The irlba Package

Bryan W. Lewis
blewis@illposed.net,

adapted from the work of:
Jim Baglama (University of Rhode Island)
and Lothar Reichel (Kent State University).

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1 Introduction

The irlba package provides a fast way to compute partial singular value decompositions (SVD) of large sparse or dense matrices. Recent additions to the package can also compute fast partial symmetric eigenvalue decompositions and principal components. The package is an R implementation of the augmented implicitly restarted Lanczos bidiagonalization algorithm of Jim Baglama and Lothar Reichel\(^1\). Source code is maintained at https://github.com/bwlewis/irlba.

The irlba package works with real- and complex-valued dense R matrices and real-valued sparse matrices from the Matrix package. It provides several easy ways to define custom matrix arithmetic that works with other matrix classes including big.matrix from the bigmemory package and others. The irlba is both faster and more memory efficient than the usual R svd function for computing a few of the largest singular vectors and corresponding singular values of a matrix. It takes advantage of available high-performance linear algebra libraries if R is compiled to use them. In particular, the package uses the same BLAS and LAPACK libraries that R uses (see https://cran.r-project.org/doc/manuals/R-admin.html#BLAS), or the CHOLMOD library from R’s Matrix package for sparse matrix problems.

A whirlwind summary of the algorithm follows, along with a few basic examples. A much more detailed description and discussion of the algorithm may be found in the cited Baglama-Reichel reference.

2 Partial Singular Value Decomposition

Let \( A \in \mathbb{R}^{\ell \times n} \) and assume \( \ell \geq n \). These notes simplify the presentation by considering only real-valued matrices and assuming without losing generality that there are at least as many rows as columns (the method works more generally). A singular value decomposition of \( A \) can be expressed as:

\[
A = \sum_{j=1}^{n} \sigma_j u_j v_j^T,
\]

where \( u_j \in \mathbb{R}^\ell \), \( v_j \in \mathbb{R}^n \), \( j = 1, 2, \ldots, n \), and \( \sigma_1 \geq \sigma_2 \geq \cdots \geq \sigma_n \geq 0 \).

Let \( 1 \leq k < n \). A rank \( k \) partial SVD of \( A \) is defined as:

\[
A_k := \sum_{j=1}^{k} \sigma_j u_j v_j^T.
\]

The following simple example shows how to use irlba to compute the five largest singular values and corresponding singular vectors of a 5000 \( \times \) 5000 matrix. We compare to the usual R svd function and report timings for our test machine, a 4-CPU core, 3.0 GHz AMD A10-7850K personal computer with 16 GB RAM, using R version 3.3.1 using the high performance AMD ACML core math library BLAS and LAPACK.

```r
> library('irlba')
> set.seed(1)
> A <- matrix(rnorm(5000*5000), 5000)
> t1 <- proc.time()
> L <- irlba(A, 5)
> print(proc.time() - t1)
user  system elapsed
17.440  0.192  4.417
> gc()
used (Mb) gc trigger (Mb) max used (Mb)
Ncells 1096734  58.6  1770749 94.6 1442291 77.1
Vcells 26685618 203.6 62229965 474.8 52110704 397.6
```

Compare with the standard svd function:
The \texttt{irlba} Package

\begin{verbatim}
> t1 <- proc.time()
> S <- svd(A, nu=5, nv=5)
> print(proc.time() - t1)
  user  system elapsed
277.092 11.552  74.425
> gc()

used (Mb) gc trigger (Mb) max used (Mb)
Ncells 1097441 58.7 1770749 94.6 1442291 77.1
Vcells 26741910 204.1 169891972 1296.2 176827295 1349.1
\end{verbatim}

The \texttt{irlba} method uses about 1/20 elapsed time as the \texttt{svd} method in this example and less than one third the peak memory. The default tolerance value yields the following relative error in the estimated singular values:

\begin{verbatim}
> sqrt(crossprod(S$d[1:5]-L$d)/crossprod(S$d[1:5]))
[,1]
[1,] 4.352641e-10
\end{verbatim}

\section{Convergence tolerance}

\texttt{IRLBA} is an iterative method that estimates a few singular values and associated singular vectors. A sketch of the algorithm is outlined in Section 3 below. The R\texttt{tol} and \texttt{svtol} arguments control when the algorithm converges with \texttt{tol} specifying subspace convergence, and \texttt{svtol} specifying convergence of estimated singular values.

Subspace convergence occurs when the algorithm iterations find estimated singular vectors that satisfy

\[ \|A V_k - U S_k\| < \texttt{tol} \cdot \|A\|, \]

where \(\cdot\) means spectral matrix norm, \(A\) is the matrix, \(V_k\) and \(U_k\) are the \textit{estimated} right and left \(k\) singular vectors computed by the algorithm, and \(\|A\|\) is the \textit{estimated} spectral norm of the matrix defined by the largest singular value computed by the algorithm. Using R notation, the algorithm stops when

\begin{verbatim}
L <- irlba(A, k, tol)
\end{verbatim}

It’s possible to encounter problems that fail to converge before the maximum number of algorithm iterations specified by the \texttt{maxit} argument.

When the largest singular values are clustered together it can be hard to detect subspace convergence. More recent versions of the \texttt{IRLBA} implementation include the \texttt{svtol} argument that specifies a maximum for the relative change in each estimated singular value from one iteration to the next.
The convergence tolerance values together help improve correct subspace detection in difficult settings when the singular values are clustered. But in the worst cases, block methods can perform better as shown in the documentation for the svdr method.

Also see the related rsvd function by N. Benjamin Erichson, https://cran.r-project.org/package=rsvd.

2.2 Differences with svd

The irlba function is designed to compute a partial singular value decomposition. It is largely compatible with the usual R svd function but there are some differences. In particular:

1. The irlba function only computes the number of singular values corresponding to the maximum of the desired singular vectors, max(nu, nv). For example, if 5 singular vectors are desired (nu=nv=5), then only the five corresponding singular values are computed. The standard R svd function always returns the total set of singular values for the matrix, regardless of how many singular vectors are specified.

2. The irlba function is an iterative method that continues until either a tolerance or maximum number of iterations is reached. Problems with difficult convergence properties are not likely to be encountered, but the method will fail with an error after the iteration limit is reached in those cases.

Watch out especially for the first difference noted above!

2.3 Principal Components

Version 2.1.0 of the package introduces optional arguments and prcomp-like function syntax for efficiently computing partial SVDs of matrices after centering and scaling their columns and other adjustments. Use the following arguments to the irlba function, or the new irlba_prcomp function for PCA:

- **center**: if center is a numeric vector with length equal to the number of columns of the matrix, then each column of the matrix has the corresponding value from center subtracted from it.

- **scale**: if 'scale' is a numeric vector with length equal to the number of columns of the matrix, then each column is divided by the corresponding value from scale.

Both centering and scaling options are performed implicitly in the algorithm and, for instance, do not affect sparsity of the input matrix or increase storage requirements. The following example compares the output of the usual prcomp function with output from irlba. Note that in general, singular vectors and principal component vectors are only unique up to sign!
> set.seed(1)
> x <- matrix(rnorm(200), nrow=20)
> p1 <- prcomp_irlba(x, n=3)
> summary(p1)

Importance of components %:

<table>
<thead>
<tr>
<th></th>
<th>PC1</th>
<th>PC2</th>
<th>PC3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Std Dev</td>
<td>1.5411</td>
<td>1.2513</td>
<td>1.1916</td>
</tr>
<tr>
<td>Var</td>
<td>0.2806</td>
<td>0.1850</td>
<td>0.1678</td>
</tr>
<tr>
<td>Cum Var</td>
<td>0.2806</td>
<td>0.4656</td>
<td>0.6334</td>
</tr>
</tbody>
</table>

> # Compare with
> p2 <- prcomp(x, tol=0.7)
> summary(p2)

Importance of components:

<table>
<thead>
<tr>
<th></th>
<th>PC1</th>
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<td>0.2806</td>
<td>0.4656</td>
<td>0.6334</td>
</tr>
</tbody>
</table>

Alternatively, you can compute principal components directly using the singular value decomposition and the center option:

> p3 <- svd(scale(x, center=colMeans(x), scale=FALSE))
> p4 <- irlba(x, 3, center=colMeans(x))

> # compare with prcomp
> sqrt(crossprod(p1$rotation[,1] - p3$v[,1]))

[1,] 9.773228e-13

> sqrt(crossprod(p1$rotation[,1] + p4$v[,1]))

[1,] 1.652423e-12

2.4 Truncated symmetric eigenvalue decomposition

Use the partial_eigen function to estimate a subset of the largest (most positive) eigenvalues and corresponding eigenvectors of a symmetric dense or sparse real-valued matrix. The function is particularly well-suited to estimating the largest eigenvalues and corresponding eigenvectors of symmetric positive semi-definite matrices of the form $A^TA$. 
2.5 User-Defined Matrix Multiplication

The `irlba` function only uses matrix vector products with the input data matrix to compute its solution. It’s easy to use R’s native object model to define custom matrix classes with user-defined matrix multiplication functions. Such functions can be used to support special matrix objects, out of core computation of large problems, or matrix-free operators.

Here is a simple example that defines a matrix product that scales the columns of the matrix to have unit norm (cf the `scale` option).

```r
> A <- matrix(runif(400), nrow=20)
> col_scale <- sqrt(apply(A, 2, crossprod))
> setClass("scaled_matrix", contains="matrix", slots=c(scale=numeric))
> setMethod("%*%", signature(x="scaled_matrix", y="numeric"),
  + function(x ,y) x@.Data %*% (y / x@scale))
> setMethod("%*%", signature(x="numeric", y="scaled_matrix"),
  + function(x ,y) (x %*% y@.Data) / y@scale)
> a <- new("scaled_matrix", A, scale=col_scale)
> irlba(a, 3)$d

[1] 3.9298391 0.9565016 0.8266859

# Compare with
> svd(sweep(A, 2, col_scale, FUN='/'))$d[1:3]

[1] 3.9298391 0.9565016 0.8266859
```

See the following link for an example that uses large-scale out of core computation: [http://bwlewis.github.io/1000genomes/examples/PCA/wholegenome.html](http://bwlewis.github.io/1000genomes/examples/PCA/wholegenome.html)

NOTE! The reference R algorithm implementation is used whenever user-defined matrix multiplication is specified (instead of the faster C code path).

3 A Quick Summary of the IRLBA Method

3.1 Partial Lanczos Bidiagonalization

Start with a given vector \( p_1 \). Compute \( m \) steps of the Lanczos process:

\[
AP_m = Q_mB_m \\
A^TQ_m = P_mD_m + r_m\epsilon_m^T,
\]
\[ B_m \in \mathbb{R}^{m \times m}, P_m \in \mathbb{R}^{n \times m}, Q_m \in \mathbb{R}^{l \times m}, \]
\[ P_m^T P_m = Q_m^T Q_m = I_m, \]
\[ r_m \in \mathbb{R}^{n}, P_m^T r_m = 0, \]
\[ P_m = [p_1, p_2, \ldots, p_m]. \]

### 3.2 Approximating Partial SVD with A Partial Lanczos bidiagonalization

\[ A^T A P_m = A^T Q_m B_m = P_m B_m^T B_m + r_m e_m^T B_m, \]
\[ A A^T Q_m = A P_m B_m^T + A r_m e_m^T = Q_m B_m B_m^T + A r_m e_m^T. \]

Compute the SVD of \( B_m \):
\[ B_m = \sum_{j=1}^{m} \sigma_j B u_j^B (v_j^B)^T. \]

(i.e., \( B_m v_j^B = \sigma_j B u_j^B \), and \( B_m^T u_j^B = \sigma_j^B v_j^B \).)

Define: \( \tilde{\sigma}_j := \sigma_j^B, \tilde{u}_j := Q_m u_j^B, \tilde{v}_j := P_m v_j^B. \)

Then:
\[ A \tilde{v}_j = A P_m v_j^B = Q_m B_m v_j^B = \sigma_j^B Q_m u_j^B = \tilde{\sigma}_j \tilde{u}_j, \]
\[ A^T \tilde{u}_j = A^T Q_m u_j^B = P_m B_m^T u_j^B + r_m e_m^T u_j^B = \sigma_j^B P_m v_j^B + r_m e_m^T u_j^B = \tilde{\sigma}_j \tilde{v}_j + r_m e_m^T u_j^B. \]
The IRLBA Package

The part in red above represents the error with respect to the exact SVD. The IRLBA strategy is to iteratively reduce the norm of that error term by augmenting and restarting.

Here is the overall method:

1. Compute the Lanczos process up to step \( m \).
2. Compute \( k < m \) approximate singular vectors.
3. Orthogonalize against the approximate singular vectors to get a new starting vector.
4. Continue the Lanczos process with the new starting vector for \( m \) more steps.
5. Check for convergence tolerance and exit if met.
6. GOTO 1.

### 3.3 Sketch of the augmented process...

\[
\bar{P}_{k+1} := [\tilde{v}_1, \tilde{v}_2, \ldots, \tilde{v}_k, p_{m+1}], \\
A\bar{P}_{k+1} = [\tilde{\sigma}_1 \tilde{u}_1, \tilde{\sigma}_1 \tilde{u}_2, \ldots, \tilde{\sigma}_k \tilde{u}_k, A p_{m+1}]
\]

Orthogonalize \( A p_{m+1} \) against \( \{\tilde{u}_j\}_{j=1}^k \): \( A p_{m+1} = \sum_{j=1}^{k} \rho_j \tilde{u}_j + r_k \).

\[
\bar{Q}_{k+1} := [\tilde{u}_1, \tilde{u}_2, \ldots, \tilde{u}_k, r_k/\|r_k\|], \\
\bar{B}_{k+1} := \begin{bmatrix}
\tilde{\sigma}_1 & \rho_1 \\
\tilde{\sigma}_2 & \rho_2 \\
\vdots & \vdots \\
\tilde{\sigma}_k & \rho_k \\
\|r_k\| & \|r_k\|
\end{bmatrix}
\]

\( A\bar{P}_{k+1} = \bar{Q}_{k+1} \bar{B}_{k+1} \).