# Gene expression GPseudoClust: deconvolution of shared pseudo-profiles at single-cell resolution

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Received on March 8, 2019; revised on September 20, 2019; editorial decision on October 4, 2019; accepted on October 9, 2019

# Abstract

**Motivation**: Many methods have been developed to cluster genes on the basis of their changes in mRNA expression over time, using bulk RNA-seq or microarray data. However, single-cell data may present a particular challenge for these algorithms, since the temporal ordering of cells is not directly observed. One way to address this is to first use pseudotime methods to order the cells, and then apply clustering techniques for time course data. However, pseudotime estimates are subject to high levels of uncertainty, and failing to account for this uncertainty is liable to lead to erroneous and/or over-confident gene clusters.

**Results:** The proposed method, GPseudoClust, is a novel approach that jointly infers pseudotemporal ordering and gene clusters, and quantifies the uncertainty in both. GPseudoClust combines a recent method for pseudotime inference with non-parametric Bayesian clustering methods, efficient Markov Chain Monte Carlo sampling and novel subsampling strategies which aid computation. We consider a broad array of simulated and experimental datasets to demonstrate the effectiveness of GPseudoClust in a range of settings.

Availability and implementation: An implementation is available on GitHub: https://github.com/magStra/ nonparametricSummaryPSM and https://github.com/magStra/GPseudoClust.

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Supplementary information: Supplementary data are available at *Bioinformatics* online.

# **1** Introduction

During response to stimulation or development, gene expression undergoes significant changes for many genes. For bulk measurements of gene expression these changes can be investigated by collecting time course data. A common analysis step for such datasets is to cluster genes on the basis of the similarities in their time course profiles. For example, Eisen et al. (1998) found that similar expression dynamics of genes are related to biological function, whereas Cooke et al. (2011) showed that clustering genes together with similar changes in expression over time can identify those likely to be coregulated by the same transcription factors. McDowell et al. (2018) emphasize that using clustering to identify shared response types helps reduce the complexity of the response, and allows the exploration of regulatory mechanisms underlying the shared response types. Most existing methods for performing such clustering analyses were developed for bulk-measurements of gene expression, and not for single-cell data.

There is clearly a need for effective clustering algorithms for genes for single-cell data, given that single-cell technologies have enabled us to obtain response and developmental trajectories with a much better resolution; see, e.g. Griffiths et al. (2018), Kunz et al. (2018) and Nestorowa et al. (2016). Single-cell RNA-seq data have been used to investigate processes of development, differentiation or immune response, with the development of pseudotemporal ordering approaches enabling researchers to order cells in terms of their progression through these processes; see Ahmed et al. (2019), Campbell and Yau (2016), Haghverdi et al. (2016), Ji and Ji (2016), Qiu et al. (2017), Reid and Wernisch (2016), Strauß et al. (2019) and Welch et al. (2016) among many others. For each gene, the ordered gene expression measurements are assumed to be noisy observations of an underlying latent trajectory characterizing the response to a stimulant or the dynamics of its expression during development. In addition to the challenge of pseudotime inference, single-cell data are also characterized by higher levels of noise, including dropout effects (see, among others, Stegle et al., 2015; Vallejos et al., 2015). Moreover, the

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number of cells in single-cell datasets typically exceeds by orders of magnitude that of time points for bulk measurements.

A number of algorithms have been developed specifically for clustering *cells* using scRNA-seq data; for instance Kiselev *et al.* (2017), Lin *et al.* (2017) and Wang *et al.* (2017), the latter method using multiple kernel learning. However, there has been far less progress on the development of methods for clustering *genes* using scRNA-seq data.

One way of clustering pseudotemporal single-cell gene expression pseudotime profiles is to adopt a two-step approach (Macaulay *et al.*, 2016): first use a pseudotime ordering method such as SLICER (Welch *et al.*, 2016) or DeLorean (Reid and Wernisch, 2016); then cluster genes using a method for time-stamped bulk data, e.g. GPclust (Hensman *et al.*, 2013, 2015). A two-step approach is also implemented in Monocle 2 (Qiu *et al.*, 2017), which uses partitioning around medoids (PAMs, Kaufman and Rousseeuw, 2008) on a distance measure between smoothed pseudotime expression profiles. However, such two-step approaches do not take into account the potential impact of the uncertainty in the inferred pseudotimes upon the identification of clusters.

The method proposed here, *GPseudoClust*, addresses this challenge by probabilistically modelling the orders of cells and cluster allocations of genes jointly, thereby accounting for dependencies between the orders of the cells and cluster allocations of the genes. GPseudoClust combines our previously developed method for modelling the uncertainty of pseudotime (Strauß *et al.*, 2019), with Bayesian clustering using Dirichlet process (DP) mixtures of hierarchical Gaussian processes (GPs) (Hensman *et al.*, 2013, 2015).

#### 2 Materials and methods

### 2.1 Cell orderings and pseudotime

We assume our data comprises pre-processed log-transformed gene expression data in the form  $\mathbf{y}_i$  of gene  $j = 1, ..., n_g$ , where  $\mathbf{y}_i$  is a vector of length *T*, the number of cells. We start with a vector of pseudo-time points  $\mathbf{\tau} = (\tau_1, ..., \tau_T)$  and seek to infer an ordering of cells as a permutation  $\mathbf{o} = (o_1, ..., o_T), o_i \in \{1, ..., T\}, o_i \neq o_i$  for  $i \neq j$ , where  $o_i$  is the index of the cell assigned to pseudotime  $\tau_i$  in the ordering. We refer the reader to our previous paper (Strauß *et al.*, 2019) and references therein for a detailed discussion of how inference of  $\mathbf{o}$  may be performed when clustering structure among the genes is ignored. An inferred ordering,  $\mathbf{o} = (o_1, ..., o_T)$ , can be mapped to pseudotimes  $\mathbf{\tau}(\mathbf{o}) = (\tau_1(\mathbf{o}), ..., \tau_T(\mathbf{o}))$  using approximate geodesic distances (Tenenbaum *et al.*, 2000) between the ordered cells.

# 2.2 Hierarchical GPs for pseudotemporal data

GPseudoClust models cluster-specific latent pseudotime profiles as well as gene-specific latent profiles which deviate from the cluster-wide profile to some extent (see Fig. 1) using hierarchical GPs. We briefly describe GPs below, and refer to Hensman *et al.* (2013, 2015) for full details of hierarchical GPs. A GP (Rasmussen and Williams, 2006) is a distribution over functions that is specified using a mean function  $\mu$ 



Fig. 1. One cluster in the hierarchical GP model. Left: cluster-wide latent mean; right: cluster-wide latent mean and latent mean for each gene in the cluster

and a covariance function  $\Sigma$ . For an input vector  $\tau(\mathbf{o}) = (\tau_1, \ldots, \tau_T)$ of pseudotime points depending on orders  $\mathbf{o}$ ,  $\mu(\tau(\mathbf{o}))$  is a vector of Tfunction evaluations of the mean function  $\mu$  and  $\Sigma(\tau(\mathbf{o}))$  is a  $T \times T$ matrix of covariance function evaluations of  $\Sigma$ . The distribution of functions  $f \sim GP(\mu(\mathbf{o}), \Sigma(\mathbf{o}))$  is described by stating that, for any vector of pseudotime points  $\tau(\mathbf{o}) = (\tau_1(\mathbf{o}), \ldots, \tau_T(\mathbf{o}))$ , the corresponding function evaluations  $f(\tau_i(\mathbf{o}))$  are distributed according to a multivariate Gaussian:  $(f(\tau_1(\mathbf{o})), \ldots, f(\tau_T(\mathbf{o}))) \sim \mathcal{N}_T(\mu(\tau(\mathbf{o})), \Sigma(\tau(\mathbf{o})))$ . Here we use a squared exponential covariance function to define  $\Sigma$ :

$$\left[\Sigma(\tau(\mathbf{o});\sigma_w^2,l)\right]_{ij} = \sigma_w^2 \exp\left(-\frac{(\tau_j - \tau_i)^2}{2l^2}\right)$$
(1)

where  $\sigma_{iv}^2$  is a scale parameter and *l* a length scale, and  $[.]_{i,j}$  refers to the element in row *i* and column *j* of a matrix.

GPs have previously been used for pseudotime ordering—see Ahmed *et al.* (2019), Campbell and Yau (2016), Reid and Wernisch (2016), Strauß *et al.* (2019) and Welch *et al.* (2017)—as well as for clustering time-stamped bulk gene expression data, see Cooke *et al.* (2011), Hensman *et al.* (2013), Kirk *et al.* (2012) and McDowell *et al.* (2018).

#### 2.3 Clustering model

We use DPs (Ferguson, 1973) as a Bayesian non-parametric way of performing model-based clustering. A DP is a distribution over discrete distributions; that is, each draw from a DP is itself a distribution. More precisely,  $G \sim DP(\alpha, G_0)$  signifies that for any partition  $B_1, \ldots, B_r$  of a parameter space  $\Theta$ , we have  $(G(B_1), \ldots, G(B_r)) \sim \text{Dirichlet}(\alpha G_0(B_1), \ldots, \alpha G_0(B_r))$ , where the Dirichlet distribution with *r* categories and concentration parameters  $(\gamma_1, \ldots, \gamma_r)$ is defined as follows:  $p(x_1, \ldots, x_r) = \frac{\Gamma(\sum_{k=1}^r \gamma_k)}{\Gamma(x_1, \ldots, x_r)} \prod_{k=1}^r x_k^{\gamma_k-1}$ 

is defined as follows: 
$$p(x_1, \dots, x_r) = \frac{\prod_{k=1}^r \gamma_k}{\prod_{k=1}^r \Gamma_{(k)}} \prod_{k=1}^r x_k^{\gamma_k-1}$$

Conditional on the order o of the cells, the allocation of genes to clusters is modelled as a DP mixture model of hierarchical GPs as follows. We model the latent cluster means  $\mu_{j}, j = 1, ..., n_g$  (see Fig. 1 (left),  $n_g$  is the number of genes) as being drawn from a DP with base distribution  $G_0$ , where:

$$\begin{array}{ll} G_0|\mathbf{o}, a, \epsilon, L \sim GP(0, \Sigma(\tau(\mathbf{o}), 3a^2 + \epsilon, L)) \\ \alpha \sim \quad \text{Gamma}(2, 4) \qquad G|G_0 \sim DP(\alpha, G_0) \qquad \mu_i|G \sim G \end{array}$$

Here, 0 represents the zero function, and  $\Sigma$  is defined as in Equation (1).  $\tau(\mathbf{o})$  is the vector of pseudotimes corresponding to cell order **o** (see Section 2.1). *L* is the length scale of the GP,  $\sigma_W^2 = 3a^2 + \epsilon$  the scale parameter corresponding to  $\sigma_w^2$  in Equation (1). This specific parametrization of the scale parameter of the latent mean profile links it to the scale and noise parameters of the deviations from the cluster-specific mean profile of the gene-specific pseudotime profiles (see Fig. 1 for an illustration).

It should be noted that while we draw a mean  $\mu_i$  for each gene  $j = 1, \ldots, n_g$ , the DP determines a number  $K \ll n_g$  and values  $\eta_1, \ldots, \eta_K$  such that for all  $j = 1, \ldots, n_g$  there is a  $k \in \{1, \ldots, K\}$  such that  $\mu_j = \eta_k$ . That is, the latent means  $\mu_j$  only take K distinct values and there are K groups of genes with identical latent means, which form a total of K clusters. The number of clusters is not fixed, but automatically determined as part of the inference for the model.

Individual gene expression pseudotime profiles are modelled by GPs with mean  $\mu_j$ ,  $j = 1, ..., n_g$  ( $n_g$  is the number of genes). GPseudoClust uses as input pre-processed log-transformed gene expression data  $y_g(\mathbf{o})$  for gene  $g = 1, ..., n_g$ . Conditional on the pseudotime ordering  $\mathbf{o}$  of the cells, the trajectory  $\mathbf{y}_j(\mathbf{o})$  of gene j is distributed as  $\mathbf{y}_i(\mathbf{o})|\mu_i, a, a_1, \mathbf{o} \sim F$ , where

$$F = GP(\mu_i, \Sigma(\tau(\mathbf{o}), a^2 \cdot a_1, 1) + a^2(1 - a_1)I_T)$$
(3)

$$a_1 \sim \text{Beta}(4, 1) \tag{4}$$

 $\Sigma$  is as in Equation (1),  $I_T$  refers to the *T*-dimensional identity function. Note that  $a_1$  represents how much variation from the cluster-wide mean is due to stochastic variation from the underlying stochastic process, while  $1 - a_1$  represents the proportion of the variation resulting from noise. By Equation (3), the ordered gene expression levels  $\mathbf{y}_i(\mathbf{o})$  of gene *j* are noisy representations of individual gene-specific latent means drawn from a GP with cluster-specific mean function.

# 2.4 Markov Chain Monte Carlo sampling and block matrix representation

We use Markov Chain Monte Carlo sampling (MCMC; Gilks et al., 1996) for inference of pseudotime orderings and cluster assignments. This allows sampling from the joint posterior probability distribution of clusters, orders and hyperparameters a, L,  $a_1$  and  $\epsilon$ . For the orders, which are sampled from the discrete space of all possible permutations of cells, we previously developed an efficient sampling strategy (Strauß et al., 2019). To reduce the number of parameters, we integrate out the cluster-specific mean profiles, and developed an efficient method for inverting the resulting block matrices, thereby reducing the computational complexity of the operations needed to obtain inverses and determinants of cluster-specific covariance matrices from  $O(n_c^3 T^3)$  to  $O(T^3)$ , where  $n_c$  is the number of genes in cluster *c* and *T* the number of cells. Although the resulting likelihood computations are similar to those derived in Hensman et al. (2013), the approach presented here additionally provides a general method for computing inverses and determinants of matrices of certain types of block structures. For details, see Supplementary Sections S1.2 and S1.3.

#### 2.5 Subsampling strategies

Sampling orders of cells and clusters of genes simultaneously is a challenging high-dimensional problem, in particular as the posterior distribution of the orders is typically highly complex (see Strauß *et al.*, 2019). We improve convergence by using parallel MCMC chains on subsets of cells. The chains are subsequently combined to a summary result approximating the posterior distribution of the cluster allocations.

#### 2.5.1 Posterior similarity matrices

A central step is the computation of posterior similarity matrices (PSMs) for each of the chains on subsets of cells. The PSM is the symmetric positive semidefinite (see Lemma 4 in Supplementary Section S2.2) matrix whose entry in the *i*th row and *j*th column is the frequency with which gene *i* and gene *j* are clustered together among the samples drawn from the posterior distribution of cluster allocations. This estimates the posterior probability of the two genes being in the same cluster (Fritsch and Ickstadt, 2009).

#### 2.5.2 Obtaining summary clusterings from PSMs

Although the uncertainty of the cluster allocations obtained for single-cell datasets does not always justify a single summary clustering, it can nevertheless sometimes be useful to compute summary clusterings for validation and comparison purposes. In addition, the methods presented below to find weights for combining the PSMs obtained from the individual subsampled MCMC chains into one joint PSM also require summary clusterings of individual PSMs. To obtain a summary clustering from a PSM, we apply hierarchical clustering to the columns of the PSM using 1 – PSM as the distance matrix (Medvedovic et al., 2004). The optimal number of clusters is determined by a method maximizing the posterior expected adjusted Rand index (PEAR) between the inferred summary clustering and the unknown true clustering structure (Fritsch and Ickstadt, 2009). The adjusted Rand index (ARI, Hubert and Arabie, 1985; Rand 1971), see also Section 2.7 and Supplementary Section S4) is a measure of agreement between two clusterings. The PEAR is therefore a measure of how well the inferred summary clustering is expected to agree with the unknown true clustering.

### 2.5.3 Combining PSMs

The following methods for combining the PSMs from the individual MCMC chains on subsampled data to obtain a joint overall PSM are proposed here:

Method 'mean psm'. The first method proposed to obtain a joint PSM is to compute the element-wise unweighted arithmetic mean of

the PSMs of the individual chains. This method is referred to as 'mean PSM' here.

Methods 'PY and PEAR', 'DPM and PEAR'. As noise levels tend to differ between subsamples of cells, an unweighted average of the PSMs may not always be the best representation of the overall posterior distribution. We propose new methods to obtain a final PSM as a weighted average of the PSMs of the individual subsampled chains. We propose a novel method using DP mixture models or Pitman-Yor process (PY, Ishwaran and James, 2001; Pitman and Yor, 1997), a generalization of the DP) mixture models with feature selection to compute the weights. We refer to the two methods as 'PY and PEAR' and 'dirichlet process mixture model (DPM) and PEAR'. For details see Supplementary Section S2.1.

Method 'lmkk'. The differences in noise for different subsampled chains may be gene-specific; to address this, this method applies localized multiple kernel k-means (lmkk, Gönen and Margolin, 2014) to obtain a summary clustering from the set of PSMs for the different chains. lmkk was first used to obtain summary clusterings from consensus clustering matrices in Cabassi and Kirk (2019). Unlike the other methods proposed in this section, the 'lmkk' method does not aim to provide a full estimate of the overall PSM, but it is an optimization method to find a summary clustering from multiple PSMs. The method proposed in this paper also finds weights for an overall summary matrix representation of posterior cluster allocation probabilities. For details on our approach see Supplementary Section S2.2.

#### 2.6 Assessment of convergence

Our approximate sampling approach using parallel MCMC chains on subsamples of cells requires us to assess convergence across subsampled chains with different cells. We consider a number of different criteria to assess convergence, which we describe in detail in Supplementary Section S3.

#### 2.7 Alternative clustering methods and assessment

We compare GPseudoClust to several widely used standard clustering methods, which we applied to the simulated and Shalek datasets: mixture of normals (mclust, Fraley and Raftery, 2002; Scrucca et al., 2017), k-medoids clustering (PAM, Kaufman and Rousseeuw, 2008; Maechler et al., 2019), hierarchical clustering and SIMLR. In addition, we applied the following two-step methods (first pseudotime ordering of cells, then clustering of genes in a second step): SLICER (Welch et al., 2016) and DeLorean (Reid and Wernisch, 2016) combined with GPclust (Hensman et al., 2013, 2015), and Monocle 2. For the simulated datasets the following measures of comparison between the true and the inferred cluster allocations are used: the ARI (Hubert and Arabie, 1985; Rand, 1971), the Fowlkes-Mallows index (FMI) and normalized mutual information (NMI; Kvalseth, 1987). For all of these measures a score of one signifies perfect agreement between true and inferred cluster allocations. For a definition of the measures and information concerning parameter settings for the methods listed above see Supplementary Section S4.

# **3 Results**

We provide details of the simulated and real datasets to which we applied GPseudoClust (see Section 3.1), followed by a summary of our results (see Sections 3.2 and 3.3), with further details in the Supplementary Material (as indicated).

#### 3.1 Datasets

#### 3.1.1 Simulated datasets

Simulation Studies 1 and 2. For each of these two simulation studies, we simulated 100 datasets with each dataset having five clusters. The specific construction of the datasets is tailored such that datasets in simulation Study 1 have very clearly separated clusters, whereas datasets in simulation Study 2 have clusters that are not easily separable (see Supplementary Fig. S1 for examples of the simulated datasets). scRNA-seq data often consist of large numbers of repeated measurements at a few capture times. To mimic this situation, we assume 3 capture times for the simulated cells: the first 20 cells have capture time 1, cells 21–40 have capture time 2 and 41–60 capture time 3. We remove information about the true order by applying a random permutation to the order of the cells within each capture time, to mimic the lack of temporal information in applications. For both simulation studies, all datasets were generated using GPs, but not the same GP model as GPseudoClust. For a detailed description of the simulation set-up see Supplementary Section S5.

Simulation studies with dropout noise. scRNA-seq data are affected by technical noise leading to zero-expression values when the gene is actually expressed in the cell. To study the robustness of the method to technical zero-inflation without the presence of any other confounders, we use one of the datasets which we used to validate the subsampling procedures (see paragraph Simulation Studies 1 and 2 above and Supplementary Fig. S1), and set non-zero values to zero at random. Note that while we could have used a dropout rate which depends on the actual gene expression level, with higher expression levels associated with lower probability of dropout (Pierson and Yau, 2015), our way of testing the robustness is more stringent since it permits larger perturbations. This additional simulation study comprises three sets of 100 datasets, to test for robustness of the GPseudoClust method and all of the proposed subsampling methods (see Section 2.5) to different levels of dropout, including a simulation study for which different groups of genes are affected by dropout to different degrees. The three dropout-related simulations were repeated 100 times each. For details, see Supplementary Section S7.1.

Additional simulations: misspecified covariance functions and large datasets. To test robustness of clustering results to covariance function misspecification, we also simulated 24 datasets each using hierarchical Matérn-3/2 and linear covariance functions, each dataset with different random parameters and cluster allocations. To assess the performance of the subsampling methods, we simulated datasets with hierarchical Matérn-3/2 covariance matrices, and 9000 simulated cells, and compared clustering results obtained by using different numbers of subsampled cells (see Supplementary Section S8 for details).

#### 3.1.2 Experimental datasets

Overview. Here we provide a brief overview of our analyses, before providing a fuller description of the datasets in the paragraphs below: (i) We apply GPseudoClust to branching data (Moignard data below), which confirms existing results and also finds new results on differences of cluster structures of genes for different branches; (ii) The effectiveness of the subsampling approach with parallel chains each run on a subset of cells is investigated by applying GPseudoClust both with and without subsampling to a dataset with 600 genes and 35 cells (Sasagawa data below); (iii) GPseudoClust is also applied to non-branching data (Shalek data); (iv) the subsampling method and the combination of weighted PSMs are used to integrate data from different cell lines (Stumpf data); and (v) the GPseudoClust method is compared with our previous method for ordering cells under uncertainty, using a small scRNA-seq dataset of lipopolysaccharide (LPS)-stimulated mouse dendritic cells (Shalek13 data) previously analysed in Strauß et al. (2019). Details on numbers of MCMC chains and subsampled cells for the different datasets are provided in Supplementary Section S6 and Supplementary Table S5.

Moignard data. Moignard *et al.* (2015) applied single-cell RTqPCR to 3934 mouse early haematopoietic cells. In an in-vivo experiment cells were captured at four time points between embryonic day 7.0 and 8.5. In Moignard *et al.* (2015), Haghverdi *et al.* (2015, 2016) diffusion maps (Coifman *et al.*, 2005) are used to identify two branches, a blood and an endothelial branch. We use the preprocessed (Haghverdi *et al.*, 2016; Moignard *et al.*, 2015) data available as Supplementary Material to Haghverdi *et al.* (2016). Before the application of GPseudoClust, branches are inferred using diffusion maps, as in Haghverdi *et al.* (2016), which leads to the identification of an endothelial and an erythroid branch. We use diffusion maps for the identification of the branches, but find cluster allocations and their uncertainties using GPseudoClust without prior pseudotime ordering.

Sasagawa data: Mouse embryonic stem cells, cell cycle-related genes. GPseudoClust is also applied to a Quartz-Seq (FPKM normalized) dataset of 35 mouse embryonic stem cells (Sasagawa *et al.*, 2013), on cell cycle-related genes. Cell cycle genes were selected by finding genes associated with GO: 0007049, as in Buettner *et al.* (2015).

Shalek data: LPS-stimulated mouse dendritic cells, scRNA-seq. Shalek *et al.* (2014) examined the response of primary mouse bone-marrow-derived dendritic cells in three different conditions using scRNA-seq. We applied GPseudoClust to a version of the Shalek dataset previously considered in an earlier paper (Reid and Wernisch, 2016) comprising 74 genes (which have the highest temporal variance relative to their noise levels) and to the 183 cells from the LPS-stimulated condition and capture times 2, 4 and 6 h, dropping the cells captured at 0 and 1 h, to focus on differences between gene expression levels in reaction to the stimulus rather than before the reaction has set in. The data were log-transformed, and an adjustment for cell size applied, according to Anders and Huber (2010) and Reid and Wernisch (2016).

**Stumpf data.** Stumpf *et al.* (2017) generated an RT-qPCR dataset for 94 genes from two cell lines following the progression of mouse embryonic stem cells along the neuronal lineage, containing 96 cells per capture time (0, 24, 48, 72, 96, 120 and 172 h). The proposed subsampling methods allow taking subsamples of cells from each cell line separately and combining the chains as described in Section 2.5. For the pre-processing, the steps described in Stumpf *et al.* (2017) were applied to each cell line separately. The raw data are available on Mendeley Data (http://dx.doi.org/10.17632/g2md5gbhz7.1).

Shalek13 data. Shalek *et al.* (2013) obtained scRNA-seq data from mouse bone-marrow-derived dendritic cells after exposure to LPS. 18 cells were captured 4 h after initial exposure. We use this dataset of smaller size to compare orders obtained by GPseudoClust without subsampling to the GPseudoRank method (Strauß *et al.*, 2019). For details, see Supplementary Section S9.



Fig. 2. Results from simulation studies 1 (top) and 2 (bottom). Performance is assessed using 3 scores (ARI, FMI and NMI), with higher values indicating better performance. Methods compared: GPs+mean = GPseudoClust and 'mean psm', GPs+DP = GPseudoClust+'DPM+PEAR', GPs+PY = GPseudoClust and 'PY+PEAR', GPs+lmkk = GPseudoClust method followed by summary clustering using lmkk, Mon 2 = Monocle 2 (two steps: ordering and then clustering), De+GCI = DeLorean & GPclust (two steps), SL+GCI = SLICER & GPclust (two steps), hier = hierarchical clustering, PAM, SIMLR, mcl = mclust. For the GPs+lmkk method there were rare cases in which the lmkk algorithm failed due to numerical issues. In these cases we set all 3 scores to 0

### 3.2 Simulation study results

# 3.2.1 The performance of GPseudoClust is robust to different levels of cluster separability in our simulation studies

Figure 2 illustrates the importance of using methods modelling the pseudotemporal nature of the data. It includes results for point estimates obtained by combining the GPseudoClust method with the proposed methods to obtain a joint PSM from several subsampled chains (see Section 2.5). Except for lmkk, where the method itself provides a summary clustering, a final summary clustering was obtained from the summary PSM by means of hierarchical clustering and the PEAR criterion.

Although for datasets with clearly separated clusters most clustering methods will perform satisfactorily (Fig. 2, top), this is not the case for datasets where the cluster structure only becomes apparent through modelling the data as a pseudotime series (see Fig. 2, bottom). In the latter case GPseudoClust, which jointly models pseudotime and cluster structures, performs best, while mclust and SIMLR perform best among those methods not incorporating the pseudotime structure.

#### 3.2.2 Robustness to dropout

Further simulation studies on a total of 300 datasets (dropout Studies 1–3, see Section 3.1.1) with different levels of dropout noise demonstrate the robustness of GPseudoClust. For details, see Supplementary Section S7.2. Supplementary Figure S3 shows high ARIs with the true clustering for summary clusterings obtained by means of GPseudoClust and the proposed subsampling methods. While all the subsampling methods have a similar level of robustness to dropout noise when all genes are affected for all cells with equal probabilities (see Supplementary Fig. S3), the 'lmkk' method is shown to be the best performing one for the case where there are groups of cells known to be less affected by dropout for a subset of the genes (dropout Study 3; see Supplementary Fig. S4). Supplementary Section S7.2 also presents comparisons of the summary PSMs obtained using the different subsampling methods (see Supplementary Figs S5–S13).

#### 3.2.3 Robustness to covariance misspecification

As illustrated by Supplementary Figures S14 and S15, GPseudoClust is robust to covariance matrix misspecification across a range of scenarios. Our results show that the degree of cluster overlap is generally more important, in terms of affecting our ability to uncover clustering structure.

# 3.2.4 Gaining efficiency and maintaining accuracy with subsampling

The simulation studies with 9000 cells and different noise levels show that 30 subsampled cells (10 per capture time) per chain permitted a good approximation of the true cluster structures both for simulated datasets with lower and higher noise levels (Supplementary Figs S16, S17, S20 and S21). The across-chain convergence measures specified in Supplementary Section S3 indicate 12 chains with 10 subsampled chains as sufficient (Supplementary Figs S18 and S22). Supplementary Figure S19 illustrates the efficiency gain obtained by subsampling. Note that without subsampling, but with efficient inversion of block matrices (see Section 2.4), computational complexity scales as the cube of the number of cells, while with subsampling it scales as the cube of the (smaller) number of subsampled cells per chain (see also Supplementary Fig, S49).

#### 3.3 Experimental data results

#### 3.3.1 Validating subsampling: Sasagawa data

The Sasagawa dataset has only 35 cells, which makes it suitable for comparing the proposed subsampling methods to applying the GPseudoClust method to all the cells.

Figure 3 illustrates good convergence of the GPseudoClust method with and without subsampling. Moreover it demonstrates that the proposed subsampling methods 'PY and PEAR' and 'DPM



GPseudoClust with subsampling: 36 subsampled chains with 15 cells each.

Fig. 3. Sasagawa data: (a) illustrates four PSMs obtained without subsampling, by applying GPseudoClust to all cells for each of the four chains (b) compares the proposed subsampling methods 'PY and PEAR', 'DPM and PEAR', and 'lmkk'

and PEAR' lead to PSMs convincingly similar to the ones obtained without the subsampling, and that a similar matrix is obtained using lmkk. For further confirmation of convergence see Supplementary Figure S48.

#### 3.3.2 Immune response genes cluster around functional profiles

The genes analysed for the Shalek data (see Section 3.1.2) are from three modules identified in Shalek *et al.* (2014) as 'peaked inflammatory module', which shows a 'rapid, yet transient induction' to LPS stimulation, 'core antiviral module, enriched for annotated antiviral and interferon response genes', and 'sustained inflammatory module; exhibiting continued rise in expression under LPS'. Although the analysis proved to be very stable with regard to the number of subsampled chains (Supplementary Fig. S45), for the following analysis the PSM obtained using the 'PY + PEAR' method with 96 subsampled chains is used. However, as illustrated by Supplementary Figure S45, for the 'PY and PEAR', 'DPM and PEAR' and 'mean PSM' methods a good approximation is achieved with only four randomly chosen chains. Supplementary Figure S46 further illustrates convergence by applying the measures presented in Section 2.6 and in Supplementary Section S3.

The PSMs allow the computation of (potentially overlapping) groups of genes with high pairwise co-clustering probabilities. We use a threshold of 80% for the identification of groups of genes with high pairwise co-clustering probability. The choice of 80% for the threshold is chosen to ensure that it is sufficiently stringent to allow meaningful groups to be identified, but low enough to allow reasonably sized groups to be identified. The word pairwise is used here to emphasize that this is not the probability of all the genes being in the same cluster, but that for any two genes in such a group the probability of these two genes being in the same cluster is above 80%. It should be noted that this approach is different from trying to find a single summary clustering, and that the groups will usually overlap.

GPseudoClust identifies four groups with pairwise co-clustering probabilities of more than 80%, three of which, however, have a large overlap. Therefore, we refer to the groups as 1, 2a, 2b and 2c. Group 1: Bcl2l11, Flrt3, Nfkbid, Ralgds, Rasgef1b and Socs3. All genes in this group belong to a 'peaked inflammatory module' identified in Shalek et al. (2014), which shows a 'rapid, yet transient induction' to LPS stimulation. Group 2: Ddx60, E030037k03rik, ligp1, Irf7, Mpa2l, Ms4a4c, Nlrc5, Nos2, Phf11 and Slco3a1 Group 2a: Group 2 and D14ertd668e, Dbx58, Il15. Except for Nos2, all genes in this group belong to a 'core antiviral module, enriched for annotated antiviral and interferon response genes' (Shalek et al., 2014). Nos2 is part of the 'sustained inflammatory module; exhibiting continued rise in expression under LPS'. Group

2b: Group 2 and D14ertd668e, Dhx58 and Procr. This group consists of genes from the 'core antiviral module', except for Nos2 and Procr. Group 2c: Group 2 and Il15, Procr. This group consists of genes from the 'core antiviral module', except for Nos2 and Procr.

We also applied the other clustering methods (listed in Section 2.7), to the Shalek dataset. The importance of quantifying the uncertainty of inferred cluster structures as done by GPseudoClust is highlighted by Figure 4, where the various clustering methods resulting in a single clustering disagree quite significantly, with most ARIs between pairs of results obtained by different methods <0.6. In addition, Figure 4 also shows that when the two-stage method of combining GPclust with a pseudotime method is used, the clustering result depends on the choice of the pseudotime method. GPseudoClust models the uncertainty in the cluster structures, which we generally represent by the summary PSMs. In Figure 4, the uncertainty is represented by eight random draws from the posterior distribution of cluster allocations.

#### 3.3.3 Detecting branch-dependent clustering structures

The analysis of the Moignard dataset shows very different clustering structures in the trunk, the endothelial and the erythroid branch (see Fig. 5), which shows summary PSMs obtained using the 'PY + PEAR' method for the different branches.

In this figure, the rows and columns of the four PSMs displayed are ordered in the same way to illustrate the differences in the clustering structures between the different branches.

For the trunk, *Fli1*, *Tal1*, *Etv2* and *Kdr* have high posterior coclustering probabilities, mirroring the fact that they are switched on early in the developmental process (see Supplementary Figs S28 and S29). Genes with very low expression levels in the trunk (*Gata1*, *Gfi1*, *Gfi1b*, *Hbbbh1*, *HoxB2*, *HoxD8*, *Ikaros*, *Itga2b*, *Mecom*, *Mitf*, *Myb*, *Nfe2* and *Sfpi1*) also have high co-clustering probabilities, see Supplementary Figure S30, similarly genes with relatively constant higher expression levels (*Ets2*, *FoxH1*, *FoxO4* and *Ldb1*; Supplementary Fig. S31).

For the endothelial branch, there is a group of genes with relatively constant higher expression level throughout the endothelial branch, which have high posterior co-clustering probabilities (*Cbfa2t3h*, *Cdh5*, *Egfl7*, *Erg*, *Ets1*, *Ets2*, *Etv6*, *Fli1*, *Hhex*, *Itga2b*, *Kdr*, *Kit*, *Ldb1*, *Lyl1*, *Mecom*, *Meis1*, *Notch1*, *Pecam1*, *Sox17*, *Sox7* and *Tal1*; Supplementary Fig. S34), and a group of genes which have very low expression levels or are not expressed (*Cdh1*, *Gata1*, *Gfi1*, *Gfi1b*, *HoxB2*, *HoxD8*, *Ikaros*, *Myb* and *Nfe2*; Supplementary Fig. S35).

For the erythroid branch GPseudoClust identifies again a group of genes with relatively constant higher expression levels (*Cbfa2t3h*, *Ets2*, *Etv6*, *FoxH1*, *FoxO4*, *Kit*, *Ldb1*, *Lyl1*, *Pecam1*, *Runx1* and *Tal1*, Supplementary Fig. S36). *Gata1* and *Nfe2* are switched on at similar pseudotimes in the erythroid branch (Supplementary Fig. S37), whereas *Cdh5*, *Ets1*, *Etv2*, *Fli1*, *Hhex*, *Kdr* and *Sox7* (Supplementary Fig. S38) have a marked decrease in expression around a similar pseudotime. For a detailed analysis and illustrations of the pseudotemporal dynamics of clusters of genes in different branches, see Supplementary Section S10 and Supplementary Figs S27–S40. Again we performed stringent convergence analysis to ensure across-chain convergence (see Supplementary Figs S41–S44).

#### 3.3.4 Combining multiple datasets

The subsampling methods proposed in Section 2.5 are also particularly useful in situations where we need to integrate data that were not obtained in exactly the same way, for instance because they were obtained from different cell lines or generally in slightly different experimental conditions. Instead of just blending the datasets, the subsampling method allows us to run chains for the different cell lines separately and then combine them in a principled way. The 'PY and PEAR' and 'DPM and PEAR' methods show particularly good agreement with each other (see Supplementary Fig. S47), but we also considered the 'mean psm' and 'lmkk' methods (see again Supplementary Fig. S47). Figure 6 illustrates the downweighting of those subsamples which are inconsistent with the integration of the two cell lines to a joint overall structure (weights close to 0 in Fig. 6), and highlights again the high level of agreement between the 'PY + PEAR' and 'DPM + PEAR' methods.



**Fig. 4.** ARI between results obtained by different clustering methods: Shalek dataset. A score of 1 shows that the two clusterings are identical, a score of 0 that they are no more related than expected by random chance. To represent the posterior distribution obtained by GPseudoClust, eight samples from the posterior distributions of cluster allocations were chosen randomly (Samples 1–8). mcl = mclust, hier. = hierarchical clustering, SL+GCl = SLICER+GPclust, De+GCl = DeLorean+GPclust, Mon 2 = Monocle 2



Fig. 5. Moignard data: PSMs for branches. PSMs for each branch are obtained by the 'PY and PEAR' method. 'Mean PSM' refers to the unweighted mean of the summary PSMs of the three branches. A summary clustering was obtained from the mean PSM to order each of the matrices in the same way



Fig. 6. Stumpf data: comparison of weights for 'PY and PEAR' (asterisks) and 'DPM and PEAR' (circles) methods. The weights are plotted along the y-axes and are sorted in the same way for both methods (ascending for 'PY and PEAR')

**3.3.5** Comparing GPseudoClust and GPseudoRank (Shalek13 data) The distributions of cell orderings obtained by the two methods are similar, but the accounting for the uncertainty in cluster allocations by GPseudoClust reveals greater uncertainty in the orderings. We have found that GPseudoClust is more likely to get stuck in local posterior modes than GPseudoRank; e.g. Supplementary Figure S23 shows that, for GPseudoClust, different MCMC chains visit different posterior modes. Although this is a drawback compared with GPseudoRank (where convergence was achieved with individual chains exploring multiple modes of complex posterior distributions), we note that it is mitigated by our subsampling strategies, which combine posterior samples across chains. In general, the existence of multiple modes highlights the benefits of adopting a Bayesian approach and running multiple chains, compared with strategies that seek a single, locally optimal result. For details see Supplementary Section S9.

## **4 Discussion**

GPseudoClust is a Bayesian non-parametric method for the clustering of genes for single-cell RNA-seq and RT-qPCR data in terms of latent shared pseudotime expression profiles. Applying the method to simulated data shows that unless the clusters are very clearly separated from each other, clustering methods not incorporating the pseudotemporal nature of the data may not be effective. Although it is possible to combine pseudotime ordering and clustering methods in a two-step process, applications to both simulated and experimental data lead to clustering results with a dependence on the pseudotime method used (see Figs 2 and 4). In an application to dendritic cells GPseudoClust identifies clusters of genes closely associated with their biological function, and shows that there is considerable uncertainty in the clustering structures. GPseudoClust captures this uncertainty by providing a distribution of posterior co-clustering probabilities rather than just one single 'point estimate' of a clustering. An application to branching data from early haematopoeitic cells demonstrates the ability of the method to identify strong differences between the clustering structures of the different branches. GPseudoClust identifies genes switched on or off at similar times in pseudotime as being co-clustered with a high probability. The uncertainty of clustering structures learned from the posterior distribution as represented by the PSM allows us to understand similarity of genes in terms of pairwise co-clustering probabilities. An application to data obtained from different cell lines illustrates the ability of the method to analyse different datasets studying the same developmental process. GPseudoClust can be used to combine studies with different experimental protocols with different levels of measurement noise. The methods for finding weighted averages from multiple PSMs proposed here are designed to discard chains inconsistent with the overall clustering structure. We note that GPseudoClust could also be used to perform meta-analyses of previous studies, thanks to its ability to integrate datasets obtained under different experimental conditions. This may be of interest beyond the study of single-cell gene expression data.

GPseudoClust, which uses MCMC methods to sample from a highly complex joint posterior distribution of both cell orders and gene clusters, was designed to infer cluster structures accounting for pseudotemporal uncertainty and not to compete, in terms of computational speed, with (e.g.) efficient variational methods for pseudotime ordering. Nevertheless, we found that computation times (see Supplementary Table S5) still reflect the efficiency of our inference method given the hugely complex inference task. GPseudoClust scales linearly with both the number of genes and the number of clusters. In terms of computation time, it is therefore feasible to apply it to larger numbers of genes. Here we have presented applications to datasets where genes were selected whose expression varies across pseudotime, for the Shalek dataset we refer to previous publications for this purpose (Reid and Wernisch, 2016; Strauß et al., 2019). An exception are two RT-qPCR datasets, the Moignard dataset (Moignard et al., 2015), which contains only 42 genes (33 transcription factors important in endothelial and haematopoietic development and 9 marker genes), and the Stumpf dataset (Stumpf et al., 2017), which contains 94 genes (excluding 2 housekeeping genes). Generally, we would recommend pre-selecting genes with pseudotemporal variation by pretesting if there are differences in gene expression across capture times; e.g. as in Strauß et al. (2019).

### Acknowledgements

We would like to thank Sascha Ott and William Astle for feedback and insightful comments and the anonymous reviewers for their helpful suggestions, which have benefitted this work.

#### Funding

This work was supported by the UK Medical Research Council [MC\_UU\_00002/1 to M.E.S., J.E.R. and L.W., and MC\_UU\_00002/10 and MC\_UU\_00002/13 to P.D.W.K].

Conflict of Interest: none declared.

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