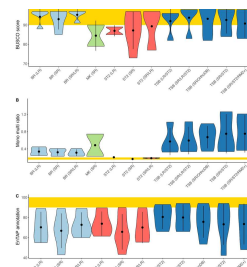


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Benchmarking protocol for plant genomes

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Protocol status: Working

We use this protocol and it's working

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Abstract

This is the protocol we used to benchmark several plant genomes. We first gather all available information of a particular species, from genome sequence to short-read and long-read RNA-seq data of the target species. Then, we annotate the genome by using *de novo* annotation software like BRAKER and MAKER.

Manuscript link: <https://www.biorxiv.org/content/10.1101/2022.10.03.510643v3>

For scripts used:

<https://gitlab.com/PlantGenomicsLab/annotationtool/-/tree/master/data-gathering>

For benchmarking stats visit:

https://docs.google.com/spreadsheets/d/1rITL0nSpJQzyq_3AepJunn_tv1QnGP67DRkTasigyIA/edit#gid=1628684765

For more stats on the runs visit:

https://docs.google.com/spreadsheets/d/1qbz8gQ1cDSFfLcFTKpfXutjOu2_eVlcO9r6pbV0hB1E/edit?copiedFromTrash#gid=1835523692

Guidelines

Access to scripts: <https://gitlab.com/PlantGenomicsLab/annotationtool/-/blob/master/data-gathering/README.md>



Materials

Visit <https://gitlab.com/PlantGenomicsLab/annotationtool/-/tree/master/data-gathering> for the scripts and a detailed summary of all the steps followed in benchmarking plant genomes

Before start

Place all files relevant to a step in the same directory. For example, all files related to genome evidence (under step 1) are placed in a directory "genome_evidence"



1 Download/gather genome resources

- Unmasked genome (for MAKER-P) pipeline
- Soft-masked genome (if available)
- gff, gtf files
- cDNA and pep files

Note

Please note the version of the genome resource.

2 Filter genome

1. filtersubmit.sh removes sequences less than 500 bp
 - The script filtersubmit.sh calls filterLen.py (custom script, no version no.)

Safety information

Make sure to filter the genome of <500 bp scaffolds before masking the genome



Overnight This step could take some time depending on the size of the genome

Soft-mask genome

Note

Check if genome is soft-masked first. If not, proceed with masking step.

1. Run repeatModeler.sh which loads RepeatModeler/2.01 to make the repeat library



Safety information

Run the genome with and without LTRStruct (these are two different RepeatModeler scripts)

Command

RepeatModeler2 generates repeat libraries which would be masked in the subsequent RepeatMasker step. The option, -LTRStruct helps to find more LTR components in the genome.

RepeatModeler2

```
BuildDatabase -name <org_dbname> <org_genome.fna>
```

#Note, LTRStruct is ON here

```
nohup RepeatModeler -database <org_dbname> -pa 20 -LTRStruct
```

#To run without LTRStruct

```
nohup RepeatModeler -database <org_dbname> -pa 20
```

1a. (optional) Run splitfasta.sh which loads anaconda and breaks the filtered genome into 100 pieces

Safety information

RepeatMasker can be run on the whole genome at once or on pieces of the genome separately using a SLURM array.
If splitting up the genome, run the splitfasta.sh script first.

2. Run repeatmasker.sh which loads RepeatMasker/4.0.6 Important: Check version before running this program.

If splitting up the genome, run **repeatmasker_array.sh**



If running on the genome as a whole, run **repeatmasker.sh**

Command

RepeatMasker actually masks the repeats generated by RepeatModeler

RepeatMasker

```
RepeatMasker -pa 12 -gff -a -noisy -low -xsmall -lib <org_dbname>-  
families.fa <org_genome.fna> -dir .
```

3. Calculate repeat content

3a. If RepeatMasker was run on the genome all at once (repeatmasker.sh), look at the repeat content value in the .tbl file

EXAMPLE:

Note

```
=====
file name: Arabidopsis_thaliana.TAIR10.dna.toplevel.fa
sequences: 7
total length:119667750 bp(119482146 bp excl N/X-runs)
GC level: 36.06 %
bases masked: 18219891 bp ( 15.23 %)
```

3b. If RepeatMasker was run as a SLURM array (split genome), use **total_pct_masked.sh** script.

Calculates correct repeat content fraction using all the .tbl files.

Provide path of directory with .tbl files as command line input with script.

Outputs FRACTION of genome that is masked.

4. Verify repeat content

rep_content.sh calculates repeat content.

Uses python script **rep_content.py**

Provide genome file as command line input with script.

Can be used to calculate repeat content for a genome that you did not mask yourself, as well as verifying repeat content seen in a RepeatMasker .tbl file



Outputs FRACTION of genome that is masked.

3 Genome quality assessment

1. Check quality of the filtered genome with Quast (quast.sh calls quast 5.0.2)

Command

QUAST

```
python /isg/shared/apps/quast/5.0.2/quast.py <org_genome.fna> -o  
quast_o
```

2. Check completeness of the filtered genome with BUSCO (busco.sh calls busco/4.1.2)

Command

BUSCO

```
busco -f -i <org_genome.fna> -l embryophyta_odb10 -o busco_o -m geno
```

Safety information

Have AUGUSTUS installed in a writeable directory

4 Creating indices for the alignment of evidence (RNA-Seq data and transcriptome) to the genome

Hisat2 and Gmap are used for aligning RNA-Seq (step 6) and transcriptome data (step 7), respectively.

Prior to the alignments, indices of the genome need to be created.

4.1 Gmap index

- gmap_index.sh calls gmap/2019-06-10 gmap_build
- -D # sets the output directory
- -d sets the index prefix

Command

gmapBuild

```
gmap_build -D <dir_name> -d <org_name> <org_genome.fna>
```

- pass in the masked, filtered genome fasta file last
- output files end in .chromosome, .chromosome.iit, .chrsubset, .contig, .contig.iit, .genomebits128, .genomecomp, .maps, .ref153offsets64meta, .ref153offsets64strm, .ref153positions, .sachildexc, .sachildguide1024, .saindex64meta, .saindex64strm, .salcpchilddc, .salcpexc, .salcpguide1024, .sarray, .version

4.2 Hisat index

- hisatBuild.sh calls hisat2/2.2.0 hisat-build
- -f sets the input .fasta file
- set the output directory/prefix last



Command

Hisat Build

```
hisat2-build -f <org_genome.fna> <org_name>.built
```

- output files are incrementing numbers like so: Arabidopsis_genome.1.ht2

5 Gather evidence

Go to NCBI to fetch short-read and long-read data from RNA-seq studies of the species

5.1 Prepare the short-read RNAseq data

1. Use the *trinityfetchSRA.sh* script to get reads from NCBI.

- Downloads FASTQ files with header format that can be used for hisat2 read alignment as well as Trinity transcriptome assembly
- `fastq-dump --define-seq '@${sn}_${rn}/${ri}' --split-files $LINE`
- Option `---split-fastq` to separate pairs into their respective left and right files while fetching.

Safety information

- at least 10M reads (spots) and >90% mapping rate for each library used for support
 - The library should be RNA-Seq based and not focused on plastid (chloroplast or mitochondria)
-
- Make a .txt file containing the SSR numbers of the reads to download.
 - *trinityfetchSRA.sh* calls *srtoolskit/2.8.1*
 - In *trinityfetchSRA.sh*, change the `FILENAME` to match your .txt.
 - output is .fastq files in the same directory.
 - move .fastq files to organism
2. Validate the retrieved reads using *validateSRA.sh*
- *validateSRA.sh* calls *srtoolskit/2.8.1*



- `FILENAME #` sets input file
- output is logged to `*.err`
- move reads to `raw_reads` dir

3. Trim reads using sickle

- `sickle.sh` calls `sickle/1.33`
- `rawreadsdir #` sets input dir
- output is `.fastq` files, two for each pair (1 and 2) that begin with *trimmed plus single_trimmed* for singles created by sickle.

4. Check quality of trimmed reads

- `fastqc.sh` loads `fastqc/0.11.7`

Safety information

Do take the time to check the FASTQC reports, this serves as a sanity check of whether the reads are good enough to use.
Pay attention to "Adaptor Content" and "Overrepresented Sequences" (may show adaptor sequences)

If there is significant adaptor content, go back to the raw read and trim this content using a tool like Scythe, Trimmomatic, or TrimGalore. Then, redo the Sickle trimming (except if using TrimGalore)

- `readsdir #` input directory
- output `*_fastqc.html` and `*_fastqc.zip` for each run

Safety information

Do hisat2 alignment (step 6) before creating Trinity *de novo* transcriptome
After hisat2 alignment, `alignRates.sh` (step 6.1.5) will show which libraries have an alignment (mapping) rate $\geq 90\%$
Only use the libraries with alignment rate $\geq 90\%$ for Trinity *de novo* transcriptome

6 Align short-read (RNA-Seq) to the genome

1. Run the hisat2 aligner

- `hisat.sh` calls `hisat2/2.2.0`
- `orgdir/trimmeddir #` set input directory (dir containing trimmed fastq)
- output is `.sam` files, one for each run (SRR file)



1.5. Run alignRates.sh

- Outputs files showing each hisat2 alignment (mapping) rate for each SRA library
- If rate $\geq 90\%$, it prints the SRA accession to good_SRAs.txt and prints the rate to good_rates.txt

Safety information

Only use libraries with alignment rate $\geq 90\%$
Only use good libraries to create BAM file with samtools.sh in next step
Only use good libraries to do Trinity transcriptome assembly

2. Convert, sort, and merge the human-readable .sam file to compressed machine-readable .bam

- samtools.sh calls samtools/1.7
- input location of .sam files
- output is one file with .bam extension

Safety information

samtools_updated.sh makes this process easier by only converting, sorting, and merging libraries in good_SRAs.txt (i.e. had a mapping rate $\geq 90\%$)

7 Assemble transcriptome and align to the genome

Safety information

Do NOT use a TSA from NCBI.
For consistency, all short-read-based transcriptomes will be created *de novo* with Trinity

1. Create *de novo* transcriptome with short reads and Trinity

- Use trimmed short-read libraries from before

Safety information

Run trinity separately on all libraries, and then concatenate



- Concatenate runs where necessary if they are from the same library, but just different lanes. Lanes are just a byproduct of the ways things are separated during the sequencing process, but it's all a part of the same sequencing library. Sometimes these are loaded into NCBI as separate runs. Library names can be found on the experiment (SRX) page. Lane info can be seen sometimes on the SRX page or by clicking on the SRR id links from the experiment page to go to the SRA Run Browser MetaData tab.
- Do not concatenate replicants. There are two kinds of replicants: sample and technical.
- Sample: Run these separately in Trinity.
- Technical: If a sample has multiple runs from the same library and same lane, it might be a technical replicant. If there is already enough read depth, we don't need multiple replicants of this kind.
- If the previous details are not clear in NCBI, referring to the published study may help clarify.

Safety information

Only use libraries with alignment rate $\geq 90\%$ (see hisat2 SRA alignment step)

trinity.sh calls trinity/2.8.5

- Run Trinity separately on all libraries - see trinity_updated.sh - runs as Slurm array on all libraries simultaneously
- PE_1 # sets the left read for each library
- PE_2 # sets the right read for each library
- out # sets the output directory
- min_contig_length is 300

Command

Trinity

```
Trinity --seqType fq --left $PE_1 --right $PE_2 --min_contig_length  
300 --output $out --full_cleanup --max_memory 150G --CPU 16
```



2. add_prefixes_trinity.sh

- Adds the SRA accession to the beginning of each header in all of the Trinity FASTA assemblies
- Gives all transcripts a unique name, prevents confusion of transcripts later
- Outputs the new FASTA files in \$org/evidence/transcriptome/trinity/trinity_prefix/

3. catTrinity.sh

- Concatenates all Trinity assemblies together

4. Frameselect transcriptome

- Input is file created before by catTrinity.sh - all Trinity assemblies concatenated together
- frameselect.sh runs TransDecoder/5.3.0
- clustered_TSA # sets directory that contains the TSA file
- filename # sets TSA file name

Command

Transdecoder finds the coding regions in transcript sequences.

Note: frameselect.sh requires the transcriptome to have a certain number of sequences to work (> 31)

Transdecoder

```
###Training with longest ORFs
```

```
TransDecoder.LongOrfs -t $TSA_dir/$filename
```

```
###generic prediction only reporting best hit
```

```
TransDecoder.Predict -t $TSA_dir/$filename --single_best_only
```

- output files are placed in directory 'bestHit' (.bed, .cds, .gff3, .pep)

5. Cluster frameselect transcriptome

- usearch.sh runs usearch/9.0.2132
- percent id is 0.98



- `concat_file #` sets the input file (.cds from previous step)
- output `centroids_$concat_file` and `centroids_$concat_file.uc`

6. Remove short genes (less than 300bps) from the centroids file

- Run `filter300.sh` which calls `filterLen.py`

Note

At this point, you have your final transcriptome (the filtered centroids file)

7. Run the gmap aligner

- `gmap.sh` calls `gmap/2019-06-10`
- `idx #` sets the location of the index
- `prefix #` sets the prefix of the index files
- `fasta` sets the location of the frameselect, clustered, filtered TSA as input
- `-a 1 #` Translate codons from given nucleotide (1-based)
- `--cross-species #` Use a more sensitive search for canonical splicing, which helps especially for cross-species alignments and other difficult cases
- `-D #` sets `idx`
- `-d #` sets `prefix`
- `-f gff3_gene #` sets the format
- `--fulllength #` Assume full-length protein, starting with Met
- `--min-trimmed-coverage=0.95 #` Do not print alignments with trimmed coverage less than this value
- `--min-identity=0.95 #` Do not print alignments with identity less than this value
- `-n1 #` set maximum number of paths to show. If set to 1, GMAP will not report chimeric alignments, since those imply two paths. If you want a single alignment plus chimeric alignments, then set this to be 0.



Command

Gmap aligns proteins to the genome

gmap

```
gmap -a 1 --cross-species -D $idx -d $prefix -f gff3_gene "$fasta" --  
fulllength --nthreads=8 --min-trimmed-coverage=0.95 --min-  
identity=0.95 -n1 > $prefix.gff3 2> $prefix.error
```

- output files are \$prefix_gmap.gff3 and \$prefix_gmap.error

8 Assemble proteome and align to the genome

1. Filter the .pep file created from frame selection

- run filterpep.sh to filter the proteins so that you only have the same sequences that correspond to the genes left after filtering in Step 7.3.
- filterpep.sh trims headers if there is additional info after id, makes a list of headers from filtered TSA, and then calls CreateFasta.py
- output is *_filtered.pep

Note

At this point, you have your final proteome
(the *_filtered.pep file)

2. Run the genomeThreader aligner

- genomeThreader.sh calls genomeThreader/1.7.1 and genomertools/1.6.1
- gt adds stop amino acids *
- gth aligns the protein sequences



Safety information

It is common for the error file to say "warning: protein sequence "some-transcript" does not end with a stop amino acid (*)" for one transcript.

Command

Genome Threader computes gene structure predictions.

Genome Threader

```
gt seqtransform -addstopamino $pep
```

```
gth -genomic $org$genome -protein $pep -gff3out -startcodon -  
gcmcoverage 80 -finalstopcodon -introncutout -dpmixonlen 20 -  
skipalignmentout -o $out -force -gcmxgapwidth 1000000
```

8.1 Prepare Stringtie proteins (using genome-guided aligner)

1. Run Stringtie to align reads
 - 0_stringtie.sh calls Stringtie and creates a gtf file in the directory
2. Extract proteins from the alignment file
 - 1_gffread.sh produces a protein file in the current directory
3. Frameselect the protein file using TransDecoder v5.5
 - 2_frameselect.sh uses TransDecoder and Egglog to frame-select proteins

9 Run genome annotation software

9.1 Genome annotation with BRAKER

Safety information

IMPORTANT STEPS BEFORE RUNNING BRAKER

Copy software to home directory

```
cp -r /labs/Wegrzyn/annotationtool/software/BRAKER_2.1.5 ~/
```

```
cp -r /labs/Wegrzyn/annotationtool/software/Augustus_3.3.3 ~/Augustus
```

```
cp -r /labs/Wegrzyn/annotationtool/software/cdbfasta ~/
```

Obtaining Genemark-ET software and license key

- http://exon.gatech.edu/GeneMark/license_download.cgi
- Select: GeneMark-ES/ET/EP ver 4.64_lic - LINUX 64
- Fill in information
- To download the software and the key, right-click on the respective link (64_bit for the key), and click "copy link address"

```
cd ~
```

Download, unzip, and rename the key

```
wget "enter-key-link-here"
```

```
gzip -d gm_key_64.gz
```

```
mv gm_key_64 local_gm_key
```

Genemark key may expire every few months, so go back and get a new one

Download and unzip the software

```
wget "enter-software-link-here"
```

```
tar -xf gmes_linux_64.tar.gz
```

```
rm gmes_linux_64.tar.gz
```

Change header in all perl scripts of the Genemark folder, fixes perl path

```
cd gmes_linux_64
```

```
module load perl/5.28.1
```

```
perl change_path_in_perl_scripts.pl "/usr/bin/env perl"
```

1. Run 1: Input- short read data

- Use a new directory for this: *annotaions/braker/braker.sh*
- genome from species of interest
- *braker.sh* calls BRAKER/2.1.5, augustus, genemark, bamtools
- augustus must be in your local directory (should be there from Step 3.3)
- make a directory in your home called 'tmp'
- softmasking 1 # indicates the genome has been softmasked
- copies of perl modules from the lab directory root are included for specific dependencies
- The merged .bam file from Step 6 is used as input.



```
braker.pl --species=yourSpecies --genome=genome.fasta \  
--bam=species.bam \  
--softmasking 1 \  
--gff3 \  
--cores 10
```

Safety information

VERY IMPORTANT

Make sure to have a separate species name for each alternate BRAKER run if running at the same time.
Otherwise, species gene data in your Augustus folder can be deleted or changed without throwing an error.

2. Run 2: Input- short read data and de novo proteins from species of interest

- Use a new directory for this: `annotaions/braker2/braker_prot.sh`
- **proteins from species of interest**
- **short reads from species of interest**
- `braker_protein.sh`
- like `braker.sh` but adds:
- `--prot_seq` : adds protein seqs,
- `--prg=gth` : this and the following param are for adding proteins of shorter evolutionary distance
- `--gth2traingen`

```
braker.pl --genome="$genome" --species="$species" --  
prot_seq="$protein" --epmode  
--softmasking --gff3
```

Following this, we run TSEBRA

```
echo `hostname`

export LC_ALL=en_US.UTF-8
export LANG=en_US.UTF-8
export LANGUAGE=en_US.UTF-8

module load python/3.6.3

tsebra=/home/FCAM/vvuruputoor/TSEBRA_1.0.3/bin
b1=/core/labs/Wegrzyn/annotationtool/testSpecies/PlantSet/arabidopsis/analysis/annotation/braker/braker/augustus.hints.gtf
b2=/core/labs/Wegrzyn/annotationtool/testSpecies/PlantSet/BRAKER+TSEBRA/arabidopsis/braker2_v2.1.6/braker/augustus.hints.gtf

config=/home/FCAM/vvuruputoor/TSEBRA_1.0.3/config/default.cfg

h1=/core/labs/Wegrzyn/annotationtool/testSpecies/PlantSet/arabidopsis/analysis/annotation/braker/braker/hintsfile.gff
h2=/core/labs/Wegrzyn/annotationtool/testSpecies/PlantSet/BRAKER+TSEBRA/arabidopsis/braker2_v2.1.6/braker/hintsfile.gff

$tsebra/tsebra.py -g $b1,$b2 -c $config -e $h1,$h2 -o
braker1+2_combined.gtf
```

3. Long Read -

- **protein from species of interest - derived from long-read data**
- **use the "--trainFromGth" flag instead of "--gth2traingenies"**

```
braker.pl --species=yourSpecies_long --genome=genome.fasta \
--prot_seq=longReadSpecies.fa --prg=gth --trainFromGth \
--softmasking 1 \
--gff3 \
--cores 10
```

4. Long and Short Read-

- combining short and long-read data for BRAKER annotation



Command

new command name

```
braker.pl --cores 14 --genome="$genome" --species="$species" --  
bam=$bam1,$bam2 --softmasking 1 --gff3
```

5. LTR masking + BRAKER-

- this uses the extra LTR masking as well as the Run 1 script

Runs 1, 3, 4, and 5 are also run with Stringtie prepared de novo proteins

9.2 Genome annotation with MAKER

Safety information

The genome for this step is the softmasked genome from RepeatModeler2, with the LTRStruct flag ON.

MAKER is run in iterations. The first step is to fill in the control files for MAKER.

Round 1: Input data includes genome, EST evidences (preferably externally aligned evidences through exonerate). The script to run MAKER is the same through all rounds:



Command

The most important step is to fill in the control files before running this step.

MAKER

```
mpiexec -n 5 maker maker_opts.ctl maker_bopts.ctl maker_exe.ctl -base  
<maker_name>
```

The next step is to run AUGUSTUS and SNAP



Command

AUGUSTUS

```
export AUGUSTUS_CONFIG_PATH=$HOME/Augustus/config
export AUGUSTUS_BIN_PATH=$HOME/Augustus/bin
export PATH=/home/FCAM/Augustus/bin:/home/FCAM/Augustus/scripts:$PATH

MAKERDIR=/home/FCAM/vvuruputoor/maker/bin
MAKERROUNDDIR=/labs/Wegrzyn/annotationtool/testSpecies/PlantSet/P_tric
hocarpa/p_trichocarpa/maker_arabi/1_round_maker
MAKERROUND=first_iter
species=arabi.augustus.maker

if [[ -d $AUGUSTUS_CONFIG_PATH/species/$species ]]; then rm -r
$AUGUSTUS_CONFIG_PATH/species/$species; fi

#take only the maker annotations
awk '{if ($2=="maker") print }' $MAKERROUNDDIR/$MAKERROUND.all.gff >
maker_rnd1.gff

/home/FCAM/vvuruputoor/Augustus/scripts/gff2gbSmallDNA.pl
maker_rnd1.gff
/labs/Wegrzyn/annotationtool/testSpecies/PlantSet/P_trichocarpa/p_tric
hocarpa_pipeline/example_run/genome/LTRStruct/P_trichocarpa.fasta.mask
ed 1000 $MAKERROUND.gb

/home/FCAM/vvuruputoor/Augustus/scripts/randomSplit.pl $MAKERROUND.gb
100

/home/FCAM/vvuruputoor/Augustus/scripts/new_species.pl --
species=$species
```

Command

SNAP

```
MAKERDIR=/home/FCAM/vvuruputoor/maker/bin
MAKERROUNDDIR=/labs/Wegrzyn/annotationtool/testSpecies/PlantSet/P_tric
hocarpa/p_trichocarpa/maker_arabi/1_round_maker
SNAPDIR=/home/FCAM/vvuruputoor/maker/exe/snap

${MAKERDIR}/maker2zff ${MAKERROUNDDIR}/first_iter.all.gff
${SNAPDIR}/fathom -categorize 1000 genome.ann genome.dna
${SNAPDIR}/fathom -export 1000 -plus uni.ann uni.dna
${SNAPDIR}/forge export.ann export.dna
${SNAPDIR}/hmm-assembler.pl first_iter . > first_iter.hmm
```

10 gFACs

1. Run transcriptome and protein alignments through gFACs - use gfacs_aln.sh (new script)
 - gfacs_aln.sh calls gFACs.pl
 - gmap and genomeThreader output is used as input
 - Options
 - -f gmap_2017_03_17_gff3 **or** genomethreader_1.6.6_gff3
 - --splice-rescue
 - --statistics
 - --statistics-at-every-step
 - --splice-table
 - --min-exon-size 3
 - --min-intron-size 3
 - --get-fasta-without-introns
 - --unique-genes-only
 - --create-gtf
 - --get-protein-fasta
 - The last option creates protein FASTA which should run through BUSCO
2. Run the output of all four braker runs through gFACs, both filtered and unfiltered

2a. Filtered

- -f braker_2.1.5_gff3
- --splice-rescue
- --statistics
- --statistics-at-every-step
- --splice-table
- --get-fasta-without-introns
- --get-fasta-with-introns
- --min-intron-size 10
- --min-exon-size 10
- --mins-CDS-size 300
- --rem-start-introns
- --rem-end-introns
- --get-protein-fasta
- --create-gtf

2b. Unfiltered

- --splice-rescue
- --statistics
- --statistics-at-every-step
- --splice-table
- --create-gtf

11 ENTAP

- database information at <https://bioinformatics.uconn.edu/databases/>
- Run EnTAP - entap.sh
- can be run with BAM/SAM file or braker output (augustus.hints.aa)
- BAM file uses flags --ontology 0, -a <BAMFILE>
- contaminants specified by -c flags, should contaminants be specific to species of interest?
- see https://entap.readthedocs.io/en/latest/basic_usage.html#id3 for additional flags
- should have multiple .dmnd files (at least 2, refseq and uniprot)