

RNA-seq Quantification



Outline

- the RNA-seq setting
- normalizing fractions
- short-read RNA-seq
- quantification:
 - read counts
 - generative models; relative abundances
 - pseudo-mapping
- error models
- analysis at log-scale
 - additive background
 - more complex models



Typical setting for bulk RNA-seq

Given a tissue or cell line:

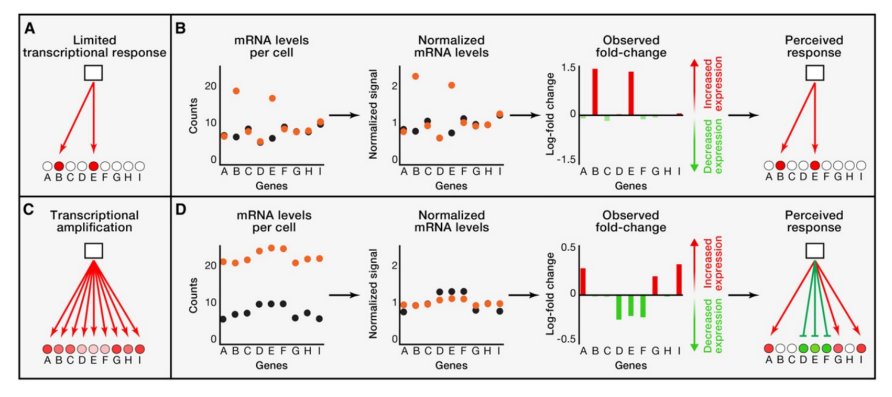
- ~25'000 genes in the genome
- \sim 10 18'000 genes expressed at any given time; expression level of the different genes differs largely

Working hypotheses:

- relative prevalence of a gene reflects activity of associated pathways; more prevalent, more active
- baseline prevalence is not known → perturbation experiments: treatment vs control
- informative quantity: relative abundance of the genes in a tissue Limitations:
- genes vs protein
- alternative explanations cell size effects; other side effects



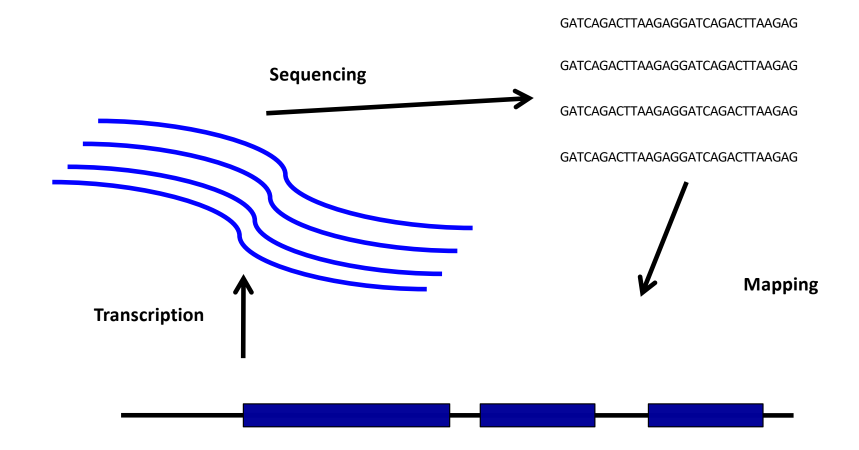
Perturbation Experiments



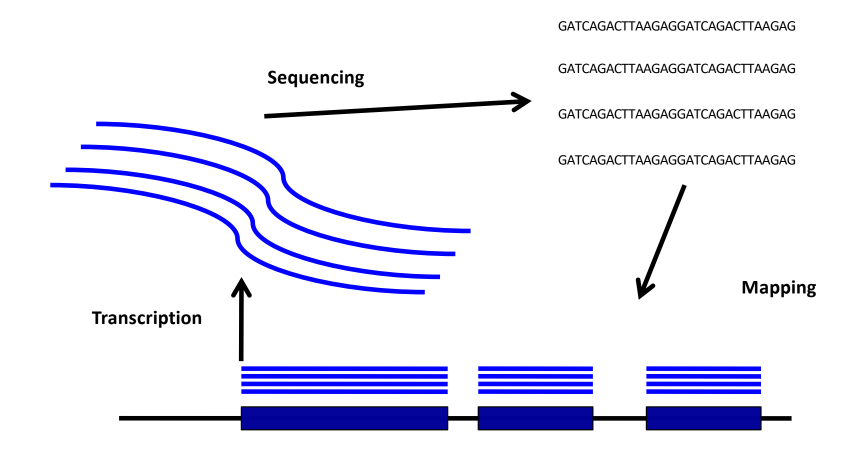
- The top row shows the assumption how a baseline expression profile (black) changes as a response (red) to a perturbation
- Note: expression counts are all around \sim 10, in practice they range from 0 1 Mio



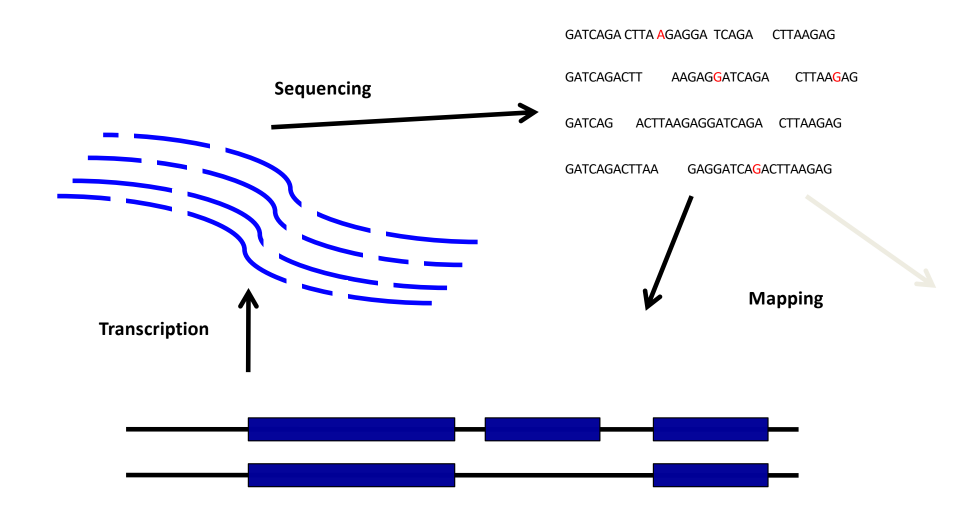
Transcript Expression with NGS



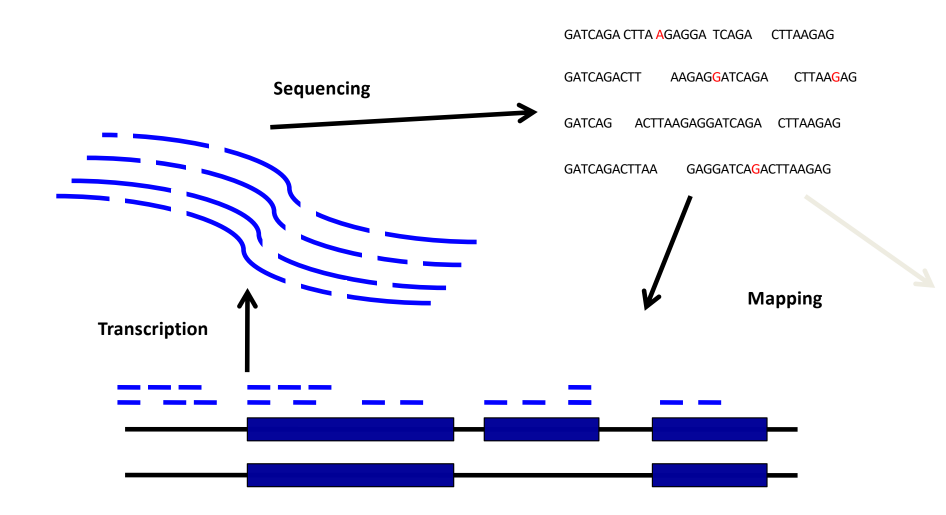
Read mapping and counting (ideal)



Read mapping and counting (today)



Read mapping and counting (today)



Abundance estimates

Abundance of what???

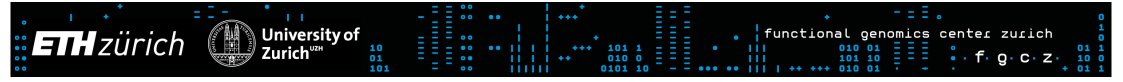
- Biologically relevant:
 - gene level:
 - # molecules transcribed from one gene locus (per cell)
 - isoform level:
 - # molecules of a specific isoform transcribed from one gene (per cell)
- Feasible with RNA-seq:
 - relative fractions that indicate the abundance relative to all other genes/isoforms

Gene-level Read Counts

- rather straightforward to compute:
 - number of reads that uniquely map to a gene locus
 - → biased by length, discards information in multi-mappers
 - number of reads that map to gene locus (including multi-mappers)
 - → disambiguation is not possible if you do not have abundance estimates of the isoforms
 - → needs to resort to heuristics to assign multi-mappers
 - randomly assign to one of the matching genes
 - do a fractional assignment with a with 1/#genes mapped

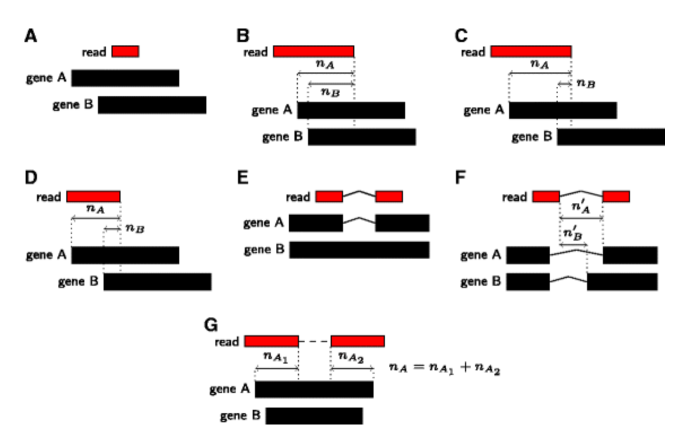
Rsubread::featureCounts – assigning reads to genes

- versatile function to count reads towards exons, transcripts, genes, ...
- implements many different counting modes
- covers different aspects of overlap situations
 - partial overlap
 - overlapping multiple features at the same alignment position
 - overlapping multiple features at different alignment positions
- Simple overlap is not sufficient, read must be compatible with exon structure



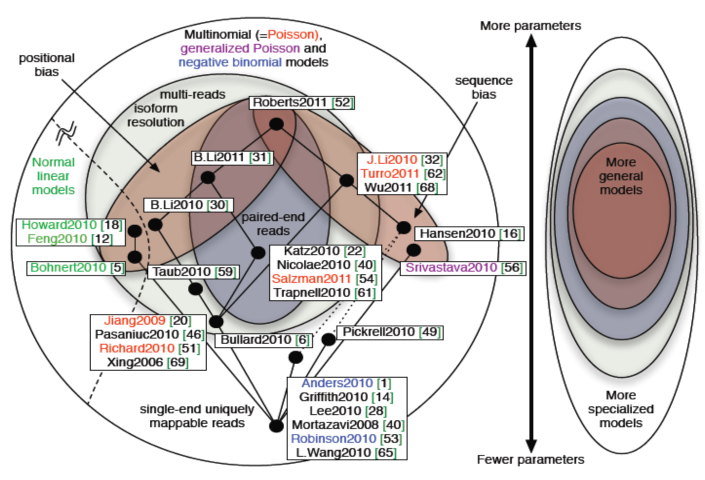
Model-free Counting of Overlapping reads – Count Modes

mmquant: resolve multi-mapping reads based on heuristics



https://bmcbioinformatics.biomedcentral.com/articles/10.1186/s12859-017-1816-4

Generative Models for RNA-seq quantification



Pachter 2011, https://doi.org/10.48550/arXiv.1104.3889

RNA-seq model

$$\alpha_t = P[\text{read from transcript } t] = \frac{1}{Z} \rho_t l_t$$

with:

$$\rho_t$$
 expression level / abundance / fraction

$$l_t$$
 transcript length

$$Z = \sum_{t} \rho_t l_t$$
 normalization factor

The normalization factor is the weighted mean length of the transcripts.

RNA-seq model

Estimation of the probability that a read is from a specific transcript:

$$\hat{\alpha}_t = \frac{X_t}{N} = \frac{\text{\#reads mapping to transcript } t}{\text{\#mappable reads in total}}$$

Abundance estimates:

$$\hat{\rho}_t \propto \frac{\hat{\alpha}_t}{l_t}$$

Maximum Likelihood Estimation

The estimated abundances represent unique MLE estimates

with
$$\alpha = \{\alpha_t\}_{t \in T}$$

$$L[\alpha] = \prod_{t \in T} \prod_{f \in F_t} P[f \in t] \frac{1}{l_t}$$

$$= \prod_{t \in T} \prod_{f \in F_t} \alpha_t \frac{1}{l_t}$$

$$= \prod_{t \in T} \left(\frac{\alpha_t}{l_t}\right)^{X_t}$$

Effective Transcript Length

 Since fragments have a non-zero length the read probabilities depend actually on an effective length:

 $l_t := \text{transcript length} - \text{fragment length} + 1$

- For simplicity we continue to use the symbol without tilde but will always assume it is the effective length
- The effective length represents the stretch of the transcript from which I can get a fragment that I can then map back to the transcript
- The effective length should also consider mappability!
- > Mappability does depend on mapping algorithm, mutations, ...

Multi-reads

- Reads that cannot be uniquely assigned to one transcript were ignored so far
- Multi-reads can occur
 - if a read aligns more than once in the genome
 - if at an alignment position there is more than one transcript defined
- Multi-reads do occur due to homology not due to pure chance

Considering Multi-reads

Define a compatibility matrix

$$\mathbf{Y} = \left\{ y_{ft} \right\}_{f \in F, t \in T}$$

with

$$y_{ft} = \begin{cases} 1 \text{ if read } f \text{ aligns to transcript } t \\ 0 \text{ else} \end{cases}$$

• The likelihood is now:

$$L[\alpha] = \prod_{f} \left(\sum_{t} y_{ft} \frac{\alpha_{t}}{l_{t}} \right)$$

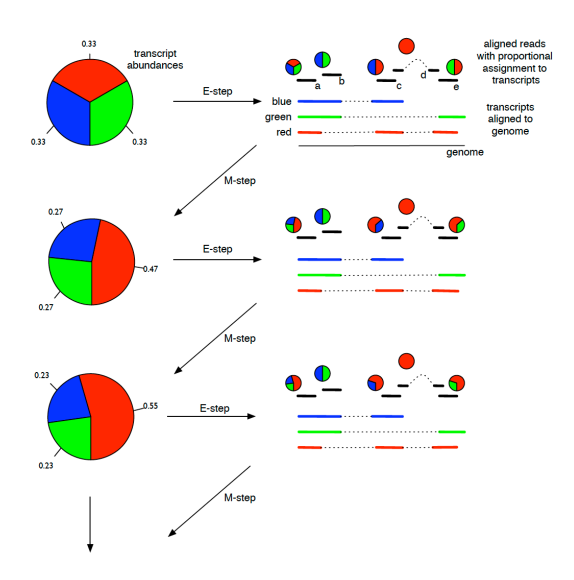
but now abundances must be estimated iteratively

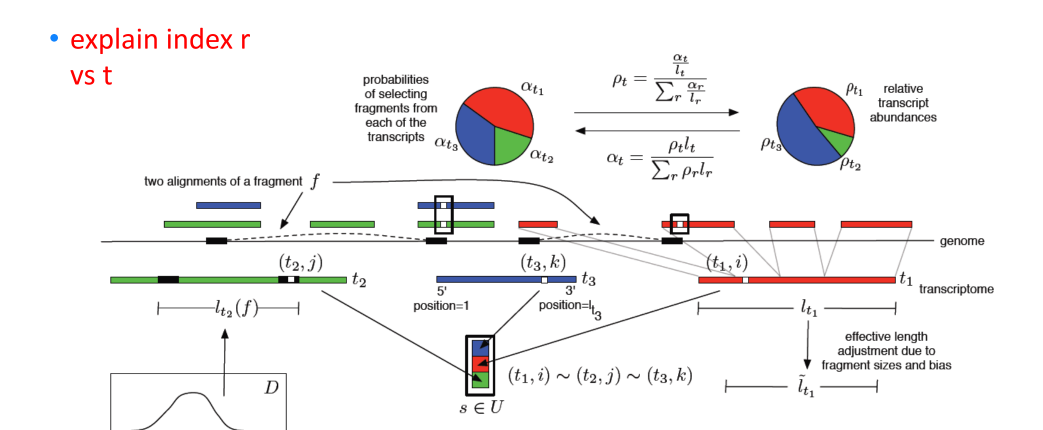
Iterative Estimation

Three step algorithm

- 1. Estimate abundances based on uniquely mapping reads only
- 2. For each multi-read, divide it between the transcripts to which it maps, proportionally to their abundances estimated in the first step
- 3. Recompute abundances based on updated counts for each transcript
- 4. Continue with Step 2

Expectation-Maximization Estimation







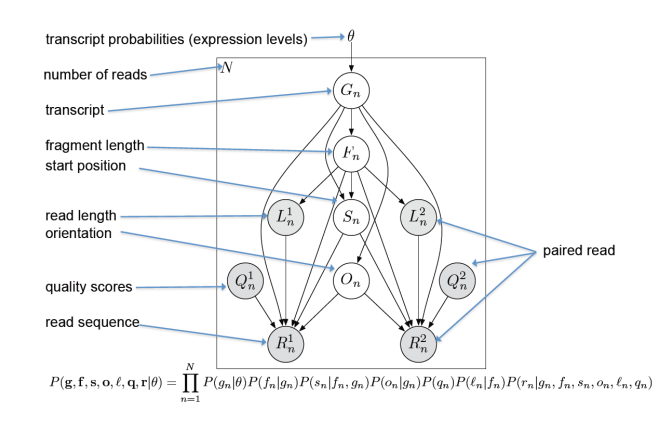
General Formulation of Abundance Estimation

A full model for the abundance estimation would ideally consider:

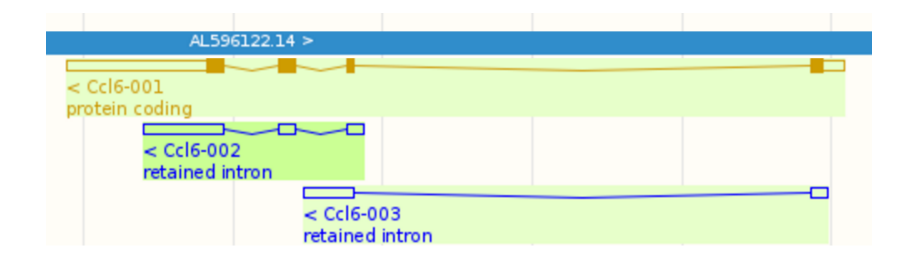
- position bias
- fragment-length distribution
- sequencing errors
- site-specific bias
- •

RSEM: Model specification

Dewey: RSEM



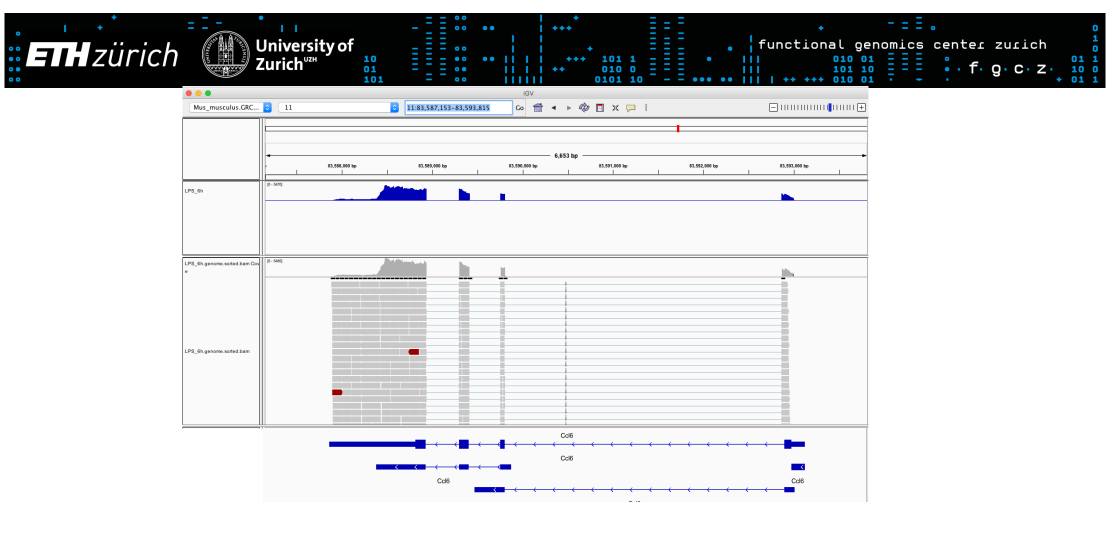
Example: RSEM



Ccl6 gene locus with 3 isoforms

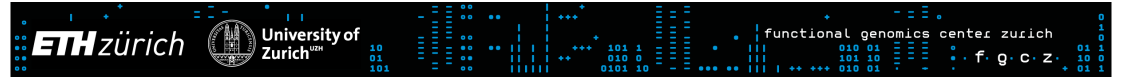
follows the example:

https://github.com/bli25broad/RSEM_tutorial

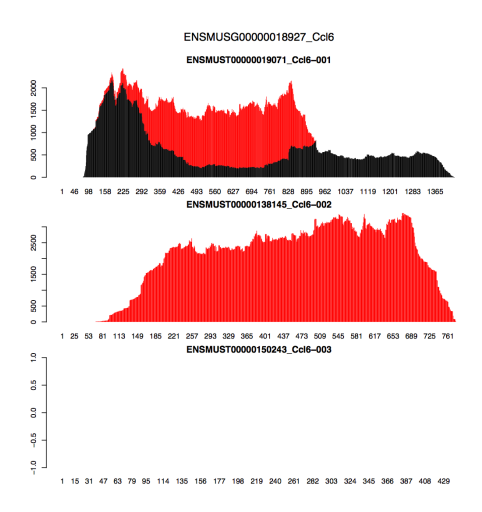


RSEM result:

transcript_id effective_length expected_count TPM **FPKM** IsoPct gene_id length ENSMUSG00000018927_Ccl6 1440 ENSMUST00000019071_Ccl6-001 1194.85 7805.95 8862.10 9334.46 31.00 ENSMUST00000138145_Ccl6-002 ENSMUSG00000018927_Ccl6 776 530.94 7719.05 19721.39 20772.55 69.00 ENSMUST00000150243_Ccl6-003 ENSMUSG00000018927_Ccl6 442 202.64 0.00 0.00 0.00 0.00



Ccl6 coverage in transcript space



- orientation is flipped because gene is on negative strand
- black: unique alignments
- red: expected depth from multi-mapping reads

Limitations of Generative Models

- Estimates can not be correct if underlying model of transcripts are incorrect or incomplete
- Abundance estimates are fractions; these can be used to get estimates of the number of reads generated by a given gene; error distribution of estimated read counts may be unclear

Implementation of Generative Models

• RSEM:

Li, B. & Dewey, C. N. RSEM: accurate transcript quantification from RNA-Seq data with or without a reference genome. *BMC Bioinformatics* **12**, 323 (2011).

• Review:

Pachter, L. Models for transcript quantification from RNA-Seq. *arXiv preprint arXiv:1104.3889* (2011).

• MISO:

Katz, Y., Wang, E. T., Airoldi, E. M. & Burge, C. B. Analysis and design of RNA sequencing experiments for identifying isoform regulation. *Nat Methods* **7**, 1009–1015 (2010)

• MMSEQ:

Turro, E. et al. Haplotype and isoform specific expression estimation using multi-mapping RNA-seq reads. *Genome Biol* **12**, R13 (2011).

NSMAP:

Xia, Z., Wen, J., Chang, C.-C. & Zhou, X. NSMAP: a method for spliced isoforms identification and quantification from RNA-Seq. *BMC Bioinformatics* **12**, 162 (2011).

Definition of expression levels

- Goal: Start from read counts and define a quantity that indicates relative molar concentration of a transcript
- Reads Per Kilobase per Million of mapped reads

RPKM for transcript
$$t = 10^6 \times 10^3 \times \frac{X_t}{l_t N}$$

Transcripts Per Million Transcripts

TPM for transcript
$$t = 10^6 \times Z \times \frac{X_t}{l_t N}$$

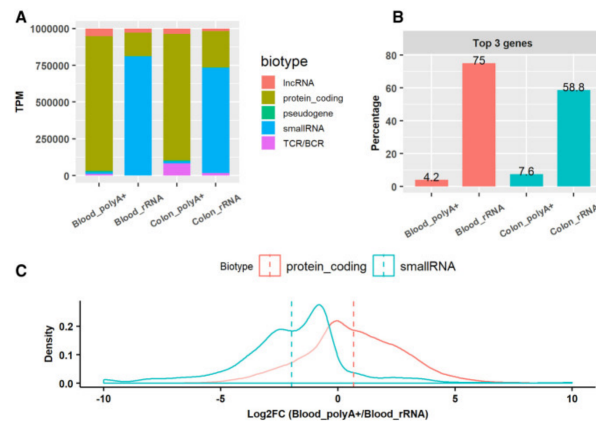
Relationship

$$TPM = 10^6 * \frac{RPKM}{Sum(RPKM)}$$

Shortcomings of RPKM and TPM

- Sum of RPKM varies from sample to sample, i.e. RPKM is not a measure of relative concentration because the measures of relative concentrations would sump up to constant
- TPM is unitless and satisfies this requirement
- Only TPM should be used!
- But: even TPM is not a suitably normalized measure that can be used to compare samples from
 - different tissues
 - different protocols

Comparing samples across protocols

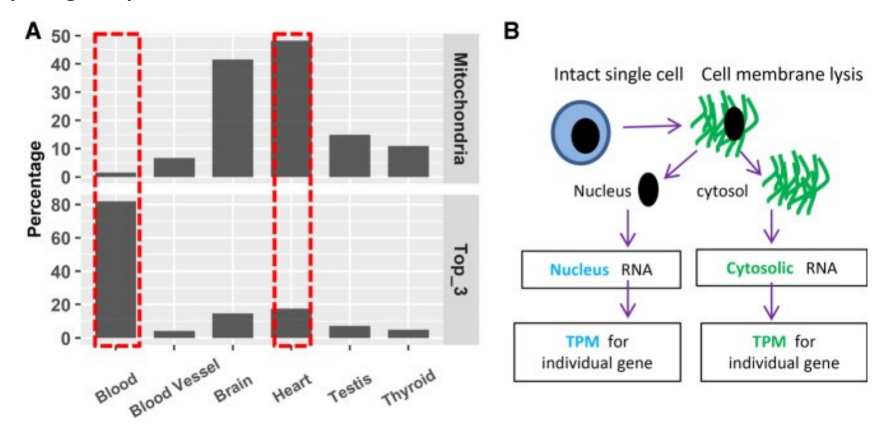


• surveyed populations are not comparable

issues:

- expression of top 3 genes will drive the TPM normalization (because it has a
- major influence on the sum of all reads)

Comparing samples across tissues



Different tissues may have different populations of genes expressed



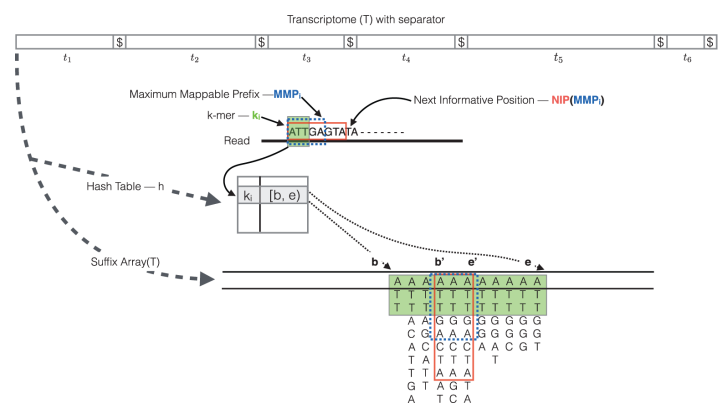
Fast approaches to get the Read-Transcript Compatibility Matrix

Salmon: quasi-mapping

kallisto: pseudo-alignments



Quasi-mapping



Default k-mer size: 31

https://hbctraining.github.io/Intro-to-rnaseq-hpc-salmon/lessons/04 quasi alignment salmon.html

Quasi-mapping

- 1. The read is scanned from left to right until a k-mer that appears in the hash table is discovered.
- 2. The k-mer is looked up in the hash table and the SA intervals are retrieved, giving all suffixes containing that k-mer
- 3. Similar to STAR, the maximal matching prefix (MMP) is identified by finding the longest read sequence that exactly matches the reference suffixes.
- 4. Salmon identifies the next informative position (NIP), by skipping ahead 1 k-mer (speedup)
- 5. Repeat above until the end of the read.
- 6. The final mappings are generated by determining the set of transcripts appearing in all MMPs for the read. The transcripts, orientation and transcript location are output for each read.
- hash table vs SA intervals

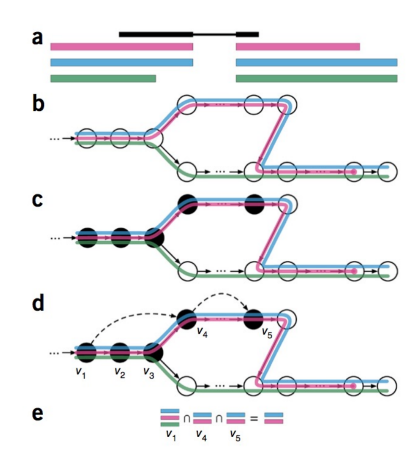
Quasi-mapping

- Result: Read-Transcript compatibility matrix
- Only based on compatibility of short k-mers
- Has an optional step to validate mappings:
 - goes through all the read-transcript associations and validates if the entire read is compatible with the transcripts by doing a base-by-base comparison
- use kingsford-regulatory-genomics-salmon.pdf or later publications ..



kallisto: Quantification with pseudo-alignments

- Instead of hashing the transcriptome build a de Bruijn graph
- Find k-mer hits in the de Bruijn graph
- Identifies only transcripts that are consistent with all k-mer hits



Performance comparison

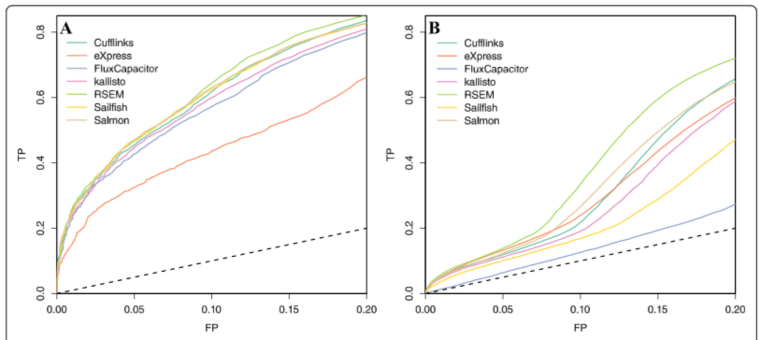
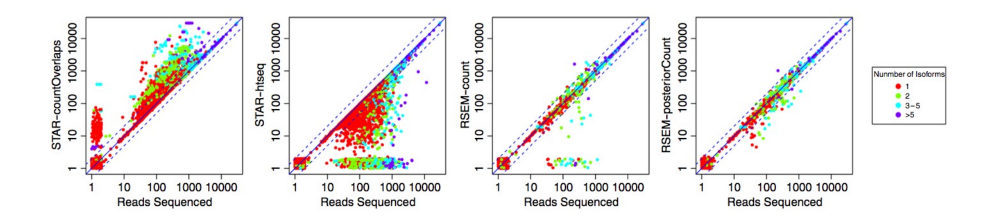


Fig. 6 ROC curves indicating performance of quantification methods based on differential expression analysis of **a** an experimental dataset and **b** a simulation dataset. Seven quantification methods are shown. *FP* false positive, *TP* true positive

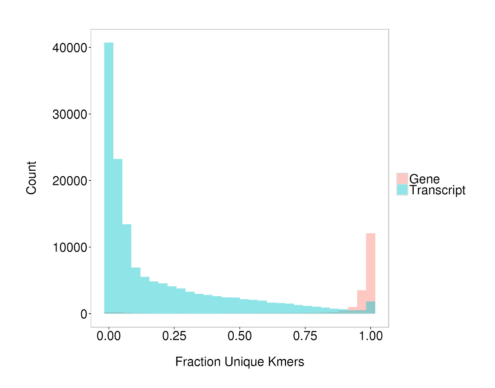
Read Counting Accuracy



remove



Uniqueness: Isoform-level vs gene-level

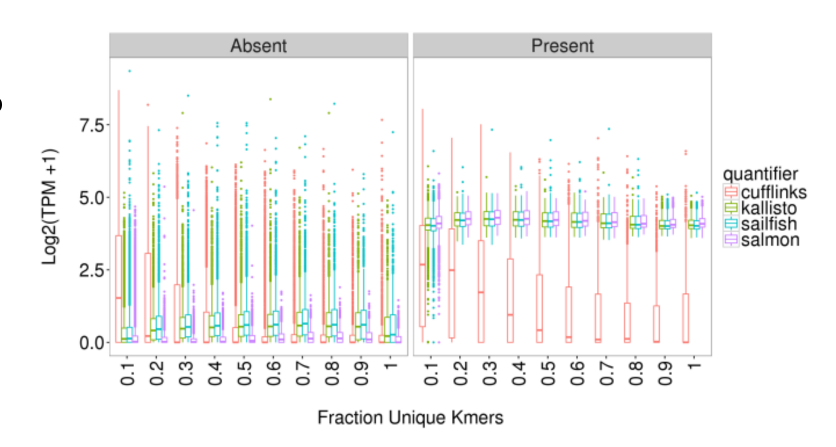


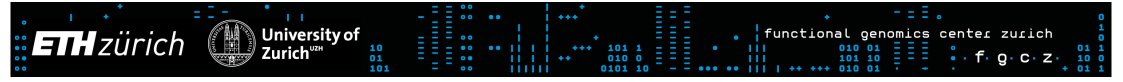
- Fraction of unique k-mer sequences for genes and transcripts
- Ambiguity is mainly between alternative transcripts from the same gene locus



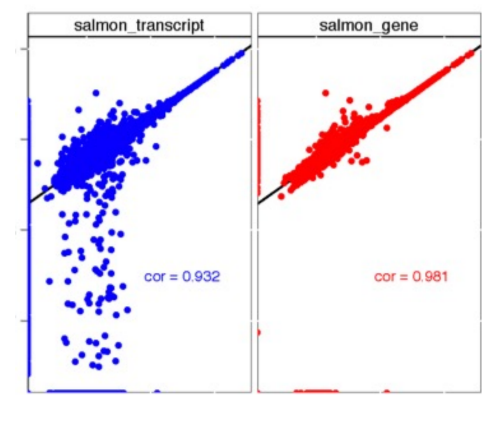
Misassignments to unexpressed transcripts

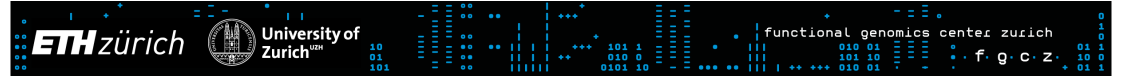
 Simulated data show that misassignments do happen



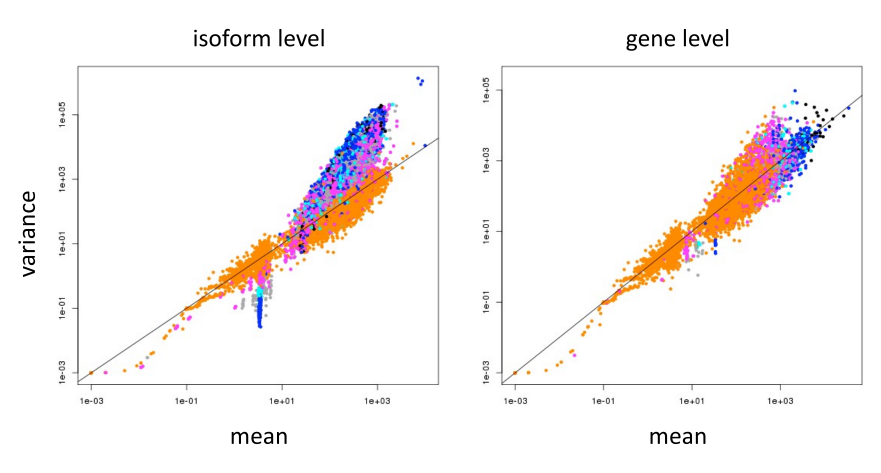


Accuracy: Isoform-level vs gene-level





Isoform level has higher variability



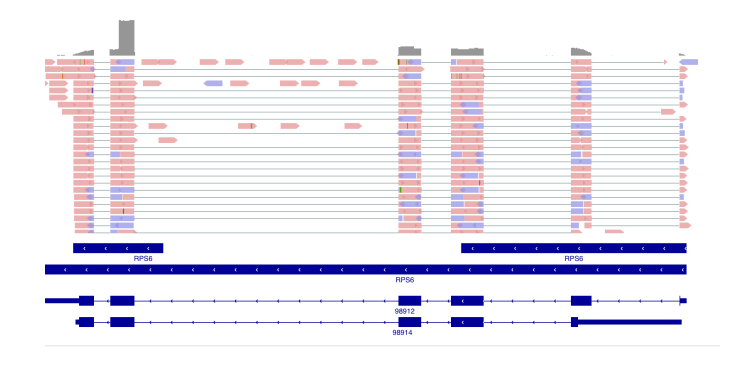
explain colors or remove

Positional bias of read distributions



Highly Multiplicated Reads

- Mainly a concern for low starting amounts
- <1ng of total RNA</p>

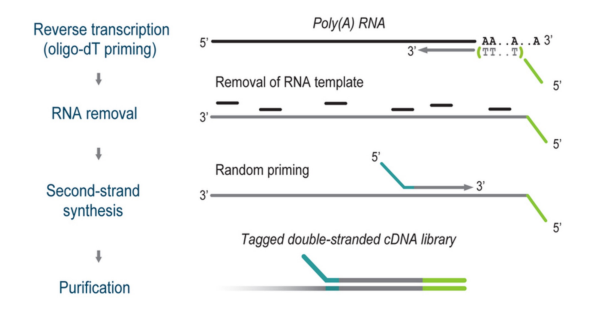


Unspliced transcripts

- Isoform quantification assumes that only spliced transcripts have been measured
- But: unspliced transcripts are also present:
 - these are transcripts from the nucleus that are not yet spliced
 - limited capturing with poly-A based protocols
 - fully captured by random-priming protocols (1 10% of mRNA is in the nucleus)

3'-Tagging

Reads are only generated near the 3'-end Isoforms can not be resolved Allows counting of the reads at the 3'-end → Assignment is to genes



Summary

- Alignment + expression estimation using generative models give good results, let you inspect the aligned reads, and can be used to discover new genes and new isoforms
- Pseudo-alignment is reliable and fast but needs as input the accurate and complete set of transcripts
- 3'-end tag sequencing provides only gene-level estimates without isoform resolution
- Full-length transcript sequencing detects isoforms accurately, sequencing depth is typically lower than for short-read sequencing

References

- Compares kallisto, salmon, featureCounts, ... https://bmcbioinformatics.biomedcentral.co m/articles/10.1186/s12859-021-04198-1
- from the salmon people, performance evaluation and tuning options (especially genomic "decoy") https://genomebiology.biomedcentral.com/articles/10.1186/s13059-020-02151-8
- from the salmon people, confirm that decoys help https://www.biorxiv.org/content/10.1101/202

 1.01.17.426996v1

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Review Open access | Published: 25 May 2021
Comparative evaluation of full-length isoform
quantification from RNA-Seq
 Dimitra Sarantopoulou, Thomas G. Brooks, Soumyashant Nayak, Antonijo Mrčela, Nicholas F.
 Lahens & Gregory R. Grant □
BMC Bioinformatics 22, Article number: 266 (2021) | Cite this article
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  Alignment and mapping methodology influence
   transcript abundance estimation
    Avi Srivastava, Laraib Malik, Hirak Sarkar, Mohsen Zakeri, Fatemeh Almodaresi, Charlotte Soneson
    Michael I. Love, Carl Kingsford & Rob Patro 

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    Genome Biology 21, Article number: 239 (2020) | Cite this article
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         quantification in RNA-seq simulations focused on increased realism
         📵 Avi Srivastava, 📵 Mohsen Zakeri, 📵 Hirak Sarkar, 📵 Charlotte Soneson, 📵 Carl Kingsford, 📵 Rob Patro
         doi: https://doi.org/10.1101/2021.01.17.426996
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