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2	Two low-lying coastal grassland species differ in
3	mechanistic response to saline flooding stress
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1 Abstract:

Coastal wetlands and low-lying coastal grasslands are increasingly susceptible to salt water 2 intrusion (SWI) influenced by storm surge and sea level rise. Our aim was to understand 3 physiological and growth responses of dominant coastal grassland species, Spartina patens (Ait.) 4 Muhl. (Poaceae) and Fimbristylis castanea (Michx.) Vahl. (Cyperaceae), to increased saline 5 flooding as expected with sea-level rise or saltwater intrusion. We hypothesized that S. patens 6 would exhibit greater resistance to salt stress due to wider distribution. Both species were 7 subjected to saltwater flooding at 0, 5, 10, and 20 ppt (parts per thousand) for 16 weeks. 8 9 Physiological, morphological, and biochemical responses were measured at the end of the experiment. Biomass and tissue chlorides were quantified above and belowground. Physiology 10 was minimally affected by increased saltwater. Relative growth rate and aboveground biomass 11 12 were reduced in both species at 20 ppt. Both species maintained belowground biomass, with increased root and leaf tissue chlorides as salinity increased. The results did not support our 13 hypothesis and found that both species are impacted by saltwater flooding, but S. patens 14 exhibited more signs of stress at 20 ppt. However, both species exhibited tolerance to long term 15 salt stress through different mechanisms. *Spartina patens* excretes salts and allocates resources 16 17 belowground, whereas F. castanea maintains root:shoot and exhibits higher relative water content. Maintaining or allocating biomass belowground with increasing salinity may be 18 important in future sea-level rise and saltwater intrusion scenarios as low-lying coastal systems 19 20 are at higher risk of erosion. Reduction in plant size (aboveground biomass) may impact function of coastal grasslands. 21

3 Additional Index Words

4	Biomass allocation; salt water intrusion; salinity; sea-level rise; Spartina patens; Fimbristylis
5	castanea
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Introduction

2	Salt water intrusion (SWI) is a natural phenomenon influenced by storm surge, relative
3	sea level rise (RSLR), and fluctuations in climate exacerbated by anthropogenic factors (White
4	and Kaplan, 2017). Sea level rise and SWI threaten vulnerable coastal ecosystems at a global
5	scale including mangrove forests, coastal wetlands, and coastal dune systems (Sasmito et al.
6	2016; White and Kaplan 2017). With expected increased RSLR and storm frequency, coastal
7	wetlands and low-lying grasslands are more susceptible to saline flooding due to SWI (Young et
8	al. 2011; Hoggart et al. 2014; White and Kaplan 2017). SWI increases the salinity of soil and
9	groundwater, and during high rainfall events causes saline inundation. In systems close to the
10	water table (<i>i.e.</i> coastal swales) SWI of the freshwater lens can lead to saline inundation.
11	Resilience of coastal plant communities to SWI is influenced by species' physiological and
12	morphological response to salinity stress. When subjected to prolonged saltwater immersion,
13	wetland vegetation diversity is reduced with an increasing dominance of a few salt tolerant
14	species (Grieger et al. 2018). The effects of SWI, RSLR, and salinity stress are well studied in
15	wetland systems (freshwater, saltwater, and forested; Hester Mendelssohn and McKee 2001;
16	Langston Kaplan and Putz 2017; Grieger et al. 2018), however little is known about vegetation
17	response to salinity stress in coastal grassland swales (i.e. low-lying elevation between dunes that
18	are dry more frequently than wetlands (Hoggart et al. 2014). Grassland swales are understudied
19	ecosystems despite their vulnerability to sea-level rise and history of dramatic landscape shifts
20	(March and Smith 2012; Zinnert et al. 2016). These areas experience intermittent flooding and
21	dry periods based on season and precipitation. They are one of the first ecosystems impacted in
22	areas where fronting dunes erode or are breached through overwash events (Young et al. 2011;
23	Brantley et al. 2014). Greenhouse experiments subjecting coastal grassland species to short term

saline waterlogging (8-24hr) found reduced growth and survival with increased salinity
 concentrations (White et al. 2014; Hanley et al. 2017).

3 Salinity stress in plants leads to morphological, physiological, and biochemical responses 4 which have implications for ion regulation and growth. Salinity affects plants through reduced osmotic potential due to high soil salinity and through ion toxicity from NaCl uptake and 5 6 accumulation (Munns and Tester 2008). An early response to salt stress (minutes to days) is 7 stomatal closure due to osmotic stress, which reduces photosynthetic potential and leaf growth (Munns and Tester 2008; Shabala and Munns 2017). Salinity induced ion toxicity caused by 8 9 NaCl uptake causes leaf death that occurs within days to weeks (Munns and Tester 2008), demonstrating the need for longer term studies (>1 month). Osmotic stress affects plant growth 10 and is often referred to as physiological drought (Munns and Tester 2008; Shabala and Munns 11 2017). Plants have adaptive responses to combat salinity stress, and range extensively along the 12 salt tolerance continuum. Plants exhibit three main mechanistic responses to salinity which 13 14 include; exclusion of toxic ions in the shoots (ion exclusion), storage of toxins in tissues (tissue tolerance), or maintenance of stomatal conductance and growth despite physiological drought 15 (osmotic tolerance) (Munns and Tester 2008). Additionally, some plants respond to salinity 16 17 stress through resource allocation from shoot to root, allowing the species greater access to water resources (Hu et al. 2015). Changes in growth allocation due to salinity stress may alter a species 18 19 role in its habitat, as greater belowground biomass could decrease erosion stress by sediment binding while greater aboveground biomass may increase photosynthetic capabilities 20 (Charbonneau et al. 2016; Zhao et al. 2017). Due to ocean proximity, coastal grassland species 21 represent a highly diverse plant community ranging in stress adaptations and salinity tolerances 22 (Marcum 2008). 23

1	The purpose of this study was to gain an understanding of the physiological and growth
2	responses of dominant coastal grassland species, Spartina patens (Ait.) Muhl. (Poaceae) and
3	Fimbristylis castanea (Michx.) Vahl. (Cyperaceae), to increased salinity flooding as expected
4	with sea-level rise or saltwater intrusion. We focused on both above and belowground metrics as
5	both have importance for future modeling with regards to sediment dynamics (<i>i.e.</i> overwash,
6	erosion; Feagin et al. 2015; Charbonneau et al. 2016; Silva et al. 2016). Studying this
7	relationship allows for predicting changes in the response of vegetation and resilience of these
8	systems due to the effects of climate change. Due to the distributions of these macrophytes, and
9	the known adaptations of S. patens (salt excretion), we hypothesized that S. patens would be
10	more tolerant to prolonged saltwater inundation with higher survival and biomass accumulation
11	relative to F. castanea.

Materials and methods

13 Studied species

Spartina patens and Fimbristylis castanea are two dominant and co-occurring grassland 14 macrophytes found in coastal ecosystems along the Atlantic and Gulf coasts of North America 15 including frontline barrier islands (Hosier and Eaton 1980; USDA NRCS 2019). Spartina patens 16 is a C₄ grass distributed throughout the Atlantic coast and Gulf of Mexico where it is often a 17 dominant species in dunes, swales, and wetlands (Hester et al. 2001). It is highly distributed 18 across the barrier island landscape, adapted to a range of coastal habitats (*i.e.* swale grasslands, 19 salt marshes, and sand dunes), and can grow in both saline and fresh waters (Silander and 20 Antonovics 1979; Wu et al. 1998; Hester et al. 2001; Visser and Peterson 2015). *Fimbristylis* 21 22 castanea is a common C₄ sedge found in dense coastal grasslands (McCaffrey and Dueser 1990).

Fimbristylis genus are found in wetland habitats (both saline and freshwater) and can tolerate
high salt stress (Zahoor et al. 2012). In the barrier island landscape, *F. castanea* is commonly
found in low-elevation swales. Both *S. patens* and *F. castanea* are able to cope with a certain
degree of salt stress due to their distributions, with *S. patens* distributed in higher saline flooded
ecosystems (wetlands), however little is known of their response to prolonged (> 1 month) saline
inundation (Pezeshki and DeLaune 1993; Hester et al. 2001).

7 Study site

Hog Island (37° 40' N; 75° 40' W) is one of fourteen barrier islands located along the 8 Eastern Shore Peninsula of Virginia of the Virginia Coast Reserve (VCR). The VCR is a Long-9 Term Ecological Research (LTER) site. The island consists of a wide north end and a slender 10 southern tip. The vegetation distribution among the island is diverse with grass dominated fore-11 dunes and shrub occurrences beyond secondary dunes. Salinity varies on Hog Island with higher 12 levels in coastal and marsh regions (> 30 ppt, parts per thousand) compared to lower levels in 13 shrub and swale regions (< 5 ppt) (Shao et al. 1996). Mean annual temperature is documented to 14 be 15°C with mean annual precipitation between 850 and 1400 mm (Aguilar et al. 2012) 15

The two dominant grassland species, *Spartina patens* and *Fimbristylis castanea* were
collected from Hog Island. Mature plants (~ 35 cm height) were collected from the interior swale
March 2018 (Figure 1). On Hog Island, *S. patens* is found in soil salinity levels between 0 – 8 ppt
while *F. castanea* is found between 0.03 – 3 ppt (Brown and Zinnert, unpublished data). *Spartina patens* is found more abundantly across the island. Plants of similar size rooting mass
and aboveground shoots were transferred into plastic pots with sand collected from Hog Island.
All pots were given equal amounts of Hoagland's nutrient solution (Hoagland and Arnon 1950)

at the beginning and middle of the experiment. The experiment was conducted for 16 weeks in a
greenhouse during the growing season (July – October). Plants were randomized between
treatment groups and control, and allowed to acclimate to growing conditions for one week prior
to treatment. At the beginning of the experiment, plants were cut to 24 cm height to standardize
initial height.

6 Experimental design

Potted plants for the control and treatment groups (n = 5 for each treatment) were placed 7 into larger containers (18 cm diameter) which were filled with water to \sim 7.5 cm below soil level 8 based on experimental design from Hester et al. (2001). Each plant was placed in a separate 9 container for each treatment application (0,5,10, and 20 ppt) which was refreshed bi-weekly. 10 11 Salinity was checked and adjusted every other day using a hand refractometer (300011, SPER scientific, Scottsdale, AZ) to maintain salinity levels. Salinity treatments were made using 12 dilutions of a commercial mixture of salts representative of ocean salinity (Instant Ocean, 13 Aquarium systems, Mentor OH). Major Cations present in the mixture are Na⁺, K⁺, Ca²⁺ and 14 Mg²⁺ (5.1, 0.18, 0.19, and 0.62 ppt respectively) and the major anion is Cl⁻ (8.9 ppt) (Atkinson 15 and Bingman 1997). Salinity treatment range was based on a sublethal salinity level for S. patens 16 of 20 ppt (Hester et al. 2001). The experiment was concluded at the end of the 16th week 17

18 Measurements

19 Physiological measurements of plant responses to salinity included electron transport rate 20 (ETR), stomatal conductance to water vapor (g_s), relative water content (RWC), water use 21 efficiency (WUE) or carboxylation capacity determined from δ^{13} C (Coplen 2011), leaf nitrogen, 22 and δ^{15} N. ETR measures photosynthetic potential, which can be an indicator of plant health.

Measurements were conducted in the 16th week of the experiment during midday conditions 1 (10:00-2.00h). Stomatal conductance measurements were quantified using a steady state 2 diffusion porometer (SC-1, Decagon Devices, Pullman, WA). ETR was measured using a mini-3 PAM fluorometer (Heinz Walz GmbH, Effeltrich, Germany). Stomatal conductance and ETR 4 5 measurements were taken in the greenhouse on multiple randomly selected grass blades, which 6 were still intact on the plants, lined side by side to increase sample area. Relative water content (RWC) was measured using one harvested leaf per plant. Fresh weight was measured 7 immediately following harvest. Leaves were soaked in water for 2 hours and blotted dry. Leaves 8 9 were weighed to obtain turgid weight. Turgid leaves were oven dried at 70 °C for 72 hours. Dried leaves were weighed to obtain dry weight. Relative water content was calculated using the 10 formula: RWC % = [(FW-DW)/(TW-DW)]*100 (Slavík, 1974), where FW = fresh weight, DW 11 = dry weight, and TW = turgid weight. Nutrient and isotope measurements ($\delta^{13}C$, $\delta^{15}N$, %N, 12 %C) were obtained from each plant to analyze nutrient and water use efficiency. Leaves were 13 oven dried at 70 °C for 72 h and ground into a fine powder using a Wiley Mill (Thomas 14 Scientific, Swedesboro NJ, USA). Nutrient and isotope analysis were conducted at the Cornell 15 University Stable Isotope Lab, Ithaca, New York, USA. The ground material was analyzed 16 17 using EA-IRMS (ThermoFischer Scientific).

Growth trait measurements included specific leaf area (SLA), shoot length, and biomass. For SLA, area measurements were made by calculating the projected area of scanned leaves using WinRHIZO software (Regent Instruments Inc., Quebec, Canada). Measured leaf samples were oven dried at 70°C for 72 hours. Dry samples were weighed to obtain dry weight. SLA was calculated as area/dry weight. Shoot length (cm) was measured from sand surface to the tip of the longest leaf during week 16 of the experiment. Shoot length is used as a common growth

metric (Price et al. 2014), but does not necessarily correspond to biomass. Aboveground biomass 1 (AGB) and belowground biomass (BGB) were separated, oven dried at 70 °C for 72 h, and 2 weighed. Biomass is a common growth metric that is a reliable estimate of plant fitness 3 (Younginger et al. 2017). Root to shoot ratio (R:S) was calculated as BGB/AGB. Leaf chlorides 4 5 (LC) and root chlorides (RC) were obtained following biomass measurements. Aboveground and 6 belowground material was oven dried at 70 °C for 72 hours. The samples were ground into a fine powder using a Wiley Mill. Samples of the ground material (~0.5 g) were measured into flasks 7 with 50 mL of deionized (DI) water and placed in a boiling water bath for 2 h. The samples were 8 9 cooled and brought up to volume of 200 mL using deionized water. To each sample, 2 mL of 5M NaNO₃ were added and leaf chloride values were determined using a chloride electrode (model 10 9617b, Orion, Boston, MA). 11

12 Statistical analysis

Survival was analyzed using logistic regression analysis. All growth parameters,
physiology, and chloride measurements were analyzed by two-factor analysis of variance
(ANOVA) with treatment and species as categorical factors. If significant interactions or
differences were found, Tukey's tests were used as post-hoc analysis. Root chlorides were log
transformed to meet assumptions of distribution normality and equal variance. Statistical analysis
was conducted using JMP software (JMP Pro 14, SAS Institute, Cary, North Carolina, USA).

19

Results

There were significant responses to salinity in both *S. patens* and *F. castanea* as seen in physiological and morphological changes. However, the two species exhibited slightly different response mechanisms to salinity stress. There was not a significant difference in survival by salinity treatment for *S. patens* (chi=2.94, p= 0.4) or *F. castanea* (chi=2.94, p= 0.4). Percent
 survival was 100% for salinity treatments of 0, 5, and 10 ppt, and 80% for 20 ppt for both
 species.

4 Physiological response

There was a significant species x treatment interaction for ETR (F=7.9, p<0.001). 5 6 ETR was significantly different between species (F=13.8, p<0.001) and treatment (F=21, p<0.0001). Fimbristylis castanea increased ETR at 10 ppt and reduced ETR at 20 ppt with a 29% 7 reduction relative to control plants. Spartina patens reduced ETR significantly at 20 ppt with a 8 9 62% reduction relative to control plants (Figure 2). There was not a significant species x treatment interaction for stomatal conductance (F=1.4, p=0.200). Stomatal conductance was 10 11 significantly different between species (F=11.4, p=0.002), but not by salinity treatment (F= 1.8, p=0.200) (Table 1). Stomatal conductance was maintained throughout treatment levels, and 12 therefore not a factor in reduced ETR. There was not a significant species x treatment interaction 13 for SLA (F=1.9, p=0.100). SLA was significantly different between species (F=67.0, p<0.001), 14 but not by salinity treatment (F=0.3, p=0.800) (Table 1). Spartina patens exhibited a higher SLA 15 compared to F. castanea. There was not a significant species x treatment interaction for RWC 16 (F=1.7, p=0.200). RWC was significantly different between species (F=6.6, p=0.010), but not by 17 salinity treatment (F=1.0, p=0.400) (Table 1). Both species exhibited the ability to maintain 18 19 RWC with increased salinity indicating potential succulence that may reduce salt stress. There was a significant species x treatment interaction for δ^{13} C (WUE) (F=3.3, p=0.030). δ^{13} C was 20 significantly different between species (F=67483.5, p<0.001), and salinity treatment (F=16.1, 21 p<0.001) (Figure 3). There was not a significant species x treatment interaction for $\delta^{15}N$ (F=0.1, 22 p=0.920). There was no significant difference in %N between species (F=4.6, p=0.040) or 23

treatment (F=7.7, p<0.001). *Fimbristylis castanea* exhibited higher δ¹⁵N than *S. patens* (Figure
 3). Both species exhibited an increase in δ¹⁵N at 5 ppt (Figure 3).

3

4 Growth trait response

5	There was a significant species x treatment interaction for shoot length (F=3.1, p=0.040)					
6	Shoot length was significantly different between species (F=13.0, p=0.001), and salinity					
7	treatment (F=22.8, p<0.001). Fimbristylis castanea reduced shoot length at 5, 10, and 20 ppt,					
8	whereas S. patens maintained until 20 ppt (Figure 4). There was not a significant species x					
9	treatment interaction in aboveground biomass (F=2.0, p=0.100). Aboveground biomass was not					
10	significantly different between species (F=0.1, p=0.700), but was significantly different between					
11	salinity treatment (F=5.3, p=0.005). Both species exhibited reductions at 20 ppt (Figure 5). There					
12	was not a significant species x treatment interaction in belowground biomass (F=2.9, p=0.050).					
13	Belowground biomass was significantly different between species (F=40.2, p<0.001), but was					
14	not significantly different between salinity treatment (F=0.3, p=0.800). Fimbristylis castanea					
15	exhibited higher belowground biomass compared to S. patens (Figure 5). There was a significant					
16	species x treatment interaction for root:shoot (F=3.3, p=0.030). Root:shoot was significantly					
17	different between species (F=4.8, p=0.040), and salinity treatment (F=3.9, p=0.020) Spartina					
18	patens increased root:shoot ratio at 5, maintained at 10 ppt, and increased at 20 ppt (Figure 5).					
19	Biochemical response					
20	There was not a significant species x treatment interaction for leaf tissue chlorides					

21 (F=0.6, p=0.600). Leaf tissue chlorides were significantly different between species (F=60.9,

22 p<0.001) and treatment (F=43.1, p<0.001). Leaf tissue chlorides linearly increased at 5, 10, and

20 ppt for both species, but *F. castanea* had higher leaf chlorides than *S. patens* (Figure 6). There
was not a significant species x treatment interaction for root tissue chlorides (F=1.7, p=0.200).
Root tissue chlorides were not significantly different between species (F=0.8, p=0.400), but were
significantly different between salinity treatment (F=67.3, p<0.001). Salinity affected root tissue
chlorides at 5, 10, and 20 ppt for both species (Figure 6). There were visible signs of salt
excretion in *S. patens* at 5, 10, and 20 ppt (Figure 7). *Fimbristylis castanea* did not show signs of
salt excretion (Figure 7).

8

Discussion

Many coastal species are well adapted to salinity (e.g. Salicornia dolichostachya, 9 Panicum heimitomon, and Spartina alterniflora; (Hester et al. 2001; Katschnig et al. 2013) and 10 11 thus, may be able to withstand long-term effects of saline flooding. Our results suggest that both S. patens and F. castanea maintain stomatal conductance and RWC with increased salinity; 12 however, there are significant changes in growth parameters within these species at 20 ppt. Our 13 results suggest that S. patens and F. castanea show partial tolerance of long-term salt stress, but 14 through different mechanisms. Spartina patens excretes salts and allocates resources to maintain 15 belowground biomass, while F. castanea maintains root:shoot and exhibits signs of leaf 16 succulence and higher RWC. 17

Many species exhibit reductions in ETR and stomatal conductance with salt stress including coastal species (Naumann et al. 2008; Geissler et al. 2015; Tounekti et al. 2018), but some saltwater marsh species exhibit high tolerance to salinity stress with levels up to 30 ppt including *Salicornia dolichostachya*, and *Spartina alterniflora* (Hester et al. 2001; Katschnig et al. 2013). Within coastal systems there is variability in salinity tolerance with decreasing

tolerance relative to distance from ocean (Oosting and Billings 1942; Young et al. 2011).
However, species that occur further inland (grassland swale) can often withstand short term
saline flooding (Tolliver et al. 1997; Naumann et al. 2008). In our study, both species were able
to maintain stomatal conductance and operational ETR at all salinity levels, and therefore exhibit
mechanisms to combat physiological salinity stress through osmotic tolerance. δ¹³C, a proxy for
WUE or carboxylation capacity (Coplen 2011), coupled with stomatal conductance and ETR
results suggest that *F. castanea* has higher carboxylation efficiency than *S. patens*.

8 Plants are known to tolerate stress by three main mechanistic responses including; ion 9 exclusion, tissue tolerance, or osmotic tolerance (Munns and Tester 2008). Spartina patens exhibits ion exclusion through salt excretion examined in our experiment and also noted in other 10 experiments (Hester et al. 2001). This coincided with lower tissue chlorides compared to F. 11 castanea across treatments. Both species exhibited a linear increase in leaf tissue chloride 12 concentrations across treatments, but an exponential increase in root tissue chlorides. Between 10 13 and 20 ppt, F. castanea more than doubled in root chlorides, while S. Patens increased more than 14 four times. Belowground appears to be a better place to store chlorides, indicating a tissue 15 tolerance mechanism. 16

17 Despite increases in leaf chlorides, the long-term construction of tissues was not affected 18 at any salinity level as observed in SLA. This is notable as a negative association between SLA 19 and increasing salinity is observed in many coastal and non-coastal species (Ishikawa and Kachi 2000; Rahimi and Biglarifard 2011; Tedeschi et al. 2017). Shoot length decreased in *F. castanea* 21 at 20 ppt, while *S. patens* maintained shoot length until 20 ppt. Relative growth rate and SLA 22 were unaffected by %N availability. As salinity increases, shifts in nitrogen occur, based on δ^{15} N 23 values, which may result from microbes being affected by salinity (Franklin et al. 2017).

1	Allocation of biomass was altered by salinity. Fimbristylis castanea maintained
2	root:shoot across treatments while S. patens increased root:shoot at higher salinity levels. Over
3	all treatments, F. castanea had higher belowground biomass. Spartina patens may also reduce
4	toxicity of chlorides to the aboveground through belowground storage. Allocation of resources
5	from shoots to roots also has implications in coastal areas experiencing erosion. Higher
6	belowground biomass may be important in stabilizing sediments of eroding coastlines
7	(Charbonneau et al. 2016) by enhancing the aggregation and strength of soil in coastal systems;
8	therefore, increasing the resistance to erosion (Sigren et al. 2014). Biomass allocation strategies
9	differ interspecifically (Visser and Peterson 2015; Charbonneau et al. 2016) which we observed
10	in our study. As salinity increased, S. patens maintained belowground biomass while reducing
11	aboveground. Similar responses have been found in coastal grasses in response to sand burial
12	(Brown and Zinnert 2018). With increases in RSLR and SWI, species maintaining resources
13	belowground may increase resistance against erosion and coastline shifts further inland.
14	Our hypothesis that S. patens would be more tolerant to prolonged salinity stress
15	compared to the less widely distributed F. castanea was not fully supported. Our findings
16	indicate similar saline flooding tolerance of both species to low levels of salinity with a reduction
17	in growth at high salinity concentrations (i.e. 20 ppt). These species are able to partially tolerate
18	salinity, but long-term saline flooding leads to a greater reduction in ETR and reallocation of
19	resources to maintain belowground biomass in S. patens. This reallocation of resources may be a
20	mechanism to reduce toxicity of aboveground leaf tissue chlorides. Spartina patens further
21	reduces chloride concentration by excreting salts. Fimbristylis castanea maintains above and
22	belowground biomass, but reduces shoot length at high salinity concentrations. It exhibits
23	succulence through high RWC which is maintained across salinity treatments. As belowground

resources are maintained, the ability to acquire nitrogen is not affected; however microbial
 associations are altered.

3 Coastal communities are subjected to RSLR, large storm events, and SWI, leading to 4 changes in species composition and community structure (Zinnert et al. 2016; Grieger et al. 2018). While some species may be lost with increasing salinity, tolerant species such as the two 5 6 studied here will still be impacted. Reduction in plant size may affect competition with more 7 salt-tolerant species, potentially altering coastal grasslands dominated by S. patens and F. castanea. Growth changes observed in our experiment have implications on coastal grassland 8 9 communities such as less aboveground sediment trapping capabilities and changes to erosion resistance. Mature plants were measured in this experiment, and younger ontogenetic stages may 10 not respond in the same way (Lum and Martin 2019). Therefore, these findings may not 11 represent newly established swales or communities with mixed life stages, but are more 12 applicable to established swales. Modeling of the dune environment is an active area of research 13 14 (e.g., Vinent and Moore 2015; Keijsers et al. 2016; Cohn et al. 2019); however, modeling areas beyond the dune that account for ecological processes is sparse (Zinnert et al. 2016). With 15 16 projected increases in storms and sea-level rise, understanding and modeling community 17 response to saline flooding in coastal systems becomes more important (e.g., Rastetter 1991; Hanley et al. 2017). 18

19 Conclusions

Salt water intrusion increasingly affects low-lying coastal grasslands through
 saline flooding due to RSLR and storm surge. Our findings indicate that *S. patens* and *F. castanea* (two coastal swale species) are able to partially tolerate long term saline

flooding (16 weeks). The two species appear to have different mechanisms for tolerating 1 2 salt stress. Spartina patens excretes salts and allocates resources to maintain belowground biomass, while F. castanea maintains root:shoot and exhibits signs of leaf succulence and 3 higher RWC. Growth is reduced for both species with increased salinity, which may have 4 implications for coastal grassland communities (i.e. shifts in dominant species, reduced 5 erosion control and sediment trapping). Future research into the response of other 6 dominant grassland species to long term saline flooding is necessary to determine the 7 impact of SWI on coastal grasslands. 8 Acknowledgments: 9 This work was supported by the National Science Foundation, Department of 10

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Table 1: Mean ± SE values for stomatal conductance (gs), specific leaf area (SLA), and relative
water content (RWC) and percent nitrogen (% N) for *Spartina patens* and *Fimbristylis castanea*at 0, 5, 10, and 20 ppt. There were no interaction or salinity treatment effects. Species differences
are notated with *.

Species	Response variable	0 ppt	5 ppt	10 ppt	20 ppt
	gs*	124 ± 10	115 ± 4	123 ± 11	101 ± 3
S natons	SLA*	93 ± 11	106 ± 6	104 ± 5	114 ± 12
S. pateris	RWC	93 ± 3	90 ± 2	95 ± 3	87 ± 5
	% N	0.7 ± 0.1	0.7 ± 0.0	0.6 ± 0.0	0.8 ± 0.1
	gs*	154 ± 12	125 ± 4	131 ± 12	139 ± 5
F. castanea	SLA*	69 ± 4	65 ± 5	56 ± 3	56 ± 6
	RWC	93 ± 1	99 ± 2	96 ± 1	96 ± 0.4
	% N	0.7 ± 0.0	0.6 ± 0.0	0.7 ± 0.0	0.6 ± 0.0

Figure 1: Map marking the location of Hog Island (A), an aerial photograph of southern Hog
 Island with the swale and dunes labeled (B), and an image of the swale ecosystem at Hog Island
 (C).

Figure 2: *Spartina patens* (black) and *Fimbristylis castanea* (grey) show a reduction in electron
transport rate (ETR) at 20 ppt salinity. Lower case letters indicate significant differences in
salinity treatment, and capital letters indicate significant differences in interaction. Differences
were determined from Tukey tests (α = 0.05).

Figure 3: *Spartina patens* (black) maintains δ^{13} C with increased salinity. *Fimbristylis castanea* (grey) reduces δ^{13} C compared to 0 ppt at 10 ppt and is therefore reduces WUE with increased salinity. *Spartina patens* (black) and *Fimbristylis castanea* (grey) increase δ^{15} N at 5ppt of salinity. Lower case letters indicate significant differences in salinity treatment, and capital letters indicate significant differences in interaction. Differences were determined from Tukey tests ($\alpha = 0.05$).

Figure 4. *Spartina patens* (black) and *Fimbristylis castanea* (grey) show a reduction in shoot length at 20 ppt salinity. Lower case letters indicate significant differences in salinity treatment, and capital letters indicate significant differences in interaction. Differences were determined from Tukey tests ($\alpha = 0.05$).

Figure 5: *Spartina patens* (black) and *Fimbristylis castanea* (grey) show a reduction in
aboveground biomass with increasing salinity, but no change in belowground biomass. There is a
significant difference between species in belowground biomass. The root:shoot ratio indicates an
increase in *S. patens* while no change in *F. castanea*. Lower case letters indicate significant

1	differences in salinity treatment, and capital letters indicate significant differences in interaction.
2	Differences were determined from Tukey tests ($\alpha = 0.05$).
3	Figure 6: There is a significant increase in leaf chlorides and root chlorides with increasing
4	salinity for both Spartina patens (black) and Fimbristylis castanea (grey). There is a difference
5	between species in leaf and root chlorides. Lower case letters indicate significant differences in
6	salinity treatment. Differences were determined from Tukey tests ($\alpha = 0.05$).
7	Figure 7: Image of <i>Spartina patens</i> and lack of salt excretion at 0 ppt (A), but seen at 10 ppt (B).
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- 2 Figure 1.























2 Figure 6.



Figure 7.