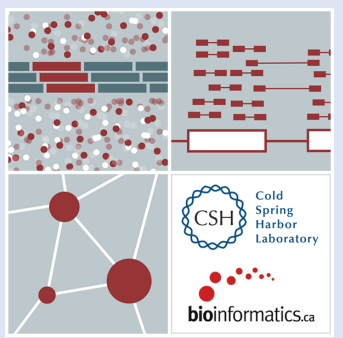
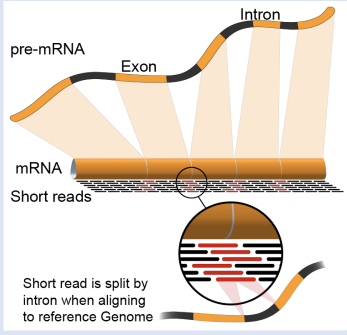


## RNA-Seq Module 3 HTSeq

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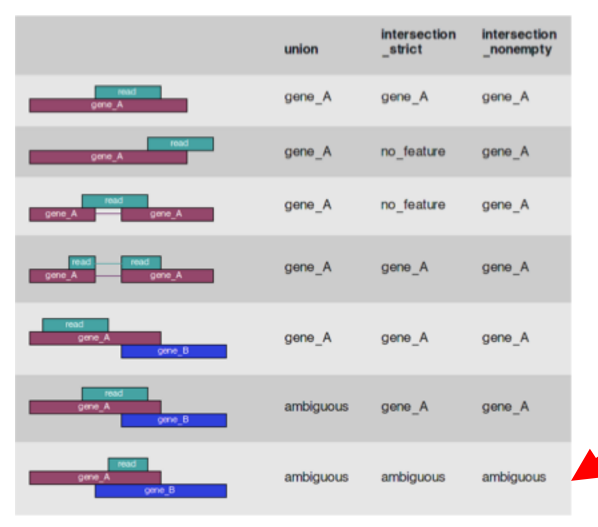
## **Alternatives to FPKM**

- Raw read counts for differential expression analysis
  - Assign reads/fragments to defined genes/transcripts, get "raw counts"
    - Transcript structures could still be defined by something like Stringtie
- HTSeq (htseq-count)
  - https://htseq.readthedocs.io/

```
htseq-count --mode intersection-strict --stranded no --minaqual 1 --type
exon --idattr transcript_id accepted_hits.sam chr22.gff >
transcript_read_counts_table.tsv
```

- Caveats of 'transcript' analysis by htseq-count:
  - Designed for genes ambiguous reads from overlapping transcripts may not be handled!
  - <a href="http://seganswers.com/forums/showthread.php?t=18068">http://seganswers.com/forums/showthread.php?t=18068</a>

## HTSeq-count basically counts reads supporting a feature (exon, gene) by assessing overlapping coordinates



Note, if gene\_A and gene\_B on opposite strands, sequence data is stranded, and correct HTSeq parameter set then this read may not be ambiguous

Whether a read is counted depends on the nature of overlap and "mode" selected

## We are on a Coffee Break & Networking Session