DREADDs PST2020

EP

03/02/2021

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1 Introduction

A behavioral study of foraging

The data were acquired by JS, CG, and CREW at Oxford univ and at Inserm Lyon. In Lyon, Donut was tested at U1028 and Homer and Dali at U1208. The animals were tested everyday with different versions of the apparatus. The testing was done in animal housing using the 1st design realized by JS:

On this board the 25 locations are numbered from 1 to 25 from the upper left corner to the lower right corner, going from left to right.

1.1 Data loading and formating

Data files include all data tested in monkeys in Oxford and in Lyon.

This Markdown deals mostly with data from HOMER and DONUT.¹

session types are initially "0" for control (transparent doors) and "1" TEST with opaque blue doors, but we use the label Clear vs BLue in figures and analyses. Codes used for target chosen and repeats or miss reflect the position of the choice (location of the hole selected from 1 TO 25 top to bottom), if negative value then it's a repeat, if 999 then it's a pause in the task, if it's a value between 100 and 2500 then it is a mistake (the animal tried but missed the reward) the number /100 giving the location.

¹The first descriptive analyses were in Graph_PST2020choices.R - now in PST2020_DREADDs.rmd]

2 Main general Plots

Let first see descriptive graphs for all sessions per monkey. The figures show for each monkey the choices (location of choice on board) selected by the animal trial by trial (chronological order from left to right). Green dots represent correct choices i.e. location chosen for the 1st time in session and with correct pick up of the reward. Blue dots represent returns to previously chosen location. These are presented as negative values to show their time course independently for correct choices. Orange is when reward is missed. Clear (transparent doors) and Blue (blue doors) sessions are presented separately First sample sessions:

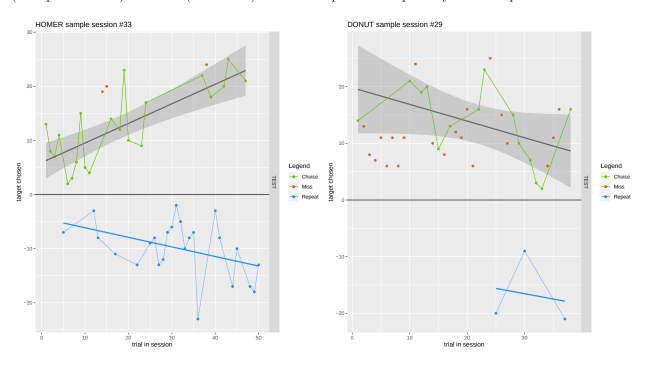
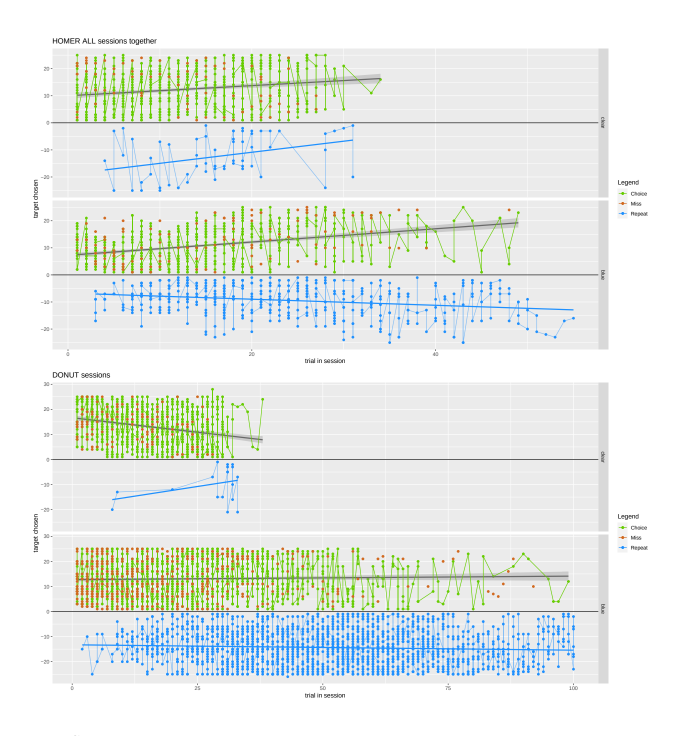


Figure 1: General figure HOMER

Then all data overlap across sessions to see the tendencies. In particular one can see the positive and negative trends that reflect monkeys choosing holes from top to bottom or bottom to top. Homer and Donut have different preferred directions but this is due to the different position of the setup in the housing.



2.1 Summary

The data are then summarized in terms of frequency of each trial (choice) type: Correct, Miss, Repeat.

Below the average number of each trial type for the different 'portes' (door) conditions.

We do not look at injections yet, this will be done later in the statistical analyses (DCZ vs Sham). See for all 8 monkeys or for Donut and Homer that there is a main effect of doors in particular on the number of repeats. Which makes sense because monkeys in blue condition (compared to clear) monkeys need to rely on memory to avoid repeating; which obviously they don't really succeed. We see later the #repeat is a very relevant parameter.

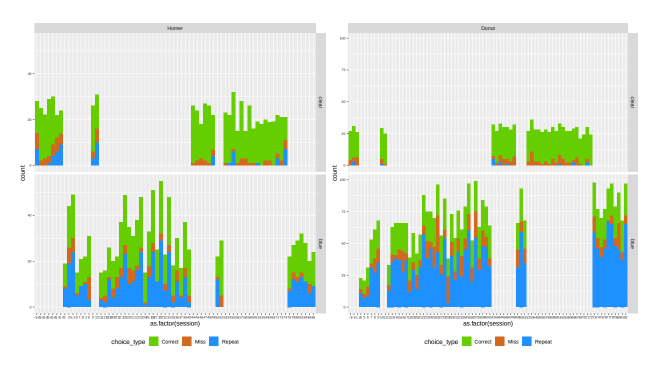
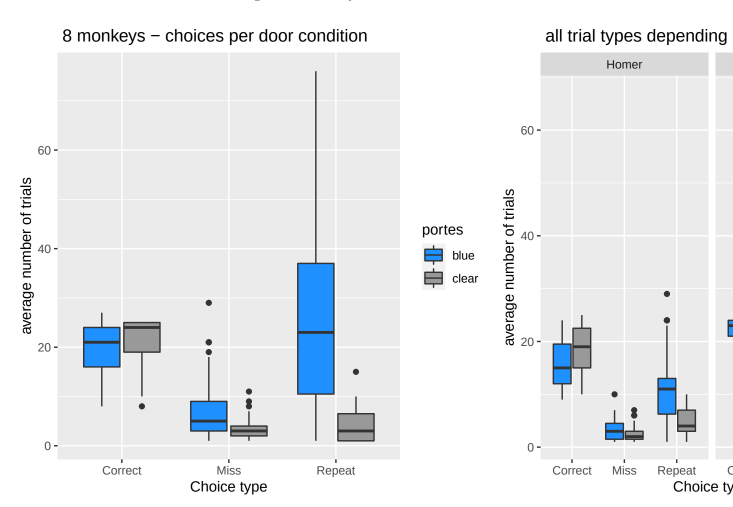


Figure 2: Summary all trials



##Stats We perform a logistic regression on the door effect for each monkey separately and test whether it influences the number of trial types - Still excluding DCZ sessions

```
##
## Call:
## glm(formula = trial ~ choice_type * portes, family = "poisson",
       data = subset(agg.data4BnoDCZ, singe = "Homer"))
##
## Deviance Residuals:
##
      Min
                     Median
                                  30
                1Q
                                          Max
## -6.4665 -1.5022 -0.1022
                              0.9405
                                        8.1527
##
## Coefficients:
##
                                Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                                 2.97972
                                            0.01729 172.361 < 2e-16 ***
                                            0.03624 -30.992 < 2e-16 ***
## choice_typeMiss
                                 -1.12302
## choice_typeRepeat
                                 0.24441
                                            0.02318 10.546
                                                             < 2e-16 ***
## portesclear
                                 0.09605
                                            0.02947
                                                      3.259 0.00112 **
## choice_typeMiss:portesclear
                                -0.79570
                                            0.07905 -10.065
                                                            < 2e-16 ***
                                            0.09901 -18.898 < 2e-16 ***
## choice_typeRepeat:portesclear -1.87109
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for poisson family taken to be 1)
##
##
      Null deviance: 6687.8 on 670 degrees of freedom
## Residual deviance: 3082.7 on 665 degrees of freedom
## AIC: 5915.2
##
## Number of Fisher Scoring iterations: 5
##
## Call:
  glm(formula = trial ~ choice_type * portes, family = "poisson",
       data = subset(agg.data4BnoDCZ, singe = "Donut"))
## Deviance Residuals:
                10
                     Median
                                  30
                                          Max
## -6.4665 -1.5022 -0.1022
                              0.9405
                                        8.1527
##
## Coefficients:
                                Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
                                 2.97972
                                            0.01729 172.361 < 2e-16 ***
## choice typeMiss
                                -1.12302
                                            0.03624 -30.992 < 2e-16 ***
## choice_typeRepeat
                                 0.24441
                                            0.02318 10.546 < 2e-16 ***
## portesclear
                                 0.09605
                                            0.02947
                                                      3.259 0.00112 **
## choice_typeMiss:portesclear
                                -0.79570
                                            0.07905 -10.065 < 2e-16 ***
## choice_typeRepeat:portesclear -1.87109
                                            0.09901 -18.898 < 2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for poisson family taken to be 1)
##
##
       Null deviance: 6687.8 on 670 degrees of freedom
## Residual deviance: 3082.7 on 665 degrees of freedom
```

AIC: 5915.2

##

Number of Fisher Scoring iterations: 5

##Exploration strategies

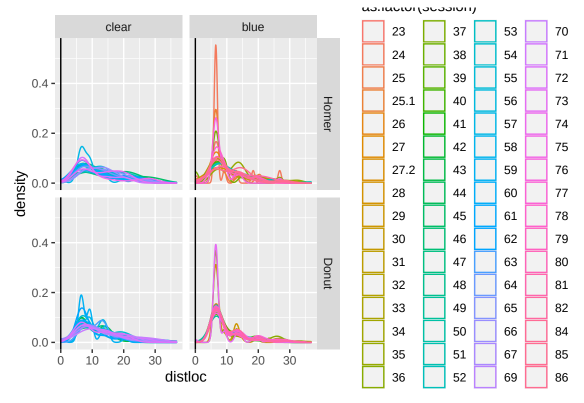
One future interesting set of analyses we can do regards the strategies of exploration: how animals scan through the setup , and then of course how they forget and repeat choices. Needs to be quantified to be used when comparing $ON/OFF\ DCZ$ sessions.

One thing we can look at is the spatial variance between successive choices (here I do not differentiate Miss, Correct or repeat).

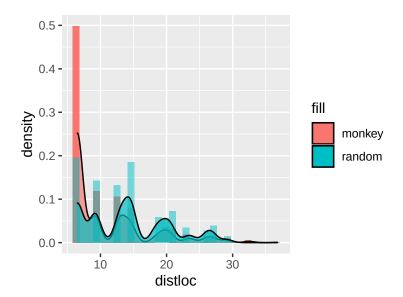
The figures below show the distributions of euclidean distances between 2 successive choices. The absolute distance between 2 holes (vertically and horizontally is 6.5cm). So we can see harmonics at 6.5 cm approximately in the distributions. Something quite obvious is that the harmonic is stronger in TEST than in CONTROL.

Note that here every choice is counted even repeats.

//////// ATTENTION: here we will select only sessions that are labelled no, sham or DCZ (i.e. >=23 for Homer and >=24 for Donut) ///////



The distribution of distances between choices is of a particular form, somewhat LogNormal. We can look at this distributions depending on conditions and also compare with a random sampling of distances. Let's first look at this accross the 2 monkeys.



The red shows distributions for monkeys, and blue shows a random sampling of 10000 distances.

Seperated for the 2 animals for Clear and Blue conditions we can see (below) that the distributions and the oscillation effects are stronger on Blue compared to Clear.

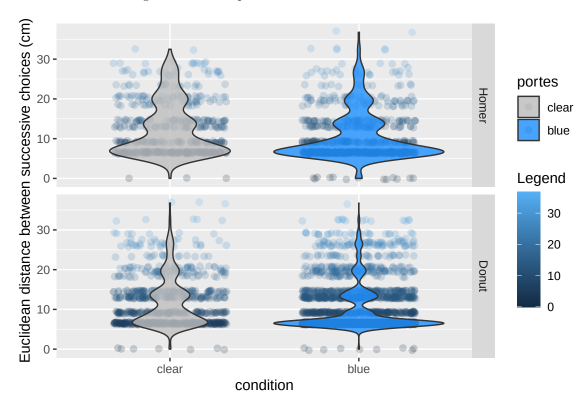


Figure 3: Spatial strategy. Distributions of euclidean distances

We test the difference in distributions between clear and blue for the 2 animals separately - we exclude DCZ sessions:

##

```
## Two-sample Kolmogorov-Smirnov test
##
## data: data4B$distloc[data4B$singe == "Homer" & data4B$portes == "clear" & data4B$Injection != "DCZ"
## D = 0.13605, p-value = 0.002588
## alternative hypothesis: two-sided
##
## Two-sample Kolmogorov-Smirnov test
##
## data: data4B$distloc[data4B$singe == "Donut" & data4B$portes == "clear" & data4B$Injection != "DCZ"
## D = 0.23376, p-value < 2.2e-16
## alternative hypothesis: two-sided</pre>
```

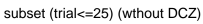
The KS tests indicate a difference between the 2 distributions (Clear and Blue) for both monkeys.

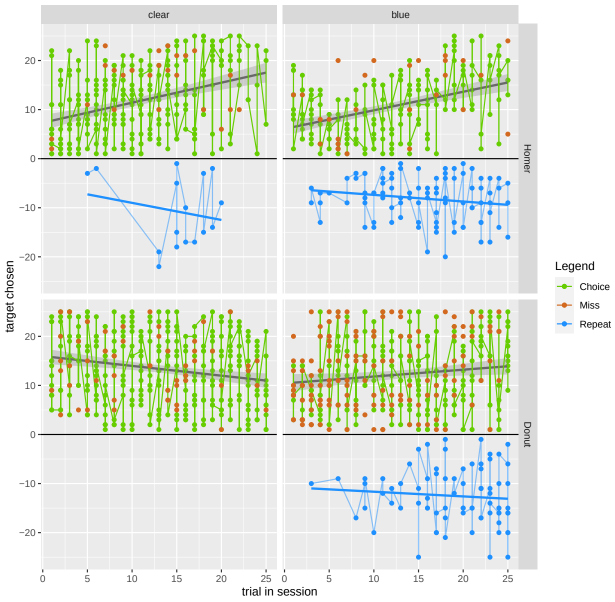
The strength of the 'harmonic' could be a marker of the two strategies used in the different sessions. The heavier harmonics in TEST could reflect an increased number of jumps between distant targets, whereas in CONTROL the animal would be more attracted to the visible reward which is just closeby to the current choice, hence proportionally more cases in which the animal choose the target just next the current one (6.5 cm distance). But we would need more control sessions to be sure.

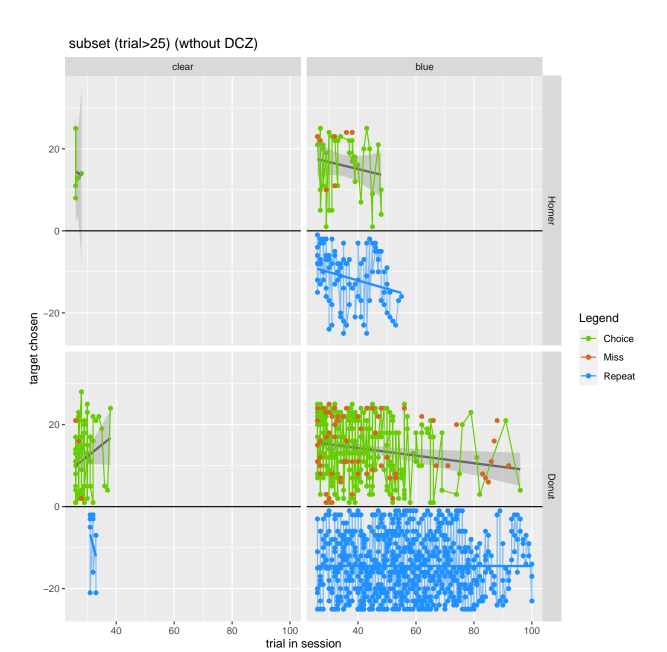
3 Subset of trials

One observation is that monkeys often behave in a somewhat more controlled, organized, manner at the beginning of a session and then choices become more dispersed. This could correlate approximately with the completion of the task (i.e. having gone through all locations). So here we separate the first 25 from the oher trials in 2 subset (25 corresponding to the number of locations on the setup).

Again here we remove DCZ (DCZ is taken into account in statistical analyses below)

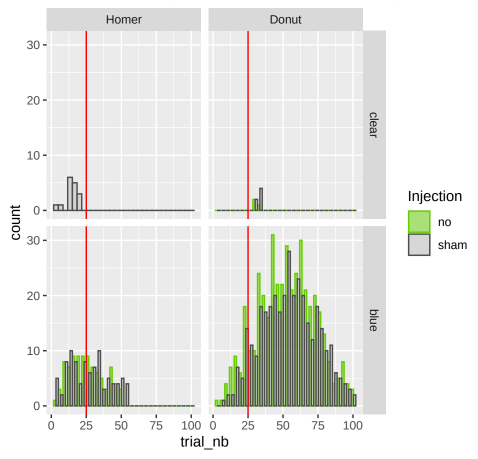






There is obviously a lot of repeats (checks?) after 25 when the animals continue trying to get rewards, and of course especially in the blue sessions. And Donut is a particularly good checker...

distribution of repeats in sessions - All trials (wthout DCZ



##The summary:

Below the average number of each trial type for the different conditions of injection (here again for the first 25 trials):

##Spatial strategy:

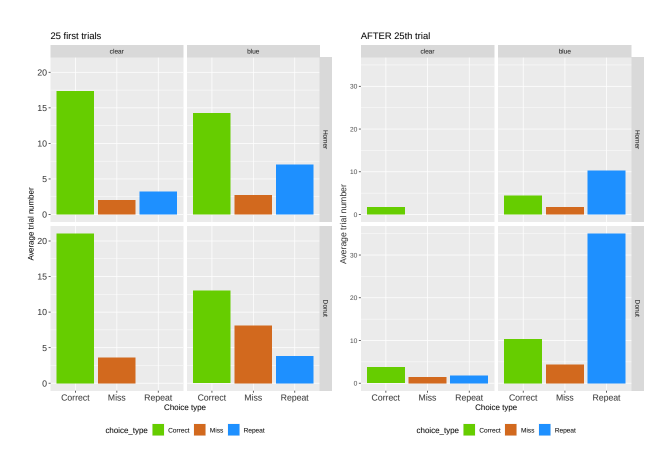
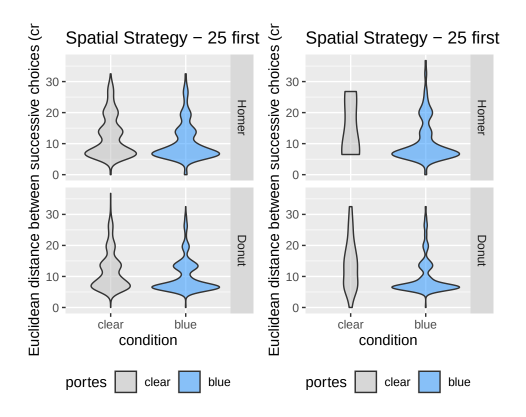
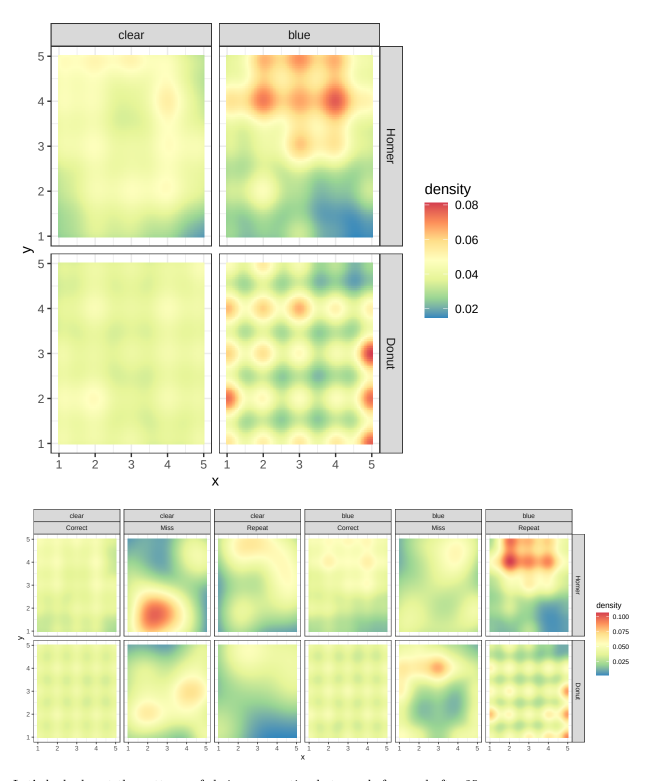


Figure 4: Injection type 25 trials

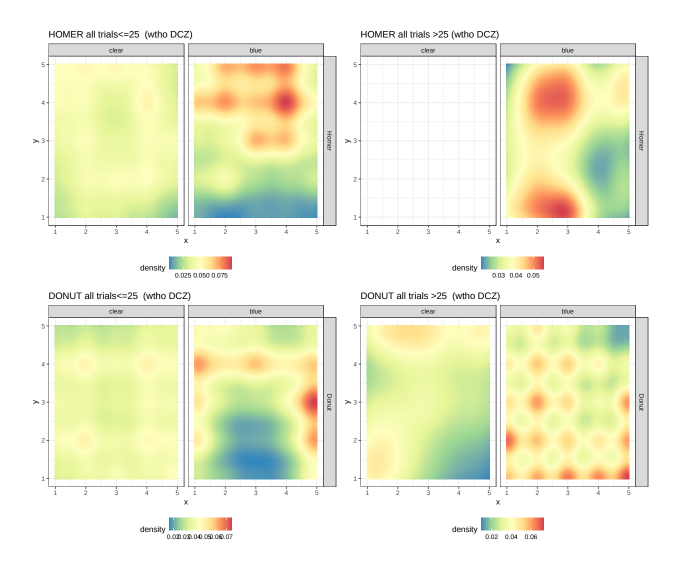


4 Spatial organization in 2D space

The tendencies for choosing some locations rather than others can reveal spatial biases. SO we use here a 2D density mapping of choices to look at that.



Let's look also at the patterns of choices separating between before and after 25:



5 STATISTICAL ANALYSES on DCZ vs Sham sessions

Here are the analyses and description of data for the 2 main types of sessions used on DCZ conditions. We subset the data for just the 2 monkeys and the 2 session types with an injection (sham and DCZ). We will also go through some more measures:

- test the numbers of repeats, and length distribution of distance to repeat
- test the number of pauses (code 99)

```
##
         = Homer
##
##
##
            sham DCZ
##
                8
     clear
                    7
                    9
##
     blue
##
##
         = Donut
##
##
```

```
## sham DCZ
## clear 7 10
## blue 10 11
```

##Descriptions of sessions

Here we answer a few general questions on the sessions, choices, repeats etc..

First, were there more trials (choices+misses+repeats) in DCZ compared to sham sessions?

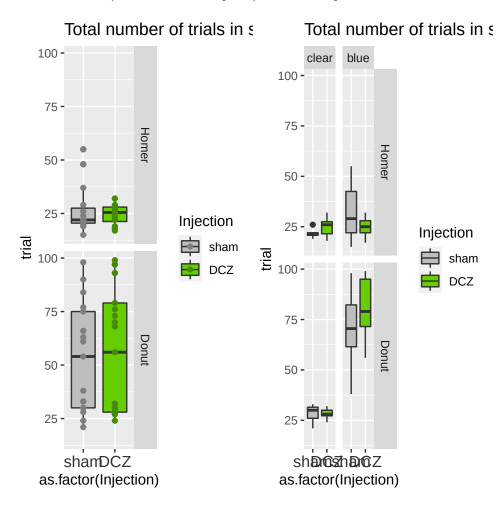
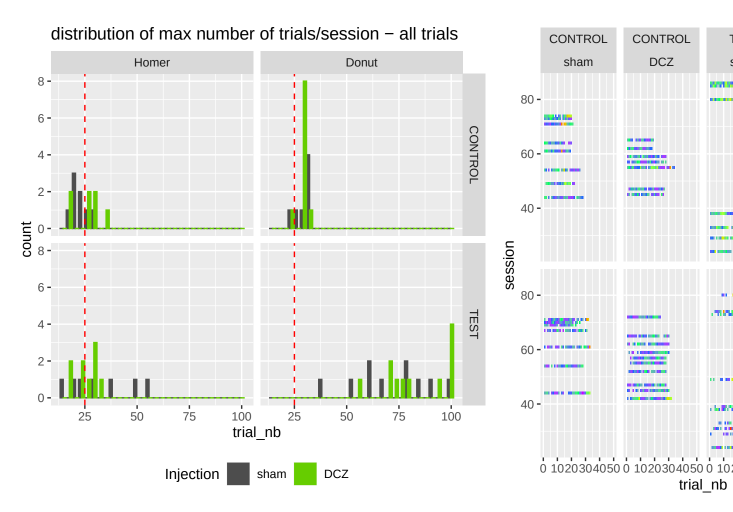


Figure 5: Summary 25 trials

```
##
## Call:
   glm(formula = trial ~ Injection * portes, family = "poisson",
       data = subset(agg.data4B, singe == "Homer"))
##
##
##
   Deviance Residuals:
                       Median
##
       Min
                                     3Q
                  1Q
                                             Max
##
   -3.4469
            -0.6071
                       0.0804
                                 0.6487
                                          3.5737
##
## Coefficients:
##
                            Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                             3.07385
                                         0.07603
                                                  40.430
                                                           < 2e-16 ***
                             0.13930
                                         0.10737
                                                    1.297
                                                           0.19450
## InjectionDCZ
```

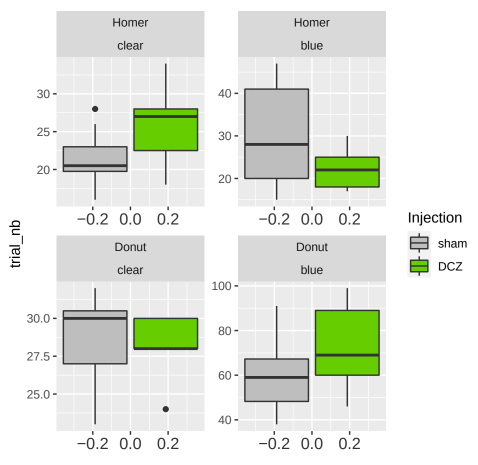
```
## portesblue
                          0.40959
                                    0.10083 4.062 4.86e-05 ***
## InjectionDCZ:portesblue -0.42179
                                    0.14296 -2.950 0.00317 **
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for poisson family taken to be 1)
      Null deviance: 74.002 on 30 degrees of freedom
##
## Residual deviance: 56.028 on 27 degrees of freedom
## AIC: 220.51
## Number of Fisher Scoring iterations: 4
##
## Call:
## glm(formula = trial ~ Injection * portes, family = "poisson",
      data = subset(agg.data4B, singe == "Donut"))
##
## Deviance Residuals:
                   Median
      Min
           1Q
                                3Q
                                        Max
## -4.2570 -0.8524 -0.0752 0.8145
                                     3.0783
##
## Coefficients:
##
                          Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                          ## InjectionDCZ
                         -0.001006 0.092446 -0.011
                                                      0.991
## portesblue
                          0.909635
                                    0.080259 11.334
                                                      <2e-16 ***
## InjectionDCZ:portesblue 0.149585
                                    0.105226
                                             1.422
                                                       0.155
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for poisson family taken to be 1)
##
      Null deviance: 507.290 on 37 degrees of freedom
## Residual deviance: 75.095 on 34 degrees of freedom
## AIC: 300.41
## Number of Fisher Scoring iterations: 4
```



No, apparently no main effect for Injection on the length of sessions. We have a significant interaction for Homer, suggesting a lower number of trials in DCZ (shorter sessions) in the blue door condition.

Then we ask whether the last correct trial performed is later in DCZ than in sham: this would mean that monkeys have more problems, or take more time, to find all or the max of rewards.

Last correct trial in sessions



```
##
## Call:
  glm(formula = trial_nb ~ Injection * portes, family = "poisson",
       data = subset(agg.maxcor, singe == "Homer"))
##
##
## Deviance Residuals:
      Min
##
                     Median
                1Q
                                   3Q
                                           Max
  -3.0810 -1.0171 -0.1082
                               0.7107
                                        2.8074
##
## Coefficients:
##
                           Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                            3.06805
                                       0.07625
                                              40.237 < 2e-16 ***
                                       0.10663
## InjectionDCZ
                            0.17899
                                                 1.679 0.093215 .
## portesblue
                            0.34262
                                       0.10262
                                                 3.339 0.000842 ***
## InjectionDCZ:portesblue -0.46877
                                       0.14487 -3.236 0.001213 **
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for poisson family taken to be 1)
##
##
      Null deviance: 67.110 on 30 degrees of freedom
## Residual deviance: 53.568 on 27 degrees of freedom
## AIC: 216.99
```

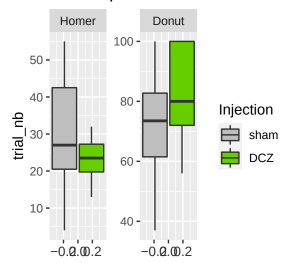
```
##
## Number of Fisher Scoring iterations: 4
##
## Call:
   glm(formula = trial_nb ~ Injection * portes, family = "poisson",
##
       data = subset(agg.maxcor, singe == "Donut"))
##
##
  Deviance Residuals:
                      Median
##
       Min
                 1Q
                                   30
                                            Max
                     -0.0752
##
   -3.3930
            -0.9278
                               0.3349
                                         3.7993
##
  Coefficients:
##
                            Estimate Std. Error z value Pr(>|z|)
  (Intercept)
                            3.352407
                                        0.070711
                                                  47.410
  InjectionDCZ
                           -0.006018
                                        0.092310
                                                  -0.065
                                                           0.9480
  portesblue
                            0.731887
                                        0.081753
                                                   8.952
                                                           <2e-16 ***
  InjectionDCZ:portesblue
                            0.212183
                                        0.107004
                                                   1.983
                                                           0.0474 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
   (Dispersion parameter for poisson family taken to be 1)
##
##
##
       Null deviance: 388.396
                               on 37
                                      degrees of freedom
## Residual deviance: 83.286
                               on 34 degrees of freedom
  AIC: 305.58
##
## Number of Fisher Scoring iterations: 4
```

Here we have an interaction for Homer and Donut (although in different directions): - Homer , an earlier last correct under DCZ trial than in sham session and clear session too maybe. - Donut , slightly later last correct in blue DCZ compared to sham.

This might mean that trials are instead repeats (since we have the same number of trials in sessions). So let see let's do the same analysis for Repeats, and then analyze the Repeats altogether

In the first analysis we take only the "blue" sessions because we have very few repeats in "clear":

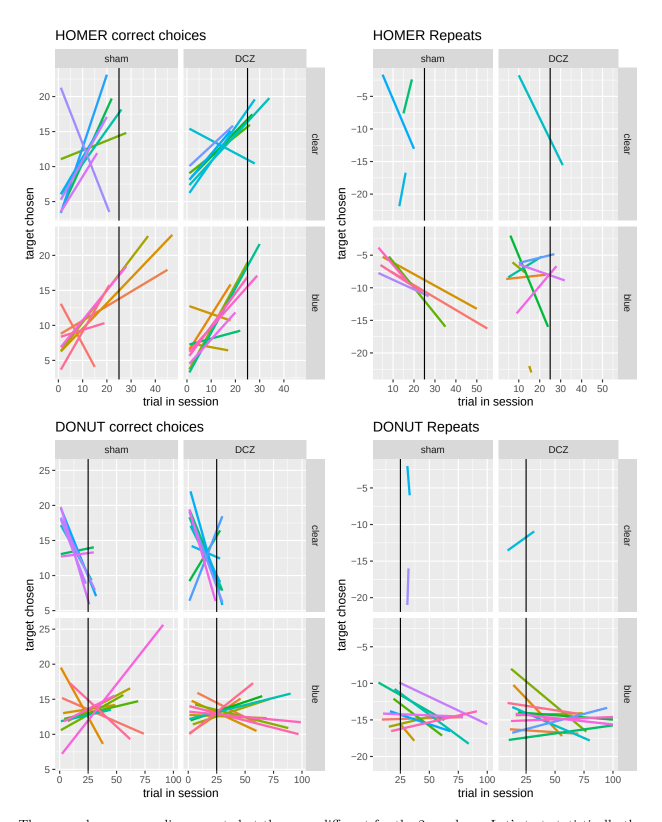
Last Repeat trial in sessions - c



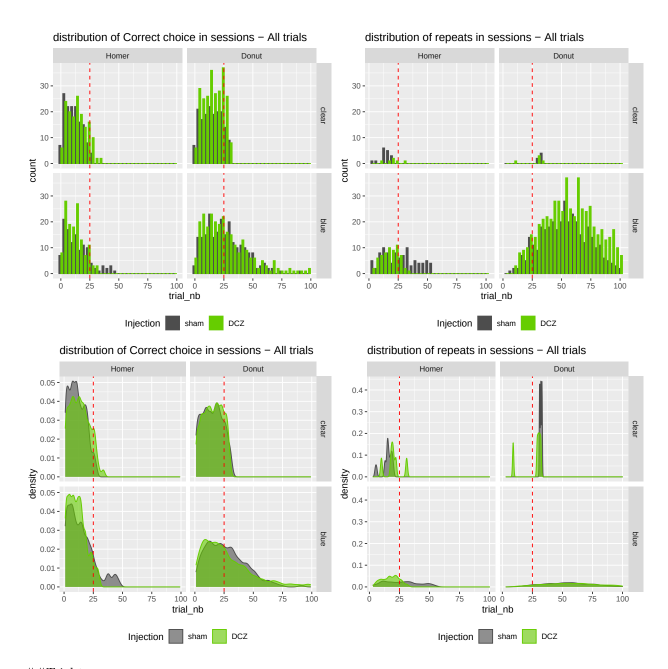
```
##
## Call:
  glm(formula = trial_nb ~ Injection, family = "poisson", data = subset(agg.maxrpt,
##
       singe == "Homer"))
##
##
## Deviance Residuals:
      Min
                 10
                     Median
                                   30
                                           Max
## -6.0313 -1.3653
                                        4.0254
                     0.0000
                               0.9218
##
## Coefficients:
                Estimate Std. Error z value Pr(>|z|)
                            0.06868 49.660 < 2e-16 ***
## (Intercept)
                3.41068
## InjectionDCZ -0.27518
                            0.10076 -2.731 0.00631 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
##
  (Dispersion parameter for poisson family taken to be 1)
##
##
      Null deviance: 92.767 on 14 degrees of freedom
## Residual deviance: 85.283 on 13 degrees of freedom
## AIC: 163.95
##
## Number of Fisher Scoring iterations: 5
##
## Call:
## glm(formula = trial_nb ~ Injection, family = "poisson", data = subset(agg.maxrpt,
##
       singe == "Donut"))
##
##
  Deviance Residuals:
##
      Min
                 1Q
                     Median
                                   3Q
                                           Max
##
   -4.5108 -1.2857 -0.4203
                               1.7148
                                        3.1647
##
## Coefficients:
##
                Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                4.27110
                            0.03737 114.287 < 2e-16 ***
## InjectionDCZ 0.15755
                            0.04981
                                      3.163 0.00156 **
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for poisson family taken to be 1)
##
##
       Null deviance: 86.851 on 20 degrees of freedom
## Residual deviance: 76.790 on 19 degrees of freedom
## AIC: 210.32
##
## Number of Fisher Scoring iterations: 4
```

Main effect for both monkeys although, again, in opposite direction; in other words the rank of the last repeat trial differ in DCZ and sham: earlier for Homer, and later for Donut.

Below we can graph the overall trends of choices across sessions. The lines represent the fit to choice patterns (as shown in the very first figures). Positive slopes means searching holes from top to bottom, and negative slopes from bottom to top. The length actually covers the number of trials.

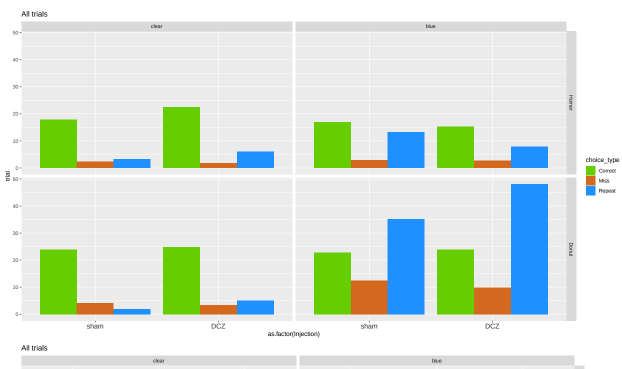


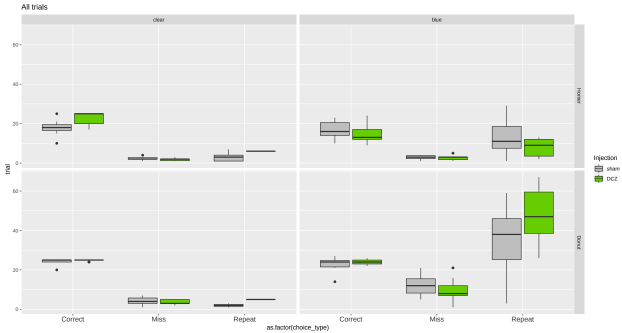
There are changes regarding repeats but these are different for the 2 monkeys. Let's test statistically the data for All trials, for <25 trials and for >25 trials:



##Trial types:

First let's look at the frequency of repeats, miss, etc. . .



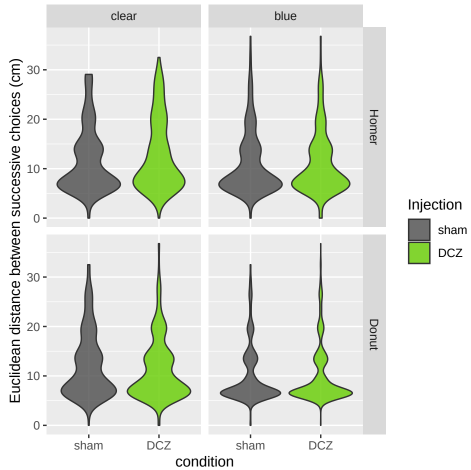


```
##
## Call:
## glm(formula = trial ~ Injection * choice_type, family = "poisson",
       data = subset(agg.data4B, singe == "Homer" & choice_type !=
##
##
           "Correct"))
##
## Deviance Residuals:
##
       Min
                 1Q
                      Median
                                   3Q
                                           Max
## -3.4284 -0.9582 -0.1931
                               0.5330
                                        5.2437
## Coefficients:
```

```
##
                                  Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                                              0.17961
                                                        5.284 1.26e-07 ***
                                   0.94908
## InjectionDCZ
                                  -0.12240
                                              0.25201
                                                       -0.486
                                                                 0.627
## choice_typeRepeat
                                   1.25736
                                              0.20355
                                                        6.177 6.53e-10 ***
## InjectionDCZ:choice_typeRepeat -0.04716
                                              0.29525
                                                       -0.160
                                                                 0.873
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for poisson family taken to be 1)
##
##
       Null deviance: 213.74 on 46 degrees of freedom
## Residual deviance: 127.80 on 43 degrees of freedom
  AIC: 281.9
##
## Number of Fisher Scoring iterations: 5
## Call:
## glm(formula = trial ~ Injection * choice_type, family = "poisson",
       data = subset(agg.data4B, singe == "Donut" & choice_type !=
##
##
           "Correct"))
##
## Deviance Residuals:
##
      Min
                1Q
                     Median
                                   3Q
                                           Max
## -7.5591 -1.7077 -0.3875
                               1.4065
                                        5.1949
##
## Coefficients:
##
                                  Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                                              0.08165
                                                      27.410 < 2e-16 ***
                                   2.23805
## InjectionDCZ
                                  -0.29214
                                              0.11751
                                                       -2.486
                                                                0.0129 *
## choice_typeRepeat
                                   1.07754
                                              0.09726
                                                       11.079 < 2e-16 ***
## InjectionDCZ:choice_typeRepeat 0.77204
                                              0.13592
                                                        5.680 1.35e-08 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for poisson family taken to be 1)
##
       Null deviance: 1096.15 on 60 degrees of freedom
## Residual deviance: 459.45 on 57 degrees of freedom
## AIC: 725.94
##
## Number of Fisher Scoring iterations: 5
```

The statistics show that there is no effect of the condition (Injection) for Homer but there is a significant increase of repeat for Donut. This is when we do not take PORTES into account. If Portes is used as an interacting fixed effect the effect size for Repeats in Donut goes down (p=0.052).

###stats 25 trials



```
\#\#\#stats spatial strategy
##
   Two-sample Kolmogorov-Smirnov test
##
##
## data: data4B$distloc[data4B$singe == "Homer" & data4B$portes == "clear" & data4B$Injection == "DCZ"
## D = 0.083018, p-value = 0.5883
  alternative hypothesis: two-sided
##
##
   Two-sample Kolmogorov-Smirnov test
##
## data: data4B$distloc[data4B$singe == "Homer" & data4B$portes == "blue" & data4B$Injection == "DCZ"]
## D = 0.038739, p-value = 0.996
## alternative hypothesis: two-sided
##
##
   Two-sample Kolmogorov-Smirnov test
##
## data: data4B$distloc[data4B$singe == "Donut" & data4B$portes == "clear" & data4B$Injection == "DCZ".
## D = 0.075606, p-value = 0.5193
## alternative hypothesis: two-sided
##
##
   Two-sample Kolmogorov-Smirnov test
```

data: data4B\$distloc[data4B\$singe == "Donut" & data4B\$portes == "blue" & data4B\$Injection == "DCZ"]

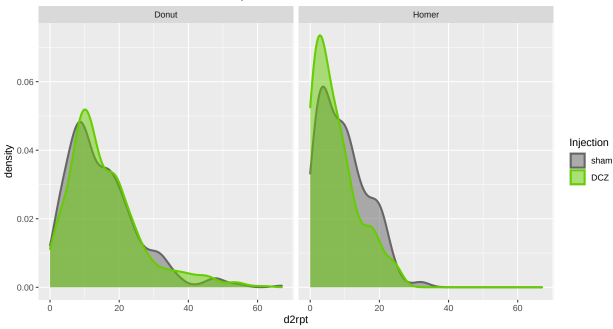
##

```
## D = 0.012805, p-value = 1
## alternative hypothesis: two-sided
```

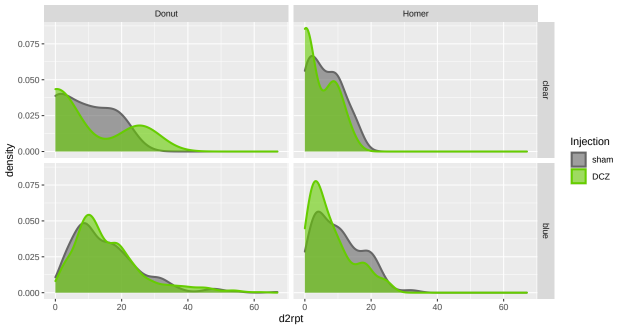
There is no difference in distribution of distances between sham and DCZ.

###distance to repeat

distance between a choice and its repetition



distance between a choice and its repetition



```
##
## Call:
## glm(formula = d2rpt ~ Injection * portes, family = "poisson",
## data = subset(stats.repeat, singe == "Homer"))
```

```
##
## Deviance Residuals:
      Min
                1Q
                     Median
                                  30
## -3.9513 -2.2065 -0.4642
                             1.4153
                                       5.4521
## Coefficients:
                          Estimate Std. Error z value Pr(>|z|)
                                      0.09535 18.985 < 2e-16 ***
## (Intercept)
                           1.81011
## InjectionDCZ
                          -0.42381
                                      0.17297
                                               -2.450
                                                        0.0143 *
## portesblue
                           0.50883
                                      0.10086
                                                5.045 4.53e-07 ***
## InjectionDCZ:portesblue 0.15983
                                      0.18184
                                                0.879
                                                        0.3794
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for poisson family taken to be 1)
##
##
       Null deviance: 1005.94 on 182 degrees of freedom
## Residual deviance: 925.47 on 179
                                      degrees of freedom
## AIC: 1582.3
##
## Number of Fisher Scoring iterations: 5
##
## Call:
## glm(formula = d2rpt ~ Injection * portes, family = "poisson",
      data = subset(stats.repeat, singe == "Donut"))
##
## Deviance Residuals:
                    Median
##
      Min
                1Q
                                  3Q
                                          Max
## -4.9180 -2.0383 -0.7408
                             1.1667
                                       9.7468
##
## Coefficients:
##
                          Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                                      0.10976 19.280 < 2e-16 ***
                           2.11626
## InjectionDCZ
                          -0.06396
                                      0.14568 -0.439
                                                         0.661
                           0.60673
## portesblue
                                      0.11062
                                                5.485 4.13e-08 ***
## InjectionDCZ:portesblue 0.10458
                                      0.14673
                                               0.713
                                                         0.476
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for poisson family taken to be 1)
##
##
       Null deviance: 6066.4 on 897 degrees of freedom
## Residual deviance: 5954.8 on 894 degrees of freedom
## AIC: 9830.7
## Number of Fisher Scoring iterations: 6
##
##
     Simultaneous Tests for General Linear Hypotheses
##
## Multiple Comparisons of Means: Tukey Contrasts
##
##
## Fit: glm(formula = d2rpt ~ -1 + BV, family = "poisson", data = subset(d,
```

```
##
       singe == "Homer"))
##
## Linear Hypotheses:
##
                               Estimate Std. Error z value Pr(>|z|)
## DCZ.clear - sham.clear == 0
                                -0.4238
                                            0.1730
                                                    -2.450
                                                              0.0598 .
## sham.blue - sham.clear == 0
                                 0.5088
                                            0.1009
                                                     5.045
                                                              <0.001 ***
## DCZ.blue - sham.clear == 0
                                                     2.318
                                 0.2448
                                            0.1056
                                                              0.0831 .
## sham.blue - DCZ.clear == 0
                                 0.9326
                                            0.1480
                                                     6.301
                                                              <0.001 ***
## DCZ.blue - DCZ.clear == 0
                                 0.6687
                                            0.1513
                                                     4.419
                                                              <0.001 ***
## DCZ.blue - sham.blue == 0
                                -0.2640
                                            0.0561 - 4.706
                                                              <0.001 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
  (Adjusted p values reported -- single-step method)
##
##
     Simultaneous Tests for General Linear Hypotheses
##
## Multiple Comparisons of Means: Tukey Contrasts
##
##
## Fit: glm(formula = d2rpt ~ -1 + BV, family = "poisson", data = subset(d,
       singe == "Donut"))
##
##
## Linear Hypotheses:
                               Estimate Std. Error z value Pr(>|z|)
##
## DCZ.clear - sham.clear == 0 -0.06396
                                           0.14568
                                                    -0.439
## sham.blue - sham.clear == 0 0.60673
                                           0.11062
                                                     5.485
                                                              <1e-04 ***
## DCZ.blue - sham.clear == 0
                                0.64735
                                           0.11031
                                                     5.868
                                                              <1e-04 ***
## sham.blue - DCZ.clear == 0
                                                              <1e-04 ***
                                0.67070
                                           0.09676
                                                     6.932
## DCZ.blue - DCZ.clear == 0
                                0.71131
                                           0.09641
                                                     7.378
                                                              <1e-04 ***
## DCZ.blue - sham.blue == 0
                                0.04062
                                           0.01755
                                                     2.314
                                                              0.0756 .
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Adjusted p values reported -- single-step method)
```

Regarding distances between a choice and a repeat, DCZ effects are absent in Donut but present in Homer. For Homer, the distances are longer under DCZ in blue conditions but much shorter in DCZ than sham in clear conditions.

##2D choices

