De novo transcriptome assembly with Trinity

Trinity de novo assembly

Now we will start the Trinity run.

First open the **QSubGen web application**.

- Choose medium time limit and reserve 6GB of memory.
- Select 'multi-thread' and choose 10 CPUs.
- In the modules field, type trinity and the select the module bioinformatics/trinity/2.6.6.
- In job specific commands, type:

```
Trinity --seqType fq
--left data/RNA_Eye_1_val_1.fastq \
--right data/RNA_Eye_2_val_2.fastq \
--max_memory 60G --CPU $NSLOTS --full_cleanup
```

- Choose a descriptive job name then click on
- Check if OK
- If it passes, either save it and upload it to Hydra, or copy the text and paste it directly into your favority text editor.

Now save your text file into your /pool/genomics/<username>/RNAseq_SMSC directory as trinity.job.

Now submit your job with the command: qsub trinity.job

Soon your transcriptome assembly will be finished!