The RJafroc Quick Start Book

Dev P. Chakraborty, PhD

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Preface

- This online book is for those already somewhat familiar running Windows JAFROC to analyze data.
- The Windows program has been replaced by RJafroc.
- This book dives into how to use RJafroc to analyze ROC/FROC data.
- It starts with explanation of the dataset structures for ROC and FROC studies.
- This is followed by an explanation of DBM and OR analyses.

0.1 Rationale and Organization

- Intended as an online update to my print book (Chakraborty, 2017).
- All references in this book to RJafroc refer to the R package with that name (case sensitive) (Chakraborty and Zhai, 2022).
- Since its publication in 2017 RJafroc, on which the R code examples in the print book depend, has evolved considerably causing many of the examples to "break" if one uses the most current version of RJafroc. The code will still run if one uses RJafroc 0.0.1 but this is inconvenient and misses out on many of the software improvements made since the print book appeared.
- This gives me the opportunity to update the print book.
- The online book has been divided into 3 books.
 - This book: RJafrocQuickStartBook.
 - The RJafrocRocBook book.
 - The RJafrocFrocBook.

0.2 Getting help on the software

- If you have installed RJafroc from GitHub:
 - ?RJafroc-package (RStudio will auto complete ...) followed by Enter.
 - Scroll down all the way and click on Index
- Regardless of where you installed from use the RJafroc help site:
 - RJafroc help site
 - Look under the Reference tab.
 - For example, for help on the function ${\tt PlotEmpiricalOperatingCharacteristics}$:
 - PlotEmpiricalOperatingCharacteristics

0.3 TBA Acknowledgements

0.3.1 Persons who have stimulated my thinking:

Harold Kundel, MD

Claudia Mello-Thoms, PhD

0.3.2 Contributors to the software

Dr. Xuetong Zhai

Dr. Peter Phillips

Online Latex Editor at

0.3.3 Dataset contributors

TBA

Quick Start

Chapter 1

JAFROC ROC data format

1.1 TBA How much finished

80% (need to add images for one reader; add one-modality dataset)

1.2 Introduction

- JAFROC data format is named after the file format adopted circa. 2006 for the input Excel file to Windows
 JAFROC software.
- The purpose of this chapter is to explain the data format of this file.
- Reading this file into a dataset object suitable for RJafroc analysis is the subject of the next chapter.
- Background on observer performance methods are in my book (Chakraborty, 2017).
- I will start with Receiver Operating Characteristic (ROC) data (Metz, 1978) as this is by far the simplest paradigm.
- In the ROC paradigm the observer assigns a rating to each image. A rating is an ordered numeric label, and, in our convention, higher values represent greater certainty or **confidence level** for presence of disease. With human observers, a 5 (or 6) point rating scale is typically used, with 1 representing highest confidence for *absence* of disease and 5 (or 6) representing highest confidence for *presence* of disease. Intermediate values represent intermediate confidence levels for presence or absence of disease.
- Note that location information, if applicable, associated with the disease, is not collected.
- There is no restriction to 5 or 6 ratings. With algorithmic observers, e.g., computer aided detection (CAD) algorithms, the rating could be a floating point number and have infinite precision. All that is required is that higher values correspond to greater confidence in presence of disease.
- The above is termed a positive-directed rating scale. If lower numbers correspond to greater confidence, termed a negative-directed rating scale, a simple transformation to max(rating) rating + 1, where max(rating) is the maximum rating, over all readers, modalities and cases, will convert a negative-directed rating scale to a positive directed rating scale.

1.3 Note to existing users

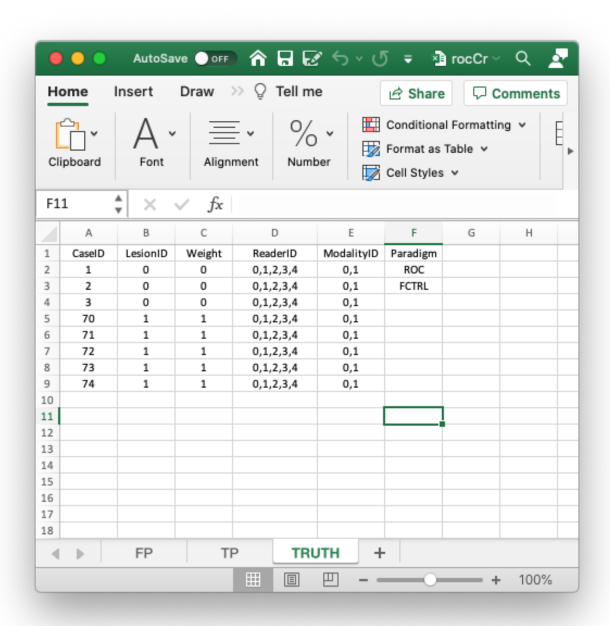
- The Excel file format has recently undergone changes, involving three additional columns in the Truth worksheet
- RJafroc will work with old format Excel files as the additional columns are ignored.
- Reasons for the change will become clearer in later chapters ¹.

¹They are needed for generalization to other data collection paradigms and for better data entry error control

1.4 Contents of Excel file

- The illustrations in this chapter correspond to Excel file R/quick-start/rocCr.xlsx in the project directory ². This is termed a *toy file*, i.e., an artificial small dataset created to illustrate essential features of the data format.
- The Excel file has three worksheets: Truth, NL (or FP) and LL (or TP).

1.5 The Truth worksheet

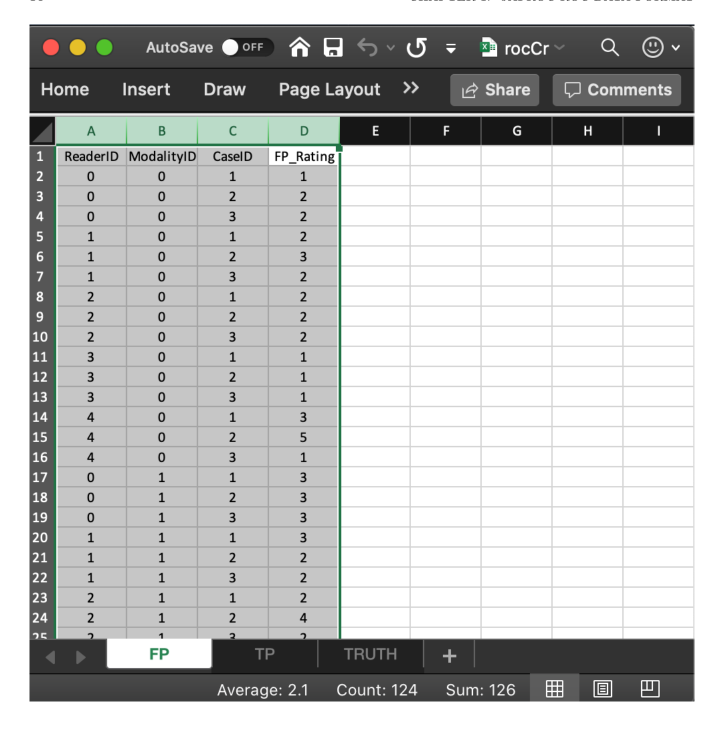


• The Truth worksheet contains 6 columns: CaseID, LesionID, Weight, ReaderID, ModalityID and Paradigm.

²To access files one needs to fork the repository, which creates, on your computer, a copy of all files used to create this document

- The first five columns contain as many rows as there are cases (images) in the dataset.
- CaseID: unique integers, one per case, representing the cases in the dataset.
- LesionID: integers 0 or 1, with each 0 representing a non-diseased case and each 1 representing a diseased case.
- In the current dataset, the non-diseased cases are labeled 1, 2 and 3, while the diseased cases are labeled 70, 71, 72, 73 and 74. The values do not have to be consecutive integers; they need not be ordered; the only requirement is that they be **unique integers**.
- Weight: A floating point value, typically filled in with 0 or 1; this field is not used for ROC data.
- ReaderID: a comma-separated listing of reader labels, each represented by a unique integer, that have interpreted the case. In the example shown below each cell has the value 0, 1, 2, 3, 4 meaning that each of these readers has interpreted all cases (hence the "factorial" design).
 - With multiple readers each cell in this column has to be text formatted as otherwise Excel will not accept it.
 - Select the worksheet, then Format Cells Number Text OK.
- ModalityID: a comma-separated listing of modalities, each represented by a **unique integer**, that are applied to each case. In the example each cell has the value 0, 1.
 - With multiple modalities each cell has to be text formatted as otherwise Excel will not accept it.
 - Format the cells as described above.
- Paradigm: this column contains two cells, ROC and factorial. It informs the software that this is an ROC dataset, and the design is factorial, meaning each reader has interpreted each case in each modality.
- There are 5 diseased cases in the dataset (the number of 1's in the LesionID column of the Truth worksheet).
- There are 3 non-diseased cases in the dataset (the number of 0's in the LesionID column).
- There are 5 readers in the dataset (each cell in the ReaderID column contains the string 0, 1, 2, 3, 4).
- There are 2 modalities in the dataset (each cell in the ModalityID column contains the string 0, 1).

1.6 The false positive (FP) ratings



- It consists of 4 columns, each of length 30 (# of modalities X number of readers X number of non-diseased cases).
- ReaderID: the reader labels: 0, 1, 2, 3 and 4. Each reader label occurs 6 times (# of modalities X number of non-diseased cases).
- ModalityID: the modality or treatment labels: 0 and 1. Each label occurs 15 times (# of readers X number of non-diseased cases).
- CaseID: the case labels for non-diseased cases: 1, 2 and 3. Each label occurs 10 times (# of modalities X # of readers).
- The label of a diseased case cannot occur in the FP worksheet. If it does the software generates an error.
- FP_Rating: the floating point ratings of non-diseased cases. Each row of this worksheet contains a rating corresponding to the values of ReaderID, ModalityID and CaseID for that row.

1.7 The true positive (TP) ratings

These are found in the TP or LL worksheet, see below.

	• •	AutoSa	ve OFF	介 5	⇔ ^ (ড ∓	rocCı	·	© v
Н	ome	Insert	Draw	Page La	ayout >	>	☆ Share	□ Com	ments
	Α	В	С	D	Е	F	G	н	- 1
1	ReaderID	ModalityID	CaseID	LesionID	TP_Rating				
2	0	0	70	1	5				
3	0	0	71	1	5				
4	0	0	72	1	5				
5	0	0	73	1	5				
6	0	0	74	1	4				
7	1	0	70	1	5				
8	1	0	71	1	3				
9	1	0	72	1	5				
10	1	0	73	1	5				
11	1	0	74	1	5				
12	2	0	70	1	5				
13	2	0	71	1	4				
14	2	0	72	1	5				
15	2	0	73	1	5				
16	2	0	74	1	5				
17	3	0	70	1	5				
18	3	0	71	1	5				
19	3	0	72	1	5				
20	3	0	73	1	5				
21	3	0	74	1	5				
22	4	0	70	1	5				
23	4	0	71	1	2				
24	4	0	72	1	5				
25	1	0	72	1	າ				
4	•	FP	TI	P	TRUTH	+			
		Average: 2	5.853333	333 Co	ount: 255	Sun	ո։ 3878		四

- It consists of 5 columns, each of length 50 (# of modalities X number of readers X number of diseased cases).
- ReaderID: the reader labels: 0, 1, 2, 3 and 4. Each reader label occurs 10 times (# of modalities X number of diseased cases).
- ModalityID: the modality or treatment labels: 0 and 1. Each label occurs 25 times (# of readers X number of diseased cases).
- LesionID: For an ROC dataset this column contains fifty 1's (each diseased case has one lesion).
- CaseID: the case labels for non-diseased cases: 70, 71, 72, 73 and 74. Each label occurs 10 times (# of modalities X # of readers). For an ROC dataset the label of a non-diseased case cannot occur in the TP worksheet. If it does the software generates an error.

• TP_Rating: the floating point ratings of diseased cases. Each row of this worksheet contains a rating corresponding to the values of ReaderID, ModalityID, LesionID and CaseID for that row.

1.8 A single reader dataset

```
rocCr1R <- "R/quick-start/rocCr1R.xlsx"</pre>
x <- DfReadDataFile(rocCr1R, newExcelFileFormat = TRUE)
str(x)
#> List of 3
#> $ ratings
                 :List of 3
   ..$ NL : num [1:2, 1, 1:8, 1] 2 3 3 2 2 ...
    ..$ LL : num [1:2, 1, 1:5, 1] 5 5 3 3 5 5 5 5 5 5
#>
#>
    ..$ LL_IL: logi NA
#> $ lesions
                :List of 3
   ..$ perCase: int [1:5] 1 1 1 1 1
    ..$ IDs : num [1:5, 1] 1 1 1 1 1
#>
#>
    ..$ weights: num [1:5, 1] 1 1 1 1 1
#> $ descriptions:List of 7
#> ..$ fileName : chr "rocCr1R"
#>
    ..$ type
                     : chr "ROC"
                : logi NA
#>
    ..$ name
#>
    ...$ truthTableStr: num [1:2, 1, 1:8, 1:2] 1 1 1 1 1 1 NA NA NA NA ...
#>
                : chr "FCTRL"
    \dots$ design
    ..$ modalityID : Named chr [1:2] "0" "1"
#>
    .. ..- attr(*, "names")= chr [1:2] "0" "1"
#> ..$ readerID
                    : Named chr "1"
#> ... - attr(*, "names")= chr "1"
```

Chapter 2

Reading the Excel data file

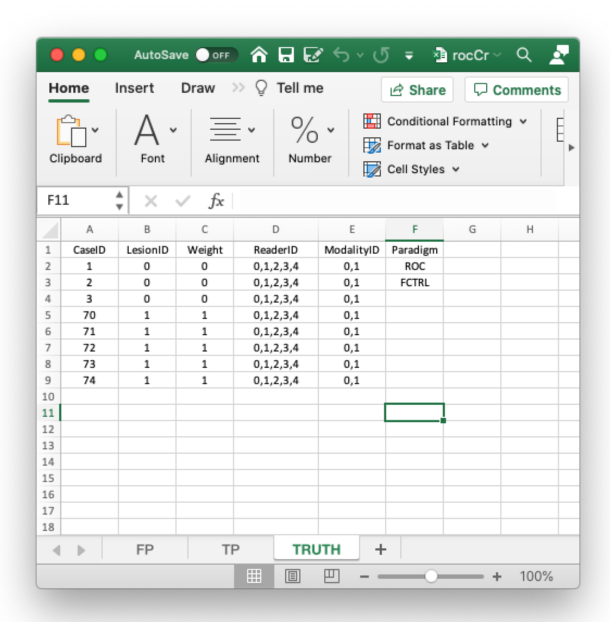
2.1 TBA How much finished

90%

2.2 Introduction

In the previous chapter I described the format of the Excel file R/quick-start/rocCr.xlsx corresponding to a small factorial ROC dataset. Described here is how to read this file in order to create an RJafroc dataset. It introduces the RJafroc function DfReadDataFile(). Also shown are the correspondences between values in the Excel file and the dataset object.

2.3 The structure of an ROC dataset



In the following code chunk the second statement reads the Excel file using the function DfReadDataFile() and saves it to object x. The third statement shows the structure of x.

```
rocCr <- "R/quick-start/rocCr.xlsx"
x <- DfReadDataFile(rocCr, newExcelFileFormat = TRUE)
str(x)
#> List of 3
#> $ ratings : List of 3
#> ..$ NL : num [1:2, 1:5, 1:8, 1] 1 3 2 3 2 2 1 2 3 2 ...
#> ..$ LL : num [1:2, 1:5, 1:5, 1] 5 5 5 5 5 5 5 5 5 ...
#> ..$ LL_IL: logi NA
```

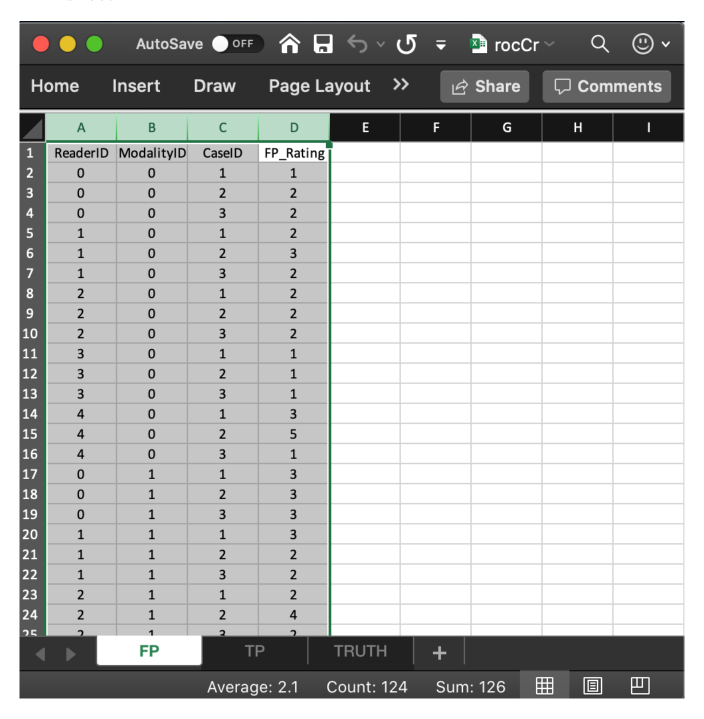
```
#>
                  :List of 3
   $ lesions
#>
    ..$ perCase: int [1:5] 1 1 1 1 1
               : num [1:5, 1] 1 1 1 1 1
#>
    ..$ IDs
    ..$ weights: num [1:5, 1] 1 1 1 1 1
#>
#>
   $ descriptions:List of 7
#>
    ..$ fileName
                    : chr "rocCr"
    ..$ type
                      : chr "ROC"
#>
                     : logi NA
#>
    ..$ name
#>
     ..$ truthTableStr: num [1:2, 1:5, 1:8, 1:2] 1 1 1 1 1 1 1 1 1 1 1 ...
#>
     ..$ design
                    : chr "FCTRL"
                    : Named chr [1:2] "0" "1"
#>
    ..$ modalityID
     ....- attr(*, "names")= chr [1:2] "0" "1"
#>
                     : Named chr [1:5] "0" "1" "2" "3" ...
    ....- attr(*, "names")= chr [1:5] "0" "1" "2" "3" ...
```

- In the above code chunk flag newExcelFileFormat is set to TRUE as otherwise columns D F in the Truth worksheet are ignored and the dataset is assumed to be factorial, with dataType "automatically" determined from the contents of the FP and TP worksheets. ¹
- Flag newExcelFileFormat = FALSE, the default, is for compatibility with older JAFROC format Excel files, which did not have columns D F in the Truth worksheet. Its usage is deprecated.
- The dataset object x is a list variable with 3 members: ratings, lesions and descriptions.
- The x\$ratings member contains 3 sub-lists.
 - The x\$ratings\$NL member, with dimension [2, 5, 8, 1], contains the ratings of normal cases. The first dimension (2) is the number of treatments, the second (5) is the number of readers and the third (8) is the total number of cases. For ROC datasets the fourth dimension is always unity. The five extra values ² in the third dimension, which are filled with NAs, are needed for compatibility with FROC datasets.
 - The x\$ratings\$LL, with dimension [2, 5, 5, 1], contains the ratings of abnormal cases. The third dimension (5) corresponds to the 5 diseased cases.
 - The x\$ratings\$LL_IL member, equal to NA'; this member is there for compatibility with LROC data,
 IL denotes incorrect-localizations.
- The x\$lesions member contains 3 sub-lists.
 - The x\$lesions\$perCase member is a vector with 5 ones representing the 5 diseased cases in the dataset.
 - The x\$lesions\$IDs member is an array with 5 ones.
 - The x\$lesions\$weights member is an array with 5 ones.
 - These are irrelevant for ROC datasets. They are there for compatibility with FROC datasets.
- The x\$descriptions member contains 7 sub-lists.
 - The x\$descriptions\$fileName member is the base name of the file that was read to create this dataset, "rocCr" in the current example, otherwise it is NA (the latter would apply, for example, for a simulated dataset).
 - The x\$descriptions\$type member indicates that this is an ROC dataset.
 - The x\$descriptions\$name member is the name of this dataset, if it is an embedded dataset, otherwise NA.
 - The x\$descriptions\$truthTableStr member, with dimension [2, 5, 8, 2], quantifies the structure of the dataset, as explained in TBA Vignette #3 (it is used to check for data entry errors).
 - The x\$descriptions\$design member specifies the dataset design, which is "FCTRL" in the present example (a factorial dataset).
 - The x\$descriptions\$modalityID member is a vector with two elements "0" and "1", naming the two
 modalities.
 - The x\$readerID member is a vector with five elements "0", "1", "2", "3" and "4", naming the five readers.

¹The assumptions underlying the "automatic" determination could be defeated by data entry errors.

 $^{^2}$ with only 3 non-diseased cases why does one need 8 values?

2.4 Correspondence between NL member of dataset and the FP worksheet



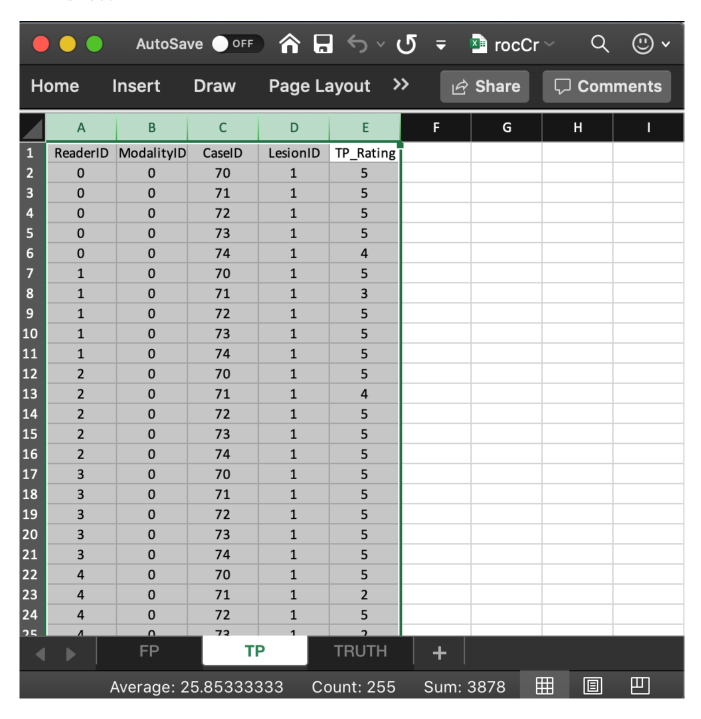
- The list member x array with dim = c(2,5,8,1).
 - The first dimension (2) comes from the number of modalities.
 - The second dimension (5) comes from the number of readers.
 - The third dimension (8) comes from the **total** number of cases.
 - The fourth dimension is alway 1 for an ROC dataset.
- The value of x\$ratings\$NL[1,5,2,1], i.e., 5, corresponds to row 15 of the FP table, i.e., to ModalityID = 0, ReaderID = 4 and CaseID = 2.
- The value of x\$ratings\$NL[2,3,2,1], i.e., 4, corresponds to row 24 of the FP table, i.e., to ModalityID 1, ReaderID 2 and CaseID 2.

• All values for case index > 3 and case index <= 8 are -Inf. For example the value of x\$ratings\$NL[2,3,4,1] is -Inf. This is because there are only 3 non-diseased cases. The extra length is needed for compatibility with FROC datasets.

2.5 Case-index vs. caseID

- Regardless of what order they occur in the worksheet, the non-diseased cases are always indexed first. In the current example the case indices are 1, 2 and 3, corresponding to the three non-diseased cases with caseIDs equal to 1, 2 and 3.
- Regardless of what order they occur in the worksheet, in the NL array the diseased cases are always indexed after the last non-diseased case. In the current example the case indices in the NL array are 4, 5, 6, 7 and 8, corresponding to the five diseased cases with caseIDs equal to 70, 71, 72, 73, and 74. In the LL array they are numbered 1, 2, 3, 4 and 5, corresponding to the five diseased cases with caseIDs equal to 70, 71, 72, 73, and 74. Some examples follow:
- x\$ratings\$NL[1,3,2,1], a FP rating, refers to ModalityID 0, ReaderID 2 and CaseID 2 (since the modality and reader IDs start with 0).
- x\$ratings\$NL[2,5,4,1], a FP rating, refers to ModalityID 1, ReaderID 4 and CaseID 70, the first diseased case; this is -Inf.
- x\$ratings\$NL[1,4,8,1], a FP rating, refers to ModalityID 0, ReaderID 3 and CaseID 74, the last diseased case; this is -Inf.
- x\$ratings\$NL[1,3,9,1], a FP rating, is an illegal value, as the third index cannot exceed 8.
- x\$ratings\$NL[1,3,8,2], a FP rating, is an illegal value, as the fourth index cannot exceed 1 for an ROC dataset.
- x\$ratings\$LL[1,3,1,1], a TP rating, refers to ModalityID 0, ReaderID 2 and CaseID 70, the first diseased case.
- x\$ratings\$LL[2,5,4,1], a TP rating, refers to ModalityID 1, ReaderID 4 and CaseID 73, the fourth diseased case.

2.6 Correspondence between LL member of dataset and the TP worksheet



- The list member x array with dim = c(2,5,5,1).
 - The first dimension (2) comes from the number of modalities.
 - The second dimension (5) comes from the number of readers.
 - The third dimension (5) comes from the number of diseased cases.
 - The fourth dimension is always 1 for an ROC dataset.
- The value of x\$ratings\$LL[1,1,5,1], i.e., 4, corresponds to row 6 of the TP table, i.e., to ModalityID = 0, ReaderID = 0 and CaseID = 74.
- The value of x\$ratings\$LL[1,2,2,1], i.e., 3, corresponds to row 8 of the TP table, i.e., to ModalityID = 0, ReaderID = 1 and CaseID = 71.

- The value of x\$ratings\$LL[1,4,4,1], i.e., 5, corresponds to row 21 of the TP table, i.e., to ModalityID = 0, ReaderID = 3 and CaseID = 74.
- The value of x\$ratings\$LL[1,5,2,1], i.e., 2, corresponds to row 23 of the TP table, i.e., to ModalityID = 0, ReaderID = 4 and CaseID = 71.
- There are no -Inf values in x\$ratings\$LL: any(x\$ratings\$LL == -Inf) = FALSE. This is true for any ROC dataset.

Chapter 3

Data format and reading FROC data

3.1 TBA How much finished

90%

3.2 Introduction

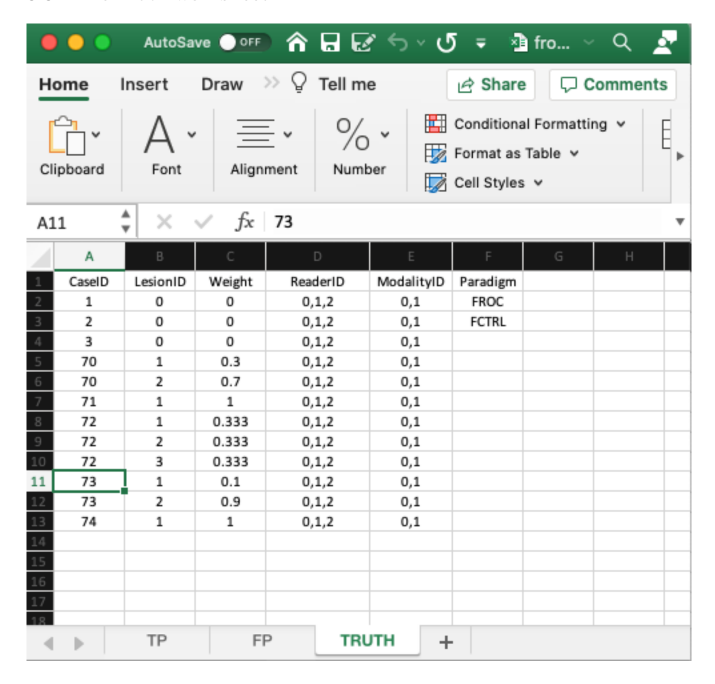
In the Free-response Receiver Operating Characteristic (FROC) paradigm the observer searches each case for signs of **localized disease** and marks and rates localized regions that are sufficiently suspicious for presence of disease. FROC data consists of **mark-rating pairs**, where each mark is a localized-region that was considered sufficiently suspicious for presence of a localized lesion and the rating is it's confidence level. As in the ROC paradigm, the rating can be an integer or quasi-continuous (e.g., 0-100), or a floating point value, as long as higher numbers represent greater confidence in presence of a lesion at the indicated region. This is termed a positive-directed confidence level scheme. By adopting a proximity criterion, the investigator classifies each mark as a lesion localization (LL) - if it is close to a real lesion - or a non-lesion localization (NL) otherwise.

The purpose of this chapter is to:

- Explain the data format of the input Excel file for FROC datasets.
- Explain the format of the FROC dataset.
- Explain the lesion distribution array returned by UtilLesionDistrVector().
- Explain the lesion weights array returned by UtilLesionWeightsMatrix().
- Details on the FROC paradigm are in my book (Chakraborty, 2017).

The chapter is illustrated with a toy data file, R/quick-start/frocCr.xlsx in which readers '0', '1' and '2' interpret 8 cases in two modalities, '0' and '1'. The design is 'factorial', abbreviated to FCTRL in the software; this is also termed a 'fully-crossed' design. The Excel file has three worksheets named Truth, NL (or FP) and LL (or TP).

3.3 The Truth worksheet



- The Truth worksheet contains 6 columns: CaseID, LesionID, Weight, ReaderID, ModalityID and Paradigm.
- Since a diseased case may have more than one lesion, the first five columns contain **at least** as many rows as there are cases (images) in the dataset. There are 8 cases in the dataset and 12 rows of data, because some of the diseased cases contain more than one lesion.
- CaseID: unique integers representing the cases in the dataset: '1', '2', '3', the 3 non-diseased cases, and '70', '71', '72', '73', '74', the 5 diseased cases. The ordering of the numbers is inconsequential. ¹
- LesionID: integers 0, 1, 2, etc.,
 - Each 0 represents a non-diseased case,
 - Each 1 represents the *first* lesion on a diseased case, 2 the *second* lesion, if present, and so on.
 - This field is zero for non-diseased cases '1', '2', '3'.

¹CaseID should not be so large that it cannot be represented in Excel by an integer; to be safe use unsigned short 8-bit integers. For example, 108057200 or 9971103254 are too large to be a valid caseID and may cause errors.

- For the first diseased case, i.e., '70', it is 1 for the first lesion and 2 for the second lesion.
- For the second diseased case i.e., '71', it is 1, as this case has only one lesion.
- For the third diseased case, i.e., '72', it is 1 for the first lesion, 2 for the second lesion and 3 for the third lesion
- For the fourth diseased case, i.e., '73', it is 1 for the first lesion and 2 for the second lesion.
- For the fifth diseased case i.e., '74', it is 1, as this case has only one lesion.
- There are 3 non-diseased cases in the dataset (the number of 0's in the LesionID column).
- There are 5 diseased cases in the dataset (the number of 1's in the LesionID column).
- Weight or clinical importance e.g., mortality associated with lesion:
 - non-negative floating point values
 - 0 for each non-diseased case
 - For each diseased case values that sum to unity.
 - A simple way to assign equal weights to all lesions in a case is to fill the Weight column with zeroes.

• LesionID

- Diseased case 70 has two lesions, with LesionIDs '1' and '2', and weights 0.3 and 0.7.
- Diseased case 71 has one lesion, with LesionID = 1, and Weight = 1.
- Diseased case 72 has three lesions, with LesionIDs 1, 2 and 3 and weights 1/3 each.
- Diseased case 73 has two lesions, with LesionIDs 1, and 2 and weights 0.1 and 0.9.
- Diseased case 74 has one lesion, with LesionID = 1 and Weight = 1.
- ReaderID: a comma-separated listing of readers, each represented by a unique **text label**, that have interpreted the case. In the example shown below each cell has the value '0, 1, 2'.
- There are 3 readers in the dataset, as each cell in the ReaderID column contains '0, 1, 2'.
- ModalityID: a comma-separated listing of modalities (or treatments), each represented by a unique integer, that apply to each case. In the example each cell has the value 0, 1. Each cell has to be text formatted.
- There are 2 modalities in the dataset, as each cell in the ModalityID column contains '0, 1'.
- Paradigm: The contents are FROC and FCTRL: this is an FROC dataset and the design is "factorial".

3.4 Reading the FROC dataset

The example shown above corresponds to file R/quick-start/frocCr.xlsx in the project directory. The next code chunk reads this file into an R object x.

```
frocCr <- "R/quick-start/frocCr.xlsx"</pre>
x <- DfReadDataFile(frocCr, newExcelFileFormat = TRUE)
str(x)
#> List of 3
#> $ ratings
                  :List of 3
    ..$ NL : num [1:2, 1:3, 1:8, 1:2] 1.02 2.89 2.21 3.01 2.14 ...
    ..$ LL : num [1:2, 1:3, 1:5, 1:3] 5.28 5.2 5.14 4.77 4.66 4.87 3.01 3.27 3.31 3.19 ...
#>
    .. $ LL_IL: logi NA
    $ lesions
#>
                  :List of 3
#>
    ..$ perCase: int [1:5] 2 1 3 2 1
#>
              : num [1:5, 1:3] 1 1 1 1 1 ...
#>
     ..$ weights: num [1:5, 1:3] 0.3 1 0.333 0.1 1 ...
   $ descriptions:List of 7
    ..$ fileName : chr "frocCr"
#>
                     : chr "FROC"
#>
    ..$ type
#>
     ..$ name
                     : logi NA
#>
     ..$ truthTableStr: num [1:2, 1:3, 1:8, 1:4] 1 1 1 1 1 1 1 1 1 1 ...
#>
                  : chr "FCTRL"
#>
     ..$ modalityID : Named chr [1:2] "0" "1"
     ....- attr(*, "names")= chr [1:2] "0" "1"
```

```
#> ..$ readerID : Named chr [1:3] "0" "1" "2" 
#> ...- attr(*, "names")= chr [1:3] "0" "1" "2"
```

This follows the general description in Chapter 1. The differences are described below.

- The x\$descriptions\$type member indicates that this is an FROC dataset.
- The x\$lesions\$perCase member is a vector whose contents reflect the number of lesions in each diseased case, i.e., 2, 1, 3, 2, 1 in the current example.
- The x\$lesions\$IDs member indicates the labeling of the lesions in each diseased case.

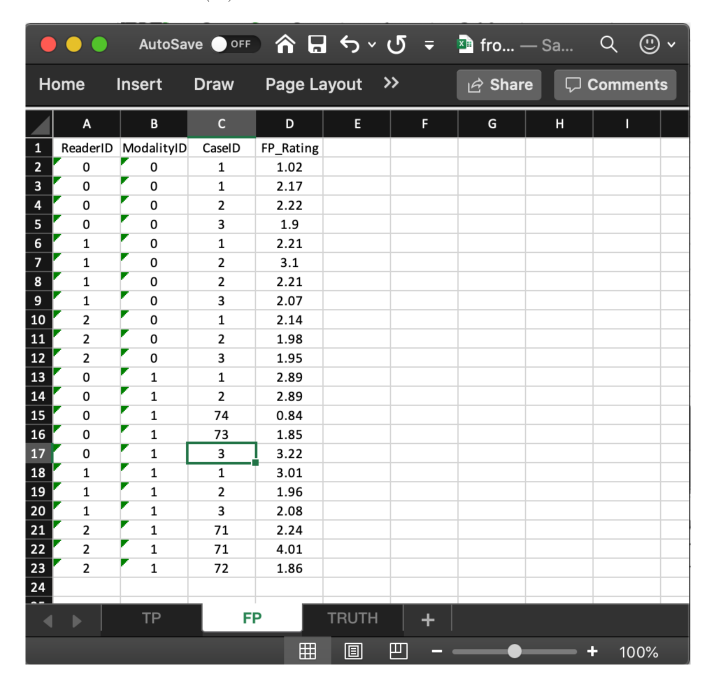
- This shows that the lesions on the first diseased case are labeled '1' and '2'. The -Inf is a filler used to denote a missing value. The second diseased case has one lesion labeled '1'. The third diseased case has three lesions labeled '1', '2' and '3', etc.
- The lesionWeight member is the clinical importance of each lesion. Lacking specific clinical reasons, the lesions should be equally weighted; this is *not* true for this toy dataset.

```
x$lesions$weights
#> [,1] [,2] [,3]
#> [1,] 0.3000000 0.7000000 -Inf
#> [2,] 1.0000000 -Inf -Inf
#> [3,] 0.3333333 0.3333333 0.3333333
#> [4,] 0.1000000 0.9000000 -Inf
#> [5,] 1.0000000 -Inf -Inf
```

- The first diseased case has two lesions, the first has weight 0.3 and the second has weight 0.7.
- The second diseased case has one lesion with weight 1.
- The third diseased case has three equally weighted lesions, each with weight 1/3. Etc.

3.5 The false positive (FP) ratings

These are found in the FP or NL worksheet.

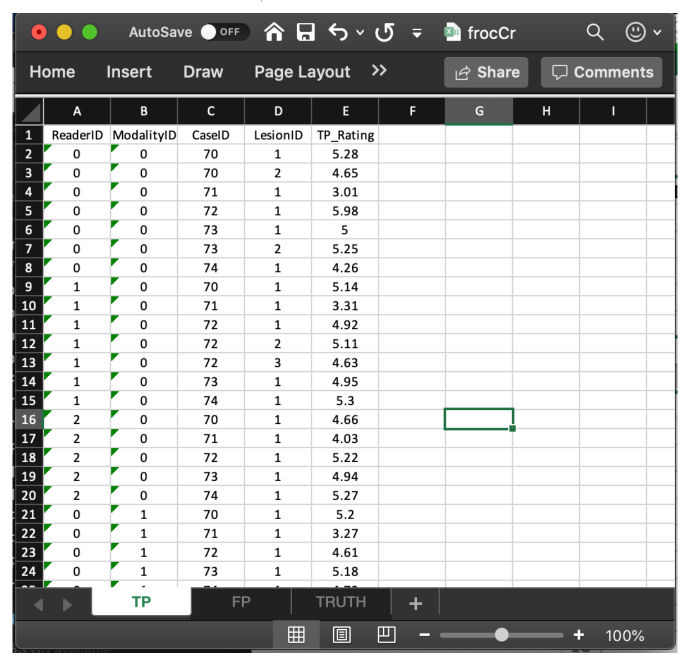


- It consists of 4 columns, of equal length. The common length is an integer random variable greater than or equal to zero. It could be zero if the dataset has no NL marks (a possibility if the lesions are very easy to find and the observer has perfect performance).
- In the example dataset, the common length is 22.
- ReaderID: the reader labels: these must be 0, 1, or 2, as declared in the Truth worksheet.
- ModalityID: the modality labels: must be 0 or 1, as declared in the Truth worksheet.
- CaseID: the labels of cases with NL marks. In the FROC paradigm NL events can occur on non-diseased and diseased cases.
- FP_Rating: the floating point ratings of NL marks. Each row of this worksheet yields a rating corresponding to the values of ReaderID, ModalityID and CaseID for that row.
- For ModalityID 0, ReaderID 0 and CaseID 1 (the first non-diseased case declared in the Truth worksheet), there is a single NL mark that was rated 1.02, corresponding to row 2 of the FP worksheet.
- Diseased cases with NL marks are also recorded in the FP worksheet. Some examples are seen at rows 15, 16 and 21, 22, 23.

- Rows 21 and 22 show that caseID = 71 got two NL marks, rated 2.24, 4.01.
- Since this is the *only* case with two NL marks, it determines the length of the fourth dimension of the x\$ratings\$NL list member, 2. Absent this case, the length would have been one.
- The case with the most NL marks determines the length of the fourth dimension of the x\$ratings\$NL list member.
- The reader should confirm that the ratings in x\$ratings\$NL reflect the contents of the FP worksheet.

3.6 The true positive (TP) ratings

These are found in the TP or LL worksheet, see below.



• This worksheet can only have diseased cases. The presence of a non-diseased case in this worksheet will generate an error.

- The common vertical length, 31 in this example, is a-priori unpredictable. The maximum possible length, assuming every lesion is marked for each modality, reader and diseased case, is 9 X 2 X 3 = 54. The 9 comes from the total number of non-zero entries in the LesionID column of the Truth worksheet, the 2 from the number of modalities and 3 from the number of readers.
- The fact that the actual length (31) is smaller than the maximum length (54) means that there are combinations of modality, reader and diseased cases on which some lesions were not marked.
- As examples, line 2 in the worksheet, the first lesion in CaseID equal to 70 was marked (and rated 5.28) in ModalityID 0 and ReaderID 0. Line 3 in the worksheet, the second lesion in CaseID equal to 70 was also marked (and rated 4.65) in ModalityID 0 and ReaderID 0. However, lesions 2 and 3 in CaseID = 72 were not marked (line 5 in the worksheet indicates that for this modality-reader-case combination only the first lesion was marked).
- The length of the fourth dimension of the x\$ratings\$LL list member, 3 in the present example, is determined by the diseased case (72) with the most lesions in the Truth worksheet.
- The reader should confirm that the ratings in x\$ratings\$LL reflect the contents of the TP worksheet.

3.7 On the distribution of numbers of lesions in diseased cases

• Consider a much larger dataset, dataset11, with structure as shown below (for descriptions of all embedded datasets the RJafroc documentation):

```
x <- dataset11
str(x)
#> List of 3
#>
   $ ratings
                 :List of 3
    ..$ NL : num [1:4, 1:5, 1:158, 1:4] -Inf -Inf -Inf -Inf -Inf ...
    ..$ LL : num [1:4, 1:5, 1:115, 1:20] -Inf -Inf -Inf -Inf -Inf ...
    ..$ LL_IL: logi NA
   $ lesions
#>
                 :List of 3
    ..$ perCase: int [1:115] 6 4 7 1 3 3 3 8 11 2 ...
#>
             : num [1:115, 1:20] 1 1 1 1 1 1 1 1 1 1 ...
    ..$ weights: num [1:115, 1:20] 0.167 0.25 0.143 1 0.333 ...
#>
    $ descriptions:List of 7
#>
#>
    ..$ fileName
                    : chr "dataset11"
    ..$ type
                     : chr "FROC"
#>
#>
    ..$ name
                    : chr "DOBBINS-1"
#>
     ..$ truthTableStr: num [1:4, 1:5, 1:158, 1:21] 1 1 1 1 1 1 1 1 1 1 ...
                 : chr "FCTRL"
#>
     ..$ design
    ..$ modalityID : Named chr [1:4] "1" "2" "3" "4"
#>
     ....- attr(*, "names")= chr [1:4] "1" "2" "3" "4"
#>
                     : Named chr [1:5] "1" "2" "3" "4"
#>
     ....- attr(*, "names")= chr [1:5] "1" "2" "3" "4"
```

- Focus for now in the 115 diseased cases.
- The numbers of lesions in these cases is contained in x\$lesions\$perCase.

```
x$lesions$perCase
               1 3 3 3 8 11 2 4 6 2 16 5 2 8 3
    [1]
       6
          4
            7
          7 3 2 5 2 2 7 6 6
                                4 10 20 12 6 4 7 12 5 1
                                                           5
   [26]
          1 2
               2
                 3 2
                                2
                                  2
                                     6
                                       3
                                          2
   [51]
        3
                      8 16 10
                             1
                                            2 4
                                                 6 10 11
                                                         1
        5
            9
                 6
                    8
                      3
                                  6
                                     3
                                        2
                                          1
                                            9
  [76]
          2
               6
                         8
                           7
                             1
                                1
                                               8 8 2 2 12
#> [101]
       1 3 1 2 2 1 1 1 1 3 1 1 1
                                        2
                                          1
```

• For example, the first diseased case contains 6 lesions, the second contains 4 lesions, the third contains 7 lesions, etc. and the last diseased case contains 1 lesion.

• To get an idea of the distribution of the numbers of lesions per diseased cases, one could interrogate this vector as shown below using the which() function:

```
for (el in 1:max(x$lesions$perCase)) cat(
  "number of diseased cases with", el, "lesions = ",
 length(which(x$lesions$perCase == el)), "\n")
#> number of diseased cases with 1 lesions = 25
#> number of diseased cases with 2 lesions = 23
#> number of diseased cases with 3 lesions = 13
#> number of diseased cases with 4 lesions = 10
#> number of diseased cases with 5 lesions = 5
#> number of diseased cases with 6 lesions = 11
#> number of diseased cases with 7 lesions = 6
#> number of diseased cases with 8 lesions = 8
#> number of diseased cases with 9 lesions = 2
#> number of diseased cases with 10 lesions = 3
#> number of diseased cases with 11 lesions = 3
#> number of diseased cases with 12 lesions = 3
#> number of diseased cases with 13 lesions = 0
#> number of diseased cases with 14 lesions = 0
#> number of diseased cases with 15 lesions = 0
\#> number of diseased cases with 16 lesions = 2
#> number of diseased cases with 17 lesions = 0
#> number of diseased cases with 18 lesions = 0
#> number of diseased cases with 19 lesions = 0
#> number of diseased cases with 20 lesions = 1
```

- This tells us that 25 cases contain 1 lesion
- Likewise, 23 cases contain 2 lesions
- Etc.

3.7.1 Definition of lesDistr array

• What is the fraction of (diseased) cases with 1 lesion, 2 lesions etc.

```
for (el in 1:max(x$lesions$perCase)) cat("fraction of diseased cases with", el, "lesions = ",
                                              length(which(x$lesions$perCase == el))/length(x$ratings$LL[1
#> fraction of diseased cases with 1 lesions = 0.2173913
#> fraction of diseased cases with 2 lesions = 0.2
#> fraction of diseased cases with 3 lesions = 0.1130435
#> fraction of diseased cases with 4 lesions = 0.08695652
#> fraction of diseased cases with 5 lesions = 0.04347826
#> fraction of diseased cases with 6 lesions = 0.09565217
#> fraction of diseased cases with 7 lesions = 0.05217391
#> fraction of diseased cases with 8 lesions = 0.06956522
#> fraction of diseased cases with 9 lesions = 0.0173913
#> fraction of diseased cases with 10 lesions = 0.02608696
#> fraction of diseased cases with 11 lesions = 0.02608696
#> fraction of diseased cases with 12 lesions = 0.02608696
#> fraction of diseased cases with 13 lesions = 0
#> fraction of diseased cases with 14 lesions = 0
#> fraction of diseased cases with 15 lesions =
#> fraction of diseased cases with 16 lesions = 0.0173913
#> fraction of diseased cases with 17 lesions = 0
\#> fraction of diseased cases with 18 lesions = 0
```

```
#> fraction of diseased cases with 19 lesions = 0
#> fraction of diseased cases with 20 lesions = 0.008695652
```

- This tells us that fraction 0.217 of (diseased) cases contain 1 lesion
- And fraction 0.2 of (diseased) cases contain 2 lesions
- Etc.
- This information is obtained using the function UtilLesionDistrVector()

```
lesDistr <- UtilLesionDistrVector(x)
lesDistr

#> [1] 0.217391304 0.200000000 0.113043478 0.086956522 0.043478261 0.095652174

#> [7] 0.052173913 0.069565217 0.017391304 0.026086957 0.026086957 0.026086957

#> [13] 0.017391304 0.008695652
```

- TBA The UtilLesionDistrVector() function returns an array with two columns and number of rows equal to the number of distinct non-zero values of lesions per case.
- The first column contains the number of distinct non-zero values of lesions per case, 14 in the current example.
- The second column contains the fraction of diseased cases with the number of lesions indicated in the first column.
- The second column must sum to unity

```
sum(UtilLesionDistrVector(x))
#> [1] 1
```

• The lesion distribution array will come in handy when it comes to predicting the operating characteristics from using the Radiological Search Model (RSM), as detailed in TBA Chapter 17.

3.8 TBA Definition of lesWghtDistr array

- This is returned by UtilLesionWeightsDistr().
- This contains the same number of rows as lesDistr.
- The number of columns is one plus the number of rows as lesDistr.
- The first column contains the number of distinct non-zero values of lesions per case, 14 in the current example.
- The second through the last columns contain the weights of cases with number of lesions per case corresponding to row 1.
- Missing values are filled with -Inf.

```
lesWghtDistr <- UtilLesionWeightsMatrixDataset(x, relWeights = 0)</pre>
cat("dim(lesDistr) =", dim(lesDistr),"\n")
#> dim(lesDistr) =
cat("dim(lesWghtDistr) =", dim(lesWghtDistr),"\n")
\#> dim(lesWghtDistr) = 14 15
cat("lesWghtDistr = \n\n")
#> lesWghtDistr =
lesWghtDistr
#>
                    [,2]
                                           [,4]
                                                      [,5]
                                                                 [,6]
                                                                             [,7]
         [,1]
                                [,3]
   [1,]
           1 1.00000000
                                -Inf
                                           -Inf
                                                      -Inf
                                                                 -Inf
                                                                             -Inf
#> [2,]
            2 0.50000000 0.50000000
                                           -Inf
                                                      -Inf
                                                                 -Inf
                                                                            -Inf
#> [3,]
            3 0.33333333 0.33333333 0.333333333
                                                      -Inf
                                                                 -Inf
                                                                            -Inf
#> [4,]
            4 0.25000000 0.25000000 0.25000000 0.25000000
                                                                 -Inf
                                                                            -Inf
#> [5,]
            5 0.20000000 0.20000000 0.20000000 0.20000000 0.20000000
                                                                            -Inf
   [6,]
            6 0.16666667 0.16666667 0.16666667 0.16666667 0.16666667
```

```
#>
   [7,]
           7 0.14285714 0.14285714 0.14285714 0.14285714 0.14285714 0.14285714
   [8,]
           8 0.12500000 0.12500000 0.12500000 0.12500000 0.12500000
#>
   [9,]
           #>
#> [10,]
          10 \ 0.10000000 \ 0.10000000 \ 0.100000000 \ 0.100000000 \ 0.100000000 \ 0.100000000
#> [11,]
          #> [12,]
          12 0.08333333 0.08333333 0.08333333 0.08333333 0.08333333
#> [13,]
          13 0.07692308 0.07692308 0.07692308 0.07692308 0.07692308 0.07692308
#> [14,]
          14 0.07142857 0.07142857 0.07142857 0.07142857 0.07142857 0.07142857
#>
              [,8]
                        [,9]
                                 [,10]
                                            [,11]
                                                      [,12]
                                                                [,13]
#>
             -Inf
   [1,]
                        -Inf
                                  -Inf
                                             -Inf
                                                       -Inf
                                                                 -Inf
#>
   [2,]
             -Inf
                        -Inf
                                  -Inf
                                             -Inf
                                                       -Inf
                                                                 -Inf
#>
   [3,]
                                             -Inf
                                                       -Inf
                                                                 -Inf
             -Inf
                        -Inf
                                  -Inf
#>
   [4,]
              -Inf
                        -Inf
                                  -Inf
                                             -Inf
                                                       -Inf
                                                                 -Inf
#>
   [5,]
              -Inf
                        -Inf
                                  -Inf
                                             -Inf
                                                       -Inf
                                                                 -Inf
#>
   [6,]
              -Inf
                        -Inf
                                  -Inf
                                             -Inf
                                                       -Inf
                                                                 -Inf
#>
   [7,] 0.14285714
                        -Inf
                                  -Inf
                                             -Inf
                                                       -Inf
                                                                 -Inf
   [8,] 0.12500000 0.12500000
                                                                 -Inf
#>
                                  -Inf
                                             -Inf
                                                       -Inf
   [9,] 0.11111111 0.11111111 0.11111111
                                             -Inf
                                                       -Inf
                                                                 -Inf
#> [10,] 0.10000000 0.10000000 0.10000000 0.10000000
                                                       -Inf
                                                                 -Inf
-Inf
#> [12,] 0.08333333 0.08333333 0.08333333 0.08333333 0.08333333 0.08333333
#> [13,] 0.07692308 0.07692308 0.07692308 0.07692308 0.07692308 0.07692308
#> [14,] 0.07142857 0.07142857 0.07142857 0.07142857 0.07142857 0.07142857
#>
             [,14]
                       [,15]
#>
   [1,]
              -Inf
                        -Inf
#>
   [2,]
             -Inf
                        -Inf
#>
   [3,]
             -Inf
                        -Inf
#>
   [4,]
              -Inf
                        -Inf
#>
   [5,]
             -Inf
                        -Inf
#>
   [6,]
             -Inf
                        -Inf
#>
   [7,]
             -Inf
                        -Inf
#>
   [8,]
             -Inf
                        -Inf
#>
   [9,]
              -Inf
                        -Inf
#> [10,]
              -Inf
                        -Inf
#> [11,]
              -Inf
                        -Inf
#> [12,]
              -Inf
                        -Inf
#> [13,] 0.07692308
                        -Inf
#> [14,] 0.07142857 0.07142857
```

- Row 3 corresponds to 3 lesions per case and the weights are 1/3, 1/3 and 1/3.
- Row 13 corresponds to 16 lesions per case and the weights are 0.06250000, 0.06250000, ..., repeated 13 times.
- Note that the number of rows is less than the maximum number of lesions per case (20).
- This is because some configurations of lesions per case (e.g., cases with 13 lesions per case) do not occur in this dataset.

Chapter 4

Data format and reading LROC data

4.1 TBA How much finished

70%

4.2 Introduction

In the Localization Receiver Operating Characteristic (LROC) paradigm (Starr et al., 1977, 1975; Swensson, 1996) the observer is restricted to at most one mark-rating pair per case. Additionally, each diseased case has *exactly* one lesion. On a diseased case and if the mark is close to the real lesion the investigator classifies the mark as a correct-localization (CL). Otherwise it is classified as an incorrect-localization (IL). On a non-diseased case the mark is always classified as a false-positive (FP).

The paradigm is illustrated with a few toy data files, R/quick-start/lroc?.xlsx, where ? is 1 or 2. These files illustrate two-modality three-reader LROC datasets with 3 non-diseased and 5 diseased cases.

File lroc1.xlsx illustrates the classic (i.e., as originally introduced) LROC paradigm where one mark per case is forced.

File lroc2.xlsx illustrates the paradigm when one mark-rating pair per case is not forced. There is some history behind this: the basic issue was what was the observer supposed to do when there there was nothing to report? Swensson initially thought that even if there was nothing to report, there must be a region, selected from the set of very low confidence regions, which was most likely to be a lesion. Most radiologists had difficulty with the forced localization requirement - if they saw nothing suspicious, why should they be forced to indicate a location. The paradigm was subsequently altered so that if the confidence level was below a certain value, say 12 percent on a 0 to 100 scale, the radiologist did not have to report a location. LROCFIT software was modified accordingly, and internal to the software the mark was assigned a random location - which ended up being classified as an incorrect-localization in most cases. The fact that there are cases with nothing to report is accounted for in the radiological search model.

4.3 Truth worksheet

Н	ome	Insert	Draw	⇒> Q Tell m	ie	🖒 Share	□ Co	omments
И	Α	В	С	D	E	F	G	Н
1	CaseID	LesionID	Weight	ReaderID	ModalityID	Paradigm		
2	1	0	0	0,1,2	0,1	LROC		
3	2	0	0	0,1,2	0,1	FCTRL		
4	3	0	0	0,1,2	0,1			
5	70	1	0	0,1,2	0,1			
6	71	1	0	0,1,2	0,1			
7	72	1	0	0,1,2	0,1			
8	73	1	0	0,1,2	0,1			
9	74	1	0	0,1,2	0,1			
10								
11								
12								
13								
14								
15								
16						<u> </u>		
17								
18								
19								
20								
21								
22								
23								
24								

- The Truth worksheet is similar to that described previously for the ROC and LROC paradigms. The only difference is the first entry in the Paradigm column, which is LROC.
- Since a diseased case has one lesion, the first five columns contain as many rows as there are cases in the dataset. There being 8 cases in the dataset, there are 8 rows of data.
- CaseID: unique integers representing the cases in the dataset: '1', '2', '3', the 3 non-diseased cases, and '70', '71', '72', '73', '74', the 5 diseased cases.
- LesionID: integers 0 or 1.
 - Each 0 represents a non-diseased case,
 - Each 1 represents the solitary lesion in the diseased case.
- There are 3 non-diseased cases in the dataset (the number of 0's in the LesionID column).
- There are 5 diseased cases in the dataset (the number of 1's in the LesionID column).

- Weight: this column is filled with zeroes. With one lesion per case, the weights are irrelevant.
- ReaderID: In the example shown each cell has the value '0, 1, 2'. There are 3 readers in the dataset, labeled 0, 1 and 2.
- ModalityID: In the example each cell has the value 0, 1. There are 2 modalities in the dataset, labeled 0 and 1.
- Paradigm: The contents are LROC and FCTRL: this is an LROC dataset and the design is "factorial".

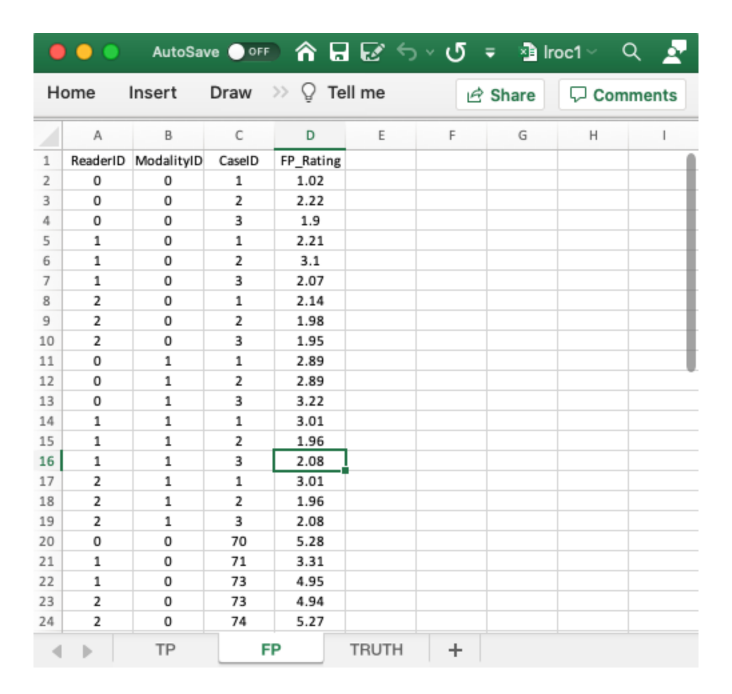
4.4 TP worksheet, forced localization true

Н	ome	Insert	Draw	>>	ell me	Ė	Share	□ Con	nments
7	A	В	С	D	Е	F	G	Н	
1	ReaderID	ModalityID	CaseID	LesionID	TP_Rating				
2	0	0	71	1	3.01				
3	0	0	72	1	5.98				
4	0	0	73	1	5				
5	0	0	74	1	4.26				
6	1	0	70	1	5.14				
7	1	0	72	1	4.92				
8	1	0	74	1	5.3				
9	2	0	70	1	4.66				
0	2	0	71	1	4.03				
1	2	0	72	1	5.22				
2	0	1	70	1	5.2				
13	0	1	72	1	4.61				
4	0	1	73	1	5.18				
5	0	1	74	1	4.72				1
6	1	1	71	1	3.19				
7	1	1	72	1	5.2				
8	1	1	74	1	5.01				
9									
0.5									
1									
2									
3									
4									

- The TP worksheet is similar to that described previously for the ROC and FROC paradigms.
- This worksheet can only have diseased cases. The presence of a non-diseased case in this worksheet will generate an error.
- The key difference is that for each modality-reader and diseased-case combination there can be at most one entry. Also, if a particular combination is missing in the TP worksheet then it must appear in the FP

- worksheet. This is because this is a forced-mark-per-case dataset.
- There can be at most 30 rows of data in this worksheet: 2 modalities times 3 readers times 5 diseased cases. Since there in fact only 17 rows of data, the missing 13 rows must occur in the FP worksheet.
- Recall that each entry in the TP worksheet represents a correct localization while each missing entry represents an incorrect localization. The incorrect localizations are recorded in the FP worksheet.

4.5 FP worksheet, forced localization true



Н	ome	Insert	Draw	>> 🔉 1	Tell me	Ė	Share	□ Con	nments
Z	A	В	С	D	E	F	G	Н	1
25	0	1	71	3.27					
26	1	1	70	4.77					
27	1	1	73	5.39					
28	2	1	70	4.87					
29	2	1	71	1.94					
30	2	1	72	5.39					
31	2	1	73	5.01					
32	2	1	74	5.01					
33									
34									
35									
36									
37									
38									
39									
40									
41									
42									
43									
44									
45									
46									
47									
48									

- The FP worksheet is similar to that described previously for the ROC and FROC paradigms.
- Because of the forced mark requirement, there are 18 rows of data corresponding to non-diseased cases: 2 modalities times 3 readers times 3 non-diseased cases. The missing 13 rows from the TP worksheet are listed next; these correspond to the incorrect localizations on diseased cases. Therefore, the total number of rows in this worksheet is 18 + 13 = 31.
- As an example, it is seen that for modalityID = 0 and readerID = 0, caseID = 70 does not appear in the TP worksheet. The lesion on this case was not localized; therefore it must appear in the FP worksheet as an incorrect localization, which is seen to be true in the FP worksheet.
- As another example, for modalityID = 0 and readerID = 1, caseID = 71 does not appear in the TP worksheet; instead it appears in the FP worksheet.
- As a final example, for modalityID = 1 and readerID = 2, none of the diseased cases appears in the TP worksheet; instead they all appear in the FP worksheet.

4.6 Reading forced localization true LROC dataset

The images shown above correspond to file R/quick-start/lroc1.xlsx. The next code chunk reads this file into an R object x1. Note the usage of the lrocForcedMark flag, which is set to TRUE, because this is a forced localization LROC dataset.

```
lroc1 <- "R/quick-start/lroc1.xlsx"</pre>
x1 <- DfReadDataFile(lroc1, newExcelFileFormat = TRUE, lrocForcedMark = T)
str(x1)
#> List of 3
#> $ ratings
                  :List of 3
     ..$ NL : num [1:2, 1:3, 1:8, 1] 1.02 2.89 2.21 3.01 2.14 3.01 2.22 2.89 3.1 1.96 ...
#>
     ..$ LL : num [1:2, 1:3, 1:5, 1] -Inf 5.2 5.14 -Inf 4.66 ...
     ..$ LL_IL: num [1:2, 1:3, 1:5, 1] 5.28 -Inf -Inf 4.77 -Inf ...
   $ lesions
                  :List of 3
#>
     ..$ perCase: int [1:5] 1 1 1 1 1
              : num [1:5, 1] 1 1 1 1 1
#>
     ..$ IDs
#>
    ..$ weights: num [1:5, 1] 1 1 1 1 1
#> $ descriptions:List of 7
    ..$ fileName : chr "lroc1"
#>
#>
    ..$ type
                    : chr "LROC"
#>
    ..$ name
                    : logi NA
#>
    ..$ truthTableStr: num [1:2, 1:3, 1:8, 1:2] 1 1 1 1 1 1 1 1 1 1 1 ...
#>
     ..$ design
                 : chr "FCTRL"
     ..$ modalityID : Named chr [1:2] "0" "1"
#>
#>
    ... - attr(*, "names") = chr [1:2] "0" "1"
     ..$ readerID
                     : Named chr [1:3] "0" "1" "2"
#>
    ....- attr(*, "names")= chr [1:3] "0" "1" "2"
```

This follows the general description in Chapter 1. The differences are described below.

• x1\$ratings\$NL is a [2,3,8,1] dimension vector. For each modality and reader, only the first three elements, corresponding to the three non-diseased cases, are finite, the rest are -Inf.

For example:

```
x1$ratings$NL[1,1,,1]
#> [1] 1.02 2.22 1.90 -Inf -Inf -Inf -Inf
```

• x1ratingsLLis a [2,3,5,1] dimension vector. For each modality and reader, only the first three elements, corresponding to the three non-diseased cases, are finite, the rest are-Inf.

For example, since none of the lesions are localized for modalityID = 1 (second modality) and readerID = 2 (third reader), the following code yields a vector consisting of five -Inf values:

```
x1$ratings$LL[2,3,,1]
#> [1] -Inf -Inf -Inf -Inf
```

• x1\$ratings\$LL_IL is a [2,3,5,1] dimension vector. These contain the ratings of incorrect localizations on diseased cases. For the just preceding modality-reader combination, this yields a vector with 5 finite values, the ratings of incorrect localizations for modalityID = 1 and readerID = 2.

```
x1$ratings$LL_IL[2,3,,1]
#> [1] 4.87 1.94 5.39 5.01 5.01
```

4.7 TP worksheet, forced localization false

Н	ome	Insert	Draw	>> Q Te	ell me		☆ Share	₽ Co	omment
И	Α	В	С	D	Е	F	G	Н	ı
1	ReaderID	ModalityID	CaseID	LesionID	TP_Rating				
2	0	0	71	1	3.01				
3	0	0	72	1	5.98				
4	0	0	73	1	5				
5	0	0	74	1	4.26				
6	1	0	70	1	5.14				
7	1	0	72	1	4.92				
8	1	0	74	1	5.3				
9	2	0	70	1	4.66				
10	2	0	71	1	4.03				
11	2	0	72	1	5.22				
12	0	1	70	1	5.2				
13	0	1	72	1	4.61				
14	0	1	73	1	5.18				
15	0	1	74	1	4.72				
16	1	1	71	1	3.19				
17	1	1	72	1	5.2				
18	1	1	74	1	5.01				
19									
20									
21									
22									
23									
2.4									
4	>	TP	F	P	TRUTH	+			

4.8 FP worksheet, forced localization false

Н	ome	Insert	Draw	>>	ell me		☆ Share	□ C	omments
/	Α	В	С	D	Е	F	G	Н	ı
1	ReaderID	ModalityID	CaseID	FP_Rating					
2	0	0	1	1.02					
3	0	0	2	2.22					
4	0	0	3	1.9					
5	1	0	1	2.21					
6	1	0	2	3.1					
7	1	0	3	2.07					
8	2	0	1	2.14					
9	2	0	2	1.98					
10	2	0	3	1.95					
11	0	1	1	2.89					
12	0	1	2	2.89					
13	0	1	3	3.22					
14	1	1	1	3.01					
15	1	1	2	1.96					
16	1	1	3	2.08					
17	2	1	1	3.01					
18	2	1	2	1.96					
19	2	1	3	2.08					
20	0	0	70	5.28					
21	1	0	71	3.31					
22	1	0	73	4.95					
23	2	0	73	4.94					
24	-	^	74	F 27					
4	▶	TP	F	P	TRUTH	+			

Н	ome	Insert	Draw	>>	ell me		☆ Share	□ Co	omments
Z	Α	В	С	D	Е	F	G	Н	ı
23	2	0	73	4.94					
24	2	0	74	5.27					
25	0	1	71	3.27					
26	1	1	70	4.77					
27	1	1	73	5.39					
28	2	1	70	4.87					
29	2	1	71	1.94					
30	2	1	72	5.39					
31	2	1	73	5.01					
32									
33									
34									
35									
36									
37									
38									
39									
40									
41									
42									
43									
44									
45									
4	•	TP	F	Р	TRUTH	+			

- If a particular modality-reader-case combination is missing in the TP worksheet then it need not appear in the FP worksheet. This is because this is not a forced-mark-per-case dataset.
- As an example, modalityID = 1, readerID = 2 and caseID = 74 does not appear in either TP or FP worksheets.

4.9 Reading forced localization false LROC dataset

The next example is for file R/quick-start/lroc2.xlsx. The following code chunk reads this file into an R object x2. Note that for this dataset one must set the lrocForcedMark flag to FALSE, because this is *not* a forced localization LROC dataset. Setting lrocForcedMark flag to TRUE will generate an error.

```
lroc2 <- "R/quick-start/lroc2.xlsx"
x2 <- DfReadDataFile(lroc2, newExcelFileFormat = TRUE, lrocForcedMark = F)
str(x2)
#> List of 3
#> $ ratings :List of 3
#> ..$ NL : num [1:2, 1:3, 1:8, 1] 1.02 2.89 2.21 3.01 2.14 3.01 2.22 2.89 3.1 1.96 ...
```

```
..$ LL : num [1:2, 1:3, 1:5, 1] -Inf 5.2 5.14 -Inf 4.66 ...
    ..$ LL_IL: num [1:2, 1:3, 1:5, 1] 5.28 -Inf -Inf 4.77 -Inf ...
#>
                 :List of 3
#>
   $ lesions
#>
    ..$ perCase: int [1:5] 1 1 1 1 1
             : num [1:5, 1] 1 1 1 1 1
#>
     ..$ IDs
#>
    ..$ weights: num [1:5, 1] 1 1 1 1 1
#>
   $ descriptions:List of 7
#>
    ..$ fileName
                    : chr "lroc2"
#>
     ..$ type
                      : chr "LROC"
#>
    ..$ name
                     : logi NA
#>
     ..$ truthTableStr: num [1:2, 1:3, 1:8, 1:2] 1 1 1 1 1 1 1 1 1 1 1 ...
#>
                  : chr "FCTRL"
     ..$ design
     ..$ modalityID : Named chr [1:2] "0" "1"
#>
     ....- attr(*, "names")= chr [1:2] "0" "1"
#>
                    : Named chr [1:3] "0" "1" "2"
#>
     ..$ readerID
     ....- attr(*, "names")= chr [1:3] "0" "1" "2"
```

• The x2\$ratings\$LL array is a [2,3,5,1] dimension vector. For each modality and reader, only the first three elements, corresponding to the three non-diseased cases, are finite, the rest are -Inf.

For example, since none of the lesions are localized for modalityID = 1 (second modality) and readerID = 2 (third reader), the following code yields a vector consisting of five -Inf values:

```
x2$ratings$LL[2,3,,1]
#> [1] -Inf -Inf -Inf -Inf
```

• The x2\$ratings\$LL_IL is a [2,3,5,1] dimension vector. These contain the ratings of incorrect localizations on diseased cases. For the just preceding modality-reader combination, this yields a vector with 4 finite values, the ratings of incorrect localizations for modalityID = 1 and readerID = 2.

```
x2$ratings$LL_IL[2,3,,1]
#> [1] 4.87 1.94 5.39 5.01 -Inf
```

For this modality-reader combination case 74 (i.e., the fifth diseased case) was unmarked. It does not appear in either the TP or the FP worksheet.

4.10 Summary

The difference from the previous data structures is the existence of LL_IL in the ratings list, which contains the ratings of incorrect localizations. Recall that for ROC and FROC paradigms this member was NA. When the data obeys forced localization, the corresponding flag should be set to TRUE, otherwise it should be set to FALSE. The default value of this flag is NA, which will work for ROC or FROC datasets. For LROC datasets it should be set to T/F.

Chapter 5

DBM analysis text output

5.1 TBA How much finished

50%

5.2 Introduction

This chapter illustrates significance testing using the DBM method.

5.3 Analyzing the ROC dataset

This illustrates the StSignificanceTesting() function. The significance testing method is specified as "DBM" and the figure of merit FOM is specified as "Wilcoxon". The embedded dataset dataset03 is used.

```
ret <- StSignificanceTesting(dataset03, FOM = "Wilcoxon", method = "DBM")</pre>
```

5.4 Explanation of the output

The function returns a list with 5 members:

- FOMs: figures of merit.
- ANOVA: ANOVA tables.
- RRRC: random-reader random-case analyses results.
- FRRC: fixed-reader random-case analyses results.
- RRFC" random-reader fixed-case analyses results.

Let us consider them individually.

```
str(ret$FOMs)
#> List of 3
#> $ foms : 'data.frame': 2 obs. of 4 variables:
#> ..$ rdrREADER_1: num [1:2] 0.853 0.85
#> ..$ rdrREADER_2: num [1:2] 0.865 0.844
#> ..$ rdrREADER_3: num [1:2] 0.857 0.84
#> ..$ rdrREADER_4: num [1:2] 0.815 0.814
```

```
#> $ trtMeans :'data.frame': 2 obs. of 1 variable:

#> ..$ Estimate: num [1:2] 0.848 0.837

#> $ trtMeanDiffs:'data.frame': 1 obs. of 1 variable:

#> ..$ Estimate: num 0.0109
```

- FOMs is a list of 3
 - foms is a [2x4] dataframe: the figure of merit for each of of the four observers in the two treatments.
 - trtMeans is a [2x1] dataframe: the average figure of merit over all readers for each treatment.
 - trtMeanDiffs a [1x1] dataframe: the difference(s) of the reader-averaged figures of merit for all different-treatment pairings. In this example, with only two treatments, there is only one different-treatment pairing.

```
ret$FOMs$foms

#> rdrREADER_1 rdrREADER_2 rdrREADER_3 rdrREADER_4

#> trtTREAT1 0.85345997 0.86499322 0.85730439 0.81524197

#> trtTREAT2 0.84961556 0.84350972 0.84011759 0.81433740

ret$FOMs$trtMeans

#> Estimate

#> trtTREAT1 0.84774989

#> trtTREAT2 0.83689507

ret$FOMs$trtMeanDiffs

#> Estimate

#> trtTREAT1 - trtTREAT2 0.010854817
```

```
str(ret$ANOVA)
#> List of 4
#>
  $ TRCanova
                   :'data.frame':
                                  8 obs. of 3 variables:
#>
    ..$ SS: num [1:8] 0.0236 0.2052 52.5284 0.0151 6.41 ...
    ..$ DF: num [1:8] 1 3 99 3 99 297 297 799
#>
    ..$ MS: num [1:8] 0.02357 0.06841 0.53059 0.00502 0.06475 ...
#> $ VarCom
                  :'data.frame': 6 obs. of 1 variable:
    ...$ Estimates: num [1:6] 3.78e-05 5.13e-02 -7.13e-04 -2.89e-03 2.79e-02 ...
#>
    $ IndividualTrt:'data.frame':
                                   3 obs. of 3 variables:
    ..$ DF
#>
                 : num [1:3] 3 99 297
#>
     ..$ TrtTREAT1: num [1:3] 0.0493 0.294 0.105
     ..$ TrtTREAT2: num [1:3] 0.0242 0.3014 0.1034
#>
#> $ IndividualRdr:'data.frame':
                                  3 obs. of 5 variables:
    ..$ DF
                   : num [1:3] 1 99 99
#>
    ..$ rdrREADER 1: num [1:3] 0.000739 0.203875 0.091559
#>
#>
     ..$ rdrREADER_2: num [1:3] 0.0231 0.2234 0.0803
#>
    ..$ rdrREADER_3: num [1:3] 0.0148 0.2142 0.0612
     ..$ rdrREADER_4: num [1:3] 4.09e-05 2.85e-01 6.06e-02
```

• ANOVA is a list of 4

- TRCanova is a [8x3] dataframe: the treatment-reader-case ANOVA table, see below, where SS is the sum of squares, DF is the denominator degrees of freedom and MS is the mean squares, and T = treatment, R = reader, C = case, TR = treatment-reader, TC = treatment-case, RC = reader-case, TRC = treatment-reader-case.
- VarCom is a [6x1] dataframe: the variance components, see below, where varR is the reader variance, varC is the case variance, varTR is the treatment-reader variance, varTC is the treatment-case variance, varRC is the reader-case variance, and varTRC is the treatment-reader-case variance.
- IndividualTrt is a [3x3] dataframe: the individual treatment variance components averaged over all readers, see below, where msR is the mean square reader, msC is the mean square case and msRC is the mean square reader-case.

IndividualRdr is a [3x5] dataframe: the individual reader variance components averaged over treatments, see below, where msT is the mean square treatment, msC is the mean square case and msTC is the mean square treatment-case.

```
ret$ANOVA$TRCanova
                   SS DF
#>
                                    MS
#> T
          0.023565410 1 0.0235654097
#> R
          0.205217999
                      3 0.0684059998
        52.528398680 99 0.5305898857
#> C
#> TR
         0.015060792
                       3 0.0050202641
#> TC
         6.410048814 99 0.0647479678
#> RC
         39.242953812 297 0.1321311576
#> TRC
         22.660077641 297 0.0762965577
#> Total 121.085323149 799
ret$ANOVA$VarCom
              Estimates
#> VarR
        3.7755679e-05
#> VarC 5.1250915e-02
#> VarTR -7.1276294e-04
#> VarTC -2.8871475e-03
#> VarRC 2.7917300e-02
#> VarErr 7.6296558e-02
ret$ANOVA$IndividualTrt
            TrtTREAT1
        DF
                         TrtTREAT2
        3 0.049266349 0.024159915
#> msC
      99 0.293967531 0.301370323
#> msRC 297 0.105047872 0.103379843
ret$ANOVA$IndividualRdr
       DF rdrREADER 1 rdrREADER 2 rdrREADER 3 rdrREADER 4
#> msT
        1 0.00073897606 0.023077021 0.014769293 0.00004091217
#> msC 99 0.20387477465 0.223441908 0.214246773 0.28541990211
#> msTC 99 0.09155873437 0.080279256 0.061228980 0.06057067104
str(ret$RRRC)
#> List of 3
#> $ FTests
                    :'data.frame': 2 obs. of 4 variables:
    ..$ DF : num [1:2] 1 3
#>
    ..$ MS : num [1:2] 0.02357 0.00502
    ..$ FStat: num [1:2] 4.69 NA
#>
#>
    ..$ p
            : num [1:2] 0.119 NA
                   :'data.frame': 1 obs. of 7 variables:
#> $ ciDiffTrt
#>
    ..$ Estimate: num 0.0109
#>
    \dots$ StdErr : num 0.00501
    ..$ DF
              : num 3
#>
#> ..$ t
               : num 2.17
    ..$ PrGTt : num 0.119
#>
#>
    ..$ CILower : num -0.00509
#>
    ..$ CIUpper : num 0.0268
#> $ ciAvgRdrEachTrt:'data.frame': 2 obs. of 5 variables:
    ..$ Estimate: num [1:2] 0.848 0.837
#>
    ..$ StdErr : num [1:2] 0.0244 0.0236
#>
#> ..$ DF
             : num [1:2] 70.1 253.6
#> ..$ CILower : num [1:2] 0.799 0.79
```

• RRRC, a list of 3 containing results of random-reader random-case analyses

..\$ CIUpper : num [1:2] 0.896 0.883

- FTtests: is a [2x4] dataframe: results of the F-tests, see below, where FStat is the F-statistic and p is the p-value. The first row is the treatment effect and the second is the error term.
- ciDiffTrt: is a [1x7] dataframe: the confidence intervals between different-treatments, see below, where StdErr is the standard error of the estimate, t is the t-statistic and PrGTt is the p-value.
- ciAvgRdrEachTrt: is a [2x5] dataframe: the confidence intervals for each treatment, averaged over all readers in the treatment, see below, where CILower is the lower 95% confidence interval and CIUpper is the upper 95% confidence interval.

```
ret$RRRC$FTests
            DF
                         MS
#>
                                FStat
                                               p
#> Treatment 1 0.0235654097 4.6940577 0.11883786
             3 0.0050202641
                               NA
ret$RRRC$ciDiffTrt
                         Estimate
                                        StdErr DF
                                                                 PrGTt
#> trtTREAT1-trtTREAT2 0.010854817 0.0050101218 3 2.1665774 0.11883786
                            CILower
                                        CIUpper
#> trtTREAT1-trtTREAT2 -0.0050896269 0.026799261
ret$RRRC$ciAvgRdrEachTrt
#>
              Estimate
                            StdErr
                                           DF
                                                            CIUpper
                                                 CILower
#> trtTREAT1 0.84774989 0.024402152 70.121788 0.79908282 0.89641696
#> trtTREAT2 0.83689507 0.023566416 253.644028 0.79048429 0.88330585
```

```
str(ret$FRRC)
#> List of 4
#> $ FTests
                     :'data.frame':
                                       2 obs. of 4 variables:
    ..$ DF : num [1:2] 1 99
    ..$ MS : num [1:2] 0.0236 0.0647
#>
    ..$ FStat: num [1:2] 0.364 NA
           : num [1:2] 0.548 NA
#>
    ..$ p
#>
   $ ciDiffTrt
                     :'data.frame': 1 obs. of 7 variables:
#>
    ..$ Estimate: num 0.0109
#>
    ..$ StdErr : num 0.018
#>
    ..$ DF
                : num 99
               : num 0.603
#>
    ..$ t
#>
    ..$ PrGTt : num 0.548
    ..$ CILower : num -0.0248
#>
#>
    ..$ CIUpper : num 0.0466
#>
   $ ciAvqRdrEachTrt :'data.frame': 2 obs. of 5 variables:
    ..$ Estimate: num [1:2] 0.848 0.837
#>
#>
    ..$ StdErr : num [1:2] 0.0271 0.0274
    ..$ DF
              : num [1:2] 99 99
#>
    ..$ CILower : num [1:2] 0.794 0.782
#>
    ..$ CIUpper : num [1:2] 0.902 0.891
   $ ciDiffTrtEachRdr:'data.frame': 4 obs. of 7 variables:
#>
    ..$ Estimate: num [1:4] 0.003844 0.021483 0.017187 0.000905
#>
    ..$ StdErr : num [1:4] 0.0428 0.0401 0.035 0.0348
#>
#>
    ..$ DF
               : num [1:4] 99 99 99 99
#>
                : num [1:4] 0.0898 0.5362 0.4911 0.026
    ..$ PrGTt : num [1:4] 0.929 0.593 0.624 0.979
#>
    ..$ CILower : num [1:4] -0.0811 -0.058 -0.0522 -0.0682
    ..$ CIUpper : num [1:4] 0.0888 0.101 0.0866 0.07
```

- FRRC, a list of 4 containing results of fixed-reader random-case analyses
 - FTtests: is a [2x4] dataframe: results of the F-tests, see below.
 - ciDiffTrt: is a [1x7] dataframe: the confidence intervals between different-treatments, see below.

- ciAvgRdrEachTrt: is a [2x5] dataframe: the confidence intervals for the average reader over each treatment
- ciDiffTrtEachRdr: is a [4x7] dataframe: the confidence intervals for each different-treatment pairing for each reader.

```
ret$FRRC$FTests
                         MS
                                 FStat
#> Treatment 1 0.023565410 0.36395597 0.54769704
           99 0.064747968
#> Error
                                    NA
ret$FRRC$ciDiffTrt
#>
                          Estimate
                                        StdErr DF
                                                           t.
                                                                  PrGTt
#> trtTREAT1-trtTREAT2 0.010854817 0.017992772 99 0.60328764 0.54769704
                            CILower
                                       CIUpper
#> trtTREAT1-trtTREAT2 -0.024846746 0.04655638
ret$FRRC$ciAvgRdrEachTrt
               Estimate
                             StdErr DF
                                          CILower
#> trtTREAT1 0.84774989 0.027109386 99 0.79395898 0.90154079
#> trtTREAT2 0.83689507 0.027448603 99 0.78243109 0.89135905
ret$FRRC$ciDiffTrtEachRdr
                                         Estimate
                                                       StdErr DF
#> rdrREADER_1::trtTREAT1-trtTREAT2 0.00384441429 0.042792227 99 0.089839080
#> rdrREADER_2::trtTREAT1-trtTREAT2 0.02148349163 0.040069753 99 0.536152334
#> rdrREADER_3::trtTREAT1-trtTREAT2 0.01718679331 0.034993994 99 0.491135520
#> rdrREADER_4::trtTREAT1-trtTREAT2 0.00090456807 0.034805365 99 0.025989329
#>
                                         PrGTt
                                                    CILower
#> rdrREADER_1::trtTREAT1-trtTREAT2 0.92859660 -0.081064648 0.088753476
#> rdrREADER 2::trtTREAT1-trtTREAT2 0.59305592 -0.058023592 0.100990575
#> rdrREADER_3::trtTREAT1-trtTREAT2 0.62441761 -0.052248882 0.086622469
#> rdrREADER_4::trtTREAT1-trtTREAT2 0.97931817 -0.068156827 0.069965963
```

```
str(ret$RRFC)
#> List of 3
#> $ FTests
                    :'data.frame': 2 obs. of 4 variables:
    ..$ DF : num [1:2] 1 3
    ..$ MS : num [1:2] 0.02357 0.00502
    ..$ FStat: num [1:2] 4.69 NA
#>
             : num [1:2] 0.119 NA
    ..$ p
                   :'data.frame': 1 obs. of 7 variables:
#> $ ciDiffTrt
    ..$ Estimate: num 0.0109
#>
    ..$ StdErr : num 0.00501
               : num 3
#>
    ..$ DF
#>
    ..$ t
               : num 2.17
#>
    ..$ PrGTt : num 0.119
    ..$ CILower : num -0.00509
#>
#>
    ..$ CIUpper : num 0.0268
#> $ ciAvgRdrEachTrt:'data.frame': 2 obs. of 5 variables:
    ..$ Estimate: num [1:2] 0.848 0.837
#>
    ..$ StdErr : num [1:2] 0.0111 0.00777
    ..$ DF
              : num [1:2] 3 3
    ..$ CILower : num [1:2] 0.812 0.812
   ..$ CIUpper : num [1:2] 0.883 0.862
```

- RRFC, a list of 3 containing results of random-reader fixed-case analyses
 - FTtests: is a [2x4] dataframe: results of the F-tests, see below.
 - ciDiffTrt: is a [1x7] dataframe: the confidence intervals between different-treatments, see below.

- ciAvgRdrEachTrt: is a [2x5] data frame: the confidence intervals for the average reader over each over each treatment.

```
ret$RRFC$FTests
                       MS
                              FStat
                                             p
#> Treatment 1 0.0235654097 4.6940577 0.11883786
#> Error
          3 0.0050202641 NA
ret$RRFC$ciDiffTrt
                        Estimate StdErr DF
#> trtTREAT1-trtTREAT2 0.010854817 0.0050101218 3 2.1665774 0.11883786
                                       {\it CIUpper}
                            CILower
#> trtTREAT1-trtTREAT2 -0.0050896269 0.026799261
ret$RRFC$ciAvgRdrEachTrt
              Estimate
                           StdErr DF
                                       {\it CILower}
                                                  CIUpper
#> trtTREAT1 0.84774989 0.011098012 3 0.81243106 0.88306871
#> trtTREAT2 0.83689507 0.007771730 3 0.81216196 0.86162818
```

Chapter 6

OR analysis text output

6.1 TBA How much finished

90%

6.2 Introduction

This chapter illustrates significance testing using the DBM and OR methods.

6.3 Analyzing the ROC dataset

The only change is to specify method = "OR" in the significance testing function. The same dataset is used as was used in the previous chapter.

```
ret <- StSignificanceTesting(dataset03, FOM = "Wilcoxon", method = "OR")
```

6.4 Explanation of the output

The function returns a list with 5 members.

- FOMs: figures of merit, identical to that in the DBM method.
- ANOVA: ANOVA tables.
- RRRC: random-reader random-case analyses results.
- FRRC: fixed-reader random-case analyses results.
- RRFC" random-reader fixed-case analyses results.

Let us consider the ones that are different from the DBM method.

- ANOVA is a list of 4
 - TRanova is a [3x3] dataframe: the treatment-reader ANOVA table, see below, where SS is the sum of squares, DF is the denominator degrees of freedom and MS is the mean squares, and T = treatment, R = reader, TR = treatment-reader.

- VarCom is a [6x2] dataframe: the variance components, see below, where varR is the reader variance, varTR is the treatment-reader variance, Cov1, Cov2,Cov3 and Var are as defined in the OR model. The second column lists the correlations defined in the OR model.
- IndividualTrt is a [2x4] dataframe: the individual treatment mean-squares, variances and Cov_2 , averaged over all readers, see below, where msREachTrt is the mean square reader, varEachTrt is the variance and cov2EachTrt is Cov2EachTrt in each treatment.
- IndividualRdr is a [2x4] dataframe: the individual reader variance components averaged over treatments, see below, where msTEachRdr is the mean square treatment, varEachRdr is the variance and cov1EachRdr is Cov_1 for each reader.

```
ret$ANOVA$TRanova
#>
                 SS DF
                                  MS
#> T 0.00023565410 1 2.3565410e-04
#> R 0.00205217999 3 6.8406000e-04
#> TR 0.00015060792 3 5.0202641e-05
ret$ANOVA$VarCom
#>
              Estimates
                              Rhos
#> VarR
         2.3319942e-05
                                NA
#> VarTR -6.8389146e-04
         7.9168215e-04 0.51887172
#> Cov1
#> Cov2
          4.8363767e-04 0.31697811
         5.1250915e-04 0.33590059
#> Cov3
          1.5257762e-03
#> Va.r
ret$ANOVA$IndividualTrt
            DF
                  msREachTrt
                                varEachTrt
                                             cov2EachTrt
#> trtTREAT1 3 0.00049266349 0.0015227779 0.00047229915
#> trtTREAT2 3 0.00024159915 0.0015287746 0.00049497620
ret$ANOVA$IndividualRdr
               DF
                     msTEachRdr
                                  varEachRdr
                                               cov1EachRdr
#> rdrREADER_1 1 7.3897606e-06 0.0014771675 0.00056158020
#> rdrREADER_2 1 2.3077021e-04 0.0015186058 0.00071581326
#> rdrREADER_3 1 1.4769293e-04 0.0013773788 0.00076508897
#> rdrREADER_4 1 4.0912170e-07 0.0017299529 0.00112424616
```

- RRRC, a list of 3 containing results of random-reader random-case analyses
 - FTtests: is a [2x4] dataframe: results of the F-tests, see below, where FStat is the F-statistic and p is the p-value. The first row is the treatment effect and the second is the error term.
 - ciDiffTrt: is a [1x7] dataframe: the confidence intervals between different treatments, see below, where StdErr is the standard error of the estimate, t is the t-statistic and PrGTt is the p-value.
 - ciAvgRdrEachTrt: is a [2x5] dataframe: the confidence intervals for the average reader over each treatment, see below, where CILower is the lower 95% confidence interval and CIUpper is the upper 95% confidence interval.

```
ret$RRRC$FTests
#>
             DF
                           MS
                                  FStat
                                                  p
#> Treatment 1 2.3565410e-04 4.6940577 0.11883786
#> Error
              3 5.0202641e-05
ret$RRRC$ciDiffTrt
                          Estimate
                                          StdErr DF
                                                                   PrGTt
#> trtTREAT1-trtTREAT2 0.010854817 0.0050101218 3 2.1665774 0.11883786
                             CILower
                                          CIUpper
#> trtTREAT1-trtTREAT2 -0.0050896269 0.026799261
ret$RRRC$ciAvgRdrEachTrt
#>
               Estimate
                             StdErr
                                             DF
                                                   CILower
                                                              CIUpper
                                                                                C0112.
#> trtTREAT1 0.84774989 0.024402152 70.121788 0.79908282 0.89641696 0.00047229915
#> trtTREAT2 0.83689507 0.023566416 253.644028 0.79048429 0.88330585 0.00049497620
```

- FRRC, a list of 5 containing results of fixed-reader random-case analyses
 - FTtests: is a [2x4] dataframe: results of the chisquare-tests, see below. Here is a difference from DBM: in the OR method for FRRC the denominator degrees of freedom of the F-statistic is infinite, and the test becomes equivalent to a chisquare test with the degrees of freedom equal to I-1, where I is the number of treatments.
 - ciDiffTrt: is a [1x6] dataframe: the confidence intervals between different treatments, see below. An additional column lists
 - ciAvgRdrEachTrt: is a [2x5] dataframe: the confidence intervals for the average reader over each treatment
 - ciDiffTrtEachRdr: is a [4x6] dataframe: the confidence intervals for each different-treatment pairing for each reader.
 - IndividualRdrVarCov1: is a [4x2] dataframe: Var and Cov_1 for individual readers.

```
ret$FRRC$FTests
#>
                      MS
                              Chisq DF
                                               p
#> Treatment 0.0002356541 0.32101347 1 0.57099922
            0.0007340941
                                 NA NA
                                              NA
ret$FRRC$ciDiffTrt
                         Estimate
                                      StdErr
                                                             PrGTz
                                                                        CILower
#> trtTREAT1-trtTREAT2 0.010854817 0.019158472 0.56658051 0.57099922 -0.026695098
#>
                          CIUpper
#> trtTREAT1-trtTREAT2 0.048404732
ret$FRRC$ciAvgRdrEachTrt
#>
              Estimate
                            StdErr\ DF
                                         CILower
                                                   CIUpper
#> trtTREAT1 0.84774989 0.027109386 99 0.79461647 0.90088331
#> trtTREAT2 0.83689507 0.027448603 99 0.78309680 0.89069334
ret$FRRC$ciDiffTrtEachRdr
#>
                                        Estimate
                                                     StdErr
#> rdrREADER_1::trtTREAT1-trtTREAT2 0.00384441429 0.042792227 0.089839080
#> rdrREADER_2::trtTREAT1-trtTREAT2 0.02148349163 0.040069753 0.536152334
#> rdrREADER_3::trtTREAT1-trtTREAT2 0.01718679331 0.034993994 0.491135520
#> rdrREADER_4::trtTREAT1-trtTREAT2 0.00090456807 0.034805365 0.025989329
#>
                                        PrGTz
                                                  CILower
                                                              CIUpper
#> rdrREADER_1::trtTREAT1-trtTREAT2 0.92841509 -0.080026809 0.087715638
#> rdrREADER_3::trtTREAT1-trtTREAT2 0.62333060 -0.051400174 0.085773761
#> rdrREADER_4::trtTREAT1-trtTREAT2 0.97926585 -0.067312693 0.069121830
ret$FRRC$IndividualRdrVarCov1
#>
                varEachRdr
                             cov1EachRdr
#> rdrREADER 1 0.0014771675 0.00056158020
#> rdrREADER_2 0.0015186058 0.00071581326
#> rdrREADER 3 0.0013773788 0.00076508897
#> rdrREADER_4 0.0017299529 0.00112424616
```

- RRFC, a list of 3 containing results of random-reader fixed-case analyses
 - FTtests: is a [2x4] dataframe: results of the F-tests, see below.
 - ciDiffTrt: is a [1x7] dataframe: the confidence intervals between different treatments, see below.
 - ciAvgRdrEachTrt: is a [2x5] dataframe: the confidence intervals for the average reader over each over each treatment.

```
#> Estimate StdErr DF t PrGTt

#> trtTREAT1-trtTREAT2 0.010854817 0.0050101218 3 2.1665774 0.11883786

#> CILower CIUpper

#> trtTREAT1-trtTREAT2 -0.0050896269 0.026799261

ret$RRFC$ciAvgRdrEachTrt

#> Estimate StdErr DF CILower CIUpper

#> TrtTREAT1 0.84774989 0.011098012 3 0.81243106 0.88306871

#> TrtTREAT2 0.83689507 0.007771730 3 0.81216196 0.86162818
```

Chapter 7

OR analysis Excel output

7.1 TBA How much finished

90%

7.2 Introduction

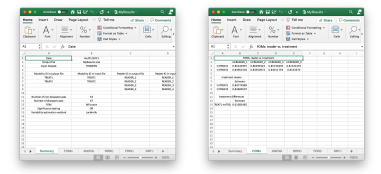
This chapter illustrates significance testing using the OR method. But, instead of the perhaps unwieldy output in Chapter 6, it generates an Excel output file containing the following worksheets:

- Summary
- FOMs
- ANOVA
- RRRC
- FRRC
- RRFC

7.3 Generating the Excel output file

This illustrates the UtilOutputReport() function. The arguments are the embedded dataset, dataset03, the same dataset as in the previous two chapters, the report file base name ReportFileBaseName is set to R/quick-start/MyResults, the report file extension ReportFileExt is set to xlsx, the FOM is set to "Wilcoxon", the method of analysis is set to "OR", and the flag overWrite = TRUE overwrites any existing file with the same name, as otherwise the program will pause for user input.

The following screen shots display the contents of the created file "R/quick-start/MyResults.xlsx".



Figure~7.1:~`Summary`~and~`FOMs`~worksheets~of~Excel~file~`R/quick-start/MyResults.xlsx``

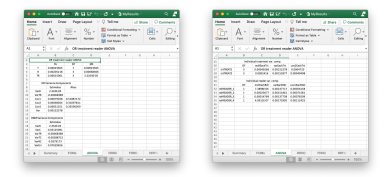


Figure 7.2: 'ANOVA' worksheet of Excel file 'R/quick-start/MyResults.xlsx'



Figure 7.3: 'RRRC', 'FRRC' and 'RRFC' worksheets of Excel file 'R/quick-start/MyResults.xlsx'

FROC analysis

Chapter 8

Analyzing FROC data

8.1 TBA How much finished

10%

8.2 Introduction

Analyzing FROC data is, apart from a single difference, very similar to analyzing ROC data. The crucial difference is the selection of an appropriate location-sensitive figure of merit. The reason is that the DBMH and ORH methods are applicable to any scalar figure of merit. Any appropriate FROC figure of merit reduces the mark rating data for a single dataset (i.e., a single treatment, a single reader and a number of cases) to a single scalar figure of merit.

The author recommends usage of the weighted AFROC figure of merit, where the lesions should be equally weighted, the default, unless there are strong clinical reasons for assigning unequal weights.

The chapter starts with analysis of a sample FROC dataset, #4 in Online Chapter 24. Any analysis should start with visualization of the relevant operating characteristic. Extensive examples are given using RJafroc implemented functions. Suggestions are made on how to report the results of a study (the suggestions apply equally to ROC studies). A method called *crossed-treatment analysis*, applicable when one has two treatment factors and their levels are crossed and one wishes to draw conclusions regarding the effect of treatments after averaging over all levels of the treatments.

8.3 Example 1

The following is a listing of file "mainAnalyzewAFROC.R". It performs both wAFROC and inferred ROC analyses of the same dataset and the results are saved to tables similar in structure to the Excel output tables shown for DBMH analysis of ROC data in §9.10.2. Empirical wAFROC-AUC and ROC-AUC for all combinations of treatments and readers, and reader-averaged AUCs for each treatment (Rdr. Avg.). The weighted AFROC results were obtained from worksheet FOMs in file FedwAfroc.xlsx. The highest rating AUC results were obtained from worksheet FOMs in file FedHrAuc.xlsx. The wAFROC-AUCs are smaller than the corresponding ROC-AUCs.

The datasets that come with this book are described in Online Chapter 24. Four of these are ROC datasets, one an LROC dataset and the rest (nine) are FROC datasets. For non-ROC datasets, the highest rating method was used to infer the corresponding ROC data. The datasets are identified in the code by strings contained in the string-array variable fileNames (line 7 - 8). Line 9 selects the dataset to be analyzed. In the example shown the "FED" dataset has been selected. It is a 5 treatment 4 radiologist FROC dataset1 acquired by Dr. Federica Zanca. Line 13 loads the dataset; this is done internal to the function loadDataFile(). Line 11 constructs the name of the wAFROC file and line 12 does the same for the ROC datafile. Line 15 which "spills over" to line 16 without the need for a special continuation character, generates an output file by performing DBMH significance testing (method = "DBMH")

using fom = "wAFROC", i.e., the wAFROC figure of merit – this is the critical change. If one changes this to fom = "HrAuc", lines 19 – 20, then inferred ROC analysis occurs. In either case the default analysis, i.e., option = "ALL" is used, i.e., random-reader random-case (RRRC), fixed-reader random-case (FRRC) and random-reader fixed-case (RRFC). Results are shown below for random-reader random-case only.

The results of wAFROC analysis are saved to FedwAfroc.xlsx and that of inferred ROC analysis are saved to FedHrAuc.xlsx. The output file names need to be explicitly stated as otherwise they would overwrite each other (as a time-saver, checks are made at lines 14 and 18 to determine if the analysis has already been performed, in which case it is skipped).

In the Excel data file the readers are named 1, 3, 4 and 5 – the software treats the reader names as labels. The author's guess is that for some reason complete data for reader 2 could not be obtained. The renumber = TRUE option has the effect of renumbering the readers 1 through 4. Without renumbering, the output would be aesthetically displeasing, but have no effect on the conclusions.

Figures of merit, empirical wAFROC-AUC and empirical ROC-AUC, and the corresponding reader averages for both analyses are summarized in Table 19.1. The weighted AFROC results were obtained by copy and paste operations from worksheet FOMs in file FedwAfroc.xlsx. The highest rating AUC results were obtained by similar operations from worksheet FOMs in Excel file FedHrAuc.xlsx. As expected, each wAFROC-AUC is smaller than the corresponding ROC-AUC.

Table 19.1: Empirical wAFROC-AUC and ROC-AUC for all combinations of treatments and readers, and reader-averaged AUCs for each treatment (Rdr. Avg.). The weighted AFROC results were obtained from worksheet FOMs in file FedWAfroc.xlsx. The highest rating AUC results were obtained from worksheet FOMs in file FedHrAuc.xlsx. The wAFROC-AUCs are smaller than the corresponding ROC-AUCs.

Table 19.2 shows results for RRRC analysis using the wAFROC-AUC FOM. The overall F-test of the null hypothesis that all treatments have the same reader-averaged FOM, rejected the NH: F(4, 36.8) = 7.8, p = 0.00012. The numerator degree of freedom ndf is I - 1 = 4. Since the null hypothesis is that all treatments have the same FOM, this implies that at least one pairing of treatments yielded a significant FOM difference. The control for multiple testing is in the formulation of the null hypothesis and no further Bonferroni-like2 correction is needed. To determine which specific pairings are significantly different one examines the p-values (listed under Pr>t) in the "95% CI's FOMs, treatment difference" portion of the table. It shows that the following differences are significant at alpha = 0.05, namely "1 - 3", "1 - 5", "2 - 3", "2 - 5", "3 - 4" and "4 - 5"; these are indicated by asterisks. The values listed under the "95% CI's FOMs, each treatment" portion of the table show that treatment 4 yielded the highest FOM (0.769) followed closely by treatments 2 and 1, while treatment 5 had the least FOM (0.714), slightly worse than treatment 3. This explains why the p-value for the difference 4 - 5 is the smallest (0.00007) of all the listed p-values in the "95% CI's FOMs, each treatment" portion of the table. Each instance where the p-value for the individual treatment comparisons yields a significant p-value is accompanied by a 95% confidence interval that does not include zero. The two statements of significance, one in terms of a p-value and one in terms of a CI, are equivalent. When it comes to presenting results for treatment FOM differences, I prefer the 95% CI but some journals insist on a p-value, even when it is not significant. Note that two sequential tests are involved, an overall F-test of the NH that all treatments have the same performance and only if this yields a significant results is one justified in looking at the p-values of individual treatment pairings.

Table 19.2: wAFROC-AUC analysis: results of random-reader random-case (RRRC) analysis, in worksheet "RRRC"". [ddf = denominator degrees of freedom of F-distribution. df = degrees of freedom of t-distribution. Stderr = standard error. CI = confidence interval. * = Significantly different at alpha = 0.05.]

Table 19.3 shows corresponding results for the inferred ROC-AUC FOM. Again the null hypothesis was rejected: F(4, 16.8) = 3.46, p = 0.032. This means at least two treatments have significantly different FOMs. Looking down the table, one sees that the same 6 pairs (as compared to wAFROC analysis) are significantly different, 1 - 3, 1 - 5, etc., as indicated by the asterisks. The last five rows of the table show that treatment 4 had the highest performance while treatment 5 had the lowest performance. At the 5% significance level, both methods yielded the same significant differences, but this is not always true. While it is incorrect to conclude from a single dataset that a smaller p-value is indicative of higher statistical power, simulation testing under controlled conditions has consistently shown higher statistical power for the wAFROC-AUC FOM3,4 as compared to the inferred ROC-AUC FOM.

Table 19.3: Inferred ROC-AUC analysis: results of random-reader random-case (RRRC) analysis, in worksheet "RRRC"". ddf = denominator degrees of freedom of F-distribution. df = degrees of freedom of t-distribution.

Stderr = standard error. CI = confidence interval; * = Significantly different at alpha = 0.05.].

8.4 Plotting wAFROC and ROC curves

It is important to display empirical wAFROC/ROC curves, not just for publication purposes, but to get a better feel for the data. Since treatments 4 and 5 showed the largest difference, the corresponding /ROC plots for them are displayed. The code is in file mainwAfrocRocPlots.R.

Sourcing this code yields Fig. 19.1. Plot (A), originating from lines 16 – 19, shows individual reader wAFROC plots for treatment 4 (solid lines) and treatment 5 (dashed lines). Running the software on one's computer best shows the color-coding. While difficult to see, examination of this plot shows that all readers performed better in treatment 4 than in treatment 5 (i.e., for each color the solid line is above the dashed line). Plot (B), originating from lines 21 – 25, shows reader-averaged wAFROC plots for treatments 4 (red line, upper curve) and 5 (blue line, lower curve). If one changes, for example, line 19 from print(plot1wAFROCPlot)toprint(plot1wAFROCPoints) the code will output the coordinates of the points describing the curve, which gives the user the option to copy and paste the operating points into alternative plotting software.

Lines 16-19 create plots for all specified treatment-reader combinations. The "trick" to creating reader-averaged curves, such as in (B) is defining two list variables, plotT and plotR, at lines 21-22, the first containing the treatments to be plotted, list(4,5), and the second, a list of equal length, containing the arrays of readers to be averaged over, list(c(1:4), c(1:4)). More examples can be found in the help page for PlotEmpiricaOperatingCharacteristics().

Meaningful operating points on the reader average curves cannot be defined. This is because ratings are treatment and reader specific labels, so one cannot for example, average bin counts over all readers to construct a table like ROC Table 4.1 or its AFROC counterpart, Table 13.3.

Instead, the following procedure is used internal to PlotEmpiricaOperatingCharacteristics(). The reader-averaged plot for a specified treatment is obtained by dividing the FPF range from 0 to 1 into finely spaced steps of 0.005. For each FPF value the wLLF values for that treatment are averaged over all readers, yielding the reader-averaged ordinate. Calculating confidence intervals on the reader-averaged curve is possible but cumbersome and unnecessary in my opinion. The relevant information, namely the 95% confidence interval on the difference in reader-averaged AUCs, is already contained in the program output, see Table 19.2, row labeled "4-5*". The difference is 0.05488 with a 95% confidence interval (0.03018, 0.07957).

Fig. 19.1: FED dataset; (A): individual reader wAFROC plots for treatments 4 and 5. While difficult to see, all readers performed better in treatment 4 as indicated by each colored solid line being above the corresponding dashed lines. (B): reader-averaged wAFROC plots for treatments 4 and 5. The performance superiority of treatment 4 is fairly obvious in this curve. The difference is significant, p = 0.00012.

Inferred ROC plots corresponding to Fig. 19.1 were generated by lines 20-24, i.e., by changing opChType = "wAFROC" to opChType = "ROC", and print(plot2wAFROCPlot)toprint(plot2ROCPlot), resulting in Fig. 19.2. From Table 19.3 it is seen that the difference in reader-averaged AUCs is 0.04219 with a 95% confidence interval (0.00727, 0.07711). The observed wAFROC effect-size, 0.05488, is larger than the corresponding inferred ROC effect-size, 0.04219. This is a common observation, but sampling variability compounded with small differences, could give different results.

Fig. 19.2: FED dataset; (A): individual reader ROC plots for treatments 4 and 5. While difficult to see, all readers performed better in treatment 4. (B): reader-averaged ROC plots for treatments 4 and 5. The performance superiority of treatment 4 is fairly obvious in this curve. The difference is significant, p = 0.03054.

8.5 Reporting an FROC study

The methods section should make it clear exactly how the study was conducted. The information should be enough to allow some one else to replicate the study. How many readers, how many cases, how many treatments were used. How was ground truth determined and if the FROC paradigm was used, how were true lesion locations determined? The instructions to the readers should be clearly stated in writing. Precautions to minimize reading order effects should be stated – usually this is accomplished by interleaving cases from different treatments so that the chances

that cases from a particular treatment is always seen first by every reader are minimized. Additionally, images from the same case, but in different treatments, should not be viewed in the same reading session. Reading sessions are usually an hour, and the different sessions should ideally be separated by at least one day. Users generally pay minimal attention to training sessions. It is recommended that at least 25% of the total number of interpretations be training cases and cases used for training should not be used in the main study. Feedback should be provided during training session to allow the reader to become familiar with the range of difficulty levels regarding diseased and non-diseased cases in the dataset. Deception, e.g., stating a higher prevalence than is actually used, is usually not a good idea. The user-interface should be explained carefully. The best user interface is intuitive, minimizes keystrokes and requires the least explanation.

In publications, the paradigm used to collect the data (ROC, FROC, etc.) and the figure of merit used for analysis should be stated. If FROC, the proximity criterion should be stated. The analysis should state the NH and the alpha of the test, and the desired generalization. The software used and appropriate references should be cited. The results of the overall F-test, the p-value, the observed F-statistic and its degrees of freedom should be stated. If the NH is not rejected, one should cite the observed inter-treatment FOM differences, confidence intervals and p-values and ideally provide preliminary sample size estimates. This information could be useful to other researchers attempting to conduct a larger study. If the NH is rejected, a table of inter-treatment FOM differences such as Table 19.3 should be summarized. Reader averaged plots of the relevant operating characteristics for each treatment should be provided. In FROC studies it is recommended to vary the proximity criterion, perhaps increasing it by a factor of 2, to test if the final conclusions (is NH rejected and if so which treatment is highest) are unaffected.

Assuming the study has been done properly and with sufficiently large number of cases, the results should be published in some form, even if the NH is not rejected. The dearth of datasets to allow reasonable sample size calculations is a real problem in this field. The dataset set should be made available, perhaps on Research Gate, or if communicated to me, they will be included in the Online Appendix material. Datasets acquired via NIH or other government funding must be made available upon request, in an easily decipherable format. Subsequent users of these datasets must cite the original source of the data. Given the high cost of publishing excess pages in some journals, an expanded version, if appropriate for clarity, should be made available using online posting avenues.

8.6 Crossed-treatment analysis

This analysis was developed for a particular application in which nodule detection in an anthropomorphic chest phantom in computed tomography (CT) was evaluated as a function of tube charge and reconstruction method. The phantom was scanned at 4 values of mAs and images were reconstructed with adaptive iterative dose reduction 3D (AIDR3D) and filtered back projection (FBP). Thus there are two treatment factors and the factors are crossed since for each value of the mAs factor there were two values of the reconstruction algorithm factor. Interest was in determining if whether performance depends on mAs and/or reconstruction method.

In a typical analysis of MRMC ROC or FROC study, treatment is considered as a single factor with I levels, where I is usually small. The figure of merit for treatment i (i =1,2,...,I) and reader j (j =1,2,...,J) is denoted; the case set index is suppressed. MRMC analysis compares the observed magnitude of the difference in reader-averaged figures of merit between treatments i and i', to the estimated standard deviation of the difference. For example, the reader-averaged difference in figures of merit is, where the dot symbol represents the average over the corresponding (reader) index. The standard deviation of the difference is estimated using the DBMH or the ORH method, using for example jackknifing to determine the variance components and/or covariances. With I levels, the number of distinct i vs. i' comparisons is I (I -1)/2. If the current study were analyzed in this manner, where I =8 (4 levels of mAs and two image reconstruction methods), then this would imply 28 comparisons. The large number of comparisons leads to loss of statistical power in detecting the effect of a specific pair of treatments, and, more importantly, does not inform one of the main points of interest: whether performance depends on mAs and/or reconstruction method. For example, in standard analysis the two reconstruction algorithms might be compared at different mAs levels, and one is in the dark as to which factor (algorithm or mAs) caused the observed significant difference.

Unlike conventional ROC type studies, the images in this study are defined by two factors. The first factor, tube charge, had four levels: 20, 40, 60 and 80 mAs. The second factor, reconstruction method, had two levels: FBP and AIDR3D. The figure of merit is represented by , where represents the levels of the first factor (mAs), and represents the levels of the second factor (reconstruction method), . Two sequential analyses were performed: (i) mAs analysis,

where the figure of merit was averaged over (the reconstruction index); and (ii) reconstruction analysis, where the figure of merit was averaged over (the mAs index). For example, the mAs analysis figure of merit is, where the dot represents the average over the reconstruction index, and the corresponding reconstruction analysis figure of merit is, where the dot represents the average over the mAs index. Thus in either analysis, the figure of merit is dependent on a single treatment factor, and therefore standard DBMH or ORH methods apply.

The mAs analysis determines whether tube charge is a significant factor and in this analysis the number of possible comparisons is only six. The reconstruction analysis determines whether AIDR3D offers any advantage over FBP and in this analysis the number of possible comparisons is only one. Multiple testing on the same dataset increases the probability of Type I error, therefore a Bonferroni correction is applied by setting the threshold for declaring significance at 0.025; this is expected to conservatively maintain the overall probability of a Type I error at = 0.05. Crossed-treatment analysis is used to describe this type of analysis of ROC/FROC data, which yields clearer answers on which of the two factors effects performance. The averaging over the other treatment has the effect of increasing the power of the study in detecting differences in each of the two factors.

Since the phantom is unique, and conclusions are only possible that are specific to this one phantom, the case (or image) factor was regarded as fixed. For this reason only results of random-reader fixed-case analyses are reported.

8.7 Discussion / Summary

An IDL (Interactive Data Language, currently marketed by Exelis Visual Information Solutions, www.exelisvis.com) version of JAFROC was first posted to a now obsolete website on 4/16/2004. This software required a license for IDL, which most users did not have. Subsequently, (9/27/2005) a version was posted which allowed analysis using the freely downloadable IDL Virtual Machine software (a method for freely distributing compiled IDL code). On 1/11/2011 the standalone Windows-compatible version was posted (4.0) and the current version is 4.2. JAFROC is windows compatible (XP, Vista and Windows 7, 8 and 10).

To our knowledge JAFROC is the only easily accessible software currently available that can analyze FROC data. Workstation software for acquiring ROC and FROC data is available from several sources 7-9. The Windows version is no longer actively supported (bugs, if pointed out, will be corrected). Current effort to conduct research and distribute software uses the R platform 10. There are several advantages to this. R is an open-source platform - we have already benefited from a bug pointed out by a user. R runs on practically any platform (Windows, OSX, Linux, etc.). Also, developing an R package benefits from other contributed R-packages, which allow easy computation of probability integrals, random number generation, and parallel computing to speed up computations, to name just a few. The drawback with R, and this has to with its open source philosophy, is that one cannot readily integrate existing ROC code, developed on other platforms and other programming languages (specifically, DLLs are not allowed in R). So useful programs like CORROC2 and CBM were coded in C++, since R allows C++ programs to be compiled and included in a package.

Due to the random number of marks per image, data entry in the FROC paradigm is inherently more complicated and error-prone than in ROC analysis, and consequently, and in response to feedback from users, much effort has gone into error checking. The users have especially liked the feature where the program indicates the Excel sheet name and line-number where an error is detected. User-feedback has also been very important in detecting program bugs and inconsistencies in the documentation and developing additional features (e.g., ROI analysis).

Interest in the FROC paradigm is evidenced by the fact that Ref. 3 describing the JAFROC method has been cited over 273 times. Over 25,000 unique visitors have viewed my website, at least 73 have downloaded the software and over 107 publications using JAFROC have appeared. The list is available on my website. JAFROC has been applied to magnetic resonance imaging, virtual computerized tomography colonoscopy, digital tomosynthesis (chest and breast), mammography dose and image processing optimization, computer aided detection (CAD), computerized tomography, and other applications.

Since confusion still appears to exist, especially among statisticians, regarding perceived neglect of intra-image correlations of ratings and how true negatives are handled in FROC analysis11, we close with a quote from respected sources 12 "(Chakraborty and Berbaum) have presented a solution to the FROC problem using a jackknife resampling approach that respects the correlation structure in the images ... their paradigm successfully passes a rigorous statistical validation test". Since 2005 the National Institutes for Health (NIH) has been generous with supporting the research and users of JAFROC have been equally generous with providing their datasets, which have resulted in several collaborations.

FROC sample size

Chapter 9

Sample size estimation for FROC studies

9.1 TBA How much finished

80%

9.2 Overview

This chapter is split into two parts.

- The first part goes into the details of FROC paradigm sample size estimation.
- The second part encapsulates most of the details in a new function SsFrocNhRsmModel(), which encapsulates some of the code in the first part (that relating to building the NH model), thereby making it easier for the user to perform FROC sample size estimation.

These parts are independently included in two RJafroc vignettes (on the GitHub master branch, not the CRAN uploaded version). These are located at:

https://dpc10ster.github.io/RJafroc/articles/Ch19Vig1FrocSampleSize.html

https://dpc10ster.github.io/RJafroc/articles/Ch19Vig2FrocSampleSize.html

9.3 Part 1

9.3.1 Introduction

FROC sample size estimation is not fundamentally different from the procedure outlined in TBA Chapter 11 for the ROC paradigm. To recapitulate, based on analysis of a pilot ROC dataset and using a specified FOM, e.g., FOM = Wilcoxon, and either method = "DBM" or method = "OR" for significance testing, one estimates the intrinsic variability of the data expressed in terms of variance components. For DBM analysis, these are the pseudovalue variance components, while for OR analysis these are the FOM treatment-reader variance component and the FOM covariances. The second step is to specify a clinically realistic effect-size, e.g., the anticipated AUC difference between the two modalities. Given these values, the sample size functions implemented in RJafroc (beginning with Ss) allow one to estimate the number of readers and cases necessary to detect (i.e., reject the null hypothesis) the modality AUC difference at specified Type II error rate β , typically chosen to be 20% - corresponding to 80% statistical power - and specified Type I error rate α , typically chosen to be 5%.

In FROC analysis the only difference, indeed the critical difference, is the choice of FOM; e.g., FOM = "wafroc" instead of the inferred ROC-AUC, FOM = "HrAuc". The FROC dataset is analyzed using either the DBM or

the OR method. This yields the necessary variance components or the covariance matrix corresponding to the wAFROC-AUC. The next step is to specify the effect-size **in wAFROC-AUC units**. The ROC-AUC has a historically well-known interpretation: the classification ability at separating diseased patients from non-diseased patients while the wAFROC-AUC does not. Needed is a way of relating the effect-size in ROC-AUC units to one in wAFROC-AUC units: as should be obvious this requires a physical model, e.g., the RSM, that predicts both ROC and wAFROC curves and the respective AUCs.

- 1. One chooses an ROC-AUC effect-size that is realistic, one that clinicians understand and can therefore participate in, in the effect-size postulation process. Lacking such information I recommend, based on past ROC studies, 0.03 as typical of a small effect size and 0.05 as typical of a moderate effect size.
- 2. One converts the ROC effect-size to a wAFROC-AUC effect-size. The method for this is described in the next section.
- 3. One uses the sample size tools in in RJafroc to determine sample size or power.

It is important to recognize is that all quantities have to be in the same units. When performing ROC analysis, everything (variance components and effect-size) has to be in units of the selected FOM, e.g., FOM = "Wilcoxon" which is identical to the empirical ROC-AUC. When doing wAFROC analysis, everything has to be in units of the wAFROC-AUC, i.e., FOM = "wAFROC". The variance components and effect-size in wAFROC-AUC units will be different from their corresponding ROC counterparts. In particular, as shown next, an ROC-AUC effect-size of 0.05 generally corresponds to a larger effect-size in wAFROC-AUC units. The reason for this is that the range over which wAFROC-AUC can vary, namely 0 to 1, is twice the corresponding ROC-AUC range.

The next section explains the steps used to implement #2 above.

9.3.2 Relating an ROC effect-size to a wAFROC effect-size

- If there are more than two treatments in the pilot dataset extract those treatments that represent null hypothesis data: for example DfExtractDataset(dataset, trts = c(1,2,4)) extracts treatments 1, 2 and 4. Save the extracted dataset to dataset. More than two treatments can be used if they have similar FOMs as the averaging/ian procedure described below benefits from more data. However, the final sample size predictions are restricted to two treatments.
- If the original data is FROC, convert it to ROC using DfFroc2Roc(dataset): this is because the RSM fits binned ROC data. The ROC dataset is rocDataset.
- If the data uses continuous scale ratings, bin the data using DfBinDataset(rocDataset, opChType = "ROC"). The default number of bins should be used. Unlike binning using arbitrarily set thresholds, the thresholds found by DfBinDataset are unique in that they maximize ROC-AUC. The binned dataset is rocDatasetB.
- For each treatment and reader the inferred ROC data is fitted by FitRsmRoc(), see example below, yielding estimates of the RSM physical (or pri) parameters (not the intrinsic values).
- The following example uses the *first two* treatments of the "FED" dataset, dataset04, which is a 5 treatment 4 radiologist FROC dataset acquired by Dr. Federica Zanca et. al. (Zanca et al., 2009). The dataset has 5 treatments and 4 readers and 200 cases and was acquired on a 5-point integer scale, i.e., it is already binned. If not one needs to bin the dataset using DfBinDataset(). I need to emphasize this point: if the dataset represents continuous ratings, as with a CAD algorithm, one must bin the dataset.
- The reason for using RSM parameter values only for the first two treatments is that these were found (Zanca et al., 2009) to be almost equivalent (more precisely, the NH could not be rejected for the first two treatments, so it makes sense to regard them as "almost" NH treatments.
- The following code block defines the pilot FROC data frocData (corresponding to dataset04, which is the "FED" dataset, but with only treatments 1 and 2 extracted, using DfExtractDataset()) and rocData, i.e., the highest-rating ROC dataset inferred from the FROC dataset using DfFroc2Roc().

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```
frocData <- DfExtractDataset(dataset04, trts = c(1,2))
rocData <- DfFroc2Roc(frocData)
rocDataB <- DfBinDataset(rocData, opChType = "ROC") # unnecessary as data is already binned
# but cant hurt</pre>
```

The next code block determines lesDistr, the lesion distribution array, which has Lmax (maximum number of lesions per diseased case over the dataset) rows and two columns. The first column contains the integers 1, 2, ..., Lmax and the second column contains the fraction of diseased cases with the number of lesions per case specified in the first column. The second column will sum to unity. The RSM fitting algorithm needs to know how lesion-rich the dataset is, as the RSM predicted ROC-AUC depends on the lesion-richness of the dataset. For reasons that will become clear below, one also needs the distribution of the lesion weights. Note that lesDist is determined from the FROC dataset as the lesion-richness information is lost upon conversion to an ROC dataset. The PlotRsmOperatingCharacteristics function used below sets relWeights = 0, which ensures equally weighted lesions: on cases with one lesion the weight of the lesion is unity, on cases with two lesions the weights of each lesion is 1/2 and on cases with three lesions the weight of each lesion is 1/3, etc.. TBA due to changes in lesDist etc.

```
lesDistr <- UtilLesionDistrVector(frocData)
print(lesDistr)</pre>
```

```
## [1] 0.69 0.20 0.11
```

For this dataset Lmax is 3, and fraction 0.69 of diseased cases have one lesion, fraction 0.2 of diseased cases have two lesions and fraction 0.11 of diseased cases have three lesions.

The next code block determines the number of treatments and readers (I and J) from the dimensions of the frocData\$ratings\$NL array. It creates an array RsmParms to hold the RSM fitted parameter values. For each treatment and reader it applies the fitting algorithm FitRsmRoc(). The first three returned values are mu, lambda and nu, corresponding to RSM parameters μ , λ' and ν' .

```
I <- dim(frocData$ratings$NL)[1]

J <- dim(frocData$ratings$NL)[2]

RsmParms <- array(dim = c(I,J,3))

for (i in 1:I) {
    for (j in 1:J) {
        x1 <- FitRsmRoc(rocDataB, trt = i, rdr = j, lesDistr)
        RsmParms[i,j,1] <- x1[[1]] # mu
        RsmParms[i,j,2] <- x1[[2]] # lambda
        RsmParms[i,j,3] <- x1[[3]] # nu
    }
}</pre>
```

I recommend taking the ian of each of the parameters, over all treatment-reader indices, as representing the average NH dataset. The ian is less sensitive to outliers than the mean.

```
mu <- median(RsmParms[,,1])
lambda <- median(RsmParms[,,2])
nu <- median(RsmParms[,,3])</pre>
```

The defining values of the fitting model are mu = 3.3121519, lambda = 1.714368 and nu = 0.7036564. Note that these obey the constraints lambda > 0 and 0 < nu < 1. We are now ready to calculate the expected NH FOMs using the ROC -AUC and the wAFROC FOM.

- The plotting function PlotRsmOperatingCharacteristics() returns a number of other objects, most importantly the plot, but here we use only the AUC, which is obtained by numerical integration of the predicted operating characteristics.
- One has aucRocNH = 0.8791542 and aucwAfrocNH = 0.7199233. Note that the wAFROC-FOM is smaller than the ROC-FOM as it includes the localization constraint.
- To induce the alternative hypothesis condition, one increments μ_{NH} by Δ_{μ} . The resulting ROC-AUC and wAFROC-AUC are calculated, again by numerical integration of the RSM predicted ROC and wAFROC curves, leading to the corresponding effect-sizes (note that in each equation below one takes the difference between the AH value minus the NH value):
- The next step is to calculate the effect size (new value minus the NH value) using ROC and wAFROC FOMs for a series of specified deltaMu values. This generates values that can be used to interpolate a wAFROC effect size for a specified ROC effect size.

```
deltaMu <- seq(0.01, 0.2, 0.01) # values of deltaMu to scan below
esRoc <- array(dim = length(deltaMu));eswAfroc <- array(dim = length(deltaMu))
for (i in 1:length(deltaMu)) {
  esRoc[i] <- PlotRsmOperatingCharacteristics(</pre>
    mu + deltaMu[i], lambda, nu, lesDistr = lesDistr, OpChType = "ROC")$aucROC - aucRocNH
  eswAfroc[i] <- PlotRsmOperatingCharacteristics(</pre>
   mu+ deltaMu[i], lambda, nu, lesDistr = lesDistr, OpChType = "wAFROC") aucwAFROC - aucwAfrocNH
  cat("ES_ROC = ", esRoc[i], ", ES_wAFROC = ", eswAfroc[i],"\n")
## ES_ROC = 0.0001500649 , ES_wAFROC = 0.000189712
## ES ROC = 0.0002978454 , ES wAFROC = 0.0003764812
## ES_ROC = 0.0004433681 , ES_wAFROC = 0.0005603432
## ES_ROC = 0.0005866597 , ES_wAFROC = 0.0007413331
## ES_ROC = 0.0007277463 , ES_wAFROC = 0.0009194859
## ES ROC = 0.0008666543 , ES wAFROC = 0.001094837
## ES_ROC = 0.001003409 , ES_wAFROC = 0.00126742
## ES_ROC = 0.001138038 , ES_wAFROC = 0.001437269
## ES_ROC = 0.001270565 , ES_wAFROC = 0.00160442
```

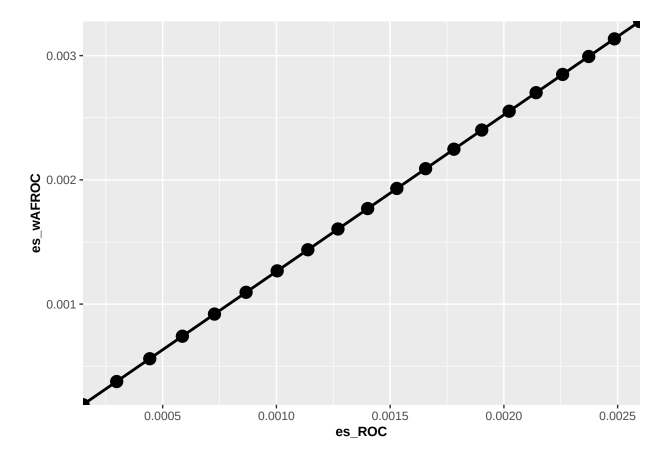
ES_ROC = 0.002485801 , ES_WAFROC = 0.00313478 ## ES_ROC = 0.002596862 , ES_WAFROC = 0.003274416

ES_ROC = 0.001401017 , ES_wAFROC = 0.001768904
ES_ROC = 0.001529418 , ES_wAFROC = 0.001930757
ES_ROC = 0.001655794 , ES_wAFROC = 0.002090012
ES_ROC = 0.00178017 , ES_wAFROC = 0.002246701
ES_ROC = 0.00190257 , ES_wAFROC = 0.002400857
ES_ROC = 0.002023021 , ES_wAFROC = 0.002552513
ES_ROC = 0.002141544 , ES_wAFROC = 0.002701702
ES_ROC = 0.002258166 , ES_wAFROC = 0.002848455
ES_ROC = 0.002372911 , ES_wAFROC = 0.002992804

Here is a plot of wAFROC effect size (y-axis) vs. ROC effect size.

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```
scale_x_continuous(expand = c(0, 0)) +
scale_y_continuous(expand = c(0, 0))
print(p)
```



The plot is very close to linear. This makes it easy to design an interpolation function. In the following code block the first line fits eswAfroc vs. esRoc using the linear model lm() function constrained to pass through the origin (the minus one): scaleFactor <- lm(eswAfroc ~ -1 + esRoc). One expects this constraint since for deltaMu = 0 the effect size must be zero no matter how it is measured.

```
scaleFactor<-lm(eswAfroc~-1+esRoc) # fit values to straight line thru origin
effectSizeROC <- seq(0.01, 0.1, 0.01)
effectSizewAFROC <- effectSizeROC*scaleFactor$coefficients[1] # r2 = summary(scaleFactor)$r.squared
```

The slope of the zero-intercept constrained straight line fit is scaleFactor = 1.2617239 and the squared correlation coefficient is R2 = 0.9999997. Therefore, the conversion from ROC to wAFROC effect size is: effectSizewAFROC = scaleFactor * effectSizeROC. The wAFROC effect size is twice the ROC effect size. All that remains is to calculate the variance components using the two FOMs.

9.3.3 Computing the respective variance components

The code block applies StSignificanceTesting() to rocData and frocData, using the appropriate FOM, and extracts the variance components.

```
temp1 <- StSignificanceTesting(rocData, FOM = "Wilcoxon", method = "DBM", analysisOption = "RRRC")
temp2 <- StSignificanceTesting(frocData, FOM = "wAFROC", method = "DBM", analysisOption = "RRRC")
varCompROC <- temp1$ANOVA$VarCom
varCompwAFROC <- temp2$ANOVA$VarCom</pre>
```

The observed wAFROC effect-size is -0.00685625. This is a very small effect size; the corresponding ROC effect-size is -0.0051; the sign does not affect the calculations, which is too small to reach 80% power. It is not surprising that the study (Zanca et al., 2009) did not find a significant difference between these two treatments

The respective variance components are:

```
print(varCompROC)
```

```
## VarR 0.00082773798
## VarC 0.03812334734
## VarTR 0.00015265067
## VarTC 0.00964432675
## VarRC 0.00354419640
## VarErr 0.09484636574
```

print(varCompwAFROC)

```
## Estimates
## VarR 0.00185422886
## VarC 0.06117804981
## VarTR -0.00044392794
## VarTC 0.01016518621
## VarRC 0.01355883396
## VarErr 0.09672559908
```

Only terms involving treatment are relevant to sample size. The wAFROC varTC and varErr values are slightly larger than the ROC ones - as expected - because the range of the wAFROC FOM is twice that of the ROC FOM.

9.3.4 Comparing ROC power to wAFROC power for equivalent effect-sizes

We are now ready to compare ROC and wAFROC powers for equivalent effect sizes. The following example is for 5 readers (JPivot) and 100 cases (KPivot) in the **pivotal study**.

```
powerROC <- array(dim = length(effectSizeROC));powerwAFROC <- array(dim = length(effectSizeROC))</pre>
JPivot <- 5; KPivot <- 100
for (i in 1:length(effectSizeROC)) {
  varYTR <- varCompROC["VarTR", "Estimates"] # these are pseudovalue based variance components assuming FOM
  varYTC <- varCompROC["VarTC","Estimates"]</pre>
  varYEps <- varCompROC["VarErr","Estimates"]</pre>
  ret <- SsPowerGivenJK(dataset = NULL, FOM = "Wilcoxon", J = JPivot, K = KPivot, analysisOption = "RRRC",
        list(VarTR = varYTR, VarTC = varYTC, VarErr = varYEps))
  powerROC[i] <- ret$powerRRRC</pre>
  varYTR <- varCompwAFROC["VarTR", "Estimates"] # these are pseudovalue based variance components assuming
  varYTC <- varCompwAFROC["VarTC", "Estimates"]</pre>
  varYEps <- varCompwAFROC["VarErr","Estimates"]</pre>
  ret <- SsPowerGivenJK(dataset = NULL, FOM = "Wilcoxon", J = JPivot, K = KPivot, analysisOption = "RRRC",
        list(VarTR = varYTR, VarTC = varYTC, VarErr = varYEps))
  powerwAFROC[i] <- ret$powerRRRC</pre>
  cat("ROC-ES = ", effectSizeROC[i], ", wAFROC-ES = ", effectSizewAFROC[i],
      ", Power-ROC = ", powerROC[i], ", Power-wAFROC = ", powerwAFROC[i], "\n")
```

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```
## ROC-ES = 0.01 , wAFROC-ES = 0.012617239 , Power-ROC = 0.064430457 , Power-wAFROC = 0.075439644

## ROC-ES = 0.02 , wAFROC-ES = 0.025234479 , Power-ROC = 0.10878897 , Power-wAFROC = 0.15449773

## ROC-ES = 0.03 , wAFROC-ES = 0.037851718 , Power-ROC = 0.18471152 , Power-wAFROC = 0.28797922

## ROC-ES = 0.04 , wAFROC-ES = 0.050468957 , Power-ROC = 0.29079274 , Power-wAFROC = 0.4612966

## ROC-ES = 0.05 , wAFROC-ES = 0.063086196 , Power-ROC = 0.41954431 , Power-wAFROC = 0.6420946

## ROC-ES = 0.06 , wAFROC-ES = 0.075703436 , Power-ROC = 0.55738123 , Power-wAFROC = 0.79495349

## ROC-ES = 0.07 , wAFROC-ES = 0.088320675 , Power-ROC = 0.68816012 , Power-wAFROC = 0.90003872

## ROC-ES = 0.08 , wAFROC-ES = 0.10093791 , Power-ROC = 0.79836108 , Power-wAFROC = 0.95891383

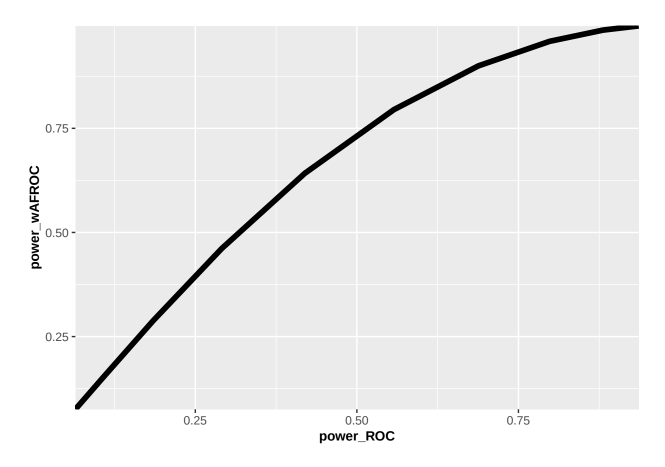
## ROC-ES = 0.09 , wAFROC-ES = 0.11355515 , Power-ROC = 0.88095077 , Power-wAFROC = 0.98585038

## ROC-ES = 0.1 , wAFROC-ES = 0.12617239 , Power-ROC = 0.93606799 , Power-wAFROC = 0.9959336
```

Since the wAFROC effect size is about a factor of two larger than the ROC effect size, wAFROC power is larger than that for ROC. The effect is magnified as the effect size enters as the square in the formula for the power (this overwhelms the slight increase in variability of wAFROC-FOM relative to ROC-FOM noted previously). The following is a plot of the respective powers.

```
## Warning: Using `size` aesthetic for lines was deprecated in ggplot2 3.4.0.
## i Please use `linewidth` instead.
```

```
print(p)
```



9.4 Part 2

9.4.1 Introduction

This example uses the FED dataset as a pilot FROC study and function SsFrocNhRsmModel() to construct the NH model (encapsulating some of the code in the first part).

9.4.2 Constructing the NH model for the dataset

One starts by extracting the first two treatments from dataset04, which represent the NH dataset, see previous part. Next one constructs the NH model - note that the lesion distribution lesDistr can be specified here independently of that in the pilot dataset. This allows some control over selection of the diseased cases in the pivotal study.

```
lesDistr <- c(0.7, 0.2, 0.1)
frocNhData <- DfExtractDataset(dataset04, trts = c(1,2))
ret <- SsFrocNhRsmModel(frocNhData, lesDistr = lesDistr)
mu <- ret$mu
lambda <- ret$lambda
nu <- ret$nu
scaleFactor <- ret$scaleFactor</pre>
```

The fitting model is defined by mu = 3.31491361, lambda = 1.6930673 and nu = 0.70649936 and lesDistr. The effect size scale factor is 1.25445833.

9.4. PART 2 75

The null hypothesis ROC AUC is 0.8790548 and the corresponding NH wAFROC AUC is 0.72320816.

9.4.3 Extracting the wAFROC variance components

The next code block applies StSignificanceTesting() to frocNhData, using FOM = "wAFROC" and extracts the variance components.

```
varCompwAFROC <- StSignificanceTesting(frocNhData, FOM = "wAFROC", method = "DBM", analysisOption = "RRRC</pre>
```

9.4.4 wAFROC power for specified ROC effect size, number of readers J and number of cases K

The following example is for ROC effect size = 0.05, 5 readers (J = 5) and 100 cases (K = 100) in the **pivotal** study.

9.4.5 wAFROC number of cases for 80% power for a given number of readers J

```
## ROC-ES = 0.05 , wAFROC-ES = 0.062722916 , K80RRRC = 123 , Power-wAFROC = 0.80210887
```

9.4.6 wAFROC-FOM power for a given number of readers J and cases K

The estimated power is close to 80% as the number of cases (ret2\$KRRRC = 123) was chosen deliberately from the previous code block.

RJafroc Vignettes

Chapter 10

F-distribution

10.1 TBA How much finished

10%

10.2 Background

A number of vignettes used to be part of the RJafroc package. These have been moved here.

10.3 Introduction

Since it plays an important role in sample size estimation, it is helpful to examine the behavior of the F-distribution. In the following ndf = numerator degrees of freedom, ddf = denominator degrees of freedom and ncp = non-centrality parameter (i.e., the Δ appearing in Eqn. (11.6) of (Chakraborty, 2017)).

The use of three R functions is demonstrated.

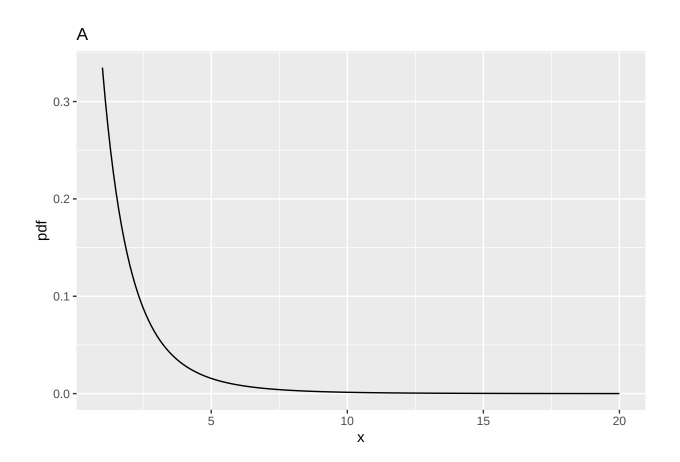
- qf(p,ndf,ddf) is the *quantile* function of the F-distribution for specified values of p, ndf and ddf, i.e., the value x such that fraction p of the area under the F-distribution lies to the right of x. Since ncp is not included as a parameter, the default value, i.e., zero, is used. This is called the *central* F-distribution.
- df(x,ndf,ddf,ncp) is the probability density function (pdf) of the F-distribution, as a function of x, for specified values of ndf, ddf and ncp.
- pf(x,ndf,ddf,ncp) is the probability (or cumulative) distribution function of the F-distribution for specified values of ndf, ddf and ncp.

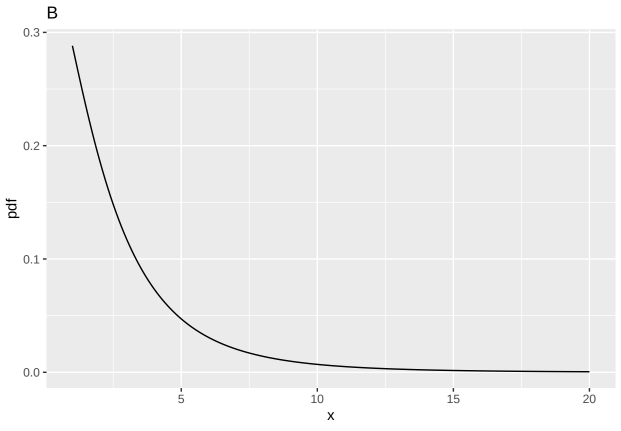
10.4 Effect of ncp for ndf = 2 and ddf = 10

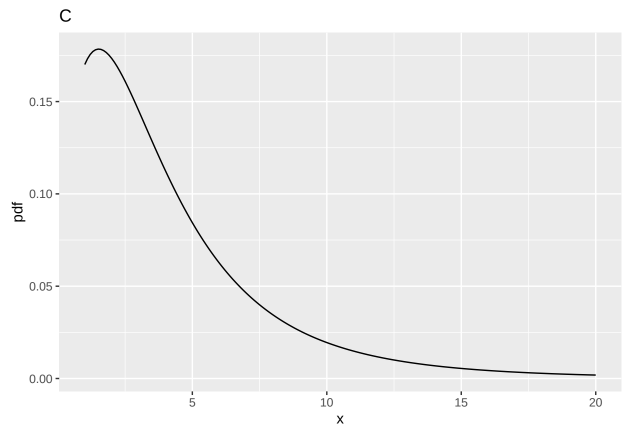
- Four values of ncp are considered (0, 2, 5, 10) for ddf = 10.
- fCrit is the critical value of the F distribution, i.e., that value such that fraction α of the area is to the right of the critical value, i.e., fCrit is identical in statistical notation to $F_{1-\alpha,ndf,ddf}$.

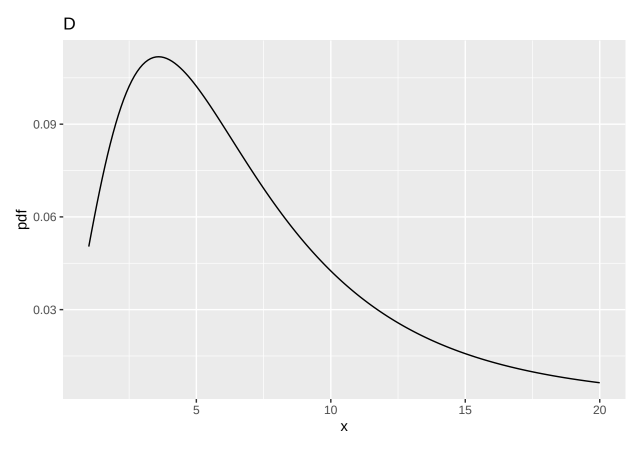
```
ndf <- 2;ddf <- 10;ncp <- c(0,2,5,10)
alpha <- 0.05
fCrit <- qf(1-alpha, ndf,ddf)</pre>
```

```
x <- seq(1, 20, 0.1)
myLabel <- c("A", "B", "C", "D")
myLabelIndx <- 1
pFgtFCrit <- NULL
for (i in 1:length(ncp))
{
    y <- df(x,ndf,ddf,ncp=ncp[i])
    pFgtFCrit <- c(pFgtFCrit, 1-pf(fCrit, ndf, ddf, ncp = ncp[i]))
}
for (i in 1:length(ncp))
{
    y <- df(x,ndf,ddf,ncp=ncp[i])
    curveData <- data.frame(x = x, pdf = y)
    curvePlot <- ggplot(data = curveData, mapping = aes(x = x, y = pdf)) +
    geom_line() +
    ggtitle(myLabel[myLabelIndx]);myLabelIndx <- myLabelIndx + 1
    print(curvePlot)
}
fCrit_2_10 <- fCrit # convention fCrit_ndf_ddf</pre>
```









	ndf	ddf	fCrit	ncp	pFgtFCrit
A	2	10	4.102821	0	0.0500000
В	2	10	4.102821	2	0.1775840
С	2	10	4.102821	5	0.3876841
D	2	10	4.102821	10	0.6769776

10.5 Comments

10.5.1 Fig. A

- This corresponds to ncp = 0, i.e., the *central* F-distribution.
- The integral under this distribution is unity (this is also true for all plots in this vignette).
- The critical value, fCrit in the above code block, is the value of x such that the probability of exceeding x is α. The corresponding parameter alpha is defined above as 0.05.
- In the current example fCrit = 4.102821. Notice the use of the quantile function qf() to determine this value, and the default value of ncp, namely zero, is used; specifically, one does not pass a 4th argument to qf().
- The decision rule for rejecting the NH uses the NH distribution of the F-statistic, i.e., reject the NH if F >= fCrit. As expected, prob > fCrit = 0.05 because this is how fCrit was defined.

10.5.2 Fig. B

- This corresponds to ncp = 2, ndf = 2 and ddf = 10.
- The distribution is slightly shifted to the right as compared to Fig. A, thereby making it more likely that the observed value of the F-statistic will exceed the critical value determined for the NH distribution.
- In fact, prob > fCrit = 0.177584, i.e., the *statistical power* (compare this to Fig. A where prob > fCrit was 0.05).

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10.5.3 Fig. C

- This corresponds to ncp = 5, ndf = 2 and ddf = 10.
- Now prob > fCrit = 0.3876841.
- Power has increased compared to Fig. B.

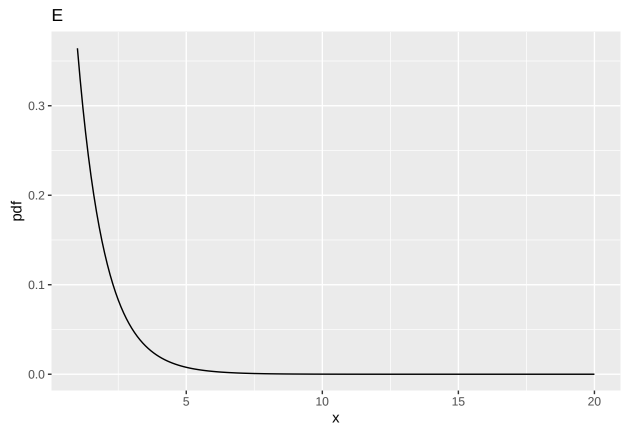
10.5.4 Fig. D

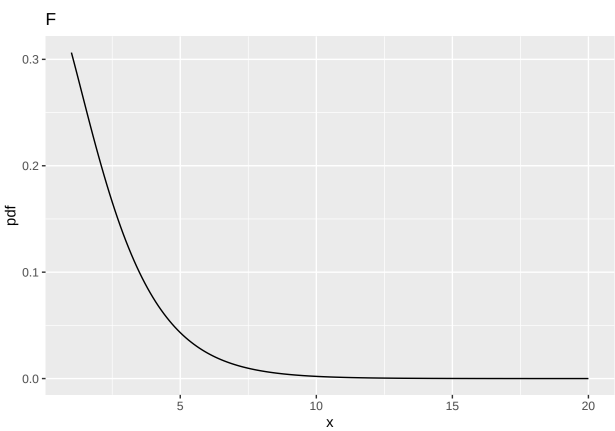
- This corresponds to ncp = 10, ndf = 2 and ddf = 10.
- Now prob > fCrit is 0.6769776.
- Power has increased compared to Fig. C.
- The effect of the shift is most obvious in Fig. C and Fig. D.
- Considering a vertical line at x=4.102821, fraction 0.6769776 of the probability distribution in Fig. D lies to the right of this line
- Therefore the NH is likely to be rejected with probability 0.6769776.

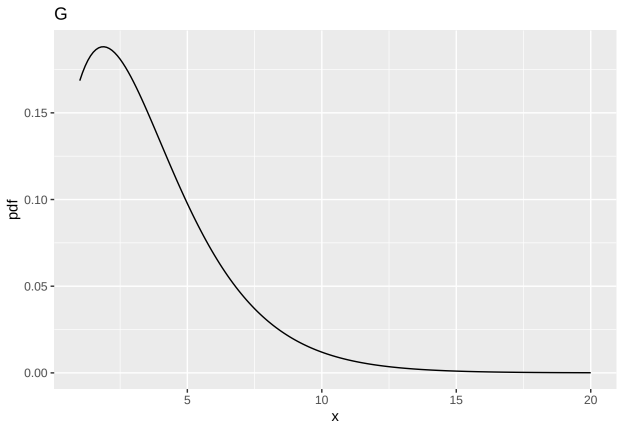
10.5.5 Summary

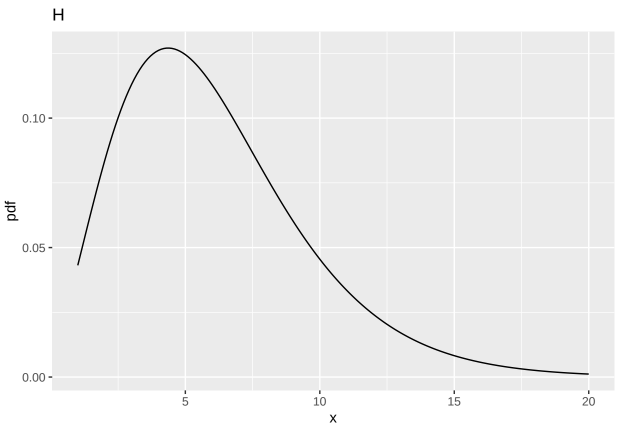
The larger that non-centrality parameter, the greater the shift to the right of the F-distribution, and the greater the statistical power.

10.6 Effect of ncp for ndf = 2 and ddf = 100









	ndf	ddf	fCrit	ncp	pFgtFCrit
A	2	10	4.102821	0	0.0500000
В	2	10	4.102821	2	0.1775840
С	2	10	4.102821	5	0.3876841
D	2	10	4.102821	10	0.6769776
\overline{E}	2	100	3.087296	0	0.0500000
F	2	100	3.087296	2	0.2199264
G	2	100	3.087296	5	0.4910802
Η	2	100	3.087296	10	0.8029764

10.7 Comments

- All comparisons in this sections are at the same values of ncp defined above.
- And between ddf = 100 and ddf = 10.

10.7.1 Fig. E

- This corresponds to ncp = 0, ndf = 2 and ddf = 100.
- The critical value is fCrit_2_100 = 3.0872959. Notice the decrease compared to the previous value for ncp = 0, i.e., 4.102821, for ddf = 10.
- One expects that increasing ddf will make it more likely that the NH will be rejected, and this is confirmed below.
- All else equal, statistical power increases with increasing ddf.

10.7.2 Fig. F

- This corresponds to ncp = 2, ndf = 2 and ddf = 100.
- The probability of exceeding the critical value is prob > fCrit_2_100 = 0.2199264, greater than the previous value, i.e., 0.177584 for ddf = 10.

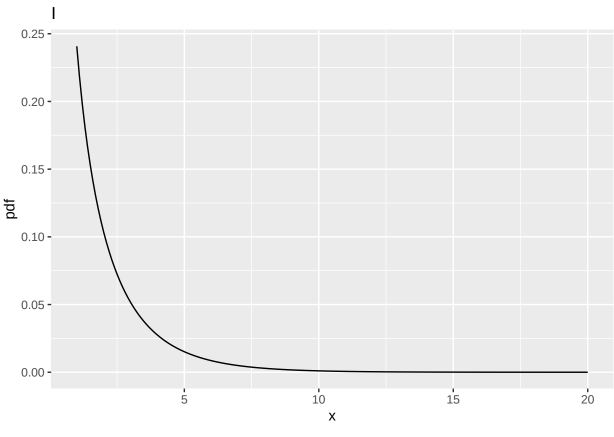
10.7.3 Fig. G

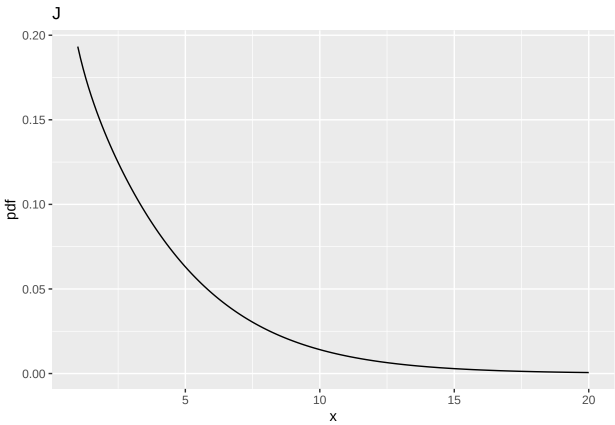
- This corresponds to ncp = 5, ndf = 2 and ddf = 100.
- The probability of exceeding the critical value is prob > fCrit_2_100 = 0.4910802.
- This is greater than the previous value, i.e., 0.3876841 for ddf = 10.

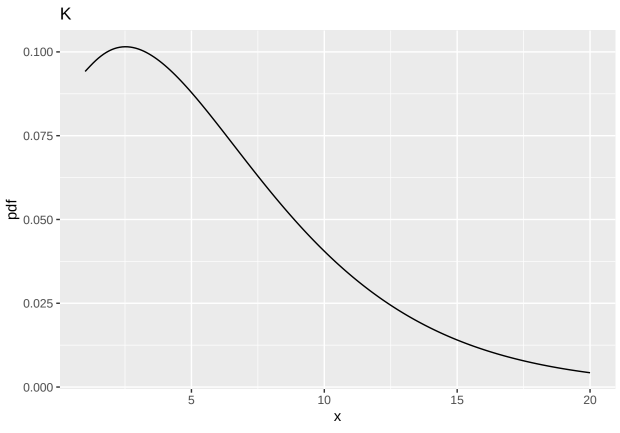
10.7.4 Fig. H

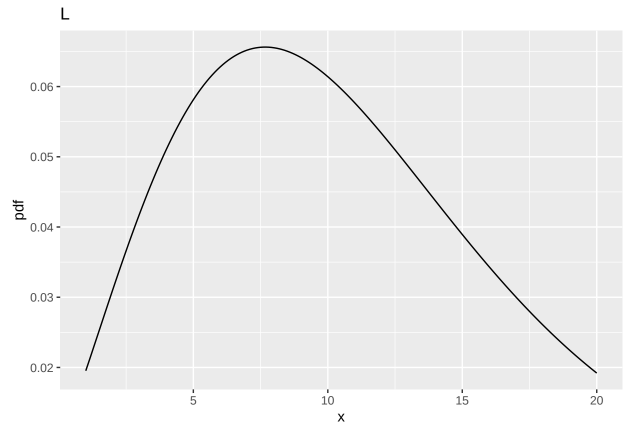
- This corresponds to ncp = 10, ndf = 2 and ddf = 100.
- The probability of exceeding the critical value is prob > fCrit_2_100 is 0.8029764.
- This is greater than the previous value, i.e., 0.6769776 for ddf = 10.

10.8 Effect of ncp for ndf =1, ddf =100









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	ndf	ddf	fCrit	ncp	pFgtFCrit
A	2	10	4.102821	0	0.0500000
В	2	10	4.102821	2	0.1775840
С	2	10	4.102821	5	0.3876841
D	2	10	4.102821	10	0.6769776
E	2	100	3.087296	0	0.0500000
F	2	100	3.087296	2	0.2199264
G	2	100	3.087296	5	0.4910802
Η	2	100	3.087296	10	0.8029764
I	1	100	3.936143	0	0.0500000
J	1	100	3.936143	2	0.2883607
K	1	100	3.936143	5	0.6004962
L	1	100	3.936143	10	0.8793619

10.9 Comments

- All comparisons in this sections are at the same values of ncp defined above and at ddf = 100.
- And between ndf = 1 and ndf = 2.

10.9.1 Fig. I

- This corresponds to ncp = 0, ndf = 1 and ddf = 100.
- The critical value is $fCrit_1_100 = 3.936143$.
- Notice the increase in the critical value as compared to the corresponding value for ndf = 2, i.e., 3.0872959.
- One expects power to decrease: the following code demonstrates that as ndf increases, the critical value fCrit
 decreases.
- In significance testing generally ndf = I 1.
- It more likely that the NH will be rejected with increasing numbers of treatments.

ndf	ddf	fCrit
1	100	3.936143
2	100	3.087296
5	100	2.305318
10	100	1.926692
12	100	1.850255
15	100	1.767530
20	100	1.676434

10.9.2 Fig. J

- This corresponds to ncp = 2, ndf = 1 and ddf = 100.
- Now prob > fCrit_1_100 = 0.2883607, 0.1351602, 0.0168844, 8.9992114×10^{-4} , 3.2584757×10^{-4} , 8.1619807×10^{-5} , 1.1084132×10^{-5} , larger than the previous value 0.2199264.
- The power has actually increased.

10.9.3 Fig. K

- This corresponds to ncp = 5, ndf = 1 and ddf = 100°,
- Now prob > fCrit_1_100 = 0.6004962, 0.3632847, 0.0699798, 0.0048836, 0.0018367, 4.6889533×10^{-4} , 6.2058692×10^{-5} , larger than the previous value 0.4910802.
- Again, the power has actually increased.

10.9.4 Fig. L

- This corresponds to ncp = 10, ndf = 1 and ddf = 100
- Now prob > fCrit_1_100 is 0.8793619, 0.7000168, 0.2459501, 0.0290856, 0.0123033, 0.0035298, 5.1213398×10^{-4} , larger than the previous value 0.8029764.
- The power has actually increased.

10.10 Summary

- Power increases with increasing ddf and ncp.
- The effect of increasing ncp is quite dramatic. This is because power depends on the square of ncp.
- As ndf increases, fCrit decreases, which makes it more likely that the NH will be rejected.
- With increasing numbers of treatments the probability is greater that the F-statistic will be large enough to exceed the critical value.

Chapter 11

RSM operating characteristics

11.1 TBA How much finished

10%

11.2 Introduction

- The purpose of this vignette is to explain the operating characteristics predicted by the RSM. It relates to Chapter 17 in my book (Chakraborty, 2017).
- This vignette is under development ...
- Also to explain the difference between dataset members (lesionID, lesionWeight) and RSM parameters (lesDistr, lesWghtDistr).

11.3 The distinction between predicted curves and empirical curves

- Operating characteristics predicted by a model have zero sampling variability.
- Empirical operating characteristics, which apply to datasets, have non-zero sampling variability.
- If the model is correct, as the numbers of cases in the dataset increases, the empirical operating characteristic asymptotically approaches the predicted curve.

11.4 The RSM model

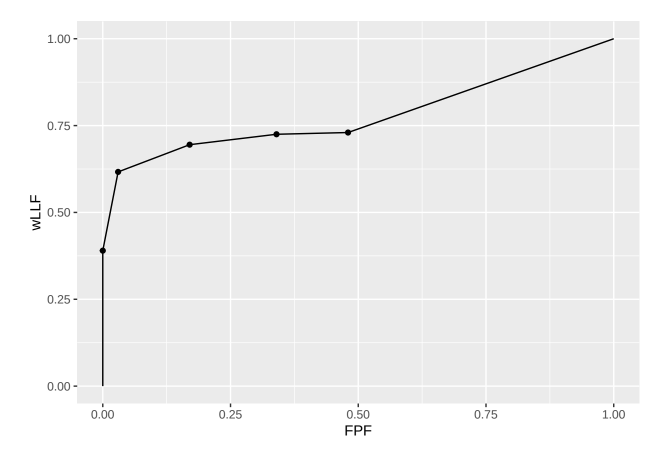
- The 3 RSM parameters and two additional parameters characterizing the dataset determine the wAFROC curve.
- The 3 RSM parameters are μ , λ and ν .
- The two dataset parameters are:
 - The distribution of number of lesions per diseased case, lesDistr.
 - The distribution of lesion weights, lesWghtDistr.
- These parameters do not apply to individual cases; rather they refer to a large population (asymptotically infinite in size) of cases.

```
str(dataset04$lesions$IDs)
#> num [1:100, 1:3] 1 1 1 1 1 1 1 1 1 1 1 ...
str(dataset04$lesions$weights)
#> num [1:100, 1:3] 1 1 1 1 1 1 1 1 1 1 ...
```

- Note that the first index of both arrays is the case index for the 100 abnormal cases in this dataset.
- With finite number of cases the empirical operating characteristic (or for that matter any fitted operating characteristic) will have sampling variability as in the following example.

11.5 The empirical wAFROC

```
p <- PlotEmpiricalOperatingCharacteristics(dataset04, opChType = "wAFROC")
p$Plot</pre>
```

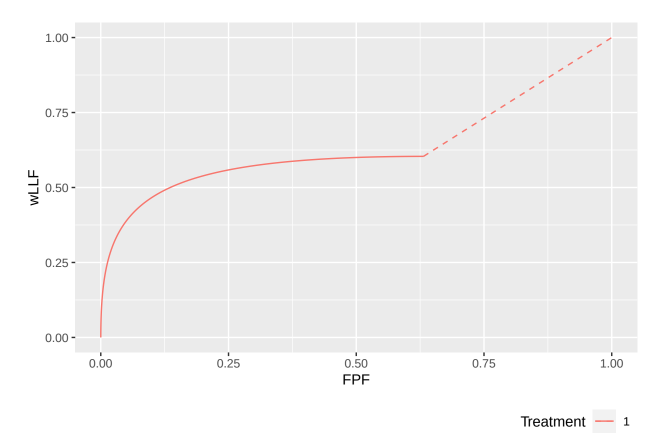


- The piecewise linear nature of the plot, with sharp breaks, indicates that this is due to a finite dataset.
- In contrast the following code shows a smooth plot, because it is a model *predicted* plot.

11.6 The predicted wAFROC

```
## Following example is for mu = 2, lambda = 1, nu = 0.6. 20% of the diseased ## cases have a single lesion, 40% have two lesions, 10% have 3 lesions, ## and 30% have 4 lesions. lesDistr <- c(0.2, 0.4, 0.1, 0.3)
## On cases with one lesion the weights are 1, on cases with 2 lesions the weights ## are 0.4 and 0.6, on cases with three lesions the weights are 0.2, 0.3 and 0.5, and ## on cases with 4 lesions the weights are 0.3, 0.4, 0.2 and 0.1:
```

```
relWeights <- c(0.3, 0.4, 0.2, 0.1)
p <- PlotRsmOperatingCharacteristics(
    mu = 2,
    lambda = 1,
    nu = 0.6,
    OpChType = "wAFROC",
    lesDistr = lesDistr,
    relWeights = relWeights,
    legendPosition = "bottom", nlfRange = c(0, 1), llfRange = c(0, 1))
p$wAFROCPlot</pre>
```



11.7 The distribution of number of lesions and weights

```
lesDistr
#> [1] 0.2 0.4 0.1 0.3
relWeights
#> [1] 0.3 0.4 0.2 0.1
```

- The lesDistr array 0.2, 0.4, 0.1, 0.3 specifies the fraction of diseased cases with the number of lesions corresponding to the column index. To specify a dataset with exactly 3 lesions per diseased case use lesDist = c(0, 0, 1, 0).
- The relWeights array 0.3, 0.4, 0.2, 0.1 specifies the relative weights.
- For cases with 1 lesion, the weight is 1.
- For cases with 2 lesions, the first lesion has weight 0.4285714 and the second lesion has weight 0.5714286, which are in the ratio 0.3: 0.4 and sum to unity.

- For cases with 3 lesions, the first lesion has weight 0.3333333, the second lesion has weight 0.4444444 and the third lesion has weight 0.2222222, which are in the ratio 0.3: 0.4: 0.2, and sum to unity.
- For cases with 4 lesions, the weights are 0.3, 0.4, 0.2 and 0.1, which are in the ratio 0.3: 0.4: 0.2: 0.1 and sum to unity.

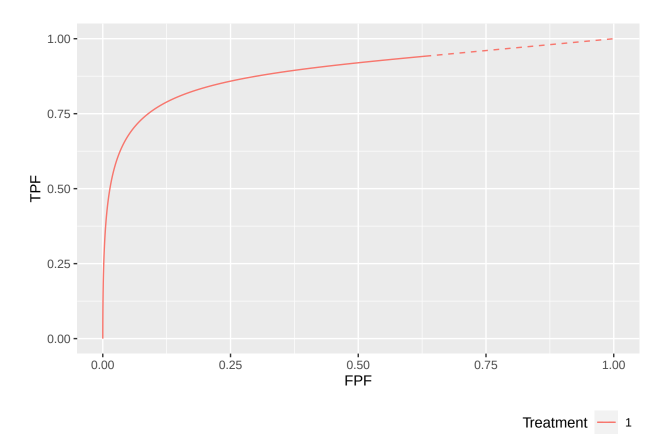
11.8 Other operating characteristics

- By changing OpChType one can generate other operating characteristics.
- Note that lesiion weights argument is not needed for ROC curves. It is only needed for wAFROC and wAFROC1 curves.

```
lesDistr <- c(0.2, 0.4, 0.1, 0.3)

p <- PlotRsmOperatingCharacteristics(
   mu = 2,
   lambda = 1,
   nu = 0.6,
   OpChType = "ROC",
   lesDistr = lesDistr,
   legendPosition = "bottom")

p$ROCPlot</pre>
```



11.9 Summary

Chapter 12

Improper ROC curves

12.1 TBA How much finished

10%

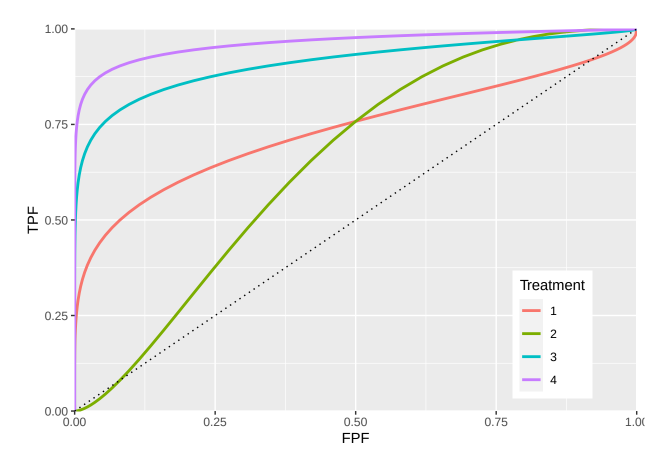
12.2 The binormal model

The binormal model has two parameters, a and b. The signal (or diseased cases) distribution has unit standard deviation. The noise (or non-diseased cases) distribution has standard deviation b. The a parameter is the separation of the two distributions.

12.3 Improper ROCs

Binormal model fits invariably lead to ROC curves that inappropriately cross the chance diagonal, leading to a prediction of a region of the ROC curve where performance is worse than chance, even for expert observers. By convention, such curves are termed *improper*. This vignette illustrates improper ROCs predicted by the binormal model.

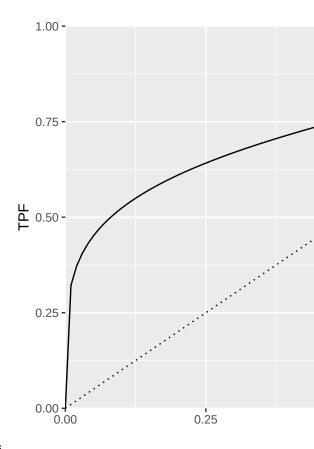
```
aArray <- c(0.7, 0.7, 1.5, 2)
bArray <- c(0.5, 1.5, 0.5, 0.5)
chance_diag <- data.frame(x = c(0,1), y = c(0,1))
p <- PlotBinormalFit(aArray, bArray) +
    scale_x_continuous(expand = c(0, 0)) +
    scale_y_continuous(expand = c(0, 0)) +
    theme(legend.position = c(0.85, 0.2))
p <- p + geom_line(data = chance_diag, aes(x = x, y = y), linetype="dotted")
print(p)</pre>
```



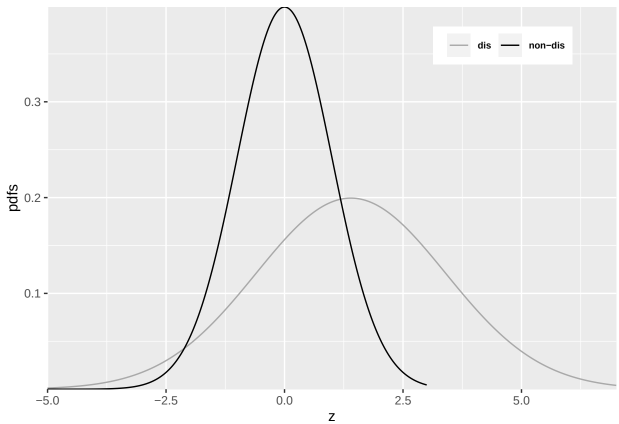
The red plot is the clearest example of an improper ROC. This type of curve occurs whenever b < 1. The chance line crossing near the upper right corner, around (0.919,0.919), and the fact that the ROC curve must eventually reach (1, 1) implies the curve must turn upwards as one approaches (1, 1), thereby displaying a "hook". Whenever b != 1 the hook is there, regardless of whether it is easily visible or not. If b < 1 the hook is near the upper right corner. If b > 1 the hook is near the origin (see green line, corresponding to b = 1.5). With increasing a the hook is less prominent (blue line corresponding to a = 1.5, b = 0.5 and purple line corresponding to a = 2, b = 0.5). But it is there.

12.3. IMPROPER ROCS 97

12.4 Reason for improper ROCs



The reason for the "hook" "becomes apparent upon examination of the pdfs.



a = 0.7 , b = 0.5

Since b < 1, the diseased pdf is broader and has a lower peak (since the integral under each distribution is unity) than the non-diseased pdf. Sliding an imaginary threshold to the left, starting from the extreme right, one sees that initially, just below z = 7, the diseased distribution starts being "picked up" while the non-diseased distribution is not "picked up", causing the ROC to start with infinite slope near the origin (because TPF is increasing while FPF is not). Around z = 2.5 the non-diseased distribution starts being "picked up", causing the ROC slope to decrease. Around z = -3, almost all of the non-diseased distribution has been "picked up" which means FPF is near unity, but since not all of the broader diseased distribution has been "picked up", TPF is less than unity. Here is a region where TPF < FPF, meaning the operating point is below the chance diagonal. As the threshold is lowered further, TPF continues to increase, as the rest of the diseased distribution is "picked up" while FPF stays almost constant at unity. In this region, the ROC curve is approaching the upper right corner with almost infinite slope (because TPF is increasing but FPF is not).

Chapter 13

Degenerate ROC curves

13.1 Two helper functions

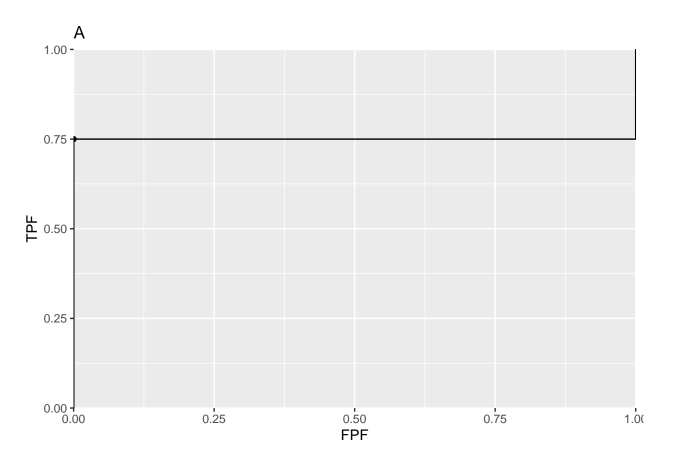
13.2 Degenerate datasets

Metz defined binormal degenerate data sets as those that result in exact-fit binormal ROC curves of inappropriate shape consisting of a series of horizontal and/or vertical line segments in which the ROC "curve" crosses the chance line. The crossing of the chance line occurs because the degenerate data sets can be fitted exactly by infinite or zero values for the model slope parameter b, and infinite values for the decision thresholds, or both.

13.3 Understanding degenerate datasets

To understand this, consider that the non-diseased distribution is a Dirac delta function centered at zero (by definition such a function integrates to unity) and the unit variance diseased distribution is centered at 0.6744898. In other words this binormal model is characterized by $\mathbf{a}=0.6744898$ and $\mathbf{b}=0$. What is the expected ROC curve? As the threshold ζ is moved from the far right, gradually to the left, TPF will increase but FPF is stuck at zero until the threshold reaches zero. Just before reaching this point, the coordinates of the ROC operating point are (0,0.75). The 0.75 is due to the fact that $\mathbf{z}=0$ is -0.6744898 units relative to the center of the diseased distribution, so the area under the diseased distribution below $\mathbf{z}=0$ is 0.25. Since pnorm is the probability below the threshold, TPF must be its complement, namely 0.75. This explains the operating point (0,0.75), which lies on the y-axis. As the threshold crosses the zero-width delta function, FPF shoots up from 0 to 1, but TPF stays constant. Therefore, the operating point has jumped from (0,0.75) to (1,0.75). When the threshold is reduced further, the operating point moves up vertically, along the right side of the ROC plot, until the threshold is so small that virtually all of diseased distribution exceeds it and the operating point reaches (1,1). The ROC curve is illustrated in plot A.

```
plotOP <- data.frame(FPF = 0, TPF = 0.75)
a <- 0.6744898; b <- 0
plotCurve <- BMPoints(a, b)
figA <- ggplot(mapping = aes(x = FPF, y = TPF)) +
    geom_line(data = plotCurve) +
    geom_point(data = plotOP) +
    scale_x_continuous(expand = c(0, 0)) +
    scale_y_continuous(expand = c(0, 0)) +
    ggtitle("A")
print(figA)</pre>
```



This is an extreme example of an ROC curve with a "hook". If the data is such that the only operating point provided by the observer is (0,0.75) then this curve will be an exact fit to the operating point.

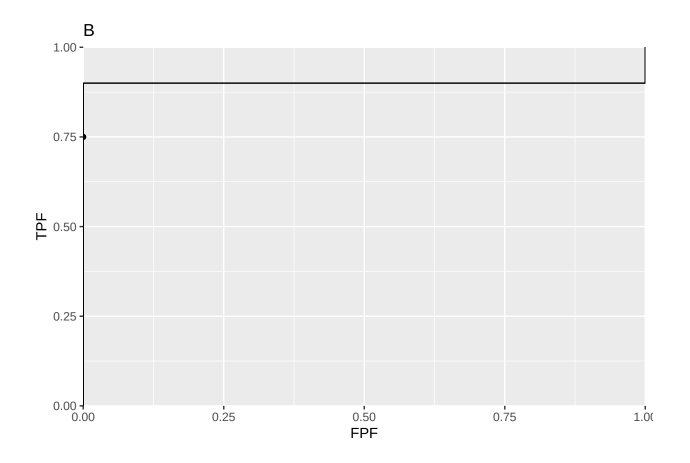
13.4 The exact fit is not unique

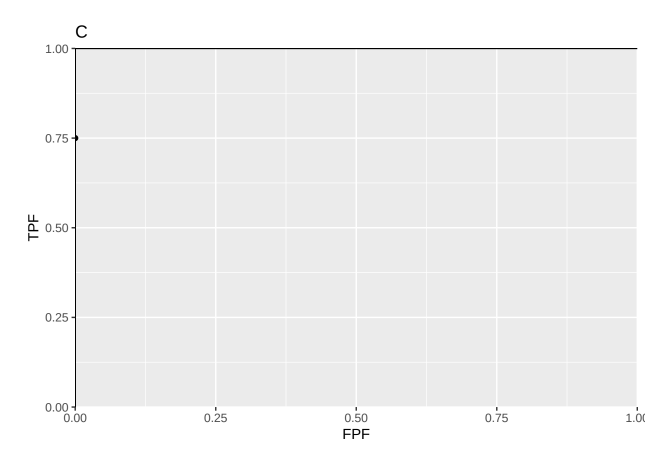
Actually, given one operating point (0, 0.75) the preceding fit is not even unique. If the diseased distribution is shifted appropriately to the right of its previous position, and one can determine the necessary value of a, then the ROC curve will shoot upwards through the operating point (0, 0.75) to (0, 0.9), as in plot B, before proceeding horizontally to (1, 0.9) and then completing the curve to (1, 1). If the diseased distribution is shifted well to the right, i.e., a is very large, then the ROC curve will shoot upwards past the operating point, as in plot C, all the way to (0,1) before proceeding horizontally to (1, 1).

```
a <- 1.281552; b <- 0
plotCurve <- BMPoints(a, b)
figB <- ggplot(mapping = aes(x = FPF, y = TPF)) +
    geom_line(data = plotCurve) +
    geom_point(data = plotOP) +
    scale_x_continuous(expand = c(0, 0)) +
    scale_y_continuous(expand = c(0, 0)) +
    ggtitle("B")

a <- Inf; b <- 0
plotCurve <- BMPoints(a, b)
figC <- ggplot(mapping = aes(x = FPF, y = TPF)) +
    geom_line(data = plotCurve) +
    geom_point(data = plotOP) +</pre>
```

```
scale_x_continuous(expand = c(0, 0)) +
scale_y_continuous(expand = c(0, 0)) +
ggtitle("C")
print(figB);print(figC)
```





All of these represent exact fits to the observed operating point, with b=0 and different values of a. Not one of them is reasonable.

13.5 Comments on degeneracy

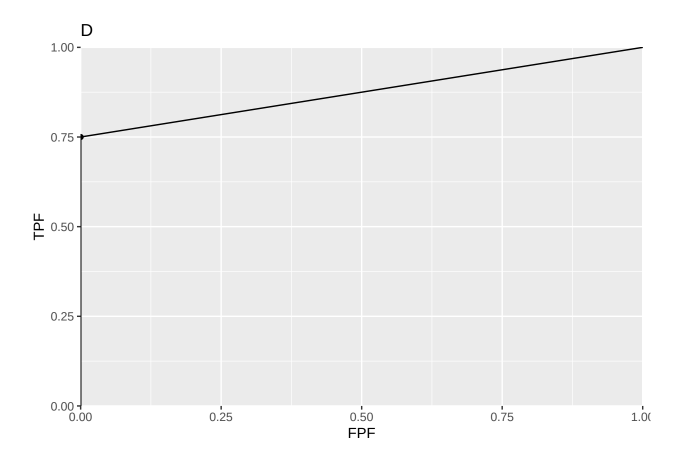
Degeneracy occurs if the observer does not provide any interior operating points. So why worry about it? Perhaps one has a non-cooperating observer, who is not heeding the instructions to spread the ratings, use all the bins. A simple example shows that the observer could if fact be cooperating fully and is still unable to provide any interior data points. Consider 100 diseased cases consisting of 75 easy cases and 25 difficult ones and 100 easy non-diseased cases. The observer is expected to rate the 75 easy diseased cases as fives, the difficult ones as ones and the 100 non-diseased cases are rated ones. No amount of coaxing please, please spread your ratings is going to convince this observer to rate with twos, threes and fours any of the 75 easy diseased cases. If the cases are obviously diseased, and that is what is meant by easy cases, they are supposed to be rated fives: definitely diseased. Forcing them to rate some of them as probably diseased or possibly diseased would be irrational and guilty of bending the reading paradigm to fit the convenience of the researcher (early in his research career, the author used to believe in the existence of non-cooperating observers, so Metz's advice to spread the ratings did not seem unreasonable at that time).

13.6 A reasonable fit to the degenerate dataset

If the dataset yields a single operating point (0, 0.75), what is a reasonable ROC plot? There is a theorem that given an observed operating point, the line connecting that point to (1, 1) represents a lower bound on achievable performance by the observer. The observer using a guessing mechanism to classify the remaining cases achieves the lower bound. Here is an explanation of this theorem. Having rated the 75 easy diseased cases as fives, the observer is left with 25 diseased cases and 100 non-diseased cases, all of which appear definitely non-diseased to the observer. Suppose the observer randomly rates 20% of the remaining cases as fours. This would pick up five of the actually

diseased cases and 20 non-diseased ones. Therefore, the total number of diseased cases rated four or higher is 80, and the corresponding number of non-diseased cases is 20. The new operating point of the observer is (0.20, 0.80). Now, one has two operating points, the original one on the y-axis at (0, 0.75) and an interior point (0.20, 0.80). Next, instead of randomly rating 20% of the remaining cases as fours, the observer rates 40% of them as fours, then the interior point would have been (0.40, 0.85). The reader can appreciate that simply by increasing the fraction of remaining cases that are randomly rated fours, the observer can move the operating point along the straight line connecting (0, 0.75) and (1, 1), as in plot D. Since a guessing mechanism is being used, this must represent a lower bound on performance. The resulting ROC curve is proper and the net AUC = 0.875.

```
mu <- Inf; alpha <- 0.75
plotCurve <- CBMPoints(mu, alpha)
figD <- ggplot(mapping = aes(x = FPF, y = TPF)) +
    geom_line(data = plotCurve) +
    geom_point(data = plotOP) +
    scale_x_continuous(expand = c(0, 0)) +
    scale_y_continuous(expand = c(0, 0)) +
    ggtitle("D")
print(figD)</pre>
```



For this dataset this is in fact the fit yielded by the contaminated binormal model (CBM) and the radiological search model (RSM). Why should one select the lowest possible performance consistent with the data? Because it yields a *unique* value for performance: any higher performance would not be unique.

Chapter 14

PROPROC ROC curves

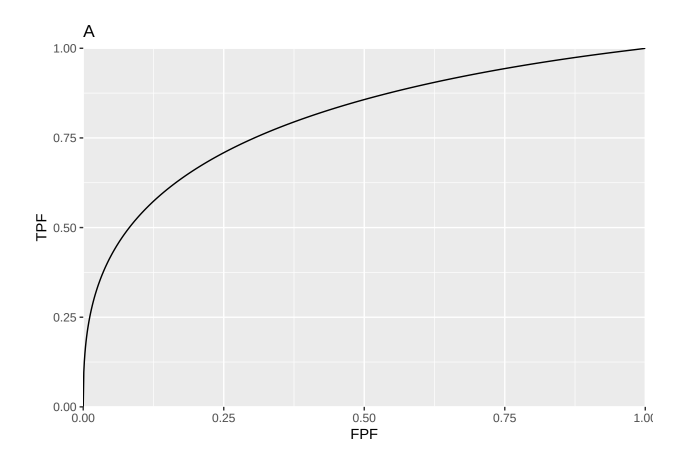
- 14.1 Helper functions
- 14.2 Definitions of PROPROC parameters in terms of binormal model parameters

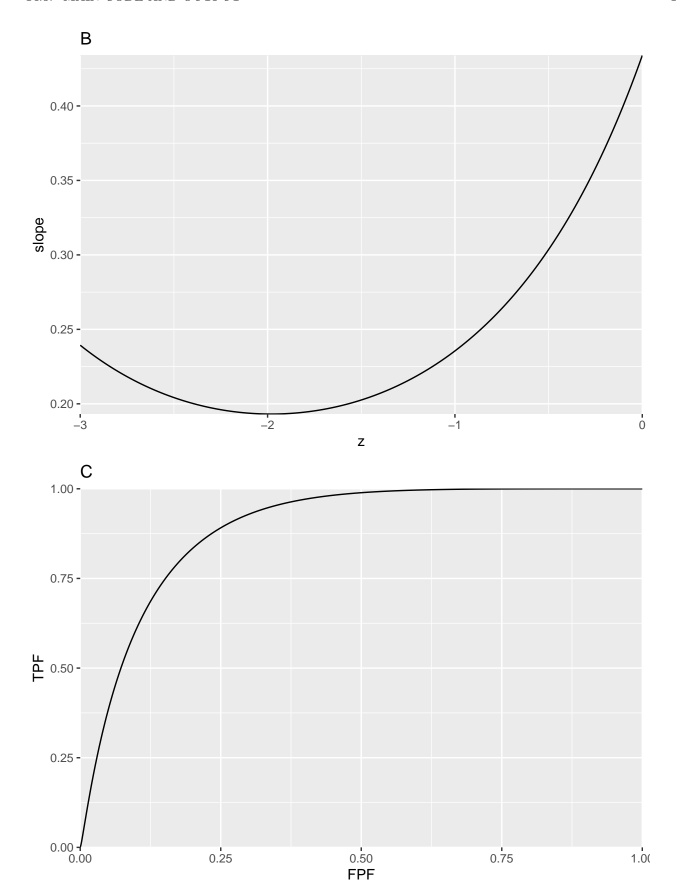
$$c = \frac{b-1}{b+1}; d_a = \frac{\sqrt{2}a}{\sqrt{1+b^2}}$$

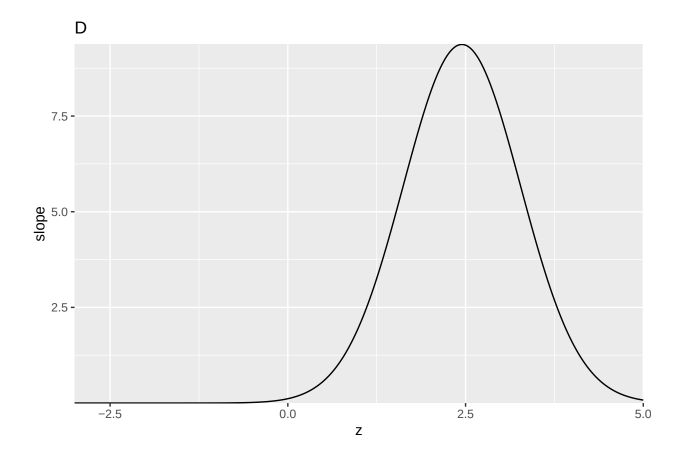
14.3 Main code and output

```
c1Arr \leftarrow c(-0.1322804, 0.2225588); daArr \leftarrow c(1.197239, 1.740157)
myLabel <- c("A", "B", "C", "D")</pre>
myLabelIndx <- 1</pre>
for (i in 1:2)
  c1 <- c1Arr[i]
  da <- daArr[i]</pre>
  ret <- Transform2ab(da, c1)
  a <- ret$a;b <- ret$b
  if (i == 1) z <- seq(-3, 0, by = 0.01) # may need to adjust limits to view detail of slope plot
  if (i == 2) z <- seq(-3, 5, by = 0.01) # may need to adjust limits to view detail of slope plot
  FPF \leftarrow seq(0.0, 1, 0.001)
  TPF <- rocY(FPF, a, b)</pre>
  rocPlot <- data.frame(FPF = FPF, TPF = TPF)</pre>
  plotRoc <- ggplot(rocPlot, aes(x = FPF, y = TPF)) +</pre>
    geom_line() +
   scale_x_continuous(expand = c(0, 0)) +
    scale_y_continuous(expand = c(0, 0))
    ggtitle(myLabel[myLabelIndx]);myLabelIndx <- myLabelIndx + 1</pre>
  slope <-b*dnorm(a-b*z)/dnorm(-z) # same as likelihood ratio</pre>
  slopePlot <- data.frame(z = z, slope = slope)</pre>
  p <- ggplot(slopePlot, aes(x = z, y = slope)) +</pre>
```

```
geom_line() +
scale_x_continuous(expand = c(0, 0)) +
scale_y_continuous(expand = c(0, 0)) +
ggtitle(myLabel[myLabelIndx]);myLabelIndx <- myLabelIndx + 1
print(plotRoc);print(p)
}</pre>
```







14.4 Discussion

Plot A is for c1 = -0.1322804, da = 1.197239 while plot C is for c1 = 0.2225588, da = 1.740157. Plots B and D are the corresponding slope plots as functions of the binormal model z-sample. In plot A, the slope is infinite near the origin and the curve approaches the upper-right corner with finite slope. The situation is reversed in plot C where the slope is finite near the origin and the curve approaches the upper-right corner with zero slope.

These two readers are from a clinical dataset, dataset01. Highest rating inferred ROC data from original FROC data, were analyzed by PROPROC and the resulting parameter values are coded here. They were chosen as they demonstrate key differences in the shapes of proper ROC plots. Plot A corresponds to a negative value of c1, which implies b < 1. The slope of the proper ROC is infinite near the origin and approaches a positive constant near the upper right corner of the ROC. Plot C is for a positive value of c1, i.e., for b > 1. Now the slope of the proper ROC is finite near the origin and approaches zero near the upper right corner.

Considering plot D, as one "cuts" the slope axis horizontally with a sliding threshold, starting with very high values and moving downwards, the slope of the ROC curve starts at the origin with a large but finite value. This corresponds to the peak in plot D. Above the peak, there are no solutions for **z**. The slope decreases monotonically to zero, corresponding to the flattening out of the slope at zero for **z** ~ -2.

The two values of **z** corresponding to each cut implies, of course, that the binormal model based proper algorithm has to do a lot of bookkeeping, since each horizontal cut splits the decision axis into 3 regions. One can think of shrinking each of plots B & D horizontally to zero width, and all that remains is the slope axis with a thick vertical line superimposed on it, corresponding to the horizontally collapsed curves. In plot B the vertical line extends from positive infinity down to about 0.1, and represents the range of decision variable samples encountered by the observer on the likelihood ratio scale. In plot D the vertical line extends from a finite value (~9.4) to zero. For the stated binormal model parameters values outside of these ranges are not possible.

Chapter 15

Metz Eqn 36 numerical check

15.1 Helper functions

15.2 Main code and output

```
npts <- 10000
for (i in 1:2) {
  for (j in 1:5) {
   C \leftarrow c1[i,j]
   da <- d_a1[i,j]
   ret <- GetLimits(da,C)
   LL <- ret$LL;UL <- ret$UL
   vc <- seq (LL, UL, length.out = npts)
   TPF <- TruePositiveFraction (vc, da, C)
    FPF <- FalsePositiveFraction (vc, da, C)
   FPF <- rev(FPF); TPF <- rev(TPF)</pre>
   df2 <- data.frame(FPF = FPF, TPF = TPF)</pre>
    # do integral numerically
   numAuc <- trapz(FPF, TPF)</pre>
    # Implement Eqn. 36 from Metz-Pan paper
   rho <-(1-C^2)/(1+C^2); sigma <- rbind(c(1, rho), c(rho, 1))
    lower \leftarrow \text{rep}(-\text{Inf}, 2); \text{upper} \leftarrow c(-\text{da/sqrt}(2), 0)
    aucProproc <- pnorm(da/sqrt(2)) + 2 * pmvnorm(lower, upper, sigma = sigma)
    aucProproc <- as.numeric(aucProproc)</pre>
    cat("i = ", i, "j = ", j, "C = ", C, ", da = ", da, "aucProproc = ", aucProproc, "Norm. Diff. = ", (aucPr
  }
#> i = 1 j = 1 C = -0.1322804 , da = 1.197239 aucProproc = 0.8014164 Norm. Diff. = 3.520017e-08
\#>i=1 j=2 C=-0.08696513 , da=1.771176 aucProproc=0.8947898 Norm. Diff.=4.741875e-08
\#>i=1 j=3 C=-0.1444419 , da=1.481935 aucProproc=0.8526605 Norm. Diff.=3.515431e-08
\#>i=1 j = 4 C = 0.08046016 , da = 1.513757 aucProproc = 0.8577776 Norm. Diff. = 4.971428e-08
\#>i=1 j=5 C=0.2225588 , da=1.740157 aucProproc=0.8909392 Norm. Diff.=2.699855e-08
\#>i=2 j=1 C=-0.08174248 , da=0.6281251 aucProproc=0.6716574 Norm. Diff.=2.801793e-08
\#>i=2 j=2 C=0.04976448 , da=0.9738786 aucProproc=0.7544739 Norm. Diff.=5.275242e-08
#> i = 2 j = 3 C = -0.1326126 , da = 1.155871 aucProproc = 0.7931787 Norm. Diff. = 3.472577e-08
\#>i=2 j=4 C=0.1182226 , da=1.620176 aucProproc=0.8740274 Norm. Diff.=3.922161e-08
\#>i=2 j = 5 C = 0.0781033 , da = 0.8928816 aucProproc = 0.7360989 Norm. Diff. = 3.798459e-08
```

15.3 Discussion

Note the close correspondence between the formula, Eqn. 36 in the Metz-Pan paper and the numerical estimate. As a historical note, Eqn. 31 and Eqn. 36 (they differ only in parameterizations) in the referenced publication are provided without proof – it was probably obvious to Prof Metz or he wanted to leave it to us "mere mortals" to figure it out, as a final parting gesture of his legacy. The author once put a significant effort into proving it and even had a bright graduate student from the biostatistics department work on it to no avail. The author has observed that these equations always yield very close to the numerical estimates, to within numerical precisions, so the theorem is correct empirically, but he has been unable to prove it analytically. It is left as an exercise for a gifted reader to prove/disprove these equations.

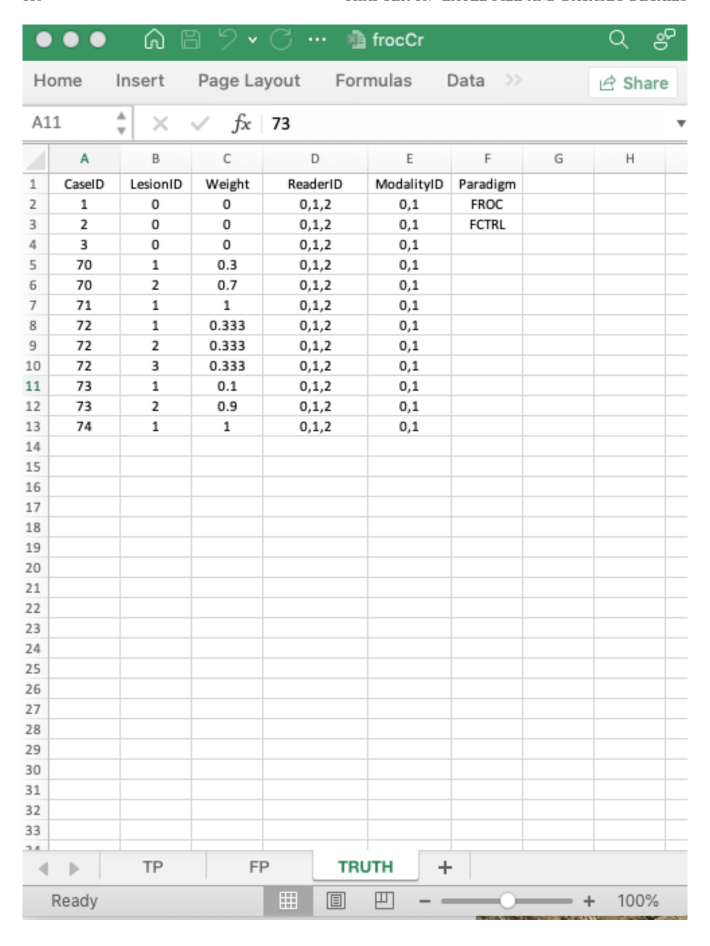
Software details

Chapter 16

Excel file and dataset details

16.1 Introduction

- This chapter is included to document recent Excel file format changes and the new dataset structure.
- We illustrate with a toy FROC data file in the RJafroc GitHub repository, inst/extdata/toyFiles/FROC/frocCr.xlsx, in which 3 readers interpret 3 non-diseased and 5 diseased cases using the FROC paradigm.
- The Excel file has three worksheets named Truth, NL (or FP) and LL (or TP).



16.2 The Truth worksheet

- The Truth worksheet contains 6 columns: CaseID, LesionID, Weight, ReaderID, ModalityID and Paradigm.
- The non-diseased cases are numbered 1,2,3; the diseased cases are numbered 70, 71, 72, 73, 74; i.e., K1 = 3, K2 = 5 and K = 8.
- The ReaderID field has three values 0, 1, 2.
- ModalityID: The first modality is labeled 0 and the second is '1'.
- Note that ReaderID and ModalityID are text formatted labels.
- Paradigm: The contents of this field are FROC and FCTRL (case insensitive, FCTRL stands for a factorial or fully crossed study design where each reader interprets all cases in all modalities).

16.3 The structure of the FROC factorial dataset

The following code reads the Excel file into a dataset object x:

Note that newExcelFileFormat must be set to TRUE to read the new Excel format dataset. The default
is FALSE which applies to the old format Excel file with only the first three columns present in the Truth
worksheet.

The structure of x is shown below.

```
str(x)
#> List of 3
#> $ ratings
                 :List of 3
    ..$ NL : num [1:2, 1:3, 1:8, 1:2] 1.02 2.89 2.21 3.01 2.14 ...
    ..$ LL : num [1:2, 1:3, 1:5, 1:3] 5.28 5.2 5.14 4.77 4.66 4.87 3.01 3.27 3.31 3.19 ...
#>
    ..$ LL_IL: logi NA
   $ lesions
                 :List of 3
#>
    ..$ perCase: int [1:5] 2 1 3 2 1
             : num [1:5, 1:3] 1 1 1 1 1 ...
    ..$ IDs
    ..$ weights: num [1:5, 1:3] 0.3 1 0.333 0.1 1 ...
   $ descriptions:List of 7
#>
    ..$ fileName : chr "frocCr"
                     : chr "FROC"
#>
    ..$ type
#>
    ..$ name
                    : logi NA
#>
    ..$ truthTableStr: num [1:2, 1:3, 1:8, 1:4] 1 1 1 1 1 1 1 1 1 1 ...
#>
    ..$ design
                 : chr "FCTRL"
#>
    ..$ modalityID : Named chr [1:2] "0" "1"
    ....- attr(*, "names")= chr [1:2] "0" "1"
#>
                    : Named chr [1:3] "0" "1" "2"
#>
    ..$ readerID
    ....- attr(*, "names")= chr [1:3] "0" "1" "2"
```

- The dataset x is a list variable with 3 members: x\$ratings, x\$lesions and x\$descriptions.
- There are K2 = 5 diseased cases (the length of the third dimension of x\$ratings\$LL) and K1 = 3 non-diseased cases (the length of the third dimension of x\$ratings\$NL minus K2).
- x\$ratings\$NL is a [2, 3, 8, 2] array containing the NL ratings on non-diseased and diseased cases.
- x\$ratings\$LL is a [2, 3, 5, 3] array containing the ratings of LLs on diseased cases.
- The maximum number of lesions per case in the dataset is 3.
- The number of lesions per diseased case is the vector x\$lesions\$perCase, i.e., 2, 1, 3, 2, 1 in this example.

- The x\$descriptions\$dataType member is FROC, which specifies the data collection method.
- The x\$descriptions\$modalityID member is a vector with two elements 0, 1, naming the two modalities.
- The x\$readerID member is a vector with three elements 0, 1, 2, naming the three readers.
- The x\$descriptions\$design member is FCTRL; specifies the study design.
- The x\$descriptions\$truthTableStr member quantifies the structure of the dataset, explained next.

16.4 The truthTableStr member

- For this dataset I = 2, J = 3 and K = 8.
- truthTableStr is a 2 x 3 x 8 x 4 array, i.e., I x J x K x (maximum number of lesions per case plus 1). The plus 1 is needed to accommodate non-diseased cases.
- Each entry in this array is either 1, meaning the corresponding interpretation happened, or NA, meaning the corresponding interpretation did not happen.

16.4.1 Explanation for non-diseased cases

Since the fourth index is set to 1, in the following code only non-diseased cases yield ones and all diseased cases yield NA.

```
all(x$descriptions$truthTableStr[,,1:3,1] ==1)
#> [1] TRUE
all(is.na(x$descriptions$truthTableStr[,,4:8,1]))
#> [1] TRUE
```

16.4.2 Explanation for diseased cases with one lesion

Since the fourth index is set to 2, in the following code all non-diseased cases yield NA and all diseased cases yield 1 as all diseased cases have at least one lesion.

```
all(is.na(x$descriptions$truthTableStr[,,1:3,2]))
#> [1] TRUE
all(x$descriptions$truthTableStr[,,4:8,2] == 1)
#> [1] TRUE
```

16.4.3 Explanation for diseased cases with two lesions

Since the fourth index is set to 3, in the following code all non-diseased cases yield NA; the first diseased case 70 yields 1 (this case contains two lesions); the second disease case 71 yields NA (this case contains only one lesion); the third disease case 72 yields NA (this case contains only two lesions); the fourth disease case 73 yields 1 (this case contains two lesions); the fifth disease case 74 yields NA (this case contains one lesion).

```
# all non diseased cases
all(is.na(x$descriptions$truthTableStr[,,1:3,3]))
#> [1] TRUE
# first diseased case
all(x$descriptions$truthTableStr[,,4,3] == 1)
#> [1] TRUE
# second diseased case
all(is.na(x$descriptions$truthTableStr[,,5,3]))
#> [1] TRUE
# third diseased case
all(x$descriptions$truthTableStr[,,6,3] == 1)
```

```
#> [1] TRUE
# fourth diseased case
all(x$descriptions$truthTableStr[,,7,3] == 1)
#> [1] TRUE
# fifth diseased case
all(is.na(x$descriptions$truthTableStr[,,8,3]))
#> [1] TRUE
```

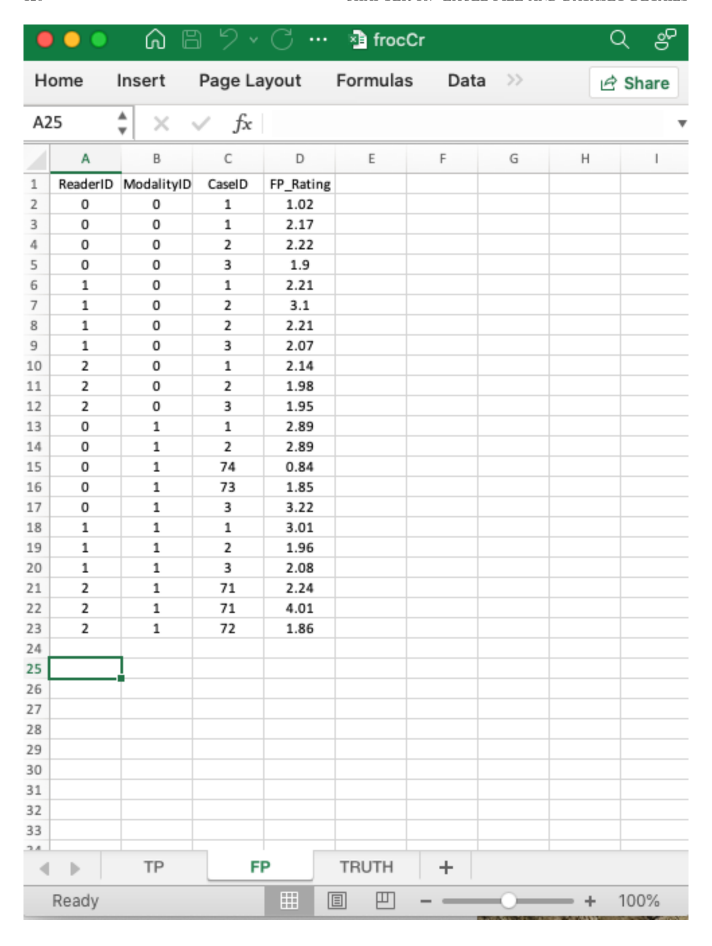
16.4.4 Explanation for diseased cases with three lesions

Since the fourth index is set to 4, in the following code all non-diseased cases yield NA; the first diseased case 70 yields NA (this case contains two lesions); the second disease case 71 yields NA (this case contains one lesion); the third disease case 72 yields NA (this case contains two lesions); the fourth disease case 73 yields 1 (this case contains three lesions); the fifth disease case 74 yields NA (this case contains one lesion).

```
# all non diseased cases
all(is.na(x$descriptions$truthTableStr[,,1:3,4]))
#> [1] TRUE
# first diseased case
all(is.na(x$descriptions$truthTableStr[,,4,4]))
#> [1] TRUE
# second diseased case
all(is.na(x$descriptions$truthTableStr[,,5,4]))
#> [1] TRUE
# third diseased case
all(x$descriptions$truthTableStr[,,6,4] == 1)
#> [1] TRUE
# fourth diseased case
all(is.na(x$descriptions$truthTableStr[,,7,4]))
#> [1] TRUE
# fifth diseased case
all(is.na(x$descriptions$truthTableStr[,,8,4]))
#> [1] TRUE
```

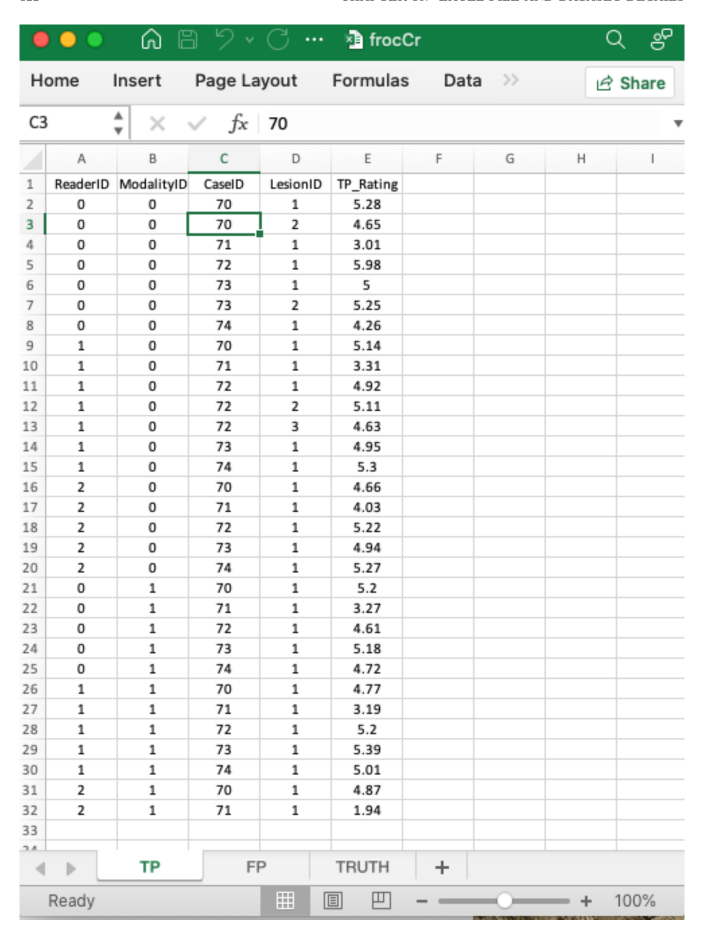
16.5 The non-lesion localization (NL or FP) ratings

These are found in the FP or NL worksheet:



- The common vertical length is 22 in this example.
- ReaderID: the reader labels: 0, 1,2, as declared in the Truth' worksheet.
- ModalityID: the modality labels: 0 or 1, as declared in the Truth worksheet.
- CaseID: 1, 2, 3, 71, 72, 73, 74, as declared in the Truth worksheet; note that not all cases have NL marks on them.
- NL_Rating: the ratings of non-diseased cases.

16.6 The lesion localization (LL or TP) ratings



- This worksheet has the ratings of diseased cases.
- ReaderID: the reader labels: these must be from 0, 1, 2, as declared in the Truth worksheet.
- ModalityID: 0 or 1, as declared in the Truth worksheet.
- CaseID: these must be from 70, 71, 72, 73, 74, as declared in the Truth worksheet; not all diseased cases have LL marks.
- LL_Rating: the ratings of diseased cases.

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