Simulate MRMC Data

There are two functions in this package that can simulate Multi-reader Multi-case (MRMC) data. sim.new.Hierarchical is the function to simulate the MRMC agreement data with no binary truth state. The sim.gRoeMetz is the function to simulate MRMC ROC data.

1. Using the new hierarchical model to simulate the MRMC agreement data

The following is how to use the sim.new.Hierachcial.R to simulate MRMC data with no truth state, the MRMC agreement data

```
# configuration
nR = 5
            #number of readers
nC = 100
            #number of cases
config <- sim.new.Hierarchical.config(nR=nR, nC=nC, modalityID = c("testA", "testB"))</pre>
# simulate MRMC study
set.seed(1, kind = "L'Ecuyer-CMRG")
dFrame.newH <- sim.new.Hierarchical(config)</pre>
# check the first and last few lines of the simulated dataframe
head(dFrame.newH)
##
     caseID readerID modalityID
                                     score
## 1 Case1 reader1
                         testA 0.6946134
## 2
     Case2 reader1
                         testA 2.9965743
## 3 Case3 reader1
                         testA 1.7412831
## 4 Case4 reader1
                         testA -1.2549331
## 5 Case5 reader1
                         testA -0.2952628
## 6 Case6 reader1
                         testA 3.0468024
tail(dFrame.newH)
##
         caseID readerID modalityID
                                         score
## 995
        Case95 reader5
                             testB -0.1549758
## 996
        Case96 reader5
                             testB -1.2040039
## 997
         Case97 reader5
                             testB 1.6544295
## 998
        Case98 reader5
                             testB -1.2125349
## 999
        Case99 reader5
                              testB -0.2017070
## 1000 Case100 reader5
                              testB 2.0605555
```

We simulated 5 readers and 100 cases for 2 modalities, so the total number of scores is 5x100x2=1000.

2. Using sim.gRoeMetz in iMRMC package to simulate MRMC ROC data

The following is how to use the sim.gRoeMetz.R to simulate MRMC data with truth state, the MRMC ROC data

```
# configuration
nR = 5  #number of readers
```

```
nC.neg = 50 #number of positive cases
nC.pos = 50 #number of negative cases
config <- sim.gRoeMetz.config(nR = nR, nC.neg = nC.neg, nC.pos = nC.neg)</pre>
# simulate MRMC study
set.seed(1, kind = "L'Ecuyer-CMRG")
dFrame.gRM <- sim.gRoeMetz(config)</pre>
# check the first and last few lines of the simulated dataframe
head(dFrame.gRM)
     readerID
               caseID modalityID score
## 1
          -1 negCase1
                             truth
## 2
           -1 negCase2
                             truth
## 3
           -1 negCase3
                             truth
## 4
           -1 negCase4
                             truth
                                        0
           -1 negCase5
## 5
                             truth
                                        0
## 6
           -1 negCase6
                             truth
                                        0
tail(dFrame.gRM)
##
        readerID
                     caseID modalityID
                                             score
## 1095 reader5 posCase45
                             testB 2.9342657
## 1096 reader5 posCase46
                                 testB 1.0639768
## 1097 reader5 posCase47
                                 testB 0.9885389
## 1098 reader5 posCase48
                                 testB -0.5130517
        reader5 posCase49
## 1099
                                 testB 0.9254049
## 1100 reader5 posCase50
                                 testB 2.4164279
The simulated data starts with truth state of each case, and followed by the reading scores from each of
the readers. Since we simulate 50 positive cases and 50 negative cases, there are 100 lines for the truth and
100x5x2 = 1000 lines for the scores from 5 readers for 2 modalities.
To combine the truth data and the reader scores and change the data to a dataframe with 5 columns:
readerID, caseID, modalityID, score, truth, we can use the function undoIMRMCdf in iMRMC pacakge:
dFrame.gRM.2 <- undoIMRMCdf(dFrame.gRM)</pre>
# check the first and last few lines of the simulated dataframe
head(dFrame.gRM.2)
##
       caseID readerID modalityID
                                          score truth
## 1 negCase1 reader1 testA 0.36795249
## 2 negCase1 reader2
                           testB 0.05607467
                                                     0
                        testB -0.46914121
testA -1.38686722
testB -0.49895246
## 3 negCase1 reader3
                                                     0
## 4 negCase1 reader4
                                                     0
## 5 negCase1 reader1
                                                     0
## 6 negCase1 reader5
                             testA -1.64836669
tail(dFrame.gRM.2)
```

```
## caseID readerID modalityID score truth
## 995 posCase9 reader5 testA 0.9233879 1
## 996 posCase9 reader4 testB 4.1614474 1
## 997 posCase9 reader4 testA 3.0874475 1
## 998 posCase9 reader2 testB 3.4945769 1
```

999 posCase9 reader1 testA 2.2259825 1 ## 1000 posCase9 reader5 testB 1.3017587 1