

GENE-SWitCH – Protocols



GENE-SWitCH

The regulatory GENomE of SWine and CHicken: functional annotation during development

Protocol WP2 Gene annotation with Isoseq sequencing using isoseq nextflow pipeline and downstream analysis.

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Workpackage: WP2

Version: 1.0

| Protocol associated with Deliverable(s): | D2.1 D2.2 | |
|--|------------|--|
| Submission date to FAANG: | 08/12/2021 | |

Research and Innovation Action, SFS-30-2018-2019-2020 Agri-Aqua Labs Duration of the project: 01 July 2019 – 30 June 2023, 48 months



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1 Summary

GENE-SWitCH aims at identifying functional elements located in the genomes of pig and chicken working on seven different tissues at three different developmental stages.

The seven tissues analysed in GENE-SWitCH are:

- Cerebellum
- Lung
- Kidney
- Dorsal skin
- Small intestine
- Liver
- Skeletal muscle

The three developmental stages are:

- Early organogenesis (E8 chick embryo and D30 pig foetuses)
- Late organogenesis (E15 chick embryo and D70 pig foetuses)
- Newborn piglets and hatched chicks

For each tissue at each time point, an Isoseq long read sequencing has been done. The raw subreads needs to be processed to generates definitive consensus sequences. The reads are mapped on the reference genome with uLTRA. The gene models are cleaned with TAMA. The resulting annotation files have to be processed to convert to the GFF format and add information (read count, annotation confidence).



2 Protocol description

The isoseq raw subreads processing, the genome mapping and the gene model annotation creation are made using the <u>isoseq nextflow pipeline</u>.

First the pipeline generates CCS using ccs Pacbio's program. The raw data are divided into batches of sequences that are processed in parallel. On each ccs batch, the program LIMA (from Pacbio) select CCS with appropriate primers pairs and removes them from the sequence. The resulting sequences are then processed by Pacbio's isoseq3 refine. It selects non-chimeric sequences with poly(A) tail, and trim it. The produces sequences are called Full Length Non Chimeric (FLNC). Before the clustering step, the sequence batches are merged using with Pacbio's pbmerge program. Next, the program isoseq3 cluster regroups similar sequences and create a consensus, called HIFI, for each cluster.

Before proceeding to the mapping, the not clustered FLNC and HIFI are merged using pbmerge and a fastq file is created using samtools fastq. The set of reads is aligned on the reference with the program uLTRA.

To accelerate the computations, the resulting alignment is split based on the chromosome with bamtools split. Each alignment is processed by TAMA collapse, for false positive and redundancy removing. To finish, the complete set of annotation bed files is merged in one unique bed file with TAMA merge.

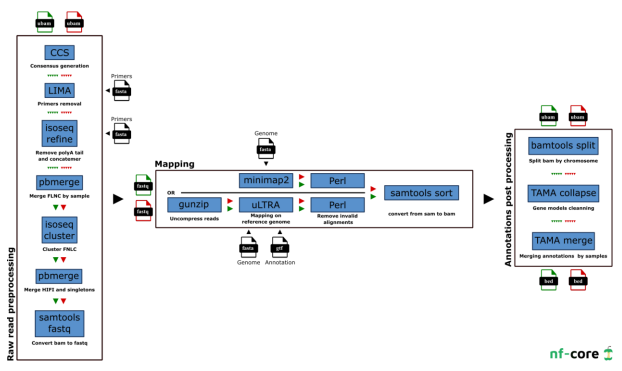


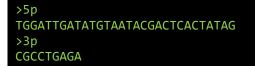
Figure 1: Isoseq nexflow pipeline

To run the pipeline, a data directory is created and the following input files are stored in it:

- Samples subreads (.bam)
- Pacbio index files (.bam.pbi)
- Genome (.fasta)
- Primer sequences (.fasta)
- Ensembl gene annotation (.gtf)



The genome used is Gallus Gallus 6 (GRCg6a) and its gene annotation from Ensembl release-104. The primers used are:



The complete command line is:



The files 31 bed files obtained for the 21 pairs of tissues/development stages have been merged together using TAMA merge.

tama_merge.py -f 31samples_filelist.tsv -d merge_dup -a 2000 -m 10 -z 2000 -p
31SamplesFiles 1> merge.log 2> merge.err

The "31samples_filelist.tsv" file is a four-column tabulation separated file. It contains the list of annotation to merge (column 1), the indication if the sequenced RNAs were capped (column 2), the priority annotation merging (column 3), and an id (column 4).



| merged_m64036_210413_213334.subreads.bed | capped | 1,1,1 | E8Brain |
|---|--------|---------|-------------|
| merged_m64036_210319_151149.subreads.bed | capped | 1,1,1 | E8Ileum |
| merged_m64036_210320_212651.subreads.bed | capped | 1,1,1 | E8Kidney |
| merged_m64036_210315_182820.subreads.bed | capped | 1,1,1 | E8Liver |
| merged_m64036_210322_035446.subreads.bed | capped | 1,1,1 | E8Lung |
| merged_m64036_210412_151836.subreads.bed | capped | 1,1,1 | E8Muscle |
| merged_m64036_210415_040208.subreads.bed | capped | 1,1,1 | E8Skin |
| merged_m64036e_210713_144608.subreads.bed | capped | 1,1,1 | E15BrainP2 |
| merged_m64036e_210719_150424.subreads.bed | capped | 1,1,1 | E15BrainP1 |
| merged_m64036_210622_160059.subreads.bed | capped | 1,1,1 | E15Ileum |
| merged_m64036_210623_220011.subreads.bed | capped | 1,1,1 | E15KidneyP2 |
| merged_m64036_210630_203419.subreads.bed | capped | 1,1,1 | E15KidneyP1 |
| merged_m64036_210418_013328.subreads.bed | capped | 1,1,1 | E15Liver |
| merged_m64036_210702_024547.subreads.bed | capped | 1,1,1 | E15LungP1 |
| merged_m64036_210625_041134.subreads.bed | capped | 1,1,1 | E15LungP2 |
| merged_m64036_210626_102305.subreads.bed | capped | 1,1,1 | E15MuscleP2 |
| merged_m64036_210703_085717.subreads.bed | capped | 1,1,1 | E15MuscleP1 |
| merged_m64036e_210714_204516.subreads.bed | capped | 1,1,1 | E15SkinP2 |
| merged_m64036e_210803_155018.subreads.bed | capped | 1,1,1 | E15SkinP1 |
| merged_m64036e_210828_025255.subreads.bed | capped | 1,1,1 | HCBrainP1 |
| merged_m64036e_210909_045748.subreads.bed | capped | 1,1,1 | HCBrainP2 |
| merged_m64036e_210717_090846.subreads.bed | capped | 1,1,1 | HCIleumP2 |
| merged_m64036e_210806_040105.subreads.bed | capped | 1,1,1 | HCIleumP1 |
| merged_m64036e_210812_170130.subreads.bed | capped | 1,1,1 | HCKidney |
| merged m64036e 210716 025706.subreads.bed | capped | 1,1,1 | HCLiverP2 |
| merged_m64036e_210804_214905.subreads.bed | capped | 1,1,1 | HCLiverP1 |
| merged_m64036e_210815_202444.subreads.bed | capped | 1, 1, 1 | HCLung |
| merged_m64036e_210907_224558.subreads.bed | capped | 1, 1, 1 | HCMuscleP2 |
| merged_m64036e_210826_204115.subreads.bed | capped | 1,1,1 | HCMuscleP1 |
| merged m64036e 210829 090412.subreads.bed | capped | 1, 1, 1 | HCSkinP1 |
| merged m64036e 210910 110947.subreads.bed | capped | 1, 1, 1 | HCSkinP2 |
| | | | |

The combined bed file has been converted is using TAMA script tama_convert_bed_gtf_ensembl_no_cds.py.

tama_convert_bed_gtf_ensembl_no_cds.py 31SamplesFiles.bed 31SamplesFiles.gtf

The GTF file has been updated to include a confidence attribute. The two possible values for this attribute are "low" and "high". If an annotation is supported by only one FLNC, the confidence is set to "low". If an annotation is supported by at least 2 FLNC or one HIFI, the confidence is set to "high". This modification has been made with two homemade R scripts: <u>get_low_high_confidence_annotations.R</u> and <u>add_confidence_annotation.R</u>.

The GTF file has been also modified to add the read count for each transcript. The read count of an annotation is the sum of FLNC supporting this annotation. If the annotation is supported by an HIFI, then the read count is incremented by the number of FLNC used to create this HIFI.

TAMA include a python script dedicated to read count computing. By running it after each TAMA collapse or TAMA merge, it's possible to keep track of read count through the cleaning/merging process. In this case, tama_read_support_levels.py has been run after pipeline's TAMA collapse (one bed file per chromosome per sample), pipeline's TAMA merge (one bed file per sample), and the last TAMA merge (one bed file for all samples).



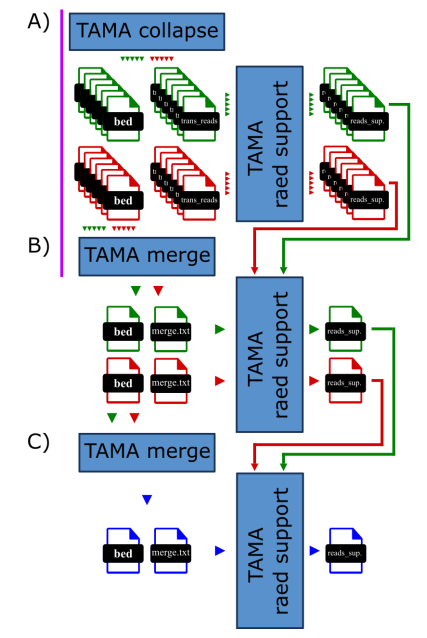


Figure 2: Read counting process – Purple bar indicates programs launch by the pipeline, others are launched manually. A) TAMA read support is launched on each trans_reads.bed files produced by TAMA collapse. B) TAMA read support is run on each sample merge.txt their associated chromosome read_support file. C) TAMA read support is run on the merge.txt of all samples with their associated sample read_support files

The first counting needs to be run on each trans_read.bed produced by TAMA collapse (Fig 2. A). For each of them, a read_support file is created. If all trans_read.bed files are gathered in the same directory, the following fish shell loop will run the script on each of them:

```
for i in (ls *.bed)
   set FILE $i
   set ID (string replace "_tc_trans_read.bed" "" $FILE)
   set FILELIST $ID"_"filelist.txt
   echo -e "$ID\t$FILE\ttrans_read" > $FILELIST
   set CMD "tama_read_support_levels.py -f $FILELIST -m no_merge -o $ID"
   echo $CMD
   eval $CMD
end
```



At each loop round, it will create a filelist.tsv input file composed of three tabulations separated columns (an ID, the trans_read.bed file and the file type) and run tama_read_support_levels.py.

Next, the generated read_support files are used to compute the read count at sample level (first TAMA merge). For each sample, a filelist.tsv file listing read_support files must be created. The first column is the ID, the second is the file path and the third one contains read_support as file type. The script can be started with following command:

```
tama_read_support_levels.py \
-f <SAMPLE>_filelist.tsv \
-m <SAMPLE>_merge.txt \
-o <SAMPLE>
```

Finally, the global read count can be computed using the read_support files generated for each sample. The filelist.tsv list sample's read_support files:

| I | | _ '' |
|-------------|------------------------------|--------------|
| E15BrainP1 | E15BrainP1_read_support.txt | read_support |
| E15BrainP2 | E15BrainP2_read_support.txt | read_support |
| E15Ileum | E15Ileum_read_support.txt | read_support |
| E15KidneyP1 | E15KidneyP1_read_support.txt | read_support |
| E15KidneyP2 | E15KidneyP2_read_support.txt | read_support |
| E15Liver | E15Liver_read_support.txt | read_support |
| E15LungP1 | E15LungP1_read_support.txt | read_support |
| E15LungP2 | E15LungP2_read_support.txt | read_support |
| E15MuscleP1 | E15MuscleP1_read_support.txt | read_support |
| E15MuscleP2 | E15MuscleP2_read_support.txt | read_support |
| E15SkinP1 | E15SkinP1_read_support.txt | read_support |
| E15SkinP2 | E15SkinP2_read_support.txt | read_support |
| E8Brain | E8Brain_read_support.txt | read_support |
| E8Ileum | E8Ileum_read_support.txt | read_support |
| E8Kidney | E8Kidney_read_support.txt | read_support |
| E8Liver | E8Liver_read_support.txt | read_support |
| E8Lung | E8Lung_read_support.txt | read_support |
| E8Muscle | E8Muscle_read_support.txt | read_support |
| E8Skin | E8Skin_read_support.txt | read_support |
| HCBrainP1 | HCBrainP1_read_support.txt | read_support |
| HCBrainP2 | HCBrainP2_read_support.txt | read_support |
| HCIleumP1 | HCIleumP1_read_support.txt | read_support |
| HCIleumP2 | HCIleumP2_read_support.txt | read_support |
| HCKidney | HCKidney_read_support.txt | read_support |
| HCLiverP1 | HCLiverP1_read_support.txt | read_support |
| HCLiverP2 | HCLiverP2_read_support.txt | read_support |
| HCLung | HCLung_read_support.txt | read_support |
| HCMuscleP1 | HCMuscleP1_read_support.txt | read_support |
| HCMuscleP2 | HCMuscleP2_read_support.txt | read_support |
| HCSkinP1 | HCSkinP1_read_support.txt | read_support |
| HCSkinP2 | HCSkinP2_read_support.txt | read_support |

The last tama_read_support_levels.py is launched with this following command:

tama_read_support_levels.py -f input_filelist.tsv -m 31SamplesFiles_merge.txt -o
31SamplesFiles

The global read count must be corrected to include the number of FLNC for each HIFI. This information can be obtained by running tama_read_support_levels.py on the cluster report generated by isoseq3 cluster. When all cluster_report.csv files are gathered in a directory, the script can be run using the following fish shell loop:





All the complete set of read_support is merged into one file before read_count correction.



The read count correction is made using a homemade R script: count_correction.R