

Genetics of Climate Adaptation Using Genome-Wide Association in Switchgrass

Unknown

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types are shown.

Why Switchgrass?

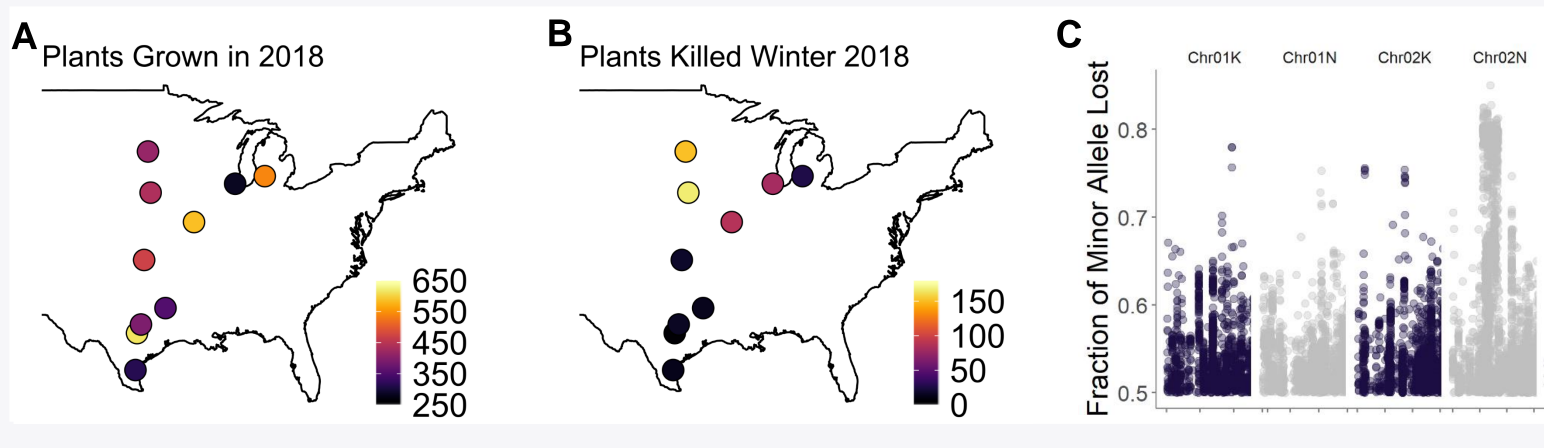
BIOFUELS: Switchgrass (*Panicum virgatum*) is the model herbaceous bioenergy crop; it has high biomass yields and is productive on marginal lands; its ethanol yield can exceed that of maize on a per acre basis.

CLIMATE ADAPTATION: Natural switchgrass populations are found across North America east of the Rocky Mountains, and span eight USDA cold hardiness zones. Natural selection to climate has shaped the genetics of these populations and shaped phenotypic groupings into *upland* (mostly northern), *lowland* (mostly southern), and *coastal* ecotypes. These ecotypes likely have different adaptations to climate.

Diversity Panel Ecotype Major phenotypic UplandCoastal and genetic LowlandUnknown groups in the tetraploid Genetic switchgrass subpopulation diversity panel. Midwest □ Atlantic Plant specimens for the five major \times Admixed

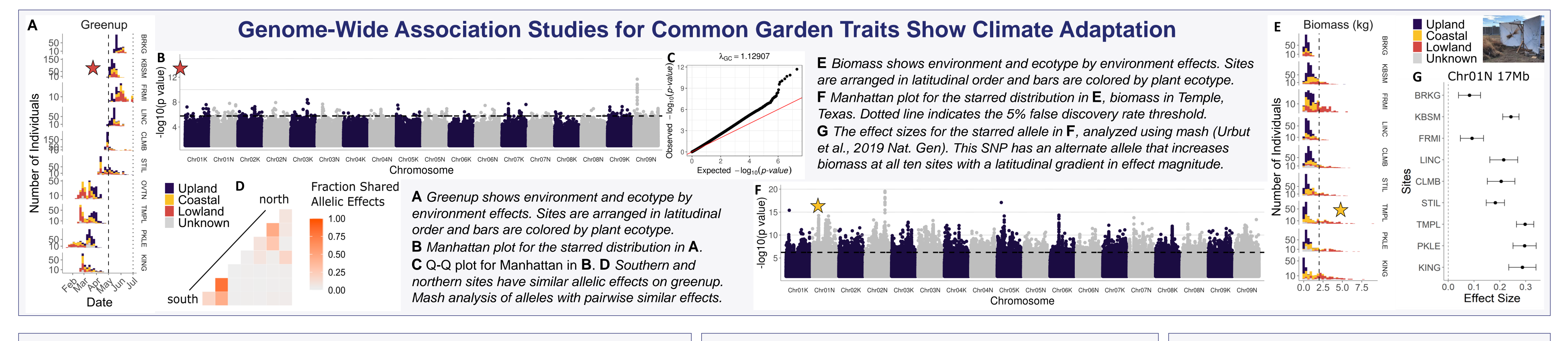
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Continent-Scale Adaptation Using Common Gardens



- A The diversity panel was planted at ten common garden locations in 2018.
- **B** Four northern common garden locations had significant overwinter plant death in winter 2018-2019. **C** Specific gulf alleles were lost at high frequency in the north in the winter of 2018/2019. The fraction of alleles lost varied across the genome (first four chromosomes are shown).

Genome-Wide Association Studies of Environments at Plant Locations of Origin Homologous to Significant Homologous to /Cold Shock Protein 1 bioclim Ethylene-responsive Homologous to variables transcription factor Suppressor of Auxin | Resistance 1 SNPs with significant associations with one or more bioclim variable. A Significant SNPs were typically associated -0.50-0.250.00 0.25 0.50 Effect Size with 14-18 bioclim variables when analyzed with mash (Urbut et al., 2019 Nat. Gen.) Three nearby genes are noted. **B** The starred allele in **A** has alternate alleles found in plants from consistent temperature, warmer climates.



Take-home message

Winter survival in the northern U.S. is a major selective agent in switchgrass.

Environmental GWAS show allelic effects that partition plants into "winter kill" and "non-winter kill" locations; e.g., warmer climates with more consistent temperatures (and more variable precipitation) versus cooler climates with more variable temperatures and more consistent precipitation.

Common garden phenotypes show genotype by environment interactions and strong signals of genetic association that vary by location.

switchgrassGWAS R Package

https://github.com/Alice-MacQueen/switchgrassGWAS

- I. Run Genome-Wide Association with `pvdiv_gwas`
- 2. Find annotations with `pvdiv_table_topsnps`
- 3. Analyze data at multiple sites using mash, with:
 - `pvdiv_bigsnp2mashr`
 - `mash_standard_run`
 - `mash_plot_manhattan_by_condition`
 - `mash_plot_effects`

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