Step-by-step user instructions to the hamlet-package

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Hamlet is an R package intended for the statistical analysis of pre-clinical studies. This document is a basic introduction to the functionality of hamlet and a general overview to the analysis workflow of preclinical studies.

This document is structured as follows: First, a general overview to inputting and processing the raw data is presented. Second, functionality is presented for the processing of pre-intervention data. Finally, functionality is presented for the post-intervention period, along with brief discussion on the differences between non-matched and matched statistical approaches. Each section comes with a list of useful functions specific for the subtask.

Latest version of hamlet is available in the Comprehensive R Archive Network (CRAN, http://cran.r-project.org/). CRAN mirrors are by default available in the installation of R, and the hamlet package is installable using the R terminal command: install.packages("hamlet"). This should prompt the user to select a nearby CRAN mirror, after which the installation of hamlet is automatically performed. After the install.packages-call, the hamlet package can be loaded with either command library("hamlet") or require("hamlet").

The following notation is used in the document: R commands, package names and function names are written in typewriter font. The notation of format pckgName::funcName indicates that the function funcName is called from the package pckgName. If only the function name is given, this indicates that it is located in the base package in R and is thus always available.

1 Analysis workflow

Two different types of case-control setups for the analysis of pre-clinical are presented in Fig. 1.

The type A experiment design in Fig. 1 is preferred, as matching is performed before allocation to the experiment groups, and therefore improves the balance and power of the experiment. The alternate experiment type B requires the bipartite matching task, where suitable pairs of individuals are identified over two or more groups that existed prior to matching. This document presents a dataset where experiment design type A was used.

2 Loading data into R

The hamlet package comes pre-installed with the VCaP dataset, which is used here to illustrate the workflow. Two different formats of the data are provided. First one is available in data(vcapwide), which includes the data in the so-called wide format. In this data format the columns are indicators for different variables available for the experimental unit (here animal). For example, the two first rows of observations are extracted with:

- > require(hamlet)
 > data(vcapwide)
- > vcapwide[1:2,]

${\tt TreatmentInitiationWeek}$	${\tt Group}$	CageAtAllocation	${\tt CastrationDate}$	
Week10	Vehicle	13489	100413	ID003
Week10	MDV	13810	170413	ID007

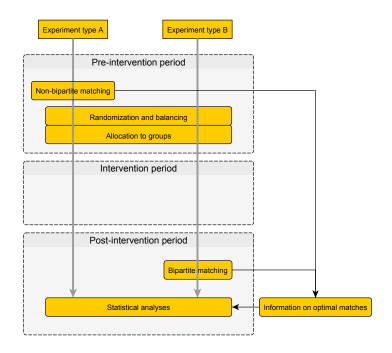


Figure 1: Analysis workflow for pre-clinical experiments

	Subma	atch II	PSAWeek2	PSAWeek3	PSAWeek4	PSAWeek5	PSAWeek6	PSAWeek7
ID003	Submate	h_1 ID003	7.67	14.76	24.78	2.03	5.97	8.16
ID007	Submatch	10 ID007	2.01	5.17	8.59	14.62	1.99	2.81
	PSAWeek8	B PSAWeek9	PSAWeek1	O PSAWeek	11 PSAWee	k12 PSAWee	k13 PSAWe	ek14
ID003	13.72	2 16.57	21.3	0 45.	69 54	.50 53	3.55 2	7.64
ID007	4.23	5.38	7.5	5 9.	70 17	.45 22	2.79 2	1.88
	BWWeek0	BWWeek1 E	BWWeek2 BW	Week3 BWW	eek4 BWWe	ek5 BWWeek	k6 BWWeek7	BWWeek8
ID003	30.5	31.7	32.6	33.8	33.9 33	2.2 32.	6 32.6	33.2
ID007	28.8	30.0	30.6	31.6	32.9 33	2.4 32.	0 31.1	30.3
	BWWeek9	BWWeek10	BWWeek11	BWWeek12	BWWeek13 I	BWWeek14		
ID003	34.2	35.0	36.1	37.9	37.5	39.7		
ID007	30.5	31.6	31.7	32.4	33.5	33.3		

An another format of the same dataset is provided in data(vcaplong). This is the data from the same experiment in the so-called *long* format, where only few column variables are available (here PSA or body weight), and the different observations belonging to a single experimental unit (here animal) are distinguished using the measurement time (variable *Week* or *DrugWeek*). Again, first few rows of the dataset:

- > data(vcaplong)
- > vcaplong[1:3,]

```
PSA log2PSA
                     {\tt BW}
                          Submatch
                                       ID Week DrugWeek
                                                           Group Vehicle ARN MDV
11 21.30 4.412782 35.0 Submatch_1 ID003
                                                      0 Vehicle
                                                                                0
                                                                            0
                                            10
                                                                        1
                                                                                0
12 45.69 5.513807 36.1 Submatch_1 ID003
                                                       1 Vehicle
                                                                        1
                                            11
                                                                            0
13 54.50 5.768184 37.9 Submatch_1 ID003
                                            12
                                                      2 Vehicle
                                                                                0
```

1	Α	В	С	D
1	Animal	PSA week 10 [ug/l]	PSA week 9 [ug/I]	Body weight week 10 [g]
2	ID003	21,3	16,57	35
3	ID007	7,55	5,38	31,6
4	ID008	23,58	17,4	33,6
5	ID009	13,17	11,14	31,7
6	ID010	9,9	9,33	34,1
7	ID016	15,05	15,29	39,6
8	ID018	13,53	12,14	34
9	ID025	13,13	10,91	33,3
10	ID027	9,59	8,79	32
11	ID031	7,04	6,95	36,6
12	ID032	8,49	8,02	34,9
13	ID037	13,74	13,38	32,4
14	ID040	23,62	19,15	35,9
15	ID045	14,27	9,8	34,8
16	ID047	6,57	6,28	31,9
17	ID054	34,72	27,14	32,1
18	ID056	28,15	22,05	32,2
19	ID058	9,74	7,68	34

Figure 2: Example Excel-format data, where rows correspond to individuals and columns to different characteristics at baseline. The single sheet data can be easily exported in a text-based format such as CSV.

The former wide format is useful for summarizing multiple variables when constructing distance matrices for the data. The latter long format is typically used for longitudinal mixed-effects modeling where observations are correlated through time.

2.1 Excel format data

An example view of a pre-clinical dataset is given in Fig. 2. Such a dataset can be saved in an R-friendly format by selecting option File > Save As and CSV (Comma delimited) as the save format in MS Excel.

2.2 CSV-files

CSV (Comma Delimited Values) is a suitable text-based format for the data to be read into R using either the function read.table or read.csv. The above presented example CSV file can be opened with the following command:

```
> ex <- read.table(file="example.csv", sep=";", dec=",", stringsAsFactors=F, header=T)
> ex
```

	Animal	PSA.week.10ug.1.	PSA.week.9ug.1.	Body.weight.week.10g.
1	ID003	21.30	16.57	35.0
2	ID007	7.55	5.38	31.6
3	ID008	23.58	17.40	33.6
4	ID009	13.17	11.14	31.7
5	ID010	9.90	9.33	34.1

6	ID016	15.05	15.29	39.6
7	ID018	13.53	12.14	34.0
8	ID025	13.13	10.91	33.3
9	ID027	9.59	8.79	32.0
10	ID031	7.04	6.95	36.6
11	ID032	8.49	8.02	34.9
12	ID037	13.74	13.38	32.4
13	ID040	23.62	19.15	35.9
14	ID045	14.27	9.80	34.8
15	ID047	6.57	6.28	31.9
16	ID054	34.72	27.14	32.1
17	ID056	28.15	22.05	32.2
18	ID058	9.74	7.68	34.0

The above presented CSV file was read into R using read.table with the following parameters: file="example.csv" is the first parameter and indicates the input file from our current working directory. The working directory may be changed using the command setwd or by including its path in the file parameter, i.e. file="D://my//current//windows//working//directory//example.csv" sep=";" indicates that the values on each line are separated with the symbol ';', as is the format defined for the CSV delimited files with ","-decimals. This could also be a value such as \tab or " " (space). dec="," indicates that the "," symbol is used for decimals. The default value for indicating decimals is "." otherwise. stringsAsFactors=F indicates that strings should not be handled as factors. Factors are an R class, where a character string may only take instances of a predetermined set of strings. As each of our animal IDs - which are read as strings - are unique, it is generally more flexible to conserve them as character strings. Lastly, header=T indicates that the text CSV file has a header row as the first row, which includes names for each column. If this value is set to header=F or header=FALSE, the first row of the text file is read as the first observation and the columns are left unnamed.

Depending on the country of origin, the CSV files may use "." decimals and "," separator, or alternatively (as assumed here) "," decimals" and ";" separators.

List of useful functions:

- read.table, read.csv
- data: data(vcaplong), data(vcapwide)

3 Distance and dissimilarity functions

A distance or dissimilarity function is used to describe the amount of (dis-)similarity between two experimental units. Common choices for computing the amount of similarity between two vectors \mathbf{x} and \mathbf{y} include:

- Euclidean distance: $d(\mathbf{x}, \mathbf{y}) = \sqrt{(x_1 y_1)^2 + (x_2 y_2)^2 + \dots + (x_n y_n)^2} = \sqrt{\sum_{i=1}^n (x_i y_i)^2}.$
- Standardized Euclidean distance: $\sqrt{\sum_{i=1}^{N} \frac{(x_i y_i)^2}{s_i^2}}$

• Mahalanobis distance: $\sqrt{(x-y)^T S^{-1}(x-y)}$

Here, \mathbf{x} and \mathbf{y} are expected to be observation vectors of length N, where each dimension describes the measured value for a particular covariate. S describes the covariance-variance matrix between covariates, and therefore incorporates inter-correlations between variables. The standard deviation s may be used to standardize differences in variation over the dimensions.

			9				7	0	0	10	11	10	1.0	1.1	15	1.0	1.77	10
	1	2	3	4	5	6		8	9	10	11	12	13	14	15	16	17	18
1	0.00	18.05	2.80	10.32	13.53	7.87	9.00	10.08	14.38	17.28	15.40	8.61	3.58	9.76	18.23	17.33	9.21	14.62
2	18.05	0.00	20.14	8.05	5.23	14.78	9.34	8.04	3.99	5.27	4.33	10.15	21.60	8.66	1.36	34.81	26.51	3.98
3	2.80	20.14	0.00	12.29	15.89	10.64	11.35	12.30	16.50	19.79	17.82	10.70	2.89	12.08	20.39	14.87	6.67	16.92
4	10.32	8.05	12.29	0.00	4.44	9.12	2.53	1.62	4.29	8.90	6.47	2.42	13.82	3.55	8.20	26.84	18.54	5.39
5	13.53	5.23	15.89	4.44	0.00	9.61	4.59	3.68	2.19	4.48	2.08	5.83	16.97	4.45	5.02	30.61	22.33	1.66
6	7.87	14.78	10.64	9.12	9.61	0.00	6.60	7.91	11.39	11.95	10.86	7.56	10.10	7.33	14.57	24.16	16.49	10.84
7	9.00	9.34	11.35	2.53	4.59	6.60	0.00	1.47	5.54	8.71	6.57	2.04	12.43	2.58	9.34	26.03	17.75	5.85
8	10.08	8.04	12.30	1.62	3.68	7.91	1.47	0.00	4.33	7.98	5.70	2.70	13.59	2.19	8.15	27.04	18.73	4.73
9	14.38	3.99	16.50	4.29	2.19	11.39	5.54	4.33	0.00	5.57	3.20	6.20	17.87	5.55	3.93	31.12	22.81	2.29
10	17.28	5.27	19.79	8.90	4.48	11.95	8.71	7.98	5.57	0.00	2.48	10.19	20.60	7.98	4.77	34.56	26.32	3.82
11	15.40	4.33	17.82	6.47	2.08	10.86	6.57	5.70	3.20	2.48	0.00	7.91	18.81	6.05	3.96	32.58	24.30	1.58
12	8.61	10.15	10.70	2.42	5.83	7.56	2.04	2.70	6.20	10.19	7.91	0.00	11.96	4.34	10.10	25.09	16.82	7.14
13	3.58	21.60	2.89	13.82	16.97	10.10	12.43	13.59	17.87	20.60	18.81	11.96	0.00	13.27	21.73	14.19	6.53	18.11
14	9.76	8.66	12.08	3.55	4.45	7.33	2.58	2.19	5.55	7.98	6.05	4.34	13.27	0.00	8.95	26.95	18.69	5.07
15	18.23	1.36	20.39	8.20	5.02	14.57	9.34	8.15	3.93	4.77	3.96	10.10	21.73	8.95	0.00	35.04	26.73	4.05
16	17.33	34.81	14.87	26.84	30.61	24.16	26.03	27.04	31.12	34.56	32.58	25.09	14.19	26.95	35.04	0.00	8.31	31.72
17	9.21	26.51	6.67	18.54	22.33	16.49	17.75	18.73	22.81	26.32	24.30	16.82	6.53	18.69	26.73	8.31	0.00	23.42
18	14.62	3.98	16.92	5.39	1.66	10.84	5.85	4.73	2.29	3.82	1.58	7.14	18.11	5.07	4.05	31.72	23.42	0.00

Table 1: Euclidean distance matrix D for 18 animals

Table 1 shows the Euclidean distance matrix for the 18 animals presented in Figure 2.

List of useful functions:

- dist includes many common distance and dissimilarity functions (Euclidean by default, others: method="manhattan", method="maximum", method="minkowski"
- cluster::daisy, daisy includes Gower's dissimilarity for mixed data (parameter metric="gower")

4 Non-bipartite optimal matching of animals at baseline

The non-bipartite optimal matching problem may be solved using the provided branch and bound algorithm:

```
> sol <- match.bb(d, g=3)
```

- [1] "Performing initial sorting for a good initial guess"
- [1] "Computing boundaries for minimum distances in possible combinations..."
- [1] "Starting branch and bound"
- [1] "Branches: 272"
- [1] "Bounds: 7140"
- [1] "Ends visited: 25"
- [1] "Solution cost 169.62"
- [1] "Solution: 5,3,5,6,4,5,6,4,3,2,2,6,1,4,3,1,1,2"

```
> submatches <- paste("Submatch_", LETTERS[1:6][sol$solution], sep="")
> names(submatches) <- names(sol$solution)</pre>
> submatches
                         2
                                      3
                                                                  5
           1
"Submatch_E" "Submatch_C" "Submatch_E" "Submatch_F" "Submatch_D"
                                                                    "Submatch_E"
           7
                        8
                                      9
                                                   10
                                                                 11
                                                                               12
"Submatch_F" "Submatch_D"
                          "Submatch_C" "Submatch_B"
                                                      "Submatch_B"
          13
                       14
                                     15
                                                   16
                                                                 17
"Submatch_A" "Submatch_D" "Submatch_C" "Submatch_A" "Submatch_A" "Submatch_B"
```

The match.bb function returns the solution to the optimal matching task. It takes as input a distance matrix d, as is indicated in the function call match.bb(d, g=3) (notice that d was defined before). Furthermore, the size of the submatches is defined using the parameter g=3. This value indicates that the optimal matching algorithm minimizes edges within triplets. Each observation has to belong to a triplet called a submatch.

List of useful functions:

- Multigroup non-bipartite matching: hamlet::match.bb
- Paired non-bipartite matching: hamlet::match.bb, nbpMatching::nonbimatch
- Paired bipartite matching: optmatch::fullmatch

5 Randomization based on matched individuals

The submatches identified in the above section should not be mistaken for the randomly allocated intervention groups. The final intervention groups are obtained by dividing members of each submatch in the found solution to a separate treatment arm. Since the within-submatch distances are minimized, this guarantees that comparable individuals are randomly divided to separate arms:

```
> ex[,"Submatch"] <- submatches
> set.seed(1) # for reproducibility
> ex[,"AllocatedGroups"] <- match.allocate(ex[,"Submatch"])</pre>
```

List of useful functions:

- Multigroup non-bipartite matching: hamlet::match.bb
- Paired non-bipartite matching: hamlet::match.bb, nbpMatching::nonbimatch
- Paired bipartite matching: optmatch::fullmatch

6 Visualizations for pre-clinical data

Various visualization functions are available to illustrate baseline balance. For example, the boxplots in respect to allocation groups can be plotted using a command such as boxplot, which is illustrated in Figure 3.

```
> boxplot(PSA.week.10..ug.1. ~ AllocatedGroups, data = ex, range=0, + xlab="Group", ylab="PSA week 10 ul/g")
```

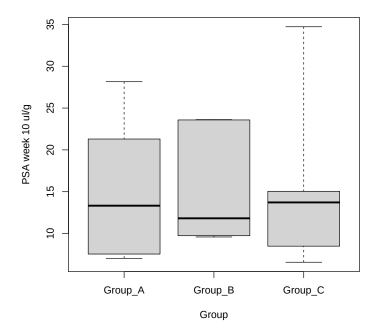


Figure 3: Boxplots for the week $10~\mathrm{PSA}$ in the example allocation

> mixplot(ex[,2:5], pch=16)

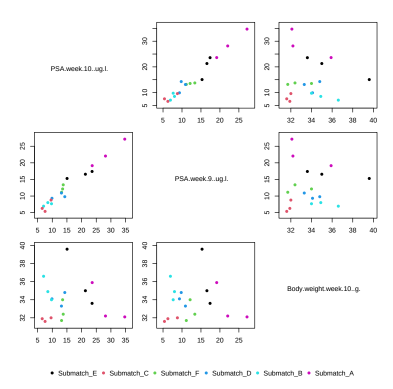


Figure 4: Test mixplot with submatch labels

> mixplot(ex[,c(2:4,6)], pch=16)

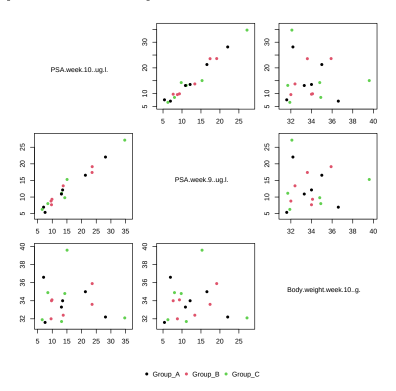


Figure 5: Test mixplot with allocation group labels

	Animal	PSA.week.10ug.l.	PSA.week.9ug.l.	Body.weight.week.10g.	Submatch	AllocatedGroups
1	ID003	21.30	16.57	35.00	Submatch E	Group A
2	ID007	7.55	5.38	31.60	Submatch C	Group A
3	ID008	23.58	17.40	33.60	Submatch E	Group B
4	ID009	13.17	11.14	31.70	Submatch_F	Group_C
5	ID010	9.90	9.33	34.10	$Submatch_D$	$Group_B$
6	ID016	15.05	15.29	39.60	$\operatorname{Submatch}_{-}\operatorname{E}$	$\operatorname{Group}_{\operatorname{\mathbb{C}}}$
7	ID018	13.53	12.14	34.00	$Submatch_F$	Group_A
8	ID025	13.13	10.91	33.30	$Submatch_D$	$\operatorname{Group}_{-}A$
9	ID027	9.59	8.79	32.00	$Submatch_C$	$Group_B$
10	ID031	7.04	6.95	36.60	$Submatch_B$	$\operatorname{Group}_{-}A$
11	ID032	8.49	8.02	34.90	$Submatch_B$	$Group_C$
12	ID037	13.74	13.38	32.40	$Submatch_F$	$Group_B$
13	ID040	23.62	19.15	35.90	$Submatch_A$	$Group_B$
14	ID045	14.27	9.80	34.80	$Submatch_D$	$Group_C$
15	ID047	6.57	6.28	31.90	$Submatch_C$	$Group_C$
16	ID054	34.72	27.14	32.10	$\operatorname{Submatch} A$	$\operatorname{Group} \overline{\ \ } \operatorname{C}$
17	ID056	28.15	22.05	32.20	$Submatch_A$	$\operatorname{Group}_{-}A$
18	ID058	9.74	7.68	34.00	$Submatch_B$	$Group_B$

Table 2: The result table in variable ex after performing the optimal matching and allocation.

Mixed variable scatterplots with annotations for the submatches or allocation groups are plotted using the function hamlet::mixplot, which can be seen in Figures 4 or 5 respectively.

A common way to illustrate distance matrices is through heatmaps, along with hierarchical clustering to connect similar individuals (Figure 6).

List of useful functions:

- Scatterplots etc: hamlet::mixplot, plot, boxplot
- Heatmaps: hamlet::hmap, heatmap, gplots::heatmap.2

7 Paired vs. non-paired testing

Pairing of samples aims to increase statistical power of tests through connecting the same measurement over two conditions. In the pre-clinical context, we incorporate the matching information from prior to interventions (Figure 1) in order to find the connecting measurements. As the submatches were used to couple measurements prior to interventions, it is natural to use this information for pairing the observations in the post-intervention tests.

If pairing is not performed, the statistical power may not be sufficient in the tests, as the populations are assumed to relatively homogeneous and no prognostic information is incorporated to the testing procedure:

> heatmap(d)

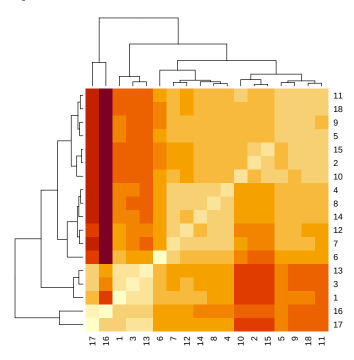


Figure 6: Heatmap for the 18x18 distance matrix

```
data: veh[, "PSAWeek14"] and mdv[, "PSAWeek14"]
t = 1.7385, df = 26.796, p-value = 0.0936
alternative hypothesis: true difference in means is not equal to 0
95 percent confidence interval:
    -3.18889 38.49813
sample estimates:
mean of x mean of y
53.39533 35.74071
```

However, the submatches are used to couple best comparable measurements. For example, in the VCaP experiment the week 10 was baseline and its PSA and body weight contain prognostic information necessary for weighting post-intervention tests. During the experiment weeks 11 to 14 interventions took place. The column "Submatch" given in the datasets vcapwide and vcaplong are the submatches that occurred in the actual matching and randomization task when the experiment itself was performed. The week 10 information in connection to the submatches were:

```
> veh <- veh[order(veh[,"Submatch"]),]
> mdv <- mdv[order(mdv[,"Submatch"]),]</pre>
```

	Veh.PSAWeek10	MDV.PSAWeek10
Submatch_1	21.30	16.25
$Submatch_10$	8.13	7.55
$Submatch_11$	8.36	13.53
$Submatch_12$	15.05	9.96
$Submatch_13$	7.03	6.57
$Submatch_14$	34.72	27.16
$Submatch_16$	9.13	10.78
$Submatch_2$	13.17	13.13
$Submatch_3$	22.29	23.58
$Submatch_4$	14.27	9.74
$Submatch_5$	13.74	9.59
$Submatch_6$	8.49	9.89
$Submatch_7$	16.29	19.51
$Submatch_8$	20.99	16.39
$_Submatch_9$	14.69	18.71

Table 3: Submatches in the real VCaP experiment, per PSA at week 10 in tumors allocated to the Vehicle and MDV groups

Tables 3 and 4 show the prognostic baseline information in terms of PSA and body weight that is connected to the submatches. These submatches are used for pairing observations in comparing MDV intervention to the vehicle, which results in a noticeable increase in statistical power in even conventional tests:

```
> t.test(veh[,"PSAWeek14"], mdv[,"PSAWeek14"], paired=TRUE)

Paired t-test

data: veh[, "PSAWeek14"] and mdv[, "PSAWeek14"]
```

	Veh.BWWeek10	MDV.BWWeek10
Submatch_1	35.00	34.80
$Submatch_10$	26.70	31.60
$Submatch_11$	35.60	34.00
$Submatch_12$	39.60	33.40
$Submatch_13$	30.00	31.90
$Submatch_14$	32.10	28.30
$Submatch_16$	29.90	29.20
$Submatch_2$	31.70	33.30
$Submatch_3$	31.10	33.60
$Submatch_4$	34.80	34.00
$Submatch_5$	32.40	32.00
$Submatch_6$	34.90	38.30
$Submatch_7$	30.00	29.90
$Submatch_8$	28.40	30.10
Submatch_9	28.30	33.30

Table 4: Submatches in the real VCaP experiment, per body weight at week 10 in tumors allocated to the Vehicle and MDV groups

For more refined statistical testing, the longitudinal profiles of a pre-clinical experiment can be modeled using the lme4-package. Functions are provided in the hamlet-package for this purpose.

List of useful functions:

- MEM-modeling packages: lme4, lmerTest, nlme
- MEM-modeling in hamlet: hamlet::mem.getcomp, hamlet::mem.plotran, hamlet::mem.plotresid