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Simulation of RNA sequencing with Oxford Nanopore Technologies

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**Introduction**

Until recently, transcriptomics applications with long reads were realized with Pacific Biosciences’ Iso-seq protocol. Pioneer works start dealing with characterization of isoforms or gene expression quantification using Oxford Nanopore Technologies (ONT) reads.

Despite the existence of long reads simulators for genomic data sets, a current lack is the possibility to adequately simulate long reads from ONT RNA protocols, which would help with the developments of new tools to handle this kind of data. The simulation of transcriptomics sequencing is a more complex task than in genomes because the gene expression and transcript variability have to be modeled.

**Methods**

**Error rates and profiles**
- learn error rates and profiles by training using real read data sets such as Nanosim [1]
- compute error rates and percentages of deletion, insertion and substitution, as well as homopolymer errors using AlignQC [2]
- pre-computed error profiles can also be input

**Simulate transcripts**
- extract template transcripts from the GTF file of the desired reference using gffreads (http://ccb.jhu.edu/software/stringtie/gff.shtml)
- extract realistic expression levels from Flux Simulator [3] results

**Integrate errors and transcripts in synthetic reads**
- novel implementation in C++ that generates the final reads
- adds the errors to the sequences extracted from the GTF
- deals with regular versus homopolymer errors
- adds supplementary characteristics such as the staircase effect

**Experiment**
- 300k reads simulation
- Genome reference: *Mus musculus* GRCm38
- Annotation: Ensembl *Mus musculus* GRCm38 release 87
- Template dataset for error training: 740k *Mus musculus* RNA ONT 1D reads
- Comparison of raw reads and simulated reads statistics using AlignQC [2]

**Results**

Comparing error rates between ONT 1D RNA reads and the simulated reads shows that the simulator produces reads with realistic error profiles:

<table>
<thead>
<tr>
<th>Metric</th>
<th>ONT 1D RNA reads</th>
<th>Simulated reads</th>
</tr>
</thead>
<tbody>
<tr>
<td>ERROR RATE</td>
<td>13.697%</td>
<td>11.592%</td>
</tr>
<tr>
<td>MISMATCHES</td>
<td>5.089%</td>
<td>5.848%</td>
</tr>
<tr>
<td>DELETION</td>
<td>7.348%</td>
<td>4.85%</td>
</tr>
<tr>
<td>INSERTION</td>
<td>1.26%</td>
<td>0.894%</td>
</tr>
<tr>
<td>NON HOMOPOLYMER DELETION</td>
<td>4.397%</td>
<td>3.069%</td>
</tr>
<tr>
<td>HOMOPOLYMER DELETION</td>
<td>2.951%</td>
<td>1.782%</td>
</tr>
<tr>
<td>NON HOMOPOLYMER INSERTION</td>
<td>0.874%</td>
<td>0.558%</td>
</tr>
<tr>
<td>HOMOPOLYMER INSERTION</td>
<td>0.387%</td>
<td>0.337%</td>
</tr>
</tbody>
</table>

The distribution of reads among genomic features is similar between the real and simulated reads:

As well as the distribution of annotated reads:

And the distribution of identified reference transcripts:

**Conclusion**

**Main messages**
- Work in progress, first tool to simulate transcriptome sequencing with ONT
- Availability: github.com/kamimrcht/RNA-long-reads-Simulator

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**References**

