# Package 'StructureMC'

May 5, 2019

Type Package

Title Structured Matrix Completion
Version 1.0
<b>Date</b> 2015-07-30
<b>Depends</b> R (>= 3.1.0), MASS, matrixcalc
Description Current literature on matrix completion focuses primarily on independent sampling models under which the individual observed entries are sampled independently. Motivated by applications in genomic data integration, we propose a new framework of structured matrix completion (SMC) to treat structured missingness by design. Specifically, our proposed method aims at efficient matrix recovery when a subset of the rows and columns of an approximately low-rank matrix are observed. The main function in our package, smc.FUN, is for recovery of the missing block A22 of an approximately low-rank matrix A given the other blocks A11, A12, A21.
URL https://github.com/celehs/StructureMC
BugReports https://github.com/celehs/StructureMC/issues
License GPL-2
Encoding UTF-8
RoxygenNote 6.1.1
NeedsCompilation no
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Structured Matrix Completion

## **Description**

Current literature on matrix completion focuses primarily on independent sampling models under which the individual observed entries are sampled independently. Motivated by applications in genomic data integration, we propose a new framework of structured matrix completion (SMC) to treat structured missingness by design. Specifically, our proposed method aims at efficient matrix recovery when a subset of the rows and columns of an approximately low-rank matrix are observed. The main function in our package, smc.FUN, is for recovery of the missing block A22 of an approximately low-rank matrix A given the other blocks A11, A12, A21.

#### Author(s)

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#### References

Cai, T., Cai, T. T., & Zhang, A. (2015). Structured Matrix Completion with Applications to Genomic Data Integration. *Journal of the American Statistical Association*.

mynorm

mynorm

## Description

This function returns the spectral norm of a real matrix if type is 2. Otherwise, it returns the matrix norm of the "norm" function using LAPACK.

#### Usage

mynorm(x, type)

## **Arguments**

x numeric matrix

type character string, specifying the type of matrix norm to be computed. Details see

norm function in R base.

## Author(s)

Yifu Liu, Anru Zhang, Tianxi Cai and T. Tony Cai,

## References

Cai, T., Cai, T. T., & Zhang, A. (2015). Structured Matrix Completion with Applications to Genomic Data Integration. *Journal of the American Statistical Association*.

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#### See Also

norm

#### **Examples**

```
A = matrix(rnorm(10, mean = 0, sd = 0.1), 10, 10)
mynorm(A, "2")
mynorm(A, "0")
```

smc.FUN

Structured Matrix Completion

## **Description**

The main function in our package, smc.FUN, is for recovery of the missing block A22 of an approximately low-rank matrix A given the other blocks A11, A12, A21.

## Usage

```
smc.FUN(A.mat, c_T, col_thresh, m1, m2)
```

#### **Arguments**

A.mat The approximately low-rank matrix that we want to recover c\_T c\_T is the thresholding level, the default value is 2.

col\_thresh is column thresholding

m1 number of rows of block A11

m2 number of columns of block A11

## Author(s)

Yifu Liu, Anru Zhang, Tianxi Cai and T. Tony Cai,

## References

Cai, T., Cai, T. T., & Zhang, A. (2015). Structured Matrix Completion with Applications to Genomic Data Integration. *Journal of the American Statistical Association*.

## **Examples**

```
##dimension of matrix A with row number p1 = 10 and column number p2 = 9 p1 = 10 p2 = 9 m1 = 6##row number of A11 m2 = 3##column number of A11 A = matrix(rnorm(10, mean = 0, sd = 0.1), p1, p2) ##find the approximation low-rank of A under singular value decomposition (SVD) j1 = svd(A)$d num<-length(which(j1!=0)) j2 = svd(A)$u
```

smc.FUN

```
j3 = svd(A)$v
D<-diag(j1)
D[num, num] = 0
app_low_rank_A = j2
##consturct the matrix Arecovery that need to be recovered
A11 = app_low_rank_A[1:m1, 1:m2]
A12 = app_low_rank_A[1:m1, (1+m2):p2]
A21 = app_low_rank_A[(1+m1):p1, 1:m2]
Arecovery = rbind(cbind(A11,A12),cbind(A21,matrix(NA,nrow=p1-m1,ncol=p2-m2)))
##recovery the block A22
A22 = smc.FUN(Arecovery, 2, "True", m1, m2)</pre>
```

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