

1 Outline

The outline for this paper (in the format of George M. Whitesides)

Title:

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2 Introduction

- *objectives of the work*: to devise a completely unsupervised method to cluster the samples (tissue or single cell samples) into biologically meaningful sub-types based on the RNA-seq gene counts data
- *justification of objectives* :
 1. People have mainly used hierarchical clustering from GTEx consortium paper to most single cell RNA seq papers I have come across. We have evidence Admixture model does better than hierarchical clustering from a biological viewpoint (see structure.beats.hierarchical.html).
 2. Hierarchical clustering does not give us directly the genes that drive the clusters, Admixture model does, and it also provides us with a log likelihood to fix how many clusters to choose, based on Bayes factor.
 3. We can predict the admixture proportions of cell types in any new sample coming in, so we can easily cluster new samples in cancer biopsy where the sub-types may involve cancer or non-cancer samples.
- *Background*
 1. The BackSpin algorithm used by Zeisel et al. Claim is it does better than hierarchical but not model based (also not convincingly proven to be better)
 2. Use of downsampling and then modified hierarchical clustering scheme as applied by Jaitin et al.
 3. Mainly, people have used hierarchical clustering scheme
 4. Population genetics uses Admixture model on a regular basis. We think we can generalize that to RNA-seq data. The only question is do we really see the tissue samples as cell type admixture, as we observe individuals as population admixture. The answer seems to be yes.
- *Guidance to the reader*
 1. The Structure plot and t-SNE plots for GTEx tissues and for Zeisel data. Much better visualization than the regular heatmaps that we tend to see in RNA-seq papers.
 2. The Structure plot analysis for Brain samples that shows 80% one cluster in cerebellum tissue samples and then from gene annotations, it is revealed this cluster is indeed associated with synaptic activities implying it must be neuronal cell types.

This is pretty cool because we have a priori knowledge from cell type specific markers that around 80% of cells in cerebellum are neurons.

3. Also the strategy is similar to the topic model strategy in natural language processing and it is a really nice technique to use for RNA-seq datasets clustering.

3 Methods and Materials

3.1 Data preprocessing

RNA-seq experiments provide us with a set of FASTQ files that contain the nucleotide sequence of each read and a quality score at each position, which can be mapped to reference genome or exome or transcriptome. The output of this mapping is usually saved in a SAM/BAM file using SAMtools [2], a task primarily accomplished by *htseq-counts* by Sanders et al 2014 [1] or *feature-Counts* [R package **Rsubread**] by Liao et al 2013 [3]. RNA-seq raw counts are the basis of all statistical workflows, be it exploration or differential expression analysis [**edgeR** [4], **limma** [5]]. There is a growing trend to make the analysis ready raw counts tables openly accessible for statistical analysis. ReCount is a online site that hosts RNA-seq gene counts datasets from 18 different studies [6] along with relevant metadata. Such gene counts datasets are the inputs for our clustering algorithm.

In the preprocessing step before applying our method, we remove the genes with 0 or same count of matched reads across all samples (non-informative genes), any sample or gene with NA values of reads and ERCC spike-in controls, as the latter may create bias due to their typical very high expression (number of reads mapped to them). For illustration, we applied our method GTEx Version 6 tissue level gene counts data [8] and on a couple of single cell data due to Zeisel *et al* [7] and Jaitin *et al* [16].

3.2 Model overview

We use a topic model approach due to Matt Taddy (package **maptpx**) to perform the clustering of the samples based on RNA-seq reads data [9]. Let us denote the gene counts matrix as $C_{N \times G}$ where N is the total number of samples (tissue/single cell) and G is the number of genes. We assume that the row vector of counts for each sample n across the genes follows a multinomial distribution.

$$c_{n*} \sim Mult(c_{n..}, p_{n*})$$

where c_{n*} is the count vector for the n th sample, $c_{n..}$ is the sum of the counts in the vector c_{n*} , and p_{n*} is the probability that a read coming from sample n would get assigned to one of the G genes. The idea here is that this read could be coming from some cell type for the tissue level expression study (or from some cell cycle phase for the single cell case study) and its probability of getting assigned to some gene g will depend on which cell type (cell cycle phase) it comes from. In general, we may assume that the read is coming from one of the several (say K) underlying classes/groups, which are not observed. Denote the probability that the sample

is coming from the k th subgroup by q_{nk} ($q_{nk} \geq 0$ and $\sum_{k=1}^K q_{nk} = 1$ for each n) and then given that the sample is coming from the k th subgroup, the probability of a read being matched to the g th gene is given by θ_{kg} ($\theta_{kg} \geq 0$ and $\sum_{g=1}^G \theta_{kg} = 1$ for k th subgroup). Then one can write

$$p_{ng} = \sum_{k=1}^K q_{nk} \theta_{kg} \quad \sum_{k=1}^K q_{nk} = 1 \quad \sum_{g=1}^G \theta_{kg} = 1$$

This model has in all $N \times (K - 1) + K \times (G - 1)$ many unconstrained parameters, which is much smaller than the $N \times G$ data values of counts. Usually $K \ll \min\{N, G\}$, N in the region of 100s to 1000s and G ranging from 10,000 to 50,000 (depending on the species the RNA-seq data is coming from and whether the set of genes recorded include non-protein genes or not). To estimate the model, a Maximum a posteriori (MAP) based approach is used (see Taddy 2012 [9]).

3.3 Visualization

For each n , q_{nk} 's which will give an idea about the relative abundance of individual subgroups (which may be driven by cell functional groups or cell types) represented in the sample (single cell or tissue respectively). If two samples n and n' are very close, say both coming from the same tissue for the tissue level data, then we expect q_{n*} and $q_{n'*}$ to be very close too. A nice way to visualize the amount of relatedness among the samples is through the Structure plot due to Pritchard Lab, which is a popular tool to visualize the admixture patterns in population genetics based on SNP/ microsatellite data [10] [11]. The Structure plot assigns a color to each of the subgroups and then presents a vertical barplot for each individual, which is fragmented by the subgroup proportions and colored accordingly. If the colored patterns of two bars are similar, then the two samples must be closely related.

Another visualizing tool we recommend is t-distributed Stochastic Neighbor Embedding (t-SNE) due to Laurens van der Maaten, which is well-suited for visualizing the high dimensional datasets on 2D, preserving the relative distance between samples in high dimension to a fair extent in 2D [12] [13]. t-SNE provides some sense about which samples are closer to each other when the data is projected on 2D. But on the flipside, it is not a clustering tool and unlike Structure plot, does not show the relative abundance patterns of different subgroups in the sample. However, both Structure plot and t-SNE give a lot more interpretable visualization of the clustering compared to the heatmap and hierarchical clustering (see Results for illustration).

3.4 Cluster annotation

A question of considerable biological interest is which genes are significantly differentially expressed across the clusters, or in other words, which genes are driving the clustering. To answer this, we fix each gene and then look at the KL divergence matrix of one cluster/subgroup k relative to other cluster/subgroup k' , which we call $KL_{K \times K}^g$. This matrix is symmetric and has all diagonal elements 0 as the divergence of a cluster with respect to itself is 0. Next we define the divergence measure for gene g as

$$Div(g) = \max_k \min_{l \neq k} KL^g[k, l]$$

$$K_{div}(g) = \arg \max_k \min_{l \neq k} KL^g[k, l]$$

The higher the divergence measure, the more significant is the role of the gene in the clustering. We choose a small subset of around 50-100 genes with highest values of $Div(g)$ and put the gene in the $K_{div}(g)$ th cluster/subgroup. Then we perform gene annotations for the top genes in each subgroup using **mygene** R Bioconductor package [20]. We observe if the significant genes in a particular subgroup/cluster are associated with some specific biological functionality. This would indicate if the subgroups are actually biologically relevant or not. For instance, for GTEx tissue sample data, if the clusters are indeed driven by cell types, then the top genes for these clusters will probably be associated with proteins related to functions for that particular cell type.

4 Results

We begin by illustrating our method on the tissue level data from the GTEx project (V6 dbGaP accession phs000424.v6.p1, release date: Oct 19, 2015, <http://www.gtexportal.org/home/>) read counts data. RNA-seq data was obtained from 8555 samples collected from 450 donors across 51 tissues and 2 cell-lines. From GTEx V4 data eQTL analysis, we extracted a set of 16407 cis-genes that fulfilled some quality criteria (gene list available in <https://github.com/stephenslab/count-clustering>).

Fig 1 presents the Structure plot for admixture model fit for $K = 15$. The Structure plot clearly highlights the similarity among the samples coming from the same tissue and also gives an idea about which tissues have similar patterns of gene expression. It is pretty nice that all the Brain tissues seem to cluster together and show very similar patterns, the same being true for the arteries (Artery-aorta, Artery-tibial and Artery-coronary). Interestingly, Muscle Skeletal and Heart tissues (Heart Left Ventricle and Heart Atrial Appendage) also seem to be very close in their clustering patterns.

As observed from the Structure plot, some tissues seem to be assigned to separate clusters for $K = 15$, (e.g: testis, whole blood), but other tissues are represented as an "admixed" version of multiple clusters (e.g: thyroid, adipose subcutaneous). But in these latter cases, the samples coming from the same tissue all seem to be cluster together as they have very similar patterns of "admixing" of different clusters. A different way of visualizing the results, which highlights the clustering and separation of the different tissues on a 2D projection space (see **Supplementary Fig 1** [url]).

In order to get a sense of the biological interpretation of the clusters in **Fig 1**, we performed cluster annotation (see Methods and Materials). In **Tab ??**, we present the gene IDs, names and a short summary of their functions, obtained from the **mygene** package in R [20]. As can be seen from the table, *PRM2*(protamine2), *PRM1* (protamine1) and *PHF7* (PHD finger protein 7) are the top three genes that drive the cluster which separates out testis from the other tissues in the admixture cluster model in **Fig 1**. Similarly, *HBB* (hemoglobin, beta),

HBA2 (hemoglobin, alpha 2) and *HBA1* (hemoglobin, alpha 1) seem to be the top three genes that distinguish the whole blood and for a separate cluster from the rest. Overall it seems from **Tab ??** that cluster annotation is highlighting tissue specific functions and pathways.

Fig 1 Structure plot and the corresponding cluster annotation mainly highlighted inter tissue variation in the clusters, which is pretty logical because the cells in different tissues are pretty differentiated. We were curious if we can extract some cell type information if we focus on samples from a particular tissue or similar tissues. **Fig 2** shows the Structure plot for $K = 4$ on just the Brain samples data. Brain Cerebellum and Cerebellar hemisphere stand out in the plot as we see one cluster (blue) explaining around 80 – 85% admixture proportion. Recent stereological approaches have shown that rat cerebellum contains $> 80\%$ neurons (Herculano-Houzel and Lent 2005) [19], much higher than other parts of the brain. We performed cluster annotation (Supplementary Table 1) and observed that the pivotal genes that separated out the blue cluster in brain cerebellum and cerebellar hemisphere were SNAP25 (synaptosomal-associated protein, 25kDa), ENO2 (enolase 2- gamma, neuronal) and CHGB (chromogranin B) all of which were associated with neuronal activities. It is definitely not true that the blue cluster represents the neuron cell type as it does not show up in other parts of the Brain, but it seems to be driven by neuronal cell type.

We next sought to demonstrate more quantitatively the utility of the model based clustering compared to other non model based clustering methods such as hierarchical clustering. In **Fig 3**, we consider every pair of tissues from the list of tissues in GTEx with number of samples > 50 . Then we generated a set of 50 samples randomly drawn from the pooled set of samples coming from these two tissues and then observed whether the hierarchical and the admixture were separating out samples coming from the two different tissues. The same remains true for data sets thinned to simulate the level of lower coverage data that might be observed in single cell experiments (the threshold parameter $p_{thin} = 0.0001$ taken on comparing the GTEx data with Jaitin *et al* data [16]). Check **Fig 6** for demonstration.

For the thinned data, though the cluster quality is poor compared to the original data, it still seems that the clustering patterns remain preserved to a great extent. **Fig 4** presents the Structure plot for $K = 12$ for the GTEx thinned data with $p_{thin} = 0.0001$. Many of the features from **Fig 1** are restored even after thinning, for instance the Brain tissues clustering together, Heart samples and Muscle Skeletal samples showing similar patterns. This implies that Admixture as a clustering technique is pretty robust to the coverage of the data.

We applied the admixture model on a couple of single cell datasets due to Jaitin *et al* [16] and Zeisel *et al* [7]. Jaitin *et al* sequenced around 4000 single cells from mouse spleen, where the cells were a heterogeneous mix enriched for expression of CD11c marker. The aim of their study was to separate out the B cells, NK cells, pDCs and monocytes. However the biological effect in their study was completely confounded with the amplification and sequencing batches. **Fig 5** (top panel) presents the Structure plot for $K = 7$ for the Jaitin *et al* data with the samples arranged by their amplification batch (which was a refinement of the sequencing batch).

Zeisel *et al* analyzed the single cell data obtained from mouse cortex and hippocampus and obtained 47 molecularly distinct subclasses, comprising all known major cell types in the region. They also identified many marker genes informative about cell types, morphology and location. **Fig 5** (bottom panel) presents the Structure plot for $K = 10$ on their data, where the samples in Structure plot are grouped by their subclass assignment and are in the same order as the data presented within each group. It was interesting that the first few samples under Oligo6 seemed to show some "admixture" due to red cluster, which was not observed in other Oligo6 samples. These samples in Oligo6 were pretty different in pattern from the rest of Oligo6 samples which had no trace of red cluster. Since within each group, the samples are ordered in the same order as reported in the dataset, there is a possibility that adjacent samples may be coming from same plate or may be sequenced in same lane etc, all of which can lead to similar patterns.

The main highlight of **Fig 5** is that one must be careful about interpreting Admixture results or any clustering results, as there is a possibility of technical effects driving the clusters instead of true biological effects. There has been a growing concern among biostatisticians today about how to deal with batch effects [17] [18].

5 Discussions

We suggest a model based clustering approach for RNA seq or scRNA seq data that takes as input the read counts matrix over the samples and genes and number of clusters to fit (K), and gives as output the admixture proportions matrix for all the samples and the relative expression profiles of the genes in each of the K clusters. It also provides us with a model log-likelihood that can be used to choose the optimal K to fit. However for genetic data, it is more recommended to observe the clustering patterns over a range of values of K to observe how the patterns change as we increase K . The clustering method is pretty fast as it uses EM algorithm along with quasi-Newton updates to speed up the iterations. The clustering proportions obtained as output can be viewed using a Structure plot or the t-SNE plot that give a much better visual representation of the clustering patterns than heatmap or PCA. As per model specifications, ideally the clusters should be driven by the cell types and we already have seen some evidence in support of that in **Fig 2** when the model was applied on brain samples. Besides the cluster proportions, the model also provides the user with the relative expression profile of all genes in each of these clusters, from which it is easy to figure out which genes have significantly high expression in one or more clusters compared to the other clusters or in other words, are informative in driving the clusters. The user can select these cluster driving genes and annotate them to get a better understanding of the biological significance of the clusters. Even purely as a clustering technique, our method outperforms hierarchical method in separating out the samples belonging to distinct classes (in case of the GTEx data, the different tissues). So, overall we feel our model has a number of advantages over the standard methods of clustering used in RNA-seq or scRNA-seq literature, like hierarchical clustering, in terms of cluster quality, biological validation, visualization and interpretation. The clustering along with the Structure plot representation based on the sample metadata is implemented in package **CountClust** available on Github (<https://github.com/kkdey/CountClust>) which is a wrapper package of **maptpx** due

to Matt Taddy [9].

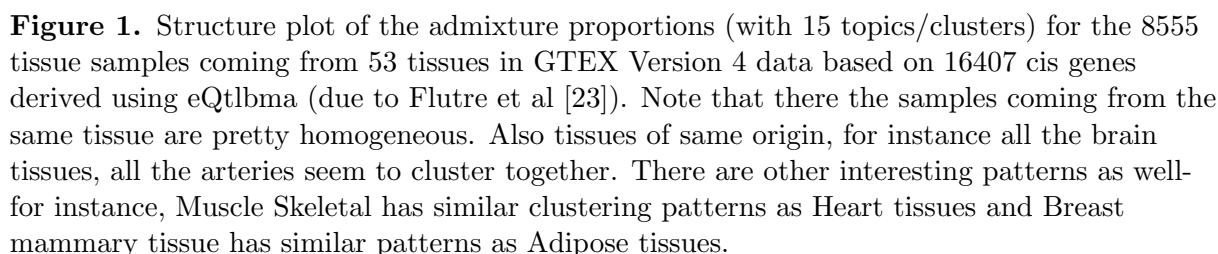
Future works

- It will be worthwhile to see if instead of finding out cluster driving genes, we can find out cluster driving gene pathways. which would have significance from a biomedical standpoint.
- Since many of the genes are not informative for the clustering, we may try to impose a variable selection preprocessing or incorporate that in our model suitably so that it will extract out only the genes that are informative about the clusters and will also speed up the model fitting.
- The admixture proportions may be useful for determining the mixing weights for the prior covariance matrices in the eQtlbma.
- We may have important metadata on the samples (for instance the individual from whom the sample came from) or on the genes (for instance the gene length, GO or KEGG annotations, GC content etc) which we have not incorporated in our clustering model so far.

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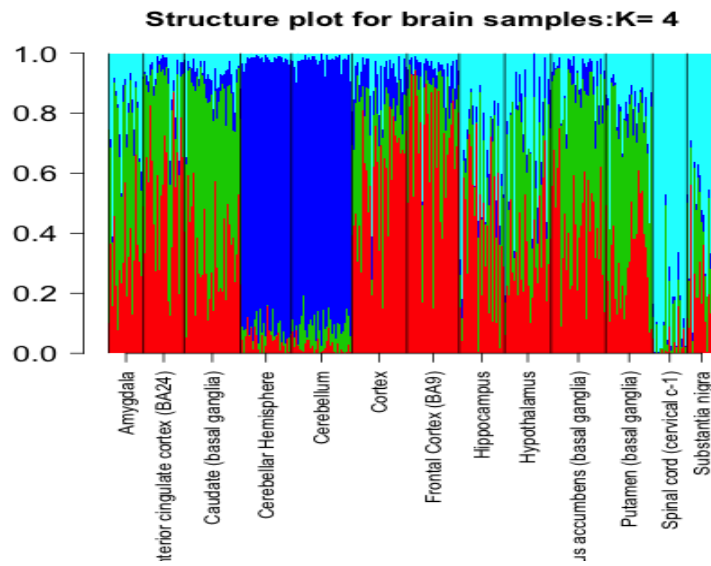


Figure 2. Structure plot of the admixture proportions (with 4 clusters) for the brain tissue samples drawn from GTEx Version 4 data. Quite clearly, brain cerebellum and cerebellar hemisphere seem to be dominated by the blue cluster while the Spinal cord and Substantia nigra by the cyan cluster. Prior marker based approaches have verified that $> 80\%$ of cells in brain cerebellum correspond to neurons [19]. So, the blue cluster seems to be driven by the neuron cell type. This fact is further attested by the gene annotations of the top genes driving the blue cluster (Supplementary Table 1).

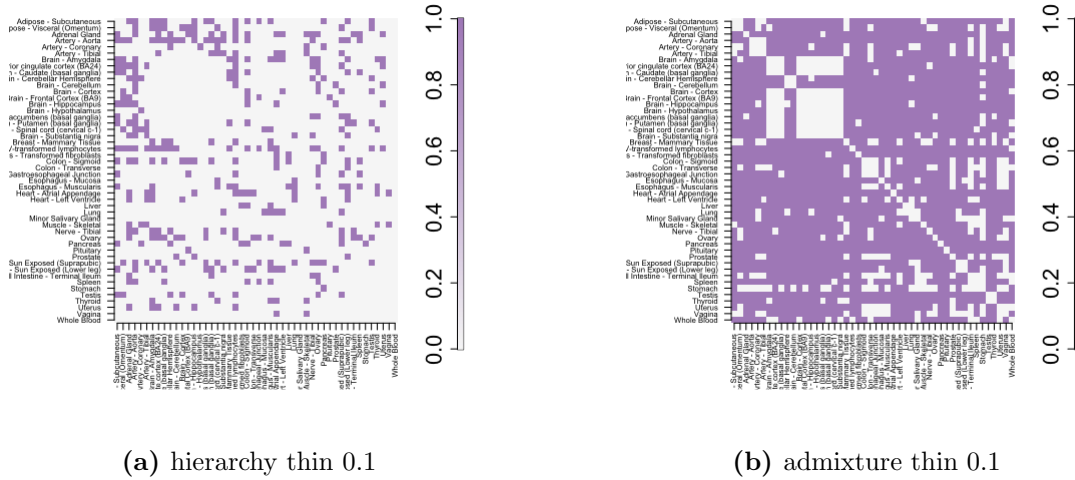


Figure 3. A comparison of the hierarchical method with the admixture method. For each pair of tissues, we selected randomly 50 samples and then on the reads data for these 50 samples, we applied the hierarchical clustering method with complete linkage and Euclidean distance and then cut the tree at $K = 2$. We then observed if it separates out the samples coming from the two tissues, in case it does, we color the cell corresponding to that pair of tissues. We apply admixture model on the same data for $K = 2$. Then we fixed one cluster, observed the proportions for that cluster, sorted the samples based on the proportions for that cluster and separated out the samples at the point of maximum jump/fall in the proportions for that cluster. If that separates out the two tissues, we color the cell, else keep it blank. From the graph it seems that the admixture model has been far more successful in separating out different tissues compared to the hierarchical method.

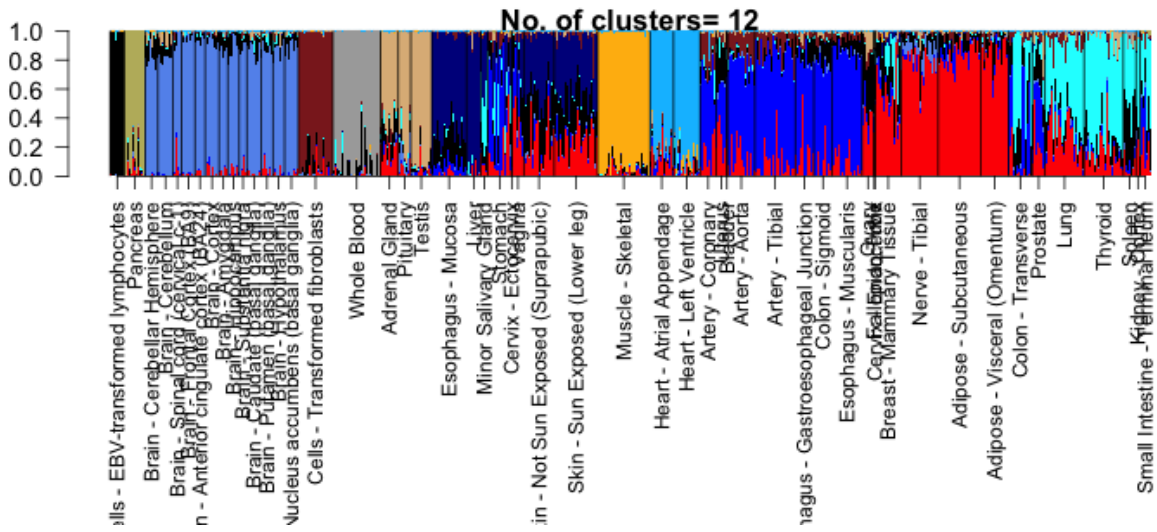


Figure 4. Structure plot of all tissue samples in GTEx version 4 data thinned data with $p_{thin} = 0.0001$ for $K=12$. The thinning parameter has been chosen so that the GTEx RNA-seq data can be interpreted at the same scale as a scRNA-seq data. It seems the results are pretty robust. This suggests that most of the structure in the clustering patterns we observed in textbfFig 1 is retained even after thinning.

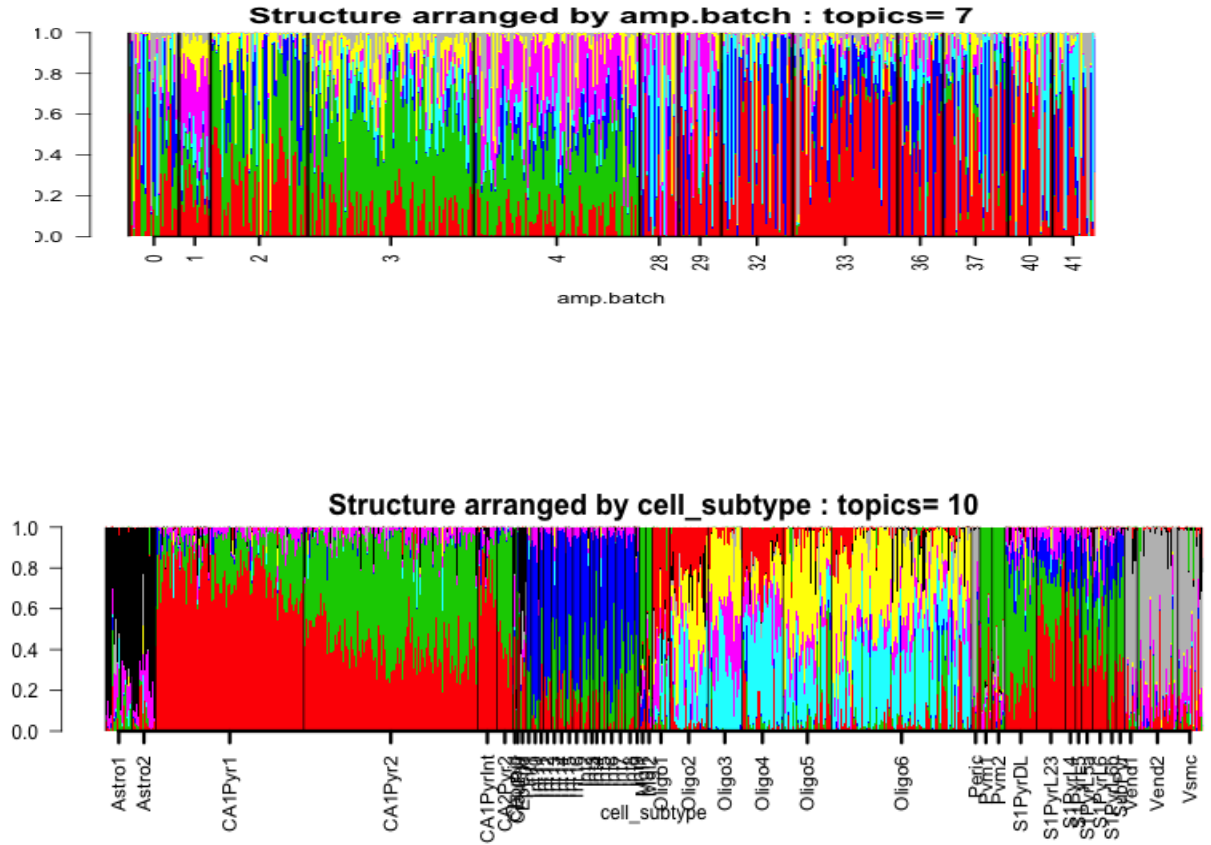
(a) $k = 10$

Figure 5. (*top panel*) Structure plot of the 1041 single cells for $K=7$ of the Jaitin *et al* data [?] arranged by the amplification batch. It is observed that the clustering patterns in each batch are pretty homogeneous and so, either the amplification batch is driving the clustering or it is confounded with the actual biological effects, making it difficult to interpret these clusters. (*bottom panel*) Structure plot of all samples for $K = 10$ of Zeisel *et al* data [7], arranged by the cell subtype labels that were determined by the authors using their BackSpin algorithm and subsequent marker gene annotations. While the admixture patterns in cell subtypes are pretty homogeneous, the first few samples in Oligo6 show mild presence of red cluster and are pretty different from the rest of the samples in Oligo6 which do not show any trace of red cluster. These first few samples of Oligo6 look similar in pattern to some Oligo4 samples with mild red cluster presence. Oligo4 samples also show some heterogeneity in terms of the proportion of red cluster present. This could either be due to misclassification of the Backspin algorithm, or some technical effects.

5.1 Supplemental figures

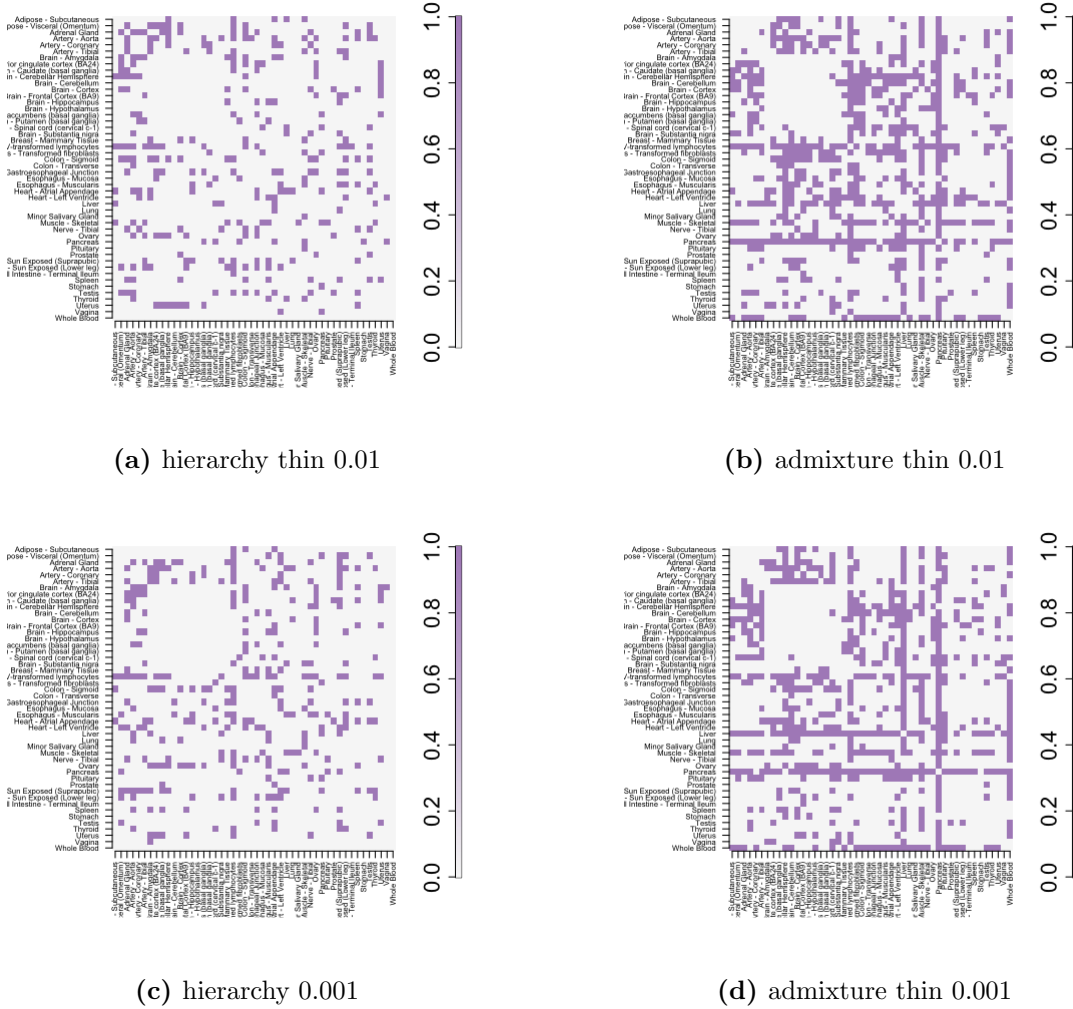


Figure 6. In this graph, we compare the hierarchical clustering method with the admixture method for thinned data with thinning parameters being $p_{thin} = 0.001$ and $p_{thin} = 0.0001$. The color coding scheme is similar to **Fig 3**. Note that the performance of the admixture indeed deteriorates from **Fig 3** in separating out the clusters as is expected. But it still outperforms the hierarchical clustering.

Cluster	Gene names	Proteins	Summary
cluster 1, red (nerve, adrenal)	ENSG00000160882	cytochrome P450, family 11, subfamily B, polypeptide 1	catalyze many reactions involved in drug metabolism and synthesis of cholesterol, steroids and other lipids, mutations cause congenital adrenal hyperplasia due to 11-beta-hydroxylase deficiency.
	ENSG00000148795	cytochrome P450, family 17, subfamily A, polypeptide 1	catalyze many reactions involved in drug metabolism and synthesis of cholesterol, steroids and other lipids, mutations associated with associated with isolated steroid-17 alpha-hydroxylase deficiency, pseudo-hermaphroditism, and adrenal hyperplasia
	ENSG00000158887	myelin protein zero	encodes a major structural protein of peripheral myelin, mutations related to autosomal dominant form of Charcot-Marie-Tooth disease type 1 and other polyneuropathies.
cluster 2, blue (adipose and lung)	ENSG00000168878	surfactant protein B	an amphipathic surfactant protein essential for lung function and homeostasis after birth, mutations cause pulmonary alveolar proteinosis, fatal respiratory distress in the neonatal period.
	ENSG00000168484	surfactant protein C	hydrophobic surfactant protein essential for lung function and homeostasis after birth, associated with pulmonary alveolar proteinosis, interstitial lung disease in older infants, children, and adults.
	ENSG00000185303	surfactant protein A2	encode pulmonary-surfactant associated proteins, mutations associated with idiopathic pulmonary fibrosis.
cluster 3, shallow blue (colon and esophagus)	ENSG00000163017	actin, gamma 2, smooth muscle, enteric	involved in various types of cell motility and maintenance of the cytoskeleton, constituent of the contractile apparatus and muscle tissues.
	ENSG00000133392	myosin, heavy chain 11, smooth muscle	functions as a major contractile protein, chromosomal rearrangement is associated with acute myeloid leukemia of the M4Eo subtype.
	ENSG00000107796	actin, alpha 2, smooth muscle, aorta	play a role in cell motility, structure and integrity, associated with aortic aneurysm familial thoracic type 6.

Cluster	Gene names	Proteins	Summary
cluster 4, black (brain)	ENSG00000259384	growth hormone 1	is expressed in the pituitary, member of the somatotropin/prolactin family of hormones, controls growth, mutations lead to short stature
	ENSG00000132639	synaptosomal-associated protein	involved in the regulation of neurotransmitter release
	ENSG00000115138	proopiomelanocortin	encodes a polypeptide hormone precursor, synthesized mainly in corticotroph cells of the anterior pituitary, hypothalamus, placenta, and epithelium, important for energy homeostasis, melanocyte stimulation, and immune modulation, associated with early onset obesity, adrenal insufficiency, and red hair pigmentation.
cluster 5, light blue (artery)	ENSG00000133392	myosin, heavy chain 11, smooth muscle	major contractile protein, converting chemical energy into mechanical energy through the hydrolysis of ATP
	ENSG00000143248	regulator of G-protein signaling 5	RGS proteins are signal transduction molecules involved in regulation of heterotrimeric G proteins by acting as GTPase activators.
	ENSG00000111341	matrix Gla protein	likely acts as an inhibitor of bone formation, defects causes Keutel syndrome.
cluster 6, deep blue (muscle heart)	ENSG00000143632	actin, alpha 1, skeletal muscle	produces highly conserved proteins that play a role in cell motility, structure and integrity, mutations cause nemaline myopathy type 3, congenital myopathy, diseases leading to muscle fibre defects
	ENSG00000104879	creatine kinase, muscle	protein encoded is cytoplasmic enzyme involved in energy homeostasis and serum marker for myocardial infarction.
	ENSG00000198125	myoglobin	encodes a member of the globin superfamily and is expressed in skeletal and cardiac muscles.
cluster 7, dark brown (brain)	ENSG00000197971	myelin basic protein	major constituent of the myelin sheath of oligodendrocytes and Schwann cells in the nervous system
	ENSG00000131095	glial fibrillary acidic protein	encodes one of the major intermediate filament proteins of mature astrocytes, mutations causes Alexander disease.
	ENSG00000180354	maturin, neural progenitor differentiation regulator homolog (Xenopus)	NA

Cluster	Gene names	Proteins	Summary
cluster 8, shallow yellow (skin stomach)	ENSG00000186395	keratin 10, type I	encodes a member of the type I (acidic) cytokeratin family, mutations associated with epidermolytic hyperkeratosis.
	ENSG00000096088	progastricsin	The protein is a digestive enzyme produced in the stomach, major component of gastric mucosa, associated with gastric cancer, Helicobacter pylori related gastritis.
	ENSG00000182333	lipase, gastric	encodes gastric lipase, responsible for fat digestion and digestion of triglycerides.
cluster 9, yellow (cell EBV)	ENSG00000211896	immunoglobulin heavy constant gamma 1 (G1m marker)	NA
	ENSG00000211893	immunoglobulin heavy constant gamma 2 (G2m marker)	NA
	ENSG00000019582	CD74 molecule, major histocompatibility complex, class II invariant chain	serves as cell surface receptor for the cytokine macrophage migration inhibitory factor (MIF)
cluster 10, grey (thyroid, small intestine)	ENSG00000042832	thyroglobulin	thyroglobulin produced predominantly in thyroid gland, synthesizes thyroxine and triiodothyronine, associated with Graves disease and Hashimoto thyroiditis.
	ENSG00000171195	mucin 7, secreted	encodes a small salivary mucin, aiding in speech, mastication, associated with asthma
	ENSG00000115705	thyroid peroxidase	plays a central role in thyroid gland function, associated with congenital hypothyroidism, congenital goiter, IIA.
cluster 11, cyan cluster (cells fibroblasts)	ENSG00000115414	fibronectin 1	Fibronectin is involved in cell adhesion, embryogenesis, blood coagulation, host defense, and metastasis
	ENSG00000108821	collagen, type I, alpha 1	Mutations in this gene associated with osteogenesis imperfecta types I-IV, Ehlers-Danlos syndrome type and Classical type, Caffey Disease
	ENSG00000164692	collagen, type I, alpha 2	Same as above

Cluster	Gene names	Proteins	Summary
cluster 12, shallow green (Whole blood)	ENSG00000244734	hemoglobin, beta	mutant beta globin causes sickle cell anemia, absence of beta chain/ reduction in beta globin leads to thalassemia
	ENSG00000188536	hemoglobin, alpha 2	deletion of alpha genes may lead to alpha thalassemia
	ENSG00000206172	hemoglobin, alpha 1	deletion of alpha genes may lead to alpha thalassemia
cluster 13, light brown (esophagus mucosa)	ENSG00000171401	keratin 13, type I	keratins are intermediate filament proteins responsible for the structural integrity of epithelial cells
	ENSG00000163209	small proline-rich protein	NA
	ENSG00000143536	cornulin	play a role in the mucosal/epithelial immune response and epidermal differentiation
cluster 14, violet (liver pancreas)	ENSG00000204983	protease, serine 1	secreted by pancreas, associated with pancreatitis
	ENSG00000091704	carboxypeptidase A1	secreted by pancreas, linked to pancreatitis and pancreatic cancer
	ENSG00000169347	glycoprotein 2 (zymogen granule membrane)	secreted from intracellular zymogen granules and associates with the plasma membrane via GPI linkage
cluster 15, salmon (testis)	ENSG00000122304	protamine 2	Protamines are the major DNA-binding proteins in the nucleus of sperm
	ENSG00000175646	protamine 1	NA
	ENSG00000010318	PHD finger protein 7	This gene is expressed in the testis in Sertoli cells but not germ cells, regulates spermatogenesis.

5.2 Supplementary Table 1

Cluster	Gene names	Proteins	Summary
cluster 1, red	ENSG00000160014	calmodulin 3 (phosphorylase kinase, delta)	is a calcium binding protein that plays a role in signaling pathways, cell cycle progression and proliferation.
	ENSG00000127585	F-box and leucine-rich repeat protein 16	Members of the F-box protein family, such as FBXL16, are characterized by an approximately 40-amino acid F-box motif.
	ENSG00000154277	ubiquitin carboxyl-terminal esterase L1	specifically expressed in the neurons and in cells of the diffuse neuroendocrine system. Mutations in this gene may be associated with Parkinson disease.
cluster 2, green	ENSG00000018625	ATPase, Na ⁺ /K ⁺ transporting, alpha 2 polypeptide	responsible for establishing and maintaining the electrochemical gradients of Na and K ions across the plasma membrane, mutations in this gene result in familial basilar or hemiplegic migraines, and in a rare syndrome known as alternating hemiplegia of childhood
	ENSG00000120885	clusterin	protein encoded by this gene is a secreted chaperone that can under some stress conditions also be found in the cell cytosol, also involved in cell death, tumor progression, and neurodegenerative disorders.
	ENSG00000132002	DnaJ (Hsp40) homolog, subfamily B, member 1	encodes a member of the DnaJ or Hsp40 (heat shock protein 40 kD) family of proteins, that stimulates the ATPase activity of Hsp70 heat-shock proteins to promote protein folding and prevent misfolded protein aggregation.
cluster 3, blue	ENSG00000132639	synaptosomal-associated protein, 25kDa	Synaptic vesicle membrane docking and fusion is mediated by SNAREs located on the vesicle membrane (v-SNAREs) and the target membrane (t-SNAREs), involved in the regulation of neurotransmitter release
	ENSG00000111674	enolase 2 (gamma, neuronal)	encodes one of the three enolase isoenzymes found in mammals, is found in mature neurons and cells of neuronal origin
	ENSG00000089199	chromogranin B	encodes a tyrosine-sulfated secretory protein abundant in peptidergic endocrine cells and neurons. This protein may serve as a precursor for regulatory peptides.
cluster 4, cyan	ENSG00000197971	myelin basic protein	protein encoded is a major constituent of the myelin sheath of oligodendrocytes and Schwann cells in the nervous system.
	ENSG00000133392	glial fibrillary acidic protein	encodes major intermediate filament proteins of mature astrocytes, a marker to distinguish astrocytes during development, mutations in this gene cause Alexander disease, a rare disorder of astrocytes in central nervous system
	ENSG00000107796	secreted protein, acidic, cysteine-rich (osteonectin)	encodes a cysteine-rich acidic matrix-associated protein, required for the collagen in bone to become calcified, in extracellular matrix synthesis and cell shape promotion, associated with tumor suppression.