Overview of the package BuyseTest

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The **BuyseTest** package contains five functions that are relevant for the user:

- the BuyseTest function to compute the net benefit/win ratio. It is the main function of the package.
- a summary function to display the results computed by the BuyseTest function.
- a confint function to extract estimates, confidence intervals, and p.values.
- the getPairScore method to extract the contribution of each pair to the net benefit/win ratio.
- the BuyseTest.options function the contain the default values for the argument of the BuyseTest function. These default values can be changed to better match the user needs.

Two additional function are presented in this document: simBuyseTest that can be used to quickly simulate data and getSurvival that is useful to reproduce the results output by BuyseTest.

Before going further we need to load the **BuyseTest** package in the R session:

library(BuyseTest)

The **BuyseTest** package is under active development. Newer package versions may include additional functionalities and fix previous bugs. To get the current version of the package use utils::packageVersion:

utils::packageVersion("BuyseTest")

[1] '1.6'

1 Example data

For this overview we will used the veteran dataset from the survival package:

```
data(veteran,package="survival")
head(veteran)
```

```
trt celltype time status karno diagtime age prior
                72
                                         7
    1 squamous
                         1
                              60
                                            69
1
2
    1 squamous 411
                         1
                              70
                                         5
                                            64
                                                  10
3
    1 squamous 228
                         1
                              60
                                         3
                                            38
                                                   0
    1 squamous 126
                         1
                                         9
4
                              60
                                            63
                                                  10
    1 squamous 118
                         1
                              70
                                            65
                                                  10
5
                                        11
                         1
6
    1 squamous
                10
                              20
                                         5
                                            49
                                                   0
```

See ?veteran for a presentation of the database.

2 Performing generalized pairwise comparisons (GPC) using the BuyseTest function

To perform generalized pairwise comparisons, the BuyseTest function needs:

• an where the data are stored - argument data

• the name of the endpoints - argument endpoint

• the type of each endpoint - argument type

• the variable defining the two treatment groups - argument treatment

The BuyseTest function has many optional arguments to specify for example:

- the threshold associated to each endpoint (default= 10^{-12}) argument threshold
- the censoring associated to each endpoint (for time to event endpoint) argument censoring
- how to compute the distribution of the statistic of interest argument method.inference

There are two equivalent ways to define the GPC:

• using a separate argument for each element¹:

Generalized Pairwise Comparisons

Settings

- management of censored survival pairs: use Kaplan Meier survival curves to compute the score

Point estimation

Gather the results in a BuyseRes object

 $^{^{1}}$ we set the argument method.inference to "none" to diseable the computation of p-values and confidence intervals. This makes the execution of BuyseTest much faster.

- or via a formula interface. In the formula interface endpoint are wrapped by parentheses. The parentheses must be preceded by their type:
 - binary (b, bin, or binary)
 - continuous (c, cont, or continuous)
 - time to event (t, tte, or timetoevent)

```
BT.f <- BuyseTest(trt ~ tte(time, threshold = 20, censoring = "status"),
data = veteran, trace = 0, method.inference = "none")
```

Here we set in addition the argument trace to 0 to force the function to be silent (i.e. no display in the terminal). We can check that the two approaches are equivalent:

```
testthat::expect_equal(BT.f,BT)
```

2.1 Displaying the results

The results of the GPC can be displayed using the summary method:

```
summary(BT)
```

Generalized pairwise comparison with 1 prioritized endpoint

To display the number of pairs instead of the percentage of pairs that are favorable/unfavorable/neutral/uniformative, set the argument percentage to FALSE:

```
summary(BT, percentage = FALSE)
```

Generalized pairwise comparison with 1 prioritized endpoint

By default summary displays results relative to the net benefit. To get results for the win ratio set the argument statistic to "winRatio":

```
summary(BT, statistic = "winRatio")
```

Generalized pairwise comparison with 1 prioritized endpoint

Since we have set the argument n.permutation to 0 (i.e. no permutation test) in the stratified analysis, we do not get confidence intervals or p.values when calling the summary method.

2.2 Using multiple endpoints

More than one endpoint can be considered by indicating a vector of endpoints, types, and thresholds. In the formula interface, just add another endpoint at then end of the formula:

```
BT.multi <- BuyseTest(trt ~ tte(time, threshold = 20, censoring = "status",) + cont( karno, threshold = 0),

data = veteran, method.inference = "none", trace = 0)
```

The hierarchy of the endpoint is defined from left (most important endpoint, here time) to right (least important endpoint, here karno).

2.3 What if smaller is better?

By default BuyseTest will always assume that higher values of an endpoint are favorable. This behavior can be changed by specifying operator = "<0" for an endpoint:

```
BTinv <- BuyseTest(trt ~ tte(time, threshold = 20, censoring = "status", operator = " <0"),

data = veteran, method.inference = "none", trace = 0)

BTinv
```

```
endpoint threshold delta Delta time 20 0.0844 0.0844
```

Internally BuyseTest will multiply by -1 the values of the endpoint to ensure that lower values are considered as favorable. A direct consequence is that BuyseTest will not accept an endpoint with different operators:

```
Error in (function (alternative, name.call, censoring, correction.uninf, :

Cannot have different operator for the same endpoint used at different priorities
```

2.4 Stratified GPC

GPC can be performed for subgroups of a categorical variable

- argument strata

For instance, the celltype may have huge influence on the survival time and the investigator would like to only compare patients that have the same celltype. In the formula interface this is achieved by adding a single variable in the right hand side of the formula:

```
BT2 <- BuyseTest(trt ~ tte(time, threshold = 20, censoring = "status") + cont(karno, threshold = 0) + celltype,

data = veteran, trace = 0, method.inference = "none")
```

The fact the it is not wrapped by bin, cont or tte indicates differentiate it from endpoint variables. When doing a stratified analysis, the summary method displays the global results as well as the results within each strata:

```
summary(BT2)
```

Generalized pairwise comparison with 2 prioritized endpoints and 4 strata

```
> statistic
                   : net benefit (delta: endpoint specific, Delta: global)
> null hypothesis : Delta == 0
> treatment groups: 1 (control) vs. 2 (treatment)
> censored pairs : use Kaplan Meier survival curves to compute the score
> results
endpoint threshold
                      strata
                              total favorable unfavorable neutral uninf
                                                                             delta
                                                                                     Delta
                      global 100.00
                                                      45.77
                                                                     0.68 -0.0971 -0.0971
    time
                20
                                         36.06
                                                              17.50
                                                       8.77
                                                               2.28
                                                                     0.00 0.2193
                              25.38
                                         14.33
                     squamous
                                                                     0.68 - 0.1792
                    smallcell
                              45.69
                                         12.69
                                                      20.88
                                                              11.44
                        adeno
                               13.71
                                          4.74
                                                       6.15
                                                               2.81
                                                                     0.00 - 0.1034
                               15.23
                                          4.30
                                                       9.97
                                                               0.96
                                                                     0.00 - 0.3722
                        large
                                          6.72
                                                       8.07
                                                               3.38
                                                                     0.00 -0.0135 -0.1106
   karno
             1e-12
                       global
                               18.17
                     squamous
                                2.28
                                          0.76
                                                       0.94
                                                               0.59
                                                                     0.00 -0.0071
                   smallcell 12.12
                                          4.33
                                                       5.75
                                                               2.03
                                                                     0.00 -0.0311
                                2.81
                                                       0.85
                                                               0.51
                                                                     0.00 0.0448
                                           1.46
                        adeno
                                                       0.54
                        large
                                0.96
                                          0.17
                                                               0.25
                                                                     0.00 -0.0241
```

Note that here the numbers in the favorable/unfavorable/neutral/uniformative columns are relative to the overall sample while the delta is only relative to the strata. The global delta is a sum of the strata specific delta weighted by the empirical proportion of pairs for each strata.

2.5 Stopping comparison for neutral pairs

In presence of neutral pairs, BuyseTest will, by default, continue the comparison on the endpoints with lower priority. For instance let consider a dataset with one observation in each treatment arm:

```
Id treatment tumor size
1: 1 Yes Yes 15
2: 2 No Yes 20
```

If we perform we GPC with tumor as the first endpoint and size as the second endpoint:

Warning message:

```
In '[.data.table'(data, , ':='(c("..rowIndex.."), 1:.N)) :
    Invalid .internal.selfref detected and fixed by taking a (shallow) copy of the data.table so th
        Generalized pairwise comparison with 2 prioritized endpoints
```

```
: net benefit (delta: endpoint specific, Delta: global)
> statistic
> null hypothesis : Delta == 0
> treatment groups: No (control) vs. Yes (treatment)
> results
endpoint threshold total favorable unfavorable neutral uninf delta Delta
               0.5
                     100
                                  0
                                              0
                                                     100
    size
             1e-12
                     100
                                100
                                              0
                                                       0
                                                             0
                                                                   1
                                                                         1
```

the outcome of the comparison is neutral for the first priority, but favorable for the second priority. If we set the argument neutral.as.uninf to FALSE, BuyseTest will stop the comparison when a pair is classified as neutral:

```
BT.pair2 <- BuyseTest(treatment ~ bin(tumor) + cont(size, operator = "<0"), data = dt .sim,

trace = 0, method.inference = "none", neutral.as.uninf = FALSE)
summary(BT.pair2)
```

```
Warning message:
```

```
In '[.data.table'(data, , ':='(c("..rowIndex.."), 1:.N)) :
    Invalid .internal.selfref detected and fixed by taking a (shallow) copy of the data.table so th
        Generalized pairwise comparison with 2 prioritized endpoints
```

```
> statistic
                  : net benefit (delta: endpoint specific, Delta: global)
> null hypothesis : Delta == 0
> treatment groups: No (control) vs. Yes (treatment)
> results
endpoint threshold total favorable unfavorable neutral uninf delta Delta
                                               0
                                                             0
   tumor
               0.5
                      100
                                  0
                                                     100
                                  0
                                               0
                                                                   0
             1e-12
                       0
                                                       0
                                                             0
                                                                          0
    size
```

So in this case no pair is analyzed at second priority.

2.6 What about p-value and confidence intervals?

P-values can be estimated via a permutation test:

```
BT.perm <- BuyseTest(trt ~ tte(time, threshold = 20, censoring = "status"),
data = veteran, trace = 0, method.inference = "permutation",
n.resampling = 10)
summary(BT.perm)
```

Generalized pairwise comparison with 1 prioritized endpoint

```
> statistic
                   : net benefit (delta: endpoint specific, Delta: global)
 > null hypothesis : Delta == 0
 > permutation test: 10 samples, confidence level 0.95
 > treatment groups: 1 (control) vs. 2 (treatment)
 > censored pairs : use Kaplan Meier survival curves to compute the score
 > results
 endpoint threshold total favorable unfavorable neutral uninf
                                                                 delta
                 20
                      100
                              37.78
                                          46.54
                                                  15.68
                                                            0 -0.0877 -0.0877
     time
 CI [2.5; 97.5] p.value
 [-0.2412; 0.1946]
NOTE: confidence intervals computed under the null hypothesis
```

The argument n.resampling indicates the permutation that will be performed. We set it to 10 to save computation time but to obtain reliable p-value/confidence intervals, n.resampling should be at least 1000. The validity of the confidence intervals obtained via a permutation test is questionable and we recommend instead to use a bootstrap approach for estimating confidence intervals. To do so, set the argument method.inference to "bootstrap" when calling BuyseTest:

Generalized pairwise comparison with 1 prioritized endpoint

```
> statistic : net benefit (delta: endpoint specific, Delta: global)
> null hypothesis : Delta == 0
```

- > bootstrap resampling: 10 samples, confidence level 0.95
- > treatment groups: 1 (control) vs. 2 (treatment)
- > censored pairs : use Kaplan Meier survival curves to compute the score
- > requilts

endpoint threshold total favorable unfavorable neutral uninf delta Delta time 20 100 37.78 46.54 15.68 0 -0.0877 -0.0877

CI [2.5; 97.5] p.value [-0.1794;0.0302] 0.5

Getting additional inside: looking at the pair level 3

So far we have looked at the overall score and probabilities. But it is also possible to extract the score relative to each pair, as well as to "manually" compute this score. This can give further inside on what the software is actually doing and what is the contribution of each individual on the evaluation of the treatment.

3.1 Extracting the contribution of each pair to the statistic

The net benefit or the win ratio statistics can be expressed as a sum of a score over all pairs of patients. The argument keep.pairScore enables to export the score relative to each pair in the output of BuyseTest:

```
form <- trt \sim tte(time, threshold = 20, censoring = "status") + cont(karno)
BT.keep <- BuyseTest(form,
             data = veteran, keep.pairScore = TRUE,
             trace = 0, method.inference = "none")
```

The method getPairScore can then be used to extract the contribution of each pair. For instance the following code extracts the contribution for the first endpoint:

```
getPairScore(BT.keep, endpoint = 1)
  strata index.1 index.2 favorable unfavorable neutral uninf weight favorableC
```

								6	
1:	1	1	70	1	0	0	0	1	1
2:	1	2	70	1	0	0	0	1	1
3:	1	3	70	1	0	0	0	1	1
4:	1	4	70	1	0	0	0	1	1
5:	1	5	70	1	0	0	0	1	1
4688:	1	65	137	0	1	0	0	1	0
4689:	1	66	137	0	1	0	0	1	0
4690:	1	67	137	0	1	0	0	1	0
4691:	1	68	137	0	1	0	0	1	0
4692:	1	69	137	0	1	0	0	1	0

4 4 4 4 unfavorableC neutralC uninfC

1:	0	0	0
2:	0	0	0
3:	0	0	0
4:	0	0	0
5:	0	0	0
4688:	1	0	0
4689:	1	0	0
4690:	1	0	0
4691:	1	0	0
4692:	1	0	0

Each line corresponds to different comparison between a pair from the control arm and the treatment arm. The column strata store to which strata the pair belongs (first, second, ...). The columns favorable, unfavorable, neutral, uninformative contains the result of the comparison, e.g. the first pair was classified as favorable while the last was classified as favorable with a weight of 1. The second and third columns indicates the rows in the original dataset corresponding to the pair:

```
veteran[c(70,1),]
```

```
trt celltype time status karno diagtime age prior 70 2 squamous 999 1 90 12 54 10 1 1 squamous 72 1 60 7 69 0
```

For the first pair, the event was observed for both observations and since 999 > 72 + 20 the pair is rated favorable. Substracting the average probability of the pair being favorable minus the average probability of the pair being unfavorable:

```
getPairScore(BT.keep, endpoint = 1)[, mean(favorable) - mean(unfavorable)]
```

[1] -0.08765836

gives the net benefit in favor of the treatment for the first endpoint:

```
BT.keep
```

```
endpoint threshold delta Delta
time 20 -0.0877 -0.0877
karno 1e-12 -0.0133 -0.1009
```

More examples and explanation can be found in the documentation of the method getPairScore.

3.2 Extracting the survival probabilities

When using method.tte equals "Peron", survival probabilities at event time, and event times +/threshold in the control and treatment arms are used to score the pair. Setting keep.survival to
TRUE in BuyseTest.options enables to export the survival probabilities in the output of BuyseTest:

```
BuyseTest.options(keep.survival = TRUE)
BT.keep2 <- BuyseTest(trt ~ tte(time, threshold = 20, censoring = "status") + cont(karno),

data = veteran, keep.pairScore = TRUE, method.tte = "Peron",
trace = 0, method.inference = "none")
```

The method **getSurvival** can then be used to extract these survival probabilities. For instance the following code extracts the survival for the first endpoint:

```
outSurv <- getSurvival(BT.keep2, endpoint = 1, strata = 1)
str(outSurv)</pre>
```

```
List of 5
 $ survTimeC: num [1:69, 1:7] 72 411 228 126 118 10 82 110 314 100 ...
  ..- attr(*, "dimnames")=List of 2
  .. ..$ : NULL
  ....$ : chr [1:7] "time" "SurvivalC-threshold" "SurvivalC_0" "SurvivalC+threshold" ...
 $ survTimeT: num [1:68, 1:7] 999 112 87 231 242 991 111 1 587 389 ...
  ..- attr(*, "dimnames")=List of 2
  .. ..$ : NULL
  ....$ : chr [1:7] "time" "SurvivalC-threshold" "SurvivalC_0" "SurvivalC+threshold" ...
 $ survJumpC: num [1:57, 1:3] 3 4 7 8 10 11 12 13 16 18 ...
  ..- attr(*, "dimnames")=List of 2
  .. ..$ : NULL
  ....$ : chr [1:3] "time" "survival" "dSurvival"
 $ survJumpT: num [1:51, 1:3] 1 2 7 8 13 15 18 19 20 21 ...
  ..- attr(*, "dimnames")=List of 2
  .. ..$ : NULL
  ....$ : chr [1:3] "time" "survival" "dSurvival"
 $ lastSurv : num [1:2] 0 0
```

3.2.1 Computation of the score with only one censored event

Let's look at pair 91:

```
getPairScore(BT.keep2, endpoint = 1, rm.withinStrata = FALSE)[91]
```

```
strata index.1 index.2 indexWithinStrata.1 indexWithinStrata.2 favorable unfavorable

1: 1 22 71 22 2 0 0.6950827

neutral uninf weight favorableC unfavorableC neutralC uninfC

1: 0.3049173 0 1 0 0.6950827 0.3049173 0
```

In the dataset this corresponds to:

```
veteran[c(22,71),]
```

```
trt celltype time status karno diagtime age prior 22 1 smallcell 97 0 60 5 67 0 71 2 squamous 112 1 80 6 60 0
```

The observation from the control group is censored at 97 while the observation from the treatment group has an event at 112. Since the threshold is 20, and (112-20)<97, we know that the pair is not in favor of the treatment. The formula for probability in favor of the control is $\frac{S_c(97)}{S_c(112+20)}$. The survival at the event time in the censoring group is stored in survTimeC. Since observation 22 is the 22th observation in the control group:

```
iSurv <- outSurv$survTimeC[22,]
iSurv</pre>
```

Since we are interested in the survival in the control arm exactly at the event time:

```
Sc97 <- iSurv["SurvivalC_0"]
Sc97
```

SurvivalC_0 0.5171924

The survival at the event time in the treatment group is stored in survTimeC. Since observation 71 is the 2nd observation in the treatment group:

```
iSurv <- outSurv$survTimeT[2,] ## survival at time 112+20
iSurv</pre>
```

```
time SurvivalC-threshold SurvivalC_0 SurvivalC+threshold 112.0000000 0.5319693 0.4549201 0.3594915 SurvivalT-threshold SurvivalT_0 SurvivalT+threshold 0.3801681 0.2827500 0.2827500
```

Since we are interested in the survival in the control arm at the event time plus threshold:

```
Sc132 <- iSurv["SurvivalC+threshold"]
Sc132
```

SurvivalC+threshold 0.3594915

The probability in favor of the control is then:

```
Sc132/Sc97
```

```
SurvivalC+threshold 0.6950827
```

3.2.2 Computation of the score with two censored events

When both observations are censored, the formula for computing the probability in favor of treatment or control involves an integral. This integral can be computed using the function calcIntegralScore_cpp that takes as argument a matrix containing the survival and the jumps in survival, e.g.:

```
head(outSurv$survJumpT)
```

time survival dSurvival

- [1,] 1 0.7681159 -0.02941176
- [2,] 2 0.7536232 -0.01470588
- [3,] 7 0.7388463 -0.02941176
- [4,] 8 0.7388463 -0.02941176
- [5,] 13 0.7092924 -0.01470588
- [6,] 15 0.6945155 -0.02941176

and the starting time of the integration time. For instance, let's look at pair 148:

```
getPairScore(BT.keep2, endpoint = 1, rm.withinStrata = FALSE)[148]
```

strata index.1 index.2 indexWithinStrata.1 indexWithinStrata.2 favorable unfavorable

1: 1 10 72 10 3 0.5058685 0.3770426

neutral uninf weight favorableC unfavorableC neutralC uninfC

1: 0.1170889 0 1 0.5058685 0.3770426 0.1170889 0

which corresponds to the observations:

trt celltype time status karno diagtime age prior

- 10 1 squamous 100 0 70 6 70 0
- 72 2 squamous 87 0 80 3 48

The probability in favor of the treatment (p_F) and control (p_{UF}) can be computed as:

$$p_F = -\frac{1}{S_T(x)S_C(y)} \int_{t>y} S_T(t+\tau) dS_C(t)$$

$$p_{UF} = -\frac{1}{S_T(x)S_C(y)} \int_{t>x} S_C(t+\tau) dS_T(t)$$

where x = 87 and y = 100. We obtain:

favorable unfavorable 0.5058685 0.3770426

4 Dealing with missing values or/and right censoring

In presence of censoring or missing values, some pairs may be classified as uninformative. This may bias the estimate of the net net benefit. Two corrections are currently proposed to correct this bias.

To illustrate the effect of these correction, we will use the following dataset:

Treatment toxicity eventtimeUncensored eventtimeCensoring eventtime status 1

1:	C	0	0.1588268	2.6268101 0.1588268	1	1
2:	C	1	1.7204676	0.2000192 0.2000192	0	1
3:	C	1	0.4900490	0.5747995 0.4900490	1	1
4:	C	0	0.1138545	1.5188001 0.1138545	1	1
5:	C	1	0.5191035	3.8340048 0.5191035	1	1
6:	C	0	0.9405830	1.9078657 0.9405830	1	1

where we have the uncensored event times as well as the censored event times. The percentage of censored observations is:

```
dt[,mean(status==0)]
```

[1] 0.317

We would like to be able to recover the net benefit estimated with the uncensored event times:

```
BuyseTest(Treatment ~ tte(eventtimeUncensored, status1, threshold = 1),
    data = dt,
    method.tte = "Gehan", method.inference = "none", trace = 0)
```

```
endpoint threshold delta Delta eventtimeUncensored 1 0.2401 0.2401
```

using the censored survival times:

```
BuyseTest(Treatment ~ tte(eventtime, status, threshold = 1),
    data = dt,
    method.tte = "Gehan", method.inference = "none", trace = 0)
```

```
endpoint threshold delta Delta eventtime 1 0.1363 0.1363
```

As we can see on this example, the net benefit is shrunk toward 0.

4.0.1 Inverse probability-of-censoring weights (IPCW)

With IPCW the weights of the non-informative pairs is redistributed to the informative pairs. This is only a good strategy when there are no neutral pairs or there are no lower priority endpoints. This gives an estimate much closer to the true net benefit:

```
BT <- BuyseTest(Treatment ~ tte(eventtime, status, threshold = 1),
data = dt, keep.pairScore = TRUE, trace = 0,
method.tte = "Gehan", method.inference = "none", correction.uninf = 2)
summary(BT)
```

Generalized pairwise comparison with 1 prioritized endpoint

```
> statistic : net benefit (delta: endpoint specific, Delta: global)
> null hypothesis : Delta == 0
> treatment groups: C (control) vs. T (treatment)
> censored pairs : uninformative pairs
> uninformative pairs: no contribution, their weight is passed to the informative pairs using IPC
> results
  endpoint threshold total favorable unfavorable neutral uninf delta Delta
eventtime    1 100 37.11 12.34 50.54 0 0.2477 0.2477
```

We can also see that no pair is finally classified as non informative. To get some inside about the correction we can look at the scores of the pairs:

```
iScore <- getPairScore(BT, endpoint = 1)
```

To get a synthetic view, we only look at the unique favorable/unfavorable/neutral/uniformative results:

```
iScore[,.SD[1], by = c("favorable", "unfavorable", "neutral", "uninformative")]
```

```
Error in eval(bysub, x, parent.frame()) :
  objet 'uninformative' introuvable
1: powerBuyseTest(sim = simBuyseTest, sample.size = c(100), n.rep = 2,
       formula = Treatment ~ tte(eventtime, censoring = status),
       method.inference = "asymptotic", trace = 4)
2: do.call(method.loop, args = list(X = 1:n.rep, FUN = function(X) {
       return(warper(i = X, envir = envirBT))
   })) at powerBuyseTest.R#310
3: (function (X, FUN, ..., cl = NULL)
   {
       FUN <- match.fun(FUN)</pre>
       if (!is.vector(X) || is.object(X))
           X <- as.list(X)</pre>
4: lapply(X[Split[[i]]], FUN, ...)
5: FUN(X[[i]], ...)
6: warper(i = X, envir = envirBT) at powerBuyseTest.R#313
```

We can see that the favorable/unfavorable/neutral pairs have seen their contribution multiplied by:

```
iScore[,1/mean(favorable + unfavorable + neutral)]
```

[1] 1.81657

i.e. the inverse probability of being informative.

4.0.2 Correction at the pair level

Another possible correction is to distribute the non-informative weight of a pair to the average favorable/unfavorable/neutral probability observed on the sample:

Generalized pairwise comparison with 1 prioritized endpoint

```
> statistic : net benefit (delta: endpoint specific, Delta: global)
> null hypothesis : Delta == 0
> treatment groups: C (control) vs. T (treatment)
> censored pairs : uninformative pairs
> uninformative pairs: score equals the averaged score of all informative pairs
> results
  endpoint threshold total favorable unfavorable neutral uninf delta Delta
eventtime    1 100 37.11 12.34 50.54 0 0.2477 0.2477
```

Looking at the scores of the pairs:

```
iScore <- getPairScore(BT, endpoint = 1)
iScore[,.SD[1], by = c("favorable", "unfavorable", "neutral", "uninformative")]</pre>
```

we can see that the corrected probability have not changed for the informative pairs, but for the non-informative they have been set to:

5 Simulating data using simBuyseTest

You can simulate data with the **simBuyseTest** function. For instance the following code simulates data for 5 individuals in the treatment arm and 5 individuals in the control arm:

```
set.seed(10)
simBuyseTest(n.T = 5, n.C = 5)
```

```
Treatment toxicity
                               score eventtime status
1:
            C
                      1
                         0.54361539 1.8252132
            С
2:
                      1 -0.70762484 2.9489056
                                                      1
            C
                      1 -0.36944577 0.7213402
                                                      0
3:
            C
                      1 -1.32197565 0.6322603
                                                      1
4:
            C
                          1.28059746 0.2212117
                                                      0
5:
6:
            Т
                         0.01874617 0.1453481
                                                      0
            Т
                      1 -0.18425254 0.4855601
                                                      0
7:
            Τ
                      0 -1.37133055 0.2547505
                                                      0
8:
9:
            Τ
                      1 -0.59916772 1.0340368
                                                      0
10:
                         0.29454513 0.3579324
                                                      1
```

By default a categorical, continuous and time to event outcome are generated independently. You can modify their distribution via the arguments argsBin, argsCont, argsTTE. For instance the following code simulates two continuous variables with mean 5 in the treatment arm and 10 in the control arm all with variance 1:

```
Treatment toxicity tumorSize
                                       score eventtime status
1:
            С
                         9.010394 10.667415 0.2729620
                                                             0
2:
            C
                         9.965152 11.691755 0.5562477
                                                             0
3:
            C
                      0 10.847160 10.001261 0.8040608
                                                             0
            C
                      0 11.525498  9.257539  1.8477048
                                                             1
4:
            C
                         9.932625 10.609684 0.3639572
                                                             1
5:
6:
            Τ
                      1
                         5.389794
                                   5.018746 0.6243732
                                                             0
7:
            Τ
                         3.791924
                                   4.815747 0.3527879
                                                             1
8:
            Τ
                        4.636324
                                   3.628669 1.7731161
                                                             0
9:
            Τ
                         3.373327
                                   4.400832 0.1055467
                                                             0
            Т
10:
                         4.743522 5.294545 0.8612402
                                                             0
```

This functionality is based on the sim function of the lava package (https://github.com/kkholst/lava)

6 Modifying default options

The <code>BuyseTest.options</code> method enable to get and set the default options of the <code>BuyseTest</code> function. For instance, the default option for trace is:

BuyseTest.options("trace")

\$trace

[1] 2

To change the default option to 0 (i.e. no output) use:

BuyseTest.options(trace = 0)

To restore the original default options do:

BuyseTest.options(reinitialise = TRUE)

7 Information about the R session used for this document

sessionInfo() R version 3.5.1 (2018-07-02) Platform: x86_64-pc-linux-gnu (64-bit) Running under: Ubuntu 16.04.5 LTS Matrix products: default BLAS: /usr/lib/libblas/libblas.so.3.6.0 LAPACK: /usr/lib/lapack/liblapack.so.3.6.0 locale: [1] LC_CTYPE=fr_FR.UTF-8 LC_NUMERIC=C LC_TIME=da_DK.UTF-8 [4] LC_COLLATE=fr_FR.UTF-8 LC_MONETARY=da_DK.UTF-8 LC_MESSAGES=fr_FR.UTF-8 [7] LC_PAPER=da_DK.UTF-8 LC_NAME=C LC_ADDRESS=C LC_MEASUREMENT=da_DK.UTF-8 LC_IDENTIFICATION=C [10] LC_TELEPHONE=C attached base packages: [1] stats graphics grDevices utils datasets methods base other attached packages: [1] BuyseTest_1.6 testthat_2.0.0 data.table_1.11.8 Rcpp_0.12.19 XML_3.98-1.16 [5] prodlim_2018.04.18 butils.base_1.1 selectr_0.4-1 [9] devtools_1.13.6 loaded via a namespace (and not attached): [1] compiler_3.5.1 pipeR_0.6.1.3 iterators_1.0.10 [4] base64enc_0.1-3 tools_3.5.1 digest_0.6.17 [7] pkgbuild_1.0.1 pkgload_1.0.0 memoise_1.1.0 [10] lattice_0.20-35 rlang_0.2.2 foreach_1.4.4 [13] Matrix_1.2-14 cli_1.0.1 commonmark_1.6 [16] parallel_3.5.1 RcppArmadillo_0.9.100.5.0 withr_2.1.2 [19] httr_1.3.1 roxygen2_6.1.0.9000 stringr_1.3.1 [22] xml2_1.2.0 desc_1.2.0 stats4_3.5.1 [25] rprojroot_1.3-2 grid_3.5.1 R6_2.3.0 [28] processx_3.2.0 pbapply_1.3-4 survival_2.42-6 [31] lava_1.6.3 purrr_0.2.5 callr_3.0.0 [34] magrittr_1.5 codetools_0.2-15 splines_3.5.1 [37] backports_1.1.2 ps_1.1.0 assertthat_0.2.0

stringi_1.2.4

zoo_1.8-4

KernSmooth_2.23-15

crayon_1.3.4

[40] rvest_0.3.2

[43] doParallel_1.0.14