The Hitchhikers Guide to Venom Gland Transcriptomics

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Introduction

The Hitchhiker's Guide to Venom Gland Transcriptomics is part of Venomancer and designed to walk you through our recommended bioinformatic pipeline. If you are new to bioinformatics, don't panic! There are some resources and information provided at the beginning of the document to get you through the basics. Overall, The topics covered in this guideline include:

- Basic bioinformatics
- Raw data processing
- Transcriptome assembly
- Transcriptome annotation
- Checking for chimeric transcripts
- Clustering redundant transcripts
- Checking for rogue/missing transcripts
- Proteomic annotation
- Estimating expression
- Post-processing visualization
- Differential expression analysis

Basic Bioinformatics

This first section is largely for people who don't have much experience with unix, python, R, or bioinformatics in general. While this guide is not designed to teach you all of these things, we do recommend you check out the resources below. If you do have experience in unix, depending on your level of experience, there might still be something new we can teach you.

Training Resources

Clemson University provides several freely-available workshops to get you started learning unix, python, R, and more:

- Clemson Computing and Information Technology Resources
 - Linux Workshop
 - Python Workshop
 - R Workshop
 - Genomics Workshop

In addition, some other useful resources might be:

- SWIRL: Learn R in R
- Learn-Bioinformatics Resources

STOP

Again. Don't panic. But from here on, we assume that you have basic knowledge of unix. If you do not, please return to the workshops above. Otherwise continue on...

Lists

Lists are your best friend in coding. A list – very simply – is a text file containing all the names of your samples or identifiers that you will want to loop through or process in parallel. You can create a text file with the command nano {file}.txt and save the file using keyboard shortcut ctrl+o and exit with ctrl+x. A list will look something like this:

```
sample_1
sample_2
...
sample_n
```

You can also create lists by simply saving the output of 1s to a file. For example:

```
ls *.fastq > list.txt
```

Once you create a list, you can provide it to a for loop or to the parallel command to process many samples the same way – and even simultaneously – without actually having to redo the same command over and over again. For example:

```
for i in 'cat list.txt'
do echo ${i}
done

# OR
parallel -a list.txt "echo {}"
```

We will be providing lists to the command parallel in nearly every step of the Hitchhiker's Guide.

Piping & Regular Expressions

Piping – initiated with a | – is simply taking the standard OUTPUT of one command and feeding it into the standard INPUT of another. This can be useful for doing several commands in a row. Below, you will see that I use piping to create 3 different lists based on the fastq files I have in a directory.

The first list I create is a list of the unique individuals which we will use throughout the pipeline. In this list, each individual has it's own line. The second two lists remove new line characters and replace them with spaces so that all individuals are on a single line.

```
 ls *.fastq.gz | cat | perl -pi -e 's/_.*.fastq.gz//g' | uniq > list.txt \\ sed "s/$/_R1.fastq/g" list.txt | tr '\n' ' ' | sed '/^\s*$/d' > list2.txt \\ sed "s/$/_R2.fastq/g" list.txt | tr '\n' ' ' | sed '/^\s*$/d' > list3.txt
```

Based on those lines of code, hopefully you understand that I am taking the output of the first command and feeding it into subsequent commands. However, you may be asking... what are perl -pi -e, sed, and tr?

Hopefully at some point, you have learned about grep and it's ability to search and print for specific text in a document. What you may not know is that grep stands for Global Regular Expression Print. grep, perl -pi -e, sed, tr, and awk all use something called Regular Expressions or regex to find and/or replace text. I've used 3 different find/replace methods to generate my lists, but they are all essentially doing the same thing. regex can take some getting used to, but they are used VERY frequently and are incredibly useful in unix scripting. So take some time to learn them!

Useful regex resources:

- https://www.rexegg.com/regex-quickstart.html
- https://regexr.com/
- https://awk.js.org/
- https://sed.js.org/

Each of the different methods have own features and intricacies; therefore, you may have to change between them. However, you will pick your favorite (whichever you learn first) and use it most frequently. My personal preference and the option I am most familiar with is perl -pi -e.

Renaming Files

Sometimes files might have something you don't want or you need to rename them in some way. If it is only one file, a simple mv command will do the trick. However, when there are a lot of files, renaming all of them can be a pain. Below, I provide three options for removing an unwanted underscore in all my file names.

Option 1: The mv strategy

```
for i in sample_*.fastq.gz
do mv "${i}" "${i/sample_/sample}"
done
```

Option 2: The find and rename strategy

```
find . -name "sample_*" -exec rename "sample_" "sample" {}\;
```

Option 3: The bash script strategy

This strategy take a little more manual work, but is useful when you have less regular patterns to change, where each individual gets a slightly different name (e.g adding the species code which is found in a spread-sheet somewhere). In this situation, let Microsoft Excel help you.

To do this, create a list of all your individuals that need renamed. You can then copy-paste your list into Microsoft Excel to use things like find-replace and VLOOKUP to add a column including the species code. You

can also create a column to copy repeat elements like mv efficiently across all your individuals. Concatenate all your columns together and move it into a text document like the example below. Once you create your text bash script, you can just run sh script.sh to do all the work for you.

```
#!/bin/bash

mv sample_001.fastq.gz sample-001.fastq.gz
mv sample-2.fastq.gz sample-002.fastq.gz
mv sample_3.fastq.gz sample-003.fastq.gz
#mv [MORE INDIVIDUALS]
```

Bash Profile

Each step of the Hitchhiker's Guide assumes that you have already installed all the programs that you need and it can be easily run by just typing in the appropriate command (i.e., trinity). Lets talk about how to make that assumption true.

First, in your home directory (cd ~) you will have a hidden file called your .bash_profile. Remember, to see hidden files you have to use the ls -a command. If you do not have a .bash_profile in your home directory, go ahead and create one with nano .bash_profile.

If you don't already know this, your .bash_profile is what is loaded every time you open a terminal/shell. It contains your specific configuration settings including your \$PATH, which is a variable set to all possible directories where your installed programs exist. My .bash_profile looks like this and may be a good starting point for you:

```
# Bash Profile

# User Profile
PS1="[Rhett@Macbook: \W] $ "

# SSH Profiles
alias remote="caffeinate ssh username@login.remote.server.edu"

# Alias Functions
alias 11="ls -lh"
alias d="conda deactivate"
alias bio="conda activate bio"
alias envs="conda info --envs"

# PATH
export PATH=/usr/local/bin:/usr/bin:/usr/sbin:/sbin
export PATH="$PATH:~/path/to/bin:~/path/to/bin/scripts"
```

From my .bash_profile, my computer knows that when I start a terminal/shell it should print out [Rhett@Macbook: ~] \$ at the beginning of each line, that each alias is a shortcut that does a specific command, and that it needs to export \$PATH as: ~/path/to/bin:~/path/to/bin/scripts:/usr/local/bin:/usr/bin:/usr/sbin:/bin:/sbin. This string represents the location of several folders on my computer where scripts, commands, and programs that I have manually installed may be found.

As you may have noticed, we still haven't installed any programs. We've only told the computer where to find programs. There's a long chunk of code that I didn't show you in my .bash_profile. What is says is unimportant, but it has to do with Anaconda. Anaconda is a package/software management toolkit which creates self-contained environments and installs programs within them. It also edits your .bash_profile

for you! Therefore, I do most program installation through Anaconda, which we will talk about in the next section...

Setup/Installation

Anaconda is a package/software management toolkit. I highly recommend you install Anaconda to make your life easier and avoid having to install many programs by hand.

When you install Anaconda, it will automatically add something to your .bash_profile which will enable the computer to find programs installed by Anaconda. The reason Anaconda is so great is that software often do not play nicely for one another. For example, while one program may want to use python v2.7 another may want to use python v3.5. Anaconda allows you to create self-contained environments within which everything works nicely! If something needs a different version of a program...simply create a new environment! Here's a cheatsheet for using Anaconda.

After you install Anaconda, we can create some environments for later use. Here's how you do that:

```
# You do not need to do this every time, but this sets up what "channels" anaconda should look into for
# Configure conda to look in certain channels for packages
conda config --add channels bioconda
conda config --add channels conda-forge
conda config --add channels defaults
# Install a couple very useful packages in your base conda environment
conda install wget
conda install parallel
# Create a bioinformatics environment
conda create -n bio # Create an environment named "bio"
conda activate bio # Activate your "bio" environment
conda install -y biopython bamtools bedtools blast bowtie2 bwa cd-hit emboss fastqc gatk4 jellyfish par
conda deactivate # Deactivate your environment to exit
# Create a Trinity environment
conda create -n trinity_env trinity parallel
# Create a Venomancer environment
conda create -n venomancer_env python=3.7 biopython perl perl-bioperl perl-mce blast parallel
# Create a ChimeraKiller environment
conda create -n chimerakiller_env python=3.6 biopython bwa samtools bedtools picard pandas matplotlib s
```

Notice that I created a separate environment for Trinity, Venomancer, and ChimeraKiller. This is because programs often depend on different versions of other programs. For example, Trinity requires a different version of samtools than most other things installed in the bio environment. Putting things in their own environment ensures that they won't interfere with each other. But how do you know when to create a new environment vs. just add to a pre-existing environment?

Anaconda will generally warn you if it needs to change the version of something and will ask for confirmation to do this before continuing. When in doubt, create a new environment. You can create as many environments as you want and if something isn't working properly, just create a new environment for it or remove the old environment and start over. All you have to do is remember to conda activate the correct environment before running an analysis.

Whenever possible, try Googling conda install {program} first. However, sometimes it is not possible to install things via Anaconda and you will have to manually download, install, and add the install location to your PATH. I recommend creating a bin folder in an easily accessible location and installing all programs in this folder. For example, I set up my bin folder in ~/Dropbox/bin so that it can be accessed by any of my computers with Dropbox connected. You just need to make sure the \$PATH is located in you .bash_profile on that computer.

Below, we download Venomancer, CodAn (a Venomancer dependency), and signalp and add them to our \$PATH. I also recommend running the tutorial for each program to ensure it is installed properly.

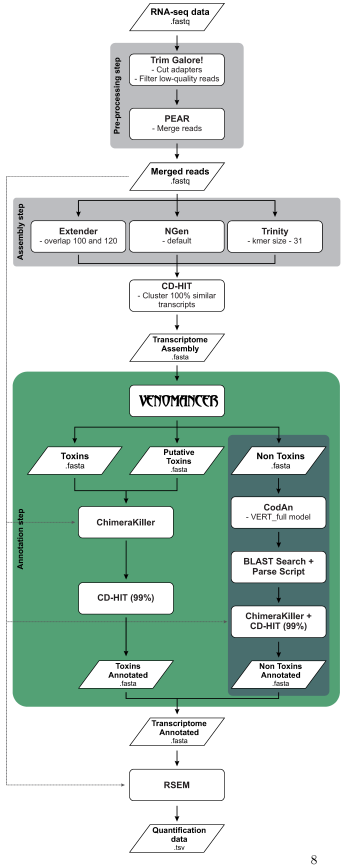
```
# Git clone the Venomancer repository and add to your PATH:
git clone https://github.com/pedronachtigall/Venomancer.git
echo 'export PATH=$PATH:path/to/Venomancer/bin/' >> ~/.bash_profile

# Git clone the CodAn repository and add to your PATH:
git clone https://github.com/pedronachtigall/CodAn.git
echo 'export PATH=$PATH:path/to/CodAn/bin/' >> ~/.bash_profile

# Download the SignalP-4.1, decompress and add it to your PATH:
tar -xzf signalp-4.1g.Linux.tar.gz
echo 'export PATH=$PATH:path/to/signalp-4.1/' >> ~/.bash_profile

# You may want to source your .bash_profile or re-open your shell before continuing
source ~/.bash_profile
```

The Hitchhiker's Guide



Downloading Data

We will use the *Bothrops alternatus* data as an example for the Hitchhiker's Guide. You can download these data with wget or use fastq-dump from the sra-toolkit.

```
conda activate bio
fastq-dump -I --split-files SRR######
```

NOTE: The -I represents a flag or argument for the command fastq-dump. We will not explain every option for every command, but an explaination can usually be found for each possible flag/argument using {program} -h or man {program}.

Pre-Processing

With the data downloaded, let's get organized by creating a folder for each sample and a folder for the raw data.

Generally our file structure will look like this:

```
|-- sample_001
   |-- 01_rawdata
   | |-- sample_001_R1.fastq.gz
   | |-- sample_001_R2.fastq.gz
   |-- 02_trimmed
      |-- sample_001_trimmed_R1.fastq.gz
       |-- sample_001_trimmed_R2.fastq.gz
    |-- 12_transcriptome
       |-- sample 001 transcriptome v1.fasta
|-- sample_002
    |-- 01 rawdata
      |-- sample_002_R1.fastq.gz
      |-- sample_002_R2.fastq.gz
    |-- 02 trimmed
      |-- sample_002_trimmed_R1.fastq.gz
      |-- sample_002_trimmed_R2.fastq.gz
    |-- 12_transcriptome
        |-- sample_002_transcriptome_v1.fasta
```

Here, I am using the parallel command to with the -a flag to provide it a list of our samples and the -j flag to tell it how many jobs to do at a single time. This may need to be modified based on your computer.

```
parallel -a list.txt -j 2 --verbose "echo {}
  mkdir {} {}/01_rawdata
  mv {}*.fastq.gz {}/01_rawdata"
```

Check Quality (fastqc)

fastqc is used to check the quality of your sequencing reads. You may want to do this several times throughout the process. Primarily before and after trimming low-quality bases and reads.

```
parallel -a list.txt -j 2 --verbose "echo {}
  cd {}/01_rawreads
  fastqc {}*.fastq.gz
  open *.html"
```

Trimming (Trim-Galore)

The sequencer provides quality scores (also known as phred scores) that indicate how good a given base call was. We obviously want to trim off reads that have poor quality. We generally use trim_galore to do this, but TRIMMOMATIC is another option. We can run fastqc right after to see the difference trimming makes in our data.

```
parallel -a list.txt -j 2 --verbose "echo {}
  cd {}
   mkdir 02_trimmed
   trim_galore -paired -phred33 -length 75 -q 5 -stringency 1 -e 0.1 -o 02_trimmed 01_rawdata/{}_R1.fa
  cd 02_trimmed
  mv {}_R1_val_1.fq.gz {}_R1_trim.fastq.gz
  mv {}_R2_val_2.fq.gz {}_R2_trim.fastq.gz
  fastqc *.fastq.gz
  open *.html"
```

Merging Reads (PEAR)

With our reads trimmed, we can merge our forward and reverse reads to create slightly longer "super reads" which will be better for assembly and easier to manage. To do this, we use the program PEAR

```
parallel -a list.txt -j 2 --verbose "echo {}
  cd {}
  mkdir 03_merged
  pear -j 4 -f 02_trimmed/{}_R1_trim.fastq.gz -r 02_trimmed/{}_R2_trim.fastq.gz -o 03_merged/{}_pear
  cd 03_merged
  pigz *.fastq
  pigz -d {}_pear.fastq.gz"
```

Transcriptome Assembly

The next step is to assemble our reads into contigs. Although the process is a little more complex than this, you can think of it as taking our short 150-250 bp fragments and lining them up end-to-end to determine what goes together. This isn't necessary if you have a reference genome as you can simply take your reads and map them back to your reference with something like HiSat2. However, if we don't have a reference genome, the process of assembly and annotation can be extremely challenging.

For de novo transcriptome assembly, we use a combination of three different assemblers to produce our transcriptomes, these include Extender, NGen, and Trinity. You are welcome to use even more assemblers or different assemblers; however, we chose these assemblers because they use three different ways to assemble the transcriptome. This pipeline and how these assemblers were chosen can be read about in Holding et al. 2018. Briefly, typical transcriptome assemblers (e.g. Trinity) use a method known as de Bruijn graphs while Extender and NGen use much simpler methods. However, these simpler methods are really good at catching longer transcripts – such as Snake Venom Metalloproteases (SVMP) – that are frequently missed by Trinity

Extender

Extender is a pretty straightforward assembler which does not use de Bruijn graphs. Developed by Rokyta et al. 2012, Extender essentially grabs 'seed' reads from the reads pile and then uses a sliding window approach to see if it can extend these seeds on either end. While we don't want to get into the mechanisms of Extender too much, we do want you to know a couple of things about the outcome of using it. The first is that Extender is really good at assembling large transcripts like metalloproteases. In fact, it is probably the best assembler we have for assembling metalloproteases right now, which is part of why it is so important. Another thing to know about Extender is that it primarily assembles really abundant transcripts, which means that it misses a lot. So Extender will give us much lower numbers of transcripts than the other assemblers, but in many cases more of what we really want: Toxins.

To run Extender we just need the merged reads from the previous step. Please contact D. Rokyta for access to Extender.

```
parallel -a list.txt -j 2 --verbose "echo {}
  cd {}
  mkdir 04_extender
  cd 04_extender
  Extender3.py -r ../03_merged/{}_pear.fastq -s 1000 -o 120 -msq 30 -mrq 20 -reps 20 -p 0.20 -e 2 -np
```

NGen

SeqMan NGen v14 (DNAStar, Inc., Madison, WI, USA; NGen14) is proprietary assembler which also does not use de Bruijn graphs. Because it is proprietary, you may opt to use several other assemblers instead. However, if you do choose to purchase or gain access to use SeqMan, we use default settings for transcriptome assembly.

Trinity

Trinity is the typical de Bruijn graph transcriptome assembler. There is plenty of information online regarding Trinity including videos that explain Trinity in detail (https://www.broadinstitute.org/broade/trinity-screencast)

NOTE: Remember that we created a separate environment for Trinity, so you will need to activate this environment

```
conda activate trinity_env
parallel -a list.txt -j 2 --verbose "cd {}
    mkdir 06_trinity
    cd 06_trinity
    Trinity --seqType fq --CPU 4 --min_contig_length 200 --max_memory 20G --full_cleanup -output {} --s
conda deactivate
```

Combining Assemblies

Now that we have our three assemblies complete, we can combine them into one before annotation. Each of the assemblers produces a fasta file of assembled contigs with complex names. We can just go ahead and rename each of those as contig1, contig2, ..., contigX. There are many ways to do this and we provide an example of how to do this with awk, but we recommend using Fasta_Renamer.py, which is a script to rename fasta headers sequentially.

Here, I'm copying the final assemblies from each assembler, renaming them, and concatenating them together into a combined assembly. After this, we run cd-hit to remove transcripts with 100% identity to reduce redundancy and computational load for annotation.

NOTE: You do NOT need to do both the awk and Fasta_Renamer.py strategies.

AWK:

```
for i in 'cat list.txt'
  do echo $i
  cd $i
  mkdir 07_assembly
  awk '/^>/{print ">extenderContig" ++i; next}{print}' < 04_extender/Final_Extender_contigs.fasta > 07
  awk '/^>/{print ">ngenContig" ++i; next}{print}' < 05_ngen/${i}_NGen14_NovelTranscripts.fas > 07_ass
  awk '/^>/{print ">trinityContig" ++i; next}{print}' < 06_trinity/${i}.Trinity.fasta > 07_assembly/${
  cd 07_assembly
  cat ${i}_extender.fasta ${i}_ngen.fasta ${i}_trinity.fasta > ${i}_assembly.fasta
  cd-hit-est -i ${i}_assembly.fasta -o ${i}_assembly_reduced.fasta -d 0 -c 1.00
  cd ../..
  done
```

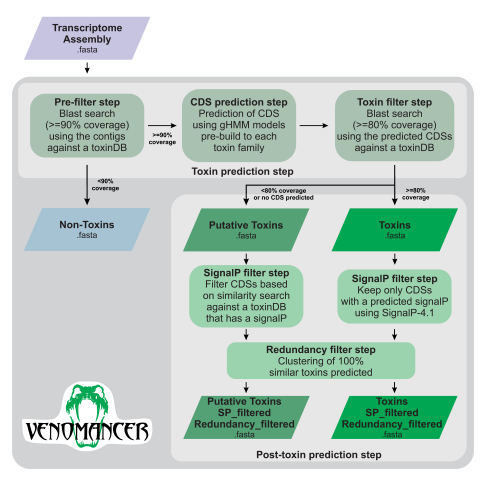
OR

Fasta_Renamer.py:

```
parallel -a list.txt -j 2 --verbose "echo {}
  cd {}
  mkdir 07_assembly
  cp 04_extender/Final_Extender_contigs.fasta 07_assembly/{}_extender.fasta
  cp 05_ngen/{}_NGen14_NovelTranscripts.fas 07_assembly/{}_ngen.fasta
  cp 06_trinity/{}.Trinity.fasta 07_assembly/{}_trinity.fasta
  cd 07_assembly
  Fasta_Renamer.py -f {}_extender.fasta -n 'extenderContig'
  Fasta_Renamer.py -f {}_ngen.fasta -n 'ngenContig'
  Fasta_Renamer.py -f {}_trinity.fasta -n 'trinityContig'
  cat {}_extender.fasta {}_ngen.fasta {}_trinity.fasta > {}_assembly.fasta
  cd-hit-est -i {}_assembly.fasta -o {}_assembly_reduced.fasta -d 0 -c 1.00"
```

Transcriptome Annotation

Venomancer



Venomancer was designed to identify and annotate toxins from a *de novo* venom gland transcriptome assembly. This program uses blast and trained generalized Hidden Markov Models (gHMM) to identify toxins in our assembly and annotated them. We will then use the resulting files Toxins_cds_Redundancyfiltered.fasta and PutativeToxins_cds_SPfiltered.fasta, combining them together and appending "TOXIN" to the beginning of the fasta names.

```
conda activate venomancer_env
parallel -a list.txt -j 2 --verbose "echo {}
  cd {}
  mkdir 08_venomancer
  venomancer.py -s {} -t 07_assembly/{}_assembly_reduced.fasta -o 08_venomancer -m /path/to/models -c 4
  cd 08_venomancer
  cat {}_Toxins_cds_Redundancyfiltered.fasta {}_PutativeToxins_cds_SPfiltered.fasta > {}_Toxins.fasta
  perl -pi -e 's/>/TOXIN_/g' {}_Toxins.fasta"
```

For nontoxin annotation, we will take the Nontoxin output from Venomancer (Nontoxins_contigs.fasta) and use Venomancer's NonToxins_annotator.py. The NonToxins_annotator.py uses CodAn to predict coding sequence (CDS) regions from the contigs followed by blast, BUSCO (optional), and Pfam (optional) to annotate the predicted CDSs, keeping uncharacterized proteins for potential novel toxin discovery.

```
parallel -a list.txt -j 2 --verbose "echo {}
  cd {}
  mkdir 09_nontoxins
  mv 08_venomancer/{}.venomancer_NonToxins_contigs.fasta 09_nontoxins/
  cd 09_nontoxins
  NonToxins_annotator.py -i {}.venomancer_NonToxins_contigs.fasta -o {}_Nontoxins.fasta"
  conda deactivate
```

With both toxins and nontoxins annotated, we can combine them together in preparation for checking for chimeras

```
parallel -a list.txt -j 2 --verbose "echo {}
  cd {}
  mkdir 10_CompleteAnnotation
  cat 08_venomancer/{}_Toxins.fasta 09_nontoxins/{}_Nontoxins.fasta > 10_CompleteAnnotation/{}_annotate
```

Now you can skip down to the section on Removing Chimeras!

Manual Annotation

If you don't think Venomancer is doing the trick or you want to be incredibly thorough and make sure you got everything, we will also describe how we manually annotate venom gland transcriptomes. If you are comfortable with Venomancer performance, you can skip down to the section on Removing Chimeras!

BLAST

First, we need to identify (to the best of our ability) each of the contigs in our assembly. One of the easiest ways to do this is with a blast search against a database of known things (e.g. GenBank, SWISSprot). We could take each of our contigs one-by-one and use the blast web interface, but that would be really slow. Therefore, we will download a database onto our local computer and then use command line blast. Depending on the size of the assembled contigs file, this may run for a few minutes or for a few hours. Afterwards though, you will have an xml file that contains the blast results of your search.

If you have not downloaded the SWISSprot database yet, go ahead and download this into a folder in your bin. It's a good idea to date this folder so you know when you downloaded it. You can then turn the fasta into a blast database with the following code.

```
makeblastdb -in uniprot_sprot.fasta -dbtype prot -out SWISSprot
```

Below, I've done a blastx search agains the full SWISSprot database. The output (-outfmt 5) is in xml format, but this can be changed.

```
parallel -a list.txt --verbose -j 2 "echo {}
  cd {}
  mkdir 08_blast
  cd 08_blast
  blastx -query ../07_assembly/{}_assembly_reduced.fasta -db /path/to/dated_SWISSprot/SWISSprot -outfmt
```

NOTES:

• blast is pretty much the best we have in many cases. However, it assumes that similar sequences to your transcriptome already exist in your database of choice. If your database is incomplete (as all databases are), you will not recieve a match.

Similarly, if you use the UniProt Toxins database, this means that only toxins are going to receive
matches and nontoxins will be unidentified.

Now we are ready for manual annotation.

Geneious Setup

We use **Geneious** to visualize our sequences and perform annotation; however, there are other softwares available capable of doing this task.

We first generate a file/folder structure in **Geneious** for each sample that resembles the following First, open **Geneious** and create the following file structure for each of your individuals.

Geneious

```
|-- sample_001
|-- Unannotated
|-- Annotated
| |-- Toxin
| |-- Nontoxin
| |-- Bad
| |-- Uncharacterized
|-- CDS
```

Then, drag your unannotated assembly contigs 07_assembly/{}_assembly_reduced.fasta into Geneious/Unannotated.

Find Open Reading Frames (ORFs)

If assembly was perfect, then each contig would represent a transcript. However, this is not the case. The transcripts are actually "hidden" in the middle of the contigs. Therefore, we need to find the start/stop codons inside our contigs. There can be multiple start/stops in each contig and these are known as the "Open Reading Frames" or ORFs. Venomancer attempts to automate the identification of the correct ORF, but now you get to experience it for yourself and gain an understanding for why automating this is so challenging.

Select all your contigs and go to the Annotate & Predict menu and and choose Find ORFs. Choose the following options:

- Minimum ORF Size: 90 bp
- Assume Start/Stop outside of Sequence

Each contig will have multiple ORFs annotated onto them. Which one is correct?! You are now ready to begin manual annotation.

Annotate

To annotate and choose the correct ORF, we compare the available ORFs to the blast results using something similar to blastxmlparser. However, it may be easier to consider alternate blast output formats. Additionally, Geneious has the option to annotate from a database. Therefore, if you have a folder of previously annotated toxins, toxin domains, or transcripts in general, this may also be useful to help you choose the correct ORF.

After identifying which ORF (if any) is the most appropriate, you will then annotate it. If you're unsure if a transcript is good, I suggest leaning on the side of keeping it since later steps (e.g. Removing Chimeras) will help eliminate bad transcripts.

If there is no match, then just throw the contig into the Bad folder you created and move onto the next contig. When you find a match do the following things:

- Double click ORF:
 - Name: CDSType: CDS
- Save
- Rename transcript to {SampleName}_{Toxin}_extender1

Export

Once you get through all the contigs, I suggest you add "TOXIN" to the name of all your toxins for easy sorting later. This can be done with the batch rename function in Geneious.

After you do this, select all of your annotated Toxins, Nontoxins, and Uncharacterized transcripts and extract the annotated regions of each contig. To do this, go to the Tools menu and select Extract Annotations. Move the extracted CDS into the CDS folder.

Now we are pretty much done in Geneious. We just need to export all those sequences into a single fasta named {}_CDS.fasta and we will put them in a new folder.

```
parallel -a list.txt -j 2 --verbose "echo {}
  cd {}
  mkdir 10_CompleteAnnotation"
```

Remove Duplicates and Ambiguities

Our fasta of annotated transcripts likely has a lot of duplicate sequences and sequences with ambiguities. We want to get rid of these. To do that we can use RemAmbRemDup.py.

```
parallel -a list.txt -j 2 --verbose "echo {}
  cd {}/10_CompleteAnnotation
  RemDupRemAmb.py -f {}_CDS.fasta -o {}_annotated"
```

Removing Chimeras

Pat yourself on the back for getting through annotation! Unfortunately, I have some bad news for you... Not everything you just annotated is real. Some of the transcripts represent artifacts of *de novo* transcriptome assembly where two or more transcripts are accidentally merged into one creating *chimeras*. We need to weed out chimeras to produce the *true* transcriptome.

ChimeraKiller

ChimeraKiller is designed to identify chimeric sequences. It starts by mapping the reads to your annotated transcripts. ChimeraKiller then looks at each site along a transcript and determines the average coverage to the left and right of that position. At chimeric sites, the reads from the two transcripts will butt up against one another. Therefore, as you near chimeric sites the discrepency in the number of bases to the left and right will increase. For example, if the average number of bases per read to the left is 10 and the average number of bases per read to the right is 200, the difference between them is 190. Using a percent difference threshold, the transcript will be removed.

```
conda activate chimerakiller_env
parallel -a list.txt -j 2 --verbose "echo {}
cd {}
```

```
mkdir 11_chimerakiller
  cd 11_chimerakiller
  ChimeraKiller_v0.7.3.py -i ../10_CompleteAnnotation/{}_annotated.fasta -r ../03_merged/{}_pear.fastq
conda deactivate
```

It is now a good idea to manually check the decisions made by ChimeraKiller. After we check the calls by ChimeraKiller we can concatenate the good fastas together. We will do this in the next section...

Clustering

Congrats!! At this point you've basically created a transcriptome. We aren't quite done, but the last few steps are just further reducing redundancy and double-checking that we've got everything annotated.

First, lets reduce more redundancy. Right now, our transcriptome contain heterozygous alleles of the same transcript. We don't need to have both alleles represented in our transcriptome. We can try to reduce this using cd-hit-est to cluster sequences at 99% identity.

```
parallel -a list.txt -j 2 --verbose "echo {}
  cd {}
  mkdir 12_transcriptome
  cat 11_chimerakiller/fastas/good/*.fasta > 12_transcriptome/{}_transcriptome_v0.fasta
  cd 12_transcriptome
  cd-hit-est -i {}_transcriptome_v0.fasta -o {}_transcriptome_v1.fasta -c 0.99"
```

NOTE: If you are creating a species-wide consensus transcriptome, you will probably want to do another round of clustering. Specifically, you will concatenate all the transcriptomes for a species together and use cd-hit-est to reduce redundancy that has resulted from general sequence variation among individuals. In this case, you may want to cluster at 98% (*i.e.* -c 0.98) or lower.

Here is an example, clustering together our two B. alternatus samples.

```
cat */12_transcriptome/*_transcriptome_v1.fasta > Balternatus_concat_transcriptome_v1.fasta
cd-hit-est -i Balternatus_concat_transcriptome_v1.fasta -o Balternatus_concensus_transcriptome_v1.fasta
```

Annotation Check (Optional)

Venomancer and NontoxinAnnotator are designed to predict CDS and annotate things potentially missed by Venomancer and even keep uncharacterized transcripts. This not only provides confidence in your transcriptome, but facilitates novel toxin discovery.

However, there are a couple other optional steps you might be able to take to make sure you don't miss anything. Primarily,

- 1. Proteomics (quantitative mass spectrometry) and
- 2. Estimating expression of all ORFs in the transcriptome assembly.

We will explain both of these briefly, but both start by requiring a fasta of all ORFs in the transcriptome assembly. We can use emboss to do this, generating both the nucleotide and protein ORFs. We will also run cd-hit to reduce redundancy and make different parts of the downstream analyses a little easier.

```
parallel -a list.txt -j 2 --verbose "echo {}
  cd {}/07_assembly
  getorf -find 1 -minsize 90 -sequence {}_assembly.fasta -outseq {}_ORFs.pro
  getorf -find 3 -minsize 90 -sequence {}_assembly.fasta -outseq {}_ORFs.fasta
  cd-hit -i {}_ORFs.pro -o {}_clusteredORFs.pro -c 1.0
  cd-hit-est -i {}_ORFs.fasta -o {}_clusteredORFs.fasta -c 1.0"
```

Proteomics

You can use quantitative mass spectrometry (qMS) to help identify important proteins in the venom. Specfically, providing the protein ORFs during qMS can help identify which ORFs are abundant in the proteome and may represent important or novel toxins. We will not talk about the generation of qMS data from the venom, but we use Scaffold to generate and export the ORFs of importance. The output of Scaffold can be placed

Scaffold will show you which ORFs had proteomic matches which you can then export ({}_Scaffold.pro), cluster to your annotated transcriptome, and annotate what remains with blast.

The script QTIBAC can help do this automatically. QTIBAC is a wrapper script which uses blast, cd-hit, and ChimeraKiller to cluster proteomically (quantitative mass spectrometry/qMS) identified ORFs to your transcriptome, blast them to a provided database (or to the NR database on Genbank), and remove chimeric ORFs.

```
parallel -a list.txt -j 2 --verbose "echo {}
  cd {}/13_qms
  QTIBAC.py -f ../07_assembly/{}_ORFs.fasta -q {}_Scaffold.pro -a ../12_transcriptome/{}_transcriptome_
```

The new transcriptome can be named {}_transcriptome_v2.fasta

ORF Expression Check

If you do not have a proteome, you can still check the expression of the ORFs to determine if you might have missed something important in your transcriptome. To do this, estimate expression of all ORFs using RSEM and use the results to make a list of the top X most highly expressed ORFs. Using grep, you can then extract those ORFs and use our proteomic friend QTIBAC in the same manner.

```
parallel -a list.txt -j 2 --verbose "echo {}
  cd {}
  mkdir 13_orf_check
  cd 13_orf_check
  rsem-prepare-reference --bowtie2 ../07_assembly/{}_clusteredORFs.fasta {}_ORF_Reference
  rsem-calculate-expression --bowtie2 ../03_merged/{}_pear.fastq {}_ORF_Reference {}
  rsem-plot-model {} {}_diagnostic.pdf"

# Make list of top X most highly expressed ORFs
parallel -a list.txt -j 2 --verbose "echo {}
  cd {}/13_orf_check
  QTIBAC.py -f ../07_assembly/{}_ORFs.fasta -q {}_Scaffold.pro -a ../12_transcriptome/{}_transcriptome_
```

The new transcriptome can be named {}_transcriptome_v2.fasta

Estimating Expression

With our final transcriptome generated, we can now estimate expression of each transcript with RSEM!

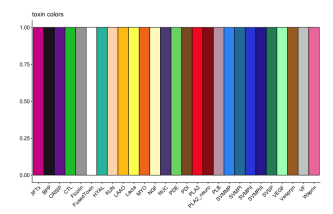
```
parallel -a list.txt -j 2 --verbose "echo {}
  cd {}/12_transcriptome
  mkdir RSEM_v2
  cd RSEM_v2
  rsem-prepare-reference --bowtie2 ../12_transcriptome/{}_transcriptome_v2.fasta {}_Reference
  rsem-calculate-expression --bowtie2 ../03_merged/{}_pear.fastq {}_Reference {}
  rsem-plot-model {} {}_diagnostic.pdf"
```

Expression Visualization

For visualization, we will move into R. Additionally, we will use data from *Crotalus cerastes* generated by Hofmann and Rautsaw et al. (2018) as our example data given their an increased sample size for differential expression analyses.

In order to visualize our data and perform downstream analyses, we need to set up a few things in R. Primarily, make sure that the appropriate R packages are installed and loaded. In addition, we will load a source script that has some useful functions and toxin color palettes.

```
source("./PlottingFunctions.R")
```



Now we can load our data into R and start to visualize expression of our toxins. There are sooooo many ways to visualize this data. I'm just going to go through a few here...

As an example, I am going to load in our data on Crotalus cerastes.

```
#setwd("PATH/TO/RSEM/RESULTS")
TPM_df<-read_excel("./example_data/RSEM.xlsx", sheet="TPM", col_types=c(rep("text",3),</pre>
                                                             rep("numeric",8),
                                                             "text"))
Counts df<-read excel("./example data/RSEM.xlsx", sheet="ExpectedCounts", col types=c(rep("text",3),
                                                                            rep("numeric",8),
                                                                            "text"))
samples <- read_excel("./example_data/samples.xlsx",sheet = "TranscriptomeSamples")</pre>
# Removing Extraneous Columns and Calculating Average TPM
TPM df <- TPM df [,1:11]
TPM_df$Average<-rowMeans(TPM_df[,4:11])</pre>
# Impute O values
TPM_df2 <- cbind(TPM_df[,1:3],t(cmultRepl(t(TPM_df[,4:12]),output = "p-counts")))</pre>
## No. corrected values:
rownames(TPM_df2)<-TPM_df2$gene_id
# Summing data by toxin class
TPM_class_df<-as.data.frame(TPM_df2 %>% group_by(toxin_class,class) %>%
                               summarize if(is numeric,sum) %>% arrange(desc(Average)))
rownames(TPM_class_df)<-TPM_class_df$Toxin_ID</pre>
Counts_class_df<-as.data.frame(Counts_df %>% group_by(toxin_class,class) %>%
                                  summarize_if(is_numeric,sum))
rownames(Counts_class_df)<-Counts_class_df$toxin_class</pre>
# Log/CLR transforming data
TPM_df2_log<-TPM_df2 %>% mutate_if(is.numeric,log) # can change to clr
TPM_df2_log<-TPM_df2_log[order(-TPM_df2_log$Average),]</pre>
rownames (TPM_df2_log) <- TPM_df2_log$gene_id
TPM_class_df_log<-TPM_class_df %>% mutate_if(is.numeric,log) # can change to clr
TPM class df log<-TPM class df log[order(-TPM class df log$Average),]
rownames(TPM_class_df_log)<-TPM_class_df_log$toxin_class</pre>
```

Fancy Figures

Fancy figures are a series of three different plots. You can plot them each separately...

```
png("figures/FullTranscriptomePlot.png", width =1080, height=720)
FullTranscriptomePlot(TPM_df2,"CLP2057",class="class")
invisible(dev.off())

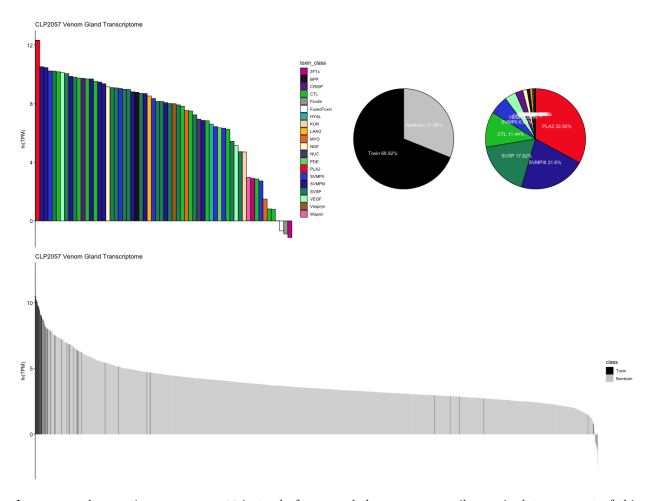
png("figures/ToxinBarplot.png", width =1080, height=720)
ToxinBarplot(TPM_df2,"CLP2057",class="class",toxin_class="toxin_class", colors=toxin_colors)
invisible(dev.off())
```

```
png("figures/ExpressionPie.png", width =1080, height=720)
ExpressionPie(TPM_df2,"CLP2057",class="class",toxin_class="toxin_class", colors=toxin_colors)
invisible(dev.off())
```

Or combine them into a Fancy Figure and loop through all your individuals.

```
png("figures/FancyFigure.png", width =1440, height=1080)
FancyFigure(TPM_df2,"CLP2057",class="class",toxin_class="toxin_class", colors=toxin_colors)
invisible(dev.off())

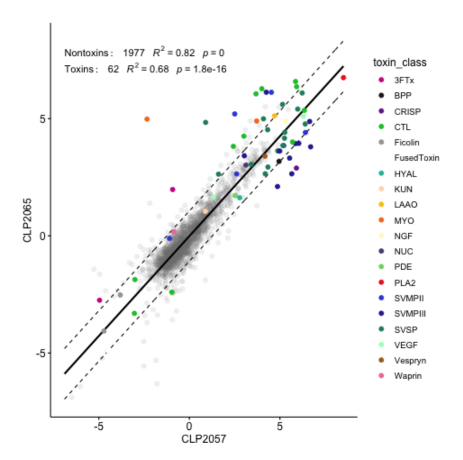
#for(i in c(samples$ID, "Average")){
# svg(paste0(i, "_FancyFigure.svg"),width=12,height=8.5)
# FancyFigure(TPM_df2,i,class="class",toxin_class="toxin_class", colors=toxin_colors)
# dev.off()
#}
```



I recommend exporting as svg or pdf instead of png, and then you can easily manipulate any part of this figure using something like Inkscape

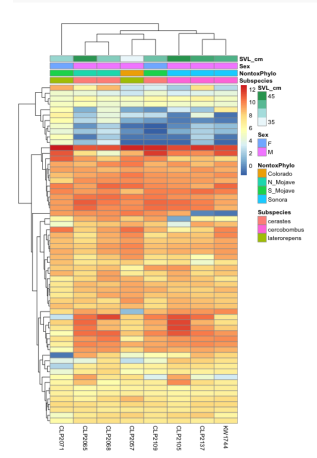
Transcriptome Comparison Plots

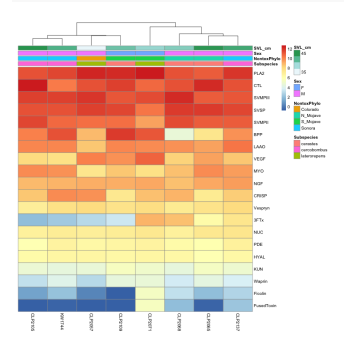
```
png("figures/TransCompPlot.png", width =480, height=480)
TransCompPlot(TPM_df2, "CLP2057", "CLP2065")
invisible(dev.off())
```



Heatmaps

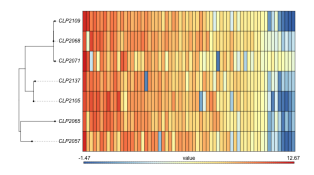
It may be helpful to visualize expression as a heatmap...

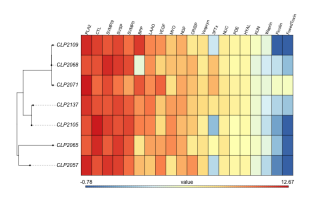




Phylogeny Heatmap

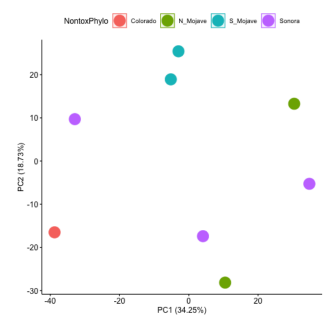
You could also use your phylogeny as the tree for the heatmap...





PCA

What about a PCA...



Differential Expression

We can also perform differential expression analysis! Make sure to use expected counts for this!!

DESeq2

```
DESeq_df<-as.data.frame(Counts_df[,4:11])</pre>
rownames(DESeq_df)<-Counts_df$gene_id</pre>
DESeq_df<-as.matrix(DESeq_df)</pre>
for (i in 1:ncol(DESeq_df)){
 DESeq_df[,i]<-as.integer(DESeq_df[,i])</pre>
## SVL Comparison
DESeq2<-DESeqDataSetFromMatrix(DESeq_df,metadata, design = ~SVL_cm)</pre>
DESeq2<-DESeq(DESeq2,fitType = "local")</pre>
DESeq2_res <- as.data.frame(results(DESeq2,alpha=0.05))</pre>
table<-DESeq2_res[DESeq2_res$padj<0.05,]</pre>
table <- head (table [complete.cases(table), c(2,6)])
                                                           log2FoldChange
#Ccera-CLP2057ext_CTL-9_ngenContig2269_-_CDS
                                                              0.29425216 0.02338078
#Ccera-CLP2105ext_MYO-2_extContiq94_-_CDS
                                                              0.39747275 0.02338078
\#Ccera-CLP2057\_SVMPIII\_extContig339\_-\_CDS
                                                             -0.28421367 0.02338078
#Ccera-CLP2057ext_SVMPIII-2_extContiq390_-_CDS
                                                             -0.17398424 0.02338078
#Ccera-CLP2057ext_ADH_ngenContig195_-_CDS
                                                             -0.06053533 0.03895370
#Ccera-CLP2065ngen_AminoAcidTrans1_ngenContig5623_-_CDS
                                                             0.12298525 0.04642247
## Phylogenetic Comparison
DESeq2<-DESeqDataSetFromMatrix(DESeq_df,metadata, design = ~NontoxPhylo)</pre>
DESeq2<-DESeq(DESeq2)
DESeq2_res <- as.data.frame(results(DESeq2,</pre>
                                     contrast=c("NontoxPhylo", "S Mojave", "N Mojave"),
                                     alpha=0.05))
table<-DESeq2_res[DESeq2_res$padj<0.05,]</pre>
table <-head(table [complete.cases(table),c(2,6)])
                                                   log2FoldChange
                                                                           padj
#Ccera-KW1744ext BPP-1a extContig132 - CDS
                                                        7.075668 5.189999e-08
#Ccera-CLP2057ext_PLA2-1a_extContig89_-_CDS
                                                        2.148142 4.467611e-02
#Ccera-CLP2105ext_SVMPII-4_extContiq172_-_CDS
                                                        -3.352636 3.900254e-02
#Ccera-CLP2105ngen_SVSP-10_ngenContig295_-_CDS
                                                        -2.698754 1.105192e-02
#Ccera-CLP2057ext_VEGF-1_extContiq1_-_CDS
                                                        2.441901 4.789369e-02
#Ccera-CLP2071ngen_39SrpL24_ngenContig6518_-_CDS
                                                      2.442785 2.608682e-07
```

edgeR

```
edgeR_df <- DGEList(counts=DESeq_df,samples=metadata)
keep <- rowSums(cpm(edgeR_df)>1) >= 2
edgeR_df <- edgeR_df[keep, , keep.lib.sizes=FALSE]
rm(keep)
edgeR_df <- calcNormFactors(edgeR_df)</pre>
```

```
# SVL Comparison
design <- model.matrix(~SVL_cm, data=edgeR_df$samples)</pre>
dispersion <- estimateDisp(edgeR_df,design)</pre>
fit <- glmFit(dispersion, design)</pre>
edgeR_res<-as.data.frame(topTags(glmLRT(fit,coef="SVL_cm"),n=Inf,sort.by="none"))</pre>
table <- edgeR res[edgeR res$FDR < 0.05,]
table <- head (table [complete.cases(table), c(1,5)])
                                                         logFC
                                                                       FDR
#Ccera-CLP2057ext_CTL-9_ngenContig2269_-_CDS
                                                   0.2943580 0.02230040
#Ccera-CLP2057ext_SVMPIII-2_extContig390_-_CDS -0.1744778 0.03014379
#Ccera-CLP2057ext_NucRec4A1_ngenContig2966_-_CDS -0.1933113 0.02230040
# Phylogenetic Comparison
design <- model.matrix(~NontoxPhylo, data=edgeR_df$samples)</pre>
dispersion <- estimateDisp(edgeR_df,design)</pre>
fit <- glmFit(dispersion, design)</pre>
edgeR_res<-as.data.frame(topTags(glmLRT(fit,</pre>
                                          coef=c("NontoxPhyloN_Mojave","NontoxPhyloS_Mojave")),
                                  n=Inf,sort.by="none"))
table <- edgeR res[edgeR res$FDR<0.05,]
table <- head (table [complete.cases (table), c(1,5)])
                                                         logFC.NontoxPhyloN_Mojave
                                                                                           PValue
#Ccera-KW1744ext_BPP-1a_extContig132_-_CDS
                                                                          -3.483285 3.416361e-06
\#Ccera-CLP2057ext\_CTL-5\_extContig12\_-\_CDS
                                                                          -2.873082 5.439565e-04
#Ccera-CLP2105ext_MYO-2_extContiq94_-_CDS
                                                                          9.956143 4.379644e-12
#Ccera-CLP2071ngen_39SrpL24_ngenContig6518_-_CDS
                                                                          -2.073622 1.091375e-06
#Ccera-CLP2057ext_3bHDDIsom_ngenContiq4815_-_CDS
                                                                          -2.327748 8.408196e-04
#Ccera-CLP2071ngen_ADPRibosyl1-2_ngenContig5670_-_CDS
                                                                           0.000000 6.485014e-34
```

That's it!

So long, and thanks for all the fish!