zUMIs A fast and flexible pipeline to process RNA sequencing data with UMIs

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Abstract

RNA sequencing is increasingly performed with less starting material and at a higher sample throughput, e.g. to analyse single-cell transcriptomes. In this context, unique molecular identifiers (UMIs) are used to reduce amplification noise and sample-specific barcodes are used to track libraries. Here, we present a fast and flexible pipeline to process data from such RNA-seq protocols.

Availability: https://github.com/sdparekh/zUMIs

1 Introduction

The recent development of sensitive protocols allows to generate RNA-seq libraries of single cells [1]. The throughput of such scRNA-seq protocols is rapidly increasing, enabling the profiling of tens of thousands of cells [2, 3] and opening exciting possibilities to analyse cellular identities [4, 5]. As the required amplification from such low starting amounts introduces substantial amounts of noise [6], many scRNA-seq protocols incorporate unique molecular identifiers (UMIs) to label individual cDNA molecules with a random nucleotide sequence before amplification [7]. This allows to computationally remove amplification noise and thus increases the power to detect expression differences [8, 9]. To increase the throughput, many protocols also incorporate sample-specific barcodes (BCs) to label all cDNA molecules of a single cell with a nucleotide sequence before library generation [10, 2]. Additionally, for cell types such as neurons it has

| Name | Reference | Open Source | Quality UMI/BC | Mapper | intron counting | Down- sampling |
|------------|-----------|----------------|-------------------|----------|--------------------|-------------------|
| CellRanger | [2] | no | no | STAR | no | yes |
| Drop-seq | [10] | no | yes | STAR | no | no |
| CEL-seq | [13] | yes | yes | bowtie2 | no | no |
| umis | [14] | yes | no | Kallisto | no | no |
| zUMIs | This work | yes | yes | STAR | yes | yes |

Table 1. Pipelines handling UMI expression data

proven to be more feasible to isolate RNA from single nuclei rather than whole cells [11, 12]. This decreases mRNA amounts further, so that it has been suggested to count intron-mapping reads as part of nascent RNAs. However, the few bioinformatic tools that process RNA-seq data with UMIs and BCs have limitations with respect to availability, mapping, quality assessment and/or can not consider intronic reads (Table 1). Here, we present zUMIs, a fast and flexible pipeline to overcome such limitations.

2 zUMIs

zUMIs is a pipeline that processes paired fastq files containing the UMI and BC in one read and the cDNA sequence in the other read, filters out reads with bad BCs or UMIs based on sequence quality, maps reads to the genome and outputs count tables of unique UMIs or reads per gene (Figure 1). To allow the quantification of intronic reads that are generated from unspliced RNAs especially when using nuclei as input material, three separate count tables for exons, introns and exon+introns are provided. Another unique feature of zUMI is that it allows for downsampling of reads before summarizing UMIs per feature, which is recommended for cases of highly different read numbers per sample [15]. zUMIs is flexible with respect to the length and sequences of the BC and UMIs, making it compatible with a large number of protocols [16, 17, 10, 13, 3, 2].

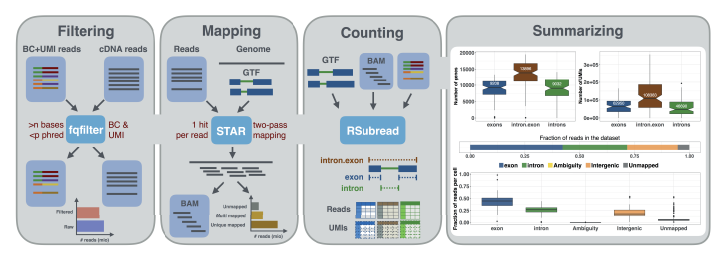


Figure 1. zUMIs schematic overview.

A Each of the grey panels from left to right depicts a step of the zUMIs pipeline. First, paired fastq files are filtered according to user-defined BC and UMI quality thresholds. Next, the remaining cDNA reads are mapped to the reference genome using STAR. Then gene-wise read and UMI count tables are generated for exon, intron and exon+intron overlapping reads. To obtain comparable library sizes, reads can be downsampled to a desired range during the counting step. Optionally, zUMIs also generates data and plots for several quality measures, such as the number of detected Genes/UMIs per barcode and distribution of reads into mapping feature categories (Supplementary Figure 3).

2.1 Processing pipeline

The input for zUMIs is a pair of fastq files, whereas one file contains the cDNA sequences and the other one the read containing the BC and UMI. The exact location and length of UMI and BC are specified by the user. Note that both fastq files need to be ordered by read name, which is usually the case if unprocessed files are used. The first step in our pipeline is to filter reads where the BC or the UMI fails a user-defined quality threshold. This helps to eliminate spurious BCs and is expected to reduce noise. The cleaned-up reads are then mapped to the genome using the splice-aware aligner STAR [18]. The user is free to adapt the STAR options to their data, however zUMIs requires that only one mapping position per read is reported. Next, reads are assigned to genes and to exons or introns based on the provided gtf file, whereas introns are defined as not overlapping with any exon. Rsubread featureCounts [19] is used to first assign reads to exons and afterwards to check whether the remaining reads fall into introns. The resulting output is then read into R using data.table [20] and count tables for UMIs and reads are generated. zUMIs tabulates the UMIs/gene either for user-specified BCs or for the *n* BCs with the highest read counts.

2.2 Output and statistics

zUMis outputs three UMI and three read count tables: one for traditional exon mapping gene-wise counts, one for intron and one for intron+exon counts. If a user chooses the downsampling option, 6 additional count-tables are provided in which samples with an excess of reads are downsampled and samples with too few reads are dismissed (Supplementary Figures 4). We highly recommend to use this option, because normalizing across samples with vastly different library sizes does not work well [15, 21]. zUMIsalso reports descriptive statistics. To evaluate library quality zUMIs summarizes the fractions of unmapped, ambiguously mapped, exon and intron mapped reads and to evaluate library complexity, the numbers of detected genes and UMIs per sample are provided (Supplementary Figures 2,3).

We processed 227 million reads with zUMIs and quantified expression levels for exonic and intronic counts on a unix machine using up to 16 threads, which took barely 3 hours. Increasing the number of reads increases the processing time approximately linearly, whereas filtering, mapping and counting each take up roughly one third of the total time (Supplementary Figure 1).

3 Conclusions

zUMIs is a fast and flexible pipeline to process raw reads to count tables for RNA-seq data using UMIs. To our knowledge it is the only open source pipeline that has a barcode and UMI quality filter, allows intron counting and has an integrated downsampling function (Table 1). These features ensure that zUMIs is applicable for most experimental designs of RNA-seq data, such as single-nuclei sequencing techniques [11, 12, 22], droplet based methods where the BC is unknown and the library sizes can vary a lot as well as plate-based UMI-methods with known BCs.

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Availability

The pipeline is freely available at https://github.com/sdparekh/zUMIs.

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