Scanpath – An R Package for Analyzing Scanpaths

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Contents

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An R package for analyzing scanpaths in eye movement data. The package includes a simple toy dataset and example code. Consult von der Malsburg & Vasishth (2011) for the details of this analysis method. The manual of the package can be found here and a PDF-version of this page here.

1 News

[2018-02-20 Tue] The new default is no normalization of scasim scores. Improved documentation and examples for find.fixation and match.scanpath. Minor improvement in functions for plotting scanpaths.

[2018-01-30 Tue] Version 1.06 doesn't logarithmize fixation durations when calculating scanpath similarities. (In previous versions, when normalize="durations" was used, the normalization was done using non-log-transformed durations, which could in some rare cases break the triangle inequality.)

2 Install

To install the latest version of the package, execute the following commands:

```
library("devtools");
install_github("tmalsburg/scanpath/scanpath", dependencies=TRUE)
```

3 Usage example

The code shown below can also be found in the file README.R. Open that file in RStudio and play with it as you read through this mini-tutorial.

Let's have a look at the toy data that is included in the package:

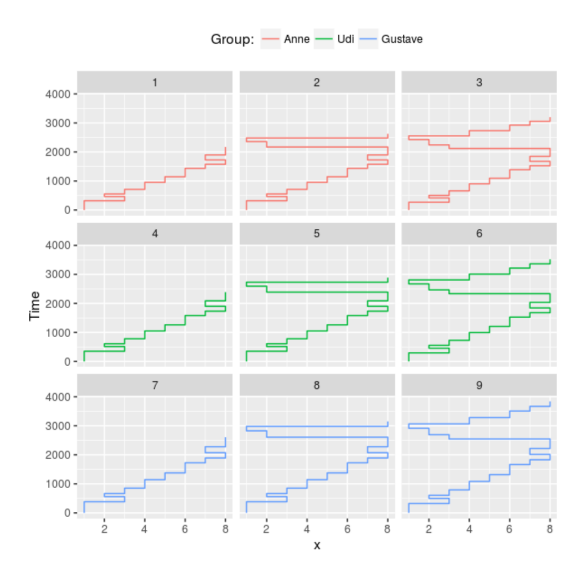
library(tidyverse)
library(magrittr)
library(scanpath)
data(eyemovements)
head(eyemovements)

subject	trial	word	Χ	у	duration
Anne	1	1	46	384	319
Anne	1	3	131	388	147
Anne	1	2	106	386	88
Anne	1	3	165	387	156
Anne	1	4	186	386	244
Anne	1	5	264	388	193

3.1 Plotting scanpaths

To get a sense of what is going on in this data set, we create a series of plots. For this purpose, we use the function plot_scanpaths from the *scanpath* package. In the first plot below, each panel shows the data from one trial. There are three participants which are coded by color. The data is from a sentence reading task. The x-axis shows words and the y-axis time within trial in milliseconds.

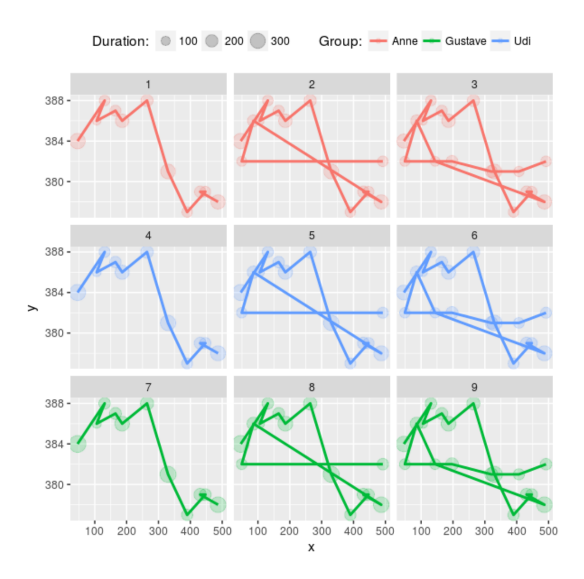
```
plot_scanpaths(duration ~ word | trial, eyemovements, subject)
```



We can see that the participants differ in their reading speed. Also we see that each participant read the sentence more or less straight from left to right (trials: 1, 4, 7), or with a short regressions from the end of the sentence to its beginning (trials: 2, 5, 8), or with a long regression from the end of the sentence (trials: 3, 6, 9).

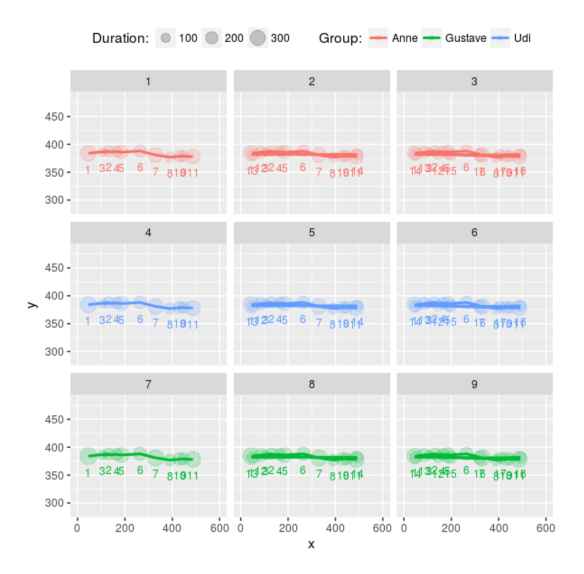
In the next plot, we use the fixations' x- and y-coordinates. Each circle is a fixation and the size of the circle represents the duration of the corresponding fixation.

plot_scanpaths(duration ~ x + y | trial, eyemovements, subject)



The function plot_scanpaths returns a *ggplot* object. This means that we add more elements to the plot before rendering. For example, we can labels the fixations with their index and change the limits of the axes:

```
plot_scanpaths(duration ~ x + y | trial, eyemovements, subject) +
  geom_text(aes(label=i), vjust=2.5, show.legend=FALSE, size=3) +
  xlim(0, 600) + ylim(284, 484)
```



3.2 Extracting subsets of fixations or sub-scanpaths

In many analyses, it is not desirable to analyze the complete scanpaths as recorded during the experiment but to analyze some subset of the fixations. For instance, in a reading experiment, we might want to investigate how readers responded to a certain word and not care about what happened earlier. The scanpath package offers two functions that can be used to easily pinpoint and extract the fixations of interest: find.fixation and match.scanpath.

The function find.fixation identifies fixations that match a set of criteria which can be specified using regular expressions. For instance, the following code finds fixations on word 6:

idx <- find.fixation(eyemovements\$word, eyemovements\$trial, "6")
eyemovements[idx,]</pre>

subject	trial	word	Χ	У	duration
Anne	1	6	330	381	290
Anne	2	6	330	381	290
Anne	3	6	330	381	290
Anne	3	6	320	381	189
Udi	4	6	330	381	319
Udi	5	6	330	381	319
Udi	6	6	330	381	319
Udi	6	6	320	381	208
Gustave	7	6	330	381	348
Gustave	8	6	330	381	348
Gustave	9	6	330	381	348
Gustave	9	6	320	381	227

Finding these fixations could also have been achieved with a subset operation. However, if have more complex criteria for the fixations we're interested in, things can get rather tricky. For instance, a subset is not enough when we're only interested in the second fixation on word 6 in each trial. The following code extracts only those:

idx <- find.fixation(eyemovements\$word, eyemovements\$trial, "6", nth=2)
eyemovements[idx,]</pre>

subject	trial	word	Х	у	duration
Anne	3	6	320	381	189
Udi	6	6	320	381	208
Gustave	9	6	320	381	227

Regular expressions also allow us to specify the context in which the fixations of interest appear. For instance the code below finds fixations on word 3 but only those that are followed by fixations on word 4:

idx <- find.fixation(eyemovements\$word, eyemovements\$trial, "34")
eyemovements[idx,]</pre>

subject	trial	word	Χ	у	duration
Anne	1	3	165	387	156
Anne	2	3	165	387	156
Anne	3	3	165	387	156
Udi	4	3	165	387	172
Udi	5	3	165	387	172
Udi	6	3	165	387	172
Gustave	7	3	165	387	187
Gustave	8	3	165	387	187
Gustave	9	3	165	387	187

Here, we find fixations on word 3 that are preceded by fixations on word 1:

idx <- find.fixation(eyemovements\$word, eyemovements\$trial, "1(3)", subpattern=1)
eyemovements[idx,]</pre>

subject	trial	word	Х	у	duration
Anne	1	3	131	388	147
Anne	2	3	131	388	147
Anne	3	3	131	388	147
Udi	4	3	131	388	162
Udi	5	3	131	388	162
Udi	6	3	131	388	162
Gustave	7	3	131	388	176
Gustave	8	3	131	388	176
Gustave	9	3	131	388	176

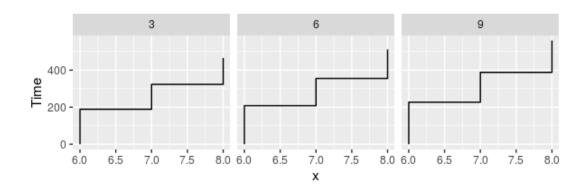
The following code finds fixations on the last word but only of those that are not directly preceded by fixations on words 4 to 7:

 $idx <- find.fixation(eyemovements$word, eyemovements$trial, "[^4-7](8)", subpattern=1 eyemovements[idx,]$

subject	trial	word	Χ	у	duration
Anne	2	8	492	382	143
Udi	5	8	492	382	157
Gustave	8	8	492	382	172

The function match.scanpath works similarly but can be used to identify not just individual fixations but sequences of fixations (let's call them scanpathlets). For example, the following code finds scanpathslets spanning words 6, 7, and 8 but only those that directly preceded by a fixation on word 4:

idx <- match.scanpath(eyemovements\$word, eyemovements\$trial, "4([678]+)", subpattern=
scanpathlets <- eyemovements[idx,]
plot_scanpaths(duration~word|trial, scanpathlets)</pre>



See the documentation of find.fixation and match.scanpath for more details and examples.

3.3 Calculating scanpath dissimilarities

Next, we calculate the pair-wise similarities of the nine scanpaths in the dataset using the *scasim* measure. A simplifying intuition is that the measure quantifies the time that was spent looking at different things (or at the same things but in different order). For a precise definition see von der Malsburg & Vasishth (2011).

d1 <- scasim(eyemovements, duration ~x + y | trial, 512, 384, 60, 1/30) round(d1)

	1	2	3	4	5	6	7	8	9
1	0	454	1129	217	717	1395	435	980	1670
2	454	0	675	671	263	941	889	526	1216
3	1129	675	0	1346	938	320	1564	1201	641
4	217	671	1346	0	500	1242	218	763	1509
5	717	263	938	500	0	742	718	263	1009
6	1395	941	320	1242	742	0	1460	1005	321
7	435	889	1564	218	718	1460	0	545	1355
8	980	526	1201	763	263	1005	545	0	810
9	1670	1216	641	1509	1009	321	1355	810	0

Like the function plot_scanpaths, the function scasim takes a formula and a data frame as parameters. The formula specifies which columns in the data frame should be used for the calculations. To account for distortion due to visual perspective, the comparison of the scanpaths is carried out in visual field coordinates (latitude and longitude). In order to transform the pixel coordinates provided by the eye-tracker to visual field coordinates, the scasim function needs some extra information. The first is the position of the gaze when the participant looked straight ahead (512, 384, in the present case), the distance of the eyes from the screen (60 cm), and the size of one pixel in the unit that was used to specify the distance from the screen (1/30). Finally, we have to specify a normalization procedure. normalize=FALSE means that we don't want to normalize. See the documentation of the scasim function for details.

The time that was spent looking at different things of course depends on the duration of the two compared trials. (total duration of the two compared scanpaths constitutes an upper bound). This means that two long scanpaths may have a larger dissimilarity than two shorter scanpaths even if they look more similar. Depending on the research question, this may be undesirable. One way to get rid of the trivial influence of total duration is to normalize the dissimilarity scores. For example, we can divide them by the total duration of the two compared scanpaths:

	1	2	3	4	5	6	7	8	9
1	0	9	21	5	14	25	9	18	28
2	9	0	12	13	5	15	17	9	19
3	21	12	0	24	15	5	27	19	9
4	5	13	24	0	9	21	4	14	24
5	14	5	15	9	0	12	13	4	15
6	25	15	5	21	12	0	24	15	4
7	9	17	27	4	13	24	0	9	21
8	18	9	19	14	4	15	9	0	12
9	28	19	9	24	15	4	21	12	0

The number are smaller now and can be interpreted as the percentage of time that was spent looking at different things.

3.4 Maps of scanpath space

The numbers in the matrix above capture a lot of information about the scanpath variance in the data set. However, dissimilarity scores are somewhat tricky to analyze. One problem is that these values have strong statistical dependencies. When we change one scanpath, this affects n dissimilarity scores. This has to be kept in mind when doing inferential stats directly on the dissimilarity scores. While there are solutions for this, it is typically more convenient to produce a representation of scanpath variance that is free from this problem. One such representation is what we call the "map of scanpath space." On such a map, every point represents a scanpath and the distances on the map reflect the dissimilarities according to our scanpath measure, i.e. the dissimilarity scores in the matrix above.

The method for calculating these maps is called multi-dimensional scaling and one simple version of the general idea is implemented in the function <code>cmdscale</code> (see also <code>isoMDS</code> in the MASS package).

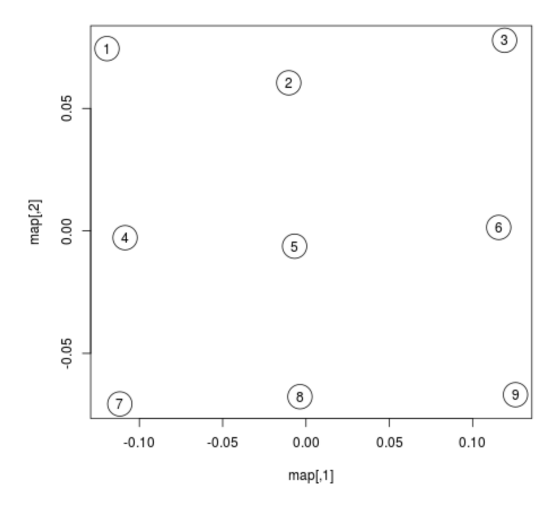
```
map <- cmdscale(d2)
round(map, 2)</pre>
```

	V1	V2
1	-0.12	-0.07
2	-0.01	-0.06
3	0.12	-0.08
4	-0.11	0
5	-0.01	0.01
6	0.12	0
7	-0.11	0.07
8	0	0.07
9	0.13	0.07

The table above contains two numbers for each scanpath in the data set. These numbers (V1 and V2) determine a scanpath's location in the two-dimensional scanpath space created by cmdscale. How many dimensions we need is an empirical question.

Below is a plot showing the map of scanpaths:

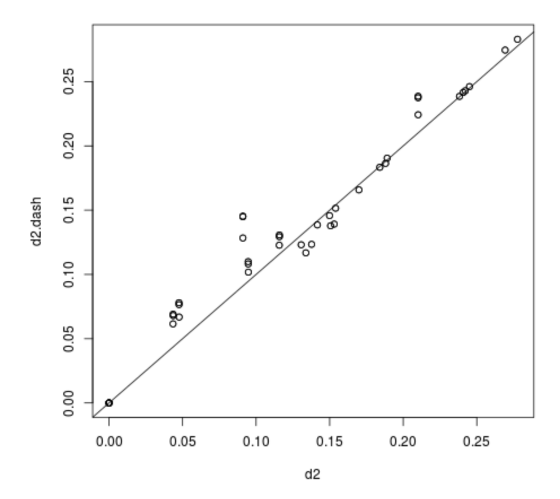
```
map <- map %*% matrix(c(1, 0, 0, -1), 2) # flip y-axis
plot(map, cex=4)
text(map, labels=rownames(map))</pre>
```



Interestingly, the scanpaths are arranged in the same way as in the plot of the data at the top. Participants are arranged vertically and reading patterns are horizontally. This suggests that *scasim* not just recovered these two different kinds of information (reading speed and reading strategy) but also that it can distinguish between them.

To test how well this map represents the original dissimilarity scores, we can calculate the pair-wise differences on the map and compare them to the pair-wise *scasim* scores:

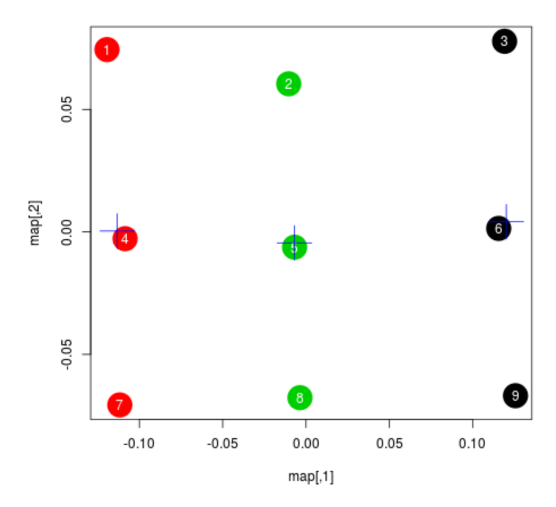
```
d2.dash <- as.matrix(dist(map))
plot(d2, d2.dash)
abline(0, 1)</pre>
```



This plot suggests that the map preserves the variance in dissimilarity scores really well. Given this very good fit of the map, it appears that two dimensions were sufficient to describe the scanpath variance that is captured by *scasim*. This is not surprising because the scanpaths in the toy data set were designed to vary with respect to two properties: 1.) The speed of the reader, and 2.) whether there was a regression back to the beginning of the sentence and how long it was.

The benefit of the map representation is that it has much weaker statistical dependencies and that it is much more suitable for all kinds of analyses. For example, we can choose among a large number of clustering algorithms to test whether there are groups of similar scanpaths in a data set. Below, we use the simple k-means algorithm to illustrate this:

```
set.seed(4)
clusters <- kmeans(map, 3, iter.max=100)
plot(map, cex=4, col=clusters$cluster, pch=19)
text(map, labels=rownames(map), col="white")
points(clusters$centers, col="blue", pch=3, cex=4)</pre>
```



In this plot, color indicates to which cluster a scanpath belongs and the crosses show the center of each cluster. We see that the clusters correspond to the different reading patterns and that participants are ordered according to their reading speed within the clusters.

Apart from cluster analyses there are many other ways to analyze scanpath variance. See the articles listed below for more details.

4 References

- von der Malsburg, T., & Vasishth, S. (2011). What is the scanpath signature of syntactic reanalysis? Journal of Memory and Language, 65(2), 109–127. http://dx.doi.org/10.1016/j.jml.2011.02.004
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