# Supplementary Information

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# About

This is supplementary material intended to be paired with files on GitHub at the repository https://github.com/avahoffman/urban-weed-genomics.

# Introduction

In this experiment, we used quaddRAD library prep to prepare the sample DNA. This means that there were both two unique outer barcodes (typical Illumina barcodes) *AND* two unique inner barcodes (random barcode bases inside the adapters) for each sample - over 1700 to be exact!

The sequencing facility demultiplexes samples based on the outer barcodes (typically called 5nn and i7nn). Once this is done, each file still contains a mix of the inner barcodes. We will refer to these as "sublibraries" because they are sort of halfway demultiplexed. We separate them out bioinformatically later.

### 0.1 Raw Data File Naming - Sublibraries

Here's a bit of information on the file name convention. The typical raw file looks like this:

AMH\_macro\_1\_1\_12px\_S1\_L001\_R1\_001.fastq.gz

• These are author initials and "macro" stands for "Macrosystems". These are on every file.

AMH\_macro

• The first number is the *i5nn* barcode for the given sublibrary. We know all these samples have a *i5nn* barcode "1", so that narrows down what they can be. The second number is the *i7nn* barcode for the given sublibrary. We know all these samples have a *i7nn* barcode "1", so that further narrows down what they can be.

1\_1

- This refers to how many samples are in the sublibrary. "12px" means 12-plexed, or 12 samples. In other words, we will use the inner barcodes to further distinguish 12 unique samples in this sublibrary. 12px
- This is a unique sublibrary name. S1 = 1 i5nn and 1 i7nn.
- This means this particular file came from lane 1 of the NovaSeq. There are four lanes. All samples should appear across all four lanes.

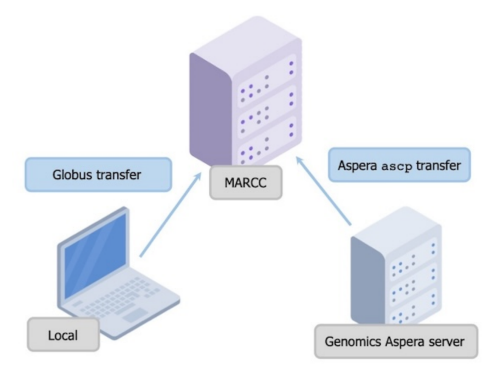
L001

- This is the first (R1) of two paired-end reads (R1 and R2).
- The last part doesn't mean anything it was just added automatically before the file suffix (fastq.gz)
   001.fastq.gz

### 0.2 A Note on File Transfers

There are three main systems at play for file transfer: the local machine, the sequencing facility's (GRCF) Aspera server, and MARCC. The Aspera server is where the data were/are stored immediately after sequencing. MARCC is where we plan to do preprocessing and analysis. Scripts and text files are easy for me to edit on my local machine. We used Globus to transfer these small files from my local machine to MARCC.

Midway through this analyses, we transitioned to another cluster, JHU's Rockfish. Scripts below, with the exception of file transfer from the Aspera server, should reflect the new filesystem, though you will have to adjust the file paths accordingly.



## 0.3 A Note on Species Names

Throughout this study, we examined 6 species. Sometimes we used abbreviations for easier file naming. These are:

- 1. Cynodon dactylon, "CD", or Bermuda grass
- 2. Digitaria sanguinalis, "DS", or crabgrass
- 3. Erigeron canadensis, "EC", or horseweed
- 4. Lactuca serriola, "LS", or prickly lettuce
- 5. Poa annua, "PA", or bluegrass
- 6. Taraxacum officinale, "TO", or dandelion

# 1 File Transfer

Referred to through files as "Step 1". Files can be found in the O1\_transfer\_files/ directory.

This directory contains files named in this convention: 01-aspera\_transfer\_n.txt. These are text files containing the names of fastq.gz files that we wanted to transfer from the sequencing facility's Aspera server to the computing cluster (MARCC). This was to maximize ease of transferring only certain files over at once, since transferring could take a long time. We definitely did this piecemeal. Possible file names shown in Aspera Transfer File Names. There are multiple of these files so that we could parallelize (replace n with the correct number in the command used below). This text file will need to be uploaded to your scratch directory in MARCC.

Files were then transferred using the following commands. Before starting, make sure you are in a data transfer node. Then, load the aspera module. Alternatively, you can install the Aspera transfer software and use that.

```
module load aspera
```

Initiate the transfer from within your scratch directory:

```
ascp -T -18G -i /software/apps/aspera/3.9.1/etc/asperaweb_id_dsa.openssh
--file-list=01-aspera_transfer_n.txt
--mode=recv --user=<aspera-user> --host=<aspera-IP> /scratch/users/<me>@jhu.edu
```

# 2 File Concatenation and Stacks Installation

Referred to through files as "Step 2". Files can be found in the 02 concatenate and check/ directory.

### 2.1 Concatenate Files for each Sublibrary

**Step 2a**. We ran my samples across the whole flow cell of the NovaSeq, so results came in 8 files for each demultiplexed sublibrary (4 lanes \* paired reads). For example, for sublibrary 1\_1, we'd see the following 8 files:

```
AMH_macro_1_1_12px_S1_L001_R1_001.fastq.gz
AMH_macro_1_1_12px_S1_L001_R2_001.fastq.gz
AMH_macro_1_1_12px_S1_L002_R1_001.fastq.gz
AMH_macro_1_1_12px_S1_L002_R2_001.fastq.gz
AMH_macro_1_1_12px_S1_L003_R1_001.fastq.gz
AMH_macro_1_1_12px_S1_L003_R2_001.fastq.gz
AMH_macro_1_1_12px_S1_L004_R1_001.fastq.gz
AMH_macro_1_1_12px_S1_L004_R2_001.fastq.gz
AMH_macro_1_1_12px_S1_L004_R2_001.fastq.gz
```

The 02\_concatendate\_and\_check/02-concat\_files\_across4lanes.sh script finds all files in the working directory with the name pattern \*\_L001\_\*.fastq.gz and then concatenates across lanes 001, 002, 003, and 004 so they can be managed further. The "L001" part of the filename is then eliminated. For example the 8 files above would become:

```
AMH_macro_1_1_12px_S1_R1.fastq.gz
AMH_macro_1_1_12px_S1_R2.fastq.gz
```

Rockfish uses slurm to manage jobs. To run the script, use the sbatch command. For example:

```
sbatch ~/code/02-concat_files_across4lanes.sh
```

This command will run the script from within the current directory, but will look for and pull the script from the code directory. This will concatenate all files within the current directory that match the loop pattern.

#### 2.2 Download and Install Stacks

**Step 2b.** On Rockfish, Stacks will need to be downloaded to each user's code directory. Stacks, and software in general, should be compiled in an interactive mode or loaded via module. For more information on interactive mode, see interact --usage.

```
interact -p debug -g 1 -n 1 -c 1
module load gcc
Now download Stacks. We used version 2.60.
wget http://catchenlab.life.illinois.edu/stacks/source/stacks-2.60.tar.gz
tar xfvz stacks-2.60.tar.gz
Next, go into the stacks-2.60 directory and run the following commands:
./configure --prefix=/home/<your_username>/code4-<PI_username>
make
make install
```

export PATH=\$PATH:/home/<your\_username>/code4-<PI\_username>/stacks-2.60

The filesystem patterns on your cluster might be different, and you should change these file paths accordingly.

# 3 PCR Clone Removal

Referred to through files as "Step 3". Files can be found in the O3\_clone\_filter/ directory.

### 3.1 Run PCR Clone Removal Script

Step 3a. The 03-clone\_filter.sh script runs clone\_filter from Stacks. The program was run with options --inline\_inline --oligo\_len\_1 4 --oligo\_len\_2 4. The --oligo\_len\_x 4 options indicate the 4-base pair degenerate sequence was included on the outside of the barcodes for detecting PCR duplicates. The script uses the file name prefixes listed for each single sub-pooled library in 03-clone\_filter\_file\_names.txt and loops to run clone\_filter on all of them. Possible file names shown in clone\_filter File Names.

#### 3.2 Parse PCR Clone Removal Results

**Step 3b**. If you want to extract descriptive statistics from the clone\_filter output, you can use the 03.5-parse\_clone\_filter.py script to do so. It can be run on your local terminal after transferring the clone\_filter.out logs to your local computer.

```
source("03_clone_filter/examine_clones.R")
make_cloneplot()
```

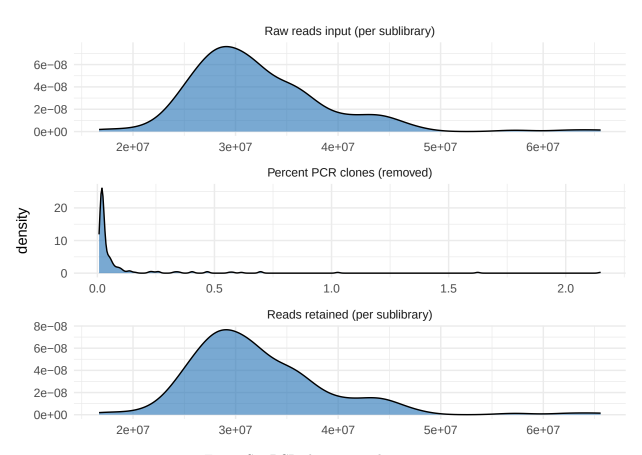


Figure S1: PCR clone removal statistics

# 4 Sample Demultiplexing and Filtering

Files can be found in the O4\_demux\_filter/ directory.

# 4.1 Demultiplex and Filter

Step 4a. The 04-process\_radtags.sh script runs process\_radtags from Stacks. The program was run with options -c -q --inline\_inline --renz\_1 pstI --renz\_2 mspI --rescue --disable\_rad\_check. The script uses the same file prefixes as Step 3 - 03-clone\_filter.sh. Each sub-pooled library has a forward and reverse read file that was filtered in the previous step. Like the above section, the script uses the file name prefixes listed for each single sub-pooled library in 04-process\_radtags\_file\_names.txt and loops to run process\_radtags on all of them. Possible file names shown in clone\_filter File Names.

Each sub-pooled library also has a demultiplexing file (04-demux/ directory) that contains the sample names and inner(i5 and i7) barcodes. For example, the sublibrary 1\_1, we'd see the following barcode file:

```
ATCACG AGTCAA DS.BA.PIK.U.1
CGATGT AGTTCC DS.BA.PIK.U.2
TTAGGC ATGTCA DS.BA.PIK.U.3
TGACCA
       CCGTCC DS.BA.PIK.U.4
ACAGTG GTCCGC DS.BA.PIK.U.5
GCCAAT GTGAAA DS.BA.DHI.U.1
CAGATC GTGGCC DS.BA.DHI.U.2
ACTTGA GTTTCG DS.BA.DHI.U.3
GATCAG CGTACG DS.BA.DHI.U.4
TAGCTT
       GAGTGG DS.BA.DHI.U.5
GGCTAC
       ACTGAT
               DS.BA.GA.U.1
CTTGTA
       ATTCCT
              DS.BA.GA.U.2
```

The 'process\_radtags' command will demultiplex the data by separating out each sublibrary into the individual samples. It will then clean the data, and will remove low quality reads and discard reads where a barcode was not found.

### 4.2 Organize files

Step 4b. In a new directory, make sure the files are organized by species. In the process\_radtags script, we specified that files be sent to ~/scratch/demux/\*sublibrary\_name\* (reasoning for this is in Step 4c), but files should manually be organized into species folders (i.e., ~/scratch/demux/\*SPP\*) after process\_radtags is performed. For example, the file "DS.MN.L01-DS.M.1.1.fq.gz" should be sent to the ~/scratch/demux/DS directory.

Note: this is not automated at this point but it would be nice to automate the file moving process so it's not forgotten at this point.

#### 4.3 Assess the raw, processed, and cleaned data

Step 4c. In the script for Step 4, we have specified that a new output folder be created for each sublibrary. The output folder is where all sample files and the log file will be dumped for each sublibrary. It is important to specify a different output folder if you have multiple sublibraries because we will be assessing the output log for each sublibrary individually (and otherwise, the log is overwritten when the script loops to a new sublibrary).

The utility stacks-dist-extract can be used to extract data from the log file. First, we examined the library-wide statistics to identify sublibraries where barcodes may have been misentered or where sequencing error may have occurred. We used:

```
stacks-dist-extract process_radtags.log total_raw_read_counts
```

to pull out data on the total number of sequences, the number of low-quality reads, whether barcodes were found or not, and the total number of retained reads per sublibary. Look over these to make sure there are no outliers or sublibraries that need to be checked and rerun.

Next, we used:

```
stacks-dist-extract process_radtags.log per_barcode_raw_read_counts
```

to analyze how well each sample performed. There are three important statistics to consider for each sample.

- 1. The proportion of reads per sample for each sublibrary indicates the proportion that each individual was processed and sequenced within the overall library. This is important to consider as cases where a single sample dominates the sublibrary may indicate contamination. (Field prop\_sample\_per\_library).
- 2. The number of reads retained for each sample can be an indicator of coverage. It is most likely a good idea to remove samples with a very low number of reads. Where you decide to place the cutoff for low coverage samples is dependent on your dataset. For example, a threshold of 1 million reads is often used but this is not universal. (Field retained\_reads).
- 3. The proportion of reads retained for each sample can also indicate low-quality samples and will give an idea of the variation in coverage across samples. (Field prop\_reads\_retained\_per\_sample).

Output for sublibraries for this step are summarized in process\_radtags-library\_output.csv.

Output for individual samples for this step are summarized in process\_radtags-sample\_output.csv.

The script O4c-process\_radtags\_stats.R was used to create many plots for easily assessing each statistic. Output from this step can be found in figures/process\_radtags/ where figures are organized by species.

The script 04c-radtags\_filter\_summary.R summarizes the filtering results from all samples.

```
source("04_demux_filter/04c-radtags_filter_summary.R")
make_filterplot()
```

# 4.4 Identify low-coverage and low-quality samples from

Step~4d. Using the same output log and the above statistics, we removed low-coverage and low-quality samples that may skew downstream analyses.

Samples were identified and removed via the following procedure:

- 1. First, samples that represented less than 1% of the sequenced sublibrary were identified and removed. These samples correlate to low-read and low-coverage samples.
- 2. Next, a threshold of **1 million retained reads per sample** was used to remove any remaining low-read samples. Low-read samples correlate to low coverage and will lack enough raw reads to contribute to downstream analyses.

Good/kept samples are listed in process\_radtags-kept\_samples.csv.

Discarded samples are listed in process\_radtags-discarded\_samples.csv.

```
source("04_demux_filter/04c-radtags_filter_summary.R")
make_manual_discard_plot()
```

Note: At this point, we started using Stacks 2.62 for its multi-threading capabilities. Functionality of the previous steps should be the same, however.

# 5 Stacks: Metapopulation Catalog Building and Parameter Search

Files can be found in the O5\_ustacks\_and\_params/ directory.

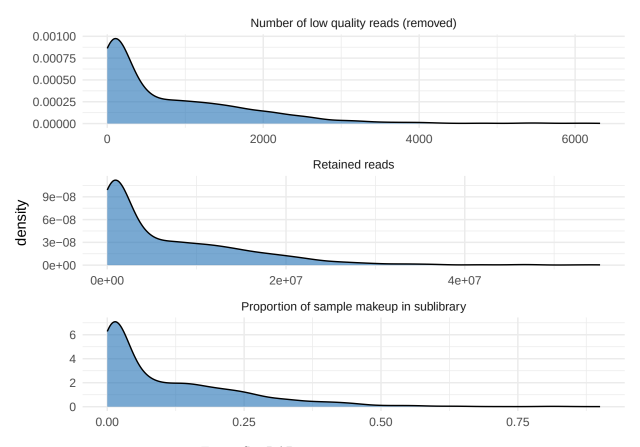


Figure S2: RAD tag processing statistics

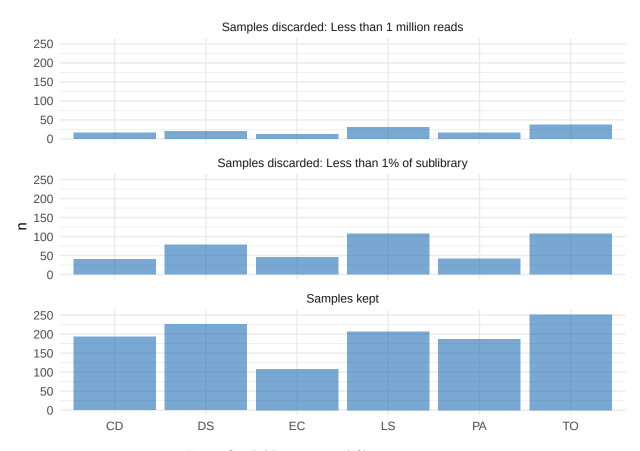


Figure S3: RAD tag manual filtering summary

Going forward, when we use the term **metapopulation**, we are referring to the collection of all samples within species among all cities where the species was present.

It is important to conduct preliminary analyses that will identify an optimal set of parameters for the dataset (see Step 5a). Following the parameter optimization, the program ustacks can be run to generate a catalog of loci.

### 5.1 Run denovo\_map.sh

Step 5a. Stack assembly will differ based on several different aspects of the dataset(such as the study species, the RAD-seq method used, and/or the quality and quantity of DNA used). So it is important to use parameters that will maximize the amount of biological data obtained from stacks.

There are three main parameters to consider when doing this:

- 1. m = controls the minimum number of raw reads required to form a stack(implemented in ustacks)
- 2. M = controls the number of mismatches between stacks to to merge them into a putative locus (implemented in ustacks)
- 3. n = controls the number of mismatches allowed between stacks to merge into the catalog (implemented in cstacks)

There are two main ways to optimize parameterization:

- 1. an iterative method were you sequentially change each parameter while keeping the other parameters fixed (described in *Paris et al. 2017*), or
- 2. an iterative method were you sequentially change the values of M and n(keeping M = n) while fixing m = 3, and then test m = 2, 4 once the optimal M = n is determined (described in *Rochette and Catchen 2017*, Catchen 2020).

We performed the second method and used the denovo\_map.sh script to run the denovo\_map.pl command to perform iterations. This script requires that we first choose a subset of samples to run the iterations on. The samples should be representative of the overall dataset; meaning they should include all populations and have similar read coverage numbers. Read coverage numbers can be assessed by looking at the descriptive statistics produced from Step 4c.

Place these samples in a text file (popmap\_test\_samples.txt) with the name of the sample and specify that all samples belong to the same population. For example, popmap\_test\_samples.txt should look like...

```
DS.BA.GA.U.1 A
DS.PX.BUF.M.5 A
DS.BO.HC4.M.1 A
```

• • •

It is important to have all representative samples treated as one population because you will assess outputs found across 80% of the individuals. The script will read this text file from the --popmap argument.

The script also requires that you specify an output directory after  $\neg o$ . This should be unique to the parameter you are testing... for example, if you are testing M=3, then you could make a subdirectory labeled stacks.M3 where all outputs from denovo\_map.sh will be placed. Otherwise, for each iteration, the outputs will be overwritten and you will lose the log from the previous iteration. The denovo\_map.sh script also requires that you direct it toward where your samples are stored, which is your directory built in Step 4b. Make sure to run the  $\neg \neg \min - samples - per - pop 0.80$  argument.

To decide which parameters to use, examine the following from each iteration:

1. the average sample coverage: This is obtained from the summary log in the ustacks section of denovo\_map.log. If samples have a coverage <10x, you will have to rethink the parameters you use here.

- 2. the number of assembled loci shared by 80% of samples: This can be found in the haplotypes.tsv by counting the number of loci: cat populations.haplotypes.tsv | grep -v ^"#" | wc -l
- 3. the number of polymorphic loci shared by 80% of samples: This can be found in populations.sumstats.tsv or by counting populations.hapstats.tsv: cat populations.hapstats.tsv | grep -v "^#" | wc -1
- 4. the number of SNPs per locus shared by 80% of samples: found in denovo\_map.log or by counting the number of SNPs in populations.sumstats.tsv: populations.sumstats.tsv | grep -v ^"#" | wc -1

The script O5a-param\_opt-figures\_script.R was used to create plots for assessing the change in shared loci across parameter iterations.

Based on this optimization step, we used the following parameters:

Table S1: Final parameter optimization values for the Stacks pipeline.

Species	M (locus mismatches)	n (catalog mismatches)	m  (minimum reads)
$\overline{\mathrm{CD}}$	8	8	3
DS	10	10	3
EC	8	8	3
LS	7	7	3
PA	5	5	3
TO	6	6	3

#### 5.2 Run ustacks

Step 5b. ustacks builds de novo loci in each individual sample. We have designed the ustacks script so that the process requires three files:

- 05-ustacks\_n.sh: the shell script that executes ustacks
- 05-ustacks\_id\_n.txt : the sample ID number
- 05-ustacks\_samples\_n.txt: the sample names that correspond to the sample IDs

The sample ID should be derived from the order\_id column(first column) on the master spreadsheet. It is unique (1-1736) across all of the samples.

The sample name is the corresponding name for each sample ID in the spreadsheet. E.g., sample ID "9" corresponds to sample name "DS.BA.DHI.U.4". Sample naming convention is species.city.site.management\_type.replicate\_plant.

05-ustacks\_n.sh should have an out\_directory (-o option) that will be used for all samples (e.g., stacks/ustacks). Files can be processed piecemeal into this directory. There should be three files for every sample in the output directory:

- <samplename>.alleles.tsv.gz
- <samplename>.snps.tsv.gz
- <samplename>.tags.tsv.gz

Multiple versions of the O5-ustacks\_n.sh script can be run in parallel (simply replace n in the three files above with the correct number).

A small number of samples (13) were discarded at this stage as the ustacks tool was unable to form any primary stacks corresponding to loci. See output/ustacks-discarded\_samples.csv.

Table S2: Summary of samples discarded at the ustacks step of the Stacks pipeline.

ustacks discarded	Retained reads	Proportion of sub-library
no	10075192	0.1608982
yes	7490510	0.1230658

### 5.3 Correct File Names

Step 5c. This step contains a script 05b-fix\_filenames.sh which uses some simple regex to fix filenames that are output in previous steps. Stacks adds an extra "1" at some point at the end of the sample name which is not meaningful. The following files:

- DS.MN.L02-DS.M.3.1.alleles.tsv.gz
- DS.MN.L03-DS.U.2.1.tags.tsv.gz
- DS.MN.L09-DS.U.1.1.snps.tsv.gz

#### become:

- $\bullet$  DS.MN.L02-DS.M.3.alleles.tsv.gz
- DS.MN.L03-DS.U.2.tags.tsv.gz
- $\bullet$  DS.MN.L09-DS.U.1.snps.tsv.gz

The script currently gives some strange log output, so it can probably be optimized/improved. The script should be run from the directory where the changes need to be made. Files that have already been fixed will not be changed.

# 5.4 Choose catalog samples/files

Step 5d. In the next step, we will choose the files we want to go into the catalog. This involves a few steps:

- 1. Create a meaningful directory name. This could be the date (e.g., stacks\_22\_01\_25).
- 2. Copy the ustacks output for all of the files you want to use in the reference from Step 5b. Remember this includes three files per sample. So if you have 20 samples you want to include in the reference catalog, you will transfer  $3 \times 20 = 60$  files into the meaningful directory name. The three files per sample should follow this convention:
- <samplename>.alleles.tsv.gz
- <samplename>.snps.tsv.gz
- <samplename>.tags.tsv.gz
- 3. Remember the meaningful directory name. You will need it in Step 6.

# 6 Stacks: Metapopulation catalog with cstacks

Files can be found in the O6\_cstacks/ directory.

cstacks builds the locus catalog from all the samples specified. The accompanying script, cstacks\_SPECIES.sh is relatively simple since it points to the directory containing all the sample files. It follows this format to point to that directory:

```
cstacks -P ~/directory ...
```

Make sure that you use the meaningful directory from Step 5c and that you have copied all the relevant files over. Otherwise this causes problems downstream. For example, you might edit the code to point to ~/scratch/stacks/stacks\_22\_01\_25.

#### cstacks -P ~/scratch/stacks/stacks\_22\_01\_25 ...

The tricky thing is ensuring enough compute memory to run the entire process successfully. There is probably space to optimize this process.

The cstacks method uses a "population map" file, which in this project is cstacks\_popmap\_SPECIES.txt. This file specifies which samples to build the catalog from and categorizes them into your 'populations', or in this case, cities using two tab-delimited columns, e.g.:

```
DS.BA.GA.U.1 Baltimore
DS.BA.GA.U.2 Baltimore
DS.BA.GA.U.3 Baltimore
DS.BA.GA.U.4 Baltimore
DS.BA.GA.U.5 Baltimore
```

Make sure the samples in this file correspond to the input files located in e.g., ~/scratch/stacks/stacks\_22\_01\_25. cstacks builds three files for use in all your samples (in this pipeline run), mirroring the sample files output byustacks:

- catalog.alleles.tsv.gz
- catalog.snps.tsv.gz
- catalog.tags.tsv.gz

Table S3: Subset of samples used in SNP catalog creation.

Sample	Species	City
DS.BA.PIK.U.1	DS	BA
DS.BA.GA.U.4	DS	BA
DS.BA.LH-1.M.4	DS	BA
DS.BA.LH-3.M.1	DS	BA
DS.BA.WB.U.2	DS	BA
DS.BA.LL-4.M.5	DS	BA
DS.BA.LH-2.M.5	DS	BA
DS.BA.TRC.U.3	DS	BA
DS.BA.W3.M.2	DS	BA
DS.BA.RG-1.M.1	DS	BA
DS.BA.LL-3.M.3	DS	BA
DS.BA.RG-2.M.4	DS	BA
DS.BO.HC1.M.3	DS	ВО
DS.BO.HC4.M.5	DS	ВО
DS.BO.LC1.M.3	DS	ВО
DS.BO.LC2.M.2	DS	ВО
DS.BO.LC3.M.5	DS	ВО
DS.BO.WL1.M.2	DS	ВО
DS.BO.WL2.M.1	DS	ВО
DS.BO.WL3.M.5	DS	ВО
DS.BO.I4.U.1	DS	ВО
DS.BO.R1.U.4	DS	ВО
DS.BO.R2.U.2	DS	ВО
DS.BO.R4.U.4	DS	ВО
DS.MN.L05-DS.M.3	DS	MN
DS.MN.L09-DS.M.3	DS	MN
DS.MN.L11-DS.M.1	DS	MN
DS.MN.L02-DS.U.1	DS	MN
DS.MN.L02-DS.M.4	DS	MN

Sample	Species	City
DS.MN.L03-DS.U.3	DS	MN
DS.MN.L04-DS.U.5	$\overline{\mathrm{DS}}$	MN
DS.MN.L06-DS.U.3	$\overline{\mathrm{DS}}$	MN
DS.MN.L07-DS.U.3	$\overline{\mathrm{DS}}$	MN
DS.MN.L09-DS.U.3	DS	MN
DS.MN.L11-DS.U.1	DS	MN
DS.MN.L11-DS.U.5	$\overline{\mathrm{DS}}$	MN
DS.PX.BUF.M.1	DS	PX
DS.PX.PIE.M.2	DS	PX
DS.PX.ALA.M.1	DS	PX
DS.PX.MTN.M.6	DS	PX
DS.PX.LAP.M.3	DS	PX
DS.PX.NUE.M.4	DS	PX
DS.PX.WES.M.2	DS	PX
DS.PX.DF1.M.1	DS	PX
DS.PX.ENC.M.1	DS	PX
DS.PX.DOW.M.1	DS	PX
DS.PX.DOW.M.4	DS	PX
DS.PX.DF2.M.3	$\overline{\mathrm{DS}}$	PX
CD.BA.LA.U.2	$\overline{\mathrm{CD}}$	BA
CD.BA.TRC.U.3	$\overline{\mathrm{CD}}$	BA
CD.BA.WGP.M.2	$\overline{\mathrm{CD}}$	BA
CD.BA.LH-2.M.2	CD	BA
CD.BA.LL-4.M.1	CD	BA
CD.BA.PIK.U.2	CD	BA
CD.BA.WB.U.2	CD	BA
CD.BA.CP.U.4	CD	BA
CD.BA.FH.U.1	CD	BA
CD.BA.PSP.M.4	CD	BA
CD.BA.AA.U.4	CD	BA
CD.BA.RG-1.M.2	CD	BA
CD.BA.W3.M.3	CD	BA
CD.BA.GA.U.3	CD	BA
CD.BA.WBO.U.5	$^{\mathrm{CD}}$	BA
CD.LA.WHI.M.3	CD	LA
CD.LA.SEP.M.3	CD	LA
CD.LA.SEP.M.4	CD	LA
CD.LA.ROS.M.5	CD	LA
CD.LA.MR2.M.2	CD	LA
CD.LA.ALL.M.2	CD	LA
CD.LA.ALL.M.5	CD	LA
CD.LA.VAL.M.5	CD	LA
CD.LA.HAR.M.4	CD	LA
CD.LA.LUB.M.3	CD	LA
CD.LA.GLO.M.4 CD.LA.ZOO.M.3	CD CD	LA
CD.LA.NWH.M.5	CD	LA LA
CD.LA.KIN.M.3	CD	LA
CD.LA.KIN.M.5	CD	LA
CD.PX.CAM.U.5	CD	PX
CD.PX.MON.U.5	CD	PX
CD.PX.PKW.U.5	CD	PX
CD.1 11.11 11 W.O.0	CD	1 11

Sample	Species	City
CD.PX.LAP.M.4	CD	PX
CD.PX.NES.U.4	CD	PX
CD.PX.PAL.M.3	CD	PX
CD.PX.ASU.M.1	CD	PX
CD.PX.NUE.M.5	CD	PX
CD.PX.WES.M.3	CD	PX
CD.PX.MAN.M.4	CD	PX
CD.PX.CLA.M.3	CD	PX
CD.PX.DF1.M.5	CD	PX
CD.PX.COY.M.5	CD	PX
CD.PX.RPC.M.3	CD	PX
CD.PX.ENC.M.2	CD	PX
EC.BA.LH-2.M.2	EC	BA
EC.BA.WBO.U.4	EC	BA
EC.BA.WB.U.5	EC	BA
EC.BA.FH.U.3	EC	BA
EC.BA.CP.U.2	EC	BA
EC.BA.TRC.U.3	EC	BA
EC.BA.LL-4.M.4	EC	BA
EC.BA.WB.U.1	EC	BA
EC.BA.PIK.U.5	EC	BA
EC.BA.PSP.M.4	EC	BA
EC.BA.GA.U.2	EC	BA
EC.BA.LL-3.M.3	EC	BA
EC.BA.ML.U.1	EC	BA
EC.BA.TRC.U.5	EC	BA
EC.BA.ML.U.3	EC	BA
EC.LA.SGB.U.2	EC	LA
EC.LA.SGB.U.5	EC	LA
EC.LA.DUR.U.2	EC	LA
EC.LA.HOW.U.2	EC	LA
EC.LA.SAN.U.2	EC	LA
EC.LA.VER.U.1	EC	LA
EC.LA.VER.U.4	EC	LA
EC.LA.VB2.U.4	EC	LA
EC.LA.AC2.U.2	EC	LA
EC.LA.AC1.U.1	EC	LA
EC.LA.VB1.U.1	EC	LA
EC.LA.VB1.U.3	EC	LA
EC.LA.SGR.U.4	EC	LA
EC.LA.SGR.U.5	EC	LA
EC.LA.HOW.U.3	EC	LA
EC.PX.BUF.M.1	EC	PX
EC.PX.BUF.M.3	EC	PX
EC.PX.ALA.M.3	EC	PX
EC.PX.MTN.M.2	EC	PX
EC.PX.WES.M.1	EC	PX
EC.PX.WES.M.2	EC	PX
EC.PX.MAN.M.1	EC	PX
EC.PX.CLA.M.1	EC	PX
EC.PX.PSC.M.1	EC	PX
EC.PX.DF1.M.1	EC	PX

Sample	Species	City
EC.PX.DOW.M.1	EC	PX
EC.PX.DOW.M.2	EC	PX
EC.PX.COY.M.2	EC	PX
EC.PX.COY.M.3	EC	PX
EC.PX.ALA.M.5	EC	PX
LS.BA.WB.U.1	LS	BA
LS.BA.WB.U.2	LS	BA
LS.BA.DHI.U.2	LS	BA
LS.BA.GA.U.1	LS	BA
LS.BA.PIK.U.3	LS	BA
LS.BA.PIK.U.5	LS	BA
LS.BA.CP.U.2	LS	BA
LS.BA.ML.U.2	LS	BA
LS.BA.WBO.U.3	LS	BA
LS.BO.WL3.M.4	LS	ВО
LS.BO.II.U.1	LS	ВО
LS.BO.II.U.1	LS	ВО
	LS LS	
LS.BO.WL2.M.2		ВО
LS.BO.R1.U.2	LS	ВО
LS.BO.R2.U.4	LS	ВО
LS.BO.R3.U.3	LS	ВО
LS.BO.HC4.M.3	LS	ВО
LS.BO.LC4.M.2	LS	ВО
LS.LA.VET.M.4	LS	LA
LS.LA.SSV.M.1	LS	LA
LS.LA.NAV.M.4	LS	LA
LS.LA.SHO.M.2	LS	LA
LS.LA.WES.M.3	LS	LA
LS.LA.GLO.M.3	LS	LA
LS.LA.HOW.U.5	LS	LA
LS.LA.SAN.U.2	LS	LA
LS.LA.ARR.U.2	LS	LA
LS.MN.L06-LS.U.2	LS	MN
LS.MN.L06-LS.U.5	LS	MN
LS.MN.L07-LS.U.4	LS	MN
LS.MN.L08-LS.U.5	LS	MN
LS.MN.L09-LS.U.3	LS	MN
LS.MN.L01-LS.M.4	LS	MN
LS.MN.L01-LS.U.3	LS	MN
LS.MN.L02-LS.U.1	LS	MN
LS.MN.L05-LS.U.2	LS	MN
LS.PX.MON.U.2	LS	PX
LS.PX.PKW.U.5	LS	PX
LS.PX.PIE.M.4	LS	PX
LS.PX.ALA.M.3	LS	PX
LS.PX.PAL.M.3	LS	PX
LS.PX.MAN.M.2	LS	PX
LS.PX.NUE.M.1	LS	PX
LS.PX.ENC.M.4	LS	PX
LS.PX.COY.M.3	LS	PX
PA.BA.PIK.U.1	PA	ВА
PA.BA.LH-3.M.2	PA	BA
	<b></b>	

Sample	Species	City
PA.BA.LH-3.M.3	PA	BA
PA.BA.WB.U.1	PA	BA
PA.BA.AA.U.1	PA	BA
PA.BA.WGP.M.3	PA	BA
PA.BA.LL-4.M.3	PA	BA
PA.BA.LA.U.2	PA	BA
PA.BA.LH-2.M.2	PA	BA
PA.BA.W3.M.3	PA	BA
PA.BA.RG-1.M.2	PA	BA
PA.BA.LL-3.M.5	PA	BA
PA.BO.I2.U.3	PA	ВО
PA.BO.HC1.M.4	PA	ВО
PA.BO.R3.U.2	PA	ВО
PA.BO.HC4.M.5	PA	ВО
PA.BO.R4.U.2	PA	ВО
PA.BO.WL2.M.5	PA	ВО
PA.BO.WL4.M.4	PA	ВО
PA.BO.LC4.M.4	PA	ВО
PA.BO.HC2.M.1	PA	ВО
PA.BO.R1.U.2	PA	ВО
PA.BO.WL1.M.1	PA	ВО
PA.BO.II.U.5	PA	ВО
PA.LA.ALL.M.5	PA	LA
PA.LA.SEP.M.1	PA	LA
PA.LA.SEP.M.5	PA	LA
PA.LA.WHI.M.2	PA	LA
PA.LA.ROS.M.5	PA	LA
PA.LA.LUB.M.2	PA	LA
PA.LA.GLO.M.2	PA	LA
PA.LA.ZOO.M.4	PA	LA
PA.LA.ZOO.M.4 PA.LA.ZOO.M.5	PA	LA
PA.LA.NWH.M.2	PA	LA LA
PA.LA.KIN.M.4	PA	LA
PA.LA.POP.M.4	PA	LA
PA.PX.BUF.M.3	PA PA	PX
PA.PX.PIE.M.4		
	PA	PX
PA.PX.LAP.M.5	PA DA	PX
PA.PX.ALA.M.1 PA.PX.PAP.M.2	PA	PX
PA.PX.PAP.M.2 PA.PX.PAP.M.5	PA DA	PX
	PA	PX
PA.PX.DF1.M.2	PA	PX
PA.PX.RPP.U.3	PA	PX
PA.PX.ENC.M.4	PA DA	PX
PA.PX.ENC.M.5	PA	PX
PA.PX.COY.M.1	PA	PX
PA.PX.BUF.M.2	PA	PX
TO.BA.WBO.U.4 TO.BA.CP.U.1	TO	BA
	TO	BA
TO.BA.FH.U.1	TO	BA
TO.BA.LH-3.M.4	TO	BA
TO.BA.WGP.M.3	TO	BA
TO.BA.GA.U.4	ТО	BA

Sample	Species	City
TO.BA.PIK.U.4	ТО	BA
TO.BA.PSP.M.1	TO	BA
TO.BA.RG-2.M.2	TO	BA
TO.BO.HC1.M.4	TO	ВО
TO.BO.HC2.M.5	TO	ВО
TO.BO.HC3.M.1	TO	ВО
TO.BO.HC4.M.5	TO	ВО
TO.BO.LC1.M.1	TO	ВО
TO.BO.LC2.M.5	TO	ВО
TO.BO.LC3.M.1	TO	ВО
TO.BO.WL2.M.1	TO	ВО
TO.BO.I2.U.3	TO	ВО
TO.LA.WHI.M.5	TO	LA
TO.LA.HAR.M.4	TO	LA
TO.LA.MR1.M.1	TO	LA
TO.LA.GLO.M.5	TO	LA
TO.LA.ZOO.M.1	TO	LA
TO.LA.NWH.M.4	TO	LA
TO.LA.VNS.M.2	TO	LA
TO.LA.PEP.M.5	TO	LA
TO.LA.COM.M.4	TO	LA
TO.MN.L11-TO.M.3	TO	MN
TO.MN.L02-TO.U.1	TO	MN
TO.MN.L04-TO.U.1	TO	MN
TO.MN.L06-TO.U.2	TO	MN
TO.MN.L08-TO.U.5	TO	MN
TO.MN.L09-TO.U.2	TO	MN
TO.MN.L11-TO.U.3	TO	MN
TO.MN.L05-TO.M.5	TO	MN
TO.MN.L08-TO.M.5	TO	MN
TO.PX.BUF.M.1	TO	PX
TO.PX.ALA.M.2	TO	PX
TO.PX.LAP.M.4	TO	PX
TO.PX.WES.M.1	TO	PX
TO.PX.CLA.M.1	TO	PX
TO.PX.DF1.M.1	TO	PX
TO.PX.DF2.M.1	TO	PX
TO.PX.COY.M.1	TO	PX
TO.PX.COY.M.6	ТО	PX

# 7 Stacks: Metapopulation locus matching with sstacks

Files can be found in the O7\_sstacks/ directory.

All samples in the population (or all samples you want to include in the analysis) are matched against the catalog produced in cstacks with sstacks, run in script stacks\_SPECIES.sh and stacks\_SPECIES\_additional.sh. It runs off of the samples based in the output directory and the listed samples in sstacks\_samples\_SPECIES.txt and sstacks\_samples\_SPECIES\_additional.txt (respectively), so make sure all your files (sample and catalog, etc.) are there and match. sstacks\_samples\_SPECIES.txt takes the form:

DS.BA.GA.U.1

```
DS.BA.GA.U.2
DS.BA.GA.U.3
DS.BA.GA.U.4
DS.BA.GA.U.5
```

There should be a new file produced at this step for every sample in the output directory:

• <samplename>.matches.tsv.gz

A small number of samples generated very few matches to the catalog (such as only 4 loci matching, obviously not enough to draw any conclusions) and therefore aren't used in the next step. See output/sstacks-discarded samples.csv.

# 8 Genotype probabilities with polyRAD

Files can be found in the O8\_polyRAD/ directory.

# 8.1 Make RADdata object

We used the polyRAD package to call genotypes because many of our species are polyploid or have historical genome duplication. PolyRAD takes the catalog output (catalog.alleles.tsv.gz) and accompanying matches to the catalog (e.g., CD.BA.AA.U.1.matches.tsv.gz) to create genotype likelihoods for species with diploidy and/or polyploidy.

We used the catalog and match files to create a RADdata object class in R for each species. We ran this on the Rockfish compute cluster at Johns Hopkins University, with the make\_polyRAD\_<spp>.R script doing the brunt of the work. The R script was wrapped by polyrad\_make\_<spp>.sh to submit the script to the SLURM scheduler.

Relevant Parameters:

- min.ind.with.reads was set to 20% of samples. This means we discarded any loci not found in at least 20% of samples for each species.
- min.ind.with.minor.allele was set to 2. This means a locus must have at least this many samples with reads for the minor allele in order to be retained.

#### Requires:

- popmap\_<spp>\_polyrad.txt, a list of samples and population
- output from sstacks

#### Outputs:

• spp>\_polyRADdata.rds, RDS object (the RADdata object)

# 8.2 Calculate overdispersion

Next, we calculated overdispersion using the polyRAD\_overdispersion\_<spp>.R script, wrapped by polyrad\_overd\_<spp>.sh to submit the script to the SLURM scheduler.

# Requires:

- $popmap_{spp}-polyrad.txt$ , a list of samples and population
- spp>\_polyRADdata.rds, RDS object (the RADdata object) output from the previous step

#### Outputs:

• spp>\_overdispersion.rds, RDS object (the overdispersion test output)

# 8.3 Estimate genotypes

Next, we calculated filtered loci based on the expected Hind/He statistic and estimated population structure/genotypes using the polyRAD\_filter\_<spp>.R script, wrapped by polyrad\_filt\_<spp>.sh to submit the script to the SLURM scheduler.

We used the table in this tutorial, which estimated an inbreeding based on the ploidy, optimal overdispersion value, and mean Hind/He. These values are hardcoded in polyRAD\_filter\_<spp>.R.

#### Requires:

- popmap\_<spp>\_polyrad.txt, a list of samples and population
- spp>\_polyRADdata.rds, RDS object (the RADdata object) output from the previous step
- spp>\_overdispersion.rds, RDS object (the overdispersion test output) output from the previous
  step

#### Outputs:

- <pp>\_filtered\_RADdata.rds, RDS object (RADdata object filtered for appropriate Hind/He)
- spp>\_IteratePopStructPCA.csv, data output from the genotype estimate PCA, suitable for plotting
- spp>\_estimatedgeno\_RADdata.rds, RDS object (RADdata object with genotype estimates)

# 8.4 Final filter and file cleanup

The output <spp>\_estimatedgeno\_RADdata.rds needs to be converted to genind and structure format for further analysis and steps. There is a little cleanup involved so the population information is retained. For example, Structure needs the population identity to be an integer, not a string. This set of functions can be run on a laptop.

At this stage, we also visually assessed the  $H_{ind}/H_e$  statistic versus the locus depth (see check\_coverage inside the convert\_genomics.R script). We removed the following samples from further analysis:

Table S4: Subset of samples discarded after genotype estimation using polyRAD.

Sample
CD.BA.PSP.M.1
CD.BA.DHI.U.2
CD.BA.DHI.U.3
CD.BA.RG-1.M.5
CD.BA.RG-1.M.4
DS.BO.WL1.M.4
DS.BO.I1.U.3
EC.BO.R4.U.1
LS.BO.HC2.M.5
LS.BO.LC4.M.3
LS.BO.R2.U.4
LS.BO.R2.U.1
PA.BA.LH-3.M.4
PA.BA.AA.U.3
PA.BA.AA.U.4
PA.PX.RPP.U.2
PA.BO.HC2.M.4
PA.PX.RPP.U.1
TO.BA.TRC.U.1
TO.BA.TRC.U.3
TO.BO.R4.U.1

Sample TO.BA.TRC.U.2 TO.BO.R4.U.2 TO.BO.R2.U.2

```
source("08_polyRAD/convert_genomics.R")
convert_all()
```

# 9 Structure Analysis

Files, inlcuding model parameters, can be found in the O9\_structure/ directory.

Structure documentation can be found here.

# 9.1 Running Structure on polyRAD output

polyRAD outputs genotype probabilities in a format suitable for Structure. These files were named as:

```
CD_estimatedgeno.structure
DS_estimatedgeno.structure
EC_estimatedgeno.structure
LS_estimatedgeno.structure
PA_estimatedgeno.structure
TO_estimatedgeno.structure
```

We ran all species using a naive approach (not using prior information) with K = 1, 2, 3, 4, 5 (MAXPOPS argument). To search for the most appropriate K, We ran Structure through 5 replicate runs for each combination of species and K, with 10000 iterations discarded as burn-in and retained 20000 iterations. These runs created files that look like:

```
structure_out_CD1_naive_f
                                  // K = 1, rep 1
                                  // K = 1, rep 2
structure_out_CD1_naive_rep2_f
structure_out_CD1_naive_rep3_f
                                  // K = 1, rep 3
                                  // K = 1, rep 4
structure_out_CD1_naive_rep4_f
                                  // K = 1, rep 5
structure_out_CD1_naive_rep5_f
structure_out_CD2_naive_f
                                  // K = 2, rep 1
structure_out_CD2_naive_rep2_f
                                  // K = 2, rep 2
structure_out_CD2_naive_rep3_f
                                  // K = 2, rep 3
```

Within each species, we compressed the result files for all K and reps and submitted to Structure Harvester to choose the optimal K using the Delta-K method (see <a href="https://link.springer.com/article/10.1007/s12686-011-9548-7">https://link.springer.com/article/10.1007/s12686-011-9548-7</a>). Once the optimal K was selected per species, we re-ran Structure using a greater number of iterations (100000) for final output and plotting.

# 10 Conceptual Figure

We made a conceptual figure to help readers understand PCA patterns we might expect.

```
source("R/10-Fig1-conceptual_fig.R")
make_fig1()
```

# 11 NLCD Data and Site Plots

NLCD is used in Fig. 2 and IBE analysis, described below.

From the USGS:

The U.S. Geological Survey (USGS), in partnership with several federal agencies, has developed and released four National Land Cover Database (NLCD) products over the past two decades: NLCD 1992, 2001, 2006, and 2011. This one is for data from 2016 and describes urban imperviousness.

https://www.mrlc.gov/data/type/urban-imperviousness

NLCD imperviousness products represent urban impervious surfaces as a percentage of developed surface over every 30-meter pixel in the United States. NLCD 2016 updates all previously released versions of impervious products for CONUS (NLCD 2001, NLCD 2006, NLCD 2011) along with a new date of impervious surface for 2016. New for NLCD 2016 is an impervious surface descriptor layer. This descriptor layer identifies types of roads, core urban areas, and energy production sites for each impervious pixel to allow deeper analysis of developed features.

# 11.1 Preparing NLCD Data

First, we trimmed the large data. This makes a smaller .rds file for each city.

```
source("R/10-trim_NLCD_spatial_data.R")
create_spatial_rds_files()
create_spatial_rds_files(spp = "CD")
```

# 11.2 Climate normals

We obtained climate normals for plotting from https://www.ncei.noaa.gov/access/us-climate-normals. We used the latest 30-year period (1991-2020): Most recent standard climatological period (2021 release); which is recommended for most purposes.

### 11.3 Maps of sampling locations

Next, we made plots for each city's sampling locations. Note that these only include sites that had viable polymorphic loci.

```
source("R/10-Fig2-plot_map_of_samples.R")
# Plot Fig 2
make_all_urban_site_plots_with_clim_normals()
```

# 12 Principal components analysis & plots

The following creates PCA plots from polyRAD data.

```
source("R/11-plot_pca.R")

# Plot Fig 4
make_pca_city_and_pctimp_all()
```

In addition to coloring points by city, we also colored points by NLCD Urban Cover %. Note that in the pdf version of this document, the figure might appear on the next pages.

```
make_pca_pctimp_only_all()
```

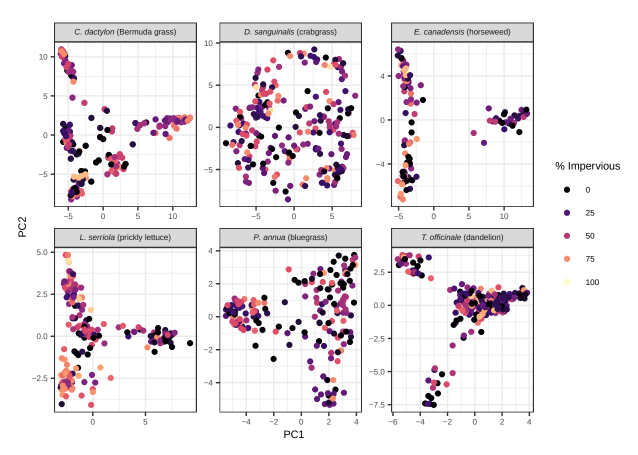


Figure S4: PCA colored by % Impervious surface.

# 13 Genetic structure using Structure and sNMF

## 13.1 Structure optimal K

Within each species, we compressed the result files for all K and reps and submitted to Structure Harvester to choose the optimal K using the Delta-K method (see <a href="https://link.springer.com/article/10.1007/s12686-011-9548-7">https://link.springer.com/article/10.1007/s12686-011-9548-7</a>).

The results were:

```
CD: K=3 DS: K=3 EC: K=2 LS: K=3 PA: K=4 TO: K=3

# This file contains output from various K from Structure..

read_csv("output/structure/structure_k_Pr.csv")
```

The code below generates plots of various K (e.g.,  $K=\{1-5\}$ ) vs likelihood, but did not end up being used in the manuscript.

```
source("R/12-structure_k.R")
```

# 13.2 Plotting Structure output

The code below generates plots for Structure results.

```
source("R/12-plot_structure.R")
make_structure_multi_plot()
```

#### 13.3 Validation of Structure results with sNMF

We ran sNMF as an alternative to Structure to validate the results. We coerced all polyploid data to diploid data to make the file types compatible with the sNMF function in R. The snmf() function computes an entropy criterion that evaluates the quality of fit of the statistical model to the data by using a cross-validation technique. We plotted the cross-entropy criterion for K=[2:10] for all species. Using the best K, we then selected the best of 10 runs in each K using the which.min() function.

```
source("R/12-sNMF.R")
```

The following runs sNMF and generates the figure. Note that in the pdf version of this document, the figure might appear on the next pages.

```
do_all_sNMF()
```

### 14 Genetic Differentiation

# 14.1 Among city - Jost's D, $G_{ST}$ , $F_{ST}$

We used polyrad::calcPopDiff() to calculate continental population statistics for each species.

```
source("R/13-calc_popdiff_stats.R")

do_all_continental_stats()

# CD as an example
read.csv("output/population_stats/popdiff_stats_CD.csv")

## X statistic value
```

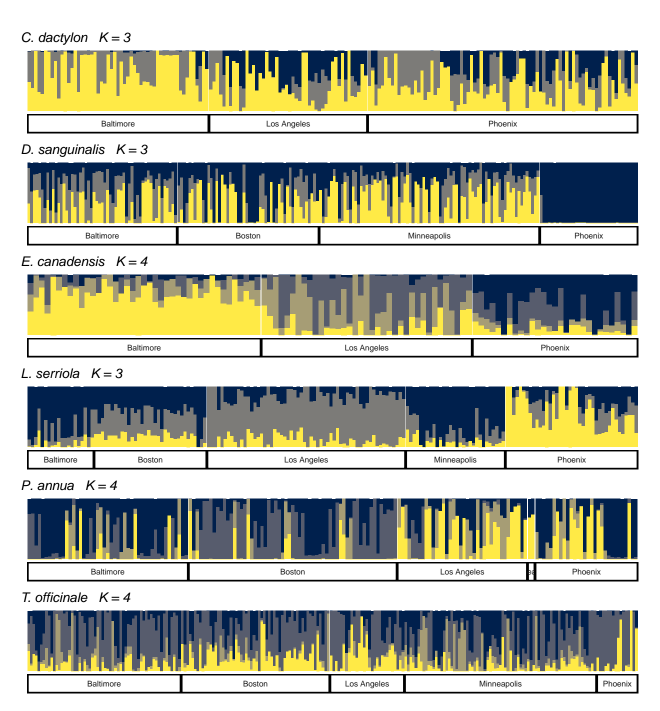


Figure S5: Ancestry coefficients obtained using snmf(). As with the Structure analysis, E. canadensis (horseweed) and L. serriola (prickly lettuce) appear to have the most population structure. D. sanguinalis, E. canadensis, and L. serriola from Phoenix appear unique. In general, sNMF produced larger K for most species, which will create more sensitivity to admixture.

# 14.2 Among city - pairwise $\rho$

We used GenoDive v. 3.0.6 to calculate pairise  $\rho$  (rho) among cities within species. Note that there is a p-value correction for testing multiple cities (species are treated as independent, however).

This can be run in GenoDive by selecting Pairwise Differentiation from the Analysis menu and selecting the "rho" statistic from the dropdown.

We used the following script to clean up the results.

```
source("R/13-rho.R")
compile_rho_table()
```

Table S5: Rho statistics for pairwise comparison between cities.

Species	City1	City2	rho	p-value	adjusted p-value
CD	PX	BA	0.050	0.001	0.001
CD	LA	BA	0.046	0.001	0.001
CD	PX	LA	0.015	0.001	0.001
DS	MN	BA	0.031	0.001	0.0015
DS	ВО	BA	0.018	0.001	0.0015
DS	PX	BA	0.012	0.001	0.0015
DS	MN	ВО	0.007	0.001	0.0015
DS	PX	ВО	-0.002	0.875	0.955
DS	PX	MN	-0.002	0.955	0.955
EC	PX	BA	0.098	0.001	0.001
EC	PX	LA	0.087	0.001	0.001
EC	LA	BA	0.038	0.001	0.001
LS	PX	BA	0.077	0.001	0.0011
LS	PX	MN	0.069	0.001	0.0011
LS	PX	LA	0.061	0.001	0.0011
LS	PX	ВО	0.056	0.001	0.0011
LS	MN	LA	0.039	0.001	0.0011
LS	LA	BA	0.038	0.001	0.0011
LS	ВО	BA	0.032	0.001	0.0011
LS	MN	ВО	0.021	0.001	0.0011
LS	LA	ВО	0.010	0.001	0.0011
LS	MN	BA	0.009	0.002	0.002
PA	PX	ВО	0.028	0.001	0.0015
PA	LA	ВО	0.024	0.001	$\overline{0.0015}$
PA	PX	BA	0.015	0.001	0.0015
PA	LA	BA	0.011	0.001	0.0015
PA	ВО	BA	0.008	0.002	0.0024
PA	PX	LA	-0.002	0.972	0.972
TO	PX	BA	0.023	0.001	0.0014
TO	PX	MN	0.015	0.001	$\overline{0.0014}$
TO	PX	ВО	0.013	0.002	0.0025
TO	LA	BA	0.011	0.001	$\overline{0.0014}$
TO	LA	ВО	0.009	0.001	$\overline{0.0014}$
TO	MN	LA	0.009	0.001	0.0014
TO	PX	LA	0.009	0.027	$\overline{0.03}$
TO	ВО	BA	0.008	0.001	$\overline{0.0014}$
TO	MN	ВО	0.008	0.001	$\overline{0.0014}$
ТО	MN	BA	0.001	0.098	0.098

# 14.3 Within city - allelic richness

We used GenoDive v. 3.0.6 to calculate several additional statistics.

This can be run in GenoDive by selecting Analysis > Genetic Diversity, and selecting "Calculate indices separately for every population" and selecting "Correct for unknown dosage of alleles" for the polyploid species.

- Num: Number of alleles
- Eff num: Effective number of alleles
- Ho: Observed Heterozygosity
- Hs: Heterozygosity within populations
- Gis: Inbreeding coefficient

```
head(read.csv("output/population_stats/genodive_genetic_diversity.csv"))
```

```
##
                  Num Eff_num
                                 Но
                                        Hs
                                             Gis
     spp city
## 1
      CD
               8.242
                        4.530 0.627 0.748 0.163
                        6.470 0.683 0.827 0.174
           LA 12.664
## 2
      CD
## 3
      CD
           PX 14.192
                        6.978 0.672 0.830 0.191
## 4
      DS
           BA 10.611
                        5.069 0.637 0.768 0.171
      DS
           BO 10.588
                        5.230 0.612 0.778 0.214
## 5
      DS
           MN 11.696
                        5.233 0.623 0.771 0.192
## 6
```

# 14.4 Within city - $F_{IS}$ (homozygosity within population)

We used GenoDive v. 3.0.6 to calculate  $F_{IS}$ . This gives a good estimate of whether there are more homozygotes than expected (positive number) or more heterozygotes than expected (negative number). Notably, GenoDive accommodates polyploids and reduces the bias on  $F_{IS}$  by performing a permutation test. By default, there are 999 permutations.

This can be run in GenoDive by selecting Analysis > Hardy-Weinberg > Heterozygosity-based (Nei) method.

```
head(read.csv("output/population_stats/genodive_output_Fis.csv"))
```

```
##
     Species Population n
                               Fis
## 1
          CD
                      BA 55 0.166
## 2
          CD
                      LA 48 0.186
## 3
          CD
                      PX 82 0.200
## 4
          CD
                 Overall NA 0.187
## 5
          DS
                      BA 55 0.208
## 6
          DS
                      BO 52 0.252
```

# 14.5 Within city - $\bar{r}_d$ - Linkage disequilibrium

We used poppr::ia() to calculate the standardized index of association of loci in the dataset ( $\bar{r}_d$  or rbard) We use the standardized index of association to avoid the influence of different sample sizes, as described by Agapow and Burt 2001.

When p.rD is small (<0.05) and rbarD is (relatively) higher, that is a sign that the population could be in linkage disequilibrium.

An interesting note from the documentation:

It has been widely used as a tool to detect clonal reproduction within populations. Populations whose members are undergoing sexual reproduction, whether it be selfing or out-crossing, will produce gametes via meiosis, and thus have a chance to shuffle alleles in the next generation. Populations whose members are undergoing clonal reproduction, however, generally do so via mitosis. This means that the most likely mechanism for a change in genotype is via mutation.

The rate of mutation varies from species to species, but it is rarely sufficiently high to approximate a random shuffling of alleles. The index of association is a calculation based on the ratio of the variance of the raw number of differences between individuals and the sum of those variances over each locus. You can also think of it as the observed variance over the expected variance.

There is a nice description here.

```
source("R/13-rbarD.R")
calc_rbarD()
head(read.csv("output/population_stats/rbarD.csv"))
     spp city n
                       Ia p.Ia
                                    rbarD p.rD
## 1
     CD
           BA 55 664.7655 0.001 0.2950534 0.001
## 2
     CD
           LA 48 470.5913 0.001 0.2070064 0.001
## 3 CD
           PX 82 634.5787 0.001 0.2792566 0.001
## 4
     DS
           BA 55 557.7334 0.001 0.2123881 0.001
## 5
     DS
           BO 52 896.0906 0.001 0.3398873 0.001
           MN 81 578.2494 0.001 0.2192197 0.001
## 6
     DS
```

## 14.6 Within city - private alleles

We used a custom script to calculate % private alleles as the percentage of total alleles unique to a particular city.

```
source("R/13-private_alleles.R")
get_all_private_alleles()
```

# 15 SessionInfo()

[9] tidyverse\_2.0.0

## [13] sp\_2.1-4

## [17] polyRAD\_2.0.0

##

```
sessionInfo()
## R version 4.4.2 (2024-10-31)
## Platform: aarch64-apple-darwin20
## Running under: macOS Sonoma 14.4.1
## Matrix products: default
           /Library/Frameworks/R.framework/Versions/4.4-arm64/Resources/lib/libRblas.0.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/4.4-arm64/Resources/lib/libRlapack.dylib;
                                                                                                LAPACK v
##
## [1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
##
## time zone: America/New_York
## tzcode source: internal
##
## attached base packages:
## [1] stats
                 graphics grDevices utils
                                               datasets methods
                                                                    base
##
## other attached packages:
  [1] polysat_1.7-7
                          LEA_3.16.0
                                            ggh4x_0.2.8
                                                               here_1.0.1
  [5] lubridate_1.9.3
                          forcats_1.0.0
                                            purrr_1.0.2
                                                               tibble_3.2.1
```

stringr\_1.5.1

magrittr\_2.0.3

viridisLite\_0.4.2 raster\_3.6-26

readr\_2.1.5

tidyr\_1.3.1

viridis\_0.6.5

cowplot\_1.2.0

dplyr\_1.1.4

```
## [21] ggplot2_4.0.0
##
## loaded via a namespace (and not attached):
   [1] fastmatch_1.1-4
                           gtable_0.3.6
                                                                   lattice_0.22-6
                                               xfun_0.52
##
    [5] tzdb_0.4.0
                           vctrs_0.6.5
                                               tools 4.4.2
                                                                   generics_0.1.3
  [9] parallel 4.4.2
                           fansi 1.0.6
                                               highr 0.11
                                                                   pkgconfig 2.0.3
##
## [13] RColorBrewer 1.1-3 S7 0.2.0
                                               lifecycle 1.0.4
                                                                   compiler 4.4.2
## [17] farver 2.1.2
                           textshaping_0.4.0
                                               tinytex_0.51
                                                                   terra 1.8-60
##
  [21] codetools 0.2-20
                           htmltools 0.5.8.1
                                               yaml_2.3.10
                                                                   pillar_1.9.0
  [25] crayon_1.5.2
                           commonmark_1.9.1
                                               tidyselect_1.2.1
                                                                   digest_0.6.35
  [29] stringi_1.8.4
                           labeling_0.4.3
                                               rprojroot_2.0.4
                                                                   fastmap_1.2.0
                                               dichromat_2.0-0.1
       grid_4.4.2
                           cli_3.6.3
                                                                   utf8_1.2.4
  [33]
  [37]
       withr_3.0.2
                           scales_1.4.0
                                               bit64_4.0.5
                                                                   timechange_0.3.0
  [41] rmarkdown_2.27
                           ggtext_0.1.2
                                               bit_4.0.5
                                                                   gridExtra_2.3
## [45] ragg_1.3.2
                           hms_1.1.3
                                               kableExtra_1.4.0
                                                                   evaluate_1.0.5
  [49] knitr_1.47
                           markdown_1.13
                                               rlang_1.1.4
                                                                   gridtext_0.1.5
  [53] Rcpp_1.0.12
                           glue_1.8.0
                                               xm12_1.3.6
                                                                   formatR_1.14
  [57] syglite 2.1.3
                           rstudioapi_0.16.0
                                               vroom_1.6.5
                                                                   R6_2.5.1
  [61] systemfonts_1.1.0
```

# **Appendix**

## 15.1 File Organization

All data files for the Macrosystems project are permanently stored under Meghan Avolio's group resources in the Johns Hopkins University Rockfish computing cluster. Files are stored under the 'data' directory under the following subdirectories:

- 01-raw\_data: This folder contains the raw, unprocessed data files that were obtained directly from the sequencing server. There are eight fastq.gz files per sublibrary that correspond to the four sequencing lanes for each read direction.
- 02-concatenated\_data: This folder contains the concatenated, unprocessed files for each sublibrary (i.e., the files containing the sequences for each lane were combined to create one file per read direction).
- 03-pcr\_filtered\_data: Here, you will find the resulting data files from the clone\_filter program,
  where pcr replicates/clones have been removed from the raw sequences. There are two fq.gz files per
  sublibrary.
- 04-process\_radtags: This folder contains various subdirectories that correspond to the process\_radtags program that demultiplexes and cleans the data. The demux\_txt\_files folder contains the .txt files used to identify barcodes and separate out the individual samples from each sublibrary. The resulting data files from the process-radtags program are separated by individual and can be found in the relevant species folder(i.e., CD, DS, EC, LS, PA, TE, TO). Each individual sample has four data files; sampleID.1.fq.gz and sampleID.2.fq.gz correspond to the forward and reverse reads for each sample and sampleID.rem,1/2.fq.gz contain the remainder reads that were cleaned and removed from the data sequence.
- 05-ustacks-denovo\_data: This folder contains species subdirectories that store the resulting data files from the ustacks program for each individual. There are three files per individual; sampleID.allelles.tsv.gz, sampleID.snps.tsv.gz, and, sampleID.tags.tsv.gz. These files should be permanently stored here and copied to a new directory for any new catalogs and/or when a group of samples are being aligned to a new catalog.
- catalogs\_by\_city: For any given species within city, there is likely to be a slightly different set of SNPs compared to the whole metapopulation of five cities. We examined 24 sets of species-city combinations. These catalogs are permanently stored here.

- catalogs\_by\_species: Metapopulation catalogs are stored within this folder for each species. The metapopulation catalog was created using samples from all populations to create a national catalog.
  - Some notes about catalog directories:
    - Catalogs contain three files; catalog.alleles.tsv.gz, catalog.snps.tsv.gz, and catalog.tafs.tsv.gz. If you would like to use the catalog on a new project, you will need to copy all three files to a new project folder.
    - You can determine which individuals were used to create the catalog by looking at the cstacks\_popmap.txt found within each folder. Specifically for the metapopulation catalogs, this information is also found in the cstacks-metapop-catalog samples-included.csv
    - You can determine which individuals were subsequently aligned to the catalog and used in the subsequent stacks analysis by looking at the popmap\*.txt found within each folder.
    - Each folder also contains the relevant ustacks and stacks pipeline scripts and output files (i.e., from cstacks, gstacks, stacks, tsv2bam, and populations),

# 15.2 Aspera Transfer File Names

See data/aspera\_transfer\_file\_names.csv. Preview:

```
readLines("data/aspera_transfer_file_names.csv", 10)
```

```
[1] "/Hoffman_macrosystems/AMH_macro_1_1_12px_S1_L001_R1_001.fastq.gz"
##
    [2] "/Hoffman_macrosystems/AMH_macro_1_1_12px_S1_L001_R2_001.fastq.gz"
    [3] "/Hoffman_macrosystems/AMH_macro_1_1_12px_S1_L002_R1_001.fastq.gz"
##
##
    [4] "/Hoffman_macrosystems/AMH_macro_1_1_12px_S1_L002_R2_001.fastq.gz"
    [5] "/Hoffman_macrosystems/AMH_macro_1_1_12px_S1_L003_R1_001.fastq.gz"
    [6] "/Hoffman_macrosystems/AMH_macro_1_1_12px_S1_L003_R2_001.fastq.gz"
##
##
    [7] "/Hoffman_macrosystems/AMH_macro_1_1_12px_S1_L004_R1_001.fastq.gz"
    [8] "/Hoffman macrosystems/AMH macro 1 1 12px S1 L004 R2 001.fastq.gz"
##
    [9] "/Hoffman_macrosystems/AMH_macro_1_10_8px_S10_L001_R1_001.fastq.gz"
## [10] "/Hoffman_macrosystems/AMH_macro_1_10_8px_S10_L001_R2_001.fastq.gz"
```

### 15.3 clone\_filter File Names

See data/clone\_filter\_file\_names.csv. Preview:

readLines("data/clone\_filter\_file\_names.csv", 10)

```
## [1] "AMH_macro_1_1_12px_S1" "AMH_macro_1_10_8px_S10" "AMH_macro_1_11_8px_S11"
## [4] "AMH_macro_1_12_8px_S12" "AMH_macro_1_13_8px_S13" "AMH_macro_1_14_8px_S14"
## [7] "AMH_macro_1_2_12px_S2" "AMH_macro_1_3_12px_S3" "AMH_macro_1_4_12px_S4"
## [10] "AMH_macro_1_5_8px_S5"
```