- A Systems-Level Analysis of Coastal Ecosystem Responses to Hurricane Impacts
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### 24 Abstract

Tropical cyclones are major disturbances for coastal systems. Hurricane Harvey made landfall in 25 Texas, USA, on 25 August 2017 as a Category 4 storm. There were two distinct disturbances 26 associated with this storm that were spatially decoupled: 1) High winds causing direct damage 27 and storm surge, and 2) High rains causing scouring floods and significant discharge of fresh 28 29 water carrying carbon and nutrients to estuaries. Here, we provide a synthesis of the effects of Hurricane Harvey on biogeochemical, hydrographic, and biotic components of freshwater and 30 estuarine systems and their comparative resistance and resilience to wind- and rain-driven 31 32 disturbances. Wind-driven disturbances were most severe along the coastal barrier islands and lower estuaries, damaging mangroves and seagrass and increasing sediment coarseness. Rain-33 driven disturbances were most pronounced within freshwater streams and the upper estuaries. 34 Large volumes of freshwater run-off reduced the abundance of riverine fauna and caused 35 hypoxic and hyposaline conditions in the estuaries for over a week. In response to this 36 37 freshwater input event, benthic fauna diversity and abundance decreased, but mobile fauna such as estuarine fishes did not markedly change. Although hydrographic and biogeochemical 38 39 components were highly perturbed, they returned to baseline conditions within days. In contrast, 40 biotic components demonstrated lower magnitude changes, but some of these organisms, particularly the sedentary flora and fauna, required weeks to months to return to pre-storm 41 42 conditions, and some did not recover within the 6 months reported here. Our synthesis illustrates 43 that resistance and resilience of system components may negatively co-vary and that structural 44 components of coastal systems may be the most vulnerable to long term changes following 45 tropical cyclones.

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## 48 Introduction

Hurricanes and tropical cyclones are one of the most destructive natural disturbances to 49 coastal environments, causing severe disruptions via storm surge, saltwater intrusion, wind 50 damage, and flooding (Paerl et al. 2001). Storm surges inundate terrestrial and freshwater 51 52 habitats with salt water, and associated waves scour estuarine benthic habitats (Mallin et al. 1999). High winds cause widespread damage to both natural and built environments, creating 53 54 significant fluxes of organic material and changing vegetation structure (Adger et al. 2005; 55 Armentano et al. 1995; Laurance and Curran 2008). Extreme precipitation mobilizes entrained sediments and nutrients and causes scouring floods in riverine ecosystems (Avery et al. 2004; 56 57 Schaefer et al. 2000).

Global climate models predict that the severity of these events will increase, and the 58 spatial distribution of regions that regularly experience hurricanes will shift poleward over the 59 60 next century (IPCC 2018; Knutson et al. 2010; Mann and Emanuel 2006). This makes it imperative to understand the factors that dictate how whole ecosystems respond to hurricanes. 61 However, the opportunistic nature of the majority of ecological studies of storm impacts has 62 63 resulted in a body of literature dominated by studies focused on particular components of study systems and often lacking adequate controls in space and time (Pruitt et al. 2019). Although 64 65 there are decades of studies documenting the effects of tropical cyclones on both developed and 66 natural systems, a conceptual understanding of factors that dictate ecosystems response to a 67 given storm is lacking (Pruitt et al. 2019).

A key concept in understanding how ecosystems respond to extreme events is ecosystemstability, which we consider separately as ecosystem resistance and resilience. We use the

following classic definitions: *resistance* is a measure of the system's ability to remain essentially 70 unchanged in the face of a perturbation, while resilience is a measure of a system's ability to 71 return to pre-perturbation condition (Harrison 1979; Pimm 1984). The resistance-resilience 72 framework can be used to enhance our understanding of the vulnerability of socio-ecological 73 coastal systems to extreme stress events (Adger et al. 2005). Taking a system level view-point, 74 75 by quantifying and comparing the resistance and resilience of different components of coastal ecosystems, we can determine where the greatest vulnerabilities lie and what intrinsic and 76 77 extrinsic factors are responsible for that vulnerability. Ultimately, this type of analysis is a step 78 towards improving understanding and enhancing predictive power about the impacts of future storms, and, ultimately, may help identify steps that can be taken to reduce impacts or to plan for 79 their effects. 80

Barring a concerted research consortium approach to studying future storms (Pruitt et al. 81 2019), meta-analysis offers us a path toward synthetic understanding of the mechanisms 82 83 controlling ecosystem responses to tropical cyclones. Two complimentary approaches to meta-84 analysis can be employed, 1) a broad survey of ecosystem responses across multiple storms 85 through space and time, or 2) an in depth comparative analysis of ecosystem responses to a 86 single event. The former approach has the advantage of potentially yielding broad generalizations, such as the observation that forest fragmentation synergistically enhances the 87 88 destructive effects of wind disturbance on tropical forests (Laurance and Curran 2008). 89 However, in depth comparative analysis of a single event can provide understanding about 90 cascading interactions driving responses. For example, Paerl et al. (2001) documented that high 91 nitrogen loads following Hurricanes Dennis, Floyd, and Irene into the Pamlico Sound, NC combined with long water residence time to drive a cascade of impacts including sustained 92

bottom water hypoxia, increased algal biomass, and increases in the instances of fish disease. 93 There are few examples of such synthetic studies of ecosystem responses to tropical cyclones 94 (Armentano et al. 1995; Greening et al. 2006; Paerl et al. 2001; Xi 2015), and fewer still that 95 contain quantitative analysis among ecosystems rather than verbal comparisons through literature 96 review. Basic questions that remain unanswered include: How does resistance and resilience to 97 98 tropical cyclones vary among ecosystem types (estuaries, barrier islands, and coastal rivers) and components (hydrographic, biogeochemical, mobile fauna, etc) within ecosystems? What role 99 does the identity of the stressor (wind force, rainfall, storm surge) play in mediating these 100 101 patterns?

In the fall of 2017, the Atlantic and Caribbean islands experienced one of the most 102 devastating hurricane seasons in recorded history, with three major storms including Irma, 103 Maria, and Harvey. Here we use one of these storms, Hurricane Harvey, as a case study. We 104 leveraged the high density of research programs evaluating the impacts of the event to document 105 106 patterns of resistance and resilience among different coastal habitats (estuaries, barrier islands, coastal plain streams and rivers) in response to different stressors. We include both a synthetic 107 108 analysis of the pattern of resistance and resilience within and among ecosystems and ecosystem 109 components and a series of analyses exploring individual responses to the disturbance event.

In addition to being one of the few such analyses of its kind, the present study has several novel features that add to our understanding of tropical cyclone affects. First, our dataset covers several different embayments and watersheds allowing for comparison among systems that experienced different levels of wind, rain, and storm surge stress. Hurricanes have a wetter side and stronger side where the storm picks up moisture and speed over warm water and then loses that energy as it rotates over land. Our region includes both the wet and dry sides of the storm,

as well as the area affected by the high rotational speeds of the eye of the storm. Second, we are not aware of any similar quantitative syntheses in sub-tropical coastal regions. Third, we employ the resistance-resilience conceptual framework an approach to quantify and comparing ecosystem responses, an approach that has not been used in other syntheses. The goal of these analyses is to provide a broad, ecosystem-level view of Hurricane Harvey's impacts on the affected coastal ecosystems.

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#### 124 Methods

### 125 Study System

The study focused on the south Texas coast, a subtropical area at the transition between 126 the mesic Coastal Plain and arid Sonoran Floristic Province (Noss et al. 2015; Sorrie and 127 Weakley 2001). The region consists of a series of lagoonal estuaries protected by barrier islands 128 129 and connected to the Gulf of Mexico by several narrow inlets (Montagna and Li 2010). Oriented from southwest to northeast, hurricane force winds impacted multiple estuaries, including the 130 131 upper Laguna Madre behind Padre Island, Corpus Christi Bay behind Mustang Island, the 132 Mission-Aransas Bay complex behind San Jose Island, and San Antonio Bay behind Matagorda Island (Fig 1). These estuaries are home to productive fish and shellfish fisheries valued by 133 134 commercial fisherman (\$196 million USD commercial catch value) and recreational anglers 135 (NOAA 2018).

At the point where Hurricane Harvey made landfall, strong winds (Category 4, gusts up to 241 km h<sup>-1</sup>) were recorded on Mustang and San Jose Islands, and within the Mission-Aransas estuarine system (Fig 1). High speed winds extended further east than west, with areas 40km to

the east experiencing 190kmph winds and areas 40km to the west experiencing 144kmph winds 139 (Fig. 1). In the area northeast of the eye, storm surge raised water levels 2-3 m above sea level 140 within Copano, Aransas, San Antonio, and Matagorda Bays. Harvey lingered over Victoria, 141 142 Texas for two days, and rainfall totals of 44 cm were recorded in Victoria County just 86 km from the landfall zone, whereas 65 km to the southwest in Nueces County (which includes the 143 city of Corpus Christi), rainfall totals were only 9 cm (Source: National Weather Service). Thus, 144 the effects of wind and rain disturbances were decoupled spatially with different bays 145 146 experiencing different combinations of wind and precipitation intensity.

147 Hurricane impact data were synthesized from freshwater, estuarine, and coastal wetland systems across the Texas coast ranging from the Upper Laguna Madre to San Antonio Bay (Fig. 148 1). Systems include nine coastal rivers, two barrier islands, and four major estuary complexes. 149 Samples consist of synoptic grab samples for water chemistry and nutrients, high frequency data 150 collections, physical habitat measurements, and quantitative surveys for flora and fauna. There 151 152 were 51 unique stations with temporal sampling of multiple responses in fixed positions in the dataset (Fig. 1), as well as repeated monthly sampling in random locations within estuaries for 153 mobile fauna (n=20 replicates per month per gear type). A complete list of station locations and 154 155 parameters measured can be found in S1 Appendix.

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#### 157 Storm Data

Wind speed data in the form of maximum 10 m wind gust during the storm and precipitation totals around the August 26<sup>th</sup> landfall date from August 24 to August 30, 2017 were provided by National Weather Service Offices in Corpus Christi, Brownsville, Austin/San

Antonio, and Houston/Galveston, TX. River discharge data were downloaded from USGS
gauging stations co-located with study sites within the affected region (USGS 2016).

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164 Hydrography

Continuous Water Quality Sampling - At one station in San Antonio Bay, Hydrolab 165 166 DS5X sondes (OTT HydroMet, Loveland, CO, USA) were deployed on the surface and bottom for continuous measurement (at 15-minute intervals) of salinity, temperature, and dissolved 167 168 oxygen from May-December of 2017. MiniDOT dissolved oxygen and temperature loggers 169 (PME, Vista, California, USA) were deployed in nine rivers from mid-August to December of 2017. Data for the entire monitoring period are reported as 24-hour averages of the high 170 frequency (15-minute interval) data. At two stations in Copano Bay, salinity, temperature (°C) 171 and turbidity (NTU) were measured continuously every 15 minutes by YSI 6600 V2 sondes 172 mounted on sampling platforms 0.5 m above the seafloor as part of the Mission-Aransas 173 174 National Estuarine Research Reserve (MANERR) system wide monitoring program. Data are available for download at the Centralized Data Management Office (CDMO, 175 http://cdmo.baruch.sc.edu). Sondes in San Antonio Bay were calibrated prior to deployment and 176 177 traded out on timescales from five days to two weeks depending on biofouling and weather conditions. Sondes in the coastal rivers were cleaned and data downloaded at two-four week 178 179 intervals. Sondes underwent rigorous post-deployment quality control check upon retrieval to 180 ensure data were not affected by biofouling or faulty probes. 181 Synoptic Water Quality Sampling – Coastal rivers were visited every two to four weeks

from August to December 2017. During each visit water quality measurements were taken at four stations within a fixed 75-m reach of stream using a YSI ProDSS Multiparameter Probe to collect turbidity (NTU), dissolved oxygen (mg L<sup>-1</sup>), temperature (°C), conductivity ( $\mu$ s cm<sup>-1</sup>), and pH. Probes were calibrated prior to use. Water quality measurements were taken at four stations in San Antonio Bay that were visited at two-week intervals from March to December 2017 to collect dissolved oxygen (mg L<sup>-1</sup>), temperature (°C), salinity, and pH. Secchi disk measurements were taken concurrently with water quality sampling in San Antonio Bay.

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## 190 Biogeochemistry

Water samples in the rivers were collected in replicates of four every two to four weeks 191 192 from August to December 2017. Samples were filtered in the field using 0.2-µm filters and frozen until analysis for DOC (dissolved organic carbon),  $NO_3^-$ ,  $NH_4^+$ , and orthophosphate. 193 Samples were analyzed for DOC as non-purgeable organic carbon (NPOC) using a Shimadzu 194 TOC-vcph with attached TNM-1 nitrogen analyzer (Shimadzu Corporation; Kyoto, Japan). 195 Samples were first acidified with 2% 2N HCl and sparged with zero CO<sub>2</sub> air for six minutes to 196 197 remove inorganic carbon, then they were analyzed for nutrients using a Lachat Flow Injection Auto-analyzer for  $NO_3^-$ ,  $NH_4^+$ , and orthophosphate. Estuarine surface water samples were 198 collected by the MANERR for monthly nutrient concentrations at two stations in Copano Bay. 199 200 Concentrations of NO<sub>3</sub><sup>-</sup>, NO<sub>2</sub><sup>-</sup>, NH<sub>4</sub><sup>+</sup>, and PO<sub>4</sub><sup>3-</sup> were determined on a nutrient autoanalyzer using standard colorimetric methods. 201

For water carbonate chemistry quantification, surface water samples were collected using a Van Dorn sampler at five system wide water quality monitoring program (SWMP) stations in the Mission-Aransas Estuary on a biweekly or monthly basis in 2017, and at seven stations in San Antonio Bay on a quarterly basis prior to June 2017 and then on a biweekly to monthly basis from June to December 2017. Total dissolved inorganic carbon (DIC) and pH were analyzed on these samples. Methods on sample collection, preservation, and analysis can be found in Yao and Hu (2017). Surface water CO<sub>2</sub> partial pressure (pCO<sub>2</sub>) at in situ temperature was calculated using lab measured DIC and pH (at 25°C) as the input variables. The carbonic acid dissociation constants were from Millero (2010), dissociation constant of bisulfate was from Dickson et al. (1990), and borate concentration was from Uppström (1974).

212 Wind speed data used in these two estuaries were downloaded from the NOAA meteorological stations located at Seadrift (Station 8773037, for San Antonio Bay) and Port 213 214 Aransas (Station 8775237, for Mission-Aransas Estuary). As the anemometers are deployed at 215 heights lower than 10 m (7.6 and 9.0 m, respectively), wind speed at 10 m was calculated using the power function in Hsu et al. (1994). Mole fraction atmospheric  $CO_2$  in dry air (xCO<sub>2</sub>) was 216 downloaded from http://www.esrl.noaa.gov/gmd/ccgg/trends. Atmospheric pCO<sub>2</sub> was then 217 calculated using xCO<sub>2</sub>, salinity and temperature (Weiss and Price 1980). CO<sub>2</sub> flux was 218 calculated using the following equation: 219

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 $F = kK_0(pCO_2, water-pCO_2, air)$ 

where k (m·d<sup>-1</sup>) is the gas transfer velocity calculated from wind speed. We used the formulation in Jiang et al. (2008); K<sub>0</sub> (mol·m<sup>-3</sup>·atm<sup>-1</sup>) is the gas solubility at in situ temperature and salinity (Weiss 1974). A positive F value indicates  $CO_2$  degassing to the atmosphere.

For estuarine chloropigment and accessory pigment measurements, surface water samples (1 L) were collected ~0.1 m below the surface and stored on ice until filtration through precombusted GF/F filters (Whatman 47 mm 0.7 μm). Pigments from GF/F filters and thawed sediments were extracted twice using acetone according to the protocol of Sun et al. (1991).
Briefly, 3-6 mL of acetone were added to the samples in centrifuge tubes, sonicated for 15 min, and centrifuged for 10 min. The supernatant was further filtered through a 0.2-μm syringe Nylon

filter. This extraction procedure was repeated again, and the filtered supernatants were 230 combined. The pigment analysis was accomplished through high performance liquid 231 chromatography (HPLC) with UV absorbance detection according to McTigue et al. (2015). The 232 233 mobile phases included 28 mM tetrabutylammonium acetate in methanol (30% : 70%; eluent A) and methanol (eluent B). After eluting through a C8 column (Agilent Eclipse XDB, 3.5 µm, 4.6 234 235 mm diameter x 150 mm length), individual pigments were detected by UV-Vis absorbance at a wavelength of 450 nm. Quantification was based on authentic standards (DHI and Sigma-236 Aldrich). The relative amounts of diatom and cyanobacteria were calculated based on established 237 algorithms from the accessory pigments (Reyna et al. 2017). 238

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#### 240 Mobile Biota

Gill nets were set during a 10-week period each fall, beginning with the 2<sup>nd</sup> full week in 241 September, 45/bay system (except 20 net sets in East Matagorda), no more than five and no less 242 243 than three net sets/week/bay. Monofilament nets (183 m long; 1.2 m deep with 45.7 m sections of 7.6, 10.2, 12.7, and 15.2 stretched mesh tied together in ascending order) were set overnight, 244 245 perpendicular to the shoreline with the 7.6 cm stretched mesh on the shoreward end, from 246 randomly selected locations. Twenty estuarine bag seines (18.33 m wide, 1.8 m deep with 1.3 cm stretched nylon multifilament mesh in the 1.88 m wide central bag and with 1.9-cm stretched 247 248 mesh in the remaining webbing) were pulled parallel to shore for 15.2 m at randomly selected 249 locations in each major bay each month. Twenty otter trawls (6.1 m wide with 3.8 cm stretched 250 nylon multifilament mesh) were pulled for 10 min in water  $\geq 1.0$  m depth at randomly selected 251 locations in each major bay each month. Date, location, water depth (m), salinity (psu), water temperature (°C), and dissolved oxygen (mg L<sup>-1</sup>) were recorded for each estuarine sample. 252

253 Organisms caught were identified to the lowest taxonomic level (typically species), counted, and 254 measured (total length in mm, TL, from the tip of the snout to the tip of the tail fully extended).

River fauna were sampled in September, October, November, and December 2017 at nine 255 256 streams in the Upper Laguna Madre, Mission-Aransas Bay, San Antonio Bay, and Matagorda Bay watersheds. Sampling was performed using a multi-pass depletion method, consisting of 257 258 three sampling passes using a backpack electrofisher (Smith Root LR-24) within a 75-meter reach bounded by block nets (McGarvey et al. 2017). Organisms caught were identified to the 259 260 lowest taxonomic level (typically species), counted, and the first 25 individuals of reach species 261 were measured (fork length in mm, FL, from the tip of the snout to the tail fork fully extended). Depletion curves were applied to the three-pass abundance data to estimate total abundance of 262 fish and crustaceans of each taxon within each reach (McGarvey et al. 2017). Fall 2017 data 263 were compared to surveys in April 2017 and October 2016 to establish baseline numbers for 264 expected faunal abundance. 265

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#### 267 Sedentary Biota

Mangroves were monitored at six sites as part of a long-term study on the mangrove-marsh 268 269 ecotone (Guo et al. 2017). Each site was dominated by black mangroves, Avicennia germinans, with ~10% cover of saltmarsh plants, including Batis maritima, Salicornia and Sarcocornia spp., 270 271 and Spartina alterniflora. At each site, a transect perpendicular to the shoreline was established, 272 extending at least 42 m from the water-vegetation interface. In  $1 \times 1$  m sub-plots along each 273 transect, we recorded either mangrove presence/absence or a visual estimate of percent cover. 274 Surveys were conducted in August 2015 and repeated after Harvey landfall in October 2017. For sites with presence/absence data, percent cover was estimated as the percent of sub-plots with 275

276 mangroves present. At other sites, mangrove cover is reported as the average across all sub-277 plots.

Seagrass percent cover was sampled at 126 fixed stations across the southern and central 278 279 Texas coast in the Mission-Aransas and Nueces estuaries in July and August 2017 prior to the impact of Hurricane Harvey as part of the Texas Statewide Seagrass Monitoring Program 280 281 (texasseagrass.org). Within 10 m of the GPS coordinate designating a station, four replicate 0.25  $m^2$  quadrats were placed. Within each quadrat, the percent cover of each seagrass species was 282 283 estimated with direct visual observation by a trained team of observers. Concurrent with each 284 sampling, in situ measurements were made of a suite of chemical and physical environmental parameters including salinity. All 126 sampling stations were re-surveyed in September to 285 October of 2017 and the absolute and proportional change in seagrass percent cover were 286 recalculated. Maximum sustained wind gust data was collected from 70 weather stations to 287 generate a raster of wind intensity for the coast. Interpolated wind speed values were extracted 288 289 for each sampling station, and linear regression was used to evaluate the relationship between 290 absolute change in seagrass cover and maximum sustained wind gusts experienced. 291 Benthic macroinfauna sampling has been performed in the Guadalupe Estuary (i.e., San 292 Antonio Bay) as part of long-term studies to identify the importance of freshwater inflow in controlling benthic dynamics (Montagna and Kalke 1992; Montagna and Kalke 1995; 293 294 VanDiggelen and Montagna 2016). Four stations are aligned along the salinity gradient within 295 the estuary from the freshest station A to the saltiest station D (Table 1). Macrofauna were 296 sampled with a 6.7-cm diameter core tube (35.4 cm<sup>2</sup> area) to a sediment depth of 10 cm. Three 297 replicates were collected per station twice prior to the storm (April and July 2017), and twice after the storm (October 2017 and January 2018). Organisms were extracted on a 0.5 mm sieve, 298

and enumerated to the lowest taxonomic level possible. Biomass was determined for higher
taxonomic groupings by drying at 55°C for 24 h.

Oyster dredges (Louisiana style 9-tooth; 46 cm wide, 25 cm tall with a 36 cm deep bag) were used to sample reefs from oyster-producing bays (Sabine Lake, Galveston, Matagorda, San Antonio and Aransas), 20 samples per month (except 30 per month in Galveston and 10 per month in Sabine). Dredges were pulled linearly at 3 knots h<sup>-1</sup> for 30 s from randomly-selected locations. Live and dead oysters were counted, and the percent live oysters were calculated for each sample. Pre-storm data were 4 mo before Harvey, May-August; post-storm data are 4 mo after Harvey, September-December.

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#### 309 Sediments

Sediment cores (8 cm I.D. × 30 cm length) were collected in the Mission-Aransas Estuary by push-corer. Triplicate cores were collected at each site. The top 5 cm of each core was sectioned and the rest was discarded; the top sections (0-5 cm from surface) from the triplicate cores were combined and homogenized in a plastic storage bag and stored in a cooler on ice until return to the lab on the same day. Sediment samples were then freeze dried and screened with a 300-µm mesh to remove large debris and shells.

Sediment grain size was measured using a laser diffraction particle size analyzer
(Beckman-Coulter LS 13 320). About 15-20 mL of hydrogen peroxide (1:2) were added to ~0.2
g of sediment in a beaker. After incubating for 24 h at 40°C to remove organic matter, sodium
hexametaphosphate [(NaPO<sub>3</sub>)<sub>6</sub>] was added to the sediment and mixed for 15 min in an ultrasonic
bath to aid disaggregation. The size distribution was measured by polarization intensity
differential scattering. The detection size range of this analyzer is from 0.02 to 2000 µm.

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## 323 Statistical Analyses

Resistance was calculated as the natural log of the maximum or minimum post-storm 324 value divided by the baseline value (hereafter referred to as the Log Response Ratio or LRR). 325 Resilience was calculated as the number of days until values returned to baseline. Baseline was 326 327 calculated as the mean value immediately before Hurricane Harvey in the case of spatially replicated data or the mean value for August in the case of temporally replicated data. In the 328 329 case of responses which did not return to baseline, all values were assigned the same value (140 330 days) to eliminate variation caused by differences in sampling frequency. All analyses were conducted using the statistical program R version 3.4.3 (R Core Team 2016). To evaluate the 331 relationship between resistance (LRR) and resilience (return time), the LRR of the absolute value 332 of the deviation from baseline was regressed against the return time using quantile regression. 333 Absolute value was chosen to focus the analysis on the magnitude rather than the direction of the 334 335 response. Quantile regression was chosen to evaluate the relationship between resistance and resilience because multiple factors may simultaneously impact resistance driving variation in 336 337 response magnitude, but there is a physical limit on the potential magnitude of a response 338 (Gotelli and Ellison 2013). Two quantile regression models, tau of 0.9 (high slope) and tau of 0.5 (no slope) were compared to evaluate the existence of the relationship using ANOVA in the 339 340 quantreg package (Koenker 2018). Differences in resistance and resilience among response 341 categories were evaluated using ANOVA followed by a Tukey-HSD post-hoc test. Tests for 342 temporal changes were evaluated as mixed effects models using the *lme()* function in the *nlme* 343 package with site identity treated as a random effect with an autocorrelation structure of order 1 (Pinheiro et al. 2017). Changes between paired time points in significant temporal series were 344

evaluated using t-tests. PERMANOVA was used to test for changes in community composition
between time points using the *vegan* library (Oksanen et al. 2014).

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348 **Results** 

## 349 *Resistance and Resilience Among Ecosystems*

We observed a negative relationship between the LRR of the absolute value of the 350 deviation from baseline (resistance) and the time to return to baseline (resilience) across response 351 variables, (slope = -0.018, F = 9.732, DF = 1, P = 0.002, Fig 2A). The response magnitudes 352 353 within ecosystem components were similar across different ecosystems (Fig 2B-2C). Hydrologic and hydrographic variables in both the rivers and estuaries displayed the highest magnitude shifts 354 from pre-storm conditions representing relatively low resistance to disturbance (Fig 2A-B). 355 However, hydrographic conditions returned to pre-storm levels significantly faster than other 356 ecosystem attributes (mean  $\pm$  SE = 18.56  $\pm$  21 d, P < 0.005, Fig 2A, 2C), indicating these system 357 358 properties had the highest resilience. Biotic variables generally displayed negative responses to wind and rain with higher resistance in sedentary biota as compared to mobile biota. The mobile 359 360 biota within the rivers and estuaries, however, were more resilient than sedentary biota (Fig 2C, 361 *P* < 0.001).

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## 363 *Hydrology*

River discharge in streams and rivers within the region increased by 264% to 103,600%, within a few days of the storm. The responses were the highest magnitude of any variable in the dataset. The highest discharge events occurred to the east of landfall, following the asymmetric pattern of precipitation associated with the storm (Fig. 1). Northeastern sites that received more rainfall took longer to return to pre-storm flow conditions (Fig. 3A), while south-western sitesthat received less rain returned to pre-storm flow conditions more rapidly (Fig. 4A).

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371 *Hydrography* 

Response magnitude and return time was highly variable among hydrographic variables. 372 373 The response magnitudes and return times were higher in the estuaries than in the rivers. This can largely be attributed to differences in water residence time, with shorter residence time 374 related to faster return time. Salinity in San Antonio Bay initially quadrupled over a 12 h period 375 376 (from 7.39.6  $\pm$  0.03 SE to 32.09 30.9) due to storm surge, then dropped over the following 5 days due to increased discharge from the Guadalupe River. Low salinity conditions (< 5 ppt) lasted 377 for over one month in San Antonio Bay (Fig. 5). Within Copano Bay, salinity dropped from 18.9 378 to 3.3 ppt from rain and river discharge (Fig. 3A), and low salinity conditions took > 7 months to 379 return to pre-hurricane levels, indicating low resilience (Fig. 3B). In Copano Bay, turbidity 380 381 increased due to resuspension of fine material from wave action and surge, peaking at >1300NTU, the highest recorded level since the sensors were installed in 2007, but this increase in 382 turbidity lasted only 2 days (Fig. 3A). Two days after the storm made landfall, there was a 383 384 secondary peak in turbidity (following a prior peak associated with storm surge) associated with riverine run-off (Fig. 3A). 385

Dissolved oxygen concentrations varied among rivers, with bottom water anoxia occurring in rivers that experienced high discharge and loss of diel cycling for 5-18 days after the hurricane passed. Following resumption of diel cycling, the amplitude of the diel oxygen cycle was reduced by 44% ( $2.3 \pm 4.7 \text{ mg L}^{-1}$ ). For example, in the Aransas River, average maximum and minimum DO dropped from 7.47 ± 1.54 to 3.27 ± 1.78 mg L<sup>-1</sup> and 1.95 ± 0.49 to 0.44 ± 0.16

mg L<sup>-1</sup> respectively, for 7 days after the storm. Within the San Antonio Bay estuary, concurrent
with the onset of high river flow and low salinity conditions (Fig. 5A), hypoxic and anoxic
conditions formed at the bottom and near the surface. Hypoxic conditions persisted for
approximately 8 days in near-surface and bottom waters simultaneously (Fig. 5B).

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### 396 Biogeochemistry

Biogeochemical responses in both the rivers and estuaries were all positive and were an 397 398 intermediate magnitude relative to other responses (Fig 2). Responses generally returned to pre-399 storm levels quickly; however, NO<sub>3</sub>- and TDN (total dissolved nitrogen) in the rivers were an exception. NO<sub>3</sub>- and TDN continuously increased post-storm, rising more than 300% in some 400 systems and remaining elevated. For example, NO<sub>3</sub><sup>-</sup> concentrations reached  $14.26 \pm 0.46$  mg L<sup>-1</sup> 401 in the Aransas River by December 2017 (df=1,4, P = 0.0005). Ammonium concentrations were 402 not affected by the storm (df=1,4, P = 0.763). Orthophosphate concentrations within rivers 403 404 increased immediately after the storm and then returned to pre-storm levels within a month (df=1,4, P = 0.0004). Concentrations of orthophosphate increased by as much as 1300% in some 405 406 systems, reaching as high as  $8.83 \pm 1.26$  mg L<sup>-1</sup>. However, nutrients including NO<sub>3</sub>-, NH<sub>4</sub><sup>+</sup>, and 407 orthophosphate remained at low levels in Copano and Aransas Bays on September 13 (first sampling date after the hurricane) until December 2017. During this period, chlorophyll-a 408 409 concentration (a proxy for total phytoplankton biomass) increased relative to pre-storm 410 conditions (June 2017; Fig. 3B).

Within the rivers, DOC (dissolved organic carbon) increased immediately after the hurricane with concentrations rising more than 250%, reaching as high as 26.35 mg L<sup>-1</sup>  $\pm$  0.412 SD in some systems before declining again (df=1,4, *P* < 0.001). Within the estuaries, CO<sub>2</sub> fluxes

to the atmosphere also shifted, and similar to DOC fluxes, the response was variable across bay 414 systems. In "normal" years, water-to-air CO<sub>2</sub> flux in San Antonio Bay is 98.4 mmol-C m<sup>-2</sup>d<sup>-1</sup> 415 (Yao et al. 2020). An extreme increase in  $pCO_2$  levels one month after the disturbance led to 416 nearly 50% increase in CO<sub>2</sub> flux on an annual scale (144.5 mmol-C m<sup>-2</sup> d<sup>-1</sup>) (Fig. 5A). San 417 Antonio Bay then quickly changed from a CO<sub>2</sub> source to a sink in October (uptake of CO<sub>2</sub> from 418 419 the atmosphere), consistent with previous multiyear observations, indicative of returning to "normal" estuarine conditions for this time of the year. In contrast, the adjacent Mission-Aransas 420 Estuary saw a decrease in CO<sub>2</sub> flux (14.7 mmol-C  $m^{-2} d^{-1}$ ) after the hurricane. 421

422

423 Mobile Biota

Mobile biota responses to and recovery from the storm differed between riverine and 424 estuarine assemblages. Across all rivers, fish and crustacean abundance was reduced by 62-95% 425 after the hurricane (Fig. 6A). In some coastal rivers, fish and crustacean biodiversity was also 426 427 reduced and composition shifted toward abundant estuarine species such as anchovy (Anchoa *mitchilli*) and daggerblade grass shrimp (*Palaemonetes pugio*). Abundances of riverine fish and 428 crustaceans recovered to pre-storm levels by November 2017 (df=1,3, P < 0.0001). In contrast, 429 430 when comparing estuarine gill net data from pre-storm to post-storm periods (Fall 2016 vs. Fall 2017), there were minor shifts in fish community structure in all Texas estuaries except the 431 432 Laguna Madre (see supplementary material). Monthly seine data from July 2017 to December 433 2017 showed a typical seasonal pattern of fish and crustacean abundance significantly declining until October and then rebounding with the arrival of the fall recruits (df = 1.5, P = 0.003, Fig. 434 435 6B, (Nelson 1992)). The reduced catch per unit effort (CPUE) from August to September 2017 436 was larger (df=9,32, P = 0.01) in Aransas (-74%), San Antonio (-66%), Corpus Christi Bay (-

437 40%), and Upper Laguna Madre (-36%) than is typical from the previous 6-year period (2010 to 438 2016 mean change in CPUE from August to September for Aransas Bay:  $-31\% \pm 11$  SE, San 439 Antonio Bay:  $-10\% \pm 23$  SE, Corpus Christi Bay:  $6\% \pm 19$ , and Upper Laguna Madre:  $-21\% \pm$ 440 13, Supplement 2G).

441

### 442 Sedentary Biota and Physical Responses

Sedentary taxa, including vegetation and sessile benthic taxa, all exhibited negative but 443 444 low-magnitude responses to the disturbance; however, return times were the highest among 445 measured responses (beyond the duration of the current study). In the coastal wetlands just south of the storm track, black mangrove (A. germinans) cover dropped 25-40%. The decline in 446 mangrove cover was largely due to loss of upper foliage, though some trees were killed after 447 being entirely uprooted and displaced. Foliage on lower branches remained largely intact, likely 448 because the lower branches were protected from the wind by submersion via storm surge. 449 450 The strongest winds were within the Mission-Aransas Estuary, which resulted in physical removal of seagrass. There was a negative relationship between absolute change in percent 451 452 cover of turtle grass (*Thalassia testudinum*) seagrass and maximum sustained wind gusts 453 experienced at a site ( $R^2 = 0.15$ , slope = -0.18). Overall, 12% of stations with T. testudinum present pre-storm lost 100% of T. testudinum cover post-storm and 30% of stations lost at least 454 455 50% of pre-storm cover. These severe declines were only in areas that experienced Category 3 or 456 Category 4 force winds (178-251 km h<sup>-1</sup> sustained wind speed). As noted by Congdon et al. 457 (2019), seagrass damage included both complete removal (roots/rhizomes ripped from the 458 sediment) and partial removal (above-ground biomass sheared off). In areas where rhizome

459 material remained intact, regrowth of above ground tissue occurred within 1-3 months (pers.460 obs., VMC).

461	Many benthic estuarine faunal assemblages exhibited negative responses to the storm
462	event. There was significant difference in percent live oysters among months in Aransas Bay
463	between July and December 2017 (df =5,104, $P = 0.0334$ ). The largest drop in live oyster cover
464	occurred between August (pre-storm; 54%) and September (post-storm; 22%). In San Antonio
465	Bay, benthic macroinfaunal diversity ( $P < 0.0001$ , df = 15,32), abundance ( $P < 0.0001$ , df =
466	15,32), and biomass ( $P < 0.0001$ , df = 15,32) all declined significantly after the storm. Four
467	months prior to the storm in April 2017, abundance was as high as 53,900 individuals m <sup>-2</sup> .
468	Following the storm, abundance dropped to 9,800 individuals m <sup>-2</sup> in October 2017 and remained
469	low (9,400 individuals m <sup>-2</sup> ) through January 2018. The biomass of the macroinfauna declined
470	from a maximum of 38.6 g m <sup>-2</sup> in July 2017 to a maximum of 6.1 g m <sup>-2</sup> in October and 1.7 g m <sup>-2</sup>
471	in January 2018. Benthic macroinfaunal species richness declined from a maximum of 10.3
472	species/core prior to the storm to 3.3 species/core in October 2017 and then increased to 6.7
473	species/core in January 2018. The declines were due to losses of 22 of the 49 species found.
474	However, the greatest losses were for the polychaetes (Mediomastus ambiseta, Streblospio
475	benedicti, and Capitella capitate), oligochaetes, and the mollusks (Rangia cuneata and Macoma
476	mitchelli). Post-storm, the benthic macroinfauna community composition shifted, and recovery
477	was driven by a recruitment event of juveniles in January and April 2018 of the mollusks
478	Texadina sphinctostoma and Mulinia lateralis, and the polychaete Spiochaetopterus costarum.
479	Several changes in the estuarine edaphic (sediment) characteristics in Mission-Aransas
480	Bay followed the storm. Median grain size in surface sediments (0-5 cm) increased as much as
481	100 microns at a sampling site near Harbor Island after the hurricane. Benthic chlorophyll-a

482 concentrations declined, possibly resulting from wind-induced scouring or burial. There was 483 also a large drop in the percentage of benthic pheophorbide from  $15 \pm 6\%$  in June 2017 to  $4 \pm$ 484 2% in October 2017. Pheophorbide concentrations approached pre-storm baseline levels by 485 January 2018 ( $12 \pm 5\%$ ).

486

### 487 Discussion

We present one of the most comprehensive syntheses of impacts of a major hurricane on 488 489 coastal systems to date, in terms of the geographic extent of systems examined and the breadth of 490 response variables. The analyses reveal an important pattern of negative covariance between resistance and resilience among ecosystem components. Hydrographic and biogeochemical 491 components of the system, while displaying some of the largest magnitude changes from pre-492 storm condition (low resistance), returned to baseline quickly (high resilience). Even though the 493 influx of freshwater and nutrients was large, rivers and estuaries are dynamic systems and thus 494 495 may be able to buffer and absorb these types of disturbances. In contrast, although structural components of ecosystems such as seagrasses, mangroves, and oyster reefs were 496 497 overwhelmingly more resistant to disturbance than factors like water chemistry and mobile 498 fauna, recovery time for these structural components can take years or even decades (Ilg et al. 2008; Levin 1984; Lytle et al. 2008). These patterns, and the observed resistance/resilience 499 500 covariance enhances our understanding of how tropical cyclones impact coastal systems. 501 Furthermore, our analysis framework is flexible and can be used in future studies to measure and 502 compare hurricane responses across abiotic and biotic variables.

503 Hurricane Harvey impacted coastal ecosystems through both the intense wind and storm 504 surge disturbance which lasted less than 24 hours within the study region, and through record-

breaking rainfall leading to large amounts of run-off, which ultimately altered salinity and 505 chemical conditions. Although there were interactions between wind- and rain-driven 506 disturbances, many responses within biological, hydrographic, and biogeochemical components 507 508 are most likely attributed to one or the other. Furthermore, there was spatial separation in the major impact zone of the physical and rain-driven disturbances, which facilitated examination of 509 the effects of each of those drivers. Many of the responses by flora and fauna can be attributed 510 to the immediate impact of physical disturbance, but the drivers differed among groups. For 511 512 example, scouring from rain induced floods likely impacted riverine mobile fauna, whereas 513 scouring resulting from high winds impacted estuarine macrofauna and seagrasses. In contrast, many of the shifts in hydrographic, hydrologic, and biogeochemical variables were likely largely 514 driven by the rainfall event. We examine these drivers and responses below. 515

516 Impacts of the storm on physicochemical responses varied both spatially and between estuarine and freshwater systems, we discuss each of these in turn. Riverine concentrations of 517 518 most dissolved solutes, in particular nutrients and DOC, increased for only a short period of time after the storm. These findings are consistent with those predicted by the Pulse-Shunt Concept, 519 which states that major hydrologic events drive the timing and flux of terrestrial DOM to aquatic 520 521 ecosystems (Raymond et al. 2016). When water tables rise during storm events, organic matter is leached from soils and flushed into streams and rivers (Boyer et al. 1997; Hornberger et al. 522 523 1994). Differences in the yields of DOC from watersheds in response to Hurricane Harvey are 524 likely driven by a combination of land use, topography, and antecedent conditions (McMillan et 525 al. 2018). Wetlands and topographic depressions are sources of DOC within watersheds that can 526 be flushed during storm events (Creed et al. 2003; Richardson et al. 2010). The flux of landscape 527 sources of DOC downstream is also driven by the degree to which a flood event increases

as Hurricane Harvey is delivered to coastal ecosystems where it enhances heterotrophic 530 microbial respiration (Crosswell et al. 2014; Watanabe and Kuwae 2015). 531 The observed impact of the storm on riverine NO<sub>3</sub>-concentrations, which increased and 532 533 remained elevated for months after the storm, was significantly different from responses of other solutes. The observed pattern mirrors the effect of Hurricane Hugo on stream water chemistry in 534 Puerto Rico, where NO3<sup>-</sup> concentrations increased 4-fold in response to the storm and remained 535 536 elevated for 12-18 months (Schaefer et al. 2000). Reduced nutrient uptake in the terrestrial environment due to loss of aboveground vegetation during the storm combined with leaching 537 from downed litter is potentially responsible for the prolonged elevation of water column 538 nutrient concentrations (Schaefer et. al. 2000). Increased nutrient loading into streams from 539 hurricanes often results in higher dissolved nutrient concentrations in the estuarine water column 540 541 (Wachnicka et al. 2019).  $NO_3^{-1}$  is particularly prone to elevated concentrations because other forms of dissolved nitrogen rapidly convert to  $NO_3^{-1}$  and positive phosphorus ions are likely to be 542 bound to negatively charged sediment particles. However, following Harvey, the receiving 543 544 estuaries showed no such prolonged elevation in dissolved nutrient concentrations despite the increased nitrogen load. This phenomenon could be attributed to rapid uptake by phytoplankton. 545 546 Overall chlorophyll-a concentrations were elevated relative to previous years, where the 547 maximum concentrations are typically observed during the summer (Reyna et al. 2017). In 548 particular, diatoms, a dominant group in Aransas and Copano Bays (Anglès et al. 2015), are 549 known to respond rapidly to elevated nutrient input (Colos 1986; Pinckney et al. 1999). Taxa 550 common in this region can bloom after freshwater inflow events (Anglès et al. 2015) and

surface water connectivity of upland DOC sources to streams and rivers (Hosen et al. 2018).

Ultimately, much of the terrestrial organic matter mobilized during extreme weather events such

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subsequently cause a rapid decrease in nutrient concentrations during blooms (Popovich et al.2008).

Differences in response magnitudes among the different estuaries can likely be attributed to 553 variation in the amount of rainfall received and/or the differences in geomorphology of the 554 basins contributing to variation in water residence time. For example, the difference in CO<sub>2</sub> flux 555 556 between Mission-Aransas Bay and San Antonio Bay can be attributed to spatial variation in rainfall. In San Antonio Bay, enhanced CO<sub>2</sub> flux post-storm was likely a combination of 557 558 enhanced respiration of river-transported terrestrial DOC and river water degassing because of 559 high DIC concentrations. The Mission-Aransas Estuary showed more of local precipitation influence, as dilution of seawater increases CO<sub>2</sub> solubility (Yao and Hu, 2017), and local runoff 560 may have flushed nutrients from the watershed into the bay, fueling primary production. 561 However, the intensity of the rainfall effects were modulated by estuary residence time. Where 562 residence time was longer, the impacts had a greater magnitude, and effects persisted for longer 563 564 periods of time. For example, Aransas Bay and Copano Bay (part of the Mission-Aransas Estuary system) received similar amounts of rainfall but had different freshwater inflow impacts. 565 566 The seven month period of low salinity in Copano Bay (compared to 1 month in Aransas Bay) 567 following the storm can be attributed to its ~1.5 year residence time (Solis and Powell 1998). High levels of precipitation combined with long water residence time may have cascading 568 569 impacts on other ecosystem properties that are less resilient. For example, bottom water hypoxia 570 (dissolved oxygen concentration less than  $2 \text{ mg } L^{-1}$ ) in conjunction with low salinities likely 571 drove losses in estuarine benthic infauna, which took longer to recover. 572 In general, fauna (mobile and sedentary) were among the more resistant components of the

573 system evaluated. This resistance may be explained by a combination of adaptation to

environmental fluctuations, and in the case of mobile fauna, the ability to move to avoid them. 574 Estuaries and rivers are inherently dynamic environments and thus associated fauna are often 575 adapted to respond to large swings in environmental conditions, like salinity or scouring floods, 576 577 making estuarine and riverine fish and crustaceans less vulnerable to disturbance events (Frid and Townsend 1989; Townsend 1989). For example, estuarine fish typically exhibit a high 578 tolerance for salinity fluctuations (Nordlie 2003). Mobile fauna showed higher resilience to 579 disturbance than sedentary fauna, suggesting dispersal/movement ability may be key in 580 581 mediating organismal responses. This is further supported by the observation that recovery 582 intervals were shorter (weeks-months) than the reproductive cycle of many of these taxa (annual). This suggests that mobile biota were able to move out of affected areas and/or 583 recolonize the systems quickly (Bell and Eggleston 2005; Massie et al. this volume). However, 584 we did not evaluate whether mobile fauna had higher tolerance to environmental fluctuations 585 than sedentary fauna, and so the mechanism behind the observed differences requires further 586 587 evaluation.

Seasonal recruitment dynamics may have also played a role in rapid recovery of mobile 588 589 fauna. For example, there is typically a drop in the abundance of estuarine fish in early fall in 590 these systems followed by an increase with the arrival of the fall recruitment class for some species (Heck et al. 2003; Reese et al. 2008). The results from nearshore seine data suggest that 591 592 the storm may have exacerbated the natural seasonal cycle, causing a larger dip than usual, but 593 the effects were short-lived (Fig. 6C). The seasonal arrival of fall recruits may have contributed 594 to the rapid recovery in estuarine fish populations; however, this was likely not the only 595 mechanism. Trawl data from these same systems within open waters showed no temporal 596 patterns (Fig. 6D), suggesting that high wave energy and tidal shifts concentrated the storm

impacts on mobile fauna along the shorelines. Therefore, recolonization to the nearshore zone 597 may have also come from populations that sought refuge in open water during or preceding the 598 storm. These observations contrast with those made in Florida Bay after Hurricane Irma, where 599 600 the relative abundance of many fish species declined by more than 50%, and pre- and post-storm assemblages were substantially dissimilar, largely due to increases in anchovies and declines in 601 602 mojarra and killifish (Zink et al. *this volume*). In the rivers, there was an increase in estuarine fauna and secondary freshwater fishes immediately after the storm before returning to a state 603 604 dominated by primary freshwater fauna. These taxa may have been pushed up into the streams 605 by the storm surge or could have colonized quickly from refugia in the downstream estuary. Sedentary biota (fauna and flora), in contrast, could not actively avoid stressful physical 606 (high wave energy) or hydrographic (low dissolved oxygen) conditions, and recolonization may 607 take longer, as it likely requires dispersal of new recruits or vegetative growth into denuded 608 areas. Dynamics in these groups were most likely a function of taxa-specific life history 609 610 characteristics and disturbance intensity. For example, impacts on estuarine benthic sedentary biota can be attributed to low salinity and anoxia such as the effects of low salinity and bottom 611 612 anoxia resulting from freshets on oyster mortality (Munroe et al. 2013), as was observed in 613 Galveston Bay following Hurricane Harvey. However, significant physical disturbance from storm surge has the potential to directly damage oyster reefs, and this is the likely explanation for 614 615 declines in live oysters in Aransas Bay following the storm, as salinities (> 5) were not low 616 enough (< 3.5) for sufficient time to elicit oyster mortality (Galtsoff 1964; La Peyre et al. 2009). 617 Saltmarsh grasses were largely resistant to physical impacts, likely because these low-stature, 618 flexible plants were either submerged or were able to lie flat when the wind was strong 619 (Armitage et al. 2019). In contrast, the taller, more rigid mangroves suffered greater defoliation;

this damage pattern is typical in many tropical mangrove species (Branoff *this volume*). This 620 pattern of damage is characteristic of major wind events, and is typically constrained to the area 621 near the landfall of the storm's eve with the highest wind speeds (Armentano et al. 1995; Smith 622 623 et al. 2009). Mangrove stands on the Texas coast are dominated by a single rapidly growing species, A. germinans (Tomlinson 2016). Storm damage to the upper branches A. germinans was 624 625 followed by rapid resprouting of leaves within two months of the storm, though complete canopy recovery may take multiple growing seasons (Armentano et al. 1995). Similar patterns of 626 627 damage and rapid recovery in A. germinans followed Hurricane Andrew in Florida in 1992 (Baldwin et al. 2001; Smith et al. 1994). Avicennia germinans is relatively resilient to these 628 types of disturbances, and even complete defoliation does not necessarily result in tree mortality 629 (Imbert et al. 2000; Roth 1992). 630

Recovery of seagrasses will likely differ substantially than that for mangroves because in 631 many cases, whole plants were uprooted (Congdon et al., 2019). One of the two dominant 632 633 seagrass species in our region, T. testudinum, has one of the slowest rhizome elongation rates among Atlantic seagrasses (Duarte 1991), which may further slow recovery of disturbed areas 634 when its rhizomes are physically disturbed or the beds are buried by storm sediment deposition 635 636 (Hernandez-Delgado et al. this volume). Recovery of T. testudinum in areas that have experienced physical disturbance from boat groundings has been observed to be 2 to 7 years 637 638 (Bourque et al. 2015; Zieman 1976), and many disturbed areas from Harvey are of an even larger 639 spatial extent (Congdon et al., 2019). However, sexual reproduction and seed recruitment may 640 enhance recovery (Whitfield et al. 2004) and sexual reproductive effort is high in T. testudinum 641 seagrass beds in South Texas (Kaldy and Dunton 2000).

Looking across mobile and sedentary biota, the results imply that the greatest long-term 642 impacts occurred on sedentary biota. This phenomenon has been observed in other systems 643 following extreme events, such as the catastrophic impact of Hurricane Agnes on Chesapeake 644 Bay in 1972, where mobile fauna recovered quickly after the storm, but submersed aquatic 645 vegetation took decades to recover (Orth and Wilcox 2009). Prolonged recovery trajectories of 646 647 sedentary biota are often linked to habitat fragmentation and small patch size. In forest responses to hurricanes, fragmentation led to negative feedback loops once extreme events occur in the 648 649 system (Laurance and Curran 2008). Looking to marine systems, seagrasses self-buffer against 650 perturbations once a critical patch size is reached (Gruber et al. 2011; Orth et al. 2017; van der Heide et al. 2011), and similar processes likely operate on oyster reefs (Moore et al. 2018) and 651 mangrove stands (Huisman et al. 2009). The integrity of structural habitat has cascading positive 652 653 effects on sediment accretion and shoreline stability, potentially promoting recovery to pre-storm conditions. The implications are clear - management actions to conserve large swathes of 654 655 structural habitat may be key to enhancing overall coastal ecosystem resilience, and management interventions to restore structural habitat may be critical to promote rapid recovery of coastal 656 657 ecosystems following major storms.

Considering all responses, we propose a general conceptual model that resistance and resilience in coastal ecosystems is a product of the relative influence of physical and biological constraints on the response variable of interest, and that these are hierarchically arranged with biological constraints following physical constraints. For example, the majority of conservative solutes (not subject to uptake by the biota) returned to pre-storm levels rapidly via settlement and open ocean exchange; however, there were noted deviations from this general pattern. Systems with high residence time remained perturbed for longer periods, reflecting the physical constraint

that basin morphology can exert on water movement. Similarly, the observation that riverine 665 NO<sub>3</sub><sup>-</sup> concentrations increased over time and remained elevated may reflect the biological control 666 that watershed vegetation, which requires months to years to re-grow post-storm, exerts on 667 riverine  $NO_3$ - concentrations through uptake (Likens et al. 1970; Schaefer et al. 2000). In 668 contrast, intermediate levels of resilience in estuarine biogeochemical responses likely reflects 669 670 the population cycling of phytoplankton, which is comparatively much vaster than terrestrial vascular plants (Laws 2013). This extends further to biological responses where, provided 671 672 environmental conditions are suitable, re-colonization and recovery must be a function of 673 dispersal ability, distance to source populations, and generation times (Levin 1984; Lundquist et al. 2010). 674

Examining the impacts of Hurricane Harvey provides us with an opportunity to forecast 675 how ecosystems along the coastline will respond to future hurricanes. We found opposing 676 spatial gradients in the two major forms of disturbance associated with the storm; rain driven 677 678 effects, which were more pronounced in rivers and within the upper estuaries to the northeast of landfall, were spatially decoupled from wind effects, which were most pronounced near the 679 landfall location on the coast. The analysis of Hurricane Harvey displays how variation in the 680 681 identity of storm stressors and ecosystem components dictates the magnitude of the immediate and long-term impacts on coastal ecosystems. The intensity, high rainfall totals, and climatic 682 683 conditions of the impact zone are all characteristics of predicted future storm scenarios for the 684 US Gulf of Mexico and Atlantic coasts (Emanuel 2017; Seager et al. 2007; Seager et al. 2013). 685 Thus, the impacts of Hurricane Harvey may be a preview of the new normal of hurricane impacts 686 and responses in these regions.

688	Supple mentary	<b>Materials</b>
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689 Supplementary methods and results, and all data used for the analyses are given in SI Appendix. 690

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Fig. 1 Spatial distribution of hurricane weather and sampling sites. Total precipitation from August 25-30 was interpolated using an inverse spline from weather stations throughout the region and is displayed as a color map (source: National Weather Service). Maximum wind speed during the storm was similarly interpolated from weather stations and displayed as contours with red indicating the zone of highest wind speeds. Sampling stations from which data were collected are depicted as black dots.



Fig. 2 Hurricane Harvey Resistance - Resilience synthesis. A) The X-axis is the return time to baseline conditions in days, representing system resilience (systems which had not returned to baseline were assigned a value of 140 days to avoid biases due to variation in sampling frequency). The Y-axis is Log Response Ratio (LRR), the natural log of the maximum response divided by the baseline value, representing system resistance. Each symbol represents a time series of a particular response for a particular ecosystem type. Symbols with error bars (Standard Error) had multiple spatial locations with the time series. Symbol shape

represents system type (Triangle = River; Circle = Estuary), and symbol color represents the ecosystem response type (dark blue: biogeochemical e.g., nutrients, carbon flux, chlorophyll-a; blue: hydrographic e.g., oxygen, temperature, pH, salinity; light blue: hydrologic e.g., river discharge; red: mobile fauna e.g., invertebrates & fish; green: sedentary fauna e.g., plants & benthic invertebrates; yellow: physical e.g., sediment grain size, TSS). The dotted lines are the quantile regression lines (tau = 0.9, P < 0.001). B) The average LRR ± SE for each response type. C) The average return time to baseline in days ± SE for each response type. For B) and C), letters over bars denote statistically significant differences among response types based on a Tukey-HSD post-hoc test following a significant one-way ANOVA.



Fig. 3. Storm impacts on the Mission-Copano Bay estuary. A) Time series of Mission River discharge (thick black) and salinity (thin black) and turbidity (dashed) in the receiving estuary.B) Time series of chl a (dashed), CO<sub>2</sub> flux (thick black), and salinity (thin black) in Copano and Mission bays. Blue bars indicate the period of the Hurricane.



Fig. 4. Storm impacts on the Aransas River. A) High frequency time series of dissolved oxygen (grey), temperature (thin black), and river discharge (thick black). B) Synoptic sampling of NO<sub>3</sub><sup>-</sup>, DOC, and orthophosphate from August to December 2017. Data show pattern of elevated nutrients and a temporary elevation in DOC following the storm. Error bars are standard error.
C) Synoptic sampling of conductivity and pH for the aforementioned period. Data show a pattern of dropping conductivity and increasing pH with the precipitation event. Blue bars indicate the period of the storm.



Fig. 5. Storm impacts on San Antonio Bay. A) Synoptic sampling for pH,  $CO_2$  flux, salinity, secchi depth, dissolved oxygen, and temperature from spring 2017 to winter 2018. Data show the increase in  $CO_2$  flux followed by rapid drop, and temporary dips in pH,

salinity, secchi depth, dissolved oxygen, and temperature. Error bars are standard error. B) Synoptic and high frequency data from mid-August to early September 2017. Salinity dropped to nearly zero for days after the storm while bottom dissolved oxygen levels became anoxic 5 days post-Hurricane and remained anoxic for another 8 days. Blue bars indicate the period of the hurricane.



Fig. 6. Mobile fauna responses. A) Natural log of normalized fish and crustacean abundance in each stream reach from September to December 2017. Black line is the mean across sites, error bars are standard error. B) Mean fish and crustacean abundance (CPUE) in seine hauls for select estuaries from July to December of 2017. C) Proportional change in fish abundance in seine hauls from August to September for select estuaries from 2010-2016 (grey bars) and 2017 (green bars). D) Mean fish and crustacean abundance (CPUE) in otter trawls for July to December of 2017.