Application of VAM to Seurat pbmc\_small scRNA-seq data using Seurat log normalization.

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### 1 Load the VAM package

> library(VAM)

## 2 Summary statistics for the pbmc\_small scRNA-seq data

This example uses the pbmc\_small data set included in the SeuratObject package and a single contrived gene set. Please see the other vignettes for more realistic examples using larger scRNA-seq data sets and gene set collections based on MSigDB.

# 3 Define gene set collection

A gene set collection containing just a single contrived set (containing the top 5 variable genes) will be used for this example.

```
message("Seurat package not available! Not executing associated vignette content.")
+ }
   "PPBP"
             "IGLL5"
                      "VDAC3"
                               "CD1C"
                                         "AKR1C3"
```

### Execute VAM method 4

Since the scRNA-seq data has been processed using Seurat, we execute VAM using the vamForSeurat() function. We have set return.dist=T so that the squared adjusted Mahalanobis distances will be returned in a "VAMdist" Assay.

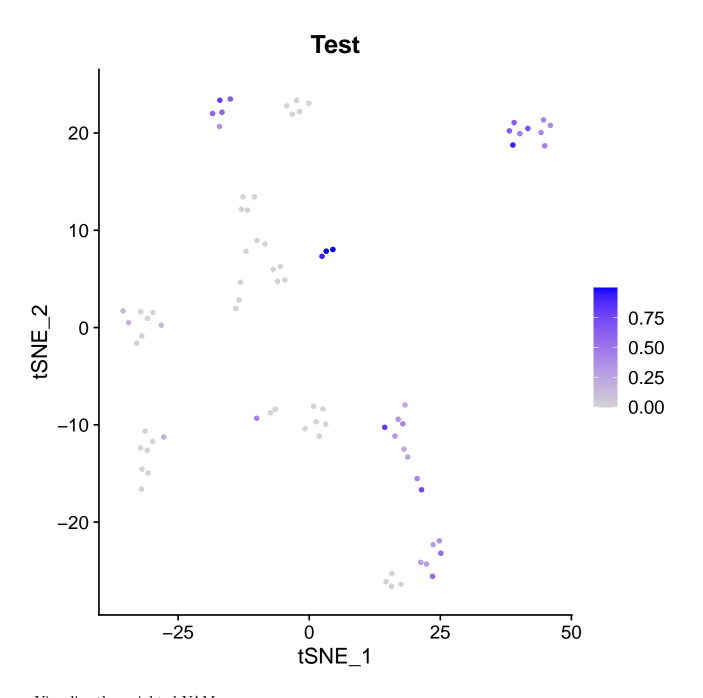
```
> if (requireNamespace("Seurat", quietly=TRUE)) {
          pbmc.vam = vamForSeurat(seurat.data=SeuratObject::pbmc_small,
              gene.set.collection=gene.set.collection,
              center=F, gamma=T, sample.cov=F, return.dist=T)
+ } else {
          message("Seurat package not available! Not executing associated vignette content.")
+ }
  Look at the first few entries in the "VAMdist" and "VAMcdf" Assays.
> if (requireNamespace("Seurat", quietly=TRUE)) {
          pbmc.vam@assays$VAMdist@data[1,1:10]
          pbmc.vam@assays$VAMcdf@data[1,1:10]
+ } else {
          message("Seurat package not available! Not executing associated vignette content.")
+ }
ATGCCAGAACGACT CATGGCCTGTGCAT GAACCTGATGAACC TGACTGGATTCTCA AGTCAGACTGCACA
     0.0000000
                    0.0000000
                                    0.1193809
                                                   0.3040717
                                                                   0.000000
TCTGATACACGTGT TGGTATCTAAACAG GCAGCTCTGTTTCT GATATAACACGCAT AATGTTGACAGTCA
     0.0000000
                                    0.0000000
                                                   0.000000
                    0.4182978
                                                                   0.3431001
```

Create gene weights that prioritize the first two genes in the set and execute VAM using the weights.

```
> if (requireNamespace("Seurat", quietly=TRUE)) {
+
          gene.weights = list(c(2,2,1,1,1))
          pbmc.vam.weights = vamForSeurat(seurat.data=SeuratObject::pbmc_small,
              gene.weights=gene.weights,
              gene.set.collection=gene.set.collection,
              center=F, gamma=T, sample.cov=F, return.dist=T)
+ } else {
          message("Seurat package not available! Not executing associated vignette content.")
+ }
```

#### Visualize VAM scores 5

Visualize VAM scores using Seurat FeaturePlot(). The default Assay must first be changed to "VAMcdf'.



Visualize the weighted VAM scores.

