### Gene regulation

Master Thesis in Biostatistics (STA495)

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# Preface

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# Abstract

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### Chapter 1

### Introduction

### 1.1 RNA sequencing (RNA-seq)

RNA sequencing (RNA-seq) is a technology for detecting and quantifying the mRNA molecules of a biological sample (Stark et al., 2019). The invention of RNA-seq was a major breakthrough in the field of bioinformatics that replaced the use of microarray technology in the late 00's. In comparison to microarrays, RNA-seq allows for full sequencing of the whole transcriptome whereas microarrays only profile predefined transcripts through hybridization (Rao et al., 2019). Further, various protocols have since been derived from the standard RNA-seq protocol e.g. single-cell RNA sequencing (Stark et al., 2019).

### 1.1.1 Bulk RNA-seq

RNA sequencing has been used with samples that are composed of a mixture of cells, which is referred to as bulk RNA-seq. There are many applications for bulk RNA-seq. For example, it can be used to study the differences of expression profiles between tissues in healthy vs disease (Stark et al., 2019). However, with bulk RNA-seq one can only estimate the average expression of each gene across a population of cells without regard for the differences between cell types. RNA-seq has several use cases. It can be used to study which genes are turned on in a cell and what their level of transcription is. This allows researchers to understand the biology of a cell on a deeper level. Further, RNA-seq allows the identification of variants and allele specific expression. It is also possible to study the patterns of alternative splicing, which are important to understand the contribution to cell differentiation and human disease.

### 1.1.2 Single-cell RNA-seq (scRNA-seq)

Single-cell RNA sequencing was developed to overcome some of the limitations of bulk RNA sequencing. With single-cell RNA sequencing it is possible to estimate the distribution of expression levels for each gene across a population of cells. This allows answering new biological questions where cell-specific characteristics are important. However, there are some caveats with scRNA-seq (Haque et al., 2017). scRNA-seq data in general is much more variable than bulk RNA-seq data due to both higher biological and technical variability at single-cell level (Haque et al., 2017). Figure 1.1 shows the typical workflow of a scRNA-seq experiment. Said workflow is broadly summarized by the following steps:

- 1. RNA extraction
- 2. Reverse transcription into cDNA
- 3. Adapted ligation

- 4. Amplification
- 5. Sequencing
- 6. Downstream analysis using bioinformatics tools



**Figure 1.1:** General workflow of single-cell RNA-sequencing experiments (Haque *et al.*, 2017)

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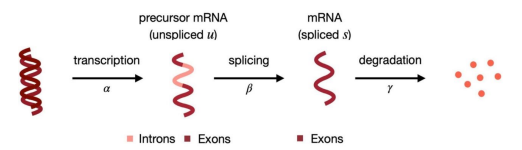
#### 1.1.3 Quantification of single-cell RNA-seq data

scRNA-seq data has distinct characteristics that prevent it from being processed by widely used tools developed for bulk RNA-seq data (He et al., 2022). Compared to bulk RNA-seq, single-cell expression data is significantly noisier due to increased biological and technical variability, such as capture inefficiency, reverse transcription efficiency and cell-size/cell-cycle stage (Haque et al., 2017). In general, quantification works by aligning the reads generated from the RNA-seq to the reference genome. There are several tools that achieve that: STAR (Dobin et al., 2013), kallisto | bustools (Melsted et al., 2021) and alevin (Srivastava et al., 2019). However, there is a difference between the first tool and the latter two. STAR is an aligner, whereas the other two tools are mapping tools (pseudo-aligners). The difference between an full-aligner and a mapping tool is that the latter does not look for the exact location of the read, as a consequence pseudoalignment is much faster than full-alignment. Here we focus on alevin-fry, and the method we have developed, which will be introduced later, has been built to work on the output of alevinfry. Alevin was developed to tackle the computational challenges that come with scRNA-seq data and to provide a tool that supports technologies other than 10x Genomics. Alevin works in two steps. First, it parses a read file which contains the cellular barcode and a unique molecule identifier to generate a frequency distribution of observed barcodes. Second, it maps the reads to the transcriptome and generates a cell-by-gene count matrix. Alevin-fry (He et al., 2022) was designed to be the successor to alevin and achieves similar accuracy at significantly lower computational costs. It generates a permit list for cellular barcodes that will be quantified in subsequent steps. By using a multi-thread approach, alevin-fry filters and collates the mapping records for permitted cellular barcodes to produce a representation optimized for quantification (He  $et \ al., 2022$ ).

### 1.2 Objective

#### 1.2.1 RNA velocity

We investigate spliced and unspliced reads from single-cell RNA-sequencing data. During transcription, DNA is decoded into precursor messenger RNA (pre-mRNA). Pre-mRNA contains both coding (exons) and non-coding regions (introns). In a next step, introns are removed from the pre-mRNA which leaves only the mature mRNA. Figure 1.2 shows the process from DNA to mature mRNA, where  $\alpha$  is the transcription rate,  $\beta$  is the splicing rate and  $\gamma$  is the degradation rate.



**Figure 1.2:** The transcription process from DNA to mature mRNA (Weiler *et al.*, 2021)

It was assumed that there is a signal (RNA velocity) detectable in scRNA-seq data that could reveal the rate and direction of change of an entire transcriptome (La Manno *et al.*, 2018). To quantify the relationship between the abundance of pre-mRNA and mature RNA, a simple

system of ordinary differential equations was assumed (1.1). The equilibrium of said system can then easily be estimated and used to explore the regulation of genes:

$$\frac{du}{dt} = \alpha - \beta u \frac{ds}{dt} = \beta u - \gamma s$$
 (1.1)

The derivative of the spliced counts is then defined as the RNA velocity of cells. Thus, the balance of spliced and unspliced counts allows for estimating whether a gene is up- or down-regulated. If more spliced counts are present than expected at equilibrium, a gene is likely upregulated. Contrarily, if more spliced counts are present at equilibrium than expected, a gene is likely downregulated.

### 1.2.2 Differential regulation

The abundance of spliced and unspliced reads is directly linked to the regulation of genes and RNA velocities. Our idea is to examine how the abundance of spliced and unspliced counts changes between experimental conditions and biological replicates. We translate this intuition into the comparison of two experimental conditions e.g. healthy vs. disease. Following the same intuitive rationale of RNA velocity, if a gene has a higher abundance of unspliced (spliced) counts in group A compared to group B, then this gene is likely being up-regulated (down-regulated) in group A compared to group B.. Thus, we want to explore the differences in abundance of spliced and unspliced counts to be able to study the differences in regulation between experimental conditions.

If the data contains multiple cell clusters (e.g. cell types), similarly to differential state analyses (Crowell et al. (2020) and Tiberi et al. (2021)) we will perform differential analyses in each cluster of cells, hence identifying cell-cluster/cell-type specific changes between conditions. The idea of performing differential analyses on the abundance of spliced and unspliced or exonic and intronic reads is not completely novel as there are at least two other methods that achieve that: eisaR and BRIE2.

### 1.2.3 Existing methods

eisaR (Gaidatzis et al., 2015) is a R package implementation that allows for the split analysis between exons and introns. It allows to measure changes in mature RNA and pre-mRNA across different experimental conditions. Ultimately, eisaR differential testing is based on edgeR (Robinson et al., 2010). edgeR is a R package that performs differential expression analyses between groups of samples. It implements statistical methods that are based on the negative binomial distribution as a model for count variability.

BRIE2 (Huang and Sanguinetti, 2021) is a Bayesian hierarchical model that is implemented in Python and supports the analysis of splicing processes between spliced and unspliced RNA. There are two modes in which the tool can be used. First, the use of differential alternative splicing (DAS), where the aim is to quantify the proportions of alternative splicing isoforms. Second, the use of differential momentum genes (DMG), where the objective is to quantify the proportions of spliced and unspliced RNA in each gene and each cell. In this thesis we focus on the DMG mode as it performs differential testing on the relative abundance of spliced and unspliced reads.

Originally, eisaR and BRIE2 were developed to analyze all cells, but can easily be adapted to perform cell-type specific differential analyses.

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#### 1.2.4 Mapping uncertainty

We can identify two main sources of mapping uncertainty concering spliced and unspliced reads: i) multi-mapping reads across spliced and unspliced versions of a gene, and ii) reads compatible with multiple genes. In fact, it has been shown that many reads (5-40%) map to multiple genes (Dharshini et al. (2020), McDermaid et al. (2018)). In our real data analyses (see Section 3), we found approximately 20-30% of such multi-mapping reads across genes. We additionally found that a significant fraction of reads (6-19%) are compatible with both S and U versions of a gene. Therefore, the estimated spliced and unspliced counts carry a substantial amount of uncertainty, which should be accounted for in downstream analyses. However, both eisaR and BRIE2 use estimated spliced and unspliced counts and neglect the mapping uncertainty. In this thesis, we propose two approaches that account for said mapping uncertainties.

### Chapter 2

### Methods

### 2.1 DEXSeq

DEXSeq (Anders et al., 2012) is a statistical method originally proposed to test for differential exon usage in RNA- seq data, which has been widely adopted in other contexts too, such as differential transcript usage (Love et al., 2018). The model is based on the negative binomial distribution and allows for covariates to be taken into account to offer reliable control of false discoveries (Anders et al., 2012). For each exon and sample it calculates how many reads map to this exon and how many reads map to any other exons of the same gene. The ratio of these two counts is compared across different conditions to infer changes in the relative exon usage. Equation (2.1) shows that the read counts follow a negative binomial distribution where  $\alpha$  is the dispersion parameter. Further, a generalized linear model is used to predict the mean via a log-linear link:

$$K_{ijl} \sim \text{NB}(\text{mean} = s_i \mu_{ijl}, \text{dispersion} = \alpha_{il})$$
 (2.1)

$$log(\mu_{ijl}) = \beta_i^G + \beta_{il}^E + \beta_{i\rho_i}^C + \beta_{i\rho_i l}^{EC}$$
(2.2)

where NB(a, b) denotes the negative binomial distribution with mean a and dispersion b,  $s_j$  is ..

The dispersion parameter allows to model over-dispersed data (i.e. higher variance than mean). Here, we propose to use DEXSeq on estimated USA counts, and perform a differential usage test between conditions. This models ambiguous reads separately from spliced and unspliced or exonic and intronic, thus eliminating one of the main sources of mapping uncertainty. However, the uncertainty related to reads mapping to multiple genes is still neglected by this approach.

### 2.2 Differential regulation

To address both sources of mapping uncertainty we propose our novel method. Similar to the idea above, we implemented a hierarchical Bayesian approach that models ambiguous counts separately from spliced and unspliced. Gene allocation is modeled as a latent state to address the gene-related mapping uncertainty. The model consists of two nested models: First, we use a Dirichlet-multinomial model for the relative abundance of the USA counts in each gene. Second, a multinomial model that models the relative abundance of genes for each sample individually.

### Chapter 3

### Results

### 3.1 Exploratory Data Analysis

### 3.1.1 Mouse kidney cells

The first data set stems from a paper that investigates potential cellular targets of kidney disease in mice (Park et al., 2018). The authors isolated and sequenced a total of 57'979 cells from whole kidney cell suspensions (one kidney per mouse) derived from seven healthy male mice using droplet-based single-cell RNA sequencing. The samples were labeled as: normal1, normal2, normal3, normal4, Ksp-cre-GFP, Scl-cre-GFP and Pod-cre-GFP. After quality controls the authors used clustering analysis on 43'745 cells to classify 16 major cell clusters. Further, cell-type specific marker genes were used to identify the cell clusters. In the end, the authors identified 21 major cell types by quantitative gene expression. For our work, we decided to use the raw data from the four samples that were labeled as categorized as normal to ensure the biological replicability between the samples.

We used the alevin-fry pipeline to quantify the raw single-cell RNA sequencing data for further use in the R programming environment. Quality control is a crucial stage in data preprocessing as low-quality libraries can contribute to misleading results in downstream analyses (Amezquita et al., 2020). The main problems that arise from neglected quality control are the following: First, low-quality libraries can form their own distinct clusters which in the worst-case scenario are generated solely based on similarities in the damage-induced expression profiles. Second, principal component analysis will capture differences between mostly on quality rather than biology. Therefore, we filtered lowly abundant genes and low-quality cells to mitigate said problems to improve interpretability of the results.

To identify low-quality cells, cell-specific QC metrics were calculated with the perCellQCMetrics function from the scater R package (McCarthy et al., 2017). These metrics include the total number of expressed genes, the overall count across all genes, and the fraction of counts assigned to control genes such as mitochondrial genes. By setting a specific threshold on per-cell QC metrics, high-quality cells can be retained. In our setting, outliers are defined as cells with library sizes more than two median absolute deviations away from the median library size. Figure 3.1 summarizes the process from unprocessed to processed single-cell experiment. After filtering, the data set consists of 23'543 cells and 18'537 genes. Next, we used the singleR function from the singleR R package (Aran et al., 2019) for cell-type annotation. Cell-type annotation is important to determine what biological state is represented by cell clusters which helps the interpretability of the results and their implications (Amezquita et al., 2020). singleR is a method that assigns labels based on the reference samples with the highest Spearman rank correlation while only using marker genes between pairs of labels to focus on the relevant differences between cell types (Aran et al., 2019).

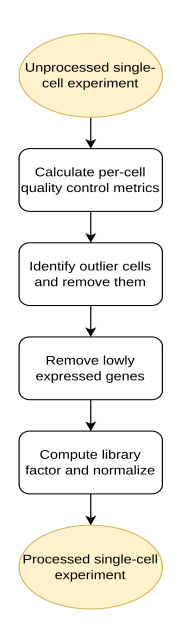
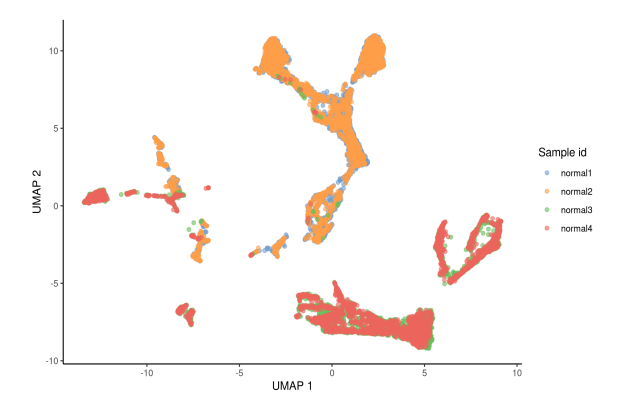


Figure 3.1: Quality control process from unprocessed, raw to processed, filtered single-cell experiment

Figure 3.2 shows that the cells largely overlap based on the UMAP representation



**Figure 3.2:** UMAP representation of the mouse kidney cells based on the sample coloured by sample id

Figure 3.3 shows

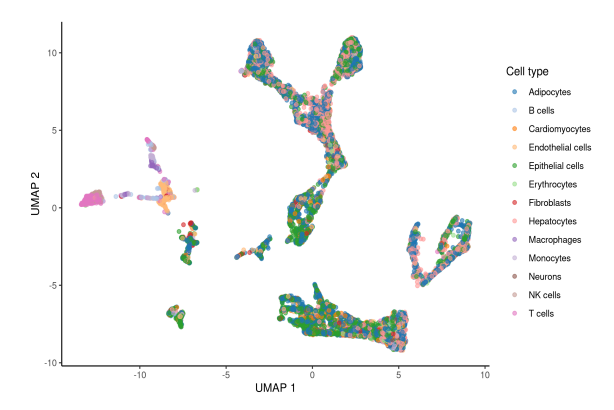


Figure 3.3: UMAP representation of the counts based on the cell type

#### 3.1.2 Human brain organoids

The second data set we investigated (Velasco *et al.*, 2019)

### 3.2 Simulation study on the mouse data

#### 3.2.1 Simulation strategy

Our initial simulation strategy was to randomly invert the spliced and unspliced counts for 10% of the genes with and without differential gene expression.

#### 3.2.2 Simulation analyses

### 3.3 Discovery analysis on the human brain organoids

### 3.4 Data availability

#### Kidney mouse cells

The raw data can be downloaded from NCBI GEO (accession number GSE107585). https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE107585

#### Human brain organoids

The raw data can be downloaded from NCBI GEO (accession number GSE129519). https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE129519

### 3.5 Code availability

All code for data cleaning and analysis associated with the current submission is available at <a href="https://github.com/joelmeili/DifferentialRegulation">https://github.com/joelmeili/DifferentialRegulation</a>. Any updates will also be published on GitHub.

## Chapter 4

### Discussion

### 4.1 Conclusion

### 4.2 Outlook

Bioconductor package DifferentialRegulation is a method for detecting differentially regulated genes between two groups of samples (e.g., healthy vs. disease, or treated vs. untreated samples), by targeting differences in the balance of spliced and unspliced mRNA abundances, obtained from single-cell RNA-sequencing (scRNA-seq) data. DifferentialRegulation accounts for the sample-to-sample variability, and embeds multiple samples in a Bayesian hierarchical model. In particular, when reads are compatible with multiple genes or multiple splicing versions of a gene (unspliced spliced or ambiguous), the method allocates these multi-mapping reads to the gene of origin and their splicing version. Parameters are inferred via Markov chain Monte Carlo (MCMC) techniques (Metropolis-within-Gibbs) (Tiberi, 2022).

# Appendix A

# Figures

Appendix B

Source code

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