# Manipulation of data-frame data with dutility functions

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# Simple data manipulation for data-frames

- Renaming variables, Deleting variables
- · Looking at the data
- Making new variales for the analysis
- Making factors (groupings)
- Working with factors
- Making a factor from existing numeric variable and vice versa

Here are some key data-manipulation moves on a data-frame which is how we typically organize our data in R. After having read the data into R it will typically be a data-frame, if not we can force it to be a data-frame. The basic idea of the utility functions is to get a simple and easy to type way of making simple data-manipulation on a data-frame much like what is possible in SAS or STATA.

The functions, say, dcut, dfactor and so on are all functions that basically does what the base R cut, factor do, but are easier to use in the context of data-frames and have additional functionality.

```
library(mets)
data(melanoma)

is.data.frame(melanoma)
melanoma=as.data.frame(melanoma)
```

Here we work on the melanoma data that is already read into R and is a data-frame.

# dUtility functions

[1] TRUE

The structure for all functions is

• dfunction(dataframe,y~x | ifcond,...)

to use the function on y in a dataframe grouped by x if condition if cond is valid. The basic functions are

• • •

A generic function daggregate, daggr, can be called with a function as the argument

- daggregate(dataframe,y~x | ifcond,fun=function,...) without the grouping variable (x)
- daggregate(dataframe,~y | ifcond,fun=function,...)

A useful feature is that y and x as well as the subset condition can be specified using regular-expressions or by wildcards (default). Here to illustrate this, we compute the means of certain variables.

```
dmean(melanoma,thick+I(log(thick))~sex|I(days>500))
dmean(melanoma,~thick+I(log(thick))|I(days>500))
dmean(melanoma,~thick+I(log(thick)))
         thick I(log(thick))
 1 0 242.9580 5.060086
    1 320.2429
                    5.353321
        thick I(log(thick))
    271.582011
                 5.168691
        thick I(log(thick))
    291.985366
                   5.223341
```

or summary of all variables starting with "s" and that contains

```
dmean(melanoma, "s*"+"*a*"~sex I(days>500))
       status
 1 0 1.831933 2399.143
    1 1.714286 2169.800
```

# Renaming, deleting, keeping, dropping variables

```
melanoma=drename(melanoma,tykkelse~thick)
names(melanoma)
 [1] "no"
                                      "ulc"
                "status"
                           "days"
                                                  "tykkelse" "sex"
  Deleting variables
data(melanoma)
melanoma=drm(melanoma,~thick+sex)
names (melanoma)
 [1] "no"
              "status" "days"
  or sas style
data(melanoma)
melanoma=ddrop(melanoma,~thick+sex)
names (melanoma)
              "status" "days"
 [1] "no"
                                 "ulc"
  alternatively we can also keep certain variables
data(melanoma)
melanoma=dkeep(melanoma,~thick+sex+status+days)
names (melanoma)
```

"status" "days"

[1] "thick" "sex"

# Looking at the data

```
data(melanoma)
dstr(melanoma)
 'data.frame':
                    205 obs. of 6 variables:
  $ no : int 789 13 97 16 21 469 685 7 932 944 ...
  $ status: int 3 3 2 3 1 1 1 1 3 1 ...
  $ days : int 10 30 35 99 185 204 210 232 232 279 ...
  $ ulc : int 10001111111...
  $ thick : int 676 65 134 290 1208 484 516 1288 322 741 ...
  $ sex : int 1 1 1 0 1 1 1 1 0 0 ...
```

The data can in Rstudio be seen as a data-table but to list certain parts of the data in output window

```
dlist(melanoma)
```

```
no status days ulc thick sex
  789 3 10 1 676 1
2
  13 3
            30 0
                    65 1
            35 0
3
    97 2
                    134 1
           99 0 290 0
4
   16 3
5
    21 1
          185 1 1208 1
201 317 2
           4492 1
                   706
                       1
202 798 2
            4668 0
                   612
                        0
203 806 2
            4688 0
                       0
                   48
204 606 2
            4926 0
                   226 0
205 328 2
           5565 0 290 0
```

```
dlist(melanoma, ~.|sex==1)
```

```
no status days ulc thick
1 789 3
         10 1
             30 0
2
   13 3
                     65
3
    97 2
             35 0
                     134
            185 1
                    1208
5
    21 1
            204 1
6 469 1
                    484
           3909 1
191 445 2
                    806
195 415 2
            4119 0
                     65
197 175 2
           4207 0
                     65
198 493 2
           4310 0
201 317 2
            4492 1
```

```
dlist(melanoma, ~ulc+days+thick+sex|sex==1)
```

```
ulc days thick sex
     10 676 1
       30
  0
           65 1
3
  0
       35
           134
  1
5
     185 1208 1
     204 484 1
      3909 806
191 1
195 0
      4119 65
               1
197 0 4207 65 1
198 0 4310 210 1
201 1 4492 706 1
```

## Getting summaries

# dsummary(melanoma)

```
ulc
                 status
                              days
                                                        thick
Min. : 2.0 Min. :1.00 Min. : 10 Min. :0.000 Min. : 10
1st Qu.:222.0 1st Qu.:1.00 1st Qu.:1525
                                      1st Qu.:0.000 1st Qu.: 97
Median :469.0
             Median :2.00
                          Median :2005
                                       Median :0.000
                                                    Median: 194
             Mean :1.79
Mean :463.9
                          Mean :2153
                                       Mean :0.439
                                                    Mean : 292
3rd Qu.:731.0
             3rd Qu.:2.00
                          3rd Qu.:3042
                                       3rd Qu.:1.000
                                                   3rd Qu.: 356
Max. :992.0 Max. :3.00 Max. :5565 Max. :1.000 Max. :1742
    sex
Min. :0.0000
1st Qu.:0.0000
Median :0.0000
Mean :0.3854
3rd Qu.:1.0000
Max. :1.0000
```

### or for specfic variables

#### dsummary(melanoma,~thick+status+sex)

```
thick
            status
                       sex
Min. : 10 Min. :1.00 Min. :0.0000
Mean : 292 Mean :1.79 Mean :0.3854
3rd Qu.: 356 3rd Qu.:2.00 3rd Qu.:1.0000
Max. :1742 Max. :3.00 Max. :1.0000
```

#### Summaries in different groups (sex)

## dsummary(melanoma,thick+days+status~sex)

```
sex: 0
  thick
                  days
                              status
Min. : 10.0 Min. : 99 Min. :1.000
1st Qu.: 97.0 1st Qu.:1636 1st Qu.:2.000
Median : 162.0
              Median:2059
                           Median :2.000
Mean : 248.6 Mean :2283 Mean :1.833
3rd Qu.: 306.0 3rd Qu.:3131 3rd Qu.:2.000
Max. :1742.0 Max. :5565 Max. :3.000
sex: 1
  thick
                  days
                              status
Min. : 16.0 Min. : 10 Min. :1.000
1st Qu.: 105.0 1st Qu.:1052 1st Qu.:1.000
Median : 258.0
              Median:1860
                           Median :2.000
Mean : 361.1 Mean :1946
                          Mean :1.722
3rd Qu.: 484.0 3rd Qu.:2784
                           3rd Qu.:2.000
Max. :1466.0 Max. :4492 Max. :3.000
```

and only among those with thin-tumours or only females (sex==1)

## dsummary(melanoma,thick+days+status~sex|thick<97)</pre>

```
sex: 0
   thick
                  days
                               status
Min. :10.00 Min. : 355 Min. :1.000
1st Qu.:32.00 1st Qu.:1762 1st Qu.:2.000
Median :64.00 Median :2227 Median :2.000
```

```
Mean :51.48 Mean :2425 Mean :2.034 3rd Qu.:65.00 3rd Qu.:3185 3rd Qu.:2.000
Max. :81.00 Max. :4688 Max. :3.000
sex: 1
    thick
                     days
                                    status
Min. :16.00 Min. : 30 Min. :1.000
1st Qu.:30.00 1st Qu.:1820 1st Qu.:2.000
Median :65.00 Median :2886
                               Median :2.000
Mean :55.75 Mean :2632 Mean :1.875 3rd Qu.:81.00 3rd Qu.:3328 3rd Qu.:2.000
Max. :81.00 Max. :4207 Max. :3.000
```

dsummary(melanoma,thick+status~+1|sex==1)

```
thick
                     status
Min. : 16.0 Min. :1.000
1st Qu.: 105.0 1st Qu.:1.000
Median : 258.0 Median :2.000
Mean : 361.1 Mean :1.722
3rd Qu.: 484.0 3rd Qu.:2.000
Max. :1466.0 Max. :3.000
```

#### Tables between variables

dtable(melanoma,~status+sex)

```
sex 0 1
status
          28 29
1
          91 43
3
          7 7
```

#### All bivariate tables

dtable(melanoma,~status+sex+ulc,level=2)

```
sex 1 2 3
 0 28 91 7
 1 29 43 7
  status
ulc 1 2 3
 0 16 92 7
 1 41 42 7
  sex
ulc 0 1
 0 79 36
 1 47 43
```

status

## All univariate tables

dtable(melanoma,~status+sex+ulc,level=1)

```
status
1 2 3
57 134 14
```

```
sex
 0
     1
126 79
ulc
 0
     1
115 90
```

# Making new variales for the analysis

To define a bunch of new covariates within a data-frame

```
melanoma= transform(melanoma,
        thick2=thick^2,
        lthick=log(thick) )
dhead(melanoma)
   no status days ulc thick sex thick2 lthick
1 789 3 10 1 676 1 456976 6.516193
2 13
        3 30 0 65 1 4225 4.174387
3 97
       2 35 0 134 1 17956 4.897840
3 99 0 290 0 84100 5.669881
 4 16
                           84100 5.669881
        1 185 1 1208 1 1459264 7.096721
5 21
```

When the above definitions are done using a condition this can be achieved using the dtransform function that extends transform with a possible condition

```
melanoma=dtransform(melanoma,ll=thick*1.05^ulc,sex==1)
melanoma=dtransform(melanoma,ll=thick,sex!=1)
dmean(melanoma,ll~sex+ulc)
  sex ulc
               11
 1 0 0 173.7342
 2 1 0 197.3611
 3 0 1 374.5532
 4 1 1 523.1198
```

# *Making factors (groupings)*

On the melanoma data the variable thick gives the thickness of the melanom tumour. For some analyses we would like to make a factor depending on the thickness. This can be done in several different ways

```
melanoma=dcut(melanoma,~thick,breaks=c(0,200,500,800,2000))
```

New variable is named thickcat.o by default. To see levels of factors in data-frame

```
dlevels(melanoma)
```

```
thickcat.0 #levels=:4
[1] "[0,200]" "(200,500]" "(500,800]" "(800,2e+03]"
```

## Checking group sizes

```
dtable(melanoma,~thickcat.0)
 thickcat.0
    [0,200]
           (200,500] (500,800] (800,2e+03]
      109
           64
                     20 12
  With adding to the data-frame directly
```

```
dcut(melanoma,breaks=c(0,200,500,800,2000)) <- gr.thick1~thick
dlevels(melanoma)
```

```
thickcat.0 #levels=:4
[1] "[0,200]" "(200,500]" "(500,800]"
                                          "(800,2e+03]"
gr.thick1 #levels=:4
[1] "[0,200]" "(200,500]" "(500,800]"
                                          "(800,2e+03]"
```

new variable is named thickcat.o (after first cut-point), or to get quartiles with default names thick.cat.4

```
dcut(melanoma) <- ~ thick ### new variable is thickcat.4
dlevels(melanoma)
thickcat.0 #levels=:4
[1] "[0,200]" "(200,500]" "(500,800]"
                                       "(800,2e+03]"
     _____
 gr.thick1 #levels=:4
 [1] "[0,200]" "(200,500]" "(500,800]"
                                       "(800,2e+03]"
 thickcat.4 #levels=:4
 [1] "[10,97]" "(97,194]" "(194,356]" "(356,1.74e+03]"
```

or median groups, here starting again with the original data,

```
data(melanoma)
dcut(melanoma,breaks=2) <- ~ thick ### new variable is thick.2</pre>
dlevels(melanoma)
 thickcat.2 #levels=:2
 [1] "[10,194]" "(194,1.74e+03]"
```

## to control new names

```
data(melanoma)
mela= dcut(melanoma,thickcat4+dayscat4~thick+days,breaks=4)
dlevels(mela)
 thickcat4 #levels=:4
 [1] "[10,97]" "(97,194]"
                                  "(194,356]"
                                                 "(356,1.74e+03]"
 dayscat4 #levels=:4
 [1] "[10,1.52e+03]" "(1.52e+03,2e+03]" "(2e+03,3.04e+03]"
 [4] "(3.04e+03,5.56e+03]"
```

or

```
data(melanoma)
dcut(melanoma,breaks=4) <- thickcat4+dayscat4~thick+days</pre>
dlevels(melanoma)
 thickcat4 #levels=:4
 [1] "[10,97]" "(97,194]" "(194,356]" "(356,1.74e+03]"
 dayscat4 #levels=:4
  [1] "[10,1.52e+03]" "(1.52e+03,2e+03]" "(2e+03,3.04e+03]"
  [4] "(3.04e+03,5.56e+03]"
   This can also be typed out more specifically
melanoma$gthick = cut(melanoma$thick,breaks=c(0,200,500,800,2000))
melanoma$gthick =

→ cut(melanoma$thick, breaks=quantile(melanoma$thick), include.lowest=TRUE)

 Working with factors
To see levels of covariates in data-frame
data(melanoma)
dcut(melanoma,breaks=4) <- thickcat4~thick</pre>
dlevels(melanoma)
 thickcat4 #levels=:4
  [1] "[10,97]" "(97,194]" "(194,356]" "(356,1.74e+03]"
   To relevel the factor
dtable (melanoma, thickcat4)
melanoma = drelevel(melanoma, "thickcat4, ref="(194,356]")
dlevels(melanoma)
 thickcat4
       [10,97] (97,194] (194,356] (356,1.74e+03]
  56 53
thickcat4 #levels=:4
[1] "[10 07]"
                                  45 51
  [1] "[10,97]" "(97,194]"
                                   "(194,356]"
                                                   "(356,1.74e+03]"
  thickcat4.(194,356] #levels=:4
                                 "(97,194]" "(356,1.74e+03]"
  [1] "(194,356]" "[10,97]"
   or to take the third level in the list of levels, same as above,
melanoma = drelevel(melanoma,~thickcat4,ref=2)
dlevels(melanoma)
  thickcat4 #levels=:4
  [1] "[10,97]" "(97,194]" "(194,356]" "(356,1.74e+03]"
 thickcat4.(194,356] #levels=:4
  [1] "(194,356]" "[10,97]"
                                    "(97,194]"
                                                    "(356,1.74e+03]"
```

"(194,356]"

"(356,1.74e+03]"

thickcat4.2 #levels=:4

[1] "(97,194]" "[10,97]"

To combine levels of a factor (first combinining first 3 groups into one)

```
melanoma = drelevel(melanoma,~thickcat4,newlevels=1:3)
dlevels(melanoma)
thickcat4 #levels=:4
 [1] "[10,97]" "(97,194]" "(194,356]" "(356,1.74e+03]"
 thickcat4.(194,356] #levels=:4
                                 "(97,194]"
                                                 "(356,1.74e+03]"
 [1] "(194,356]" "[10,97]"
 thickcat4.2 #levels=:4
 [1] "(97,194]" "[10,97]" "(194,356]"
                                                 "(356.1.74e+031"
thickcat4.1:3 #levels=:2
 [1] "[10,97]-(194,356]" "(356,1.74e+03]"
```

or to combine groups 1 and 2 into one group and 3 and 4 into another

```
dkeep(melanoma) <- ~thick+thickcat4</pre>
melanoma = drelevel(melanoma,gthick2~thickcat4,newlevels=list(1:2,3:4))
dlevels(melanoma)
thickcat4 #levels=:4
 [1] "[10,97]" "(97,194]" "(194,356]" "(356,1.74e+03]"
 gthick2 #levels=:2
 [1] "[10,97]-(97,194]" "(194,356]-(356,1.74e+03]"
```

# Changing order of factor levels

```
dfactor(melanoma,levels=c(3,1,2,4)) <- thickcat4.2~thickcat4
dlevel(melanoma,~ "thickcat4*")
dtable(melanoma,~thickcat4+thickcat4.2)
thickcat4 #levels=:4
 [1] "[10,97]" "(97,194]" "(194,356]" "(356,1.74e+03]"
 thickcat4.2 #levels=:4
 [1] "(194,356]" "[10,97]" "(97,194]" "(356,1.74e+03]"
            thickcat4.2 (194,356] [10,97] (97,194] (356,1.74e+03]
 thickcat4
 [10,97]
                                   56
                             0 0 53
 (97,194]
                                                        0
                                    0
                                          0
                                                         0
 (194,356]
                              45
 (356,1.74e+03]
                                    0
```

#### Combine levels but now control factor-level names

```
melanoma=drelevel(melanoma,gthick3~thickcat4,newlevels=list(group1.2=1:2,group3.4=3:4))
dlevels(melanoma)
 thickcat4 #levels=:4
 [1] "[10,97]" "(97,194]" "(194,356]" "(356,1.74e+03]"
 gthick2 #levels=:2
```

```
[1] "[10,97]-(97,194]" "(194,356]-(356,1.74e+03]"
 _____
thickcat4.2 #levels=:4
[1] "(194,356]" "[10,97]" "(97,194]" "(356,1.74e+03]"
gthick3 #levels=:2
[1] "group1.2" "group3.4"
```

Making a factor from existing numeric variable and vice versa

A numeric variable "status" with values 1,2,3 into a factor by

```
data(melanoma)
melanoma = dfactor(melanoma,~status,

→ labels=c("malignant-melanoma", "censoring", "dead-other"))
melanoma = dfactor(melanoma,sexl~sex,labels=c("females","males"))
 dtable(melanoma,~sexl+status.f)
          status.f malignant-melanoma censoring dead-other
  sexl
  females
                                  28
                                           91
                                                      7
                                  29
                                           43
  males
```

A gender factor with values "M", "F" can be converted into numerics by

```
melanoma = dnumeric(melanoma,~sexl)
dstr(melanoma, "sex*")
dtable(melanoma,~'sex*',level=2)
 'data.frame':
                  205 obs. of 3 variables:
 $ sex : int 1 1 1 0 1 1 1 1 0 0 ...
  $ sexl : Factor w/ 2 levels "females", "males": 2 2 2 1 2 2 2 1 1 ...
 $ sexl.n: num 2 2 2 1 2 2 2 2 1 1 ...
        sex
 sexl
         0 1
  females 126 0
  males 0 79
     sex
 sexl.n 0 1
     1 126 0
     2 0 79
     sexl
 sexl.n females males
    1 126 0
2 0 79
```