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Spatial habitat structure assembles willow-dependent communities across the primary successional watersheds of Mount St. Helens

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Abstract: Major disturbances fundamentally alter ecosystems through the transformation of both biotic and abiotic factors. The eruption of Mount St. Helens in 1980 resulted in a cataclysmic restructuring of its surrounding landscapes. The Pumice Plain is one of these landscapes, where tree species such as Sitka willow (*Salix sitchensis*) and their dependent communities have established along newly-formed streams. Thus, the study of these dependent communities provides a unique opportunity to investigate factors influencing metacommunity assembly during true primary succession. We analyzed the influence of landscape connectivity on metacommunity assembly through a novel application of circuit theory, alongside the effects of other factors such as stream locations, willow leaf chemistry, and leaf area. We found that landscape connectivity structures community composition on willows across the Pumice Plain, while our other factors had varied effects. Most importantly, multiple levels of spatial habitat structure linked via landscape connectivity can predict the presence of organisms lacking high rates of dispersal, such as the invasive stem-boring poplar weevil (*Cryptorhynchus lapathi*). This is critical for management as we show that the maintenance of a heterogeneous mixture of landscape connectivities and resource locations can facilitate metacommunity dynamics to promote ecosystem function and mitigate the influences of invasive species.

Keywords: dioecious; landscape connectivity; plant-insect interactions; metacommunity assembly; Mount St. Helens; willow; invasive species

1. Introduction

Major disturbances such as hurricanes, wildfires, and human activity can fundamentally alter ecosystems through the removal of organisms from the landscape [1-4]. Some disturbances can have even more pronounced effects on organisms through the modification of abiotic factors such as topography and soil chemistry (e.g., landslides, volcanic eruptions)[5-6]. Ecological consequences of disturbances include the severance of landscape connectivity [7-8], population bottlenecks that reduce genetic diversity [9], and the disruption of nestedness within ecological communities (i.e., metacommunities)[10-12]. These consequences can have both positive and negative effects on post-disturbance communities depending on their prior compositions, such as degree of invasibility versus more “natural” compositions [13-14], resulting in a context-specific understanding of how major disturbance processes shape the world’s landscapes [15-17]. Biological assembly of a landscape after major disturbance events is often referred to as primary succession, wherein the re-establishment of organismal populations and communities over time is governed by environmental conditions and resource availability [8, 18-19]. The regeneration of metacommunity dynamics is an important component of primary succession that

can often take many years [20], yet is critical to understanding current and future trends for the function of ecosystems post-disturbance [10, 21-22].

Landscape connectivity, measured as how easily organisms can disperse across a landscape, structures the geospatial arrangement of plant species and can aid in the regeneration of metacommunities following disturbance [19, 23-24]. The resulting distributions of host plants, a nested degree of biological landscape connectivity, often have direct influences on dependent communities of arthropods and microbes [25-27]. Landscape connectivity as a result of host plant location can thus assemble plant-associated communities based on dispersal ability and proximity of habitat corridors [28-30]. In many landscapes, topography, established vegetation patches, and the presence of water interact to influence where plant communities can establish [31-33]. Specifically, water availability limits where plants establish, especially for many tree species in the western US [34], as decreased water availability results in decreased recruitment of trees and greater early-age mortality [35-36]. Furthermore, tree community regeneration can be determined based on accessibility of water following major disturbances given that light, through the removal of competitors, and nutrient accessibility, through the deposition of organic material, are not typically limiting [37-40]. Thus, the ability of primary successional tree communities to influence connectivity is reliant on where water is found on the landscape, how much water is available, and how these characteristics change over time [41-44]. The connectivity of these tree communities, and by proxy the connectivity of water availability, should then be directly related to associated arthropod and microbial communities that establish on trees post-disturbance [45-46].

Many attempts have been made to model landscape connectivity [47-49] across different regions of the world and for markedly different ecosystem types [50-52]. The most powerful methods currently used, such as circuit theory [53], rely on the stochastic dispersal of a target population or community over space and time directed by some set of structuring parameters [54]. While circuit theory can require substantial computational resources [55], it has been consistently found to outperform other techniques for modeling landscape connectivity for both plants and animals [53, 56-57]. However, circuit theory has almost exclusively been used to model gene flow across populations or metapopulations of one species rather than modeling the establishment of community members in primary successional landscapes [58-59]. Furthermore, applications of circuit theory typically focus on instantaneous connectivity between known individuals on a landscape and how movements are facilitated among them [53, 58-60], rather than on possible pathways for a new individual to find habitat in a novel landscape [61]. Despite limited use by community ecologists, circuit theory may be useful for understanding landscape patterns of species establishment, especially in primary successional landscapes, as it relies on simple yet robust assumptions of what geographical, chemical, or spatial parameters resist or facilitate the movement of certain species [48, 54, 60-61].

Primary successional riparian habitats in the western US tend to follow a sequence whereby willow species (*Salix* spp.) establish first, reaching maximum productivity at approximately 10 years, followed by replacement by alder (*Alnus* spp.) until around 40 years [62]. The establishment of these pioneer trees creates corridors on the landscape and may allow for increased habitat connectivity, especially across otherwise harsh environments. As riparian plants establish, they create a landscape mosaic of patches that may vary based on within-species genetic variation [63-65] and other plant traits, such as plant sex in dioecious species like willow [66]. Genetic variation, in addition to species-level differences, may create a landscape that varies in terms of plant biomass, leaf chemistry, leaf shape, leaf size, and individual susceptibility to herbivory [67-69]. Additionally, interactions between these riparian host plants and leaf-modifiers, herbivores, and endophytes can further alter the chemical and structural landscape of these developing riparian forest ecosystems [70-73]. As a result, other species which depend upon certain plants for their habitat may disproportionately utilize different plant species or even individuals of the same plant species which exhibit varying chemical characteristics, fundamentally altering communities and metacommunity compositions across the landscape [74-76].

Extreme volcanic eruptions, alongside the creation of new lava fields and massive landslide events, can result in the complete removal of all biotic material from a landscape, and thus are an example of true primary succession [77-80]. During the eruption of Mount St. Helens (Lawetlat'la in the Cowlitz language; Washington, USA) in 1980, all three of these disturbance types occurred simultaneously and their interaction transformed vast areas by incinerating pre-existing forests and depositing >100 m of mostly inert material onto the landscape. These disturbances generated a unique primary successional landscape feature called the Pumice Plain, a relatively flat area, where springs and seeps created several streams along which Sitka willow (*Salix sitchensis*) and green alder (*Alnus viridis* (Chaix) DC) have begun to establish. Previous research has demonstrated an overall 2:1 female:male ratio of willows across the Pumice Plain [81], and that female willows establish closer to stream edges [82]. Additionally, it has been demonstrated that female willows are more susceptible to attack by the stem-boring poplar weevil, *Cryptorhynchus lapathi* (Curculionidae, Coleoptera) [83]. Attack by the weevil causes premature branch death and alters leaf litter chemistry [84-86], possibly influencing other plant-associated species. In addition, plant sex in *S. sitchensis* has been shown to alter leaf chemistry [82, 86] and potentially plant size and leaf size, which may both influence plant-associated organisms [87-88].

Few studies have used a connectivity perspective to understand patterns of true primary succession, or the assembly of plant-associated communities across landscapes. Here we propose two specific hypotheses: 1) Species richness of willow-dependent communities across the Pumice Plain will be influenced broadly by landscape connectivity, followed by nuance through stream identity, changes in willow chemistry, willow sex, and leaf size. Specifically, we predict that greater connectivity, leaf size, nitrogen content, and available tannins will lead to greater species richness. Furthermore, we predict that willows of different sexes and sampled from different streams across the Pumice Plain will host different levels of species richness. 2) Organisms with different life histories will depend differentially upon landscape connectivity, stream identity, willow chemistry, sex, and leaf size across the Pumice Plain. We predict that organisms with limited ability to disperse, such as weevils, galling mites, and fungal rust, will require more connected landscapes. We predict that organisms with more symbiotic relationships will be more influenced by female willows given the noted higher rate of attack above (e.g., stem-boring weevils vs. generalist chewers). Finally, we predict that intrinsic plant characteristics such as leaf chemistry and leaf size will differentially influence the presence and absence of willow-dependent organisms.

2. Materials and Methods

2.1. Site Description

Mount St. Helens erupted in 1980 in a massive lateral blast that transformed over 600 km² of forests, lakes, and streams. A 15 km² area in the main blast zone, called the Pumice Plain, was first buried by over 100 m of sterile pumice, ash, and sand in a 2.8 km³ debris avalanche [89]. It was then hit with a hot lateral blast of flying rock debris, covered in 0.3 km³ of lava which was as much as 40 m thick in places, and subjected to ash and tephra fall [90]. In the years following the 1980 eruption, springs, seeps, and run-off from snow-melt across the Pumice Plain created five new watersheds (Figure 1) which are slowly seeing establishment of riparian vegetation. Dominant riparian species include green alder (*Alnus viridis*) and Sitka willow (*Salix sitchensis*; hereafter simply willow), the latter of which has been regularly attacked by a stem-boring weevil (*Cryptorhynchus lapathi*, Curculionidae, Coleoptera) since its introduction in 1989 [83].

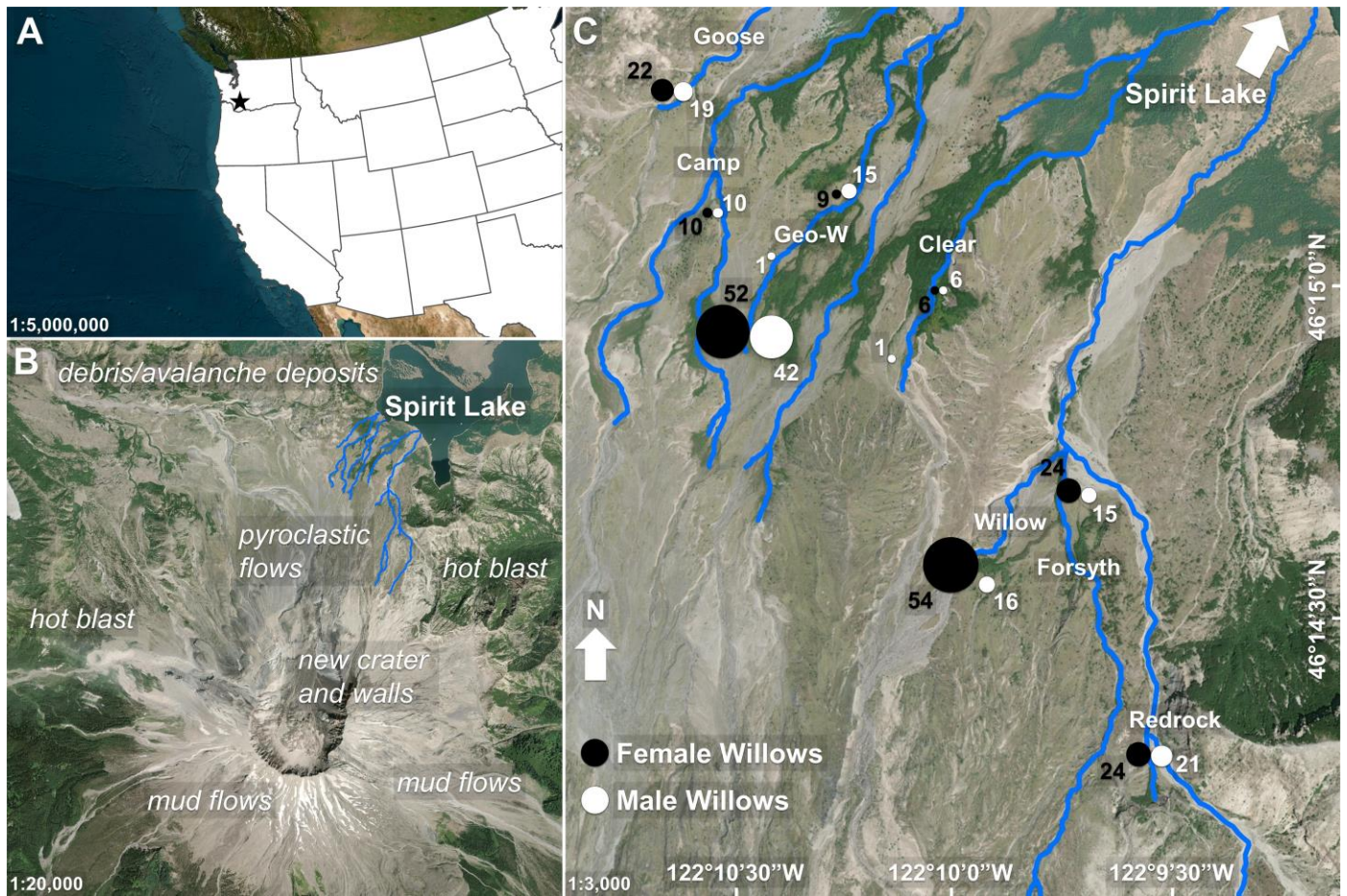


Figure 1. Site and study descriptions for the Pumice Plain of Mount St. Helens. A) Location of Mount St. Helens in Washington state, USA (represented by the black star). B) Dominant geomorphological processes structuring landscape processes post-eruption [91]. Blue lines are stream features on the Pumice Plain, from which willow trees were sampled. C) Sampling locations of willow trees across the Pumice Plain. Sampling of female v. male willows was uneven due to uneven field access and sampling of the most available trees. Nearest streams to sampling locations are labeled (Goose, Camp, Geo-W, Clear, Willow, Forsyth, and Redrock). For analyses, Forsyth and Redrock Creeks were separate from Willow Creek, even though they are tributaries. Projection: WGS 84 / UTM Zone 10N. Figure created in ArcGIS Pro (version 3.0.0)[92].

2.2. Field Data Collection

Between May-June of 2018, we haphazardly tagged 348 individual willows across the Pumice Plain and identified each as either male or female based on the presence of reproductive structures [93]. Individual trees were tagged with unique identification numbers and colored beads to represent sex, and their locations were recorded by GPS. We also recorded the identity of the nearest stream (Camp, Clear, Forsyth, Geo-West (Geo-W), Goose, Redrock, and Willow Creeks). In the summer of 2019 (26 June to 11 July), we surveyed all willows once for a suite of willow-dependent organisms. These species are varied in their life histories and plant-usage strategies, yet all leave identifiable markings on their willow hosts. Thus, the resulting data, despite a potential temporal mismatch of sampling and organism presence, are not skewed as these markings were maintained by willows to the time of sampling. We surveyed for the presence of attack by the stem-boring poplar weevil, *Cryptorhynchus lapathi*, based on evidence of frass at the base of stems, dead branches, and/or the presence of adult weevils. We surveyed for the presence of western tent caterpillars, *Malacosoma fragilis* (Lasiocampidae, Lepidoptera), through evidence of webs, caterpillars, and/or cocoons. We surveyed for evidence of the Pacific willow leaf beetle, *Pyrrhalta decora carbo* (Chrysomelidae, Coleoptera), based on the presence of skeletonized leaves. We noted other types of leaf chewing (chewed edges) caused by generalist herbivores, as well as the presence of ants, aphids, and other unidentified caterpillars. We surveyed for the presence of the willow leaf gall sawfly, *Pontania pacifica* (Tenthredinidae,

Hymenoptera), based on the presence of round, red galls. We also noted the presence of a second type of gall, most likely mites of the Eriophyidae family, that presented a range of colors (red, orange, and yellow) on the top surface of the leaf with white spindle growths on the underside. We noted the presence of *Melampsora* spp. rust (an orange fungus common on members of the Salicaceae family), and unidentified endophytes based on the presence of discrete black stromata. Species richness for each tagged tree was determined based on their unique dependent community.

2.3. Leaf Litter Chemistry

Leaf litter was collected from a random subset of the tagged willows ($n = 36$) from across the Pumice Plain to support related leaf litter decomposition studies. Weevil-induced male and female willow litter was collected in July 2019, as damage due to weevils was more obvious during the growing season, and naturally abscised male and female willow litter was collected in August 2019 [86]. All litter was stored in individual paper sacks and air-dried in the laboratory. Subsamples (0.50 g) were freeze-dried (Millrock Technology, Kingston, NY), ground to a homogeneous consistency (KRUPS Type F203), and used to measure a variety of litter chemical traits (condensed tannins, carbon [C], nitrogen [N], and C:N). First, subsamples of ground material (25 mg) were extracted for soluble condensed tannins with 70% acetone and 10 mM ascorbic acid. We used the butanol-HCl method to determine soluble condensed tannin concentrations [94], with standards purified from local *S. sitchensis* [95]. Condensed tannin concentrations were determined by measuring absorbance at 550 nm on a spectrophotometer (Spectramax 384, Molecular Devices, San Jose, CA) and comparing samples to a standard curve, before being converted to % condensed tannins (%CT). Second, ground subsamples (2 mg) were weighed into tin capsules (5×8 mm) to determine %C and %N on an elemental analyzer (2400 CHNS/O Series II System, Perkin Elmer, Waltham, MA). The molar element ratio of C:N was calculated as the ratio of %C divided by the atomic mass of C over %N divided by the atomic mass of N.

2.4. Leaf Area Measurements

Leaf area can influence associated insect and microbial community members [96-97], thus fresh leaves were collected from a subset of the tagged willows ($n = 122$). Roughly five to ten leaves (depending on size) were collected from each willow and either preserved in silica gel or laid onto individual paper sheets in a plant press. Both sample types were dried for several months, weighed to the nearest mg, then scanned on a flatbed scanner. Images were converted to black and white, and leaf area was calculated in ImageJ [98] using a batch process. Specific leaf area (SLA) was calculated by dividing the total leaf area by the mass of the leaves, then by the number of leaves in each sample.

2.5. Landscape Connectivity

Landscape connectivity often determines plant arrangements on a landscape as well as dependent communities [23, 61]. Thus, we used the Circuitscape connectivity package in Julia [56] with a high-performance computing cluster to generate continuous landscape connectivity measures for the Pumice Plain based on resistance maps produced in ArcGIS Pro (version 3.0.0)[92]. Primary stream channels were mapped in-field using GPS. Updated and ground-truthed geospatial data was required due to the dynamic nature of our study streams. In this analysis, the greater distance from stream features resulted in greater resistance on the landscape, given that water availability is often the most important limiting factor in post-disturbance community regeneration in the western US [37-40]. We tested a combination of stream locations and physical parameters such as slope and elevation to predict landscape resistance [99], but landscape resistance changed little compared to only using stream locations. Thus, we used resistance maps created by linearly extrapolating resistance from mapped stream locations to maintain simplicity in understanding landscape connectivity for the Pumice Plain. Circuitscape runs were based

on randomized node locations that produced pairwise current walks through our resistance maps rather than simply producing instantaneous connectivity for known individuals ($n = 20$ to reduce compute time and capture within-run variability). We then recreated the Circuitscape output with 20 randomized node repetitions to ensure that sufficient variation based on different arrangements of focal nodes was captured. From each model output, we sampled current for all of our tagged trees, thus producing a mean and variance of connectivity at each location.

2.6. Statistical Analysis

All statistical analyses were performed in R version 4.2.0 [100]. We specified three suites of predictor parameters for use in different analytical methods based on varying sample sizes. Willow sex was included in all three suites given its influence on ecological communities at each scale and potential interactions with all tested parameters. Prior to analysis, some entries were removed on which we observed no dependent organisms. The first suite contained the following landscape-level parameters: stream identity, landscape connectivity, and willow sex ($n = 335$). Stream identity served as both a measure of unique hydrological inputs as well as a proxy for relative geographical location given that willow trees typically clustered along streams. The second suite contained leaf litter chemistry variables: %C, %N, C:N molar ratio, %CT, willow sex, and weevil-induction ($n = 31$). The third and final suite contained SLA and willow sex ($n = 109$). We then used each of these parameter suites in generalized linear models (GLMs) versus willow community species richness ('glms' function in the 'nlme' package)[101] and in permutational MANOVAs (PERMANOVAs) versus willow community dissimilarity (999 permutations, 'adonis2' function in the 'vegan' package)[102]. Community dissimilarities for our PERMANOVAs were determined using the Bray-Curtis dissimilarity index for the leaf-dependent communities found on each tagged tree [103]. To determine the GLMs and PERMANOVAs with the greatest explanatory power, we performed Akaike's Information Criterion (AIC) model selection for all parameter suites [104]. In this method, model parameters were reduced from the full suite of parameters and their most likely interactions to the simplest two-term models to maximize simplicity and ensure effects were adequately captured. The AICs of each combination of parameters for each parameter suite were then ranked and the best overall GLMs and PERMANOVAs were selected.

We analyzed pairwise post-hoc stream comparisons for our species richness GLMs using estimated marginal means ('emmeans' function in the 'emmeans' package)[105] and Tukey's HSD letters ('cld' function in the 'lsmeans' package) [106]. We performed NMDS analyses ('metaMDS' function in the 'vegan' package)[102] to understand community dissimilarity in the parameter space for differences observed in our PERMANOVAs (stress <0.1, 999 iterations, and 200 random starts). We then used a pairwise PERMANOVA for stream differences based on landscape community dissimilarities ('pairwise.adonis2' function in the 'pairwiseAdonis' package)[107] and similarity percentage (SIMPER) analyses to understand which community members contributed the most to differences found between streams as well as between male and female willows ('simper' function in the 'vegan' package)[102]. Using the results of our SIMPER analyses, we generated a heatmap to highlight the most important community members that drove observed differences ('heatmap' function in base R).

3. Results

Overall taxa richness for willow-associated organisms (willow communities) ranged from 1 to 7, with a slightly higher average of 3.27 ± 0.08 taxa on female willows and 3.08 ± 0.09 taxa on male willows (Table S1). Additionally, taxa richness was highest on willows growing along Forsyth Creek, followed by Camp, Clear, Willow, and Goose, respectively, with lowest taxa richness on willows growing along Redrock and Geo-W. A mosaic of outputs for all randomized Circuitscape runs can be found in Figure 2. Connectivity values ranged from 0.05 to 1.25 across the Pumice Plain, with average values slightly higher

for female willows (0.30 ± 0.02) compared to male willows (0.22 ± 0.02). Connectivity was highest for willows sampled along Willow, followed by Clear, Forsyth, and Geo-W, while the lowest connectivity values were found along Camp, Goose, and Redrock Creeks. Litter %C ranged from 44.80 to 48.80, %N ranged from 0.41 to 2.79, C:N molar ratio ranged from 19.80 to 134.50, %CT ranged from 1.10 to 28.60, and specific leaf area (SLA) ranged from 2.15 to 25.57 mm² mg⁻¹ across the Pumice Plain. Litter %C, %N, %CT, and SLA were similar between willow males and females, while C:N was slightly higher for male willows (71.69 ± 10.27) compared to female willows (61.91 ± 8.89). Litter %C was highly similar across willows from all streams, while %N was highest for Forsyth and Willow and lowest for Geo-W and Goose. C:N molar ratio was highest for willows from Geo-W and Goose, both nearly three times higher than Camp, Forsyth, and Willow Creek. Litter %CT was also highest for willows from Geo-W and Goose, moderately low for Willow and Forsyth, and the lowest for Camp Creek. Finally, SLA was highest for willows from Goose and moderately high for Geo-W, which were over 4 times higher than for Willow Creek.

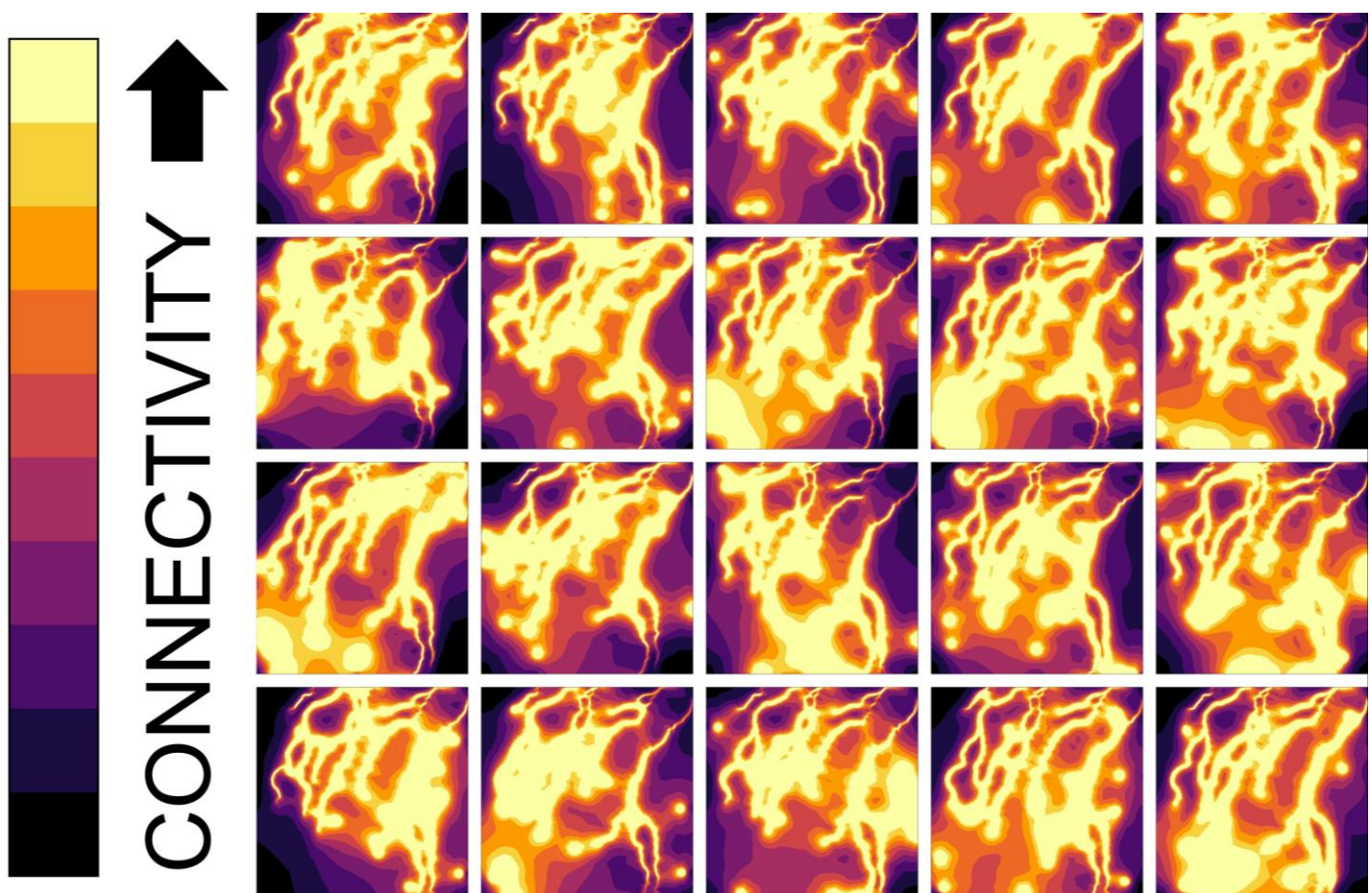


Figure 2. Electrical current outputs from 20 randomized start locations using Circuitscape in Julia [56]. Panes depict relative landscape connectivity based on landscape resistance dictated by distance from nearest water source. Elevation and slope were tested as potential modifiers of resistance, but added little to understanding landscape resistance across the Pumice Plain versus the simpler landscape resistance with distance from water. The size and extent of the geospatial area in this figure are the same as those for Figure 1, Pane C. Projection: WGS 84 / UTM Zone 10N. Figure created in ArcGIS Pro (version 3.0.0)[92].

3.1. Landscape-level Models

We found that landscape connectivity and stream identity with no interaction produced the best landscape-level GLM through model selection ($L = -498.826$, $AIC = 1015.653$; Table S2). With this model (Table 1), connectivity ($\hat{\beta} = -0.718$, $\sigma = 0.263$, $t_{327} = -2.731$, $p = 0.007$) explained differences in willow community species richness where less connected trees were associated with higher species richness. The effects of stream identity were, however, varied given that each stream was only tested in-turn against the first

level of the stream factor (Camp) where only Geo-W ($\hat{\beta} = -0.808$, $\sigma = 0.264$, $t_{327} = -3.059$, $p = 0.002$) and Redrock ($\hat{\beta} = -0.936$, $\sigma = 0.294$, $t_{327} = -3.187$, $p = 0.002$) produced differences in species richness. Thus, from our pairwise post-hoc investigation (Table S3), we found three stream groupings for willows in terms of richness that differed from each other: a) Redrock, Geo-W, Goose, and Clear; b) Goose, Clear, Willow, and Camp; and c) Clear, Willow, Camp, and Forsyth. Overlaps of stream identity for these groupings can be considered as not different from any other stream in either grouping, while the absolute differences between groupings are maintained.

Table 1. Results of three generalized linear models for community richness as a response to changes in landscape, chemistry, and leaf area factors (AIC model selection in Table S2). Factors not present in this table were not part of the best model. Three separate models were used due to differences in number of observations. The stream factor was categorical and thus tested against its respective first level. A pairwise estimated marginal means analysis can be found for the stream factor in Table S3. Bolded rows indicate significant effects at $\alpha = 0.05$. Abbreviations: SLA is specific leaf area, df is degrees of freedom, $\hat{\beta}$ is the effect estimate, σ is the standard error of the effect estimate, t is the test statistic, and p is the null model probability parameter.

Factor	df	$\hat{\beta}$	σ	t	p
Landscape Model (n = 335)					
Connectivity	1	-0.718	0.263	-2.731	0.007
Stream - Clear	1	-0.188	0.394	-0.477	0.633
Stream - Forsyth	1	0.423	0.301	1.406	0.161
Stream - Geo-W	1	-0.808	0.264	-3.059	0.002
Stream - Goose	1	-0.488	0.304	-1.608	0.109
Stream - Redrock	1	-0.936	0.294	-3.187	0.002
Stream - Willow	1	-0.025	0.293	-0.086	0.931
Residuals	327	0.114	1.065	-	-
Litter Chemistry Model (n = 31)					
%N	1	1.492	0.657	2.269	0.031
%CT	1	0.150	0.053	2.842	0.008
Weevils	1	0.985	0.996	0.988	0.332
Residuals	28	0.095	0.867	-	-
Leaf Area Model (n = 109)					
SLA	1	>-0.001	<0.001	-5.417	<0.001
Willow Sex	1	-0.289	0.186	-1.555	0.123
Residuals	107	0.054	0.969	-	-

We found that connectivity, stream identity, and willow sex with no interaction produced the best landscape-level PERMANOVA through model selection ($L = -909.464$, AIC = 1046.270; Table S4). All three parameters predicted differences in community dissimilarity on willow trees in the final model (Table 2; connectivity: SS = 0.361, $F_{1,326} = 5.298$, $p = 0.003$; stream identity: SS = 0.386, $F_{6,326} = 13.716$, $p = 0.003$; and willow sex: SS = 5.600, $F_{1,326} = 5.670$, $p = 0.001$). Our landscape-level NMDS analysis (Figure 3) revealed that weevils,

fungal rust, and galling mites were strongly associated with both female willows and with landscape connectivity. On the other hand, we found that sawflies and endophytes were negatively associated with female willows (i.e., associated with males) and landscape connectivity. Chrysomelid beetles and tent caterpillars were weakly negatively associated with female willows and landscape connectivity, while all other community members (aphids, other caterpillars, and other chewers) were not associated with either factor. Our pairwise stream analysis indicated that willow-dependent community compositions for Redrock and Clear were different from all other streams and each other (Table S5). Furthermore, the compositions for Geo-W were different from all other streams with the exception of Forsyth.

Table 2. Results of three permutational MANOVA (PERMANOVA) tests for community dissimilarity as a response to changes in overall landscape, chemistry, and leaf area factors (AIC model selection in Table S4). Factors not present in this table were not part of the best model. Three separate models were used due to differences in number of observations for each subset of data. Differences found for continuous variables were analyzed using NMDS analyses in Figures 3 and S1. Abbreviations: df is degrees of freedom, SS is sum of squares, R² is the correlation coefficient, F is a pseudo-F test statistic, and *p* is the null model probability parameter. Bolded rows indicate significant effects at $\alpha = 0.05$.

Factor	df	SS	R ²	F	<i>p</i>
Landscape Model (n = 335)					
Connectivity	1	0.361	0.013	5.298	0.003
Willow Sex	1	0.386	0.014	5.670	0.003
Stream	6	5.600	0.196	13.716	0.001
Residual	326	22.183	0.778	-	-
Litter Chemistry Model (n = 31)					
%N	1	0.097	0.056	1.748	0.166
%CT	1	0.080	0.046	1.444	0.237
Residual	28	1.559	0.898	-	-
Leaf Area Model (n = 109)					
SLA	1	0.353	0.041	4.575	0.015
Willow Sex	1	0.064	0.007	0.825	0.501
Residual	107	8.179	0.952	-	-

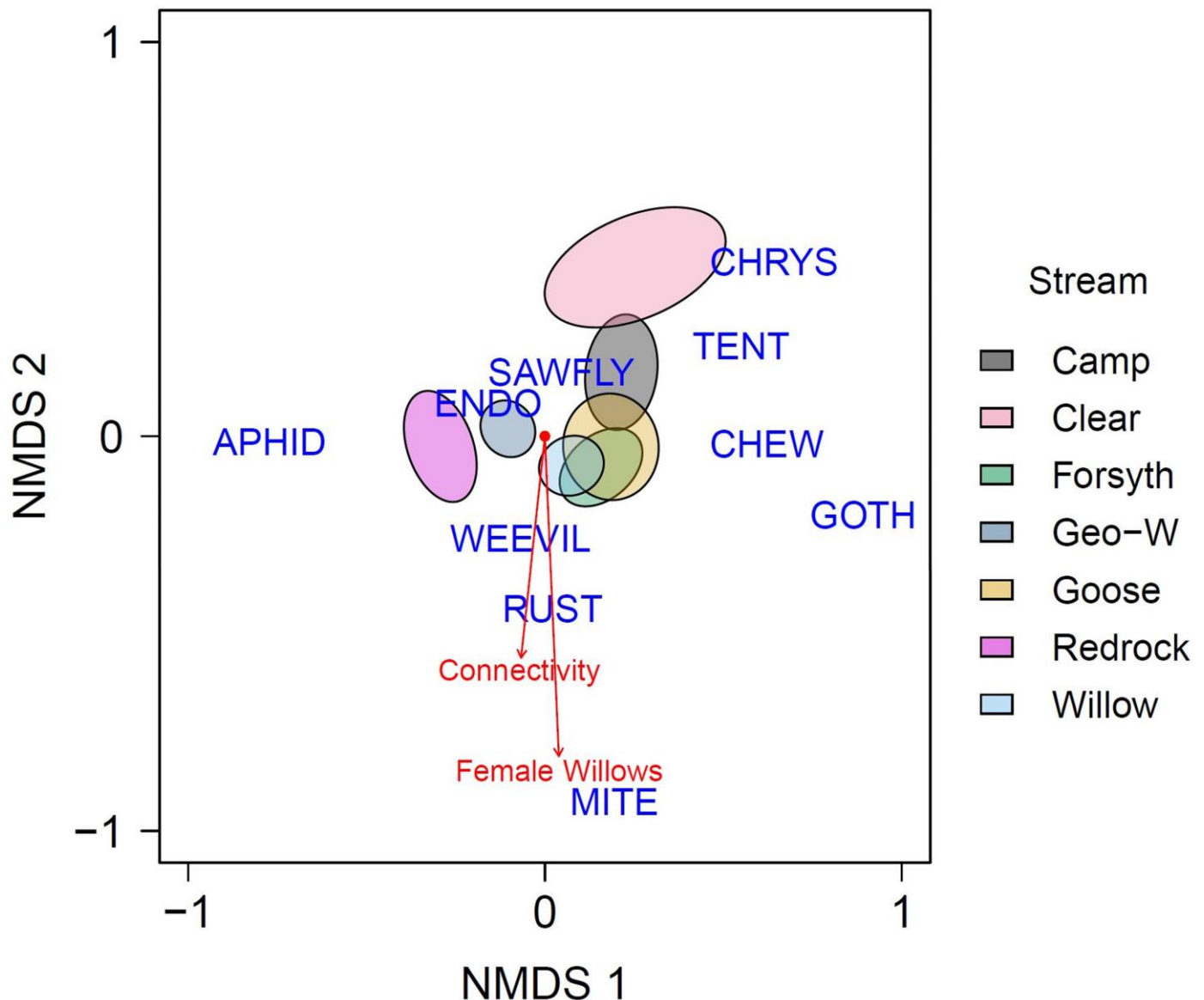


Figure 3. NMDS analysis (four-dimensional) examining dependent-community dissimilarity on tagged willow trees for our landscape-level parameters. Points omitted to focus on community member differences versus parameters of assembly. Blue words are centroids for the 10 observed community members. Stream ellipses are 95% confidence intervals for standard error. Relative length of connectivity and female willow vectors indicate strength of effect of that parameter on related community members. To find the best NMDS outputs, we sequentially increased the number of dimensions until stress was reduced to <0.1 , based on 999 iterations and 200 random starts. Abbreviations: CHEW represents unidentified chewing guild arthropods, CHRYYS represents chrysomelid beetles, ENDO represents endosymbiont organisms on willows, GOTH represents unidentified caterpillars, RUST represents fungi creating leaf rust on willow leaves, and TENT represents tent caterpillars.

Our SIMPER analysis (Figure 4 and Table S6) revealed that all stream-based differences in the willow communities above were driven in some way by tent caterpillars (15/15 comparisons) and nearly all by weevils (14/15 comparisons; only Redrock v. Camp was not). In general, community differences between streams that were closer together as well as farther away tended to be driven mostly by tent caterpillars while those at intermediate distances from one another were driven primarily by weevils, chrysomelid beetles, and other chewers. Additionally, sawflies and chewers were important drivers for the majority of observed differences (12/15 and 11/15 respectively), but were not as important as tent caterpillars or weevils overall. Chrysomelid beetles were very important drivers for all differences found between Clear and other streams despite driving differences in fewer than half of the stream comparisons (6/15). The presence of endophytes, rust, aphids, and other caterpillars did not drive any community differences and the

presence of galling mites only drove differences in two comparisons (Redrock v. Forsyth and Willow v. Geo-W). Finally, we found that community differences between female and male willows were driven primarily by tent caterpillars and weevils, but also in some manner by sawflies, mites, and other chewers.

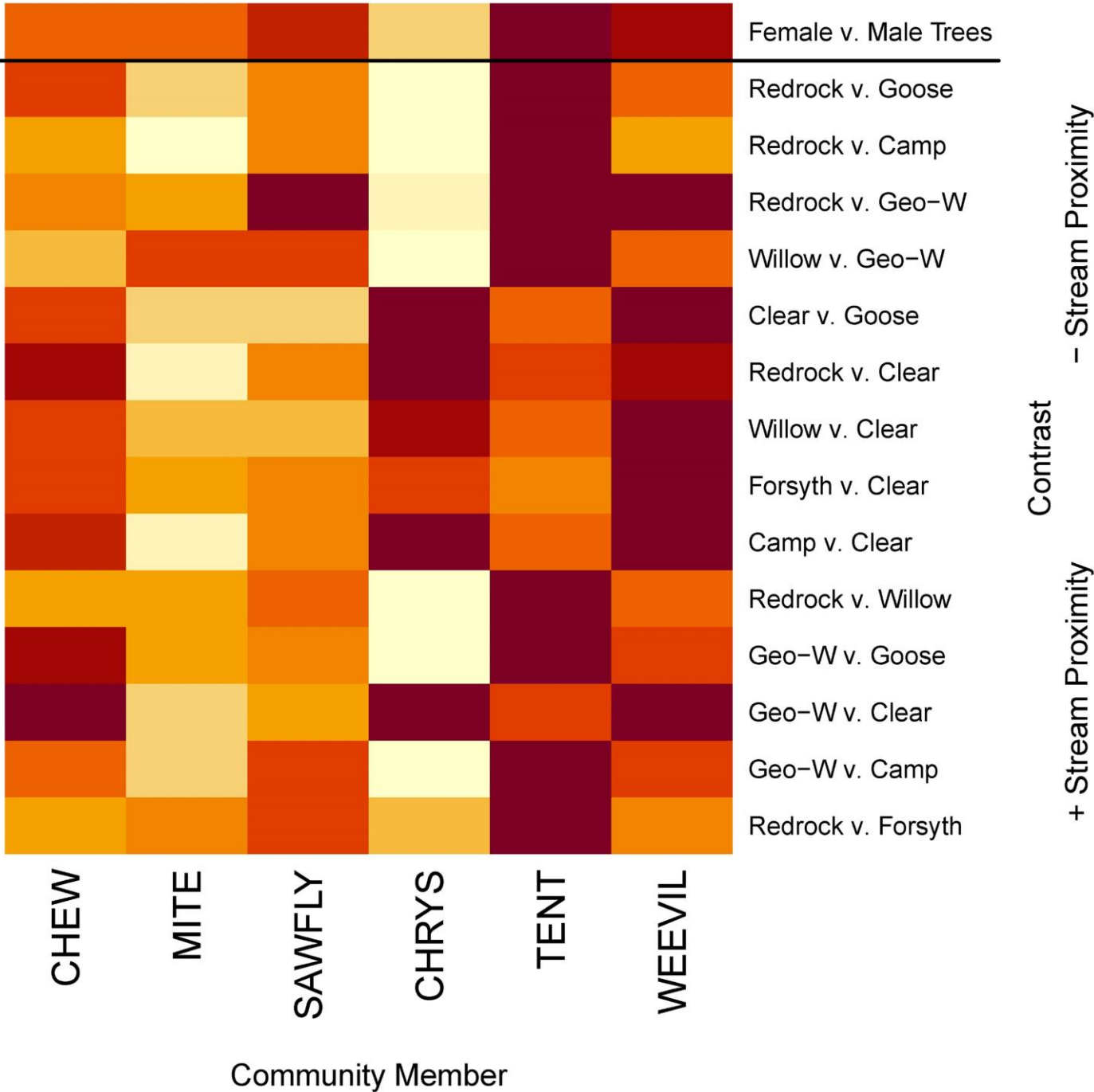


Figure 4. Heatmap highlighting the most important willow-dependent community members that drove observed differences for willow sex and stream comparisons (contrast above) from our SIMPER analyses (Table S6). Darker red colors represent more important community members for a given comparison. “+ Stream Proximity” indicates stream comparisons that are geographically closer together, while “- Stream Proximity” indicates those farther apart. Abbreviations: CHEW represents unidentified chewing guild arthropods, CHRYS represents chrysomelid beetles, and TENT represents tent caterpillars.

3.2. Leaf Litter Chemistry Models

We found that %N, %CT, and weevil presence with no interaction produced the best leaf litter chemistry GLM through model selection ($L = -40.386$, $AIC = 90.771$; Table S2). In

this model (Table 1), both %N ($\beta = 1.492$, $\sigma = 0.657$, $t_{28} = 2.269$, $p = 0.031$) and %CT ($\beta = 0.150$, $\sigma = 0.053$, $t_{28} = 2.842$, $p = 0.008$) explained differences in willow-dependent community species richness, where species richness was positively associated with %N and %CT. Weevil presence did not drive differences in species richness ($\beta = 0.985$, $\sigma = 0.996$, $t_{28} = 0.988$, $p = 0.332$). We also found that %N and %CT with no interaction produced the best leaf litter chemistry PERMANOVA through model selection ($L = -92.689$, AIC = 19.765; Table S4). However, neither %N (SS = 0.097, $F_{1,28} = 1.748$, $p = 0.166$) nor %CT (SS = 0.080, $F_{1,28} = 1.444$, $p = 0.237$) explained differences in willow community dissimilarity in the final model (Table 2).

3.3. Leaf Area Models

We found that SLA and willow sex with no interaction produced the best leaf area GLM through model selection ($L = -162.167$, AIC = 332.333; Table S2). In this model (Table 1), only SLA ($\beta = >-0.001$, $\sigma = <0.001$, $t_{107} = -5.417$, $p = <0.001$) explained differences in willow community species richness, where species richness was negatively associated with SLA. Willow sex did not drive differences in species richness in this context ($\beta = -0.289$, $\sigma = 0.186$, $t_{107} = -1.555$, $p = 0.123$). We also found that SLA and willow sex with no interaction produced the best leaf area PERMANOVA through model selection ($L = -282.286$, AIC = 235.071; Table S4). Only SLA (SS = 0.353, $F_{1,107} = 4.575$, $p = 0.015$) and not willow sex in this model (SS = 0.064, $F_{1,107} = 0.825$, $p = 0.501$) explained differences found in willow-dependent community dissimilarity (Table 2). Our leaf area NMDS analysis (Figure S1) revealed that only endophytes were positively associated with SLA. Weevils and galling mites were strongly negatively associated with SLA, while fungal rust and chewers were weakly negatively associated. Tent caterpillars, sawflies, and chrysomelid beetles were not associated with SLA.

4. Discussion

Given its relative simplicity, species richness is often a fundamental yet coarse first investigation into how geospatial, structural, or chemical parameters influence ecological communities [108-109]. Contrary to our expectations, landscape connectivity and specific leaf area (SLA) were negatively related to species richness such that less-connected willow trees and those with smaller leaves housed more species-rich communities. Landscape connectivity as well as SLA typically facilitate the establishment of plant-dependent species given that higher connectivity makes target plants more accessible across the landscape and greater SLA presents as a larger food source. However, our results indicate that connectivity and SLA may have unintended consequences for a species-poor primary successional community. Mechanistically, larger willow leaves could produce a tradeoff with palatability in which certain species may avoid leaves with unfavorable chemical traits [87-88], be that tradeoff created by willow age or sex [82, 86]. In addition, larger willow leaves could facilitate competition which leads to an imbalance of species present or alters microbial establishment, which would further impact leaf palatability [110-112]. For connectivity, species that require greater connectivity of willows to establish across a landscape may possess the ability to outcompete others, resulting in the reduced species richness we observe. Furthermore, species that can utilize less-connected willows farther from stream edges may simply avoid over-utilized trees with higher connectivity. In conjunction with sex-ratio dynamics where willows farther from streams are more likely to be male [82], community compositions are likely to be different on willows experiencing different levels of connectivity. We also found that species richness was differentially related to identity of streams nearest willow samples and that those streams organized into three separate groupings. Interestingly, species richness appeared to be randomly distributed across the Pumice Plain as none of the three groupings shared distinct geospatial placements. This potentially indicates that proximity of streams does not facilitate mutual levels of species richness. For chemistry of willow leaf litter, we found that %N and %CT positively influenced species richness while no other parameter explained differences or

improved model power. The positive influence of nitrogen is not unexpected as nitrogen is often limiting in many plant-dependent communities and higher nitrogen often results in more energetically favorable food sources [113-114]. However, tannins are typically astringent and bitter components of plant chemistry, with the unique ability to deter many types of herbivores [70]. Their positive influence on species richness could be the result of a food-limited environment where willow trees are often simply the best or only food source available to the species found on the Pumice Plain. Willow sex and interactions with other parameters did not influence species richness for the leaf area model and did not add to the explanatory power of our landscape and litter chemistry models. This result is contrary to previous findings that female willows typically facilitate species presence, particularly stem-boring weevils [83]. However, with such a species poor community, the effects of willow sex on richness may not be as prominent as they are on explicit community structure.

While species richness can provide a broad-scale view into the assembly of ecological communities, understanding influences on community composition can provide critical deeper insights into life history and dispersal characteristics that could affect establishment across the landscape [115-116]. Parallel to richness, we found that landscape connectivity, SLA, and stream identity drove differences in community structures between individual willow trees. Furthermore, willow sex predicted community dissimilarity despite no overall influence on species richness. Landscape connectivity was positively associated with weevils, fungal rust, and galling mites, meaning that willows which were more connected experienced higher rates of attack from these community members. These organisms share a common life history based on passive physical dispersal ability in which they utilize more connected habitats to move from host to host. For example, fungal rust is notably spread through spores, and potentially spreads more easily in denser tree communities with higher water availability [117], while galling mites disperse after adult eclosure through ballooning into the air and passively transferring to nearby plants [118-119]. Weevil adults seldom fly and often lay eggs on the same individual from which they emerged as larvae [83]. Directional flyers which exhibit the ability to disperse actively across the Pumice Plain and select hosts [120], such as tent caterpillars and chrysomelid beetles, were less associated with landscape connectivity alongside aphids, which have been noted to travel large distances along the ground in search of living plant material [121]. Thus, our findings indicate that passive host use is directly related to increased landscape connectivity while the ability to actively select hosts mitigates the necessity of more connected habitats. We also observed that endophytes were negatively related to connectivity, which is contrary to expectations of low dispersal ability for symbiotic microbes. Upon further investigation, however, endophytes were present on nearly every willow, and their ubiquitous life history in communities of the Pumice Plain is a finding mirrored in other studies [122]. Results for willow sex were similar to the above results for landscape connectivity, where female willows were associated with the same species as increased landscape connectivity. These results indicate that certain willow-dependent organisms differentially utilize willow trees of different sexes, meaning that placement of willow sex on the landscape can be a fundamental predictor of dependent community assembly. However, we found no association with leaf chemistry and community dissimilarity, suggesting that the influence of willow sex is more complex than potential differences in leaf chemistry. We also predicted that endophytic symbionts would prefer female willows, yet their ubiquity on all willows indicates that there may be no mechanism for endophytes to preferentially utilize females. We did not explicitly test landscape connectivity of female versus male willows, but our results suggest that female willows were associated with higher overall connectivity. However, it is important to note that we did not sample all willow individuals in the Pumice Plain and that this distinction could simply be caused by the locations where certain willows were sampled (i.e., distances from the nearest stream).

Landscape connectivity is a broad-scale metric of habitat structure that is essential in understanding how certain organisms can disperse between habitat locations [123], but it

is also important to understand how nested levels of habitat structure can assemble communities and predict dissimilarity. Stream identity represents habitat structure at a finer scale than connectivity as a proxy for physical landscape location and allows us to investigate how stream heterogeneity influences community identity. Stream identity also allowed us to test geographic location in a clearer way than with 2-dimensional geographic coordinates, given that individual sampling coordinates for willows at different streams overlapped and that differences in willow-dependent communities should be more associated with proximity to the same stream (given proximity and connectivity to each other) than with absolute coordinate locations. Furthermore, a single categorical value for each willow location was critical for testing our hypotheses with a suite of assembly characteristics (landscape, chemistry, and leaf area) rather than with more complex pairwise distances across the Pumice Plain. Overall, we observed differences in 14/21 comparisons of willow-dependent community dissimilarity between varying stream locations. These differences were driven mostly by comparisons of other streams to Redrock, Clear, and Geo-W as well as comparison to each other. These streams are spread out across the Pumice Plain and are not part of an interconnected network like Forsyth, Redrock, and Willow, thus relative geographic location appears not to influence dependent community assembly. However, willows from each stream alone harbored unique subsets of the overall dependent community (e.g., Redrock willows with aphids, Clear with chrysomelid beetles, Camp with sawflies, etc.), suggesting that specific locations of willows near individual streams (i.e., independent geographic locations) do drive community assembly. We found that the presence of tent caterpillars and weevils drove the majority of comparisons, where weevils were more important in stream comparisons of moderate distance and tent caterpillars were more important in all others. Thus, tent caterpillars may avoid connected trees that have more weevils, since the outer comparisons in both directions are streams with higher landscape connectivity overall. Tent caterpillars may also not expend energy in finding connected trees given their lack of mobility constraints or they may simply avoid willow branches established by weevils, as those branches are more likely to die and collapse. These findings enlighten our previous species richness results as the absences of other community members such as chrysomelid beetles and chewing guild herbivores also drive comparisons in which weevils are dominant, meaning that these community members are disassociated with the presence of weevils. In a species-poor community such as this, the lack of certain community members can have a powerful effect on the services and resilience of the resulting ecosystem [124-127]. Leaf area is a much finer metric of habitat structure given its role in potential food availability, but can also be important in a willow's ability to house different types of species. We found that SLA was not positively associated with any community member, while it was negatively associated with several organisms such as weevils, galling mites, and rust. This finding elucidates a potential mechanism of lower species richness associated with higher SLA, in that the presence of no one organism led to a lower presence of others on larger leaves. Rather, all community members either preferred smaller leaves or indifferently utilized willows regardless of leaf size, indicating that some composition (e.g., energy density or secondary chemicals) within larger leaves was less preferable to smaller leaves [128-129]. As a habitat metric, this finding has important implications for landscape management as leaf size can be a proxy for leaf health and thus habitat carrying capacity [130-131], and targeting certain habitat structures (e.g., willow ages or sexes) in this population can facilitate willow-dependent community establishment.

Our findings indicate that community assembly on the Pumice Plain can be directed by a variety of different factors. However, the scales at which each of these factors produced an effect were markedly different. For example, we deliberately tested willow sex with each suite of model parameters to understand parameter interactions (landscape, chemical, and leaf area), but found that willow sex variably contributed to model explanatory power and only produced differences for richness via the leaf area dataset and community structure at the landscape-level. Additionally, all chemistry, and leaf area, parameters were performed for smaller subsets of the overall data due to sampling limitations.

Considering the varied effect of willow sex, it is possible that an expanded set of data, or even for the full set of surveyed willows for this research, could elucidate heterogeneous patterns of community response to different factors. Despite this, we have illustrated that the metacommunity across willows in the Pumice Plain is regenerating based on this array of community-structuring parameters. Specifically, we found that willows distinct by different streams, leaf area, leaf chemistry, sex, and connectivity at the landscape-scale are habitat for different subsets of the full community. This heterogeneity of use by different community members is essential for facilitating metacommunity interactions among different species [10, 132], promoting ecosystem resilience [133], and establishing a diversity of ecosystem services [134]. Thus, our findings are critical for understanding community regeneration as a function of primary succession following a major disturbance that effectively reset landscape processes and dynamics. Furthermore, our work is important to direct future research focused over longer periods of time to understand the true trajectory of metacommunity regeneration, given this study was conducted at one point in time. This is especially important when considering the dynamics of newly developing streams, of which our target streams are, in that different streams may be present year-in and year-out [135]. Establishment of new trees on the landscape could be highly dependent on seasonal availability of water [136-137], and landscape connectivity will be dependent on future stream locations and their variability. Furthermore, landscape connectivity for currently established trees such as those sampled for this study will change based on shifting hydrodynamics. Thus, mapping the same suites of parameters over time will allow us to understand how the community changes over time based on these parameters and how the landscape may change in the future.

Landscape connectivity and habitat structure, in association with dispersal ability, often dictate how organisms will spread throughout a landscape [23-24]. However, effectively capturing a measure of landscape connectivity, and thus dispersal potential for organisms requiring high connectivity, is difficult. In this study we utilize circuit theory in Circuitscape, a modeling method that measures landscape connectivity not simply based on absolute distribution of a species, but as a process of varying resistance inherent to the landscape itself. The majority of other studies utilizing circuit theory analyze the connectivity between specific individuals of the same species [58-59], while we treat the connectivity produced as a community assembly process independent of willow sampling locations (i.e., using randomized starts). In this manner, we tested hypotheses of community assembly versus potential landscape connectivity and were able to make judgements about what level of connectivity was influential for specific community members. Classic measures of landscape connectivity simply do not capture these dynamics as they typically produce one output for a given pairwise comparison rather than a range of data and a testable central tendency [47-49, 53]. In our method, we effectively simulate how an organism might experience the landscape: randomness and variance are much more important given that migrants rarely enter or disperse from the same places, especially for a primary successional landscape. To our knowledge, this application of circuit theory is novel and could present new opportunities for other habitats experiencing rapid change or those which are critically understudied, given that the production of resistance maps could be readily achieved with adequate remote-sensing data.

5. Conclusions

Landscape connectivity can be a powerful tool for the management of primary successional landscapes; in our study, we found that variations in landscape connectivity can sustain variations in organismal life histories such that patchy mosaics of connectivity can support a wider range of potential community members. Furthermore, simply targeting more connected landscapes has the potential to facilitate the expansion of invasive species. If maintenance and further regeneration of metacommunity dynamics is a target for restoration and management, understanding potential pathways of invasibility is vital given the influences species like the stem-boring poplar weevil can have on the communities to

which they are introduced [84-86]. Studies in ecology often experience difficulty in extrapolating results from areas too fine-scaled to larger landscapes, but also with broader parameters such as physical location applied to more fine-scale processes. We successfully capture dynamics across three separate resolutions of community structure: leaf area, physical location via nearest streams, and overall landscape connectivity. Utilizing these dynamics, in conjunction with the maintenance of heterogeneity in community-structuring parameters (e.g., physical and chemical factors), will be key in facilitating further development and regeneration of willow-dependent communities across the Pumice Plain.

Supplementary Materials: The following supporting information can be downloaded online, Tables S1-S6; Figure S1.

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