NON-PARAMETRIC MODELLING OF TEMPORAL AND SPATIAL COUNTS DATA FROM RNA-SEQ EXPERIMENTS

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July 29, 2020

ABSTRACT

**Motivation:** The negative binomial distribution is a good model for counts data from both bulk and single-cell RNA-sequencing (RNA-seq). Gaussian process (GP) regression provides a useful non-parametric approach for modeling temporal or spatial changes in gene expression. However, currently available GP regression methods that implement negative binomial likelihood models do not scale to the increasingly large datasets being produced by single-cell and spatial transcriptomics.

**Results:** The GPcounts package implements GP regression methods for modelling counts data using negative binomial likelihood functions. Computational efficiency is achieved through the use of variational Bayesian inference. The GP function models changes in the mean of the negative binomial likelihood through a logarithmic link function and the dispersion parameter is fitted by maximum likelihood. We also provide the option of modelling additional dropout using a zero-inflated negative binomial likelihood. We validate the method on simulated time course data, showing that it is better able to identify changes in over-dispersed counts data than methods based on Gaussian or Poisson likelihoods. To demonstrate temporal inference, we apply GPcounts to single-cell RNA-seq datasets after pseudotime and branching inference. To demonstrate spatial inference, we apply GPcounts to data from the mouse olfactory bulb to identify spatially variable genes and compare to a published GP method with a Gaussian likelihood function. Our results show that GPcounts can be used to model temporal and spatial counts data in cases where simpler Gaussian and Poisson likelihoods are unrealistic.

**Availability:** GPcounts is implemented using the GPflow library in Python and is available at https://github.com/ManchesterBioinference/GPcounts along with the data, code and notebooks required to reproduce the results presented here.
1 Introduction

Biological data are often summarised as counts, e.g. high-throughput sequencing allows us to count the number of sequence reads aligning to a genomic region of interest, while single molecule fluorescence in-situ hybridization (smFISH) allows us to count the number of RNA molecules in a cell or within a cellular compartment. For datasets that are collected with temporal or spatial resolution, it can be useful to model changes in time or space using a Gaussian process (GP). GPs provide a flexible framework for non-parametric Bayesian modelling, allowing for non-linear regression while quantifying the uncertainty associated with the estimated latent function and data measurement process (Rasmussen and Williams, 2006). GPs are useful methods for analysing gene expression time series data, e.g. to identify differentially expressed genes, model temporal changes across conditions, cluster genes or model branching dynamics (Stegle et al., 2010; Kalaitzis and Lawrence, 2011; Aijö et al., 2014; Yang et al., 2016; Ahmed et al., 2019; Hensman et al., 2013; McDowell et al., 2018; Boukouvalas et al., 2018). GPs have also been applied to spatial gene expression data as a method to discover spatially varying genes (Svensson et al., 2018; Sun et al., 2020). In many previous applications of GP regression to counts, the log-transformed counts data are modelled using a Gaussian noise assumption. Such Gaussian likelihood models do not properly capture the typical characteristics of counts data, in particular substantial probability mass at zero counts and heteroscedastic noise. Alternative data transformations attempt to match the mean/variance relationship of counts data (Anscombe, 1948) but cannot model all relevant aspects of the distribution, e.g. the probability mass at zero. More suitable distributions for modelling counts data include the negative binomial distribution, which can model over-dispersion beyond a Poisson model, and zero-inflated extensions that have been developed for single-cell data exhibiting an excess of zero counts (Pierson and Yau, 2015; Risso et al., 2017). A negative binomial likelihood was implemented for GP regression using Markov chain Monte Carlo (MCMC) inference (Aijö et al., 2014). However, this approach does not scale to large datasets, e.g. single-cell RNA-seq (scRNA-seq) data after pseudotime inference or from spatial transcriptomics assays.

Various statistical methods have been proposed for spatially resolved omics data (Svensson et al., 2018; Sun et al., 2020; Arnol et al., 2019; Edsgärd et al., 2018). The SpatialDE approach uses a GP to model the expression variability of each gene with a spatial and non-spatial component (Svensson et al., 2018). The latter is modelled as observation noise, while the former is modelled using a covariance function that depends on the pairwise distance between the cells. The ratio of these components is used to measure the spatial variability of each gene. A Gaussian likelihood is used for inference and the raw counts data is transformed using Anscombe’s transformation (Anscombe, 1948) to reduce heteroscedasticity. Statistical significance is assessed by a likelihood ratio test against a spatially homogeneous model, with p-values calculated under the assumption of a $\chi^2$-distribution. Trendseek (Edsgärd et al., 2018) assesses spatial expression of each gene by modelling normalised counts data as a marked point process, where the points represent the spatial locations and marks represent the expression levels, with a permuted null used to assess significance. SPARK (Sun et al., 2020) is a recently introduced GP-based method that relies on a variety of spatial kernels and a penalized quasi-likelihood algorithm. SPARK models counts using a Poisson likelihood function and captures over-dispersion through a nugget (white noise) term in the underlying GP covariance function.

Here, we introduce an alternative GP regression model (GPcounts) to model temporal or spatial counts data with negative binomial (NB) or zero-inflated negative binomial (ZINB) likelihoods. GPcounts can be used for a variety of tasks, e.g. to infer temporal trajectories, identify differentially expressed genes using one-sample or two-sample tests, infer branching genes from scRNA-seq after pseudotime inference, or to infer spatially varying genes. We use a GP with a logarithmic link function to model variation in the mean of the counts data distribution across time or space. As an example, in Figure 1 we show how GPcounts captures the distribution of a short RNA-seq time course dataset from Leong et al. (2014). In this example we use a two-sample test to determine whether time-series measured under different conditions have differing trajectories. The GP with Gaussian likelihood fails to model the data distribution well, leading to a less plausible inferred dynamics and overly broad credible regions. Although the trajectories of the two samples clearly differ, the Gaussian likelihood model provides a poor fit and the single trajectory model is preferred in the two-sample test.

Our package is developed using the GPyflow library (De G. Matthews et al., 2017) which we have extended to include the DEtime kernel (Yang et al., 2016) for branching genes and to include NB and ZINB likelihoods with an efficient variational approximate inference method.

2 Methods

2.1 Gaussian Process Regression

A Gaussian Process (GP) is a stochastic process over real valued functions and defines a probability distribution over function space (Rasmussen and Williams, 2006). GPs provide a non-linear and non-parametric framework for inference.
Figure 1: Example of data from a two-sample time course RNA-seq experiment (Leong et al., 2014). With a Negative Binomial likelihood we are able to identify this differentially expressed gene based on a likelihood ratio statistic. With a Gaussian noise model we obtain a poor fit and the likelihood ratio does not identify the gene as differentially expressed.

Consider a set of $N$ temporal or spatial locations $\mathbf{x} = [x_1, x_2, ..., x_N]$ associated with observations $\mathbf{y} = [y_1, y_2, ..., y_N]$. As this is a continuous model, spacings between data points can vary. Each observation $y_n$ is modelled as a noisy observation of the function evaluated at $x_n$, $f_n = f(x_n)$, through some likelihood function $p(y_n | f_n)$. The function $f$ is a latent function sampled from a GP prior and $p(y_n | f_n)$ models the data distribution at $x = x_n$. The simplest and most popular choice of likelihood function is i.i.d. Gaussian noise centred at $f$, in which case $y_n | f_n \sim N(f, \sigma^2 I)$, where $f = [f_1, f_2, ..., f_N]$ is the GP function evaluated at the times or locations in $\mathbf{x}$. In this work we use likelihood functions that are more suitable for counts data, which we introduce below.

To indicate that $f$ is drawn from a GP we write,

$$f \sim \mathcal{GP}(\mu, k).$$

Here, $\mu(x) = \mathbb{E}[f(x)]$ is the mean function of the GP. The covariance function $k$ (also known as the kernel function) is a positive semidefinite function $k(x, \tilde{x}) = \mathbb{E}[f(x)f(\tilde{x})] = \mathbb{E}[f(x)]\mathbb{E}[f(\tilde{x})] - \mathbb{E}[f(x)f(\tilde{x})]$ that determines the covariance of $f$ at any two locations $x$ and $\tilde{x}$. The covariance function describes our beliefs when modelling the function, e.g. whether it is stationary or non-stationary, rough or smooth etc. A popular choice is the Radial Basis Function (RBF) kernel which leads to smooth and infinitely differentiable functions, defined as:

$$k(x, \tilde{x}) = \sigma^2_l \exp\left(-\frac{||x - \tilde{x}||^2}{2l^2}\right)$$

where the hyper-parameters are the lengthscale $l$, controlling spatial or temporal variability, and the amplitude $\sigma^2_l$, controlling the marginal variance of the function. The RBF kernel is also known as the squared exponential, exponentiated quadratic or Gaussian kernel (Rasmussen and Williams, 2006). Below we describe an alternative kernel appropriate for branching functions.

The kernel hyper-parameters and parameters of the likelihood function can be learned from the data by optimising the log marginal likelihood function of the GP. The marginal likelihood is given by the probability of the data $\mathbf{y}$ after integrating out the GP function,

$$p(\mathbf{y}) = \int p(\mathbf{y}|\mathbf{f}) \mathcal{N}(\mathbf{f}|0, K(\mathbf{x}, \mathbf{x})) d\mathbf{f},$$

where $K(\mathbf{x}, \mathbf{x})$ is the covariance matrix of the GP evaluated at the times or locations in $\mathbf{x}$.
where we have set the GP prior mean to be zero and $K(x, x)$ is the covariance matrix of the GP function evaluated at the data locations $x$. For the special case of a Gaussian likelihood function this integral is tractable but for the non-Gaussian likelihoods below we use a variational approximation introduced by [Opper and Archambeau (2009)].

### 2.2 Negative binomial likelihood

The Negative Binomial (NB) distribution is a popular model for bulk RNA-seq counts data ([Robinson and Smyth, 2007](#fn1){ref#}) and has been shown to be suitable for modelling scRNA-seq count data with UMI (unique molecular identifier) normalisation ([Svensson, 2020](#fn2){ref#} [Townes et al., 2019](#fn3){ref#}). It can be parametrised by a fixed number of failures $r$ and mean $\mu$:

$$
\text{NB}(y; \mu, r) = \frac{\Gamma(y + r)}{\Gamma(y + 1)\Gamma(r)} \left( \frac{r}{r + \mu} \right)^r \left( \frac{\mu}{r + \mu} \right)^y, \quad \forall y \in \mathbb{N}
$$

(4)

where $\Gamma$ is the Gamma function. It is convenient to parametrise the NB distribution by a dispersion parameter $\alpha = r^{-1}$ which captures excess variance relative to a Poisson distribution, since $\text{Var}[y] = \mu + \alpha\mu^2$. A logarithmic link function is used to model the mean of the NB as a transformation of the GP function $f(x) = \log \mu(x)$.

### 2.3 Zero-inflated negative binomial likelihood

In some cases the model may benefit from an excess of zeros beyond the NB distribution assumption. For example, the NB distribution may not adequately model non-UMI scRNA-seq counts data since an excess of zeros may lead to an overestimate of the dispersion parameter and a biased mean estimate. Even in the case of UMI scRNA-seq data, the model may benefit from some additional flexibility when fitting data to an inferred pseudotime trajectory.

A zero-inflated distribution can be used to model count data with excessive zeros ([Pierson and Yau, 2015](#fn4){ref#} [Risso et al., 2017](#fn5){ref#}). These extra zeros are assumed to be generated by a separate process different from the negative binomial process that generates counts including zeros. The ZINB distribution is parametrised by $\Psi$ (the excess proportion of zeros) and the NB model parameters:

$$
\text{ZINB}(y; \mu, \alpha, \Psi) = \begin{cases} 
\Psi \cdot \text{NB}(y = 0) & \text{for } y = 0, \\
(1 - \Psi) \cdot \text{NB}(y) & \text{for } y \geq 1.
\end{cases}
$$

(5)

We parametrise the probability of dropout $\Psi$ using a Michaelis–Menten (MM) equation relating the probability of dropout to the mean of the NB distribution used to model the non-dropout data $\mu = e^{f(x)}$ (using the same link function as above for the NB likelihood),

$$
\Psi = 1 - \frac{e^{f(x)}}{K_M + e^{f(x)}}
$$

(6)

where the Michaelis constant $K_M$ is estimated by our model. The MM model was previously found to provide a similar fit to a logistic function and better fit than using a double exponential function ([Andrews and Hemberg, 2016](#fn6){ref#}).

### 2.4 Tests and credible regions

We provide one-sample and two-sample likelihood-ratio statistics in GPcounts to identify differentially expressed genes. In the one-sample case the null hypothesis assumes there is no difference in gene expression across time or space ([Kalaitzis and Lawrence, 2011](#fn7){ref#} [Svensson et al., 2018](#fn8){ref#}) and we compute the ratio of the GP model marginal likelihood versus a constant model likelihood. The models are nested since the constant model is equivalent to the GP with an infinite lengthscale parameter. In the two-sample case the null hypothesis assumes there is no difference in gene expression under two different conditions and the alternative hypothesis is that two different GP functions are required to model each sample ([Stegle et al., 2010](#fn9){ref#}). Rejecting the null hypothesis in each case indicates that a gene is differentially expressed. In the two-sample case, we fit three GPs: one for each dataset separately and a shared GP where the datasets are treated as replicates.

For spatial transcriptomics data we follow the approach of [Svensson et al., 2018](#fn10){ref#}, with $p$-values estimated according to a $\chi^2$-distribution and 5% FDR threshold estimated using the approach of [Storey and Tibshirani, 2003](#fn11){ref#}.

To plot [5%-95%] credible regions we draw 100 random samples from the GP at 100 equally spaced times. We exponentiate each GP sample to set the mean of the count distribution (NB, ZINB and Poisson) and draw counts at each time. We use the mean and percentiles to plot the predictive distribution with the associated credible regions.
2.5 Scale normalisation for spatial gene expression data

In some cases there may be confounding variation that will dominate the temporal or spatial trends in the data. For example, Svensson et al. (2018) point out that there may be spatial patterns in cell size that can lead to almost all genes being identified as spatially variable. In this case it is necessary to normalise away such confounding variation in order to model other sources of spatial variation. Svensson et al. (2018) used ordinary least-squares regression of Anscombe-normalised spatial expression data against logged total counts to remove this confounding variation. We use NB regression with an identity link function to learn location-specific normalisation factors for spatial counts data. We subsequently model the data using a modified GP likelihood evaluated at each grid point, \( y_i \sim \text{NB}(k_i \mu_i, r) \) for \( i = 1, \ldots, N \) with \( \mu_i = e^{f(x_i)} \) and \( f \sim \mathcal{GP}(0, k) \). The multiplicative normalisation factor \( k_i \) is calculated as \( k_i = \beta T_i \) for \( i = 1, \ldots, N \), where \( \beta \) is the slope calculated by fitting for each gene a NB regression model with intercept zero and \( T_i \) corresponds to the total counts at the \( i \)th spatial location.

2.6 Modelling branching dynamics

In the two-sample time course setting it can also be useful to identify the time at which individual genes begin to follow different trajectories. This can be useful in bulk time course data in a two-sample testing scenario (Yang et al. 2016) or for modelling branching dynamics in single-cell data after pseudotime inference (Boukouvalas et al. 2018). We first define a joint covariance function for two GPs with the same covariance function \( f \sim \mathcal{GP}(0, k) \) and \( g \sim \mathcal{GP}(0, k) \) constrained to cross at a branching point \( x_b \):

\[
\begin{pmatrix}
k_{ff} & k_{fg} \\
k_{gf} & k_{gg}
\end{pmatrix} = 
\begin{pmatrix}
\frac{k(x, x)}{k(x, x) + k(x, x)} & \frac{k(x, x)}{k(x, x) + k(x, x)} \\
\frac{k(x, x)}{k(x, x) + k(x, x)} & \frac{k(x, x)}{k(x, x) + k(x, x)}
\end{pmatrix}. 
\]

(7)

Now consider data from two lineages \( y^i \) and \( y^b \) representing noise-corrupted measurements of a baseline (trunk) and diverging (branch) time course respectively. Before the divergence point \( x_b \) the data are distributed around the trunk function \( f(x) \) according to some likelihood function \( p(y|f) \),

\[
y_i(x_n) \sim p(y|f(x_n)) \quad \text{for } x_n \leq x_b. 
\]

(8)

After the branching point \( x_b \), the mean function of the trunk continues to follow \( f \) while the mean function of the diverging branch trajectory changes to follow \( g \),

\[
y_i(x_n) \sim p(y|f(x_n)), \\
y^b(x_n) \sim p(y|g(x_n)) \quad \text{for } x_n \geq x_b. 
\]

(9)

The branching point \( x_b \) is a hyper-parameter of the joint covariance function of this model along with the hyper-parameters of the GP functions (lengthscale \( l \) and amplitude \( \sigma_f^2 \)). The lengthscale and amplitude are shared by both functions and are fitted (along with the likelihood model parameters) by fitting two separate GP regression models to the data from both conditions or lineages and estimated using maximum likelihood (Yang et al. 2016). This leaves the problem of inferring the branching time \( x_b \) only. As this is a one-dimensional problem, the posterior distribution of \( x_b \) is estimated using a simple histogram approach. We have used a simple discretisation \( x_b \in [x_{\text{min}}, x_{\text{min}} + \delta, x_{\text{min}} + 2\delta, \ldots, x_{\text{max}}] \) as in Yang et al. (2016) and estimate the posterior by using the normalised likelihood evaluated at each grid point,

\[
p(x_b|y^i, y^b) \propto \frac{p(y^i, y^b|x_b)}{\sum_{x=x_{\text{min}}}^{x_{\text{max}}} p(y^i, y^b|x)},
\]

(10)

which avoids the need for complex optimisation or integration schemes.

2.7 Efficient approximate inference implementation

The integral in Eqn. (3) is intractable for non-Gaussian likelihood functions. Therefore, we need to approximate the posterior function using variational inference (Opper and Archambeau 2009, Bauer et al. 2016). Full variational Bayesian inference is computationally intensive and requires \( O(N^2) \) memory and \( O(N^3) \) computation time so we use a sparse approximation to reduce the computational requirements. In sparse inference, we choose \( M < N \) inducing points \( z \) defined in the same space of regressors \( x \). Using inducing points reduces the time complexity to \( O(NM^2) \) and reduces the size of the covariance matrix from \( N \times N \) to \( M \times M \). In GPcounts the default is to set the number of inducing points to \( M = 5\% (N) \) and we apply a k-means clustering algorithm to select the candidate inducing points \( z \). For very large datasets a smaller number of inducing points may be required.
2.8 Practical considerations

Practical limitations of the inference framework include local optima in the hyperparameter search and occasional numerical instabilities. We have therefore incorporated some checks to detect numerical errors such as failure of Cholesky decomposition or failure of optimization. Local optima are a likely issue if the GP posterior predictive is consistently higher or lower than the observations. Where we suspect local optima or numerical errors we restart the optimisation from new random values for the hyper-parameters.

3 Results and discussion

Below we apply GPcounts on simulated counts time course data, real scRNA-seq after pseudotime inference and spatial transcriptomics data.

3.1 Assessment on synthetic time course data

We simulated four counts time course datasets with two levels of mean expression (high/low) and two levels of dispersion (high/low) to assess the performance of GPcounts in identifying differentially expressed genes using a one-sample test. Each dataset has 600 genes with time-series measurements at 11 equally spaced time points \( x = [0, 0.1, 0.2, \ldots, 1.0] \) and two replicates at each time. Half of the genes are differentially expressed across time. We use two classes of generative functions \( f(x) \), sine functions and cubic splines, to simulate data from time-varying genes. The sine functions are of the form \( f(x) = a \sin(xb + d) + e \) where parameters are drawn from uniform distributions. Specifically, \( b \sim U[\pi/4, 2\pi], d \sim U[0, 2\pi] \) while the \( a \) and \( e \) distributions are chosen to alter the signal amplitude and mean ranges respectively. The cubic spline function has the form \( f(x) \in C^2[0, 1] \) passing through two control points \((x, y)\) where \( x \) and \( y \) are drawn from uniform distributions. For non-differentially expressed genes we choose constant values set to the median value of each dynamic function in the dataset. The low and high dispersion values are drawn from uniform distributions \( \alpha_{\text{low}} \sim U[0.01, 0.1] \) and \( \alpha_{\text{high}} \sim U[1, 3] \) respectively. An exponential inverse-link function is used to determine the mean of count data at each time \( \mu(x) = e^{f(x)} \) and we use the SciPy library (Millman and Aivazis, 2011) to sample counts from the negative binomial distribution. Specific simulation parameters and final datasets are provided with the supporting code.

In Figure 2 we compare the performance of GPcounts using the NB likelihood with a Poisson likelihood and the Gaussian GP model. We ran DESeq2 with time as a covariate and used the adjusted \( p \)-values as a score to label differentially expressed genes. We then compared the performance of one-sample tests using the NB likelihood and Gaussian likelihood. We consider different adjusted \( p \)-value thresholds to identify DE genes according to DESeq2 and look at the concordance with using a GP with NB and Gaussian likelihoods. Figure 4 shows precision–recall curves to improve the performance of the Gaussian likelihood model and typically makes it worse.

3.2 Modelling scRNA-seq pseudotime-series

We apply GPcounts on mouse pancreatic \( \alpha \) cell data from scRNA-seq experiments without UMI normalisation (Qiu et al., 2017). We use the pseudotime inference results from Qiu et al. (2017) based on PCA. Figure 3 shows the inferred trajectory for two genes with many zero count measurements: Fam184b with 86% zero counts and Pde1a with 68% zero counts. From left to right we show the GP regression fit with Gaussian, NB and ZINB likelihoods respectively. For Fam184b we see that the Gaussian and NB models do not capture most of the non-zero data within their credible regions. The ZINB model has 21/324 (6.4%) cells above the 95% upper credible region while the NB model has 68/324 (21.1%) suggesting the ZINB model is a little better calibrated. For both genes the Gaussian model is unable to effectively model the high probability region at zero counts, due to the symmetric nature of the distribution. The fits for the NB and ZINB likelihood are very similar for Pde1a with only a subtle difference in the credible regions (4% of cells above 95%) and we find that ZINB and NB give similar fits for most genes. Qiu et al. (2017) identify 611 differentially expressed genes in mouse pancreatic \( \alpha \) cells using DESeq2 (Love et al., 2014) which they applied to two distinctive clusters of cells along the above pseudotime dimension. Since DESeq2 also assumes an NB model we examined whether the results from GPcounts would be closer to DESeq2 than to a Gaussian GP model. We ran DESeq2 with time as a covariate and used the adjusted \( p \)-values as a score to label differentially expressed genes. We then compared the performance of one-sample tests using the NB likelihood and Gaussian likelihood. We consider different adjusted \( p \)-value thresholds to identify DE genes according to DESeq2 and look at the concordance with using a GP with NB and Gaussian likelihoods. Figure 4 shows precision–recall curves to...
Figure 2: ROC curves for a one-sample test using a Gaussian process to identify dynamic time-series from synthetic counts data. Negative binomial, Gaussian and Poisson likelihoods are used to identify differentially expressed genes with a one-sample test. Genes are ranked by likelihood ratio between a dynamic and constant model. Results are shown for (a) low counts/low dispersion (b) low counts/high dispersion (c) high counts/low dispersion and (d) high counts/high dispersion datasets.

Figure 3: GP models of gene expression against pseudotime (Qiu et al., 2017). We show two genes with a large number of zero-counts: Fam184b with 86% zeros in the first row and Pde1a with 68% zeros in the second row. We show the inferred mean trajectory and credible regions using (a) Gaussian likelihood, (b) NB likelihood and (c) ZINB likelihood.
explore how similarly GPcounts performs compared to DESeq2. We see that with the NB likelihood the genes obtained are very similar to those found by DESeq2. With a Gaussian likelihood the GP identifies very similar genes among its top-ranked DE genes but has much less concordance further down the list. This result is also reflected in the higher rank correlation between the DESeq and NB likelihood GP results than with the Gaussian likelihood GP (Figure 5). This suggests that the test statistic is more influenced by the form of likelihood term rather than the form of regression model in this example.

For scRNA-seq data the number of cells can be very large and computational efficiency becomes important. We have checked that the above results are very similar when using the sparse inference approximation (see notebook accompanying the package) with computation time reduced by 60% with the NB and ZINB likelihoods.

### 3.3 Identification of spatially variable genes

We applied GPcounts to spatial transcriptomics sequencing counts data from mouse olfactory bulb (Ståhl et al., 2016) consisting of measurements of 16,218 genes at 262 spatial locations. We compare performance to SpatialDE which uses a GP with a Gaussian likelihood (Svensson et al., 2018) and we use as similar a set-up as possible in order to make the comparison fair. We filter out genes with less than 3 total counts and spatial locations with less than 10 total counts and ended up with 14,859 genes across 260 spatial locations. To assess statistical significance we used the $q$-value method (Storey and Tibshirani, 2003) to determine an FDR cut-off (0.05) based on $p$-values assuming a $\chi^2$-distribution of log likelihood ratios. We compare our findings in figure 6(a), where GPcounts identified 468 spatially variable (SV) genes in total, whilst SpatialDE identified 67, with 62 of them overlapping between the two methods. The SPARK and Trendsceek methods, which consider calibrated $p$-values under a permuted null, identified 772 and 0 SV genes respectively (Sun et al., 2020). In figure 6(b) we plot the GPcounts log likelihood ratio versus SpatialDE log likelihood.
Figure 6: (a) Venn diagram shows the overlap between GPcounts and SpatialDE spatially variable (SV) genes with the green area showing the 5 SV genes that are only identified by SpatialDE. (b) GPcounts Log likelihood ratio versus SpatialDE Log Likelihood ratio. The dashed horizontal and vertical lines correspond to 5% FDR threshold. (c) Relative spatially resolved expression profiles for four selected SV genes, with their $q$-value in the parenthesis. (d) In-situ hybridization images of the selected SV genes in (c). The images are taken from Ståhl et al. (2016).

3.4 Identifying gene-specific branching locations

We used scRNA-seq of haematopoietic stem cells (HSCs) from mouse (Paul et al., 2015) to demonstrate the method’s effectiveness in identifying branching locations for individual genes. The data contain cells that are differentiated into myeloid and erythroid precursor cell types. Paul et al. (2015) analysed changes in gene expression for myeloid progenitors and created a reference compendium of marker genes related to the development of erythrocytes and several other types of leukocytes from myeloid progenitors. The Slingshot algorithm is used to get trajectory-specific pseudotimes as well as assignment of cells to different branches. After removing two small outlier cell clusters related to the dendritic and eosinophyl cell types, Slingshot infers two lineages for this dataset. These two outlier cell clusters do not belong to any particular lineage and have previously been excluded from trajectory inference and downstream analysis (Van den Berge et al., 2019), which leaves us 2660 cells under consideration.

Figure 7 shows examples of GPcounts model fits with associated credible regions (upper sub-panels) as well as the posterior probability distribution over branching time (bottom sub-panels) for an early branching known bio-marker MPO (upper row) and for a late branching gene LY6E (bottom row). Here, we have sub-sampled the data and larger markers in Figure 7 represent the cells used in the inference process. Figure 7 (a) and (c) show the results with a
Gaussian likelihood, equivalent to the model in Yang et al. (2016), while Figure 7 (b) and (d) show the results with the NB likelihood. In all cases, the bottom sub-panels reflect the significant amount of uncertainty associated with the identification of precise branching points. Both models provide a reasonably similar posterior probability of the branching time. However, looking at the credible region of the data we find that the model with NB likelihood better models the data. In the case of the Gaussian likelihood the credible regions are wide but they still miss some points that have zero values. In the case of NB likelihood the credible regions can adequately model the points having zero values. Further example fits are shown in the accompanying notebook.

![Figure 7](image_url)

Figure 7: Mouse haematopoietic stem cells (Paul et al., 2015): Identifying the gene-specific branching location for an early branching gene MPO (a and b) and a later branching gene LY6E (c and d). Cell assignments and pseudotime are estimated using the Slingshot algorithm. The upper sub-panels shows the GP regression fit based on the MAP estimate of the gene-specific differentiating time and the bottom sub-panels shows the posterior distribution over the differentiating or branching times. Bigger markers used in the upper sub-panels indicate the sub-sampled cells used in the inference. Results on the left are for the Gaussian likelihood fit (a,c) and on the right for the NB likelihood fit (b,d).

4 Conclusion

We have developed a Gaussian process (GP) regression method, GPcounts, implementing negative binomial (NB) and zero-inflated negative binomial (ZINB) likelihoods. This provides a useful tool for bulk and single-cell RNA-seq data in time-series and spatial transcriptomics experiments. Although the ZINB likelihood may be more suitable in some cases, especially for non-UMI normalised scRNA-seq data, in practice the differences are relatively minor even when the number of zeros is very high for a gene. For most practical purposes we have therefore applied the simpler NB likelihood. This is in line with a recent study showing that although zero-inflation certainly exists in scRNA-seq data, there may be little benefit in modelling it since the additional zero-inflation parameter can be difficult to identify (Choi et al., 2020).
Our results show that the NB likelihood can provide a substantial improvement over a Gaussian likelihood when modelling counts data. In the data from [Qui et al., 2017] we found that the NB GP identifies DE gene lists more similar to DESeq2 than to a Gaussian GP. This suggests that the likelihood function plays a more important role than the regression model in some problems. Similarly, we found a major difference in spatial inference using the NB likelihood compared to the SpatialDE method that is based on a Gaussian likelihood GP. Using a similar normalisation and testing set-up, we found a much larger set of spatially variable (SV) genes than SpatialDE. However, we found less genes than the over-dispersed Poisson method, SPARK, which also uses GP inference but with differences in both the modelling and testing set-up. In particular, SPARK uses a permutation approach to compute p-values and therefore the number of SV genes is not directly comparable with our approach. It will be useful to consider alternative testing set-ups and a broader range of spatial inference problems in order to compare these approaches more carefully. Finally, we applied a branching kernel on a Mouse bone marrow scRNA-seq dataset [Paul et al., 2015] to infer the initial point when the two gene expression time profiles begin to diverge. For genes with strong branching evidence the NB and Gaussian likelihood provided similar inference results.

To improve the practical performance of GPcounts we implement a heuristic to detect locally optimal solutions and to detect numerical instability. Since the naive GP scales cubically with number of time points we improve the computational requirements through a sparse inference algorithm from the GPflow library (De G. Matthews et al., 2017). Our implementation of GPcounts is flexible and can easily be extended to work with any kernel or likelihood compatible with the GPflow library. A natural next step would be to better parallelise model fitting for each gene. Improving the null models to produce better calibrated p-values could also improve FDR calibration in DE and SE calling.

Acknowledgements

Funding

NB was supported by King Saud University funded by Saudi Government Scholarship (KSU1546). MR is supported by a Wellcome Trust Investigator Award (204832/B/16/Z). MR and AB were supported by the MRC (MR/M008908/1).

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