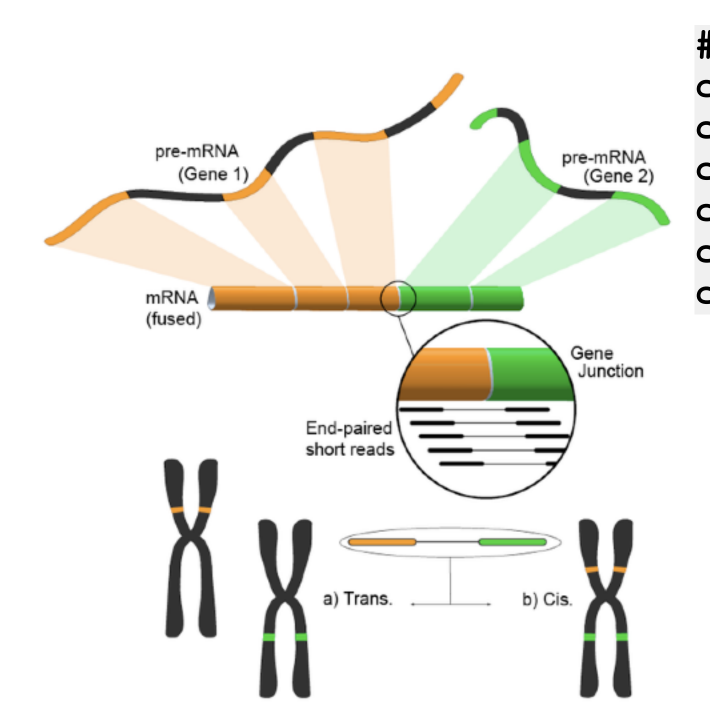
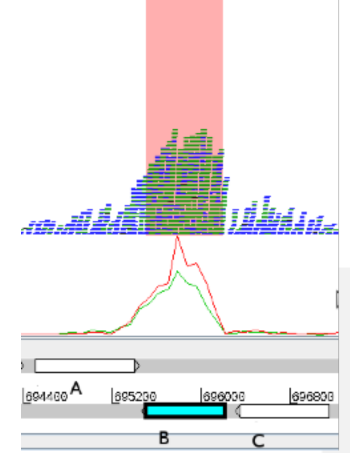
## Contar o que?



- A medida base é o número de reads.
- Contam-se o número de reads mapeadas a uma feature.
- Reads curtos
- Single read (quantificação) paired-end (splice)



##gff-v	ei	rsion 3							
ctg123	•	mRNA	13	300	9000	•	+	•	ID=mrna0001; Name=sonichedgehog
ctg123	•	exon	13	300 :	1500	•	+	•	<pre>ID=exon00001;Parent=mrna0001</pre>
ctg123	•	exon	10	050	1500	•	+	•	<pre>ID=exon00002;Parent=mrna0001</pre>
ctg123	•	exon	3	000	3902	•	+	•	<pre>ID=exon00003;Parent=mrna0001</pre>
ctg123	•	exon	5	000	5500	•	+	•	<pre>ID=exon00004;Parent=mrna0001</pre>
ctg123	•	exon	7	000	9000	•	+	•	<pre>ID=exon00005;Parent=mrna0001</pre>



## Quem expressa mais?



	Exp1	Exp2	Exp3	Ctrl1	Ctrl2	Ctrl3
Gene A	0.010%	0.015%	0.012%	0.02%	0.030%	0.029%
Gene B	0.0072%	0.0062%	0.0095%	0.0031%	0.0040%	0.0015%
Gene C	1%	1.5%	1.2%	2%	3%	2.9%
Gene D	72%	62%	95%	31%	40%	15%