

# 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.

```
> if (requireNamespace("Seurat", quietly=TRUE)) {  
+     SeuratObject::pbmc_small  
+     gene.names = rownames(SeuratObject::pbmc_small)  
+     gene.names[1:5]  
+     Seurat::VariableFeatures(SeuratObject::pbmc_small)[1:5]  
+ } else {  
+     message("Seurat package not available! Not executing associated vignette content.")  
+ }  
  
[1] "PPBP"    "IGLL5"   "VDAC3"   "CD1C"    "AKR1C3"
```

## 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.

```
> if (requireNamespace("Seurat", quietly=TRUE)) {  
+     gene.set.name = "Test"  
+     gene.ids = c("PPBP", "IGLL5", "VDAC3", "CD1C", "AKR1C3")  
+     # Create a collection list for this gene set  
+     gene.set.id.list = list()  
+     gene.set.id.list[[1]] = gene.ids  
+     names(gene.set.id.list)[1] = gene.set.name  
+     gene.set.id.list  
+     # Create the list of gene indices required by vamForSeurat()  
+     (gene.set.collection = createGeneSetCollection(gene.ids=gene.names,  
+             gene.set.collection=gene.set.id.list))  
+     gene.indices = gene.set.collection[[1]]  
+     (gene.names = gene.names[gene.indices])  
+ } else {
```

```

+           message("Seurat package not available! Not executing associated vignette content.")
+ }

[1] "PPBP"    "IGLL5"    "VDAC3"    "CD1C"     "AKR1C3"

```

## 4 Execute VAM method

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	0.0000000
TCTGATACACGTGT	TGGTATCTAACAG	GCAGCTCTGTTCT	GATATAAACACGGCAT	AATGTTGACAGTCA
0.0000000	0.4182978	0.0000000	0.0000000	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.")
+ }

```

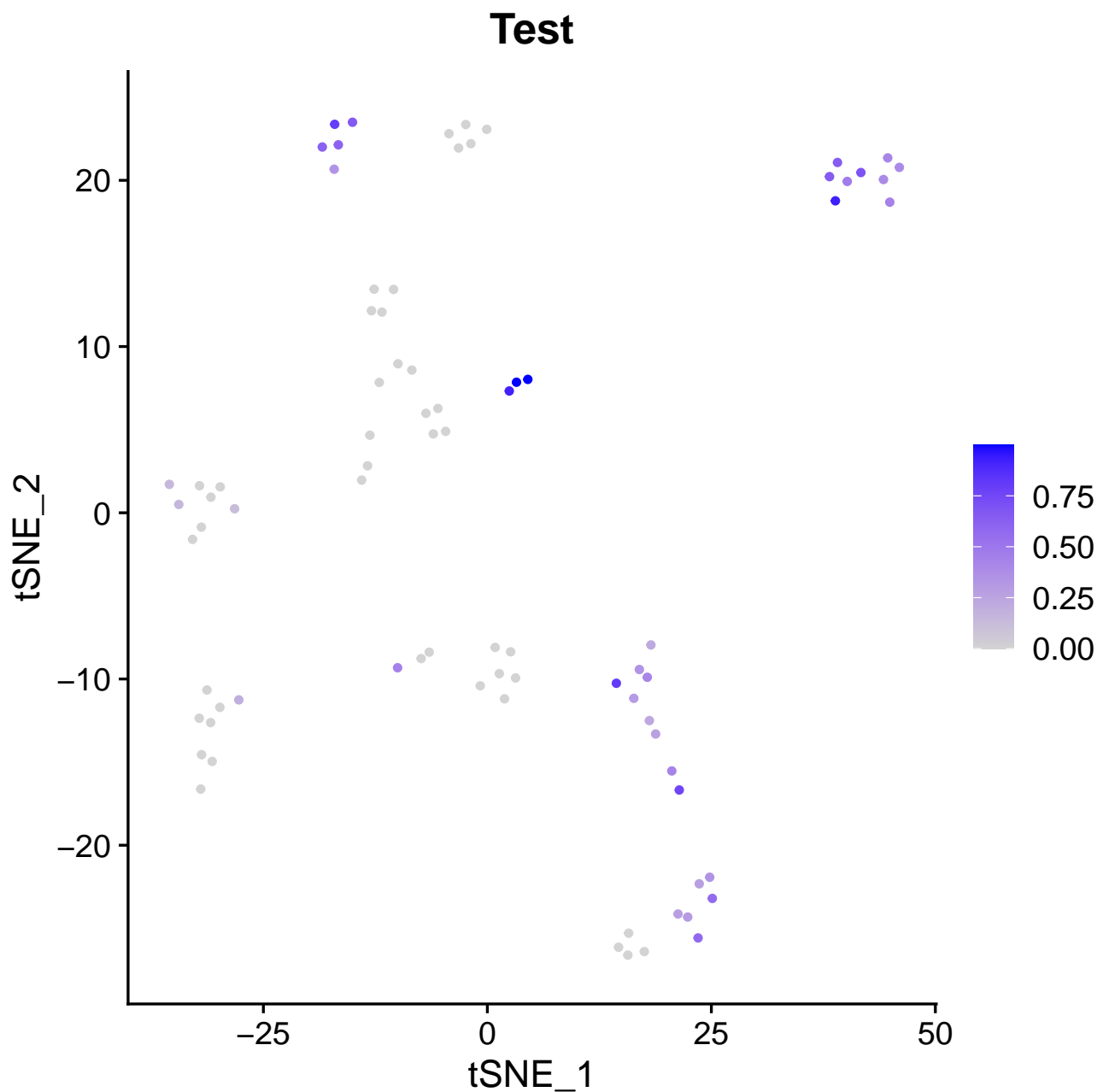
## 5 Visualize VAM scores

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

```

> if (requireNamespace("Seurat", quietly=TRUE)) {
+     Seurat::DefaultAssay(object = pbmc.vam) = "VAMcdf"
+     Seurat::FeaturePlot(pbmc.vam, reduction="tsne", features=gene.set.name)
+ } else {
+     message("Seurat package not available! Not executing associated vignette content.")
+     par(mar = c(0,0,0,0))
+     plot(c(0, 1), c(0, 1), ann = F, bty = 'n', type = 'n', xaxt = 'n', yaxt = 'n')
+     text(x = 0.5, y = 0.5,paste("Seurat package not available!\n",
+                                   "FeaturePlot not generated."),
+          cex = 1.6, col = "black")
+ }

```



Visualize the weighted VAM scores.

```

> if (requireNamespace("Seurat", quietly=TRUE)) {
+     Seurat::DefaultAssay(object = pbmc.vam.weights) = "VAMcdf"
+     Seurat::FeaturePlot(pbmc.vam.weights, reduction="tsne", features=gene.set.name)
+ } else {
+     message("Seurat package not available! Not executing associated vignette content.")
+     par(mar = c(0,0,0,0))
+     plot(c(0, 1), c(0, 1), ann = F, bty = 'n', type = 'n', xaxt = 'n', yaxt = 'n')
+     text(x = 0.5, y = 0.5,paste("Seurat package not available!\n",
+                                   "FeaturePlot not generated."),
+          cex = 1.6, col = "black")
+ }

```

