# Making Individual Plot-Year Maps From Long-Term Data

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# 1 Introduction

I need to make individual plot maps as they were in Year X (each of their measurement years, up to 7), which accurately represent the number of trees currently present in the plot, alongside their DBH measures if necessary. We have a few different types of trees that are present in the dataset.

- 1. Trees that were present before plot establishment as seedlings
- 2. Trees that were present before plot establishment as adults
- 3. Trees that were present before plot establishment as seedlings that died
- 4. Trees that were present before plot establishment as adults that died
- 5. Trees that appeared after plot establishment as seedlings that died
- 6. Trees that appeared after plot establishment as seedlings that survived
- 7. Seedlings that established and then died before DBH measure could be taken
- 8. Trees (saplings) that established then got too small for DBH before death

So, I think I need to do a loop for each of the 13,000 rows in ssdAllTrees. I should split them into groups, so that I know all NA's in DBH are expected because the tree is too small to measure, instead of it being measurements not taken yet. Also, let's switch the order of columns so that the xy columns are NOT last. That way, we have a definite number of columns before the dbh columns start.

```
> ## load library
> library(disperseR)
> ##subsetting by years sampled
> sevenyears <- ssdAllTrees[ssdAllTrees$plot=="bellow" |
+ ssdAllTrees$plot=="palate" |</pre>
```

```
ssdAllTrees$plot=="reclusive" |
                               ssdAllTrees$plot=="octane" |
                               ssdAllTrees$plot=="artist",]
 sixyears <- ssdAllTrees[ssdAllTrees$plot=="trigger" |</pre>
>
                               ssdAllTrees$plot=="rigid",]
 fiveyears <- ssdAllTrees[ssdAllTrees$plot=="crackers" |</pre>
>
                               ssdAllTrees$plot=="trinity" |
                               ssdAllTrees$plot=="realtor" |
                               ssdAllTrees$plot=="sodium" |
                               ssdAllTrees$plot=="distress" |
                               ssdAllTrees$plot=="gravy" |
                               ssdAllTrees$plot=="chestnut",]
> ## check to make sure all trees are accounted for
> (nrow(sevenyears) + nrow(sixyears) + nrow(fiveyears) == nrow(ssdAllTrees))
[1] TRUE
> ## delete the extraneous columns from sixyears and fiveyears
> fiveyears <- fiveyears[,-c(15,16)]</pre>
> sixyears <- sixyears[,-16]</pre>
```

Okay, so I now have three dataframes with DBH values that can be acted upon for each year. Let's get to writing this function. It should take a vector, which is a row of the dataframe. We know the identities of each of the first 9 columns for this vector, and the rest are to be dealt with in a loop. This function should output a data.frame that can be bound to a response data.frame. This response data.frame should be organized like so: Plot Name, Subplot, Tag, Species, Measure Year, Ingrowth Year, First Record Year, Death Year, x, y, DBH, Class

We need the plot, to look up sampling years. The other early columns just need to tag along, we don't need any information from them. If the ingrowth year is NA, that means that the tree was present in the plot before it was established. Now, in these plots, trees are checked yearly for establishment and death, but only measured every few years. What we know from ingrowth year, then, is the year that a seedling germinated and established. If ingrowth year is NOT NA, then we know the tree's entire life history, from the moment it was "born", and we can build the rows based on that.

If ingrowth year is NA, it gets a bit trickier. To deal with this, we can do a check. If there's no ingrowthyear, that means that the tree was alive before plot establishment. Therefore, we can make an assumption that if its first dbh value is NA, then it was too small to measure, e.g., a seedling. Otherwise, it's already a tree.

Wrapping all of this into a giant for loop, we can expand our ssdAllTrees data.frame to a df better suited for dispersal analysis

```
> ## get values where ingrowthyear has a value, for testing
> samplerows <- head(ssdAllTrees[!is.na(ssdAllTrees$ingrowthyear),])</pre>
```

```
> ## add in rows that have no ingrowth year
> samplerows <- rbind(samplerows,
                      head(ssdAllTrees[is.na(ssdAllTrees$ingrowthyear),]))
> ##add in rows that have no ingrowthyear but DO have a mortality year
> samplerows <- rbind(samplerows,
                      head(ssdAllTrees[is.na(ssdAllTrees$ingrowthyear) &
                                          !is.na(ssdAllTrees$mortalityyear),]))
> ## make the function
> processTreeRows <- function(samplerows, ssdPlotDesc, ssdAllTrees){
+ i <- 1
+ j <- 1
      finalrows <- data.frame(plot="dummy",
                               subplot="dummy",
                               treeid="dummy",
                               species="dummy"
                               ingrowth="dummy",
                               firstrec="dummy",
                               deathyear="dummy",
                               x="dummy",
                               v="dummy",
                               measyear="dummy",
                               dbh="dummy",
                               stage="dummy",
                               stringsAsFactors=FALSE
+ for(i in 1:nrow(samplerows)){
    ##get plot measurement years
      plotYears <- (ssdPlotDesc[ssdPlotDesc$plot==samplerows[i, "plot"], 5:11])</pre>
    ## make a dataframe of dbh1-7 and year values to search later
     rPlotYears <- as.data.frame(cbind(paste("dbh", 1:7, sep=""),
                                         as.numeric(plotYears[1,])),
                                   stringsAsFactors=FALSE,
                                   na.string="NA")
    ## clean it up a bit
      colnames(rPlotYears) <- c("colname", "year")</pre>
      ##get only years where there are acutally values
        years <- plotYears[!is.na(plotYears)]</pre>
    ## if there is an ingrowth year in this particular row...
    if(!is.na(samplerows[i,"ingrowthyear"])){
      ## remove any years that are less than that.
        years <- years[years >= samplerows[i, "ingrowthyear"]]
      ## remember that this row is a seedling at least once
        class <- "seedling"
```

```
} else{
  ## if it was NA, mark it as seedling
    if(is.na(samplerows[i, "dbh1"])){
      class <- "seedling"</pre>
    } else{
      class <- "tree"
    } ## end else
} ##end else
## is there a mortalityyear
if(!is.na(samplerows[i, "mortalityyear"])){
  ## remove any years greater than mortality year
    years <- years[years < samplerows[i, "mortalityyear"]]</pre>
  ## remember that this tree died
  ## this is currently unused, but we may need it later
    dead <- TRUE
    ##if a seedling died before it could be officially measured
  if(length(years)==0){
        addedrow <- NULL
        addedrow <- data.frame(plot=samplerows[i, "plot"],</pre>
                        subplot=samplerows[i, "subplot"],
                        treeid=samplerows[i, "treeid"],
                        species=samplerows[i, "sppcode"],
                        ingrowth=samplerows[i, "ingrowthyear"],
                       firstrec=samplerows[i, "yearfirstrecorded"],
deathyear=samplerows[i, "mortalityyear"],
                        x=samplerows[i, "xcoord"],
                        y=samplerows[i, "ycoord"],
                       measyear=samplerows[i, "yearfirstrecorded"],
                        dbh=NA,
                       stage="seedling",
                        stringsAsFactors=FALSE
        finalrows <- rbind(finalrows, addedrow)</pre>
    ## so, for each year that we have measurements...
    for(j in 1:length(years)){
      ##find which dbh column to use
        colname <- rPlotYears[rPlotYears$year==years[i],"colname"]</pre>
      ## clean it up because R sucks
        colname <- colname[!is.na(colname)]</pre>
      ## select that dbh value
```

```
## make sure we don't label a tree as a seedling more than once
            if(j > 1 & class=="seedling"){
              class <- "tree"
            }
          ## build row for new table.
          addedrow <- NULL
            addedrow <- data.frame(plot=samplerows[i, "plot"],</pre>
                           subplot=samplerows[i, "subplot"],
                           treeid=samplerows[i, "treeid"],
                           species=samplerows[i, "sppcode"],
                           ingrowth=samplerows[i, "ingrowthyear"],
                           firstrec=samplerows[i, "yearfirstrecorded"],
                           deathyear=samplerows[i, "mortalityyear"],
                           x=samplerows[i, "xcoord"],
                           y=samplerows[i, "ycoord"],
                          measyear=years[j],
                           dbh=dbh,
                           stage=class,
                           stringsAsFactors=FALSE
            #print(addedrow)
            ## do the rbind
            finalrows <- rbind(finalrows, addedrow)</pre>
        } ## end j loop
      }##end seedling loop
+ }## end i loop
+ return(finalrows[-1,])
+ }## end function
> ## use the function
> finalrows <- processTreeRows(samplerows, ssdPlotDesc, ssdAllTrees)
> head(finalrows)
      plot subplot treeid species ingrowth firstrec deathyear
2 crackers
                             PILA
                 1
                       71
                                       2004
                                                2004
                                                          <NA> 89893.515
                             PILA
                                                2004
3 crackers
                 1
                       71
                                       2004
                                                          <NA> 89893.515
                       72
                                                2000
4 crackers
                 1
                             PILA
                                       2000
                                                          <NA> 89893.072
5 crackers
                 1
                       72
                             PILA
                                       2000
                                                2000
                                                          <NA> 89893.072
6 crackers
                 1
                       72
                             PILA
                                       2000
                                                2000
                                                          <NA> 89893.072
7 crackers
                       73
                             PILA
                                       1997
                                                1997
                                                          <NA> 89900.691
                 1
                 y measyear dbh
                                    stage
2 48118.8599999999
                       2007 0.8 seedling
3 48118.8599999999
                       2012 1.3
                                     tree
4 48146.4199999999
                       2002 0.1 seedling
5 48146.4199999999
                       2007 1.8
                                    tree
6 48146.4199999999
                       2012 2.7
                                     tree
```

dbh <- as.numeric(samplerows[i, colname])</pre>

```
7 48127.4100000001 1997 0.9 seedling
```

### > tail(finalrows)

```
plot subplot treeid species ingrowth firstrec deathyear
55 crackers
                        26
                              CADE
                                        <NA>
                                                 1992
                                                           2009 89901.767
56 crackers
                        26
                              CADE
                                        <NA>
                                                 1992
                                                           2009 89901.767
                              CADE
                                        <NA>
                                                 1992
57 crackers
                        26
                                                           2009 89901.767
                  1
                              CADE
                                                 1992
                                                           2009 89901.767
58 crackers
                  1
                        26
                                        < NA >
59 crackers
                        27
                              CADE
                                        <NA>
                                                 1992
                                                           1993 89894.712
                  1
60 crackers
                  1
                        34
                              CADE
                                        <NA>
                                                 1992
                                                           1993 89888.256
                  y measyear
                              dbh stage
55 48135.6699999999
                        1992 8.1 tree
56 48135.6699999999
                        1997 8.2 tree
57 48135.6699999999
                        2002 8.3 tree
58 48135.6699999999
                        2007
                              8.2
                                   tree
59 48120.569999998
                        1992 10.6 tree
60 48139.319999998
                        1992 6.1 tree
```

#### > nrow(finalrows)

#### [1] 59

Ok, so the tests seem to work, in that the stage value seems to be correct in all cases; that we don't have records for trees before they were alive; and that we don't have records for trees after they were dead. Let's go ahead and do it to a larger dataframe.

> palate <- processTreeRows(ssdAllTrees[ssdAllTrees\$plot=="palate",], ssdPlotDesc, ssdAllTrees
> str(palate)

```
'data.frame':
                    277 obs. of 12 variables:
                  "palate" "palate" "palate" ...
$ plot
           : chr
                  "5" "5" "5" "5" ...
$ subplot : chr
                  "7342" "7342" "7343" ...
$ treeid
           : chr
$ species : chr
                  "PIMO" "PIMO" "PIMO" ...
$ ingrowth : chr
                  NA NA NA NA ...
                  "1996" "1996" "1996" "1993" ...
$ firstrec : chr
                  "1998" "1998" "1998" "1998" ...
$ deathyear: chr
                  "100211.022" "100211.022" "100211.022" "100211.09" ...
$ x
           : chr
                  "51357.0600000001" "51357.0600000001" "51357.0600000001" "51356.950000000
$ у
           : chr
$ measyear : chr
                  "1983" "1988" "1993" "1983" ...
$ dbh
           : chr
                  NA NA NA NA ...
                  "seedling" "tree" "tree" "seedling" ...
$ stage
           : chr
```

# [1] 277

> nrow(palate)

# > head(palate)

```
plot subplot treeid species ingrowth firstrec deathyear
                                                                          х
                                                                                            У
2 palate
                    7342
                            PIMO
                                       < NA >
                                                1996
                                                           1998 100211.022 51357.0600000001
3 palate
                5
                    7342
                            PIMO
                                       <NA>
                                                1996
                                                           1998 100211.022 51357.0600000001
4 palate
                5
                            PIMO
                                                           1998 100211.022 51357.0600000001
                    7342
                                       <NA>
                                                1996
                5
5 palate
                    7343
                            PIMO
                                       <NA>
                                                1993
                                                           1998
                                                                 100211.09 51356.9500000002
6 palate
                5
                    7343
                             PIMO
                                       <NA>
                                                1993
                                                                 100211.09 51356.9500000002
7 palate
                5
                    7343
                             PIMO
                                       <NA>
                                                           1998 100211.09 51356.9500000002
                                                1993
  measyear
            dbh
                    stage
      1983 <NA> seedling
2
3
      1988 <NA>
                     tree
4
      1993 <NA>
                     tree
5
      1983 <NA> seedling
6
      1988 <NA>
                     tree
7
      1993 17.3
                     tree
```

## > tail(palate)

plot	subplot	treeid	species	ingrowth	firstrec	deathyear	х
273 palate	16	7398	PIMO	<na></na>	1998	<na></na>	100201.144
274 palate	16	7398	PIMO	<na></na>	1998	<na></na>	100201.144
275 palate	16	7398	PIMO	<na></na>	1998	<na></na>	100201.144
276 palate	16	7398	PIMO	<na></na>	1998	<na></na>	100201.144
277 palate	16	7398	PIMO	<na></na>	1998	<na></na>	100201.144
278 palate	16	7398	PIMO	<na></na>	1998	<na></na>	100201.144
		y measy	ear dbl	n stage			
273 51445.8	350000000	1 1	1988 <na< td=""><td>&gt; tree</td><td></td><td></td><td></td></na<>	> tree			
274 51445.8	350000000	1 1	1993 <na< td=""><td>&gt; tree</td><td></td><td></td><td></td></na<>	> tree			
275 51445.8	350000000	1 1	1998	1 tree			
276 51445.8	350000000	1 2	2003 2.5	2 tree			
277 51445.8	350000000	1 2	2008 2.	1 tree			
278 51445.8	350000000	1 2	2013 2.	1 tree			

The run time on this is not great, so we need to do it outside of the Sweave document. I'll save it all as an R object for the package. The R-object is entitled "expandedTrees" and can be loaded directly after the package loads. Keep in mind that the messy x/y coordinates are just because of a conversion to character, and I cleaned those up in the final dataset. I also converted everything back to numeric where necessary.

```
> ## responses <- list()
> ## for(i in 1:length(plotnames)){ responses[[plotnames[i]]] <- processTreeRows(ssdAllTrees
> ## expandedTrees <- rbind(responses[[1]], responses[[2]], responses[[3]], responses[[4]],
> ##
> head(expandedTrees)
```

```
plot subplot treeid species ingrowth firstrec deathyear
                                                                          Х
                               ABCO
                                                   1992
                                                               NA 89902.54 48132.50
2 crackers
                                           NA
                  1
                          1
3 crackers
                  1
                          1
                               ABCO
                                           NA
                                                   1992
                                                               NA 89902.54 48132.50
                               ABCO
4 crackers
                  1
                          1
                                           NA
                                                   1992
                                                               NA 89902.54 48132.50
5 crackers
                  1
                          1
                               ABCO
                                           NA
                                                   1992
                                                               NA 89902.54 48132.50
6 crackers
                          1
                               ABCO
                                           NA
                                                   1992
                                                               NA 89902.54 48132.50
                  1
                                                               NA 89908.76 48133.72
7 crackers
                  1
                               ABCO
                                           NA
                                                   1992
  measyear dbh stage
      1992 4.4
2
                 tree
3
      1997 5.0
                 tree
      2002 5.6
4
                 tree
5
      2007 6.0
                 tree
6
      2012 6.2
                 tree
7
      1992 8.7
```

# > str(expandedTrees)

```
'data.frame':
                    51493 obs. of 12 variables:
$ plot
                  "crackers" "crackers" "crackers" ...
           : chr
$ subplot
                 1 1 1 1 1 1 1 1 1 1 ...
          : num
$ treeid
           : num
                  1 1 1 1 1 2 2 2 2 2 ...
$ species : chr
                  "ABCO" "ABCO" "ABCO" "ABCO" ...
$ ingrowth : num
                 NA NA NA NA NA NA NA NA NA ...
$ firstrec : num
                 1992 1992 1992 1992 ...
$ deathyear: num
                 NA NA NA NA NA NA NA NA NA ...
$ x
           : num
                 89903 89903 89903 89903 ...
                 48132 48132 48132 48132 ...
           : num
                 1992 1997 2002 2007 2012 ...
$ measyear : num
$ dbh
           : num
                 4.4 5 5.6 6 6.2 8.7 9.1 9.8 10.4 11.1 ...
$ stage
                  "tree" "tree" "tree" ...
           : chr
```

I'm not going to incorporate the script above into the package itself, because the type of code that an individual needs to do to get their plots into the appropriate format will differ based on what they start with. In other words, I don't think the monstrosity of code above is really reusable. It will, however, serve as a record of how I generated the expandedTrees table.