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Rapid sea level rise causes loss of seagrass meadows

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ABSTRACT

As global declines in seagrass populations continue to cause great concern, long-term assessment of seagrass meadows show promise in providing valuable insights into fundamental causes of seagrass loss and drivers of environmental change. Here we report two long-term records of seagrass presence in western Gulf of Mexico coastal waters that provided insight into their rapid decline in a relatively pristine ecosystem. Coincident with unprecedented increases in water depth starting in 2014 (14-25 mm y⁻¹), monthly measurements at a “deep edge” fixed station revealed that two ubiquitous seagrass species (*Halodule wrightii* and *Syringodium filiforme*) vanished altogether in just five years; a subsequent basin wide assessment revealed that seagrasses disappeared at 23% of 144 sentinel stations. Models that incorporate differing sea level rise scenarios and water depth thresholds reveal additional global losses of seagrass habitat (14,000 km²), with expansion into newly created shallow habitats further exacerbated by hardened shorelines.

Introduction

Seagrasses, a polyphyletic group of angiosperms containing over 60 species, possess specialized adaptations for aquatic life that have facilitated colonization across 160,000 km² of coastal ecosystems^{1,2}. Furthermore, they function as foundation species and provide numerous ecosystem services valued at \$19,000 ha⁻¹, including nursery habitat for fisheries, food for numerous marine organisms, shoreline protection, ecotourism, and educational opportunities^{3,4}. The role of seagrass within earth's global carbon biochemical cycle is becoming increasingly recognized due to their high productivity, sequestration of allochthonous and autochthonous organic “blue” and inorganic carbon in sediments⁵⁻⁷. Seagrasses contain a disproportionately high carbon stock (0.025 Tg km⁻²) compared to high-biomass terrestrial forests (0.014 Tg km⁻²) and are able to potentially store this carbon for thousands of years if undisturbed⁸⁻¹⁰.

Despite the global importance of seagrass meadows, their persistence is significantly threatened due to a variety of factors¹¹. Since the mid-1700s, the global coverage of seagrasses has decreased by about 29% (51,000 km²) with 10 species at risk of extinction^{12,13}. Furthermore, annual losses of seagrass habitat return about 300 Tg y⁻¹ to the active global carbon pool⁸. Major anthropogenic and environmental drivers of seagrass loss include algal blooms¹⁴, chemical pollution¹⁵, disease¹⁶, drought^{17,18}, eutrophication¹⁹, heat waves²⁰, invasive species²¹, mechanical damage²², and storm action¹⁸. Mechanistically, these drivers disrupt the plant's photosynthetic balance between carbon fixation and respiration by 1) limiting the light available for photosynthesis, 2) altering the basal metabolic costs of the plant, and/or 3) damaging the plant's photosynthetic apparatus. Seagrasses have amongst the highest light requirements of aquatic photoautotrophs owing to the high respiratory demand of their root/rhizome system, which must maintain an oxidized rhizosphere in an otherwise anoxic environment^{23,24}.

Seagrasses meadows persist at a variety of depths (0 - 90 m), but species-specific tolerances to light availability control community composition along this steep gradient²³. Plants living near the edge of their physiological light limit maintain a neutral to slightly positive carbon balance and are known as deep edge populations. In aquatic environments, irradiance at depth (I_z) is modelled using the equation $I_z = I_0 e^{-K_d z}$, where I_0 is surface irradiance, z is depth, and K_d is the extinction coefficient. Typical values of K_d in pristine western Gulf of Mexico estuaries (e.g., Laguna Madre²⁵) range between 1 and 2, with a change of 0.1 m sufficient to reduce bottom irradiance by 10 - 20%. Consequently, slight changes in water depth may impact the ability of the populations to persist, which allows them to act as sentinel species for long-term shifts in water depth.

Over the past century global mean sea level has increased by 1 - 3 mm y⁻¹ depending on a suite of complex factors including subsidence, crustal uplift, isostatic rebound, thermal expansion, and glacial melt runoff²⁶. In contrast, natural cycles of Rossby waves have compounded sea level rise in the Gulf of Mexico²⁷, which is rising 2-3 times faster than the global average²⁸ and has increased to > 10 mm y⁻¹ since 2010. Numerous local factors can amplify these effects as exemplified by the weight of water displaced during Hurricane Harvey, which depressed coastal regions of Texas by up to 20 mm for weeks after the storm²⁹. Likewise, water and hydrocarbon extraction are causing land subsidence of up to 9.5 mm y⁻¹ within the Texas Coastal Bend³⁰. As water depths increase, seagrass distribution may shift away from deeper waters and begin to colonize newly

submerged lands. However, an increasing fraction (14% and growing) of US shorelines are hardened³¹ to protect from coastal hazards (e.g., groins, jetties, breakwaters), preventing seagrass proliferation into shallow waters and leading to a net loss of habitat.

Here we examine two long-term records of seagrass presence in the Upper Laguna Madre (TX, USA) to investigate the rapid decline of seagrasses in a relatively pristine ecosystem: a fixed deep edge station (LM-151) and basin wide sampling of 144 sentinel “Tier-2” stations (Figure 1). Our monitoring at deep edge station LM-151 documented the recent disappearance of seagrasses from this area after 30 years of monitoring. By combining data from these two programs, we noted that vegetation shifts at LM-151 (sampled at monthly intervals for three decades) were confirmed on a landscape scale at our 144 sentinel stations sampled annually over the period 2011-2022. We hypothesize that seagrass losses at the deep edge site were caused by long-term increases in water depth and that rises in water level were also responsible for seagrass loss on a landscape scale at deeper edge populations throughout the Upper Laguna Madre.

Results

Environmental Conditions

Water temperature varies seasonally at LM-151 with hot summers (max = 34°C) and cooler winters (min = 7.8°C) with relatively low interannual variability (Figure 2a). In contrast, average daytime underwater irradiance, water depth, and salinity also vary seasonally, but show differing trajectories across years (2a,c). Irradiance was lowest in the early 1990s, increased and plateaued from 2000 - 2015 before beginning to decline again (Figure 2b). Salinity varied among years, but rapidly switched in the early 2010s from a hyposaline (17) to a hypersaline environment (62.6). Water depth increased in 1990s, decreased in the 2000s, and has significantly ($p = 0.02$) increased since 2014 by 14 mm yr⁻¹ (Figure 2c). Across the Texas coast similar patterns were found with low rates of sea level change pre-2014 (up to 9 mm yr⁻¹), followed by a significant increase afterwards (3 - 21 mm yr⁻¹; Figure 2d). Low water depths in the 2010s corresponded with increased Secchi depth, decreased TSS, and decreased pH within the water column (Figure S1). Dissolved oxygen, ammonium and dissolved inorganic nitrogen remained relatively constant until the late 2010s where DO decreased and DIN and NH₄⁺ significantly increased (Figure S1).

Tier-2 sentinel sites formed five distinct clusters based on water depth measurements during annual monitoring from 2011 - 2022 (Figure 3). Cluster 4 had the lowest average water depth (0.45 m) whereas Cluster 5 had the highest (1.66 m). There was relatively low variability in depth between Cluster 1 (0.75 m), Cluster 2 (0.96 m), and Cluster 3 (1.22 m). Overall, the interannual water depth variability at each site was fairly low, but many sites in Cluster 1 were characterized by high variability.

Biological Parameters

From 1989 - 2005 LM-151 was nearly a monotypic *Halodule wrightii* meadow, but in 2005 *Syringodium filiforme* recruited, and by the early 2010s had greater biomass than *Halodule* and equal density (Figure 4a,b,d). *Syringodium* then retreated and was completely absent by 2013, but *Halodule* did not return to its pre-2010 levels. By 2018 *Halodule* also disappeared from the site. Biomass ratios revealed *Syringodium* lost aboveground tissues before belowground, whereas the opposite was true for *Halodule* (Figure 4c).

Seagrass was present during annual Tier-2 monitoring from 2011 - 2022 at most monitoring stations (left panel Figure 3), but varied amongst sites grouped by water depth. Cluster 5 (deepest sites) lost most seagrass. In 2013 seagrass was present at 80% of Cluster 5 sites, dropping to only 4% by 2022 (Figure 3). Overall, Cluster 2 (intermediate depths) had the highest proportion of seagrass presence during sampling (259/261 samplings), whereas Cluster 5 had the lowest (130/261 samplings).

Relationship Between Seagrass and Environment

LM 151

Water temperature and bottom irradiance were significant predictors of both *Syringodium* ($R^2 = 0.68$) and *Halodule* ($R^2 = 0.31$) aboveground biomass (Table S1), but water depth and irradiance best explained the presence of *Syringodium* ($R^2 = 0.22$). *Syringodium* biomass had narrower optimal environmental conditions (25-30°C, $\geq 365 \mu\text{mol photons m}^{-2} \text{s}^{-1}$; Figure S2a) than *Halodule* ($\geq 25^\circ\text{C}$, $\geq 50 \mu\text{mol photons m}^{-2} \text{s}^{-1}$; Figure S2c). However, both *Syringodium* and *Halodule* were negatively influenced by high light under cooler conditions (Figure S2a). Overall, *Syringodium* was less likely to be present at LM-151 under dim, deep conditions ($\leq 200 \mu\text{mol photons m}^{-2} \text{s}^{-1}$ and $\geq 1.5\text{m}$)

Tier-2

Seagrass presence at Tier-2 sampling sites was significantly affected by water depth, year, and the site's geographic coordinates ($R^2 = 0.62$; Table S1). Generally, water depths greater than 1.5 m negatively affected the probability of seagrass presence, but the strength of this effect varied among years (e.g., 2017 - 2018; Figure S2d). Spatially, seagrasses were more likely to be found in northeastern Laguna Madre, and least likely in the southern region (Figure S3).

Future Scenarios

Seagrass meadows have historically covered up to 200 km² within Upper Laguna Madre, but by 2018 suitable seagrass habitat decreased by more than 41 km² with the increased prevalence of water depths ≥ 1.5 m (Figure 5). Under varying sea-level rise scenarios, this unsuitable habitat will expand to 73-113 km², especially within the northern reaches of the estuary. However, newly submerged habitat is predicted to vary between 90 - 135 km², creating a potential net gain of 59 - 65 km² in suitable seagrass habitat.

Discussion

This study synthesizes decades of monitoring efforts (LM-151 & sentinel Tier-2 stations) to assess the long-term response of seagrass populations at local and landscape scales to environmental changes within the Upper Laguna Madre. Over the past 30 years *Syringodium* and *Halodule* populations showed a typical successional pattern at a deep edge monitoring station (LM-151), but overall seagrass cover began to decline significantly after 2014 as sea level continued to rapidly rise at a rate of 14 mm y⁻¹. Statistical modeling revealed that seagrass biomass was sensitive to changes in temperature and bottom irradiance, whereas seagrass presence was controlled by water depth and irradiance. Since 2018 seagrass has vanished from the LM-151 deep edge site and from deep Tier-2 monitoring locations across the Upper Laguna Madre.

Halodule wrightii and *Syringodium filiforme* are both common subtropical Atlantic species but have varying biological characteristics that lead them to represent different successional stages. At LM-151, *Halodule* populations thrived until 2007; at that time *Syringodium* began to invade and shade the shorter *Halodule* canopy. *Halodule* is typically a pioneer species that has faster growth rates, higher nutrient requirements, and is more tolerant of hypersaline conditions than *Syringodium*^{32,33}. The sudden decline of *Syringodium* in 2012 was caused by drought-induced hypersalinity³⁴. We would expect *S. filiforme* biomass to have a stronger correlation with the environment than *H. wrightii* because it has more specific growth requirements. Indeed, we found a greater link with *Syringodium* ($R^2 = 0.68$) and environmental conditions than *Halodule* ($R^2 = 0.31$). However, both species were tied to the same environmental variables: water temperature and irradiance. Seagrasses often show strong annual biomass cycles linked with seasonal conditions (namely light and temperature)^{34,35}. In the absence of no significant long-term and consistent trends in temperature or salinity, we believe that low irradiance coupled with deeper waters is responsible for the failure of *Syringodium* to recolonize as salinities moderated, the slow disappearance of *Halodule* at LM-151, and the overall loss of seagrasses throughout the Upper Laguna Madre.

Partial effect plots generated through general additive modeling (Figure S2) are valuable tools for identifying physiologically relevant drivers of seagrass distribution. For example, we identified optimum *S. filiforme* conditions between 25-30°C and ≥ 365 $\mu\text{mol photons m}^{-2} \text{ s}^{-1}$. Previous research identified that peak productivity occurred between 23 - 29 °C³⁶ and photosynthesis became saturated³⁷ at 370 $\mu\text{mol photons m}^{-2}$. Likewise, *Halodule* biomass increased at $\geq 25^\circ\text{C}$ water temperature across all measured irradiances. These values agree with regional measurements of optimal water temperatures (25 - 30°C)³⁸ and saturation (100 - 600 $\mu\text{mol photons m}^{-2}$ depending on the season)³⁵. Together, these results suggest that the individual physiologies of the plants are more important in shaping the communities than physical disruption from local climatic events (e.g., storms or freezing).

Changes in the below:aboveground biomass ratio yield insight into the drivers of seagrass loss. The ratio rapidly rises as *Syringodium* photosynthetic tissues are lost to the system, indicating an acute stress. Depending on the duration, recovery from belowground tissues (roots/rhizomes) is possible. Earlier work described widescale *Syringodium* loss in Laguna Madre in 2011 and 2012 because of drought-induced hypersalinity³⁴, but *Syringodium* has not returned to LM-151 despite salinity decreases. Conversely, the below:aboveground biomass ratio in *Halodule* is known to decline slightly in response to chronic stressors that reduces the plant photosynthetic output or increases their basal metabolic costs³⁹. Likewise, *H. uninervis* had a significant reduction in biomass, but maintained a constant root:shoot ratio during a year of shading⁴⁰. The long-term decline of seagrasses in Indian River lagoon shows a similar pattern⁴¹. Therefore, an added stressor beyond salinity likely prevented recolonization of *Syringodium* and caused a *Halodule* decline. We found a strong negative relationship between water depth and seagrass biomass in Laguna Madre. Long-term changes in water level would function as a chronic stressor that integrates various environmental factors (e.g., stratification, temperature), but especially light penetration.

Contributions from thermal expansion of rapidly warming waters as well as large removals of groundwater produce a rate of relative sea level rise (rSLR) along Texas' Gulf Coast that is one of the highest in the world^{27,28}. Water depth data collected for over 100 yr in Galveston (TX, USA) revealed a rSLR rate of 6.6 mm y⁻¹, nearly 3x faster than the global average²⁸. Here we report contemporary (post-2014) water-depth increases far exceeding this average at 28 out of 31 NOAA stations (up to 21 mm y⁻¹ in Nueces Bay) and our long-term LM-151 site. In contrast, 14 out of 20 stations had rates lower than 6.6 mm y⁻¹ from 1990 - 2013. These data are consistent with recently described sea level rise acceleration caused by the compounding of wind driven Rossby waves over the Caribbean^{26,27}. This recent acceleration will likely be short-lived²⁷, but these rapid effects provide valuable empirical evidence of long-term effects of sea level rise on seagrasses rather than projected impacts alone^{42,43}.

The 0.25 m increase in water depth observed at LM-151 would decrease available light for photosynthesis by 25-50%, sufficient to cause local extirpation and/or habitat shifts at this deep edge site.

Previous research has noted seagrasses as a potential "winner" compared to other coastal habitats under localized water depth changes⁴⁴. Likewise, under realistic future sea level rise scenarios⁴⁵ we predict that seagrass habitat can expand by up to 65 km² (Figure 5). This value is an upper-bound and actual colonization may be significantly less because of factors like poor environmental conditions and shoreline hardening. Upper Laguna Madre has the largest potential for seagrass expansion into newly submerged lands in the Gulf of Mexico because it is bounded by undeveloped shorelines suitable for seagrass growth (53% of shorelines are suitable; Table S2). But the favorable situation in Laguna Madre is not reflected on most shorelines in the US because they have become increasingly artificially hardened³¹ or are otherwise unsuitable for seagrass colonization. Currently, marshes represent a large source of uncertainty as potential seagrass habitat because they may or may not keep pace with sea level rise by maintaining sufficiently high accretion rates⁴⁶. In Texas, 41-64% (242 - 378 km) of the shoreline that incorporates seven large estuarine systems is unsuitable for seagrass colonization. Across the United States, hardened structures have removed 21,400 km of potential seagrass habitat and this number will continue to grow³¹.

Globally, the trend of shoreline hardening is expected to continue and become exacerbated under future scenarios of stronger storms, higher erosion, and deeper waters³¹. Although seagrasses within Laguna Madre are likely more resilient to continued sea level rise despite localized-rapid losses, but the fate of other meadows within the Western Gulf of Mexico and elsewhere remains unknown. Worldwide, we estimate that over 14,000 km² of seagrass meadows live in waters of lower transparency equivalent to that of the Upper Laguna Madre ($K_d > 1$), suggesting they may also be sensitive to sea level rise (Table 1). These include over 50% of the seagrass meadows in the Netherlands, Poland, Sweden, Denmark, Canada, Senegal, Germany, and Norway. Seagrasses that occur in waters of high water transparency will also be lost because deep edge populations in these systems will switch from sufficient to insufficient light availability. Global seagrass populations are already threatened by the increased frequency of major storm events, eutrophication, fungal infections, algal blooms, and temperature extremes^{11,12}. Accelerated rises in sea level are yet another perturbation that displaces functional seagrass communities with unvegetated sediments that further compromise the productivity and integrity of coastal ecosystems worldwide through loss of both habitat and carbon sequestration potential is lost.

Methods

Site and Monitoring Description

The western Gulf of Mexico contains numerous estuarine systems including Laguna Madre, a unique hypersaline lagoon, which contains over 70% of the Texas's 900 km² of seagrass meadows⁴⁷. Broadly, the system is divided into two regions (Upper and Lower) by a large tidal flat. Here, we are conducting seagrass monitoring efforts within the Upper Laguna Madre (ULM) that is bordered by Corpus Christi (TX) on the western shore and Padre Island National Seashore on the eastern (Figure 1). Three seagrasses are found within ULM: (*Halodule wrightii*, *Halophila engelmannii*, and *Syringodium filiforme*). ULM is bisected by the Gulf Intercoastal Waterway system, but either side is dominated by shallow (< 2m) seagrass meadows. A long-term seagrass monitoring station "LM-151" (27.35 N, 97.37 W) was established in 1989 within Padre Island National Seashore to track changes in a deep edge population of *Halodule*⁴⁸. Annual Tier-2 sampling of ULM began in 2011 across 144 stations in the ULM to assess broad-scale changes in seagrass populations. Details of the LM-151 sampling are provided below, whereas Tier-2 methods have been described previously³⁴. Data for the Tier-2 and LM-151 monitoring programs are available online⁴⁹.

Water Quality Measurements

Near-continuous underwater measurements of canopy-level photosynthetically active radiation (PAR) began at LM-151 in 1989 using a LI-193SA spherical (4π) quantum sensor connected to a LI-1000 datalogger (LI-COR Inc., Lincoln, Nebraska, USA)⁴⁸. Measurements were taken every minute and integrated over the hour before logging. Average daytime irradiance (ADI) was calculated by taking the monthly mean of daily averaged irradiance values between sunrise and sunset. This metric was chosen for analysis to minimize daily light variability and reflect growing conditions. A thin, optically-clear plastic covering was placed over the sensor and replaced ever 2 - 4 wk to reduce biofouling. During these maintenance trips, water temperature, conductivity (converted to salinity), pH, and dissolved oxygen were typically measured using a YSI-6920 datasonde (YSI, Yellow Springs, OH). Total suspended solids in the water column were quantified by measuring the dry filtrate from a 1L seawater sample retained on a pre-combusted 0.7 μ m glass fiber filter. Water depth was measured using a marked PVC pole. Additionally, secchi depth was measured by lowering a black and white secchi disk until no longer visible or the bottom was reached. Multiple measurements of environmental parameters taken within a month were averaged for analysis. Concentrations of ammonium and dissolved inorganic nitrogen were measured from replicate water samples following standard protocols⁵⁰.

Biological Sampling

At LM-151, four biomass samples were taken haphazardly every 3 - 6 months. At each position, a PVC corer was pressed 30 cm into the sediment to collect aboveground and belowground tissues. The number of shoots in each sample were counted for each species before tissues were segmented into photosynthetic (aboveground) and non-photosynthetic (belowground) portions in the lab, dried, and massed. All values were averaged, extrapolated, and standardized to m^{-2} coverage.

Data Analysis

Data analysis and visualization primarily used R (v 4.2.1), but ArcGIS Pro was used for mapping, inverse distance weighted interpolation, and areal calculations of spatial data (v 3.0.2). Scripts and data for replication of data analysis are available online (DOI PENDING). Locally estimated scatterplot smoothing (LOESS) line was added to plots of environmental variables to aid in visualization of long-term trends. The relationship between water depth and the presence of seagrass at Tier-2 sampling locations was explored through hierarchical clustering of yearly water depth measurements. Data were scaled such that $\bar{x} = 0, \sigma = 1$ before a pairwise euclidean distance matrix was generated for clustering through ward's method. Seagrass presence data were sorted according to match this clustering scheme.

Statistical Modeling

General additive models (GAMs) were constructed using the mgcv package to evaluate the statistical relationship between biological and environmental parameters because of the non-linear nature of the parameters. The model basis functions were adjusted from their default values until smoothing was adequate, p-values were non-significant, and effective degrees of freedom did not significantly change. Various model families were tested, but assessment of model diagnostics led us to use the gamma distribution to assess environmental effects on seagrass biomass and binomial with a logit link function for seagrass presence. Aboveground biomass was chosen as the response variable because of greater data coverage than shoot density and strong correlation to belowground biomass. Water depth was the only explanatory variable tested for Tier-2 data because it is less sensitive to variations in weather or time of day and directly links to our *a priori* hypotheses. Biomass models were limited by the number of explanatory variables that could be added, so non-significant factors or those with low power were removed.

Coastwide Sea level Rise Assessment

To determine statewide rates of relative sea level rise, we acquired data from 54 stations across the coast referenced to NAVD88⁵¹. We queried data from 1/1/1990 - 12/31/2022, but data coverage varied by station. Data was split into pre- and post-2014 sections for analysis because to understand how rates may have changed following the loss of *Syringodium* in 2014. For each section, we used linear regression of daily maximum tidal heights over time to calculate relative sea level rise if at least 5 years of data existed. Non-significant slopes ($p > 0.05$) were assumed to be 0. For comparison, monthly averaged water depth measurements from LM-151 were analyzed in the same fashion.

Seagrass Habitat Change

To evaluate seagrass habitat change under future sea level rise we considered three scenarios of rSLR by 2050 for the Western Gulf of Mexico⁴⁵: best-case (+0.29m), intermediate/current-case (+0.37m), and worst-case (+0.49m). Under each scenario, we calculated the areal extent of land that would be submerged by each water depth increase and seafloor that would become unsuitable for seagrass growth. To estimate newly submerged habitat, 1-m resolution digital elevation models (DEM) were used to construct high-resolution shoreline topographic maps⁵¹. Next, the current shoreline position was estimated by calculating the mean sea level at Bird Island Basin tidal gauge (#8776139) during DEM acquisition⁵². Increases in water depth were assessed using the 2018 water depth data generated by Tier-2 sampling to be specific for seagrass meadows, have moderate spatial resolution, and align with the DEM acquisition year. We chose a threshold of 1.5 m to represent unsuitable seagrass habitat, because meadows at this depth within Laguna Madre have less biomass and reduced presence probability (see results).

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Author contributions statement

K.D. conceived the monitoring programs and acquired funding, K.D. and K.C. conducted the monitoring, K.C. analysed the results. Both authors wrote and reviewed the manuscript.

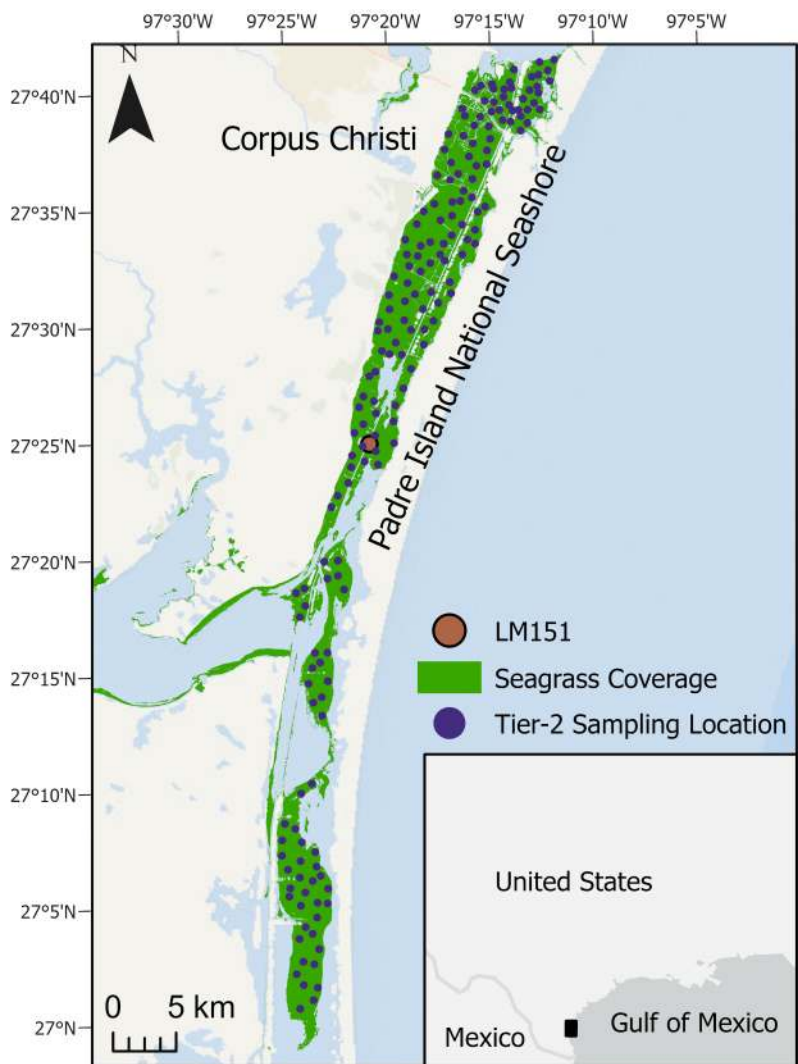


Figure 1. Map of seagrass monitoring locations within the Upper Laguna Madre (TX, USA). Seagrass coverage represents historically known meadows within the system⁴⁷.

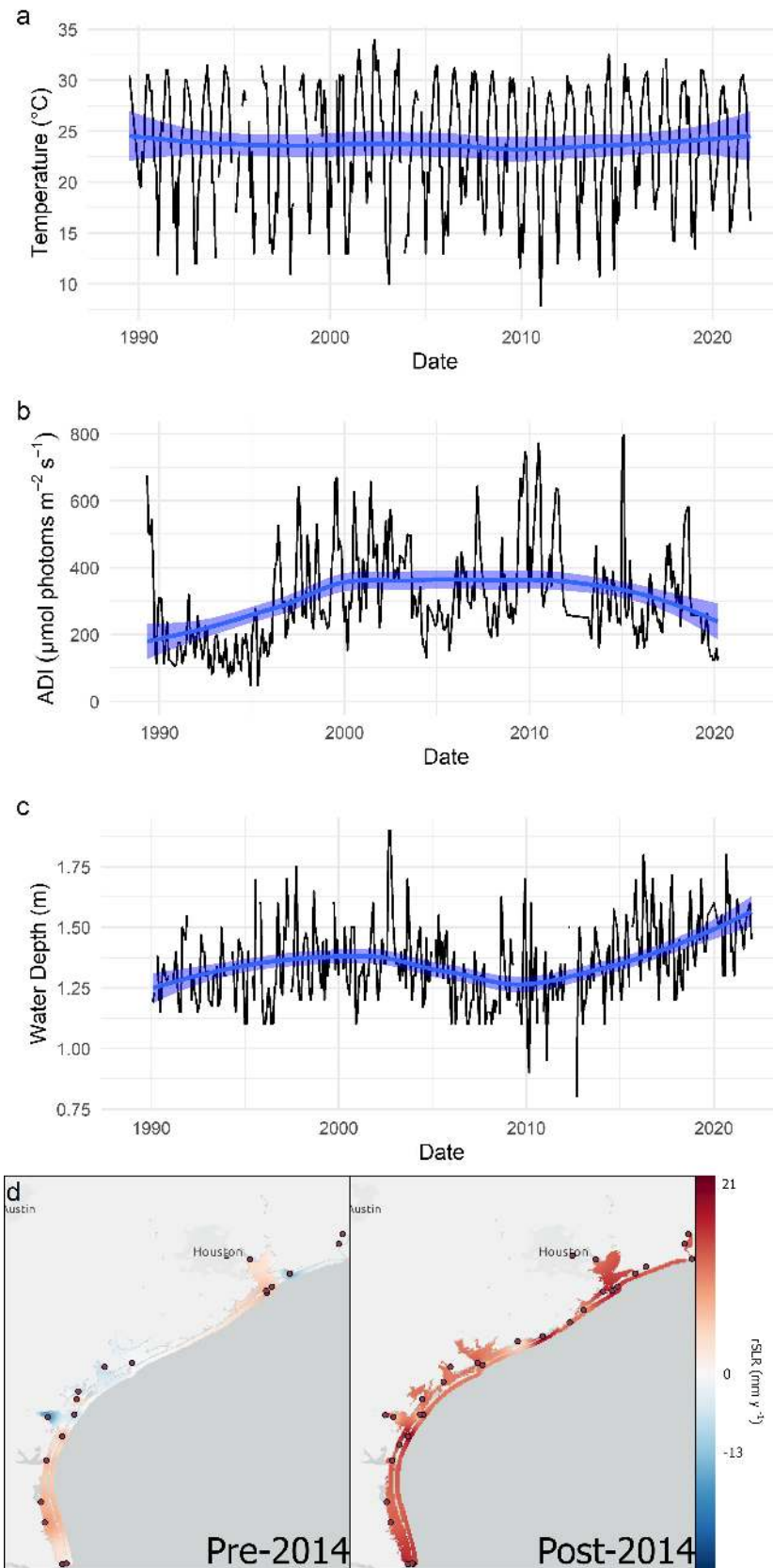


Figure 2. a-c) Environmental variables from LM-151. d) Calculated rates of relative sea level rise from NOAA buoys before or after 2014. ADI = Average daytime irradiance.

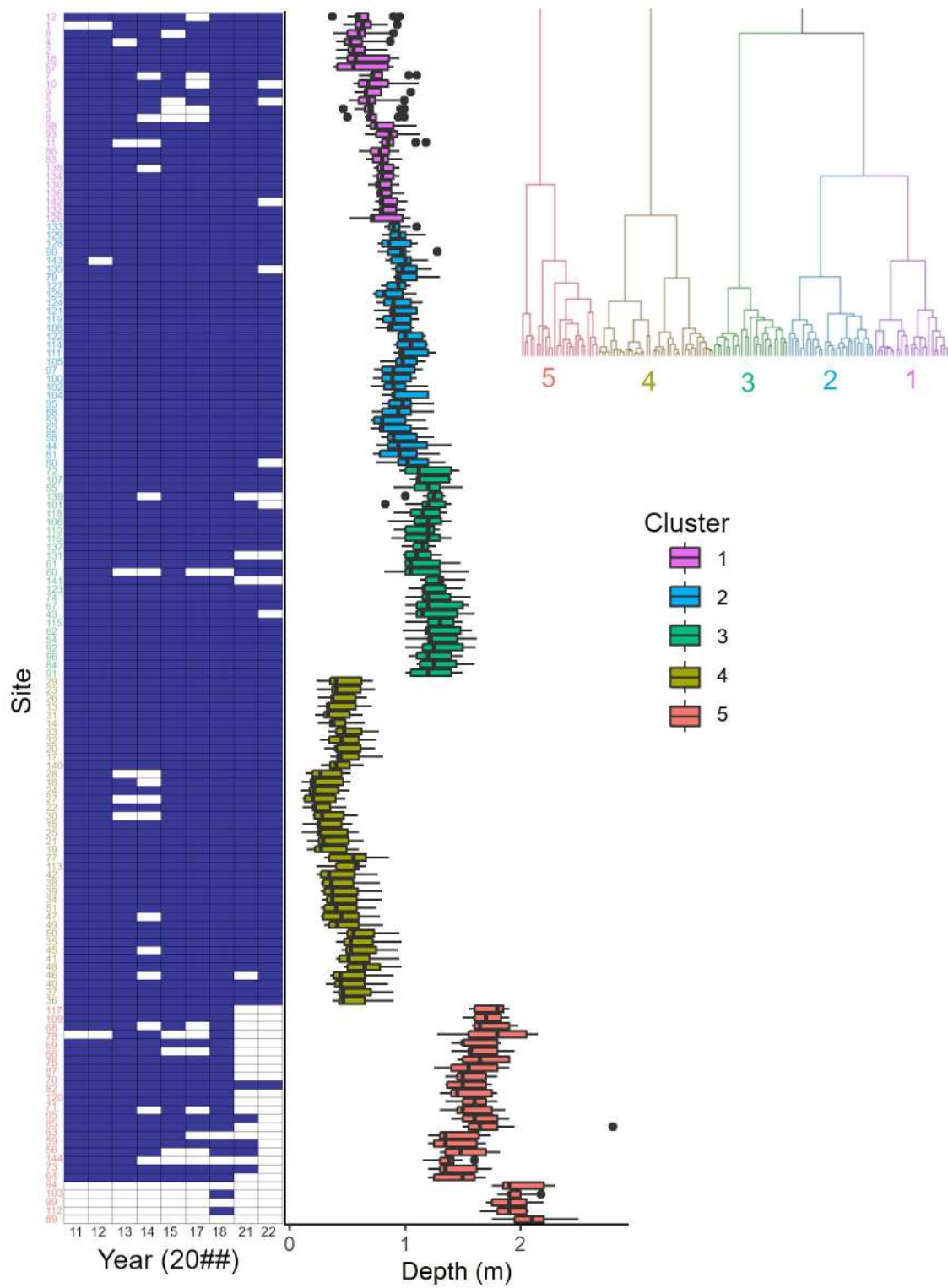


Figure 3. The presence (blue) or absence (white) of seagrass at each Tier-2 monitoring station over time. Sites are aligned based off hierarchical cluster of water depth measurements. The inset shows the overall clustering scheme with clusters 4 + 5 different from 1-3.

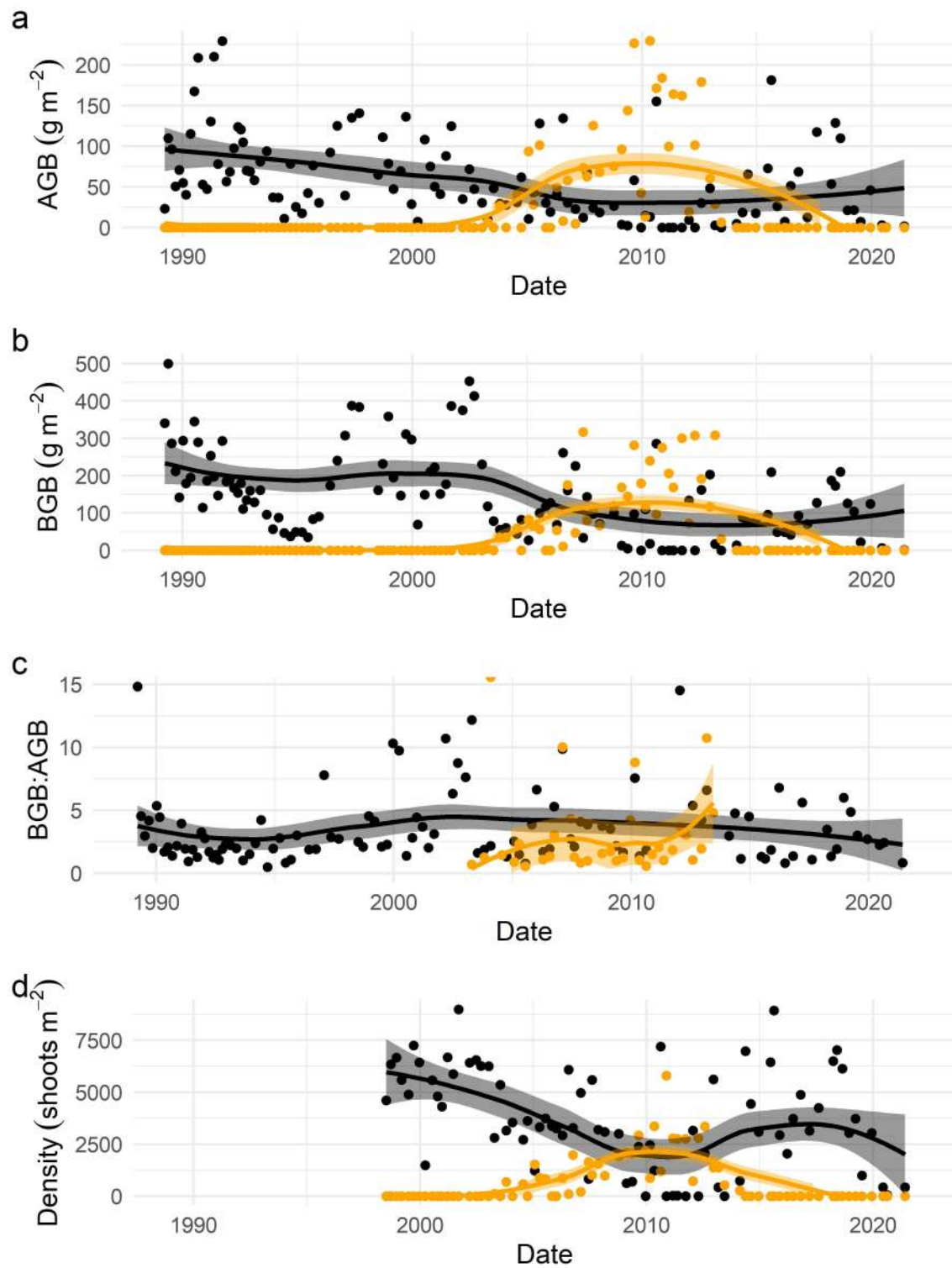


Figure 4. Seagrass community metrics measured at LM-151 from 1998 - 2022 include A) Aboveground biomass, B) Belowground biomass, C) Belowground:aboveground biomass ratios, and D) Shoot density.

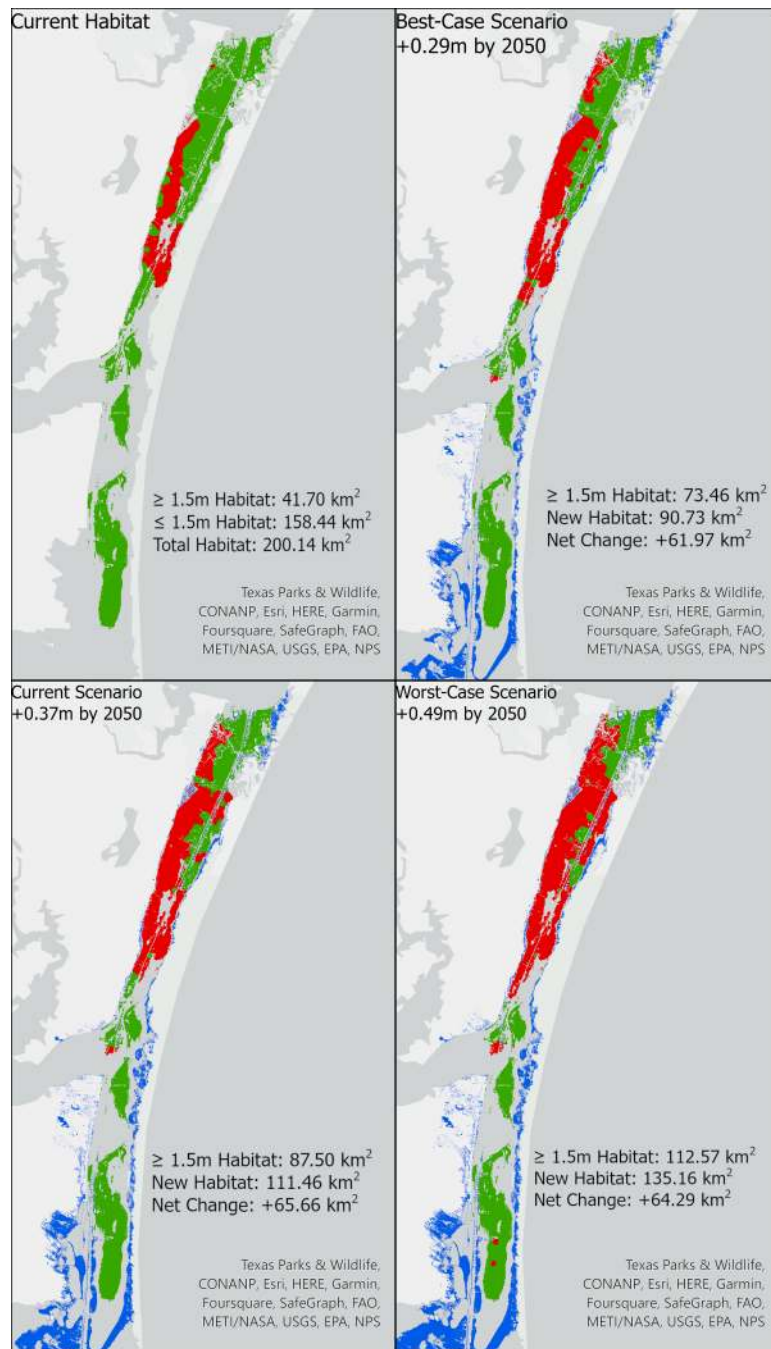


Figure 5. Potential seagrass distribution under current and future sea-level rise scenarios. Green represents areas with < 1.5 m water depth, red with ≥ 1.5 m water depth, and blue as newly submerged seafloor.

Country	$K_d < 1$ (km ²)	$K_d > 1$ (km ²)	% $K_d < 1$	% $K_d > 1$
TOTAL	622017	14773	98	2
United States	11829	2852	81	19
Mexico	24101	2161	92	8
Senegal	1174	1775	40	60
Ukraine	9878	1727	85	15
Nigeria	16035	1317	92	8
Cuba	11662	1305	90	10
Australia	59546	881	99	1
Denmark	57	630	8	92
Philippines	13612	629	96	4
Germany	464	489	49	51
Sierra Leone	8192	278	97	3
Canada	62	158	28	72
OTHER	465405	572	-	-

Table 1. Areal coverage of global seagrass meadows⁵³ classified using remotely-sensed K_d values⁵⁴ based on susceptibility to sea level rise ($K_d > 1$).

Supplementary Files

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