

Visualizing structure in RNA-seq Expression Data using Grade of Membership Models

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Abstract

Grade of membership models, also known as "admixture models", "topic models" or "Latent Dirichlet Allocation", are a generalization of cluster models that allow each sample to have membership in multiple clusters. These models are widely used in population genetics to model admixed individuals who have ancestry from multiple "populations", and in natural language processing to model documents having words from multiple "topics". Here we illustrate the potential for these models to cluster samples of RNA-seq gene expression data, measured on either bulk samples or single cells. We also provide methods to help interpret the clusters, by identifying genes that are distinctively expressed in each cluster. By applying these methods to several example RNA-seq applications we demonstrate their utility in identifying and summarizing structure and heterogeneity. Applied to data from the GTEx project on 51 human tissues, the approach highlights similarities among biologically-related tissues and identifies distinctively-expressed genes that recapitulate known biology. Applied to single-cell expression data from mouse preimplantation embryos, the approach highlights both discrete and continuous variation through early embryonic development stages, and highlights genes involved in a variety of relevant processes - from germ cell development, through compaction and morula formation, to the formation of inner cell mass and trophoblast at the blastocyte stage. The methods are implemented in the Bioconductor package CountClust.

Author Summary

The gene expression profile of a biological sample (either tissue or single cells) results from a complex interplay between many biological processes. Consequently, for example, distal tissue samples may be similar in their gene expression profiles because they share biological processes. Our goal here is to illustrate that grade of membership (GoM) models – an approach widely used in population genetics to cluster admixed individuals who have ancestry from multiple populations – provide an attractive approach for clustering biological samples of RNA sequencing data. The GoM model allows each biological sample to have partial memberships in multiple biological-distinct clusters, in contrast to traditional clustering methods that partition samples into distinct subgroups. We also provide methods for identifying genes that are distinctively expressed in each cluster to help biologically interpret the results. Applied to a dataset of 51 human tissues, the GoM approach highlights similarities among biologically-related tissues and identifies distinctively-expressed genes that recapitulate known biology. Applied to gene

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expression data of single cells from mouse preimplantation embryos, the approach highlights both discrete and continuous variation through early embryonic development stages, and genes involved in a variety of relevant processes. Our study highlights the potential of GoM models for elucidating biological structure in RNA-seq gene expression data.

Introduction 19

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Ever since large-scale gene expression measurements have been possible, clustering – of both genes and samples – has played a major role in their analysis [3–5]. For example, clustering of genes can identify genes that are working together or are co-regulated, and clustering of samples is useful for quality control as well as identifying biologically-distinct subgroups. A wide range of clustering methods have therefore been employed in this context, including distance-based hierarchical clustering, k-means clustering, and self-organizing maps (SOMs); see for example [6,7] for reviews.

Here we focus on cluster analysis of samples, rather than clustering of genes (although our methods do highlight sets of genes that distinguish each cluster). Traditional clustering methods for this problem attempt to partition samples into distinct groups that show "similar" expression patterns. While partitioning samples in this way has intuitive appeal, it seems likely that the structure of a typical gene expression data set will be too complex to be fully captured by such a partitioning. Motivated by this, here we analyse expression data using grade of membership (GoM) models [8], which generalize clustering models to allow each sample to have partial membership in multiple clusters. That is, they allow that each sample has a proportion, or "grade" of membership in each cluster. Such models are widely used in population genetics to model admixture, where individuals can have ancestry from multiple populations [16], and in document clustering [34,35] where each document can have membership in multiple topics. In these fields GoM models are often known as "admixture models", and "topic models" or "Latent Dirichlet Allocation" [34]. GoM models have also recently been applied to detect mutation signatures in cancer samples [33].

Although we are not the first to apply GoM-like models to gene expression data, previous applications have been primarily motivated by a specific goal, "cell type deconvolution", which involves using cell-type-specific expression profiles of marker genes to estimate the proportions of different cell types in a mixture [40, 42, 43]. Specifically, the GoM model we use here is analogous to – although different in detail from – blind deconvolution approaches [38, 39, 41] which estimate cell type proportions and cell type signatures jointly (see also [36, 37] for semi-supervised approaches). Also, the GoM model is similar in spirit to the Non-negative Matrix Factorization (NMF) approach, although differs in choice of prior and has been shown to produce more coherent topics than NMF [9] [10].

Our goal here is to demonstrate that GoM models can be useful much more broadly for understanding structure in RNA-seq data – not only to deconvolve mixtures of cell types. For example, in our analysis of human tissue samples from the GTEx project below, the GoM model usefully captures biological heterogeneity among samples even though the inferred grades of membership are unlikely to correspond precisely to proportions of specific cell types. And in our analyses of single-cell expression data the GoM model highlights interesting structure, even though interpreting the grades of membership as "proportions of cell types" is clearly inappropriate because each sample is a single cell! Here we are exploiting the GoM as a flexible extension of traditional cluster models, which can capture "continuous" variation among cells as well as the more "discrete" variation captured by cluster models. Indeed, the extent to which

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variation among cells can be described in terms of discrete clusters versus more continuous populations is a fundamental question that, when combined with appropriate single-cell RNA-seq data, the GoM models used here may ultimately help address.

Methods Overview

We assume that the RNA-seq data on N samples has been summarized by a table of counts $C_{N\times G}=(c_{ng})$, where c_{ng} is the number of reads from sample n mapped to gene g (or other unit, such as transcript or exon) [14]. The GoM model is a generalization of a cluster model, which allows that each sample has some proportion ("grade") of membership, in each cluster. For RNA-seq data this corresponds to assuming that each sample n has some proportion of its reads, q_{nk} coming from cluster k. In addition, each cluster k is characterized by a probability vector, θ_{k} , whose gth element represents the relative expression of gene g in cluster k. The GoM model is then

$$(c_{n1}, c_{n2}, \cdots, c_{nG}) \sim \text{Multinomial}(c_{n+}, p_{n1}, p_{n2}, \cdots, p_{nG}),$$
 (1)

where

$$p_{ng} := \sum_{k=1}^{K} q_{nk} \theta_{kg}. \tag{2}$$

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Here c_{n+} is the library size, or number of reads mapped to the genome, for sample n. The number of clusters K is set by the analyst, and it can be helpful to explore multiple values of K (see Discussion).

To fit this model to RNA-seq data, we exploit the fact that exactly the same GoM model is commonly used for document clustering [34]. This is because, just as RNA-seq samples can be summarized by counts of reads mapping to each possible gene in the genome, document data can be summarized by counts of each possible word in a dictionary. Recognizing this allows existing methods and software for document clustering to be applied directly to RNA-seq data. Here we use the R package maptpx [15] to fit the GoM model.

Fitting the GoM model results in estimated membership proportions q for each sample, and estimated expression values θ for each cluster. We visualize the membership proportions for each sample using a "Structure plot" [17], which is named for its widespread use in visualizing the results of the *Structure* software [16] in population genetics. The Structure plot represents the estimated membership proportions of each sample as a stacked barchart, with bars of different colors representing different clusters. Consequently, samples that have similar membership proportions have similar amounts of each color. See Fig 1 for example.

To help biologically interpret the clusters inferred by the GoM model we also implemented methods to identify, for each cluster, which genes are most distinctively differentially expressed in that cluster (see Methods). Functions for fitting the GoM model, plotting the structure plots, and identifying the distinctive ("driving") genes in each cluster, are included in our R package CountClust [46] available through Bioconductor [31].

Results

Clustering human tissue samples using bulk RNA-seq

We begin by illustrating the GoM model on bulk RNA expression measurements from the GTEx project (V6 dbGaP accession phs000424.v6.p1, release date: Oct 19, 2015,

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http://www.gtexportal.org/home/). These data consist of per-gene read counts from RNA-seq performed on 8,555 samples collected from 450 human donors across 51 tissues, lymphoblastoid cell lines, and transformed fibroblast cell-lines. We analyzed 16,069 genes that satisfied filters (e.g. exceeding certain minimum expression levels) that were used during eQTL analyses by the GTEx project (gene list available in http://stephenslab.github.io/count-clustering/project/utilities/gene_names_all_gtex.txt).

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We fit the GoM model to these data, with number of clusters K=5,10,15,20. For each K we ran the fitting algorithm three times and kept the result with the highest log-likelihood. Fig 1(a) shows the Structure plot for K=20, with results for other K in S1 Fig. (See also http:

//stephenslab.github.io/count-clustering/project/src/tissues_tSNE_2.html for an alternative visualization using a 2-dimensional projection with t-SNE [22, 23].)

In all cases results reflect the known division of samples into tissues: that is, samples from the same tissue tend to have similar cluster membership proportions. As might be expected, increasing K highlights finer structure in the data, with tissues that cluster together with smaller K being subdivided into distinct subgroups for larger K. For brevity we focus on results for K=20 (Fig 1(a)). Here some tissues are represented by essentially a single cluster (e.g. Pancreas, Liver), whereas other tissues are represented as a mixture of multiple clusters (e.g. Thyroid, Spleen). Furthermore, the results highlight biological similarity among some tissues by assigning similar membership proportions to samples from those tissues. For example, samples from several different parts of the brain often have similar memberships, as do the arteries (aorta, tibial and coronary) and skin samples (sun-exposed and un-exposed).

To help biologically interpret results we implemented methods to identify the genes and genetic processes that characterize each cluster (see Methods). Table 1 summarizes results for the GTEx results in Fig 1a (see also S1 Table). Reassuringly, many results align with known biology. For example, the purple cluster (cluster 18), which distinguishes Pancreas from other tissues, is enriched for genes responsible for digestion and proteolysis, (e.g. PRSS1, CPA1, PNLIP). Similarly the yellow cluster (cluster 12), which primarily distinguishes Cell EBV Lymphocytes from other tissues, is enriched with genes responsible for immune responses (e.g. IGHM, IGHG1) and the pink cluster (cluster 19) which mainly shows up in Whole Blood, is enriched with genes related hemoglobin complex and oxygen transport (e.g. HBB, HBA1, HBA2). Further, Keratin-related genes characterize the skin cluster (cluster 6, light denim), Myosin-related genes characterize the muscle skeletal cluster (cluster 7, orange), etc. The royal purple cluster (cluster 1) has memberships in most tissues and the genes distinguishing the cluster seem to be responsible for nucleus and nucleoplasm related functionality. In cases where a cluster occurs in multiple tissues these biological annotations may be particularly helpful for understanding what is driving this co-membership. For example, the top genes in the red cluster (cluster 3), which is common to Breast Mammary tissue, Adipose Subcutaneous and Adipose Visceral, are related to adipocytes and/or fatty acid synthesis; and the top genes in the salmon cluster (cluster 4), which is common to the Gastroesophageal Junction, Esophagus Muscularis and Colon Sigmoid, are related to smooth muscle.

Although global analysis of all tissues is useful for highlighting major structure in the data, it may miss finer-scale structure within tissues or among similar tissues. For example, here the global analysis allocated only three clusters to all brain tissues (clusters 1,2 and 9 in Fig 1(a)), and we suspected that additional substructure might be uncovered by analyzing the brain samples separately with larger K. Fig 1(b) shows the Structure plot for K=6 on only the Brain samples. The results highlight much finer-scale structure compared with the global analysis. Brain Cerebellum and

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Cerebellar hemisphere are essentially assigned to a separate cluster (lime green), which is enriched with genes related to cell periphery and communication (e.g. PKD1, CBLN3) as well as genes expressed largely in neuronal cells and playing a role in neuron differentiation (e.g. CHGB). The spinal cord samples also show consistently strong membership in a single cluster (yellow-orange), the top defining gene for the cluster being MBP which is involved in myelination of nerves in the nervous system [44]. Another driving gene, GFAP, participates in system development by acting as a marker to distinguish astrocytes during development [2].

The remaining samples all show membership in multiple clusters. Samples from the putamen, caudate and nucleus accumbens show similar profiles, and are distinguished by strong membership in a cluster (cluster 4, bright red) whose top driving gene is PPP1R1B, a target for dopamine. And cortex samples are distinguished from others by stronger membership in a cluster (cluster 2, turquoise in Fig 1(b)) whose distinctive genes include ENC1, which interacts with actin and contributes to the organisation of the cytoskeleton during the specification of neural fate [1].

Quantitative comparison with hierarchical clustering

We used the GTEx data to test whether the GoM model is more accurate in detecting substructure than distance-based clustering methods. Specifically, for each pair of tissues in the GTEx data we assessed whether or not each clustering method correctly partitioned samples into the two tissue groups (see Methods). The GoM model was substantially more accurate in this test, succeeding in 81% of comparisons, compared with 29% for the distance-based method (Fig 2). This presumably reflects the general tendency for model-based approaches to be more efficient that distance-based approaches, provided that the model is sufficiently accurate.

Clustering of single-cell RNA-seq data

Recently RNA-sequencing has become viable for single cells [11], and this technology has the promise to revolutionize understanding of intra-cellular variation in expression, and regulation more generally [12]. Although it is traditional to describe and categorize cells in terms of distinct cell-types, the actual architecture of cell heterogeneity may be more complex, and in some cases perhaps better captured by the more "continuous" GoM model. In this section we illustrate the potential for the GoM model to be applied to single cell data.

To be applicable to single-cell RNA-seq data, methods must be able to deal with lower sequencing depth than in bulk RNA experiments: single-cell RNA-seq data typically involve substantially lower effective sequencing depth compared with bulk experiments, due to the relatively small number of molecules available to sequence in a single cell. Therefore, as a first step towards demonstrating its potential for single cell analysis, we checked robustness of the GoM model to sequencing depth. Specifically, we repeated the analyses above after thinning the GTEx data by a factor of 100 and 10,000 to mimic the lower sequencing depth of a typical single cell experiment. For the thinned GTEx data the Structure plot for K=20 preserves most of the major features of the original analysis on unthinned data (S2 Fig). For the accuracy comparisons with distance-based methods, both methods suffer reduced accuracy in thinned data, but the GoM model remains superior (S3 Fig). For example, when thinning by a factor of 10,000, the success rate in separating pairs of tissues is 0.32 for the GoM model vs. 0.10 for hierarchical clustering.

Having established its robustness to sequencing depth, we now illustrate the GoM model on two single cell RNA-seq datasets, from Jaitin *et al* [24] and Deng *et al* [25].

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Jaitin et al, 2014

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Jaitin et al sequenced over 4,000 single cells from mouse spleen. Here we analyze 1,041 of these cells that were categorized as CD11c+ in the sorting markers column of their data (http://compgenomics.weizmann.ac.il/tanay/?page_id=519), and which had total number of reads mapping to non-ERCC genes greater than 600. We believe these cells correspond roughly to the 1,040 cells in their Fig S7. Our hope was that applying our method to these data would identify, and perhaps refine, the cluster structure evident in [24] (their Fig 2A and 2B). However, our method yielded rather different results (Fig 3), where most cells were assigned to have membership in several clusters. Further, the cluster membership vectors showed systematic differences among amplification batches (which in these data is also strongly correlated with sequencing batch). For example, cells in batch 1 are characterized by strong membership in the orange cluster (cluster 5) while those in batch 4 are characterized by strong membership in both the blue and yellow clusters (2 and 6). Some adjacent batches show similar patterns - for example batches 28 and 29 have a similar visual "palette", as do batches 32-45. And, more generally, these later batches are collectively more similar to one another than they are to the earlier batches (0-4).

The fact that batch effects are detectable in these data is not particularly surprising: there is a growing recognition of the importance of batch effects in high-throughput data generally [28] and in single cell data specifically [29]. And indeed, both clustering methods and the GoM model can be viewed as dimension reduction methods, and such methods can be helpful in controlling for batch effects [26,27]. However, why these batch effects are not evident in Fig 2A and 2B of [24] is unclear.

Deng et al, 2014

Deng et al collected single-cell expression data of mouse preimplantation embryos from the zygote to blastocyst stage [25], with cells from four different embryos sequenced at each stage. The original analysis [25] focusses on trends of allele-specific expression in early embryo development. Here we use the GoM model to assess the primary structure in these data without regard to allele-specific effects (i.e. combining counts of the two alleles). Visual inspection of the Principal Components Analysis in [25] suggested perhaps 6-7 clusters, and we focus here on results with K=6.

The results from the GoM model (Fig 4) clearly highlight changes in expression profiles that occur through early embryonic development stages, and enrichment analysis of the driving genes in each cluster (Table 3, S3 Table) indicate that many of these expression changes reflect important biological processes during embryonic preimplantation development.

In more detail: Initially, at the zygote and early 2-cell stages, the embryos are represented by a single cluster (blue in Fig 4) that is enriched with genes responsible for germ cell development (e.g., Bcl2l10 [54], Spin1 [55]). Moving through subsequent stages the grades of membership evolve to a mixture of blue and magenta clusters (mid 2-cell), a mixture of magenta and yellow clusters (late 2-cell) and a mixture of yellow and green (4-cell stage). The green cluster then becomes more prominent in the 8-cell and 16-cell stages, before dropping substantially in the early and mid-blastocyst stages. That is, we see a progression in the importance of different clusters through these stages, from the blue cluster, moving through magenta and yellow to green. By examining the genes distinguishing each cluster we see that this progression reflects the changing relative importance of several fundamental biological processes. The magenta cluster is driven by genes responsible for the beginning of transcription of zygotic genes (e.g., Zscan4c-f show up in the list of top 100 driving genes: see https://stephenslab.github.io/count-clustering/project/src/deng_cluster_annotations.html),

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which takes place in the late 2-cell stage of early mouse embryonic development [57]. The yellow cluster is enriched for genes responsible for heterochromation Smarcc1 [58] and chromosome stability Cenpe [59] (see S3 Table). And the green cluster is enriched for cytoskeletal genes (e.g., Fbxo15) and cytoplasm genes (e.g., Tceb1, Hsp90ab1), all of which are essential for compaction at the 8-cell stage and morula formation at the 16-cell stage.

Finally, during the blastocyst stages two new clusters (purple and orange in Fig 4) dominate. The orange cluster is enriched with genes involved in the formation of outer trophoblast cells (e.g., Tspan8, Krt8, Id2 [52]), while the purple cluster is enriched with genes responsible for the formation of inner cell mass (e.g., Pdgfra, Pyy [53]). Thus these two clusters are consistent with the two cell lineages, the trophectoderm and the primitive endoderm, that make up the majority of the cells of the blastocyst [56]. Interestingly, however, the cells do not appear to fall into two distinct and clearly-separated populations – at least, not in terms of their expression patterns – but rather show a continuous range of memberships in these two clusters, even in the late blastocyst stage.

In addition to these trends across development stages, the GoM results also highlight some embryo-level effects in the early stages (Fig 4). Specifically, cells from the same embryo sometimes show greater similarity than cells from different embryos. For example, while all cells from the 16-cell stage have high memberships in the green cluster, cells from two of the embryos at this stage have memberships in both the purple and yellow clusters, while the other two embryos have memberships only in the yellow cluster.

Finally, we note that, like clustering methods, the GoM model can be helpful in exploratory data analysis and quality control. Indeed, the GoM results highlight a few single cells as outliers. For example, a cell from a 16-cell embryo is represented by the blue cluster - a cluster that represents cells at the zygote and early 2-cell stage. Also, a cell from an 8-stage embryo has strong membership in the purple cluster - a cluster that represents cells from the blastocyst stage. It would seem prudent to consider excluding these cells from subsequent analyses of these data.

0.1 Comparison with existing methods

Common methods for visualizing the structure of gene expression data include hierarchical clustering, PCA, multidimensional scaling (MDS), and t-Distributed Stochastic Neighbor Embedding (t-SNE). All of these methods, except for hierarchical clustering, are tools for dimension reduction of multivariate data. The visualizations produced by these methods may vary and depend on the structure of sample variation in the dataset. To highlight their differences and similarities, we provide visualizations produced by these methods for all the example datasets in the supplementary materials.

PCA and MDS map multivariate gene expression data to lower dimensions that aim to maximize global variation between samples. The sample coordinates in PCA/MDS dimensions are traditionally presented in two-dimensional scatter plots. However to extract the full structure from the data, one would need to perform a series of bi-scatter plots for different PC dimensions. For the GTEx data, the PCA plot of the first versus the second dimension, presented in ??, reveals that the whole blood samples are the most distinct among all the GTEx tissue samples. The higher PCA/MDS dimensions further reveal that xx tissue samples are separated from xx tissue samples (when plotting first versus second dimension) and xx tissue samples are separated from xx tissue samples (xx versus xx dimension) (TO DO: check PCA/MDS results to see if this is true and also up to which dimension). Nevertheless, sometimes only the first two PC dimensions provide a lot of information. For the mouse embryo data due to Deng et al [25], it is still pretty useful as it seems to identify the developmental phase trajectory

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from zygote to the blastocyst stages. Unlike PCA/MDS which projects the data effectively in 2 or maximum 3 dimensions at a time, the Structure plot representation by the GoM model is capable of representing the data in a lot more dimensions (same as the number of clusters) at a time, which leads to more effective representation of the latent structure.

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t-SNE was specifically developed for compact visualization of local behavior in a data using two-dimensional plots. For the GTEx data, the solution provided by t-SNE appears to produce nice clusters of samples, correspond to their tissue labels, evenly spread out in space, see ??. However, these visualizations provide little insight into the similarity between GTEx tissues. Also unlike PCA, the projections of t-SNE for different dimensions give drastically different results. Applied to the Deng et al mouse embryo data, t-SNE fails to represent the developmental phase trajectory unlike PCA/MDS. Applied to the GTEx brain data, t-SNE indicates no structure among the brain samples and the visualization does not group brain samples by subregions. Interestingly, the t-SNE visualization of the GTEx brain data is similar to the dendrogram produced by hierarchical clustering.

Hierarchical clustering is a distance-based clustering method that is frequently used to detect substructure in gene expression data. The method computes pairwise distance between samples and applies agglomerative clustering technique to produce a dendrogram visualization. The number of distances to be computed increases exponentially with the number of samples. For instance, the GTEx visualization using hierarchical clustering requires nearly 36 million distance computations. We performed hierarchical clustering and plotted the results in the form of a dendogram for the GTEx all tissues data, GTEx Brain tissue data and the Deng et al single cell developmental phase data in ??, ?? and ?? respectively. For the GTEx all tissues data, the dendogram revealed that the samples from same tissue seem to cluster together, however further patterns on how close different tissues are to one another is unclear. Also in this case, the dendogram is congested for so many samples, which makes it difficult to see patterns. For the GTEx brain data, the visualization revealed that Brain cerebellum and cerebellar hemisphere together form one outgroup, whereas the Spinal cord and Substantia nigra form another outgroup. These results are consistent with the PCA/MDS visualization. For the Deng et al developmental phase data, the three main clusters seem to be zygote to 4-cell stage, 8 cell and 16 cell and the blastocyst stages, but there is a fair amount of mixing among the cells from early stages of development which makes it difficult to see the trajectory of development.

Furthermore, we used the GTEx data to test whether the GoM model is more accurate in detecting substructure than distance-based clustering methods. Specifically, for each pair of tissues in the GTEx data we assessed whether or not each clustering method correctly partitioned samples into the two tissue groups (see Methods). The GoM model was substantially more accurate in this test, succeeding in 81% of comparisons, compared with 29% for the distance-based method (Fig 2). This presumably reflects the general tendency for model-based approaches to be more efficient that distance-based approaches, provided that the model is sufficiently accurate.

Another method that has close connection to the GoM models is sparse factor analysis (SFA). See [21] for discussion of relationships among these methods in the context of inferring population genetic structure. The GoM model requires all factors to be positive and their loadings summing up to 1. While, SFA is more flexible than the GoM model and allows the factors to be sparse as well as to hold either positive or negative loadings. However, the flexibility of SFA comes at the cost of making the results harder to interpret or to visualize. We have developed methods to visualize results from SFA for this particular application, but we feel that the visualization produced by SFA does not have the immediate appeal of the Structure plot. On the

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other hand, the flexibility of SFA in modeling both positive and negative contribution of underlying factors might allow the model to capture features that cannot be easily captured by the GoM models. Future work is needed to investigate the application of SFAT for detecting the structure of gene expression data.

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[Say something about NMF here?] Although people have applied NMF to expression data, they never used it to interpret the results and look at visualization as we have. In summary, our examples illustrate that the GoM models provide a more visually and biologically appealing summary of the data than the other dimension reduction techniques. In particular, for the GTEx data, the GoM models combined with Structure plot provides a convenient visualization of tissue samples in 20 dimensions corresponding to 20 clusters. While, PCA, MDS and t-SNE provide solutions that are difficult to visualize more than two dimensions at one time. For the GTEx brain tissue samples, PCA and MDS produce a solution in which most of the brain regions are not well distinguished except for Cerebellum and Cerebellar Hemisphere, and t-SNE fails to uncover a clustering pattern of the brain tissue samples (??). While, the GoM model revealed possibly four cell types underlying the brain subregions that are suggested as similar in PCA and MDS. Finally, applying to embryonic cell samples collected at each developmental stage, PCA and MDS recover a pattern of sample variation that is consistent with the developmental stages from which the embryos were collected. t-SNE groups cell samples by their embryonic stage. At the same time, the GoM models revealed a rich admixture of cell types that reflect important biological processes during embryonic preimplantation development.

Discussion

Our goal here is to highlight the potential for GoM models to elucidate structure in RNA-seq data from both single cell sequencing and bulk sequencing of pooled cells. We also provide tools to identify which genes are most distinctively expressed in each cluster, to aid interpretation of results. As our applications illustrate, these methods have the potential to highlight biological processes underlying the cluster structure identified.

The GoM model has several advantages over distance-based hierarchical methods of clustering. At the most basic level model-based methods are often more accurate than distance-based methods. Indeed, in our simple test on the GTEx data the model-based GoM approach more accurately separated samples into "known" clusters. However, there are also other subtler benefits of the GoM model. Because the GoM model does not assume a strict "discrete cluster" structure, but rather allows that each sample has a proportion of membership in each cluster, it can provide insights into how well a particular dataset really fits a "discrete cluster" model. For example, consider the results for the data from Jaitin et al [24] and Deng et al [25]: in both cases most samples are assigned to multiple clusters, although the results are closer to "discrete" for the latter than the former. The GoM model is also better able to represent the situation where there is not really a single clustering of the samples, but where samples may cluster differently at different genes. For example, in the GTEx data, the stomach samples share memberships in common with both the pancreas (purple) and the adrenal gland (light green). This pattern can be seen in the Structure plot (Fig 1) but would be hard to discern from a standard hierarchical clustering.

GoM models also have close connections with dimension reduction techniques such as factor analysis, principal components analysis and non-negative matrix factorization. All of these methods can also be used for RNA-seq data, and may often be useful. See [21] for discussion of relationships among these methods in the context of inferring population genetic structure. While not arguing that the GoM model is uniformly superior to these other methods, we believe our examples illustrate the appeals of the

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approach. In particular, we would argue that for the GTEx data, the Structure plot (Fig 1) combined with the cluster annotations (Table 1) provide a more visually and biologically appealing summary of the data than would a principal components analysis.

Fitting GoM models can be computationally-intensive for large data sets. For the datasets we considered here the computation time ranged from 12 minutes for the data from [25] (n=259;K=6), through 33 minutes for the data from [24] (n=1,041;K=7) to 3,370 minutes for the GTEx data (n=8,555;K=20). Computation time can be reduced by fitting the model to only the most highly expressed genes, and we often use this strategy to get quick initial results for a dataset. Because these methods are widely used for clustering very large document datasets there is considerable ongoing interest in computational speed-ups for very large datasets, with "on-line" (sequential) approaches capable of dealing with millions of documents [49] that could be useful in the future for very large RNA-seq datasets.

A thorny issue that arises when fitting these types of model is how to select the number of clusters, K. Like many software packages for fitting these models, the maptpx package implements a measure of model fit that provides one useful guide. However, it is worth remembering that in practice there is unlikely to be a "true" value of K, and results from different values of K may complement one another rather than merely competing with one another. For example, seeing how the fitted model evolves as K increases is one way to capture some notion of hierarchy in the clusters identified [17]. More generally it is often fruitful to analyse data in multiple ways using the same tool: for example our GTEx analyses illustrate how analysis of subsets of the data (in this case the brain samples) can complement analyses of the entire data.

The version of the GoM model fitted here is relatively simple, and could certainly be embellished. For example, the model allows the expression of each gene in each cluster to be a free parameter, whereas we might expect expression of most genes to be "similar" across clusters. This is analogous to the idea in population genetics applications that allele frequencies in different populations may be similar to one another [20], or in document clustering applications that most words may not differ appreciably in frequency in different topics. In population genetics applications incorporating this idea into the model, by using a correlated prior distribution on these frequencies, can help improve identification of subtle structure [20] and we would expect the same to happen here for RNA-seq data.

Methods and Materials

Model Fitting

We use the maptpx R package [15] to fit the GoM model (1,2), which is also known as "Latent Dirichlet Allocation" (LDA). The maptpx package fits this model using an EM algorithm to perform Maximum a posteriori (MAP) estimation of the parameters q and θ . See [15] for details.

Visualizing Results

In addition to the Structure plot, we have also found it useful to visualize results using t-distributed Stochastic Neighbor Embedding (t-SNE), which is a method for visualizing high dimensional datasets by placing them in a two dimensional space, attempting to preserve the relative distance between nearby samples [22,23]. Compared with the Structure plot our t-SNE plots contain less information, but can better emphasise clustering of samples that have similar membership proportions in many clusters. Specifically, t-SNE tends to place samples with similar membership proportions together

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in the two-dimensional plot, forming visual "clusters" that can be identified by eye (e.g. http://stephenslab.github.io/count-clustering/project/src/tissues_tSNE_ 2.html). This may be particularly helpful in settings where no external information is available to aid in making an informative Structure plot.

Cluster annotation

To help biologically interpret the clusters, we developed a method to identify which genes are most distinctively differentially expressed in each cluster. (This is analogous to identifying "ancestry informative markers" in population genetics applications [18].) Specifically, for each cluster k we measure the distinctiveness of gene g with respect to any other cluster l using

$$KL^{g}[k,l] := \theta_{kg} \log \frac{\theta_{kg}}{\theta_{lg}} + \theta_{lg} - \theta_{kg}, \tag{3}$$

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which is the Kullback-Leibler divergence of the Poisson distribution with parameter θ_{kg} to the Poisson distribution with parameter θ_{lg} . For each cluster k, we then define the distinctiveness of gene g as

$$D^{g}[k] = \min_{l \neq k} KL^{g}[k, l]. \tag{4}$$

The higher $D^g[k]$, the larger the role of gene g in distinguishing cluster k from all other clusters. Thus, for each cluster k we identify the genes with highest $D^g[k]$ as the genes driving the cluster k. We annotate the biological functions of these individual genes using the mygene R Bioconductor package [30].

For each cluster k, we filter out a number of genes (top 100 for the Deng $et\ al$ data [25] and GTEx V6 data [13]) with highest $D^g[k]$ value and perform a gene set over-representation analysis of these genes against all the other genes in the data representing the background. To do this, we used ConsensusPathDB database (http://cpdb.molgen.mpg.de/) [50] [51]. See Table 1-2 and Table 3 for the top significant gene ontologies driving each cluster in the GTEx V6 data and the Deng $et\ al$ data respectively.

Comparison with hierarchical clustering

We compared the GoM model with a distance-based hierarchical clustering algorithm by applying both methods to samples from pairs of tissues from the GTEx project, and assessed their accuracy in separating samples according to tissue. For each pair of tissues we randomly selected 50 samples from the pool of all samples coming from these tissues. For the hierarchical clustering approach we cut the dendogram at K=2, and checked whether or not this cut partitions the samples into the two tissue groups. (We applied hierarchical clustering using Euclidean distance, with both complete and average linkage; results were similar and so we showed results only for complete linkage.)

For the GoM model we analysed the data with K=2, and sorted the samples by their membership in cluster 1. We then partitioned the samples at the point of the steepest fall in this membership, and again we checked whether this cut partitions the samples into the two tissue groups.

Fig 2 shows, for each pair of tissues, whether each method successfully partitioned the samples into the two tissue groups.

Thinning 494

We used "thinning" to simulate lower-coverage data from the original higher-coverage data.. Specifically, if c_{ng} is the counts of number of reads mapping to gene g for sample

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n for the original data, we simulated thinned counts t_{nq} using

$$t_{ng} \sim Bin(c_{ng}, p_{thin}) \tag{5}$$

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where p_{thin} is a specified thinning parameter.

Code Availability

Our methods are implemented in an R package CountClust, available as part of the Bioconductor project at

 $https://www.bioconductor.org/packages/3.3/bioc/html/CountClust.html. \ The development version of the package is also available at$

https://github.com/kkdey/CountClust.

Code for reproducing results reported here is available at http://stephenslab.github.io/count-clustering/.

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Disclosure Declaration

The authors have no conflict of interest.

Supporting Information

S1 Fig. Structure plot of GTEx V6 all tissue samples for (a) K = 5, (b) K = 10, (c) K = 15, (d) K = 20. Some tissues form a separate cluster from the rest of the tissues from K = 5 onwards (for example: Whole Blood, Skin), whereas some tissues only form a distinctive subgroup only at K = 20 (for example: Arteries).

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S2 Fig. Structure plot of GTEx V6 all tissue samples K=20 in 2 runs under the thinning parameters settings (a) $p_{thin} = 0.01$ and (b) $p_{thin} = 0.0001$. The patterns in two plots closely correspond to the plot in Fig. 1 (a), though there are a few differences from the unthinned version.

S3 Fig. A comparison of "accuracy" of hierarchical vs. model-based clustering on thinned GTEx data, with thinning parameter $p_{thin} = 0.01$ and $p_{thin} = 0.0001$. For each pair of tissues from the GTEx data we assessed whether or not each clustering method (with K=2 clusters) separated the samples according to their actual tissue of origin, with successful separation indicated by a filled square. Thinning deteriorates accuracy compared with the unthinned data (Fig 2), but even then the model-based method remains more successful than the hierarchical clustering in separating the samples by tissue or origin.

S1 Table. Cluster Annotations GTEx V6 data with top driving gene summaries.

S2 Table. Cluster Annotations GTEx V6 Brain data with top driving gene summaries.

S3 Table. Deng et al (2014) Cluster GO annotations of top driving genes.

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Table 1. Cluster Annotations GTEx V6 data (with GO annotations).

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Cluster	Top 5 Driving Genes	Top significant GO terms (function)[q-value]	
1. Royal purple	NEAT1, IGFBP5, CCLN2, SRSF5, PNISR	GO:0005654 (nucleoplasm)[2e-10], GO:0044822 (poly-A RNA binding)[3e-09], GO:0044428 (nuclear part)[1e-09], GO:0043233 (organelle lumen)[2e-08]	
2. Light purple	SNAP25, FBXL16, NCDN, SNCB, SLC17A7	GO:0097458 (neuron part)[2e-25], GO:0007268 (synaptic transmission)[9e-18], GO:0030182 (neuron differentiation)[2e-14], GO:0022008 (neurogenesis)[1e-13], GO:0007267 (cell-cell signaling)[3e-13]	
3. Red	FABP4, PLIN1, FASN, GPX3, LIPE	GO:0044255 (cellular lipid metabolism)[1e-09], GO:0006629 (lipid metabolism)[1e-09], GO:0006639 (acyl-glycerol metabolism)[3e-08], GO:0045765 (angiogenesis regulation)[4e-08], GO:0019915 (lipid storage)[4e-08]	
4. Salmon	ACTG2, MYH11, SYNM, MYLK, CSRP1	GO:0043292 (contractile fiber)[3e-13], GO:0006936 (muscle contraction)[5e-12], GO:0030016 (myofibril)[5e-12], GO:0015629 (actin cytoskeleton)[2e-12], GO:0005925 (focal adhesion)[6e-11]	
5. Denim	RGS5, MGP, AEBP1, IGFBP7, MFGE8	GO:0005578 (proteinaceous extracellular matrix)[4e-20], GO:0030198 (extracellular matrix)[2e-18], GO:0007155 (cell adhesion)[4e-14], GO:0001568 (blood vessel development)[4e-13]	
6. Light denim	KRT10, KRT1, KRT2, LOR, KRT14	GO:0008544 (epidermis development)[3e-12], GO:0043588 (skin development)[5e-10], GO:0042303 (molting cycle)[8e-06], GO:0042633 (hair cycle)[7e-06], GO:0048513 (organ development)[6e-05]	
7. Orange	NEB, MYH1, MYH2, MYBPC1, ACTA1	GO:0043292 (contractile fiber)[6e-52], GO:0030016 (myofibril)[1e-51], GO:0030017 (sarcomere)[5e-40], GO:0003012 (muscle system process)[2e-25], GO:0015629 (actin cytoskeleton)[1e-25]	
8. Light orange	FN1, COL1A1, COL1A2, COL3A1, COL6A3	GO:0030198 (extracellular matrix)[6e-29], GO:0043062 (extracellular structure)[4e-29], GO:0032963 (collagen metabolism)[3e-16], GO:0030199 (collagen fibril organization)[1e-14], GO:0030574 (collagen catabolism)[1e-14]	
9. Green	MBP, GFAP, MTURN, HIPK2, CARNS1	GO:0043209 (myelin sheath)[4e-07], GO:0007399 (nervous system development)[4e-05], GO:0008366 (axon ensheathment)[9e-05], GO:0044430 (cytoskeletal part)[1e-04], GO:0005874 (microtubule)[3e-04]	
10. Light green	CYP17A1, CYP11B1, PIGR, GKN1, STAR	GO:0006694 (steroid biosynthesis)[2e-13], GO:0008202 (steroid metabolism)[1e-12], GO:0016125 (steroid metabolism)[1e-11], GO:0042446 (hormone biosynthesis)[1e-10], GO:0008207 (C21-steroid hormone metabolism)[3e-10]	
11. Turquoise	MPZ, APOD, PMP22, PRX, NGFR	GO:0007272 (ensheathment of neurons)[4e-07], GO:0008366 (axon ensheathment)[7e-07], GO:0042552 (myelination)[7e-06], GO:0048856 (anatomical structure development)[1e-06], GO:0005578 (proteinaceous extracellular matrix)[1e-06]	
12. Yellow	IGHM, IGHG1, IGHG2, IGHG4, CD74	GO:0006955 (immune response)[1e-18], GO:0002252 (immune effector process)[7e-18], GO:0003823 (antigen binding)[1e-15], GO:0019724 (B-cell mediated immunity)[5e-13], GO:0002684 (positive regulation immune system)[6e-13]	
13. Sky blue	TG, PRL, GH1, PRM2, PRM1	GO:0019953 (sexual reproduction)[8e-10], GO:0048232 (male gamete generation)[2e-08], GO:0035686 (sperm fibrous sheath)[4e-06], GO:0005179 (hormone activity)[6e-05], GO:0042403 (thyroid hormone metabolism)[2e-04]	
14. Light pink	NPPA, MYH6, TNNT2, ACTC1, MYBPC3	GO:0045333 (cellular respiration)[2e-34], GO:0022904 (respiratory electron transport)[8e-33], GO:0015980 (energy derivation by oxidation of organic compounds)[4e-30], GO:0031966 (mitochondrial membrane)[5e-26]	
15. Light gray	KRT13, KRT4, MUC7, CRNN, KRT6A	GO:0070062 (extracellular exosome)[2e-23], GO:0043230 (extracellular organelle)[3e-23], GO:0031982 (vesicle)[3e-20], GO:0008544 (epidermis development)[2e-18], GO:0043588 (skin development)[1e-13]	
16. Gray	$SFTPB\beta$, $SFTPA1$, $SFTPA2$, $SFTPC$, $A2M$	GO:0001525 (angiogenesis)[5e-08], GO:0001944 (vasculature development)[2e-07], GO:0048514 (blood vessel morphogenesis)[2e-07], GO:0040012 (locomotion regulation)[4e-06], GO:2000145 (cell motility)[1e-05]	
17. Brown	CSF3R, MMP25, IL1R2, SELL, VNN2	GO:0006955 (immune response)[8e-22], GO:0006952 (defense response)[9e-16], GO:0071944 (cell periphery)[7e-15], GO:0005886 (plasma membrane)[7e-15], GO:0050776 (regulation of immune response)[2e-12]	
18. Purple	PRSS1, CPA1, PNLIP, CELA3A, GP2	GO:0007586 (digestion)[3e-14], GO:0004252 (serine-type endopeptidase activity)[4e-08], GO:0006508 (proteolysis)[6e-06], GO:0016787 (hydrolase activity)[6e-05], GO:0044241 (lipid digestion)[1e-04]	
19. Pink	HBB, HBA2, HBA1, FKBP8, HBD	GO:0005833 (hemoglobin complex)[1e-13], GO:0015669 (gas transport)[4e-11], GO:0020037 (hemoglobin binding)[3e-07], GO:0031720 (haptoglobin binding)[3e-06], GO:0006950 (response to stress)[6e-04]	
20. Dark gray	ALB, HP, FGB, FGA, ORM1	GO:0072562 (blood microparticle)[1e-27], GO:0043230 (extracellular organelle)[1e-24], GO:0044710 (single organism metabolism)[7e-20], GO:0019752 (carboxylic acid metabolism)[1e-18], GO:0034364 (high density lipoprotein)[3e-16]	

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Table 2. Cluster Annotations for GTEx V6 Brain data.

Cluster	Top 5 Driving Genes	Top significant GO terms (function)[q-value]	
1. Royal blue	CLU, OXT, GLUL, NDRG2, CST3	GO:0043230 (extracellular organelle)[1e-14], GO:1903561 (extracellular vesicle)[2e-14], GO:0031988 (membrane bound vesicle)[3e-14], GO:0070062 (extracellular exosome)[4e-13], GO:0006950 (response to stress)[2e-12]	
2. Turquoise	ENC1, NCALD, YWHAH, KIF5A, NPTXR	IAH, KIF5A, ron projection development)[7e-09], GO:0007268 (synaptic transmission)[1e-08], GO:0030182 (neuron	
3. Lime green	PKD1, CBLN3, CHGB, COL27A1, ABLIM1	GO:0005089 (Rho guanyl-nucleotide exchange factor activity)[1e-03], GO:0016604 (nuclear body)[1e-03], GO:0022008 (neurogenesis)[2e-02], GO:0035239 (tube morphogenesis)[8e-02], GO:0007269 (neurotransmitter secretion)[0.13]	
4. Red	PPP1R1B, RGS14, NCDN, PDE1B, RAP1GAP	$N_{ m N} = PDE1B_{ m N} = 05$], GO:0007268 (synaptic transmission)[1e-03], GO:0010646 (cell communication regulation)[1e-03]	
5. Yellow orange	MBP, GFAP, TF, MTURN, SCD	GO:0043209 (myelin sheath)[2e-09], GO:0007399 (nervous system development)[1e-05], GO:0005737 (cytoplasm)[1e-04], GO:0007272 (ensheathment of neurons)[1e-03], GO:0008366 (axon ensheathment)[1e-03]	
6. Yellow	IQGAP1, A2M, C3, MYH7, TG	GO:0072562 (blood microparticle)[1e-10], GO:0044449 (contractile fiber part)[1e-10], GO:0043230 (extra cellular organelle)[7e-10], GO:0030017 (sarcomere)[1e-09], GO:0072376 (protein activation cascade)[1e-08]	

Table 3. Cluster Annotations for Deng et al (2014) data.

Cluster	Top 10 Driving Genes	Top significant GO terms (function)[q-value]
1. Blue	Bcl2l10, E330034G19Rik, Tcl1,LOC100502936, Oas1d, AU022751, Spin1, Khdc1b, D6Ertd527e, Btg4	$\label{eq:G0:0007276} Go:0007276 \ (gamete \ generation)[7e-06], \ GO:0032504 \ (multicellular \ organism \ reproduction)[3e-06], \ GO:0044702 \ (single \ organism \ reproduction)[2e-05], \ GO:0048477 \ (oogenesis)[5e-04], \ GO:0048599 \ (oocyte \ development)[9e-04], \ GO:0009994 \ (oocyte \ differentiation)[1e-03]$
2. Magenta	Obox3, Zfp352, Gm8300, Usp17l5, BB287469, Rfpl4b, Gm2022, Gm5662, Gm11544, Gm4850	$\label{eq:GO:0016604} GO:0016604 \ (nuclear \ body)[1e-04], \ GO:0005814 \ (centriole)[4e-03], \ GO:0044450 \ (microtubule \ organizing \ center \ part)[8e-03]$
3. Yellow	Rtn2, Ebna1bp2, Zfp259, Nasp, Cenpe, Rnf216, Ctsl, Tor1b, Ankrd10, Lamp2	GO:0044428 (nuclear part)[1e-08], GO:0031981 (nuclear lumen)[3e-08], GO:0070013 (intracellular organelle lumen)[9e-08], GO:0005730 (nucleolus)[5e-07], GO:0005654 (nucleoplasm)[4e-05]
4. Green	Timd2, Isyna1, Alppl2, Prame15, Fbxo15, Tceb1, Gpd1l, Pemt, Hsp90aa1, Hsp90ab1	GO:0005829 (cytosol)[4e-10], GO:0044444 (cytoplasmic part)[2e-05], GO:1901575 (organic substance catabolic process)[9e-05], GO:0000151 (ubiquitin ligase com- plex)[1e-04], GO:0023026 (MHC class II protein complex binding)[4e-04], GO:0051082 (unfolded protein binding)[9e-04]
5. Purple	Upp1, Tdgf1, Aqp8, Fabp5, Tat, Pdgfra, Pyy, Prdx1, Col4a1, Spp1	GO:0070062 (extracellular exosome)[1e-05], GO:0043230 (extracellular organelle)[2e-05], GO:1903561 (extracellular vesicle)[1e-05], GO:0006950 (response to stress)[1e-05], GO:0048514 (blood vessel morphogenesis)[7e-04], GO:0001944 (vasculature development)[3e-03]
6. Orange	Actb, Krt18, Fabp3, Id2, Tspan8, Gm2a, Lgals1, Adh1, Lrp2, BC051665	GO:0065010 (extracellular membrane-bounded organelle)[3e-23], GO:1903561 (extracellular vesicle)[3e-23], GO:0070062 (extracellular exosome)[4e-23], GO:0043230 (extracellular organelle)[5e-23], GO:0032432 (actin filament bundle)[1e-09], GO:0030036 (actin cytoskeleton and organization)[5e-09]

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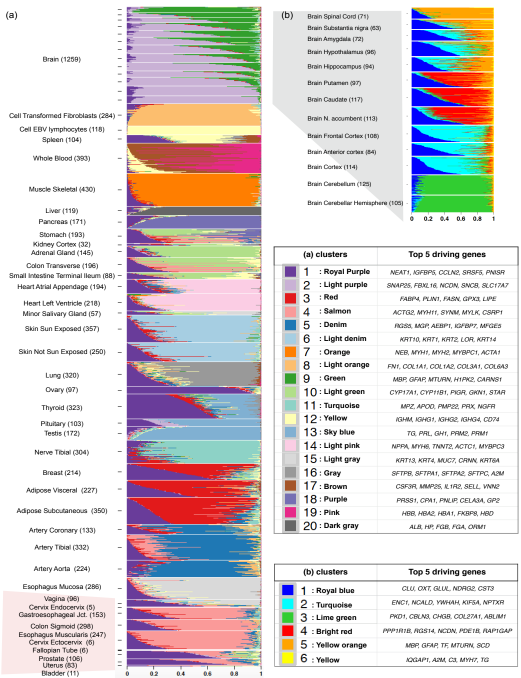
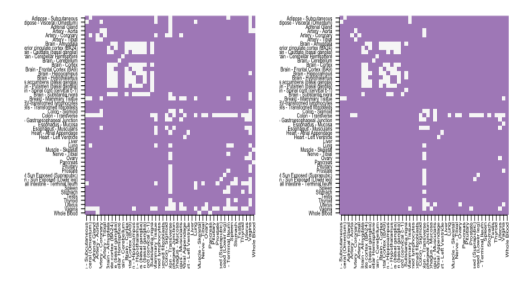


Fig 1. (a): Structure plot of estimated membership proportions for GoM model with K=20 clusters fit to 8555 tissue samples from 53 tissues in GTEx data. Each horizontal bar shows the cluster membership proportions for a single sample, ordered so that samples from the same tissue are adjacent to one another. Within each tissue, the samples are sorted by the proportional representation of the underlying clusters. (b): Structure plot of estimated membership proportions for K=4 clusters fit to only the brain tissue samples. This analysis highlights finer-scale structure among the brain samples that is missed by the global analysis in (a).

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(a) hierarchy method $\,$

(b) GoM method

Fig 2. A comparison of accuracy of GoM model vs hierarchical clustering. The hierarchical clustering was performed on the log-normalized RPKM (reads per kilobase per million) data, that adjusts for the library size of the samples as well as gene lengths. For each pair of tissues from the GTEx data we assessed whether or not each method (with K=2 clusters) separated the samples precisely according to their actual tissue of origin, with successful separation indicated by a filled square. Some pairs of tissues (e.g. pairs of brain tissues) are more difficult to distinguish than others. Overall the GoM model is successful in 86% comparisons and the hierarchical clustering in 39% comparisons.

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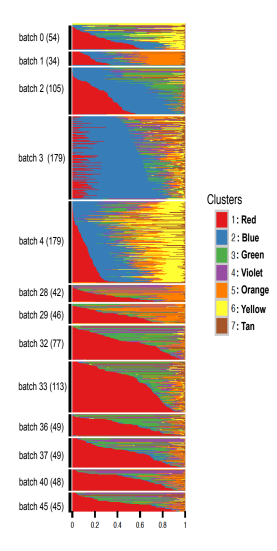


Fig 3. Structure plot of estimated membership proportions for GoM model with K=7 clusters fit to 1,041 single cells from [24]. The samples (cells) are ordered so that samples from the same amplification batch are adjacent and within each batch, the samples are sorted by the proportional representation of the underlying clusters. In this analysis the samples do not appear to form clearly-defined clusters, with each sample being allocated membership in several "clusters". Membership proportions are correlated with batch, and some groups of batches (e.g. 28-29; 32-45) show similar palettes. These results suggest that batch effects are likely influencing the inferred structure in these data.

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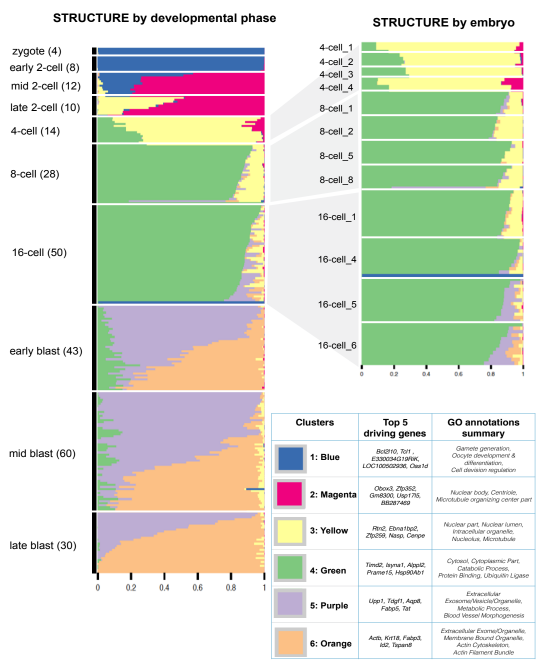


Fig 4. Structure plot of estimated membership proportions for GoM model with K=6 clusters fit to 259 single cells from [25]. The cells are ordered by their preimplantation development phase (and within each phase, sorted by the proportional representation of the clusters). While the very earliest developmental phases (zygote and early 2-cell) are essentially assigned to a single cluster, others have membership in multiple clusters. Each cluster is annotated by the genes that are most distinctively expressed in that cluster, and by the gene ontology categories for which these distinctive genes are most enriched (see Table 3 for more extensive annotation results). See text for discussion of biological processes driving these results.

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