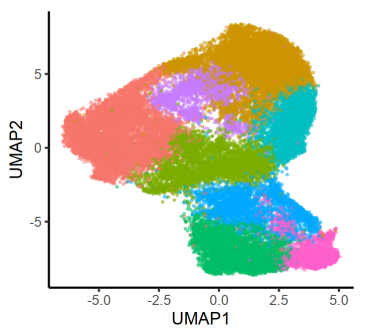
PBMC vs Tissue Figures

```
In [1]: source('jupyterFunctions_perCellType.R')
In [2]: data_prefix <- paste(sep='','../data/PBMCvsTissue/')
    Tcell_PvT <- readRDS(paste(sep='',data_prefix,'Tcell_PBMCvsTissue_df.rd s'))
    Bcell_PvT <- readRDS(paste(sep='',data_prefix,'Bcell_PBMCvsTissue_df.rd s'))
    myeloid_PvT <- readRDS(paste(sep='',data_prefix,'myeloid_PBMCvsTissue_d f.rds'))
In [3]: tissue_color <- 'slateblue3'
    PBMC_color <- 'brown3'</pre>
In [24]: save_dir <- NA #'../output/' #or NA if don't want to save
```

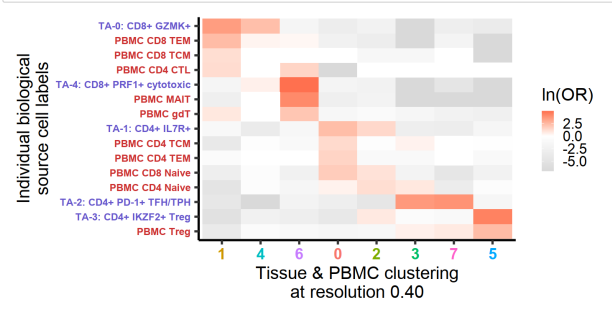
T cell

In [9]: #Fig S7a left Tcell_cluster_colors <- hue pal()(length(unique(Tcell_PvT[,Tcell_res])))</pre> names(Tcell_cluster_colors) <- sort(unique(Tcell_PvT[,Tcell_res]))</pre> options(repr.plot.height=6,repr.plot.width=6) g <- ggplot(Tcell_PvT[which(!(Tcell_PvT\$bio_src_cellType %in% Tcell_tooS mall)),], aes_string(x='UMAP1',y='UMAP2',color=Tcell_res)) + geom_point(size=1,alpha=0.5) + theme_classic(base_size=20) + sca le color manual(values=Tcell cluster colors) + theme(legend.position="none") + ggtitle('Tissue & PBMC T cells') + theme(plot.title = element_te xt(hjust = 0.5))print(g) if(!is.na(save_dir)) ggsave(file=paste(sep='',save_dir,'Tcell_PBMCvsTiss ue_UMAP.png'), plot=g,units='in',height=6,width=6,dpi=600)

Tissue & PBMC T cells



```
In [10]: #Fig 7a right
         Tcell fisher_df <- calc OR(Tcell PvT[which(!(Tcell PvT$bio src cellType
         %in% Tcell_tooSmall)),],
                                      Tcell_res, 'bio_src_cellType')
         g <- plot OR(Tcell fisher df, Tcell res, 'bio src cellType',
                       paste('Tissue & PBMC clustering\nat resolution',str split f
         ixed(Tcell_res,'_',2)[,2]),
                       'Individual biological\nsource cell labels',
                       Tcell PvT order, Tcell original order, clustColors=c(Tcell c
         olors, Tcell_cluster_colors))
         g <- g + theme(axis.text.x=element_text(size=22))</pre>
         options(repr.plot.height=6,repr.plot.width=12)
         print(g)
         if(!is.na(save dir)) ggsave(file=paste(sep='',save dir,save dir,'Tcell P
         BMCvsTissue OR heatmap.png'),
                                      plot=q,units='in',height=6,width=12,dpi=600)
```



Myeloid

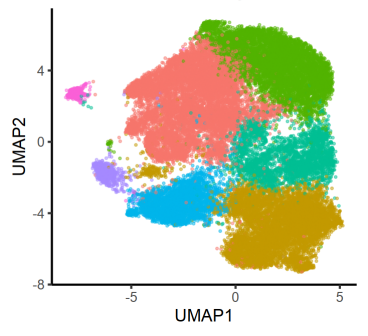
```
In [11]: myeloid_res <- 'hres_0.20'
    if(!(myeloid_res %in% colnames(myeloid_PvT))) stop('cluster resolution n
    ot in df')</pre>
```

```
myeloid_tissue_clusters <- sort(unique(myeloid_PvT[which(myeloid_PvT$bio</pre>
In [12]:
          src=='tissue'),'bio_src_cellType']))
         myeloid PBMC clusters <- sort(unique(myeloid PvT[which(myeloid PvT$bio s
         rc=='PBMC'),'bio_src_cellType']))
         myeloid colors <- c(rep(tissue color,length(myeloid tissue clusters)),re</pre>
         p(PBMC color,length(myeloid PBMC clusters)))
         names(myeloid colors) <- c(myeloid tissue clusters, myeloid PBMC cluster</pre>
         s)
In [13]: | 11 <- table(myeloid_PvT[which(myeloid_PvT$bio_src=='PBMC'),'bio_src_cell</pre>
         Type'])
         myeloid_tooSmall <- names(ll[ll<10])</pre>
In [14]: myeloid original order <- c('MA-0: F13A1+ MARCKS+ TRM', 'PBMC CD16 Mon
         o', 'PBMC CD14 Mono',
                                       'MA-1: FCN1+ SAMSN1+ infiltrating monocyte
         s',
                                       'MA-2: LYVE1+ TIMD4+ TRM',
                                       'MA-4: SPP1+ FABP5+ intermediate',
                                       'MA-3: CD1C+ AFF3+ DC', 'PBMC cDC1', 'PBMC cDC
          2', 'PBMC pDC')
         if(!all(myeloid original order %in% unique(myeloid PvT$bio src cellTyp
         e))) stop('original order insufficient')
         myeloid PvT order <-c(0,6,1,2,3,4,5)
         if(!all(myeloid PvT order %in% unique(myeloid PvT[,myeloid res]))) stop
```

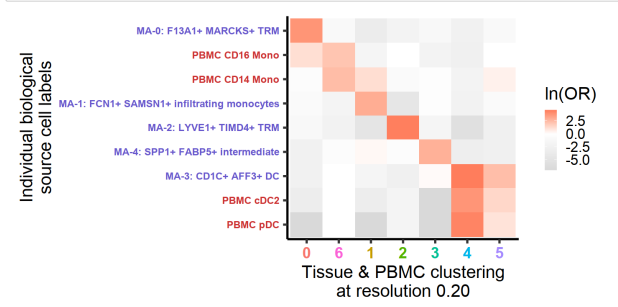
('cluster order insufficient')

```
In [15]: #Fig S7b left
         myeloid cluster_colors <- hue pal()(length(unique(myeloid PvT[,myeloid r</pre>
         names(myeloid_cluster_colors) <- sort(unique(myeloid_PvT[,myeloid_res]))</pre>
         options(repr.plot.height=6,repr.plot.width=6)
         g <- ggplot(myeloid PvT[which(!(myeloid PvT$bio src_cellType %in% myeloi
         d_tooSmall)),],
                     aes_string(x='UMAP1',y='UMAP2',color=myeloid_res)) +
                 geom point(size=1,alpha=0.5) + theme classic(base size=20) + sca
         le color manual(values=myeloid cluster colors) +
                 theme(legend.position="none") +
                 ggtitle('Tissue & PBMC myeloid cells') + theme(plot.title = elem
         ent_text(hjust = 0.5))
         print(g)
         if(!is.na(save dir)) ggsave(file=paste(sep='',save dir,'myeloid PBMCvsTi
         ssue UMAP.png'),
                                      plot=q,units='in',height=6,width=6,dpi=600)
```

Tissue & PBMC myeloid cells



```
In [16]: #Fig 7b right
         myeloid fisher_df <- calc_OR(myeloid PvT[which(!(myeloid PvT$bio src_cel
         lType %in% myeloid_tooSmall)),],
                                      myeloid res, 'bio src cellType')
         g <- plot OR(myeloid fisher df, myeloid res, 'bio src cellType',
                      paste('Tissue & PBMC clustering\nat resolution',str split f
         ixed(myeloid_res,'_',2)[,2]),
         'Individual biological\nsource cell labels',
                       myeloid PvT_order, myeloid original order,clustColors=c(mye
         loid colors, myeloid cluster colors))
         g <- g + theme(axis.text.x=element_text(size=22))</pre>
         options(repr.plot.height=6,repr.plot.width=12)
         print(g)
         if(!is.na(save_dir)) ggsave(file=paste(sep='',save_dir,save_dir,'myeloid
         _PBMCvsTissue_OR_heatmap.png'),
                                      plot=q,units='in',height=6,width=12,dpi=600)
```



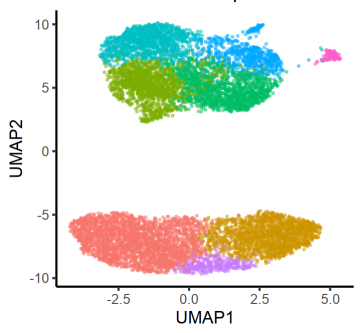
B cell

```
In [17]: Bcell_res <- 'hres_0.60'
    if(!(Bcell_res %in% colnames(Bcell_PvT))) stop('cluster resolution not i
        n df')</pre>
```

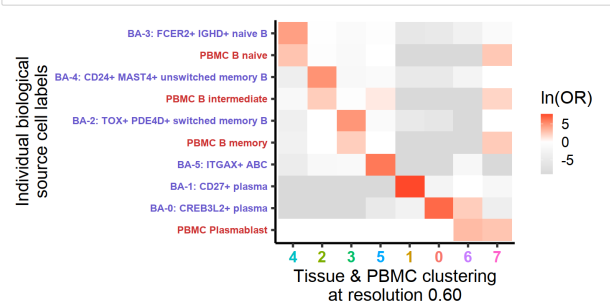
```
In [18]: Bcell tissue_clusters <- sort(unique(Bcell_PvT[which(Bcell_PvT$bio_src=</pre>
          ='tissue'), 'bio src cellType']))
          Bcell PBMC clusters <- sort(unique(Bcell PvT[which(Bcell PvT$bio src=='P</pre>
          BMC'),'bio_src_cellType']))
          Bcell_colors <- c(rep(tissue_color,length(Bcell_tissue_clusters)),rep(PB</pre>
         MC_color,length(Bcell_PBMC_clusters)))
          names(Bcell colors) <- c(Bcell tissue clusters, Bcell PBMC clusters)</pre>
In [19]:
         11 <- table(Bcell_PvT[which(Bcell_PvT$bio_src=='PBMC'),'bio_src_cellTyp</pre>
          e'])
         Bcell_tooSmall <- names(ll[ll<10])</pre>
In [20]: Bcell_original_order <- c('BA-3: FCER2+ IGHD+ naive B','PBMC B naive',</pre>
                                     'BA-4: CD24+ MAST4+ unswitched memory B', 'PBMC
         B intermediate',
                                     'BA-2: TOX+ PDE4D+ switched memory B', 'PBMC B
         memory',
                                     'BA-5: ITGAX+ ABC',
                                     'BA-1: CD27+ plasma', 'BA-0: CREB3L2+ plasm
          a', 'PBMC Plasmablast')
          if(!all(Bcell_original_order %in% unique(Bcell_PvT$bio_src_cellType))) s
          top('original order insufficient')
         Bcell_PvT_order <- c(4,2,3,5,1,0,6,7)
          if(!all(Bcell PvT order %in% unique(Bcell PvT[,Bcell res]))) stop('clust
          er order insufficient')
```

In [21]: #Fig S7c left Bcell_cluster_colors <- hue pal()(length(unique(Bcell_PvT[,Bcell_res])))</pre> names(Bcell_cluster_colors) <- sort(unique(Bcell_PvT[,Bcell_res]))</pre> options(repr.plot.height=6,repr.plot.width=6) g <- ggplot(Bcell_PvT[which(!(Bcell_PvT\$bio_src_cellType %in% Bcell_tooS mall)),], aes_string(x='UMAP1',y='UMAP2',color=Bcell_res)) + geom_point(size=1,alpha=0.5) + theme_classic(base_size=20) + sca le color manual(values=Bcell cluster colors) + theme(legend.position="none") + ggtitle('Tissue & PBMC B/plasma cells') + theme(plot.title = ele ment_text(hjust = 0.5)) print(g) if(!is.na(save_dir)) ggsave(file=paste(sep='',save_dir,'Bcell_PBMCvsTiss ue_UMAP.png'), plot=g,units='in',height=6,width=6,dpi=600)

Tissue & PBMC B/plasma cells



```
In [22]: #Fig 7c right
         Bcell fisher df <- calc OR(Bcell PvT[which(!(Bcell PvT$bio src cellType
         %in% Bcell_tooSmall)),],
                                      Bcell_res, 'bio_src_cellType')
         g <- plot OR(Bcell fisher df, Bcell res, 'bio src cellType',
                       paste('Tissue & PBMC clustering\nat resolution',str split f
         ixed(Bcell_res, '_',2)[,2]),
                       'Individual biological\nsource cell labels',
                       Bcell PvT order, Bcell original order, clustColors=c(Bcell c
         olors, Bcell_cluster_colors))
         g <- g + theme(axis.text.x=element_text(size=22))</pre>
         options(repr.plot.height=6,repr.plot.width=12)
         print(g)
         if(!is.na(save dir)) ggsave(file=paste(sep='',save dir,save dir,'Bcell P
         BMCvsTissue OR heatmap.png'),
                                      plot=q,units='in',height=6,width=12,dpi=600)
```



Session Info

```
In [23]:
         sessionInfo()
         R version 3.6.1 (2019-07-05)
         Platform: x86_64-conda_cos6-linux-gnu (64-bit)
         Running under: Red Hat Enterprise Linux Server release 6.5 (Santiago)
         Matrix products: default
         BLAS/LAPACK: /PHShome/kew47/miniconda3/lib/R/lib/libRblas.so
         locale:
         [1] en_US.UTF-8
         attached base packages:
         [1] grid
                                  graphics grDevices utils
                                                                 datasets methods
                        stats
         [8] base
         other attached packages:
                                                   scales_1.1.1
                                                                      viridis_0.5.
          [1] repr_1.0.1
                                 gridExtra_2.3
         1
          [5] viridisLite_0.3.0 ggrepel_0.8.2
                                                   ggrastr_0.2.3
                                                                      ggplot2_3.3.
          [9] tidyr_1.0.3
                                 stringr 1.4.0
                                                   ROCR 1.0-7
                                                                      gplots 3.0.
         1.1
                                 plyr_1.8.6
                                                                      gtools 3.8.2
         [13] Rmisc 1.5.1
                                                   lattice 0.20-41
         [17] Matrix 1.2-18
         loaded via a namespace (and not attached):
          [1] Rcpp 1.0.4.6
                                   vipor 0.4.5
                                                       pillar 1.4.4
          [4] compiler 3.6.1
                                   bitops 1.0-6
                                                        base64enc 0.1-3
          [7] tools 3.6.1
                                   digest_0.6.25
                                                        uuid 0.1-2
         [10] gtable 0.3.0
                                   jsonlite 1.7.1
                                                        evaluate 0.14
         [13] lifecycle 0.2.0
                                   tibble 3.0.1
                                                        pkgconfig 2.0.3
         [16] rlang 0.4.8
                                   IRdisplay_0.7.0
                                                        IRkernel 1.0.2.9000
         [19] beeswarm 0.2.3
                                   withr 2.2.0
                                                       dplyr 1.0.2
         [22] generics 0.0.2
                                   vctrs 0.3.5
                                                       caTools 1.18.0
         [25] tidyselect 1.1.0
                                   glue_1.4.0
                                                       R6 2.4.1
         [28] ggbeeswarm 0.6.0
                                   gdata 2.18.0
                                                       pbdZMQ 0.3-3
         [31] farver 2.0.3
                                   purrr 0.3.4
                                                       magrittr 1.5
         [34] htmltools 0.4.0
                                   ellipsis 0.3.1
                                                       colorspace 1.4-1
         [37] labeling 0.3
                                   KernSmooth 2.23-15
                                                       stringi 1.4.6
         [40] munsell 0.5.0
                                   crayon 1.3.4
                                                       Cairo 1.5-10
```

In []: