

The gstudio Package

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Preface

This document is intended to be a more in-depth overview of the functionality contained in the `gstudio` package. This package is released under the GPL so if you have particular additions you would like to make to it, feel free to submit them to rjdye@vcu.edu.

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Chapter 1

Getting Genetic Data Into R

1.1 Synopsis

Here you will learn to get genetic data files into the R environment using the `gstudio` package. This package was designed to handle marker-based genetic data (e.g., not sequences *per se* though it can use SNP's and haplotypes) as well as additional data that is typically collected along with individuals.

To get started, first import the `gstudio` package as:

```
> require(gstudio)
> options(warn=-1)
> options(verbose=FALSE)
```

1.2 The Locus Class

The locus class is the fundamental class that handles marker-based genetic data. At present it can handle dominant and co-dominant marker types at any ploidy level. Internally, alleles are stored as a character vector and by default they are not sorted so that the alleles will be presented in the order that you import them (e.g., a 3:1 locus instead of a 1:3 locus). I do not sort these because it may be necessary to know the phase of the alleles in a locus and sorting them would remove that information. If you abhor the sight of a genotype 3:1 then sort it earlier and then try to figure out why you have this affliction.

```
> loc1 <- Locus( c(120,122) )
> loc1

120:122

> loc2 <- Locus( c("A","T") )
> loc2

A:T
```

Note, that internally the alleles are translated into character objects. In all the functions dealing with alleles both integer and character arguments are accepted. There are several methods associated with the Locus, the main ones that you will be working with are shown below by example. See `help("Locus-class")` for a complete discussion.

```
> loc3 <- Locus( c(122,122) )
> loc3

122:122

> is.heterozygote( loc3 )

[1] FALSE
```

```

> loc3[2]
[1] "122"
> loc3[2] <- "124"
> is.heterozygote( loc3 )
[1] TRUE
> length( loc3 )
[1] 2
> summary( loc3 )
Class : Locus
Ploidy : 2
Alleles : 122,124

```

Another useful method of the Locus class is the `as.multivariate` function. This translates the locus into a multivariate coding vector so you can do some real statistics with it. Here is an example:

```

> loc4 <- Locus( c("A", "C") )
> loc4
A:C
> all.alleles <- c("A", "G", "C", "T")
> all.alleles
[1] "A" "G" "C" "T"
> as.vector( loc4, all.alleles )
[1] 1 0 1 0

```

Given that our interaction with SNP data is only going to increase, the Locus class can also handle these in a novel way. Obviously, if snp genotypes are given as nucleotides, then the previous example is perfectly valid. However in a lot of cases (e.g., simulations) we can encode SNP data as the number of minor alleles, and in doing so get away with a little memory savings. The Locus class can input these directly without you having to change the 0/1/2 into alleles with the optional flag `as.snp.minor` (it defaults to `FALSE`). The constructor assigns the generic alleles "A" and "B" (where "B" is the minor allele).

```

> loc5 <- Locus( 0, as.snp.minor=TRUE )
> loc5
A:A
> loc6 <- Locus( 1, as.snp.minor=TRUE )
> loc6
A:B
> loc7 <- Locus( 2, as.snp.minor=TRUE )
> loc7
B:B

```

1.3 The Population Class

You can think of a Population as a collection of one or more individuals. While no man is an island, an individual is just a population of $N = 1$. Each individual, can have any number of Locus objects along with other non-genetic information associated with them (e.g., latitude, longitude, dbh, hair color, etc.). You create a population by passing it data columns in much the same way as how you create a `data.frame` (in fact, the Population class is just a `data.frame` that knows how to deal with Locus objects and how to give you population genetic summaries).


```

> strata <- c("A","A","B","B","B")
> TPI <- c(Locus(c(1,2)),Locus(c(2,3)),Locus(c(2,2)),Locus(c(2,2)),Locus(c(1,3)))
> PGM <- c(Locus(c(4,4)),Locus(c(4,3)),Locus(c(4,4)),Locus(c(3,4)),Locus(c(3,3)))
> Env <- c(12,20,14,18,10)
> thePop <- Population( Pop=strata, Env=Env, TPI=TPI, PGM=PGM )
> thePop

  Pop Env TPI PGM
1   A  12 1:2 4:4
2   A  20 2:3 3:4
3   B  14 2:2 4:4
4   B  18 2:2 3:4
5   B  10 1:3 3:3

> summary(thePop)

      Pop           Env      TPI      PGM
Length:5      Min.    :10.0   1:2:1   3:3:1
Class :character 1st Qu.:12.0   1:3:1   3:4:2
Mode  :character Median :14.0   2:2:2   4:4:2
                        Mean  :14.8   2:3:1
                        3rd Qu.:18.0
                        Max.   :20.0

> names(thePop)

[1] "Pop" "Env" "TPI" "PGM"

```

1.3.1 Accessing Population Elements

You can also add data to a Population or remove it

```

> WXY <- c(Locus(c(122,124)),Locus(c(124,126)),Locus(c(124,124)),Locus(c(122,124)),Locus(c(126,126)))
> thePop$WXY <- WXY
> thePop

  Pop Env TPI PGM   WXY
1   A  12 1:2 4:4 122:124
2   A  20 2:3 3:4 124:126
3   B  14 2:2 4:4 124:124
4   B  18 2:2 3:4 122:124
5   B  10 1:3 3:3 126:126

> thePop$WXY <- NULL
> thePop

  Pop Env TPI PGM
1   A  12 1:2 4:4
2   A  20 2:3 3:4
3   B  14 2:2 4:4
4   B  18 2:2 3:4
5   B  10 1:3 3:3

```

Similar to the previous constructs, you can access elements within a Population using either numerical indexes, slices, or names.

```

> ind3 <- thePop[3,]
> ind3

  Pop Env TPI PGM
1   B  14 2:2 4:4

```

```
> thePop[ thePop$Pop=="B", ]
```

```
  Pop Env TPI PGM
1   B  14 2:2 4:4
2   B  18 2:2 3:4
3   B  10 1:3 3:3
```

```
> thePop[ thePop$Env<15 , ]
```

```
  Pop Env TPI PGM
1   A  12 1:2 4:4
2   B  14 2:2 4:4
3   B  10 1:3 3:3
```

```
> TPI <- thePop[,3]
```

```
> print(TPI)
```

```
[[1]]
1:2
```

```
[[2]]
2:3
```

```
[[3]]
2:2
```

```
[[4]]
2:2
```

```
[[5]]
1:3
```

1.3.2 Getting Data Types within Population Objects

Since a Population can hold several types of data and the main way to get data from one is to know its name, the method `column.names` can provide you quick access to all the data names of a specific R class.

```
> strata <- column.names(thePop,"character")
```

```
> strata
```

```
[1] "Pop"
```

```
> column.names(thePop,"Locus")
```

```
[1] "TPI" "PGM"
```

```
> column.names(thePop,"numeric")
```

```
[1] "Env"
```

1.3.3 Partitioning Population Objects

A Population object can contain individuals with several other categorical data variables (e.g., population, region, habitat, etc.) and it is relatively easy to get single elements (as shown in the slicing above) as well as complete partitions. It should be pointed out that when you partition a Population on some stratum, it will remove that stratum from all the partitions though it will leave the other partitions in the subpopulations.

```
> subpops <- partition(thePop,stratum="Pop")
```

```
> print(subpops)
```

```
$A
  Env TPI PGM
1  12 1:2 4:4
2  20 2:3 3:4
```

```
$B
  Env TPI PGM
1  14 2:2 4:4
2  18 2:2 3:4
3  10 1:3 3:3
```

1.3.4 Generic Population Functions

The following generic functions are available for the `Population` class and work just like they do using other data structures.

length The number of `Individual` objects (rows) in the `Population`.

dim The number of row and columns in the `Population`.

names The data column names.

summary A summary of the data columns in the `Population`.

show Dumps the `Population` to the terminal.

row.names Returns the names of the rows (they are integers so this isn't too exciting).

1.4 Importing Data

OK, so typing all this stuff in is rather monotonous and will be a total pain if you have a real data set with hundreds or thousands of individuals and a righteous amount of loci.

The main function for importing data from a text file into a `Population` object is `read.population` and assumes the following about your data:

1. You have your data in a TEXT file that is comma separated (*.csv).
2. You have a header row on your file with the names of each column of data. Headers should not have spaces in them, R will replace them with a period.
3. Genetic marker that have more than one allele are encoded using a colon ":" separating alleles. This means that the diploid microsatellite locus with alleles 122 & 128 would be in a single column as 122:128. This allows you to have triploid, tetraploid, etc markers with not other encoding.
4. Haploid markers are do not need a ":", just put in the haplotype. With haploid data, searching for ":" won't work so you need to pass the number of haploid loci as the optional parameter `num.single.digit` to `read.population`. The haploid loci *must* be the last `num.single.digit` right-most columns in your data set.
5. All alleles will be treated internally as a character string (except for in a few cases such as estimating ladder-distance). So you can use all alphanumeric characters for alleles but stay away from punctuation.
6. Missing data should be encoded as NA (for the whole genotype NA:NA is just silly).
7. If you have a mixture of genetic data types, columns with ":" will be automatically interpreted as `Locus` objects. You can mix in haploid data types by putting them in the last, right-most, columns and pass the optional parameter `num.single.digit` with the number columns to put as haploid.

1.4.1 Reading From a Text File

An example data file may look like:

```
Population,Lat,Lon,PGM,TPI
Loreto,22.25,-102.01,120:122,A:T
Loreto,22.25,-102.01,122:124,A:C
Cabo,22.88,-109.9,120:A:A
Cabo,22.88,-109.9,NA,A:T
```

This file can be loaded as (assuming `getwd()` contains the file)

```
> pop <- read.population(file="testData.csv")
```

In general, if you can open your file using `read.table`, then `read.population` should work.

For SNP data sets that are encoded as 0/1/2 (# minor alleles), there is an optional switch `num.single.digit` that will allow you to indicate the last X columns of data as SNP loci with the minor allele encoding.

1.4.2 Using Google Spreadsheets To Share Data

One of the really great things about google docs is that you can use it to share information and documents with others and here we will be examining how to use it to keep public data available for analysis in R.

The first step is to provide a bit of data to share. The following example uses the shared *Cornus florida* data set. This consists of adults and offspring.

To share a document, click the "Share" button and you will be presented with a popup window giving you options on what to do similar to Figure 1.4.2.

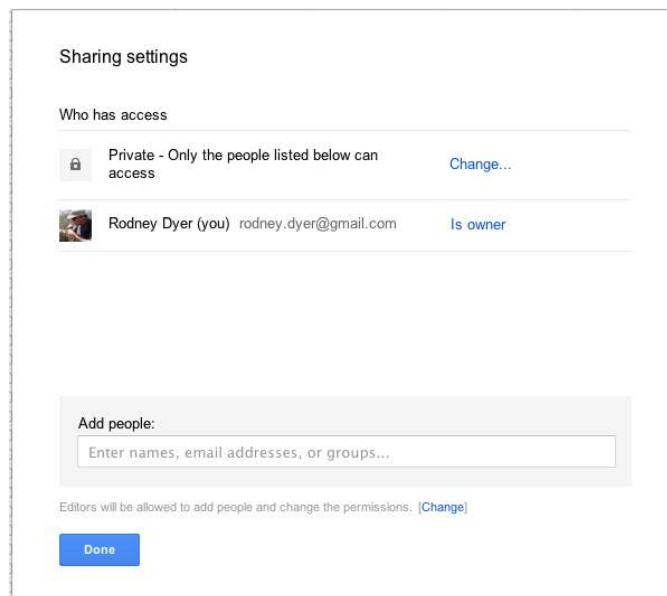


Figure 1.1: Settings to adjust sharing options for google document.

Where it says Private select the "Change..." option and change the Visibility Options to "Anyone with the link" and hit save. It will then return to the Sharing Settings (Figure 1.4.2) page and provide you a unique link to the document.

This gives individuals access to the spreadsheet as a whole, but what we would like to do is to get to the contents of it as a *.csv file. In the spreadsheet, select File → Publish to the Web and select the following options in the dialog:

1. Sheets to Publish → All sheets
2. Check the box Automatically republish when changes are made
3. Select Start publishing.

This will make the bottom part of the dialog active and you'll need to make the following changes:

1. Change type from Web → CSV

2. Change All Sheets → Sheet1
3. Change All Cells → the range that you want to share. Here you need to use Excel-like notation such as A1:I63 for the box from column A, first row to column I, 63nd row.

The dialog provides a URL for these data, the one above is:

```
https://docs.google.com/spreadsheet/pub?hl=en_US&hl=en_US&key=0Aq-1sUWPDuZtdF9xMXZGQWNtbk1FNTVWd3F3U0FDdXc&single=true&gid=0&range=A1%3AG63&output=csv
```

1.5 Getting Data Into R from GoogleDocs

Now we have a data set that is available on the web and we can get to it from within R using the `getURL`, `read.csv`, and `textConnection` functions as follows (n.b. I truncated the URL as it goes off the end of the page, it is the one from above.)

```
> spreadsheetURL <- "https://docs.google.com/spreadsheet/pub?hl=en_US&hl=en_US&key=0Aq-1sUWPDuZtdF9xMXZGQWNtbk1FNTVWd3F3U0FDdXc&single=true&gid=0&range=A1%3AG63&output=csv"
> dogwood <- read.population( googleURL=dogwoodURL )
```

And there you go, you have now used your Google Account to host data that is available to everyone... No go forth and share.

1.5.1 Example Data Sets

The `gstudio` package comes with some example data sets already loaded. To access these data sets, use the `data` function and they will be put into your workspace (already formatted as `Population` objects).

```
> data(araptus_attenuatus)
> summary(araptus_attenuatus)
```

Species	Cluster	Pop	Individual	Lat
CladeA: 75	CBP-C :150	32 : 19	101_10A: 1	Min. :23.08
CladeB: 36	NBP-C : 84	75 : 11	101_1A : 1	1st Qu.:24.59
CladeC:252	SBP-C : 18	Const : 11	101_2A : 1	Median :26.25
	SCBP-A: 75	12 : 10	101_3A : 1	Mean :26.25
	SON-B : 36	153 : 10	101_4A : 1	3rd Qu.:27.53
		157 : 10	101_5A : 1	Max. :29.33
		(Other):292	(Other):357	
Long	LTRS	WNT	EN	EF
Min. :-114.3	01:01:147	03:03 :108	01:01 :225	01:01:219
1st Qu.: -113.0	01:02: 86	01:01 : 82	01:02 : 52	01:02: 52
Median :-111.5	02:02:130	01:03 : 77	02:02 : 38	02:02: 90
Mean :-111.7		02:02 : 62	03:03 : 22	NA : 2
3rd Qu.: -110.5		NA : 11	01:03 : 7	
Max. :-109.1		03:04 : 8	03:04 : 6	
		(Other): 15	(Other): 13	
ZMP	AML	ATPS	MP20	
01:01: 46	08:08 : 51	05:05 :155	05:07 : 64	
01:02: 51	07:07 : 42	03:03 : 69	07:07 : 53	
02:02:233	07:08 : 42	09:09 : 66	18:18 : 52	
NA : 33	04:04 : 41	02:02 : 30	05:05 : 48	
	NA : 23	07:09 : 14	05:06 : 22	
	07:09 : 22	08:08 : 9	11:11 : 12	
	(Other):142	(Other): 20	(Other):112	

Chapter 2

Summarizing Genetic Data

2.1 Synopsis

There are several ways you can summarize genetic data and here we will cover some simple approaches and introduce another class that aids in the analysis of population genetic data.

2.2 The Frequencies Class

The `Frequencies` class was designed to help out with allele frequency issues and provide a single interface from which you can extract frequency-related information. At its most basic level, a new `Frequencies` object is created from a list of `Locus` objects.

```
> loc1 <- Locus( c(1,2) )
> loc2 <- Locus( c(2,2) )
> loc3 <- Locus( c(2,2) )
> freqs <- Frequencies( c( loc1, loc2, loc3) )
> freqs
```

Allele Frequencies:

```
1 = 0.1666667
2 = 0.8333333
```

Estimates of allele frequencies can be extracted from the `Frequencies` class using the `get.frequencies` method. This method needs to have the object and an optional list of alleles you are interested in getting frequencies for. If you do not pass the second parameter, it will give you the frequencies for all the alleles it currently has. If you do, it will give you the observed frequency of each (notice the value for the '42' allele)

```
> names(freqs)
[1] "1" "2"
> length(freqs)
[1] 2
> get.frequencies( freqs )
      1      2
0.1666667 0.8333333
> get.frequencies( freqs, c("1","42") )
      1      42
0.1666667 0.0000000
```

2.3 Heterozygosities

A fundamental component of many population genetic analysis is the estimation of heterozygosity. There are two basic types of heterozygosity, that which is expected under Hardy-Weinberg Equilibrium and that which was observed. For simplicity, these are denoted as H_e and H_o in many common texts.

Observed heterozygosity is probably the simplest of the two and it is simply the fraction of genotypes in the group you are looking at (could be a population or a region or a site) that are heterozygotes. In terms of the Locus class, the function `is.heterozygote` returns TRUE if the locus has at least two alleles (allowing for ploidy levels in excess of 2) and at least two different alleles are present. As part of the data accumulation process in the construction of an `AlleleFrequency` object, observed heterozygosity is recorded.

Expected heterozygosity requires an assumption of equilibrium (in the most simple case). For a diploid locus with alleles A & B and frequencies of each allele denoted as p_A & p_B , genotypes are expected to occur at a frequency of:

$$\begin{aligned} AA &\rightarrow p_A^2 \\ AB &\rightarrow 2 * p_A * p_B \\ BB &\rightarrow p_B^2 \end{aligned}$$

From the example set of loci we used above, the observed and expected frequencies are:

```
> ho( freqs )
      ho
0.3333333
> he( freqs )
      he
0.2777778
```

2.4 Allele Frequencies

The estimation of allele frequencies for a single site or population is probably one of the least informative summary approaches available. It is the differences among sites & populations and the various evolutionary and demographic processes that create these differences that are often of interest.

There are several helper functions and methods that can be used to examine allele frequencies across strata.

2.4.1 Getting Frequencies from Populations

The `Population` class has a method for returning an `AlleleFrequency` object for a particular locus. This is mostly a convenience method that goes through all the `Individual` objects in the `Population` and creates a new `AlleleFrequency` object for you. As a single population you can grab it using the `allele.frequencies` routine.

```
> data(araptus_attenuatus)
> araptus.ltrs.freq <- allele.frequencies(araptus_attenuatus, "LTRS")
> araptus.ltrs.freq

$LTRS
Allele Frequencies:
 01 = 0.523416
 02 = 0.476584
```

If you do not pass `get.frequencies` the optional `loci` parameter, it will return a list of `Frequency` objects for all loci.

```
> all.freqs <- allele.frequencies(araptus_attenuatus)
> print(all.freqs[1:2])
```

```
$LTRS
Allele Frequencies:
  01 = 0.523416
  02 = 0.476584
```

```
$WNT
Allele Frequencies:
  01 = 0.3579545
  03 = 0.4303977
  04 = 0.02698864
  02 = 0.1818182
  05 = 0.002840909
```

With the partition method, you can take the entire data set and easily find allele frequencies for subsets of data.

```
> clades <- partition(araptus_attenuatus,"Species")
> names(clades)
```

```
[1] "CladeA" "CladeB" "CladeC"
```

```
> cladeC.freqs <- allele.frequencies(clades$CladeC)
> summary(cladeC.freqs)
```

	Length	Class	Mode
LTRS	2	Frequencies	S4
WNT	4	Frequencies	S4
EN	5	Frequencies	S4
EF	2	Frequencies	S4
ZMP	2	Frequencies	S4
AML	10	Frequencies	S4
ATPS	6	Frequencies	S4
MP20	8	Frequencies	S4

```
> summary(cladeC.freqs$AML)
```

```
Class : Frequencies
N : 252
A : { 01, 02, 05, 06, 07, 08, 09, 10, 11, 13 }
ho : 0.4677419
he : 0.7284242
```

```
> get.frequencies(cladeC.freqs$AML, 11)
```

```
11
0.002016129
```

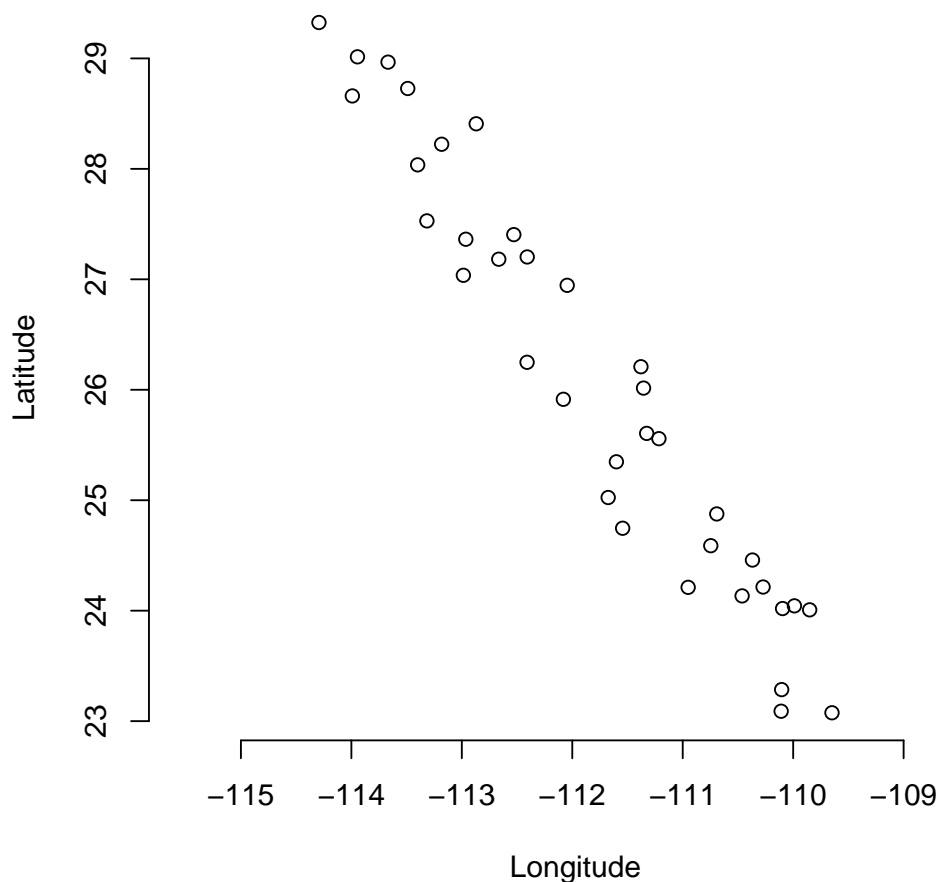
```
> allele.frequencies( araptus_attenuatus[ araptus_attenuatus$Lat > 26.3 ,], loci="AML" )
```

```
$AML
Allele Frequencies:
  08 = 0.308642
  09 = 0.2592593
  07 = 0.2407407
  10 = 0.02469136
  06 = 0.03703704
  11 = 0.08333333
  02 = 0.00308642
  13 = 0.00308642
  05 = 0.00308642
  01 = 0.00308642
  12 = 0.03395062
```


2.4.2 Plotting Frequencies

The combination of Population and Frequencies can easily be used to explore population structure. In the next snippet, we partition the dataset into populations along the Baja Peninsula and plot their locations (n.b., the `bty` option to `plot` removes the box around the image and the `asp` makes the axes equal).

```
> baja <- araptus_attenuatus[araptus_attenuatus$Species!="CladeB",]  
> pop.coords <- unique( cbind( baja$Long, baja$Lat ) )  
> plot(pop.coords, bty="n", xlab="Longitude", ylab="Latitude",asp=1)
```



Next, we can adjust the size of the symbol by diversity at any locus (below `LTRS` is used). Here the `lapply` function is used to apply a function to the elements of the `baja.pops` list. If you are not familiar with this function, you should look it up. The resulting heterozyosity estimates are scaled and used as symbol size (via `cex`; Figure 2.1).

```
> baja.pops <- partition( baja, "Pop" )  
> pop.he <- lapply( baja.pops, function(x) he( Frequencies( x$LTRS ) ) )  
> summary( unlist(pop.he) )
```

Min.	1st Qu.	Median	Mean	3rd Qu.	Max.
0.0000	0.0000	0.1800	0.2036	0.3457	0.4800

```
> plot(pop.coords, bty="n", xlab="Longitude", ylab="Latitude",asp=1,cex=2*unlist(pop.he)+1, main="Heterozygosity")
```

```

> baja.pops <- partition( baja, "Pop" )
> pop.he <- lapply( baja.pops, function(x) he( Frequencies( x$LTRS ) ) )
> summary( unlist(pop.he) )
  Min. 1st Qu.  Median    Mean 3rd Qu.    Max.
0.0000 0.0000 0.1800 0.2036 0.3457 0.4800
> plot(pop.coords, bty="n", xlab="Longitude", ylab="Latitude",asp=1,cex=2*unlist(pop.he)+1, main="Heterozygosity of LTRS

```

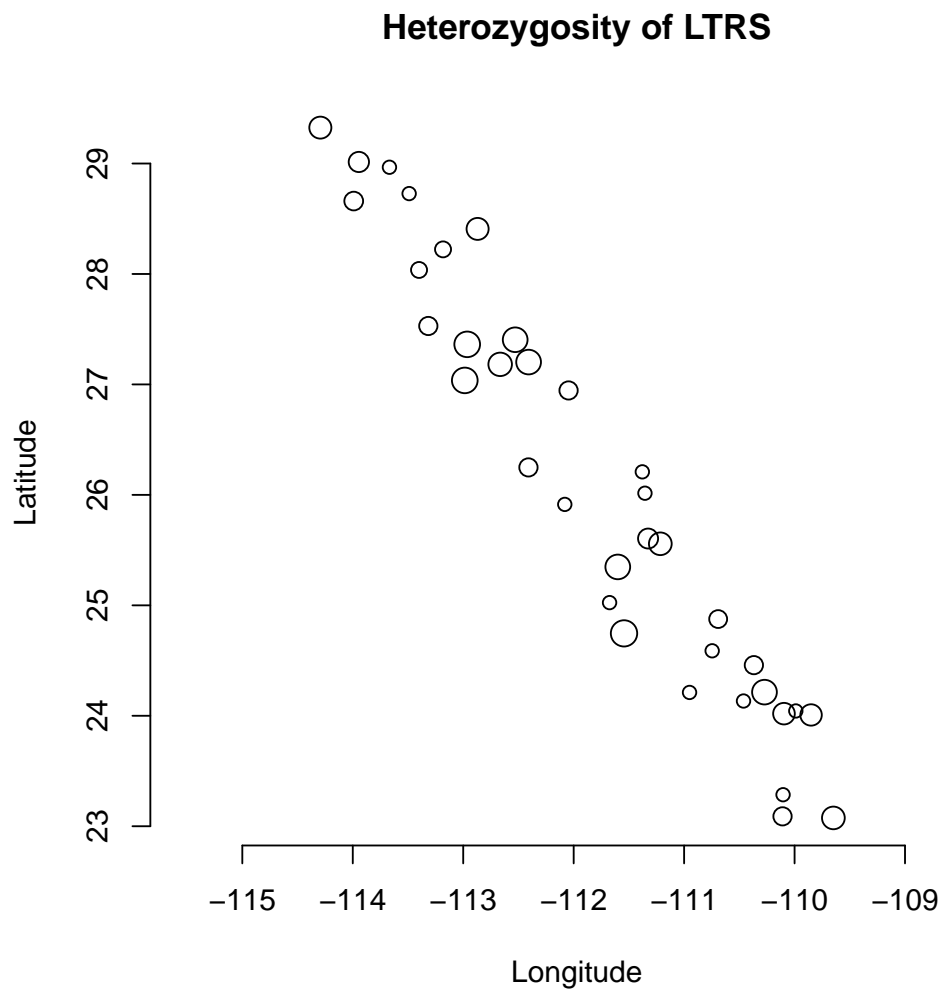


Figure 2.1: Heterozygosity of *Araptus attenuatus* populations (depicted by symbol size) on the peninsula of Baja California.

Chapter 3

Genetic Diversity

3.1 Synopsis

Genetic diversity is measure of within stratum variance and there are several methods available for the estimation of diversity. In a general sense, we will be using measures of allelic richness from the Baja California data set, which can easily be found by examining the Frequencies of the loci.

```
> data(araptus_attenuatus)
> baja <- araptus_attenuatus[araptus_attenuatus$Species != "CladeB",]
> freqs <- allele.frequencies(baja)
> freqs$LTRS
```

Allele Frequencies:

```
01 = 0.5519878
02 = 0.4480122
```

```
> freqs$MP20
```

Allele Frequencies:

```
07 = 0.2892308
05 = 0.2969231
15 = 0.001538462
08 = 0.02769231
06 = 0.08923077
04 = 0.009230769
18 = 0.1784615
19 = 0.009230769
17 = 0.04153846
10 = 0.01384615
11 = 0.04153846
16 = 0.001538462
```

In this data set, the raw allelic diversity across all the samples range from 2 - 12 alleles. However, using a base approach such as this falls short for several reasons:

1. We are only looking at the number of alleles across the entire data set and there are many cases where it may be of interest to look at allelic diversity within substrata. It is possible to use the partition function along with `allele.frequencies` to get to the number of alleles at partitions but the problem with that is:
2. The raw number of alleles depends upon the number of individuals sampled. It is not statistically sound to compare raw diversity of stratum with different numbers of individuals. This is where *rarefaction* comes in.
3. The sole number of alleles present may not be as important as other measures of genetic diversity such as the diversity of non-rare alleles, or the average 'effective' number of alleles.

To overcome both of these issues, the `genetic.diversity` function is used.

3.1.1 Rarefaction

Before we get into the nitty-gritty, the basic concept of rarefaction should be examined. Rarefaction is a permutation technique that can be used to standardize samples based upon sample allocation and is an old friend to ecologists.

For our purposes, we will consider rarefaction as a subsampling of alleles in strata standardized by the size of the smallest stratum. So if we have one population with 10 individuals (20 alleles if the locus is diploid) and the rest of the populations have 50 individuals (100 alleles), a rarefied comparison of diversity should be based upon sampling of 20 alleles.

The function `genetic.diversity` takes random samples of the alleles within each population and recomputes the requested allelic diversity statistic. While in many ecological studies, rarefaction is depicted as an accumulation curve (they are generally interested in sampling intensity), `genetic.diversity` only reports the distribution at the largest size where all strata are equal (e.g., the number of alleles present in the smallest population).

3.2 Allelic Diversity: A

The parameter A is solely a measure of the number of alleles at a locus. If a population has a single individual with a single copy of allele A and everyone else has allele C , $A = 2$, which is the same case as if half the population was homozygous for A and the remaining individuals were homozygous for C . The function `genetic.diversity` returns an object that can be both printed and examined in plot fashion (by default it is a boxplot)

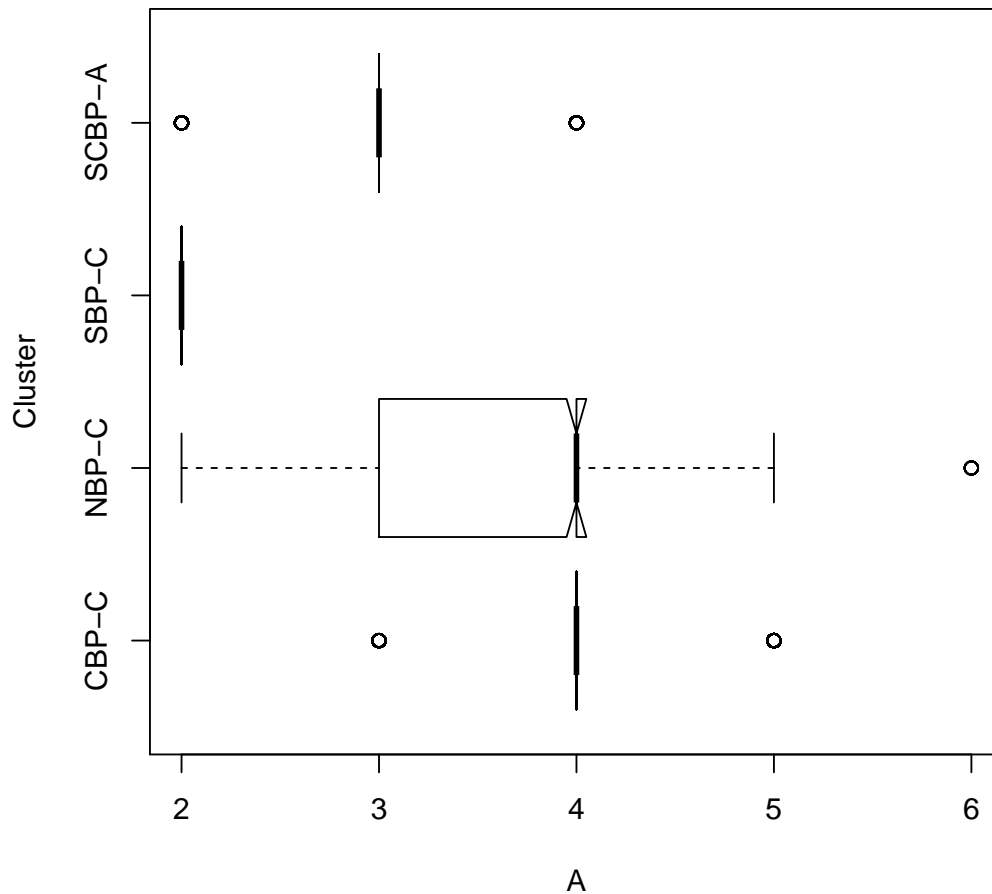
```
> A <- genetic.diversity(baja, stratum="Cluster", loci="MP20", mode="A")
> A
```

Geneic Diversity:

```
Estimator: A
Stratum: Cluster
Loci: { MP20 }
Locus = MP20
CBP-C A = 5 ; Rarefaction A = 4.15115115115115
NBP-C A = 6 ; Rarefaction A = 3.54454454454454
SBP-C A = 2 ; Rarefaction A = 2
SCBP-A A = 4 ; Rarefaction A = 2.96796796796797
```

```
> plot(A)
```

Genetic Diversity MP20



The plot itself is a horizontal boxplot. If you conduct the analysis with either the `loci` missing or as a list of loci, the results from each locus will be displayed in the terminal and the plotting will cycle through each locus requiring some input from the keyboard. It is also possible to plot just a single locus by passing the locus name as a second parameter to the `plot` command.

3.3 Allelic Diversity of Non-Rare Alleles: A_{95}

The parameter A_{95} ignores rare alleles by not counting those whose frequencies are below 95% *within* the stratum. So alleles locally rare will not be counted and in general $A \geq A_{95}$.

```
> A95 <- genetic.diversity(baja, stratum="Cluster", loci="MP20", mode="A95")
> A95
```

Genetic Diversity:

Estimator: A95

Stratum: Cluster

Loci: { MP20 }

Locus = MP20

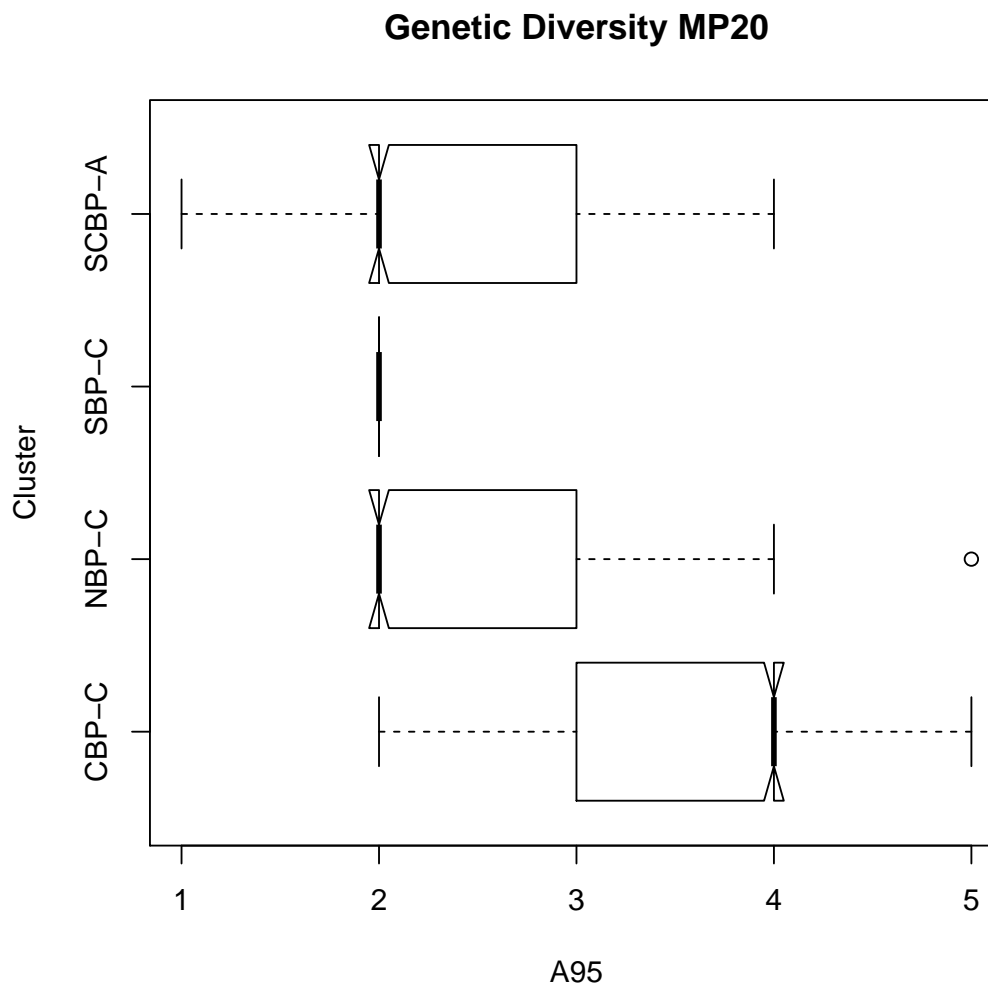
CBP-C A95 = 4 ; Rarefaction A95 = 3.61961961961962

NBP-C A95 = 2 ; Rarefaction A95 = 2.4034034034034

SBP-C A95 = 2 ; Rarefaction A95 = 2

SCBP-A A95 = 2 ; Rarefaction A95 = 2.44044044044044

```
> plot(A95)
```



3.4 Effective Allelic Diversity: A_e

The last diversity statistic is A_e , which is another frequency corrected allelic diversity statistic. For a locus with ℓ alleles, each of which occurs at a frequency of p_i , the effective number of alleles is:

$$A_e = \frac{1}{\sum_{i=1}^{\ell} p_i^2} \quad (3.1)$$

And for the example data:

```
> Ae <- genetic.diversity(baja,stratum="Cluster",loci="MP20",mode="Ae")
> Ae
```

Geneic Diversity:

Estimator: Ae

Stratum: Cluster

Loci: { MP20 }

Locus = MP20

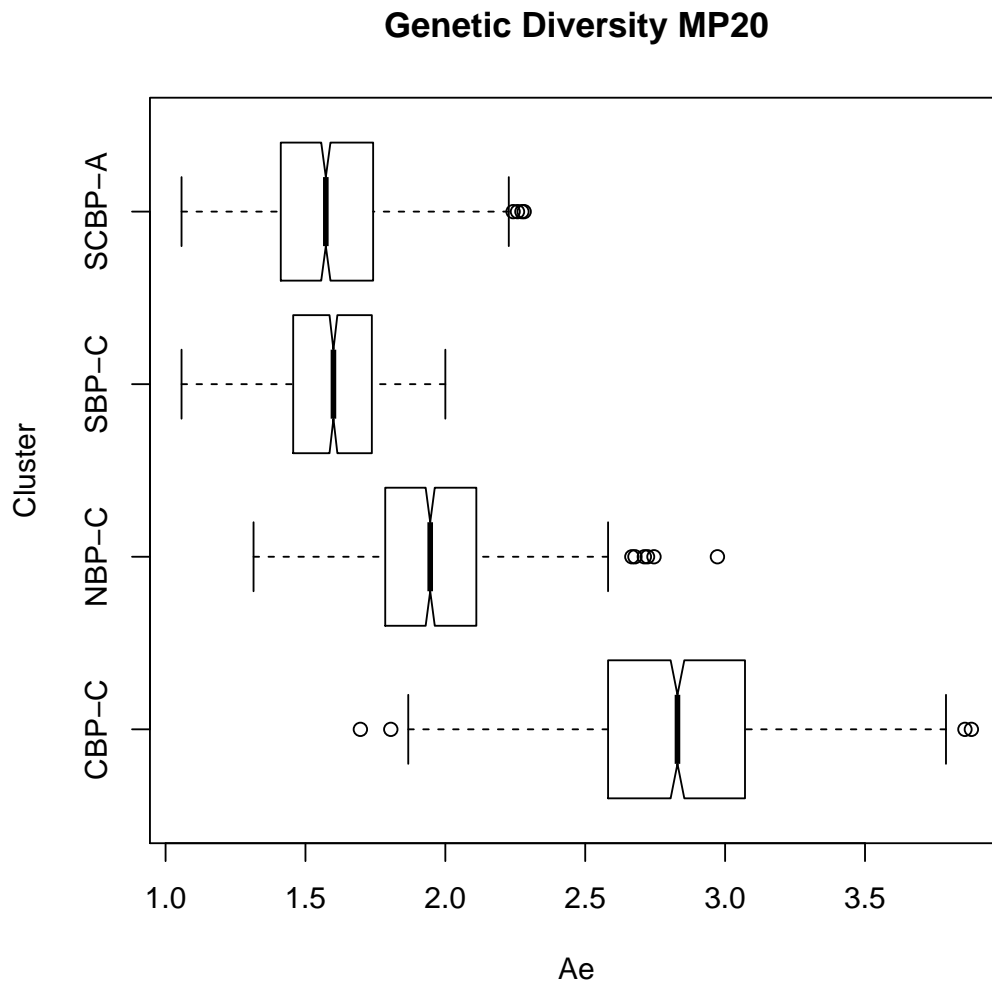
CBP-C Ae = 2.93481610504455 ; Rarefaction Ae = 2.83608876964118

NBP-C Ae = 1.97536394176932 ; Rarefaction Ae = 1.95049522307667

SBP-C $A_e = 1.6$; Rarefaction $A_e = 1.59774826523195$

SCBP-A $A_e = 1.58205596962453$; Rarefaction $A_e = 1.59294272219415$

```
> plot(Ae)
```



One obvious difference in A_e from the others is that it is not an integer value (both A and A_{95} are integers) and as such can show a bit more granularity.

Chapter 4

Genetic Distance

4.1 Synopsis

The analysis of genetic data is largely an analysis of distances; distances among frequencies, distances among centroids of populations, etc.

4.2 Genetic Distances Among Individuals

In these examples, the data from *Araptus attenuatus* will be used again but this time we'll use the subset of individuals from "CladeB" (mainland populations).

```
> data(araptus_attenuatus)
> sonora <- araptus_attenuatus[ araptus_attenuatus$Species=="CladeB" , ]
> summary(sonora)
```

Species	Cluster	Pop	Individual	Lat	Long	
CladeB:36	SON-B:36	101: 9	101_10A: 1	Min. :26.38	Min. : -110.6	
		102: 8	101_1A : 1	1st Qu.:26.64	1st Qu.: -109.6	
		32 :19	101_2A : 1	Median :26.64	Median : -109.3	
			101_3A : 1	Mean :26.90	Mean : -109.6	
			101_4A : 1	3rd Qu.:26.95	3rd Qu.: -109.3	
			101_5A : 1	Max. :27.91	Max. : -109.1	
			(Other):30			
LTRS	WNT	EN	EF	ZMP	AML	ATPS
01:01: 1	01:01:29	01:01: 7	01:01:23	01:01: 1	08:08: 1	02:02:28
01:02:17	01:03: 1	01:03: 2	01:02:11	02:02:19	08:11: 1	02:03: 1
02:02:18	NA : 6	03:03:19	NA : 2	NA :16	08:12: 1	02:04: 2
		03:04: 6			10:11: 1	02:09: 3
		04:04: 1			11:11:12	04:04: 1
		NA : 1			12:12: 5	09:09: 1
					NA :15	
MP20						
12:12 : 6						
03:13 : 4						
11:12 : 3						
13:13 : 3						
NA : 3						
02:10 : 2						
(Other):15						

4.2.1 Jaccard Distance

Jaccard distance is a set-theoretic distance quantifying dissimilarity. Assuming that loci are sets of alleles, the Jaccard dissimilarity between genotypes A and B is given by:

$$J_{\delta}(A, B) = \frac{|A \cup B| - |A \cap B|}{|A \cup B|} \quad (4.1)$$

Using the LTRS locus, we compute this distance as:

```
> d.jaccard <- genetic.distance(sonora, stratum="Pop", loci="EN", mode="Jaccard")
> dim(d.jaccard$LTRS)
```

NULL

YOU can look at the elements of the LTRS matrix (it is 36x36 so I am not printing it out here). With `mode="Jaccard"`, missing genotypes will result in NA rows and columns in the distance matrix. It is not entirely clear how this metric can easily handle missing genotypes.

4.2.2 Bray-Curtis Distance

Bray-Curtis Distance (Bray & Curtis 1957) has been primarily used to quantify differences in species composition. It is defined as the total number of species that are unique to either of the two sites standardized by the number of species in both sites.

$$BC_{\delta} = \frac{S_i + S_j - 2S_{ij}}{S_i + S_j} \quad (4.2)$$

where S_x is the species count and S_{ij} is the sum of minimum abundances. Lately, this has seen considerable use within individual-based landscape genetic studies. Missing genotypes are set to average allele frequencies, that is to say that every missing genotype is considered to have all the alleles present in the entire population, but with probability equal to their global frequencies. Essentially, this removes the NA problem like in the `mode="Jaccard"` situation and does so by taking the non-missing genotype's genetic distance from the global genetic centroid (it's cosmic man!). Here is the estimation using two loci.

```
> d.bray <- genetic.distance(sonora, stratum="Pop", loci=c("LTRS", "EN"), mode="Bray")
> summary(d.bray)
```

	Length	Class	Mode
LTRS	1296	-none-	numeric
EN	1296	-none-	numeric

4.2.3 AMOVA Distance

The final individual-based approach is based upon the Analysis of Molecular Variance (AMOVA) analysis. A geometric interpretation of this genetic distance is given in Figure 4.1 indicating distances among diploid genotypes.

Algebraically, we can define an individual locus using a multivariate vector as an allele coding vector. The Locus class has a method, `as.multivariate`, that does the translation. The distance between the two alleles is defined as:

$$\delta_{ij}^2 = 2(p_i - p_j)^2 \quad (4.3)$$

as shown below.

The amova distance is simply the vector distance between these two vectors as demonstrated below

```

> locAA <- Locus( c("A","A") )
> locBB <- Locus( c("B","B") )
> locAB <- Locus( c("A","B") )
> locBC <- Locus( c("B","C") )
> vAA <- as.vector( locAA, c("A","B","C") )
> vBB <- as.vector( locBB, c("A","B","C") )
> vAB <- as.vector( locAB, c("A","B","C") )
> vBC <- as.vector( locBC, c("A","B","C") )
> dist.AA.BB <- 2*( (vAA - vBB) %*% (vAA - vBB) )
> dist.AA.BB

```

```

      [,1]
[1,]    16

```

```

> dist.AA.AB <- 2*( (vAA - vAB) %*% (vAA - vAB) )
> dist.AA.AB

```

```

      [,1]
[1,]     4

```

```

> dist.AA.BC <- 2*( (vAA - vBC) %*% (vAA - vBC) )
> dist.AA.BC

```

```

      [,1]
[1,]    12

```

While we will deal more with the AMOVA analysis in the section on Genetic Structure, the AMOVA genetic distance matrix can be estimated as follows, this time using *all* the loci. This metric is additive across loci, so only a single distance matrix is returned. The list key for the multilocus parameters is a list of the locus names, joined using a period.

```

> d.amova <- genetic.distance(sonora,stratum="Pop",mode="AMOVA",loci="EN")
> summary(d.amova)

```

```

      Length Class  Mode
EN 1296    -none- numeric

```

There are several other measures of individual-to-individual distance such as relatedness and coancestry. These are not currently implemented in R but may become available in the near future. That being said, it is probably something not too difficult for someone to extend these functions with their own code.

4.2.4 Differences Between Distances

These three distances are correlated, and here we can look at how close they are for this three allele locus in *Euphorbia lomelii*. They will be transformed from a dist matrix object into columns within a data.frame and then their relationship can be tested using cor.test.

```

> df <- data.frame( jaccard = d.jaccard$EN[lower.tri(d.jaccard$EN)],bray = d.bray$EN[lower.tri(d.bray$EN)],
> summary(df)

```

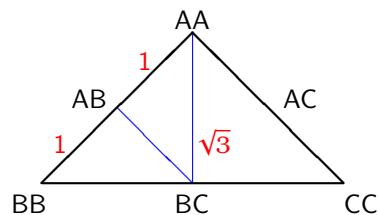


Figure 4.1: Geometry of AMOVA distances. The resulting squared distance is the square of the geometric distance.

jaccard	bray	amova
Min. :0.000	Min. :0.0000	Min. :0.000
1st Qu.:0.000	1st Qu.:0.0000	1st Qu.:0.000
Median :0.500	Median :0.5000	Median :1.000
Mean :0.527	Mean :0.5238	Mean :1.568
3rd Qu.:1.000	3rd Qu.:1.0000	3rd Qu.:4.000
Max. :1.000	Max. :1.0000	Max. :4.000

```
> cor(df)
```

	jaccard	bray	amova
jaccard	1.0000000	0.9985311	0.8883334
bray	0.9985311	1.0000000	0.8919370
amova	0.8883334	0.8919370	1.0000000

```
> pairs(df)
```

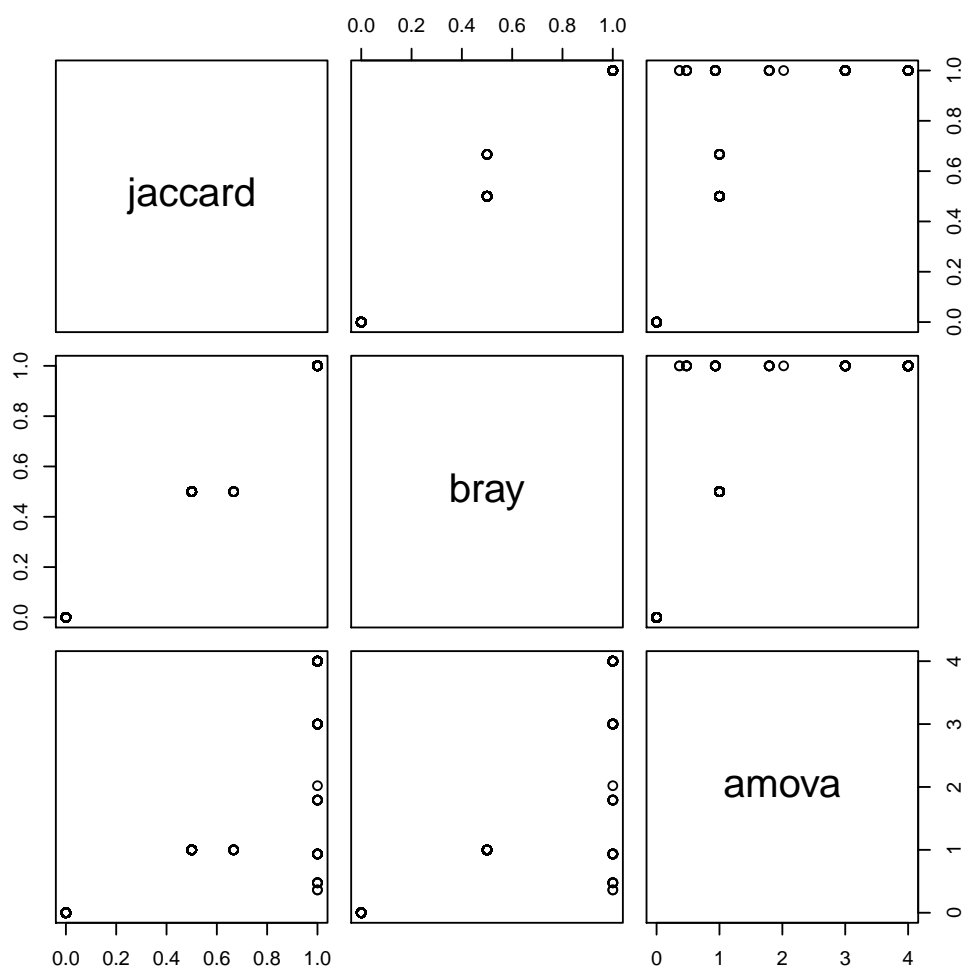


Figure 4.2: Relationship among three individual genetic distance metrics estimated for individual *Araptus attenuatus* individuals in Sonora & Sinoloa, Mexico.

4.3 Genetic Distance Among Strata

Genetic distances can also be estimated among groups of individuals. The same data will be used here but since there are only three populations, we'll be able to see the whole distance matrix.

4.3.1 Euclidean Distance

Euclidean distance is the most straight-forward distance metric available as it is essentially straight-line distance based upon the allele frequencies in each population. It is given by:

$$d_{eucl} = \sqrt{\sum_{j=1}^L (p_{ij} - p_{kj})^2}$$

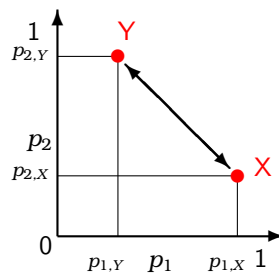


Figure 4.3: Geometry of euclidean distance based upon a two-allele locus denoted as frequencies p_1 & p_2 .

where p_{ij} and p_{kj} are the frequencies of the j^{th} allele in both the i^{th} and j^{th} population. In this and the following distance examples, I am going to take the resulting distance matrix among all pairs of populations and put them into a Neighbor joining tree (via the `nj` function from the `ape` package) as it may be easier to see differences in topologies rather than matrices.

It is perhaps easiest to think of Euclidean distance in x,y coordinate space (Figure 4.3). This distance can be estimated by `stratum.distance` using the optional parameter `method='eucl'` and it will return a `dist` matrix.

Once the matrix has been estimated, you can visualize it in many ways. One of the most straight-forward approaches it to visualizing the relationships among rows and columns is to put it into a bifurcating tree.

```
> d.eucl <- genetic.distance(sonora, stratum="Pop", loci="EN", mode="Euclidean")
> d.eucl
```

```
$EN
      [,1]      [,2]      [,3]
[1,] 0.0000000 0.5611959 0.6908633
[2,] 0.5611959 0.0000000 0.2698923
[3,] 0.6908633 0.2698923 0.0000000
```

4.3.2 Cavalli-Sforza Distance

Another distance approach that is commonly used for microsatellite loci is Cavalli-Sforza distance, D_C (Cavalli-Sforza and Edwards, 1967). Here population allele frequencies are plot on the surface of a sphere (radius=1) using the square root of the allele frequencies.

$$D_C = \frac{2}{\pi} \sqrt{(2 - 2\cos\theta)}$$

The genetic distance, D_C is measured as the chord distance as indicated in Figure ?? . The resulting Neighbor joining tree from this distance is shown in Figure 4.4

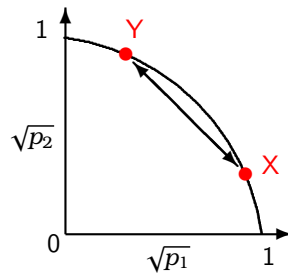


Figure 4.4: Geometry of Cavalli-Sforza distance. Population allele frequencies at two loci are plot at $\sqrt{p_1}$ and $\sqrt{p_2}$ and D_C is the chord between the populations.

```
> d.cavalli <- genetic.distance(sonora,"Pop","EN","Cavalli")
> d.cavalli
```

```
$EN
      [,1]      [,2]      [,3]
[1,] 0.0000000 0.4131725 0.7554523
[2,] 0.4131725 0.0000000 0.5155875
[3,] 0.7554523 0.5155875 0.0000000
```

4.3.3 Nei's Genetic Distance

Nei's genetic distance is based upon mutation drift equilibrium therefore you should be reasonably comfortable with the notion that your populations have been separated a sufficient period of time such that drift and mutation may have played a significant role in their structure.

The formula for Nei's distance that is used here is:

$$D_{Nei} = -\ln \left(\frac{(2N-1) \sum_{i=1}^L \sum_{j=1}^{\ell} p_{ij,x} p_{ij,y}}{\sqrt{\sum_{i=1}^L (2N \sum_{j=1}^{\ell} p_{ij,x} - 1)(2N \sum_{j=1}^{\ell} p_{ij,y} - 1)}} \right)$$

where the summation L is across loci and ℓ is across alleles at each locus in population x and y .

```
> d.nei <- genetic.distance(sonora,"Pop","EN","Nei")
> d.nei
```

```
$EN
      [,1]      [,2]      [,3]
[1,] 0.0000000 1.200027 0.5848357
[2,] 1.2000270 0.000000 2.8444285
[3,] 0.5848357 2.844428      Inf
```

4.3.4 Conditional Genetic Distance

Conditional genetic distance (cGD, Dyer et al. 2010) is a graph-theoretic genetic distance derived from Population Graphs (Dyer and Nason 2004). In some cases it has been shown to be more sensitive to landscape features and heterogeneity in dispersal than structure statistics and other distance metrics (see Dyer et al. 2010).

```
> d.cgd <- genetic.distance(sonora,"Pop","EN","cGD")
```

```

transforming data... done
Rotating mv genos and partitioning... done
[1] "*****"
[1] "matrix"
Estimating conditional genetic covariance... done
Making graph... done

> d.cgd

$EN
      [,1]      [,2]      [,3]
[1,] 0.000000 4.154197 2.477791
[2,] 4.154197 0.000000 2.010668
[3,] 2.477791 2.010668 0.000000

```

4.4 Isolation-By-Distance

Under models with restrictions in gene flow, there is an expectation that genetic distance should increase with physical separation. Using populations found along the Baja Peninsula, it is pretty easy to see which one of these among-strata distance approaches provides a better fit to the data.

```

> baja <- araptus_attenuatus[araptus_attenuatus$Species != "CladeB", ]
> euc <- genetic.distance(baja,"Pop","EN","Euclidean")$EN
> cav <- genetic.distance(baja,"Pop","EN","Cavalli")$EN
> nei <- genetic.distance(baja,"Pop","EN","Nei")$EN
> cgd <- genetic.distance(sonora,"Pop","EN","cGD")$EN

transforming data... done
Rotating mv genos and partitioning... done
[1] "*****"
[1] "matrix"
Estimating conditional genetic covariance... done
Making graph... done

> phys <- stratum.distance(baja,"Pop",lat="Lat",lon="Long")
> df <- data.frame(Euclidean=euc[lower.tri(euc)], Cavalli=cav[lower.tri(cav)], Nei=nei[lower.tri(nei)], cGD
> pairs( df )
> cor(df)

```

	Euclidean	Cavalli	Nei	cGD	Physical.Dist
Euclidean	1.00000000	0.94146219	NaN	0.09490699	0.29356944
Cavalli	0.94146219	1.00000000	NaN	0.08888958	0.27044432
Nei	NaN	NaN	1	NaN	NaN
cGD	0.09490699	0.08888958	NaN	1.00000000	0.02110107
Physical.Dist	0.29356944	0.27044432	NaN	0.02110107	1.00000000

```

> baja <- araptus_attenuatus[araptus_attenuatus$Species != "CladeB", ]
> euc <- genetic.distance(baja,"Pop","EN","Euclidean")$EN
> cav <- genetic.distance(baja,"Pop","EN","Cavalli")$EN
> nei <- genetic.distance(baja,"Pop","EN","Nei")$EN
> cgd <- genetic.distance(sonora,"Pop","EN","cGD")$EN
transforming data... done
Rotating mv genos and partitioning... done
[1] "*****"
[1] "matrix"
Estimating conditional genetic covariance... done
Making graph... done
> phys <- stratum.distance(baja,"Pop",lat="Lat",lon="Long")
> df <- data.frame(Euclidean=euc[lower.tri(euc)], Cavalli=cav[lower.tri(cav)], Nei=nei[lower.tri(nei)], cGD
> pairs( df )
> cor(df)

```

	Euclidean	Cavalli	Nei	cGD	Physical.Dist
Euclidean	1.00000000	0.94146219	NaN	0.09490699	0.29356944
Cavalli	0.94146219	1.00000000	NaN	0.08888958	0.27044432
Nei	NaN	NaN	1	NaN	NaN
cGD	0.09490699	0.08888958	NaN	1.00000000	0.02110107
Physical.Dist	0.29356944	0.27044432	NaN	0.02110107	1.00000000

Figure 4.5: Relationship among strata genetic distance metrics estimated for *Araptus attenuatus* sites in Baja California along with physical distance.

Chapter 5

Genetic Structure

5.1 Synopsis

Estimation of genetic structure is a fundamental process in population genetic analyses. Broadly defined, structure can be defined as the non-random association of genotypes and alleles in populations due to evolutionary processes such as gene flow, drift, selection, and inbreeding. For this, the *Araptus attenuatus* data set and will be used again.

```
> data(araptus_attenuatus)
> baja <- araptus_attenuatus[araptus_attenuatus$Species != "CladeB",]
```

5.2 Genotype Frequencies

The manner by which alleles are arranged into genotypes tells us a lot about the history of a species. The structure statistic that are presented below all rely upon estimation of genotype frequencies so a brief digression to talk about genotype frequencies is in order.

Under a model of random mating, a locus with ℓ alleles whose frequencies are denoted by p_1, p_2, \dots, p_ℓ , homozygotes for the i^{th} allele are expected to occur at a frequency of p_i^2 and ij -heterozygotes are expected at $2p_i p_j$.

The expected frequencies are estimated from the allele frequencies assuming Hardy-Weinberg Equilibrium. If you were only interested in the proportion of heterozygotes, you can use the `ho` and `he` functions.

```
> freq.ltrs <- allele.frequencies(baja, "LTRS")
> he(freq.ltrs$LTRS)*length(baja$LTRS)
```

```
he
161.7324
```

```
> ho(freq.ltrs$LTRS)*length(baja$LTRS)
```

```
ho
69
```

However, at times, it is of interest to look at all genotypes. If you use the `as.character` method for Locus objects, you can easily tabulate the counts of each genotypic state¹.

```
> obs <- genotype.counts(araptus_attenuatus, "LTRS")
> obs
```

```
01:01 01:02 02:02
147    86   130
```

¹This function does take into consideration the non-sorting nature of the Locus object so that a 3:4 locus and a 4:3 locus will be counted as the same heterozygote.


```
> obs/sum(obs)

      01:01      01:02      02:02
0.4049587 0.2369146 0.3581267
```

Below they are denoted as a matrix, the values on the diagonal of `exp` are the expected number of homozygotes and off-diagonal estimates are the expected frequency of heterozygotes.

```
> p <- get.frequencies( freq.ltrs$LTRS )
> p

      01      02
0.5519878 0.4480122

> exp.freq <- p %*% t(p)
> row.names(exp.freq) <- colnames(exp.freq)
> exp <- exp.freq * length(baja$LTRS)
> exp

      01      02
01 99.63379 80.86621
02 80.86621 65.63379
```

As you can see there are fewer heterozygotes than expected ($N_{hets; exp} = 162$, $N_{hets; obs} = 86$).

5.3 Hardy-Weinberg Equilibrium

While the `gstudio` package provides the basic units for population genetic analyses, there are already some very good packages that conduct analyses like testing for Hardy-Weinberg Equilibrium².

```
> require(HardyWeinberg)
> ltrs.genotypes <- genotype.counts( araptus_attenuatus, "LTRS" )
> HWChisq(ltrs.genotypes, verbose=T)
```

Chi-square test with continuity correction for Hardy-Weinberg equilibrium

```
Chi2 = 98.52808 p-value = 0 D = -47.55096
```

```
$chisq
[1] 98.52808
```

```
$pval
[1] 0
```

```
$D
      01:02
-47.55096
```

```
$p
      01:01
0.523416
```

5.4 Structure Parameters

Population structure parameters are fundamental tools for population genetics and have been perhaps, the most poorly understood and misused as well. At the end of this section, some examples of the differences between the parameters is given.

²There are many other functional packages on cran.r-project.org and you should always make sure someone hasn't already solved a problem for you before you try to code up a solution.

These structure parameters are estimated using the function `genetic.structure` and requires a `Population` object, a `stratum`, the loci you want to estimate parameters from, and a `mode` (the parameter you want). If you leave off the `loci` parameter, all loci will be used. There is also an optional parameter, `num.perm` that is used to test significance.

Finally, of note here is that all these parameters use a sample-size corrected estimates of heterozygosity.

$$\hat{H}_S = \frac{2\mu}{2\mu - 1} H_S$$

$$\hat{H}_T = H_T + \frac{\hat{H}_S}{2k\mu}$$

Where μ is the harmonic mean strata size and k is the number of stratum. As you can see as μ gets larger $\hat{H}_S \rightarrow H_S$, which translates to "if you have more samples, you can get a better estimate of the average heterozygosity" and as k get larger, $\hat{H}_T \rightarrow H_T$ which says the same thing about the number of populations. The take-home here is that you need many samples from many places.

5.4.1 The G_{ST} Parameter

The parameter G_{ST} is an estimate of the reduction in heterozygosity due to individuals being in different populations. It is functionally equivalent to F_{ST} from Wright and as he points out, it is not a measure of differentiation in the way that we think of differentiation. Rather it is a measure of the extent to which populations have gone to fixation. It is estimated as:

$$G_{ST} = 1 - \frac{\hat{H}_S}{\hat{H}_T}$$

where H_S is the average expected heterozygosity at each stratum $[1 - \sum_{i=1}^{\ell} p_i^2]/K$ and H_T is the expected heterozygosity across the entire dataset.

For the *EN* locus in the Baja California dataset, G_{ST} is estimated by:

```
> gst.baja <- genetic.structure(baja, stratum="Pop", loci="EN", mode="Gst", num.perm=999)
> print(gst.baja)
```

Geneic Structure Analysis:

```
Estimator: Gst
Stratum: Pop
Loci: { EN }
- EN ; Gst = 0.345786963051191 ; P = 0.001
```

5.4.2 The G'_{ST} Parameter

The parameter G'_{ST} was introduced by Hedrick (20XX) in response to the observation that the parameter G_{ST} is not insensitive to the number of alleles at a locus. Fixing this is done by standardizing the estimate of G_{ST} by the maximal is can be given the number of alleles present, essential a restandardization to the [0,1] range. This is done by:

$$G'_{ST} = \frac{G_{ST}(k - 1 + \mu)}{(k - 1)(1 - \hat{H}_S)}$$

For the same locus, we get a larger

```
> gst.prime.baja <- genetic.structure(baja, stratum="Pop", "EN", mode="Gst.prime", num.perm=999)
> print(gst.prime.baja)
```

Geneic Structure Analysis:

```
Estimator: Gst.prime
Stratum: Pop
Loci: { EN }
- EN ; Gst.prime = 0.459618108931204 ; P = 0.001
```

5.4.3 The D_{EST} Parameter

It has been pointed out that even with the corrections for large numbers of alleles, G'_{ST} may not be acting like a statistic of "differentiation" in the way that we think of differentiation. For example consider the following code where I make three populations, the first one fixed for the "1" allele and the next fixed for the "2" (sure this is an extreme point, but Wright originally made it and it should be repeated).

```
> locus1 <- list()
> for(i in 1:50)
+   locus1[i] <- Locus( c(1,1) )
> for(i in 51:150)
+   locus1[i] <- Locus( c(2,2) )
> strata <- c(rep("Pop-A",50), rep("Pop-B",50), rep("Pop-C",50) )
> pop <- Population(strata=strata, loci=locus1)
> summary(pop)

      strata      loci
Length:150      1:1: 50
Class :character 2:2:100
Mode  :character
```

When we estimate either G_{ST} or G'_{ST} on these data we get:

```
> genetic.structure(pop,"strata","loci",mode="Gst")
```

Geneic Structure Analysis:

```
Estimator: Gst
Stratum: strata
Loci: { loci }
- loci ; Gst = 1
```

```
> genetic.structure(pop,"strata","loci",mode="Gst.prime")
```

Geneic Structure Analysis:

```
Estimator: Gst.prime
Stratum: strata
Loci: { loci }
- loci ; Gst.prime = 1
```

Now, intuitively, if it were just "Pop-A" and "Pop-B" then this would make sense but look at the differences between "Pop-B" and "Pop-C", this should be $G_{ST} = G'_{ST} = 0$! In fact, if you had only one population fixed for the "1" allele and a thousand populations fixed for the other, these parameters would still equal unity. This is because, as Wright originally pointed out, these population parameters are not meant to measure differentiation but fixation. The parameter D_{est} was introduced by Joost (20XX) to address this issue (n.b., Gregorius proposed this back in the 80's but was not taken serious about it then, perhaps Joost can have better luck).

The parameter is defined as:

$$D_{est} = \frac{k-1}{k} \frac{\hat{H}_T - \hat{H}_S}{1 - \hat{H}_S}$$

For the contrived data set, it gives:

```
> genetic.structure(pop,"strata","loci","Dest")
```

Geneic Structure Analysis:

```
Estimator: Dest
Stratum: strata
Loci: { loci }
- loci ; Dest = 0.296296296296296
```

Which is what would be expected, roughly a third, if we have two populations that are identical and one that is differentiated from the rest. I recommend looking at the several papers that go over these issues for more clarity.

For completeness, the results of the Baja California data set, under D_{est} are:

```
> Dest.baja <- genetic.structure(baja,stratum="Pop","EN",mode="Dest",num.perm=999)
> print(Dest.baja)
```

Geneic Structure Analysis:

```
Estimator: Dest
Stratum: Pop
Loci: { EN }
- EN ; Dest = 0.164464814867075 ; P = 0.001
```

5.5 Pairwise Structure

The `genetic.structure` function can also be used to estimate pairwise estimates of each parameter using the optional `pairwise` flag.

```
> sonora <- araptus_attenuatus[araptus_attenuatus$Species=="CladeB",]
> genetic.structure(sonora,"Pop",loci="EN",mode="Gst.prime", pairwise=TRUE)
```

	101	32	102
101	0.0000000	0.4136727	0.3661894
32	0.4136727	0.0000000	0.1245850
102	0.3661894	0.1245850	0.0000000

Chapter 6

Parent Offspring Data

6.1 Synopsis

There are several cases where you have data that consists of both adults and offspring. With these kinds of data, there are some interesting kinds of analyses available for examining structure and diversity. The functionality that `gstudio` provides focuses on translations of offspring data into common formats that can be analyzed using regular routines.

6.2 Getting Data

The use of GoogleDocs as a repository for your data is not unique to parent-offspring data and is used here to demonstrate how to utilize this options. There is a more detailed discussion of how to set up your GoogleSpreadsheets so that you can access them in the `DataImport` vignette. In what follows, I will use the *Cornus florida* data. I split URL (see Data Import) because it was so long it trailed off the page...

```
> data(cornus_florida)
> cornus <- cornus_florida
```

The structure of adult/offspring data is just like any other kind of data and can consist of covariates such as physical location, size, etc. along with strata and loci. The distinction is that there must be *at least* two strata columns:

Individual ID There should a column in the dataset that has identification number or names that are unique to adults. Every adult *must* have a unique identification numver.

Offspring ID To differentiate offspring from adults, the Offspring ID column should have the maternal individual (or paternal if you like) equal to 0. Offspring from this individual have non-zero values for the Offspring ID column. Offspring do not need to all have unique Offspring ID designations, just unique ones within the set of offspring with the same Individual ID.

Here is an example to show the distinctions.

```
> family <- offspring.array(cornus,474)
> family

$mom
  IndID OffID    X    Y      G8      H18      N5      N10      O5
1   474     0 1545 2234 156:164 104:112 126:126 198:200 185:193

$offspring
  IndID OffID    X    Y      G8      H18      N5      N10      O5
1   474     1 1545 2234 164:168 104:112 126:126 198:202 185:195
2   474     2 1545 2234 156:156 102:112 126:126 198:198 179:185
3   474     3 1545 2234 162:164 112:114 124:126 192:198 185:193
4   474     4 1545 2234 164:188 110:112 126:126 194:198 185:193
```

5	474	5	1545	2234	156:158	112:112	126:128	192:198	185:193
6	474	6	1545	2234	164:180	108:112	126:126	188:198	177:193
7	474	7	1545	2234	164:188	110:112	126:126	190:198	177:185
8	474	8	1545	2234	164:168	104:112	126:126	200:202	193:193
9	474	9	1545	2234	156:164	112:112	126:126	190:198	185:193
10	474	10	1545	2234	164:188	110:112	126:126	190:198	177:185
11	474	11	1545	2234	164:168	110:112	126:126	188:200	181:185
12	474	12	1545	2234	164:180	112:112	126:126	190:198	179:193
13	474	13	1545	2234	164:164	112:114	126:126	188:198	181:185
14	474	14	1545	2234	156:180	112:118	126:126	188:200	185:193
15	474	15	1545	2234	156:180	112:114	126:126	190:200	179:185
16	474	16	1545	2234	156:168	104:112	126:126	192:198	193:193
17	474	17	1545	2234	164:164	112:112	126:126	198:200	193:193
18	474	18	1545	2234	156:164	104:114	126:126	198:202	181:193
19	474	19	1545	2234	164:164	112:112	126:128	198:200	185:193
20	474	20	1545	2234	164:168	112:112	126:126	192:198	193:193

Notice that all the offspring from mom '474' have the same IndID and she is differentiated from the offspring by having 'OffID=0'. In fact, all the adults in the dataset can be found as:

```
> adults <- cornus[cornus$OffID==0,]
> adults
```

	IndID	OffID	X	Y	G8	H18	N5	N10	O5
1	226	0	1392	3534	162:180	114:114	124:126	192:192	185:195
2	232	0	1656	3414	158:180	112:112	124:126	184:192	185:185
3	234	0	1718	3330	158:180	112:96	128:128	184:192	185:185
4	300	0	1175	3114	180:188	112:116	126:126	198:200	191:195
5	305	0	1529	3237	154:170	122:124	124:126	188:192	181:195
6	432	0	1336	2748	164:180	114:116	124:126	198:202	185:193
7	433	0	1337	2749	180:188	112:114	126:126	198:202	179:193
8	468	0	1588	2233	164:164	110:116	124:124	198:202	181:193
9	474	0	1545	2234	156:164	104:112	126:126	198:200	185:193
10	484	0	1514	2302	160:168	112:116	126:126	192:192	193:193
11	487	0	1517	2305	164:176	110:112	126:126	192:202	179:181
12	489	0	1519	2307	160:164	104:112	126:126	192:202	179:181
13	490	0	1520	2308	164:176	112:112	128:128	192:202	179:181
14	493	0	1523	2311	168:168	104:112	124:126	192:202	193:195
15	512	0	1174	2279	156:174	108:114	126:126	200:202	179:195
16	513	0	1239	2276	156:180	102:114	126:126	190:198	179:179
17	516	0	1299	2135	156:180	102:114	124:126	190:198	179:179
18	519	0	1357	2148	156:180	104:114	126:126	182:188	181:181
19	520	0	1412	2041	164:172	114:124	126:126	188:198	195:195
20	521	0	1511	1949	160:164	112:112	128:128	198:200	185:193
21	590	0	1880	1040	164:168	112:118	124:126	192:198	177:193
22	607	0	2286	2888	154:164	114:114	126:126	188:202	181:181

6.3 Pollen Pools

Since my research is primarily focused on the analysis of plant populations and mother/offspring combinations provide information about pollen donors, naturally these kinds of analyses will be the first kind to have functionality.

6.3.1 Minus Mom

If you have the collection of offspring and a mother, you can estimate pollen pool allele frequencies as by subtracting the maternal contribution to each genotype and then estimating the allele frequencies of the paternal components (this could

be reversed if you have father/offspring data and need to estimate maternal genotype frequencies just as easily).

```
> offs <- minus.mom( cornus )
> offs
```

	IndID	OffID	X	Y	G8	H18	N5	N10	O5
1	468	1	1588	2233	156	112	126	200	185
2	468	2	1588	2233	180	112	126	198:202	193
3	468	3	1588	2233	188	114	126	198	179
4	468	4	1588	2233	180	112	126	192	185
5	468	5	1588	2233	154	104	126	202	195
6	468	6	1588	2233	154	104	124	188	193
7	468	7	1588	2233	162	114	124	192	185
8	468	8	1588	2233	180	114	126	192	195
9	468	9	1588	2233	180	114	126	202	193
10	468	10	1588	2233	182	114	124	184	195
11	468	11	1588	2233	158	112	128	192	185
12	468	12	1588	2233	158	112	128	192	185
13	468	13	1588	2233	164	108	126	188	191
14	468	14	1588	2233	182	108	126	198:202	181:193
15	468	15	1588	2233	180	112	126	198:202	181:193
16	468	16	1588	2233	164	102	126	198	181:193
17	468	17	1588	2233	164	104	126	188	193
18	468	18	1588	2233	182	108	126	198:202	181:193
19	468	19	1588	2233	182	104	126	202	181:193
20	468	20	1588	2233	182	110	126	198:202	179
21	474	1	1545	2234	168	104:112	126	202	195
22	474	2	1545	2234	156	102	126	198	179
23	474	3	1545	2234	162	114	124	192	185:193
24	474	4	1545	2234	188	110	126	194	185:193
25	474	5	1545	2234	158	112	128	192	185:193
26	474	6	1545	2234	180	108	126	188	177
27	474	7	1545	2234	188	110	126	190	177
28	474	8	1545	2234	168	104:112	126	202	193
29	474	9	1545	2234	156:164	112	126	190	185:193
30	474	10	1545	2234	188	110	126	190	177
31	474	11	1545	2234	168	110	126	188	181
32	474	12	1545	2234	180	112	126	190	179
33	474	13	1545	2234	164	114	126	188	181
34	474	14	1545	2234	180	118	126	188	185:193
35	474	15	1545	2234	180	114	126	190	179
36	474	16	1545	2234	168	104:112	126	192	193
37	474	17	1545	2234	164	112	126	198:200	193
38	474	18	1545	2234	156:164	114	126	202	181
39	474	19	1545	2234	164	112	128	198:200	185:193
40	474	20	1545	2234	168	112	126	192	193

```
> freqs.G8 <- Frequencies(offs$G8)
> freqs.G8
```

Allele Frequencies:

```
156 = 0.0952381
180 = 0.2142857
188 = 0.0952381
154 = 0.04761905
162 = 0.04761905
182 = 0.1190476
158 = 0.07142857
164 = 0.1904762
```

```
168 = 0.1190476
```

Now the distinction should be made that these are the pollen donor allele frequencies since the contribution of the maternal individual has been removed from each offspring, the differences you can see as by comparing the above to:

```
> unreduced.offss <- cornus[cornus$OffID!=0,]  
> freqs.unreduced.G8 <- Frequencies( unreduced.offss$G8 )  
> freqs.unreduced.G8
```

Allele Frequencies:

```
156 = 0.1125  
164 = 0.5125  
180 = 0.1125  
188 = 0.05  
154 = 0.025  
162 = 0.025  
182 = 0.0625  
158 = 0.0375  
168 = 0.0625
```

where the genotype of each offspring has 50% of the mother's genotype.

6.3.2 Genetic Distances and Structure (e.g., 2Gener)

The reduced genotypes can be used in traditional genetic analyses as any other type of genetic data. For example, the Two-Generation Analysis of Pollen Structure (hereafter 2Gener; Smouse *et al.* 2001, Dyer *et al.* 2004) is essentially an AMOVA analysis on pollen donor genotypes. This is a bit of a manual version of it but it can be conducted as (in the next version I'll add a the AMOVA/2Gener options to the `genetic.structure` function).

```
> require(pegas,quietly=TRUE,warn.conflicts=FALSE)  
> D <- genetic.distance(offss,mode="AMOVA")[[1]]  
> D <- as.dist(D)  
> Moms <- as.factor( offss$IndID )  
> amova(D ~ Moms)
```

Analysis of Molecular Variance

Call: `amova(formula = D ~ Moms)`

	SSD	MSD	df
Moms	8.10625	8.106250	1
Error	324.48750	8.539145	38
Total	332.59375	8.528045	39

Variance components:

	sigma2	P.value
Moms	-0.021645	0.486
Error	8.539145	

Variance coefficients:

```
a  
20
```

6.4 Paternity

The `gstudio` package has some basic functionality regarding estimating paternity (or maternity if you have those kinds of data). Thus far, only fractional paternity is implemented and only basically.

Initially,

```
> pollen.freqs <- allele.frequencies( offs )  
> Pexcl <- lapply( pollen.freqs, exclusion.probability )  
> Pexcl
```

\$G8

Pe

0.7225813

\$H18

Pe

0.5868707

\$N5

Pe

0.17702

\$N10

Pe

0.6501088

\$O5

Pe

0.5702125

The multilocus exclusion probability is given by:

$$P_{excl} = 1 - \prod_{i=1}^{\ell} (1 - P_{excl,i})$$

which in R can be found as:

```
> 1- prod((1-unlist(Pexcl)))  
[1] 0.985816
```

Which means that on average, these loci are expected to be able to exclude 98.6% of potential fathers for an mother/offspring pair.

The function `paternity` estimates fractional paternity for a particular mother and set of offspring. Fractional paternity is estimated using multilocus Mendelian transition probabilities for triplet of male parent (MP), female parent (FP), and offspring (O) standardized by the likelihood of all potential fathers.

$$\hat{\pi}_i = \frac{T(O|FP, MP_i)}{\sum_k T(O|FP, MP)_k}$$

This ensures that $\sum \hat{\pi} = 1$. The function `paternity` estimates this for all the offspring within a single family providing the subset of offspring that have potential fathers in the population, the identity of each father, and the fractional likelihood of each father.

```
> pat <- paternity(cornus,474)  
> print(pat)
```

Paternity Analysis:

Family ID: 474

Number of Offspring: 20

Offspring Assigned Paternity: 12

Fractional Paternity (off: dad(prob)):

1:493(1)

13:607(1)

```
> paternity.spiderplot(cornus,pat,X="X",Y="Y", bty="n", xlab="X", ylab="Y")
```

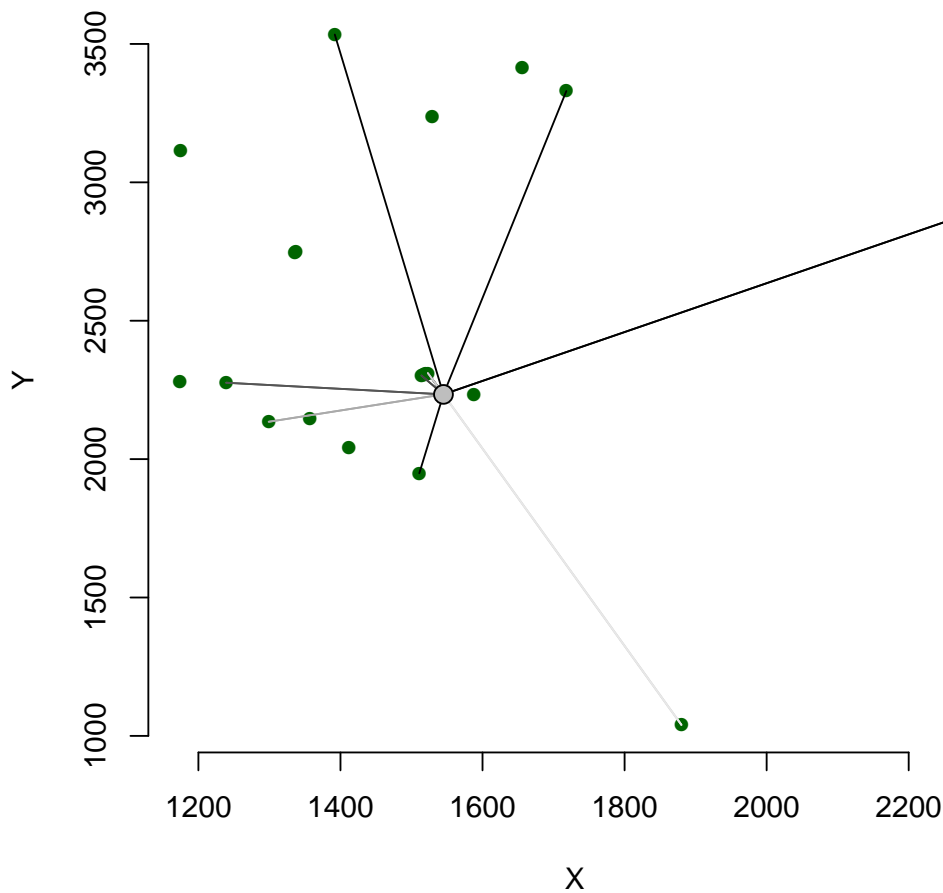


Figure 6.1: Spatial pattern of parentage for family 474 in the *Cornus florida* dataset. Darkness of the lines indicated fractional paternity (light=less, darker=greater).

```
15:513(0.666666666666667) 516(0.333333333333333)
16:484(0.615384615384615) 493(0.307692307692308) 590(0.0769230769230769)
17:474(0.8) 590(0.2)
18:607(1)
19:521(1)
2:513(0.666666666666667) 516(0.333333333333333)
20:484(0.727272727272727) 493(0.181818181818182) 590(0.0909090909090909)
3:226(1)
5:234(1)
8:493(1)
```

You can visualize the results using the `paternity.spiderplot` function that plots the location of all the individuals and indicates putative paternity by connecting mothers and indicated fathers.

```
> paternity.spiderplot(cornus,pat,X="X",Y="Y", bty="n", xlab="X", ylab="Y")
```

Chapter 7

Population Graphs

7.1 Synopsis

A population graph is a topological representation of within and among population genetic variance first introduced by Dyer & Nason (2004). It is particularly well suited to characterizing how spatial genetic variation is distributed among sites.

```
> data(araptus_attenuatus)
> baja <- araptus_attenuatus[araptus_attenuatus$Species != "CladeB",]
```

7.2 Simple Population Graphs

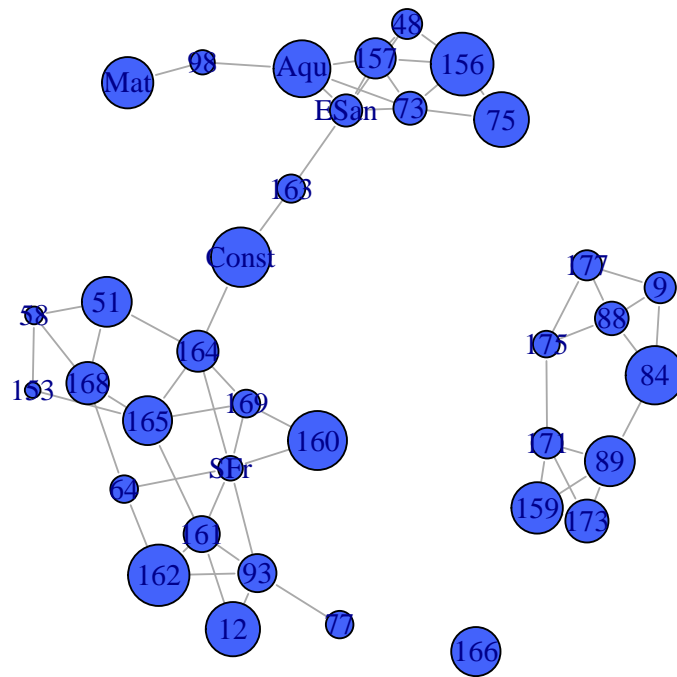
```
> graph <- population.graph(baja, "Pop")

transforming data... done
Rotating mv genos and partitioning... done
[1] "*****"
[1] "matrix"
Estimating conditional genetic covariance... done
Making graph... done

> summary(graph)

Vertices: 36
Edges: 59
Directed: FALSE
No graph attributes.
Vertex attributes: name, size, color.
Edge attributes: weight.

> l <- layout.fruchterman.reingold(graph)
> plot(graph, layout=l, vertex.label=V(graph)$name)
```



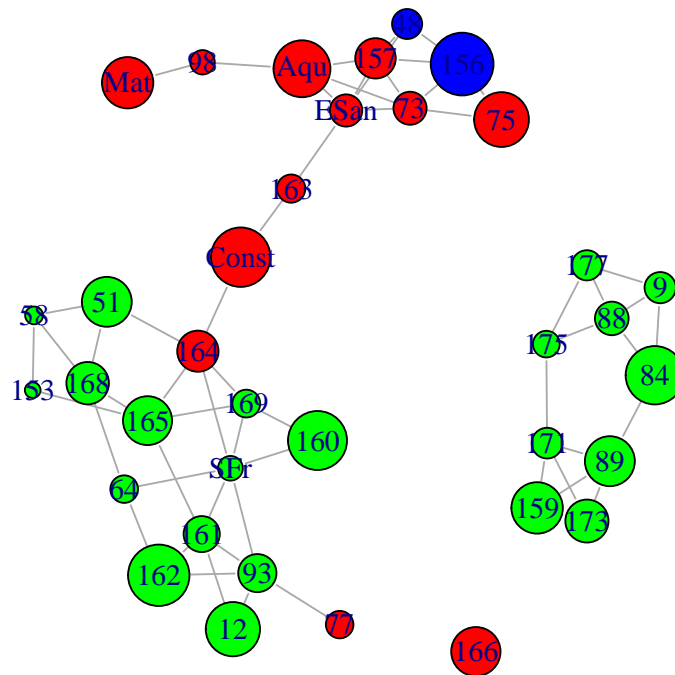
We know that these data are a mixture of two putative species denoted as CladeA and CladeC.

```
> table(baja$Species)
```

```
CladeA CladeC
    75    252
```

We can color the nodes depending upon the identity of clade representation at the node-level. If there is a mixture of species, you would expect to find that the mixed populations would be topologically intermediate between populations made up of pure samples.

```
> getCladeColor <- function(pop,data) {
+   inds <- data$Species[data$Pop==pop]
+   levels.inds <- levels(as.factor(as.character(inds)))
+   if(length(levels.inds)==2) return("red")
+   else if( levels.inds=="CladeA" ) return("blue")
+   else return("green")
+ }
> colors <- unlist(lapply(V(graph)$name, function(x) getCladeColor(x,baja)))
> plot(graph,layout=1,vertex.label=V(graph)$name,vertex.color=colors)
```



So if we only use the samples from CladeC we may be actually analyzing the data in a way that makes sense. Do this by:

1. Use only the CladeC individuals
2. Get rid of the populations with say $N < 5$ individuals
3. Make graph and examine the topology

```
> baja.cladeC <- baja[baja$Species=="CladeC",]
> inds.per.pop <- lapply( partition(baja.cladeC,"Pop"), function(x) dim(x)[1] )
> ## Examine inds per pop to figure out which have <5 individuals save in smPops
> smPops <- c("Const","ESan","157","73","Aqu","Mat","98","75")
> baja.cladeC <- baja.cladeC[ !(baja.cladeC$Pop %in% smPops) , ]
> graph.cladeC <- population.graph(baja.cladeC,"Pop")
```

transforming data... done

Rotating mv genos and partitioning... done

```
[1] "*****"
```

```
[1] "matrix"
```

Estimating conditional genetic covariance... done

Making graph... done

```
> summary(graph.cladeC)
```

Vertices: 26

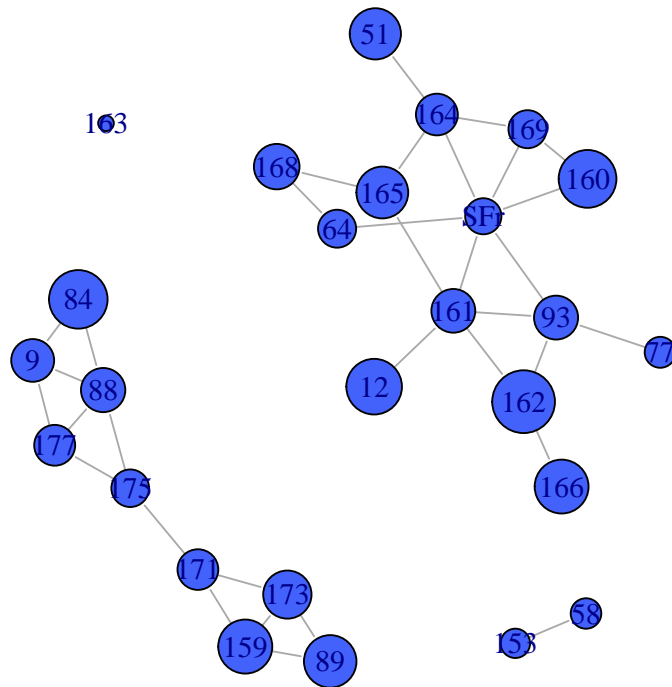
Edges: 33

```

Directed: FALSE
No graph attributes.
Vertex attributes: name, size, color.
Edge attributes: weight.

> l <- layout.fruchterman.reingold(graph.cladeC)
> plot(graph.cladeC,layout=l,vertex.label=V(graph.cladeC)$name)

```



From this plot, you can see even when we only focus on the true CladeC individuals, there is still partitioning of genetic covariance!

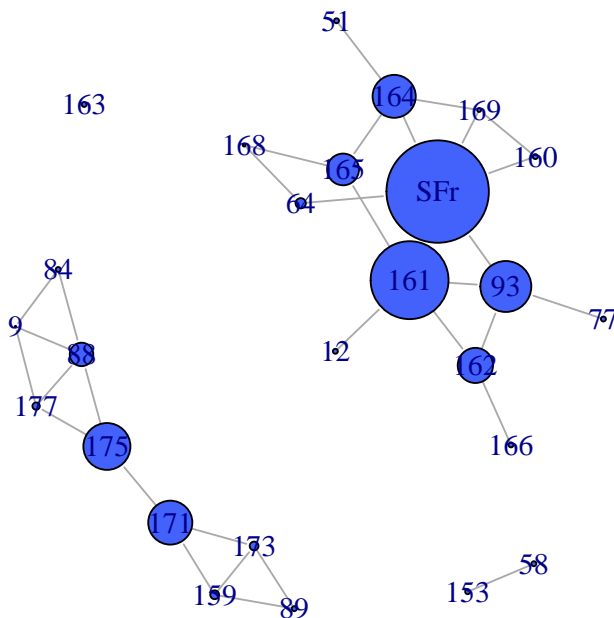
7.3 Node Position

Both node and edge position in the topology can easily be determined using common network analysis tools. The *igraph* package has some as does the most excellent *sna* package. Here is a quick example where the size of the node is depicting the node's betweenness (e.g., the number of shortest paths that go through that node).

```

> pop.betweenness <- betweenness(graph.cladeC,directed=F)
> plot(graph.cladeC,layout=l,vertex.label=V(graph.cladeC)$name,vertex.size=pop.betweenness)

```



Which is rather interesting since betweenness can be used to classify relative population importance. Presently, it is common to use genetic diversity as a surrogate to identify populations of high conservation importance, but betweenness relates to the connectivity of the gene flow topology on the landscape and is not necessarily correlated with genetic diversity.

```
> cor.test(V(graph.cladeC)$size, pop.betweenness, method="spearman")
```

Spearman's rank correlation rho

data: V(graph.cladeC)\$size and pop.betweenness

S = 3259.634, p-value = 0.5779

alternative hypothesis: true rho is not equal to 0

sample estimates:

rho

-0.1144049

7.4 Conditional Genetic Distance

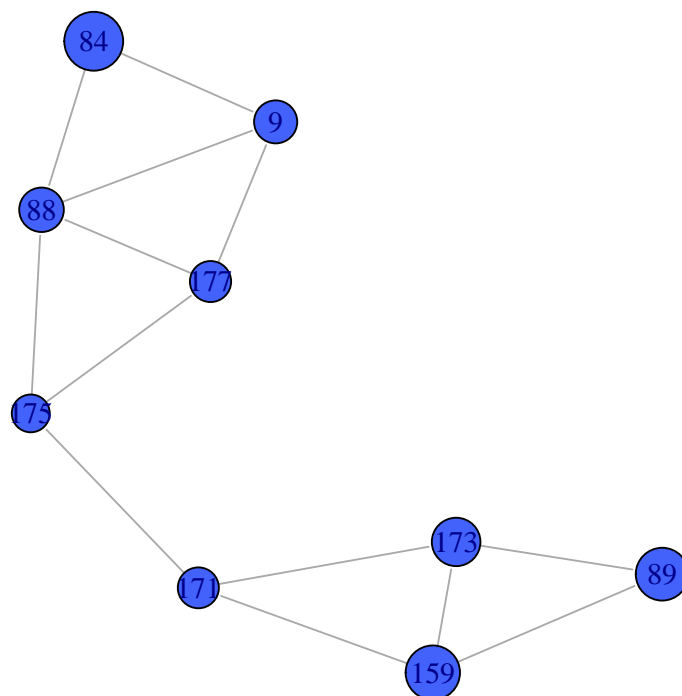
In Dyer *et al.* (2010) we showed that graph distance (e.g., the shortest path connecting points in the topology) was more powerful than pair-wise structure and distance approaches. We denoted the among population distance as *cGD* for conditional graph distance.

Since this topology is disconnected, we'll just focus on the medium sized component, the one with 84 in it.

```

> connected.to.84 <- subcomponent(graph.cladeC,v="84")
> med.graph <- subgraph(graph.cladeC,v=connected.to.84)
> med.layout <- layout.fruchterman.reingold(med.graph)
> plot(med.graph,layout=med.layout,vertex.label=V(med.graph)$name)
> D <- shortest.paths(med.graph)

```

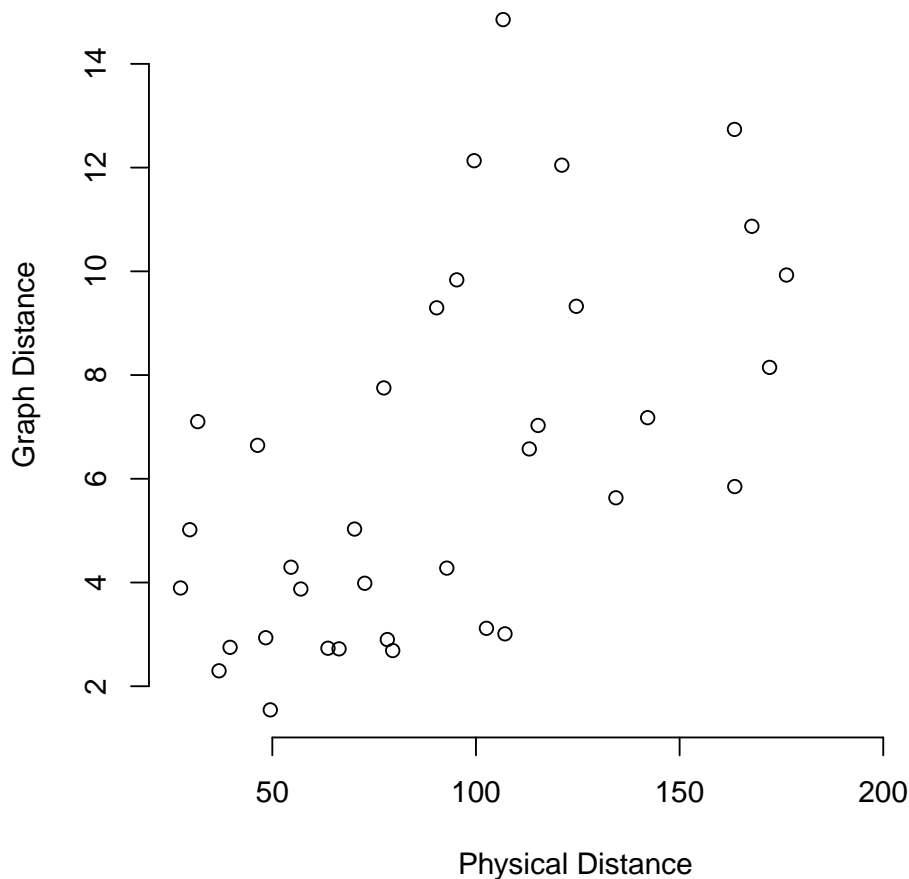


As discussed previously, we can also get the pair-wise physical distance and then examine "Isolation by Graph Distance" (IBGD), which has some nice properties that make it perhaps more precise than IBD based upon pair-wise structure estimates.

```

> pops <- V(med.graph)$name
> P <- stratum.distance(baja.cladeC,"Pop",lat="Lat",lon="Long",subset=pops)
> plot(D[lower.tri(D)] ~ P[lower.tri(P)], bty="n",xlab="Physical Distance",ylab="Graph Distance")

```

We can use a Mantel test to see if there is a correlation between graph and physical distance for this subcomponent.

```
> require(ecodist,quietly=T)
> mantel(as.dist(D)~as.dist(P)) ##pval3 is Ho: Mantel-R=0

      mantelr      pval1      pval2      pval3      llim.2.5%      ulim.97.5%
0.5687480  0.0100000  0.9910000  0.0100000  0.4855738  0.7048877
```

The pval3 is the probability of $H_0 : Mantel\rho = 0$.

7.5 Graph Partitions

A very important point needs to be made here regarding subgraphs and partitions of the whole data set. The disconnected subgraph in the previous section is not necessarily the same graph you would get if you partitioned the genotypes into only those populations and then make the graph. Compare the previous network topology to this one.

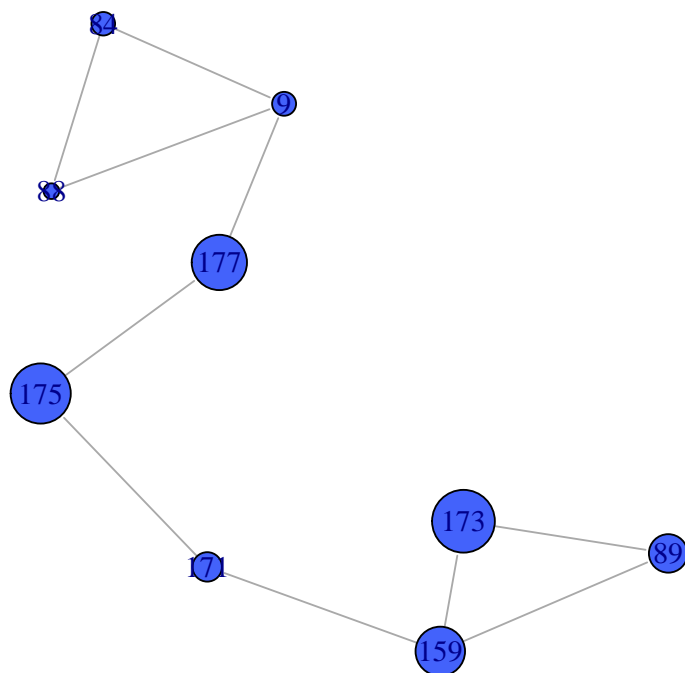
```
> tmp.pop <- baja[baja$Pop %in% c("9","84","88","89","159","171","173","175","177")]
> tmp.graph <- population.graph(tmp.pop,"Pop")
```

transforming data... done

Rotating mv genos and partitioning... done

```
[1] "*****"
[1] "matrix"
```

```
Estimating conditional genetic covariance... done
Making graph... done
> plot(tmp.graph,layout=med.layout,vertex.label=V(tmp.graph)$name)
```



This is because Population Graphs are constructed using *Conditional Genetic Covariance*. The genetic covariance between populations 173 & 171 is conditional on the their covariance with all the other data in the data set. In the first graph this includes the populations in this subgraph as well as the populations outside the subgraph.

Chapter 8

Mapping Population Genetic Data

8.1 Synopsis

This vignette goes over some of the methodologies available for creating google KML files to display aspects of genetic data in either Google Earth, Google Maps, or even as an import into Arc. These functions are part of an extension package `gstudio-sp` that extends the `gstudio` package by adding spatial components. They are kept separate from the rest of the `gstudio` package because one may not need to use the spatial components every time.

Here the *Araptus attenuatus* data set will be used and in particular the subset of populations that formed the disconnected subgraph in the *Population Graphs* vignette from the `gstudio` package.

```
> data(araptus_attenuatus)
> popsToKeep <- c("88", "9", "84", "177", "175", "173", "171", "89", "159")
> baja <- araptus_attenuatus[araptus_attenuatus$Pop %in% popsToKeep]
```

8.2 Pies On Maps

Often it is of interest to look at global changes in allele frequencies. While it is true that the frequency of an allele or set of alleles can be plot as a function of latitude or longitude, there is also value in putting it on a map. The function `pies.on.map` takes a Population file, a stratum, a list of loci, and some coordinate names in the population. In most of the functions in `spgen` if you have your latitude and longitude variables labeled "Latitude" and "Longitude", you do not need to specify them in the function call.

```
> pies.on.map(filename="~/Desktop/Baja.pies.kml",pop=baja,stratum="Pop",loci=c("EN", "LTRS"),lat="Lat",lon="
```

This creates a KML file that you can open in GoogleEarth and looks something like Figure 8.1

8.3 Population Graphs On Maps

It is also helpful to put graph topologies on a map. Here a population graph is created using the wrapper function `spatial.population.graph`. This function adds latitude, longitude, and colors as properties to a normal population graph and is required for spatial plotting. You can add these properties yourself if you like (use the `list.vertex.properties` function to see what is different) to a normal graph or you can just make the graph using this function.

```
> graph <- spatial.population.graph(pop="baja",stratum="Pop",lat="Lat",lon="Long")  
> popgraph.on.map(graph,filename="~/Desktop/popgraph.on.map.kml")
```



Figure 8.2: Population graph for the northern group of *Araptus attenuatus* populations.

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