Anaquin TransQuin Report

Read Alignment

```
Summary for dataset: K_RMXA1v2.accepted_hits.sorted.bam
  Unmapped:
             0 reads
  Experiment: 36484961 reads
  Synthetic: 11440146 reads
  Reference: 1190 exons
  Reference: 1028 introns
  Reference: 149219 bases
  Query: 84739030 exons
Query: 27056077 introns
Query: 153438 bases
  Dilution: 0.238709
  *** The following statistics are computed at the exon, intron and base level.
  *** Exon level is defined by performance per exon. An alignment that
  *** is not mapped entirely within an exon is considered as a FP. The
  *** intron level is similar.
  *** Base level is defined by performance per nucleotide. A partial
  *** mapped read will have FP and TP.
  ***
  ----- Exon level -----
  Sensitivity: 0.997479
  Accuarcy: 0.984404
  Detection: 0.0590086 (R2_33)
  ----- Intron level -----
  Sensitivity: 0.993191
  Accuarcy: 1
  Detection: 0.0590086 (R2_33)
  ----- Base level -----
  Sensitivity: 0.703341
  Accuarcy:
             1
  Detection: 0.0590086 (R2_33)
```

Transctiptome Assembly

Expression Analysis

Differential Analysis