# EHR Vignette for Extract-Med and Pro-Med-NLP

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# Introduction

The EHR package provides several modules to perform diverse medication-related studies using data from electronic health record (EHR) databases. Especially, the package includes modules to perform pharmacokinetic/pharmacodynamic (PK/PD) analyses using EHRs, as outlined in Choi, et al.<sup>1</sup>, and additional modules will be added in future. This vignette describes two modules in the system, when drug dosing information should be obtained from unstructured clinical notes. The process starts with data extraction (Extract-Med), then moves on to data processing (Pro-Med-NLP), from which a final data set can be built.

library(EHR)

# **Extract-Med**

The Extract-Med module uses a natural language processing (NLP) system called medExtractR.<sup>2</sup> The medExtractR system is a medication extraction system that uses regular expressions and rule-based approaches to identify key dosing information including drug name, strength, dose amount, frequency or intake time, dose change, and last dose time. Function arguments can be specified to allow the user to tailor the medExtractR system to the particular drug or dataset of interest, improving the quality of extracted information.

#### Setup of extractMed

The function extractMed will run the Extract-Med module. In order to run the Extract-Med module with medExtractR, certain function arguments must be specified, including:

- note\_fn: The file name of the note on which to run the system. This can be either a single file name (e.g., "clinical\_note01.txt") or a vector or list of file names (e.g., c("clinical\_note01.txt", "clinical\_note02.txt")).
- drugnames: Names of the drugs for which we want to extract medication dosing information. This can include any way in which the drug name might be represented in the clinical note, such as generic name (e.g., "lamotrigine"), brand name (e.g., "Lamictal"), or an abbreviation (e.g., "LTG").
- drgunit: The unit of the drug(s) listed in drugnames, for example "mg".
- windowlength: Length of the search window around each found drug name in which to search for dosing information. There is no default for this argument, requiring the user to carefully consider its value through tuning (see tuning section below).
- max\_edit\_dist: The maximum edit distance allowed when identifying drugnames. Maximum edit distance determines the difference between two strings, and is defined as the number of insertions, deletions, or substitutions required to change one string into the other. This allows us to capture

misspellings in the drug names we are searching for, and its value should be carefully considered through tuning (see tuning section below).

- The default value is '0', or exact spelling matches to drugnames. A value of 0 is always used for drug names with less than 5 characters regardless of the value set by max\_edit\_dist.
- A value of 1 would capture mistakes such as a single missing or extra letter, e.g., "tacrlimus" or "tacroolimus" instead of "tacrolimus"
- A value of 2 would capture these mistakes or a single transposition, e.g. "tcarolimus" instead of "tacrolimus"
- Higher values (3 or above) would capture increasingly more severe mistakes, though setting the value too high can cause similar words to be mistaken as the drug name.

Generally, the function call to extractMed is

```
extractMed(note_fn, drugnames, drgunit, windowlength, max_edit_dist, ...)
```

where ... refers to additional arguments to medExtractR (see package documentation for details<sup>3</sup>). Examples of additional arguments include:

- strength\_sep, where users can specify special characters to separate doses administered at different times of day. For example, consider the drug name "lamotrigine" and the phrase "Patient is on lamotrigine 200-300", indicating that the patient takes 200 mg of the drug in the morning and 300 mg in the evening. Setting strength\_sep = c('-') would allow extractMed to identify the expression 200-300 as "Dose" (i.e., dose given intake) since they are separated by the special character "-". The default value is NULL.
- lastdose, a logical input specifying whether or not the last dose time entity should be extracted. Default value is FALSE.

As mentioned above, some arguments to extractMed should be specified through a tuning process. In a later section, we briefly describe the process by which a user could tune the medExtractR system (and by extension the extractMed function) using a validated gold standard dataset.

### Running extractMed

Below, we demonstrate how to execute the Extract-Med module using sample notes for two drugs: tacrolimus (simpler prescription patterns, used to prevent rejection after organ transplant) and lamotrigine (more complex prescription patterns, used to treat epilepsy). The arguments specified for each drug here were determined based on training sets of 60 notes for each drug.<sup>2</sup> Note that for tacrolimus we enter the file names as a list, and for lamotrigine we enter the file names as a character vector; either form of input is acceptable to the extractMed function and will produce the same output. We also specify lastdose=TRUE for tacrolimus to extract information about time of last dose, and strength\_sep="-" for lamotrigine.

```
max_edit_dist = 2,
lastdose=TRUE)
```

## running notes 1-3 in batch 1 of 1 (100%)

```
# lamotrigine note file name
lam fn <- c(
  system.file("examples", "lampid1_2016-02-05_note4_1.txt", package = "EHR"),
  system.file("examples", "lampid1_2016-02-05_note5_1.txt", package = "EHR"),
  system.file("examples", "lampid2_2008-07-20_note6_1.txt", package = "EHR"),
  system.file("examples", "lampid2_2012-04-15_note7_1.txt", package = "EHR")
)
# execute module
lam_mxr <- extractMed(lam_fn,</pre>
                       drugnames = c("lamotrigine", "lamotrigine XR",
                                      "lamictal", "lamictal XR",
                                      "LTG", "LTG XR"),
                       drgunit = "mg",
                       windowlength = 130,
                       max_edit_dist = 1,
                       strength sep="-")
```

#### ## running notes 1-4 in batch 1 of 1 (100%)

The format of output from the Extract-Med module is a data.frame with 4 columns:

- filename: The file name of the corresponding clinical note, to label results.
- entity: The label of the entity for the extracted expression.
- expr: Expression extracted from the clinical note.
- pos: Position of the extracted expression in the note, in the format startPosition:stopPosition
- ## tacrolimus medExtractR output:

```
##
                           filename
                                        entity
                                                      expr
                                                                  pos
## 1 tacpid1_2008-06-26_note1_1.txt
                                      DrugName
                                                   Prograf
                                                              930:937
## 2 tacpid1_2008-06-26_note1_1.txt
                                      Strength
                                                      1 mg
                                                              951:955
## 3 tacpid1 2008-06-26 note1 1.txt
                                       DoseAmt
                                                              956:957
                                                          3
## 4 tacpid1_2008-06-26_note1_1.txt Frequency twice a day
                                                              976:987
## 5 tacpid1_2008-06-26_note1_1.txt
                                                   prograf 2709:2716
                                      DrugName
## 6 tacpid1_2008-06-26_note1_1.txt
                                          Dose
                                                        3mg 2717:2720
## lamotrigine medExtractR output:
##
                           filename
                                        entity
                                                      expr
                                                                pos
## 1 lampid1_2016-02-05_note4_1.txt
                                      DrugName
                                                  Lamictal 810:818
## 2 lampid1_2016-02-05_note4_1.txt
                                          Dose
                                                    300 mg 819:825
## 3 lampid1_2016-02-05_note4_1.txt Frequency
                                                       BID 826:829
## 4 lampid1_2016-02-05_note4_1.txt
                                      DrugName Lamotrigine 847:858
## 5 lampid1_2016-02-05_note4_1.txt
                                                     200mg 859:864
                                      Strength
## 6 lampid1_2016-02-05_note4_1.txt
                                                       1.5 865:868
                                       DoseAmt
```

The lamotrigine results can be immediately put into the parse step of the Pro-Med-NLP module. For the tacrolimus output, we chose to also extract the last dose time entity by specifying lastdose=TRUE. The last dose time entity is extracted as raw character expressions from the clinical note, and must first be converted

to a standardized datetime format, which is addressed in a later section (see "Handling lastdose" section). The output of extractMed must be saved as a CSV file, the filename of which will serve as the first input to the Pro-Med-NLP module.

```
# save as csv files
write.csv(tac_mxr, file='tac_mxr.csv', row.names=FALSE)
write.csv(lam_mxr, file='lam_mxr.csv', row.names=FALSE)
```

#### Tuning the medExtractR system

In a previous section, we mentioned that parameters within the Extract-Med module should be tuned in order to ensure higher quality of extracted drug information. This section describes how we go about tuning medExtractR, the NLP system underlying the Extract-Med module.

In order to tune medExtractR, we recommend selecting a small set of tuning notes, from which the parameter values can be selected. Below, we describe this process with a set of three notes (note that these notes were chosen for the purpose of demonstration, and we recommend using tuning sets of at least 10 notes).

Once a set of tuning notes has been curated, they must be manually annotated by reviewers to identify the information that should be extracted. This process produces a gold standard set of annotations, which identify the correct drug information of interest. This includes entities like the drug name, strength, and frequency. For example, in the phrase

Patient is taking **lamotrigine** 300 mg in the morning and 200 mg in the evening

bolded, italicized, and underlined phrases represent annotated drug names, dose (i.e., dose given intake), and intake times, respectively. These annotations are stored as a dataset.

First, we read in the annotation files for three example tuning notes, which can be generated using an annotation tool, such as the Brat Rapid Annotation Tool (BRAT) software.<sup>4</sup> By default, the output file from BRAT is tab delimited with 3 columns: an annotation identifier, a column with labelling information in the format "label startPosition stopPosition", and the annotation itself, as shown in the example below:

```
##
     id
                  entity
                          annotation
## 1 T1
          DrugName 19 30 lamotrigine
              Dose 31 37
## 2 T2
                               300 mg
## 3 T3 IntakeTime 45 52
                              morning
## 4 T4
              Dose 57 63
                               200 mg
## 5 T5 IntakeTime 71 78
                              evening
```

In order to compare with the medExtractR output, the format of the annotation dataset should be four columns with:

- 1. The file name of the corresponding clinical note
- 2. The entity label of the annotated expression
- 3. The annotated expression
- 4. The start and stop position of the annotated expression in the format "start:stop"

The exact formatting performed below is specific to the format of the annotation files, and may vary if an annotation software other than BRAT is used.

```
tune_ann <- do.call(rbind, lapply(ann_filenames, function(fn){</pre>
  annotations <- read.delim(fn,
                             header = FALSE, sep = "\t", stringsAsFactors = FALSE,
                             col.names = c("id", "entity", "annotation"))
  # Label with file name
  annotations filename <- sub(".ann", ".txt", sub(".+/", "", fn), fixed=TRUE)
  # Separate entity information into entity label and start:stop position
  # Format is "entity start stop"
  ent_info <- strsplit(as.character(annotations$entity), split="\\s")</pre>
  annotations$entity <- unlist(lapply(ent_info, '[[', 1))</pre>
  annotations$pos <- paste(lapply(ent_info, '[[', 2),</pre>
                            lapply(ent_info, '[[', 3), sep=":")
  annotations <- annotations[,c("filename", "entity", "annotation", "pos")]
  return(annotations)
}))
head(tune ann)
```

```
##
          filename
                      entity annotation
                                                pos
## 1 tune note1.txt DrugName
                                  Prograf 1219:1226
## 2 tune_note1.txt Strength
                                     1 mg 1227:1231
## 3 tune_note1.txt
                     DoseAmt
                                        3 1236:1237
## 4 tune note1.txt Frequency twice a day 1256:1267
## 5 tune note1.txt DrugName
                                 porgraf 3912:3919
                                      3mg 3920:3923
## 6 tune_note1.txt
                         Dose
```

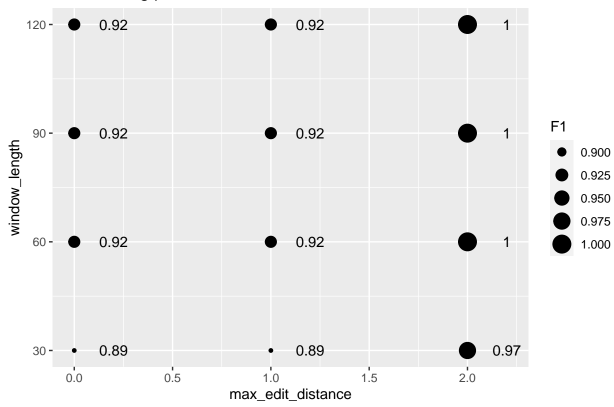
To select appropriate tuning parameters, we identify a range of possible values for each of the windowlength and max\_edit\_dist parameters. Here, we allow windowlength to vary from 30 to 120 characters in increments of 30, and max\_edit\_dist to take a value of 0, 1, or 2. We then obtain the extractMed results for each combination. We set progress=FALSE to avoid printing the progress message for each combination of arguments.

```
max_edit_dist = tune_pick$max_edit_distance[i],
progress = FALSE)
}
```

Finally, we determine which parameter combination yielded the highest performance, quantified by some metric. For our purpose, we used the F1-measure (F1), the harmonic mean of precision  $\left(\frac{\text{true positives}}{\text{true positives} + \text{false positives}}\right)$  and recall  $\left(\frac{\text{true positives}}{\text{true positives} + \text{false negatives}}\right)$ . Tuning parameters were selected based on which combination maximized F1 performance within the tuning set. The code below determines true positives as well as false positives and negatives, used to compute precision, recall, and F1.

```
# Functions to compute true positive, false positive, and false negatives
# number of true positives - how many annotations were correctly identified by extractMed
Tpos <- function(df){</pre>
  sum(df$annotation == df$expr, na.rm=TRUE)
# number of false positive (identified by extractMed but not annotated)
Fpos <- function(df){</pre>
  sum(is.na(df$annotation))
}
# number of false negatives (annotated but not identified by extractMed)
Fneg <- function(df){</pre>
  # keep only rows with annotation
  df_ann <- subset(df, !is.na(annotation))</pre>
  sum(is.na(df$expr))
}
prf <- function(df){</pre>
  tp <- Tpos(df)</pre>
  fp <- Fpos(df)</pre>
  fn <- Fneg(df)
  precision <- tp/(tp + fp)</pre>
  recall <- tp/(tp + fn)
  f1 <- (2*precision*recall)/(precision + recall)</pre>
  return(f1)
}
tune_pick$F1 <- sapply(mxr_tune, function(x){</pre>
  compare <- merge(x, tune_ann,</pre>
                    by = c("filename", "entity", "pos"), all = TRUE)
  prf(compare)
})
ggplot(tune pick) + geom point(aes(max edit distance, window length, size = F1)) +
  scale_y_continuous(breaks=seq(30,120,30)) +
  annotate("text", x = tune_pick$max_edit_distance+.2, y = tune_pick$window_length,
           label = round(tune_pick$F1, 2)) +
  ggtitle("F1 for tuning parameter values")
```

# F1 for tuning parameter values



The plot shows that the highest F1 achieved was 1, and occurred for three different combinations of parameter values: a maximum edit distance of 2 and a window length of 60, 90, or 120 characters. The relatively small number of unique F1 values is likely the result of only using 3 tuning notes. In this case, we would typically err on the side of allowing a larger search window and decide to use a maximum edit distance of 2 and a window length of 120 characters. In a real-world tuning scenario and with a larger tuning set, we would also want to test longer window lengths since the best case scenario occurred at the longest window length we used.

## Pro-Med-NLP

The *Pro-Med-NLP* module uses a dose data building algorithm to generate longitudinal medication dose data from the raw output of an NLP system.<sup>5</sup> The algorithm is divided into two parts. Part I parses the raw output and pairs entities together, and Part II calculates dose intake and daily dose and removes redundant information.

#### Part I

Two main functions are used in this part of the algorithm, a parse function (parseMedExtractR, parseMedXN, parseCLAMP, or parseMedEx) and buildDose.

Parse functions (parseMedExtractR, parseMedXN, parseCLAMP, parseMedEx) Parse functions are available for the medExtractR, MedXN, CLAMP, and MedEx systems (parseMedExtractR, parseMedXN, parseCLAMP, parseMedEx). If the user has output from another NLP system, the user can write code to standardize the output before calling the buildDose function.

The parse functions require the following argument:

• filename: file name for a file containing the raw output from the NLP system

The following are the function calls for the parse functions.

```
parseMedExtractR(filename)
parseMedXN(filename, begText = "^[ID0-9]+_[0-9-]+_[Note0-9]")
parseCLAMP(filename)
parseMedEx(filename)
```

The parseMedXN function also has an argument specifying a regular expression pattern which indicates the start of each row (i.e., drug mention) in the raw MedXN output. In the example above, the begText argument identifies how the file names are structured for a set of clinical notes in a Vanderbilt University dataset (e.g., "ID111\_2000-01-01\_Note001" where ID111 is the patient ID, 2000-01-01 is the date of the note, and Note001 is the note ID). Thus, each row beginning with a file name in this format indicates a new drug mention.

The parse functions output a standardized form of the data that includes a row for each drug mention and columns for all entities anchored to that drug mention.

Running parseMedExtractR Below we demonstrate how to run parseMedExtractR using the example medExtractR output from above. The file names here correspond to the files generated from the final output of the Extract-Med module.

```
tac_mxr_fn <- system.file("examples", "tac_mxr.csv", package = "EHR")
lam_mxr_fn <- system.file("examples", "lam_mxr.csv", package = "EHR")
lam_mxn_fn <- system.file("examples", "lam_medxn.csv", package = "EHR")
tac_mxr_parsed <- parseMedExtractR(tac_mxr_fn)
lam_mxr_parsed <- parseMedExtractR(lam_mxr_fn)</pre>
```

Running parseMedXN Below is an example using the parseMedXN function.

```
## MedXN output:
## ID1_2012-11-22_Note1.txt|lamotrigine::255::266|28439::196502|100 mg::278::284|
## 2::288::289`1.5::310::313|tabs::290::294`tabs::314::318`tabs::350::354||
## morning::298::305`evening::322::329|2 weeks::334::341
## ID1_2012-11-22_Note1.txt|Vimpat::1086::1092|809979|200mg::1093::1098||tab::1099::1102|
## mouth::1106::1111|twice daily::1112::1123
## ID1_2012-11-22_Note1.txt|lamotrigine::1172::1183|28439|100 mg::1184::1190||
## tablet::1191::1197
## ID1_2012-11-22_Note1.txt|Lamictal::1213::1221|196502||1.5::1223::1226|
## tablets::1227::1234|mouth::1238::1243|twice a day::1244::1255
```

The output from all systems, once parsed, has the same structure as the example parsed MedXN output below.

```
lam_mxn_parsed <- parseMedXN(lam_mxn_fn, begText = "^[ID0-9]+_[0-9-]+_[Note0-9]")</pre>
```

```
##
                     filename
                                              drugname
                                                                  strength
## 1 ID1_2012-11-22_Note1.txt
                                 lamotrigine::255::266
                                                         100 mg::278::284
## 2 ID1_2012-11-22_Note1.txt
                                    Vimpat::1086::1092
                                                        200mg::1093::1098
## 3 ID1_2012-11-22_Note1.txt lamotrigine::1172::1183 100 mg::1184::1190
## 4 ID1_2012-11-22_Note1.txt
                                 Lamictal::1213::1221
##
##
                                dose
                                                 route
##
        1 2::288::289`1.5::310::313
```

```
2
##
                                       mouth::1106::1111
##
        3
                                                     <NA>
##
        4
                     1.5::1223::1226 mouth::1238::1243
##
##
                                            freq
                                                           duration
##
        1 morning::298::305`evening::322::329 2 weeks::334::341
##
                       twice daily::1112::1123
                                                                <NA>
        3
##
                                                                <NA>
##
        4
                       twice a day::1244::1255
                                                                <NA>
```

buildDose After the NLP output is parsed, the buildDose function is run to pair the parsed entities. The main buildDose function arguments are as follows:

- dat: standardized form of the data (i.e., output from one of the parse functions)
- dn: argument to specify drug names of interest (generally not used with medExtractR since medExtractR is a targeted system and only includes the drug of interest in the raw output)

The general function call is:

```
buildDose(dat, dn = NULL)
```

Other notable arguments that can be specified by buildDose include:

- preserve: columns which should be preserved in the final output, but not considered for pairing with other entities. The default is NULL, though for medExtractR the dose change entity is always preserved.
- dist\_method: which distance method calculation to use in cost calculation for pairing entities. For more details see the supplementary material in McNeer, et al. (2020). The default from getOption('ehr.dist\_method') is minEntEnd, or using the minimum distance between the end of one entity and beginning of the next).
- checkForRare: a logical argument, indicates whether or not the output should return rare occurrences of entities (default is FALSE)

The output of the buildDose function is a dataset with a column for each entity and a row for each pairing.

Running buildDose In our medExtractR example from above, the output of the buildDose function is the following:

```
(tac_part_i_out <- buildDose(tac_mxr_parsed))</pre>
```

```
##
                                        drugname strength dose route
                             filename
                                                                               freq
## 1 tacpid1_2008-06-26_note1_1.txt
                                                               3
                                         Prograf
                                                      1 mg
                                                                    NA twice a day
## 2 tacpid1_2008-06-26_note1_1.txt
                                         prograf
                                                      <NA> <NA>
                                                                    NA
                                                                                bid
## 3 tacpid1_2008-06-26_note2_1.txt
                                         Prograf
                                                      1 mg
                                                               3
                                                                    NA twice a day
## 4 tacpid1_2008-12-16_note3_1.txt Tacrolimus
                                                      <NA> <NA>
                                                                    NA
                                                                               <NA>
## 5 tacpid1_2008-12-16_note3_1.txt
                                         Prograf
                                                               3
                                                                    NA twice a day
                                                      1 mg
## 6 tacpid1_2008-12-16_note3_1.txt
                                         Prograf
                                                      <NA> <NA>
                                                                    NA
                                                                                bid
## 7 tacpid1_2008-12-16_note3_1.txt
                                         Prograf
                                                      <NA> <NA>
                                                                    NA
                                                                               <NA>
##
     dosestr dosechange lastdose drugname_start
## 1
        <NA>
                    <NA>
                              <NA>
                                               930
## 2
                                              2709
         3mg
                    <NA>
                            8:30pm
## 3
        <NA>
                    <NA>
                             14 hr
                                               618
## 4
        <NA>
                    <NA>
                              <NA>
                                               722
## 5
        <NA>
                    <NA>
                              <NA>
                                               761
## 6
         2mg
                              <NA>
                                              2179
                decrease
                    <NA> 10:30 pm
## 7
                                              2205
        <NA>
```

# (lam\_part\_i\_out <- buildDose(lam\_mxr\_parsed))</pre>

```
##
                                               drugname strength dose route
                              filename
## 1
      lampid1_2016-02-05_note4_1.txt
                                               Lamictal
                                                             <NA> <NA>
                                                                         <NA>
## 2
      lampid1_2016-02-05_note4_1.txt
                                            Lamotrigine
                                                            200mg
                                                                    1.5
                                                                         <NA>
## 3
      lampid1_2016-02-05_note4_1.txt Lamotrigine XR
                                                           100 mg
                                                                      3
                                                                         <NA>
## 4
      lampid1_2016-02-05_note4_1.txt Lamotrigine XR
                                                           100 mg
                                                                      2
                                                                         <NA>
## 5
      lampid1_2016-02-05_note5_1.txt
                                                    ltg
                                                           200 mg
                                                                    1.5
                                                                         <NA>
      lampid1_2016-02-05_note5_1.txt
                                                                      3
##
  6
                                                 ltg xr
                                                           100 mg
                                                                         <NA>
                                                                      2
##
      lampid1 2016-02-05 note5 1.txt
                                                 ltg xr
                                                           100 mg
                                                                         <NA>
##
  8
      lampid2_2008-07-20_note6_1.txt
                                            lamotrigine
                                                             <NA> <NA>
                                                                         <NA>
      lampid2 2008-07-20 note6 1.txt
                                               lamictal
                                                             <NA> <NA>
                                                                         <NA>
## 10 lampid2_2008-07-20_note6_1.txt
                                                             <NA> <NA>
                                               Lamictal
                                                                         <NA>
## 11 lampid2_2012-04-15_note7_1.txt
                                            lamotrigine
                                                                  <NA>
                                                                         <NA>
                                                           150 mg
  12 lampid2_2012-04-15_note7_1.txt
                                               Lamictal
                                                             <NA>
                                                                         <NA>
##
              freq dosestr dosechange
                                        lastdose drugname start
##
               BID
                    300 mg
                                   <NA>
                                             < NA >
                                                              810
  1
## 2
      twice daily
                       <NA>
                                   <NA>
                                             <NA>
                                                              847
## 3
                       <NA>
                                   <NA>
                                                              954
          morning
                                             <NA>
## 4
           evening
                       <NA>
                                   <NA>
                                             <NA>
                                                              954
## 5
             daily
                       <NA>
                                   <NA>
                                             <NA>
                                                              442
## 6
             in am
                       <NA>
                                   <NA>
                                             <NA>
                                                              465
## 7
             in pm
                       <NA>
                                   <NA>
                                             <NA>
                                                              465
## 8
              <NA>
                       <NA>
                                   <NA>
                                             <NA>
                                                             1267
## 9
              q12h
                    150 mg
                                   <NA>
                                             <NA>
                                                             1280
## 10
               BID
                                                             2273
                      200mg
                              Increase
                                             <NA>
## 11
              <NA>
                       <NA>
                                   <NA>
                                             <NA>
                                                              103
## 12 twice a day
                       <NA>
                                   <NA>
                                             <NA>
                                                              141
```

If the checkForRare argument is set to TRUE, any extracted expressions with a proportion of occurrence less than 0.2 are returned as rare values. When rare values are identified, a warning is printed to notify the user. The var column indicates the entity (note that dose in this output refers to dose amount, while dosestr would indicate dose given intake). This can be used as a quick check for potentially inaccurate information and allow the user to remove incorrect extractions before applying the Pro-Med-NLP module as incorrect extractions would reduce accuracy of the dose building data. Note that these values may still be correct extractions even though they are rare, as is the case for our output below.

```
lam_checkForRare <- buildDose(lam_mxr_parsed, checkForRare=TRUE)</pre>
```

#### Part II

In Part II of the algorithm, we form the final analysis datasets containing computed dosing information at the note and date level for each patient. This process requires more detailed meta data associated with each clinical note file, the format of which is described below. In this part, we also discuss how time of last dose should be incorporated when present before producing the final output dataset. noteMetaData The meta data argument is required by the functions processLastDose and collapseDose, and requires four columns: filename, pid, date, note. In our example data, pid (patient ID), date, and note can all be extracted from the filename. Take the filename "tacpid1\_2008-06-26\_note1\_1.txt" for example. It contains information in the form "[PID]\_[date]\_[note]", where PID = "tacpid1", date = "2008-06-26" and note = "note1". The function below can build our meta data from each of the filenames.

```
bmd <- function(x) {</pre>
  fns <- strsplit(x, '_')</pre>
  pid <- sapply(fns, `[`, 1)</pre>
  date <- as.Date(sapply(fns, \tilde{}[, 2), format = \frac{1}{2}Y-\frac{1}{2}m-\frac{1}{2}d')
  note <- sapply(fns, `[`, 3)</pre>
  data.frame(filename = x, pid, date, note, stringsAsFactors = FALSE)
}
bmd("tacpid1_2008-06-26_note1_1.txt")
                                           pid
                             filename
                                                      date note
## 1 tacpid1_2008-06-26_note1_1.txt tacpid1 2008-06-26 note1
(tac_metadata <- bmd(tac_part_i_out[['filename']]))</pre>
##
                             filename
                                           pid
                                                      date note
## 1 tacpid1_2008-06-26_note1_1.txt tacpid1 2008-06-26 note1
## 2 tacpid1_2008-06-26_note1_1.txt tacpid1 2008-06-26 note1
## 3 tacpid1_2008-06-26_note2_1.txt tacpid1 2008-06-26 note2
## 4 tacpid1_2008-12-16_note3_1.txt tacpid1 2008-12-16 note3
## 5 tacpid1_2008-12-16_note3_1.txt tacpid1 2008-12-16 note3
## 6 tacpid1_2008-12-16_note3_1.txt tacpid1 2008-12-16 note3
## 7 tacpid1_2008-12-16_note3_1.txt tacpid1 2008-12-16 note3
```

Handling lastdose In this section, we cover how incorporation of the last dose entity should be handled if it was extracted during the Extract-Med module. From the buildDose output above, we see the raw last dose time extractions for the tacrolimus dataset. Using the functions processLastDose and addLastDose, we convert the extracted times into a processed and standardized datetime variable, and add the processed times to the buildDose output.

The processLastDose function requires the following arguments:

- mxrData: raw output from the extractMed function
- noteMetaData: note meta data for each file name in mxrData
- labData: a data frame containing lab dates and times associated with the file names within mxrData. This must contain at a minimum the columns pid and date (in the same format as noteMetaData), as well as labtime, a POSIXct variable indicating the date and time of a laboratory drug measurement

Extracted last dose times can fall into two categories: a time expression (e.g., "10am", "22:00", "7 last night") or a duration expression (e.g. "14 hour" level), where the "time" of last dose indicates the number of hours since the last dose was taken relative to the time of the clinical visit. In the latter case, the lab time (from the labData argument) is needed in order to convert the extracted duration expression into a datetime variable. Below is an example lab dataset for our sample tacrolimus data.

```
data(tac_lab)
tac_lab

## pid date labtime
## 1 tacpid1 2008-06-26 2008-06-26 10:42:00
## 2 tacpid1 2008-12-16 2008-12-16 12:11:00
```

Within processLastDose, extracted times are converted to time expressions of the format "HH:MM:SS" and assigned a date based on the date of the corresponding note. When the last dose time is after 12pm, it is assumed to have been taken on the previous date.

```
(tac_ld <- processLastDose(mxrData = tac_mxr, noteMetaData = tac_metadata, labData = tac_lab))</pre>
```

```
filename
                                                lastdose
                                                            ld_pos
## 1 tacpid1_2008-06-26_note1_1.txt 2008-06-25 20:30:00 2740:2746 tacpid1
## 2 tacpid1_2008-06-26_note2_1.txt 2008-06-25 20:42:00
                                                           678:683 tacpid1
## 3 tacpid1_2008-12-16_note3_1.txt 2008-12-15 22:30:00 2231:2239 tacpid1
##
           date raw time ld start time type
                                                         labtime
                                        time 2008-06-26 10:42:00
## 1 2008-06-26
                  8:30pm
                             2740
## 2 2008-06-26
                   14 hr
                              678
                                    duration 2008-06-26 10:42:00
## 3 2008-12-16 10:30 pm
                             2231
                                        time 2008-12-16 12:11:00
```

The function output contains the processed and standardized last dose time (lastdose), the original extracted expression (raw\_time), whether the raw expression was a time or duration (time\_type), as well as position information for the last dose time (ld\_start) for appropriate pairing with dosing information in addLastDose. The labtime column in the output above corresponds to the information provided in the labData argument.

The addLastDose function requires the following arguments:

- buildData: output from buildDose
- lastdoseData: dataset containing last dose time information for the file names in buildData. This
  should include columns for filename and lastdose, with lastdose being a processed POSIXct datetime
  variable.

In the case where last dose information was extracted from clinical notes using medExtractR, the lastdoseData input should be output from the processLastDose function containing the last dose start positions, as demonstrated below. It is possible for multiple times to be extracted from a clinical note. For extracted times within a 2 hour window of one another, addLastDose treats these as equivalent and extracts the last dose time. Note that this may be context-dependent, and this rule was determined based on drugs administered every 12 hours and assuming a trough drug level. For time differences of more than two hours, the last dose start position is used to pair the extracted time with the closest drug mention. Alternatively, if the user has a separate dataset with validated last dose times, they can provide their own dataset. When providing a validated dataset, there should be only one last dose time per patient ID and date.

```
(tac_part_i_out_lastdose <- addLastDose(buildData = tac_part_i_out, lastdoseData = tac_ld))</pre>
```

```
##
                            filename
                                        drugname strength dose route
                                                      1 mg
## 1 tacpid1_2008-06-26_note1_1.txt
                                         Prograf
                                                               3
                                                                    NA twice a day
## 2 tacpid1_2008-06-26_note1_1.txt
                                         prograf
                                                      <NA> <NA>
                                                                    NA
                                                                                bid
## 3 tacpid1_2008-06-26_note2_1.txt
                                         Prograf
                                                               3
                                                                    NA twice a day
                                                      1 mg
## 4 tacpid1 2008-12-16 note3 1.txt Tacrolimus
                                                      <NA> <NA>
                                                                               <NA>
                                                                    NA
## 5 tacpid1_2008-12-16_note3_1.txt
                                         Prograf
                                                               3
                                                                    NA twice a day
                                                      1 mg
## 6 tacpid1 2008-12-16 note3 1.txt
                                         Prograf
                                                      <NA> <NA>
                                                                    NA
                                                                                bid
## 7 tacpid1_2008-12-16_note3_1.txt
                                         Prograf
                                                                    NΔ
                                                                               <NA>
                                                      <NA> <NA>
##
     dosestr dosechange
                                     lastdose drugname start
## 1
        <NA>
                    <NA>
                                          <NA>
                                                           930
## 2
                    <NA> 2008-06-25 20:30:00
                                                          2709
         3mg
## 3
                    <NA> 2008-06-25 20:42:00
        <NA>
                                                           618
## 4
        <NA>
                    <NA>
                                          <NA>
                                                           722
## 5
        <NA>
                    <NA>
                                          <NA>
                                                           761
## 6
         2mg
                decrease
                                          <NA>
                                                          2179
## 7
                    <NA> 2008-12-15 22:30:00
                                                          2205
        <NA>
```

Note that in the lastdose columns, we now have standardized datetime objects instead of the raw extracted expressions.

collapseDose The main function used in Part II of the algorithm is the collapseDose function, which relies on the underlying makeDose function. collapseDose allows the user to split the data using drug names given by regular expressions (...) and run makeDose separately on each subset of the data. For example, if the data includes multiple drugs, regular expressions can be specified for each drug. Another use of this function is to split the data by different formulations of the drug, such as separating immediate release formulations from extended release formulations, which are often written using "XR" or "ER" in the drug name.

The collapseDose function requires the following arguments:

- x: output from the buildDose function, or from addLastDose if last dose information is incorporated
- noteMetaData: a data.frame with columns for filename, pid (patient id), date, and note
- naFreq: method to use when assigning missing frequencies; the default is to assign the most common frequency

The general function call is:

```
collapseDose(x, noteMetaData, naFreq = 'most', ...)
```

The main function underlying collapseDose is makeDose, which standardizes entities, imputes missing values, calculates dose intake and daily dose, removes redundancies, and generates the final dose data. Two data.frames are generated from the makeDose function, one with redundancies removed at the note level and one at the date level (see McNeer et al. (2020) for details). The collapseDose function serves as a wrapper to makeDose to ensure results are collated and formatted properly for analysis.

Running collapseDose For our tacrolimus example above, the output of this function is below. Note that we use the output from addLastDose rather than directly from buildDose.

```
tac_part_ii <- collapseDose(tac_part_i_out_lastdose, tac_metadata, naFreq = 'most')</pre>
```

Note level collapsing:

```
tac_part_ii$note
```

```
##
                            filename drugname strength dose
                                                              route freq dosestr
## 1 tacpid1_2008-06-26_note1_1.txt
                                       Prograf
                                                    1 mg
                                                            3 orally
                                                                       bid
                                                                               <NA>
## 2 tacpid1_2008-06-26_note2_1.txt
                                                                               <NA>
                                       Prograf
                                                    1 mg
                                                            3 orally
                                                                       bid
## 3 tacpid1_2008-12-16_note3_1.txt
                                       Prograf
                                                            3 orally
                                                                               <NA>
                                                    1 mg
                                                                       bid
## 4 tacpid1_2008-12-16_note3_1.txt
                                       Prograf
                                                    <NA> <NA> orally
                                                                       bid
                                                                                2mg
##
     dosechange
                            lastdose drugname_start dosestr.num strength.num
## 1
           <NA> 2008-06-25 20:30:00
                                                  930
                                                                NA
                                                                               1
## 2
           <NA> 2008-06-25 20:42:00
                                                                NA
                                                  618
                                                                               1
## 3
           <NA> 2008-12-15 22:30:00
                                                  761
                                                                NA
                                                                               1
                                                                 2
## 4
       decrease 2008-12-15 22:30:00
                                                 2179
                                                                             NA
##
     doseamt.num freq.num dose.intake intaketime dose.seq dose.daily
## 1
                         2
                                      3
                                               <NA>
                                                          NA
## 2
               3
                         2
                                      3
                                               <NA>
                                                          NA
                                                                       6
## 3
               3
                         2
                                      3
                                               <NA>
                                                                       6
                                                          NA
                                      2
## 4
                         2
                                               <NA>
                                                                       4
              NΑ
                                                          NA
```

Date level collapsing:

```
tac_part_ii$date
```

```
##
                           filename drugname strength dose
                                                            route freq dosestr
## 1 tacpid1_2008-06-26_note1_1.txt
                                     Prograf
                                                  1 mg
                                                          3 orally
                                                                    bid
                                                                            <NA>
## 2 tacpid1_2008-06-26_note2_1.txt
                                     Prograf
                                                  1 mg
                                                           3 orally
                                                                     bid
                                                                            <NA>
## 3 tacpid1_2008-12-16_note3_1.txt
                                     Prograf
                                                          3 orally
                                                                            <NA>
                                                  1 mg
                                                                    bid
```

```
## 4 tacpid1_2008-12-16_note3_1.txt Prograf
                                                    <NA> <NA> orally bid
     dosechange
##
                             lastdose drugname_start dosestr.num strength.num
           <NA> 2008-06-25 20:30:00
## 1
                                                  930
                                                                NA
## 2
            <NA> 2008-06-25 20:42:00
                                                  618
                                                                NA
                                                                               1
## 3
            <NA> 2008-12-15 22:30:00
                                                  761
                                                                NA
                                                                               1
##
       decrease 2008-12-15 22:30:00
                                                 2179
                                                                 2
                                                                              NA
     doseamt.num freq.num dose.intake intaketime dose.seq dose.daily
##
## 1
                3
                         2
                                      3
                                               <NA>
                                                           NA
## 2
                3
                         2
                                      3
                                               <NA>
                                                           NA
                                                                        6
## 3
                3
                         2
                                      3
                                               <NA>
                                                           NA
                                                                        6
## 4
               NA
                         2
                                      2
                                               <NA>
                                                           NA
                                                                        4
```

Below, we demonstrate collapseDose using our lamotrigine example. In the function call, we supply an additional argument 'xr|er' to indicate that we want to separately consider extended release formulations of lamotrigine, (usually denoted by "XR" or "ER"). This prevents regular lamotrigine mentions from being collapsed with lamotrigine XR mentions, even if the dosage is identical.

```
data(lam_metadata)
lam_part_ii <- collapseDose(lam_part_i_out, lam_metadata, naFreq = 'most', 'xr|er')</pre>
```

Note level collapsing:

```
lam_part_ii$note
```

```
##
                             filename
                                             drugname strength dose route
                                                            <NA> <NA> orally
## 1 lampid1 2016-02-05 note4 1.txt
                                             Lamictal
## 2 lampid1 2016-02-05 note4 1.txt Lamotrigine XR
                                                         100 mg
                                                                    3 orally
                                                                                 am
## 3 lampid1_2016-02-05_note4_1.txt Lamotrigine XR
                                                         100 mg
                                                                    2 orally
                                                                                 pm
## 4 lampid1_2016-02-05_note5_1.txt
                                                   ltg
                                                         200 mg
                                                                  1.5 orally daily
## 5 lampid1_2016-02-05_note5_1.txt
                                               ltg xr
                                                         100 mg
                                                                    3 orally
                                                                                 am
## 6 lampid1 2016-02-05 note5 1.txt
                                               ltg xr
                                                         100 mg
                                                                    2 orally
                                                                                 рm
## 7 lampid2 2008-07-20 note6 1.txt
                                                            <NA> <NA> orally
                                             lamictal
                                                                                bid
## 8 lampid2 2008-07-20 note6 1.txt
                                             Lamictal
                                                            <NA>
                                                                 <NA> orally
                                                                                bid
## 9 lampid2_2012-04-15_note7_1.txt
                                             Lamictal
                                                            <NA>
                                                                    1 orally
                                                                                bid
##
     dosestr dosechange lastdose drugname_start dosestr.num strength.num
      300 mg
## 1
                    <NA>
                              <NA>
                                                             300
                                               810
                                                                            NA
## 2
        <NA>
                    <NA>
                              <NA>
                                               954
                                                              NA
                                                                           100
## 3
        <NA>
                    <NA>
                              <NA>
                                               954
                                                              NA
                                                                           100
## 4
        <NA>
                    <NA>
                              <NA>
                                               442
                                                              NA
                                                                           200
## 5
        <NA>
                                                                           100
                    <NA>
                              <NA>
                                               465
                                                              NA
        <NA>
                                                                           100
## 6
                    <NA>
                              <NA>
                                               465
                                                              NA
      150 mg
## 7
                    <NA>
                              <NA>
                                               1280
                                                             150
                                                                            NA
       200mg
                              <NA>
                                               2273
                                                             200
## 8
                Increase
                                                                            NA
## 9
        <NA>
                    <NA>
                              <NA>
                                               141
                                                              NA
                                                                           150
##
     doseamt.num freq.num dose.intake intaketime dose.seq dose.daily
## 1
               NA
                          2
                                                            NA
                                     300
                                               <NA>
                                                                      600
## 2
              3.0
                          1
                                     300
                                                             1
                                                                      500
                                                  am
## 3
              2.0
                                                            2
                                                                      500
                          1
                                     200
                                                  pm
## 4
              1.5
                          1
                                     300
                                               <NA>
                                                            NA
                                                                      300
## 5
              3.0
                          1
                                     300
                                                  am
                                                             1
                                                                      500
## 6
              2.0
                          1
                                     200
                                                            2
                                                                      500
                                                  pm
                          2
                                                                      300
## 7
               NA
                                     150
                                                <NA>
                                                            NA
## 8
                          2
                                     200
                                                                      400
               NA
                                               <NA>
                                                            NA
## 9
              1.0
                          2
                                     150
                                               <NA>
                                                            NA
                                                                      300
```

Date level collapsing:

```
lam_part_ii$date
                                             drugname strength dose route
                             filename
                                                                               freq
## 1 lampid1 2016-02-05 note4 1.txt
                                                           <NA> <NA> orally
                                             Lamictal
                                                                                bid
## 2 lampid1 2016-02-05 note4 1.txt Lamotrigine XR
                                                         100 mg
                                                                    3 orally
                                                                                 am
## 3 lampid1_2016-02-05_note4_1.txt Lamotrigine XR
                                                         100 mg
                                                                    2 orally
                                                                                 pm
## 4 lampid1_2016-02-05_note5_1.txt
                                                         200 mg
                                                                 1.5 orally daily
                                                   ltg
## 5 lampid2_2008-07-20_note6_1.txt
                                             lamictal
                                                           <NA> <NA> orally
                                                                                bid
## 6 lampid2 2008-07-20 note6 1.txt
                                                           <NA> <NA> orally
                                             Lamictal
                                                                                bid
## 7 lampid2 2012-04-15 note7 1.txt
                                             Lamictal
                                                           <NA>
                                                                    1 orally
                                                                                bid
##
     dosestr dosechange lastdose drugname_start dosestr.num strength.num
## 1
      300 mg
                    <NA>
                              <NA>
                                               810
                                                            300
                                                                            NA
## 2
        <NA>
                    <NA>
                              <NA>
                                               954
                                                                          100
                                                             NΑ
## 3
        <NA>
                    <NA>
                              <NA>
                                               954
                                                             NA
                                                                          100
                                                                          200
## 4
        <NA>
                    <NA>
                                               442
                                                             NA
                              <NA>
## 5
      150 mg
                    <NA>
                              <NA>
                                              1280
                                                             150
                                                                            NA
## 6
       200mg
                              <NA>
                                              2273
                                                            200
                                                                           NA
                Increase
##
        <NA>
                    <NA>
                              <NA>
                                               141
                                                             NA
                                                                          150
##
     doseamt.num freq.num dose.intake intaketime dose.seq dose.daily
                                    300
## 1
               NA
                          2
                                               <NA>
                                                           NA
## 2
              3.0
                                                                      500
                          1
                                    300
                                                  am
                                                            1
## 3
              2.0
                          1
                                    200
                                                            2
                                                                      500
                                                 pm
## 4
              1.5
                          1
                                    300
                                                                      300
                                               <NA>
                                                           NA
## 5
              NA
                          2
                                    150
                                               <NA>
                                                           NA
                                                                      300
                          2
## 6
               NA
                                    200
                                               <NA>
                                                           NA
                                                                      400
## 7
              1.0
                          2
                                    150
                                               <NA>
                                                           NA
                                                                      300
```

**Additional collapsing** Collapsing by date or note produces observations at the daily intake level. It is possible to further collapse data to the daily level, though you may want to drop the *dose.intake* variable as it would potentially lose meaning.

```
x <- lam_part_ii[['note']]</pre>
# retrieve metadata for each filename
k1 <- lam_metadata[match(x[,'filename'], lam_metadata[,'filename']), c('pid','date','note')]
# select additional key data
k2 <- cbind(k1, x[,c('dose.daily','drugname_start')])</pre>
# turn keys into character string
chk <- do.call(paste, c(k2, sep = '|'))
# keep first instance of each chk key
lam_part_iii_note <- x[!duplicated(chk),]</pre>
lam_part_iii_note[,c('filename','drugname','drugname_start','dose.daily')]
##
                                            drugname drugname_start dose.daily
## 1 lampid1_2016-02-05_note4_1.txt
                                                                             600
                                            Lamictal
                                                                 810
## 2 lampid1_2016-02-05_note4_1.txt Lamotrigine XR
                                                                 954
                                                                             500
## 4 lampid1_2016-02-05_note5_1.txt
                                                                 442
                                                                             300
                                                 ltg
## 5 lampid1_2016-02-05_note5_1.txt
                                              ltg xr
                                                                 465
                                                                             500
## 7 lampid2_2008-07-20_note6_1.txt
                                                                             300
                                                                1280
                                            lamictal
## 8 lampid2_2008-07-20_note6_1.txt
                                            Lamictal
                                                                2273
                                                                             400
## 9 lampid2_2012-04-15_note7_1.txt
                                                                             300
                                            Lamictal
                                                                 141
x <- lam_part_ii[['date']]</pre>
# ignore note for date level collapsing
k1 <- lam_metadata[match(x[,'filename'], lam_metadata[,'filename']), c('pid','date')]
```

```
k2 <- cbind(k1, x[,c('dose.daily','drugname_start')])
chk <- do.call(paste, c(k2, sep = '|'))
lam_part_iii_date <- x[!duplicated(chk),]
lam_part_iii_date[,c('filename','drugname','drugname_start','dose.daily')]</pre>
```

```
##
                            filename
                                            drugname drugname_start dose.daily
## 1 lampid1_2016-02-05_note4_1.txt
                                           Lamictal
                                                                 810
                                                                            600
## 2 lampid1_2016-02-05_note4_1.txt Lamotrigine XR
                                                                 954
                                                                            500
## 4 lampid1 2016-02-05 note5 1.txt
                                                                 442
                                                                            300
                                                 ltg
## 5 lampid2 2008-07-20 note6 1.txt
                                            lamictal
                                                                1280
                                                                            300
## 6 lampid2_2008-07-20_note6_1.txt
                                           Lamictal
                                                                2273
                                                                            400
## 7 lampid2 2012-04-15 note7 1.txt
                                           Lamictal
                                                                 141
                                                                            300
```

## References

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