Bayesian Clustering and Topic Discovery: Adventures with Gene Expression Data

Karren Dai Yang, Skanda Koppula

{KARREN, SKOPPULA}@MIT.EDU

Massachusetts Institute of Technology, Cambridge, MA 02139 USA

1. Introduction

Tumors cell lines are composed of different subpopulations of cells which often exhibit shared patterns of gene expression. Biologists are interested in two key questions: given gene expressions values, (1) can we identify biologically meaningful gene modules, and (2) can we identify cell sub-populations and find gene modules that illuminate their differences? Our goal with this project was to address these questions using Bayesian methods.

To address the first question, we explored the use of Latent Dirichlet Allocation (LDA), two non-parametric topic models, and a dynamic-topic LDA. To address the second question, we explored the use of a finite mixture model as well as a clustering topic model.

1.1. Description of Data

To do our analysis, we used a single-cell RNAsequencing (scRNA-seq) dataset obtained from human melanoma samples (Tirosh, 2016). The data consisted of the expression values of 22,712 genes for each of 4645 cells. In total, the dataset amounted to 0.86 GB, presenting significant computational challenges when attempting posterior inference. Apart from a few computational tricks (online inference, multi-core parallelization), for most methods, we pruned noninformative genes from the dataset, ranking based on the deviation in value across cells ¹. The Seurat biological toolkit in R was also used for the purpose of lowvariance feature selection (Satija, 2012). We eschewed dimensionality reduction techniques such as PCA because of loss of its direct feature interpretability. Prior researchers have labeled the cells in our dataset; there are a total of 9 cell categories (Tirosh, 2016). The complete dataset, as well our preprocessed and pruned versions, are openly available (Yang & Koppula).

1.2. Prior work

Prior research has explored the use of various computational techniques to analyze gene expression data. Most commonly, Spearman and Pearson correlation metrics are frequently used infer sets of genes that cluster together (Xie, 2015; Borenszstein, 2017). Other techniques, including PCA followed by linear regression, has been used for expression-based cell clustering (Stegle, 2016). Yu et al. propose an unsupervised classifier ensemble as another approach to cell clustering (Yu, 2016).

More Bayesian approaches have also been tried in prior work from the Pe'er lab. (Prabhakaran, 2016) uses a Heirarchical Dirichlet Mixture Model to learn cell clusterings. (Azizi, 2017) builds on this to jointly learn optimal normalization pre-processing of the data. Bayesian networks have also been used in an attempt to learn gene dependencies from expression data (Pe'er, 2000). Our work uses different models to explore gene expression data, but where appropriate (e.g. LDA vs. non-parametric models), we compare results.

1.3. Structure of Report

We first discuss our experiments using Bayesian topic models to discover topics in scRNA-seq data: LDA in Section 2, non-parametric topic models in Section 3 and Dynamic Time Models in Section 4. Then, we discuss our experiments in clustering: Finite Mixtures in Section 5, and Integrated Topic-Clustering in Section 6. We conclude our paper with our observations from across all our studies.

For the purpose of reproducibility, all code can be found at https://github.com/skoppula/882.

2. Latent Dirichlet Allocation

2.1. Model Description

In the generative process for LDA, the topic assigned to each word is a drawn from the document's topic

¹We recognize that this can bias towards noisy genes. We favor this method because it is simple and easy to implement, and a practice used in literature (Ling, 2012)

distribution. The identity of the word is drawn from topic's word distribution. Assuming the reader is familiar with LDA, we relegate further details and formalization of the model to (Blei, 2003).

In the context of analyzing gene expression data, we are interested in discovering 'topics' that comprise of a set of top-N genes within the topic distribution that are biologically related. For example, together the genes may direct a specific chemical function in a cell. Biologists denote such sets of genes as 'gene modules' which can be cross-referenced with existing gene module databases.

2.2. Implementation

A first attempt using the built-in Python 1da package resulting in early memory overflows during what we suspect was pre-allocation of per-document variables. The source code was not available, so we had few clues.

We switched to two open-source implementations: an online mean-field variational Bayes for posterior estimation (Nothman, 2017), and a broken C++ Gibbs sampler for LDA (OpenDataGroup, 2015).

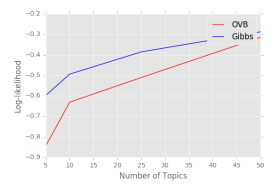
We fixed portions of the sampler to compile properly and extended the sampler to run across four cores. Details of our sampling procedure can be found in Appendix 9.2. We compared these two posterior estimation approaches using our entire dataset, using a 10% held-out testing partition. We experimented using k=5,10,25,50 topics. We did not require dataset pruning after these optimizations.

2.3. Experiments

We evaluated the interpretability of our topics (ranked lists of genes) using hypergeometric tests (Wagner, 2015). In brief, this determines the enrichment of our topics for any existing collections of genes catalogued by biologists in the gene module database MSigDB (Broad-Institute, 2015). More details about the score can be found in the Appendix.

Figure 9 in the Appendix shows gene module matches in MSigDB for which the p-value of the match is at least less than 0.3. Notice that models with more topics tended to have more matches with higher significance. The p-values in our tests do not factor for multiple hypothesis corrections, so at the moment we are only using them as relative measures of model quality.

It is validating to see that many of the similar modules, such as the BRCA1 and DOXORUBICIN, are cancer related, given that our expression assay was from tumor cells.



 $Figure\ 1.$ Log-likelihood of the held-out testing set across numbers of topics. Higher topic sizes are explored in later models.

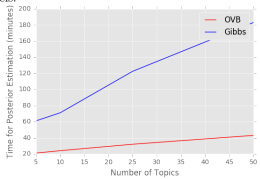


Figure 2. Comparison of the running times of each of the posterior estimation methods across various numbers of topics. Termination conditions across all runs were constant, and described in Appendix 9.2

Extracting the parameters of each model, we calculated log-likelihood on our held-out test set. This is shown in Figure 1. We found that for this dataset increasing the topic count increases log-likelihood, roughly linear from 5 to 50. Further experiments will need to test at what point log-likelihood drops off; this is one way to optimize the number of topics.

As an interesting aside, Figure 2 shows the time until inference completion (collected during our experiments). Gibbs appears to scale poorly with the parameter dimensionality, in contrast to online variational Bayes ².

Posterior predictive checks to test the mutual information between the each cell and its words' topics is something that we did not have time, but would be interesting to examine. It's not clear that this inde-

²It is important to note that comparing the magnitude of the time to complete inference in Figure 2 is not particularly meaningful; as described in Appendix 9.2, we chose a reasonable guess for the number of iterations and optimization threshold after which to stop estimation

pendence assumption is true in the context of gene expression data, because of cross-talk and regulatory mechanisms between gene modules.

3. Non-Parametric Topic Models

3.1. Model Description

One limitation of LDA is the need to define the number of topics a priori. Non-parametric models overcome this limitation by assuming a countably infinite number of topics. We trained and compared two nonparametric topic models with LDA: the Hierarchical Dirichlet Process (HDP) and the Indian Buffet Process Compound Dirichlet Process (IDP-DP). On a high level, HDP is very similar to LDA, with the main difference being that the draws from a Dirichlet distribution in LDA are replaced with draws from a Dirichlet process in HDP (Blei, 2005). One drawback to the HDP is that there tends to be a correlation between how frequently a topic appears across all documents and how prevalent this topic is within documents that it appears in. This may be undesirable in circumstances where there are 'rare' topics with high prevalence in a small number of documents. IBP-DP tries to overcome this problem by separately modeling the frequency of documents that contain a topic and the prevalence of this topic within a document where it is present (Blei, 2005). Both generative models are described in greater detail in the Appendix (sections 9.3 and 9.4).

3.2. Experiment: LDA vs. HDP

We trained an HDP topic model and compared it to LDA models with similar numbers of topics, using held-out perplexity and gene set enrichment analysis as described above as our evaluation metrics. We used an existing C++ implementation of the Gibbs sampler provided by the authors of the original paper (Blei, 2005). Since the available implementation of the posterior inference algorithm for HDP was too slow to run on the entire dataset, we did feature selection using the Seurat toolkit in R (Satija, 2012) to reduce the number of genes in our dataset with low variance across cells, and trained all models on this subset of data. We tuned the concentration parameters in HDP to obtain a reasonable number of topics (i.e. 150); subsequently, we trained LDA models with similar numbers of topics for comparison. We found that the HDP model had higher perplexity on a held-out dataset (Figure 11). Moreover, the topics in the HDP model tended to have less significant enrichment for known gene sets from MSigDB, and we found that this difference to be significant (Figure 12, Mann-Whitney-U test, p < 0.0001). Overall, the LDA model proved superior to the HDP model in this experiment.

3.3. Experiment: LDA vs. IBP-DP

One drawback to the HDP is that there tends to be a correlation between how frequently a topic appears across all documents and how prevalent this topic is within documents that it appears in. Williamson et al. (Blei, 2010) proposed the 'focused topic model' to overcome this drawback. We implemented their model to compare it with LDA on the Reuters-21578 dataset. Since the code from the original IBP-DP paper was not available, we implemented an inference algorithm using collapsed Gibbs sampling (Blei, 2010). Due to the non-conjugacy of the model, sampling each latent variable from its full conditional required using another sampling method. To sample the topics parameters, the number of which changes depending on how many topics are represented in the dataset, we used slice sampling based on the semi-ordered stick-breaking representation of the model (Teh, 2007). For more details about implementation, please refer to those papers.

We tested our code on a subset of the Reuters-21578 dataset, using several different values of the concentration hyper-parameter α , which influences the number of clusters. Although higher values of α yielded better log-likelihood values, we found that it resulted in a large number of very small topics, which are not very useful (Figure 13). Qualitatively, we did not find the topics from IBP-DP (Figure 14) to be more coherent than topics than LDA (Figure 15). The most prevalent topics from the IBP-DP each corresponded to similar topics from LDA; less prevalent topics tended to consist of a few unrelated words. These results discouraged us from optimizing the code to train this model on our scRNA-seq dataset, as we do not think it would yield more coherent gene modules than LDA. We emphasize that these results are not completely unexpected, as the authors of the IBP-DP paper did not show any topics from their model, nor did they assess the quality of their model with metrics other than perplexity.

4. Dynamic Topic Model

4.1. Model Description

As tumor cells proliferate, they undergo a process called *differentiation*. The set of cells types during tumor emergence may differ from the set of types expressed days later. Correspondingly, patterns of gene

expression may also change over time. The set of gene modules expressed in the cells, or the composition of each gene module (topic), may also change.

To capture these evolving topics, we look to dynamic topic models (Blei & Lafferty, 2006). In brief, dynamic topic models establish a conditional distribution over the hyperparameters α and β , that govern the document's topic distribution and topic's word distribution, respectively. The distribution over each hyperparameter is conditioned on the prior in the previous time step, allowing changes to topic composition and assignment. This can be seen graphically in the plate model in Figure 3. For convenience, we provide a concise description of the generative process for the implemented heirarchical model for a time slice t in Appendix 9.5.

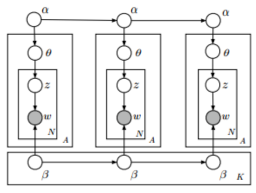


Figure 3. Plate diagram for the Dynamic Topic Model. Reproduced from (Blei & Lafferty, 2006).

Note that in contrast to LDA, dynamic topic models use a logistic-normal to express proportion uncertainty. This is stated more explicitly in Appendix 9.5.

4.2. Implementation

Unfortunately, due to the non-conjugacy of the Gaussian/Multinomial in our logistic-normal setup, integrating out parameters for any sort of Gibbs sampling becomes hard. Instead, the original DTM paper uses a variational approximation based on a Kalman filter, that preserves time dependencies, unlike a mean-field approximation.

Re-implementing the variational approximation, while educational and interesting, would be a project of itself, so our first attempt in employing dynamic time models to our gene expression data re-used the Kalman Filter-based variational inference used by the original authors, published recently by the Blei Lab (Blei & Gerrish, 2015). The results we show in the subsequent section are using a wrapper we've implemented around the group's inference code.

The authors stumbled upon a recent arXiv pre-print

that proposed a set of sampler update rules to create a correct DTM Gibbs sampler (Bhadury, 2016). Using the probabalistic programming framework Edward, and modifying the in-built sampler to follow these updates, we were able to obtain a working Gibbs-sampler for sample time-sliced data for a dynamic topic model defined in Edward (Tran et al., 2016). While we don't have time to conduct benchmarks or re-run our results right now, after verifying that results are consistent, the authors hope to submit a merge for this Edward sketch into the mainstream Edward examples library this coming summer.

4.3. Experiments

We subdivided our data into three separate time slices, and pruned each slice, retaining the top 20% of high-variance genes. We experimented using $k=15,\ 30,$ and 50 topics.

Similar to our analysis in Section 2, we used the hypergeometric test to evaluate interpretability.

Figure 10 in the Appendix shows gene module matches for which the *p*-value is less than 0.3, across the three time slices and the three topic count sizes. Interestingly, we notice a very different set of matched gene modules, with exception of the KLF1 module³. Like before, we notice that some of the identified topics are relevant to tumor cells: ST_TUMOR_NECROSIS_FACTOR_PATHWAY and WEST_ADRENOCORTICAL_TUMOR_MARKERS_UP to name the two most prominent topics.

We do not see that much variation across time-slices, demonstrating that our time-slice partitions are largely uniform. For example, for model k=50, we see that topic 20 consistently matches with MSigDB module PILON_KLF1_TARGETS_UP, suggesting that its gene composition is not varying.

Models with more topics tended to have more matches with higher significance. But again, the *p*-values in our tests do *not* factor for multiple hypothesis corrections, so at the moment we are only using them as relative measures of model quality.

After extracting the mode of learned parameters, we evaluated the log-likelihood of our held-out test set, also partitioned by time. This is shown in Figure 4. We find a leveling of log-likelihood after k=30.

With more time, we would ideally repeat the experiments to obtain confidence intervals on these experiments, and examine the gene-module/cell/time-slice

³This could very well be a result of our dataset pruning, which we did to keep inference runtimes manageable

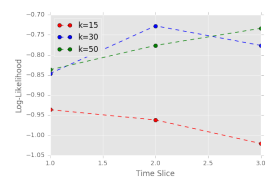


Figure 4. Log-likelihood of the held-out testing set across our time slices for varying numbers of topics.

mutual information.

5. Mixture Model

5.1. Model Description

The second question – whether we could learn cell groupings – we tackled using a canonical Gaussian mixture model. In brief, the model assigns every data point to a cluster; every cluster is Gaussian whose parameters are learned through posterior inference. A plate diagram can be found at (Ben Wing, 2016). This model is different from the previously mentioned Pe'er papers which use Dirichlet Processes Mixtures to study expression data. By plotting histograms of whitened gene expression values, we've found that for many genes the distribution is Gaussian across our data points. This hints that the generative process in a Gaussian mixture model is plausible.

5.2. Implementation

We implemented the mixture model in **Edward**, the probabalistic programming framework on top of Tensorflow. This allowed us to experiment with different samplers and variational inference approximations. The results we show below are for our runs using a Gibbs sampler, with a fixed burn-in of number of samples (200), a fixed number of sampling iterations (500).

5.3. Experiments

Figure 6 shows an estimated cluster assignment for one-hundred sampled data point. We see that while our learned assignments do capture some of the clustering, there is significant bias toward one group (the burgundy cluster).

Following this trend, we examined the set of unique clusters assigned in the posterior. The results are shown in Table 5.3. Interestingly, the model consis-

tently uses less clusters than is allocated in the model. We are unsure why this is the case.

Finally, we examine the log-likelihood of a held-out test set three different runs of posterior estimation with our sampler. This is shown in Figure 5. The runs are consistent with each other, and the likelihood is largely invariant across the five different cluster counts we tried, $k = \{10, 25, 50, 75, 50\}$. There are an estimated 9 true cell types in the original dataset, which could explain this result.

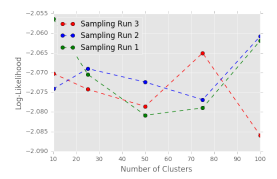


Figure 5. Log-likelihood of held-out test set, on three seperate runs of Gibbs sampling procedure, described above

# of Clusters	# of Clusters Used
10	3
25	9
50	16
75	18
100	20

Table 1. The number of clusters defined in the model appears to be more than the actual number of unique clusters assigned to the data points after posterior estimation. The estimated true number of categories is 9.

6. Integrated Topic-Clustering Model

6.1. Model Description

LDA is suitable for modeling corpuses with no special structure, but this is not the case of scRNA-seq data. Since the cells within a tissue sample may belong to many different sub-populations, we would expect cells within the same sub-population to have more similar topic proportions. We capture this additional structure using the clustering topic model (CTM), a unified framework for clustering and topic modeling based on LDA. In this model, which is similar to the one proposed by (Wallach, 2008), the following generative process is assumed:

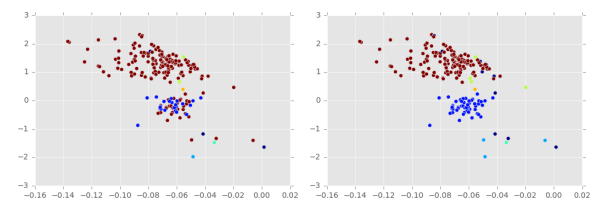


Figure 6. 2D-projection of sampled gene expression data points, colored by predicted cluster (left) and by true cluster assignments (right). Axis are the two learned PCA dimensions.

- 1. Choose a distribution over the clusters, $\pi \sim \text{Dir}(\pi_0/C)$, where π_0 is a hyperparameter and C is the number of clusters.
- 2. Choose a global distribution over the topics, $\phi_0 \sim \text{Dir}(\alpha_0/T)$, where α_0 is a hyperparameter and T is the number of topics.
- 3. For each cluster c, choose a cluster distribution over the topics, $\phi_c \sim \text{Dir}(\alpha_1 \phi_0)$, where α_1 is a hyperparameter
- 4. For each topic k, choose a distribution over the vocabulary, $\beta_k \sim \text{Dir}(\eta)$, where η is a hyperparameter.
- 5. For each document d,
 - (a) Choose a cluster assignment $\xi_d \sim \text{Cat}(\pi)$
 - (b) Choose a distribution over topics based on the cluster assignment, $\theta_d \sim \text{Dir}(\alpha \phi_{xi_d})$, where α is a hyperparameter.
 - (c) Choose the number of words in this document, $N_d \sim \text{Poisson}(\kappa)$, where κ is a hyperparameter.
 - (d) For the *i*th word in the *d*th document,
 - i. Choose a topic, $z_{di} \sim \text{Categorical}(\theta_d)$, based on the distribution over topics
 - ii. Choose a word, $w_{di} \sim \text{Categorical}(\beta_{z_{di}})$, based on the distribution over the vocabulary for topic z_{di}

This model is very similar to LDA, with the addition of cluster assignments that influence document topic proportions. Note that the documents in each cluster have more similar topic proportions, particularly if we choose a reasonably large value of α .

6.2. Inference

We implemented a collapsed Gibbs sampling in MATLAB to perform inference for the CTM, integrating out all latent variables except the topic assignments z and the cluster assignments ξ to facilitate mixing. First, we establish the prior distribution of z_{di} :

$$p(z_{di} = k | z_{-di}, \xi_d) = \frac{N_{d,k} + \alpha \frac{N_{\xi_d,k} + \alpha_1 \frac{N_k + \alpha_0 / K}{N + \alpha_0}}{N_{\xi_d}} + \alpha_1}{N_d + \alpha}$$
(1)

where z_{-di} refers to topic assignments other than that for word i in document d, $N_{d,k}$ is the number of words in document d assigned to topic k, N_d is the total number of words in document d, $N_{\xi_d,k}$ is the number of words in the documents from cluster ξ_d assigned to topic k, N_{ξ_d} is the total number of words in the documents from cluster ξ_d , N_k is the number of words in the corpus assigned to topic k, and N is the total number of words in the corpus k. It follows that the full conditional for each k is:

$$p(z_{di} = k|w_{di}, w_{-di}, z_{-di}, \xi_d)$$

$$\propto p(w_{di}|z_{di} = k, w_{-di})p(z_{di} = k|z_{-di}, \xi_d)$$

$$= \frac{N_{w_{di},k}}{N_{l\cdot}}p(z_{di} = k|z_{-di}, \xi_d)$$
(2)

where $N_{w_{di},k}$ is the number of times the word w_{di} appears in topic k, N_k is the total number of words assigned to topic k, and the prior of z_{di} is given by equation 1. Finally, the full conditional for each ξ_d is:

$$p(\xi_{d} = c | \xi_{-d}, z_{d}, z_{-d}; \pi)$$

$$\propto p(z_{d} | \xi_{d} = c, z_{-d}) p(\xi_{d} = c | \xi_{-d})$$

$$= Multi(z_{d}, p(z_{d} | z_{-d}, \xi_{d} = c)) \frac{M_{d,c} + \pi_{0}/C}{M_{d} + \pi_{0}}$$
(3)

⁴When counting the number of words, we always exclude the word currently being considered, but we omit this in the equation to avoid clutter.

where M_c is the number of documents assigned to topic c and M is the total number of documents in the corpus⁵. $Multi(z_d, p(z_d|\cdots))$ is the probability under the multinomial distribution of getting topic assignments z_d when the topic proportions are given by $p(z_d|\cdots)$. The topic proportions are obtained essentially as described in equation 1, except we set the number of words in document d to be 0 and adjust all the other counts accordingly.

6.3. Experiments

We trained the CTM on a subset of 500 cells and 3000 genes from the human melanoma scRNA-seq dataset using the collapsed Gibbs sampler until convergence (about 500 iterations). All hyperparameters were set to 1 with the exception of α , which we set to 10 to encourage similarity of topic proportions for cells assigned to the same cluster, and the numbers of clusters and topics were set to 10 and 20 respectively. To determine the efficacy of the clustering, we generated a tSNE plot of the cells using the Euclidean distance between their topic proportions (Figure 7). Compared to a tSNE plot based on the Euclidean distance between the counts of all 3000 genes (Figure 16), in which there was no obvious grouping of cells, we found that the topic proportions were nicely grouped based on their learned cluster assignments. To determine if our clusters had any biological significance, we compared them with cell labels published by the original authors (Figure 8)⁶. Interestingly, our clusters corresponded nearly exactly to the different cell types, and we were even able to classify the cells with unknown labels.

Subsequently, we wanted to determine if the learned topics were meaningful. We first used the biological relevance score described earlier and found that the topics were significantly enriched for some known gene sets from MSigDB (Figure 17). We then did some examination of the specific genes appearing in select topics using a combination of the MSigDB analysis and GO-enrichment analysis⁷. We found that topic 18, which

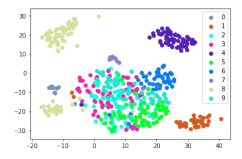


Figure 7. tSNE plot of cells from human melanoma, plotted using Euclidean distances between topic proportions, with cluster assignments shown in color. The cells nicely group into their cluster assignments, suggesting that the clustering topic model inference algorithm worked as intended. Moreover, the cluster assignments correpond to the cell labels found by the group that published the dataset, see Figure 8

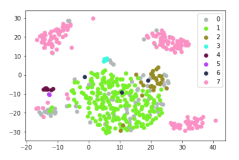


Figure 8. tSNE plot of cells from human melanoma, plotted using Euclidean distances between topic proportions, with labels published by the group that collected the dataset. 0 - label unknown; 1 - T-cell; 2 - B-cell; 3 - Macrophage; 4 - Endoderm; 5 - Cancer-associated fibroblast; 6 - Natural Killer; 7 - Malignant. The labels correpond to the cluster assignments learned using the CTM model.

⁵When counting the number of documents, we exclude the document currently being considered, but omit this in the equation to avoid clutter

⁶The authors published the cell labels based on prior biological knowledge, i.e. thresholding the cells based on known gene markers of different cell types. Due to noise and dropout in scRNA-seq data, they were not able to label all cells using this approach.

⁷GO enrichment analysis looks for enrichment of genes associated with cell processes. Whereas MSigDB is a database of various published gene sets, the GO database contains only those well-characterized gene modules that

was prevalent in the T-cells and not in other cells (Figure 19), was highly enriched for genes involved in "myeloid dendritic cell activation involved in immune response". Topic 3, which was prevalent in the B-cells and not elsewhere (Figure 20), was highly enriched for genes involved in "lymphocyte activation" and "inflammatory response". Topic 4, which was highly prevalent in the T-cells and B-cells only (Figures 19 and 20), was highly enriched for genes involved in "somatic diversification of immune receptors". Finally, topic 8 which was mainly present in the tumor cells (18), was enriched for genes involved in "G1 to S transition of mitotic cell cycle" and "cell division". The correspondence between the genes enriched in the topics and the cell sub-populations with those topics suggests that our learned topics have biological significance. As a result, the highly-ranked genes in some of these topics may be of interest to experimental biologists as candidate gene markers for specific cell subpopulations.

7. Discussion

In this paper, we sought to address the following two questions using Bayesian topic models: given gene expression data, (1) can we identify biologically meaningful gene modules, and (2) can we identify cell sub-populations and find gene modules that illuminate their differences.

7.1. Can we identify biologically meaningful gene modules?

We addressed the first question by trying to model scRNA-seq data with LDA as well as two nonparametric topic models, the HDP and the IBP-DP. The purpose of doing this was mainly to determine if topic models based on LDA are suitable for modeling gene expression; as far as we are aware, there are no publications applying LDA to scRNA-seq data. Our results from section 2 suggest that LDA could potentially be useful for scRNA-seq, since our topics were enriched for previously discovered biological gene sets. One major consideration is computational cost: although scRNA-seq data is sparse, it is not nearly as sparse as the bag-of-words representation of a corpus, so most inference algorithms take a long time to run on the entire dataset. Based on our results, however, it appears that this problem can be addressed either by using a faster posterior inference method such as online variational inference or by pre-processing the

are generally accepted as being involved in a particular cell process

dataset for those genes with the greatest variability.

Compared to LDA, we found that HDP and IBP-DP were not as suitable for modeling scRNA-seq data (Section 3). The HDP underperformed in comparison to LDA in both log-likelihood on a held-out dataset and the biological significance scores of its topics. The IBP-DP did not seem to work better than LDA on a subset of the Reuters-21578 corpus, based on a qualitative analysis of the topics, so we did not try it on the gene expression data, as we do not expect it to perform better than LDA. The underperformance of the IBP-DP compared to LDA on the Reuters-21578 dataset does not surprise us, as the authors of the IBP-DP paper did not show any topics from their model, even though they claimed it may produce more 'focused' topics.

After we were confident that LDA could be useful for modeling scRNA-seq data, we wanted to try models that could capture higher-level structure of gene expression data - for instance, time. As cells proliferate, they may differentiate into different cell types. A useful topic model might learn not just the topics at a given time point, but also how the topics are evolving over time. We have presented some preliminary results applying a dynamic topic model to our scRNA-seq dataset and selecting an appropriate number of topics (Section 4). To further test this model, experiments could be done on simulated time-series scRNA-seq data with known evolving topics, and one could evaluate how well those evolving topics are picked up by the model.

7.2. Can we identify cell sub-populations and find gene modules that illuminate their differences?

In addition to time, another source of structure in the scRNA-seq dataset is the clustering of cells into sub-populations. In biology, there is much interest in identifying these sub-populations and finding gene modules or markers that are characteristic of different sub-populations. We first wanted to determine if a Bayesian finite mixture model would be appropriate for separating the cells into clusters (Section 5). We found that we were approximately able to learn the cell type labels published by (Tirosh, 2016)

Finally, we wanted to create a unified framework for cell clustering and topic modeling. We implemented the clustering topic model (Section 6) and found that it was able to correctly separate the cells into subpopulations. It appeared to outperform the vanilla mixture model without clustering, suggesting that knowledge of the topics helps the model learn better clusters. We also found that several topics corresponded to the cell types that they were prevalent in; we believe these results suggest that biologists can use the topics - which are ranked lists of genes - as candidate gene markers for cell sub-populations of interest within the melanoma, such as malignant cells, T-cells and B-cells. Whether these candidate gene markers are truly useful must be confirmed by biologists experimentally. Looking forward, it would be very interesting to combine the dynamic topic model and the clustering topic model, so that we can capture the time-evolution of the cell sub-populations within a scRNA-seq dataset.

8. Author Contributions

The implementations and experiments involving LDA, Gaussian Mixtures, and the Dynamic Topic Model was completed by SK (Section 2, Section 4, and Section 5). KY completed the implementations and experiments involved non-parametric topic models, the clustering topic model, and the gene set enrichment test implementation (Section 3, Section 6, and Section 9.1). The authors contributed equally in this work.

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9. Appendix

9.1. Biological Significance Score / Gene Set Enrichment

In general, it is difficult to evaluate the 'coherence' of topics of genes, since unlike topics of words, we cannot just read them and get a sense of how similar they are. Therefore, nothing like the intrusion test exists for evaluating topics of genes. To overcome this, we developed our own test for the biological significance of gene topics. The MSigDB contains thousands of gene sets discovered through previous research, catalogued by biologists from the Broad Institute. We deduced that if a topic is 'coherent', then it is likely significantly enriched for some of these gene sets. For each topic, our biological significance scoring method iterates through the MSigDB database and tests each gene set for enrichment using the minimum hyper-geometric test (Wagner, 2015). The most significant p-values for each topic are recorded and used as a proxy for its biological coherence.

9.2. Latent Dirichlet Allocation

For our Gibb's sampler, we had a fixed burn-in of number of samples (200), a fixed number of sampling iterations after that (500). We didn't extensively explore varying these values, but trying out significantly more iterations (700) didn't seem to change the topic's word distributions significantly. There was one sampling chain on each of four cores.

9.3. HDP

HDP is an infinite topic extension of LDA based on the Dirichlet process (Blei, 2005). Briefly, words in a corpus are assumed to be generated as follows:

- 1. Sample the global distribution over topics, $G_0 \sim \mathrm{DP}(\gamma, H)$, from a Dirichlet process with concentration γ and base Dirichlet distribution H^8 .
- 2. For each document d,
 - (a) Sample the local distribution over topics, $G_d \sim \mathrm{DP}(\alpha_0, G_0)$, from a Dirichlet process with concentration α_0 and base distribution G_0 .
 - (b) Choose the number of words in this document, $N_d \sim \text{Poisson}(\xi)$, where ξ is a hyperparameter.
 - (c) For the *i*th word in the *d*th document,
 - i. Choose a topic (i.e. distribution over words in vocab), $\beta_{di} \sim G_d$, based on the local distribution over topics
 - ii. Choose a word, $w_{di} \sim \text{Categorical}(\beta_{di})$, based on the distribution over the vocabulary in the topic

9.4. IBP-DP

One drawback to the HDP is that there tends to be a correlation between how frequently a topic appears across all documents and how prevalent this topic is within documents that it appears in. Williamson et al. (Blei, 2010) proposed the 'focused topic model' to overcome this drawback. In their model, the frequency of a topic k depends on two separate variables, its relative prevalence within a document (ϕ_k) and the probability that a given document contains this topic (π_k) , thus reducing correlation between the two. Briefly, the words in the corpus are assumed to be generated as follows:

⁸More specifically, H is a Dirichlet distribution over the (V-1)-dimensional simplex, where V is the size of the vocabulary, and G_0 is a countably infinite set of point masses over this simplex whose weights sum to 1. G_0 in HDP plays a role analogous to α in LDA.

- 1. For each topic k = 1, 2, ...
 - (a) Choose a population frequency (i.e. probability that a document contains topic) $\pi_k = \prod_{j=1}^k \mu_k$, where each $\mu_k \sim Beta(\alpha, 1)$, where α is a hyperparameter.
 - (b) Choose a relative prevalence (i.e. how often the topic appears within a document containing the topic) $\phi_k \sim Gamma(\gamma, 1)$, where γ is a hyperparameter.
 - (c) Choose a distribution over the vocabulary, $\beta_k \sim \text{Dir}(\eta)$, where η is a hyperparameter.
- 2. For each document d,
 - (a) Choose the topics that will appear as a binary vector, $b_d \sim Bernoulli(\pi)$
 - (b) Choose the topic proportion as $\theta_d \sim Dirichlet(b_d \cdot \phi)$
 - (c) Choose the number of words in this document, $N_d \sim \text{NegativeBin}(\sum_k b_{dk} phi_k, \delta)$, where δ is a hyperparameter.
 - (d) For the *i*th word in the *d*th document,
 - i. Choose a topic, $z_{di} \sim \text{Categorical}(\theta_d)$, based on the distribution over topics
 - ii. Choose a word, $w_{di} \sim \text{Categorical}(\beta_{z_{di}})$, based on the distribution over the vocabulary for topic z_{di}

9.5. Dynamic Topic Model

Here is a description of the generative process for our dynamic topic model:

- 1. Draw a new topic composition hyperparameter $\beta \leftarrow \mathbb{N}(\beta_{t-1}, \sigma I^2)$
- 2. Draw a new document composition hyperparameter $\alpha \leftarrow \mathbb{N}(\alpha_{t-1}, \delta I^2)$
- 3. For each document:
 - (a) Draw a new document topic distribution $\theta \leftarrow \pi(\mathbb{N}(\alpha_t))$
 - (b) For every word in the document:
 - i. Draw the word topic assignment $Z \leftarrow \text{Mult}(\theta)$
 - ii. Draw the word identity $Z \leftarrow \text{Mult}(\pi(\beta_{t,z}))$

Here, $\pi(x_i)$ is the softmax function $\frac{exp(x_i)}{\sum_k exp(x_k)}$ (Blei & Lafferty, 2006).

```
GIBBS TOPICS
        cluster size: 5
        cluster size: 10
                match: topic 1 , KINSEY_TARGETS_OF_EWSR1_FLII_FUSION_UP , pval: 0.041958041958
                match: topic 1 , PILON_KLF1_TARGETS_DN , pval: 0.137062937063
                match: topic 6 , GRAESSMANN_APOPTOSIS_BY_DOXORUBICIN_DN , pval: 0.263736263736
        cluster size: 25
                match: topic 8 , PILON KLF1 TARGETS DN , pval: 0.263736263736
                match: topic 22 , KINSEY_TARGETS_OF_EWSR1_FLII_FUSION_UP , pval: 0.041958041958
                match: topic 22 ,
                                    PILON KLF1 TARGETS DN , pval: 0.137062937063
                match: topic 22 , GRAESSMANN APOPTOSIS BY DOXORUBICIN DN , pval: 0.263736263736
        cluster size: 50
                match: topic 2 , ROME_INSULIN_TARGETS_IN_MUSCLE_UP , pval: 0.193006993007
                match: topic 3 , PILON_KLF1_TARGETS_DN , pval: 0.00699300699301
                match: topic 6 , PILON_KLF1_TARGETS_DN , pval: 0.263736263736
                match: topic 7 , BLALOCK_ALZHEIMERS_DISEASE_UP , pval: 0.018648018648 match: topic 17 , KINSEY_TARGETS_OF_EWSR1_FLII_FUSION_UP , pval: 0.041958041958
                match: topic 17 , PILON_KLF1_TARGETS_DN , pval: 0.193006993007
                match: topic 22 , PILON_KLF1_TARGETS_DN , pval: 0.153846153846
                match: topic 24 , WEI MYCN TARGETS WITH E BOX , pval: 0.0839160839161
OVB TOPICS
        cluster size: 5
        cluster size: 10
                match: topic 5 , PILON KLF1 TARGETS DN , pval: 0.263736263736
        cluster_size: 25
                match: topic 5 , DIAZ_CHRONIC_MEYLOGENOUS_LEUKEMIA_UP , pval: 0.263736263736
                match: topic 8 , KINSEY_TARGETS_OF_EWSR1_FLII_FUSION_UP , pval: 0.041958041958
                match: topic 15 , PUJANA_ATM_PCC_NETWORK , pval: 0.263736263736
                match: topic 15 ,
                                    PUJANA BRCA1 PCC NETWORK , pval: 0.263736263736
        cluster_size: 50
                match: topic 6 , GRAESSMANN APOPTOSIS BY DOXORUBICIN DN , pval: 0.263736263736
                match: topic 15 , PUJANA_ATM_PCC_NETWORK , pval: 0.263736263736
                                    MARSON BOUND BY FOXP3 UNSTIMULATED , pval: 0.263736263736
                match: topic 22 ,
                match: topic 28 ,
                                    PILON KLF1 TARGETS_DN , pval: 0.201398601399
                match: topic 38 ,
                                    PUJANA BRCA1 PCC NETWORK , pval: 0.263736263736
                match: topic 41,
                                    MARSON_BOUND_BY_FOXP3_UNSTIMULATED , pval: 0.263736263736
                match: topic 42,
                                    KINSEY TARGETS OF EWSR1 FLII FUSION UP , pval: 0.193006993007
```

Figure 9. Matches between the gene collections found in LDA topics and published gene sets in MSigDB. 'Cluster size' refers to the number of topics in the model

```
k=15
        t=1
                 match: topic 10 , PILON_KLF1_TARGETS_UP , pval: 0.05484091166
                 match: topic 11 , ST_TUMOR_NECROSIS_FACTOR_PATHWAY , pval: 0.13159861000
        t=2
        t=3
                 match: topic 2 , ST_INTEGRIN_SIGNALING_PATHWAY , pval: 0.22324501174
                 match: topic 10 , PILON_KLF1_TARGETS_UP , pval: 0.25017876663
k=30
        t=1
                 match: topic 26 , WEST ADRENOCORTICAL TUMOR MARKERS UP , pval: 0.04716140093
                 match: topic 27 , ST_TUMOR_NECROSIS_FACTOR_PATHWAY , pval: 0.15884551563
        t=2
                 match: topic 12 , KEGG_PARKINSONS_DISEASE , pval: 0.19929937855
                 match: topic 26 , WEST_ADRENOCORTICAL_TUMOR_MARKERS_UP , pval: 0.08559859059
        t=3
                 match: topic 11 , TIMOFEEVA GROWTH STRESS VIA STAT1 DN , pval: 0.08405900408
                 match: topic 27 , ST_TUMOR_NECROSIS_FACTOR_PATHWAY , pval: 0.09390276030
k=50
        t=1
                 match: topic 13 , WEST_ADRENOCORTICAL_TUMOR_MARKERS_UP , pval: 0.00922042410
                 match: topic 20 , HOLLMANN_APOPTOSIS_VIA_CD40_DN , pval: 0.05797811105
                match: topic 20 , PILON_KLF1_TARGETS_UP , pval: 0.05483747800 match: topic 20 , WEST_ADRENOCORTICAL_TUMOR_MARKERS_UP , pval: 0.02003797479
        t=2
                 match: topic 15 , ST_P38_MAPK_PATHWAY , pval: 0.19462717810
        t=3
                 match: topic 7 , ST_TUMOR_NECROSIS_FACTOR_PATHWAY , pval: 0.20037158939
                 match: topic 13 , WEST_ADRENOCORTICAL_TUMOR_MARKERS_UP , pval: 0.05761453734
                 match: topic 20 , PILON_KLF1_TARGETS_UP , pval: 0.09429082710
```

Figure 10. Matches between the gene collections found in DTM topics and published gene sets in MSigDB. k refers to the number of topics in the model, t refers to one of three time slices.

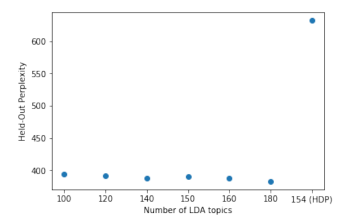
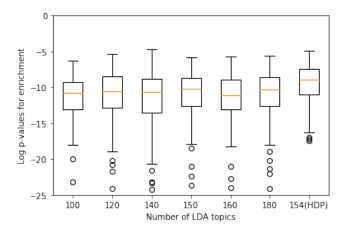


Figure 11. Comparison of log-likelihood of the held-out testing set, under various LDA models and the HDP model.



Figure~12. Comparison of distributions of p-values from gene set enrichment analysis between LDA models and the HDP model.

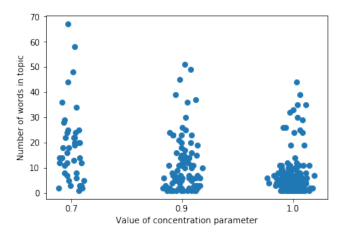


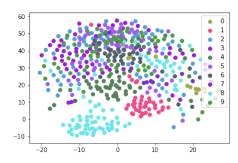
Figure 13. Beeswarm plot of number of words per topic, for 3 different IBP-DP models with different concentration parameters. Each point represents one topic from its model

```
Topic 0: charles diana prince royal parker bowles camilla queen family marriage public princess love britain england
Topic 1: church told during year time world very years made life saying last became take while
Topic 2: catholic n't bishop son women father love mother television day cardinal britain woman leaders days
Topic 3: sunday including against day won group held known last police official roman few come late
Topic 4: pope france visit john french paul first both pontiff trip church religious catholics king against
Topic 5: mother teresa doctors home heart charity hospital order tuesday people work told peace world house
Topic 6: set years ceremony led german germany second made called around rights married time few south
Topic 7: pope health vatican mass trip reporters surgery during saturday death left past spokesman people monday
Topic 8: people president state last later took around government good since say percent age won head Topic 9: media later catholic former own head newspaper next known leader against wednesday taken n't called
Topic 10: official added first health ago under among around wednesday paul few minister monday children mother
Topic 11: former end throne century taken reports children newspaper england home french paris 1992 international state
Topic 12: french members visit leaders family germany national statement say three work against thursday year
Topic 13: four heart clinton percent age
Topic 14: life say wednesday left while month tuesday times later official take peace paris doctors including
Topic 15: says 1992 princess love left government political under german married took since saturday house around
Topic 16: bernardin cardinal among death u.s own surgery told minister doctors great several news end until
Topic 17: reports roman us told president times
Topic 18: paul white show reports end son union month monday bishop long
Topic 19:
             service end million since wednesday roman home wife spokesman son reports city
Topic 20: first local place princess british peace white saturday taken expected several united married made century Topic 21: part say sunday members year leaders church days
Topic 22: former ago n't three century rome funeral group year saying led family
Topic 23: france home news south whose president work east take first both united women country officials
Topic 24: four week throne saying became former died members camilla made long show
Topic 25: statement french week thursday war christian born vatican house heart leader britain 1992 set three
Topic 26: minister prime expected group officials union died children times michael whose off around american church
Topic 27: years president
Topic 28: diana funeral service princess reuters hospital son died
Topic 29: monday u.s around several four children throne year since john tuesday members churchill told statement Topic 30: day friday party private wednesday later british officials former family until throne white
Topic 31: against
Topic 32: city international good prize second won paris take since died house off local years
Topic 33: times children world
Topic 34: paris work
Topic 35: michael king paul local father political show ceremony next part private war german week whose
Topic 36: rights government prize service million become officials held police head reuters political us party
Topic 37: television reporters show n't political times clinton several off own government son wednesday very during
Topic 38: known news
Topic 39: part great mass later women past says
Topic 40: told charles called off born
Topic 41: taken ceremony president
Topic 42: thursday long leaders
```

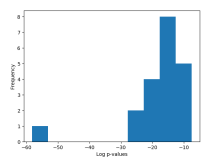
Figure 14. Topics from IBP-DP model trained on subset of Reuters dataset.

```
Topic 0 : france home news years work mother president during take whose women east country love told
                church years cardinal bishop take england against million vatican past news british told sunday ceremony pope health mass during visit saturday trip told john paul pontiff people church service spokesman mother teresa heart sunday home hospital tuesday told doctors order people catholic charity peace house
Topic 1
Topic 2
Topic 3
Topic 4
                 pope france french visit church trip first paul catholic pontiff both john state including paris
                 teresa mother doctors charity official hospital home work first around told world during under saying
Topic 5
                 church media michael paul former marriage princess england never n't love very told public years bishop church catholic son father n't told women love mother woman roman years leaders ago royal family queen prince charles throne church princess century britain first british media 1992 head
Topic 6
Topic 7
Topic 8
                 order day city friday group during own doctors monday very reuters prize last people roman
Topic 9
              television told show n't reporters president later day own times political clinton off year years:
president rights government people last church says state life died told political country group catholic:
diana charles princess britain time wednesday ago family monday million camilla church newspaper bowles parker
charles parker bowles prince camilla diana royal marriage public queen love king church woman family
Topic 10
Topic 11
Topic 12
Topic 13:
Topic 14 : pope bernardin vatican church surgery health year time left told life say death cardinal made
```

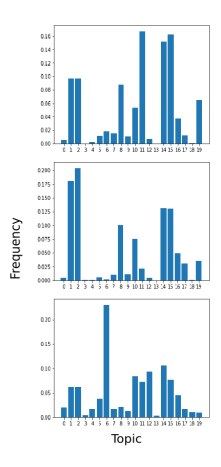
Figure 15. Topics from LDA model with 15 topics trained on subset of Reuters dataset.



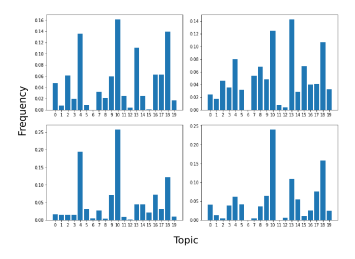
 $Figure~16.~{\rm tSNE~plot~of~cells~from~human~melanoma,~plotted~using~Euclidean~distances~between~gene~count~vectors,~with~cluster~assignments~shown~in~color.}$



 $Figure~17.~{\rm Histogram~of~topics~from~CTM~model~based~on~p-value~of~enrichment~of~MSigDB~gene~sets.}$



 $Figure~18.~{
m Bar~charts}$ of topic proportions for malignant cell clusters.



 $Figure~19.~{
m Bar~charts}$ of topic proportions for T-cell clusters.

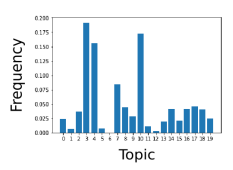


Figure 20. Bar charts of topic proportions for B-cell clusters