EVIDENCE OF LOCAL ADAPTATION IN TEMPERATURE TOLERANCE TRAITS OF THE GAMETOPHYTIC AND SPOROPHYTIC STAGES IN *SOLANUM CAROLINENSE* (HORSENETTLE)

Abstract

Climate change is rapidly altering local temperature regimes and in different ways across the landscape. To cope with these rapid changes, plants species must have the capacity to respond to changes in temperature stress or risk extinction. We compared temperature tolerance traits in *Solanum carolinense* populations from Texas and Minnesota to understand how a species adapts or acclimates to extreme temperature stress. We included traits in both the gametophytic and sporophytic stages to distinguish between these distinct phases of selection. We found that mechanisms in temperature tolerance differ between populations of the south that face extreme heat regularly in Texas and northern populations that do not, in both the sporophyte and gametophyte. Our results are consistent with local adaptation and divergence of thermotolerance traits between northern and southern populations. These findings suggest that populations have the potential to adapt to rising temperatures due to climate change in the future.

Introduction

Climate change is rapidly altering environmental conditions at the local level and in particular, temperature and precipitation regimes and the severity of weather events. How will plants, a mostly sessile taxonomic group, cope with these rapid changes? Given the rapid change in local conditions, there are three ways plants can respond while avoiding extinction; quickly adapt, tolerate changing conditions through plasticity in phenotype that allows acclimation to the new conditions, or shift ranges (Janzen 1967; Molina-Montenegro and Naya 2012; Schlichting 1986). We conducted a study that focuses on the variation within populations and addresses the potential for the first two of these options in a widespread, weedy species.

The conditions across a species’ range are almost always heterogeneous and can have a variety of selective pressures that act on the populations differently. Divergent selection in two different locations can result in differing trait optima in separate populations, leading to local adaptation (Kawecki and Ebert 2004). How a species adapts or acclimates to separate locations provides a clue to how a species in one location might respond as global warming changes local conditions. Temperature is a variable that can determine species distributions and can vary greatly in both severity and consistency with geographic region (e.g. von Bϋren and Hiltbrunner, 2022). There have been many adaptations in different species that improve survival in extreme temperatures, but how do populations of the same species persist in different temperature regimes? To understand local adaptation to diverging temperature regimes, we must understand the biology of plants and how they are vulnerable to extreme temperatures.

Temperature can impact plant physiology and cell structure in a few ways. Temperature stress changes the fluidity of phospholipid bilayers. Heat increases fluidity and dissociation of membrane components from one another (Zhu et al. 2018), while cold decreases lipid adhesion and increases rigidity (Valitova et al. 2019). Both heat and cold stress results in cytoplasm leaking from the cell membrane. Plants that are more tolerant of temperature stress should have the capacity to maintain optimal cell membrane fluidity and reduce cytoplasm leakage across a wider range of temperatures. The incorporation of sterols in membranes can maintain fluidity and expand temperature range for plants (Dufourc 2008a, 2008b; Valitova et al. 2019). On the other hand, saturated fatty acids can be incorporated in the cell membrane to reduce fluidity and are often associated heat tolerance (Knight and Ackerly 2001; Zhu et al. 2018).

High temperatures also affect photosynthesis via the decreased affinity of Rubisco (enzyme responsible for carbon fixation in photosynthesis) to CO2 and increase in its affinity to O2 (Bauwe et al. 2010; Zhu et al. 2018). The fixation of O2 produces compounds that are needless and requires photorespiration to recycle components necessary for photosynthesis, in the process, reducing the efficiency of photosynthesis (Bauwe et al. 2010). In extreme temperatures, hot and cold, proteins and enzymes can be damaged or rendered inactive. This can have an immense effect on photosynthesis because the protein complexes in photosystem II and the electron transport chain can unfold (Zhu et al. 2018). The degradation of integral proteins and enzymes can also lead to the production of reactive oxygen species (ROS) through the excess absorption of light energy and prolonged excitation of chlorophyll molecules (Mishra et al. 2019; Wahid 2007; Wahid et al. 2007). Chlorophyll excitation isn’t exclusively in the thylakoid membrane, where the light reaction typically takes place. Temperature stress can damage thylakoid membranes resulting in a release of chlorophyll. Chlorophyll will continue to absorb light energy even when free from the membrane. Without a source to receive the light energy (normally photosystems in the thylakoid membrane), excess energy forms free radicals that are typically donated to oxygen molecules forming ROS, which are highly reactive and damaging to cellular components. Plants typically degrade free chlorophyll or transform chlorophyll into alternative configurations quickly, and as a result chlorophyll fluorescence decreases (Kariola et al. 2005). Plants that are capable of tolerating temperature stress have less chlorophyll degradation in the context of relatively high temperature. Oxidative stress due to ROS hinders physiological mechanisms such as photosynthesis, metabolism, and cellular structure directly or indirectly by reacting with metabolites or damaging macromolecules. Some of these cellular processes are not unique to diploid cells of the plants (sporophyte), but also occur in the haploid cells such as pollen and ovules (gametophytes). Extreme temperatures can limit pollen production, tube growth rate, and viability (Gajanayake et al. 2011; Kakani et al. 2002; Singh et al. 2008).

There is variation in the sensitivity to temperature stress and thus adaptations do lead to populations that are less sensitive. For example, cell membrane stability can be maintained in high or low temperature stress with the incorporation of fatty acids (Zhu et al. 2018) or sterols (Dufourc 2008a, 2008b; Valitova et al. 2019). The production of heat shock proteins, a chaperone protein, also reduces temperature stress by preserving the shape of other proteins and enzymes required for normal function (Frank et al. 2009; Goswami et al. 2010; Knight and Ackerly 2001; Lin et al. 2018; Liu et al. 2016; Nurminsky et al. 2018; Rhoads et al. 2005). For these adaptations to occur, temperature must be a selective pressure that influences the survival or reproduction of the species.

In angiosperms, selection can act independently in the two life stages, the sporophyte (diploid; full plants, vegetation) and the gametophyte (haploid; ovules, pollen). It has been shown that there is a substantial overlap in genes and gene expression between the two stages (Beaudry et al. 2020; Pedersen et al. 1987; Tanksley et al. 1981; Willing and Mascarenhas 1984). There is also evidence of a correlation between the gametophytic and sporophytic stages in temperature tolerance traits (Hedhly et al. 2005; Poudyal et al. 2019).

In this study, we compared plants from Minnesota and Texas and estimated temperature tolerance to extreme hot and cold conditions. Since temperature-based selection in the two life stages has the potential for inter-generational adaptations (thermotolerant pollen yields progeny with thermotolerant leaves), we incorporated variables from both the sporophyte and gametophyte. Sporophytic tolerance was measured using leaf measurements ( net photosynthesis, chlorophyll content, and cell membrane stability). The gametophytic variables were pollen germination (viability) and pollen tube growth rate. The first objective was to determine if local thermal conditions have divergently selected for temperature tolerance traits and led to adaptations reflecting regional climate regimes. We hypothesized that if the temperature regimes in the north and south have resulted in divergent selection and local adaptation of temperature tolerance, then the plants in the north would be more tolerant of cold stress and plants from the south would be more tolerant of heat stress. The second objective was to determine if there is a correlation between temperature tolerance in the gametophyte and sporophyte. If temperature stress is similar in both stages and gene expression patterns in the gametophyte and sporophyte overlap, then there is the potential for positive correlation for temperature tolerance in the two life stages.

Methods

Species Description

*Solanum carolinense* L.(Solanaceae), commonly known as horsenettle, is a weedy, herbaceous perennial that originated in southeastern North America. *Solanum carolinense* is in the Carolinense clade of the subgroup Leptostemonum characterized by abundant prickles and spines on the calyx of the flowers (Wahlert et al. 2014). Since all other species in this clade are neotropical, this species likely arose through dispersal to North America and independent diversification. Recently, this species has been reported in states across the United States, along both coasts, as far south as Texas and Florida and as far north as Minnesota and Idaho (Figure 1.1). *Solanum carolinense* reproduces both sexually and asexually. Asexually, this species utilizes clonal recruitment by growth from rhizomes. Sexual reproduction in *Solanum carolinense* is complex. First, *Solanum carolinense* is indeterminate and andromonoecious, producing mostly hermaphroditic flowers with some staminate flowers (Connolly and Anderson 2003). Second, *Solanum carolinense* has a self-incompatibility system that reduces the occurrence of self-fertilization through multiple alleles at the S-locus (Mena-Ali and Stephenson 2007; Mena-Ali et al. 2009). S-RNases are produced within the style of a flower in accordance with the S-allele and degrade RNA of pollen tubes with like S-alleles. However, the self-incompatibility system is plastic and degrades with flower age (Travers et al. 2004). This species is buzz-pollinated, meaning that a certain frequency of vibration must be applied to the anthers for pollen to release. The primary pollinators for this species are bumble bees. Once ovules are fertilized, a small, round, green to yellow tomato-like fruit develops on a truss and is dispersed by small mammals, such as skunks, and birds (Cipollini and Levey 1997).

Plant Collection

*Solanum carolinense* plants from three populations in Texas and two populations in Minnesota were collected between October 2019 and August 2020 (Figure 1.1). The three southern populations were from Collin County, Texas near McKinney (Oil Patch: 33.173465 N, -96.615402 W; Reserve: 33.159962 N, -96.619011 W; and Cemetery: 33.173672 N, -96.615096 W). At the time of collection, each population consisted of between 10 and 50 mature plants. The Reserve population was located approximately 1.5 km from the Oil Patch and Cemetery populations which were adjacent to each other (Figure 1.2). The two populations from the north were from Houston County, Minnesota and from here on will be referred to as plants from the northern region or Prairie Island (44.07959 N, -91.684545 W) and Frontenac (44.523056 N, -92.338611 W). These populations are separated by approximately 80 Km (Figure 1.3). In Colin County TX, the average monthly low temperature is 18°C (65°F) and the average monthly high is 43°C (111°F). In Houston County, MN, the average monthly low temperature is -14°C (7°F) and the average monthly high is 29°C (85°F).

Since *Solanum carolinense* reproduces asexually by growing individual plants from rhizome material, plants in close proximity may be genetically identical or ramets of the same genet. To avoid sampling two plants of the same genotype, plants with a minimum inter-plant distance of 1 meter were collected. Collections involved digging up and cutting rhizome of at least 10 cm in length and placing them in ziplock bags. Rhizomes were stored in a cooler with blue ice and shipped to Fargo, ND, where the collections were stored in a 4°C refrigerator. The rhizomes were potted in one-gallon containers with a standard potting mix and grown throughout the summer of 2020. In October, all above ground matter was cut and the rhizomes were again stored in a 4°C refrigerator to induce a period of dormancy.

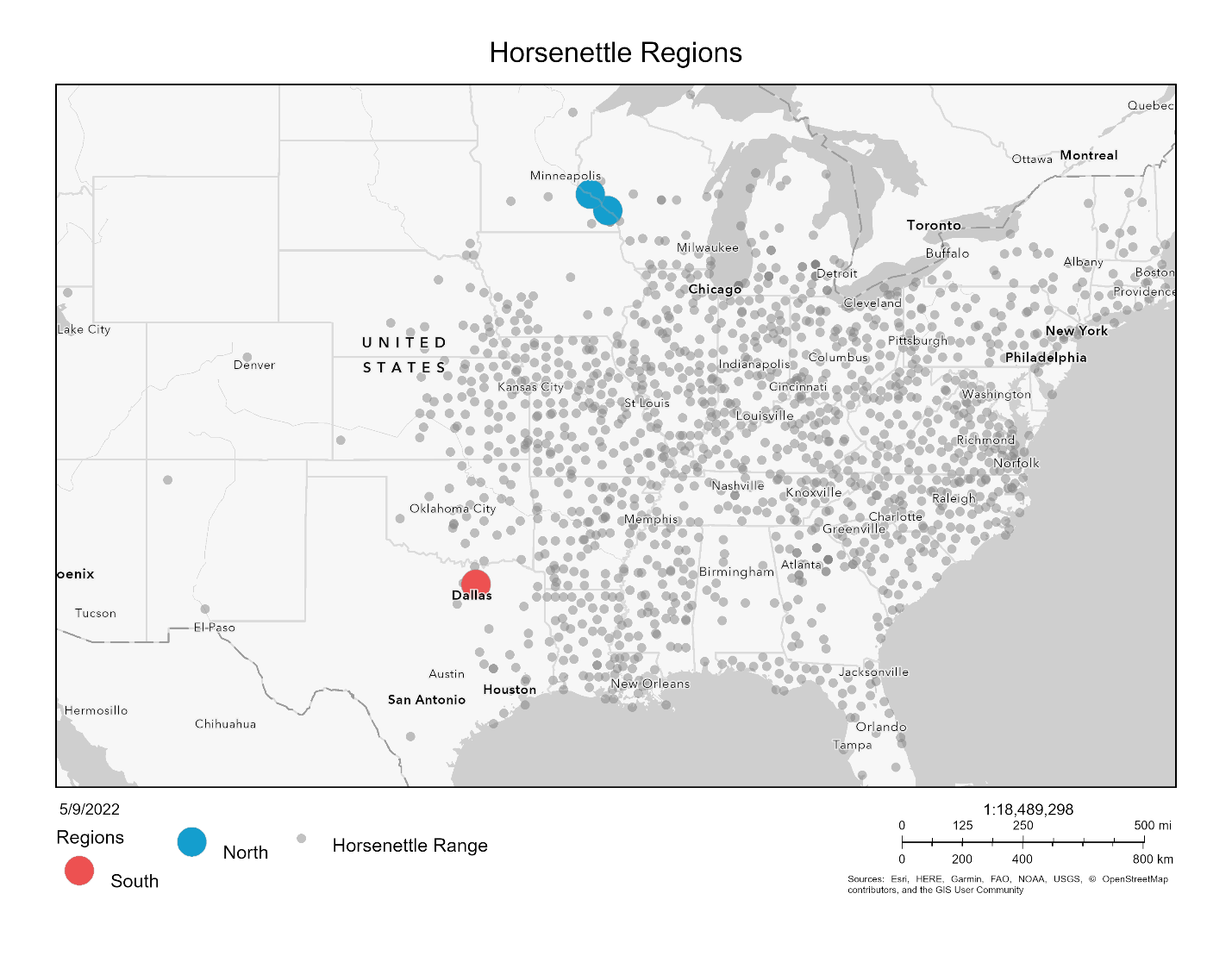
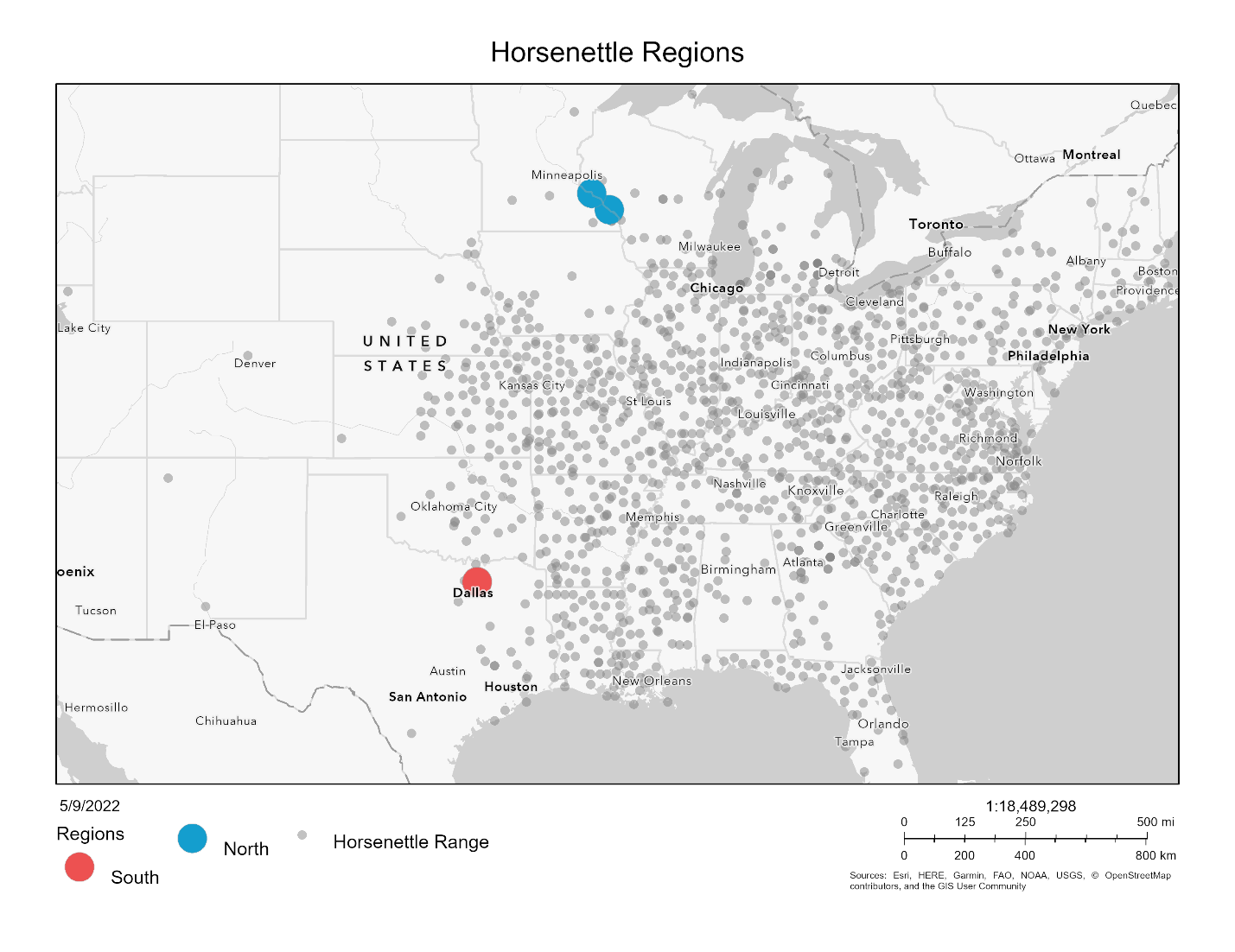
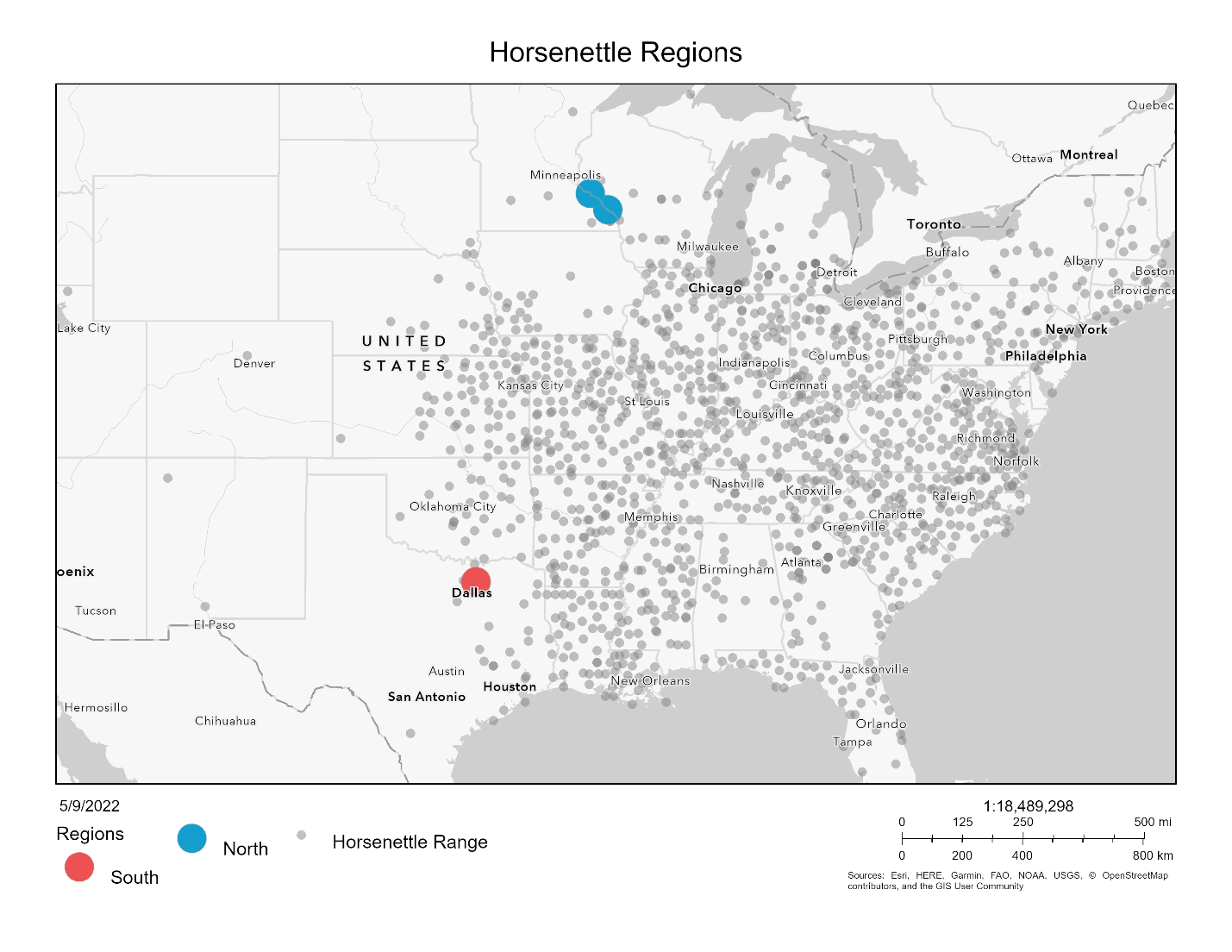


Figure 1.1. Map with collection site. Northern sites in blue and southern sites in red. Grey points indicate sites where *Solanum carolinense* was observed (EDDMapS 2022).

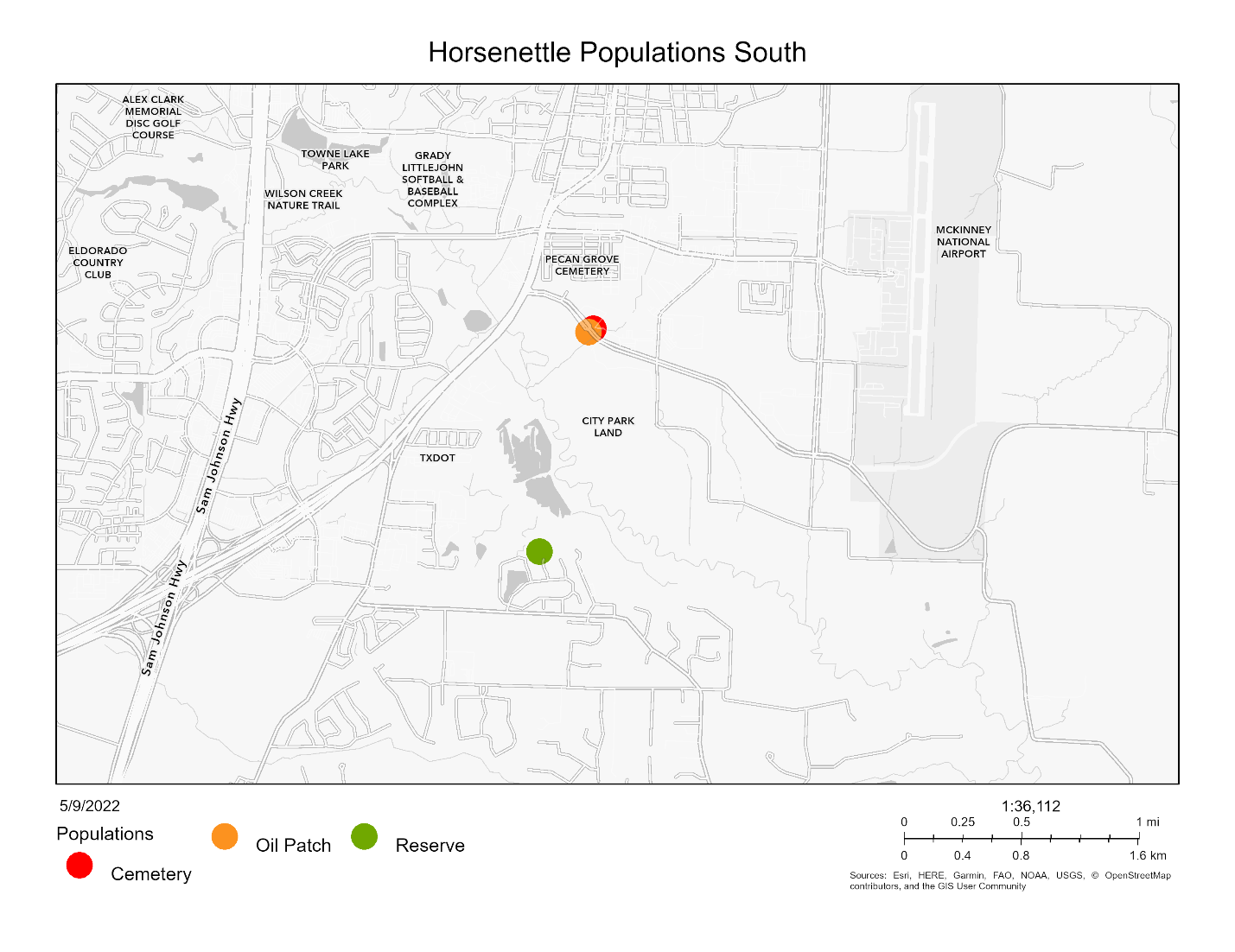
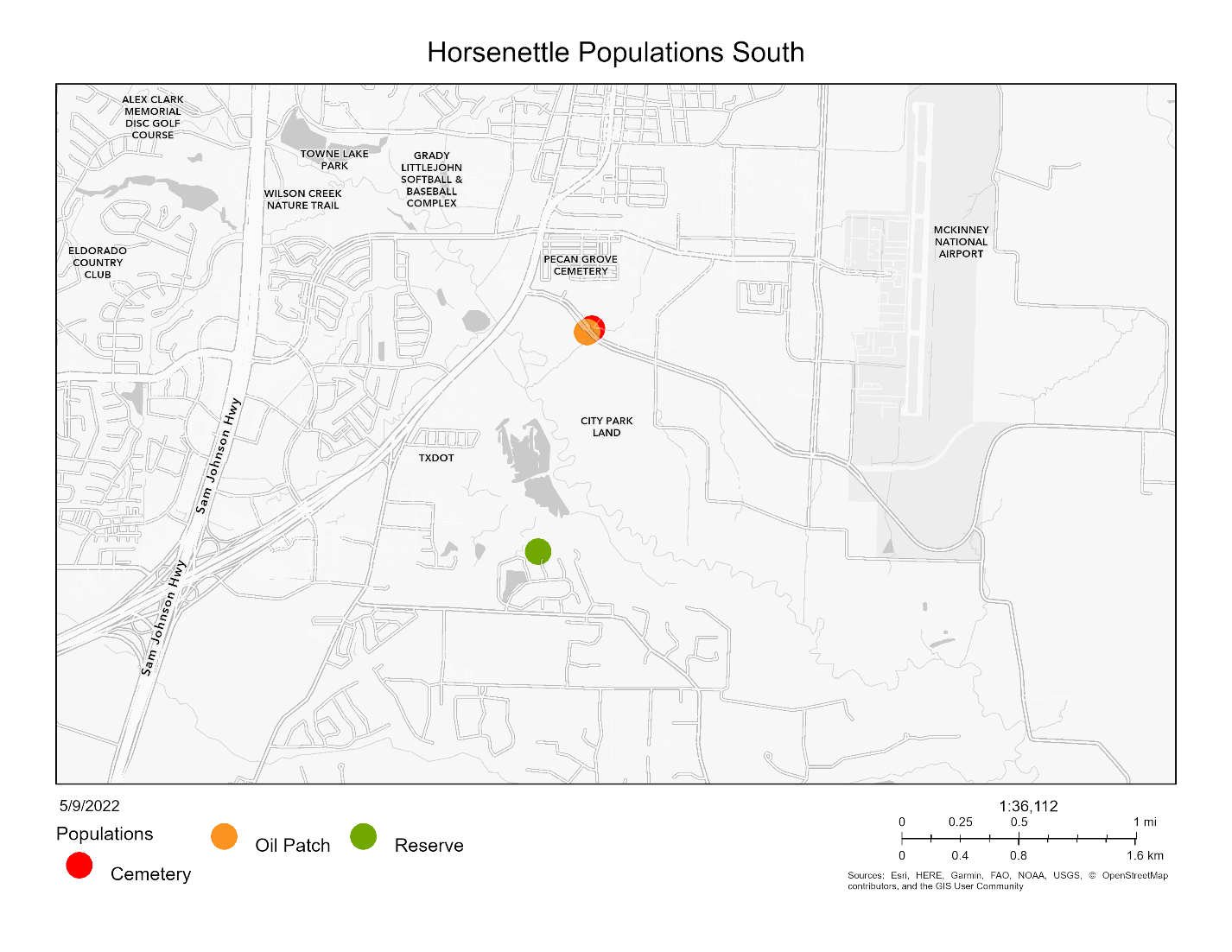
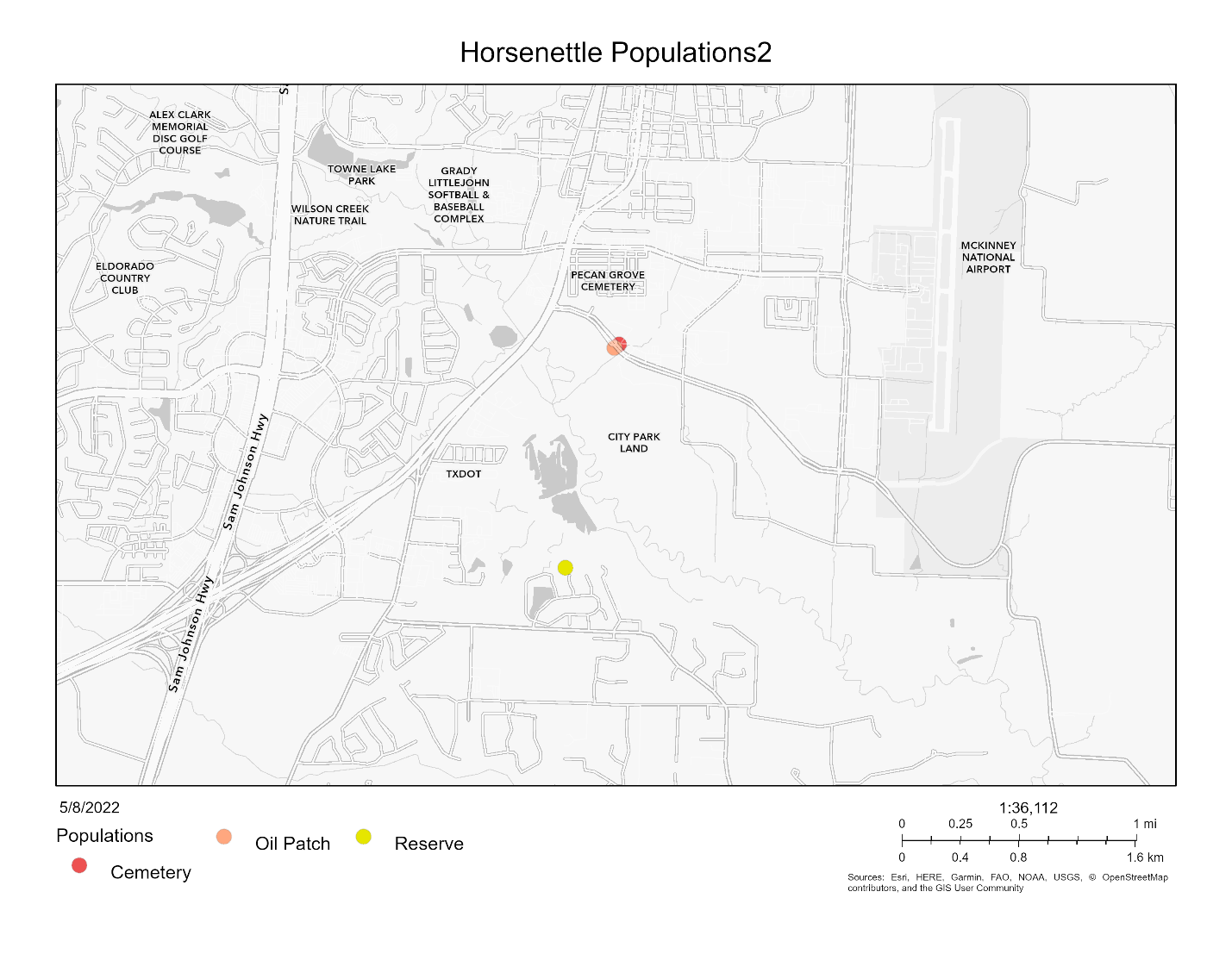
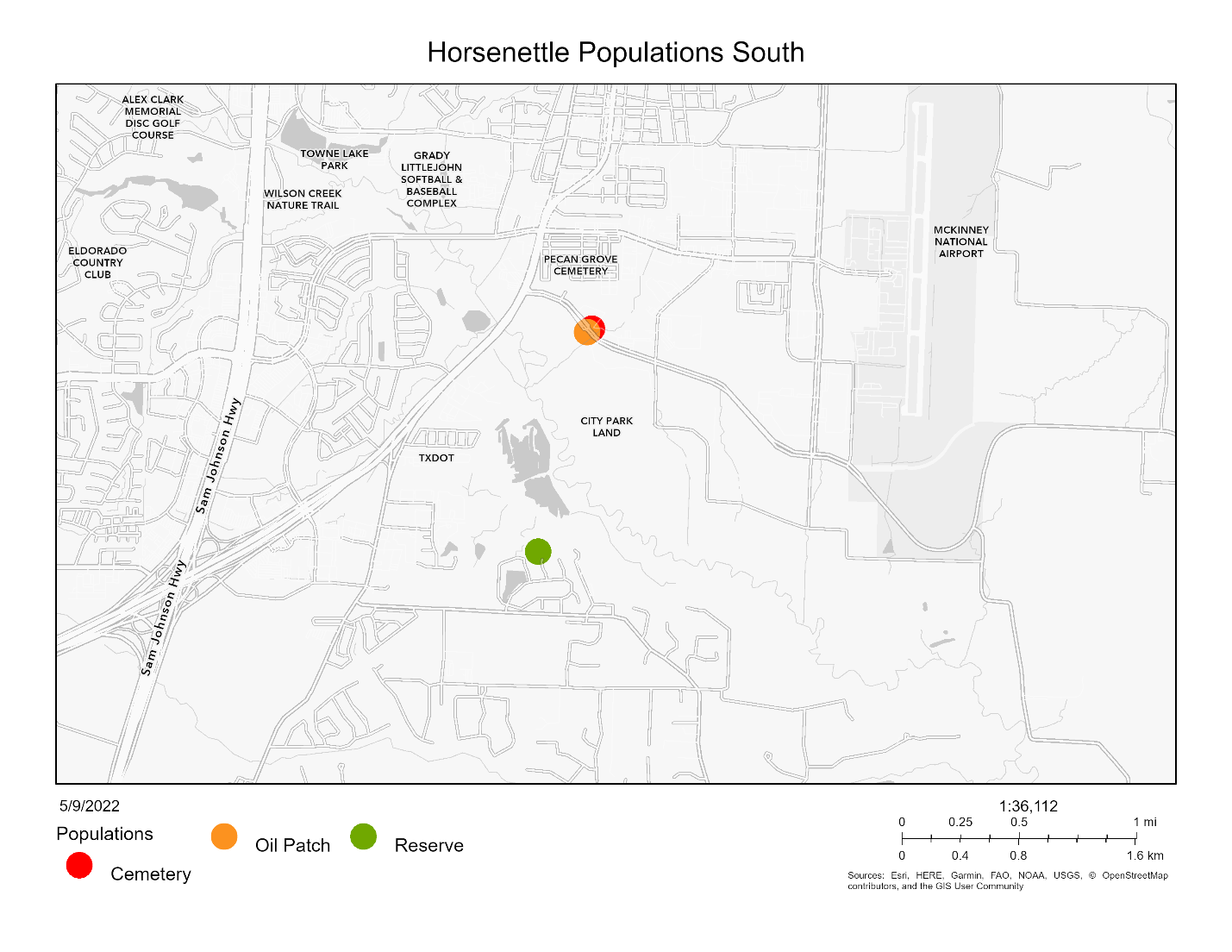


Figure 1.2. Populations in the southern region. Cemetery in red, Oil Patch in orange, and Reserve in green.

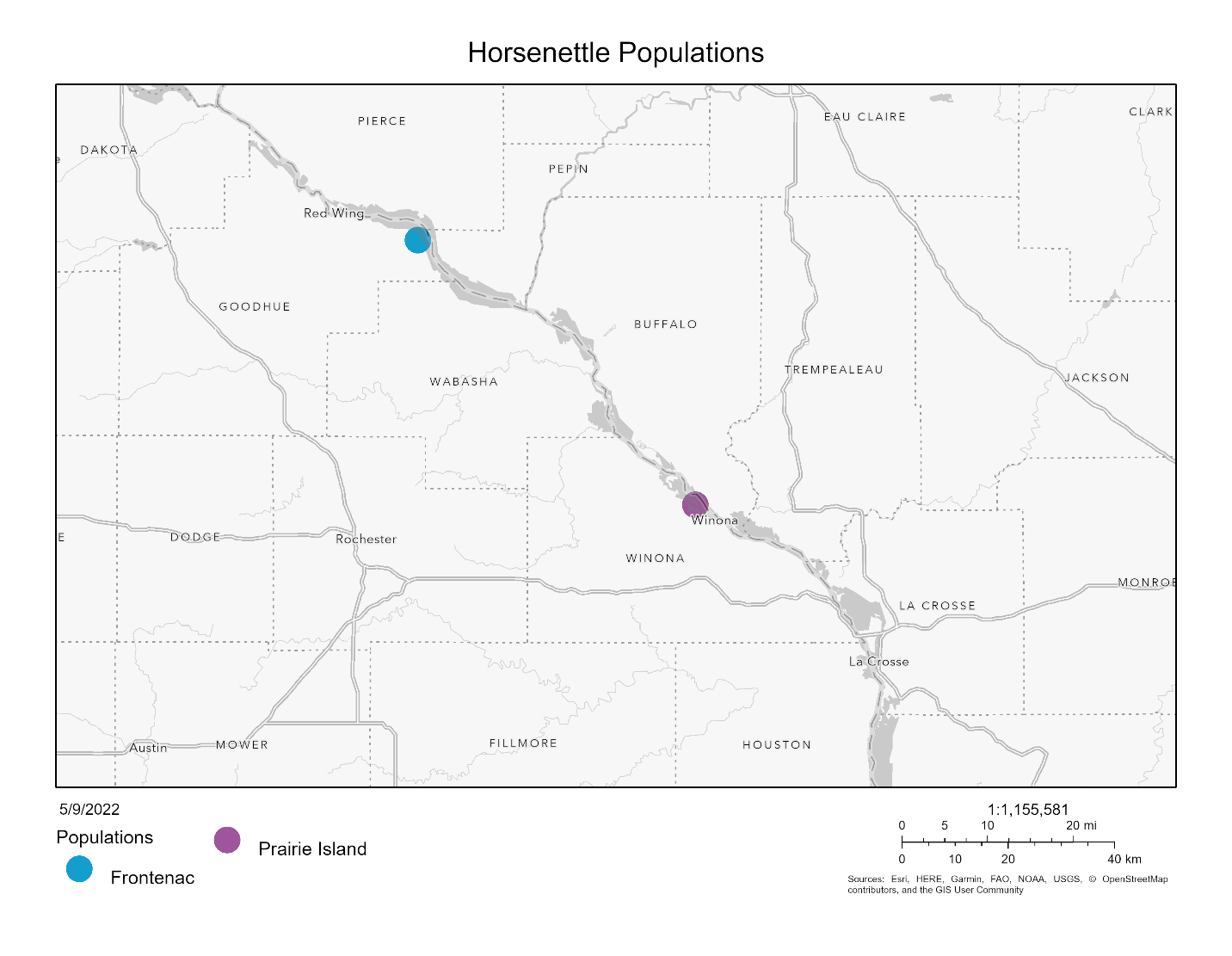
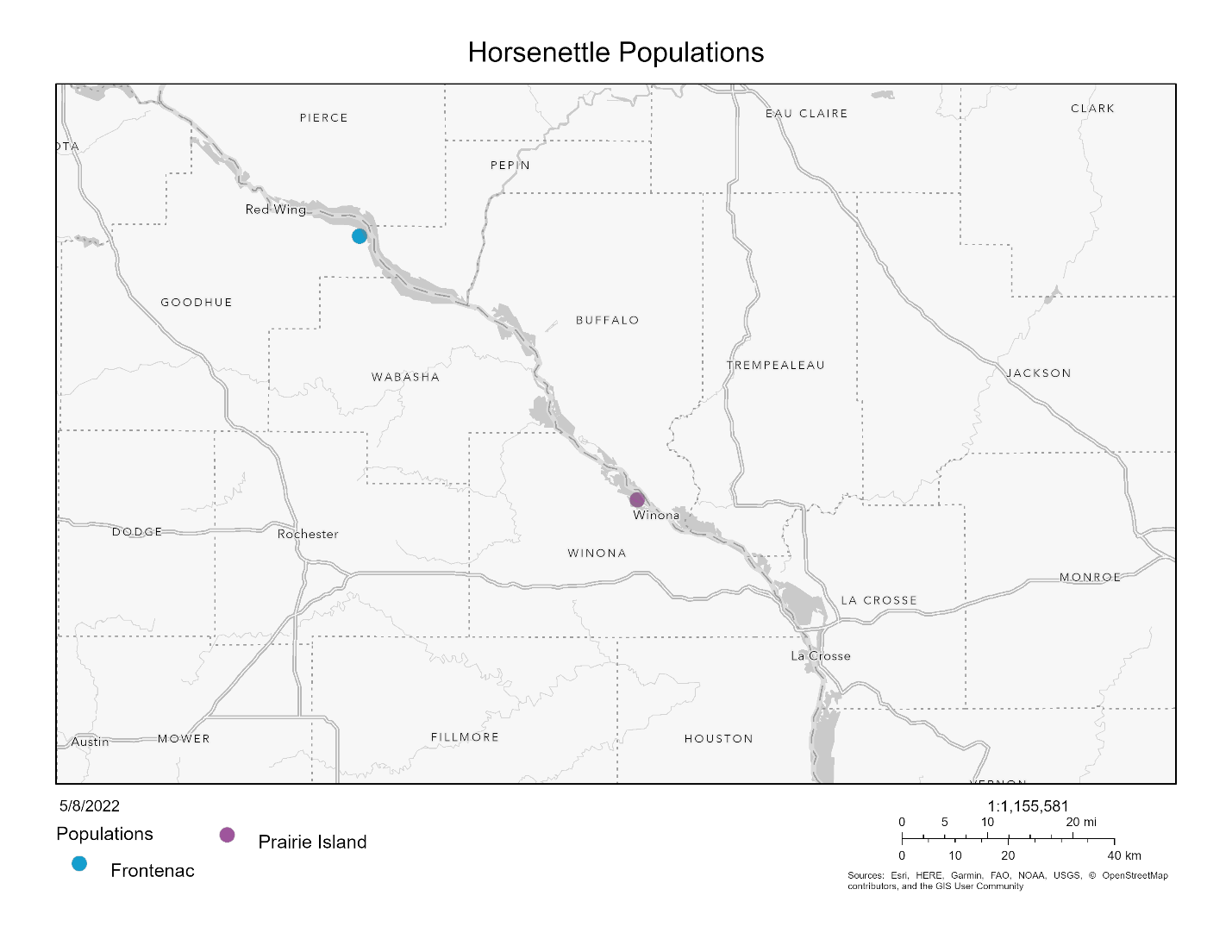


Figure 1.3. Populations in the northern region. Frontenac in blue and Prairie Island in purple.

Greenhouse Experiment

After the dormancy period (3 months), equal sections of rhizome (at least 2 cm for thick rhizomes and increased lengths for thinner rhizomes) were cut to grow ramets (genetically identical copies) in 3.8 cm diameter cone-shaped containers in the greenhouse. In total, four ramets (blocks A, B, C, and D) were grown from each genetat separate times. We started 10 or 12 ramets each week (sub-block 1-20), randomly selected from the 52 genets. Of the ramets planted each week, half were from the southern region and half were from the northern region. All ramets in block A were planted over five weeks prior to the planting of the ramets in block B and so on. Northern plants were paired with a southern plant and these pairs were randomly located on greenhouse benches. The plants were fertilized regularly with 10-10-10 fertilizer and transplanted to larger, 4.5 L containers when they outgrew the small cone-shaped containers. After approximately X weeks, we began collecting sporophytic measurements from one sub-block per week. Gametophytic data were measured when plants began flowering.

Sporophytic Traits

Cell Membrane Stability

In order to estimate tolerance of leaves to both heat and cold, we examined the cellular stability of leaf material when exposed to relatively high and low temperatures. We used a handheld conductivity meter to measure cell membrane stability (CMS) of leaves after a temperature treatment following the protocol of Gajanayake et al. (2011) and Fang and To (2016). Two large, intact leaves were removed from the middle of a plant and rinsed with deionized water. One leaf was used for the high temperature treatment and the second leaf was used for the cold temperature treatment. Twenty rounds per leaf were punched from each leaf with a hole puncher. Ten of the 20 leaf rounds were placed in a test tube for each temperature treatment (high or low) and 10 were placed in a test tube for a control treatment.

Prior to the high temperature treatment, 10 mL of deionized water was added to the control and temperature treatment test tubes. The high temperature treatment test tubes were placed in a water bath at 55°C for 20 minutes, while the control test tubes were left at room temperature. After exposure to heat, the heat treatment tube was moved to room temperature for 10 minutes prior to the first conductivity measurement.

The low temperature treatment test tubes were placed without water at 10°C for 24 hours followed by 24 hours at 4°C to acclimate the leaf rounds to cooler temperatures. The treatment tubes were then placed at -18°C for 1 hour. The control treatment tubes remained at room temperature for the total 49 hours. After the temperature treatment, 10 mL of deionized water were added to all tubes for both the treatment and control. The tubes were placed at room temperature for 1 hour prior to the first conductivity measurement.

All tubes were then subjected to a maximum damage treatment after the first conductivity measurements to quantify maximum conductivity for each sample. All test tubes were placed in a water bath at 98°C for 1 hour and then left to cool at room temperature for 15 minutes before the second conductivity measurement.

The cell membrane stability value (CMS) used for data analysis was calculated as one minus the proportion of treatment final conductivity to treatment group maximum conductivity divided by one minus the proportion of control final conductivity to control group maximum conductivity. Thus, larger values correspond with higher tolerance to temperature stress (Gajanayake et al. 2011).

Chlorophyll Content

Mishra et al. (2011) reported on the use of chlorophyll fluorescence as a measure of cold tolerance and Wahid et al. (2007) discussed the correlation between chlorophyll fluorescence and heat tolerance. We were interested in both cold and heat tolerance in this study. We used a chlorophyll meter (Opti-Sciences CCM-300) to measure chlorophyll content. The chlorophyll meter measures the fluorescence emitted at 735nm/700nm and uses a ratio based on experiments by Gittelson et al. (1998) to measure chlorophyll content in mg/m2. Two intact leaves were removed from the middle of the plant. One leaf was used for the heat treatment and the other was used for the cold treatment. Each leaf was cut in half. One half was placed in the treatment temperature and the other half was placed in a control setting at room temperature. The chlorophyll content was measured for both halves before and after the temperature treatment.

The high temperature treatment was 60°C for 1 hour. The leaf halves in the cold treatment were subjected to 4°C for 1 hour followed by 1 hour in -18°C. The leaf halves were moved to room temperature for two hours prior to the second cold treatment measurement. Leaves in all treatments were kept in complete darkness.

To incorporate the control and treatment groups in one measurement, the chlorophyll content ratio (CHPL) was calculated as the compliment of the difference between the proportions of the final treatment chlorophyll content to the initial treatment chlorophyll content and final control chlorophyll content to initial control chlorophyll content. Thus, larger values correspond with higher temperature tolerance (citation).

Photosynthesis

We used a LI-6400 infrared gas analyzer with a red/blue light source to measure net photosynthetic rate (µmol CO2m-2s-1) on leaves before and after the whole plant was exposed to the temperature treatment. The following settings were used for photosynthesis measurements: flow rate 500 μmol s-1, reference CO2 420 μmol CO2 mol-1, reference H2O 0 mmol H2O mol-1, ParIn\_μml 400 μmol m-2 s-1.

The high temperature treatment was 33°C and the low temperature treatment was 10°C.. All four ramets, if alive, for the 52 genets were subjected to both treatments with a rest period of one week between them. Several plants died or lost leaves by the time net photosynthetic rate was measured and thus were not included. Ramets A and C were subjected to the high temperature treatment first; ramets B and D were subjected to the low temperature treatment first. The proportion of the photosynthetic rate measurement after the treatment to before was calculated as our measure of photosynthetic temperature tolerance (PS). Any value below zero and above one was omitted prior to analysis.

Gametophytic Traits

We measured two pollen traits as estimates of male thermotolerance during the gametophytic stage: 1) the propensity for pollen grains to germinate (pollen germination) and 2) the growth rate of pollen tubes while exposed to a range of temperatures. Once a plant from the north and from the south flowered, we removed a mature flower from both plants. Since *Solanum carolinense* is buzz-pollinated, a device crafted from a nose hair trimmer and a paper clip was used to mimic the vibrations needed to release pollen from the anther. Pollen from each flower was thus dispersed over five petri dishes containing 3% Bacto-Agar based growth medium (sucrose, Ca(NO3)2, MgSO4, KNO3, H3BO3) following the protocol of Reddy and Kakani (2007). The dusted plates were each placed at one of the five temperature treatments (10°C, 20°C, 25°C, 30°C, 40°C) for 16 hours in a refrigerator (10°C), Conviron E7/2 environmental chamber (20°C), or three drying ovens (25°C, 30°C, 40°C). After the temperature treatments, each plate was covered with a thin layer of ethanol to halt further pollen tube growth and stored at 4°C until data collection could begin. Four pictures of each plate were taken using a microscope (Leica DM500 microscope, Leica ICC50 HD camera) and the LAS EZ 2.1.0 software. Pollen did not evenly cover petri dishes; therefore, pictures were taken in locations where pollen was visible. The petri dish was positioned so pollen visible to the naked eye (miniscule white spots) was under the objective. The petri dish was not repositioned once pollen grains were viewed magnified to avoid sampling bias when taking the pictures.

Pollen germination (Germ) was measured by counting the number of pollen grains that produced pollen tubes and dividing that by the total number of pollen grains observed. All pollen grains in an image were counted until at least 100 pollen grains were observed. Pollen was considered germinated if it produced a tube that was at least half the diameter of the pollen grain. We used the percent of pollen grains with tubes out of the total number of pollen grains as our measure of pollen germination.

Pollen tube growth rate (PTGR) was determined by first measuring the 10 longest pollen tubes in each of the 4 images using the software ImageJ (Schneider et al. 2012). Pollen tubes were only included if they were completely visible in the picture. The actual length of each tube was calculated by tracing the length of each tube, calculating length in pixels and then calibrating each measurement with a stage micrometer. We calculated the mean of the 20 longest tubes out of the 40 measured per plate and estimated growth rate by dividing the mean length by the time allowed for growth (16 hours).

Data Analysis

All data were analyzed in R 4.1.2 (R Core Team 2020). In order to measure differences in sporophytic traits between plant origins and among genets, we fit linear mixed effects models using the lmer function from the *lmerTest* package (Kuznetsova et al. 2017). Region (north vs. south) was considered the fixed effect and block (A, B, C, D) and genet nested in population as random effects. We dropped the genet nested in population term for cell membrane stability and both random effects terms for heat, net photosynthetic rate to avoid overfitting the model. Since the genet nested in population term was significant for some variables, we compared population and genets independently. Populations were compared using a linear mixed effects model (*lmerTest*; function lmer) with population as the fixed effect and block as the random effect. We used an analysis of variance model in the *stats* package (R Core Team 2020), to determine if there were differences between genets for each of the sporophytic variables. Since there was a significant block effect in some of the variables, we compared plants from the north and south within block using a paired t-test (*stats*; function t.test). To determine if variation within the northern and southern regions differed, we used the Bartlett’s test of homogeneity of variance (*stats*; function bartlett.test).

For the gametophytic variables, we fit temperature performance curves to the multiple temperature measurements taken for each plant that flowered using the nls.multstart function in the *rTPC* package (Padfield and O'Sullivan 2021). Of the 25 temperature performance curves available in the *rTPC* package, the quadratic\_2008 and the weibull\_1995 models had the lowest AIC values. The weibull\_1995 model was eliminated from our analyses because maximum values extracted by the weibull\_1995 model were infinite for some of the northern plants. From the quadratic curves of each plant that flowered, we extracted three key values for both pollen germination and pollen tube growth rate: the temperature minimum, temperature optimum, and temperature maximum. We then used the key values in an analysis of variance (*stats*; function aov) to determine if there were differences between region and among genets. One outlier was identified using the Grubbs’ test for outliers, grubbs.test function in the *outliers* package (Komsta 2011), and subsequently dropped from the analysis.

We used correlation analysis (*stats*; function cor) using Pearson’s method to determine if there were any correlations between sporophytic and gametophytic variables. We conducted correlation analysis for all plants together and then the northern and southern plants separately. The Holm-Bonferroni method (*stats*; function p.adjust) was used to adjust p-values to account for multiple correlations. To incorporate relationships between all the variables and examine amalgamated differences among regions and populations, we conducted principal component analysis (PCA) (*stats*; function prcomp). We first conducted PCA on all the sporophytic variables and all gametophytic variables separately and then all variables collectively. Photosynthetic rate was not included in the collective PCA because of limited sample size. We extracted the eigenvalues for the first three principal components for all three PCAs. The eigenvalues for each principal component were compared for the two regions using t-tests (*stats*; function t.test).

Results

Sporophytic Variables

Cell Membrane Stability

When *Solanum carolinense* plants from the north were compared to the south, we found no significant difference in cell membrane stability following the hot treatment (HCMS), but there was a significant difference in the cold treatment (CCMS; Figure 1.4, Table 1.1). Southern plants had significantly higher CCMS values than northern plants. We found a significant difference among genotypes in the hot treatment, but not in the cold treatment (Figure 1.5, Table 1.1). For both the hot and cold treatments, there were significant differences between the populations. Population differences mostly followed the regional patterns in CCMS. For HCMS, one population (Oil Patch) from the southern region was less tolerant than all other populations (Appendix Figure A1, Table A1).

Because we could not grow all the experimental plants at the same time due to lack of space, we made the above comparisons among regions and genotypes in five different temporal blocks over the course of the spring and summer. To avoid confounding treatments with temporal effects, plants from different regions were paired with each other within blocks. When we tested for the presence of block effects, we found significant effects for both hot and cold CMS (Figure 1.6). Plants grown at different times in the greenhouse had different CMS ratios. We started growing the plants in the winter and early spring and outside temperatures gradually rose during that time (Appendix Figure A2). Acclimation to higher temperatures later in the year could account for the block differences observed. To remove block effects, we conducted paired t-tests of northern versus southern plants for each of the variables. When plants from the north and south were compared for HCMS, there was a significant difference between the regions (Figure 1.6) but only in the first block. In that block, northern plants had a higher HCMS than those from the south. For CCMS, there was a significant difference between regions for blocks B and C (Figure 1.6). In both cases, southern plants were more tolerant of the cold temperatures than northern plants.

Chlorophyll Content Stability

Chlorophyll content was measured before and after either a heat stress (HCHPL) or cold stress (CCHPL) and the calculated value that incorporates the two measurements was used as a proxy for temperature tolerance. As the CHPL increases, the individual sporophyte is more tolerant of the temperature treatment (Gajanayake et al. 2011). There was a significant difference between plants originating in the north and south for both the hot and cold treatments (Figure 1.4). Northern plants were more tolerant of both heat and cold than were southern plants regardless of block (Table 1.1). We found a significant difference among individual genotypes (Figure 1.5, Table 1.1) and populations (Appendix Figure A1, Table A1) for the cold treatment, but not for the hot treatment. The two regions also differed in variation for HCHPL. In the hot treatment, northern plants had significantly more variation than southern plants (Bartlett’s test p-value = 1.68E-4; Figure 1.7).

Net Photosynthetic Rate

We used net photosynthetic rate after thermal stress as a physiological indicator of temperature tolerance. PS is the ratio of net photosynthetic rate after the treatment (heat or cold) divided by net photosynthetic rate before the treatment. Increasing PS indicates increasing temperature tolerance of either hot or cold thermal stress (Poudyal et al. 2019). For both the cold (CPS) and hot (HPS) treatments, there was no significant difference between north and south (Figure 1.4, Table 1.1). There were also no significant differences among blocks and genotypes for both the hot and cold treatments. There was a significant difference between populations for CPS (Appendix Figure A1, Table A1).

Table 1.1. Results from the mixed linear model for the difference in region (north vs south) and the one-way analysis of variance results for the difference between individual genets. Red font color highlights observed outcomes when the result was different from the expected pattern. Asterisk indicates analysis with one outlier removed determined using Grubbs’ test for one outlier. Bolded values represent relationships that were statistically significant. Statistic values reported in the Appendix (Table A2), as well as results from a mixed model using only control values (Appendix Table A3).

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
|  |  | Region | | | Genet | |
|  | Variable | Expected | Observed | p-value | Observed | p-value |
| Sporophyte | Cell Membrane Stability (Heat) | S > N | - | 0.06102 | **Yes** | **0.013** |
| Cell Membrane Stability (Cold) | N > S | **S > N** | **0.0117** | No | 0.886 |
| Chlorophyll Content (Heat) | S > N | **N > S** | **0.0405** | No | 0.38 |
| Chlorophyll Content (Cold) | N > S | N > S | **9.96E-11** | **Yes** | **1.05E-07** |
| Photosynthetic Rate (Heat) | S > N | - | 0.997 | No | 0.127 |
| Photosynthetic Rate (Cold) | N > S | - | 0.77 | No | 0.883 |
| Gametophyte | Pollen Germination (Tmax) | S > N | **N > S** | **0.00037** | **Yes** | **0.0251** |
| Pollen Germination (Topt) | S > N | **N > S** | **0.000685** | **Yes** | **0.0351** |
| Pollen Germination (Tmin) | S > N | - | 0.331 | **Yes** | **\*0.0135** |
| Pollen Tube Growth Rate (Tmax) | S > N | - | 0.568 | No | 0.418 |
| Pollen Tube Growth Rate (Topt) | S > N | - | 0.77 | No | 0.608 |
| Pollen Tube Growth Rate (Tmin) | S > N | - | 0.683 | No | 0.496 |

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Figure 1.4. Regional differences for temperature tolerance traits including hot and cold cell membrane stability (HCMS, CCMS), hot and cold chlorophyll content stability (HCHPL, CCHPL), hot and cold net photosynthetic rate (HPS, CPS). Center line of boxplot is the median value for the region. Shared letters represent statistically non-significant differences between regions. Variables with significant differences denoted with asterisks: CCMS (F1,50 = 7.792, p = 0.006), HCHPL (F1,51 = 4.334, p = 0.043), and CCHPL (F1,50 = 64.652, p = 1.6e-10).

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Figure 1.5. Genotype differences for temperature tolerance traits including hot cell membrane stability (HCMS) and cold chlorophyll content stability (CCHPL). Genets are ordered by increasing ratio for each variable. The center line in boxplot is the median of the measurements among the ramets of one genet. There is a significant difference among the genets for HCMS (F = 1.5, p = 0.029) and CCHPL (F = 3.341, p = 6.1e-9). Plots of genet effect for other variables are in appendix (Figure A3).

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Figure 1.6. Cell membrane stability across temporally independent blocks and colored by region. The center line of the boxplot is the median of the measurements taken for each region within a ramet. There is a significant difference between blocks for hot cell membrane stability (HCMS, p = 0.0297) and cold cell membrane stability (CCMS, p = 7.30e-05). Asterisks indicate a significant difference between regions from a paired t-test of regions for each block independently. There was a significant difference between regions for HCMS block A (t = -2.910, p = 0.015), CMS block B (t = 2.190, p = 0.040), and CMS block C (t = 2.073, p = 0.049). Results from paired t-tests between blocks for each variable located in the appendix (Table A4).

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Figure 1.7. Hot chlorophyll content (HCHPL) vs cold chlorophyll content (CCHPL) for plants from the north and south. Ellipse indicating 95% confidence interval for multivariate T distribution. Results from Bartlett’s test for heterogeneity of variance between regions for all variables located in the appendix (Table A5).

Gametophytic Traits

Pollen Germination

Of all genets included in this study, 20 from the north flowered and 10 from the south flowered. The number of ramets that flowered for each genet differed, so the total number of plants that flowered were 32 from the north and 29 from the south. We fit quadratic curves (Appendix Figure A4) to temperature performance profiles of each plant for pollen germination at five temperatures (Figure 1.8). From the quadratic fit, we calculated the minimum (Tmin), maximum (Tmax), and optimal (Topt) temperature of pollen germination for each individual. There was a significant difference between regions for Tmax and Topt (Figure 1.8, Figure 1.9). Plants from the north germinated more readily at high temperatures and had higher thermal optima than plants from the south. There was no significant difference between the two regions for Tmin. The genets were significantly different from one another for Tmin, Tmax, and Topt (Figure 1.10, Appendix Figure A5). One outlier was identified using the Grubbs’ test for outliers and subsequently dropped from the analysis.

Pollen Tube Growth Rate

The pollen tube growth rates for each individual were also fit with a quadratic curve to estimate the Tmin, Tmax, and Topt. There were no significant differences between plants from the north and south for any of the three variables (Appendix Figure A6). There were also no significant differences among genets (Appendix Figure A7, Figure A8).

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Figure 1.8. Percent germination and mean pollen tube growth rate (PTGR) for *Solanum carolinense* pollen grains from the north (blue) and south (red) across a temperature gradient (10°C, 20°C, 25°C, 30°C, 40°C). Thin lines and points represent each individual plant that flowered. Thick lines indicate the mean value for the region at each temperature.



A

B

\*

A

B

\*

A

A

Figure 1.9. Estimates for the maximum (Tmax), optimal (Topt), and minimum (Tmin) germination temperature extracted from quadratic fits of the germination data for each individual. Asterisks and different letters indicate significant differences. There was a significant difference between regions for Tmax (F = 14.28, p = 3.7E-4) and Topt (F = 12.85, p = 6.85E-4).

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Figure 1.10. Boxplots of the maximum (Tmax), optimal (Topt), and minimum (Tmin) pollen germination temperatures by genet. There was a significant difference between the genets for Tmax (F = 2.064, p = 0.025), Topt (F = 1.952, p = 0.035), and Tmin (F = 2.284, p = 0.0135). Asterisk indicates the outlier removed for analysis.

Correlations

We used correlation analysis to identify relationships between hot and cold tolerance for the sporophytic and gametophytic variables. Pearson’s correlations were determined for all pairings of variables. When all plants were included, there were no significant correlations between the gametophytic and sporophytic variables (Figure 1.11, Appendix Table A6). However, there were two significant correlation coefficients between gametophytic variables. Maximum and minimum pollen tube growth rate were positively correlated (r = 0.46). Maximum pollen tube growth rate and maximum pollen germination were positively correlated (r = 0.3).

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Figure 1.11. Correlation matrix of all plants. Gametophytic (labels blue font) and sporophytic variables (labels red font) with significant Pearson’s correlations for all study plants.

When the correlation analysis was performed on all variables for the regions separately, there were different results. For the northern plants, there were no significant correlations. The southern plant had one significant correlation between Tmin germination and Tmax germination (r = -0.63; Appendix Figure A9).

Principal Component Analysis

We conducted principal component analysis to further explore relationships among all variables and the sporophytic and gametophytic variables separately. For the full PCA, we included all gametophytic and sporophytic variables, except HPS and CPS due to inadequate sample size. The first three principal components accounted for 57% of the variation (full PCA plots and loadings in the appendix Figure A11, Table A7). There was little divergence between regions. When the eigenvalues of the principal components were compared between regions, PC2 was the only principal component that showed a significant difference (t58 = -2.69, p = 0.0092). Chlorophyll content (HCHPL and CCHPL) loads primarily on PC2 and is likely driving the divergence between northern and southern plants.

Sporophytic PCA

In the sporophytic variables PCA, the first three principal components explained 60% of the variation. The variables HCMS and HPS primarily loaded on PC1 (Table 1.2, Figure 1.13). The second and third principal components were mostly influenced by CCHPL and HCHPL respectively. There was a significant difference between the regions for the eigenvalues extracted from both PC2 (t78 = -5.09, p = 2.39e-06) and PC3 (t101 = 2.38, p = 0.019). The divergence in PC2 can be explained by the opposite responses we observed for CCMS and both chlorophyll content treatments. Northern plants have a higher chlorophyll content ratio for both treatments, while southern plants had less cell membrane damage in the cold treatment. PC1 did divide HCMS and CCMS, suggesting an antagonistic relationship between the two variables, though there was no correlation between the two that was statistically significant. Hot and cold treatment variables were also divided in PC3. HPS and HCHPL were opposite in direction to CPS and CCHPL.

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Figure 1.12. Plots of the results of principal component analysis for the sporophytic variables. A) PC1 and PC2, B) PC2 and PC3, C) PC1 and PC3. Ellipsoid indicating 95% confidence interval. PC1 explains 22.38% of the variance, PC2 explains 21.55% of the variance, and PC3 explains 16.79% of the variance. Tables with principal component importance for PC1 through PC6 in the Appendix (Table A8).

Table 1.2. Results from principal component analysis of only sporophytic variables. Loadings for each of the variables on the principal components

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
|  | PC1 | PC2 | PC3 | PC4 | PC5 | PC6 |
| HCMS | 0.613999 | 0.02998 | 0.344975 | 0.147492 | 0.573797 | 0.390002 |
| CCMS | -0.35207 | 0.435008 | 0.204072 | -0.57534 | 0.520509 | -0.20789 |
| HCHPL | 0.284794 | -0.36797 | -0.33836 | -0.75803 | -0.08744 | 0.294536 |
| CCHPL | 0.117118 | -0.57752 | 0.579996 | -0.14114 | -0.03843 | -0.5431 |
| HPS | 0.577968 | 0.302596 | -0.40867 | 0.000171 | 0.044159 | -0.63673 |
| CPS | 0.264909 | 0.499375 | 0.470594 | -0.22955 | -0.6235 | 0.13244 |

Gametophytic PCA

In the gametophytic PCA, the first three components explained 92.5% of the variance. Pollen germination variables divided the northern and southern plants (Figure 1.14). Tmax and Topt loaded evenly in the opposite direction of Tmin for both PC1 and PC2 (Table 1.3, Figure 1.14). There was a significant difference between north and south for the eigenvalues extracted from PC2 (t46 = -3.17, p = 0.0025). PTGR variables loaded evenly on the first two principal components, indicated by the common diagonal direction among the PTGR variables (Table 1.3, Figure 1.14).

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Figure 1.13. Plot of the results of principal component analysis of the gametophytic variables. PC1 describes 48% of the variation and PC2 explains 27%. A table of importance of principle components 1 through 6 is in the Appendix (Table A9).

Table 1.3. Results from principal component analysis of only gametophytic variables. Loadings for each of the variables on the principal components

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
|  | PC1 | PC2 | PC3 | PC4 | PC5 | PC6 |
| Germ.Tmin | -0.01334 | 0.405643 | 0.812217 | 0.376238 | -0.00058 | -0.18446 |
| Germ.Topt | 0.418665 | -0.45436 | 0.390064 | -0.00262 | 0.001723 | 0.682727 |
| Germ.Tmax | 0.407763 | -0.5446 | 0.164764 | -0.10069 | -0.00179 | -0.707 |
| PTGR.Tmin | 0.367661 | 0.452838 | 0.127813 | -0.75119 | -0.28131 | 0.00071 |
| PTGR.Topt | 0.523981 | 0.308129 | -0.20261 | 0.123985 | 0.757676 | -0.00193 |
| PTGR.Tmax | 0.498538 | 0.180049 | -0.3219 | 0.518304 | -0.58888 | 0.001496 |

Discussion

Regional Differences

If *Solanum carolinense* has locally adapted to the respective temperature regimes in TX and MN, we would expect that plants from the north would be more tolerant of cold temperatures and plants from the south would be more tolerant of hot temperatures. Rather than a clear-cut difference between north and south for hot and cold treatments, there were mixed results that suggest divergence between regions in ways we had not anticipated. In contrast to our expectations, northern plants were generally more tolerant of extreme heat than southern plants, but also had more variation in certain trait values. Northern plants had higher chlorophyll content (HCHPL) and baseline cell membrane stability (HCMS; Figure 1.6) under hot conditions, as well as higher maximum and optimal temperatures for pollen germination in comparison to southern plants (Table 1.1). Conversely, southern plants had increased tolerance for cell membrane stability in cold conditions (CCMS). These results suggest that adaptation to extreme temperatures is complex and may reflect avoidance strategies rather than physiological mechanisms to withstand thermal stress.

There was no significant difference between regions for HCMS for all study plants together, but there was a significant difference for plants in block A. Temperatures in the greenhouse progressively rose throughout the spring and summer leading to a block effect in both the hot and cold treatments of CMS. In block A, northern plants had a higher HCMS, but this difference degraded in the later blocks during the times when greenhouse temperatures during plant development increased. A possible explanation for the block effect on CMS is that plants have the capacity to induce heat tolerance as they acclimate to warmer conditions (Clarke et al. 2004). Block A is the best representative measurement of baseline heat tolerance for HCMS, and later blocks likely represent induced heat tolerance, which may be more dramatic in southern plants. Conversely, plants from the south had more stable cell membranes when exposed to an extreme cold treatment. The counter gradient pattern we measured for northern and southern plants may be due to constraints of adaptation to extreme heat or cold. Adapting to match the extreme environmental conditions may not be advantageous or possible, reducing the variation in a population for tolerance in extreme conditions. Thus, populations in locations that do not experience extreme temperatures on one end of the spectrum may have more variation than those that do experience extreme temperatures, leading to the counter gradient results we attained for CMS.

Plants from the north had more stable chlorophyll content in both the hot (HCHPL) and cold treatments (CCHPL; Table 1.1). More stable chlorophyll content may be explained by northern plants experiencing a larger range of temperatures. Between 2018 and 2021, temperatures during the growing season (March to September) in Houston County, MN ranged from -28°C to 34°C (62°C difference), while in Collin County, TX they ranged from -7°C to 42°C (49°C difference). Since the temperate conditions of Minnesota are more variable and rarely exceed temperatures likely to stop plant growth (Hatfield et al. 2011), populations in the north may have evolved to acclimate to temperature stress, while plants in the south do not. Furthermore, northern plants also had significantly more variation in HCHPL than southern plants. This may suggest that there is stabilizing selection occurring in the southern region for heat tolerance in chlorophyll content. Less variation in HCHPL in the south may contribute to the counter-gradient results we attained. If northern plants experience less heat stress selection and have greater variation, then there may be more potential to have individuals with high HCHPL.

Pollen from the north had a higher propensity to produce pollen tubes (Germ) at high temperatures than their southern counterparts. Pollen germination was higher in pollen grains from the northern plants than those in the south for both Tmax and Topt (Table 1.1). The distinct difference between north and south suggests that there might be sensitivity to high temperatures and an adaptive response occurring. Since southern populations experience extremely high temperatures more regularly than northern plants, there may be an avoidance strategy in southern populations whereby pollen grains remain dormant at high temperatures. In contrast, there is no selection for dormancy at high temperatures in the north . Rutley et al. (2022) proposed the two-baskets model categorizing pollen, which states that there are high-ROS and low-ROS subpopulations of pollen within anthers of flowering species. The low-ROS pollen have a lower metabolic rate than high-ROS pollen due to partial dehydration during development. The two subpopulations of pollen are adaptive and beneficial under different conditions as they allow for asynchrony in pollen germination, permitting some pollen to remain dormant in a stressful environment, such as extreme heat or drought, and grow pollen tubes later in more favorable conditions. Keller and Simm (2018) compared the transcriptome and proteome in *Solanum lycopersicum* (tomato) and determined that pollen have two responses during heat stress – direct and delayed translation. Luria et al. (2019) later showed that *Solanum lycopersicum* has pollen that fall in the low-ROS and high-ROS groups, supporting the two-basket model in a species closely related to *Solanum carolinense*. We hypothesize that *Solanum carolinense* populations in the south have higher proportions of low-ROS to high-ROS pollen grains than those in the north due to stronger selection from increased exposure to extreme heat in the south. Low-ROS pollen that remains dormant would not be adaptive in northern populations, with little exposure to high temperature stress.

There was a significant negative correlation between Tmax and Tmin germination in southern plants. This correlation supports the two-basket model. The negative correlation indicates that plants with pollen that germinate readily at high temperatures also germinate at low temperatures, while those that have a lower Tmax have a higher Tmin. \ High-ROS pollen would germinate in any condition (extreme heat and cold stress). Low-ROS pollen would not germinate as freely during stressful conditions. Since plants of the south may have evolved to have the dual pollen types, there may be more variation in pollen activity driving this correlation.

There was no significant difference between northern and southern plants for net photosynthetic rate in both the hot and cold treatments. Net photosynthesis was the only sporophytic variable where the whole plant was placed in a temperature treatment and leaves were measured on the plant. The plant may compensate for temperature stress through physiological mechanisms, such as increasing transpiration. Therefore, the temperature treatments may not have stressed the plants to the extent that temperature tolerance for the northern and southern plants was distinguishable.

The response of plants from the two regions to extreme cold were considerably more mixed. There was no significant difference between northern and southern populations for Tmin of either pollen germination or pollen tube growth rate. Of all cold traits only two sporophytic traits (CCMS and CCHPL) differed between regions and were not consistent. Pollen may have a low temperature limit on physiological processes necessary for pollen tube growth that are consistent across all populations.

Inter-Generational Relationships

Tanksley et al. (1981) highlited the association between selection in the gametophyte and sporophyte when they found a correlation between allozyme genes expressed in both stages. Based on their findings and several studies that followed (Hedhly et al. 2005; Pedersen et al. 1987; Poudyal et al. 2019; Willing and Mascarenhas 1984), including studies on temperature tolerance (Hedhly et al. 2005; Poudyal et al. 2019), we hypothesized that there would be a correlation between temperature tolerance in the sporophyte and the gametophyte. Correlations between the two life stages have implications for the rate of temperature tolerance evolution. Selection in either stage for similar traits that are expressed independently would rapidly increase or decrease the allele frequency of associated genes in a population. Furthermore, in the gametophyte, there is a lack of dominance allowing selection to act on one allele (Haldane 1932, Beaudry et al. 2020). The alleles selected for in the gametophyte can then affect traits in the sporophyte.

There were no significant correlations between any of the gametophytic and sporophytic variables in our study, suggesting that there are different mechanisms mitigating temperature stress in the two stages. This is not the first study to find differences in patterns for extreme temperature tolerance in the sporophyte and gametophyte. Dominguez et al. (2005) conducted a study to determine if pollen selection can be used to improve cold tolerance in the gametophyte by selecting pollen from cold tolerant plants (sporophyte). They found that pollen selection did not improve pollen viability and formation in cold and explained their results by describing how the genes mediating cold stress may be expressed in the sporophyte tissue surrounding the site of pollen formation, rather than the pollen grains themselves.

Another explanation for the lack of coordinated response to temperature stress between the two life stages is that horsenettle has not been located in MN and TX long enough for selection to act on the populations. All populations included in this study were located toward the edge of the range for this species. Time for selective pressures to act on the populations may be insufficient for local adaptation to occur. The first record of *Solanum carolinense* in Minnesota is from 1939 and in Houston County 1975 (Bell Museum Plants, Minnesota Biodiversity Atlas; The University of Minnesota). The first record in Texas is from 1917 and the closest record of horsenettle to Collin County is from 2011 (Lundell Herbarium, Billie L. Turner Plant Resources Center; The University of Texas at Austin).

Conclusion

Our results are consistent with a process of local adaptation due to temperature acting as a selective pressure. The results of this study do not completely support our original predictions based on the assumption that northern latitudes are simply cooler than southern latitudes. The measurements of chlorophyll content did provide some evidence that populations from areas with larger thermal ranges, such as those in higher latitudes, have more variation and possibly more phenotypic plasticity, which is consistent with the climate variability hypothesis. The block effects observed in both HCMS and CCMS also suggest that there is plasticity in the phenotype when exposed to long-term changes in ambient temperature. Lastly, we found evidence of southern plants avoiding pollen germination in high temperatures by increasing the proportion of low-ROS to high-ROS pollen.

These results could inform restoration efforts by changing the way we think about seed sourcing and adaptive potential in a rapidly changing environment. Seeds from the south may have evolved stress responses to temperature that are lacking in northern populations or vice versa. The evidence for the two-basket model in a wild species is also a novel finding that could add to our perception of the influence gametophytic traits have on species persistence in extreme environments.

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