Eigenvectors from Eigenvalues Sparse Principal Component Analysis (EESPCA) R logic for simple example results

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The following R logic was used to generate the results associated with the running example in the EESPCA manuscript and is included as a vignette in the EESPCA R package. This example is based on a data set simulated according to a 10-dimensional multivariate normal (MVN) distribution.

0.1 Load required R packages

The MASS package is used to simulate the MVN data, the PMA package contains the implementation of the Witten et al. LASSO-based sparse principal components method [2], the rifle package contains the implementation the rifle method by Tan et al. [1], and the EESPCA package contains the implementation of the EESPCA method and TPower method by Yuan and Zhang [3] as well as logic to support sparsity parameter selection for both TPower and rifle via cross-validation.

- > library(MASS)
- > library(PMA)
- > library(rifle)
- > library(EESPCA)

0.2 Set random seed

Set the random seed so that simulation is reproducible.

> set.seed(2)

0.3 Set parameters that control the simulation

- > p =10 # number of variables
- > prop.info = 0.4 # proportion of variables that have non-zero loadings on the 1st PC
- > rho = 0.5 # covariance between informative variables
- > n = 100 # simulated sample size

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0.4 Define MVN distribution

The population mean for data is set to the zero vector and the population covariance matrix is given a block covariance structure with one block including the first 4 variables and one block including the last two variables. To simplify the example, we set the population variance for all variables to 1, which aligns with the common practice of performing PCA after standardization.

```
> S = matrix(0, nrow=p, ncol=p)
> num.info = p*prop.info
> S[1:num.info, 1:num.info] = rho
> S[p,p-1] = S[p-1,p] = rho
> diag(S) = 1
> S
       [,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8] [,9] [,10]
 [1,]
       1.0
             0.5
                   0.5
                        0.5
                                0
                                      0
                                            0
                                                 0
                                                     0.0
                                                            0.0
 [2,]
       0.5
             1.0
                   0.5
                        0.5
                                0
                                      0
                                            0
                                                 0
                                                     0.0
                                                            0.0
 [3,]
       0.5
             0.5
                   1.0
                        0.5
                                      0
                                                 0
                                                     0.0
                                0
                                            0
                                                            0.0
 [4,]
       0.5
             0.5
                   0.5
                        1.0
                                0
                                      0
                                            0
                                                 0
                                                     0.0
                                                            0.0
 [5,]
       0.0
             0.0
                   0.0
                        0.0
                                1
                                      0
                                            0
                                                 0
                                                     0.0
                                                            0.0
 [6,]
       0.0
             0.0
                   0.0
                        0.0
                                                 0
                                                     0.0
                                0
                                      1
                                            0
                                                            0.0
 [7,]
       0.0
             0.0
                   0.0
                        0.0
                                0
                                      0
                                            1
                                                 0
                                                     0.0
                                                            0.0
 [8,]
                                                 1
       0.0
             0.0
                   0.0
                        0.0
                                0
                                      0
                                            0
                                                     0.0
                                                            0.0
 [9,]
       0.0
             0.0
                   0.0
                        0.0
                                0
                                      0
                                            0
                                                 0
                                                     1.0
                                                            0.5
[10,]
       0.0
             0.0
                   0.0
                                                  0
                                                     0.5
                        0.0
                                0
                                      0
                                            0
                                                            1.0
```

0.5 Compute population PCs

For this population covariance matrix, the first population PC has equal non-zero loadings (-0.5) for just the first four variables and the second population PC has equal non-zero loadings (0.7071068) for just the last two variables. The variances of these population PCs are 2.5 and 1.5.

```
> (eigen.out = eigen(S))
eigen() decomposition
$values
 [1] 2.5 1.5 1.0 1.0 1.0 1.0 0.5 0.5 0.5 0.5
$vectors
      [,1]
                 [,2] [,3] [,4] [,5] [,6]
                                                 [,7]
                                                             [,8]
                                                                         [,9]
 [1,] -0.5 0.0000000
                         0
                              0
                                   0
                                            0.0000000
                                                       0.8660254
                                                                  0.0000000
 [2,] -0.5 0.0000000
                         0
                              0
                                   0
                                         0
                                            0.0000000 -0.2886751 -0.5773503
 [3,] -0.5 0.0000000
                              0
                         0
                                   0
                                         0
                                            0.0000000 -0.2886751 -0.2113249
 [4,] -0.5 0.0000000
                         0
                              0
                                   0
                                         0
                                            0.0000000 -0.2886751 0.7886751
```

```
0.0 0.0000000
[5,]
                       0
                            0
                                 0
                                      1 0.0000000 0.0000000
                                                               0.0000000
[6,]
      0.0 0.0000000
                            0
                                 1
                                      0 0.0000000 0.0000000
                       0
                                                               0.0000000
[7,]
      0.0 0.0000000
                       0
                            1
                                 0
                                      0.0000000
                                                    0.0000000
                                                               0.000000
[8,]
      0.0 0.0000000
                       1
                            0
                                 0
                                      0.0000000
                                                    0.0000000
                                                               0.000000
[9,]
      0.0 0.7071068
                       0
                            0
                                 0
                                      0 0.7071068
                                                    0.0000000
                                                               0.000000
                                      0 -0.7071068 0.0000000
[10,]
      0.0 0.7071068
                       0
                            0
                                 0
                                                               0.000000
          [,10]
[1,]
      0.000000
[2,] -0.5773503
[3,]
      0.7886751
[4,] -0.2113249
[5,]
      0.0000000
[6,]
      0.0000000
[7,]
      0.0000000
[8,]
      0.000000
[9,]
      0.000000
[10,]
      0.0000000
```

0.6 Compute approximate normed squared loadings for the first population PC.

The approximate normed squared loadings for the first population PC according the formula used by the EESPCA method are:

```
> v1 = eigen.out$vectors[,1]
> lambda1 = eigen.out$values[1]
> (approx.v1.sq = computeApproxNormSquaredEigenvector(S, v1, lambda1, trace=T))
[1] 1.7 1.7 1.7 1.7 1.5 1.5 1.5 1.5 1.5
```

0.7 Simulated MVN data

Simulate one set of 100 independent samples are drawn from this MVN distribution and compute the sample covariance matrix.

```
> X = mvrnorm(n=n, mu=rep(0,p), Sigma=S)
> S.hat = cov(X)
```

0.8 Perform standard PCA

```
> prcomp.out = prcomp(X)
> (V.2 = prcomp.out$rotation[,1:2])

PC1 PC2
[1,] -0.540731693 0.113426021
```

0.9 Compute PCA reconstruction error

Compute the rank 2 reconstruction based on the first two PCs (computed as the squared Frobenius norm of the residual matrix). By definition, this is the minimum rank 2 reconstruction error. It is important to note that this reconstruction error is distinct from the out-of-sample reconstruction error used for comparative analysis in the EESPCA paper and used as the objective function for cross-validation based selection of sparsity parameters.

```
> (pca.error = reconstructionError(X, V.2))
[1] 574.8416
```

Calculate the Euclidean distance between the true PC loadings and the estimated loadings for the first PC.

```
> computeEuclideanDistance = function(x1, x2) {
+  # To account for the arbitrary sign of eigenvectors, taking absolute value first.
+  return (sqrt(sum((abs(x1) - abs(x2))^2)))
+ }
> (pca.L2 = computeEuclideanDistance(v1, prcomp.out$rotation[,1]))
```

[1] 0.108386

0.10 Compute approximate normed squared loadings for the first sample PC.

```
> v1 = prcomp.out$rotation[,1]
> v1^2

[1] 2.923908e-01 1.985940e-01 2.559105e-01 2.460219e-01 1.068926e-03
[6] 4.702457e-04 7.594391e-07 5.170148e-04 2.372812e-03 2.653075e-03
```

```
> lambda1 = prcomp.out$sdev[1]^2
> (approx.v1.sq = computeApproxNormSquaredEigenvector(S.hat, v1, lambda1, trace=F))
[1] 2.639643 2.567531 2.604235 2.596694 2.393685 2.393267 2.392936 2.393280
[9] 2.394510 2.394809
> (ratio = sqrt(approx.v1.sq)/abs(v1))
[1] 3.004628 3.595628 3.190040 3.248803 47.321619 71.340010
```

30.044201

0.11 Compute the first two sparse PCs using EESPCA method

31.767034

```
> eespca.out = eespcaForK(X, k=2, trace=F, compute.sparse.lambda=T)
> eespca.out
```

68.037025

\$V

[7] 1775.084836

```
PC_1 PC_2
[1,] 0.5036013 0.0000000
[2,] 0.4966747 0.0000000
[3,] 0.5002122 0.0000000
[4,] 0.4994874 0.0000000
[5,] 0.0000000 0.0000000
[6,] 0.0000000 0.0000000
[7,] 0.0000000 0.0000000
[8,] 0.0000000 0.7289421
[10,] 0.0000000 0.6845754
```

\$lambdas

[1] 3.366955 1.459852

Compute the rank 2 reconstruction error using the EESPCA sparse PCs:

```
> (eespca.error = reconstructionError(X, eespca.out$V))
```

[1] 583.4963

Calculate the L2 distance between the true PC loadings and the estimated sparse loadings for the first PC.

```
> (eespca.L2 = computeEuclideanDistance(v1, eespca.out$V[,1]))
```

[1] 0.1054057

0.12 Compute the first two sparse PCs using EESPCA.cv method

Find optimal penalty via 5-fold cross-validation for first PC:

```
> p = ncol(X)
> default.thresh = 1/sqrt(p)
> sparse.threshold.values=seq(from=0.75*default.thresh, to=1.25*default.thresh,
+ length.out=21)
> cv.out = eespcaCV(X, sparse.threshold.values=sparse.threshold.values)

Compute sparse PC 1 using penalty that generated minimum error on fist PC
> eespca.cv.v1 = eespca(X, sparse.threshold=cv.out$best.sparsity, compute.sparse.lambd
    Find optimal penalty for second PC using cross-validation on residual matrix. Use that
to compute sparse PC 2.
```

- > X.resid = computeResidualMatrix(X, eespca.cv.v1)
- > cv.out = eespcaCV(X.resid, sparse.threshold.values=sparse.threshold.values)
- > eespca.cv.v2 = eespca(X.resid, sparse.threshold=cv.out\$best.sparsity, compute.spars
- > (V = cbind(eespca.cv.v1,eespca.cv.v2))

```
eespca.cv.v1 eespca.cv.v2
[1,]
        0.5036013
                     0.000000
[2,]
        0.4966747
                     0.000000
[3,]
        0.5002122
                     0.000000
[4,]
        0.4994874
                     0.000000
[5,]
       0.0000000
                     0.0000000
[6,]
        0.0000000
                     0.0000000
[7,]
        0.0000000
                     0.000000
[8,]
        0.0000000
                     0.000000
[9,]
        0.0000000
                     0.7289421
[10,]
        0.0000000
                     0.6845754
```

Compute the rank 2 reconstruction error using the EESPCA.cv sparse PCs:

```
> (eespca.cv.error = reconstructionError(X, V))
```

[1] 583.4963

Calculate the L2 distance between the true PC loadings and the estimated sparse loadings for the first PC.

```
> (eespca.cv.L2 = computeEuclideanDistance(v1, eespca.cv.v1))
```

[1] 0.1054057

0.13Use the Witten et al. method [2] to compute the first two sparse PCs

Find optimal penalty via 5-fold cross-validation for first PC:

```
> cv.out = SPC.cv(X, sumabsv=seq(1, sqrt(p), len=20), niter=10, trace=F)
```

Look at solution at penalty that generated minimum error on fist PC (this applies the same penalty to both the first and second PCs and does not enforce orthogonality):

```
> spc.out=SPC(X, sumabsv=cv.out$bestsumabsv, K=2, trace=F)
> spc.out$v
             [,1]
                         [,2]
 [1,] -0.54059796 0.13111568
 [2,] -0.44564166 -0.05782282
 [3,] -0.50865986 -0.02326875
 [4,] -0.49999193 -0.09621028
 [5,] 0.00000000 0.14463543
 [6,] 0.00000000 -0.03042624
 [7,]
      0.0000000 0.0000000
 [8,] 0.00000000 -0.19873804
 [9,] -0.01472116 0.73352834
[10,] -0.01462424 0.60849121
> (spc.error = reconstructionError(X, spc.out$v))
```

[1] 575.4584

Look at solution at penalty 1 SE from value that generated minimum error:

```
> spc.1se.out =SPC(X, sumabsv=cv.out$bestsumabsv1se, K=2, trace=F)
> spc.1se.out$v
```

```
[,1]
                         [,2]
 [1,] -0.7432266 0.068410686
 [2,] -0.1235048 -0.013146635
 [3,] -0.4631567 -0.002846501
 [4,] -0.4667404 -0.086864933
 [5,]
      0.0000000 0.120546022
 [6,]
      0.000000 0.00000000
 [7,]
      0.000000 0.00000000
 [8,]
      0.0000000 - 0.126978432
 [9,]
      0.0000000 0.751060194
[10,]
      0.0000000 0.626775275
> (spc.1se.error = reconstructionError(X, spc.1se.out$v))
```

```
[1] 620.0697
```

Calculate the L2 distance between the true PC loadings and the estimated sparse loadings for the first PC.

```
> (spc.L2 = computeEuclideanDistance(v1, spc.out$v[,1]))
[1] 0.06779645
> (spc.1se.L2 = computeEuclideanDistance(v1, spc.1se.out$v[,1]))
[1] 0.3931142
```

0.14 Use the Yuan and Zhang TPower method [3] to compute the first two sparse PCs

Find optimal k-value via 5-fold cross-validation for first PC:

```
> k.values = round(seq(1, p, len=20))
> (optimal.k = tpowerPCACV(X=X, k.values=k.values, nfolds=5))
[1] 4
```

Generate initial PC loadings using non-truncated powerIteration:

```
> v.init = powerIteration(X=S.hat)$v1
```

Generate first and second sparse PC loadings using TPower method (second sparse PC is generated on residual matrix using new optimal sparsity value):

- [2,] 0.4479383 0.0000000
- [3,] 0.5070190 0.0000000
- [4,] 0.4997253 0.0000000
- [5,] 0.0000000 0.0000000
- [6,] 0.0000000 0.0000000
- [7,] 0.0000000 0.0000000
- [8,] 0.0000000 0.0000000
- [9,] 0.0000000 0.7556217
- [10,] 0.0000000 0.6550083

```
> (tpower.error = reconstructionError(X, v))
```

[1] 582.2063

Calculate the L2 distance between the true PC loadings and the estimated sparse loadings for the first PC.

```
> (tpower.L2 = computeEuclideanDistance(v1, tpower.v1))
```

[1] 0.08428099

0.15 Use the Tan et al. rifle method [1] to compute the first two sparse PCs

Find optimal k-value via 5-fold cross-validation for first PC:

```
> k.values = round(seq(1, p, len=20))
> (optimal.k = riflePCACV(X=X, k.values=k.values, nfolds=5))
[1] 4
```

Generate initial PC loadings using recommended rifle approach (as implemented in rifle.init() method):

```
> v.init = rifleInit(X)
```

[10,] 0.0000000 0.6323178

Generate first and second sparse PC loadings using rifle method (second sparse PC is generated on residual matrix using new optimal sparsity value):

```
> rifle.v1 = rifle(A=S.hat, B=diag(p), init=v.init, k=optimal.k)
> X.resid = computeResidualMatrix(X, rifle.v1)
> (optimal.k2 = riflePCACV(X=X.resid, k.values=k.values, nfolds=5))
[1] 2
> X.resid.cov = cov(X.resid)
> v2.init = rifleInit(X.resid)
> rifle.v2 = rifle(A=X.resid.cov, B=diag(p), init=v2.init, k=optimal.k2)
> (v = cbind(rifle.v1, rifle.v2))
           [,1]
                     [,2]
 [1,] 0.5507995 0.0000000
 [2,] 0.4347170 0.0000000
 [3,] 0.5081865 0.0000000
 [4,] 0.4993871 0.0000000
 [5,] 0.0000000 0.0000000
 [6,] 0.0000000 0.0000000
 [7,] 0.0000000 0.0000000
 [8,] 0.0000000 0.0000000
 [9,] 0.0000000 0.7747091
```

```
> (rifle.error = reconstructionError(X, v))
[1] 582.377
```

Calculate the L2 distance between the true PC loadings and the estimated sparse loadings for the first PC.

```
> (rifle.L2 = computeEuclideanDistance(v1, -rifle.v1))
```

[1] 0.08555848

0.16 Summary of reconstruction errors and estimation distances

```
> estimation.summary = data.frame(pca=c(pca.error, pca.L2),
          spc=c(spc.error, spc.L2),
          spc.1se=c(spc.1se.error, spc.1se.L2),
          eespca=c(eespca.error, eespca.L2),
          tpower=c(tpower.error, tpower.L2),
          rifle=c(rifle.error, rifle.L2))
> rownames(estimation.summary) = c("Reconstruction Error", "Loadings distance")
> estimation.summary
                                                 spc.1se
                            pca
                                         spc
                                                               eespca
Reconstruction Error 574.841592 575.45837865 620.0696917 583.4962900
                       0.108386
                                  0.06779645
                                               0.3931142
Loadings distance
                                                           0.1054057
                           tpower
Reconstruction Error 582.20628324 582.37695461
Loadings distance
                       0.08428099
                                    0.08555848
```

References

- [1] Kean Ming Tan, Zhaoran Wang, Han Liu, and Tong Zhang. Sparse generalized eigenvalue problem: optimal statistical rates via truncated rayleigh flow. *Journal of the Royal Statistical Society: Series B (Statistical Methodology)*, 80(5):1057–1086, 2018.
- [2] Daniela M. Witten, Robert Tibshirani, and Trevor Hastie. A penalized matrix decomposition, with applications to sparse principal components and canonical correlation analysis. *Biostatistics*, 10(3):515–534, July 2009.
- [3] Xiao-Tong Yuan and Tong Zhang. Truncated power method for sparse eigenvalue problems. J. Mach. Learn. Res., 14(1):899–925, April 2013.