

Multi-Scaled Grassland-Woody Plant Dynamics in the Heterogeneous Marl Prairies of the Southern Everglades

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ABSTRACT

The Everglades freshwater marl prairie is a dynamic and spatially heterogeneous landscape, containing thousands of tree islands nested within a marsh matrix. Spatial processes underlie population and community dynamics across the mosaic, especially the balance between woody and graminoid components, and landscape patterns reßect interactions among multiple biotic and abiotic drivers. To better understand these complex, multi-scaled relationships we employed a three-tiered hierarchical design to investigate the effects of seed source, hydrology, and more indirectly bre on the establishment of new woody recruits in the marsh, and to assess current tree island patterning across the landscape. Our analyses were conducted at the ground level at two scales, which we term the micro- and meso-scapes, and results were related to remotely detected tree island distributions assessed in the broader landscape, that is, the macro-scape. Seed source and hydrologic effects on recruitment in the micro- and meso-scapes were analyzed via

logistic regression, and spatial aggregation in the macro-scape was evaluated using a grid-based univariate O-ring function. Results varied among regions and scales but several general trends were observed. The patterning of adult populations was the strongest driver of recruitment in the micro-and meso-scape prairies, with recruits frequently aggregating around adults or tree islands. However in the macro-scape biologically associated (second order) aggregation was rare, suggesting that emergent woody patches are heavily controlled by underlying physical and environmental factors such as topography, hydrology, and Þre.

Key words: woody plant invasion; recruitment; dispersal; hierarchical patch dynamics; univariate grid-based O-ring function; spatial patterning; community and landscape structure; shifting boundaries; tree islands; Cape Sable seaside sparrow habitat.

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INTRODUCTION

A century of drainage, compartmentalization, and Pre suppression has irrevocably reduced the Everglades to half its original size, and despite recent restoration efforts, more than 2500 km of canals, levees, and water control structures threaten to slowly deteriorate what remains. Preservation of extant communities may require a restructuring of hydrologic systems and a revision of Pre management practices to better emulate natural conditions. However, natural environmental conditions are not always clearly debned and the historical distribution of landscape patches is not well documented. Although we know that both historical and present day landscapes contained thousands of tree islands embedded within marsh/grassland matrices, we do not clearly understand the extent to which management has altered the scale and pattern of this woody-graminoid mosaic, nor do we understand the sensitivity of patch dynamics to variation in management scenarios. Researchers suspect that the freshwater marl prairie landscapes comprising the outer regions of Everglades National Park (ENP) may be vulnerable to woody plant expansion (Jenkins and others 2003a, b) due to alterations in natural disturbance mechanisms (namely, ßood and Þre management). However, potential encroachment patterns and processes have not been examined in detail.

Spatial processes play an integral role in the dynamic balance between woody and graminoid cover in the Everglades marl prairie landscape, and these processes vary across scales. Although locallevel studies provide valuable information on the distribution and abundance of interacting species within patches, neighborhood processes also inevitably inßuence within-patch conditions (Levin 1992; Wu and Levin 1994; Chase and Leibold 2002). Because increases in woody plant cover may be obscured or exaggerated in studies that focus specibcally on patch-scale phenomena (With and King 2001), multi-scaled spatial detail is indispensable for conservation biologists who wish to effectively manage the unique habitat requirements of multiple interacting species. Preservation of habitat complexity within the Everglades heterogeneous marl prairies for example, necessarily requires a better understanding of the spatially explicit effects of seed source, hydrology and bre on woody plant recruitment and encroachment.

Marl prairies contain species-rich grasslands and tree islands that vary in size, density, and composition. The resident ßora and fauna depend upon the persistence and connectivity of both grassland and woody community types, maintained through intertwining biotic and physical feedback mechanisms. The endemic Cape Sable seaside sparrow (CSSS;Ammodramus maritimus mirabilis) in particular requires extensive and well-connected prairie habitat patches (Nott and others 1998; Browder and Ogden 2000; Pimm and others 2002; Jenkins and others 2003a, b), whereas several wading bird species such as the woody stork (Mycteria

Americana) and white ibis (Eudocimus albus) utilize prairie regions for foraging, but require intermittent woody patches for nesting and roosting (Hoffman and others 1994). Cyclical shifts in the spatial patterning of habitat types are driven to various degrees by environmental dynamics and the physiological adaptations of local biota, such as ßood and/or Pre tolerance, life history traits, and phenological patterns. Woody plant establishment in the marsh may be the leading spatial process inßuencing habitat shifts across the landscape. The distribution of new woody recruits provides a template for successive processes that promote or inhibit woody patch nucleation and persistence (Nathan and Muller-Landau 2000).

Nucleation around woody stems involves multiple positive feedback mechanisms, which are limited by hydrology and disturbance. Tree island development frequently begins with soil accretion around newly established individuals or surface anomalies. As woody plants colonize these topographic rises, concomitant increases in peat production further alleviate Gooding stress in support of continued patch expansion (Rietkerk and others 2004). In addition to altering the local topography, established individuals serve as magnets for seed-dispersing animals, which results in the development of mixed species assemblages. These assemblages exhibit higher transpiration rates than surrounding marshes, which increase the concentration of nutrients below their developing canopies. Transpirative nutrient accumulation and increased faunal input also contribute to the competitive advantage of woody plants over grasses (McCarthy and others 1993; Wetzel and others 2005; Ross and others 2006a; Givnish and others 2008; Hanan and Ross 2010). Newly developed tree islands promote woody plant expansion at landscape scales through seed production and physiochemical interactions with neighboring communities. Meanwhile, disturbance in the form of periodic Pre serves to slow and set back tree island establishment and nucleation (Lockwood and others 2003).

The patterning of habitat patches within a land-scape affects the ultimate size and distribution of emerging patches (Wu and Levin 1994). Therefore, to understand habitat change, it is necessary to assess both community-level processes such as those affecting recruitment, and landscape patterns. The Hierarchical Patch Dynamics Paradigm (HPDP) combines patch dynamics perspective (Watt 1947) and hierarchy theory to structure knowledge on multi-scaled heterogeneous systems (Wu and Loucks 1995). Within the HPDP framework, we integrated ground-level and remote

sensing data to investigate the effects of seed source, hydrology and more indirectly, Pre on the recruitment of new woody stems in the marsh. We also mapped and analyzed the spatial patterning of existing forested area across the study region to test the following hypotheses:

- (1) Source effects (i.e., size and proximity of adult trees and tree islands) are a strong determinant of recruitment in the prairie. SpeciPcally, establishment probability increases with proximity to conspeciPc adults and/or tree islands due to gradients in dispersal and/or habitat suitability.
- (2) Hydrology impacts establishment across scales and differing hydrologic scenarios favoring the establishment of some species over others at each location.
- (3) Tree islands are aggregated at multiple scales, due to biological and biogeochemical feedbacks (i.e., biogenesis).

Our ground-level analyses were conducted at two scales and levels of detail, which we term the microscape (i.e., single tree islands and their immediate surrounding environments) and the meso-scape (i.e., intermediately-sized prairie regions that include multiple tree islands). We also related our ground-level observations to remotely detected woody plant distributions assessed at the macroscape (i.e., large landscapes that include a heterogeneous mosaic of high- and low-density patches of tree islands).

METHODS

Study Site

Our study was conducted within the short-hydroperiod marl prairies located along the edges of the Shark and Taylor Sloughs in Everglades National Park. Sampling took place between 2006 and 2008 in six regions designated as subpopulations for the endangered Cape Sable seaside sparrow (subpopulations AĐF; Figure 1). Soils range from calcitic marls classibed as ßuvaquents in the open marsh, to histosols found within tree islands and/or other densely vegetated patches. Marl prairies exhibit characteristically high graminoid diversity maintained by relatively short hydroperiods (inundated for approximately 3D7 months per year). In this part of the Everglades, tree islands typically form in conjunction with limestone bedrock highs, which reduce Booding stress that might otherwise inhibit woody growth.

Hydroperiods in the Everglades are inßuenced by climatic drivers, and modibed by operations of the

South Florida Water Management DistrictÖs (SFWMD) network of canals, levees, and water control structures. The primary controls of water redistribution to the study area are channeled through the S12 structures, located along Tamiami Trail, north of Shark Slough, and the S332 detention basins, located along the L31W and L31N canals (Figure 1). The South Florida wet season typically extends from May through early October, whereas the dry season occurs from the end of October through April. Mean annual precipitation ranges from 120 to 160 cm, and mean daily temperatures range from 17 to 25°C (Obeysekera and others 1999). A record-setting drought between 2006 and 2008 impacted hydroperiods across the southern Everglades (Meeker 2008).

Extensive burns (consuming more than 93 km² of vegetation) occur approximately every 10Đ15 years in the larger Everglades ecosystem, typically in late spring when conditions are still relatively dry (Gunderson and Snyder 1994). Smaller Þres that occur more frequently tend to spark during wet season thunderstorms typical of June and July. The natural Pre return interval is complicated by Pre management practices, which both suppress large natural Þres and ignite controlled burns to reduce fuel loads, maintain early successional stages, and inhibit the proliferation of invasive species (Lockwood and others 2003; La Puma and others 2007). The role of Pre in the maintenance of species composition and structure within the marl prairie landscape is not fully understood.

Sampling Design

Our study was designed to analyze the woody plant encroachment process broadly across the marl prairie landscape and more intensively at specibc invasion fronts within each CSSS subpopulation. Six study areas were established, one within each subpopulation (Figure 1), and sampling followed a hierarchical design (Figure 2). One macro-scape (i.e., a 3.2 km \times 3.2 km cell comprising a single South Florida Water Management Model grid cell) was selected at random within each subpopulation. and then divided into nine 1 km \times 1 km plots (leaving a buffer of 100 m on all sides). From these, a single 1-km² meso-scape was randomly selected in each subpopulation (Figure 1). To examine the demography of individual woody species in the vicinity of tree islands, one 100 m \times 100 m (1 ha) micro-scape was centered on a randomly selected tree island within each meso-scape (Figure 1). Each micro-scape was partitioned into

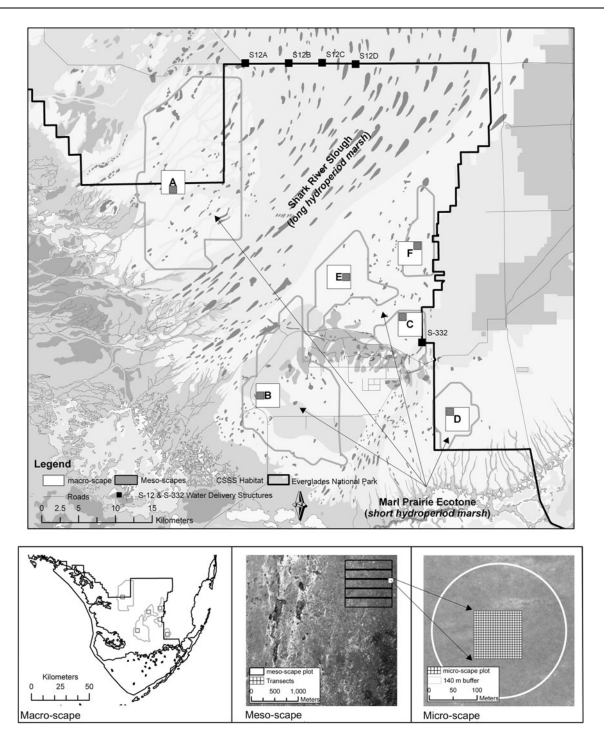


Figure 1. Location of the six macro-scape plots within the habitat of the Cape Sable seaside sparrow (CSSS) in Everglades National Park, FL, and sampling design for meso- and micro-scapes.

100 10 m x 10 m subplots and those subplots were each further partitioned into four 5 m \times 5 m grid cells (400 cells total per micro-scape; Figure 1).

In each micro-scape, all adult (height > 2 m) trees and recruits (height < 2 m) present outside the central tree island were identibed to species, mapped to nearest 0.5 m, tagged, and measured by

height. Trees within the central tree islands were mapped and measured in similar fashion, but recruits were not. Elevations and soil depths were determined at the corners of each $10~\text{m} \times 10~\text{m}$ subplot, by auto-level surveys from benchmarks of known elevation. Additional elevations were determined at random points within the central

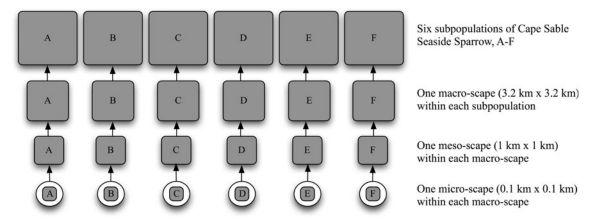


Figure 2. Schematic of studyÕs nested scale hierarchy within the Everglades marl prairie landscape.

tree islands, and at the base of each woody recruit in the marsh. Elevation surfaces were created via ordinary kriging in ESRI [®] ArcMapTM 9.3 using a spherical model, and elevations were extracted for each of the 400 5 m × 5 m cells within the 1-ha plot, based on the center of each cellÕs geometric axis. Radial distances between cell centroids and each adult tree were calculated and subsequently used to relate size and proximity of potential seed sources to the distribution of recruits outside the tree island. To identify additional potential seed sources, all tree islands and individual woody plants located within a 140-m buffer of each plotÕs centroid (Figure 1) were also identiÞed, mapped, and measured.

Four 1-km long, EĐW oriented strip transects were established across each meso-scape, equally spaced at 200-m intervals (Figure 1). Individual woody plants present within 10 m of the transect center were identibed to species, mapped, and measured for height. GPS coordinates were recorded every 100 m along the transect. For analytical purposes, transects were divided into ten 90-m long segments (with 20-m unsampled spaces between segments to prevent overlap resulting from compass error), and each segment was divided into two rows of 10 m \times 10 m cells (one on either side of the transect; 18 total per segment). Presence/absence and counts of woody plants within each 10 m \times 10 m cell were determined. Radial distances between cell centroids and remotely digitized tree islands within 200 m of each cell were calculated and subsequently used to relate size and proximity of potential seed sources to mapped individuals found along each transect. Also, marsh vegetation community types were visually recorded every 20 m along each transect using designations outlined by Ross and others (2006b).

All landscape features within the six macroscapes were identiPed and mapped by soft-copy remote sensing techniques using a set of 2004 NAPP color infrared aerial photographs with a 1-m spatial resolution (1 \times 1 m pixel). Because of the quality and spatial resolution of the imagery, a 5 m² minimum mapping unit (mmu) was considered to be the lowest resolution by which landscape features could be properly discerned and accurately classiPed. Features larger than the mmu were digitized on screen, using ESR ArcMap 9.3. Landscape features that extended beyond the perimeter of the macro-scape were truncated at the macro-scape boundary.

Data Analysis

To analyze source and hydrologic effects on the establishment of woody species within each microscape, Species Source Factors (SSFs) and hydroperiods were calculated for all plot cells, and subsequently evaluated using stepwise logistic regressions. SSFs were calculated as follows:

$$SSF_{ij} = \sum \left(\frac{h_j}{r_{ii}}\right) \tag{1}$$

where h represents adult tree heights for speciesj, and r_{ij} is the radial distance from the centroid of cell i to potential source trees of speciesj within the 140-m buffer. Adult tree heights were normalized by calculating their percent of the maximum height for each species. Assumptions underlying SSF include (1) seed outputs of individual trees are proportional to their size (Thomas 1996; Nathan and Muller-Landau 2000), and (2) because the number of cells at radial distance r from a source tree is proportional to $2\pi r$, the probability for recruitment within a given cell is proportional to 1/r.

Hydroperiods (mean days per year ßooded) were calculated for each cell using elevation estimates, coupled with Everglades Depth Estimation Network (EDEN 2008) daily water stage records for the 6-year period 2000Đ2006.

Using presence/absence of recruits within plot cells (outside tree islands), the effects of SSF and hydroperiod were evaluated by stepwise logistic regression using the following model:

$$P(m) = \frac{1}{1 + e^{-(\beta_0 + \beta_1 SSF_{ij} + \beta_2 Hydroperiod_i)}}$$
(2)

where P(m) is the probability of species j establishment within cell i. In our stepwise logistic regression, variables were selected in the order of their contribution to the model. If a variable did not contribute signipcantly, it was removed and a reduced model was then run. Logistic regressions were carried out for all species present in at least two cells in 2007 that had at least one potential parent tree within the 140-m buffer. To avoid potential confounding of hydroperiod with SSF we (1) omitted cells that were completely or partially within tree island boundaries and (2) tested for collinearity between hydroperiod and SSF.

Species-specibc observations of saplings and adults were made along transects, but individual regressions were not possible in the meso-scape analysis because the species compositions of all tree islands within the 1 km ² were not known. Likewise, the precise hydrology associated with each stem could not be estimated, due to the lack of a detailed and comprehensive topography for the region. Therefore, to analyze source and hydrologic effects on the establishment of woody plants within each meso-scape, we calculated a non-specibc Woody Plant Source Factor (WPSF) and a vegetation-based Prairie Wetness Index (PWI) for each transect cell. WPSFs were calculated as follows:

$$WPSF_{ij} = \sum \left(\frac{a_j}{r_{ij}}\right) \tag{3}$$

where WPSF_{ij} is the Woody Plant Source Factor for cell i, and j represents tree islands within 100 m of cell i. Assumptions underlying WPSF are similar to those for SSF, namely that (1) seed outputs of tree islands are proportional to their area, and (2) the probability for recruitment within a given transect cell is proportional to 1/ r, where r is the distance between the source tree islandÕs centroid and the centroid of each cell.

PWI was calculated from vegetation community observations made at 20-m intervals along each transect. For each herbaceous community

observed, a relativized wetness index ranging from 1 to 10 was derived from vegetation-inferred hydroperiods outlined in Ross and others (2006b). Communities at the lower end of the scale were less Bood-tolerant than communities at the upper end of the scale; for example, the PWI of Muhlenbergia Wet Prairie and Eleocharis-Rhynchospora Marsh were 1 and 10, respectively.

Using presence/absence of individual woody recruits and/or adults within transect cells (outside tree islands), the effects of WPSF and PWI were evaluated by stepwise logistic regression using the following model:

$$P(m) = \frac{1}{1 + e^{-(\beta_0 + \beta_3 WPSF_{ij} + \beta_4 PWI_i)}}$$
(4)

where P(m) is the probability of species j establishment within cell i. Logistic regressions were carried out for all species present in at least two cells along the transects. As with the micro-scape, we avoided potential confounding of PWI with WPSF by (1) omitting cells that were completely or partially inside tree islands and (2) testing for collinearity between PWI and WPSF. All logistic regressions were run in SPSS version 14.0 (SPSS Inc.) and plotted using Statistica (Statsoft Inc.).

The spatial patterning of tree islands within each of the six macro-scapes was analyzed using an O-ring statistic derived in Programita (Wiegand and Moloney 2004). The O-ring statistic is a second-order probability density function, similar to the more commonly used RipleyÕs K; however, the O-ring statistic uses annuli instead of circles, thus identifying the levels of aggregation or dispersion that occur at speciPc distance classes (Wiegand and Moloney 2004). In this study, second-order aggregation may be indicative of tree island biogenesis via positive feedback mechanisms.

Because tree island dimensions varied signib-cantly across our macro-scapes, non-dimensional point patterns, which are frequently used in spatial analysis, were not suitable to represent their distribution and inßuence. Instead we converted our vector-based maps to ASCII grid-Ples (cell size = 5 m), which allowed us to randomize entire islands rather than just object centroids, thereby conserving their dimensional properties. Features that were not tree islands (for example, pine forest, roads, and so on) and tree islands that were truncated by the macro-scape boundary were masked from the analyses.

Complete spatial randomness (CSR) is the simplest and most commonly used null model in spatial pattern analyses (Wiegand and others 2006b).

CSR assumes that objects within a given study area follow a Poisson distribution and as such, the intensity λ of these objects is constant and no interactions occur between them. However, ecological systems rarely meet this assumption because observed spatial trends can result from both broadscale Þrst-order phenomena, such as hydrologic, topographic, and edaphic gradients, or from smaller-scale second-order processes, such as dispersal and establishment. The simplest alternative to CSR is the heterogeneous Poisson null model (Wiegand and Moloney 2004). Like CSR, the heterogeneous Poisson null model assumes that the occurrence of any object is independent of any other object within the landscape; however, the distribution of these objects follows an intensity function $\lambda(x, y)$ which varies by location (x, y; Wiegand and Moloney 2004). Because the heterogeneous Poisson null model accounts for large-scale (Prst-order) aggregation, it is more suitable for identifying second-order patterns in non-homogeneous landscapes (Wiegand and Moloney 2004).

We tested each macro-scape for homogeneity using the two-step grid-based approach outlined by Wiegand and Moloney (2004). Step 1 utilized a large moving window, R set to 150 (i.e., 750 m; one-quarter the width of each macro-scape) to remove all sizeable gaps or regions with extremely low point densities within each macro-scape. Step 2 calculated a smaller moving window, R_{min} that could discriminate between gaps and possible empty plots to further rebne each study region. R_{min} values were 10 (i.e., 50 m) for macro-scapes A, B, C, and F, 12 (i.e., 60 m) for macro-scape D and 15 (i.e., 75 m) for macro-scape E. The heterogeneous Poisson null model then randomized tree islands locally within their given neighborhood R_{min} in each macro-scape. In other words, the heterogeneous Poisson null model applied CSR to newly delineated homogenous sub-regions, thereby conserving large-scale (Prst order) patterns.

A Monte Carlo approach was used to construct conbdence limits for our null model. Ninety-nine Monte Carlo simulations were used to determine the general broad-scale patterning of tree islands within each macro-scape at a maximum distance of 750 m. This distance was approximately one-quarter the width of each macro-scape and extended well beyond the scale at which the observed spatial pattern of tree islands deviated from our null model. Once general patterns were established, we re-ran the analyses using 299 Monte Carlo simulations at a maximum distance of 325 m in each macro-scape. We used the 5th lowest and highest values of the simulation envelope to

determine whether the observed patterns deviated from our null model. Because Type I errors can occur when the observed values fall particularly close to the simulation envelopes, a Goodness of Fit test (Loosmore and Ford 2006; Wiegand and others 2007) was also applied.

Lastly, Þre histories for all three landscape scales (macro, meso, and micro) were constructed from a geodatabase populated with Everglades National Park 1980 to 2008 Þre records. This spatial dataset allowed for the identiÞcation and extraction of all Þres that occurred within the boundaries of the study areas during the last 29 years. For each plot at each scale, a Þre recurrence interval (i.e., Þre frequency) was calculated using the following equation:

Fire recurrence =
$$\frac{1}{A_b/Y/A_t}$$
 (5)

where A_b is the sum total area of all burns, Y is the number of years being analyzed (29), and A_t is the total plot area (1036, 100, and 1 ha for the macro, meso, and micro-scapes, respectively).

Because of the error associated with Pre boundary locations it was not practical to apply Pre recurrence effects to our spatially explicit logistic regressions. However, general trends were derived for each plot at each scale and discussed within the context of our other analyses.

RESULTS

Thirty-Pve woody plant species were identiPed in varying densities across the micro- and meso-scapes (Tables 1, 2). Micro and meso-scapes also varied in their hydrology (Table 3), creating suitable conditions for the habitat preferences and dispersal mechanisms of each woody species (Table4).

Recruitment in the Micro-Scape

Highest recruit densities typically occurred immediately adjacent to central tree islands in the microscape (Figure 3) and declined exponentially with distance. Some species such as nona glabra and Salix caroliniana however were more broadly distributed into the prairies (Figure 3). A positive source effect was identibed for seven of the eight species-site combinations that contained enough individuals to justify a logistic regression (Tables 5, 6; Figure 4A). Signibcant hydroperiod effects were observed in only three cases; contrary to expectation, higher recruitment probability was associated with longer periods of sooding in each case (Tables 5, 6; Figure 4B).

Table 1. Density (stems/ha) of Individuals Present in Micro-scapes

| Species | A | | В | | С | | D | | Е | | F | |
|---------------------------------------|-------|---------|-------------|---------|--------------|---------|-------|---------|-------|---------|-------|---------|
| | Adult | Recruit | Adult | Recruit | Adult | Recruit | Adult | Recruit | Adult | Recruit | Adult | Recruit |
| Annona glabra | 1.6 | 41 | | | | 1 | | | | 23 | | 1 |
| Baccharis halimifolia | | | 0.2 | 1 | | | | | | | 0.5 | |
| Baccharis spp. | | | | | | | | | | | 0.6 | 2 |
| Bursera simaruba | | | | | | | | | | | 0.8 | |
| Cephalanthus occidentalis | 0.2 | | | | | | | | | | | |
| Chiococca parvifolia | | | | | | 1 | | | | | | |
| Chrysobalanus icaco | 0.3 | 5 | | | | | | | | 2 | | |
| Coccoloba diversifolia | | | | | | | | | | | 0.2 | |
| Conocarpus erectus | | | | | 0.2 | 1 | | | | | 0.2 | |
| Eugenia axillaris | | | | | | | | | | | 1.1 | |
| Ficus aurea | 0.2 | | | | 0.2 | | | 6 | | | 1.8 | 6 |
| llex cassine | 0.2 | | | | 0.3 | 3 | | | | 3 | 0.8 | |
| Magnolia virginiana | 0.2 | | | | | 1 | 2.0 | 59 | 0.3 | 3 | 1.6 | |
| Melaleuca quinquenervia ¹ | _ | | | | | | _ | | | - | 0.2 | |
| Morella cerifera | 3.9 | 2 | 0.8 | 11 | 2.1 | 4 | 4.1 | 34 | 1.6 | 6 | 18.7 | |
| Myrsine floridana | | | 0.2 | 1 | 8.0 | | | | | | 5.7 | 24 |
| Persea borbonia | 2.6 | 66 | 0.2 | 9 | 2.8 | 29 | 5.9 | 423 | 1.5 | 105 | 17.1 | |
| Psychotria nervosa | | | - | - | | | | | - | | 0.2 | |
| Randia aculeata | | | | | | 3 | | | | | | |
| Sabal palmetto | 0.7 | | | | 0.2 | 9 | | | 0.5 | 9 | 7.1 | 4 |
| Salix caroliniana | | 2 | | | 0.3 | • | | | 0.2 | Ţ | 7.1 | 3 |
| Schinus terebinthifolius ¹ | | | | | | | | | | | 8.6 | 1 |
| Serenoa repens | | | 0.2 | | | | | | | | 0.0 | • |
| Sideroxylon reclinatum | | | v. <u> </u> | | | 1 | | | | | 0.2 | |
| Sideroxylon salicifolium | | | | | 0.2 | - | | | | | 1.1 | |
| Solanum donianum | | | | | - | 1 | | 1 | | | | 155 |
| Taxodium distichum | | | | | 0.3 | • | 21.4 | 391 | | | | |
| Subtotals | 9.7 | 116 | 1.5 | 22 | 7.3 | 54 | 33.3 | 914 | 4.1 | 151 | 73.6 | 211 |

Adults were greater than 2 m in height, found within 140-m buffer including central tree island. Recruits were less than 2 m in height, found within 1-ha plot. Recruit densities do not include those inside central tree island.

¹Invasive exotic species.

Recruitment in the Meso-Scape

In meso-scapes A, B, C, E, and F, a reduced model that included only Woody Plant Source Factor as a predictor of recruitment produced the strongest results (Tables 7, 8, Figure 5). In each case, the probability of establishment increased signibcantly with WPSF, that is, the area and proximity of tree islands. In meso-scape D, the probability of establishment increased with both PWI and WPSF (Tables 7, 8; Figure 5), namely, in wet locations near tree islands.

Tree Island Patterning in the Macro-Scape

Homogeneity tests revealed that tree islands in all six macro-scapes were heterogeneously distributed (i.e., aggregated at very large [>200 m] scales). Using the heterogeneous Poisson null model, our

O-ring function showed that tree islands in macroscapes A, B, D, E, and F were more or less randomly structured with minor dispersion at some scales (Figure6). SpeciPcally, tree islands in macroscapes A, B, and D were over-dispersed at intermediate scales (i.e., 30Đ140, 20Đ150, and 10Đ125 m, respectively), and tree islands in macroscapes E and F were over-dispersed at smaller scales (i.e., below 95 and 5Đ25 m, respectively). Tree islands in macro-scape C were also over-dispersed at small scales (i.e., 5Đ25 m), and uniquely, they displayed aggregation at intermediate scales (i.e., 55Đ165 m).

Fire Patterns Across Micro-, Meso-, and Macro-Scapes

Seventy-eight Pres burned 36,951 ha of marl prairie habitat across our study region over the last

Table 2. Density (stems/ha) of Individuals Present in Meso-scapes

| Species | A | | | В | | C | | | D | | | E | | F | | | |
|--------------------------------------|-------|-----|-----|-------|--------|----------|-------|------|-------|------|------|-------|-----|--------|-----|------|-----|
| | 0.5-1 | 1-2 | >2 | 0.5-1 | 1–2 > | 2 0.5–1 | 1-2 | >2 | 0.5-1 | 1-2 | >2 | 0.5-1 | 1-2 | >2 0.5 | -1 | 1–2 | >2 |
| Acacia pinetorum | | | | | | 1.1 | | | | | | | | | | | |
| Annona glabra | 5.0 | 0.2 | | 0.6 | 0.2 | 1.7 | • | | | | | 1.1 | 0.4 | ; | 3.9 | 1.6 | |
| Baccharis spp. | | | | | 0.0 | 2.8 | 3 1.8 | | | | | | | | | | |
| Casuarina equisitifolia ¹ | | | | | | | 0.2 | | | | | | | 30 | 0.6 | 24.7 | 2.4 |
| Cephalanthus occidentalis | | | | | | 6.7 | 5.8 | | | | | | 0.2 | (| 3.9 | 1.6 | |
| Conocarpus erectus | | | | 1.1 | 0.7 | 3.3 | 6.2 | 2.4 | | | | | | | | 0.4 | |
| Eugenia axillaris | | | | | | | | | | | | | | • | .7 | | |
| Ficus aurea | | | | 2.8 | 1.3 | 0.6 | 0.4 | | 2.2 | | | 3.9 | 0.4 | | 1.1 | 2.0 | |
| Ilex cassine | | | | 0.6 | 0.4 | 16.7 | 1.8 | 0.2 | 0.6 | | | 2.8 | | | 1.1 | | |
| Magnolia virginiana | 1.1 | 0.9 | | 16.7 | 18.4 | 5.0 | 6.4 | 0.2 | 0.6 | 0.2 | 1.6 | ; | | | 1.1 | 0.9 | |
| Melaleuca quinquenervia ¹ | | | | | | | | | | | | | 0.2 | ţ | 5.0 | 3.3 | 0.2 |
| Metopium toxiferum | | | | 2.2 | 2.7 | | 0.2 | 0.9 | 0.6 | 0.9 | 0.4 | | | | | 0.7 | |
| Morella cerifera | 4.4 | 3.3 | 1.6 | 7.8 | 4.4 2 | .7 8.3 | 4.4 | | 5.0 | 0.7 | | 1.1 | 3.6 | : | 2.8 | 3.1 | 1.1 |
| Myrsine floridana | | 0.4 | | 6.7 | 2.9 | 10.0 | 2.4 | | | 0.2 | | 3.9 | | 12 | 2.8 | 6.4 | |
| Persea borbonia | 0.6 | 2.0 | | 12.2 | 13.3 0 | .2 20.6 | 5.1 | 1.6 | 5.0 | 2.4 | 0.2 | 1.7 | 2.2 | 10 | 0.6 | 4.9 | |
| Pinus elliotitii var. densa | | | | 1.1 | 2.2 | | | | | | | | | | | | |
| Randia aculeata | | | | 12.8 | 2.0 | | | | | | | 1.1 | | | | | |
| Sabal palmetto | 0.6 | 0.4 | 0.7 | 3.9 | 2.4 | 7.2 | 14.0 | 1.3 | | 1.3 | | 3.3 | 1.8 | 1 | 5.0 | 10.9 | 2.9 |
| Salix caroliniana | | | | | | 2.2 | 19.1 | 14.9 | | | | | | į | 5.6 | 21.3 | 9.8 |
| Sideroxylon reclinatum | | | | 10.6 | 0.4 | 5.0 | 8.2 | 3.1 | | | | | 0.2 | | | | |
| Sideroxylon salicifolium | | | | 1.1 | 0.9 | | 0.9 | | | | | | | | | | |
| Solanum donianum | | | | 0.0 | | 21.1 | 0.4 | | | | | | | 72 | 2.2 | 0.7 | |
| Stillingia sylvatica | | | | 0.6 | | | | | | | | | | | | | |
| Swietenia mahogani | | | | | 0.2 | | | | | | | | | | | | |
| Taxodium distichum | | | | | | | | | 56.1 | 77.3 | 82.9 | | | | | | |
| Subtotals | 11.7 | 7.3 | 2.2 | 80.6 | 52.7 2 | .9 112.2 | 77.6 | 24.7 | 70.0 | 83.1 | 85.1 | 18.9 | 9.1 | 16 | 7.2 | 82.4 | 16. |

¹Invasive exotic species.

Table 3. Mean Hydroperiod in Micro-Scapes, and Mean PWI in Meso-Scapes

| Plot | Micro-scape | | Meso-scape |
|------|---|------------------------------------|--------------|
| | Average center tree island hydroperiod \pm SD | Average marsh hydroperiod \pm SD | PWI |
| A | 173.3 ± 11.2 | 219.7 ± 8.8 | 8.4 ± 1.5 |
| В | 47.4 ± 3.6 | 78.8 ± 13.8 | 2.91 ± 1.7 |
| С | 126.7 ± 5.3 | 171.4 ± 24.0 | 2.48 ± 1.8 |
| D | 90.2 ± 10.0 | 164.4 ± 23.7 | 6.12 ± 1.3 |
| E | 168.3 ± 18.6 | 179.2 \pm 12.4 | 3.38 ± 1.8 |
| F | 166.3 ± 15.5 | 180.0 ± 16.4 | 2.80 ± 1.7 |

29 years. For the most part, <code>Pre</code> occurrences were consistent across the micro-, meso-, and macro-scapes in each subpopulation (Table9), suggesting that the processes observed at the smallest scales are representative of the general landscape. Fire occurrence intervals varied across the six micro-, meso-, and macro-scapes, with the longest <code>Pre</code> occurrence interval in macro-scape A (one <code>Pre</code> every 366.3 years), and <code>Pre</code> occurrence was

undetectable within the meso - and micro-scapes (at least over the last 400 years using our 29 year dataset; Table9). By contrast, macro-scapes BĐF averaged one Pre approximately every 14.5 years.

DISCUSSION

Vegetation in the marl prairie landscape is heterogeneous over time and space, and multiple factors

Table 4. Habitat Preference and Dispersal Mechanisms for Species in Micro- and Meso-Scapes

| Species | Habitat | Dispersal |
|---------------------------------------|---------|---------------------|
| Acacia pinetorum | Upland | Bird, mammal |
| Annona glabra | Wetland | Mammal, water |
| Baccharis halimifolia | Upland | Wind |
| Baccharis spp. | Upland | Wind |
| Bursera simaruba | Upland | Bird |
| Casuarina equisitifolia ¹ | Exotic | Wind |
| Cephalanthus occidentalis | Wetland | Water |
| Chiococca parvifolia | Upland | Bird, mammal |
| Chrysobalanus icaco | Wetland | Mammal |
| Coccoloba diversifolia | Upland | Bird, mammal |
| Conocarpus erectus | Wetland | Water |
| Eugenia axillaris | Upland | Bird, mammal |
| Ficus aurea | Upland | Bird, mammal |
| Ilex cassine | Wetland | Bird, mammal |
| Magnolia virginiana | Wetland | Bird, mammal |
| Melaleuca quinquenervia ¹ | Exotic | Wind, Water |
| Metopium toxiferum | Upland | Bird, mammal |
| Morella cerifera | Wetland | Bird, mammal, water |
| Myrsine floridana | Upland | Bird, mammal |
| Persea borbonia | Wetland | Bird, mammal |
| Pinus elliotitii var. densa | Upland | Wind |
| Psychotria nervosa | Upland | Bird, mammal |
| Randia aculeata | Upland | Bird, mammal |
| Sabal palmetto | Upland | Bird, mammal, water |
| Salix caroliniana | Wetland | Wind |
| Schinus terebinthifolius ¹ | Exotic | Bird, mammal |
| Serenoa repens | Upland | Bird, mammal |
| Sideroxylon reclinatum | Upland | Bird, mammal |
| Sideroxylon salicifolium | Upland | Bird, mammal |
| Solanum donianum | Upland | Bird, mammal |
| Taxodium distichum | Wetland | Water |

Dispersal for each species inferred from Wunderlin (1998) and/or Tomlinson (2001).

Invasive exotic species.

interact to drive the emergence, expansion, and contraction of woody assemblages at each scale. At local and intermediate scales recruits tend to aggregate around existing adults and/or tree islands, due to gradients in dispersal and/or smaller-scale gradients in habitat suitability. Early marsh invaders may eventually contribute to tree island growth, or serve as nucleation points for new neighboring communities. However, long-term nucleation and survival around recruits appears to be powerfully constrained by abiotic factors, so much so that tree islands at larger scales exhibit random and over-dispersed distributions in most macro-scapes.

Recruitment Patterns and Processes at Small and Intermediate Scales

Woody patch establishment begins with seedling recruitment in the marsh. Establishment success

depends upon several variables, including the spatial distribution of adult populations, spatial and temporal variation in seed vectors, and ambient environmental conditions (Bazzaz 1991; Schupp 1995). Logistic regressions at the micro and mesoscales uniformly support the premise that the size and distribution of tree islands in the landscape play a formative role in the patterning of germinant populations. In each model, source size and proximity increased the likelihood of establishment in the marsh. Seed densities are known to be highest in close proximity to parent trees across many ecosystems (Hughes and Fahey 1988; Campbell and others 1990; Clark and others 1999), although seedfall patterning can vary widely among species (Ribbens and others 1994). The interspecibe variation observed in our logistic recruitment curves may result from factors known broadly to affect dispersal, including parental fecundity (Herrera

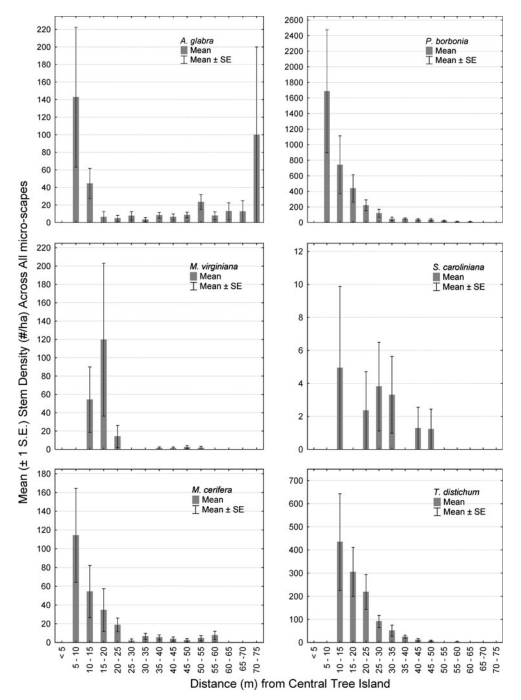


Figure 3. Mean (±1 SE) A. glabra, M. virginiana, M. cerifera, P. borbonia, S. caroliniana, and T. distichum stem densities (#/ha) for all micro-scapes combined as a function of distance from tree island centroid. Seedlings were not sampled below tree island canopies.

1998), diaspore morphology (Howe and Smallwood 1982; Bazzaz 1996), disperser behavior (Nathan and Muller-Landau 2000), and substrate conditions (Fort and Richards 1998; van der Wall and Joyner 1998). Seed shadows serve as the template on which subsequent environmental processes further constrain germination, establishment, and survival (Nathan and Muller-Landau 2000). Such constraints include light, nutrient and water availability, seed predation, herbivory, competition, and disease. In addition to biotic and

abiotic source effects, environmental factors outside the zone of adult inßuence such as ßooding and Þre likely inßuence early recruitment success.

Hydrology can impact seedling establishment in several ways, and evidence for hydrologic impacts may depend on which metric is analyzed. For example, long ßooding durations may favor one species over another during average hydrologic years; however, the same seedling populations may respond differently when hydrologic timing is altered (for example, inundation early in the

Table 5. Logistic Regression Results for Source and Hydroperiod Effects in Micro-Scapes

| | Species | Model P | Source effect (β_1) | SE | Wald | P | Exp (β_1) | Hydroperiod (β_2) | SE | Wald | P | Exp (β_2) |
|-----|--------------|---------|---------------------------|------|-------|-------|-----------------|---------------------------|------|-------|-------|-----------------|
| Α | A. glabra | 0.00* | 44.82 | 8.54 | 27.55 | 0.00* | 2.90E19 | (-) | (-) | (-) | (-) | (-) |
| Α | P. borbonia | 0.00* | 5.61 | 1.07 | 27.46 | 0.00* | 273.46 | (-) | (-) | (-) | (-) | (-) |
| В | M. cerifera | 0.01* | 4.72 | 1.54 | 9.45 | 0.00* | 112.03 | (-) | (-) | (-) | (-) | (-) |
| D I | M. cerifera | 0.01* | (-) | (-) | (-) | (-) | (-) | 0.03 | 0.01 | 6.97 | 0.01* | 1.03 |
| D | P. borbonia | 0.01* | 0.70 | 0.22 | 9.69 | 0.00* | 2.01 | 0.02 | 0.01 | 7.09 | 0.01* | 1.02 |
| D . | T. distichum | 0.00* | 0.29 | 0.10 | 9.13 | 0.00* | 1.33 | 0.03 | 0.01 | 27.17 | 0.00* | 1.03 |
| E | M. cerifera | 0.03* | 4.36 | 1.66 | 6.91 | 0.01* | 78.07 | (-) | (-) | (-) | (-) | (-) |
| E | P. borbonia | 0.00* | 5.57 | 1.85 | 9.08 | 0.00* | 262.56 | (-) | (-) | (-) | (-) | (-) |

⁽⁻⁾ indicates term was removed from model.

Table 6. Meso-Scape Logistic Regression Model Assessments

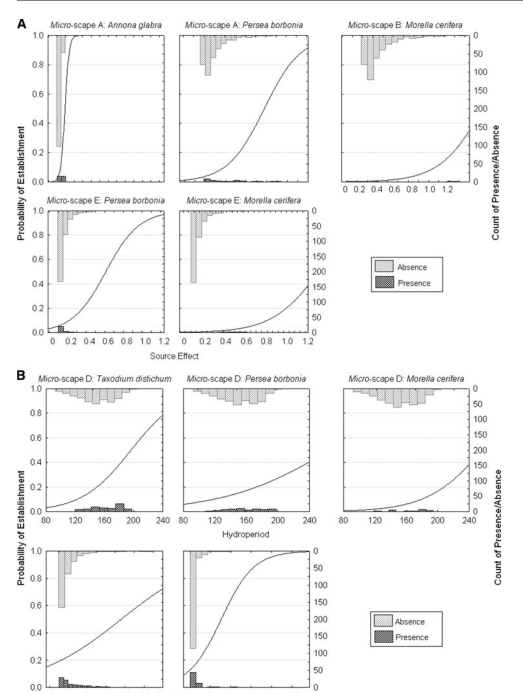
| Plot | Species | Omnibus test (<i>P</i>) | Classification accuracy (%) | Tolerance | Variance inflation factor | Occupied cells | Adults |
|------|--------------|------------------------------|-----------------------------|-----------|------------------------------|----------------|--------|
| A | A. glabra | 0.00 | 92.1 | 1.00 | 1.00 | 36 | 10 |
| Α | P. borbonia | 0.00 | 92.4 | 0.97 | 1.04 | 41 | 16 |
| В | M. cerifera | 0.01 | 98.7 | 0.99 | 1.01 | 6 | 27 |
| D | M. cerifera | 0.01 | 94.3 | 1.00 | 1.00 | 24 | 25 |
| D | P. borbonia | 0.00 | 83.8 | 1.00 | 1.00 | 73 | 36 |
| D | T. distichum | 0.00 | 77.5 | 0.99 | 1.01 | 97 | 132 |
| Е | M. cerifera | 0.03 | 99.0 | 1.00 | 1.00 | 5 | 10 |
| Е | P. borbonia | 0.00 | 92.3 | 1.00 | 1.00 | 34 | 9 |

growing season; Reed 2007). In this study, we focused on annual duration of inundation (hydroperiod) in the micro-scape and a vegetationderived wetness index (PWI) in the meso-scape. However, variation in hydroperiod and PWI at these scales showed weak and non-signibcant correlations with establishment in all but one area. High densities of hydrophilic species in our wettest populations, and positive effects of Booding on establishment in micro-scape D do suggest that surface hydrology plays an important role in large-scale woody species distributions. The lack of evidence for Þner-scale hydrologic impacts could result from a scale mismatch between species response and hydrologic cell size (5 \times 5 or 10 \times 10 m for micro- and meso-scapes, respectively); although, source effects clearly play a stronger role in recruitment patterns in the marsh. Furthermore, source effects may incorporate some hydrologic impacts, where metrics such as duration, depth, and timing are most optimal in close proximity to adult populations.

Local scale interactions between hydrology, community composition, and ignition sources promote a heterogeneous Pre regime across the

southern Everglades (Wade and others 1980). As one might expect, Pre frequency was highest in our dry populations (i.e., C and F) and decreased with Booding duration. Effects of the heterogeneous Pre regime may be magniped by the differential responses of woody species, which depend upon their physiological traits and local environmental conditions. R-strategists and species with clonal and vegetative growth patterns typically recover quickly, whereas woody stems that can only invade under a narrow set of conditions (for example, dry periods such as 2006D2008) may be removed for decades by Pre. Ross and others 2009) found that the combination of Pre and Booding can drastically alter community composition in the marl prairies. A prime example occurred in micro-scape B, which burned intensively in 2005, followed within days by ßooding associated with Hurricane Katrina. In 2007, only four new recruits were found in the 1-ha plot, suggesting that the PreDßood interaction interrupted the annual recruitment process surrounding the small tree island. Very few reproductive trees survived the Pre and Booding inhibited the establishment and survival of the propagules that remained.

^{*}Significant at $\alpha = 0.05$.



6 8 10 12

Figure 4. A Signibcant source effects for all reduced model logistic regressions in microscapes (A, B and E). B Source and hydroperiod effects for all signibcant logistic regressions in micro-scape D. M. cerifera establishment was best expressed with a reduced model using hydroperiod only. Both source effects and hydrology signibcantly inßuence the establishment of P. borbonia and T. distichum.

Large-Scale Patterning of Tree Islands in the Marl Prairie Landscape

0 2

6 8 10 12 0 2

According to the HPDP patterns observed at large scales, such as tree island distributions across the macro-scape, impose constraints on smaller-scale processes such as recruitment in the prairie. At the same time, lower-level processes give rise to higher-level patterns (Wu 1999). Using this logic, one

Source Effect

might expect woody plants to aggregate at multiple scales across the marl prairie landscape, resulting from (1) Þrst order, abiotic-driven association of tree islands at very large scales, which limits dispersal to regions in which tree islands are present, and (2) smaller-scale dispersal patterns, where short distance dispersal contributes to the growth of existing tree islands, and longer distance dispersal

Table 7. Logistic Regression Results for Source and Hydrology Effects in Meso-Scapes

| Plot | Model P | Source effect (β_3) | SE | Wald | P | Exp (β_3) | Hydroperiod (β_4) | SE | Wald | P | Exp (β ₄) |
|------|---------|-----------------------------|------|-------|-------|-----------------|---------------------------|------|-------|-------|-----------------------|
| Α | 0.00* | 0.08 | 0.01 | 42.97 | 0.00* | 1.09 | (-) | (-) | (-) | (-) | (-) |
| В | 0.00* | 0.02 | 0.00 | 25.55 | 0.00* | 1.02 | (-) | (-) | (-) | (-) | (-) |
| С | 0.00* | 0.05 | 0.01 | 30.25 | 0.00* | 1.05 | (-) | (-) | (-) | (-) | (-) |
| D | 0.00* | 0.31 | 0.00 | 49.72 | 0.00* | 1.03 | 0.29 | 0.08 | 12.25 | 0.00* | 1.34 |
| E | 0.03* | 0.06 | 0.02 | 6.39 | 0.00* | 1.06 | (-) | (-) | (-) | (-) | (-) |
| F | 0.00* | 0.03 | 0.01 | 19.80 | 0.00* | 1.04 | (-) | (-) | (-) | (-) | (-) |

Plots A, B, C, E, and F employ a reduced model; plot D employs a complete model. (–) indicates term was removed from model. *Significant at $\alpha = 0.05$.

Table 8. Meso-Scape Logistic Regression Model Assessments

| Plot | Omnibus test for model coefficients (P) | Classification accuracy (%) | Tolerance | Variance inflation factor | Occupied cells | Islands |
|------|---|-----------------------------|-----------|------------------------------|----------------|---------|
| A | 0.00 | 99.2 | 0.90 | 1.12 | 15 | 19 |
| В | 0.00 | 93.8 | 0.95 | 1.05 | 38 | 69 |
| С | 0.00 | 84.2 | 0.99 | 1.01 | 117 | 50 |
| D | 0.00 | 86.0 | 0.99 | 1.02 | 105 | 18 |
| E | 0.03 | 97.9 | 0.93 | 1.07 | 15 | 27 |
| F | 0.00 | 87.2 | 0.99 | 1.01 | 92 | 33 |

creates nucleation points for new, spatially associated communities. Much to our surprise however tree islands in the macro-scape were mostly randomly distributed, with over-dispersion at small scales. The lack of second-order aggregation suggests that biological feedbacks, such as those promoting the aggregation of new recruits around adult populations are not sufpciently strong to promote nucleation and subsequent tree island aggregation over time in the face of external constraints.

Over-dispersion or ÔÔrepulsionÕÕ is usually observed with individual organisms, and is normally the result of biological processes such as competition or allelopathy. It is not easy to imagine a biological process that would cause one tree island to be repelled by another. Instead, we suggest that the observed over-dispersion may result from interplay between environmental constraints such as Þre, ßooding, and existing tree island patterning across the landscape.

Hydrologic patterns in the marl prairies are depend by long-term variables including climate and geology, and modiped by anthropogenic alterations such as compartmentalization and drainage. Under natural hydrologic regimes, ßooding stress may limit tree island establishment to micro-topographic highs where hydroperiods are much more abbreviated than elsewhere in the marsh. Bedrock

outcrops in the marsh are often precursors to surface topographic rises, though it remains unclear to what degree tree island patterning depends upon bedrock topography. Observed recruitment patterns in the micro- and meso-scapes and tree island distributions across the macro-scape suggest that although second-order (biological) feedbacks might allow woody stems to colonize new environmentally favorable habitats throughout the prairie, tree island establishment, a much more prolonged process, could be limited to topographic outcrops, which may or may not exhibit aggregation at different scales. However, tree island size, distribution, and overall landscape complexity may change or have changed rapidly in regions where hydrologic conditions have been drastically altered.

Fire is another important constraint on woody plant patterning in mixed landscapes (Bond and others 2005), and may be particularly important in the Þre-prone marl prairies of the southern Everglades. As tree islands form and increase in size they develop more humid microclimates, and become increasingly resistant to Þre (van der Valk and Sklar 2004), whereas nearby recruits and small woody plant aggregates remain extremely vulnerable. In fast-moving head-Þres, ßow paths may diverge on approach to large islands, intensifying along their ßanks. Such an effect on Þre behavior could serve to eliminate previously established recruits from an

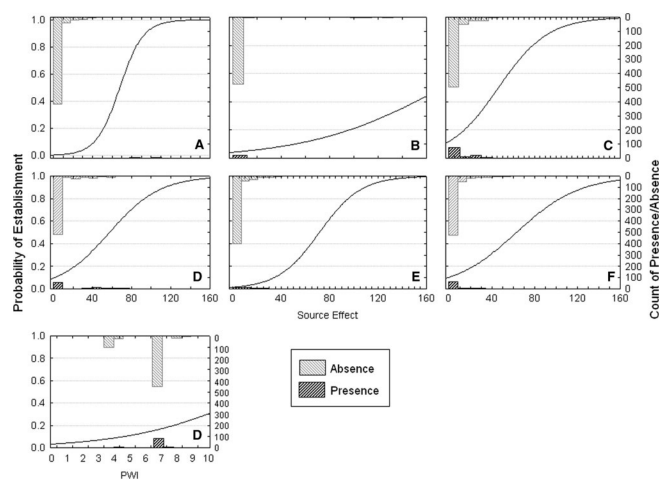


Figure 5. Source and PWI effects for all signibcant logistic regressions across all meso-scapes.

islandÕs vicinity, short-circuit fuel buildup nearby, and limit island size. Over time, it could perhaps lead to over-dispersion of islands at small scales.

Paradoxically, the only area in which local tree island patterning indicated aggregation instead of over-dispersion was the area in which Pre frequency was greatest. Second-order tree island aggregation in the relatively dry macro-scape C may have progressed similarly to invasion processes that sometimes occur in open savannas. Mass recruitment of woody seedlings can occur in savannas when Þre and subsequent rain storms reduce the graminoid layer and moisten the soil (Danthu and others 2003; Wiegand and others 2006a). These new woody patches can subsequently expand and aggregate over time because reduced grass layers can no longer sustain intense burns (Menaut and others 1990; Wiegand and others 2006a). Similarly in macro-scape C, Pre and reduced ßooding may have created optimal environments for tree island establishment, expansion, and aggregation. Perhaps, when recruitment occurs

in localized mass events (due to Pre-drought interplay) longer-term tree island growth and survival is more probable. Furthermore, frequent Pres in population C likely prevent fuel accumulation, thus reducing the probability of intense burns that would otherwise consume more woody resources. Lastly, large aggregates of tree islands are known to deßect Pres, and interfere with their movement through the prairie (personal communication, Rick Anderson, ENP Fire Management OfPcer), so in dry regions, frequent low-intensity Pres may allow aggregation to become an autogenic process.

In their discussion of multi-scaled patch and boundary dynamics, Peters and others (2006) debned three types of biotic transitions: directional, stationary, and shifting. Directional transitions such as those associated with woody plant invasion involve patch succession from one end state to another. These shifts are strongly inßuenced by abiotic drivers such as changes in climate, in addition to biotic and abiotic feedbacks. Over time, directional transitions create larger and more

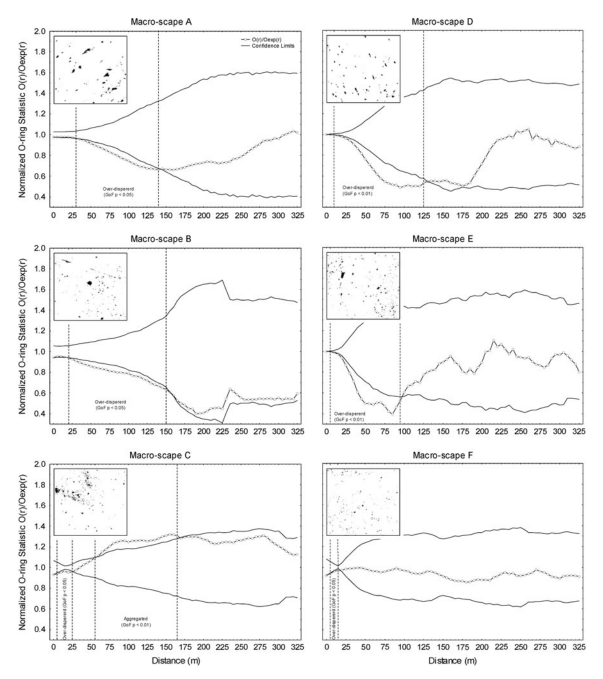


Figure 6. Univariate analysis of tree island aggregation within each of the six macro-scapes using the O-ring function, results from goodness of Pt tests and inlay map of all macro-scapes. To better see the positive or negative departures from the null model Pt0 is normalized by Pt0 under the homogeneous Pt0 is normalized by Pt0 under the homogeneous Pt0 is the average Pt0 of the simulations of the null model. Black features in inlay represent tree islands that were used in our o-ring analyses, whereasgray features represent other communities (for example, pine forests) or islands truncated by the plot boundary, and were masked from the analyses.

spatially aggregated patches, which eventually may coalesce (Peters and others 2006). Stationary transitions remain somewhat stable over decades with little patch movement from one state to another. Stationary boundaries are controlled by intrinsic abiotic constraints including topographic anomalies, and are strengthened over time by

biotic feedbacks. Shifting transitions involve no net long-term change in boundary locations, though patch boundaries shift over shorter timescales. Shifting transitions are strongly affected by abiotic drivers such as ßooding, drought and Þre, and are not reinforced over time by biotic or abiotic feedback mechanisms (Peters and others2006).

Table 9. Fire Occurrence Interval for All Three Landscape Scales

| CSSS | Fire occurrence interval (years) | | | | | | | |
|----------------|----------------------------------|------------|-------------|--|--|--|--|--|
| subpopulations | Macro-scape | Meso-scape | Micro-scape | | | | | |
| A | 366.3 | >400 | >400 | | | | | |
| В | 10.5 | 17.9 | 14.5 | | | | | |
| С | 8.3 | 10.9 | 9.7 | | | | | |
| D | 21.5 | 11.0 | 7.3 | | | | | |
| E | 16.9 | 11.2 | 9.7 | | | | | |
| F | 15.5 | 7.3 | 4.9 | | | | | |

In interpreting this study, multiple transition types likely exist across the Everglades marl prairie landscape. The observed disconnect between aggregation of recruits around tree islands and tree island over-dispersion at larger scales however suggests that directional transitions are rare in most macro-scapes. Although nucleation feedbacks promote the recruitment of woody stems near conspecibe adults in populations A, B, D, E, and F, they do not appear to be sufbeintly strong to reinforce directional patch aggregation over time. Tree island over dispersion may result from the patterning of underlying bedrock highs and/or the distribution of

Pre and water through the prairie. Such factors are likely associated with stationary and shifting boundaries, respectively. However, when natural hydrologic and disturbance regimes are directionally altered (for example, ßow diversion and long-term Pre suppression), shifting transitions might become directional (Peters and others 2006). Large-scale tree island aggregation in population C may result from directional shifts, triggered by dry conditions.

CONCLUSION

The Everglades grassland-woody plant mosaic is heterogeneous and dynamic across multiple scales. Despite the adaptive capacities of resident biota, landscape patches are perpetually subject to expansion, contraction, or replacement. Similar to other grasslands worldwide, woody encroachment into Everglades prairies may involve a chain of spatially dependent processes, beginning with the dispersal and establishment of individual propagules, the subsequent aggregation of individuals into tree islands and Þnally the potential coalescence of tree islands into more extensive forests (Peters and others 2006; Allen and Breshears 1998). Conversely, woody patch contraction may

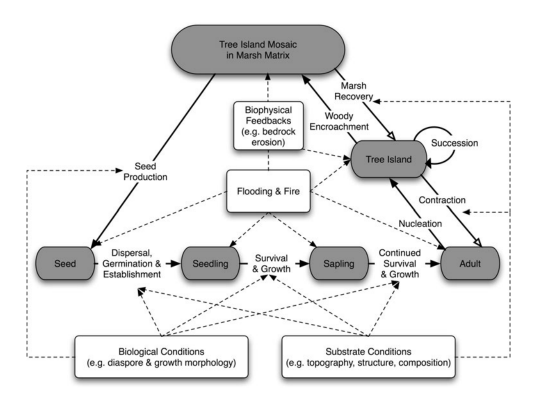


Figure 7. The major controls and feedbacks inßuencing patterns among woody plants at various life stages. Solid boxes indicate stages of woody encroachment/contraction. Hollow boxes indicate environmental factors inßuencing the two processes. Solid filled arrows represent encroachment processes, whereashollow arrows denote marsh recovery. Dashed arrows indicate inßuences upon each process and/or stage.

result from environmental constraints such as nutrient limitation, ßooding, and Þre (Lockwood et al 2003; Figure 7). Therefore, a proper balance between environmental feedbacks and stressors is required to maintain a landscape mosaic suitable to the diverse requirements of its component species.

The patterning of adult populations within the Everglades marl prairies appears to be the most signibcant driver of tree island development and growth. However, following initial recruitment in the marsh, establishment of new tree islands may be extremely slow and relatively rare, with most early marsh invaders being eliminated over time by disturbance. Under this scenario, general tree island distributions may date back to the origins of grassland-woody mosaic, approximately 2,000 years (Willard and others 2006). Nevertheless, pronounced changes in hydrology or Pre regime, such as those resulting from development and drainage, might dramatically restructure this legacy tree island pattern.

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REFERENCES

- Allen CD, Breshears DD. 1998. Drought-induced shift of a forest-woodland ecotone: rapid landscape response to climate variation. Proc Natl Acad Sci USA 95:14839Đ42.
- Bazzaz FA. 1991. Habitat selection in plants. Am Nat (Suppl) 137:S116Đ30.

- Bazzaz FA. 1996. Plants in changing environments: linking physiological, population, and community ecology. New York: Cambridge University Press. pp 82D107.
- Bond WJ, Woodward FI, Midgley GF. 2005. The global distribution of ecosystems in a world without Pre. New Phytol 165:525D38.
- Browder JA, Ogden JC. 2000. The natural South Florida system II: predrainage ecology. Urban Ecosyst 3:245Đ77.
- Campbell BM, Lynam T, Hatton JC. 1990. Small-scale patterning in the recruitment of forest species during succession in tropical dry forest, Mozambique. Vegetatio 87:51Đ7.
- Chase JM, Leibold MA. 2002. Spatial scale dictates the productivity-biodiversity relationship. Nature 416:427Đ30.
- Clark JS, Silman M, Kern R, Macklin E, HilleRisLambers J. 1999. Seed dispersal near and far: patterns across temperate and tropical forests. Ecology 80(5):1475Đ94.
- Danthu P, Ndongo M, Diaou M, Thiam O, Sarr A, Dedhiou B, Ould Mohamed Vall A. 2003. Impact of bush Pre on germination of some West African Acacias. For Ecol Manag 173:1Đ10.
- EDEN Everglades Depth Estimation Network. 2008. South Florida Information Access (SoÞa). http://soÞa.usgs.gov/eden. Cited 12 May 2008.
- Fort KP, Richards JH. 1998. Does seed dispersal limit initiation of primary succession in desert playas? Am J Bot 85:1722Đ31.
- Givnish TJ, Volin JC, Owen VD, Volin VC, Muss JD, Glaser PH. 2008. Vegetation differentiation in the patterned landscape of the central Everglades: importance of local and landscape drivers. Glob Ecol Biogeogr 17(3):384Đ402.
- Gunderson LH, Snyder JR. 1994. Fire patterns in the southern Everglades. In: Davis SM, Ogden JC, Eds. Everglades: the ecosystem and its restoration. Delray Beach: St. Lucie Press. p 291Đ306.
- Hanan EJ, Ross MS. 2010. Across-scale patterning of plant-soil-water interactions surrounding tree islands in southern Everglades Landscapes. Landscape Ecol. ddi0.1007/s10980-009-9426-9.
- Herrera CM. 1998. Population-level estimates of interannual variability in seed production: what do they actually tell us? Oikos 82:612D16.
- Hoffman W, Bancroft GT, Sawicki RJ. 1994. Foraging habitat of wading birds in the water conservation areas of the Everglades. In: Davis SM, Ogden JC, Eds. Everglades: the ecosystem and its restoration. Boca Raton: St. Lucie Press. p 323Đ40.
- Howe HF, Smallwood J. 1982. Ecology of seed dispersal. Annu Rev Ecol Syst 13:201Đ28.
- Hughes JW, Fahey TJ. 1988. Seed dispersal and colonization in a disturbed northern hardwood forest. Bull Torrey Bot Club 115:89D99.
- Jenkins CN, Powell RD, Bass OL Jr, Pimm SL. 2003a. Demonstrating the destruction of the habitat of the Cape Sable seaside sparrow (Ammodramus maritimus mirabilis). Anim Conserv 6:29D38.
- Jenkins CN, Powell RD, Bass OL Jr, Pimm SL. 2003b. Why sparrow distributions do not match model predictions. Anim Conserv 6:39Đ46.
- La Puma DA, Lockwood JL, Davis MJ. 2007. Endangered species management requires a new look at the benebt of bre: the Cape Sable seaside sparrow in the Everglades ecosystem. Biol Conserv 136:398b407.
- Levin SA. 1992. The problem of pattern and scale in ecology. Ecology 73:1943Đ67.

- Lockwood JL, Ross MS, Sah JP. 2003. Smoke on the water: the interplay of Pre and water ßow on Everglades restoration. Front Ecol Environ 1:462Đ8.
- Loosmore NB, Ford ED. 2006. Statistical inference using the G or K point pattern spatial statistics. Ecology 87:1925D31.
- McCarthy TS, Ellery WN, Ellery K. 1993. Vegetation-induced, subsurface precipitation of carbonate as an aggradational process in the permanent swamps of the Okavango (delta) fan, Botswana. Chem Geol 107:111Đ31.
- Meeker M. 2008. Weathering the challenge of drought management in south Florida. Proceedings from the American Water Resources Association 2008 annual water resources conference, New Orleans, LA.
- Menaut JC, Gignoux J, Prado C, Clobert J. 1990. Tree community dynamics in a humid savanna off the Cote DÕlvoire: modeling the effects of Pre and competition with grass and neighbours. J Biogeogr 17:471Đ81.
- Nathan R, Muller-Landau HC. 2000. Spatial patterns of seed dispersal, their determinants and consequences for recruitment. TREE 15(7):278Đ85.
- Nott MP, Bass OL Jr, Fleming DM et al. 1998. Water levels, rapid vegetational changes, and the endangered Cape Sable Seaside Sparrow. Anim Conserv 1:23Đ32.
- Obeysekera J, Browder J, Hornung L, Harwell MA. 1999. The natural South Florida system I: climate, geology, and hydrology. Urban Ecosyst 3:223Đ44.
- Peters DPC, Yao J, Gosz JR. 2006. Woody plant invasion at a semi-arid/arid transition zone: importance of ecosystem type to colonization and patch expansion. J Veg Sci 27:389Đ96.
- Pimm SL, Lockwood JL, Jenkins CN, Curnutt JL, Nott MP, Powell RD, Bass Jr. OL. 2002. Sparrow in the Grass. A report on the Prst ten years of research on the Cape Sable Seaside Sparrow, Unpublished Report.
- Reed DL. 2007. Effects of hydrology and light on seedling establishment and growth of four wetland tree species in tree islands of northern Shark Slough, Everglades National Park. Florida International University, Miami, FL, Unpublished Masters Thesis.
- Ribbens E, Silander JA Jr, Pacala SW. 1994. Seedling recruitment in forests: calibrating models to predict patterns of tree seedling dispersion. Ecology 75(6):1794D806.
- Rietkerk M, Dekker SC, Wassen MJ, Verkroost AWM, Bierkens MFP. 2004. A putative mechanism for bog patterning. Am Nat 163:699D708.
- Ross MS, Sah JP, Ruiz PL, et al. 2009. Effect of hydrologic restoration on the habitat of the Cape Sable seaside sparrow: unpublished annual report of 2008D2009.
- Ross MS, Mitchell DBruker S, Sah JP, et al. 2006a. Interaction of hydrology and nutrient limitation in the Ridge and Slough landscape of the southern Everglades. Hydrobiologia 569:37D59.
- Ross MS, Sah JP, Ruiz PL, et al. 2006b. Effect of hydrologic restoration on the habitat of the Cape Sable seaside sparrow: unpublished annual report of 2005D2006.

- Schupp EW. 1995. Seed-seedling conßicts, habitat choice, and patterns of plant recruitment. Am J Bot 82(3):399£409.
- Thomas SC. 1996. Reproductive allometry in Malaysian rain forest trees; biomechanics versus optimal allocation. Evol Ecol 10:517Đ30.
- Tomlinson PB. 2001. The biology of trees native to tropical Florida. 2nd edn. Allston: Harvard Printing and Publication Services.
- van der Wall SB, Joyner JW. 1998. Secondary dispersal by the wind of winged pine seeds across the ground surface. Am Midl Nat J 139:365Đ73.
- van der Valk A, Sklar FA. 2002. What we know and should know about tree islands. In: Tree islands of the Everglades. Kluwer, Dordrecht. pp 311Đ336
- Wade D, Ewel J, Hofstetter R. 1980. Fire in South Florida ecosystems. US Forest Service General Technical Report SE-17. Ashville, NC: Southeast Forest Experiment station.
- Watt AS. 1947. Pattern and process in the plant community. J Ecol 35:1Đ22.
- Wetzel PR, Van Der Valk AG, Newman S et al. 2005. Maintaining tree islands in the Florida Everglades: nutrient redistribution is the key. Front Ecol Environ 3(7):370Đ6.
- Wiegand K, Saltz D, Ward D. 2006a. A patch-dynamics approach to savanna dynamics and woody plant encroachmentDInsights from an arid savanna. Perspect Plant Ecol Evol Syst 7:229D42.
- Wiegand T, Moloney K. 2004. Rings, circles, and null-models for point pattern analysis in ecology. Oikos 104:209D29.
- Wiegand T, Kissling D, Cipriotti PA, Aguiar MA. 2006b. Extending point pattern analysis for objects of Pnite size and irregular shape. J Ecol 94(4):825Đ37.
- Wiegand T, Gunatilleke S, Gunatilleke N, Okuda T. 2007. Analyzing the spatial structure of a Sri Lankan tree species with multiple scales of clustering. Ecology 88(12):3088Đ102.
- Willard DA, Bernhardt CE, Holmes CW, Landacre B, Marot M. 2006. Response of Everglades tree islands to environmental change. Ecol Monogr 76(4):565Đ83.
- With KA, King AW. 2001. Analysis of landscape sources and sinks: the effect of spatial pattern on avian demography. Biological Conservation 100:75Đ88.
- Wu J. 1999. Hierarchy and scaling: extrapolating information along a scaling ladder. Can J Remote Sens 25:367Đ80.
- Wu J, Levin SA. 1994. A spatial patch dynamics modeling approach to pattern and process in an annual grassland. Ecol Monogr 64(4):447Đ64.
- Wu J, Loucks OL. 1995. From balance-of-nature to hierarchical patch dynamics: a paradigm shift in ecology. Q Rev Biol 70:439Đ66.
- Wunderlin RP. 1998. Guide to the vascular plants of Florida. Jacksonville: University of Florida Press.