

Reading and Writing

- Input data from files
- Read data from a program
- Print data into a file or print data out to a program

Input/Ouput

```
1 open(IN, "input.txt") || die $!;
2 # read a line in
3 my $line = <IN>;
4 # read the whole file
5 while(<IN>) {
6  my ($col1, $col2) = split;
7 }
8 close(IN);
```

Filehandles

Filehandles can also be stored in variables

```
1 my $fh;
2 open($fh => "gene.dat") || die $!;
3 while(<$fh>) {
4  print $_;
5 }
6 #I like to use this in one line
7 open(my $fh2 => "gene2.dat") || die $!;
8 while(<$fh2>) {
9  print $_;
10 }
```

Writing to a file

One can also write data out to a file

```
1 open(my $fh => ">outputfile.txt") || die $!;
2 print $fh join("\t", qw(NAME RANK TOWN)), "\n";
3 print $fh join("\t", qw(GRIFFITH SHERIFF MAYBERRY)), "\n";
4 print $fh join("\t", qw(RAWLS DEPUTY BALTIMORE)), "\n";
5 close($fh);
```

Data embedded in a script

```
1 while(<DATA>) {
2 my ($col1,$col2) = split(/\s+/,$_);
3 }
4
5 __DATA__
6 Color Size Model
7 red 10 Jumbo
8 yellow 8 Large
9 pink 2 Mini
```

Pipes for processes

You can combine operations that are on the command line with the I operator in UNIX. This can also be used in Perl when specifying an open command to actually run a program and have the output sent back to the Perl program. This can be used to obtain data from a program. For example here we run the grep cmmand to find lines in a file that have the string 'gene'. Only those lines will be returned and thus be seen by the Perl program.

```
1 open(IN,"grep 'gene' gene.dat | ") || die $!;
2 while(my $line = <IN>) {
3  print "line is $line\n";
4 }
```

Or it can be used to send data to a program. For example this script will print out data to a program which will then compress the data. Notice how the pipe comes at the beginning because we plan to send data into the program.

```
1 open(my $fh => "| gzip -c > file.gz") || die $!;
2 print "hello there\n";
3 print "can you see this?\n";
```

Now look at the file in your directory. It is compressed – you can read it with more and see it is binary file. However you can read it with zmore or you can uncompress it with gunzip.

Pipe trick

Can use it to open a compressed file on the fly.

```
1 open(my $fh => "zcat file.gz |") || die $!;
2 while(<$fh>) {
3  # process data in a file that was compressed, without making a new copy of the file as un 4 }
```

Read data from the web with cmdline

```
1 my $url = 'http://eutils.ncbi.nlm.nih.gov/entrez/eutils/efetch.fcgi?db=nucleotide&id=1636
2 # -S option will not print any statistics
3 open(my $fh => "curl -S '$url' |") || die $!;
4 while(<$fh>) {
5    print $_;
6 }
7
8 open($fh => "GET '$url' |") || die $!;
9 while(<$fh>) {
10    print $_;
11 }
```

Let's try this together

Login to biocluster, Download data files. Data are in this http://courses.stajich.org/public/gen220/data/ which represent some time points in growth for a fungus.

http://courses.stajich.org/public/gen220/data/Nc20H.expr.tab http://courses.stajich.org/public/gen220/data/Nc3H.expr.tab

```
(on biocluster)
wget http://courses.stajich.org/public/gen220/data/Nc3H.expr.tab
```

Write a script to read in the Nc20H and Nc3H data into a hash (one hash for each datafile). Store in the hash the gene name (the 1st column) and the FPKM data. Each gene will appear once in each file.

- Print out a new file which has the gene name, the expression in 3H and the expression in 20H.
- Extra print it out so that it is sorted by the 3HR timepoint

A solution

```
1 use strict;
 2 use warnings;
 3 my (%expr3H,%expr20H);
 4 open(my $fh => 'Nc3H.expr.tab') || die $!;
 5 while(<$fh>) {
6 my @row = split("\t",$_);
7 next if $row[0] eq 'gene_id'; # skip when it is the header line
 8 \frac{9}{9} $\text{expr3H}\frac{1}{9} = \frac{1}{9}
 9 }
10
11 open($fh => 'Nc20H.expr.tab') || die $!;
12 while(<$fh>) {
13 my @row = split("\t", \$_);
14 next if $row[0] eq 'gene_id'; # skip when it is the header line
15 \frac{9}{7} = \frac{9}{7} = \frac{15}{7}
16 }
17
18 open(my $outfh => ">Combined.tab") || die $!;
19 my $gene;
20 for $gene ( keys %expr3H) {
21 print $outfh join("\t", $gene, $expr3H{$gene}, $expr20H{$gene}), "\n";
22 }
23
24 open($outfh => ">Combined_sorted.tab") || die $!;
25 for my qene (sort { $expr3H{$b} <=> $expr3H{$a} } keys %expr3H) {
26 print $outfh join("\t", $gene, $expr3H{$gene}, $expr20H{$gene}), "\n";
27 }
```

References

Reference are ways to refer to a complicated data structures as a single, scalar value. This lets one pass around multiple arrays and they stay separate. We also primarily use reference to store multiple things in a slot in an array.

- Reference to an array is done with \ or []
- Reference to a hash is done with \ or \{\}

For example this lets one pass around multiple arrays and they aren't flattened into one. Consider this code.

```
1 my @array1 = qw(A B C D);
2 my @array2 = qw(W X Y Z);
3 my @array3 = (@array1, @array2);
4
5 print join(",", @array3), "\n";
```

Storing multiple items in an Array

```
1 my $url = 'http://courses.stajich.org/public/gen220/data/Ncrassa_OR74A_InterproDomains.ta
2 open(my $fh => "GET $url |") || die $!;
3 my %genes;
4 while(<$fh>) {
       my ($gene,$domain, $domain_name, $start,$end,$score) = split;
6 # store an array as the value for each key by making it a reference to an array
7 # using the @{$genes{$gene}} which is forcing what is the value
8 # to be an array reference. Then we use push to add something to
9 # this array
10 # Because perl will automatically initialize the value, based on the context
11 # we DON'T need to do this, but it is what is happening under the hood
12 # if this is the first time accessing this key
13 # qenes{qene} = \Gamma;
14 push @{$genes{$gene}}, $domain_name;
15 }
16 # now unpack to print this out
17 for my $gene ( keys %genes ) {
       my @domains = @{$qenes{$qene}};
19 print join("\t", $gene, join(",", @domains)), "\n";
20 }
```

Subroutines

```
1 sub a_routine {
2 my @args = @_; # the arguments passed in are avaialable as @_;
3 print "the arguments are ", join(",", @args), "\n";
4 }
5 &a_routine('a','b','c');
```

Command line arguments

```
$ perl myscript.pl A B C

!perl
print join(",", @ARGS), "\n";
print "the first argument is ", $ARGS[0], "\n";

A,B,C
```

Use this to specify a data file to read in, or specific options you want to run.

Practice Write a script that will open and print out the first 5 lines of a file. The name of the file to open should be passed in on the command line as the first argument.