Benchmarking methods for alignment of scRNA-sea and scATAC-seq data

# Introduction

Following technological advances and discoveries with single-cell transcriptomics, new single-cell sequencing strategies are emerging to profile other molecular layers in thousands of cells, from chromatin accessibility, to expression of surface proteins, to methylation. These methods have already been employed to discover patterns of epigenetic heterogeneity in a variety of tissues [ref]. However, at present these methods produce noisier data and at lower throughput. Integration of such datasets with more detailed and comprehensively annotated scRNA-seq datasets can allow denoising of biological signals, guide cell type annotation and disentangle causal relationships between biological layers of information and how these co-determine complex phenotypes.

Methods for simoultaneous profiling of biological layers are starting to emerge [refs], but these are mostly low-throughput and labor intensive. In most cases, molecular profiles will be measured in parallel from cells sampled from the same tissue/cellular population. Consequently, aligning different datasets requires an at least partial correspondence between profiled features across omics. This is true when integrating different scRNA-seq datasets or scRNA-seq data with spatial transcriptomics. For omic types that measure molecular features that are different than genes, data is usually preprocessed to generate a matrix of gene level features e.g. measuring gene activity from ATAC accessibility peaks [<https://www.ncbi.nlm.nih.gov/pubmed/30078726>]. Different integration methods use different inference algorithms for the latent space projection (e.g. canonical correlation analysis, non-negative matrix factorization, variational autoencoders), but all allow the mapping of a gene expression (or other molecular feature) vector x on a latent space vector z, via a vector of feature loadings w. Inspection of loadings can distinguish features that allow alignment across datasets and potentially indicate cell and omic specific contributions to the overall data variation.

* Finding which method works best
* but also comparing different preprocessing strategies

Here we implemented an analysis workflow to optimize the performance of methods to transfer labels inferred from a single-cell gene expression dataset to a single-cell accessibility dataset. We compared three published methods based on robustness and performance on a range of metrics. We then applied our workflow to optimize integration of scRNA-seq and scATAC-seq datasets generated from developing human thymus.

# Results

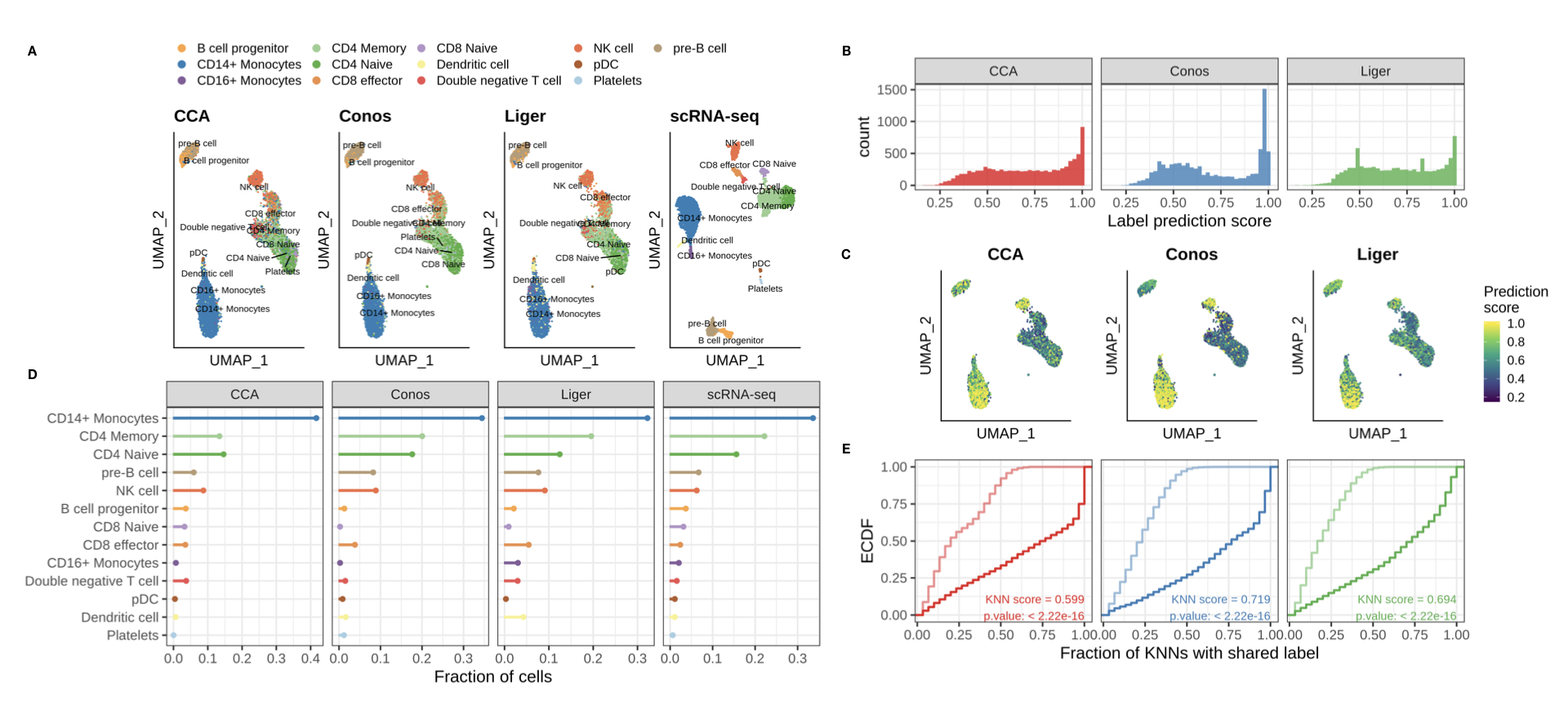
## Designing a bechmark for label transfer methods: PBMC dataset

We run our initial benchmark on a publicly available dataset of Peripheral Blood Mononuclear Cells (PBMC) downloaded from the 10XGenomics website.

We first evaluate the ability of different methods to transfer cell type annotations derived from scRNA-seq data to scATAC-seq data of the same tissue (See Methods 4.5.1 for details about label transfer procedure for each method).

Visualizing the predicted label assignments on the embedding of ATAC cells based on the bin x cells matrix, we find that for all the methods the label assignment have some coincidence with the clusters from ATAC data alone.

All integration methods measure the uncertainty of their assignment (Fig.??A). Setting a threshold on the label prediction score allows to remove from downstream analysis cells with a low confidence prediction label, and mark them as unasssigned. We found that at the cutoff of 0.5, suggested by (**???**), Liger excludes the least amount of cells, while Conos scores the most cells with higher confidence (Fig.??B).

Taking a closer look at predicted label composition and confidence score for each cell type, we found that the different methods called similar fractions of cells for the same labels. On average Liger score with lower confidence smaller cell populations, such as pDCs, platelets, CD8+ naive cells. 

We wanted to evaluate if the predicted annotations maintain the structure found when clustering cells based on bin accessibility alone, i.e. if cells that have a similar bin accessibility profile also get annotated with the same cell type. For each cell we measure the fraction of the K nearest neighbors that share the same predicted label (KNN score). Looking at the distributions of KNN scores, we find that Liger maintains the original connectivity structure better, followed by Conos and finally CCA, even if the differences are not that substantial. Also the KNN score is dependent on the assigned cell type.

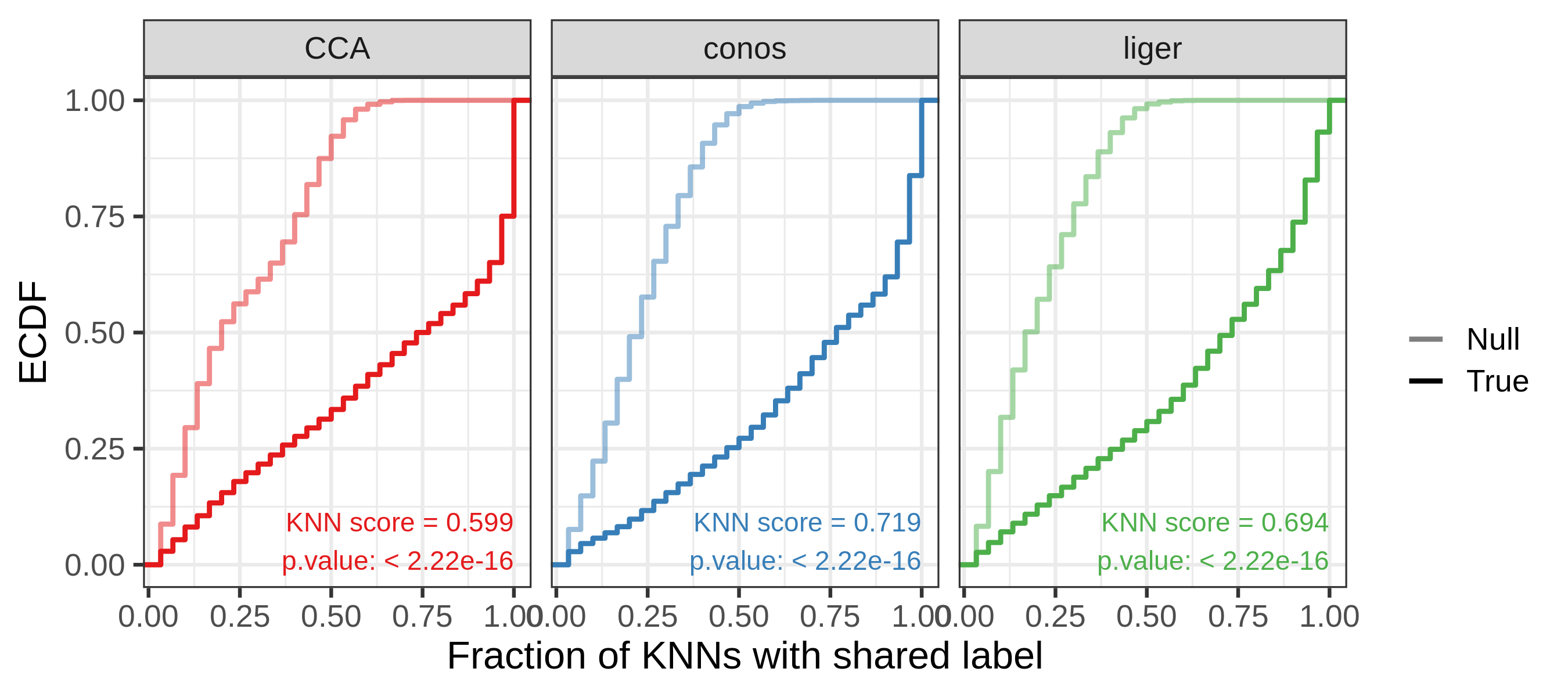


Figure 2: Cumulative distributions of KNN scores

We checked accessibility of PBMC marker genes in the called cell types. We found that cell populations called with CCA show accessibility of expected marker genes from transcriptomics, for NK cells, monocytes, CD8+ cells, and the platelets even.

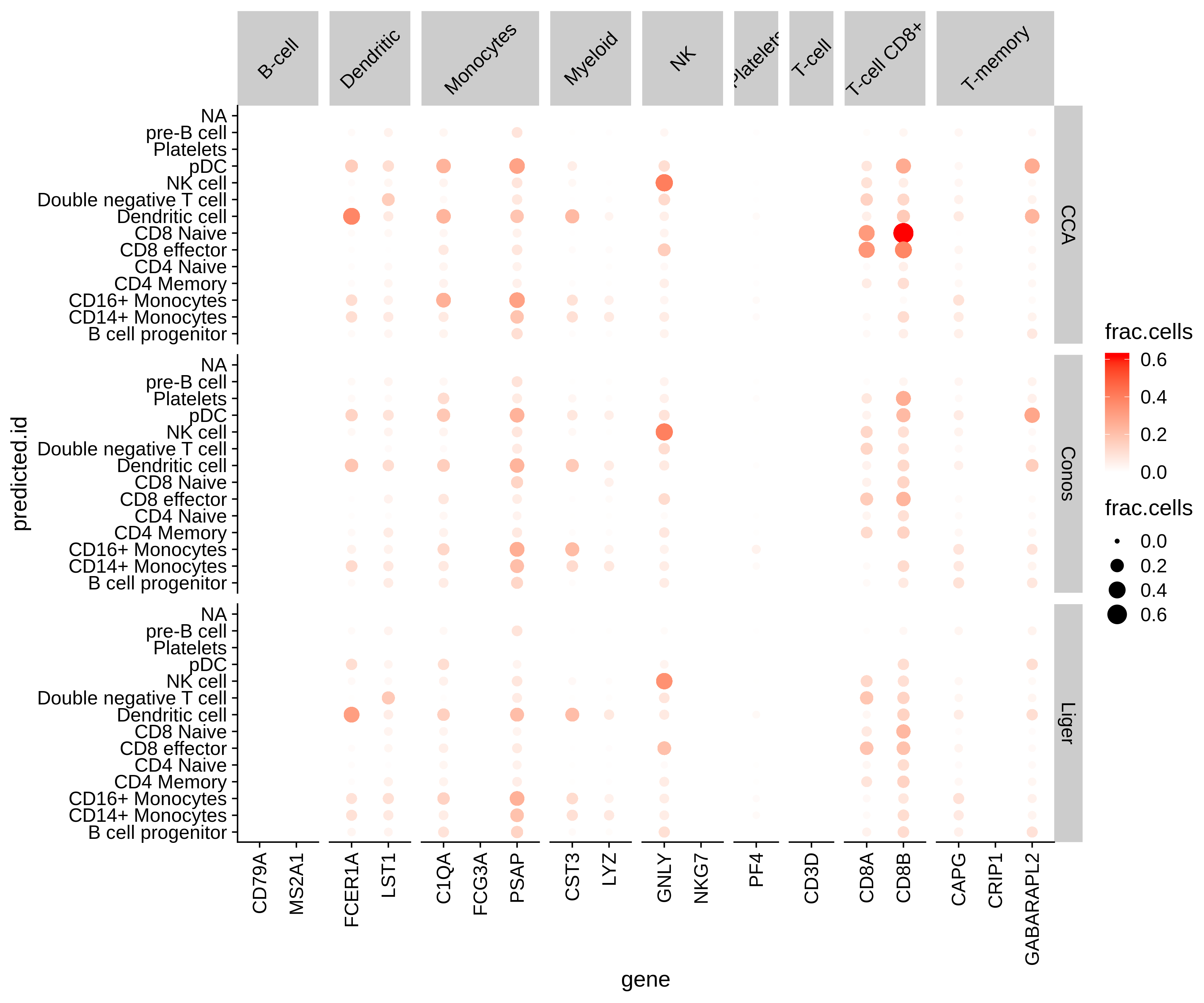


Figure 3: Accessibility of markers in predicted cell clusters

## Optimizing label transfer on Thymus dataset

### Label transfer on count gmat

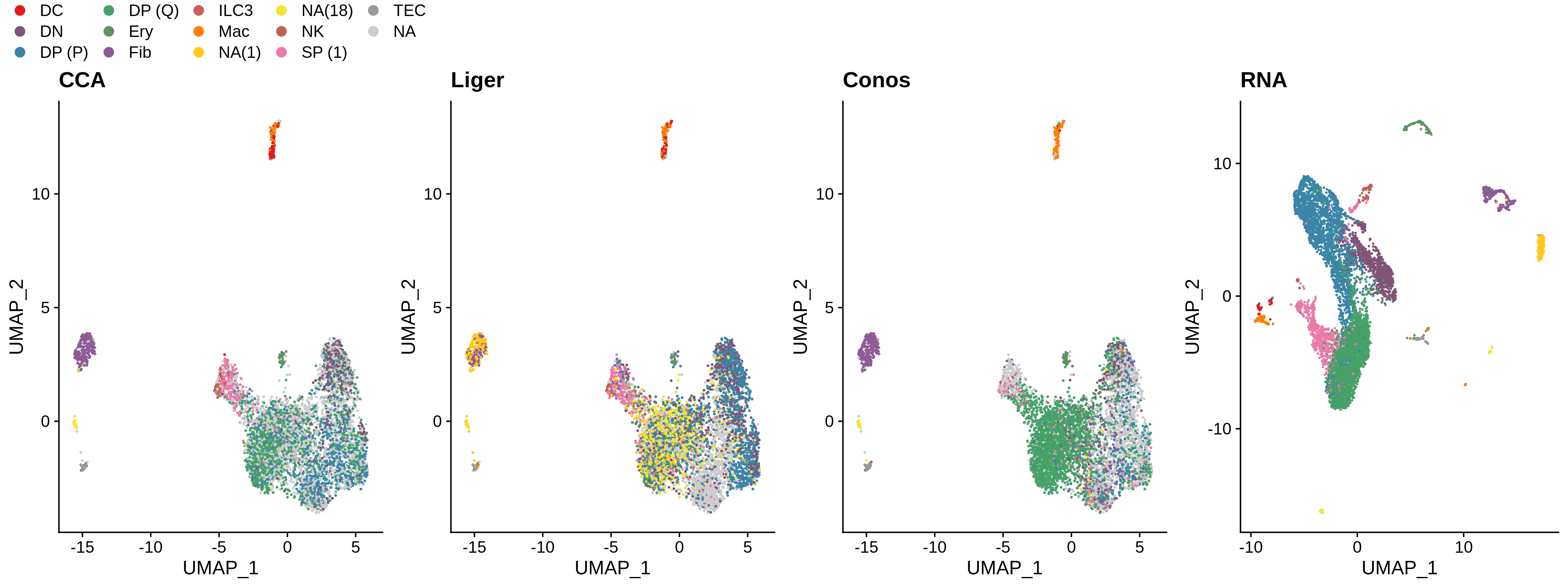
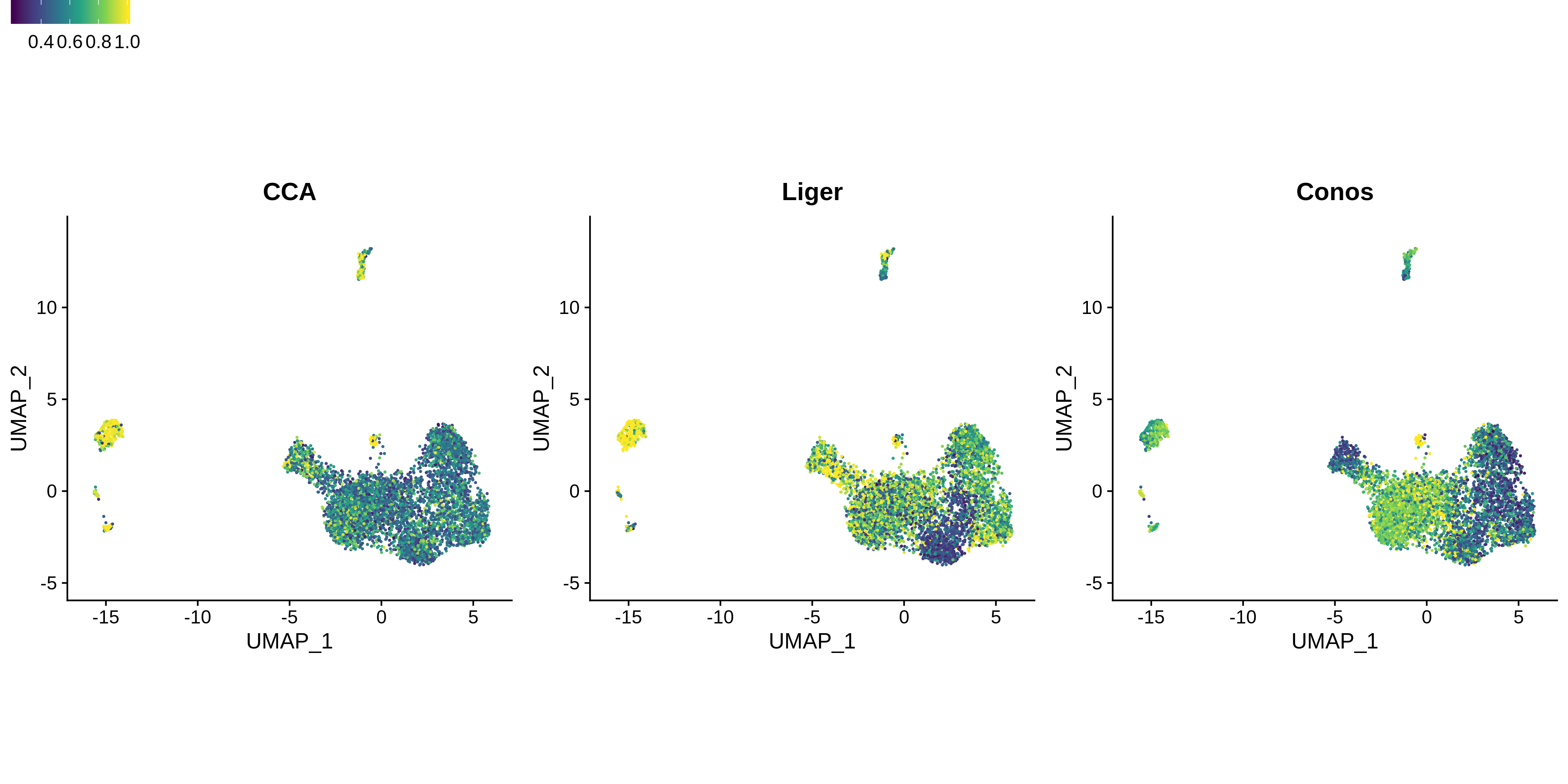
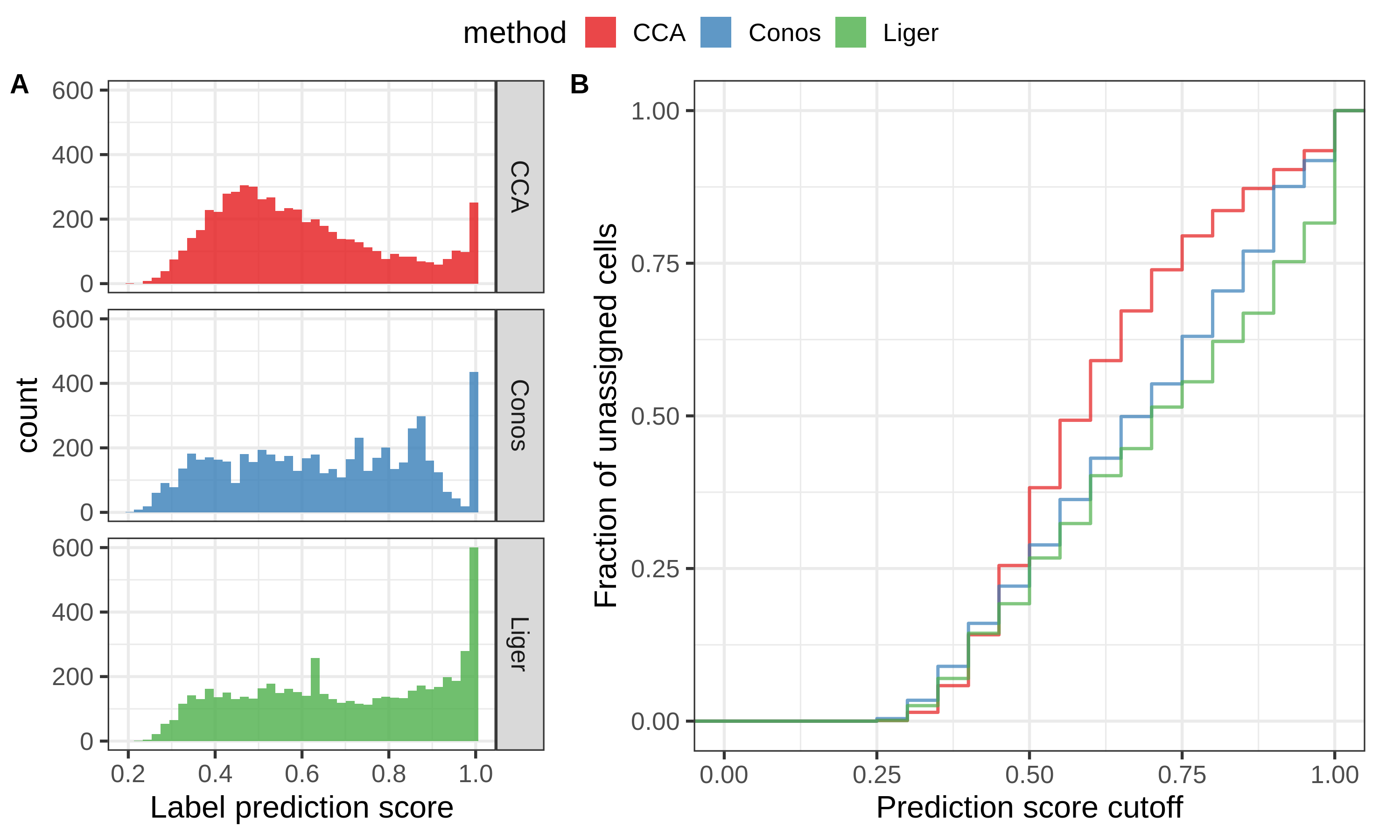


Figure 4: UMAP of predicted labels for thymus dataset



Cell type composition

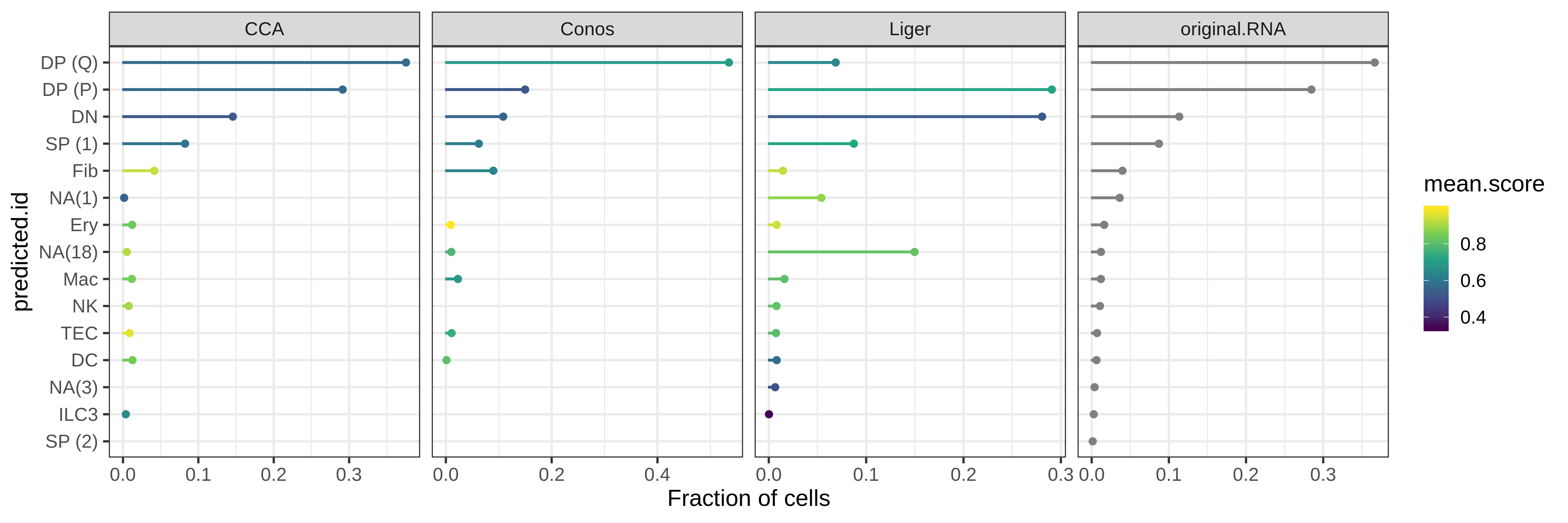


Figure 7: Predicted cell type compositions for thymus dataset

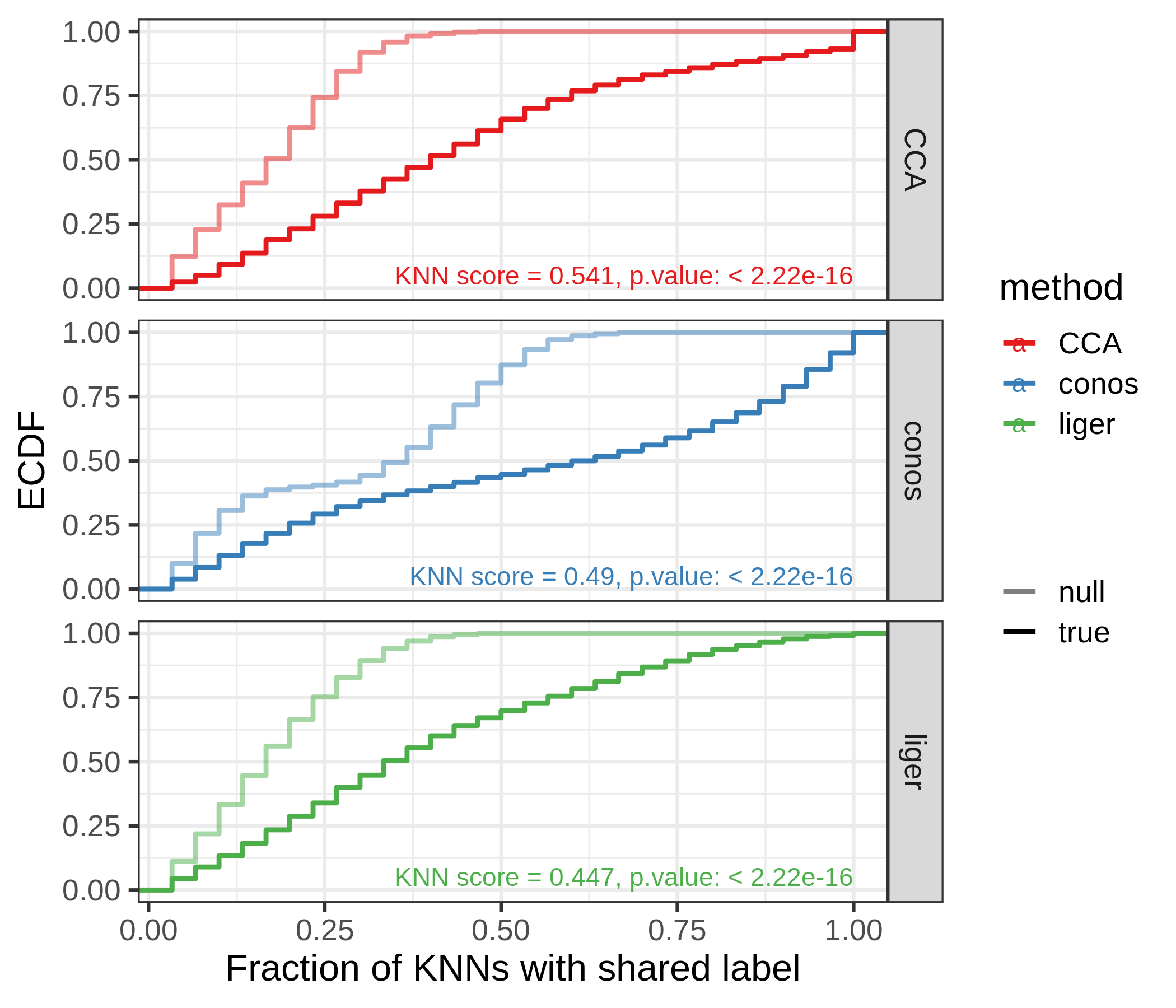
CCA is also better at maintaining the nearest neighboorhood structure from the ATAC alone in its label assignments 



Figure 9: Accessibility of known thymic markers in predicted cell types

Trying to reproduce accessibility profiles of ordered T-cell populations (see Fig. 2H of JP’s manuscript).

* Good: TOX2, ST18, SATB1 on the entry cells, cyclin D genes,
* Weird: CD8 accessibility for CD4 cells (possibly wrongly assigned double positives?)

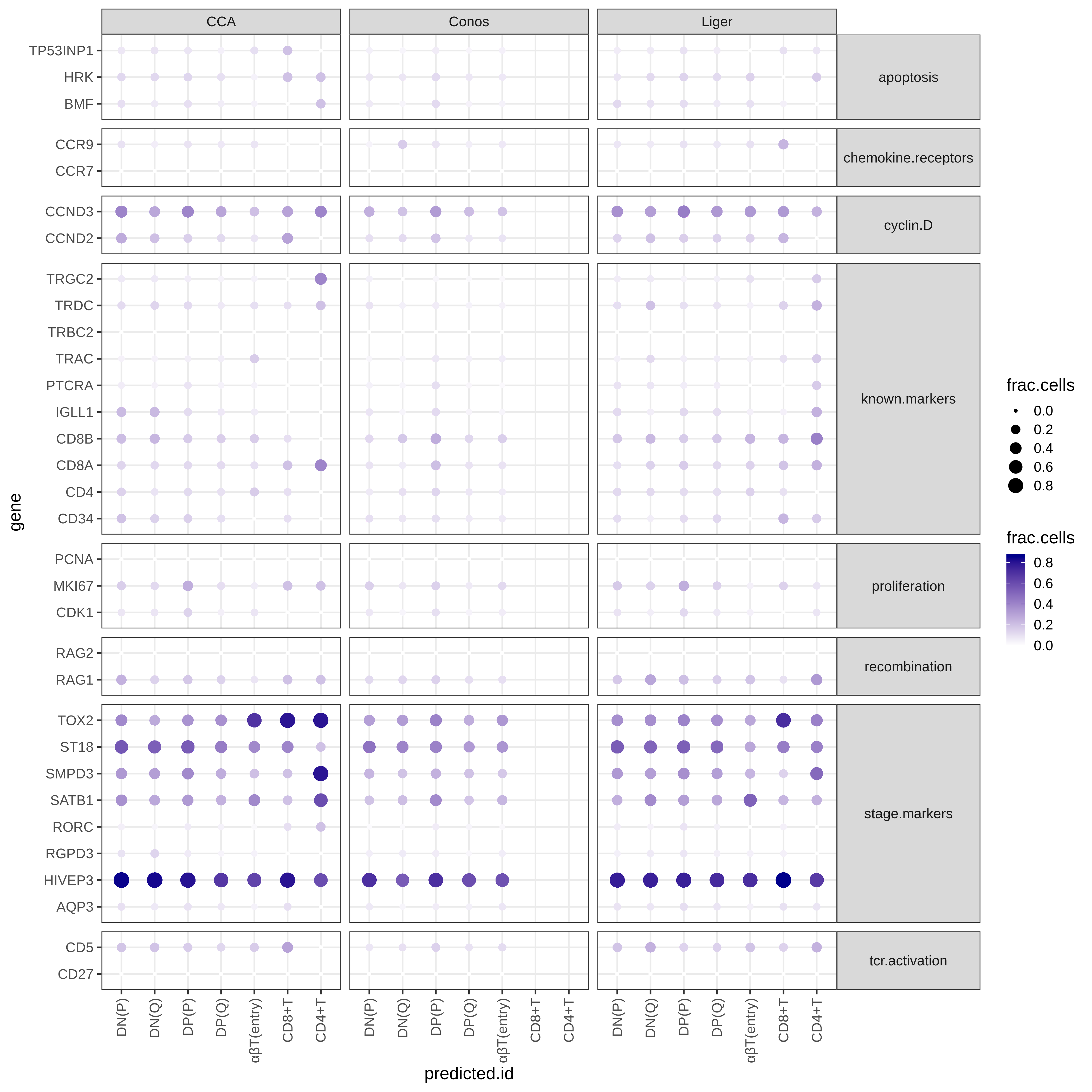


Figure 10: Accessibility of known T-cells markers in predicted cell types

## What next?

* Evaluate label transfer for different cluster sizes
* Label transfer w bigger clusters provided by Cecilia
* MATCHER comparison: CCA seems able to reconstruct the main differences between differentiating T-cells. Will MATCHER find the same relationships between cells?
* Integration w imputation instead of label transfer
* Performance comparison on matched joint profilinf of ATAC - RNA on the same cells

# Discussion

* Does it make a difference how do you reduce ATAC to gene - features? Scoring that takes into account enhancer activity (e.g. Cicero) might be more informative to align differentiating cells.
* Testing performance on datasets from joint profiling protocols

# Methods

## Dataset details

Incl. stats about dataset quality (median no. of fragments per cell, perc. of fragments mapping to peaks, median no. of fragments in peaks per cell)

### PBMC dataset

I used datasets of Peripheral Blood Mononuclear Cells (PBMCs) provided by 10X genomics. Raw scRNA-seq counts were extracted from the R object provided by the Seurat package [cit] (downloaded [here](https://www.dropbox.com/s/3f3p5nxrn5b3y4y/pbmc_10k_v3.rds?dl=1)). Raw scATAC-seq fragments were converted to a snap file using snaptools (downloaded from [here](http://renlab.sdsc.edu/r3fang//share/github/PBMC_ATAC_RNA/atac_pbmc_10k_nextgem.snap)).

### Thymus dataset

I used a dataset of developing thymus at *n* weeks post conception. scATAC-seq and scRNA-seq profiles were measured from tissue samples of the same donor.

## Data preprocessing

**ATAC-seq:**

ScATAC-seq reads were aligned and preprocessed using CellRanger (10X genomics). For preprocessing and quality control of scATAC-seq data we used the SnapATAC pipeline (**???**). Briefly, we selected high-quality cell barcodes based on the number of unique fragments and the ratio of fragments in gene promoters. Then, we binned the genome into fixed-size windows (selected bin size: 5 kb) and generated a bin x cell binary matrix estimating coverage for each bin. We filtered out bins that overlap ENCODE-defined blacklist regions, as well as bins with excidingly high or low z-scored coverage.

We generated cell-by-gene count matrices by aggregating bin coverage over the gene bodies and promoters (2kb upstream of transcriptional start site). We then generated a second cell-by-gene matrix binarizing the count matrix.

**RNA-seq:** cells with high expression of mitochondrial genes were filtered out. Raw counts were normalized by total cell coverage and converted to as a variance stabilizing transformation.

**Feature selection** For label transfer, we select informative genes by taking the union of the 2000 most highly variable genes (HVGs) in scRNA-seq and scATAC-seq datasets (using the Seurat function FindVariableGenes). Other studies have used only the HVGs from the reference dataset (in most cases the scRNA-seq). We found that these two feature selection strategies gave similar label transfer results and we opted for the union to avoid selecting only genes that have very low coverage in the ATAC-seq data.

## Dimensionality reduction and clustering

We used principal component analysis as a low dimensional representation of scRNA-seq datasets. Unless otherwise stated, we used Latent Semantic Indexing for dimensionality reduction of accessibility matrices, as proposed by Cusanovich et al. ([2015](#ref-cusanovichMultiplexSinglecellProfiling2015)). For data visualization, we used the Uniform Manifold Approximation and Projection (UMAP) algorithm (McInnes, Healy, and Melville [2018](#ref-mcinnesUMAPUniformManifold2018)) on low-dimensional data representation.

## scATAC-seq data embedding

We performed dimensionality reduction on the bin x cell accessibility matrix using Latent Semantic Indexing as implemented in Seurat. After selection of signification dimensions, we embedded cells for visualization in 2D using the UMAP algorithm.

## Tested integration methods

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| Method | Reference | Description | Included in benchmark | Reason for Excluding |
| Seurat CCA | Stuart et al. ([2019](#ref-stuartComprehensiveIntegrationSingleCell2019)) | Pairing of cells based on Canonical Correlation analysis | Yes | / |
| LIGER | Welch et al. ([2019](#ref-welchSingleCellMultiomicIntegration2019)) | Joint Non-Negative Matrix factorization | Yes | / |
| Conos | Barkas et al. ([2019](#ref-barkasJointAnalysisHeterogeneous2019)) | Joint neighboorhood graph construction | Yes | / |
| scGen | Lotfollahi, Wolf, and Theis ([2019](#ref-lotfollahiScGenPredictsSinglecell2019)) | Joint embedding with Variational Autoencoder | No | Requires cell type annotation in both datasets |
| totalVI | Gayoso et al. ([2019](#ref-gayosoJointModelRNA2019)) | generative model + variational inference | No | Requires multi-omic data from the same single-cells |
| BBKNN | Pola’nski et al. ([n.d.](#ref-polanskiBBKNNFastBatch)) | Joint neighboorhood graph construction | No | Bad alignment during testing |
| Cusanovich2018 | Cusanovich et al. ([2018](#ref-cusanovichSingleCellAtlasVivo2018)) | Joint neighboorhood graph construction | No | Code unavailable |
| gimVI | Lopez et al. ([2019](#ref-lopezJointModelUnpaired2019)) | joint generative model + variational inference | No | No implementation for right log-likelihood distribution |

### Label transfer

One of the key tasks for integration methods is to be able to transfer cell type annotations learnt from a reference to a query dataset. This is especially useful if the query is a scATAC-seq dataset, where calling of cell types based on prior knownledge on marker genes is often not possible. Different models are adapted to transfer discrete cell state labels derived from gene expression to cells measured with scATAC-seq.

##### Seurat CCA

Identified anchor pairs are weighted based on the query cell local neighboorhood (the k nearest anchors) and the anchor score. The obtained reference cells x query cells weight matrix is then multiplied by the matrix of annotation x reference cells, to generate a query cell x annotation matrix. This returns a prediction score for each class for every cell in the query dataset, ranging from 0 to 1 (**???**).

##### LIGER

While the authors do not describe a method for transferring discrete labels, I adapted their strategy for feature imputation. I build a cross-dataset KNN graph in the aligned factor space, then I assign each query cell to the most abundant label between the k nearest neighbors in the reference dataset (k=30). The prediction score for label is computed as the fraction of nearest neighbors that have the predicted label.

##### Conos

Label transfer is treated as a general problem of information propagation between vertices of the common graph (detailed in Barkas et al. ([2019](#ref-barkasJointAnalysisHeterogeneous2019))). The label score is the label probability updating during the diffusion process.

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