gClinBiomarker: example use cases

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1 Loading packages and data
library(knitr) library(devtools)
Warning: package 'devtools' was built under R version 3.3.2
<pre>install_github("RPackages/gClinBiomarker", host="https://github.roche.com/api/v3") library(gClinBiomarker)</pre>
Warning: package 'survival' was built under R version 3.3.2

2 Example data set

library(ggplot2)
data(input)

sample.data <- input</pre>

```
head(sample.data)
     Patient.ID Sample.ID Arm Age Weight ECOG Sex Baseline.SLD
                                                                          Country
                    SID001 TRT
## 1
         PID001
                                                   F
                                      61.0
                                                         550.4970 Western Europe
## 2
         PID002
                    SID002 TRT
                                      87.0
                                                   F
                                                         549.7751
                                 60
                                               1
## 3
         PID003
                    SID003 TRT
                                 53
                                      57.6
                                               1
                                                   М
                                                         548.8829 Western Europe
## 4
         PID004
                                                   F
                                                         549.6050
                    SID004 TRT
                                      57.0
                                                                               USA
## 5
         PID005
                    SID005 TRT
                                 49
                                      72.5
                                               2
                                                   F
                                                         551.5498
                                                                             Other
## 6
         PID006
                    SID006 TRT
                                      76.0
                                               0
                                                         549.2565
                                                                             Other
##
     CD8.ihc KRAS.mutant KRAS.exprs
                                            0S
                                                     PFS
                                                          Response BEP OS.event
## 1
           3
                   Mutant
                               171.25 9.691992 5.782341
                                                                 PR
## 2
           2
                Wild Type
                                 2.77 4.435318 1.412731
                                                                 PD
                                                                      1
                                                                                1
## 3
           3
                Wild Type
                                61.82 9.691992 1.215606
                                                                 PD
                                                                                0
## 4
           1
                                71.51 1.905544 1.905544
                                                                 PD
                                                                      1
                                                                                0
                   Mutant
## 5
          NA
                     <NA>
                                   NA 2.628337 2.628337
                                                                 SD
                                                                                1
## 6
           2
                                44.63 4.829569 4.139630 NON CR/PD
                                                                                0
                   Mutant
##
     PFS.event Lab_ontrt
## 1
             0 28.993922
## 2
             1 9.890427
## 3
             1 26.896020
## 4
             1 5.421392
## 5
             1 14.741382
## 6
             0 16.957035
```

```
str(sample.data)
## 'data.frame':
                  550 obs. of 19 variables:
## $ Patient.ID : chr "PID001" "PID002" "PID003" "PID004" ...
## $ Sample.ID : chr "SID001" "SID002" "SID003" "SID004" ...
## $ Arm
                : Factor w/ 2 levels "CTRL", "TRT": 2 2 2 2 2 2 2 2 1 ...
## $ Age
                : int 40 60 53 44 49 62 70 49 62 45 ...
## $ Weight
               : num 61 87 57.6 57 72.5 76 46 69.3 65 54 ...
## $ ECOG
                : int 0 1 1 0 2 0 0 1 0 1 ...
                : chr "F" "F" "M" "F" ...
## $ Sex
## $ Baseline.SLD: num 550 550 549 550 552 ...
## $ Country : Factor w/ 3 levels "Other", "USA",..: 3 2 3 2 1 1 3 3 3 1 ...
## $ CD8.ihc : num 3 2 3 1 NA 2 NA 1 NA NA ...
## $ KRAS.mutant : Factor w/ 2 levels "Mutant", "Wild Type": 1 2 2 1 NA 1 NA 2 NA NA ...
## $ KRAS.exprs : num 171.25 2.77 61.82 71.51 NA ...
## $ OS
                : num 9.69 4.44 9.69 1.91 2.63 ...
## $ PFS
                : num 5.78 1.41 1.22 1.91 2.63 ...
## $ Response : Factor w/ 6 levels "CR", "NE", "NON CR/PD",...: 5 4 4 4 6 3 4 3 4 6 ...
## $ BEP
               : num 1 1 1 1 0 1 0 1 0 0 ...
## $ OS.event : int 0 1 0 0 1 0 0 0 1 0 ...
## $ PFS.event : int 0 1 1 1 1 0 1 0 1 0 ...
## $ Lab ontrt : num 28.99 9.89 26.9 5.42 14.74 ...
```

3 SummaryVars(): Summarize demographics variables (check demographics imbalance)

3.1 Default

	All(CTRL)	BEP(CTRL)	All(TRT)	BEP(TRT)
Age				
N	182	81	368	176
Mean	52.54	53.1	54.03	54.42
Median	51.5	52	54	54
Min-Max	2785	3285	3089	3389
NA's	0	0	0	0
Sex				
Total	182	81	368	176
NA's	0	0	0	0
\mathbf{F}	89 (48.9%)	39 (48.15%)	184 (50%)	90 (51.14%)
M	93 (51.1%)	42 (51.85%)	184 (50%)	86 (48.86%)

3.2 Compare to non-BEP instead of ITT

	BEP_1(CTRL)	BEP_0(CTRL)	BEP_1(TRT)	BEP_0(TRT)
Age				
N	81	101	176	192
Mean	53.1	52.09	54.42	53.67
Median	52	50	54	54
Min-Max	3285	2782	3389	3079
NA's	0	0	0	0
Sex				
Total	81	101	176	192
NA's	0	0	0	0
F	39 (48.15%)	50 (49.5%)	90 (51.14%)	94 (48.96%)
M	42 (51.85%)	51 (50.5%)	86 (48.86%)	98 (51.04%)

3.3 Allow testing

	All(CTRL)	BEP(CTRL)	pvalue(CTRL)	All(TRT)	BEP(TRT)	pvalue(TRT)
Age						
N	182	81	0.5	368	176	0.69
Mean	52.54	53.1		54.03	54.42	
Median	51.5	52		54	54	
Min-Max	2785	3285		3089	3389	
NA's	0	0		0	0	
Sex						
Total	182	81	0.88	368	176	0.75
NA's	0	0		0	0	
F	89 (48.9%)	39 (48.15%)		184 (50%)	90 (51.14%)	
M	93 (51.1%)	42 (51.85%)		184 (50%)	86 (48.86%)	

3.4 Reorder TRT/CTRL arm in display

	All(TRT)	BEP(TRT)	All(CTRL)	BEP(CTRL)
	All(TRT)	BEP(TRT)	All(CTRL)	BEP(CTRL)
Age				
N	368	176	182	81
Mean	54.03	54.42	52.54	53.1
Median	54	54	51.5	52
Min-Max	3089	3389	2785	3285
NA's	0	0	0	0
Sex				
Total	368	176	182	81
NA's	0	0	0	0
F	184 (50%)	90 (51.14%)	89 (48.9%)	39 (48.15%)
M	184 (50%)	86 (48.86%)	93 (51.1%)	42 (51.85%)

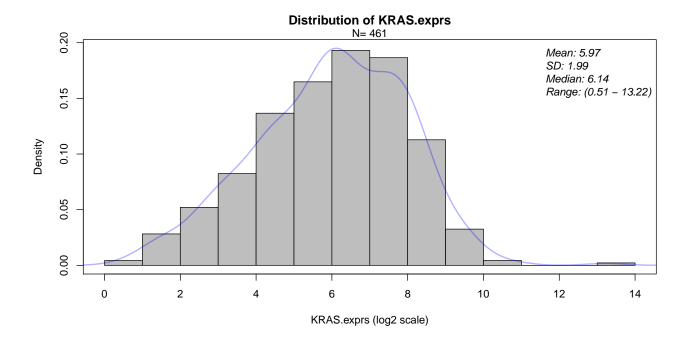
3.5 Combine trt arms

	All	BEP
Age		
N	550	257
Mean	53.53	54
Median	53	54
Min-Max	2789	3289
NA's	0	0
Sex		
Total	550	257
NA's	0	0
F	273 (49.64%)	129 (50.19%)
M	277 (50.36%)	128 (49.81%)

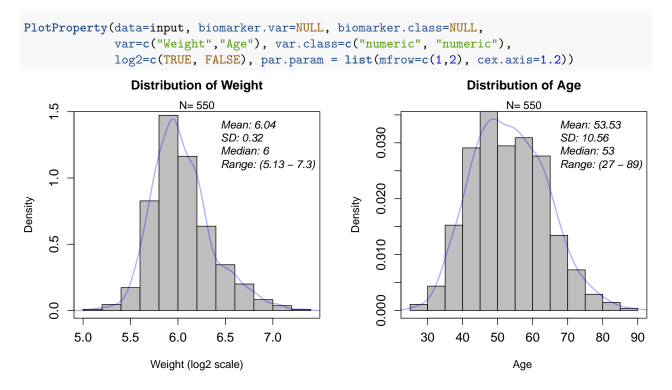
4 PlotProperty(): Plot biomarker, clinical covariate property and their association

 $4.1\,$ Numeric biomarker variable. No clinical variables. Log transformation.

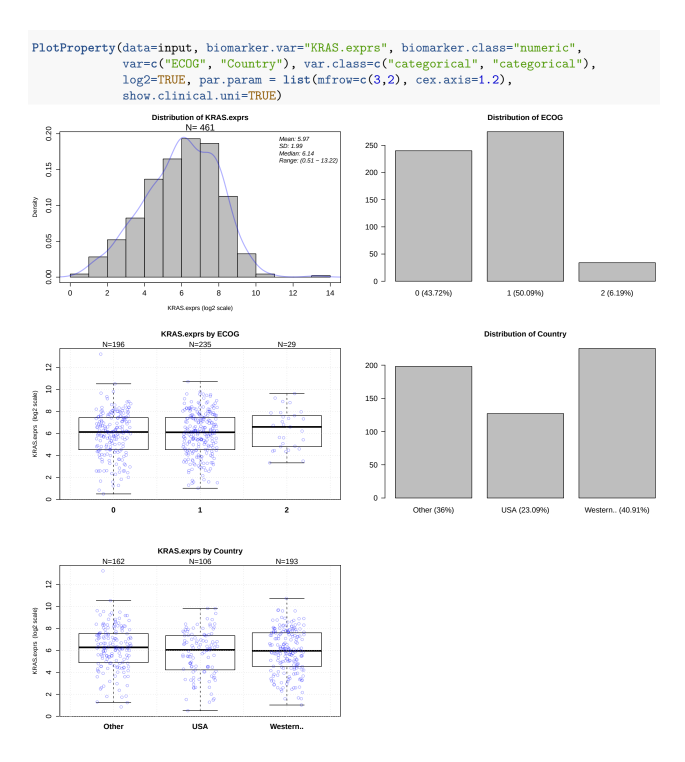
```
PlotProperty(data=input, biomarker.var="KRAS.exprs", biomarker.class="numeric", log2=TRUE)
```



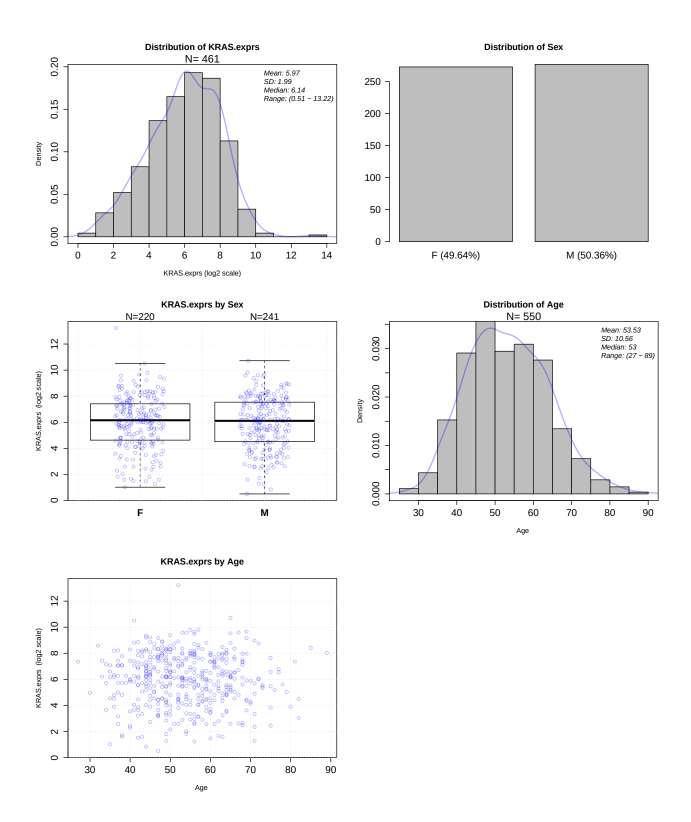
4.2 No biomarker variable. Two numeric clinical variables. Log transformation for one varible.



4.3 Numeric biomarker variable. Two clinical categorical variables. Log transformation for numeric variable.



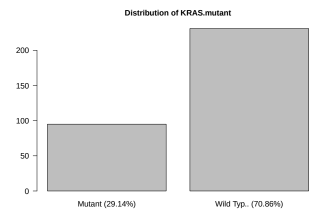
4.4 Numeric biomarker variable. Two clinical variables: one is categorocal, second is numeric. Log transformation for biomarker (numeric) variable.

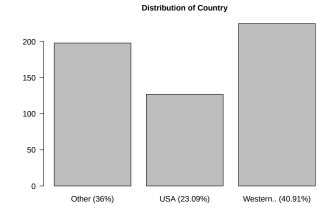


4.5 Categorical biomarker variable. Categorical clinical variable.

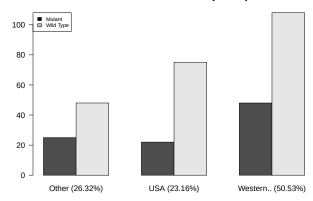
```
PlotProperty(data=input, biomarker.var="KRAS.mutant", biomarker.class="categorical", var="Country", var.class="categorical",
```



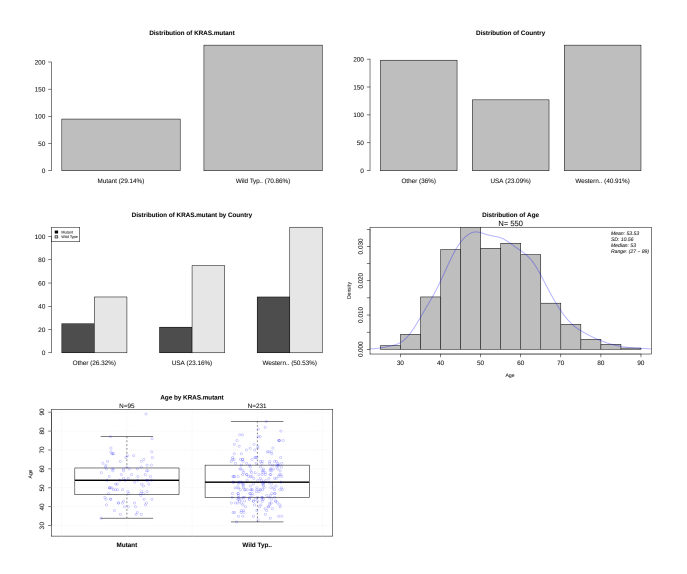




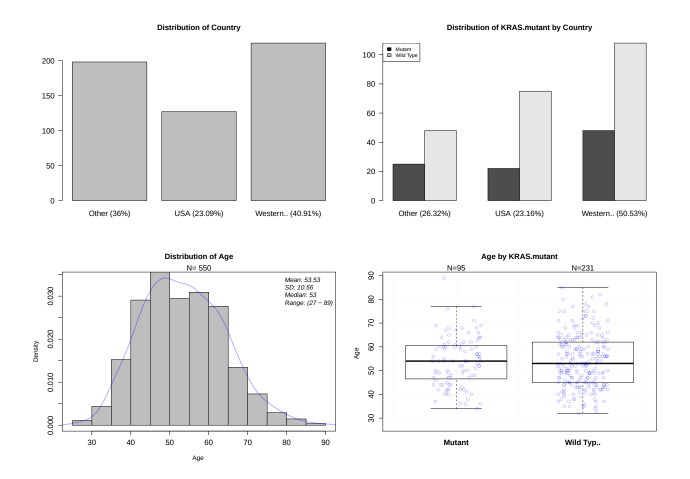
Distribution of KRAS.mutant by Country



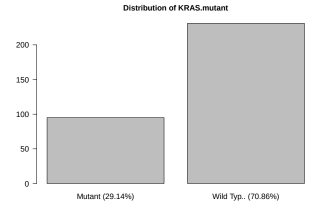
4.6 Categorial biomarker variable. Categorical and numeric clinical variables. No log transformation.

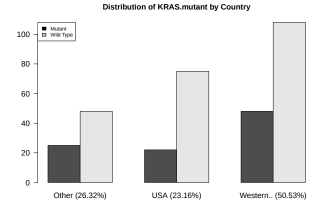


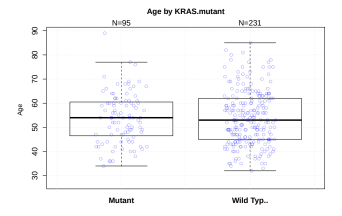
4.7 Categorial biomarker variable. Categorical and numeric clinical variables. No log transformation. Don't plot biomarker variable but plot clinical variables and association between biomarker and clinical variables.



4.8 Categorial biomarker variable. Categorical and numeric clinical variables. No log transformation. Don't plot clinical variables but plot biomarker variable and association between biomarker and clinical variables.

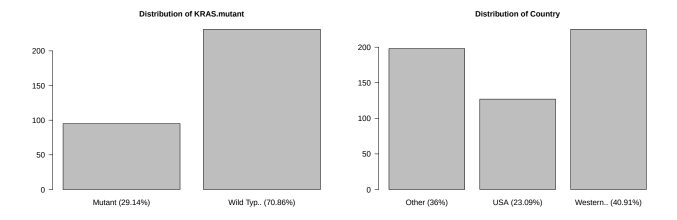


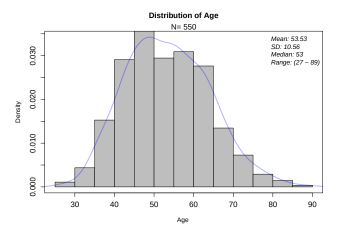




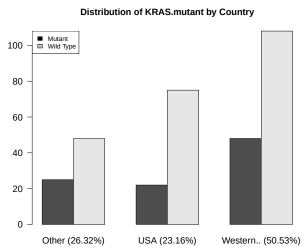
4.9 Categorial biomarker variable. Categorical and numeric clinical variables. No log transformation. Don't plot association between variables but plot biomarker and clinical variables.

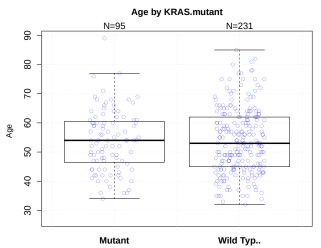
```
PlotProperty(data=input, biomarker.var="KRAS.mutant", biomarker.class="categorical", var=c("Country", "Age"), var.class=c("categorical", "numeric"), par.param = list(mfrow=c(2,2), cex.axis=1.2), show.association = FALSE, show.clinical.uni=TRUE)
```



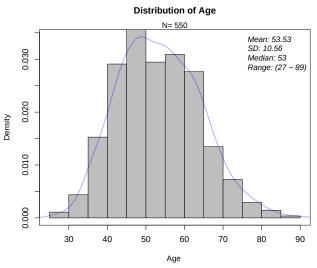


4.10 Categorial biomarker variable. Categorical and numeric clinical variables. No log transformation. Plot only association between variables.



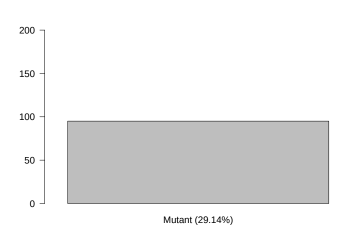


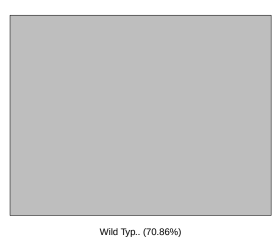
4.11 Categorial biomarker variable. Categorical and numeric clinical variables. No log transformation. Plot only clinical variables.



Categorial biomarker variable. Categorical and numeric clinical variables. No log transformation. Plot only biomarker variable.

Distribution of KRAS.mutant



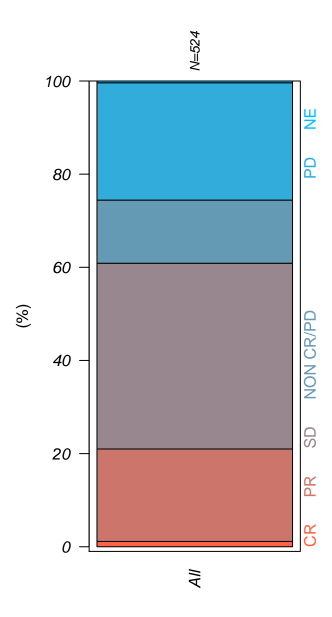


5 PlotRspBar(): Compare response outcome across different population

5.1 Plot response of one population

entries with missing outcome.var are removed! 524 entries left

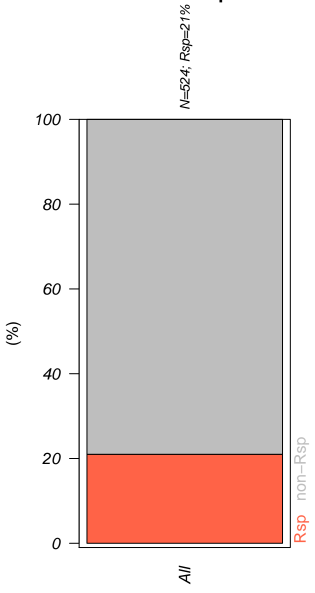
Association of response rate



5.2 Plot response of one population, binarize outcome (classify classes to responder vs non responder)

entries with missing outcome.var are removed! 524 entries left

Association of response rate

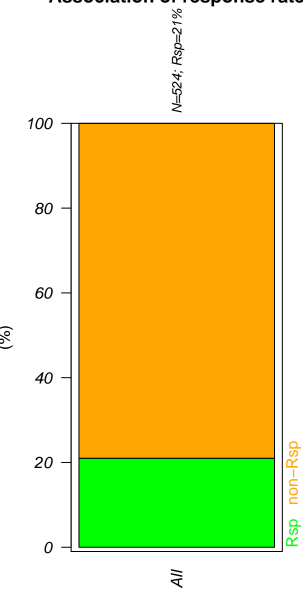


5.3 Change color

```
rsp.response = c("CR","PR"),
rsp.nonresponse = c("SD", "PD","NON CR/PD","NE",NA),
col=c("green","orange"))
```

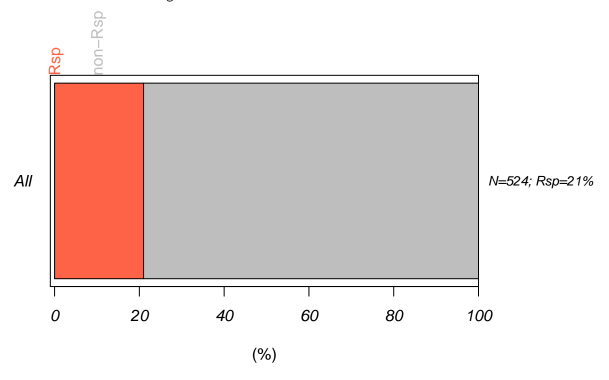
entries with missing outcome.var are removed! 524 entries left

Association of response rate



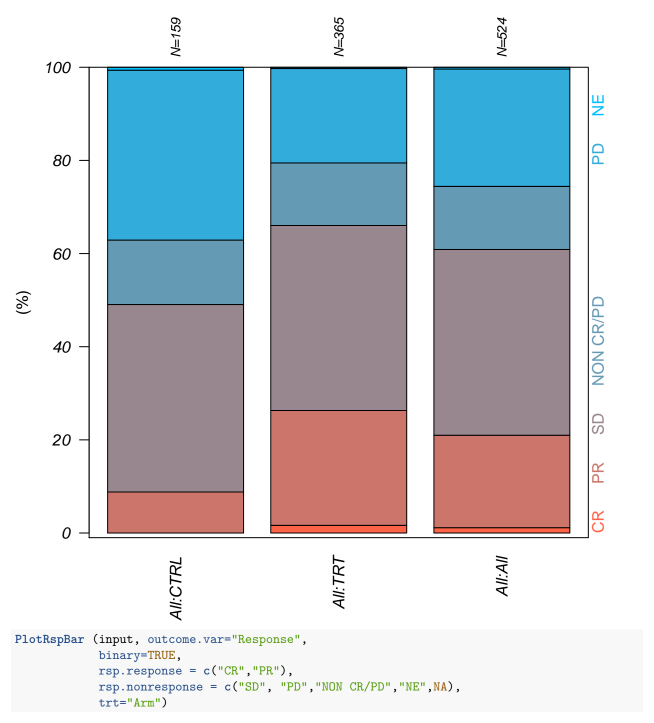
5.4 Plot horizontally

entries with missing outcome.var are removed! 524 entries left

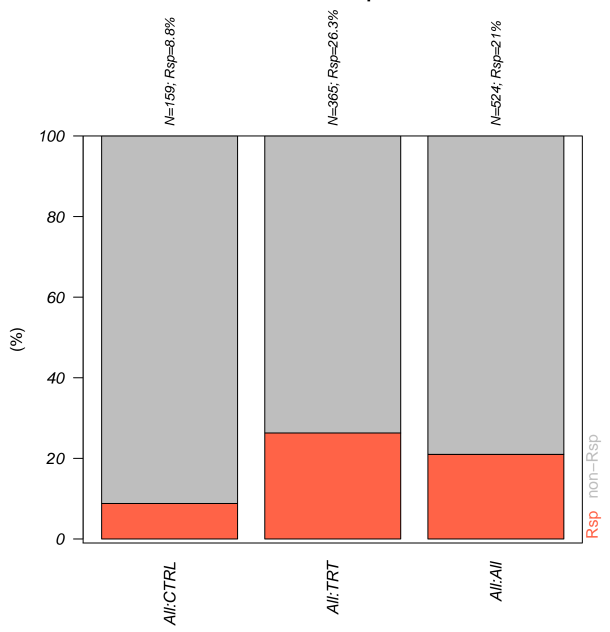


5.5 By arm

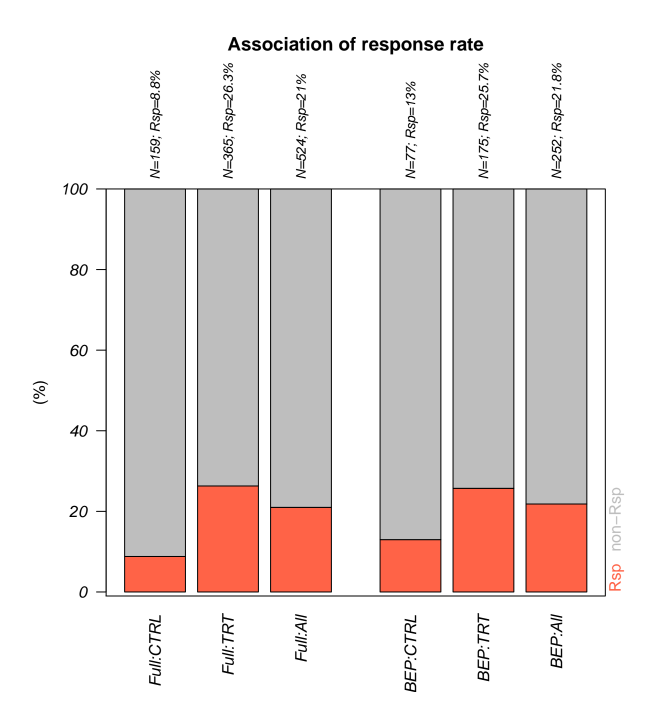
Association of response rate



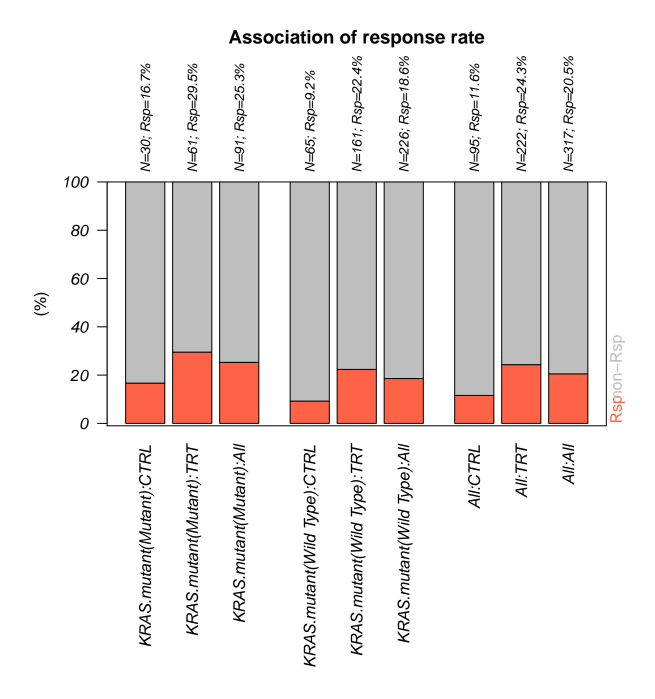




5.6 Compare full population vs BEP

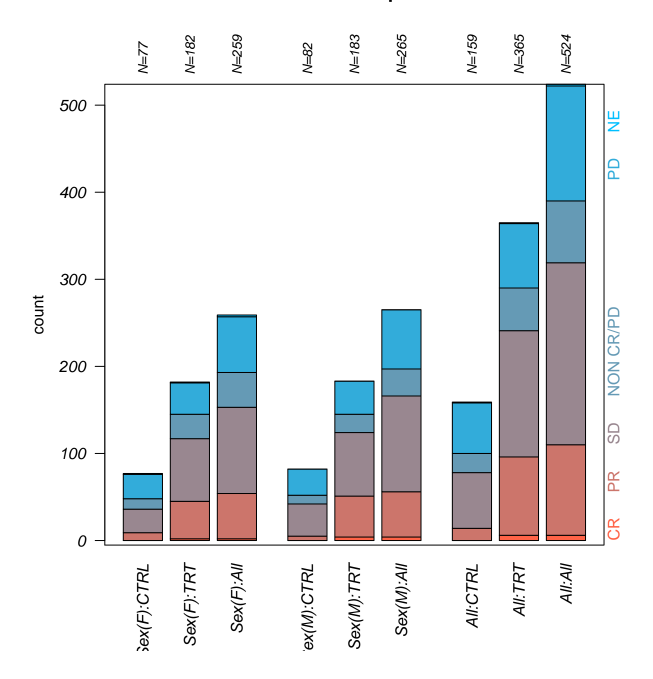


5.7 Compare subgroups



5.8 Plot count instead of percentage

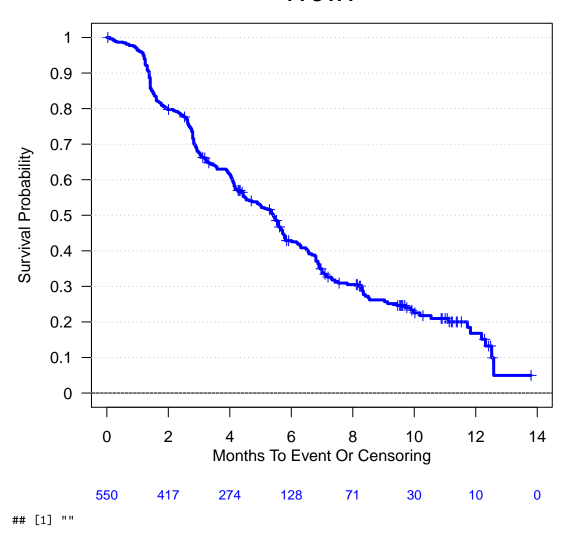
Association of response rate



6 PlotKM(): Plot KM curves for subpopulations

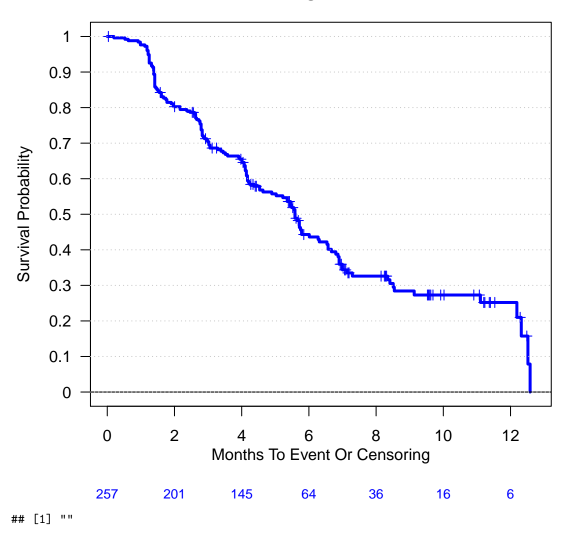
6.1 Without subgroup

PFS ITT



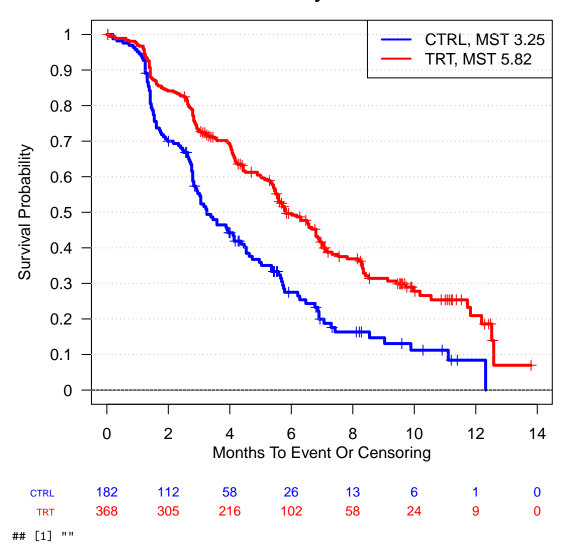
6.2 Without subgroup, BEP

PFS BEP



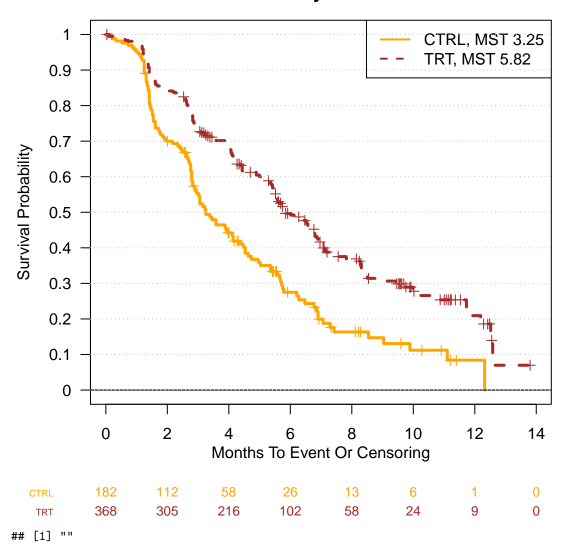
6.3 By TRT

PFS ITT by treatment



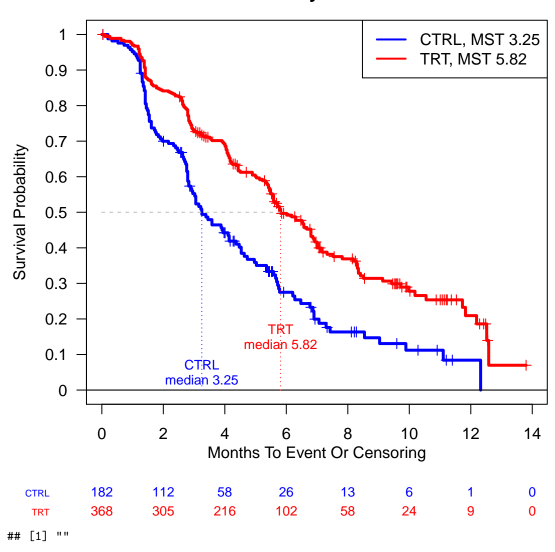
6.4 By TRT, change color, line type

PFS ITT by treatment



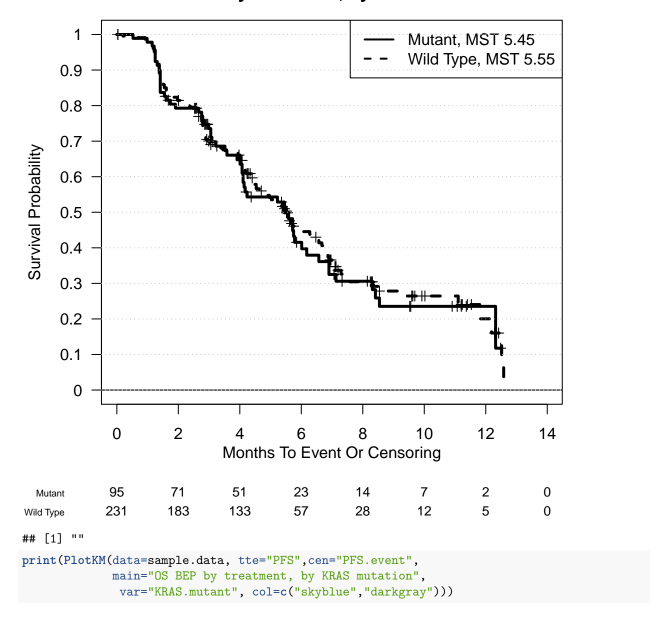
6.5 Mark median PFS, no grid

PFS ITT by treatment

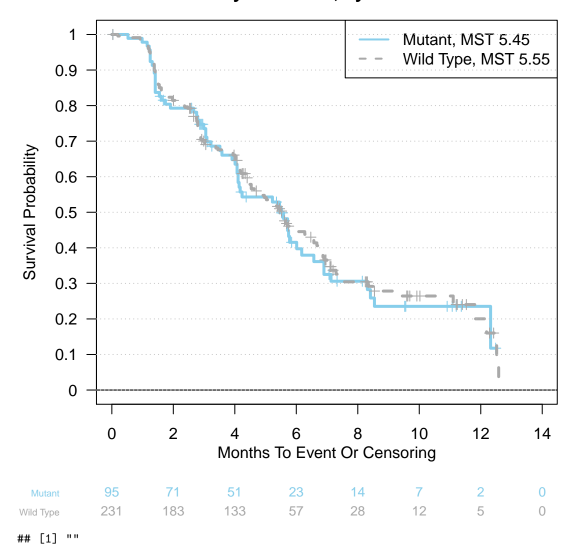


6.6 By KRAS.mutant only

OS BEP by treatment, by KRAS mutation

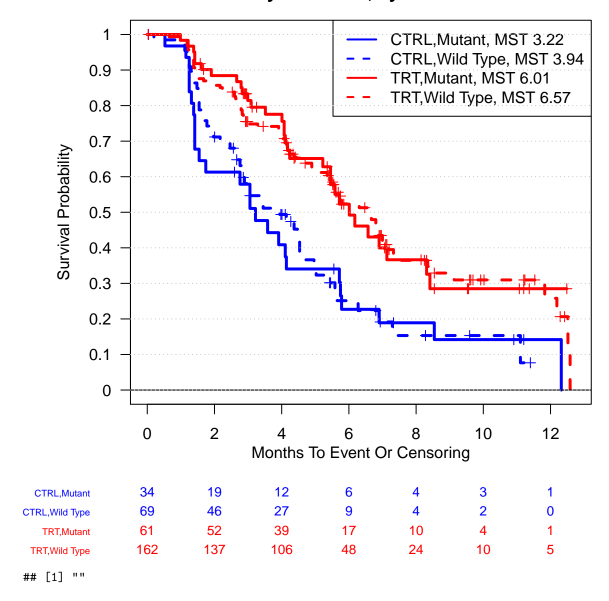


OS BEP by treatment, by KRAS mutation



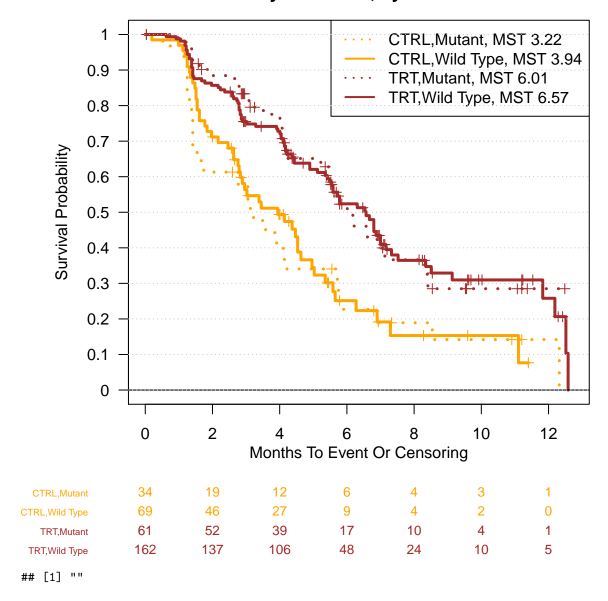
6.7 By TRT and KRAS.mutant

OS BEP by treatment, by KRAS mutation



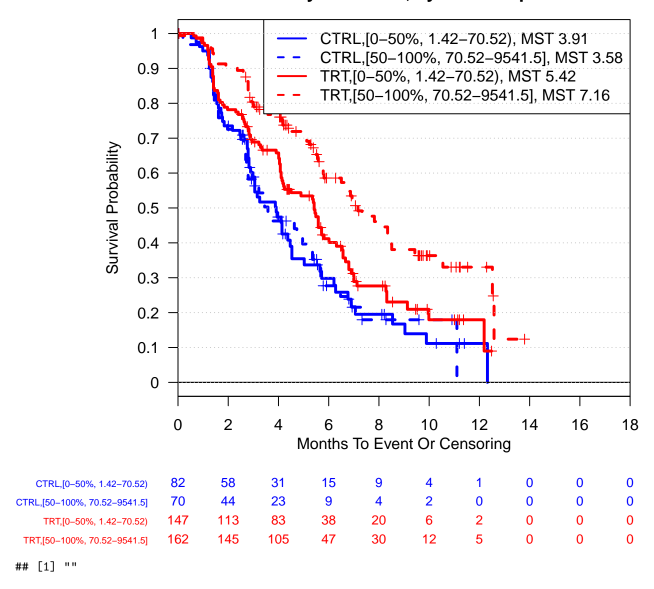
6.8 By TRT and KRAS.mutant, change color and line type

PFS BEP by treatment, by KRAS mutation



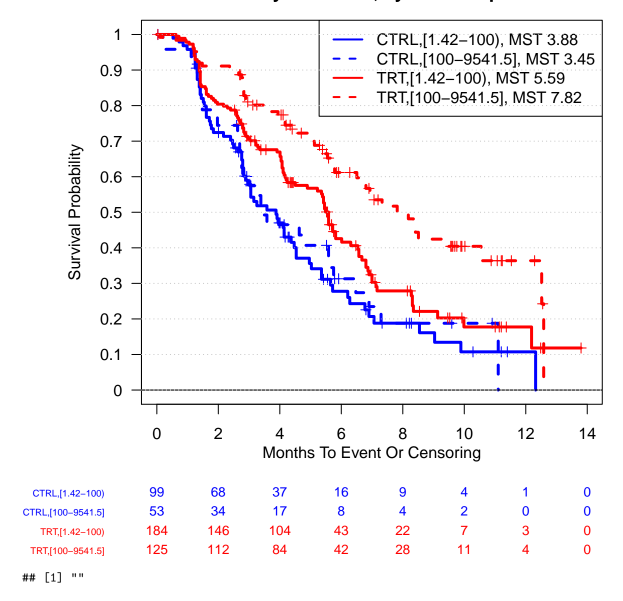
6.9 By TRT and KRAS.expression (continuous biomarker, cut at median)

PFS BEP by treatment, by KRAS expression



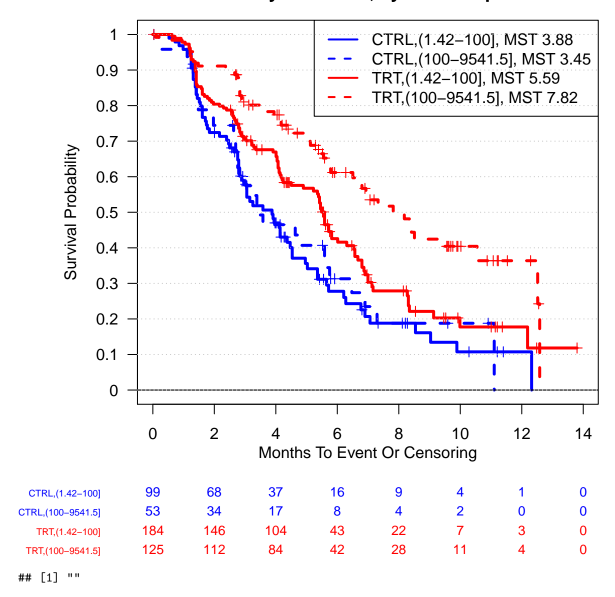
6.10 By TRT and KRAS.expression (continuous biomarker, cut at a numerical cutoff)

PFS BEP by treatment, by KRAS expression



6.11 By TRT and KRAS.expression (continuous biomarker, cut at a numerical cutoff). High group defined as "greater than", instead of "greater than or equal to"

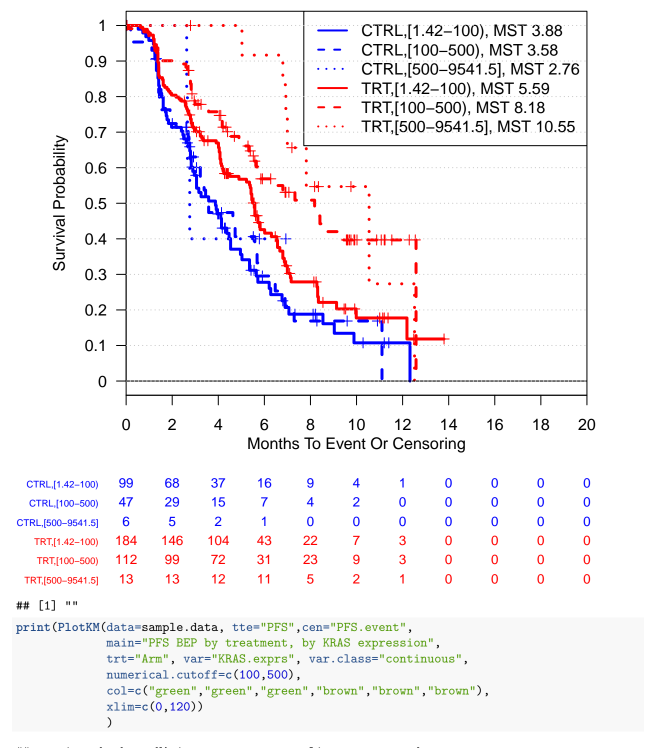
PFS BEP by treatment, by KRAS expression



6.12 By TRT and KRAS.expression, more than 2 groups

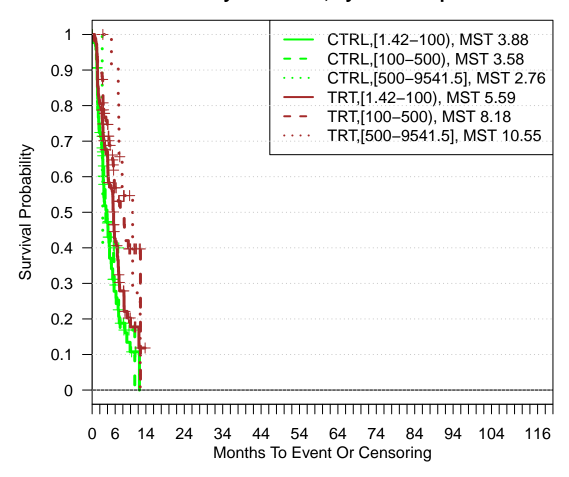
entries who have NA in trt, var, or varlist are removed

PFS BEP by treatment, by KRAS expression



entries who have NA in trt, var, or varlist are removed

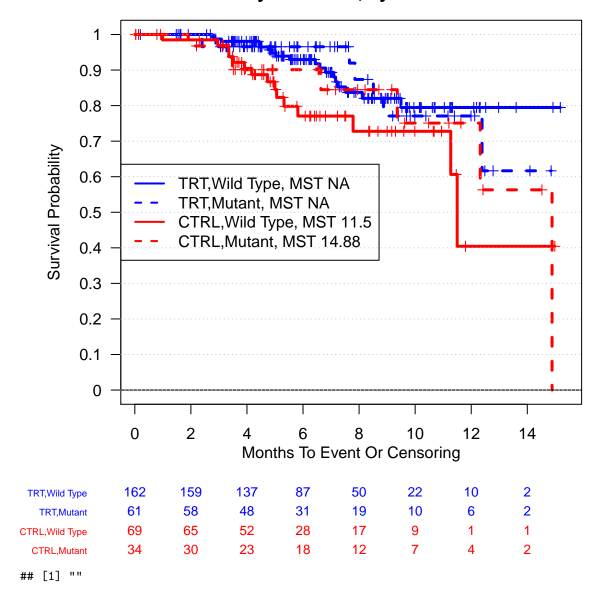
PFS BEP by treatment, by KRAS expression



6.13 More flexibility, reorder subgroups

'varlist' is specified, trt and var parameters will be ignored
entries who have NA in trt, var, or varlist are removed

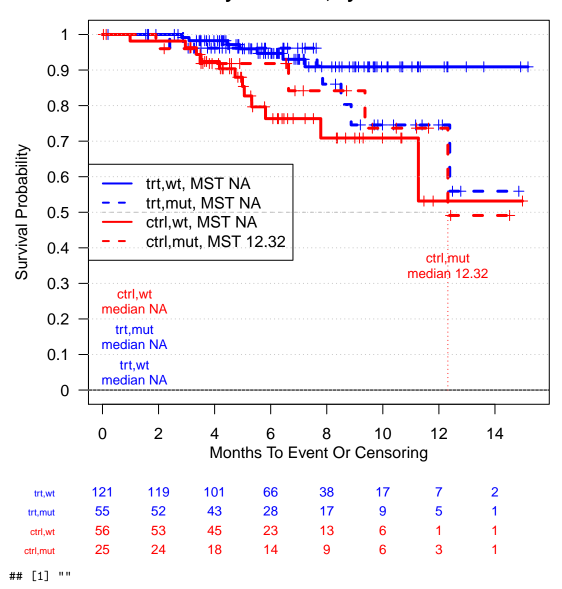
OS BEP by treatment, by KRAS mutation



6.14 By TRT and KRAS.mutant, reorder and rename

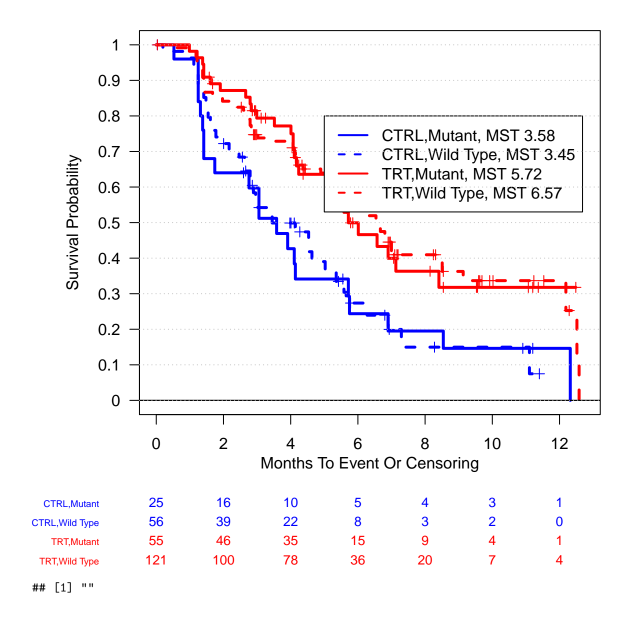
'varlist' is specified, trt and var parameters will be ignored

OS BEP by treatment, by KRAS mutation



6.15 Legend location

more than one elements in 'var', trt parameter will be ignored
'varlist' is specified, trt and var parameters will be ignored



- 7 PlotTabForestBiomarker(): forest plot and summary statistics table for a single biomarker (include cutoff exploration for cont. biomarker)
- 7.1 Survival outcome, 2-arm, categorical variable

by default BEP will be defined as patients non-NA biomarker measurement

Across-arm Effect of Biomarker PFS, KRAS.mutant Unadjusted, unstratified analysis

Subgroup	Group	Event/N	MST	HR	(CI)	raw P		
All	CTRL	120 / 182	3.25					
	TRT	202 / 368	5.82	0.54	(0.43 – 0.67)	7.8e-08	-	
BEP	CTRL	73 / 103	3.58					
	TRT	118 / 223	6.54	0.5	(0.37 – 0.66)	2.8e-06	-	
KRAS.mutant(Mutant)	CTRL	26 / 34	3.22					
	TRT	31 / 61	6.01	0.48	(0.28 - 0.8)	0.0055		
KRAS.mutant(Wild Type)	CTRL	47 / 69	3.94					
	TRT	87 / 162	6.57	0.5	(0.35 – 0.72)	0.00019	_	

0.28 0.53 1.0 1.89 3.57 --- TRT better [HR] CTRL bette * Unadjusted Interaction P = 0.

```
##
                                   code.v
##
    [1,] "Subgroup"
                                   "Group" "Event/N"
                                                        "MST"
                                            "120 / 182" "3.25" ""
##
    [2,] "All"
                                   "CTRL"
    [3,] ""
##
                                   "TRT"
                                            "202 / 368" "5.82" "0.54"
##
    [4,] "BEP"
                                   "CTRL"
                                            "73 / 103"
                                                        "3.58" ""
    [5,] ""
                                    "TRT"
                                            "118 / 223" "6.54" "0.5"
##
    [6,] "KRAS.mutant(Mutant)"
                                    "CTRL"
                                            "26 / 34"
                                                        "3.22" ""
##
    [7,] ""
                                   "TRT"
                                            "31 / 61"
                                                        "6.01" "0.48"
##
                                   "CTRL"
                                            "47 / 69"
                                                        "3.94" ""
##
    [8,] "KRAS.mutant(Wild Type)"
    [9,] ""
                                    "TRT"
                                            "87 / 162"
                                                        "6.57" "0.5"
##
##
##
    [1,] "CI"
                        "raw P"
    [2,] ""
##
    [3,] "0.43 - 0.67" "7.8e-08"
##
    [4,] ""
##
    [5,] "0.37 - 0.66" "2.8e-06"
##
    [6,] ""
    [7,] "0.28 - 0.8"
                        "0.0055"
##
   [8,] ""
##
   [9,] "0.35 - 0.72" "0.00019"
```

7.2 Survival outcome, another format

```
PlotTabForestBiomarker(data=input,

outcome.class="survival",
outcome.var=c("PFS","PFS.event"),
trt="Arm",
var="KRAS.mutant",
var.class="categorical",
tabforest = TRUE)
```

Some NAs in var column, will define the non NA entries as BEP

Across-arm Effect of Biomarker PFS, KRAS.mutant

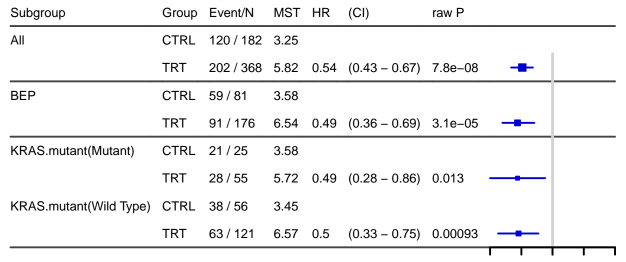
Subgroup	Group	Event/N	MST	HR	CI	raw P	Fore	st plot
							TRT better	CTRL better
All	CTRL	120 / 182	3.25					1
	TRT	202 / 368	5.82	0.54	0.43 - 0.67	7.8e-08	-	
BEP	CTRL	73 / 103	3.58					1
	TRT	118 / 223	6.54	0.5	0.37 - 0.66	2.8e-06		i I I
KRAS.mutant(Mutant)	CTRL	26 / 34	3.22					i ! !
	TRT	31 / 61	6.01	0.48	0.28 - 0.8	0.0055 -	•	
KRAS.mutant(Wild Type)	CTRL	47 / 69	3.94					i !
	TRT	87 / 162	6.57	0.5	0.35 - 0.72	0.00019	-	i ! !
Unadjusted Interaction P = 0.87		•	-					
	Unadjus	ted, unstratif	ied analy	/sis		0.2	8 0.53	1 1.89

```
##
                                  code.v
   [1,] "Subgroup"
                                  "Group" "Event/N"
                                                      "MST" "HR"
##
   [2,] "All"
                                  "CTRL"
                                          "120 / 182" "3.25" ""
##
##
   [3,] ""
                                  "TRT"
                                          "202 / 368" "5.82" "0.54"
                                          "73 / 103" "3.58" ""
##
   [4,] "BEP"
                                  "CTRL"
   [5,] ""
                                  "TRT"
                                          "118 / 223" "6.54" "0.5"
                                                      "3.22" ""
   [6,] "KRAS.mutant(Mutant)"
                                  "CTRL"
                                          "26 / 34"
##
   [7,] ""
                                  "TRT"
                                          "31 / 61"
                                                      "6.01" "0.48"
##
   [8,] "KRAS.mutant(Wild Type)" "CTRL"
                                          "47 / 69"
                                                      "3.94" ""
##
   [9,] ""
                                  "TRT"
                                          "87 / 162" "6.57" "0.5"
##
##
##
   [1,] "CI"
                       "raw P"
##
   [2,] ""
   [3,] "0.43 - 0.67" "7.8e-08"
##
##
   [4,] ""
##
  [5,] "0.37 - 0.66" "2.8e-06"
## [6,] ""
                       11 11
## [7,] "0.28 - 0.8" "0.0055"
## [8,] ""
## [9,] "0.35 - 0.72" "0.00019"
```

7.3 Survival outcome, 2-arm, categorical variable, don't show ITT, BEP, rename variable name in display

user can also define BEP column

Across-arm Effect of Biomarker PFS, KRAS.mutant Unadjusted, unstratified analysis



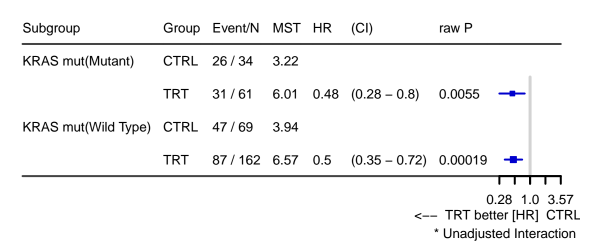
0.28 0.53 1.0 1.89 3.57 --- TRT better [HR] CTRL bette * Unadjusted Interaction P = 0.

```
##
                                   code.v
##
    [1,] "Subgroup"
                                   "Group" "Event/N"
                                                        "MST"
                                                               "HR"
    [2,] "All"
                                   "CTRL"
                                            "120 / 182" "3.25" ""
##
    [3,] ""
                                   "TRT"
                                            "202 / 368" "5.82" "0.54"
##
                                                        "3.58" ""
   [4,] "BEP"
                                   "CTRL"
                                           "59 / 81"
##
                                                        "6.54" "0.49"
   [5,] ""
                                   "TRT"
                                            "91 / 176"
##
                                           "21 / 25"
                                                        "3.58" ""
##
    [6,] "KRAS.mutant(Mutant)"
                                   "CTRL"
                                                        "5.72" "0.49"
   [7,] ""
                                   "TRT"
                                           "28 / 55"
##
                                                        "3.45" ""
                                   "CTRL"
                                           "38 / 56"
##
   [8,] "KRAS.mutant(Wild Type)"
                                   "TRT"
                                            "63 / 121" "6.57" "0.5"
   [9,] ""
##
##
  [1,] "CI"
##
                        "raw P"
##
  [2,] ""
   [3,] "0.43 - 0.67" "7.8e-08"
##
```

```
[4,] ""
##
   [5,] "0.36 - 0.69" "3.1e-05"
##
   [6,] ""
  [7,] "0.28 - 0.86" "0.013"
##
   [8,] ""
## [9,] "0.33 - 0.75" "0.00093"
PlotTabForestBiomarker(data=input,
                                  outcome.class="survival",
                                  outcome.var=c("PFS","PFS.event"),
                                  trt="Arm",
                                  var="KRAS.mutant",
                                  var.class="categorical",
                                  var.name="KRAS mut",
                                  show.itt=FALSE,
                                  show.bep=FALSE)
```

Some NAs in var column, will define the non NA entries as BEP

Across-arm Effect of Biomarker PFS, KRAS mut Unadjusted, unstratified analysis



```
##
                               code.v
## [1,] "Subgroup"
                               "Group" "Event/N"
                                                   "MST"
                                                                  "CI"
                               "CTRL"
                                       "26 / 34"
                                                  "3.22" ""
                                                                  11 11
## [2,] "KRAS mut(Mutant)"
                                                  "6.01" "0.48" "0.28 - 0.8"
## [3,] ""
                               "TRT"
                                       "31 / 61"
                                       "47 / 69"
                                                  "3.94" ""
                                                                  11 11
## [4,] "KRAS mut(Wild Type)"
                               "CTRL"
## [5,] ""
                               "TRT"
                                       "87 / 162" "6.57" "0.5" "0.35 - 0.72"
##
## [1,] "raw P"
## [2,] ""
## [3,] "0.0055"
## [4,] ""
## [5,] "0.00019"
```

7.4 Survival outcome, 2-arm, continuous variable, greater than some percentage cutoffs

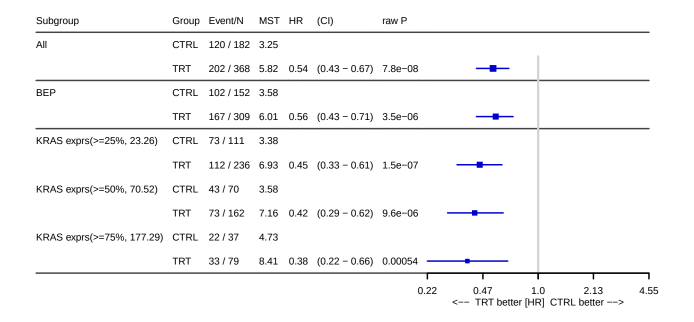
```
PlotTabForestBiomarker(data=input,

outcome.class="survival",
outcome.var=c("PFS","PFS.event"),
trt="Arm",
var="KRAS.exprs",
var.class="numeric", var.name="KRAS exprs",
percentile.cutoff=c(.25,.5,.75),

# cols=c("black","black","darkgreen","darkgreen"),
numerical.cutoff=NULL,
greater=TRUE, less=FALSE,
within.bin=FALSE,
show.itt=TRUE, show.bep=TRUE)
```

Some NAs in var column, will define the non NA entries as BEP

Across-arm Effect of Biomarker PFS, KRAS exprs Unadjusted, unstratified analysis



```
##
                                      code.v
##
    [1,] "Subgroup"
                                      "Group" "Event/N"
                                                           "MST"
                                      "CTRL"
                                              "120 / 182" "3.25" ""
##
   [2,] "All"
   [3,] ""
                                      "TRT"
##
                                              "202 / 368" "5.82" "0.54"
##
   [4,] "BEP"
                                      "CTRL"
                                              "102 / 152" "3.58" ""
##
    [5,] ""
                                      "TRT"
                                               "167 / 309" "6.01" "0.56"
##
   [6,] "KRAS exprs(>=25%, 23.26)"
                                      "CTRL"
                                              "73 / 111" "3.38" ""
   [7,] ""
                                      "TRT"
                                              "112 / 236" "6.93" "0.45"
                                                           "3.58" ""
    [8,] "KRAS exprs(>=50%, 70.52)"
                                      "CTRL"
                                              "43 / 70"
##
    [9,] ""
                                      "TRT"
                                              "73 / 162" "7.16" "0.42"
```

```
## [10,] "KRAS exprs(>=75%, 177.29)" "CTRL" "22 / 37" "4.73" ""
## [11,] ""
                                    "TRT"
                                           "33 / 79" "8.41" "0.38"
##
## [1,] "CI"
                      "raw P"
   [2,] ""
##
## [3,] "0.43 - 0.67" "7.8e-08"
## [4,] ""
## [5,] "0.43 - 0.71" "3.5e-06"
##
   [6,] ""
## [7,] "0.33 - 0.61" "1.5e-07"
## [8,] ""
## [9,] "0.29 - 0.62" "9.6e-06"
## [10,] ""
## [11,] "0.22 - 0.66" "0.00054"
```

7.5 Survival outcome, 2-arm, continuous variable, less than some percentage cutoffs

```
PlotTabForestBiomarker(data=input,

outcome.class="survival",
outcome.var=c("PFS","PFS.event"),
trt="Arm",
var="KRAS.exprs",
var.class="numeric", var.name="KRAS exprs",
percentile.cutoff=c(.25,.5,.75),
numerical.cutoff=NULL,
greater=FALSE, less=TRUE,
within.bin=FALSE,
show.itt=TRUE, show.bep=TRUE,
bep = 'BEP', bep.indicator=1)
```

Across-arm Effect of Biomarker PFS, KRAS exprs Unadjusted, unstratified analysis

Subgroup	Group	Event/N	MST	HR	(CI)	raw P					
All	CTRL	120 / 182	3.25								_
	TRT	202 / 368	5.82	0.54	(0.43 – 0.67)	7.8e-0	8 -	-			
BEP	CTRL	59 / 81	3.58								
	TRT	91 / 176	6.54	0.49	(0.36 – 0.69)	3.1e-0	5 —				
KRAS exprs(<25%, 22.63)	CTRL	16 / 22	4.14								
	TRT	30 / 42	5.45	0.7	(0.37 – 1.32)	0.27		•	+		
KRAS exprs(<50%, 63.56)	CTRL	32 / 44	4.11								
	TRT	52 / 84	5.72	0.68	(0.43 – 1.06)	0.09	-	-	+		
KRAS exprs(<75%, 170.07)	CTRL	44 / 61	3.91								
	TRT	75 / 131	5.59	0.61	(0.42 – 0.89)	0.0098	_	-			
							0.36	0.60 < TRT better	1.0 [HR] CTR	1.67 L better>	2.78

```
##
                                     code.v
                                                         "MST" "HR"
##
    [1,] "Subgroup"
                                     "Group" "Event/N"
    [2,] "All"
                                     "CTRL"
                                            "120 / 182" "3.25" ""
   [3,] ""
                                     "TRT"
                                             "202 / 368" "5.82" "0.54"
##
                                                         "3.58" ""
    [4,] "BEP"
                                            "59 / 81"
                                     "CTRL"
##
   [5,] ""
##
                                     "TRT"
                                             "91 / 176" "6.54" "0.49"
                                    "CTRL"
                                            "16 / 22"
                                                         "4.14" ""
##
   [6,] "KRAS exprs(<25%, 22.63)"
                                             "30 / 42"
##
   [7,] ""
                                     "TRT"
                                                         "5.45" "0.7"
                                             "32 / 44"
                                                         "4.11" ""
##
    [8,] "KRAS exprs(<50%, 63.56)"
                                     "CTRL"
                                             "52 / 84"
                                                         "5.72" "0.68"
##
   [9,] ""
                                     "TRT"
   [10,] "KRAS exprs(<75%, 170.07)" "CTRL"
                                             "44 / 61"
                                                         "3.91" ""
   [11,] ""
                                     "TRT"
                                             "75 / 131" "5.59" "0.61"
##
##
##
   [1,] "CI"
                       "raw P"
##
   [2,] ""
    [3,] "0.43 - 0.67" "7.8e-08"
##
##
   [4,] ""
   [5,] "0.36 - 0.69" "3.1e-05"
##
##
   [6,] ""
   [7,] "0.37 - 1.32" "0.27"
##
  [8,] ""
##
## [9,] "0.43 - 1.06" "0.09"
## [10,] ""
## [11,] "0.42 - 0.89" "0.0098"
```

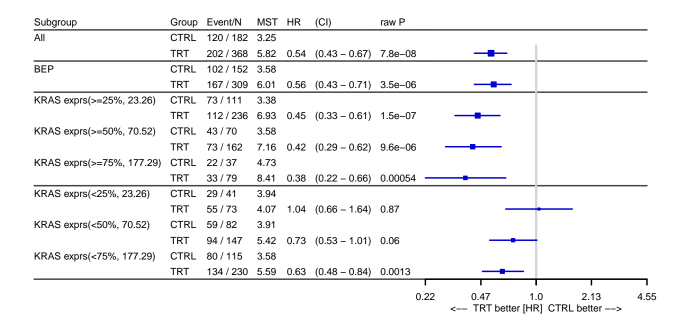
7.6 Survival outcome, 2-arm, continuous variable, greater and less than some percentage cutoffs

```
PlotTabForestBiomarker(data=input,

outcome.class="survival",
outcome.var=c("PFS","PFS.event"),
trt="Arm",
var="KRAS.exprs",
var.class="numeric", var.name="KRAS exprs",
percentile.cutoff=c(.25,.5,.75),
numerical.cutoff=NULL,
greater=TRUE, less=TRUE,
within.bin=FALSE,
show.itt=TRUE, show.bep=TRUE)
```

Some NAs in var column, will define the non NA entries as BEP

Across-arm Effect of Biomarker PFS, KRAS exprs Unadjusted, unstratified analysis



```
##
                                       code.v
                                       "Group"
                                               "Event/N"
                                                            "MST"
##
    [1,] "Subgroup"
##
    [2,] "All"
                                       "CTRL"
                                               "120 / 182" "3.25" ""
    [3,] ""
                                               "202 / 368" "5.82" "0.54"
                                       "TRT"
##
##
   [4,] "BEP"
                                       "CTRL"
                                               "102 / 152" "3.58" ""
##
   [5,] ""
                                       "TRT"
                                               "167 / 309" "6.01" "0.56"
    [6,] "KRAS exprs(>=25%, 23.26)"
                                       "CTRL"
                                               "73 / 111"
                                                            "3.38" ""
##
   [7,] ""
                                       "TRT"
                                               "112 / 236" "6.93" "0.45"
##
   [8,] "KRAS exprs(>=50%, 70.52)"
                                       "CTRL"
                                               "43 / 70"
                                                            "3.58" ""
   [9,] ""
                                               "73 / 162"
                                                            "7.16" "0.42"
##
                                       "TRT"
## [10,] "KRAS exprs(>=75%, 177.29)" "CTRL"
                                               "22 / 37"
                                                            "4.73" ""
```

```
## [11,] ""
                                     "TRT"
                                             "33 / 79"
                                                         "8.41" "0.38"
                                     "CTRL"
## [12,] "KRAS exprs(<25%, 23.26)"
                                             "29 / 41"
                                                         "3.94" ""
                                             "55 / 73"
                                                         "4.07" "1.04"
## [13,] ""
                                     "TRT"
                                                          "3.91" ""
## [14,] "KRAS exprs(<50%, 70.52)"
                                     "CTRL"
                                             "59 / 82"
                                             "94 / 147" "5.42" "0.73"
## [15,] ""
                                     "TRT"
## [16,] "KRAS exprs(<75%, 177.29)"
                                     "CTRL" "80 / 115" "3.58" ""
## [17,] ""
                                     "TRT"
                                             "134 / 230" "5.59" "0.63"
##
## [1,] "CI"
                       "raw P"
## [2,] ""
## [3,] "0.43 - 0.67" "7.8e-08"
## [4,] ""
## [5,] "0.43 - 0.71" "3.5e-06"
## [6,] ""
## [7,] "0.33 - 0.61" "1.5e-07"
## [8,] ""
## [9,] "0.29 - 0.62" "9.6e-06"
## [10,] ""
## [11,] "0.22 - 0.66" "0.00054"
## [12,] ""
## [13,] "0.66 - 1.64" "0.87"
## [14,] ""
                       11 11
## [15,] "0.53 - 1.01" "0.06"
## [16.] ""
## [17,] "0.48 - 0.84" "0.0013"
"Less" rows next to "Greater" rows:
PlotTabForestBiomarker(data=input,
                                  outcome.class="survival",
                                  outcome.var=c("PFS","PFS.event"),
                                  trt="Arm",
                                  var="KRAS.exprs",
                                  var.class="numeric", var.name="KRAS exprs",
                                  percentile.cutoff=c(.25,.5,.75),
                                  numerical.cutoff=NULL,
                                  greater=TRUE, less=TRUE, greater.by.less = TRUE,
                                  within.bin=FALSE,
                                  show.itt=TRUE, show.bep=TRUE)
```

Some NAs in var column, will define the non NA entries as BEP

Across-arm Effect of Biomarker PFS, KRAS exprs Unadjusted, unstratified analysis

Subgroup	Group	Event/N	MST	HR	(CI)	raw P		
All	CTRL	120 / 182	3.25					_
	TRT	202 / 368	5.82	0.54	(0.43 - 0.67)	7.8e-08	_	
BEP	CTRL	102 / 152	3.58					
	TRT	167 / 309	6.01	0.56	(0.43 - 0.71)	3.5e-06		
KRAS exprs(>=25%, 23.26)	CTRL	73 / 111	3.38					
	TRT	112 / 236	6.93	0.45	(0.33 – 0.61)	1.5e-07		
KRAS exprs(<25%, 23.26)	CTRL	29 / 41	3.94					
	TRT	55 / 73	4.07	1.04	(0.66 – 1.64)	0.87		
KRAS exprs(>=50%, 70.52)	CTRL	43 / 70	3.58					
	TRT	73 / 162	7.16	0.42	(0.29 - 0.62)	9.6e-06		
KRAS exprs(<50%, 70.52)	CTRL	59 / 82	3.91					
	TRT	94 / 147	5.42	0.73	(0.53 – 1.01)	0.06		
KRAS exprs(>=75%, 177.29)	CTRL	22 / 37	4.73					
	TRT	33 / 79	8.41	0.38	(0.22 - 0.66)	0.00054		
KRAS exprs(<75%, 177.29)	CTRL	80 / 115	3.58					
	TRT	134 / 230	5.59	0.63	(0.48 - 0.84)	0.0013		
						0		.0 2.13 4.5 R] CTRL better>

```
##
                                      code.v
##
   [1,] "Subgroup"
                                      "Group" "Event/N"
                                                          "MST" "HR"
                                      "CTRL"
                                              "120 / 182" "3.25" ""
   [2,] "All"
   [3,] ""
                                      "TRT"
                                              "202 / 368" "5.82" "0.54"
##
    [4,] "BEP"
                                              "102 / 152" "3.58" ""
                                      "CTRL"
##
##
   [5,] ""
                                      "TRT"
                                              "167 / 309" "6.01" "0.56"
                                      "CTRL"
                                              "73 / 111" "3.38" ""
##
   [6,] "KRAS exprs(>=25%, 23.26)"
##
   [7,] ""
                                      "TRT"
                                              "112 / 236" "6.93" "0.45"
                                              "29 / 41"
##
   [8,] "KRAS exprs(<25%, 23.26)"
                                      "CTRL"
                                                          "3.94" ""
                                      "TRT"
                                              "55 / 73"
                                                          "4.07" "1.04"
##
   [9,] ""
## [10,] "KRAS exprs(>=50%, 70.52)"
                                      "CTRL"
                                              "43 / 70"
                                                          "3.58" ""
## [11,] ""
                                      "TRT"
                                              "73 / 162"
                                                          "7.16" "0.42"
## [12,] "KRAS exprs(<50%, 70.52)"
                                      "CTRL"
                                              "59 / 82"
                                                          "3.91" ""
## [13,] ""
                                      "TRT"
                                              "94 / 147"
                                                          "5.42" "0.73"
## [14,] "KRAS exprs(>=75%, 177.29)"
                                      "CTRL"
                                              "22 / 37"
                                                          "4.73" ""
                                              "33 / 79"
## [15,] ""
                                      "TRT"
                                                          "8.41" "0.38"
  [16,] "KRAS exprs(<75%, 177.29)"
                                      "CTRL"
                                              "80 / 115"
                                                          "3.58" ""
                                      "TRT"
                                              "134 / 230" "5.59" "0.63"
## [17,] ""
##
##
   [1,] "CI"
                       "raw P"
##
   [2,] ""
   [3,] "0.43 - 0.67" "7.8e-08"
##
##
   [4,] ""
##
    [5,] "0.43 - 0.71" "3.5e-06"
   [6,] ""
##
   [7,] "0.33 - 0.61" "1.5e-07"
   [8,] ""
##
   [9,] "0.66 - 1.64" "0.87"
```

```
## [10,] "" ""
## [11,] "0.29 - 0.62" "9.6e-06"
## [12,] "" ""
## [13,] "0.53 - 1.01" "0.06"
## [14,] "" ""
## [15,] "0.22 - 0.66" "0.00054"
## [16,] "" ""
## [17,] "0.48 - 0.84" "0.0013"
```

7.7 Survival outcome, 2-arm, continuous variable, within cutoff bin

```
PlotTabForestBiomarker(data=input,

outcome.class="survival",
outcome.var=c("PFS","PFS.event"),
trt="Arm",
var="KRAS.exprs",
var.class="numeric", var.name="KRAS exprs",
percentile.cutoff=c(.25,.5,.75),
numerical.cutoff=NULL,
greater=FALSE, less=FALSE,
within.bin=TRUE,
show.itt=TRUE, show.bep=TRUE)
```

Some NAs in var column, will define the non NA entries as BEP

Across-arm Effect of Biomarker PFS, KRAS exprs Unadjusted, unstratified analysis

Subgroup	Group	Event/N	MST	HR	(CI)	raw P	
All	CTRL	120 / 182	3.25				_
	TRT	202 / 368	5.82	0.54	(0.43 – 0.67)	7.8e-0	98 —
BEP	CTRL	102 / 152	3.58				
	TRT	167 / 309	6.01	0.56	(0.43 – 0.71)	3.5e-0	06 —
KRAS exprs[0-25%, 1.41-23.26)	CTRL	29 / 41	3.94				
	TRT	55 / 73	4.07	1.04	(0.66 – 1.64)	0.87	-
KRAS exprs[25-50%, 23.26-70.52)	CTRL	30 / 41	3.15				
	TRT	39 / 74	6.67	0.51	(0.31 – 0.82)	0.0057	· _ •
KRAS exprs[50-75%, 70.52-177.29)	CTRL	21 / 33	3.22				
	TRT	40 / 83	5.82	0.46	(0.27 – 0.79)	0.0049	·
KRAS exprs[75-100%, 177.29-9541.51]	CTRL	22 / 37	4.73				
	TRT	33 / 79	8.41	0.38	(0.22 – 0.66)	0.0005	4
							0.22 0.47 1.0 2.13 4.55 TRT better [HR] CTRL better>

```
## code.v

## [1,] "Subgroup" "Group" "Event/N" "MST"

## [2,] "All" "CTRL" "120 / 182" "3.25"
```

```
[3,] ""
                                                "TRT"
                                                        "202 / 368" "5.82"
##
##
   [4,] "BEP"
                                                "CTRL"
                                                        "102 / 152" "3.58"
  [5,] ""
##
                                                "TRT"
                                                        "167 / 309" "6.01"
## [6,] "KRAS exprs[0-25%, 1.41-23.26)"
                                                "CTRL"
                                                        "29 / 41"
                                                                     "3.94"
                                                        "55 / 73"
   [7,] ""
                                                "TRT"
                                                                     "4.07"
## [8,] "KRAS exprs[25-50%, 23.26-70.52)"
                                                "CTRL"
                                                        "30 / 41"
                                                                     "3.15"
## [9.] ""
                                                "TRT"
                                                        "39 / 74"
                                                                     "6.67"
                                                        "21 / 33"
## [10,] "KRAS exprs[50-75%, 70.52-177.29)"
                                                "CTRL"
                                                                     "3.22"
## [11,] ""
                                                "TRT"
                                                        "40 / 83"
                                                                     "5.82"
## [12,] "KRAS exprs[75-100%, 177.29-9541.51]" "CTRL"
                                                        "22 / 37"
                                                                     "4.73"
## [13,] ""
                                                "TRT"
                                                        "33 / 79"
                                                                     "8.41"
##
## [1,] "HR"
                "CI"
                              "raw P"
## [2,] ""
                11 11
## [3,] "0.54" "0.43 - 0.67" "7.8e-08"
   [4,] ""
                11 11
##
## [5,] "0.56" "0.43 - 0.71" "3.5e-06"
              11 11
## [6,] ""
## [7,] "1.04" "0.66 - 1.64" "0.87"
## [8,] ""
                11 11
## [9,] "0.51" "0.31 - 0.82" "0.0057"
## [10,] ""
                11 11
## [11,] "0.46" "0.27 - 0.79" "0.0049"
## [12.] ""
                11 11
## [13,] "0.38" "0.22 - 0.66" "0.00054"
```

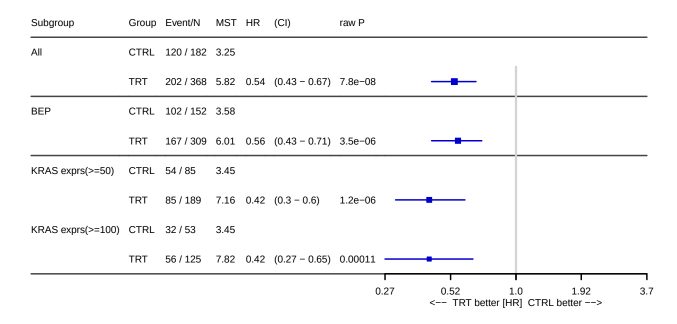
7.8 Survival outcome, 2-arm, continuous variable, greater than some numerical cutoffs

```
PlotTabForestBiomarker(data=input,

outcome.class="survival",
outcome.var=c("PFS","PFS.event"),
trt="Arm",
var="KRAS.exprs",
var.class="numeric", var.name="KRAS exprs",
percentile.cutoff=NULL,
numerical.cutoff=c(50,100),
greater=TRUE, less=FALSE,
within.bin=FALSE,
show.itt=TRUE, show.bep=TRUE)
```

Some NAs in var column, will define the non NA entries as BEP

Across-arm Effect of Biomarker PFS, KRAS exprs Unadjusted, unstratified analysis

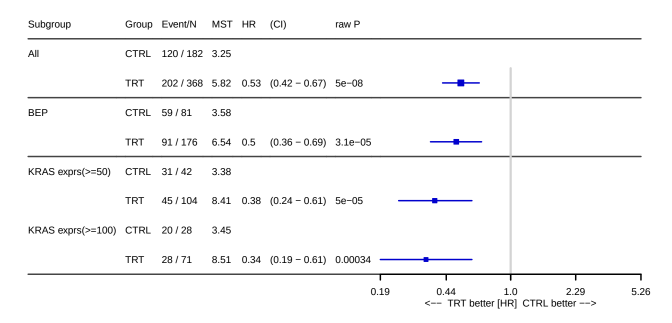


```
##
                              code.v
                                                                   "CI"
##
    [1,] "Subgroup"
                              "Group" "Event/N"
                                                   "MST"
                                                           "HR"
    [2,] "All"
                                                                   11 11
                              "CTRL"
                                       "120 / 182" "3.25" ""
##
    [3,] ""
##
                              "TRT"
                                       "202 / 368" "5.82" "0.54" "0.43 - 0.67"
    [4,] "BEP"
                              "CTRL" "102 / 152" "3.58" ""
##
##
    [5,] ""
                              "TRT"
                                       "167 / 309" "6.01" "0.56" "0.43 - 0.71"
                              "CTRL"
                                       "54 / 85"
                                                   "3.45" ""
    [6,] "KRAS exprs(>=50)"
##
    [7,] ""
                               "TRT"
                                       "85 / 189"
                                                   "7.16" "0.42" "0.3 - 0.6"
##
                                                   "3.45" ""
    [8,] "KRAS exprs(>=100)" "CTRL"
                                       "32 / 53"
##
    [9,] ""
                               "TRT"
                                       "56 / 125"
                                                   "7.82" "0.42" "0.27 - 0.65"
##
##
    [1,] "raw P"
##
   [2,] ""
##
   [3,] "7.8e-08"
##
    [4,] ""
##
##
    [5,] "3.5e-06"
    [6,] ""
##
##
    [7,] "1.2e-06"
    [8,] ""
##
    [9,] "0.00011"
##
```

7.9 Survival outcome, 2-arm, continuous variable, adjust for covariates

```
var="KRAS.exprs",
var.class="numeric", var.name="KRAS exprs",
percentile.cutoff=NULL,
numerical.cutoff=c(50,100),
greater=TRUE, less=FALSE,
within.bin=FALSE,
show.itt=TRUE, show.bep=TRUE,
bep = 'BEP', bep.indicator=1, covariate="Age")
```

Across-arm Effect of Biomarker PFS, KRAS exprs Results adjusted by Age;



```
##
                               code.v
                                                     "MST"
                                                                    "CI"
##
    [1,] "Subgroup"
                               "Group" "Event/N"
                                                            "HR"
    [2,] "All"
                               "CTRL"
                                        "120 / 182" "3.25" ""
                                                                    11 11
##
    [3,] ""
                               "TRT"
                                        "202 / 368" "5.82" "0.53" "0.42 - 0.67"
##
    [4,] "BEP"
                               "CTRL"
                                        "59 / 81"
                                                     "3.58" ""
                                                                    11 11
##
    [5,] ""
                                        "91 / 176"
                               "TRT"
                                                    "6.54" "0.5"
                                                                   "0.36 - 0.69"
##
    [6,] "KRAS exprs(>=50)"
##
                               "CTRL"
                                        "31 / 42"
                                                     "3.38" ""
                                                                    11 11
##
    [7,] ""
                               "TRT"
                                        "45 / 104"
                                                    "8.41" "0.38" "0.24 - 0.61"
    [8,] "KRAS exprs(>=100)" "CTRL"
                                        "20 / 28"
                                                     "3.45" ""
                                                                    11 11
##
                                        "28 / 71"
    [9,] ""
                                                     "8.51" "0.34" "0.19 - 0.61"
                               "TRT"
##
##
##
    [1,] "raw P"
##
    [2,] ""
##
    [3,] "5e-08"
    [4,] ""
##
##
   [5,] "3.1e-05"
    [6,] ""
##
##
    [7,] "5e-05"
##
    [8,] ""
```

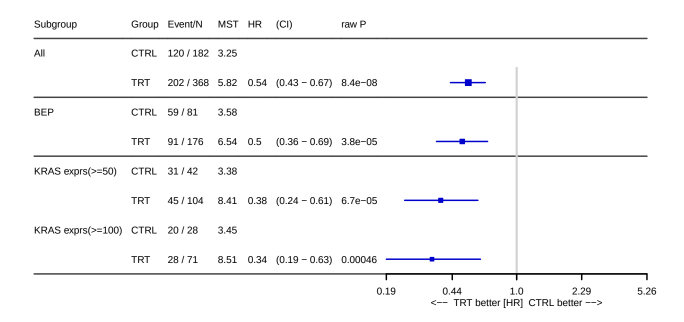
```
## [9,] "0.00034"
```

7.10 Survival outcome, 2-arm, continous variable, stratification

```
PlotTabForestBiomarker(data=input,

outcome.class="survival",
outcome.var=c("PFS","PFS.event"),
trt="Arm",
var="KRAS.exprs",
var.class="numeric", var.name="KRAS exprs",
percentile.cutoff=NULL,
numerical.cutoff=c(50,100),
greater=TRUE, less=FALSE,
within.bin=FALSE,
show.itt=TRUE, show.bep=TRUE,
bep = 'BEP', bep.indicator=1, strata="Sex")
```

Across-arm Effect of Biomarker PFS, KRAS exprs ;Results stratified by Sex



```
##
                              code.v
                              "Group" "Event/N"
    [1,] "Subgroup"
                                                    "MST"
                                                                   "CI"
##
                                                           "HR"
                              "CTRL" "120 / 182" "3.25" ""
   [2,] "All"
##
##
   [3,] ""
                              "TRT"
                                       "202 / 368" "5.82" "0.54" "0.43 - 0.67"
    [4,] "BEP"
                               "CTRL"
                                       "59 / 81"
                                                    "3.58" ""
                                                                   11 11
##
   [5,] ""
                                       "91 / 176"
                                                   "6.54" "0.5"
                                                                  "0.36 - 0.69"
##
                              "TRT"
                                                    "3.38" ""
                                                                   11 11
   [6,] "KRAS exprs(>=50)"
                                       "31 / 42"
##
                              "CTRL"
                                                    "8.41" "0.38" "0.24 - 0.61"
   [7,] ""
                              "TRT"
                                       "45 / 104"
##
                                                    "3.45" ""
                                                                   11 11
    [8,] "KRAS exprs(>=100)" "CTRL"
                                       "20 / 28"
##
   [9,] ""
##
                               "TRT"
                                       "28 / 71"
                                                   "8.51" "0.34" "0.19 - 0.63"
```

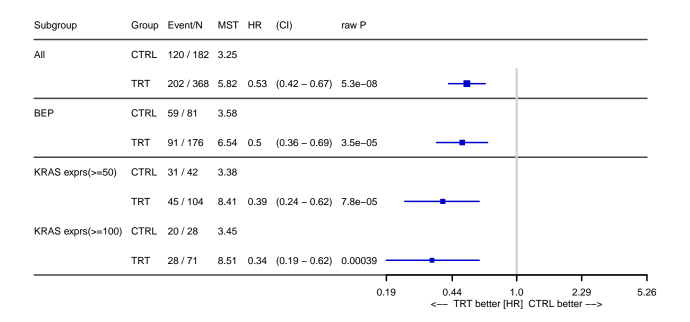
```
##
    [1,] "raw P"
##
    [2,] ""
##
   [3,] "8.4e-08"
##
    [4,] ""
##
##
   [5,] "3.8e-05"
##
   [6,] ""
   [7,] "6.7e-05"
##
##
   [8,] ""
   [9,] "0.00046"
```

7.11 Survival outcome, 2-arm, continuous variable, adjust for covariates and stratified by strata

```
PlotTabForestBiomarker(data=input,

outcome.class="survival",
outcome.var=c("PFS","PFS.event"),
trt="Arm",
var="KRAS.exprs",
var.class="numeric", var.name="KRAS exprs",
percentile.cutoff=NULL,
numerical.cutoff=c(50,100),
greater=TRUE, less=FALSE,
within.bin=FALSE,
show.itt=TRUE, show.bep=TRUE,
bep = 'BEP', bep.indicator=1,covariate="Age",strata="Sex")
```

Across-arm Effect of Biomarker PFS, KRAS exprs Results adjusted by Age;Results stratified by Sex



```
##
                              code.v
##
   [1,] "Subgroup"
                              "Group" "Event/N"
                                                   "MST" "HR"
                                                                  "CT"
                                                                  11 11
                              "CTRL"
                                      "120 / 182" "3.25" ""
##
   [2,] "All"
   [3,] ""
                              "TRT"
                                      "202 / 368" "5.82" "0.53" "0.42 - 0.67"
##
    [4,] "BEP"
                                                   "3.58" ""
                                                                  11 11
##
                              "CTRL"
                                      "59 / 81"
##
   [5,] ""
                              "TRT"
                                      "91 / 176"
                                                   "6.54" "0.5"
                                                                 "0.36 - 0.69"
   [6,] "KRAS exprs(>=50)"
                              "CTRL"
                                      "31 / 42"
                                                   "3.38" ""
                                                                  11 11
##
                                       "45 / 104"
    [7,] ""
                              "TRT"
                                                   "8.41" "0.39" "0.24 - 0.62"
##
##
    [8,] "KRAS exprs(>=100)" "CTRL"
                                      "20 / 28"
                                                   "3.45" ""
                                                                  11 11
   [9,] ""
##
                              "TRT"
                                      "28 / 71"
                                                   "8.51" "0.34" "0.19 - 0.62"
##
##
   [1,] "raw P"
   [2,] ""
##
##
   [3,] "5.3e-08"
  [4,] ""
##
##
   [5,] "3.5e-05"
##
  [6,] ""
## [7,] "7.8e-05"
## [8,] ""
## [9,] "0.00039"
```

7.12 Survival outcome, 1arm

within.bin, show.itt, show.bep will be ignored

Within-arm Effect of Biomarker PFS, KRAS exprs Results adjusted by Age; Results stratified by Sex

Subgroup	Group	Event/N	MST	HR	(CI)	rav	v P	
KRAS exprs(>=50)	Less	82 / 120	5.39					
	Greater	85 / 189	7.16	0.54	(0.4 – 0.7	4) 9.5	ie-05 -	-
KRAS exprs(>=100)	Less	111 / 184	5.59					
	Greater	56 / 125	7.82	0.57	(0.41 – 0.	78) 0.0	00062 -	-
							0.37	1.0 2.72 HR
##	s(>=50)"		"82 ' "85 "111	/ 184	"7.16" "5.59"	"0.54" ""	"CI" "0.4 - 0	

```
## [4,] KRAS exprs(>-100) Less 111 / 164 5.59

## [5,] "" "Greater" "56 / 125" "7.82" "0.57" "0.41 - 0.78"

## ## [1,] "raw P"

## [2,] ""

## [3,] "9.5e-05"

## [4,] ""

## [5,] "0.00062"
```

7.13 Survival outcome, 2arm, flip TRT/CTRL order (calculate ctrl over trt HR)

Across-arm Effect of Biomarker PFS, KRAS exprs Unadjusted, unstratified analysis

Subgroup	Group	Event/N	MST	HR	(CI)	raw P	
All	TRT	202 / 368	5.82				
	CTRL	120 / 182	3.25	1.87	(1.49 – 2.34)	7.8e-08	-
BEP	TRT	167 / 309	6.01				
	CTRL	102 / 152	3.58	1.8	(1.4 – 2.3)	3.5e-06	
KRAS exprs(>=25%, 23.26)	TRT	112 / 236	6.93				
	CTRL	73 / 111	3.38	2.23	(1.65 – 3)	1.5e-07	
KRAS exprs(>=50%, 70.52)	TRT	73 / 162	7.16				
	CTRL	43 / 70	3.58	2.37	(1.62 – 3.48)	9.6e-06	
KRAS exprs(>=75%, 177.29)	TRT	33 / 79	8.41				
	CTRL	22 / 37	4.73	2.64	(1.52 – 4.57)	0.00054	
						0.22	1.0 2.14 4.57 [HR] TRT better>

```
##
                                      code.v
##
    [1,] "Subgroup"
                                      "Group" "Event/N"
                                                           "MST" "HR"
##
    [2,] "All"
                                      "TRT"
                                               "202 / 368" "5.82" ""
##
    [3,] ""
                                      "CTRL"
                                               "120 / 182" "3.25" "1.87"
    [4,] "BEP"
                                      "TRT"
                                               "167 / 309" "6.01" ""
##
    [5,] ""
                                       "CTRL"
                                               "102 / 152" "3.58" "1.8"
##
                                               "112 / 236" "6.93" ""
    [6,] "KRAS exprs(>=25%, 23.26)"
                                      "TRT"
##
    [7,] ""
                                               "73 / 111"
##
                                      "CTRL"
                                                           "3.38" "2.23"
                                      "TRT"
                                               "73 / 162"
                                                           "7.16" ""
    [8,] "KRAS exprs(>=50%, 70.52)"
    [9,] ""
                                       "CTRL"
                                               "43 / 70"
                                                           "3.58" "2.37"
##
                                               "33 / 79"
                                                           "8.41" ""
##
   [10,] "KRAS exprs(>=75%, 177.29)" "TRT"
   [11,] ""
                                      "CTRL"
                                               "22 / 37"
                                                           "4.73" "2.64"
##
##
##
    [1,] "CI"
                        "raw P"
##
    [2,] ""
   [3,] "1.49 - 2.34" "7.8e-08"
##
   [4,] ""
                        11 11
##
                        "3.5e-06"
##
    [5,] "1.4 - 2.3"
##
    [6,] ""
                        11 11
   [7,] "1.65 - 3"
                        "1.5e-07"
##
   [8,] ""
##
                        11 11
   [9,] "1.62 - 3.48" "9.6e-06"
##
## [10,] ""
## [11,] "1.52 - 4.57" "0.00054"
```

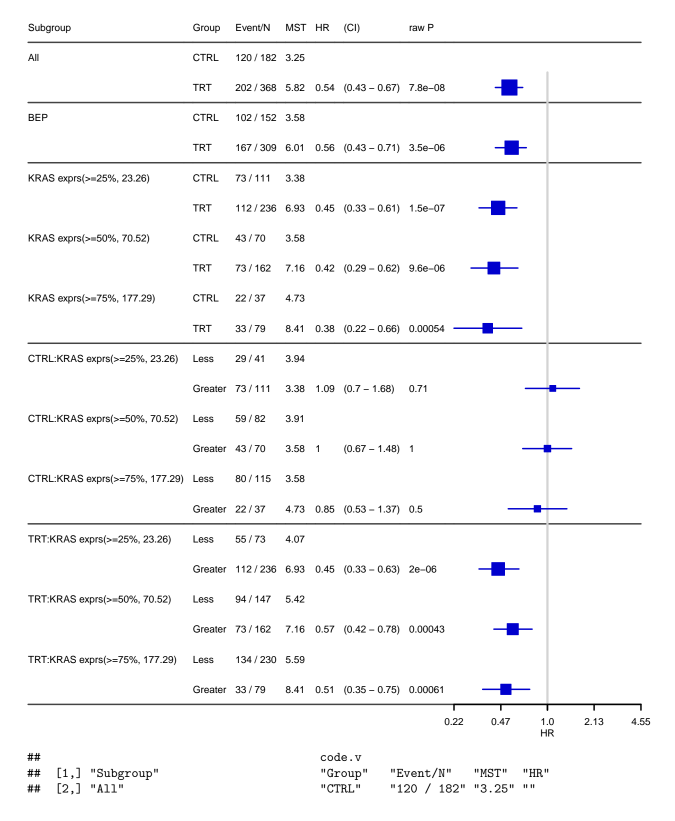
7.14 Survival outcome, 2-arm, continuous variable, greater than some percentage cutoffs, show across-arm and within-arm results in the same plot

```
PlotTabForestBiomarker(data=input,

outcome.class="survival",
outcome.var=c("PFS","PFS.event"),
trt="Arm",
var="KRAS.exprs",
var.class="numeric", var.name="KRAS exprs",
percentile.cutoff=c(.25,.5,.75),
numerical.cutoff=NULL,
greater=TRUE, less=FALSE,
within.bin=FALSE,
show.itt=TRUE, show.bep=TRUE,
across.and.within = TRUE)
```

Some NAs in var column, will define the non NA entries as BEP

Across-arm and Within-arm Effect of Biomarker PFS, KRAS exprs Unadjusted, unstratified analysis



```
[3,] ""
                                                      "202 / 368" "5.82" "0.54"
##
                                           "TRT"
##
    [4,] "BEP"
                                           "CTRL"
                                                      "102 / 152" "3.58" ""
    [5,] ""
                                           "TRT"
                                                      "167 / 309" "6.01" "0.56"
   [6,] "KRAS exprs(>=25%, 23.26)"
                                           "CTRL"
                                                      "73 / 111"
                                                                  "3.38" ""
##
##
    [7,] ""
                                           "TRT"
                                                      "112 / 236" "6.93" "0.45"
                                                                  "3.58" ""
   [8,] "KRAS exprs(>=50%, 70.52)"
                                           "CTRL"
                                                      "43 / 70"
##
                                           "TRT"
                                                      "73 / 162"
                                                                  "7.16" "0.42"
   [9.] ""
                                                      "22 / 37"
                                                                  "4.73" ""
## [10,] "KRAS exprs(>=75%, 177.29)"
                                           "CTRL"
## [11.] ""
                                           "TRT"
                                                      "33 / 79"
                                                                  "8.41" "0.38"
                                           "Less"
                                                      "29 / 41"
                                                                  "3.94" ""
## [12,] "CTRL:KRAS exprs(>=25%, 23.26)"
## [13,] ""
                                           "Greater" "73 / 111"
                                                                  "3.38" "1.09"
## [14,] "CTRL:KRAS exprs(>=50%, 70.52)"
                                                      "59 / 82"
                                                                  "3.91" ""
                                           "Less"
                                           "Greater" "43 / 70"
                                                                  "3.58" "1"
## [15,] ""
                                                                  "3.58" ""
                                           "Less"
                                                      "80 / 115"
## [16,] "CTRL:KRAS exprs(>=75%, 177.29)"
                                           "Greater" "22 / 37"
                                                                  "4.73" "0.85"
## [17,] ""
                                                      "55 / 73"
                                                                  "4.07" ""
## [18,] "TRT:KRAS exprs(>=25%, 23.26)"
                                           "Less"
## [19,] ""
                                           "Greater" "112 / 236" "6.93" "0.45"
                                                                  "5.42" ""
  [20,] "TRT:KRAS exprs(>=50%, 70.52)"
                                           "Less"
                                                      "94 / 147"
## [21,] ""
                                           "Greater" "73 / 162"
                                                                  "7.16" "0.57"
                                                      "134 / 230" "5.59" ""
  [22,] "TRT:KRAS exprs(>=75%, 177.29)"
                                           "Less"
## [23,] ""
                                           "Greater" "33 / 79"
                                                                  "8.41" "0.51"
##
##
    [1,] "CI"
                        "raw P"
    [2,] ""
##
   [3,] "0.43 - 0.67" "7.8e-08"
##
   [4,] ""
    [5,] "0.43 - 0.71" "3.5e-06"
##
    [6,] ""
##
   [7,] "0.33 - 0.61" "1.5e-07"
##
   [8,] ""
   [9,] "0.29 - 0.62" "9.6e-06"
##
## [10,] ""
## [11,] "0.22 - 0.66" "0.00054"
## [12,] ""
## [13,] "0.7 - 1.68"
                        "0.71"
## [14.] ""
## [15,] "0.67 - 1.48" "1"
## [16,] ""
## [17,] "0.53 - 1.37" "0.5"
## [18,] ""
## [19,] "0.33 - 0.63" "2e-06"
## [20,] ""
## [21,] "0.42 - 0.78" "0.00043"
## [22,] ""
## [23,] "0.35 - 0.75" "0.00061"
```

7.15 Response outcome, 2-arm, categorical variable

by default BEP will be defined as patients non-NA biomarker measurement

```
rsp.cat = TRUE,
rsp.response = c("CR","PR"),
rsp.nonresponse = c("SD", "PD","NON CR/PD","NE",NA),
trt="Arm",
var="KRAS.mutant",
var.class="categorical")
```

Covariate adjustment and stratification are not supported for binary outcome

Some NAs in var column, will define the non NA entries as BEP

Across-arm Effect of Biomarker Response, KRAS.mutant Unadjusted, unstratified analysis

Subgroup	Group	nRsp/N	Rsp Rate	deltaRR	(CI)	raw P	
All	CTRL	14 / 182	0.08				
	TRT	96 / 368	0.26	0.18	(0.12 – 0.25)	7e-07	-
BEP	CTRL	11 / 103	0.11				
	TRT	54 / 223	0.24	0.14	(0.05 – 0.22)	0.007	-
KRAS.mutant(Mutant)	CTRL	5 / 34	0.15				
	TRT	18 / 61	0.3	0.15	(-0.04 - 0.34)	0.17	-
KRAS.mutant(Wild Type)	CTRL	6 / 69	0.09				
	TRT	36 / 162	0.22	0.14	(0.03 – 0.24)	0.024	-
·							

<-- CTRL better [deltaRR] TRT * Unadjusted Interaction P =

0 0.17

-0.34

```
##
                                   code.v
##
   [1,] "Subgroup"
                                   "Group" "nRsp/N"
                                                       "Rsp Rate" "deltaRR"
##
   [2,] "All"
                                   "CTRL"
                                           "14 / 182" "0.08"
##
   [3,] ""
                                   "TRT"
                                           "96 / 368" "0.26"
                                                                  "0.18"
   [4,] "BEP"
                                   "CTRL"
                                           "11 / 103" "0.11"
                                                                  11 11
##
                                   "TRT"
                                                                  "0.14"
##
   [5,] ""
                                           "54 / 223" "0.24"
   [6,] "KRAS.mutant(Mutant)"
                                   "CTRL"
                                           "5 / 34"
                                                       "0.15"
##
##
    [7,] ""
                                   "TRT"
                                           "18 / 61"
                                                       "0.3"
                                                                  "0.15"
    [8,] "KRAS.mutant(Wild Type)" "CTRL"
                                           "6 / 69"
                                                       "0.09"
                                                                  11 11
##
                                   "TRT"
##
   [9,] ""
                                           "36 / 162" "0.22"
                                                                  "0.14"
##
##
   [1,] "CI"
                         "raw P"
##
   [2,] ""
   [3,] "0.12 - 0.25"
                        "7e-07"
##
   [4,] ""
##
                        "0.007"
##
   [5,] "0.05 - 0.22"
  [6,] ""
##
```

```
## [7,] "-0.04 - 0.34" "0.17"
## [8,] "" ""
## [9,] "0.03 - 0.24" "0.024"
```

7.16 Response outcome, another format

Covariate adjustment and stratification are not supported for binary outcome

Some NAs in var column, will define the non NA entries as BEP

Across-arm Effect of Biomarker Response, KRAS.mutant

Subgroup	Group	nRsp/N	Rsp Rate	deltaRR	CI	raw P		Fo	rest plo	t	
					_		CTRL	better		TRT better	
All	CTRL	14 / 182	0.08								
	TRT	96 / 368	0.26	0.18	0.12 - 0.25	7e-07				-	
BEP	CTRL	11 / 103	0.11			•					
	TRT	54 / 223	0.24	0.14	0.05 - 0.22	0.007			_	•	
KRAS.mutant(Mutant)	CTRL	5/34	0.15								
	TRT	18 / 61	0.3	0.15	-0.04 - 0.34	0.17				•	
KRAS.mutant(Wild Type)	CTRL	6 / 69	0.09								
	TRT	36 / 162	0.22	0.14	0.03 - 0.24	0.024			_	•	
* Unadjusted Interaction P = 0.77	Unadjus	ted, unstrati	ified analy	sis		-0.:	34 –0).17	0	0.17	0.34

```
##
                                   code.v
    [1,] "Subgroup"
                                   "Group" "nRsp/N"
                                                       "Rsp Rate" "deltaRR"
##
    [2,] "All"
                                   "CTRL"
                                            "14 / 182" "0.08"
##
   [3,] ""
                                   "TRT"
                                            "96 / 368" "0.26"
                                                                   "0.18"
##
                                                                   11 11
   [4,] "BEP"
                                   "CTRL"
                                            "11 / 103" "0.11"
                                                                   "0.14"
##
    [5,] ""
                                   "TRT"
                                            "54 / 223" "0.24"
    [6,] "KRAS.mutant(Mutant)"
                                   "CTRL"
                                            "5 / 34"
                                                       "0.15"
                                                                   11 11
##
   [7,] ""
                                   "TRT"
                                            "18 / 61" "0.3"
                                                                   "0.15"
##
                                                                   11 11
   [8,] "KRAS.mutant(Wild Type)" "CTRL"
                                            "6 / 69"
                                                       "0.09"
                                            "36 / 162" "0.22"
                                                                   "0.14"
   [9,] ""
                                   "TRT"
##
##
```

```
## [1,] "CI" "raw P"

## [2,] "" ""

## [3,] "0.12 - 0.25" "7e-07"

## [4,] "" ""

## [5,] "0.05 - 0.22" "0.007"

## [6,] "" ""

## [7,] "-0.04 - 0.34" "0.17"

## [8,] "" ""

## [9,] "0.03 - 0.24" "0.024"
```

7.17 Continuous outcome, 2-arm, categorical variable

Stratification is not supported for continuous outcome

Some NAs in var column, will define the non NA entries as BEP

Across-arm Effect of Biomarker Lab_ontrt, KRAS.mutant Unadjusted, unstratified analysis

Subgroup	Group	Mean	delta	(CI)	raw P
All	CTRL	19.22			
	TRT	22.17	2.95	(1.64 – 4.26)	1.2e-05
BEP	CTRL	19.67			
	TRT	22.42	2.75	(1.06 – 4.44)	0.0015
KRAS.mutant(Mutant)	CTRL	19.54			
	TRT	23.22	3.68	(0.56 - 6.8)	0.021
KRAS.mutant(Wild Type)	CTRL	19.73			
	TRT	22.11	2.38	(0.36 – 4.41)	0.021
##			code.ī	1	-6.8 -3.4 0 3.4 6.8 < CTRL better [delta] TRT better: * Unadjusted Interaction P = 0.48
## [1] Cubamaun				all IIMaanii II.	1-1+- CT

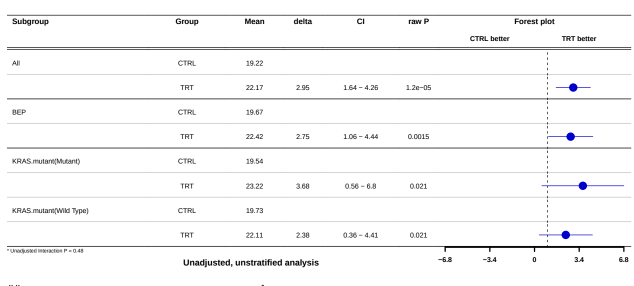
```
## [1,] "Subgroup" "Group" "Mean" "delta" "CI"
## [2,] "All" "CTRL" "19.22" "" ""
## [3,] "" "TRT" "22.17" "2.95" "1.64 - 4.26"
```

```
[4,] "BEP"
                                   "CTRL"
                                                             11 11
##
                                            "19.67" ""
    [5,] ""
##
                                   "TRT"
                                            "22.42" "2.75"
                                                             "1.06 - 4.44"
                                            "19.54" ""
##
    [6,] "KRAS.mutant(Mutant)"
                                   "CTRL"
   [7,] ""
                                    "TRT"
                                            "23.22" "3.68"
                                                             "0.56 - 6.8"
##
                                            "19.73" ""
                                                             11 11
##
    [8,] "KRAS.mutant(Wild Type)" "CTRL"
    [9,] ""
##
                                    "TRT"
                                            "22.11" "2.38"
                                                             "0.36 - 4.41"
##
   [1,] "raw P"
##
##
    [2,] ""
   [3,] "1.2e-05"
##
   [4,] ""
   [5,] "0.0015"
##
   [6,] ""
##
##
   [7,] "0.021"
  [8,] ""
##
##
   [9,] "0.021"
PlotTabForestBiomarker(data=input,
                                   outcome.class="continuous",
                                   outcome.var=c("Lab_ontrt"),
                                   trt="Arm",
                                   var="KRAS.mutant",
                                   var.class="categorical",tabforest=T)
```

Stratification is not supported for continuous outcome

Some NAs in var column, will define the non NA entries as BEP

Across-arm Effect of Biomarker Lab ontrt, KRAS.mutant



```
##
                                    code.v
   [1,] "Subgroup"
                                    "Group" "Mean" "delta" "CI"
##
                                            "19.22" ""
                                                             11 11
##
   [2,] "All"
                                    "CTRL"
   [3,] ""
                                            "22.17" "2.95"
##
                                    "TRT"
                                                             "1.64 - 4.26"
   [4,] "BEP"
                                            "19.67" ""
                                                             11 11
##
                                    "CTRL"
   [5,] ""
                                            "22.42" "2.75"
##
                                    "TRT"
                                                             "1.06 - 4.44"
##
   [6,] "KRAS.mutant(Mutant)"
                                    "CTRL"
                                            "19.54" ""
                                                             11 11
   [7,] ""
                                    "TRT"
                                            "23.22" "3.68"
                                                             "0.56 - 6.8"
##
```

```
[8,] "KRAS.mutant(Wild Type)" "CTRL" "19.73" ""
   [9,] ""
                                 "TRT"
                                         "22.11" "2.38" "0.36 - 4.41"
##
##
   [1,] "raw P"
##
   [2,] ""
##
##
  [3,] "1.2e-05"
  [4,] ""
   [5,] "0.0015"
##
##
   [6,] ""
##
  [7,] "0.021"
  [8,] ""
## [9,] "0.021"
```

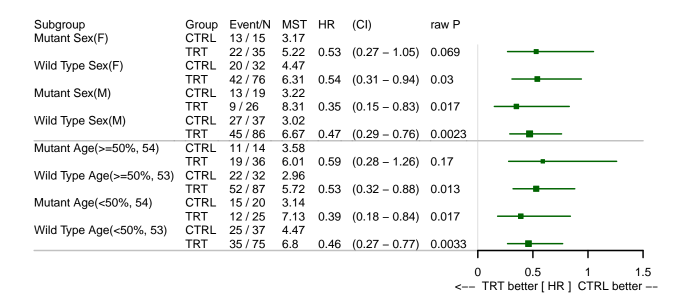
- 8 PlotTabForestMulti(): Forest plots to compare multiple variates' effect in ITT vs BEP, or compare multiple variates' effect in subpopulations
- 8.1 2-arm, compare BEP vs ITT

Across arm, Compare BEP vs. All PFS Unadjusted, unstratified analysis

Subgroup	Group	Event/N	MST	HR	(CI)	raw P	
ITT Sex(F)	CTRL	57 / 89	3.38				
	TRT	105 / 184	5.72	0.54	(0.39 – 0.74)	0.00018	-
BEP Sex(F)	CTRL	28 / 39	4.14				
	TRT	52 / 90	5.49	0.58	(0.36 – 0.92)	0.021	
ITT Sex(M)	CTRL	63 / 93	3.22				
	TRT	97 / 184	6.54	0.54	(0.39 – 0.74)	0.00012	-
BEP Sex(M)	CTRL	31 / 42	3.02				
	TRT	39 / 86	6.8	0.42	(0.26 – 0.68)	0.00043	
ITT Age(>=50%, 53)	CTRL	56 / 80	3.06				
	TRT	120 / 207	5.59	0.56	(0.41 – 0.77)	0.00035	-
BEP Age(>=50%, 54)	CTRL	26 / 36	3.38				
	TRT	53 / 93	5.72	0.63	(0.39 – 1.01)	0.053	
ITT Age(<50%, 53)	CTRL	64 / 102	3.94				
	TRT	82 / 161	6.8	0.48	(0.35 – 0.67)	1.9e-05	-
BEP Age(<50%, 54)	CTRL	33 / 45	3.91				
	TRT	38 / 83	6.8	0.37	(0.23 – 0.6)	4.8e-05	-
							0 0.5 1 - TRT better [HR] CTRL better>

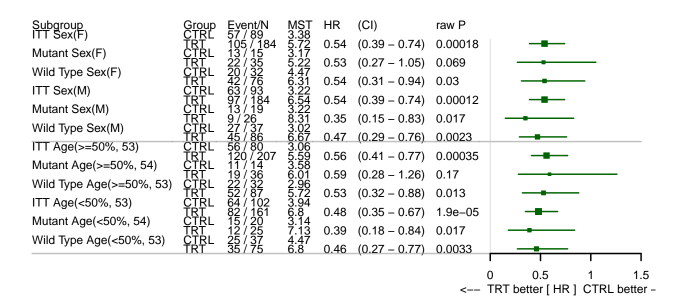
8.2 2-arm, compare subgroups defined by biomarker

Across arm, Compare KRAS.mutant subgroup PFS Unadjusted, unstratified analysis



8.3 2-arm, compare subgroups defined by biomarker, also show ITT

Across arm, Compare KRAS.mutant subgroup PFS Unadjusted, unstratified analysis



8.4 2-arm, compare subgroups defined by biomarker, also show BEP

Across arm, Compare KRAS.mutant subgroup PFS Unadjusted, unstratified analysis

Subgroup	Group	Event/N	MST	HR	(CI)	raw P	
ITT Sex(F)	CTRL	57 / 89	3.38				1
	TRT	105 / 184	5.72	0.54	(0.39 - 0.74)	0.00018	-
BEP Sex(F)	CTRL	28 / 39	4.14				
	TRT	52 / 90	5.49	0.58	(0.36 - 0.92)	0.021	-
Mutant Sex(F)	CTRL	13 / 15	3.17				
	TRT	22 / 35	5.22	0.53	(0.27 – 1.05)	0.069	
Wild Type Sex(F)	CTRL	20 / 32	4.47				
	TRT	42 / 76	6.31	0.54	(0.31 – 0.94)	0.03	_
ITT Sex(M)	CTRL	63 / 93	3.22				
	TRT	97 / 184	6.54	0.54	(0.39 - 0.74)	0.00012	-
BEP Sex(M)	CTRL	31 / 42	3.02				
	TRT	39 / 86	6.8	0.42	(0.26 - 0.68)	0.00043	-
Mutant Sex(M)	CTRL	13 / 19	3.22				
	TRT	9 / 26	8.31	0.35	(0.15 – 0.83)	0.017	-
Wild Type Sex(M)	CTRL	27 / 37	3.02				
	TRT	45 / 86	6.67	0.47	(0.29 - 0.76)	0.0023	-
ITT Age(>=50%, 53)	CTRL	56 / 80	3.06				
	TRT	120 / 207	5.59	0.56	(0.41 – 0.77)	0.00035	-
BEP Age(>=50%, 54)	CTRL	26 / 36	3.38				
	TRT	53 / 93	5.72	0.63	(0.39 – 1.01)	0.053	
Mutant Age(>=50%, 54)	CTRL	11 / 14	3.58				
	TRT	19 / 36	6.01	0.59	(0.28 – 1.26)	0.17	
Wild Type Age(>=50%, 53)	CTRL	22 / 32	2.96				
	TRT	52 / 87	5.72	0.53	(0.32 - 0.88)	0.013	_
ITT Age(<50%, 53)	CTRL	64 / 102	3.94				
	TRT	82 / 161	6.8	0.48	(0.35 - 0.67)	1.9e-05	-
BEP Age(<50%, 54)	CTRL	33 / 45	3.91				
	TRT	38 / 83	6.8	0.37	(0.23 - 0.6)	4.8e-05	-
Mutant Age(<50%, 54)	CTRL	15 / 20	3.14				
	TRT	12 / 25	7.13	0.39	(0.18 - 0.84)	0.017	
Wild Type Age(<50%, 53)	CTRL	25 / 37	4.47		,		
	TRT	35 / 75	6.8	0.46	(0.27 - 0.77)	0.0033	_
					. ,		
							0 0.5 1 1.5 TRT better [HR] CTRL better -
						_ _	The better [This] OTHE better -

If bep column is not defined, the program will take the non NA entries in subgroups column as BEP

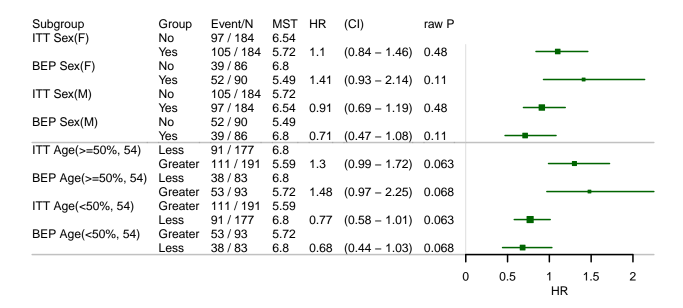
show.bep is TRUE but bep is not specified, will define the non NA entries in subgroup column as BEP

Across arm, Compare KRAS.mutant subgroup PFS Unadjusted, unstratified analysis

Subgroup	-	Event/N	MST	HR	(CI)	raw P	
ITT Sex(F)	CTRL	57 / 89	3.38	0.54	(0.00 0.74)	0.00040	
DED 0 (E)	TRT	105 / 184	5.72	0.54	(0.39 - 0.74)	0.00018	-
BEP Sex(F)	CTRL	33 / 47	4.11	0.50	(0.05 0.04)	0.0007	_
	TRT	64 / 111	5.59	0.53	(0.35 - 0.81)	0.0037	
Mutant Sex(F)	CTRL	13 / 15	3.17		(0.07		_
	TRT	22 / 35	5.22	0.53	(0.27 - 1.05)	0.069	
Wild Type Sex(F)	CTRL	20 / 32	4.47				_
	TRT	42 / 76	6.31	0.54	(0.31 - 0.94)	0.03	
ITT Sex(M)	CTRL	63 / 93	3.22				_
	TRT	97 / 184	6.54	0.54	(0.39 - 0.74)	0.00012	-
BEP Sex(M)	CTRL	40 / 56	3.06				_
	TRT	54 / 112	6.8	0.46	(0.3 - 0.7)	0.00024	-
Mutant Sex(M)	CTRL	13 / 19	3.22				
	TRT	9 / 26	8.31	0.35	(0.15 - 0.83)	0.017	
Wild Type Sex(M)	CTRL	27 / 37	3.02				
	TRT	45 / 86	6.67	0.47	(0.29 - 0.76)	0.0023	
ITT Age(>=50%, 53)	CTRL	56 / 80	3.06				
	TRT	120 / 207	5.59	0.56	(0.41 - 0.77)	0.00035	-
BEP Age(>=50%, 53)	CTRL	34 / 47	3.02				
	TRT	71 / 123	6.01	0.52	(0.34 - 0.79)	0.0019	-
Mutant Age(>=50%, 54)	CTRL	11 / 14	3.58				
	TRT	19 / 36	6.01	0.59	(0.28 - 1.26)	0.17	
Wild Type Age(>=50%, 53)	CTRL	22 / 32	2.96				
	TRT	52 / 87	5.72	0.53	(0.32 - 0.88)	0.013	-
ITT Age(<50%, 53)	CTRL	64 / 102	3.94				
	TRT	82 / 161	6.8	0.48	(0.35 - 0.67)	1.9e-05	-
BEP Age(<50%, 53)	CTRL	39 / 56	4.11				
	TRT	47 / 100	6.8	0.44	(0.29 - 0.67)	0.00017	-
Mutant Age(<50%, 54)	CTRL	15 / 20	3.14				
	TRT	12 / 25	7.13	0.39	(0.18 - 0.84)	0.017	
Wild Type Age(<50%, 53)	CTRL	25 / 37	4.47				
	TRT	35 / 75	6.8	0.46	(0.27 – 0.77)	0.0033	_
							0 0.5 1 1.5 TRT better [HR] CTRL better -

8.5 1-arm, compare ITT vs BEP

Within arm, Compare BEP vs. All PFS
Unadjusted, unstratified analysis



8.6 2-arm, compare subgroups defined by biomarker, multiple cutoffs for continuous biomarker

```
PlotTabForestMulti(data=input,

outcome.class="survival",
outcome.var=c("PFS","PFS.event"),
trt="Arm", percentile.cutoff=c(.33,.66),
var=c("Sex","Age"),
bep="BEP",bep.indicator=1, compare.bep.itt=FALSE,
compare.subgroup=TRUE, subgroup="KRAS.mutant"
)
```

Across arm, Compare KRAS.mutant subgroup PFS Unadjusted, unstratified analysis

Subgroup	Group	Event/N	MST	HR	(CI)	raw P	
Mutant Sex(F)	CTRL	13 / 15	3.17				
	TRT	22 / 35	5.22	0.53	(0.27 – 1.05)	0.069	_
Wild Type Sex(F)	CTRL	20 / 32	4.47				
	TRT	42 / 76	6.31	0.54	(0.31 – 0.94)	0.03	-
Mutant Sex(M)	CTRL	13 / 19	3.22				
	TRT	9 / 26	8.31	0.35	(0.15 – 0.83)	0.017	-
Wild Type Sex(M)	CTRL	27 / 37	3.02				
	TRT	45 / 86	6.67	0.47	(0.29 – 0.76)	0.0023	-
Mutant Age(>=33%, 49)	CTRL	19 / 22	3.06				
	TRT	23 / 42	5.59	0.53	(0.29 – 0.98)	0.043	-
Wild Type Age(>=33%, 48)	CTRL	30 / 44	3.94				
	TRT	61 / 111	6.31	0.49	(0.31 – 0.76)	0.0017	-
Mutant Age(>=66%, 59)	CTRL	7/9	5.78				
	TRT	13 / 25	6.18	0.87	(0.32 – 2.38)	0.79	
Wild Type Age(>=66%, 58)	CTRL	16 / 25	3.38				
	TRT	35 / 57	5.49	0.63	(0.35 – 1.16)	0.14	-
Mutant Age(<33%, 49)	CTRL	7 / 12	3.56				
	TRT	8 / 19	8.31	0.4	(0.15 – 1.13)	0.083	-
Wild Type Age(<33%, 48)	CTRL	17 / 25	4.37				
	TRT	26 / 51	6.8	0.55	(0.29 – 1.02)	0.057	-
Mutant Age(<66%, 59)	CTRL	19 / 25	3.14				
	TRT	18 / 36	5.72	0.4	(0.21 – 0.76)	0.0052	-
Wild Type Age(<66%, 58)	CTRL	31 / 44	4.37				
	TRT	52 / 105	6.8	0.43	(0.27 – 0.68)	0.00028	
							0 0.5 1 1.5 2 2.5 TRT better [HR] CTRL better

8.7 2-arm, compare subgroups defined by biomarker, multiple cutoffs for continuous biomarker, calculate "< cutoff"

Across arm, Compare KRAS.mutant subgroup PFS Unadjusted, unstratified analysis

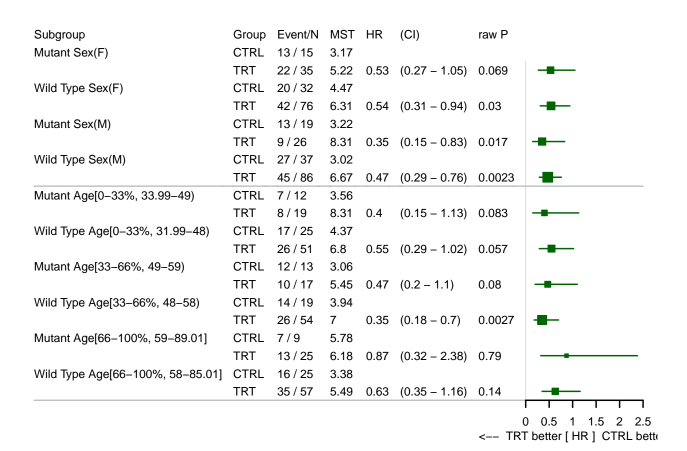
Subgroup	Group	Event/N	MST	HR	(CI)	raw P	
Mutant Sex(F)	CTRL	13 / 15	3.17				
	TRT	22 / 35	5.22	0.53	(0.27 – 1.05)	0.069	_
Wild Type Sex(F)	CTRL	20 / 32	4.47				
	TRT	42 / 76	6.31	0.54	(0.31 – 0.94)	0.03	-
Mutant Sex(M)	CTRL	13 / 19	3.22				
	TRT	9 / 26	8.31	0.35	(0.15 – 0.83)	0.017	-
Wild Type Sex(M)	CTRL	27 / 37	3.02				
	TRT	45 / 86	6.67	0.47	(0.29 – 0.76)	0.0023	-
Mutant Age(>=33%, 49)	CTRL	19 / 22	3.06				
	TRT	23 / 42	5.59	0.53	(0.29 – 0.98)	0.043	-
Wild Type Age(>=33%, 48)	CTRL	30 / 44	3.94				
	TRT	61 / 111	6.31	0.49	(0.31 – 0.76)	0.0017	-
Mutant Age(>=66%, 59)	CTRL	7/9	5.78				
	TRT	13 / 25	6.18	0.87	(0.32 – 2.38)	0.79	
Wild Type Age(>=66%, 58)	CTRL	16 / 25	3.38				
	TRT	35 / 57	5.49	0.63	(0.35 – 1.16)	0.14	-
Mutant Age(<33%, 49)	CTRL	7 / 12	3.56				
	TRT	8 / 19	8.31	0.4	(0.15 – 1.13)	0.083	-
Wild Type Age(<33%, 48)	CTRL	17 / 25	4.37				
	TRT	26 / 51	6.8	0.55	(0.29 – 1.02)	0.057	-
Mutant Age(<66%, 59)	CTRL	19 / 25	3.14				
	TRT	18 / 36	5.72	0.4	(0.21 – 0.76)	0.0052	-
Wild Type Age(<66%, 58)	CTRL	31 / 44	4.37				
	TRT	52 / 105	6.8	0.43	(0.27 – 0.68)	0.00028	
							0 0.5 1 1.5 2 2.5 TRT better [HR] CTRL better

8.8 2-arm, compare subgroups defined by biomarker, multiple cutoffs for continuous biomarker, calculate within bin statistics

Across arm, Compare KRAS.mutant subgroup PFS

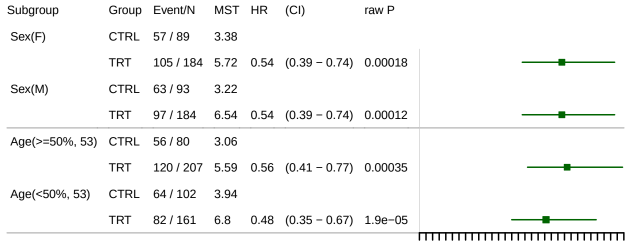
within.bin is TRUE, greater and less will be ignored
within.bin is TRUE, greater and less will be ignored

Unadjusted, unstratified analysis



8.9 Basic forest plot without comparison

Across arm PFS Unadjusted, unstratified analysis



00.075 0.2 0.3 0.4 0.5 0.6 0.7 <-- TRT better [HR] CTRL better -->

8.10 Response outcome

Covariate adjustment and stratification are not supported for binary outcome ## Covariate adjustment and stratification are not supported for binary outcome

Across arm Response Unadjusted, unstratified analysis

Subgroup	Group	nRsp/N	Rsp Rate	deltaRR	(CI)	raw P	
Sex(F)	CTRL	9 / 89	0.1				
	TRT	45 / 184	0.24	0.14	(0.05 – 0.24)	0.0086	
Sex(M)	CTRL	5/93	0.05				
	TRT	51 / 184	0.28	0.22	(0.14 – 0.31)	2.5e-05	
Age(>=50%, 53)	CTRL	6 / 80	0.08				
	TRT	50 / 207	0.24	0.17	(0.08 – 0.26)	0.0025	
Age(<50%, 53)	CTRL	8 / 102	0.08				
	TRT	46 / 161	0.29	0.21	(0.11 – 0.3)	9.7e-05	

0.044 0.088 0.177 0.354 0.707 <-- CTRL better [deltaRR] TRT better

- ## Covariate adjustment and stratification are not supported for binary outcome ## Covariate adjustment and stratification are not supported for binary outcome
- ## Warning in log(clip): NaNs produced

Across arm Response

Subgroup	Group	nRsp/N	Rsp Rate	deltaRR	CI	raw P	Fores	st plot
							TRT better	CTRL better
Sex(F)	CTRL	9 / 89	0.1					
	TRT	45 / 184	0.24	0.14	0.05 – 0.24	0.0086 —		•
Sex(M)	CTRL	5 / 93	0.05					
	TRT	51 / 184	0.28	0.22	0.14 – 0.31	2.5e-05		
Age(>=50%, 53)	CTRL	6 / 80	0.08					
	TRT	50 / 207	0.24	0.17	0.08 - 0.26	0.0025		•
Age(<50%, 53)	CTRL	8 / 102	0.08					
	TRT	46 / 161	0.29	0.21	0.11 – 0.3	9.7e-05	_	•
		Unadjusted,	unstratified	d analysis			0.1	0.2

8.11 Continuous endpoint

Stratification is not supported for continuous outcome
Stratification is not supported for continuous outcome

Across arm Lab_ontrt Unadjusted, unstratified analysis

Subgroup	Group	Mean	delta	(CI)	raw P		
Sex(F)	CTRL	18.86					
	TRT	22.11	3.25	(1.38 – 5.11)	0.00073		
Sex(M)	CTRL	19.57					
	TRT	22.24	2.67	(0.82 – 4.51)	0.0048		
Age(>=50%, 53)	CTRL	19.1					
	TRT	22.05	2.95	(1.07 – 4.83)	0.0022	-	-
Age(<50%, 53)	CTRL	19.32					
	TRT	22.33	3.01	(1.14 – 4.87)	0.0017	-	
							\neg
						0 0.5 1 1.5 2 2.5 3 3.5 4 4.5 TRT better [delta] CTRL better>	_

 $\ensuremath{\mbox{\#\#}}$ Stratification is not supported for continuous outcome

Stratification is not supported for continuous outcome

Across arm Lab ontrt

Subgroup	Group	Mean	delta	CI	raw P	Forest plot	
						TRT be@aiRL bette	
Sex(F)	CTRL	18.86	·	1		 	
	TRT	22.11	3.25	1.38 – 5.11	0.00073	•	
Sex(M)	CTRL	19.57					
	TRT	22.24	2.67	0.82 – 4.51	0.0048	•	
Age(>=50%, 53)	CTRL	19.1					
	TRT	22.05	2.95	1.07 – 4.83	0.0022	•	
Age(<50%, 53)	CTRL	19.32					
	TRT	22.33	3.01	1.14 – 4.87	0.0017	•	
	Unadjusted, unstr	atified analysis				-4 0 2 4	

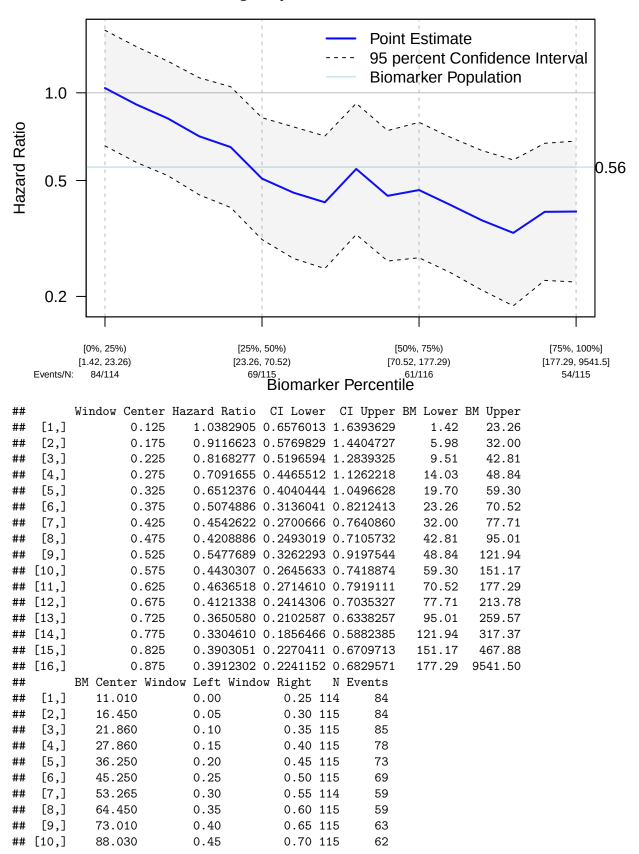
PlotSTEPP(): STEPP (Subpopulation Treatment Effect Pattern Plot) from the given point estimates and confidence intervals at desired percentiles.

9.1 Survival outcome

```
PlotSTEPP(data = input,
          outcome.var = c("PFS", "PFS.event"),
          outcome.class = "survival",
          trt = "Arm",
          var = "KRAS.exprs",
          placebo.code = "CTRL",
          active.code = "TRT",
          csv.name = NULL,
          pdf.name = NULL
```

some NA in var column, will ignore NA entries



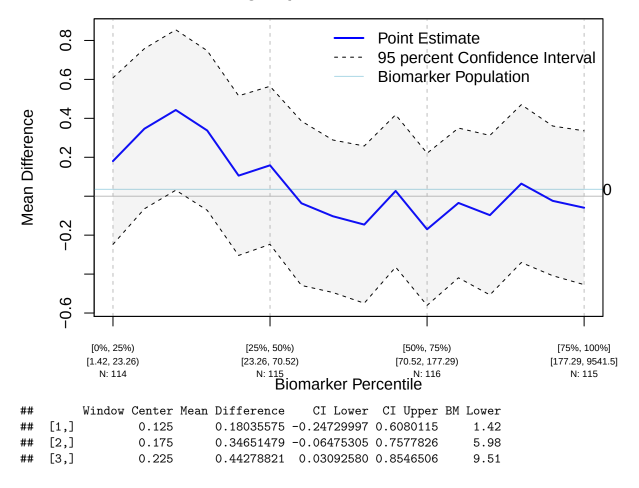


```
## [11,]
                           0.50
                                         0.75 116
           101.480
                                                       61
## [12,]
           133.900
                           0.55
                                         0.80 116
                                                       61
                                         0.85 115
## [13,]
           165.420
                           0.60
                                                       55
## [14,]
           195.360
                           0.65
                                         0.90 115
                                                       51
## [15,]
           238.860
                           0.70
                                         0.95 115
                                                       56
## [16,]
           292.040
                           0.75
                                         1.00 115
                                                       54
```

9.2 Continuous outcome

some NA in var column, will ignore NA entries

STEPP: Subgroup Treatment Effect Pattern Plot

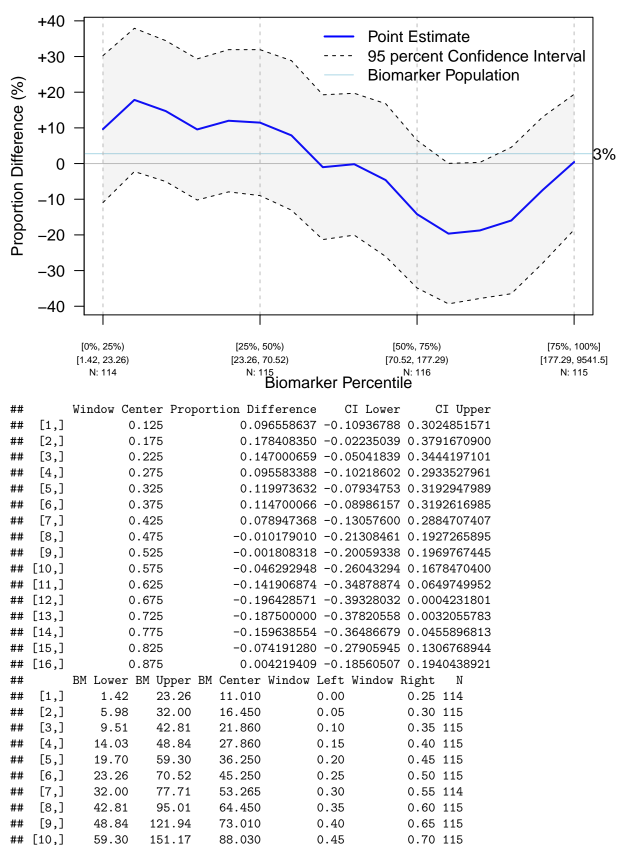


```
[4,]
##
                 0.275
                             0.33749063 -0.07236992 0.7473512
                                                                  14.03
##
   [5,]
                 0.325
                             0.10572728 -0.30412158 0.5155761
                                                                  19.70
  [6,]
##
                 0.375
                             0.15866042 -0.24724256 0.5645634
                                                                  23.26
## [7,]
                 0.425
                            -0.03630432 -0.45831952 0.3857109
                                                                  32.00
##
   [8,]
                 0.475
                            -0.10286329 -0.49436652 0.2886399
                                                                  42.81
## [9,]
                            -0.14583073 -0.54953031 0.2578689
                                                                  48.84
                 0.525
## [10,]
                             0.02704796 -0.36419706 0.4182930
                                                                  59.30
                 0.575
## [11,]
                            -0.16994763 -0.56071364 0.2208184
                 0.625
                                                                  70.52
## [12,]
                 0.675
                            -0.03456171 -0.41907553 0.3499521
                                                                  77.71
## [13,]
                            -0.09734050 -0.50684711 0.3121661
                 0.725
                                                                  95.01
## [14,]
                 0.775
                             0.06465414 -0.34072771 0.4700360
                                                                 121.94
                            -0.02383926 -0.40821143 0.3605329
## [15,]
                 0.825
                                                                 151.17
                            -0.05926090 -0.45420790 0.3356861
## [16,]
                 0.875
                                                                 177.29
##
         BM Upper BM Center Window Left Window Right
##
   [1,]
            23.26
                     11.010
                                    0.00
                                                 0.25 114
##
   [2,]
            32.00
                     16.450
                                    0.05
                                                 0.30 115
## [3,]
            42.81
                     21.860
                                    0.10
                                                 0.35 115
## [4,]
            48.84
                     27.860
                                    0.15
                                                 0.40 115
                                                 0.45 115
## [5,]
            59.30
                     36.250
                                    0.20
## [6,]
            70.52
                     45.250
                                    0.25
                                                 0.50 115
## [7,]
            77.71
                     53.265
                                    0.30
                                                 0.55 114
## [8,]
            95.01
                     64.450
                                    0.35
                                                 0.60 115
## [9,]
           121.94
                     73.010
                                    0.40
                                                 0.65 115
## [10,]
           151.17
                     88.030
                                    0.45
                                                 0.70 115
## [11,]
                                                 0.75 116
           177.29
                    101.480
                                    0.50
## [12,]
           213.78
                    133.900
                                    0.55
                                                 0.80 116
## [13,]
           259.57
                    165.420
                                    0.60
                                                 0.85 115
## [14,]
           317.37
                    195.360
                                    0.65
                                                 0.90 115
## [15,]
           467.88
                    238.860
                                    0.70
                                                 0.95 115
## [16,]
          9541.50
                    292.040
                                    0.75
                                                 1.00 115
```

9.3 Binary outcome variable

some NA in var column, will ignore NA entries





```
## [11,]
           70.52
                   177.29
                                          0.50
                                                      0.75 116
                           101.480
                                                      0.80 116
## [12,]
           77.71
                   213.78
                           133.900
                                          0.55
## [13,]
          95.01
                                          0.60
                                                      0.85 115
                   259.57
                           165.420
## [14,]
          121.94
                   317.37
                           195.360
                                          0.65
                                                      0.90 115
## [15,]
          151.17
                   467.88
                           238.860
                                          0.70
                                                      0.95 115
## [16,]
          177.29 9541.50
                           292.040
                                          0.75
                                                      1.00 115
```

10 CoxTab(): fit cox proportional model for multiple covariates (additive model or separate models)

10.1 single covariate

```
CoxTab(data=sample.data, tte="OS", cens="OS.event",bep='BEP', var='Sex')

## HR CI.low CI.high p-value n.trt n.ref

## Sex (M/F) "0.85" "0.42" "1.72" "0.65" "128" "129"
```

10.2 multiple covariates, ITT

Additive model will be fitted.

	HR	CI.low	CI.high	p-value	n.trt	n.ref
Sex (M/F)	0.72	0.48	1.08	0.11	277	273
Country (USA/Other)	0.65	0.36	1.2	0.17	127	198
Country (Western Europe/Other)	0.77	0.5	1.2	0.24	225	198
Age	1.01	0.99	1.03	0.43		

Simplify: automatically learn var.class from column class:

```
kable(
CoxTab(data=sample.data,tte="OS", cens="OS.event", var=c('Sex',"Country","Age"))
)
```

	HR	CI.low	CI.high	p-value	n.trt	n.ref
Sex (M/F)	0.72	0.48	1.08	0.11	277	273
Country (USA/Other)	0.65	0.36	1.2	0.17	127	198
Country (Western Europe/Other)	0.77	0.5	1.2	0.24	225	198
Age	1.01	0.99	1.03	0.43		

10.3 BEP

	HR	CI.low	CI.high	p-value	n.trt	n.ref
Sex (M/F)	0.85	0.42	1.73	0.65	128	129
Country (USA/Other)	1.17	0.4	3.38	0.77	76	57
Country (Western Europe/Other)	1.11	0.44	2.83	0.83	124	57
Age	1.01	0.98	1.04	0.48		

10.4 Reorder Sex: M as reference

	HR	CI.low	CI.high	p-value	n.trt	n.ref
Sex (F/M)	1.12	0.68	1.85	0.65	129	128
Country (USA/Other)	1.17	0.4	3.38	0.77	76	57
Country (Western Europe/Other)	1.11	0.44	2.83	0.83	124	57
Age	1.01	0.98	1.04	0.48		

10.5 Fit separate models for each variable instead of fitting an additive model

	HR	CI.low	CI.high	p-value	n.trt	n.ref
Sex (M/F)	0.71	0.48	1.07	0.099	277	273
Country (USA/Other)	0.67	0.37	1.22	0.19	127	198
Country (Western Europe/Other)	0.77	0.5	1.2	0.25	225	198
Age	1.01	0.99	1.02	0.54		

11 StatSummary()

11.1 Contunuous outcome

11.1.1 Create summary statistics

11.1.2 Add a covariate variable

11.1.3 Add return.fit = TRUE to return a table of summary statistics

```
StatSummary(outcome.var = input$OS, treatment.var = input$Arm,
            placebo.code = "CTRL", active.code = "TRT",
            outcome.class = "continuous", covariate.var = input$Sex,
           return.fit = TRUE)
## [[1]]
## Effect.Size
                       Lower
                                    Upper
                                                     P Mean.Placebo
## 0.795751256 0.199865806 1.391636707 0.008954882 5.563721382
## Mean.Active
## 6.359472638
##
## [[2]]
##
## Call:
## lm(formula = outcome.var[subgroup.var] ~ covariate.var[subgroup.var] +
      treatment.var[subgroup.var])
##
##
## Coefficients:
##
                      (Intercept)
                                     covariate.var[subgroup.var]M
##
                                                           0.4204
                           5.5637
## treatment.var[subgroup.var]TRT
                           0.7958
##
```

11.2 Binary outcome

11.2.1 Create summary statistics

```
StatSummary(outcome.var = input$OS.event, treatment.var = input$Arm,
            placebo.code = "CTRL", active.code = "TRT",
            outcome.class = "binary")
## Effect.Size
                      Lower
                                   Upper
                                                     Ρ
                                                       Rsp.Placebo
   -0.04846512 -0.12251455
##
                              0.02558430
                                           0.19901328
                                                         0.20879121
                  N.Placebo
                                N.Active nRsp.Placebo nRsp.Active
##
    Rsp.Active
    0.16032609 182.00000000 368.00000000 38.00000000 59.00000000
##
```

11.3 Survival variable

11.3.1 Create summary statistics

```
StatSummary(outcome.var = input[, c("OS", "OS.event")],
            treatment.var = input$Arm, placebo.code = "CTRL",
            active.code = "TRT", outcome.class = "survival")
## CTRL.events
                   CTRL.n
                             CTRL.MST TRT.events
                                                        TRT.n
                                                                  TRT.MST
## 38.0000000 182.0000000 14.8829569 59.0000000 368.0000000
## Effect.Size
                    Lower
                                Upper
    0.6667451
                0.4432032
                            1.0030365
                                       0.0517253
```

11.3.2 Add a covariate variable

```
StatSummary(outcome.var = input[, c("OS", "OS.event")],
            treatment.var = input$Arm, placebo.code = "CTRL",
            active.code = "TRT", outcome.class = "survival",
            covariate.var = input$Sex)
## CTRL.events
                      CTRL.n
                                 CTRL.MST
                                            TRT.events
                                                              TRT.n
##
   38.00000000 182.00000000 14.88295688
                                           59.0000000 368.0000000
##
        TRT.MST Effect.Size
                                    Lower
                                                 Upper
                 0.63926120
##
             NA
                               0.42390908
                                            0.96401541
                                                         0.03277593
```

11.3.3 Add a stratification variable

```
StatSummary(outcome.var = input[, c("OS", "OS.event")],
            treatment.var = input$Arm, placebo.code = "CTRL",
            active.code = "TRT", outcome.class = "survival",
            covariate.var = input$Sex, strat.factor.var = input$Age)
   CTRL.events
                      CTRL.n
                                 CTRL.MST
                                            TRT.events
                                                               TRT.n
##
   38.00000000 182.00000000 14.88295688
                                           59.00000000 368.00000000
        TRT.MST Effect.Size
##
                                    Lower
                                                 Upper
##
                  0.60116855
                               0.37830923
                                            0.95531273
                                                         0.03128551
            NA
```

12 LogRankTab(): log rank test for subgroup analysis

```
kable(
   LogRankTab(data=input,tte="PFS",cens="PFS.event",var="Arm")
)
```

	CTRL	TRT
	N=182	N = 368
Patients with event	120~(65.9%)	202 (54.9%)
Patients without event	62 (34.1%)	166 (45.1%)
Time to event (month)		
Median (KM)	3.25	5.82
95% CI Median	(2.83;4.14)	(5.49;6.8)
25% and $75%$ -ile	1.61;6.47	2.89;11.73
Range (inc. cens.)	0.03 to 12.32	0.03 to 13.8
p-value (Log-Rank Test)		0
Hazard Ratio		0.54
95% CI		(0.43;0.67)

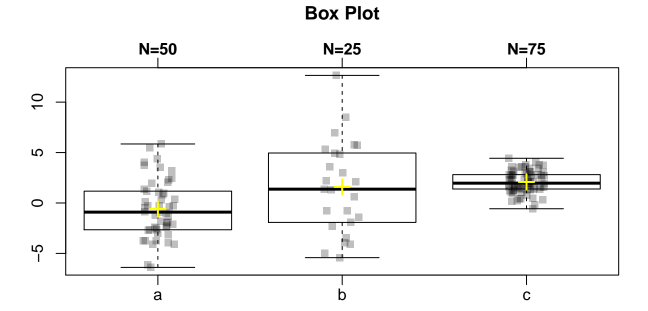
13 BoxPlot(): advanced box plot function

13.1 Generate example dataset

```
example \leftarrow data.frame( y=c(rnorm(30)+10, rnorm(4)+20, rnorm(15)+15, NA),
                           time=c(rep("t2", 30), rep("t4",4), rep("t1", 15), "t3"),
                           grp=sample(1:3, 50, TRUE), sex=sample(1:2, 50, TRUE))
head(example)
            y time grp sex
## 1 9.453643
               t2
## 2 9.176470
                t2
                         1
               t2 2
                         2
## 3 10.103796
## 4 8.487022
                t2 2
                         1
## 5 9.599060
                t2 3
                         1
                t2
## 6 9.589478
                         1
str(example)
## 'data.frame':
                   50 obs. of 4 variables:
## $ y : num 9.45 9.18 10.1 8.49 9.6 ...
## $ time: Factor w/ 4 levels "t1","t2","t3",...: 2 2 2 2 2 2 2 2 2 2 ...
## $ grp : int 3 3 2 2 3 3 2 3 3 2 ...
## $ sex : int 2 1 2 1 1 1 1 2 2 2 ...
```

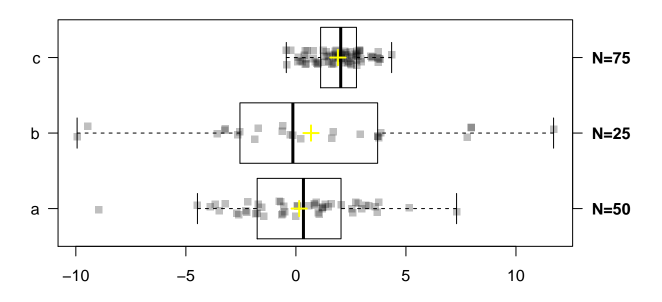
13.2 Specify data as named list

```
BoxPlot(list(a=rnorm(50,,3), b=rnorm(25,1,4), c=rnorm(75,2,1)))
```



13.3 Same plot, now horizontally plotted

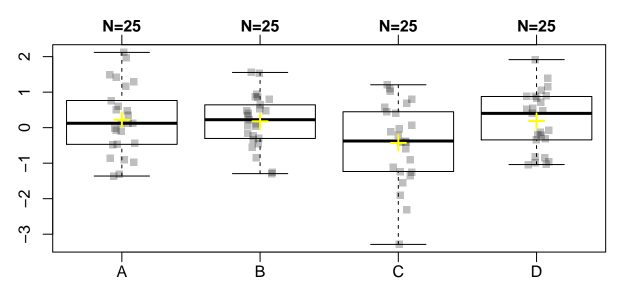
Box Plot



13.4 Specify data as numeric matrix

```
BoxPlot(matrix(rnorm(100), ncol=4, dimnames=list(NULL, LETTERS[1:4])) )
```

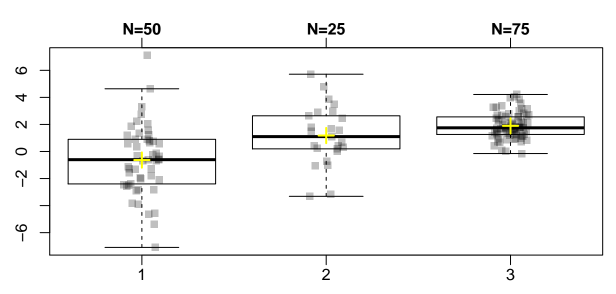
Box Plot



13.5 Specify data as numeric (unnamed) vectors

BoxPlot(rnorm(50,,3), rnorm(25,1,2), rnorm(75,2,1))

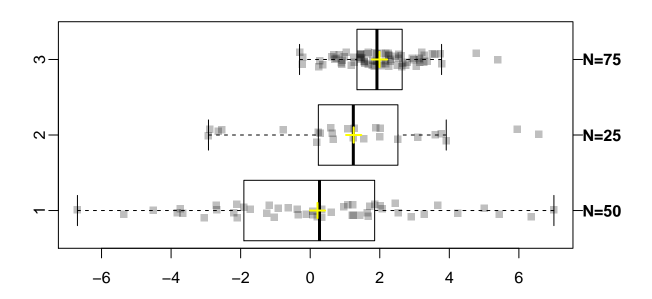
Box Plot



13.6 ... horizontally plotted (no axis-label rotation done here)

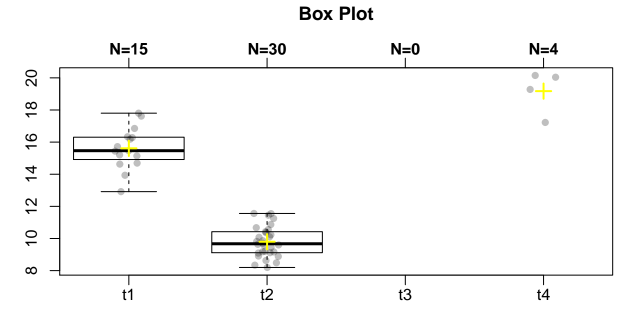
BoxPlot(rnorm(50,,3), rnorm(25,1,2), rnorm(75,2,1), horizontal=TRUE)

Box Plot



13.7 Plot values 'y' according to time 'time' (factor levels are automatically ordered as e.g. function sort does)

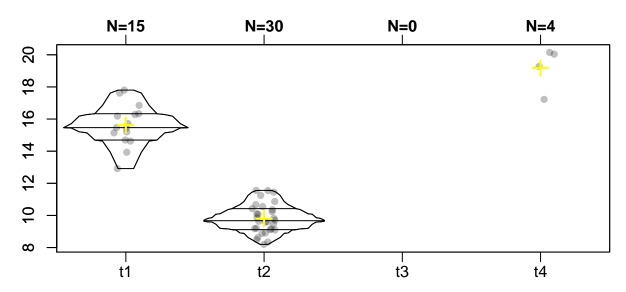
BoxPlot(example, y~time, sc.pch=16)



13.8 Now as box-percentile plot

BoxPlot(example, y~time, sc.pch=16, box.type="bp")

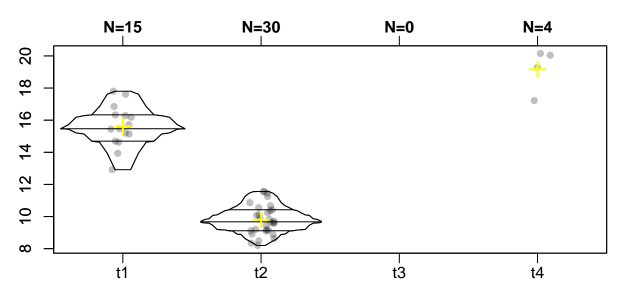
Box-Percentile Plot



13.9 With custom main title

BoxPlot(example, y~time, sc.pch=16, box.type="bp", Title=list(main="Custom Main Title"))

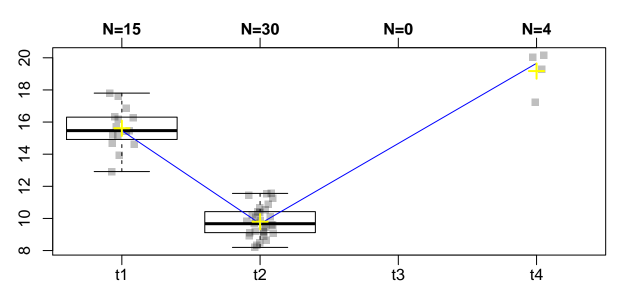
Custom Main Title



13.10 The same plot with trend line (connects either means or medians) and Y-axis label

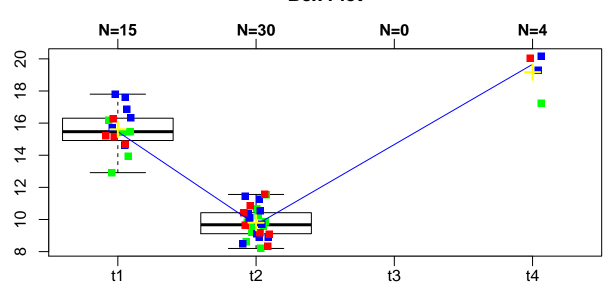
BoxPlot(example, y~time, trend="median", ylab="Y-Axis Label")

Box Plot



13.11 Use an addition grouping variable to color points in the stripchart

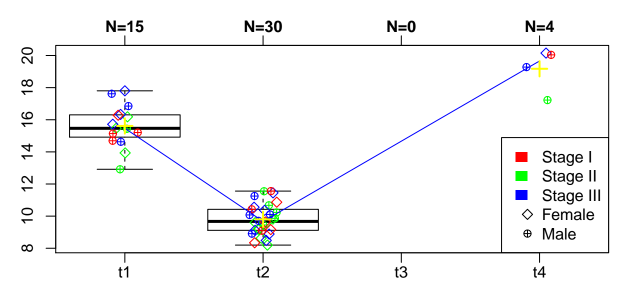
Box Plot



13.12 Use yet another grouping factor for plotting symbols in the stripchart

```
legend=c("Stage I", "Stage II", "Stage III", "Female", "Male"),
pch=c(-1, -1, -1, 5, 10), border=NA)
```

Box Plot



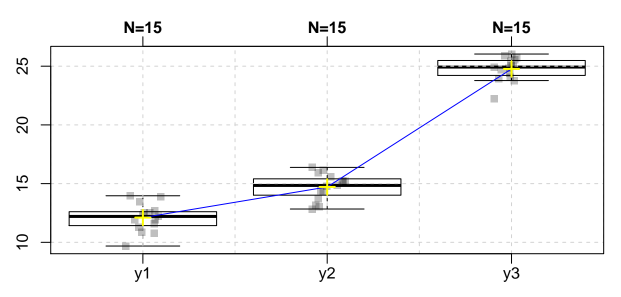
13.13 Generate new dataset with different stucture

```
example2 <- data.frame(y1=12+rnorm(15), y2=15+rnorm(15), y3=25+rnorm(15))
```

13.14 Boxplot with trend lines (mean) and grid added to the plot

```
BoxPlot(example2, var=c("y1", "y2", "y3"), Grid=TRUE, trend="mean")
```

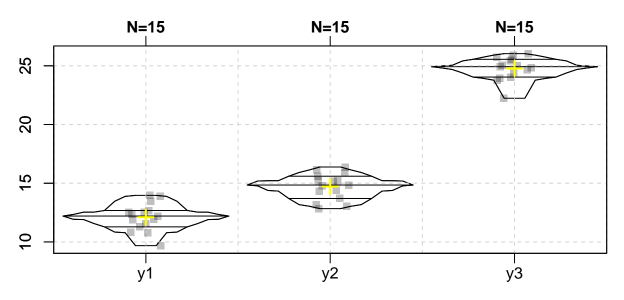
Box Plot



13.15 Now as box-percentile plot

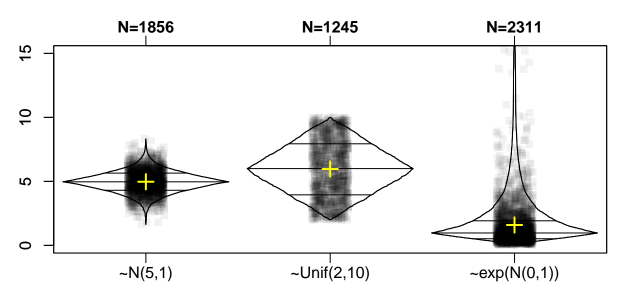
```
BoxPlot(example2, var=c("y1", "y2", "y3"), box.type="bp", Grid=TRUE)
```

Box-Percentile Plot



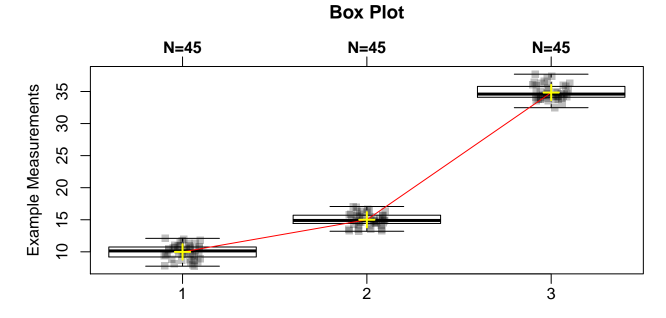
13.16 Use yet another way to specify the data

Box-Percentile Plot



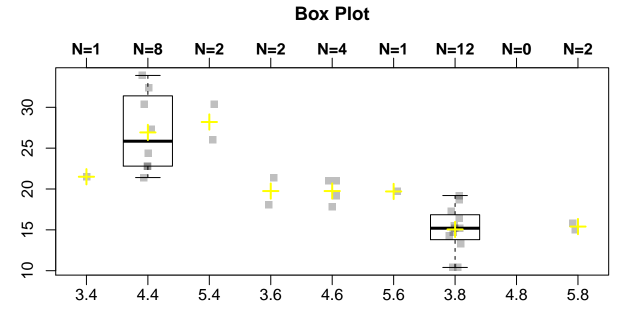
13.17 Specifying data as numeric matrix and using a trend-line connecting the means

```
mat <- matrix(c(rep(10,45), rep(15,45), rep(35,45))+rnorm(135), ncol=3)
BoxPlot(mat, trend="mean", trend.col="red", Ylabel=list(text="Example Measurements"))</pre>
```



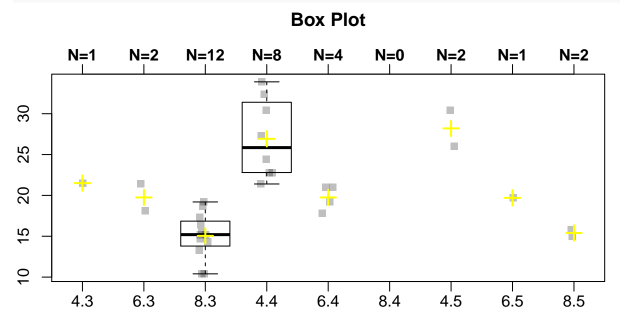
13.18 Multiple grouping factors can be specified via the formula interface which is exemplified using the mtcars dataset, of interest is miles per gallon (mgp) depending on number of gears and on the number of cylinders

```
data(mtcars)
BoxPlot(mtcars, mpg~gear:cyl)
```



13.19 Now 'cyl' is explicitly nested within 'gear' which changes the ordering of combined grouping factors (which is identical to using formula 'mpg~cyl:gear').

BoxPlot(mtcars, mpg~cyl %in% gear)



13.20 More meaningful group-labels are best specified using custom factor-level names (increase width of the plot window)

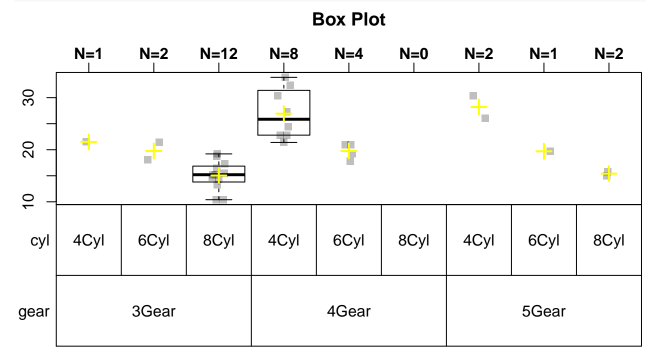
```
dat <- mtcars
dat$cyl <- factor(dat$cyl, levels=c(4,6,8), labels=c("4Cyl", "6Cyl", "8Cyl"))
dat$gear <- factor(dat$gear, levels=c(3,4,5), labels=c("3Gear", "4Gear", "5Gear"))
BoxPlot(dat, mpg~cyl %in% gear)</pre>
```

Box Plot

N=12 N=1 N=2 N=4 N=2 N=1 N=2 N=8 N=030 25 20 15 10 8Cyl.3Gear 6Cyl.4Gear 4Cyl.5Gear 4Cyl.3Gear 8Cyl.5Gear

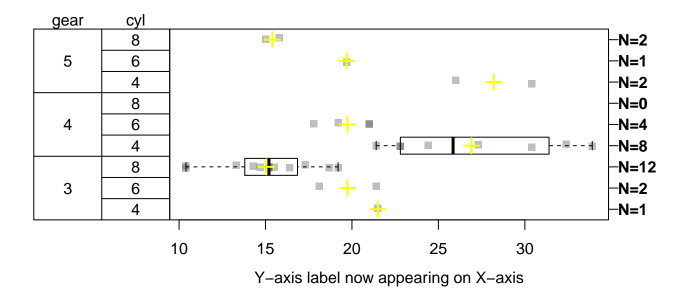
13.21 One can use a table as Xaxis label representing the factor-level combination defining sub-classes





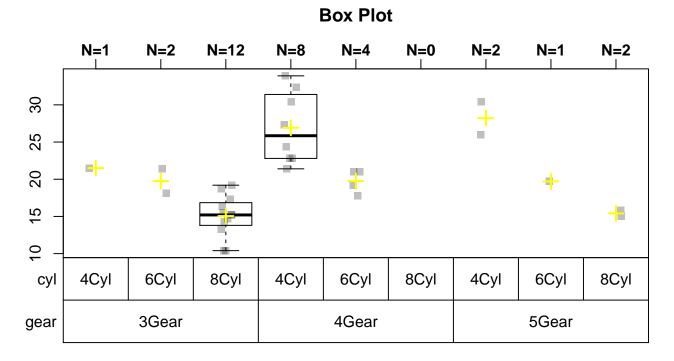
13.22 With the original factor levels and as horizontal plot

Box Plot

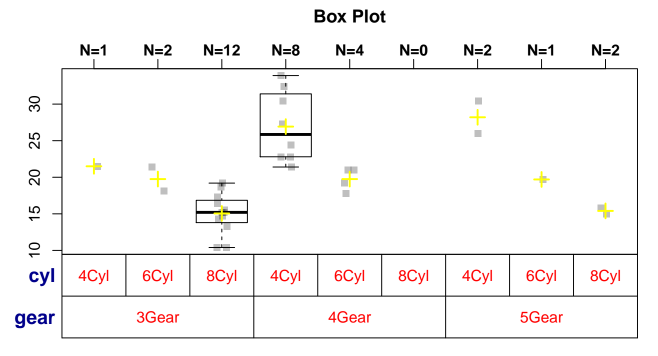


13.23 Using smaller bottom margin will result in smaller table height

BoxPlot(dat, mpg~cyl:gear, XaxisTab=list(font=2, col="darkblue", cex=1.25), mar=c(5,3,5,1))

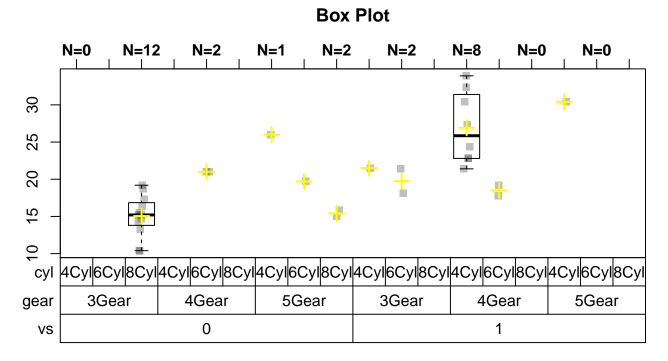


13.24 One can use different font-settings for rownames and cells of the table



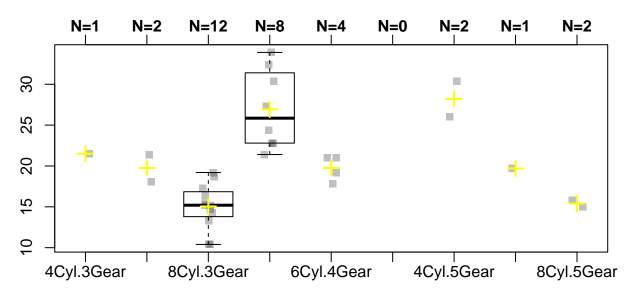
13.25 Use more crossed factors

```
BoxPlot(dat, mpg~cyl:gear:vs, XaxisTab=list(), mar=c(5,3,5,1))
```



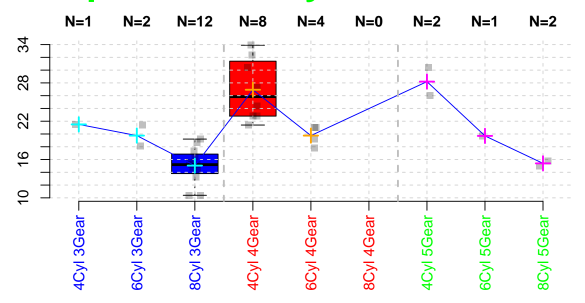
13.26 Alternatively one can use the 'Xaxis' argument, but the ordering of these labels is not checked which is not important for automatically generated group-labels as shown in the previous example

Box Plot



13.27 The same plot with some fancy options

Miles per Gallon by Number of Gea

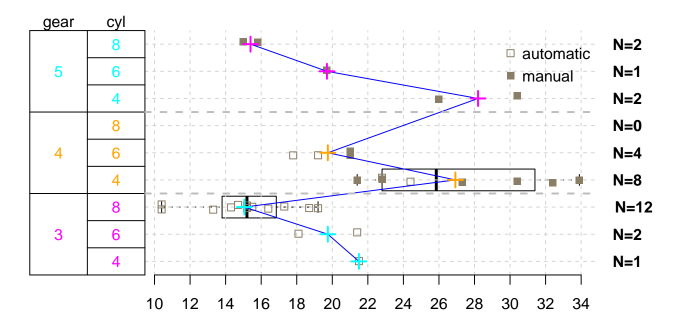


13.28 Horizontal fancy plot

```
BoxPlot(mtcars, mpg~cyl %in% gear, Title=list(main="Miles per Gallon by Number of Gears", col.main="#84
    vline=c(3.5, 6.5), vl.lty=2, vl.col="gray", vl.lwd=2,
    Xaxis2=list(tick=FALSE, las=2, hadj=-.25), Yaxis=list(at=seq(10,34,2)), Grid=list(x=1:9, y=seq(
    mean.col=c(rep("cyan", 3), rep("orange", 3), rep("magenta", 3)), Box=FALSE, trend="mean",
    mar=c(3, 7, 4, 4), horizontal=TRUE, sc.pch=c(0, 15)[dat$am+1], sc.col="wheat4",
    XaxisTab=list(Text=list(col=c(rep("cyan", 3), rep("orange", 3), rep("magenta", 3)))))

legend(x="topright", pch=c(0, 15), legend=c("automatic", "manual"), box.lty=0, col="wheat4")
```

Miles per Gallon by Number of Gears



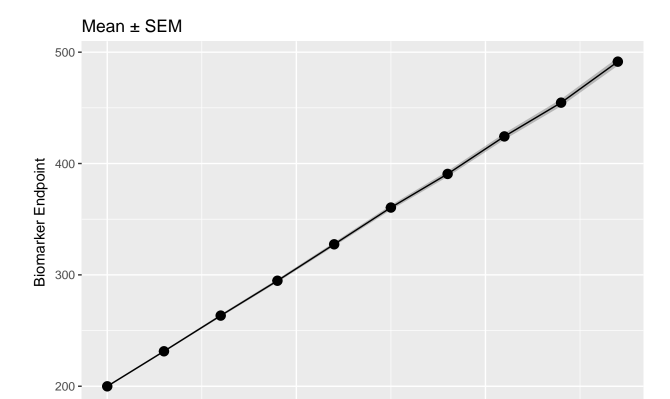
14 Longitudinal analysis

In the context of biomarker analysis, the PlotLong function can be used to plot the longitudinal measurement of a biomarker endpoint for one or more patient subpopulations. Here we'll use the packaged dataset bmkr (biomarker) which has columns for patient ID (pid), treatment (trt), gender (sex), visitation month (vm) and endpoint reading (ep)

14.1 Biomarker Endpoint Progression

14.1.1 Mean \pm SEM Timecourse

Here we plot the mean and standard error of the mean over time. We can see clearly that the mean biomarker endpoint increases throughout the study.



14.1.2 Tukey Hinges and Whiskers

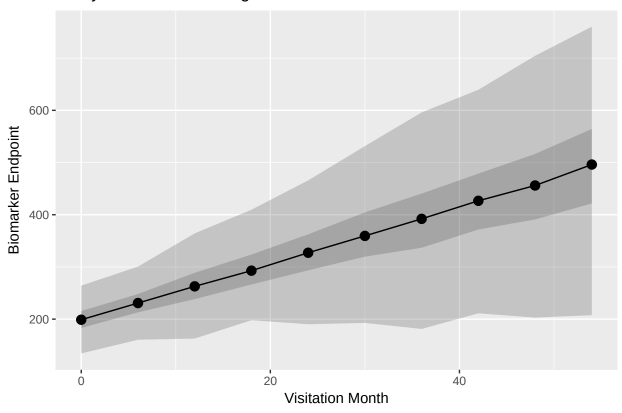
However, we may be more interested in the population distribution progression where we'd want to plot the Tukey boxplot hinges and whiskers over time. From this plot, we can see that although there is a slow rise in population mean, the patient response also becomes more distributed.

Visitation Month

40

20

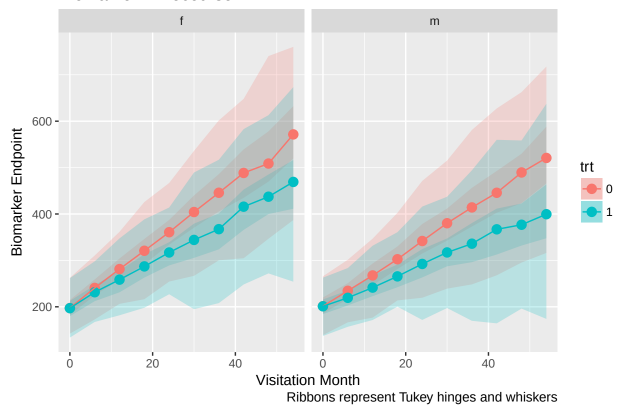
Tukey Whiskers and Hinges



14.2 Plotting Subpopulations

```
PlotLong(longbmkr, aes(x=vm, y=ep, group=trt, color=trt, fill=trt),
    fun.data = 'tukey', facet.fun = . ~ sex,
    xlab = 'Visitation Month',
    ylab = 'Biomarker Endpoint',
    labs.title = 'Biomarker Timecourse',
    labs.caption = 'Ribbons represent Tukey hinges and whiskers')
```

Biomarker Timecourse



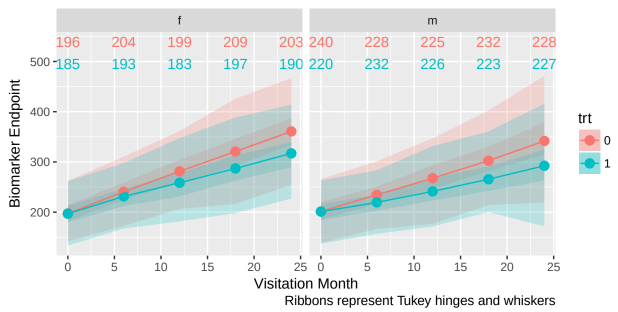
14.3 Including sample counts

14.3.1 Sample counts as table above plot

```
library(dplyr)
## Warning: package 'dplyr' was built under R version 3.3.2
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
##
##
       filter, lag
  The following objects are masked from 'package:base':
##
##
       intersect, setdiff, setequal, union
##
PlotLong(longbmkr %>% filter(vm <= 24),</pre>
         aes(x=vm, y=ep, group=trt, color=trt, fill=trt),
         fun.data = 'tukey', facet.fun = . ~ sex,
         show.counts = 'table',
         xlab = 'Visitation Month',
         ylab = 'Biomarker Endpoint',
         labs.title = 'Biomarker Timecourse',
         labs.caption = 'Ribbons represent Tukey hinges and whiskers')
```

Warning: package 'bindrcpp' was built under R version 3.3.2

Biomarker Timecourse



14.4 Plotting Deviations as Errorbars

Biomarker Timecourse

