradients in environmental conditions and oceanographic features that support a 216 diverse assemblage of species and habitats⁴⁵. We divided the California Bight into four sub-217 regions: southern California, northern and central Baja California, and Guadalupe Island. These 218 four sub-regions represent geographic borders (the USA-Mexico border) and distinct 219 biogeographic areas where species composition varies because of environmental conditions⁴⁶. 220

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222 Developing and integrating biophysical guidelines

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We developed biophysical guidelines (Table 1) for designing transboundary networks of climate-smart marine reserves for biodiversity conservation, fisheries replenishment, and climate-adaptation objectives in the California Bight. We compiled, adapted, and refined the guidelines using criteria developed for California, Mexico, and other regions^{9,10,40,47}. Our work does not provide an extensive review of the proposed guidelines or their ecological rationale, as this has been addressed in previous work^{9,10}. Instead, we summarise our findings in the context of transboundary and climate-smart spatial planning.

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Our work builds on a workshop held in 2017, which developed biophysical design guidelines for the Pacific region of Baja California⁴². Following this first workshop, five working groups comprising researchers from Mexico, the USA and Australia, fishing cooperatives, governmental agencies, and non-profit organizations in Mexico and the USA further developed the principles and their application to the California Bight (Table 1). Finally, following a

second workshop held in 2019, we conducted spatial analysis and developed maps that 237 integrate and meet some of the proposed guidelines for the California Bight, focussing on giant 238 kelp (M. pyrifera) forests (henceforth "giant kelp") and six focal species of commercial and 239 ecological importance associated with giant kelp. Focal species included California Sheephead 240 (Semicossyphus pulcher), sea basses (Paralabrax clathratus and P. nebulifer), whitefish 241 (Caulolatilus princeps), spiny lobster (Panulirus interruptus), abalone (Haliotis spp.), sea 242 urchins (Mesocentrotus franciscanus and Strongylocentrotus purpuratus), and sea cucumbers 243 (Apostichopus spp. And Parastichopus spp.). We did not always use the same focal species 244 245 among principles and analysis (see, Supplementary Table 1) because of differences in the availability of data, and in some cases, we used data from multiple taxa as composite focal 246 species. 247

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The proposed transboundary climate-smart guidelines fall into five major categories: (1) habitat 249 representation and replication; (2) protecting critical and unique areas; (3) incorporating 250 connectivity; (4) allowing time for recovery; and (5) minimizing and avoiding threats (Fig. 1). 251 Marine habitats are broad indicators of the distribution of biodiversity⁴⁸, which are relatively 252 easy to map, and if adequately represented (e.g., protect enough of each habitat) can effectively 253 protect biodiversity and replenish overfished populations^{9,10}. Moreover, each habitat should be 254 replicated in the network in case a large-scale disturbance impacts part of the system⁴⁹. It is 255 256 also important to represent and replicate habitats or areas more resistant and resilient to climate stressors (climate refugia), which can replenish impacted populations and habitats^{9,10}. 257

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To facilitate recovery of focal species populations, the network should also protect areas critical 259 for their life cycles, such as those required for spawning, nesting, or breeding^{9,10}. Notably, 260 because most marine species are structured as metapopulations, connected through movements 261 262 of adults, juveniles, or larval dispersal, ensuring the metapopulation persistence of focal species requires incorporating ecological connectivity^{9,10,47,50}. It is also crucial to consider changes in 263 ocean conditions that could modify patterns of larval dispersal and recruitment and precipitate 264 range shifts among species. Determining how much time overexploited species need to recover 265 is essential to assessing the network's effectiveness, especially if this is done while evaluating 266 the projected impacts of climate change for vulnerable species. Finally, we need to consider 267 threats that marine reserves can abate, versus those that reserves cannot abate, therefore 268 requiring alternative strategies to support ecosystem resilience^{19,51}. 269

271 Binational habitat mapping

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We mapped the distribution of intertidal, estuarine, subtidal, and deep-sea habitats in the 273 territorial seas (within 12 nautical miles of the coast) of the California Bight (Supplementary 274 Table 2 and 3). We extracted depth contours (30, 100, and 200 m) from the General 275 Bathymetric Chart of the Oceans⁵² and the California Department of Fish and Wildlife (CDFW, 276 2010 version) using ESRI ArcGIS Pro v10.8. We then used these contours to classify subtidal 277 habitats based on depth categories (0-30, 30-100, 100-200, over 200 m). We obtained intertidal, 278 279 estuarine and subtidal habitat polygons for southern California (CDFW, 2010 version) and northern Baja California from published work⁵³. There was no available coastline or habitat 280 mapping in the sub-regions of central Baja California and Guadalupe Island. We followed 281 Arafeh-Dalmau, et al. ⁵³ and digitized the coastline and mapped intertidal and subtidal habitats 282 by visualizing Google Earth historical images. 283

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We then combined existing maps of giant kelp distribution for California (CDFW, 2010 285 version) and Baja California⁵³ with a 35-year satellite time series that maps the distribution and 286 persistence of giant kelp at 30-m² grid resolution, and classifies giant kelp persistence into three 287 classes (high, mid, and low). For more details see Arafeh-Dalmau, et al.²⁰. Highly persistent 288 giant kelp can be a good proxy for climate refugia because they have endured through time 289 despite multiple cycles of marine climate oscillations²⁰. We classified giant kelp polygons that 290 did not overlap with persistence maps as having low persistence. We also collected information 291 and mapped the distribution of tidal flats in southern California⁵⁴ and eelgrass (*Zostera marina*) 292 for northern and central Baja California from existing information (Pronatura Noroeste A.C.) 293 294 and in situ presence-absence surveys combined with low-altitude drone imagery (for more details, see Supplementary Table 3). Finally, we catalogued geomorphic features data⁵⁵ to map 295 296 submarine canyons.

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298 Vulnerability of habitats and focal species to human threats

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We conducted an expert-knowledge survey to quantify the vulnerability of six marine habitat types (estuaries, rocky intertidal, seagrass, kelp forest/rocky reefs, deep sea and pelagic) to eight major threats (marine heatwaves, ocean acidification, hypoxia, sea-level rise, storms, resource extraction, pollution, and physical habitat destruction) for the California Bight. The survey allowed experts to score the vulnerability of each habitat to each threat. Options

included no threat, low, moderate, and high threat (coded 0, 1, 2, and 3, respectively; for moredetails see Supplementary Methods).

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308 We complemented and expanded the expert-knowledge survey through a review of published studies for the California Bight that empirically examined the vulnerability and recovery rate 309 of five focal species (California Sheephead, lobster, abalone, sea urchin, giant kelp; 310 Supplementary Table 1) following marine heatwaves, hypoxia, and ocean acidification. We 311 scored vulnerability to climate stressors that a species experiences as high for lethal effects, 312 313 medium for extensive sublethal effects, and low for limited sublethal effects. We scored a species recovery rate from climatic stresses as rapid (< 1 year), moderate (2-5 years), or slow 314 (> 5 years) (for more details, see Supplementary Methods). 315

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317 Sea surface temperature variability as a proxy for climate refugia

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To identify potential climate refugia, we used two proxies: temperature variability³⁰ and giant 319 kelp persistence²⁰. We analyzed patterns in their spatial variation and their correlation across a 320 1-km² grid in the California Bight. We conducted this analysis to ascertain whether sea surface 321 322 temperature variability is a comprehensive proxy of climate refugia for each sub-region in the California Bight. We obtained 17 years (2003-2019) of daily SST data from the Aqua-MODIS 323 satellite at 1-km² grid resolution, accessed through the NOAA-ERDDAP data repository 324 (https://coastwatch.pfeg.noaa.gov/erddap/index.html). We estimated an annual cycle using a 325 326 30-day weighted moving average smoothing window for each year to create an annual climatology. We then used these annual cycles to estimate SST variability by computing the 327 328 variance relative to the moving mean of the annual cycle (see Supplementary Fig. 1, for example). This SST variability metric represents high-frequency (24-h or faster) temperature 329 variability. We then computed the mean SST variance across the 1-km² grid to evaluate the 330 accuracy of our model for SST variability with in situ sensors (a series of MiniDOT 331 temperature/dissolved-oxygen sensors [PME Inc.] across Baja California⁵⁶) and compared 332 variability at sub-daily intervals. We then computed the mean giant kelp persistence²⁰ over the 333 same 1-km² grid cells. We examined the correlation (linear least-squares regression) between 334 these datasets for each sub-region, excluding Guadalupe Island, which does not have giant kelp. 335 336

337 Transboundary movement of juveniles and adults into critical areas

To assess transboundary connectivity, we reviewed and mapped the transboundary movements 339 of sharks, bony fishes, seabirds, marine mammals, and sea turtles between southern California 340 and the three sub-regions in Baja California. We reviewed data from published papers and 341 public databases (e.g., Tagging of Pacific Predators (TOPP) animal tracking dataset) for the 342 period 1999-2019 using a combination of keywords: (1) movement or migration, (2) adults or 343 344 juvenile, and (3) shark, bony fish, sea bird, marine mammals, or sea turtles. Because most studies tagged animals in USA waters, we considered connectivity only for organisms released 345 in USA waters that moved into Mexican juvenile or adult habitats (e.g., spawning, breeding, 346 347 nursery, resting, or foraging habitats). We aggregated the number of tagged studies that released organisms in southern California that were subsequently recorded in the three sub-348 regions in Baja California into 0.25° grid cells. Grid cells with high numbers of tagged animals 349 recorded in Baja California were identified as essential areas for juvenile and adult 350 transboundary connectivity. 351

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353 Transboundary larval dispersal model and impact of climate change

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To further explore transboundary connectivity, we implemented the AGRIF version of the Regional Ocean Modelling System (ROMS; for more details, see Supplementary Methods, Supplementary Fig. 2) to simulate passive spores and larval dispersal (henceforth "larval dispersal") for four focal species (California Sheephead, abalone, sea urchin and giant kelp; Supplementary Table 1) in the California Bight. We obtained information about spawning time and planktonic larval duration (PLD) from the literature (Supplementary Table 4), and for simplicity rounded to the nearest week when PLD exceeded 7 days.

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To measure larval connectivity, we divided the coast into 54 polygons, each covering 20 km 363 of latitude and limited by the 200 m isobath (Supplementary Fig. 3). This isobath represents 364 the edge of the continental shelf that is a limit for coastal environments, where most fishing 365 takes place⁴³. In the centroid of each polygon, we released 1,000 virtual larvae at the start of 366 each month of the year and followed their trajectories hourly for 60 days (2 months). We 367 368 imported hourly coordinates for each modelled particle into MATLAB (Mathworks). We identified the intersection between particles and each polygon at the end of the PLD with a 369 selection-by-location function. We generated connectivity matrices reflecting the proportion of 370 larvae that settled in each polygon relative to the total number of larvae released at each site. 371 372 We averaged matrices for the larval release dates within each month during each species'

spawning season. We calculated local retention as the proportion of larvae released within a 373 polygon remaining within the natal area at the end of the PLD for each species. We explored 374 connectivity matrices for each season using graph theory and a spatial network approach using 375 the software GEPHI⁵⁷, where nodes represent larval release sites and links represent directional 376 377 larval dispersal probabilities. We estimated network density to compare changes in cohesiveness or saturation that relate to functional attributes, such as resilience⁵⁸. We defined 378 379 density as the number of links observed divided by the maximum number of possible links⁵⁸, representing the probability that any given tie between two random nodes is present⁵⁹. 380

381

We simulated two contrasting scenarios to investigate the potential effect of climate change on 382 larval connectivity due to reduction of PLD with increased temperatures and the reduction of 383 recruitment habitat due to climate change, since both could significantly alter metapopulation 384 dynamics^{21,27}. In the first or "Current" scenario, we downscaled the larval connectivity matrices 385 to the polygon unit (following the approach described by Alvarez-Romero, et al. ²¹) based on 386 two factors: probability of connections between two polygons according to the connectivity 387 matrix based on the PLD reported for each species in the literature, and the total area with giant 388 kelp found within each polygon. The second or "Future" scenario employed shortened 389 390 connectivity matrices due to warming and consequent restriction of giant kelp to highly persistent habitats, defined as potential climate refugia²⁰. We calculated the reduction in PLD 391 in fish and invertebrates following a 3°C increment in SST using previously described 392 methods^{21,60} (Supplementary Table 4). This increase in temperature is extreme, yet not unlikely 393 for the end of the 21st century under the highest IPCC Shared Socio-Economic Pathway (SSP5-394 $(8.5)^{61}$. 395

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397 Simulating recovery time for focal species

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To project the expected effectiveness of transboundary marine reserve networks for recovery of exploited populations, in the absence of climate-change impacts, we simulated the effect of marine reserves on six focal species within the region (sea bass, whitefish, lobster, abalone, sea cucumber and sea urchin; Supplementary Table 1) using a deterministic, discrete-time logistic growth model with spatially implicit reserve and fishing zones:

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$$X_{t+1} = X_t + \left(rX_t \left(1 - \frac{X_t}{K} \right) \right) - \left((1 - R)X_t F_{MSY} \right),$$

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where X_t represents biomass at time t, r is the intrinsic growth rate, and K is the carrying 407 capacity. The last term represents harvesting of biomass outside the reserve, where the (1 - R)408 parameterization corresponds to the portion of biomass outside the reserve. We test three 409 different scenarios of reserve coverage: R = (10%, 30%, 100%). The first two scenarios are 410 representative of commonly cited protection targets, while the third scenario provides an upper 411 bound of maximum attainable protection. We estimated population parameters and fishing 412 mortality (F_{MSY}) by applying a catch-only data-limited stock assessment method⁶² to catch data 413 from 2000-2015 in Baja California, from Comisión Nacional de Acuacultura y Pesca 414 (CONAPESCA) (Supplementary Table 5, Supplementary Fig. 4). We ran all simulations for 415 50 years, with initial biomass set at 20% of carrying capacity and fishing mortality outside the 416 reserve held constant at F_{MSY} . We considered a population "recovered" when the population 417 size was within 90% of the theoretical equilibrium size (\overline{X}): 418

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420
$$\bar{X} = \frac{K[r - (1-R)F_{msy}]}{r}$$

421

422 Considering the impact of marine heatwaves on focal species biomass and recovery423

For a subset of focal species of invertebrates with limited movement (abalone, sea cucumber 424 and sea urchin; Supplementary Table 1, 6) vulnerable to marine heatwaves, we explored 425 recovery for three climate-change scenarios by running a stochastic version of the model. We 426 simulated the impact on the biomass of species based on the probability of a year experiencing 427 marine heatwaves with a cumulative intensity at least as strong as those that impacted the 428 California Bight in 2014-2015^{34,36-38}. We used 17 years of giant kelp forest community data 429 (1999-2015), which integrate four different monitoring programs for the California Bight³⁷, to 430 model the rate of change of the density of the focal species following the 2014-2015 extreme 431 marine heatwaves (henceforth "marine heatwaves"). We excluded monitoring data north of 432 latitude 33.8°N and west of longitude 118.7°W because this area is subject to colder average 433 temperatures, forms a separate sub-bioregion⁶³, and giant kelp forest communities there are 434 less impacted by marine heatwaves⁶⁴ (for more details, see Supplementary Methods). 435

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We used 0.25° grid cell Optimum Interpolation Sea Surface Temperature data set⁶⁵ to estimate
the average annual cumulative intensities registered in 2014-2015 marine heatwaves, based on

the climatology for 1983-2012. Then we estimated projected marine heatwaves for the 20202100 period for three climate scenarios generated under each of three SPPs⁶⁶ – SSP1-2.6,
SSP2-4.5, SSP5-8.5 – from a multi-model ensemble mean derived from 11 Earth-System
models from the Coupled Model Intercomparison Project Phase 6 (for more details see
Supplementary Methods).

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We derived the probability of a marine heatwave year using a generalized linear model with a binomial link function, fitted to marine heatwave years for the three climate scenarios. Then we used the empirically derived density change effect of a marine heatwave on abalone, sea urchins, and sea cucumber to yield the model:

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450
$$X_{t+1} = (1 - \delta \Phi_t) X_t + \left(r X_t \left(1 - \frac{X_t}{K} \right) \right) - ((1 - R) X_t F_{MSY}),$$

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where $\delta \in [0, 1]$ represents the density-reducing effect of the MHW, and $\Phi_t = 1$ if the year *t* has a MHW and $\Phi_t = 0$, otherwise. We ran 10,000 independent simulations of this model for each of the three species and three climate scenarios. All data analyses and simulations were conducted in R 4.0.4⁶⁷.

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457 Mapping threats

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We mapped climate and land-based threats that marine reserves cannot directly abate but can build resilience to (henceforth "threats") at a 1-km² grid resolution in the California Bight using the most comprehensive dataset available for the cumulative human impact (see Halpern, et al. ⁶⁸). We classified those cells with high cumulative impacts (top 10%) for each sub-region as "highly threatened".

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For each of the three sub-regions in Baja California, we mapped two extractive activities that marine reserves can directly abate: small-scale commercial fishing and recreational fishing (henceforth "fishing"). We prioritized the analysis for these regions because they have less spatial information available. We mapped the total catch of small-scale fisheries (tonnes) at one-hectare grid resolution (Supplementary Table 7) inside authorized fishing polygons using 19 years (2000-2018) of catch information referenced to each fishing cooperative and individual permit holder (hereafter "concessions") from CONAPESCA. Each fishery has a

designated concession where cooperatives can extract specific resources (for more details see
Supplementary Methods). Then, we estimated the probability of recreational fishing sites or
banks in a 100-m² grid resolution based on georeferenced commercial and non-commercial
sport fishing maps (e.g., FISH.n.MAP CO., Baja California North, Sportfishing Atlas Baja
California Edition) for the three sub-regions in Baja California (for more details see
Supplementary Methods).

- 478
- 479 **Results**
- 480

We provide 16 biophysical guidelines for designing transboundary networks of climate-smart 481 marine reserves adapted for the California Bight and recommendations for their application 482 (Table 1). The guidelines provide recommendations for habitat representation and replication 483 (Supplementary Table 8), protecting critical and unique areas, incorporating connectivity, 484 allowing time for recovery, and minimizing or avoiding local threats. Instead of addressing 485 measures for adapting to changes in climate and ocean chemistry separately^{9,10}, we integrate 486 climate-adaptation strategies within each of the other principles, to achieve a climate-smart 487 network. 488

489

490 Habitat mapping

491

We produced maps of the distribution of 31 coastal and island habitats from intertidal to deep-492 493 sea habitats for the four sub-regions (Fig. 2, Supplementary Table 3). Although southern 494 California covers fewer degrees of latitude than northern and central Baja California, it 495 represents almost half (~46%) of the area of the California Bight. We mapped rocky intertidal and sandy beaches for ~2700 km of coastal and island coastline. Our habitat mapping includes 496 497 biogenic habitats such as eelgrass and surfgrass (*Phyllospadix* spp.) and different levels of giant kelp persistence. We also mapped subtidal sandy and rocky habitats at different depths and 498 submarine canyons in all regions. We found no seamounts, guyots or other geomorphic features 499 of importance for biodiversity in the territorial sea of the California Bight. We could not map 500 surfgrass and rocky habitats deeper than 30 m for the three regions in Baja California. Finally, 501 we found no giant kelp or estuarine habitats in Guadalupe Island. 502

503

504 Vulnerability of habitats and focal species to human threats

We received 55 responses to our expert-knowledge survey. Respondents ranked estuaries and rocky intertidal habitats as the most threatened habitats, whereas pelagic and deep-sea habitats ranked as the least threatened (Fig. 3a). Across habitats, marine heatwaves and resource extraction were identified as the most serious threats, while storm events and sea-level rise were the least threatening (Fig. 3b). Moreover, although the respondents ranked warming as the most threatening stressor, non-climate related threats, on average, scored higher (2.23 \pm 0.17) than climate threats (1.94 \pm 0.18).

513

514 Our risk assessment indicated that mobile species, such as California Sheephead and lobsters, are least vulnerable and recover fastest because they are least sensitive to acute climate 515 stressors. In contrast, sessile species like giant kelp, or species with limited mobility, such as 516 abalone and sea urchins, exhibit high and intermediate vulnerability to at least one climate 517 stressor (Fig 3c). The recovery from climate stressors is intermediate for giant kelp, but slower 518 for abalone and sea urchins. The combination of higher vulnerability and slow recovery renders 519 these sessile or limited mobility benthic invertebrates most vulnerable to climate stressors (see 520 Supplementary Results for detailed justification). 521

522

523 Sea surface temperature variability as a proxy for climate refugia

524

Analysis of remote sensing SST and giant kelp canopy data revealed that high kelp persistence 525 coincides with low temperature variability in southern California (p = 0.042), has no relation 526 with temperature variability in northern Baja California (p = 0.745), and correlates with high 527 temperature variability in central Baja California (p = 0.007) (Fig. 4). We also found that SST 528 529 variability performs well compared to the observed high frequency bottom temperature variance ($R^2 = 0.87$; Supplementary Fig. 5). Even when we excluded an outlier site (Punta 530 531 Prieta) which has the highest temperature variability of all sites, we found a correlation coefficient $R^2 > 0.6$. Our results suggest that in the absence of other metrics, SST variability 532 may identify potential refugia for giant kelp and possibly kelp-associated species in the 533 southern portion of the region (central Baja California). 534

535

536 Transboundary movement of juveniles and adults

537

We found movement data for juvenile and adult individuals for seven shark, three bony fish,four seabird, four marine mammal, and one sea turtle species (Supplementary Table 9). Focal

species are relatively large and migratory, with reported connectivity distances ranging from 540 hundreds (e.g., white seabass, yellowtail) to thousands of kilometers (e.g., white sharks, blue 541 whales; Supplementary Table 9) and temporal scales of two weeks (e.g., bottlenose dolphin, 542 Black-vented shearwater) to over two years (white seabass). Greater connectivity was reported 543 in northern Baja California, from north of Ensenada (31.8°N) and near the USA/Mexico border, 544 followed by Punta Colonet (30.9°N), and areas in central Baja California, with important 545 locations near the southern limit of the California Bight in Bahía Asunción (27.1°N). We also 546 found areas of high connectivity between southern California and Guadalupe Island (Fig. 5). 547 548

Tagging studies highlight transboundary nursery and foraging areas for white sharks in near-549 shore regions of central Baja California (Supplementary Table 9) and spawning areas for 550 California Scorpionfish and Vermilion Rockfish in the USA-Mexico border. Breeding areas 551 are also reported for Laysan albatross in Guadalupe Island, post-breeding dispersal areas for 552 black-footed albatross offshore of San Quintín (30.5°N), in northern Baja California, and 553 habitat use for the red knot in Guerrero Negro and Ojo de Liebre Lagoons (27.7°N) in central 554 555 Baja California. In general, most studies did not report specific transboundary feeding, spawning, or foraging locations for most species, but rather broad general distributions along 556 557 the California Bight (Supplementary Table 9).

558

559 Transboundary larval dispersal model

560

Results of the larval dispersal models reveal that the California Current transports larvae 561 southward throughout the year from California to Baja California, while a coastal undercurrent 562 also transports larvae northward from Mexico towards California during summer and winter 563 (Supplementary Movie 1). In the Current scenario, we observed transboundary larval 564 connectivity along a stretch of coast covering ~800 km, where larvae from Mexico travel ~400 565 km north, reaching Point Conception (34.5°N) at the northern limit of the California Bight, 566 while larvae from the USA travel ~400 km south to El Rosario (29.8°N) in northern Baja 567 California (Fig. 6a). 568

569

570 The strength of larval connectivity is a function of the geographic distance between sites, which 571 generally decreases as the distance between sites increases. Connectivity patterns are strongly 572 influenced by the life history of species, including planktonic larval duration (PLD) and 573 spawning time. For example, propagules of the giant kelp (with a PLD of a few days to hours)

move almost exclusively between adjacent sites on scales of < 25-50 km throughout the year, 574 while larval dispersal of abalone (PLD = 7 days) occurs between sites 25-100 km apart, year-575 round. For these species, we found high levels of local larval retention (45% and 25% on 576 average, respectively) compared to species with longer PLDs (average < 7%). For all species, 577 connections extend from southern California to central Baja California, but we found some 578 579 isolated sites (with null probabilities of connectivity to other sites) for short PLD, such as in the offshore islands in southern California (henceforth "Channel Islands") and areas in central 580 581 Baja California.

582

In contrast, larvae of the California Sheephead (PLD = 42 days) can travel 75-500 km during 583 the fall, and larvae of sea urchins (PLD = 56 days) can be transported 100-700 km during winter 584 (Fig. 6a). Thus, transboundary connectivity is more important for species with longer PLD. 585 The importance of transboundary connections varied by commercial species and country. For 586 example, for sea urchin, 16% of all the larvae that settled within each country originated from 587 the other country. On the other hand, for California Sheephead, 20% of all larvae in the USA 588 589 came from Mexico and only 3% in Mexico from the USA. Although transboundary connectivity is more important for species with longer PLD in any single generation, long-term 590 591 resilience over multiple generations depends on sites across the border that are tens to hundreds of kilometers away, even for species with short PLD. 592

593

Climate change is predicted to cause a significant decline in the number and strength of 594 connections for all species. In the Future scenario, which considers the effects of climate 595 change, the number of connections present and consequently the average density of larvae 596 dispersing in the network decreased by half (range from -24% on giant kelp to -63% on 597 abalone), the average probability of the connections reduced by an order of magnitude (range 598 599 from -90.7% in California Sheephead to -96.0% in giant kelp), while local retention improved for all species except giant kelp (Supplementary Table 4). We found that binational connections 600 were significantly reduced (61% and 52% lost for California Sheephead and sea urchins, 601 respectively) or completely lost (giant kelp and abalone) either because larvae cannot reach as 602 603 far or because stepping-stone connections disappeared due to the loss of giant kelp with low and intermediate persistence. For sea urchin and California Sheephead, the number of larvae 604 crossing the border dropped to \leq 3%. For species with short PLD, some sites become 605 completely disconnected, forming independent sub-networks, or sites become only loosely 606 607 connected through a few key sites, especially around the Channel Islands and in northern and

central Baja California (Fig. 6b). Importantly, the Future scenario identified some sites (e.g.,
around the northern Channel Islands and Vizcaino Bay in latitudes 28-29°N) that may be
crucial to avoid the collapse of connectivity in the region and should be prioritized for
protection from additional impacts.

612

Time for recovery of focal species and reserve coverage

614

The deterministic recovery model revealed that, overall, increased protection results in faster 615 616 recovery and higher equilibrium biomasses. For all protection levels in the region (i.e., 10, 30, and 100%), sea cucumber showed the fastest recovery from fishing, at less than 10 years (Fig. 617 7a). Abalone and lobster showed the slowest rates of recovery, requiring between 31 and 47 618 years. Importantly, our results show that protecting 10, 20, or even 30% of abalone populations 619 is not enough to allow recovery (Fig. 7b), highlighting that increasing reserve coverage and 620 621 combining with other management actions is needed to facilitate population recoveries for slow-growing populations such as abalone. All fish and sea urchin species showed an 622 623 intermediate recovery, requiring 15-20 years, regardless of reserve coverage. All populations reach recovery status fastest under a 100% protection, followed by 30% and then 10% (Fig. 624 625 7b).

626

627 Considering impacts of marine heatwaves on focal species recovery

628

We found an average cumulative marine heatwave intensity for 2014 and 2015 of 465.6 °C days and 684.5 °C days, respectively, for the nine pixels with monitoring data. When modelling future marine heatwaves, we found that by 2100 the probability of any given year experiencing a marine heatwave of this magnitude will be 0.46, 0.88, and 0.99 for scenarios SSP1-2.6, SSP2-4.5 and SSP5-8.5, respectively (Fig. 8). Importantly, if greenhouse gas emissions are not mitigated, the California Bight could be in a permanently extreme marine heatwave within the next 50 years (Fig. 8).

636

In linear models of yearly density change, we found significant differences between years with (2014-2015) and without (1999-2013) marine heatwaves for abalone, sea urchin, and sea cucumber (Supplementary Table 10; p < 0.01). Simulated yearly density of abalone, sea urchin, and sea cucumber species decreased during marine heatwaves (2014-2015) by 59.1%, 67.3%, and 72.4%, respectively.

642

When accounting for potential impacts due to marine heatwaves for vulnerable species, we found that no species reaches recovery status under any combination of emission scenario and reserve coverage, with abalone being particularly vulnerable (Fig. 7a). While not reaching equilibrium, sea cucumbers show the largest population sizes across climate and protection scenarios, but there was great uncertainty about those changes. Our simulations suggest that even rapidly growing species such as sea cucumber may not reach equilibrium biomass within the next 50 years even under 100% protection (Fig. 7a).

650

651 Mapping threats

652

We found highly threatened areas that require management to support resilience, mainly near 653 cities in southern California and northern Baja California (e.g., Los Angeles, San Diego, 654 655 Tijuana, and Ensenada; Fig. 9a). However, in central Baja California and Guadalupe Island, we found highly threatened areas also in remote offshore sites. Both regions are isolated and 656 657 scarcely populated, with Guadalupe Island located ~250 km from the mainland of Baja California Peninsula. We also found an overlap of highly threatened areas with areas of high 658 659 fishing pressure in northern Baja California but also in less populated areas further south at San Quintín (mainly recreational fishing) and El Rosario (primarily commercial fishing) (Fig. 9b-660 c). On the other hand, fishing pressure in central Baja California was lower and mainly 661 concentrated near Punta Eugenia and Bahía Tortugas for commercial fishing and south of 662 Cedros Island for recreational fishing. Sea urchin and sea cucumber are the most important 663 commercial fishing catch in northern Baja California, lobster and abalone in central Baja 664 California, and abalone in Isla Guadalupe (Supplementary Table 7). 665

666

667 Discussion

668

669 Our results show that although southern California occupies ~25% of the California Bight 670 extension, spanning the USA-Mexico border, it contains almost half of the marine habitats and 671 supports strong ecological connections with northern Baja California. This transboundary 672 connectivity is already significant for populations with long planktonic larval duration (PLD) 673 and highly mobile adults and juveniles, and is likely to become more important as species shift 674 their distributions in response to changing environmental conditions. Given that many 675 populations of ecologically and commercially important species depend on both countries, it is

essential to protect important transboundary areas for migratory species and larval dispersal to
 recover and maintain populations, communities, and genetic flows^{10,69} and benefit both
 conservation and fisheries^{47,50,70,71}.

679

Under a future climate scenario, we found that many areas in the California Bight become 680 isolated and binational connections will diminish or be lost, especially for species with short 681 PLD. However, local retention improves for most species, suggesting that establishing large 682 marine reserves in areas that will become more isolated is critical to maintaining self-683 replenishment and supporting local populations 61,72 . Like other studies 21,27 , we found that the 684 strength of connections weakens, the overall larval recruitment decreases, and that some nodes 685 for species with short PLD may become disconnected. Under this future climate scenario, 686 687 networks of marine reserves will need to prioritize the protection of key stepping-stone nodes to avoid the collapse and fragmentation of larval dispersal in the region. If not adequately 688 protected from fishing, depleted populations' limited supply of larvae could lead to genetic 689 bottlenecks⁷³ and local population collapse⁷⁴, with economic implications. 690

691

Our results indicate that SST variability is a good proxy for climate refugia, particularly for 692 central Baja California. Areas with high-frequency daily variability in ocean conditions (24 693 hours or less) can provide refuge^{30,75,76}, similarly to deeper nearby habitats where vulnerable 694 sessile species can survive adverse conditions and mobile species can retreat^{77,78}. The observed 695 lack of relationship between SST variability and giant kelp persistence in southern California 696 and northern Baja California may imply that giant kelp there are less limited by nutrient 697 availability and temperature extremes than giant kelp near their southern distribution limit in 698 central Baja California⁷⁹. Therefore, it is a research priority to assess whether highly persistent 699 700 giant kelp provide refuge for vulnerable species to climate impacts.

701

The probability that the California Bight will be subject to similar marine heatwaves as extreme as those in 2014-2015 in the following decades is high, and it becomes more likely every year while carbon emissions continue to rise, consistent with global analysis⁸⁰. Even if carbon emissions can be reduced in the next decades, the California Bight will most likely face new extreme events. Along with marine heatwaves, resource extractions are the main threats in the region, which can have interactive and cumulative effects and degrade the resilience of marine ecosystems⁸¹. Notably, when we include the potential impacts of future marine heatwaves on the recovery of vulnerable species (sessile or limited mobility), our modelling results suggest that even with high levels of protection, these species of commercial and ecological importance will not fully recover in subsequent decades. Results from the expert-opinion survey and literature review corroborate the notion that species with limited movement and with slow recovery (e.g., abalone) are more vulnerable than mobile, fast-recovering species. Protecting climate refugia from extractive activities might be the best available climate-adaptation strategy¹¹ to buffer the impacts of future marine heatwaves on vulnerable species.

716

Coordinated conservation efforts for entire ecoregions will support climate-smart designs for 717 biodiversity conservation and fisheries management to a greater degree than networks 718 developed separately by each nation^{50,71,82,83}. We linked climate-smart strategies^{11,17} (e.g., 719 assessing climate impacts and vulnerabilities, representing climate refugia, incorporating 720 connectivity, forecasting effectiveness of protection) using kelp forest ecosystems and 721 722 associated focal species considering future threats from climate change. Notably, the climatesmart focus of our work addresses the growing need to meet post-2020 conservation targets 723 and protect 30% of the oceans by 2030 while adapting to climate change^{23,24}. 724

725

We developed biophysical guidelines to ensure that a transboundary network of marine 726 reserves in the California Bight would complement the existing network in California⁶³, by 727 adapting and refining existing best practice guidelines from California, Mexico, and other 728 regions^{9,10,40}. Setting representation targets at 30% of the distribution of each habitat in each 729 730 biogeographic sub-region will protect ecological communities and processes in the California Bight^{9,10,22} and support adequate recovery for a range of focal species⁸⁴. However, we found 731 certain habitats (e.g., seagrass, giant kelp forests) to be more vulnerable to threats and may 732 require higher levels of protection⁴⁹. Habitat representation targets can be adjusted if robust 733 fisheries measures are in place^{9,10}, but will need to consider the between-country differences 734 in catch and management policies^{53,85}. 735

736

Many focal migratory species in the California Bight can move for hundreds to thousands of
kilometres⁸⁵. Protecting these species requires a combination of marine reserves and other
management tools, such as gear restrictions, catch quotas, seasonal bans, or species moratoria,
among other measures^{9,70}. Although marine reserves cannot fully protect species with longdistance (>10s km) movements that migrate outside their borders, they can protect areas critical

for their life history (e.g., nursery, foraging, spawning)^{9,10,70}. For these actions to effectively be in place, protecting critical areas across the border will require increased investment in international cooperation and governance⁸⁶. Moreover, we found little information about critical transboundary sites, highlighting the need to develop research programs and collaborations to generate this information. Notably, setting adequate protection and management strategies will also require a better understanding of the populations of focal migratory species within and across borders in the California Bight⁸⁵.

749

750 Our findings that transboundary connectivity is more important for species with longer PLD agree with empirical data on the genetic structure of these species. For example, studies found 751 strong genetic differentiation for neutral genetic markers for species with short dispersal 752 distances, such as giant kelp⁸⁷ and pink abalone⁷³, between southern California and populations 753 from Baja California. In contrast, studies found lack of genetic structure for California 754 Sheephead between California and Baja California⁸⁸ and purple and red sea urchin from 755 Washington to southern California^{89,90}. To meet international commitments²⁴, fully protected 756 areas will need expansion in California (currently ~10%) and in Baja California (currently < 757 1%). This provides an opportunity for both regions to coordinate efforts and maximize 758 759 conservation and fisheries benefits based on biophysical principles and modelling of larval dispersal. 760

761

We recommend establishing marine reserves in areas where threats can be managed effectively 762 within reserves in the California Bight. Areas with high cumulative impacts (e.g., coastal 763 development, pollution, runoffs) are likely degrading ecosystem health, fisheries productivity, 764 and resilience to climate change (reviewed by Green, et al. ⁹), preventing marine reserves from 765 producing the expected benefits^{9,10}. However, these are general recommendations since 766 reducing overfishing inside marine reserves, combined with restoration actions and other 767 management strategies that directly address those threats, can build resilience^{3,5,7} to threats not 768 directly managed by marine reserves and contribute to the recovery of degraded areas. 769 Therefore, the decision to protect or restore highly threatened areas requires cost-benefit 770 analysis on a site-specific basis⁹¹ and other considerations such as ecological connectivity. 771

772

Prior to our effort, there were no comprehensive high-resolution maps of small-scale fisheries
catch in the Peninsula of Baja California, limiting our capacity to conduct cost-effective marine
spatial planning. Although information on recreational fishing existed⁵³, we updated and

extended the mapping to cover the three sub-regions in Baja California. These maps, combined with existing data from socioeconomic activities from California, provides the unique opportunity to guide marine spatial planning and identify priority areas for conservation that meet our biophysical design principles while minimizing the potential conflict from implementation to stakeholders' activities⁴⁸.

781

We note that our findings are subject to some caveats. We simulated passive larval dispersal 782 and did not consider other biological traits (e.g., larval mortality, behavior, and settlement 783 competency)⁹². However, we did tailor our model to the availability of giant kelp. Successful 784 recruitment of larvae is linked to the quality and suitability of available habitats^{22,93,94}. 785 Although our climate scenario is a realistic expectation if carbon emissions are not mitigated 786 in the 21st century⁶¹ (decrease in PLD following a 3°C warming tailored to the availability of 787 only highly persistent giant kelp habitat), it is an extreme scenario. Thus, if our model was to 788 be used for identifying climate-smart networks of marine reserves, we recommend adjusting 789 790 the model to consider multiple emission scenarios (e.g., SSP1-2.6, SSP2-4.5, SSP5-8.5) based on projected multi-model ensemble means for the California Bight. 791

792

793 While it is currently unfeasible to obtain information for all habitats and species in a region, we chose giant kelp forests because of existing information on their persistence²⁰ and the 794 availability of long-term information for associated focal species³⁷. Also, kelp forests are 795 threatened by marine heatwaves globally^{35,95}, acting as early indicators of climate-change 796 797 impacts to other ecosystems. However, as more information becomes available, similar assessments for other vulnerable ecosystems, such estuaries, and rocky intertidal habitats 798 799 should be conducted. Although our analysis found no positive correlation between SST variability and persistence of giant kelp forests for southern California and northern Baja 800 801 California, SST variability may still be a good proxy for climate refugia for these regions, but anthropogenic activities may be eroding the persistence of giant kelp forests and masking the 802 potential relationship. For example, many of the kelp forests in mainland southern California 803 and northern Baja California experience high human impacts (e.g., run-offs, overfishing), while 804 805 central Baja California is sparsely populated and less impacted. Also, our analysis compared SST variability with giant kelp persistence, and we did not include other species vulnerable to 806 climate change such as sessile invertebrates. Therefore, it is a research priority to understand 807 808 the drivers and synergies controlling species and habitat resilience to extreme events.

810 We also acknowledge caveats in our recovery model which did not include the potential benefit of larval dispersal from climate refugia, where populations may be less impacted by marine 811 heatwaves. Also, we simulated the loss of biomass based on empirical data of the impacts of 812 the 2014-2015 marine heatwaves from giant kelp forest monitoring surveys³⁷ at shallow depths 813 (typically less than 15 m). Thus, our results need to be taken with due care, as some deeper 814 populations of invertebrate and sessile (or limited mobility) species might be less impacted^{77,78}, 815 survive, and support the replenishment of nearby affected areas. Finally, we used catch-only 816 methods to estimate population parameters and fishing mortality. Recent work has shown that 817 these methods may produce biased estimates⁹⁶, however our parameter estimates are similar to 818 those reported in other studies⁹⁷. Regardless of the potential limitations, our results are 819 consistent with other work that suggest the need to increase the coverage of marine reserves to 820 rebuild marine life, including protection of climate-refugia^{5,98-100}. Notably, other management 821 actions such as fishing moratoria, catch quotas, and repopulation of vulnerable species will be 822 needed to complement marine reserves. 823

824

Fully protecting 30%²⁴ of the California Bight by 2030 will require national and transboundary 825 policies and political will. Importantly, there is a need to integrate biophysical guidelines with 826 827 socio-economic and governance principles to produce effective, equitable and robust policies and practices^{42,101} while considering cultural and management differences across the border. 828 829 Unfortunately, despite the scientific capacity and established collaboration among institutions and research groups between the USA and Mexico³¹, existing political cooperation matches 830 neither the level of ecological connectivity observed⁸⁵, nor the needs identified under current 831 and projected climate impacts. Urgent, coordinated binational action needs to be taken to 832 preserve fisheries and conserve biodiversity in the region. It is a grand challenge, given the 833 strong asymmetries in economic wellbeing, governance, implementation capacity, resources, 834 835 and language, among other barriers.

836

Marine reserves in Baja California will require co-management that includes local fishing cooperatives, complemented with other effective management strategies⁴². Some wellmanaged fishing concessions may need less protection, and in some cases, coordination with improved management and restoration actions may achieve biodiversity, fisheries, and climateadaptation objectives. On the other hand, California has the legislative infrastructure for expanding its existing network of marine reserves and the experience in marine spatial planning to create synergies across the border. This collaborative and socio-ecological setting creates a

unique opportunity for the California Bight to implement transboundary and climate-smart 844 marine spatial planning and influence marine conservation worldwide. 845

846

Here, we provide a case study that links biophysical design principles for transboundary 847 climate-smart networks of marine reserves. Our analysis suggests that achieving climate-smart 848 status requires integrating multiple adaptation strategies such as protecting climate refugia and 849 850 considering the implications of climate change for ecological connectivity and protection efficiency. Given that many marine ecoregions worldwide are shared by nations⁴³, our 851 biophysical guidelines and recommendations can inform other regions' aspirations to achieve 852 post-2020^{23,24} protection targets. These regions will need to develop biophysical dispersal 853 models to understand patterns of connectivity, identify potential climate refugia and levels of 854 protection needed to maximize biodiversity, fisheries, and climate adaptation outcomes. 855 Notably, to design climate-smart networks of marine reserves, they will need to coordinate 856 research programs and policies while considering cultural, governance, and management 857 differences across borders. 858

859

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861

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877 **Table 1.** Biophysical guidelines and transboundary and climate-smart considerations for the

design of networks of marine reserves in the Southern California Bight, compiled, adapted, and

refined from California, Mexico, and other regions 9,10,40 .

880

Biophysical guidelines	Transboundary and climate-smart considerations for application
Habitat representation and replication	
1. Represent at least 30% of each habitat type in each biogeographic sub-region. Habitat representation targets should consider habitat rarity and vulnerability, and fishing pressure and management outside reserves	Habitat types include intertidal, subtidal, biogenic (e.g., kelp forests, seagrass beds), and deep-sea habitats.
	The four biogeographic sub-regions include southern California, northern Baja California, central Baja California, and Guadalupe Island ⁴⁶ . Ensure representing the variation in biodiversity across geographic gradients.
	Consider higher levels of protection for vulnerable habitats (e.g., rocky intertidal, estuaries), rare habitats (e.g., eelgrass beds, island habitats), and overfished or poorly managed habitats ^{9,10} for each sub-region.
	Consider differences in fisheries management across borders ⁸⁵ .
2. Represent at least three examples of each habitat type in widely separated reserves to reduce the chance that they will all be impacted by a large-scale disturbance	Replicate habitats should meet a minimum size required to encompass 90% of the biodiversity associated with each habitat ⁴⁰ (Supplementary Table 6).
3. Represent and replicate climate-refugia habitats in each biogeographic sub-region	Adjust and increase representation targets for each biogeographic sub-region to protect highly persistent giant kelp ²⁰ . Represent deeper habitats used by vulnerable species (i.e., sessile species) or areas with high temperature variability as these populations might support recovery of nearby populations after an extreme event ^{30,77,78} .
4. Represent habitats used by focal species for ecosystem resilience	Represent habitat attributes (e.g., steepness, rugosity) known to favour the biomass recovery of species ¹⁰² that enhance the resilience of ecosystems to adapt to climate change ¹⁰³ . Examples include predatory species that stabilize sea urchin populations, allowing giant kelp to persist ^{7,103} .
Protect critical and unique areas	
5. Protect critical areas in the life history of focal species in marine reserves	Critical areas include spawning, nesting, or breeding areas, nursery habitats (e.g., estuaries and seagrass beds), resting and feeding areas ^{9,10} . If necessary, combine protection with other measures such as temporal fisheries closures during spawning season or regulations to protect migratory species like cetaceans and large sharks.
	Coordinate binational protection and other measures when focal migratory species use transboundary critical and unique areas, particularly near the USA/Mexico border where there is higher transboundary movement.
6. Protect areas with special and unique biodiversity in marine reserves	Protect special and unique features including areas with remaining populations of rare species, protected species, unique habitats, healthy habitats, high species richness, and endemic species ^{9,10} .

Incorporate connectivity

7. Consider movement patterns of adult and juvenile organisms when determining the size of marine reserves. They should be more than twice the size of the home range of species and consider other management tools for species with long-distance movements (>10 km).

8. Consider larval dispersal to replenish populations within marine reserves and in adjacent areas, enhance metapopulation persistence, and support fisheries in adjacent areas. Base the size for protection on the habitats that species use, rather than the overall size of the marine reserve^{9,10,70}. Ensure marine reserves extend from intertidal (minimum linear extension of 5 km) to deeper habitats (will vary depending on the slope)⁴⁰. Species movement can vary from short distances of 100 m for abalone to long distances of 100 to 1000 km for sharks and tuna species^{9,10,40}.

There is strong transboundary connectivity in the region with larval dispersal patterns driven by seasonal north-south currents, which reverse twice a year. Marine reserves should be separated by no more than 25-100 km to ensure connectivity of species with short dispersal distances (e.g., abalone).

Consider large marine reserves for isolated reas to support larval self-replenishment	Species with short dispersal distances require binational coordination near the USA/Mexico border, whereas species with long dispersal distances (e.g., lobster), require coordination for the entire region.
 Consider changes in larval duration and labitat availability due to changes in climate and ocean chemistry. Facilitate range shifts of species driven by 	Consider larger self-sustained marine reserves ⁷² when an area is isolated. Isolation can a function of distance to nearby suitable habitat or short planktonic larval duration for certain focal species, like abalone and giant kelp. There are isolated areas in the entire region, especially in central Baja California from 28-29°N.
	Simulations suggest that a decrease in planktonic larval duration and giant kelp availability due to climate change will weaken the number and geographic scale connections, decreasing transboundary connectivity and increasing isolation. The Channel Islands and areas in central Baja California are expected to become more isolated.
limate change	Distribute reserves across geographic, latitudinal and depth gradients to facilitate the latitudinal and depth shifts of species in response to climate change.
llow time for recovery	
2. Establish marine reserves for the long-term >25 years), preferably permanently, to allow opulations of focal species to recover and eplenish adjacent areas and maintain cosystem functioning and resilience	Populations of focal species recover at different rates in marine reserves and adjacent areas in the California Bight ⁹⁶ . Populations of some focal species, such as sea-cucumbe are likely to recover within a decade, while others, like abalone, will take over 40 years of protection. When possible, establish permanent marine reserves to allow full population recovery and maximum biomass export to adjacent areas.
	Establish permanent marine reserves near the USA/Mexico border where there is strong transboundary connectivity to maximize national conservation efforts.
3. Short-term or seasonal closures should be sed in addition to, rather than instead of, ermanent marine reserves.	Short term (<5-10 years) closures have limited benefits for enhancing fisheries, conserving biodiversity, or building ecosystem resilience. The exception is seasonal closures that can be used to protect critical areas (e.g., spawning or nursery areas), whic can be very important to protect or restore populations of focal fisheries species.
4. Establish permanent and increase the overage of marine reserves when the objective s to protect focal species that are sensitive to narine heatwaves	Some species that are vulnerable to marine heatwaves (sessile or low-mobility species) will likely not reach exploited equilibrium in the next 50 years (at least for shallow populations), which could be achieved only by permanent reserves and increase the are of protection. This is particularly important in Baja California, where average temperatures are higher.
Inimize and avoid local threats	
5. Establish marine reserves in areas with ower levels of cumulative threats for each iogeographic region	Consider the cumulative effects of multiple threats in each location and distinguish among threats that marine reserves cannot directly abate ⁵¹ — but can build resilience to — and threats marine reserves can directly abate.
6. Preferably, establish marine reserves in	Threats not directly abated by marine reserves include climate and land-based stressors While in southern California and northern Baja California areas with higher threats are mostly located near urban areas, in central Baja California and Guadalupe Island they a mostly found off the coast.
	Threats directly abated by marine reserves are those related to extractive uses, such as

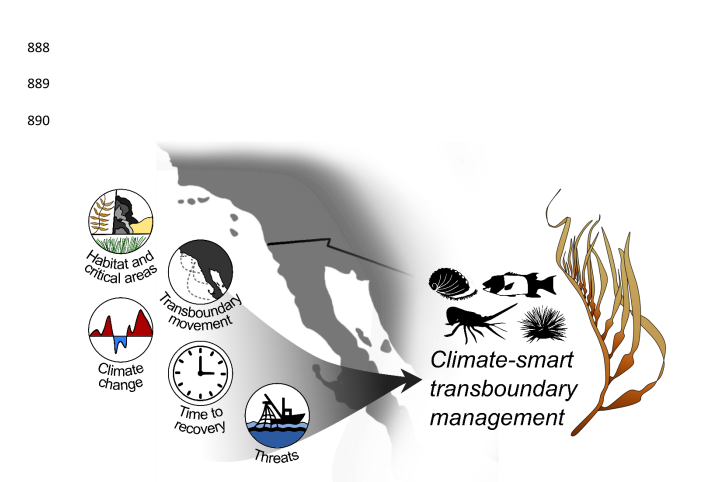


Fig. 1 Graphical framework. Represents the biophysical principles used to develop spatial
maps and analyses based on giant kelp forest ecosystems and associated species for
transboundary networks of climate-smart marine reserves in the California Bight.

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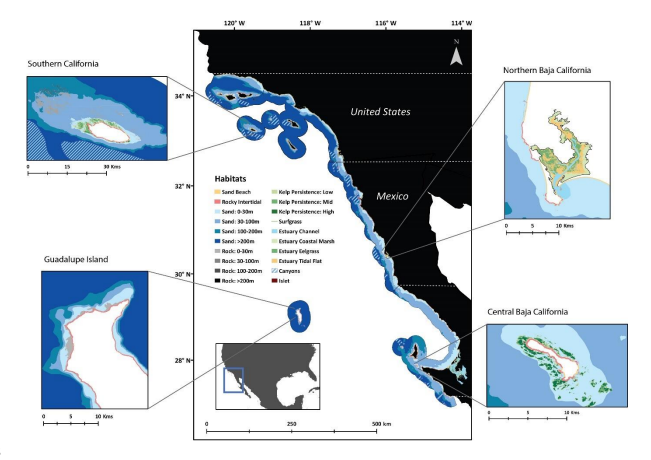


Fig. 2 Distribution of marine habitats for the Southern California Bight. Inserts represent examples of intertidal, subtidal, estuary, and deep-sea habitats for each sub region. Southern California for San Nicolas Island with giant kelp forests, subtidal shallow and deeper habitats, northern Baja California for San Quintin Estuary, central Baja California for Isla Natividad intertidal and shallow habitats, and for Guadalupe Island intertidal and subtidal habitats at different depths. Dashed white lines represent the limits of each sub-region.

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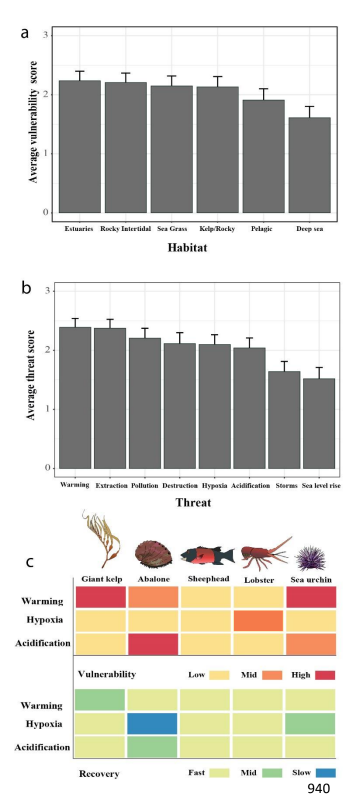


Fig 3. Vulnerability of habitats and focal species to human threats. Average score (± 95%
confidence interval) for a vulnerability of marine habitats to human threats and b threat of
individual stressors across marine ecosystems based on expert opinion surveys. c Ranking of
focal species vulnerability to climate threats and their recovery rates based on scientific
literature. Images credit: Katherine E. Dale.



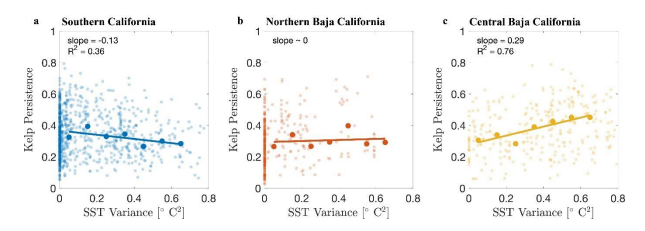




Fig 4. Sea surface temperature variability as a proxy for climate-change refugia.
Relationship between giant kelp persistence and SST variability for a southern California, b
northern Baja California, and c central Baja California. Raw data (light circles), binnedaveraged values (large circles), and best fit to bin-averaged data (line) are shown.

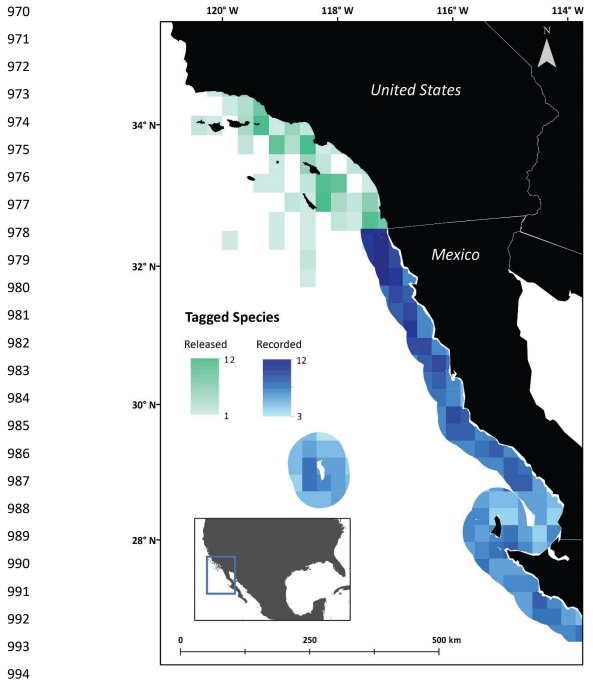
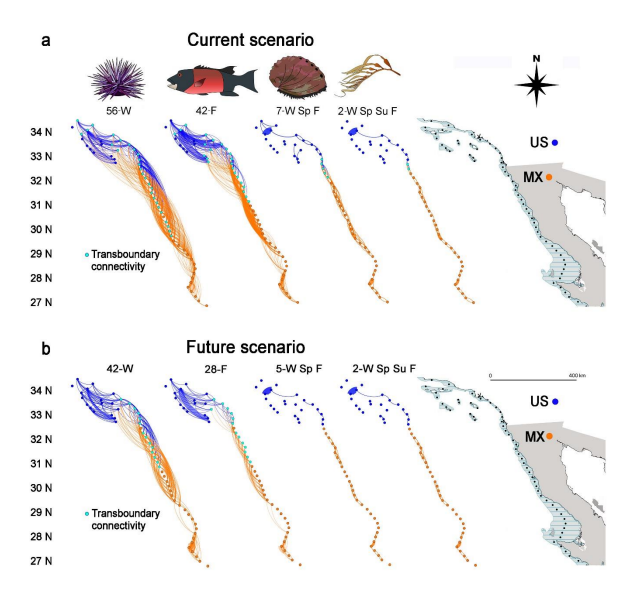


Fig 5. Transboundary connectivity of species moving from California to the Peninsula of
Baja California. Green colors represent the number of studies that released species in southern
California, and blue colors the number of studies that record species in Baja California. We
consider blue quadrants with higher number of studies as quadrants with higher transboundary
connectivity.



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1002 Fig 6. Networks of larval connectivity in the California Bight. Spatial networks of modelled larval dispersal for focal species (from left to right, sea urchin, California Sheephead, abalone, 1003 and giant kelp) between nodes delimited by the 200-m isobath. Connectivity polygons are 1004 shown on the far right. Line width represents the probability of larval dispersal (thicker lines 1005 have higher probability) and line colour the country of origin. Blue-coloured nodes and lines 1006 represent the USA and orange Mexico as larval origins, respectively. Sites involved in 1007 transboundary connectivity are highlighted in light blue color. For each focal species, we 1008 indicate the planktonic larval duration (PLD, in days), followed by a letter representing the 1009 1010 spawning season (Spring = Sp, Summer = Su, Fall = F, Winter = W, See Supplementary Table 4). Images credit: Katherine E. Dale. a Current scenario, considering the PLD reported in the 1011 literature and the total area with giant kelp found within each polygon. **b** Future scenario, 1012 accounting for reduction of PLD and giant kelp habitat due to climate change. Sites involved 1013 in transboundary connectivity are highlighted in light blue color. 1014

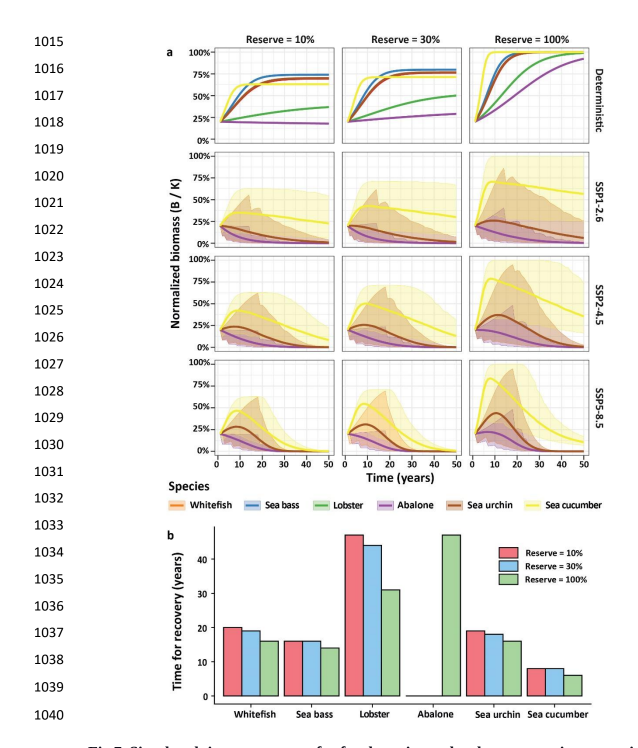


Fig 7. Simulated time to recovery for focal species under three protection scenarios in the 1041 California Bight. Recovery pathway of six species (mean ± standard deviation), a for 1042 scenarios of protection (10, 30, or 100%) and models. The first row of panel shows the 1043 deterministic model for all six species. Rows 2-4 show results from the stochastic models for 1044 vulnerable species (abalone, sea urchin, sea cucumber) to marine heatwaves under three 1045 climate scenarios (SSP1-2.6, SSP2-4.5, SSP5-8.5). b Time to reach 90% of equilibrium 1046 1047 biomass for each of the seven species across three protection scenarios. Missing bars indicate no recovery within the simulated 50 years. 1048

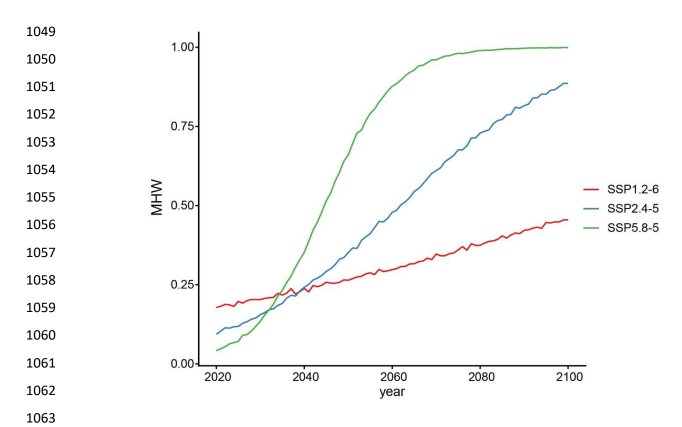


Fig. 8.- Probability of projected marine heatwaves for 2020-2100. Marine heatwaves under climate scenarios SSP1-2.6, 2-4.5, and 5-8.5 that match or exceed the cumulative intensities registered in 2014-2015 in the California Bight.

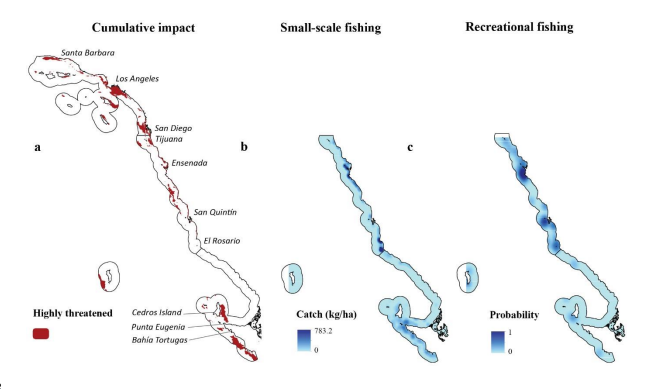


Fig. 9.- Distribution of threats and fishing in the California Bight. Maps of a highly
threatened areas (top 10% of cumulative impacts of 13 threats), b cumulative catch for six
small-scale fisheries and c probability of recreational fishing in a 100-m grid square in the three
regions of Baja California.

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