Influence of Soil Biogeochemical Properties on the Invasiveness of Old World Climbing Fern (Lygodium microphyllum)

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INFLUENCE OF SOIL BIOGEOCHEMICAL PROPERTIES ON THE INVASIVENESS OF OLD WORLD CLIMBING FERN (*LYGODIUM MICROPHYLLUM*)

A dissertation submitted in partial fulfillment of the requirements for the degree of DOCTOR OF PHILOSOPHY in GEOSCIENCES by Pushpa Gautam Soti

2013
To: Dean Kenneth G. Furton  
College of Arts and Sciences  

This dissertation, written by Pushpa Gautam Soti, and entitled Influence of Soil Biogeochemical Properties on the Invasiveness of Old World Climbing Fern (*Lygodium microphyllum*), having been approved in respect to style and intellectual content, is referred to you for judgment.

We have read this dissertation and recommend that it be approved.

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Date of Defense: October 31, 2013  

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Florida International University, 2013
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ABSTRACT OF THE DISSERTATION

INFLUENCE OF SOIL BIOGEOCHEMICAL PROPERTIES ON THE
INVASIVENESS OF OLD WORLD CLIMBING FERN (*LYGODIUM
MICROPHYLLUM*)

by

Pushpa Gautam Soti

Florida International University, 2013

Miami, Florida

Professor Krishnaswamy Jayachandran, Major Professor

The state of Florida has one of the most severe exotic species invasion problems
in the United States, but little is known about their influence on soil biogeochemistry. My
dissertation research includes a cross-continental field study in Australia, Florida, and
greenhouse and growth chamber experiments, focused on the soil-plant interactions of
one of the most problematic weeds introduced in south Florida, *Lygodium microphyllum*
(Old World climbing fern). Analysis of field samples from the ferns introduced and their
native range indicate that *L. microphyllum* is highly dependent on arbucular mycorrhizal
fungi (AMF) for phosphorus uptake and biomass accumulation. Relationship with AMF
is stronger in relatively dry conditions, which are commonly found in some Florida sites,
compared to more common wet sites where the fern is found in its native Australia. In the
field, *L. microphyllum* is found to thrive in a wide range of soil pH, texture, and nutrient
conditions, with strongly acidic soils in Australia and slightly acidic soils in Florida.
Soils with pH 5.5 - 6.5 provide the most optimal growth conditions for *L. microphyllum*,
and the growth declines significantly at soil pH 8.0, indicating that further reduction
could happen in more alkaline soils. Comparison of invaded and uninvaded soil characteristics demonstrates that *L. microphyllum* can change the belowground soil environment, with more conspicuous impact on nutrient-poor sandy soils, to its own benefit by enhancing the soil nutrient status. Additionally, the nitrogen concentration in the leaves, which has a significant influence in the relative growth rate and photosynthesis, was significantly higher in Florida plants compared to Australian plants. Given that *L. microphyllum* allocates up to 40% of the total biomass to rhizomes, which aid in rapid regeneration after burning, cutting or chemical spray, hence management techniques targeting the rhizomes look promising. Over all, my results reveal for the first time that soil pH, texture, and AMF are major factors facilitating the invasive success of *L. microphyllum*. Finally, herbicide treatments targeting rhizomes will most likely become the widely used technique to control invasiveness of *L. microphyllum* in the future. However, a complete understanding of the soil ecosystem is necessary before adding any chemicals to the soil to achieve a successful long-term invasive species management strategy.
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## ABBREVIATIONS AND ACRONYMS

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<tr>
<td>Al</td>
<td>Aluminum</td>
</tr>
<tr>
<td>AMF</td>
<td>Arbuscular mycorrhizal fungi</td>
</tr>
<tr>
<td>C</td>
<td>Carbon</td>
</tr>
<tr>
<td>Ca</td>
<td>Calcium</td>
</tr>
<tr>
<td>Cu</td>
<td>Copper</td>
</tr>
<tr>
<td>CFU</td>
<td>Colony forming units</td>
</tr>
<tr>
<td>DI</td>
<td>Deionized</td>
</tr>
<tr>
<td>Fe</td>
<td>Iron</td>
</tr>
<tr>
<td>FLEPPC</td>
<td>Florida Exotic Pest Plant Council</td>
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<tr>
<td>K</td>
<td>Potassium</td>
</tr>
<tr>
<td>LMR</td>
<td>Leaf mass ratio</td>
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<tr>
<td>Mg</td>
<td>Magnesium</td>
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<tr>
<td>Mn</td>
<td>Manganese</td>
</tr>
<tr>
<td>N</td>
<td>Nitrogen</td>
</tr>
<tr>
<td>OM</td>
<td>Organic matter</td>
</tr>
<tr>
<td>P</td>
<td>Phosphorus</td>
</tr>
<tr>
<td>RPM</td>
<td>Rounds per minute</td>
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<tr>
<td>RGR</td>
<td>Relative growth rate</td>
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<tr>
<td>RhiMR</td>
<td>Rhizome mass ratio</td>
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<tr>
<td>RMR</td>
<td>Root mass ratio</td>
</tr>
<tr>
<td>SLA</td>
<td>Specific leaf area</td>
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<tr>
<td>SMR</td>
<td>Stem mass ratio</td>
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<tr>
<td>Zn</td>
<td>Zinc</td>
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INTRODUCTION

1.1 Background

Invasive species are among the major factors affecting the integrity and function of ecosystems worldwide (Reid and Miller 1989; Williamson 1996; Luken and Thieret 1997). Invasive species are characterized by their adaptability to various habitats, increased competitiveness, higher reproductive potential and lack of natural predators to limit their extensive growth. Along with economic loss, invasive species cause significant loss of ecosystem stability, functional complexity and biodiversity (Gordon 1998; Williams and West 2000) because the biological interactions in the rhizosphere play a significant role in plant growth, ecosystem productivity and vegetation dynamics (Brussaard et al., 2001). Individual species have been shown to affect a variety of components of the carbon (C) and nutrient cycles, including pools of aboveground and belowground C, nitrogen (N), and other elements; net primary productivity and plant growth rates; chemical quality and rates of litter fall; and nutrient and C mineralization rates (Tilman et al., 1997; Hooper and Vitousek 1998; van Breeman 1998; Hector et al., others 1999; Chapin et al., 2000). This body of evidence strongly suggests that when the species composition of a community changes because of the invasion and spread of an exotic species, there are likely to be consequent changes in nutrient cycling processes.

The state of Florida has one of the most severe invasive exotic species problems in the United States. According to Wunderlin (1998), non-native species make up 33% of all the plants found growing out of cultivation in Florida, and the
Florida Exotic Pest Plant Council (FLEPPC) considers 152 of these naturalized plants to be invasive to some degree as they are altering native plant communities by displacing native plant species, changing community structures or ecological functions or hybridizing with native plant species (Langeland and Hutchinson 2013).

1.2 Study Species

*Lygodium microphyllum* (Cav.) R. Br., native to Asia and Australia (Fig. 1.1) invades many freshwater and moist habitats in Florida, and is common in cypress swamp, pine flatlands, wet prairies, sawgrass marshes, mangrove communities, and Everglades tree islands (Pemberton and Ferriter, 1998). *Lygodium microphyllum* with its ability to form dense mats, spreads very rapidly and dominates both understory and overstory native wetland habitats. It has the ability to grow in varying hydrological (Gandiaga et al., 2009) and light gradients (Pemberton et al., 2002). Fire, which is not a very effective method (Maithana et al., 1986), is the most commonly used technique to control *L. microphyllum*. It is reported that has the ability to alter the fire ecology in the Everglades, prescribed burning which normally stop at the margins of flooded cypress sloughs, but in *L. microphyllum* infested areas enters the tree canopy through the mats of *L. microphyllum* (Ferriter et al., 2005) thus causing the loss of some canopy trees as well as a loss of native epiphytes and bromeliads residing on tree trunk (Roberts, 1996). *Lygodium microphyllum* spreads quickly in the South Florida’s landscape because of its ability to reproduce through the three mating systems possible in a fern: intra- and inter-gametophytic selfing and out-crossing (Lott et al., 2003). Spores can germinate in six to
seven days (Brown, 1984). *Lygodium microphyllum* is now spreading rapidly throughout Southern Florida (Fig. 1.2). By 1997 it had covered 15,800 hectares and by 1999 it spread to more than 43,000 hectares (Pemberton and Ferriter, 1998; Ferriter et al., 2005). According to a model developed by Volin et al., (2004), the landscape coverage of *L. microphyllum* infestations could exceed the current combined coverage of the top five invasive species by 2014. Volin et al., (2010) have suggested that the belowground microbial community can influence the establishment of *L. microphyllum* in south Florida and contribute to its invasiveness. Thus the role of belowground ecology in invasions by exotic plants cannot be overlooked and need to be addressed to achieve sustainable management of exotic invasive species in the Everglades.

1.3 Significance of the Study

The current management techniques: fire, herbicides, and mechanical removal, are inadequate and the prospect of developing a method that targets the rhizomes of *L. microphyllum* is compelling; however, the soil biogeochemical characteristics of *L. microphyllum* infested sites remain unexplored. Before we target the rhizomes, it is imperative to elucidate the soil characteristics of the sites (both infested and uninfested) to obtain baseline information. Additionally, it is also important to gain information of the soil characteristics of the native range where this plant has its origin and adapted habitat. Thus, I did a detailed cross continental study on the soil biogeochemical properties of the recipient habitat in Florida and native range in Australia. The results from my study will assist in better understanding of the complex feedbacks between exotic invasive plants, soil microbial community and soil elements. It will further provide
opportunities to land managers and researchers to develop a successful integrated management technique.
Figure 0.1 Native, introduced, and predicted distribution of *Lygodium microphyllum* (Adapted from Volin et al., 2009, with permission from Springer).
Figure 0.2 Increase in the distribution of *Lygodium microphyllum* in Florida from 1993 to 2010, data source: Florida Exotic Pest Plant Council (FLEPPC).

1.4 Dissertation Outline

My study combined two experimental approaches: i) cross continental field study to compare the growth environment faced by *L. microphyllum* in its native range in Australia and the recipient community in Florida; ii) growth rate experiment conducted in a greenhouse and a growth chamber to examine phenotypic plasticity and growth response of *L. microphyllum* with various soil treatments. The dissertation is organized in six chapters. The first chapter provides a basic introduction to the problem and focus of the study. Chapter 2, 3, 4 and 5 are independent chapters addressing the interaction of *L. microphyllum* with soil chemical, biological and physical parameters. Finally, chapter 6 presents the major findings of the dissertation presented in the previous four chapters.
Additionally it includes the conclusion of the study, directions for future research, and recommendations towards a development of an integrated management technique to control *L. microphyllum* in the invaded regions of Florida.

In chapter two, analysis of field root and soil samples from the ferns introduced and native range as well as a seven-week growth chamber experiment were done to determine the level of mycorrhizal colonization in the roots of *L. microphyllum* and the dependency on mycorrhizal fungi for growth and phosphorus (P) uptake. The field root samples showed that *L. microphyllum* was heavily colonized by arbuscular mycorrhizal fungi (AMF) in relatively dry conditions, which are commonly found in some Florida sites compared to more common wet sites where the fern is found in its native Australia. The results from the growth chamber experiment showed that the mycorrhizal treatment plants had significantly higher relative growth rate and biomass compared to the non-mycorrhizal plants. Similarly, *L. microphyllum* was highly dependent on the mycorrhizal fungi for growth and P uptake. Chapter two highlights the role of AMF in the vegetative reproduction and enhanced invasive success of *L. microphyllum* in south Florida natural areas.

Chapter three investigates the effects of soil pH on the growth, nutrient uptake and degree of mycorrhizal colonization of *L. microphyllum*. I conducted a 60 day greenhouse experiment by growing this plant in pots filled with pH adjusted soils to a range of 4.8 to 8.0. *Lygodium microphyllum* was able to survive and grow at all soil pH levels; however, final biomass, relative growth rate, photosynthesis, and specific leaf area were all significantly greater in soil pH 5.5 - 6.5 compared to the other treatments. Correspondingly, leaf nitrogen concentration was also had a significant influence on
these four plant parameters. Root colonization by mycorrhizal fungi was significantly higher in soil pH 5.5-7.5 than in lower or higher pH soil, and was significantly correlated with plant growth parameters as well as elemental concentration in the leaves. In its native Australia, *L. microphyllum* responds robustly following fire. Fire is also known to commonly raise soil pH, and given the treatment response to soil pH in this study and the plant’s known fire tolerance in its home range, this management option should likely be reconsidered.

In chapter four, I compared the soil characteristics six invaded and adjacent uninvaded plots in three different locations. The results from this study show that the fern can grow and thrive in a wide range of soil types and the impact on the soil was site specific with effects being more prominent in sites with low nutrient status. Additionally, there was significant difference in the soil nutrient status in the invaded and uninvaded sites. Sites with Old World climbing fern had significantly higher nutrient concentration with the corresponding differences in the soil organic matter. Overall my study highlights that this exotic pest plant can alter its belowground environment to its own benefit by enhancing the soil nutrient status with the added soil organic matter.

In chapter five, I conducted a cross continent comparison of soil characteristics associated with *L. microphyllum*. Here, I present evidence that the invasion by *L. microphyllum* in south Florida is not only facilitated by the soil microbial community but also by the soil chemical characteristics. My results indicate that aluminum, which is considered phytotoxic in acidic soil condition, was significantly higher in Australian soils compared to the Florida soils. I suggest that invasive plants not only escape from their natural herbivores but also the toxic soil environment in their native habitats and
conclude that a successful management technique and the future invasion prediction model should consider the soil elemental status.

Finally in chapter six, the major findings of the dissertation presented in the previous four chapters are presented. Along with discussion on the conclusions, recommendations are presented for the development of a successful integrated management technique for *L. microphyllum* in Florida and the restoration of the previously invaded habitats.

**References**


MYCORRHIZAL SYMBIOSIS AND LYGODIUM MICROPHYLLUM INVASION IN SOUTH FLORIDA – A BIOGEOGRAPHIC COMPARISON


Abstract

*Lygodium microphyllum* (Old World climbing fern) is one of the most problematic weeds in south Florida, invading numerous habitats from mangroves to pine flatwoods natural ecosystems. Much of the research efforts on *L. microphyllum* has been focused on reproductive potential, spore release, growth under different environmental conditions, belowground rhizome dormancy and survival strategies that describes its invasiveness. However, the role of an important mutualistic association with arbuscular mycorrhizal fungi (AMF) in the competitive ability and successful invasion of *L. microphyllum* by enhancing nutrient uptake has not been previously considered. Analysis of field root and soil samples from the ferns introduced and native range as well as a seven-week growth chamber experiment were done to determine the level of mycorrhizal colonization in the roots of *L. microphyllum* and the dependency on mycorrhizal fungi for growth and phosphorus (P) uptake. The field root samples showed that *L. microphyllum* was heavily colonized by AMF in relatively dry conditions, which are commonly found on some Florida sites compared to wet sites where the fern is found in its native Australia. The results from the growth chamber experiment showed that the mycorrhizal treatment plants had significantly higher relative growth rate and biomass compared to the non-mycorrhizal plants. Similarly, *L. microphyllum* was highly dependent on the
mycorrhizal fungi for growth and P uptake. My results suggest that AMF play a significant role in vegetative reproduction and likely enhance the invasiveness of *L. microphyllum* in south Florida natural areas.

Key words: arbuscular mycorrhizal fungi (AMF), inorganic phosphorus, relative growth rate (RGR), mycorrhizal dependency, exotic pest plant.

2.1 Introduction

*Lygodium microphyllum* (Old World Climbing Fern) is one of the most problematic weeds in south Florida. It invades many freshwater and moist habitats and is common in cypress swamp, pine flatlands, wet prairies, sawgrass marshes, mangrove communities, and the Everglades tree islands (Pemberton and Ferriter, 1998). *L. microphyllum*, with its ability to form dense mats, spreads very rapidly and dominates both understory and overstory native wetland habitats. It has the ability to grow in varying hydrological (Gandiaga et al. 2009), nutrient (Volin et al. 2010) and light gradients (Volin et al. 2004). It is estimated to occupy 183,080 acres across the South/Central Florida region (Ferriter and Pernas 2006) and a model developed by Volin et al. (2004) shows that, in the absence of aggressive control measures, *L. microphyllum*'s infestations could exceed the current combined coverage of the top five most invasive species in Florida by 2014. Managing *L. microphyllum* has been a significant challenge for land resource managers and researchers because of its extensive rapid invasion in natural areas of south Florida. Much of the research work previously performed on *L. microphyllum* focuses on reproductive potential, spore release, belowground rhizome
dormancy and survival strategies investigating its invasiveness (Lott et al. 2003). Volin et al. (2010) have suggested that the belowground microbial community can influence the establishment of *L. microphyllum* in south Florida and contribute to its invasiveness. Thus the role of belowground biota in invasions by exotic plants cannot be overlooked, in particular, the role of arbuscular mycorrhizal fungi (AMF) deserves consideration.

Most vascular plants form symbiotic associations with AMF, and many plants are highly dependent on this association for their growth and survival (Smith and Read 1997). Arbuscular mycorrhizal fungi are obligate symbionts of plants; approximately 95% of all vascular plants can form AMF associations (Fitter and Moyersoen 1996). Read (1991) stated that the mycorrhizal association is the most ubiquitous and abundant form of terrestrial symbiosis, and AMF are considered the most common type of mycorrhizae which dominates grasslands, croplands, tropical forests, and desert communities. They occur naturally in most soils and their important ecosystem function is to assist in the acquisition of soil mineral nutrients (Dighton 2003). Arbuscular mycorrhizal fungi are known to benefit plants by improving plant phosphorus (P) uptake (Fitter, 1990; Gao et al. 2007) and also potentially enhance defense against soil born pathogens (Azcón-Aguilar and Barea 1997). South Florida soils are poor in P because of the binding of P with Ca in alkaline soils and to certain extent Al or Fe in acidic soils. Arbuscular mycorrhizal fungi can facilitate P uptake by increasing 1) diffusion rate into plant roots; 2) P concentration at the root surface; and 3) the rate of P dissociation from the surface of soil particles (Bolan 1991). Elements other than P, such as N, Cu and Zn, also experience enhanced uptake through AMF (Gildon and Tinker 1983; Gao et al. 2007). It has been estimated that external hyphae of AMF can contribute up to 80% of the
P, 10% of the N, 10% of the K, 25% of the Zn, and 60% of the Cu absorbed by plants (Li et al. 1991; Marschner and Dell 1994; DeLuca et al. 2002). Mycorrhizal fungi help overcome the nutrient deficiency by extending their external hyphae to areas of soil beyond the depletion zone and increasing the absorptive surface of the root. However, Smith and Read (1997) have reported that host plant species do not equally benefit from AMF and some plants acquire more nutrients from AMF than others. Furthermore if the symbiotic relationship is non-host specific competing plant species could be interconnected by AMF hyphal networks (Grime et al. 1987; Newman 1988) thus creating an imbalance in the nutrient distribution in the soil. This imbalance in nutrient distribution could interfere the competitive interaction between native and exotic species by promoting growth of the invasive species and inhibiting growth of the native plant species (Fumanal et al. 2006; Callaway et al. 2008).

Populations of AMF are highly influenced by the various environmental factors including climatic conditions, soil physico-chemical properties, age of the host plant species etc. In southeastern Queensland, Australia, the native range of *L. microphyllum* is climatically similar to the *L. microphyllum* invaded areas of south Florida (Volin et al. 2010) however the soil physico-chemical properties are significantly different in these two locations. Thus in my study, I characterized the root colonization by AMF in *L. microphyllum* under field conditions in both its native Australia and its introduced environment in Florida. I further explored the influence of AMF on the growth and biomass allocation strategy of *L. microphyllum* in growth chambers. The objectives of this study were to: 1) evaluate the mycorrhizal status of natural populations of *L. microphyllum* in both Australia and Florida; 2) determine the effect of AMF on the
reproductive and biomass allocation strategy of *L. microphyllum*; and, 3) evaluate the dependency of *L. microphyllum* on AMF for growth and phosphorus uptake. I hypothesized that the south Florida population of *L. microphyllum* would have a higher degree of mycorrhizal colonization compared to the Australian population because of the difference in soil characteristics. I also hypothesized that *L. microphyllum* is highly dependent on AMF for increased biomass accumulation and P uptake.

### 2.2 Methods

#### 2.2.1 Experiment 1: Degree of mycorrhizal colonization in *L. microphyllum*

Roots and rhizosphere soil samples (for nutrient analysis and spore extraction) of wild *L. microphyllum* were collected from different locations in south Florida and Australia to assess the mycorrhizal fungal root colonization and presence of AMF spores. To estimate the AMF colonization, root samples were collected from fully grown adult plants from populations that were known to be at least five years old in both Australia and south Florida. The locations, sampling date and the dominant vegetation in each of the sites are given in Table 1. Root tip samples were placed in 70% ethanol immediately until further processing following the method described in McGonigle et al. (1990). At least 10 cm of fine roots from multiple plants were collected at seven different locations. These roots were cut into 1.5 cm fragments, cleared in 15% KOH at 70°C for 4 hours, rinsed twice with water, bleached with ammoniated H₂O₂, and acidified with 1 N HCl. Once the roots were cleared, staining was done using 0.05% Trypan blue in acidic glycerol at 80°C for 20 minutes. The stained roots were examined with a dissecting microscope at 30–60 X magnification; the portions that showed the presence of
mycorrhizal fungi were mounted on slides in lactic acid and further examined at 100–400 X magnification to analyze the presence of mycorrhizal structures (hyphae, vesicles, and arbuscules). At least 50 root fragments were selected randomly for each site and the percentage of colonization was estimated.

The wet sieving and decanting technique was used to enumerate the mycorrhizal spores in the soil (Gerdemann et al. 1963). The soil samples were mixed to homogeneity. Fifty grams of the soil sample were then mixed with water and passed through a series of sieves allowing heavy soil particles to settle for a few seconds. The sievate retained on the sieves was washed and centrifuged with water to remove floating organic debris and the supernatant was discarded. The pellet in the bottom was re-suspended in a 50% sucrose solution, and centrifuged for one minute at 2000 RPM to separate the spores from denser soil components. Immediately after centrifugation, spores in the sucrose supernatant were rinsed in a fine sieve to remove the sucrose. The spores were then washed into a filter paper for vacuum filtration. The spores on the filter paper were counted under a stereo microscope.

Analysis of soil properties

At each sampling site, six 1 m x 1 m plots were selected randomly and soil sample was collected from the 10-15 cm deep zone at all four corners and the center of each plot with a soil corer (core diameter: 18 mm) and mixed homogeneously into one bulk sample for each plot. The samples were collected to a depth of 10-15 cm. The soil samples from south Florida were transported to the laboratory in a cooler; the Australian samples were stored at 4°C and were shipped overnight. The soil samples for the
chemical and physical properties were air dried and passed through a 2 mm sieve. They were then ground to fine powder with a mortar and pestle, and stored at room temperature in air-tight containers for further analysis of nutrients. The soil pH was measured with a pH meter, (soil solution ratio 1:1 in water), texture was measured by the hydrometer method, percentage of carbon and nitrogen was measured with a TruSpec Carbon/Nitrogen Analyzer (Leco Corporation, USA), total organic matter was measured based on the standard loss on ignition method (500°C, 5 hours; Storer 1984), for the measurement of total P, soil samples (0.25 grams, finely ground) were ashed (500 °C), digested in 2 ml HCL (6N) and 10 ml HNO₃, and then analyzed with an UV spectrophotometer (Shimadzu Scientific Instruments, ) (total P and some other variables were not measured for samples collected in 2006).

2.2.2 Experiment 2: Mycorrhizal dependency of L. microphyllum

Plant Material

Experimental plants were grown from spores collected from an infestation in Jonathan Dickinson State Park, Florida, following the method used by Lott et al. (2003). The spores of L. microphyllum were disinfected with 1% bleach and transferred to Petri dishes that contained Parker-Thomson Medium. The plates were placed in an incubator set at 25-27°C for ten weeks and were watered with sterile DI water every week. After ten weeks, individual gametophytes were transferred to fresh Petri dishes. When the sporelings’ roots and leaves developed, 50 plants were transplanted to small pots previously filled with sterile sand. These 50 plants were placed in a growth chamber for approximately four weeks. The plants were kept very moist, and were watered with half
strength Hoagland’s nutrient solution as needed. Plants were then transferred to 2.5 L pots filled with the top soil collected from a *L. microphyllum* infested site located in the Tree Tops County Park, Davie, Florida. The potting soil was sterilized in an autoclave to kill mycorrhizal fungal spores to ensure the experimental plants remained free of mycorrhizal fungi.

**Growth Chamber Experiment**

A seven-week growth chamber (Percival Scientific, with irradiance = 500 \( \mu \)mol m\(^{-2}\) s\(^{-1}\), photoperiod = 12 h and temperature 27°C) experiment was done to determine the mycorrhizal dependency of *L. microphyllum*. The experiment consisted of two treatments: mycorrhizal treatment and non-mycorrhizal treatment with eight replicate pots per treatment. In the mycorrhizal treatment plants received mycorrhizal inoculum; the top soil collected from the field directly under *L. microphyllum* while a systemic fungicide was added every three weeks in the non-mycorrhizal treatment plants to prevent any kind of mycorrhizal contamination during the experiment. The non-mycorrhizal plants received 50 ml of microbial wash to provide similar microflora except the mycorrhizal fungi. The microbial wash was prepared by filtering the field soil slurry through a 25\( \mu \)m filter paper, which removed the mycorrhizal fungi spores in the soil but allowed the other soil microorganisms to pass through (Johnson 1993). Three hundred mg of the systemic fungicide Benomyl was applied in 100 ml of water per pot (50 mg/kg growth medium); this fungicide is reported to effectively reduce the mycorrhizal colonization in roots without significant impact on the plants (Fitter and Nichols 1988; Hetrick et al. 1992). Plants were watered to saturation once per week and received 250 ml
of half strength Hoagland's solution weekly, modified by the addition of phosphorus as inositol hexaphosphate (Marler et al. 1999). This form of phosphorus is not directly available to plants for uptake, and requires alteration in the soil by mycorrhizal fungi, soil microbes, or root exudates (DeLucia et al. 1997).

**Measurements**

Two harvests were conducted during this study: at time 0 (the transplanting day), and after 50 days. The allometric relationship between stem length and total mass \((R^2 = 0.87)\) from the time 0 harvest was developed to estimate the initial plant mass of the experimental plants and to calculate the relative growth rate (RGR) (see Gandiaga et al. 2009). The RGR \((\text{mg g}^{-1} \text{ d}^{-1})\) was calculated for individual plants used for the experiment, where \(RGR = [\ln (\text{final dry mass})-\ln (\text{initial dry mass})]/\text{days}\) (Evans, 1972). After each harvest, roots, stem, and leaves (pinnae) were separated from each plant and the leaf area was measured with the leaf area meter to calculate the specific leaf area (SLA). The separated plant parts were oven-dried (at 70°C) to constant mass and weighed to determine the leaf mass ratio (LMR), stem mass ratio (SMR), rhizome mass ratio (RhiMR), and root mass ratio (RMR); differences in the mean growth parameters between the treatments and relative growth rate (RGR). The roots were washed with water. Twenty 1-cm root pieces were collected from each plant before drying to quantify the AMF colonization in the roots. Root and shoot dry mass were measured after oven-drying for one week at 65°C. Leaf samples (0.25 grams, finely ground) were ashed (500°C), digested in 2 ml HCL (6N) and 10 ml HNO₃, and then analyzed with an UV spectrophotometer (Shimadzu Scientific Instruments, US) for total phosphorus (P).
concentration. The percentage of carbon and nitrogen in the leaves was measured with a TruSpec Carbon/Nitrogen Analyzer (Leco Corporation, US) and the C/N ratio was calculated. Dependency of shoot P uptake and growth of plants on AMF was calculated using the formulae from Plenchette et al. (1989), where +M represents inoculated plants and –M, fungicide treated plants:

Dependency of P uptake = \( \frac{P\text{ content (+M)} - P\text{ content (-M)}}{P\text{ content (+M)}} \times 100 \)

Dependency of growth = \( \frac{\text{Total dry mass (+M)} - \text{Total dry mass (-M)}}{\text{Total dry mass (+M)}} \times 100 \)
2.2.3 Statistical Analyses

Data for the soil and mycorrhizal colonization (Experiment 1) were analyzed with one-way analysis of variance (ANOVA) to compare the means of the pH, total P, total N and soil organic matter from different sites. Correlation analysis determined the association of soil pH and organic matter on the degree of mycorrhizal colonization.

For the greenhouse experiment (Experiment 2), after the harvest at 50 days, regression analysis examined the influence of initial plant mass on RGR and its morphological, allocational and physiological determinants as it has been reported frequently (e.g., Mcconnoughay and Coleman 1999; Volin et al. 2002; Kruger & Volin 2006). Regression analysis indicated that RGR was negatively correlated to initial plant mass (p<0.001). Additionally, RhMR and SLA at final harvest were all significantly related (P < 0.05) to final plant mass. Therefore, each was normalized for variation in plant mass using analysis of covariance. All of the variables in mycorrhizal and non-mycorrhizal treatments were then compared with two-treatment t-test for significance at p ≤ 0.05. Regression analysis was also used to assess relationships between RGR and its principal determinants. All the parameters were analyzed with SAS Version 9.2 software.

2.3 Results

2.3.1 Degree of mycorrhizal colonization and influence of soil factors

A wide variety of different fungal structures such as extraradical hyphae, vesicles, and arbuscules were visible in root samples from all sites. The total percentage of roots colonized by arbuscular mycorrhizal fungi in the south Florida plants in some cases were up to three times that of the Australian plants (Table 1.1). In Florida, the range was from
31-79%, while in Australia the range was much narrower, 27-29%. Mycorrhizal spores were present in the rhizosphere soil samples from all sites; spore abundance was highest in Tree Tops Park in south Florida, while the lowest abundance of spores at Nudgee in Australia.

Tree Tops Park, the south Florida site had slightly higher total P in the soil but this difference was not significantly different (Fig. 1.1a). However there was a significant difference in the total C and N among the different sites (Fig. 1.1 a, b). Tree Tops Park in Florida and Logan in Australia had significantly higher percentage of C and N compared to the other sites. The south Florida soil samples were slightly acidic, ranging from 5.49 at Tree Tops Park to 6.22 at Jonathan Dickinson Park. However, the Australian soil samples were highly acidic ranging from 3.97 at Nudgee to 4.7 at Logan. Soil organic matter was significantly higher in the Tree Tops and Logan than at Jonathan Dickinson, Nudgee and Daintree (Table 2.1).
Figure 0.1 Means (± SE) of soil nutrients: a) total soil phosphorus (mg g⁻¹); b) total soil nitrogen %; c) total soil carbon (%).
Figure 0.2 Arbuscular mycorrhizal fungal structures (a), vesicles and fungal hyphae (b), in root cortex region of South Florida Plants, vesicles and fungal hyphae in the root cortex region of Australian plants (c).
Table 0.1 Sampling site information with the number of AMF spores and mycorrhizal colonization.

<table>
<thead>
<tr>
<th>Site</th>
<th>Coordinates</th>
<th>Dominant vegetation</th>
<th>Soil texture</th>
<th>pH</th>
<th>SOM % ± standard error</th>
<th>No. of spores/10g dry soil</th>
<th>Sampling Date</th>
<th>Deg. of colonization (%) [min–max]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tree Tops Park, FL, US</td>
<td>26° 4'0.04&quot;N, 80° 16' 5.88&quot;W</td>
<td>Royal fern</td>
<td>Sandy loam</td>
<td>5.56</td>
<td>39.7±1.6</td>
<td>29±7</td>
<td>Dec, 2010</td>
<td>79 ± 3.0 [65 – 85]</td>
</tr>
<tr>
<td>Jonathan Dickinson, FL, US</td>
<td>27°0'37.33&quot;N, 80°7'20.28&quot;W</td>
<td>Slush pine</td>
<td>Sand</td>
<td>6.02</td>
<td>4.30±0.9</td>
<td>19±5</td>
<td>Dec, 2010</td>
<td>74 ± 2.2 [67 – 88]</td>
</tr>
<tr>
<td>Big Cypress Seminole Indian Reservation, FL, US</td>
<td>Approximately, 26°17'N, 80°54'W</td>
<td>Bald cypress</td>
<td>Sand</td>
<td>4.99</td>
<td>*</td>
<td>*</td>
<td>October, 2006</td>
<td>31 ± 7.8 [16.0 – 49.5]</td>
</tr>
<tr>
<td>Daintree Ferry, Queensland, AU</td>
<td>16°15'25.57&quot;S, 145°24'3.94&quot;E</td>
<td>Drynaria</td>
<td>Silt loam</td>
<td>4.43</td>
<td>8.07±1.2</td>
<td>12±6</td>
<td>June, 2011</td>
<td>27 ± 1.4 [24 – 33]</td>
</tr>
<tr>
<td>Logan Reserve, Queensland, AU</td>
<td>27°40'4.16&quot;S, 153°16'0.44&quot;E</td>
<td>Bungwall fern</td>
<td>Sandy clay loam</td>
<td>4.55</td>
<td>35.5±2.9</td>
<td>15±3</td>
<td>June, 2011</td>
<td>28 ± 1.3 [25 – 32]</td>
</tr>
<tr>
<td>Nudgee, Queensland, AU</td>
<td>27°22'31.12&quot;S, 153° 5'39.42&quot;E</td>
<td>Melaleuca</td>
<td>Loam</td>
<td>4.01</td>
<td>11.45±1.9</td>
<td>10±5</td>
<td>June, 2011</td>
<td>27 ± 0.6 [25 – 29]</td>
</tr>
<tr>
<td>SE Brisbane, Queensland, AU</td>
<td>27°40.36'S, 153°16.60'E</td>
<td>Melaleuca</td>
<td>Sandy loam</td>
<td>5.55</td>
<td>6.83</td>
<td>*</td>
<td>February, 2006</td>
<td>24.0 ± 2.7 (3) [19.5 – 28.9]</td>
</tr>
<tr>
<td>Near Amity Point, Stradbroke Island, Queensland, AU</td>
<td>Approximately, 27.4°S, 153.4°E</td>
<td>Melaleuca</td>
<td>Sand</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>February, 2006</td>
<td>30.2 ± 6.5 (5) [7.4 – 44.7]</td>
</tr>
</tbody>
</table>

Note: Mean values ± standard error, Soil organic matter (SOM), number of samples (N), * values not determined.
Correlation analysis shows that the degree of colonization was significantly positively correlated with soil pH ($r = 0.86$, $p<0.001$); N% ($r= 0.45$, $p=0.026$); and C ($r = 0.43$, $p = 0.0177$) while there was no significant correlation with soil organic matter ($p = 0.02$, $p=0.35$).

2.3.2 Mycorrhizal dependency

As expected, in the non-mycorrhizal treatment, benomyl application significantly suppressed the mycorrhizal colonization of *L. microphyllum* roots. After 50 days, colonization rate was high in the mycorrhizal treatment (>75%), but low (<5%) in the non-mycorrhizal treatment (Fig. 2.2). Different fungal structures were observed in the plant roots including hyphae, vesicles as well as arbuscules.

![Figure 2.3](image)

Figure 2.3 High colonization of the roots by arbuscular mycorrhizal fungi in the mycorrhizal treatment plants (a), no colonization in the non-mycorrhizal treatment (b), at the end of 50 days.
The mycorrhizal plants had greater total P uptake in shoots compared to the non-mycorrhizal plants. The mycorrhizal dependency index for growth was 67%, and 64% for P uptake. Relative growth rate of mycorrhizal treated plants was 29% greater (P=0.01) than the RGR of the non-mycorrhizal plants (Table 2). Correspondingly, the mean biomass of the *L. microphyllum* plants inoculated with mycorrhizal fungi was also significantly greater (by 66%) (P=0.001) than the non-mycorrhizal plants (Table 2).

There was no significant difference in SMR (P=0.15) or LMR (P=0.99) between the two treatments (Table 2.2). On the other hand, allocation to rhizomes (RhiMR) (P=0.001) was significantly different between treatments, resulting in greater rhizome allocation for mycorrhizal plants compared to untreated plants. In contrast, mycorrhizal plants tended to allocate less to roots than non-mycorrhizal plants (P=0.08).

The mycorrhizal treatment increased the leaf area of the plants nearly four-fold (P=0.0001) and SLA was significantly greater (P=0.003) (24%) in the presence of mycorrhizal fungi compared to non-mycorrhizal plants. There was no significant difference (P=0.83) in the average P concentration in the leaves of the mycorrhizal and non-mycorrhizal plants. However, the total P per plant (3,291 µg) of the mycorrhizal plants was significantly higher (P=0.0001) than the non-mycorrhizal plants.
Table 0.2 Effect of arbuscular mycorrhizal fungi (AMF) inoculation on plant growth parameters and on leaf content of biolimiting elements (P and N).

<table>
<thead>
<tr>
<th>Variable</th>
<th>Mycorrhizal (g)</th>
<th>Non-mycorrhizal (g)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Final biomass(g)</td>
<td>1.99 ± 0.37</td>
<td>0.68 ± 0.17</td>
<td>0.0001</td>
</tr>
<tr>
<td>RGR mg g⁻¹ day⁻¹</td>
<td>75.13 ± 0.01</td>
<td>53.88 ± 0.01</td>
<td>0.000</td>
</tr>
<tr>
<td>SMR</td>
<td>0.17 ± 0.03</td>
<td>0.20 ± 0.05</td>
<td>0.15</td>
</tr>
<tr>
<td>RMR</td>
<td>0.18 ± 0.05</td>
<td>0.24 ± 0.08</td>
<td>0.08</td>
</tr>
<tr>
<td>RhiMR</td>
<td>0.24 ± 0.04</td>
<td>0.14 ± 0.04</td>
<td>0.001</td>
</tr>
<tr>
<td>LMR</td>
<td>0.42 ± 0.02</td>
<td>0.42 ± 0.02</td>
<td>0.99</td>
</tr>
<tr>
<td>SLA</td>
<td>505 ± 64</td>
<td>383.6 ± 85</td>
<td>0.003</td>
</tr>
<tr>
<td>Leaf area (cm²)</td>
<td>414 ± 112</td>
<td>109.7 ± 40.7</td>
<td>0.0001</td>
</tr>
<tr>
<td>Total P per plant (µg)</td>
<td>3,291 ± 615</td>
<td>1,129 ± 279</td>
<td>0.0001</td>
</tr>
<tr>
<td>C/N ratio</td>
<td>1.7 ± 0.1</td>
<td>1.6 ± 0.1</td>
<td>0.21</td>
</tr>
</tbody>
</table>

Note: mean values (± SD) of the study variables: relative growth rate (RGR), stem mass ratio (SMR), root mass ratio (RMR), rhizome mass ratio (RhiMR), leaf mass ratio (LMR), specific leaf area (SLA) after growing 7 weeks in a growth chamber: analyzed by (two sample t test, p<0.05).

2.4 Discussion and Conclusion

The results of this study suggest that *L. micorphyllum* likely has a strong symbiotic relationship with mycorrhizal fungi, and the degree of mycorrhizal colonization is generally higher in the invaded regions of south Florida than in the plant’s native range in Australia. Detailed information on the mycorrhizal status of the coexisting species is not available but the high degree of mycorrhizal colonization may assist in
absorption and competition for nutrients in the fern’s introduced environment, especially on sandy sites with low water holding capacity. This could in turn provide a competitive advantage over native Florida plants. However, others have found that that many invasive plants do not associate with mycorrhizal fungi (see Pringle et al. 2009). Plants such as Alliaria petiolata, Centaurea diffusa, etc. have been found to use alternative mechanisms to disrupt existing symbiotic relationships by secreting lethal biochemicals in the introduced range, resulting in the reduced growth and competitiveness of native plants (Callway and Aschehoug 2000; Callaway et al. 2008).

My results show that soil pH is positively correlated with mycorrhizal colonization, indicating that the low soil pH of the Australian soil may influence the degree of mycorrhizal colonization and the number of spores associated with L. microphyllum, other research have shown species-specific responses of AMF to soil pH (Wang et al. 1985; Porter et al. 1987; Gemma et al. 1989). It is widely reported that abundance of mycorrhizal fungi declines in response to N and P fertilization (see Treseder 2004). In contrast, My results indicate a positive correlation between mycorrhizal colonization and soil nitrogen. Similar results have also been reported by (Persson & Ahlstrom 1991; Heijne et al. 1992). I did not determine the mycorrhizal fungal species in my study but there is substantial variation in the environmental effects on different mycorrhizal fungal species, this variation could explain in part the contradictory results seen in my study. Likewise, there was a significantly positive correlation between the root mycorrhizal status and soil C%, this result supports the existing assumption that mycorrhizal fungi have a significant contribution to soil carbon storage (Treseder and Allen 2000).
Interestingly, the two sites in Florida with high AMF colonization appear to be drier than the four Australian sites and one Florida Big Cypress Seminole Indian Reservation site. These latter sites are characterized by periodic inundation that may or may not occur on an annual basis, while the two dry Florida sites are not inundated for any appreciable amount of time. Relationship between flooding and AMF colonization seen in my study could explain in part the lowered growth rate of *L. microphyllum* in flooded conditions compared to the drought and field conditions seen by Gandiaga et al. (2009). Additionally, Rickerl et al. (1994); Stevens and Peterson (1996); and Miller (2000) have found a strong relationship between mychorrizal colonization and site hydrology, and this potential relationship for *L. microphyllum* needs to be explored further.

We found that *L. microphyllum* can attain high RGR under suitable environmental conditions. A high RGR for invasive species has been reported for many different species (Burns 2004; James and Drenovsky 2007; Soti and Volin 2010). From an ecological perspective, high RGR can lead to the rapid occupation of a large space, which could be advantageous for exotic invasive plants (Grime and Hunt 1975). In my study, RGR in *L. microphyllum* was highly enhanced by the presence of mycorrhizal fungi. Mycorrhizal plants produced almost three times more biomass than non-mycorrhizal plants. Increased growth and development in mycorrhizal plants compared to non-mycorrhizal plants has also been found in several different species (Gupta and Janardhanan 1991; Smith and Read 1997; Guadarrama et al. 2004; Liu et al. 2005; Pezzani et al. 2006). On the other hand, Philip et al. (2001) observed that colonization by AMF of *Lythrum salicaria* decreased plant biomass both aboveground and belowground. Likewise Botham et al.
(2009) observed that the AMF inoculated *Fragaria virginiana* plants showed no difference in biomass accumulation and growth rate compared to control plants.

The mycorrhizal treatment plants had significantly higher SLA compared to the non-mycorrhizal plants. The difference in the SLA between the two treatments could lead to higher RGR in the mycorrhizal plants. Although we did not measure photosynthesis in our study, Gandiaga et al. (2009) found that SLA together with photosynthesis were the major determinants of growth in *L. microphyllum* plants grown under different hydrological conditions. Other studies, using different species, have also found that mycorrhizal plants had higher leaf area, leaf area ratio and SLA compared with control plants (Waschkies et al. 1994; Caglar and Bayram, 2006; Vega-Frutis et al. 2011).

The enhanced ability of a plant to take up phosphorus from low P soils is considered to be the major contributing factor for mycorrhizal dependency (Hall 1975; Smith & Read 1997). Increased P uptake by the extraradical mycelia of mycorrhizal fungi in the roots may allow *L. microphyllum* to absorb more nutrients leading to larger shoots and more extensive roots, compared to non-mycorrhizal plants. This relationship would potentially convey a competitive growth advantage in the fern’s introduced range in Florida as this region is conspicuous for its P-limiting growth environment (McCormic et al. 1999). Previous research has shown that AMF increase plant uptake of phosphate (Bolan 1991), micronutrients (Burkert and Robson 1994), nitrogen (Barea et al. 1991), and act as antagonists against some plant pathogens (Duponnois et al. 2005). Moreover, it has been demonstrated that plants inoculated with AMF utilize more soluble phosphate from rock phosphate than non-inoculated plants (Antunes and Cardoso 1991). The
current study supports these results since the mean P uptake per plant was significantly greater in the mycorrhizal plants than in the non-mycorrhizal plants.

In conclusion, it is clear that *L. microphyllum* can form a very strong symbiotic relationship with AMF in its introduced environment in Florida. It is likely that this relationship is strongly influenced by site hydrological conditions, but this hypothesis will need to be tested in future research, especially when the Florida Everglades is undergoing a major hydrological shift as an effort for restoration. The enhanced mycorrhizal fungi are also likely responsible for the greater P uptake and biomass accumulation in the control study. Symbiotic relationships such as found in my study, are highly beneficial, and likely enhance the aggressive growth characteristic of this exotic pest plant in its *de novo* environment. Further field experiments are necessary to better evaluate the potential role of mycorrhizal fungi in the growth of this highly invasive species in south Florida natural areas.

**Acknowledgements**

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EFFECT OF SOIL pH ON GROWTH, NUTRIENT UPTAKE, AND MYCORRHIZAL COLONIZATION IN EXOTIC INVASIVE *LYGODIUM MICROPHYLLUM*


Abstract

*Lygodium microphyllum* is an invasive exotic plant species taking over many sites in freshwater and moist habitats in Florida. Managing it has been a significant challenge for land resource managers and researchers because of its extensive rapid invasion. To assess the effects of soil pH on growth, nutrient uptake and mycorrhizal colonization in the roots of *L. microphyllum*, we conducted a 60-day greenhouse experiment by growing the fern in pots filled with pH adjusted soils to a range from 4.5 to 8.0. *Lygodium microphyllum* was able to survive and grow at all soil pH levels; however, final biomass, relative growth rate, photosynthesis and specific leaf area were all significantly greater in soil pH 5.5 - 6.5 compared to other treatments. Correspondingly, nitrogen concentration was also significantly related to these four plant parameters. Root colonization by mycorrhizal fungi was significantly higher in soil pH 5.5-7.5 and lowest for plants growing in 4.5 or 8.0, and was significantly correlated with plant growth parameters as well as elemental concentration in the leaves. In its native Australia, *L. microphyllum* responds robustly following fire. Fire increases soil pH, and given the treatment response to soil pH in my study, increased pH may help partially explain *L. microphyllum*’s response to burning. Recently, fire has been used as a potential management control option in its introduced range in Florida, given the results of this study and the plants
known fire tolerance in its home range, this management option should be reconsidered. Key words: element toxicity, invasive species management, plant-soil interactions, relative growth rate.

3.1 Introduction

Soil pH is an important factor for plant growth, as it affects nutrient availability, nutrient toxicity, and has a direct effect on the protoplasm of plant root cells (Rorison 1980; Alam et al. 1999). It also affects the abundance and activity of the soil organisms (from microorganisms to arthropods) responsible for the transformations of nutrients (De Boer and Kowalchuk, 2001; Nicol et al. 2008). Since most mineral nutrients are readily available to plants when soil pH is near neutral (pH = 7.0), species richness is high in such neutral soils, declining in both acidic and alkaline soils (Grime 1973; Gould and Walker 1999; Pausas and Austin 2001). Soil pH further influences the fate of chemicals, nutrients, and pesticides/herbicides added to the soil (Liu et al. 2001). Past research has shown that the species diversity is low in most acidic soils (Dupre et al. 2002) as essential nutrients (such as calcium, magnesium, potassium, phosphorus, and molybdenum) exist in unavailable forms to plants causing nutrient deficiency (Larcher 1980). Likewise, because to the inhibition of nitrification processes, nitrite, which can be toxic to plant and microorganisms, accumulates in acidic soils (Black 1968; Shen et al. 2003). In strongly acidic soils, certain ions (Al$^{3+}$, Cu$^{2+}$, Fe$^{3+}$, Mn$^{2+}$) rise to levels toxic for the majority of plants (Foy 1984; Kinraide 1993). Additionally, acidic soils have high cation exchange capacity, and promote leaching of nutrients resulting in soil unfavorable for plant growth (Johnson 2002). At the other extreme, alkaline soils tend to be deficient in iron,
manganese and phosphate (Marschner 1986; Tyler 1999). Marschner (1986) suggest that in alkaline soils, boron can rise to phytotoxic concentrations.

Plants differ enormously in their degree of tolerance to changes in soil characteristics (pH, moisture content, etc.): some have a narrow tolerance for one variable but a wide tolerance for others (Hill & Ramsay 1977). Weedy species collected from different climate zones show large growth differences when planted in soils with pH ranging from 4.8 to 6.4 (Buchanan et al. 1975). Stephenson & Rechcigl (1991) found that many weedy species grew significantly better when soil pH increased from 4.5 to 5.4, with good growth maintained at pH of 5.5 and above. Since invasive species have an affinity for disturbed areas, can reproduce sexually as well as asexually and yield a high number of seeds, they have a greater ability to adapt to changing conditions, potentially displacing native species through competitive exclusion (Baker 1974; Mooney and Cleland 2001; Prentis et al. 2008).

*Lygodium microphyllum* is an invasive exotic plant species taking over many sites in freshwater and moist habitats in Florida. It has the ability to grow in varying hydrological (Gandiaga et al. 2009), nutrient (Volin et al. 2010), and light conditions (Volin et al. 2004). Analysis of soil samples from both its native range and invaded region have shown that although *L. microphyllum* grows in highly acidic soils in its native range in Australia, it has adapted to thriving in close-to-neutral soils in Florida (Chapter 2). The roots of *L. microphyllum* are heavily colonized by mycorrhizal fungi that absorb nutrients, specifically P; biomass accumulation in mycorrhizal plants was almost three times that of non-mycorrhizal plants (Chapter 2). Furthermore, the level of mycorrhizal colonization was related with the soil pH: a higher degree of mycorrhizal
colonization is present in plants from the slightly acidic soils in the invaded regions compared with those from the highly acidic soil in the native regions (Chapter 2). Since mycorrhizal fungi have strong associations with *L. microphyllum*, supporting nutrient uptake in both its invaded regions as well as in the native regions (Chapter 2), the response of mycorrhizal fungi to variation in soil pH should be considered, especially if the manipulation of soil pH is integrated in the management plan for this invasive pest plant.

The aim of this study was to compare the degree of mycorrhizal colonization, nutrient uptake, biomass accumulation, and growth rate of *L. microphyllum* at different soil pH levels. Since the existing chemical control method is not very efficient in controlling *L. microphyllum*, this information may be useful in developing an integrated weed management technique. I hypothesized that plant growth and mycorrhizal colonization will be highest in slightly acidic soils with growth highly reduced (or the plants not surviving) in alkaline soils. I predicted that changing the soil pH can reduce the competitive ability of *L. microphyllum*. 
3.2 Methods

To test the hypothesis, I undertook a greenhouse experiment to investigate the effects of soil pH on various aspects of growth of *L. microphyllum*. Plants were maintained in pots in the Florida International University greenhouses until they began to sporulate, at which time the experiment was concluded.

3.2.1 Potting soil

Soil from plots in Tree Tops County Park at Davie, Florida was collected for this study; this site was not yet invaded by *L. microphyllum*. Soil was then passed through a 2 mm sieve and air dried in room temperature. Quartz sand was added to this soil to form a 1:1 soil/sand ratio. A sub-sample of the soil was analyzed to determine the initial soil characteristics. The soil pH was measured with a pH meter, (soil solution ratio 1:2 in water), texture was measured by the hydrometer method, total organic matter was measured on the basis of standard loss on ignition method (500°C, 5 hours; Storer 1984). To generate a soil neutralization curve (Kellog et al. 1957), 150 g of air dried soil samples were placed in 120 ml plastic containers and mixed with Ca(OH)$_2$ at rates 0, 1, 2, 3, 4, 5, 6, 7, 8 Mg Ha$^{-1}$, elemental S was added at rates 0.35, 0.40, 0.45 0.50, 0.55 and 0.60 Mg Ha$^{-1}$, with 5 replicates for each treatment. These soil samples were watered with DI water and incubated for 28 days and the soil pH was measured (soil solution ratio 1:2 in water). The amount of lime or sulfur required to raise the experimental soil pH to the desired level was determined based on the regression equation resulting from pH measurement of the incubated soils. The rate of Ca(OH)$_2$ and S added is given in Table 3.1.
The soil was sandy loam with organic matter 37%, 1.10 mg/g total P, 1.2 % total N and pH 5.5 ± 0.2. The soil was divided into 5 subsamples and elemental sulfur was added to lower the pH to 4.5 in one set; no treatment was done in the 5.5 pots; and Ca(OH)$_2$ was added to increase the pH to 6.5, 7.5 and 8.0. The soil samples with elemental sodium or Ca(OH)$_2$ were thoroughly mixed, added to the pots and watered with DI water. The soils were allowed to equilibrate for eight weeks with frequent mixing. Soil pH was measured weekly and after eight weeks all the pH measurements were within ± 0.3 of the targeted pH value and remained constant throughout the experiment time (measured every week in 1:2 water).

Table 0.1 Rates of application of Ca(OH)$_2$ and elemental S for pH adjustment of the experimental soils

<table>
<thead>
<tr>
<th>Soil original pH</th>
<th>Final pH</th>
<th>Rates of S or Ca(OH)$_2$ Application (Mg Ha$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.5</td>
<td>0.522</td>
<td></td>
</tr>
<tr>
<td>5.5</td>
<td>0.000</td>
<td></td>
</tr>
<tr>
<td>5.5</td>
<td>6.5</td>
<td>2.215</td>
</tr>
<tr>
<td>7.5</td>
<td>4.255</td>
<td></td>
</tr>
<tr>
<td>8.0</td>
<td>5.275</td>
<td></td>
</tr>
</tbody>
</table>

3.2.2 Plant material

Experimental plants were grown from spores following the method used by Lott et al. (2003). Spores of *L. microphyllum* were disinfected with 1% bleach and transferred
to Petri dishes that contained Parker-Thomson Medium. The plates were placed in an incubator set at 25-27°C for ten weeks and were watered with sterile DI water every week. After ten weeks, individual gametophytes were transferred to fresh Petri dishes. When the sporelings’ roots and leaves developed, 60 plants were transplanted to small pots previously filled with sterile sand. These 60 plants were placed in a growth chamber for approximately four weeks. The plants were kept moist, and were watered with half strength Hoagland’s nutrient solution as needed. Plants were then transferred to 2.5 L pots filled pH modified soil. The plants were grown in the green house for 60 days. Plants were watered to saturation biweekly and received 250 ml of half strength Hoagland's solution weekly. Before the plants were harvested, photosynthesis was measured using a Li-Cor 6400 Portable Photosynthesis System (Li-Cor Biosciences) on two fully grown leaves per plant in all the treatments. Measurements were taken at leaf temperatures ranging from 34°C to 38°C, CO₂ concentration of 400 µmol mol⁻¹ and photosynthetic photon flux was at 600 µmol m⁻² s⁻¹.

3.3.3 Harvest and plant nutrient analysis

Two harvests were conducted during this study: at time 0 (the beginning of the treatment/transplant date), and after 60 days. The allometric relationship between stem length and total mass (R² = 0.92) from the initial harvest was developed to estimate the initial plant mass of the experimental plants and to calculate the relative growth rate (RGR) (see Gandiaga et al. 2009). The RGR (mg g⁻¹ d⁻¹) was calculated for each individual plant used for the experiment, where RGR = [ln (final dry mass)-ln (initial dry mass)]/days (Evans 1972). After each harvest, individual plants’ roots, stem, and leaves
(pinnae) were separated and leaf area was measured with a leaf area meter to calculate the specific leaf area (SLA). The separated plant parts were oven-dried (one week at 65°C) to constant mass and weighed to determine the leaf mass ratio (LMR), stem (rachis) mass ratio (SMR), rhizome mass ratio (RhiMR), and root mass ratio (RMR); and relative growth rate (RGR).

The aboveground parts of the plants (shoot tissue) were analyzed for nutrient content. The oven dried tissues were carefully ground by hand using a mortar and a pestle. Samples underwent acid digestion using Method 3050B (USEPA 1996), summarized here: One gram of finely ground plant tissue sample was transferred to a large glass tube and mixed with 10 ml of 30% HNO₃. The tubes were covered with a vapor recovery system and heated to 95±5°C and refluxed for 10 minutes without boiling under the hood in a heating block maintained with a Partlow Mic 6000 Profile Process Controller. After cooling to 40°C, 5 ml of concentrated HNO₃ was added and the sample was heated again until no brown fumes were emitted. After cooling to 40°C, 2 ml of DI water and 3 ml of 30% H₂O₂ was added and heated until the effervescence subsided. The samples were cooled and diluted to 100 ml with DI water, centrifuged at 2000 rpm for 10 minutes and filtered with Whatman No. 41 filter paper and analyzed with an ICP-MS at USDA ARS, Homestead, Florida.
3.3.4 Mycorrhizal colonization

Before drying, forty-five 1.5 cm root fragments were collected from each plant, and the colonization of AMF was quantified following a modified method described by McGoingle et al. (1990). Roots were cleared in 15% KOH at 70°C for 4 hours, rinsed twice with water, bleached with ammoniated H₂O₂, and acidified with 1 N HCl. Staining was done using Trypan blue in acidic glycerol at 80°C for 20 minutes. The stained roots were examined with a dissecting microscope at 30–60 X magnification; the portions that showed the presence of mycorrhizal fungi were mounted on slides in lactic acid and examined at 100–400 X magnification.

3.3.5 Experimental design and data analysis

The experimental design was a randomized complete block with five pH treatments and six replicates. It was a single factor experiment investigating the effects of pH on plant growth, nutrient accumulation and level of mycorrhizal colonization. After the harvest at 60 days, regression analysis examined the influence of initial plant mass on RGR and its morphological, allocational and physiological determinants (e.g., Mcconnaughay and Coleman 1999; Volin et al. 2002; Kruger and Volin 2006). Regression analysis indicated that RGR was negatively correlated to the natural log (ln) of initial plant mass (p<0.001). Additionally, RMR final harvest was significantly related (P < 0.05) to final plant mass. Therefore, variation in plant mass was normalized using analysis of covariance. All of the variables in the four pH treatments were then compared with one-way ANOVA for significance at p ≤ 0.05. Correlation analysis between total biomass, RGR SLA, and leaf concentration of Al, Ca, P, N, and Fe were conducted to
determine the effects of leaf elemental status on plant growth. Regression analysis analyzed the relationship between the plant growth parameters and N concentration in the leaves. Regression analysis examined the relationship between RGR and its determinants. All analyses were performed with SAS Version 9.2 software (SAS Institute 2009).

3.3 Results

A significant effect of soil pH was visible on L. microphyllum growth, nutrient uptake, and degree of mycorrhizal colonization in its roots, despite the small sample size (n=6) and short duration (60 days) of this experiment (Fig. 3.1a & b, Table 3.2, Fig. 3.2). Relative growth rate and biomass allocation patterns were significantly different among the pH treatments (Fig. 3.1b). The growth of L. microphyllum was significantly greater in pH 5.5 and 6.5 compared to the strongly acidic and alkaline soils (Fig. 3.3). Total final plant mass was greatest in plants grown in soil with pH 5.5 and 6.5 and these were more than twice the biomass of plants grown in pH 8.0 (Fig. 3.1a). Correspondingly, this significant pattern was found for RGR, which increased with increasing soil pH from 4.5 to 5.5, remained unchanged at 6.5, and gradually declined with increasing soil pH, with lowest RGR at soil pH 8.0 (Fig. 3.1b).

Surprisingly, biomass allocation to the above ground parts was not influenced by soil pH (data not shown). There was, however, significant difference in biomass allocation to the belowground parts. Plants growing in soil pH 4.5 had significantly
Figure 0.1 Mean (±SE) final biomass (a); mean (±SE) relative growth rate (RGR) (b); mean (±SE) root mass ratio (RMR) (c) and mean (±SE) rhizome mass ratio (RhiMR) (d), measured at the end of 60 days in four different soil pH levels. Similar letters represent no significant difference at $P < 0.05$. 
higher biomass allocation to the roots compared to the other plants (Fig. 3.1c), while plants grown in soil pH 4.5 had significantly lower biomass allocation to the rhizomes compared to the plants grown in soil pH 5.5 and 6.5, though not significantly different from pH 7.5 and 8.0. Plants in soil pH 5.5 and 6.5 had significantly higher SLA than the other plants (Fig. 3.3). The influence of soil pH also was strongly reflected in the photosynthetic rates, which showed the same response as SLA. In other words, plants grown in pH 5.5 and 6.5 had significantly higher area based photosynthetic rates than plants grown in lower or higher pH soils, but there were no significant differences among the remaining three treatments (Fig. 3.4a). As a result of higher SLA and area based photosynthesis, mass based photosynthesis was also higher in plants grown in pH 5.5 and 6.5 (Fig. 3.4b).

### 3.3.1 Element concentration and uptake

Soil pH significantly affected the concentrations of Al, Ca, Fe, and N in the leaf tissue of *L. microphyllum* (Table 3.2), while it did not have any influence on the leaf concentration of P, K, Mg, Mn, and Zn. Plants grown in soil pH 4.5 had significantly higher concentration of Al and Fe and significantly lower concentrations of N. Similarly, plants grown in pH 8.0 had significantly high concentration of Ca. There was a strong relationship between biomass (p<0.0001), RGR (p<0.0001), SLA (p<0.0001) photosynthesis (p<0.0001)) and RhiMR (p<0.0001) and leaf concentration of N when all the treatments were pooled (Fig. 3.5). However, there were no significant correlations between the plant growth parameters and leaf concentration of Al, Ca, Fe, Mg, Mn, P, and Zn.
Table 0.2 Effect of soil pH on element concentration (mean ± std. dev.) in the leaf tissue of *L. microphyllum*

<table>
<thead>
<tr>
<th>Element</th>
<th>Concentration mg g⁻¹</th>
<th>pH 4.5</th>
<th>pH 5.5</th>
<th>pH 6.5</th>
<th>pH 7.5</th>
<th>pH 8.5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al</td>
<td>0.19 ± 0.02a</td>
<td>0.14 ± 0.01b</td>
<td>0.14 ± 0.03b</td>
<td>0.12 ± 0.02b</td>
<td>0.13 ± 0.02b</td>
<td></td>
</tr>
<tr>
<td>Ca</td>
<td>5.44 ± 0.40a</td>
<td>5.53 ± 0.33 a</td>
<td>5.53 ± 0.99a</td>
<td>5.52 ± 0.48a</td>
<td>7.01 ± 0.5 b</td>
<td></td>
</tr>
<tr>
<td>Fe</td>
<td>0.18 ± 0.05a</td>
<td>0.15 ± 0.03ab</td>
<td>0.12 ± 0.01b</td>
<td>0.13 ± 0.02b</td>
<td>0.13 ± 0.01b</td>
<td></td>
</tr>
<tr>
<td>K</td>
<td>22.01 ± 2.1a</td>
<td>21.75 ± 4.56a</td>
<td>21.1 ± 4.9a</td>
<td>21.16 ± 4.39a</td>
<td>23.9 ± 3.0a</td>
<td></td>
</tr>
<tr>
<td>Mg</td>
<td>2.05 ± 0.29a</td>
<td>2.08 ± 0.45a</td>
<td>2.13 ± 0.4a</td>
<td>1.98 ± 0.14a</td>
<td>2.12 ± 0.29a</td>
<td></td>
</tr>
<tr>
<td>Mn</td>
<td>0.11 ± 0.01a</td>
<td>0.10 ± 0.01a</td>
<td>0.09 ± 0.05a</td>
<td>0.08 ± 0.01a</td>
<td>0.08 ± 0.01a</td>
<td></td>
</tr>
<tr>
<td>P</td>
<td>3.72 ± 0.30a</td>
<td>4.25 ± 0.53a</td>
<td>3.73 ± 0.85a</td>
<td>3.78 ± 0.54a</td>
<td>3.46 ± 0.28a</td>
<td></td>
</tr>
<tr>
<td>N</td>
<td>1.94 ± 0.32a</td>
<td>3.34 ± 0.31b</td>
<td>3.35 ± 0.26b</td>
<td>2.83 ± 0.18c</td>
<td>1.99 ± 0.23a</td>
<td></td>
</tr>
<tr>
<td>Zn</td>
<td>0.09 ± 0.01a</td>
<td>0.09 ± 0.01a</td>
<td>0.07 ± 0.02a</td>
<td>0.07 ± 0.01a</td>
<td>0.08 ± 0.02a</td>
<td></td>
</tr>
</tbody>
</table>

Values in a row followed by the same letter are not significantly different at p ≤ 0.05.
Table 0.2 Comparison of the topsoil characteristics (means with standard deviations in parentheses) at the three sites with and without *L. microphyllum*.

<table>
<thead>
<tr>
<th>Site</th>
<th>Al (mg/g)</th>
<th>C (%)</th>
<th>Ca (mg/g)</th>
<th>N (%)</th>
<th>P (mg/g)</th>
<th>Zn (µg/g)</th>
<th>OM (%)</th>
<th>pH (H₂O)</th>
<th>TBC</th>
<th>TFC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Central FL</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Native</td>
<td>2.62 (0.64)</td>
<td>2.18 (0.50)</td>
<td>0.68 (0.34)</td>
<td>0.04 (0.11)</td>
<td>0.67 (0.10)</td>
<td>21.46 (9.94)</td>
<td>5.18 (0.80)</td>
<td>4.95 (0.45)</td>
<td>152.5</td>
<td>61.16</td>
</tr>
<tr>
<td>Invasive</td>
<td>5.07 (0.68)</td>
<td>4.03 (0.84)</td>
<td>0.41 (0.13)</td>
<td>0.195 (0.27)</td>
<td>1.03 (0.26)</td>
<td>15.93 (8.25)</td>
<td>8.65 (5.78)</td>
<td>5.78 (138.33)</td>
<td>88.83</td>
<td></td>
</tr>
<tr>
<td>P Levela</td>
<td>**</td>
<td>**</td>
<td>*</td>
<td>ns</td>
<td>**</td>
<td>ns</td>
<td>**</td>
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<tr>
<td>Jonathan</td>
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</tr>
<tr>
<td>Dickinson</td>
<td>0.43 (0.07)</td>
<td>3.13 (1.10)</td>
<td>1.19 (0.21)</td>
<td>0.26 (0.08)</td>
<td>1.02 (0.06)</td>
<td>7.24 (1.26)</td>
<td>1.08 (0.41)</td>
<td>(0.09)</td>
<td>(7.25)</td>
<td>(4.84)</td>
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<tr>
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<td>7.02 (1.88)</td>
<td>3.35 (1.77)</td>
<td>0.44 (0.12)</td>
<td>1.15 (0.09)</td>
<td>8.77 (1.86)</td>
<td>4.32 (0.90)</td>
<td>(0.1)</td>
<td>(4.28)</td>
<td>(7.76)</td>
</tr>
<tr>
<td>Invasive</td>
<td>1.88 (0.30)</td>
<td>22.43 (1.77)</td>
<td>17.21 (0.17)</td>
<td>1.27 (0.16)</td>
<td>1.22 (0.16)</td>
<td>23.21 (4.53)</td>
<td>44.42 (5.60)</td>
<td>5.60 (143.66)</td>
<td>73.83</td>
<td></td>
</tr>
<tr>
<td>P Levela</td>
<td>*</td>
<td>**</td>
<td>*</td>
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<tr>
<td>Tree Tops</td>
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<td></td>
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<tr>
<td>Native</td>
<td>1.62 (0.14)</td>
<td>16.55 (3.02)</td>
<td>9.11 (1.15)</td>
<td>1.31 (0.40)</td>
<td>1.11 (0.04)</td>
<td>17.48 (5.17)</td>
<td>36.75 (0.10)</td>
<td>(0.04)</td>
<td>(11.07)</td>
<td>(5.68)</td>
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<tr>
<td>Invasive</td>
<td>1.88 (0.25)</td>
<td>22.43 (4.15)</td>
<td>17.21 (6.30)</td>
<td>1.27 (0.17)</td>
<td>1.22 (0.16)</td>
<td>23.21 (4.53)</td>
<td>44.42 (5.60)</td>
<td>5.60 (143.66)</td>
<td>73.83</td>
<td></td>
</tr>
<tr>
<td>P Levela</td>
<td>*</td>
<td>**</td>
<td>*</td>
<td>ns</td>
<td>*</td>
<td>ns</td>
<td>ns</td>
<td>**</td>
<td>ns</td>
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</tr>
</tbody>
</table>

* Significance for paired t-test, ns: not significant; Probability levels: *: P<0.05; ** P<0.01; ***P<0.0001.

OM: soil organic matter; TBC: total bacterial count (count x 10⁷); TFC: total fungal count (count x 10³).
Table 0.3 Results of two-way analyses of variance (ANOVA) with degree of freedom (DF), F-value and probability levels for the effects of site, plant type and the interaction of the two on the soil characteristics.

<table>
<thead>
<tr>
<th>Source</th>
<th>DF</th>
<th>Al</th>
<th>C</th>
<th>Ca</th>
<th>Cu</th>
<th>Fe</th>
<th>K</th>
<th>Mg</th>
<th>Mn</th>
<th>N</th>
<th>P</th>
<th>OM</th>
<th>Zn</th>
<th>pH</th>
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<td>Site 2</td>
<td></td>
<td>195.3</td>
<td>179.89</td>
<td>75.91</td>
<td>46.87</td>
<td>19.5</td>
<td>10.09</td>
<td>28.59</td>
<td>3.47</td>
<td>99.65</td>
<td>16.78</td>
<td>2132.23</td>
<td>14.73</td>
<td>102.88</td>
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<tr>
<td>Plant 1</td>
<td></td>
<td>65.24</td>
<td>25.33</td>
<td>13.52</td>
<td>0.14</td>
<td>2.55</td>
<td>14.74</td>
<td>12.16</td>
<td>2.54</td>
<td>1.9</td>
<td>19.04</td>
<td>85.22</td>
<td>0.08</td>
<td>30.13</td>
</tr>
<tr>
<td></td>
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<td>***</td>
<td>**</td>
<td>ns</td>
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<td>Site × Plant 2</td>
<td>2</td>
<td>27.05</td>
<td>2.29</td>
<td>7.53</td>
<td>0.52</td>
<td>0.55</td>
<td>0.83</td>
<td>4.19</td>
<td>3.31</td>
<td>0.94</td>
<td>2.8</td>
<td>7.68</td>
<td>2.66</td>
<td>12.29</td>
</tr>
<tr>
<td></td>
<td>***</td>
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<td>**</td>
<td>ns</td>
<td>ns</td>
<td>ns</td>
<td>*</td>
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<td>***</td>
<td>ns</td>
<td>***</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Source</th>
<th>DF</th>
<th>TBC</th>
<th>TFC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Site 2</td>
<td></td>
<td>660.62</td>
<td>60.40</td>
</tr>
<tr>
<td></td>
<td>***</td>
<td>***</td>
<td>***</td>
</tr>
<tr>
<td>Plant 1</td>
<td></td>
<td>235.15</td>
<td>63.65</td>
</tr>
<tr>
<td></td>
<td>***</td>
<td>***</td>
<td>***</td>
</tr>
<tr>
<td>Site × Plant 2</td>
<td>2</td>
<td>120.48</td>
<td>2.58</td>
</tr>
<tr>
<td></td>
<td>***</td>
<td>ns</td>
<td>ns</td>
</tr>
</tbody>
</table>

Probability levels: *: P<0.05; ** P<0.01; ***P<0.0001. OM: soil organic matter; TBC: total bacterial count; TFC: total fungal count.
3.3.2 Mycorrhizal colonization

As expected, soil pH also had a significant effect on the degree of mycorrhizal colonization (Fig. 2). The degree of colonization was highest at pH 5.5 with no significant difference at pH 6.5 and 7.5, while the degree of colonization was significantly lower at both pH 4.5 and pH 8.0. There was no significant difference in the mycorrhizal structures such as vesicles, arbuscules, and hyphae among the four soil pH levels (data not shown).

When the pH treatments were analyzed independently there was no strong correlation between the root colonization by mycorrhizal fungi and plant growth parameters or the leaf concentration of elements. However when the samples were pooled there was a strong correlation between the degree of mycorrhizal colonization and plant growth parameters as well as the leaf element status (Table 3.3). Additionally, there was a significant correlation between the degree of mycorrhizal colonization and element uptake by *L. microphyllum*. 
Figure 0.2 Mean value (± SE) of degree of mycorrhizal colonization at different pH levels. Similar letters represent no significant difference at $p \leq 0.05$.

Table 0.3 Correlation coefficients of plant growth parameters and leaf element concentration with the degree of mycorrhizal colonization.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Pearson Correlation Coefficients</th>
<th>P value</th>
</tr>
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<tbody>
<tr>
<td>Biomass</td>
<td>0.64</td>
<td>0.0002</td>
</tr>
<tr>
<td>RGR</td>
<td>0.65</td>
<td>&lt;.0001</td>
</tr>
<tr>
<td>SLA</td>
<td>0.67</td>
<td>&lt;.0001</td>
</tr>
<tr>
<td>RhiMR</td>
<td>0.78</td>
<td>&lt;.0001</td>
</tr>
<tr>
<td>RMR</td>
<td>-0.65</td>
<td>0.0001</td>
</tr>
<tr>
<td>Photosynthesis</td>
<td>0.64</td>
<td>0.0001</td>
</tr>
<tr>
<td>Al</td>
<td>-0.43</td>
<td>0.0191</td>
</tr>
<tr>
<td>Fe</td>
<td>-0.45</td>
<td>0.0136</td>
</tr>
<tr>
<td>N</td>
<td>0.87</td>
<td>&lt;.0001</td>
</tr>
</tbody>
</table>
Figure 0.3 Mean (±SE) value specific leaf area (SLA) at different pH levels. Similar letters represent no significant difference at $p \leq 0.05$.

Figure 0.4 Mean (±SE) value of area based photosynthesis (a), mass based photosynthesis (b) at different pH levels. Similar letters represent no significant difference at $p \leq 0.05$. 
Figure 0.5 Linear regression of leaf nitrogen concentration and (a) total biomass, (b) specific leaf area (SLA); (c) relative growth rate (RGR); (d) area based photosynthesis ($\mu$mol m$^{-2}$ s$^{-1}$)
3.4 Discussion

Different soil pH levels were selected to include a wide range of soil pH where *L. microphyllum* has been reported to grow in its native range in Australia and the invaded regions in Florida. At soil pH 4.5 and 8.0 the plants were noticeably smaller and grew less vigorously compared to the other treatments. Soil pH 5.5 and 6.5 provided the most favorable conditions for the nutrient uptake, growth and biomass accumulation. This result was expected for *L. microphyllum* because extensive growth occurs in slightly acidic soils of Florida. Few other ferns in Florida have also been reported to prefer soil pH close to 6.0; field study by Van Loan (2006) showed that *Lygodium japonicum*, another pest plant species in Florida, was present in sites with soil pH 6.0. Similarly, Mathur (1980) reported that the fern *Rumohra adiantiformis* requires soil pH between 5.5 and 6.0 for optimal growth. However, as opposed to my expectation the plants grown in alkaline soils survived, maintained a fair growth rate and produced fertile fronds.

Past research has shown that invasive species have a higher tolerance to low soil pH and have superior ability to assimilate nutrients (Thompson et al. 1987; Emery and Perry 1995). In its native range in Australia, *L. microphyllum* grows in highly acidic soils (soil pH range 3.9-4.7) (Chapter 2), this adaptive capability of *L. microphyllum* to acidic soils was visible in my study. Contrary to my expectation plants grown in soil pH 8.0 maintained a fair growth rate, but the decline in the RGR was visible with the increasing soil pH thus further increasing the soil pH could provide a desired outcome, although its potential negative impact on native flora (both plants and microorganisms) would need to be assessed. The high RGR of plants grown in soil pH 5.5 and 6.5 plants corresponded to the higher mass-based photosynthesis, which resulted from the increased SLA as well as
area-based photosynthesis. My results indicate that variation in RGR was explained by
the variation in photosynthetic capacity of L. microphyllum and SLA.

My results show that the biomass allocation to the belowground structures was
different across the different pH treatments. An unexpected result was that plants grown
in strongly acidic soils allocated the highest biomass to the roots compared to the other
plants at the cost of biomass allocation to the rhizomes. The higher biomass allocation to
the roots in acidic soils contradict the common assumption that Al toxicity in acidic soil
causes a significant reduction in root growth by inhibiting cell division in the root apical
meristem (Farid 1991; Ryan et al. 1993; Crawford & Wilkens 1998) resulting in reduced
water and nutrient uptake. Abhramhamsen (1983) suggested that certain plant species
have the ability to translocate the Al absorbed from roots to other parts of plant to avoid
Al toxicity, element concentration in the roots were not measured in my study but this
may be one explanation for the extensive root growth in the plants in soil pH 4.5 and
would need to be substantiated in future research. Additionally, there is a possibility of
root exudates secretion by L. microphyllum as a defense mechanism to Al toxicity. The
ability to avoid Al toxicity in acidic soils could in part explain the extensive growth of L.
microphyllum in the sand mine spoils with toxic levels of Al and Fe in central Florida
(Soti, pers. obser.). The pH of soil in direct contact with the roots was not possible to
measure without disturbing the plants so I do not know if the L. microphyllum plants had
any influence on the soil in direct contact with the roots.

Another possibility is that extensive root growth is necessary for the acquisition of
water and nutrients for plants; in my study water was not a limiting factor, but nutrients
could have been limiting resource for plants growing in acidic soil. Schindelbeck and
Riha (1988) and Kidd & Proctor (2001), have found that decrease in soil pH caused an increased biomass allocation to roots; Bates et al. (2002) found that when the soil pH was lower than 4.4 the root: shoot ratio increased in Vitis labruscana L. plants. Phenotypic plasticity is one of the key characteristic of invasive plants which allows them to adapt to a wide range of habitat types (Claridge & Franklin, 2002). Previous studies have shown that L. microphyllum is extremely plastic in its ability to respond to myriad environmental conditions, including plasticity in reproduction, physiology, allocation, and morphology (see: Lott et al. 2003, Gandiaga et al. 2009, Volin et al. 2004, 2010 and 2013). In the present study, my results show that L. microphyllum adapts to low nutrient conditions in acidic soil by increasing biomass allocation to the roots. Phenotypic plasticity in response to environmental conditions has been reported in Melaleuca quinquenervia, a flowering tree which shares habitat with L. microphyllum in its native range in Australia and is invasive in most of the regions in south Florida, where 97% of its variation was accounted for soil pH (Kaufman & Smouse, 2001).

Soil pH had a significant effect on the element uptake by L. microphyllum. Higher concentration of Al in the leaves of plants growing in acidic soils did not substantially alter the growth parameters of L. microphyllum. It is reported that Al and Mn toxicity occur in soil when the pH is below 4.8 (Slattery et al. 1999), but L. microphyllum plants grown in strongly acidic soils did not show any sign of toxicity. Marschner (1995) suggested that N acts as growth regulator itself and is tied into plant growth allocation by direct involvement with plant growth regulators as well. In my study, the major element influencing plant growth was N. Its concentration was significantly higher in the plants grown in soil pH 5.5 and 6.5, and strongly correlated with the RGR, photosynthesis,
RhiMR, and SLA. A positive correlation between leaf N concentration and RGR, SLA, and photosynthetic capacity have reported for a wide range of plants (Poorter et al. 1990; Grime, 1991; Poorter and Bergkotte 1992; Reich et al. 1994; Nielsen et al. 1996; Cornelissen et al. 1997; Reich et al. 1998). Phosphorus is reported to form insoluble compounds under high soil pH conditions, causing P deficiency in plants (Shen et al. 2011). In my study there was no significant difference in the leaf concentration of P among the various soil pH treatments; this could have been in part influenced by arbuscular mycorrhizal fungi (AMF). Root colonization by AMF colonization is reported to be most positive when the soil is P-limited (Hoeksema et al. 2010), but in my study the plants were not limited by nutrients. However, AMF did have a significant effect on the P accumulation in the leaves of *L. microphyllum*.

*L. microphyllum* is reported to have most of the traits of an aggressive invader, including its reproductive characteristics, and its lack of a significant pathogens or herbivores in its introduced range. My results show that *L. microphyllum* can maintain a fair growth rate over a wide range of soil pH, indicating a continuing threat to most uninvaded sites. Soil pH levels 5.5-6.5 were optimal for rapid growth and biomass accumulation.

Burning and application of herbicides are the most commonly used methods to control *L. microphyllum*. Loveless (1959) found that burning raises the soil pH in tree islands of the northern Everglades. Furthermore, the fate of the chemical herbicides used to control *L. microphyllum* and other exotic invasive species depends upon soil pH. I found that increasing soil pH from highly acidic to near-neutral pH provides a more favorable condition for *L. microphyllum* growth. Prescribed burning, which causes a
temporary increase in soil pH, is a widely used method to control *L. microphyllum*, but my study shows that *L. microphyllum* could be benefitting from the slight increase in soil pH resulting from fire as well as the release of nutrients that are associated with burning. In my study, *L. microphyllum* had highest growth at neutral soil pH and began to show a significant decrease at a soil pH of 8.0, likely further growth reductions would happen in even more alkaline soils. Thus, raising soil pH may be a possible management option to explore in the future, but increasing the soil pH would need to be studied carefully for its potential adverse effects to native flora as well, including both native plants and soil microorganisms.

**Acknowledgements**

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Black CA (1968) Soil-plant relationships. Soil-plant relationships.


Hill SB, Ramsay J (1977) Weeds as indicators of soil conditions. The McDonald Journal 38:8-12


ALTERED SOIL BIOGEOCHEMICAL PROPERTIES BY EXOTIC INVASIVE OLD WORLD CLIMBING FERN (*LYGODIUM MICROPHYLLUM*) IN FLORIDA: IMPLICATIONS FOR MANAGEMENT AND RESTORATION


**Abstract**

Invasion by exotic species threatens ecosystems not only by competing with native species for resources, but also by having a substantial impact on the structure, function, and composition of the belowground ecosystem by modifying the physical, chemical, and biological properties of the soil. Old World climbing fern (*Lygodium microphyllum*) is one of the worst non-native plant species and has become a serious threat to the greater Everglades ecosystem of south Florida. In the present study, I analyzed the effects of Old World climbing fern on surface soil characteristics at invaded sites in Florida. I compared soil characteristics of six invaded and adjacent uninvaded plots at three different locations. My results show that the fern can grow and thrive in a wide range of soil types and the impact on the soil was site specific with effects being more prominent in sites with low nutrient status. Additionally, there were significant differences in the soil nutrient status and microbial population in the invaded and uninvaded sites. Sites with Old World climbing fern had significantly higher nutrient concentrations that correlated with higher soil organic matter. Overall my results indicate that this exotic pest plant can alter its belowground environment to its own benefit by enhancing the soil nutrient status by adding soil organic matter.
Nomenclature: Old World climbing fern, *Lygodium microphyllum*

Key words: invasive plants, nutrient cycle, pH, soil organic matter

4.1 Introduction

Invasion by exotic invasive plants has a substantial impact on the structure, function, and composition of the native communities (Evans et al. 2001; Ehrenfeld 2003; Rice and Emery 2003; Vila et al. 2011). Existing literature provides evidence that invasive plant species can modify physical, chemical, and biological properties of the soil including inputs and cycling of nutrients (Ehrenfeld 2003; Hawkes et al. 2005; Sperry et al. 2006), soil pH (Kourtev et al. 2003), soil organic matter and aggregation (Saggar et al. 1999). Invasive plants also modify the biotic composition of the soil by affecting the soil food web (Duda et al. 2003), total microbial communities (Kourtev et al. 2003), and fungal communities (Hawkes et al. 2006). Some invasive plants are also reported to exude allelochemicals which could inhibit soil borne pathogens, defend against disease, and repel insects (Yuan et al. 2012; and references there in). However, the documented impacts of invasive species on soil characteristics are diverse. While most of the studies have reported increased soil nutrient stock in invaded sites compared to non-invaded sites creating a positive feedback benefiting invasive species (Duda et al. 2003; Vanderhoeven et al. 2005; Liao et al. 2008; Perkins 2011), some other studies have shown negative feedback (Ley and D’Antonio 1998; Mack and D’Antonio 1998; Leary et al. 2006). A meta-analysis of litter decomposition rates by Liao et al. (2008) showed that the litter decomposition rate of the invasive plants was on average, 117% faster than the co-occurring native species. However, Ehrenfeld (2010) demonstrated slower rates of leaf
decay in exotic species compared with the native plant species. The species-specific variation was attributed to be the most important factor in determining the decomposition rates (Hoorens et al. 2003), as each plant species has a unique biochemical composition, including the nitrogen concentration and carbon-to-nitrogen ratio. Gordon (1998) in her meta-analysis has shown that out of 31 species considered most invasive, 12–20 (39–64%) potentially alter the ecosystem properties of geomorphology, hydrology, biogeochemistry, and disturbance in Florida.

*Lygodium microphyllum* (Old World Climbing Fern) is a highly invasive species distributed throughout the freshwater and moist habitats of south Florida. It is common in cypress swamp, pine flatlands, wet prairies, sawgrass marshes, mangrove communities, and Everglades tree islands (Pemberton and Ferriter, 1998). Once established, *L. microphyllum* dominates both understory and overstory native wetland habitats. It has the ability to grow in varying hydrological (Gandiaga et al. 2009), nutrient (Volin et al. 2010), soil pH (Chapter 2), and light gradients (Volin et al. 2004). Results from Chapter 2 also show that *L. microphyllum* is highly dependent on mycorrhizal fungi for growth and phosphorus uptake; thus, fungi could highly enhance its invasiveness. According to an estimate by (Ferriter and Pernas 2006), *L. microphyllum* covers 183,080 acres in the entire South/Central Florida region. Managing *L. microphyllum* has been a significant challenge for land resource managers and researchers as a consequence of its extensive rapid invasion in natural areas of south Florida.

Aboveground changes caused by *L. microphyllum* in south Florida natural areas are obvious and have remained the focus of land managers and researchers, but belowground changes caused by the plant-soil feedback have not gained much research
interest so far. However, for the management of this exotic invasive and restoration of invaded sites, it is essential to understand the direct and indirect impacts of *L. microphyllum* on soil processes and how these changes can influence the successful management of invaded areas. If *L. microphyllum* can successfully modify soil processes such as nutrient cycling, litter decomposition, and soil microbial communities, simply removing it may not be an effective management strategy. Furthermore, it is necessary to assess the effects of soil modification by *L. microphyllum* on the invasability of plant communities, whether this modification facilitates other invasive species and if it has a negative impact on the native species. The aim of the present is to obtain baseline information on how *L. microphyllum* alters the physical, chemical, and microbial characteristics of the invaded sites, which would help to better understand and interpret effects of soil additives to control this pest plant species. I compared an invaded and a nearby non-invaded site to examine the impacts of *L. microphyllum* invasion on the topsoil chemistry and microbial populations at three sites with different soil characteristics.

4.2 Methods

4.2.1 Sampling sites

Two sites in south Florida were selected for the comparison of soil characteristics: Tree Tops Park (Broward County) and Jonathan Dickinson Park (Martin County), and one site in central Florida, the Trustcorp/Tiedtke property (Lake County). These sites were paired to include one plot with *L. microphyllum* and another with native plants. The sites were selected on the basis of the following criteria: 1) a well-established
monospecific population; 2) sites that had not undergone any management activities for at least the last five years; 3) homogeneous soil type under both the native and invasive plants; and 4) adjacent plots with native plants. Site location and dominant vegetation in the sampling sites are given in Table 4.1.

Table 0.1 Site locations, vegetation, and site type.

<table>
<thead>
<tr>
<th>Site</th>
<th>Coordinates</th>
<th>Dominant species in the invaded sites</th>
<th>Site Type</th>
</tr>
</thead>
<tbody>
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<td>26° 4'0.04&quot;N,</td>
<td><em>Chrysobalanus icaco</em>, <em>Osmunda regalis</em> var. <em>spectabilis</em>, <em>Annona glabra</em></td>
<td>County Park, disturbed habitat</td>
</tr>
<tr>
<td>Park</td>
<td>80° 16' 5.88&quot;W</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Jonathan</td>
<td>27°0’37.33”N,</td>
<td><em>Pinus elliotii</em>, <em>Myrica cerifera</em>, <em>Ilex cassine</em>, <em>Serenoa repens</em></td>
<td>State Park, undisturbed habitat</td>
</tr>
<tr>
<td>Dickinson</td>
<td>80°7’20.28”W</td>
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<td></td>
</tr>
<tr>
<td>Central</td>
<td>28° 23’ 4.03&quot; N,</td>
<td><em>Pinus elliotii</em>, <em>Quercus geminata</em>, <em>Quercus nigra</em>, <em>Serenoa repens</em></td>
<td>Private property, sand mine spoil</td>
</tr>
<tr>
<td>Florida</td>
<td>81° 44' 41.30&quot; W</td>
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</table>
4.2.2 Soil sampling and analysis

At each sampling site, six 1 m × 1 m plots were selected randomly and soil from the 10-15 cm deep zone was collected from each of the four corners and the center of each plot with a soil corer (ø 18 mm) and mixed homogeneously into one bulk sample for each plot. The soil samples from south Florida were transported to the laboratory in a cooler and the samples from central Florida were cooled to 4ºC and shipped overnight. A portion of the soil samples from all sites were stored in a 4ºC refrigerator until analysis for biological measurements. A small portion of each soil sample was air dried and passed through a 2 mm sieve for analysis of physicochemical properties. Those subsamples were then ground to fine powder with a mortar and pestle, and stored at room temperature in air-tight containers for further analysis of nutrients and trace elements. The soil pH was measured with a pH meter, (soil: solution ratio 1:2 in water), texture was measured by the hydrometer method, and total organic matter was measured based on the standard loss-on-ignition method (500ºC, 5 hours; Storer 1984). Total C and N were measured with a Truspec CN analyzer. Total Ca, Fe, Al, Mg, K, Mn, and P were measured with an ICP–MS at USDA, ARS Laboratory, Homestead, Florida after following the acid digestion Method 3050 (USEPA 1996). One gram of each finely ground soil sample was transferred to a large glass tube and mixed with 10 ml of 30% HNO₃. The tubes were covered with a vapor recovery system and heated to 95±5ºC and refluxed for 10 minutes without boiling under the hood in a heating block maintained with a Partlow Mic 6000 Profile Process Controller. After cooling to 40ºC, 5 ml of concentrated HNO₃ was added and the sample was heated again until no brown fumes were given off. After cooling to 40ºC, 2 ml of DI water and 3 ml of 30% H₂O₂ was added
to each tube and heated until the effervescence subsided. The samples were cooled and
diluted to 50 ml with DI water, centrifuged at 2000 rpm for 10 minutes and filtered with
Whatman No. 41 filter paper.

The total colony-forming-units (CFU) of bacteria and fungi were determined by
the standard dilution spread plate method as described by Seely and VanDemark (1981).
The dry equivalent of one gram soil was mixed in 9 ml sterile water (autoclaved) and was
diluted serially. Samples were vigorously mixed during dilution to assist in dislodging the
bacteria from the soil particles. A serial dilution of $10^{-2}$, $10^{-3}$, $10^{-4}$, and $10^{-5}$ was made for
fungi and $10^{-4}$, $10^{-5}$, $10^{-6}$, and $10^{-7}$ for bacteria. A total of 100 µl of diluted soil suspension
was spread on three plates per soil sample for both bacteria and fungi at each dilution
level. Nutrient agar containing cycloheximide solution (to prevent fungal growth) was
used for bacteria and Rose Bengal Agar (RBA) with streptomycin sulphate (to prevent
bacteria growth) was used for the estimation of fungal colonization. Sterilized water was
spread on the agar plates that were used as controls.

**4.2.3 Data Analysis**

Differences in soil characteristics between the invaded and uninvaded plots were
compared by means of paired t-tests. Additionally, a two-way ANOVA was done with
site and vegetation type (invaded and uninvaded) as the fixed main effects for selected
soil parameters. Pearson’s correlation analysis was done with all sites pooled to
determine relationships between the measured soil variables. Differences are reported as
significant for tests with $P$-values $\leq 0.05$. All the parameters were analyzed with SAS
Version 9.2 software.
4.3 Results and Discussion

Despite increasing evidence for positive feedback between the exotic invasive species and soil, along with the ongoing challenges in successful management of exotic species in the south Florida Everglades, there are very few studies quantifying the impacts of exotic invasive species on ecosystem processes (See Gordon, 1998). The goal of this paper is to determine the soil factors influenced by *L. microphyllum* and to test whether the competitive advantage of *L. microphyllum* was due in part to nutrient sequestration and soil factors that might contribute to the suppression of native plant species. I compared soil from three different sites with contrasting soil characteristics and land use history; thus, the differences observed in soil measured soil parameters among these sites are in large part explained by this.

There was a clear contrast between the rhizosphere soils from the three sites (Table 4.2; Table 4.3). All the study sites were acidic (soil pH 4.95-6.36), but the sites in central Florida and at Tree Tops Park had lower soil pH compared to the sites at Jonathan Dickinson Park. Likewise, the soil texture differed among the sites; Jonathan Dickinson sites had sandy soils, while the soils in central Florida were clayey, and the soil at Tree
Tops Park was sandy loam. Soil organic matter was highest at Tree Tops Park and was lowest at Jonathan Dickinson Park.

Impacts on the nutrient cycling process are considered the most prominent effects of invasive species in the ecosystem. In my study, several soil characteristics were significantly different between the invaded plots and non-invaded plots in all three sampling sites. Most notably, there were significant differences in the total soil Al, C, Ca, and OM% at all three sites (Table 4.2). At all the sites, soils invaded by *L. microphyllum* had higher concentrations of Al, Ca, C and organic matter. Differences in the concentrations of N, P, Zn, and pH were site-specific. Soils under native vegetation were generally more acidic; this was statistically significant at the central Florida and Jonathan Dickinson sites but was not statistically significant at the Tree Tops Park site. This indicates that the effect of *L. microphyllum* on soil characteristics is site-specific and depends on existing soil conditions. There was no significant difference in the Cu, Fe, K, Mg, and Mn concentrations under native vegetation and *L. microphyllum* at all the three sites (data not shown).

Plant species can change the soil microbial community structure and function with varying amount and quality of litter deposition (Ehrenfeld 2003; Kurtev et al. 2002), which, in turn, could influence the soil nutrient concentration under the native and invasive species. The bacteria population was significantly higher under the native species compared to *L. microphyllum* in two sites; the difference was not statistically significant at the central Florida site. On the other hand, the fungal population was significantly higher under *L. microphyllum* compared to the native plants at all three sites (Table 2). Further analysis on the type of microbes and the role of allelochemicals, which
are reported to be an important determinant for invasive success of exotic plants (Bais et al. 2003), in regulating the soil microbial community structure is necessary, but my results provide some evidence that *L. microphyllum* could regulate the structure of soil microbial communities in its rhizosphere.

My results corroborate the results of Ehernfield (2003) and Callaway et al. (2004), where invasive plants enhance productivity and nutrient availability in invaded soils via an abundant litter deposition thus increasing their own success. Soil organic matter was strongly correlated to the available soil nutrients (Table 4.4) which indicates that the difference in the organic matter inputs to the soil under the natives and *L. microphyllum* could influence the difference in the nutrient availability. The most significant effects on soil characteristics were seen at sites with the lowest nutrient concentrations (Table 4.2). Additionally, the site effect was highly significant for all the soil parameters analyzed, indicating that *L. microphyllum* can adapt to and thrive in sites with a significant variation in nutrients as well as other soil characteristics.
Table 0.4 Pearson’s correlation coefficients between the selected soil parameters with all sites pooled.

<table>
<thead>
<tr>
<th></th>
<th>Al</th>
<th>C</th>
<th>Ca</th>
<th>N</th>
<th>OM</th>
<th>P</th>
<th>Zn</th>
<th>pH</th>
<th>TBC</th>
<th>TFC</th>
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<tbody>
<tr>
<td>Al</td>
<td>-</td>
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<td>C</td>
<td>-0.171</td>
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<tr>
<td>Ca</td>
<td>-0.173</td>
<td>0.892</td>
<td>-</td>
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<tr>
<td>N</td>
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<td>0.894</td>
<td>0.792</td>
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<tr>
<td>OM</td>
<td>-0.030</td>
<td>0.928</td>
<td>0.874</td>
<td>0.875</td>
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<tr>
<td>P</td>
<td>-0.099</td>
<td>0.542</td>
<td>0.571</td>
<td>0.524</td>
<td>0.433</td>
<td>-</td>
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<td>Zn</td>
<td>0.255</td>
<td>0.389</td>
<td>0.447</td>
<td>0.325</td>
<td>0.470</td>
<td>-0.151</td>
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<tr>
<td>pH</td>
<td>-0.385</td>
<td>-0.127</td>
<td>-0.142</td>
<td>-0.069</td>
<td>-0.321</td>
<td>0.457</td>
<td>-0.716</td>
<td>-</td>
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<tr>
<td>TBC</td>
<td>0.254</td>
<td>0.463</td>
<td>0.339</td>
<td>0.533</td>
<td>0.651</td>
<td>-0.057</td>
<td>0.469</td>
<td>-0.632</td>
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<td>ns</td>
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<tr>
<td>TFC</td>
<td>0.776</td>
<td>0.190</td>
<td>0.135</td>
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<td>0.270</td>
<td>-0.035</td>
<td>0.524</td>
<td>-0.432</td>
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<td>-</td>
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Probability levels: *: P<0.05; ** P<0.01; ***P<0.0001. Coefficients higher than 0.75 are in bold.

OM: soil organic matter; TBC: total bacterial count; TFC: total fungal count.
Overall, my results show increased mineral nutrient concentrations in the rhizosphere of *L. microphyllum* compared to the rhizosphere of adjacent native species. These results follow the general trend reported by various researchers, where the nutrient pools in the invasive species rhizosphere are significantly increased compared to the coexisting natives (Duda et al. 2003; Vanderhoven 2005; Dassonville et al. 2008; Liao et al. 2008; Perkins 2011). This effect was most evident at sites with lowest nutrient concentration. As reported by Ehrenfeld (2003) and Liao et al. (2008), this may be the direct effect of increased amounts of C and N added to the soil with higher litter input. Dassonville et al. (2008), have reported an opposite impact of invasive species in nutrient-poor versus nutrient-rich sites. Although any pre-existing differences in the plots with and without *L. microphyllum* cannot be disregarded with complete certainty, I believe that the differences in the soil characteristics between the invaded and uninvaded plots could be the result of difference in plant species in them. *L. microphyllum* invasion is still expanding and pre- and post invasion comparison could provide a better insight. Successful management of habitats invaded by exotic plant species requires a prior knowledge of whether the invaders have significantly altered the ecosystem (Walker and Smith 1997) because soil properties such as texture, pH, and organic matter content influence herbicide efficiency and therefore control success. Along with the added organic matter, nutrients, and changes in the pH of the soil, other specific ecosystem process as outlined by Gordon (1998) could also be influenced by various exotic invasive species that create positive feedback for themselves and future invaders. Additionally, mechanisms such as production of allelochemicals, and changes in the microbial
communities merit future research to achieve successful control of *L. microphyllum* and other exotic invasive species in the Everglades.

**Acknowledgement**

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INFLUENCE OF SOIL BIOGEOCHEMICAL PROPERTIES ON EXOTIC INVASIVE *LYGODIUM MICROPHYLLUM*: A CROSS CONTINENT COMPARISON OF SOIL CHARACTERISTICS TO INVASION SUCCESS

Abstract

With the influence in the plant’s ability to extract water and nutrients, soil characteristics play an important role in the distribution of plant species. The objective of this research was to analyze the soil characteristics associated with exotic invasive, *Lygodium microphyllum*, in its native range in Australia and the recipient habitat in south Florida. Rhizosphere soil samples from both the continents were analyzed for the soil physical, chemical and biological characteristics. The results from this study indicate that rhizosphere soil characteristics were very different in the two regions. Likewise, leaf nutrient status of this plant also varied in the two continents. The composition of mycorrhizal fungi, which is believed to aid this plant in the recipient habitat, was also very different with higher diversity in the disturbed sites compared to the undisturbed sites. The most important result was the Australian sites had a high concentration of aluminum and zinc which are phytotoxic in a highly acidic soil conditions compared to the Florida sites. Overall, my results indicate that *L. microphyllum* could be growing poorly in its native range in Australia because of the soil toxic effects associated with strong soil acidity and low foliar nitrogen concentration which in turn could affect the photosynthetic capacity of the plant. On the other hand, Jonathan Dickinson Park, which has the worst case of *L. microphyllum* infestation in Florida, provides a more favorable growth environment for this plant with well drained sandy, slightly acidic soils with low concentration of soil elements. This study highlights that along with the characteristics of
exotic plant species and native plant community, the understanding of invasive success of exotic plants needs the understanding of belowground community and ecology.

Key words: soil toxicity, habitat restoration, exotic invasive species, mycorrhizal fungi.

5.1 Introduction

*Lygodium microphyllum* (Old World climbing fern) is an invasive exotic plant species taking over many sites in freshwater and moist habitats across southern and central Florida. It is reported to have reached a “critical mass” of coverage and begun exponential rate of expansion, where the spread rate is higher than the management effort. Biannual surveys conducted by South Florida Water Management District estimate that this fern had doubled its coverage in just two years (FNPS 2005). *Lygodium microphyllum* does not have a high economic value in its native range, thus there is very little information available about its native ecology and the available information is mostly on its native herbivores. Very little is known about why this plant is invasive outside its native range (Ferriter 2001). It is reported to be found in a variety of habitats including freshwater creeks, perennial creeks, coastal depression wetlands, upland rainforests and sheltered canyons near permanent springs in its native range in Australia (Goolsby et al. 2003). It Asia, the fern is reported to occur in lowland rainforests in peat soils, coastal wetlands and in habitats dominated by wet clay soils (Goolsby et al. 2003).

In the invaded regions of south Florida *L. microphyllum* displays most of the ecological characteristic associated with successful invasive plants (Westbrooks 1998): it has the ability to grow in varying hydrological (Gandiaga et al. 2009), nutrient (Volin et al. 2010), and light conditions (Volin et al. 2004). It produces millions of spores all year
round which can be transported by wind up to 30 miles; it tolerates a wide range of soil pH (Chapter 3); has a symbiotic relationship with arbuscular mycorrhizal fungi (AMF) (Chapter 2). Comparative analysis of soil samples from both its native range and invaded region have shown that *L. microphyllum*, which had adapted to close-to-neutral soils in Florida, grows in highly acidic soils in its native range in Australia (Chapter 2). Additionally the Food and Agriculture Organization (FAO) world soil distribution map shows the soil in the native habitats, identified by Goolsby et al. (2006), is a highly acidic region (Fig 5.1).

Soil pH has complex effect on plant growth leading to the variation in the distribution of plant species in acidic or calcareous soils. Diekmann and Lawesson (1999) reported that pH is one of the major underlying variables determining the floristic variation within forests. The major growth limiting factors associated with acid soil infertility include toxicities of aluminum and manganese, and deficiencies or low availability of certain essential elements including calcium, magnesium, phosphorus and molybdenum (Foy 1984). These factors may directly restrict plant growth or indirectly restrict plant growth through interference in the development and functioning of symbiotic associations with rhizobia, mycorrhizas and actinomycetes. It is reported that Al toxicity as a result of strong acidic soils have a detrimental effect in plant growth by lowering rooting depth, increasing susceptibility to drought and decreased uptake of subsoil nutrients. Plants exposed to Al toxicity are reported to have stunted growth, small dark green leaves, and late maturity, the root tips and lateral roots are thickened. They have many stubby lateral roots but lack in fine branching, thus, inefficient in absorbing nutrients and water (Kochian et al. 2004, and references there in). Similarly, soil texture
is also an important factor influencing the distribution of minerals, organic matter and microbial community and other soil properties (Scott and Robert 2006). Along with soil pH, soil texture plays an important role in controlling the mobility of elements in the soil, with the mobility of metals being highest in acidic coarse-textured soils (McBride 1994). Likewise, soils which are sandy or better drained have extensive fine roots compared to clay soils. Volin et al. (2010) indicated that the growth of *L. microphyllum* was highest in sandy soils which indicate that this plant prefers well drained sandy soils. Furthermore, the root and rhizome growth of *L. microphyllum* was highest in the sandy soils of south Florida compared to the native Australian soils (Volin et al. unpublished data).

Most of the research on exotic invasive species is focused on the traits that enhance the probability such as high growth rate, short lifecycle, high levels of resource allocation to reproduction, and flexible utilization of available environmental resources, of a particular species being a successful invader in a recipient community. There are several studies with biogeographic comparison of invasive species in their native range and invaded range focusing on the impact of variable soil microbes on plant performance (Callaway and Aschehoug 2000; Hierro et al 2005; Vermeij et al. 2009; Volin et al. 2010). However, there are no studies conducted comparing the soil element status and its effect on plant growth in the native and recipient habitat.

I conducted a cross continent comparison of soil characteristics associated with *L. microphyllum*. Here, I present evidence that the invasion by *L. microphyllum* in south Florida is not only facilitated by the soil microbial community but also by the soil chemical characteristics. In this paper I suggest that invasive plants not only escape from their natural herbivores but also the toxic soil environment in their native habitats.
Figure 0.1 World soil pH map data source ISRIC-Wise world dataset
5.2 Methods

5.2.1 Sites and sample collection

I compared soil samples collected from three different sites in Australia, where the plant is native, and the invaded sites in Florida. Sampling dates and site information are given in Table 5.1. The soil samples were collected during the dry season in both the continents except for the central Florida site, which was added later, because of its unique characteristics. At each sampling site, six 1 m × 1 m plots were selected randomly and soil from the 10-15 cm deep zone was collected from all four corners and the center of each plot with a soil corer (diameter: 18 mm) and mixed homogeneously into one bulk sample for each plot. The soil samples from south Florida were transported to the laboratory in a cooler. Samples from Australia and central Florida were stored in 4°C and shipped overnight.

5.2.2 Soil nutrient analysis

Small portion of each soil sample was air dried, passed through a 2 mm sieve for analysis of physicochemical properties. They were then ground to fine powder with a mortar and pestle, and stored at room temperature in air-tight containers for further analysis of nutrients and trace elements. The soil pH was measured with a pH meter, (soil solution ratio 1:2 in water), texture was measured by the hydrometer method, total organic matter was measured based on the standard loss on ignition method (500°C, 5 hours; Storer 1984). Total C and N in soil and leaves were measured with a Truspec CN analyzer. Total Ca, Fe, Al, Mg, K, Mn and P in soil were measured with an ICP –MS at USDA, ARS Laboratory, Miami, Florida after following the acid digestion Method
3050B (USEPA 1996). One gram of finely ground soil samples were transferred to large glass tubes and mixed with 10 ml of 30% HNO₃. The tubes were covered with a vapor recovery system and heated to 95±5°C and refluxed for 10 minutes without boiling under the hood in a heating block maintained with a Partlow Mic 6000 Profile Process Controller.

Table 0.1 Sampling sites, dominant vegetation and date of sample collection for the two continents.

<table>
<thead>
<tr>
<th>Site</th>
<th>Coordinates</th>
<th>Dominant vegetation</th>
<th>Sample date</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tree Tops Park, FL, US</td>
<td>26° 4'0.04&quot;N, 80° 16' 5.88&quot;W</td>
<td>Royal fern</td>
<td>Dec, 2010</td>
</tr>
<tr>
<td>Central Florida, US</td>
<td>28° 23' 4.03&quot; N, 81° 44' 41.30&quot; W</td>
<td>Royal fern</td>
<td>June 2012</td>
</tr>
<tr>
<td>Jonathan Dickinson, FL, US</td>
<td>27°0'37.33&quot;N, 80°7'20.28&quot;W</td>
<td>Slush pine</td>
<td>Dec, 2010</td>
</tr>
<tr>
<td>Daintree Ferry, Queensland, AU</td>
<td>16°15'25.57&quot;S, 145°24'3.94&quot;E</td>
<td>Drynaria</td>
<td>June, 2011</td>
</tr>
<tr>
<td>Logan Reserve, Queensland, AU</td>
<td>27°40'4.16&quot;S, 153°16'0.44&quot;E</td>
<td>Bungwall fern</td>
<td>June, 2011</td>
</tr>
</tbody>
</table>

After cooling to 40°C 5 ml of concentrated HNO₃ was added and the sample was heated again until no brown fumes were given off. After cooling to 40°C, 2 ml of DI
water and 3 ml of 30% H₂O₂ was added and heated until the effervescence subsided. The samples were cooled and diluted to 50 ml with DI water, centrifuged at 2000 rpm for 10 minutes and filtered with a Whatman No. 41 filter paper.

5.2.3 Microbial analysis: bacteria and fungi population

The total colony forming units (CFU) of bacteria and fungi was determined by the standard spread plate dilution method as described by Seely and VanDemark (1981). Dry equivalent of one gram soil was mixed in 9 ml sterile water (autoclaved) and was diluted serially. Samples were vigorously mixed during dilution to assist in dislodging the bacteria from the soil particles. A serial dilution of 10⁻², 10⁻³, 10⁻⁴, and 10⁻⁵ was made for fungi and 10⁻⁴, 10⁻⁵, 10⁻⁶, and 10⁻⁷ for bacteria. A total of 100 µl of diluted soil suspension was spread on three plates per soil sample for both bacteria and fungi at each dilution level. Nutrient agar containing cycloheximide solution (to prevent fungal growth) was used for bacteria and Rose Bengal Agar (RBA) with streptomycin sulphate (to prevent bacteria growth) was used for the estimation fungal colonization. Sterilized water was spread on the agar plates were used as control. Inoculated plates were incubated at 26°C for 3 days before the colonies were counted. Dilution plates with 100 to 300 colonies per plate were counted.

5.2.4 Mycorrhizal spore identification

Results from previous chapters indicate that the mycorrhizal root colonization in L. microphyllum is significantly higher in the invaded regions compared to the native regions in Australia. I further identified the mycorrhizal spores in the rhizosphere soil of
L. microphyllum in both the regions following the wet sieving technique (Gerdemann and Nicolson 1963). 100 ml of DI water was added to dry equivalent of 50 g of soil from each site. It was then mixed vigorously to separate the spores from soil aggregates. The mixture was washed through a series of sieves (2 mm, 100 µm and 32 µm). Washing was done until the water flowing through the sieves was clear. The sievate retained on the sieves was washed and centrifuged with water to remove floating organic debris and the supernatant was discarded. The pellet in the bottom was re-suspended in a 50% sucrose solution, and centrifuged for one minute at 2000 RPM to separate the spores from denser soil components. Immediately after centrifugation, spores in the sucrose supernatant were rinsed in a fine sieve to remove the sucrose. The spores were then washed into a filter paper for vacuum filtration. The fungal spores were then mounted on slides for taxonomic identification to the genus level based on the spore morphology and wall characteristics, using the descriptions by the International Culture Collection of Arbuscular and Vesicular-Arbuscular Mycorrhizal Fungi (http://invam.caf.wvu.edu). The genus that was dominant was taken as the representative mycorrhizal AMF type for each site.
5.2.5 Statistical analysis

Analysis of variance (ANOVA) was done to compare the difference in the soil and leaf nutrient status of *L. microphyllum* among the different sites in the two continents. Linear regression with Pearson’s correlation analysis and was done with all sites pooled to determine relationship between the measured soil variables. Differences are reported as significant for tests with $P$-values $\leq 0.05$. All the parameters were analyzed with SAS Version 9.2 software.

5.3 Results

5.3.1 Soil texture

Soil texture, shown in Table 5.2, varied significantly among the different sites. Soil in Jonathan Dickenson was dominantly sand (98% sand), sandy loam in Tree Tops, sandy clay loam in Logan, loam in Nudgee, silt loam in Daintree and clay in Central Florida.

5.3.2 Nutrient analysis

My results indicate a significant variation in the soil properties in the recipient habitat in Florida and native range in Australia (Table 5.3). The native range in Australia had strong acidic soil ranging from pH = 4.1 at Nudgee to pH = 4.55 at Logan. The Florida sites had significantly higher soil pH ranging from 5.60 at Tree Tops Park to 6.57 at Jonathan Dickinson Park. Soil Al concentration was highest at the Central Florida site followed by Logan and Nudgee in Australia. Soil C% and organic matter % was highest in Tree Tops park followed by Logan, C% was lowest in Daintree while organic matter %
was lowest in Jonathan Dickinson. Total Cu ranged from 43.94 µg g⁻¹ in Daintree to 1.84 µg g⁻¹ in Jonathan Dickinson, there was no significant difference among the other four sites. Total Fe was also highest in Daintree in Australia: 15.86 mg g⁻¹ and lowest in Jonathan Dickinson 0.54 mg g⁻¹. Total K in soil was also highest in Daintree (1.77 mgg⁻¹) and Logan (1.49) while there was no significant difference among the other sites. The Australian sites and Central Florida site had significantly lower level of N compared to the Tree Tops and Jonathan Dickinson sites. Total Mg in soil was highest in Logan (1.27 mg g⁻¹) followed by Tree Tops Park (0.68 mg g⁻¹) and was lowest in Central Florida (0.20 mg g⁻¹). Total soil Mn was also highest in Daintree (0.27 mg g⁻¹) while there was no significant difference among other five sites. N% in the soil was highest in Tree Tops (1.27%) followed by Logan (0.71%), and Jonathan Dickinson (0.49%). There was no significant difference in the total P level among all the sites. Logan had highest concentration of Zn in the soil (37.80 µg g⁻¹) followed by Daintree (27.91 µg g⁻¹) and Tree Tops (23.21 µg g⁻¹)

5.3.3 Bacteria and fungi population

The average counts of bacteria and fungi, colony forming units CFU per gram of 1 g dry soil, was significantly different in all the sites (Table 5.3). The CFU of bacteria and fungi was influenced by soil texture. Total colony forming units of bacteria was highest in Daintree \((288 \times 10^7)\) and lowest in Jonathan Dickinson \((43.83 \times 10^7)\) and there was no significant difference in the population in the other four sites. Likewise, the total colony forming units of fungi was also highest in Daintree \((123.5 \times 10^3)\) while there was no significant difference among the other five sites. Correlation analysis indicated that,
there was no relationship between the soil organic matter, total carbon on the soil bacteria and fungal population, however there a strong relationship with the soil texture. There was a strong negative relationship between the sand content in the soil and the CFU of bacteria ($r = -0.62; p < 0.0001$) and fungi ($r= -0.67; p< 0.0001$) on the other hand there was strong positive relationship with the silt content in the soil (bacteria: $r = 0.76; p< 0.0001$ and fungi: $r=0.75; p<0.0001$). Surprisingly the bacteria population had a negative relationship with the soil pH ($r= -0.57; p=0.0003$) while the fungi had no relationship with the soil pH.

5.3.4 AMF spores

The spore composition based on the morphology was different among the six sampling sites. Spores of different sizes and colors were present in all sites. Highest morphological diversity was a seen in Tree Tops followed by Central Florida, and the lowest diversity was seen in Jonathan Dickinson. *Glomus* was found in all the locations but was dominant in Nudgee, Logan and central Florida; *Scutellospora* was dominant in Tree Tops, and Jonathan Dickinson; and *Gigaspora* in Daintree (Fig. 5.2).
Figure 0. 2 AMF morphotypes extracted from the rhizosphere soil of *L. microphyllum* in the two continents. *Glomus* sp. spore (a, e, f); *Gigaspora* sp. with the bulbous sporangeneous cell (h, j); *Scutellospora* sp. showing the germination shield (b, d, g, i).
Table 0.2 Soil texture and mean ± standard error of the means of total colony forming units (counts × 10^7) of bacteria; (count × 10^3) of fungi per gram of soil in the invaded and native sites.

<table>
<thead>
<tr>
<th>Site</th>
<th>Soil Texture</th>
<th>Sand %</th>
<th>Silt %</th>
<th>Clay %</th>
<th>CFU of Bacteria g⁻¹ soil</th>
<th>CFU of Fungi g⁻¹ soil</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tree Tops Park, FL, US</td>
<td>Sandy loam</td>
<td>78</td>
<td>16.5</td>
<td>5.5</td>
<td>143.66 (13.00)</td>
<td>73.83 (12.27)</td>
</tr>
<tr>
<td>Central Florida US</td>
<td>Clay</td>
<td>13</td>
<td>40</td>
<td>47</td>
<td>152.5 (16.42)</td>
<td>61.16 (9.82)</td>
</tr>
<tr>
<td>Jonathan Dickinson, FL, US</td>
<td>Sand</td>
<td>98</td>
<td>2</td>
<td>0</td>
<td>43.83 (8.13)</td>
<td>46.66 (7.76)</td>
</tr>
<tr>
<td>Daintree Ferry, Queensland, AU</td>
<td>Silt loam</td>
<td>25</td>
<td>57.5</td>
<td>17.5</td>
<td>288 (16.56)</td>
<td>123.5 (13.63)</td>
</tr>
<tr>
<td>Logan Reserve, Queensland, AU</td>
<td>Sandy clay loam</td>
<td>67.5</td>
<td>20</td>
<td>12.5</td>
<td>103.83 (10.12)</td>
<td>50.5 (8.75)</td>
</tr>
<tr>
<td>Nudgee, Queensland, AU</td>
<td>Loam</td>
<td>50</td>
<td>32.5</td>
<td>17.5</td>
<td>153.5 (6.15)</td>
<td>49 (4.60)</td>
</tr>
</tbody>
</table>
Table 5.3 Mean (Std. Dev.) of the selected soil chemical characteristics in the native sites in Australia and invaded sites in Florida

<table>
<thead>
<tr>
<th>Variable</th>
<th>Central Florida (FL)</th>
<th>Daintree (AU)</th>
<th>Jonathan Dickinson (FL)</th>
<th>Logan (AU)</th>
<th>Nudgee (AU)</th>
<th>Tree Tops (FL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al (mg g⁻¹)</td>
<td>5.07 (0.61)a</td>
<td>2.14 (0.30)b</td>
<td>0.8 (0.07)c</td>
<td>2.55 (0.07)b</td>
<td>2.35 (0.11)b</td>
<td>1.38 (0.37)d</td>
</tr>
<tr>
<td>C %</td>
<td>4.03 (0.84)a</td>
<td>2.70 (0.44)a</td>
<td>3.02 (1.88)a</td>
<td>12.90 (2.43)b</td>
<td>4.28 (1.19)a</td>
<td>22.43 (4.15)c</td>
</tr>
<tr>
<td>Ca (mg g⁻¹)</td>
<td>0.41 (0.13)a</td>
<td>0.47 (0.11)a</td>
<td>3.35 (1.77)a</td>
<td>0.43 (0.25)a</td>
<td>0.09 (0.01)a</td>
<td>17.21 (6.31)b</td>
</tr>
<tr>
<td>Cu (µg g⁻¹)</td>
<td>8.42 (2.81)a</td>
<td>43.94 (12.31)b</td>
<td>1.84 (0.54)c</td>
<td>10.16 (0.57)a</td>
<td>6.74 (1.57)a</td>
<td>14.76 (4.02)a</td>
</tr>
<tr>
<td>Fe (mg g⁻¹)</td>
<td>2.92 (1.78)ad</td>
<td>15.86 (2.96)b</td>
<td>0.54 (0.25)a</td>
<td>8.34 (1.10)c</td>
<td>5.38 (1.86)ac</td>
<td>4.46 (1.91)d</td>
</tr>
<tr>
<td>K (mg g⁻¹)</td>
<td>0.16 (0.10)a</td>
<td>1.77 (0.35)b</td>
<td>0.09 (0.05)a</td>
<td>1.49 (0.30)b</td>
<td>0.12 (0.02)a</td>
<td>0.18 (0.05)a</td>
</tr>
<tr>
<td>Mg (mg g⁻¹)</td>
<td>0.20 (0.10)a</td>
<td>0.55 (0.18)b</td>
<td>0.21 (0.11)a</td>
<td>1.27 (0.17)c</td>
<td>0.25 (0.05)a</td>
<td>0.68 (0.27)b</td>
</tr>
<tr>
<td>Mn (mg g⁻¹)</td>
<td>0.02 (0.01)a</td>
<td>0.27 (0.14)b</td>
<td>0.02 (0.01)a</td>
<td>0.02 (0.00)a</td>
<td>0.04 (0.00)a</td>
<td>0.05 (0.03)a</td>
</tr>
<tr>
<td>N%</td>
<td>0.20 (0.26)a</td>
<td>0.18 (0.03)a</td>
<td>0.49 (0.10)b</td>
<td>0.71 (0.04)b</td>
<td>0.26 (0.10)a</td>
<td>1.27 (0.17)c</td>
</tr>
<tr>
<td>OM%</td>
<td>8.65 (1.09)a</td>
<td>8.07 (2.89)a</td>
<td>4.32 (0.90)a</td>
<td>35.50 (7.04)b</td>
<td>11.45 (4.71)a</td>
<td>44.42 (2.71)c</td>
</tr>
<tr>
<td>P (mg g⁻¹)</td>
<td>1.03 (0.25)</td>
<td>0.91 (0.16)</td>
<td>1.15 (0.09)</td>
<td>1.16 (0.05)</td>
<td>0.97 (0.06)</td>
<td>1.22 (0.16)</td>
</tr>
<tr>
<td>Zn (µg g⁻¹)</td>
<td>15.93 (8.25)a</td>
<td>27.91 (5.63)a</td>
<td>8.77 (1.86)b</td>
<td>37.80 (12.43)c</td>
<td>14.28 (3.77)a</td>
<td>23.21 (4.53)a</td>
</tr>
<tr>
<td>pH</td>
<td>5.77 (0.12)a</td>
<td>4.24 (0.15)b</td>
<td>6.57 (0.10)c</td>
<td>4.55 (0.14)d</td>
<td>4.01 (0.05)e</td>
<td>5.60 (0.06)a</td>
</tr>
</tbody>
</table>

Note: values in the same row followed by different letters represent significant difference at p<0.05.
5.3.5 Leaf nutrient status

Figure 0.3 Nutrient concentration in the leaves (Mean ± S E) of *L. microphyllum* collected from different sites. For N and P separately, different letters indicate significant differences in leaf concentration of N and P (P < 0.05; Tukey's test).

There was a significant positive correlation between the leaf N concentration and soil C (r = 0.48; p=0.003); Ca (r=0.63; p<0.0001); N (r=0.50; p=0.002); P (r = pH (r=0.71; p<0.0001); sand% (r=0.65; p<0.0001). Likewise leaf P concentration had a negative correlation with soil Al (r= -0.47; p = 0.1013); Cu (r= -0.34; p = 0.04); Fe (r = -0.49; p = 0.002); K (r= -0.40; p=0.01); silt (r = -0.65; p<0.0001) and clay % ( r = -0.55;
There was a significant positive correlation between leaf P concentration and soil C% (r = 0.75; p<0.0001); Ca (r = 0.82; p<0.0001); N% (r=0.81; p<0.0001); OM% (r=0.54; p=0.0006); P (r=0.54; p=0.0006); pH (r=0.64; p<0.0001) and sand % (r=0.73; p<0.0001).

5.4 Discussion

My goal was to determine if there was significant difference in the soil characteristics in the native and recipient habitats of *L. microphyllum*. There was a significant difference in the soil chemical, biological as well as physical characteristics in the two regions. These soil characteristics can, on their own or in association with other habitat features, promote the extensive growth of *L. microphyllum* in its recipient habitat in Florida. There was also a significant difference in the foliar nutrient concentration among the sites. My results show soil texture and pH to be the major factors influencing *L. microphyllum* growth.

My results show that *L. microphyllum* has adapted to nutrient poor highly acidic soils in its native range and invades slightly acidic soils in Florida. Along with strongly acidic conditions, the Australian soils have high concentration of Al, which is considered phytotoxic in strongly acidic soils. Acidic soil conditions are reported to enhance the presence of trivalent cation (Al$^{3+}$), the most toxic form of Al available to plants (Delhaaize and Ryan 1995; Lidon and Barreiro 2002; Kochian et al. 2005). My Central Florida site, where *L. microphyllum* was growing over sand mine spoil, had the highest concentration of Al. However this was not a restricting factor for *L. microphyllum* which could be because of the soil pH. Al toxicity in plants is widely studied in crop plants,
according to Kochian (1995), Al toxicity causes alterations of physiological and biochemical process of plants and consequently in their productivity. Plant species differ in their Al tolerance, but given that *L. microphyllum* grown in Australian soil had lower biomass allocation to the belowground structures (rhizomes and roots) compared to the plants grown in Florida soils (Volin et al. unpublished data), my results indicate the possibility of the “evolution of increased competitive ability hypothesis”. When *L. microphyllum* escaped the highly acidic soil environment to the sites in Jonathan Dickinson sandy sites, the plants could have evolved with lowered investment cost to defense and reallocation of the resources to growth and reproduction, increasing their colonizing success.

Even with varying soil nitrogen status, foliar N concentration was significantly higher in all south Florida plants compared to the Australian plants. Leaf N concentration is directly related to increase in relative growth rate (RGR) and photosynthetic capacity leading to increased plant productivity and litter decomposition in *L. microphyllum* (Chapter 3) and several other plant species (Vitousek 2004; Treseder 2008; Vitousek 2010; Chen et al. 2011). This could in part explain the higher growth of *L. microphyllum* in south Florida invaded areas compared to its native range in Australia. Similar results of higher nutrient concentration in the Florida plants compared to the Australian plants have been reported by Goolsby et al. (2006). *Lygodium microphyllum* is reported to be highly mycorrhizal in south Florida compared to its native range in Australia (Chapter 2), this higher degree of mycorrhizal colonization could play a role in the increased foliar N concentration.
Various soil and plant factors cause a significant influence in the soil microbial community, which have a fundamental role in nutrient cycling, plant growth and root health. It is widely reported that the rhizosphere community of different plant species growing in the same soil are distinct. Likewise an individual plant species can harbor different microbial communities in different soil types. A strong effect of soil texture on bacteria and fungi population was evident in this study. Daintree, in Australia with the highest percentage of silt harbored highest CFUs of bacteria and fungi, while Jonathan Dickinson in Florida with 98% sand had the lowest CFUs of bacteria and fungi. This kind of influence of soil texture on the structure of microbial population has been reported previously (Garbeva et al. 2004; Fang et al. 2005). An unexpected result was: the bacteria and fungi population remained uninfluenced by the soil organic matter, C% or soil pH which indicates a possible difference in the litter quality and secondary metabolites produced by the plant in its native range and invaded community.

My results indicate that *L. microphyllum* had a symbiotic relationship with multiple species of AMF depending on the site conditions. My two sites, Central Florida and Tree Tops which had higher diversities of spores are relatively disturbed sites compared to the other sites which had lower diversity of spores. This is an expected result and is in line with the Intermediate Disturbance Hypothesis (IDH) (Huston, 1979) which suggests that a less disturbed healthy ecosystem has lower diversity of arbuscular mycorrhizal fungi. I found that *Glomus*, which is reported to be the dominant and most abundant genus of AMF, was present in all sites but dominated in Logan and Nudgee in Australia and Central Florida sites. *Scutellospora* was dominant in Jonathan Dickinson, which has the worst case of *L. microphyllum* infestation in Florida and in the Tree Tops
Park. This indicates the influence of soil texture in the composition of AMF species in the soil. Even though spore morphology has been used to identify AMF species, it has been reported that this technique is not sufficient (Kruger et al. 2009). This study provides evidence that the mycorrhizal fungi composition is different in the native and recipient habitats along with the soil characteristics, but an in-depth analysis with the use of molecular technique is necessary to identify the AMF species and their relationship with *L. microphyllum*.

Overall, this study provides baseline information on the variation in the rhizosphere soil characteristics of *L. microphyllum* in its native and recipient habitat. My results indicate that Al sensitivity could be a determining factor that restricts the growth of *L. microphyllum* in the highly acidic soils rich in Al in its native range in Australia. Further research is necessary to gain an insight on the Al tolerance level of *L. microphyllum* and the role of Al concentration in soil in the growth limitation of *L. microphyllum*. I found that there was significant difference in the microbial populations and types in the regions, but I was not able to determine the specific roles of these microbes. However, these results are in line with the conclusion of Volin et al. (2010) who reported that escape from pathogenic soil microbial community could in part explain the extensive growth of *L. microphyllum* in the recipient habitats of Florida. My results indicate that *L. microphyllum* can be growing poorly in its native range in Australia because of the soil toxic effects associated with soil acidity and low foliar nitrogen concentration which in turn could affect the photosynthetic capacity of the plant.
References


Ferriter A 2001 Lygodium management plan for Florida: a report from the Florida Exotic Pest Plant Council's Lygodium Task Force. Florida Exotic Pest Plant Council,


CONCLUSIONS AND RECOMMENDATIONS

This study was conducted to determine why *Lygodium microphyllum* becomes a strong competitor in all hydrological, nutrient, light gradient habitats in south Florida compared to its native range in Australia. The studies presented in this dissertation provide baseline information and help to understand the complex feedbacks between exotic invasive species, soil microbial community and soil elements. I tested the hypothesis that the biogeochemical properties of Florida soils provide a more favorable condition for this species in the invaded region in Florida than in its native range. I did a cross-continent comparison of rhizosphere soil properties to determine if there were any specific characteristics in the soil which promote the invasiveness of this plant species in Florida.

Based on the results presented in the second chapter, *L. microphyllum* appears to be a strong host for arbuscular mycorrhizal fungi. However, this relationship with AMF varied with location, most probably by site hydrological conditions. *Lygodium microphyllum* had a stronger association with AMF in the dry areas of invaded regions in Florida compared to the flooded sites in Florida as well as its native range in Australia. The enhanced mycorrhizal fungi are also likely responsible for the greater P uptake and biomass accumulation in the control study. This strong association with mycorrhizae and an extensive belowground rhizome growth could in part explain efficient nutrient uptake leading to the competitiveness of *L. microphyllum* in nutrient poor Florida soils.

The green house study in the third chapter indicates that *L. microphyllum* is able to survive and grow in a wide range of soil pH; however, final biomass, relative growth rate, photosynthesis and specific leaf area were all significantly greater in soil pH 5.5 -
6.5 compared to the other treatments. Correspondingly, nitrogen concentration was also significantly related to these four plant parameters. Additionally root colonization by mycorrhizal fungi was significantly higher in soil pH 5.5-7.5 and lowest in plants growing in 4.5 or 8.0. Arbuscular mycorrhizal fungi colonization in roots was significantly correlated with plant growth parameters and nutrient concentration in the leaves. Comparison of soil characteristics in the invaded and uninvaded sites in three different locations in the fourth chapter shows that even after removal the effect of *L. microphyllum* may persist leaving behind a “legacy” influencing the belowground ecology. It causes a slight increase in soil pH, increase in soil organic matter and changes the ratio of bacteria and fungi population in the soil. This can have long-term effects on the restoration of the invaded sites or sites difficult or challenging for management. Cross continent soil characteristics comparison in the fourth chapter show that *L. microphyllum* can be growing poorly in its native range in Australia because of the soil toxic effects associated with soil acidity and low foliar nitrogen concentration which in turn could affect the photosynthetic capacity of the plant. Plant species differ in their Al tolerance, my results indicate the possibility of the “evolution of increased competitive ability hypothesis”. When *L. microphyllum* escaped the highly acidic soil environment to the sites in Jonathan Dickinson sandy sites the plants could have evolved with lowered investment cost to defense and reallocation of the resources to growth and reproduction, increasing their colonizing success. This study documents that that *L. microphyllum* can allocate up to 40% of the total biomass to the rhizomes, which remain unaffected by the different control techniques. Thus, *L. microphyllum* immediately regenerates from the
rhizomes after the use of management techniques such as fire, chemical spray, burning or cutting.

Overall, the results of this study could provide broader understanding in the ability of *L. microphyllum* being equally competitive in different habitat types in Florida. This study highlights that along with the characteristics of exotic plant species and native plant community, the understanding of invasive success of exotic plants needs the understanding of belowground community and ecology. It also provides information applicable for land managers responsible for protecting the Everglades, developing a sustainable control program towards minimizing the impacts of *L. microphyllum* as well as other exotic invasive species. Based on this study I recommend the following issues to be addressed in future studies:

- *Lygodium microphyllum* can form a very strong symbiotic relationship with AMF in its introduced environment in Florida. It is likely that this relationship is strongly influenced by site hydrological conditions, but this hypothesis needs to be tested in future research, especially when the Florida Everglades is undergoing a major hydrological shift as an effort for restoration. Future studies should also look into the possibilities of developing an integrated management plan which targets the micorrhizal fungi in the roots and rhizosphere of *L. microphyllum*.

- Prescribed burning, which causes a temporary rise in soil pH, is a widely used method to control *L. microphyllum*, but my study shows that *L. microphyllum* could be benefiting from the slight increase in soil pH resulting from fire as well as the release of nutrients that are associated with burning. Further
research should be done to determine the effect of fire on the mycorrhizal fungi and in soil with various pH levels.

- In my study, *L. microphyllum* had highest growth at neutral soil pH’s and began to show a significant decrease at a soil pH of 8.0, likely further growth reductions would happen in even more alkaline soils. Thus, raising soil pH may be a possible management option to explore in the future, but increasing the soil pH would need to be studied carefully for its potential adverse effects to native flora as well, including both native plants and soil microorganisms.

- My results indicate that *L. microphyllum* recruits different species of AMF in different sites. This relationship of *L. microphyllum* with AMF merits further research. In-depth analysis with the use of molecular technique is necessary to identify the AMF species and their relationship with *L. microphyllum*.

Finally, exotic species invasion will be a continuous threat to the Everglades ecosystem and will continuously challenge land managers and researchers. With the increased rate and number of exotic species invasion, herbicide treatment will most likely become the widely used technique to control invasive species in the future. But, the effort of invasive species control should not ignore the belowground effects of invasive plants in the Florida Everglades ecosystem, especially when it is undergoing a major hydrological shift as an effort for restoration. Understanding the soil nutrient and microbial dynamics will provide opportunities to develop a successful integrated management technique. I believe a complete understanding of the soil ecosystem is necessary before adopting a management technique to achieve a successful long-term invasive species management strategy in the south Florida Everglades.
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