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Sex-specific Fungal Communities of the Dioicous Moss Ceratodon purpureus

by

Mehmet Ali Balkan

A thesis submitted in partial fulfillment of the requirements for the degree of

Master of Science in Biology

Thesis Committee: Todd N. Rosenstiel, Chair Sarah M. Eppley Daniel J. Ballhorn Kenneth M. Stedman

Portland State University 2015

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Abstract

Mosses display a number of hallmark life history traits that influence their ecology at the population and community level. The long lived separation of sexes observed in the haploid gametophyte (dioicy) is one such feature of particular importance, as it is observed in the majority of bryophytes and creates intraspecific specialization of male and female individuals.

The prevalence of sexually dimorphic mosses raises the possibility of sex-specific interactions with fungi as observed in some vascular plants. Here I investigated how moss sex shapes fungal communities associated with gametophytic tissues of the ubiquitous moss, *Ceratodon purpureus*. Using greenhouse populations of *C. purpureus* grown in a common garden, I examined fungal community structure and overall abundance of fungal biomass associated with male and female individuals from multiple populations. I hypothesized that individual mosses would harbor unique fungal communities based on their sex, and that overall fungal biomass associated with host tissues would differ significantly due to differences in morphological and physiological characteristics between the sexes. I found that fungal community composition and overall abundance (i.e. biomass) differ between male and female individuals of C. *purpureus*, and that sex-specific patterns are retained across individuals from three different populations. This work provides a first glance at how genetically based sexual systems in early land plants influence affiliated fungal community composition.

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Chapter 1: Introduction

Patterns in Ecological Communities

All life on earth coexists not only within populations of the same species, but with many other species groups forming interactive living collectives known as communities. Communities operate based on complex, intersecting, and rarely discrete principles referred to as assembly rules (Diamond, 1975; Keddy, 1992; Weiher & Keddy, 2001) that influence their functionality, stability, and longevity. These governing principles should not be viewed simply as black and white rules of engagement, but as theoretical frameworks to describe and measure community dynamics that are based on commonly observed phenomena. In the modern approach, four main processes make up the umbrella under which assembly rules unfold: selection, drift, speciation, and dispersal (Vellend, 2010). These four pillars of community ecology may be familiar as they are reflections of "the big four" in population genetics. Consideration of each of the four processes is important but here a primary focus on selection will be pursued. In particular, a suite of selective elements defined in part by external species interactions called ecological filters is considered.

The structure and functional attributes of biological communities are governed in part by ecological or environmental filters (biotic and abiotic) that collectively act to shape assembly trajectories (Keddy, 1992; Kelt *et al.*, 1995;

Diaz et al., 1998). Species interactions are among the most important filters shaping communities and are often hierarchically structured (Baumeister & Callaway, 2006; Kunstler et al., 2012), reflecting their relative strength in influencing community outcomes. Within this continuum, foundation species are those members of a community whose disproportionately large effect on community processes can cause shifts in diversity and structure (Ellison et al., 2005; Bangert et al., 2006; Angelini et al., 2011; Schöb et al., 2012). In tandem with the robust development of our understanding of foundation species, a new movement has begun in which the synergy between evolutionary biology and ecology has come to a new level (Johnson & Stinchcombe, 2007). The genes-toecosystem or community genetics approach seeks to understand how genetic properties of individuals and populations can manifest as phenotypic traits that emanate through different organizational levels up to the ecosystem level (Shuster et al., 2006; Whitham et al., 2006; Bangert et al., 2006). In this emerging field, the traditional *individual phenotype* is expanded to encompass a predictable and heritable *community phenotype*: the result of certain critical traits of the foundation species that have a strong genetic basis. Perhaps the best documented example of heritable community phenotypes in action is in the *Populus* system (Whitham *et al.*, 2006). In this system the poplar tree acts as a foundation for multitrophic communities, and variation in certain aspects of leaf chemistry-controlled by a single quantitative trait locus-across individual poplar genotypes have been shown to create cascading effects across each level of the

community examined. In this particular case fungal endophyte, arthropod, and aquatic macroinvertebrate communities associated with poplars vary predictably in their composition based on condensed tannin levels in the leaves of different host genotypes (Wimp et al., 2005; Bailey et al., 2005). These studies and others (Dungey et al., 2000; Whitham et al., 2003; Hochwender & Fritz, 2004; Johnson & Agrawal, 2005; Brändle & Brandl, 2006; Tovar-Sánchez & Oyama, 2006), highlight the power of community genetics to uncover how the genetic basis of foundation species traits can influence community dynamics at different levels. and thus aid in the characterization of assembly rules. However, it is critical to continue developing this approach with consideration for other traits and further experimentation in other systems. Bryophytes present one such promising model system for the study of foundational traits in communities. Certain features of this group such as their reproductive systems and unique genetic architecture, in addition to their prominent roles within ecological communities, support this prospect and are further discussed in the following sections.

Bryophytes: A Brief Introduction

Bryophytes are an ancient lineage of plants that represent the closest extant relatives of the first terrestrial plants and are widely distributed across most ecosystems on earth (Herzog, 1926; Kenrick & Crane, 1997). Thought to have originated as early as the mid Ordovician period (Wellman & Gray, 2000;

Rubinstein *et al.*, 2010), the bryophytes are composed of three major groups; the liverworts (Marchantiophyta), the hornworts (Anthocerotophyta), and the mosses (Bryophyta), the later of which boasts the greatest diversity with over 12,000 described species (Shaw et al., 2005). Bryophytes exhibit a number of unique morphological and physiological traits that truly set them apart from other embryophytes. These traits include extreme desiccation tolerance (poikilohydry), lack of complex vasculature and true roots, a haploid-dominant life cycle, and a large proportion of dioicous (dioecious haploids) species, among others. Bryophytes have maintained their ancient lifestyle across many evolutionary epochs with little deviation, embodying evidence for truly successful strategies that have persisted alongside the explosion of vascular plant diversity in later periods (Shaw *et al.*, 2011). Additionally, over the course of their evolutionary history bryophytes have developed highly complex relationships with other biota such as mutualistic associations with nitrogen-fixing cyanobacteria (Lindo et al., 2013), fungi (Davey & Currah, 2006), and a potential pollinator like syndrome with collembola (Rosenstiel et al., 2012), among others. Through some of these relationships and their own biophysical and physiological properties, bryophytes can have a major influence on their surrounding environment, in some cases on a global scale. The section that follows details some of the ways in which bryophytes 'engineer' abiotic and biotic elements of communities and ecosystems, further underscoring their utility as a model system in community ecology.

Bryophytes as Foundation Species

Owing to their distinctive lifestyles among land plants, bryophytes fulfill unique ecological roles in ecosystems; acting as pioneer species that are some of the first plants to colonize disturbed habitats, and often growing in places and under conditions that other plants cannot (Vanderpoorten & Goffinet, 2009). Bryophytes, and in particular mosses, contribute a number of critical ecosystem services to the environs they inhabit (Turetsky, 2003; Cornelissen et al., 2007; Lindo & Gonzalez, 2010; Turetsky et al., 2012). Among the most notable contributions of mosses are their capacity to support diverse, multitrophic, biological communities (Lindo & Gonzalez, 2010), influence nitrogen and phosphorus cycling at a landscape scale (Chapin III *et al.*, 1987; Turetsky, 2003; Cornelissen *et al.*, 2007; Rousk *et al.*, 2014), augment abiotic soil conditions such as moisture, temperature, and redox potential, and make significant contributions to net primary productivity in many terrestrial systems (Longton, 1992; Gornall et al., 2007; Blok et al., 2011; Rousk et al., 2014). The role of mosses in global change biology is perhaps best documented in peatland systems where the genus *Sphagnum* dominates and is responsible for the storage of more than one third of the worlds terrestrial carbon (Loisel et al., 2012). Though peatlands have long served as the model system for studying 'bryogeochemistry' (Damman, 1986; Gorham, 1991; Dunfield et al., 1993; Kang et al., 2001; Turetsky et al., 2002; Loisel et al., 2012), an ever-growing body of work is highlighting the global importance of the Bryosphere in many other

systems including Antarctica (Pressel, 2009), the Arctic (Longton, 1997), and boreal and temperate forests (DeLucia *et al.*, 2003; Turetsky *et al.*, 2012).

Due to their disproportionately large ecological influence mosses can be considered as foundation species (Whitham et al., 2006; Lindo & Gonzalez, 2010; Tuba *et al.*, 2011). This foundational capacity of the mosses is especially evident in a community ecology context, whereby the moss acts as critical host to a multitude of other life forms ranging from bacteria, fungi, and protists, to tardigrades, rotifers, mites, springtails and other invertebrates. This biologically rich, and dynamic matrix is collectively referred to as the Bryosphere, and is an integral component of global detrital systems that drive biogeochemical cycling (Lindo & Gonzalez, 2010). Microbial constituents of the bryosphere are of particular interest as they represent a more basal level of the trophic web, and have far reaching functional influence. Though some microbial components of the bryosphere such as cyanobacteria (Adams & Duggan, 2008; Lindo et al., 2013; Arróniz-Crespo et al., 2014; Stuiver et al., 2015) have been relatively well studied, the factors that drive relationships with other major players-specifically fungi-are still largely unknown. The prevalence of the fungi in the bryosphere (Davey & Currah, 2006) beckons more attention to the study of this relationship. What follows is a brief survey of what we know about moss-fungi interactions.

Fungal Components of the Bryosphere

The bryosphere is an integrated detrital network driven by multitrophic communities that operate on a host bryophyte framework. Like other detrital systems, microbial activity is at the core of the bryosphere. In particular, taxonomically and functionally diverse fungal components have been observed to influence the bryosphere through a spectrum of interactions with host mosses ranging from seemingly mutualistic to parasitic, though the nature of interaction between the vast majority of bryophillous fungi and their hosts has yet to be revealed (Davey & Currah, 2006; Kauserud et al., 2008; Davey et al., 2009, 2012b). Members of all five major fungal lineages have been observed inhabiting the moss phyllosphere, constituting functionally diverse assemblages (Davey et al., 2012b) that have the propensity to drive ecosystem processes (Christensen, 1989). Within the vast diversity of bryophilous fungi, members of the phylum Ascomycota appear to be the dominant group in most reports (Felix, 1988; Döbbeler, 2002; Davey & Currah, 2006; Kauserud et al., 2008; Ptaszyńska et al., 2009; Davey et al., 2012a,b; Döbbeler & Hertel, 2013). Furthermore, many bryophilous ascomycetes are highly specialized to specific parts of moss hosts (e.g. rhizoids, hyaline hair points, apical stems)(Döbbeler, 2002; Ptaszyńska et al., 2009; Döbbeler & Hertel, 2013). A study by Döbbeler et al. (2009) found phylogenetic evidence for multiple origins of specialized bryophyte-ascomycte associations, underscoring the intimate coevolution of bryophytes and fungi. Another example of a highly specialized, ancient bryophyte-fungi interaction is

the relationship between arbuscular mycorrhizal fungi (AMF) and some bryophytes (Read et al., 2000; Ligrone et al., 2007; Wang et al., 2010; Desirò et al., 2013). As in vascular plants, AMF form a symbiotic association with bryophytes that enables enhanced nutrient uptake by the host through fungal hyphae infecting the host's tissues in exchange for photosynthetically derived carbon that is translocated to the fungus (Read *et al.*, 2000; Ligrone *et al.*, 2007; Desirò et al., 2013; Pressel et al., 2014). These mutualisms have been well documented in the liverworts (Marchantiophyta) and hornworts (Anthocerotophyta), and although AMF have been previously observed in the tissues of mosses (Rabatin, 1980; Parke & Linderman, 1980; Zhang & Guo, 2007), there is as of yet no evidence that functional mutualisms exist between AMF and mosses suggesting that mosses may have unique interactions with fungi relative to other bryophytes. Despite a growing body of knowledge regarding the interactions between bryophytes and fungi, investigations of whole fungal communities associated with bryophytes are scarce (Kauserud et al., 2008; Davey et al., 2012a,b). A handful of recent studies have provided insight into some of the factors influencing the structural characteristics of mossassociated fungal communities (Davey et al., 2012a,b), but further investigation is needed to refine our understanding of how unique biological features of bryophytes influence their interactions with fungi in a fundamentally different way than vascular plants. The reproductive biology of mosses offers an exceptional opportunity to examine how a distinctive, ecologically important set of traits in

bryophytes can be a formative force in shaping interactions with fungi in the bryosphere. An exploration of these reproductive traits will reveal their potential to act as biotic filters for community assembly.

Bryophyte Reproductive Ecology

Given the complexities of bryophyte reproductive ecology, first I will discuss some fundamental concepts in bryophyte reproduction and how some special features make for unique biotic interactions. Bryophyte reproduction involves a dominant stage that is opposite of their vascular cousins. The haploid gametophyte dominates the majority of the life cycle and at maturity bears sexual structures called archegonia (females) and antheridia (males). In dioicous bryophytes, these structures appear on separate individuals. When sexual structures reach maturity, biflagellate sperm produced in the antheridia swim across water films on the surface of gametophytes in search of the female ovum. Sperm cells are guided towards their female counterparts by sucrose gradients released from the archegonia (Pfeffer, 1884). When fertilization is successful, a diploid sporophyte begins to develop and is largely nurtured by the maternal gametophyte. It is within the developing sporophyte that meiosis occurs producing haploid spores of distinct sex. Spores are released from the mature sporophyte and viable individuals germinate to produce the primary stage of bryophyte life: chlorotic, filamentous networks called protonema. The protonemal networks function in a fashion similar to a fungal mycelium or vascular plant's

root mass: branching out and extending in many directions to seek out nutrition and water. When ideal conditions are met the protonemal network begins to restructure into a denser, often initially thalate, gametophytic tissue matrix, returning once again to the beginning of the bryophyte life cycle (Vanderpoorten & Goffinet, 2009).

As with most other species that are unisexual, male and female bryophytes display a host of sex-specific traits that influence their relationships with the external environment and biota. The occurrence of sexually dimorphic ecology is widespread and well documented in the animal kingdom (Darwin, 1871; Lailvaux & Vincent, 2007), but its implications for plants have only recently begun to be appreciated (Eppley et al., 2009; Varga, 2010; Petry et al., 2013; Vega-Frutis et al., 2013), and have likely been mostly overlooked due to the small proportion of vascular plants that maintain separate sexes (Renner & Ricklefs, 1995). Sexual dimorphisms present a unique form of intraspecific trait diversity that is inherently tied to reproductive fitness, but whose influence can extend beyond the context of reproduction. The literature describing the dimorphic ecology of dioecious angiosperms demonstrates that male and female individuals interact with and change their biotic and abiotic environment in different ways. For example, males and females typically have different resource requirements and allocation patterns which can directly and specifically influence their physical distribution and success in heterogeneous environments (Bierzychudek & Eckhart, 1988). Furthermore, sex-specific differences in

beneficial biotic interactions such as those involving mycorrhizal fungi (Eppley *et al.*, 2009; Varga, 2010), as well as antagonistic interactions, such as those involving herbivores and pathogens (Ågren *et al.*, 1999; Vega-Frutis *et al.*, 2013), have the potential to not only shape populations of the plants in question but also that of the interacting biota. In contrast to their vascular relatives, bryophytes exhibit a much greater degree of dioicy with over 60% of taxa producing separate sexes (Wyatt & Anderson, 1984), and thus the potential for sex-specific biotic interactions is much higher in bryophytes.

Bryophytes are typically sexually dimorphic and can have skewed distributions in natural populations (Fuselier & Stark, 2004; Stark *et al.*, 2010). Important sex-specific physiological and morphological differences such as size (Shaw & Beer, 1999), prezygotic investment (Horsley *et al.*, 2011), time to sexual maturity (Stark, 2002), and emission of volatile organic compounds (VOCs) (Rosenstiel *et al.*, 2012), have been observed in bryophytes. Furthermore, bryophytes possess unique sex chromosomes on which crossing over is completely suppressed (Bachtrog *et al.*, 2011); contributing greatly to the accumulation of sexually antagonistic traits in the haploid gametophyte stage predominant in the bryophyte life cycle (McDaniel & Perroud, 2012). Such extreme sex-based genetic partitioning supports the prospect of intraspecific differences in ecological interactions based on sex. Recently, Rosenstiel *et al.* (2012) found that common microinvertebrate inhabitants of the moss *Ceratodon purpureus* were disproportionately attracted to volatile compounds emitted by

female individuals. Further, the study revealed that overall volatile organic compound (VOC) emissions formed distinct profiles among males and females. These findings coupled with others showing increased fertilization events with the addition of collembolans to patches of *C. purpureus* suggest a possible pollinator-like syndrome in the mosses, adding an additional layer of complexity to bryophyte reproduction

With the interest of exploring the prospect of other sexually dimorphic ecological interactions, the moss *Ceratodon purpureus* is an excellent candidate species to be used as a model. In addition to the sex-specific biotic interactions already observed in this species mentioned above, a number of other critical features bring *C. purpureus* to center stage as an ideal subject for further elucidating how a genetically based like sex may influence other community components such as fungi.

Ceratodon purpureus

Ceratodon purpureus is a dioicous, acrocarpous moss in the family Ditrichaceae with a cosmopolitan distribution across all continents including Antarctica (McDaniel & Shaw, 2005). *C. purpureus* is often found growing in urban environments, poor soils, and recently burned areas, and evidently is tolerant of a wide range of environmental conditions including soils laden with heavy metals (Jules & Shaw, 1994) and extreme UV (Clarke & Robinson, 2008).

For many years *C. purpureus* has been used a model to study physiology (Algarra *et al.*, 1993; Kohn *et al.*, 1994), population genetics (Skotnicki *et al.*, 2004; McDaniel, 2005; McDaniel *et al.*, 2007), and sexual ecology (Shaw & Gaughan, 1993; Shaw & Beer, 1999; Rosenstiel *et al.*, 2012). Along with the recent effort to sequence its genome (http://jgi.doe.gov/why-sequence-ceratodonpurpureus-moss/), and an increasing understanding of its general ecology, *C. purpureus* is increasingly becoming a robust model system for studying community dynamics in the bryosphere.

Goals of this thesis

Expanding our understanding of the nature of biological communities and the forces that shape them is a central goal of ecology that has far reaching implications in many other fields. The genes to ecosystem approach, which is focused on tracking how the genomic patterns of foundation species can have cascading effects across different levels of a community, shows great promise in untangling the basis of assembly rules in a manner in which traditional approaches to community ecology can not. Bryophytes present an exceptional model for the study of community genetics because of their foundational roles within the complex, multitrophic communities, as well as unique genomic features that allow an unobscured view into traits that influence these communities. Specifically, the dominant haploid gametophyte life stage coupled with a distinctive sex chromosomal system in dioicous species permits the observation

of genetically rooted, sex-specific traits directly in the individual phenotype. While a small but growing number of studies have begun to elucidate how plant sex influences biotic interactions in vascular systems, remarkably few focus on other prolific groups such as the bryophytes. Furthermore, to my knowledge, there are no studies investigating how plant sex structures microbial *communities* (as opposed to individual interactions) in any part of the kingdom. Given the wellestablished importance of dynamics in the plant-microbe continuum in global processes, unveiling the basis of these dynamics remains a critical task in community ecology.

The goals of the research presented herein are to investigate how sex as a concrete, genetically governed trait in a haplodioicous foundation species influences the composition and overall biomass of associated fungal communities. I predict that both community composition and biomass will differ between fungal communities associated with males and females due to sexually dimorphic characteristics.

Chapter 2: Materials and Methods

Study System

Greenhouse populations of *C. purpureus* were generated using tissue collected from three geographically isolated populations. Two of the populations were within the urban boundaries of Portland, OR: Neuberger Hall (NH) (45°30'N, 122°41'W), located on the downtown campus of Portland State University, and Northeast 35th (35th) (45°32'N, 122°37'W), located 8 km northeast of downtown. The third population (UCUT) (45°36'N, 123°0'W) was located 32 km west of the Portland metropolitan area. Climatic conditions were similar in all three sites where *C. purpureus* was found growing on moderately heavy clay soils with a shallow organic horizon. Single intact shoots (~2 cm) from 8 sexually expressing males and females from each population were vegetatively propagated in 10 cm² pots on growth media composed of 2:1 peat/sand (Sun Gro). Pots containing males and females were organized in blocks with ~2 cm between pots within two separate trays, and rotated regularly to avoid tray effects. A plastic barrier was placed between benches housing males and females to prevent fertilization events. Mosses were grown for ~2 years in common garden under semi-exposed greenhouse conditions that were not completely isolated from the external environment, but likely had more limited exposure to fungi than mosses growing outdoors. Temperatures were typically within 2°C of outside temperatures. Pots were misted from above with tap water

four times per day for 2 minutes, at 6-hour intervals by an automated irrigation system.

DNA Extraction and PCR

Total DNA was extracted from single shoots of *C. purpureus* using the REDExtract-N-Amp Tissue PCR Kit (Sigma-Aldrich) according to the manufacturer's protocol with the following changes: moss shoots were submerged in 20 µl of 1:4 tissue preparation and extraction solutions and incubated at RT for 10 min followed by 10 min at 95°C, and 10 min at 65°C. Following incubation, 20 µl of neutralization solution was added to each sample and mixed by pipetting. All residual plant material was physically removed from finished extractions using sterile pipette tips. Finished extractions were subsequently stored at 4°C. The fungal ITS region was amplified by PCR using primers ITS1F and ITS4 in duplicate 20 µl reactions containing 10 µl REDExtract-N-Amp PCR Reaction Mix (Sigma-Aldrich), 0.4 µl 6-FAM labeled forward primer ITS1F (CTTGGTCATTTAGAGGAAGTAA), 0.4 µl reverse primer, ITS4 (TCCTCCGCTTATTGATATGC), 1 μ l template DNA, and 8.2 μ l PCR H₂0). Reactions were run in a Bio-Rad DNA Engine[®] thermal cycler (Bio-Rad Laboratories) under the following cycling conditions; 94°C for 1 min, 35 cycles of 94°C for 1 min, 53°C for 1 min, 72°C for 1 min, and 1 cycle at 72°C for 8 min. PCR products from duplicate reactions were combined and purified using an

UltraClean PCR Clean-Up Kit (Mo Bio Laboratories), and clean PCR products stored at 4°C until further use.

T-RFLP Analysis

Restriction digests were prepared in 20 μ l reactions containing 12 μ l purified PCR products, 1 μ l restriction enzymes Hae III or Hinf I (Thermo Fisher Scientific), 2 μ l digest buffer, and 5 μ l PCR H₂O. Digests were incubated at 37°C for 4 hours, followed by a 20 minute deactivation period at 80°C. Digest products were purified using an UltraClean PCR Clean-Up Kit (Mo Bio Laboratories) and stored at 4°C. Clean digest products were transferred to a 96-well plate, concentrated in a Savant SpeedVac (Holbrook, NY), and shipped to OSU CGRB Core Labs for fragment analysis. Samples were resuspended in .9 μ l Hi-Di formamide and .1 μ l MapMarker 1000 fluorescent standard and analyzed on an ABI 3730 sequencer. Resulting fluorescent terminal restriction fragment (fTRF) lengths were determined using GeneMapper 5 software (Life Technologies).

T-RFLP Data Processing

All fungal community profiles (electrophoretograms) were manually inspected and peaks smaller than 50bp and larger than 800bp were removed to reflect the range in which the MapMarker 1000 standard can be used to accurately predict fragment sizes. In order to distinguish true peaks representing fungal OTUs from noise, raw fragment analysis data were refined using the

constant percentage threshold technique whereby all peaks whose percent area contribution to the total area of a profile fell below 0.5% were discarded prior to further analysis. I found that using a percentage threshold below 1% yielded more accurate results by increasing the detection of relatively less abundant fTRFs (see Blackwood *et al.*, 2007). All remaining peaks were aligned to 1bp using the Treeflap macro

(http://www.sci.monash.edu.au/wsc/staff/walsh/treeflap.xls) in Excel. Global singletons (fTRFs occurring only once within the entire dataset) were removed prior to ordination and similarity analyses to minimize the inclusion of PCR artifacts and miscalled electrophoretic reads (fTRF drift) (Culman *et al.*, 2008). All qualified fTRFs were compiled into presence-absence matrices based only on occurrence, since methodological limits of T-RFLP do not allow for accurate estimation of OTU abundance based on fluorescence intensity (Schütte *et al.*, 2008).

Fungal Biomass Estimation by Ergosterol Quantitation

Whole gametophytes of each *C. purpureus* individual were collected, washed thoroughly in deionized water to remove soil and coarse debris, frozen at -80°C for 48 hrs, and lyophilized. 100-200 \pm 0.1 mg of lyophilized tissue was placed in 2 ml microcentrifuge tubes with one 3 mm zirconium bead and ground to a fine powder using a bead beater at 2500 RPM for 40 seconds. Ergosterol was extracted from powdered tissues following the methods of Dahlman *et al.* (2002) with the following changes. Samples were suspended in 1 ml HPLC grade MeOH, agitated in an orbital shaker for 1 hr at 320 RPM in darkness, and subsequently allowed to precipitate overnight at 4°C. Extractions were then centrifuged for 1 hr at 14,000 RPM and 4°C in an Eppendorf Centrifuge 5402. Following centrifugation, the supernatant from each extraction was transferred to a fresh 2 ml microcentrifuge tube and centrifuged for an additional 10 min at 14,000 RPM and 4°C (Eppendorf Centrifuge 5402). The supernatants from this secondary centrifugation were filtered through 0.2 μm acrodiscs into amber autoanalyzer vials and stored at 4°C until HPLC analysis. Extractions were analyzed following Davey *et al.* 2012 on an 1100 Series HPLC (Agilent Technologies, Waldbronn, Germany). The limit of detection was 5.59 μg.

Statistical Analysis

The compositional similarity between individual communities was estimated by calculating Bray-Curtis dissimilarity values (Bray & Curtis, 1957), which were consequently ordinated by non-metric multidimensional scaling (NMDS). Compositional differences between fungal communities associated with male and female *C. purpureus* individuals were statistically compared by analysis of similarity (ANOSIM) and permutational multivariate analysis of variance (PERMANOVA). OTU richness estimates and extrapolation were performed using Mao Tau, Chao2, and ICE estimators. The Mao Tau algorithm allows the calculation of estimated species richness based on an infinite number of randomized resampling schemes during sample-based rarefaction (Colwell et al., 2004). The non-parametric asymptotic species richness estimators Chao2 and ICE, were chosen based on their sensitivity to rare species (Chao, 1987; Lee & Chao, 1994), and because they do not assume any particular species abundance distributions making them suited for incidence based data. To assess the robustness and adequacy of sampling efforts to capture overall diversity of the sampled communities, sample completeness, sample-size-based, and coveragebased rarefaction and extrapolation curves were generated using Hill numbers according to Chao et al. (2014) using the iNEXT package for R. The influence of sex, population, and the interaction between sex and population on ergosterol concentration in moss tissues was assessed by two-way ANOVA. A post-hoc Tukey HSD test (p<0.05) was performed in order to determine significant differences between populations. All statistical analyses were performed using R statistical environment v. 3.0.2 (R Development Core Team, 2008) and EstimateS v. 8 software (Colwell, 2013).

Chapter 3: Results

Fungal Community Composition

Fungal community composition was statistically significantly different

based on C. purpureus sex in TRFLP community profiles generated by both Hae

III and Hinf I restriction enzymes (ANOSIM: R=0.134, p=0.003; R=0.102,

p=0.006, respectively). PERMANOVA analyses further support statistically

significant differences between fungal community composition based on sex and

population (Tables 1a, 1b).

Table 1a. Permutational multivariate analysis of variance between fungal community profiles generated by Hae III. Differences between communities associated with male (n=23) and female (n=24) individuals from three different populations (NH, 35th, UCUT). Values for degrees of freedom (df), sum of squares (SS), mean squares (Mean Sq), F-ratio (F), r-squared (R2), and p-value (p) are displayed. Bold values indicate statistical significance.

Hae III	df	SS	Mean Sq	F	R2	р
Sex	1	0.83	0.83	3.48	0.07	0.002**
Population	2	0.88	0.44	1.85	0.07	0.012*
Sex x Population	2	0.52	0.26	1.11	0.04	0.318
Residuals	40	9.52	0.23		0.80	
Total	45	11.76			1	

Significance tests are shown in bold, and *0.01 $\leq p < 0.05$; **0.001 $\leq p < 0.01$.

Table 1b. Permutational multivariate analysis of variance between fungal community profiles generated by Hinf I. Differences between communities associated with male (n=23) and female (n=24) individuals from three different populations (NH, 35th, UCUT). Values for degrees of freedom (df), sum of squares (SS), mean squares (Mean Sq), F-ratio (F), r-squared (R2), and p-value (p) are displayed. Bold values indicate statistical significance.

Hinf I	df	SS	Mean Sq	F	R2	р
Sex	1	0.66	0.66	3.42	0.06	0.004**
Population	2	1.33	0.66	3.43	0.12	0.001**
Sex x Population	2	0.66	0.33	1.71	0.06	0.047*
Residuals	41	7.95	0.19		0.74	
Total	46	10.62			1	

Significance tests are shown in bold, and *0.01 $\leq p < 0.05$; **0.001 $\leq p < 0.01$.

NMDS ordination indicates that this sex-based difference was due primarily to aggregate differences in overall fungal composition rather than high specificity of fungal communities on either sex (Fig. 1a, 2a).

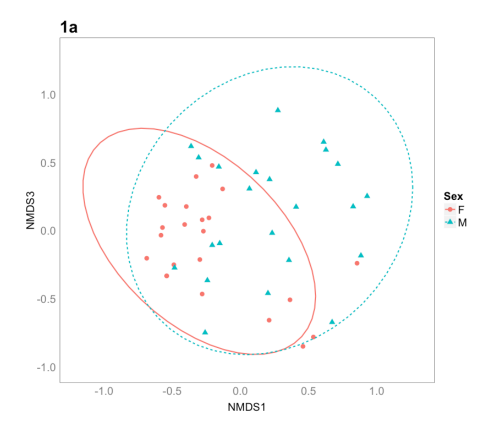


Figure 1a. NMDS ordination of Hae III fungal community profiles by sex (3D Stress = .16). Fungal communities associated with individuals from all three *C. purpureus* populations examined are represented. Red circles and blue triangles represent female and male associated fungal communities respectively. Solid and dashed line ellipses indicate 95% confidence intervals for female and male associated communities respectively.

When ordinations are coded by both sex and *C. purpureus* populations, the same sex-based pattern is evident across all three populations examined (Fig. 3b, 4b).

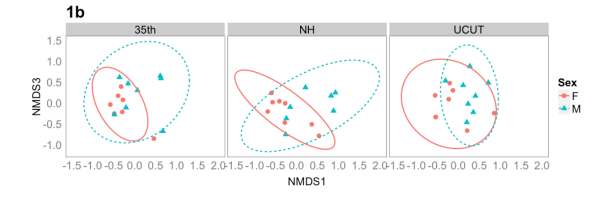
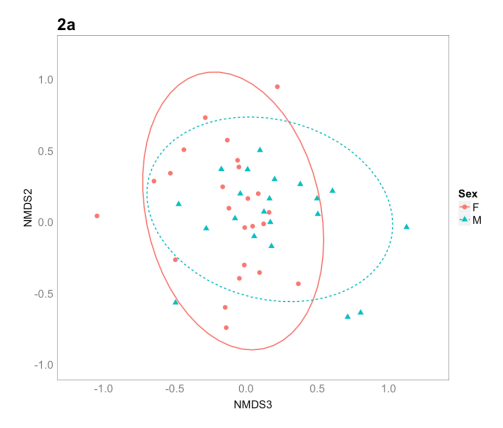
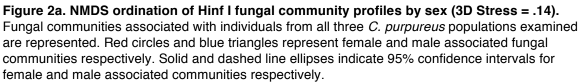


Figure 1b. NMDS ordination of Hae III fungal community profiles by sex and population (3D Stress = .16). Fungal communities associated with individuals from all three *C. purpureus* populations examined are represented. Circles and triangles represent female and male associated fungal communities respectively. Each of three panels represents fungal communities associated with *C. purpureus* individuals from 35th, NH, and UCUT populations respectively. Solid and dashed line ellipses indicate 95% confidence intervals for female and male associated communities respectively.





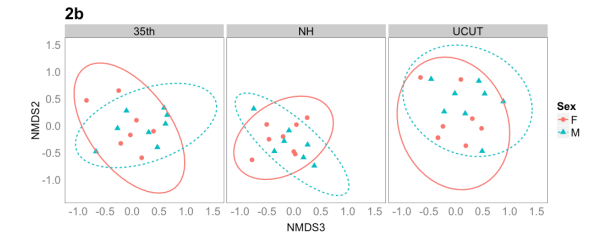
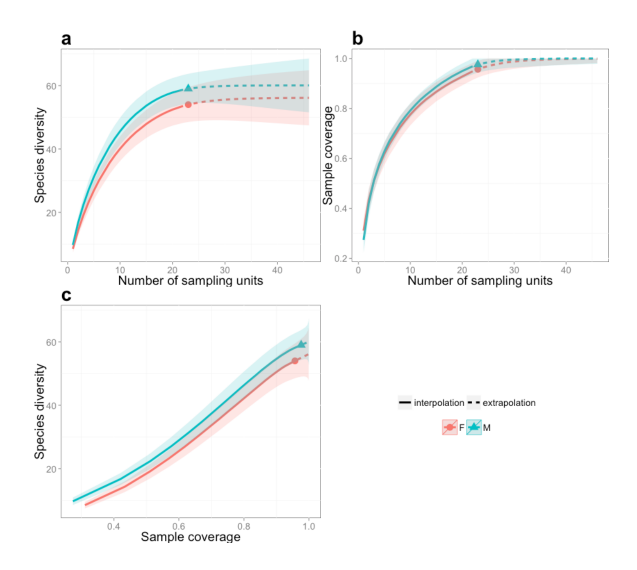
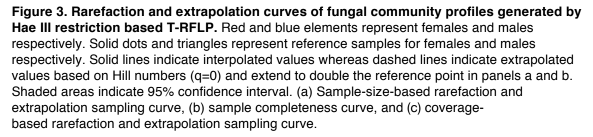


Figure 2b. NMDS ordination of Hinf I fungal community profiles by sex and population (3D Stress = .14). Fungal communities associated with individuals from all three *C. purpureus* populations examined are represented. Circles and triangles represent female and male associated fungal communities respectively. Each of three panels represents fungal communities associated with *C. purpureus* individuals from 35th, NH, and UCUT populations respectively. Solid and dashed line ellipses indicate 95% confidence intervals for female and male associated communities respectively.

Sample-based rarefaction and extrapolation curves for community profiles generated by both enzymes form asymptotes just beyond reference points, indicating that sampling efforts were adequate to accurately capture the diversity of each community without further sampling (Figs. 5a, 6a). Sample completeness curves, which asses the degree of completeness (based on coverage) for a given sampling effort, also form asymptotes close to the reference points indicating that further sampling efforts will not yield a much greater level of sample completeness (Figs. 5b, 6b). Finally, coverage-based rarefaction and extrapolation curves show increased sample coverage with increased sampling effort up to the reference points for each set of community profiles (Figs. 5c, 6c). Since there is only a slight increase in coverage when the reference sample is doubled, as indicated by the sample completeness curves (Figs. 5b, 6b), extrapolated portions of the coverage-based sampling curves are almost invisible signifying that almost no increase in diversity would be expected from further sampling.





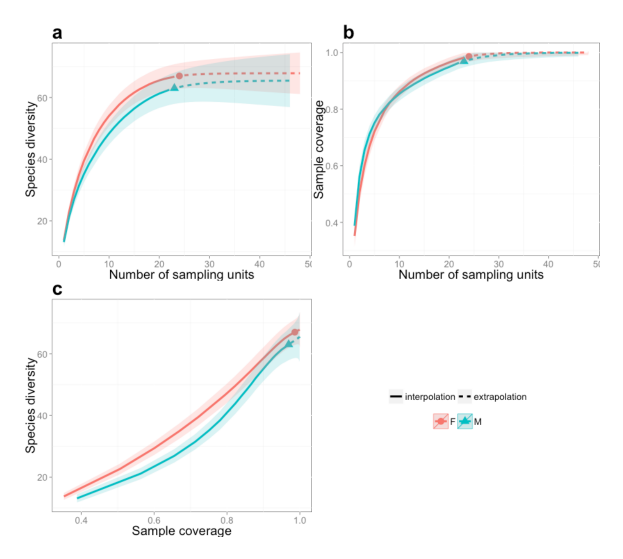


Figure 4. Rarefaction and extrapolation curves of fungal community profiles generated by Hinf I restriction based T-RFLP. Red and blue elements represent females and males respectively. Solid dots and triangles represent reference samples for females and males respectively. Solid lines indicate interpolated values whereas dashed lines indicate extrapolated values based on Hill numbers (q=0) and extend to double the reference point in panels a and b. Shaded areas indicate 95% confidence interval. (a) Sample-size-based rarefaction and extrapolation sampling curve, (b) sample completeness curve, and (c) coveragebased rarefaction and extrapolation sampling curve.

All non-parametric, asymptotic richness estimates indicate greater OTU richness in male associated fungal communities for profiles generated by both enzymes

(Table 2).

Table 2. Non-parametric diversity estimators of fungal diversity. Sample size (n), number of observed OTUs (S_{obs}), Mao Tau expected OTU richness (S_{est}), and estimated OTU richness using Chao2 estimator (Chao2) and incidence-based coverage estimator (ICE), ± standard deviation for fungal community profile generated by Hae III and Hinf I.

Hae III		n	S_{obs}	S _{est}	Chao2	ICE
	Females	23	54	56.5±2.18	56.49 ± 0.0	56.49±2.5
	Males	23	59	66± 1.37	67.78± 1.6	83.18±0.0
Hinf I						
	Females	24	67	65± 2.55	65.01±0.0	65.01±2.0
	Males	23	63	78± 1.77	81.07±2.18	104± 0.0

Fungal Biomass Estimates

Overall ergosterol concentrations were statistically significantly greater (p=0.02; df=1; SS=1.0) in male tissues than in female tissues of *C. purpureus* (Fig. 7a, Table 2), a pattern that was consistent across all three populations sampled (Fig. 7b).

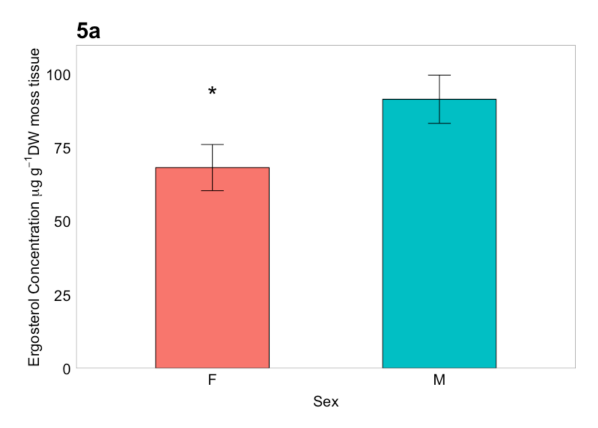
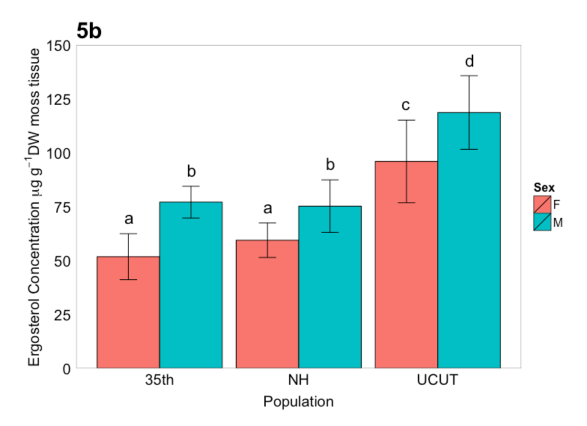
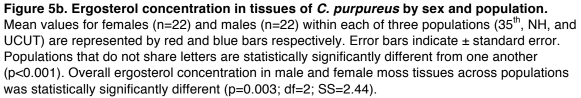


Figure 5a. Ergosterol concentration in tissues of *C. purpureus* by sex. Mean values for all females (n=22) and all males (n=22) are represented by red and blue bars respectively. Error bars indicate \pm standard error. Ergosterol concentration in male and female moss tissues was statistically significantly different (p=0.02; df=1; SS=1.0).





Populations also differed in overall ergosterol concentrations (p=0.003;

df=2; SS=2.44), with individuals from the UCUT population having statistically

significantly higher levels than individuals in either the 35th or NH populations

(Fig. 7b, Table 4). The interaction between plant sex and population was not

statistically significant (p=0.6; df=2; SS=0.18).

Table 3. Results of two-way ANOVA for ergosterol concentration in moss tissues as influenced by host sex (male and female) and population (NH, 35th, and UCUT). Values for degrees of freedom (df), sum of squares (SS), mean squares (Mean Sq), F-ratio (F), and p-value (p) are displayed. Bold values indicate statistical significance.

	df	SS	Mean Sq	F	р
Population	2	2.44	1.22	6.80	0.003**
Sex	1	1.00	1.00	5.59	0.023*
Sex x Population	2	0.18	0.09	0.50	0.613
Residuals	38	6.82	0.18		

Significance tests are shown in bold, and *0.01 $\leq p < 0.05$; **0.001 $\leq p < 0.01$.

Table 4. Results of post-hoc Tukey HSD test showing pairwise differences in mean ergosterol concentration between populations (NH, 35th, and UCUT). Differences between means (Difference), lower (Lower) and upper (Upper) bounds of confidence, and p-value after adjustment for multiple comparisons (p adj) are displayed. Bold values indicate significant differences between means.

	Difference	Lower	Upper	p adj
NH-35 th	0.10	-0.28	0.48	0.792
UCUT-35 th	0.54	0.15	0.92	0.003**
UCUT-NH	0.43	0.06	0.81	0.019*
<u> </u>		1+0.01	A A - ++A A A +	

Significance tests are shown in bold, and *0.01 $\leq p < 0.05$; **0.001 $\leq p < 0.01$.

Chapter 4: Discussion

The goal of the present study was to investigate how moss sex influences associated fungal community composition and biomass under common garden conditions. I used a rapid and robust molecular fingerprinting method (T-RFLP) that boasts high accuracy in assessing microbial community structure (Pilloni *et al.*, 2012), but is limited by its inability to resolve species identity. My results show evidence of host sex-specific structuring of fungal community composition associated with the moss *C. purpureus* (Figs. 3a, 3b). Differences in fungal composition across the three *C. purpureus* populations examined were also evident (Figs. 3b, 4b), though sex-specific patterns were maintained and female associated fungal communities generally appear to be more similar to one another as evidenced by tighter clustering relative to male associated using all three non-parametric, asymptotic species richness estimators (Mao Tau, Chao2, ICE) indicate greater overall species richness in male associated fungal communities.

Furthermore, I observed significantly greater amounts of fungal biomass associated with the tissues of male individuals than females, suggesting that host sex is an important factor influencing the extent of fungal abundance. The quantities of ergosterol measured in greenhouse populations of *C*. purpureus were comparable (mean female = 69.3, mean male = 91.5, mean total = 79.9 μ g ergosterol g⁻¹ dry weight plant material) to those previously measured in forest

mosses (48.5 μ g ergosterol g⁻¹ dry weight plant material; Davey *et al.*, 2009), which indicates that fungal loads can be just as high (and higher in this study) in mosses that prefer disturbed habitats such as *C. purpureus*, and likewise under greenhouse conditions. I realize that although the synthetic nature of common garden approaches may not accurately reflect the ecological dynamics in natural populations, they are an indispensible tool for examining genetically rooted features of populations under controlled environmental conditions.

Mosses are considered foundation species owing to their disproportionately large influence on ecosystem processes and community structure in the biomes they inhabit (Lindo & Gonzalez, 2010; Turetsky et al., 2012). A number of the critical ecological roles of mosses are manifestations of their intimate relationships with other organisms, for example, the symbiotic association of dinitrogen-fixing cyanobacteria with feather mosses (Turetsky, 2003; Lindo & Gonzalez, 2010). The sheer magnitude of the fungal presence generally observed within the bryosphere is indicative, at the very least, of a major source of fungal propagules (Davey & Currah, 2006), but undoubtedly extends functional importance at the ecosystem level (Christensen, 1989), similar to other moss-associated biota. The separation of sexes is an ancient reproductive strategy that has emerged many times in the bryophytes across evolutionary time (McDaniel et al., 2011). Sexual dimorphisms are widespread in dioicous bryophytes (Fuselier & Stark, 2004) and are the basis of differential fitness optima for each sex. Females are almost always larger in size (Shaw &

Beer, 1999; Glime & Bisang, 2014a) which is likely a consequence of the resources required by the maternal gametophyte to provide sole sustenance to the maturing sporophyte. Female sex structures (archegonia) typically initiate later and develop faster than male structures (antheridia), which also appear to have a narrower range of environmental conditions in which they are expressed (Stark, 2002; Glime & Bisang, 2014b). The unique life history and physiological traits of the sexes create distinct physical niches that are expected to be suitable for different fungal taxa accordingly. Davey *et al.* (2012a) observed significant seasonal differences in fungal community composition associated with three common boreal mosses, all of which are dioicous, though host moss sex was not considered in their study.

Recent work measuring leaf, cell, and physiological traits in *C. purpureus* from the same populations used in this study revealed that there were significant sex-specific differences in characteristics such as leaf thickness, canopy mass area, and photosynthetic capacity, among others. (Slate *et al.* unpublished). Interestingly females had larger, thicker leaves with thicker cell walls than males that may provide greater protection from invading fungi who often inhabit specific microniches within the cells of the host moss (Döbbeler, 2002; Ptaszyńska *et al.*, 2009); perhaps explaining to some extent the reduced fungal richness and biomass associated with female individuals of *C. purpureus* observed in the current study. Slate *et al.* also found that females had significantly greater dark-adapted F_v/F_m values which is indicative of the maximum potential quantum

efficiency of Photosystem II (i.e. photosynthetic performance) (Maxwell & Johnson, 2000), which coupled with other leaf traits that likely enhance CO₂ assimilation in females compared to males, may allow for greater net assimilation by females. If this is the case, then females may have a greater resource pool to invest in other traits such as chemical defenses. Better-defended females are common among dioecious angiosperms (Jing & Coley, 1990; Fritz & Simms, 1992; Ågren *et al.*, 1999; Cornelissen & Stiling, 2005; Cepeda-Cornejo & Dirzo, 2010; Barrett & Hough, 2013) and may also explain the more refined fungal communities and lower fungal biomass observed in female individuals of *C. purpureus*.

Another factor that likely contributes to sex-specific patterns in fungal affiliations is resource availability associated with sexual expression. Stark (2002) highlights the different lines of evidence that suggest that male antheridia are more energetically expensive to produce than female archegonia. This is further supported by the findings of McDaniel (2005) who showed that there is a trade-off between allocation to reproductive and vegetative tissue in *C. purpureus* males, but not in females. A greater physiological investment to male sexual structures, such as lipid and starch rich sperm (Paolillo, 1979; Renzaglia & Garbary, 2001) and mucilage exuded by paraphyses (Harvey-Gibson & Miller-Brown, 1927) and may provide a richer storehouse of resources for fungi to exploit, which may in turn explain the significantly greater fungal biomass and richness associated with male tissues in this study. These components have

been observed to serve as attractants and food for arthropods (Harvey-Gibson & Miller-Brown, 1927) supporting the possibility of them being a food source for microbiota as well. Likewise, the chemical properties of female sexual structures may also influence the distribution of fungi in the moss phyllosphere. Early work by Pfeffer (1884) and later Kaiser et al. (1985) demonstrated that moss archegonia produce sugars (namely sucrose) that were shown to attract chemotactic male sperm (Cronberg, 2012). Subsequent investigations revealed that a suite of sugars accumulate in specialized cells of the archegonia during the development of Bryum capillare, and are released shortly before fertilization creating a gradient that sperm can follow (Kaiser *et al.*, 1985). Such accretion of sugars may be an important factor effecting the composition of microbial inhabitants as observed in the floral nectar of vascular plants (Alvarez-Pérez & Herrera, 2013). However, owing to the actually low relative concentration of sucrose (0.086%; Ziegler et al., 1988) in archegonia exudates compared to nectar found in angiosperm flowers (7-70%; Nicolson et al., 2007; Cronberg, 2012), perhaps a more qualitative (i.e. specific community composition) rather than quantitative (i.e. species abundance) effect on microbial communities might be excepted. The overall greater richness together with the greater dissimilarity of community composition exhibited by male associated fungal communities relative to female communities: may be indicative of a more specific assembly or narrower realized niche for fungi associated with females.

The potential for multi-trophic community feedbacks associated with genetic variation in foundation species has been previously demonstrated in various angiosperm systems (see review by Whitham *et al.*, 2006), and was recently shown to be driven by plant sex (Petry *et al.*, 2013). In their study, Petry and collaborators found that female plants of the species Valeriana edulis attracted a disproportionate abundance of aphid, aphid predators, and aphidtending ants, showing 4-fold, 1.5-fold, and 4-fold higher densities respectively compared to their male counterparts. Interestingly, differences in floral nectar between males and females were found in large part to explain the sexual dimorphisms in arthropod communities observed. This finding suggests that a resource-based partitioning of community structure across sexes is indeed a strong possibility as I have speculated for *C. purpureus*, though instead with a male bias. Other studies have also observed sex-specific biotic interactions with between plants and fungi. In a study by Eppley *et al.* (2009), higher levels of colonization (1.62-2.58 times greater) by arbuscular mycorhizal fungi were observed in the roots of females versus males of the dioecious grass *Distichlis* spicata under field conditions. The authors suggest that these observed differences may cause antagonisms between males and females through intersexual competition for resources. Though the functional capacities of the fungal communities examined in my study remain to be resolved, the likelihood of some level of sexual antagonism or possibly synergism (see below) based on fungal interactions remains. Given that such complex communities are also

common in the bryosphere, I predict that similar differential structuring is likely occurring as a result of moss sexual identity. In a recent study, Rosenstiel *et al.* (2013) found that *Collembolan* microarthropods commonly occurring in the *C. purpureus* phyllosphere were differentially attracted to complex VOC profiles emitted from wild-collected female gametophytic tissue. As the VOCs originating from moss tissues were not distinguished from those emitted from phyllosphere microbes, it is possible that specific fungal volatiles may be functioning as attractants for the largely fungivorous microarthropods.

The milieu of sexually dimorphic traits exhibited by dioicous bryophytes provides a robust foundation for divergent structuring of resident fungal communities, but further work is necessary to link the complex physiological and phenological events that define them and their population and community level effects. This work provides a first glance at how genetically based sexual systems in early land plants influence affiliated fungal composition.

Future Directions

While this research takes initial steps towards understanding sex-specific structuring of community phenotypes in a moss based system, there is a fruitful path ahead for further work in this area. Utilizing next-generation sequencing techniques will allow for the identification of microbial elements, and coupled with phylogenetic methods, can greatly strengthen our knowledge concerning the evolutionary trajectories that build community interactions. In the same vein,

including other moss species featuring dioicous and other sexual systems in such studies will reveal the extent to which sex-specific community structuring exists in this group. Furthermore, identifying and incorporating quantitative trait loci that govern moss sex-specific characteristics influential across community levels (e.g. concentration of phenolics in tissues, generation of nutritive exudates) will open the door to testing for the relative strength of continuous traits at different levels of expression, as well as their heritability across generations. Such techniques have been applied to the study of community genetics in the *Populus* model system (Shuster *et al.*, 2006; Whitham *et al.*, 2006) and no doubt are applicable to moss systems with recently available genomic resources for *Physcomitrella patens* and *C. purpureus*.

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