

# **Computational Biology** (BIOSC 1540)

## Lecture 07B

Quantification

Methodology

Feb 20, 2025



# Announcements

**Assignments** • P02A is due March 14

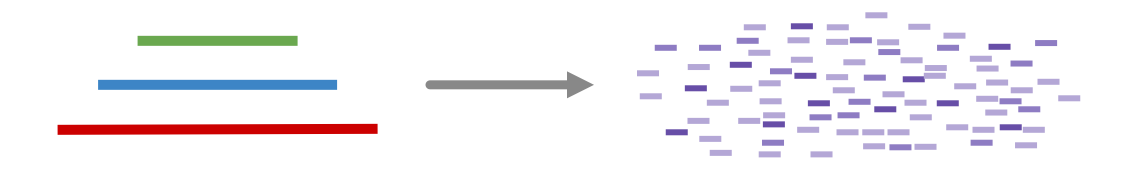
• Quiz 03 is on Mar 18 and will cover L06B to L08B

• César will provide optional Python recitations on Fridays from 2 - 3 pm (Located in Clapp Hall, room TBD).

#### **RNA** quantification problem formulation

### The RNA quantification problem statement

Given the sequencing reads that were sampled from these transcripts



Transcriptome

Unknown quantity

#### **Reads/Fragments**

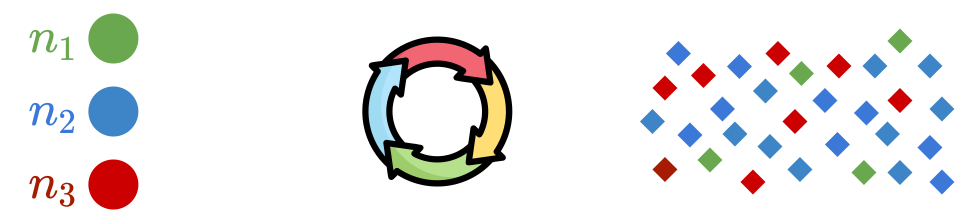
Experimental biases and errors

How many copies of each transcript were in my original sample?

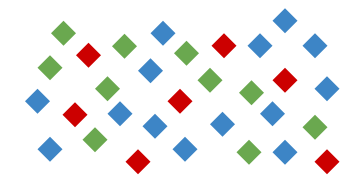
We need to maximize the probability that our generative model and parameters explain our observations

**1.** Estimate transcript abundance

**2.** Randomly sample *n* fragments

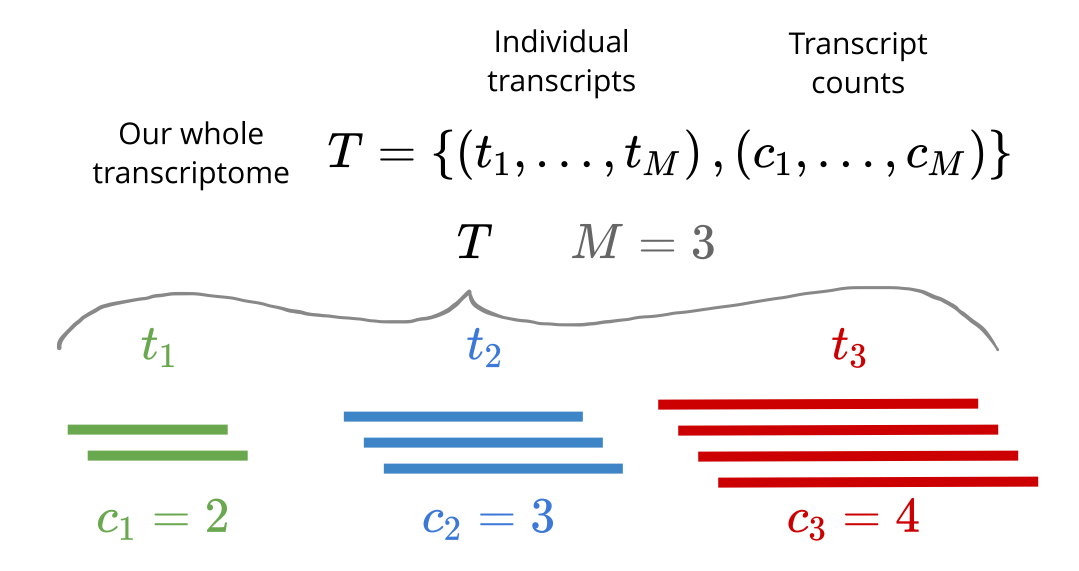


We iteratively optimize our transcript abundances until our generated reads look very similar to our observed reads

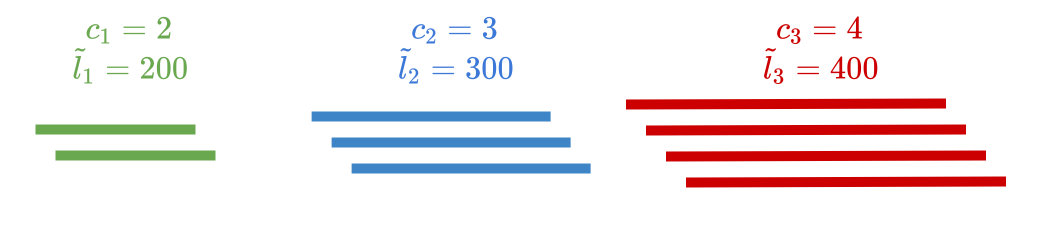


#### Generative models for RNA quantification

#### Salmon's mathematical definition of a transcriptome



## Salmon's formulation of transcript abundance



So far, we have been talking about transcript fractions

$$f_i = rac{c_i}{\sum_j^M c_j}$$

$$\eta_i = rac{c_i ilde{l}_i}{\sum_j^M c_j ilde{l}_j} \quad \eta = egin{bmatrix} \eta_1 \ \eta_2 \ \eta_3 \end{bmatrix}$$

We can also take nucleotide fractions by taking into account the effective length of each transcript

This tells us how much of the total RNA pool comes from each transcript

I will explain the effective length later. For now, think of it as a "corrected" length

#### **Converting to relative abundances**

 $\tau_i \quad \begin{array}{l} \text{The transcript fraction normalizes} \\ \text{nucleotide fraction by the effective length} \end{array}$ 

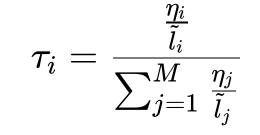
Adjusts for the fact that longer transcripts generate more reads

This gives the relative abundance of each transcript *i* 

 $\mathrm{TPM}_i = au_i \cdot 10^6$ 

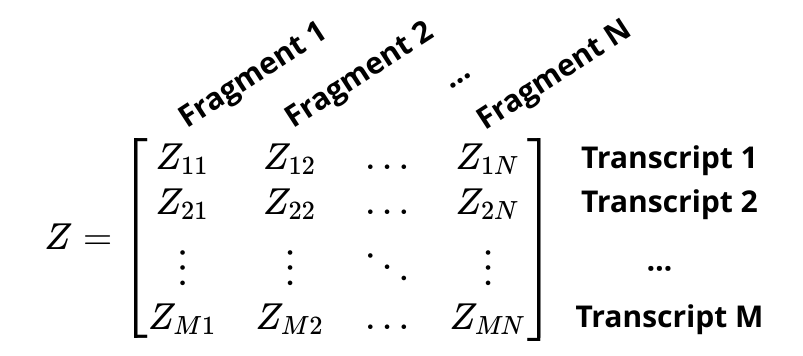
The **transcript fraction** tells us the proportion of total RNA molecules in the sample that come from transcript *i* 

**TPM** is "Transcripts per million"



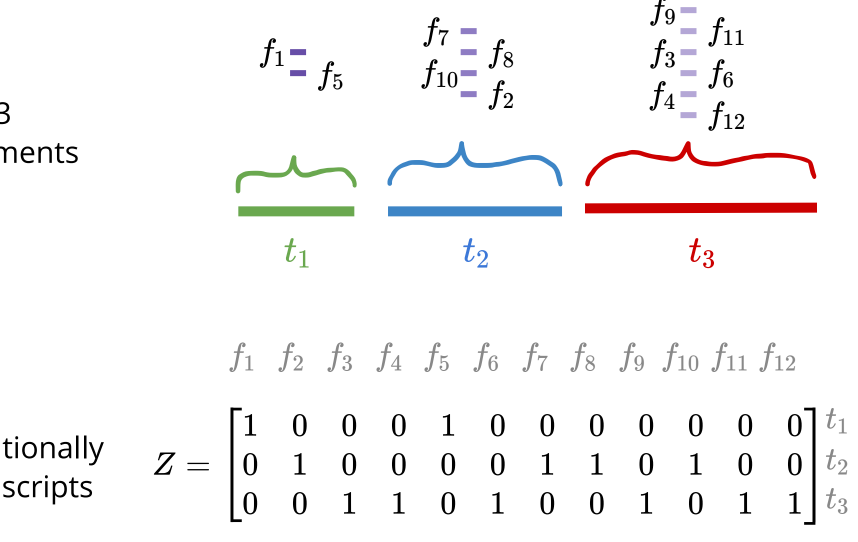
#### **Transcript-Fragment Assignment Matrix**

*Z* is a binary matrix (i.e., all values are 0 or 1) of *M* transcripts (rows) and *N* fragments (columns)



 $Z_{i,j} = 1$  if fragment j is assigned to transcript i

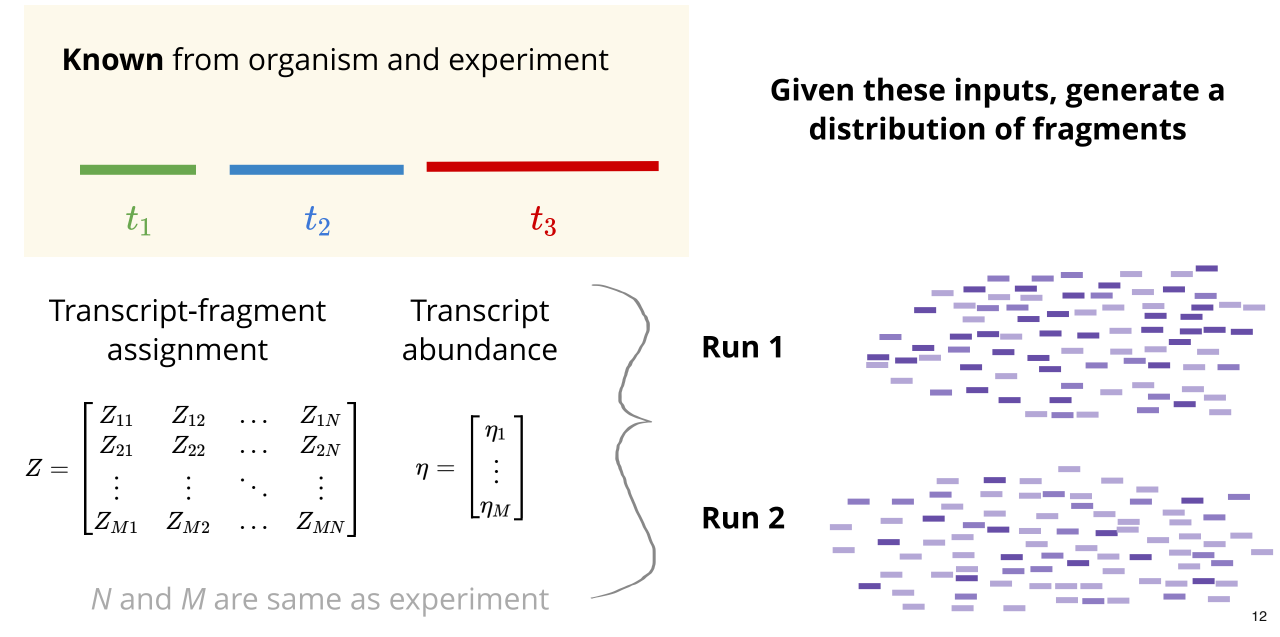
#### Z example



# Suppose we have 3 transcripts and 12 fragments

*Z* is just how we computationally assign fragments to transcripts

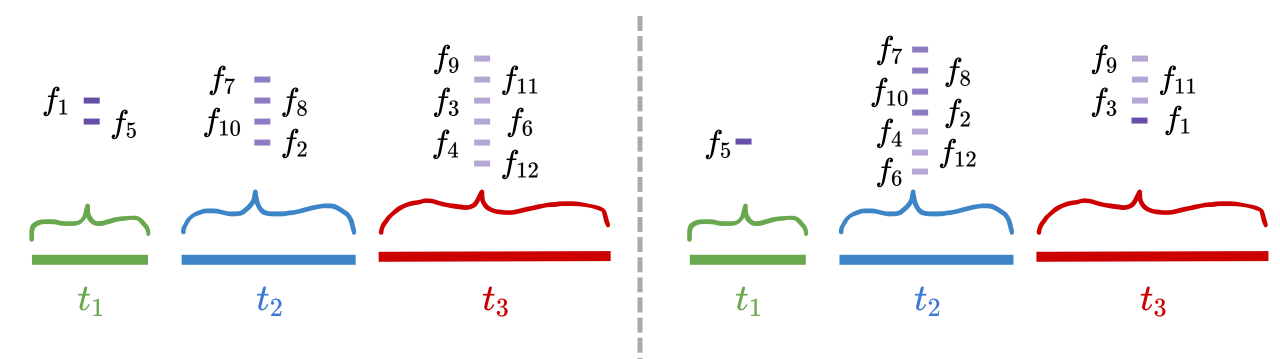
## **Generative model inference**



#### Probability of observing the sequence fragments

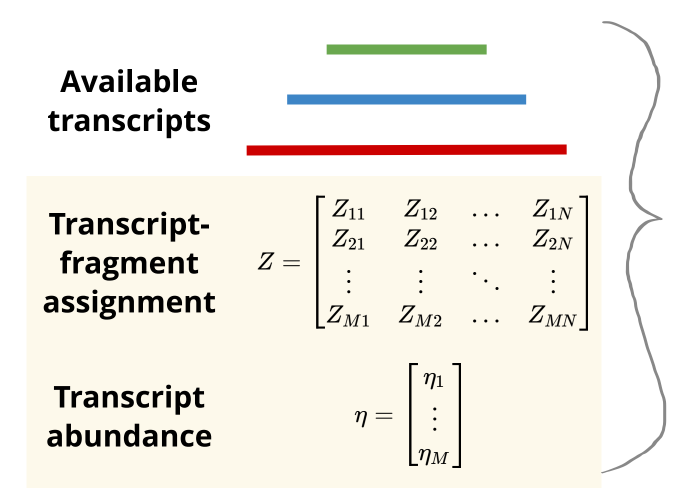
Which scenario is more likely, given our generative model?

We can use probabilistic methods to find parameters that explain our observed distirbution

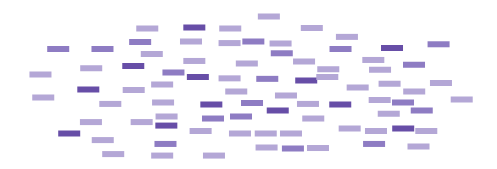


### Probability of observing the sequenced fragments

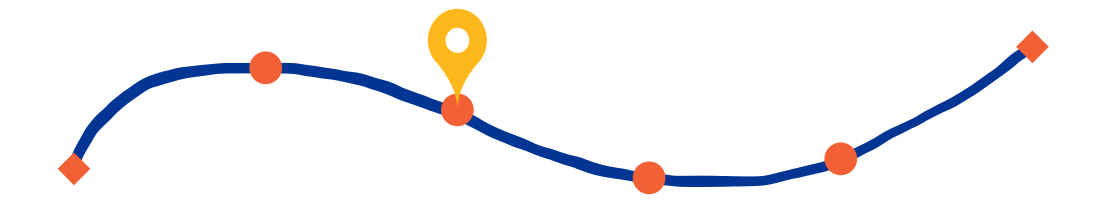
 $P\left(F|T,\eta,Z
ight)$ 



Given these **parameters**, how probable is it that our experiment generated these observed reads?



Optimize these values until we get the highest probability



#### Probability optimization instead of generation

## Probability of observing the sequenced fragments

We can now compute the probability of observing: Set of fragments  $\,F\,$ 

Given: Transcriptome T Transcript assignment Z Transcript abundance  $~\eta$ 

 $P\left(F|\eta,Z,T
ight) = \prod_{j=1}^{N}\sum_{i=1}^{M}\eta_{i}P\left(f_{j}|t_{i}
ight)$ 

 $P\left(f_{j}|t_{i}
ight)$ 

Probability of observing fragment  $f_j$ 

given that it comes from transcript  $t_i$ 

This expression accounts for all possible transcripts a fragment might come from, weighted by how likely that fragment is to come from each transcript **Fragment probabilities** 

 $P\left(f_{j}|t_{i}
ight)$ 

is a conditional probability that depends on the **position** of the fragment within the transcript, the **length** of the fragment, and any technical biases

In Salmon's quasi-mapping approach, this probability is approximated based on transcript compatibility rather than exact positions.

 $P(f_j|t_i) = P(\text{fragment length}, \text{position}, \text{GC content}, \ldots)$ 

## **Positional bias**

Fragments that include transcript ends might be too short

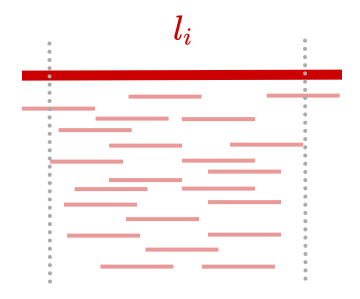
Fragments from central regions are more likely to be of optimal length for sequencing reads

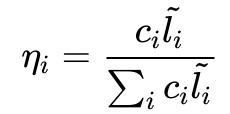
A transcript's **effective length** adjusts for the fact that fragments near the ends of a transcript are less likely to be sampled

$$ilde{l}_i = l_i - \mu_i \qquad \quad ilde{l}_i < l_i$$

Mean of the truncated empirical fragment length distribution

 $\mu_i$ 





18

#### Probability maximization with inference

## Two-phase inference in salmon

**Inference** refers to the process of estimating transcript abundances from observed RNA-seq reads using statistical models.

Salmon processes reads in **two stages** 

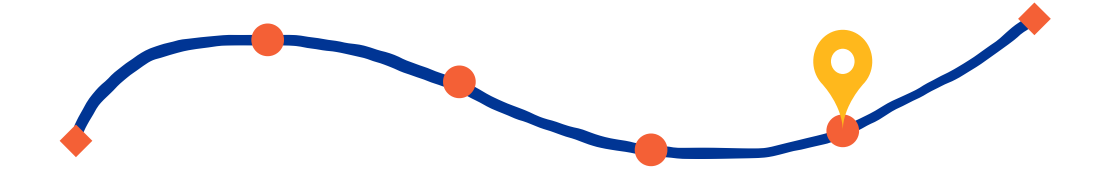
#### **Online phase**

Makes fast, initial estimates of transcript abundances as the reads are processed

#### **Offline phase**

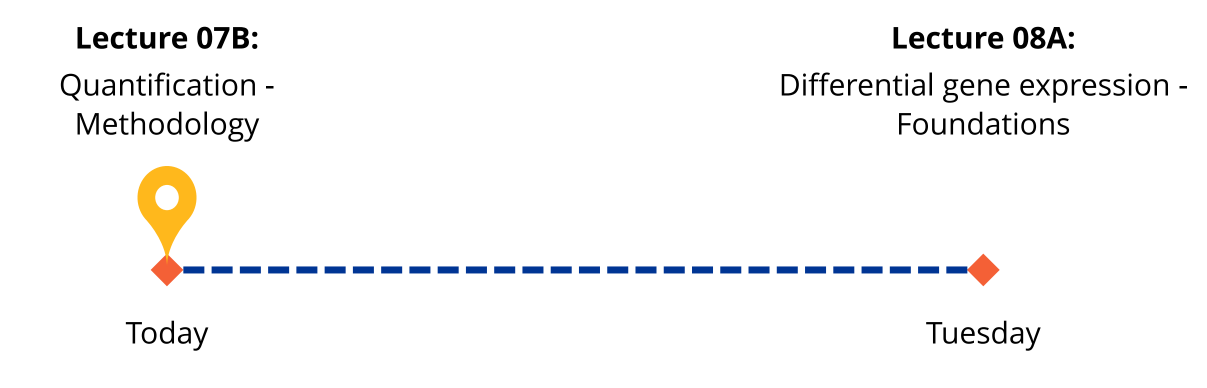
Refines these initial estimates using more complex optimization techniques

This two-phase approach balances **speed** (in the online phase) with **accuracy** (in the offline phase)



#### Methodology with a Python implementation

## Before the next class, you should



• Work on P02A (due Mar 14)