

# Portfolio Assignment 1, Methods 3, 2021, autumn semester

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29/09/2021

## Assignment 1: Using mixed effects modelling to model hierarchical data

In this assignment we will be investigating the *politeness* dataset of Winter and Grawunder (2012) and apply basic methods of multilevel modelling.

```
#this is the dataset we will be exploring in this exercise  
politeness <- read.csv('politeness.csv') ## read in data
```

### Exercise 1 - describing the dataset and making some initial plots

1. Describe the dataset, such that someone who happened upon this dataset could understand the variables and what they contain

#### Exercise 1, part 1– (EH)

The politeness dataset contains the data obtained from the study of Korean formal and informal speech (<https://doi.org/10.1016/j.wocn.2012.08.006>) which investigated the fundamental frequency of male and female participants' speech in a variety of formal and informal scenarios.

The following table describes the variables in the dataset:

Variable	Description
subject	participant ID
gender	participant's gender
scenario	the experimental scenario from 1 to 7 such as "asking a favour"
attitude	either 'inf' for informal stimuli or 'pol' for formal stimuli
total_duration	duration of participant's response in seconds
f0mn	mean fundamental frequency (f0) of the participant's speech
hiss_count	number of times the participants made a noisy breath intake

Remark: The gender , scenario and attitude variables should be encoded as factors as they show a categorical function withing this dataset . In addition, these variables have non-unique values across participants, and are not ordered.

```
#Encoding some of the variables as factors (gender, attitude, and scenario)
```

```
politeness$attitude <- as.factor(politeness$attitude)
politeness$gender <- as.factor(politeness$gender)
politeness$scenario <- as.factor(politeness$scenario)
```

## Exercise 1, part 2–(EH)

2. Create a new data frame that just contains the subject *F1* and run two linear models; one that expresses *f0mn* as dependent on *scenario* as an integer; and one that expresses *f0mn* as dependent on *scenario* encoded as a factor
  - i. Include the model matrices, *X* from the General Linear Model, for these two models in your report and describe the different interpretations of *scenario* that these entail
  - ii. Which coding of *scenario*, as a factor or not, is more fitting?

```
# Create a subset dataframe for subject F1 only
pf1 <- politeness[politeness$subject == "F1", ]
pf1
```

```
##   subject gender scenario attitude total_duration f0mn hiss_count
## 1      F1      F       1     pol    18.392  214.6     2
## 2      F1      F       1     inf    13.551  210.9     0
## 3      F1      F       2     pol     5.217  284.7     0
## 4      F1      F       2     inf     4.247  265.6     0
## 5      F1      F       3     pol     6.791  210.6     0
## 6      F1      F       3     inf     4.126  285.6     0
## 7      F1      F       4     pol     6.244  251.5     1
## 8      F1      F       4     inf     3.245  281.5     0
## 9      F1      F       5     pol     5.625  229.6     1
## 10     F1      F       5     inf     3.950  250.5     0
## 11     F1      F       6     pol    28.508  181.1     1
## 12     F1      F       6     inf    55.159  229.3     0
## 13     F1      F       7     inf    60.309  219.8     2
## 14     F1      F       7     pol    40.825  175.8     0
```

```
# make model predicting f0mn by scenario (integer)
m1<- lm(f0mn ~ as.integer(scenario), data = pf1)
```

```
# get model matrix
mm1 <- model.matrix(m1)
```

```
# make model predicting f0mn by scenario (factor)
m2 <- lm(f0mn ~ as.factor(scenario), data = pf1)
```

```
# get model matrix
mm2 <- model.matrix(m2)
```

Here is the model using "scenario" encoded as an integer

```
summary(m1)
```

```

## 
## Call:
## lm(formula = f0mn ~ as.integer(scenario), data = pf1)
##
## Residuals:
##    Min     1Q Median     3Q    Max
## -44.836 -36.807   6.686  20.918  46.421
##
## Coefficients:
##                               Estimate Std. Error t value Pr(>|t| )
## (Intercept)             262.621    20.616 12.738 2.48e-08 ***
## as.integer(scenario)   -6.886     4.610  -1.494    0.161
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 34.5 on 12 degrees of freedom
## Multiple R-squared:  0.1568, Adjusted R-squared:  0.0865
## F-statistic: 2.231 on 1 and 12 DF,  p-value: 0.1611

```

mm1

```

## (Intercept) as.integer(scenario)
## 1           1                 1
## 2           1                 1
## 3           1                 2
## 4           1                 2
## 5           1                 3
## 6           1                 3
## 7           1                 4
## 8           1                 4
## 9           1                 5
## 10          1                 5
## 11          1                 6
## 12          1                 6
## 13          1                 7
## 14          1                 7
## attr(,"assign")
## [1] 0 1

```

And here is the model using “scenario” encoded as a factor

```
summary(m2)
```

```
##  
## Call:  
## lm(formula = f0mn ~ as.factor(scenario), data = pf1)  
##  
## Residuals:  
##      Min     1Q Median     3Q    Max  
## -37.50 -13.86   0.00  13.86  37.50  
##  
## Coefficients:  
##                               Estimate Std. Error t value Pr(>|t| )  
## (Intercept)                212.75    20.35 10.453 1.6e-05 ***  
## as.factor(scenario)2       62.40    28.78  2.168  0.0668 .  
## as.factor(scenario)3       35.35    28.78  1.228  0.2591  
## as.factor(scenario)4       53.75    28.78  1.867  0.1041  
## as.factor(scenario)5       27.30    28.78  0.948  0.3745  
## as.factor(scenario)6      -7.55    28.78 -0.262  0.8006  
## as.factor(scenario)7      -14.95   28.78 -0.519  0.6195  
## ---  
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1  
##  
## Residual standard error: 28.78 on 7 degrees of freedom  
## Multiple R-squared:  0.6576, Adjusted R-squared:  0.364  
## F-statistic:  2.24 on 6 and 7 DF,  p-value: 0.1576
```

mm2

```

##      (Intercept) as.factor(scenario)2 as.factor(scenario)3 as.factor(scenario)4
## 1             1                 0                 0                 0
## 2             1                 0                 0                 0
## 3             1                 1                 0                 0
## 4             1                 1                 0                 0
## 5             1                 0                 1                 0
## 6             1                 0                 1                 0
## 7             1                 0                 0                 1
## 8             1                 0                 0                 1
## 9             1                 0                 0                 0
## 10            1                 0                 0                 0
## 11            1                 0                 0                 0
## 12            1                 0                 0                 0
## 13            1                 0                 0                 0
## 14            1                 0                 0                 0
##      as.factor(scenario)5 as.factor(scenario)6 as.factor(scenario)7
## 1             0                 0                 0
## 2             0                 0                 0
## 3             0                 0                 0
## 4             0                 0                 0
## 5             0                 0                 0
## 6             0                 0                 0
## 7             0                 0                 0
## 8             0                 0                 0
## 9             1                 0                 0
## 10            1                 0                 0
## 11            0                 1                 0
## 12            0                 1                 0
## 13            0                 0                 1
## 14            0                 0                 1
## attr(,"assign")
## [1] 0 1 1 1 1 1
## attr(,"contrasts")
## attr(,"contrasts")$`as.factor(scenario)`
## [1] "contr.treatment"

```

Conclusion: The above output shows the difference in model matrices between scenario encoded as an integer and factor. The integer version treats scenario as a continuous variable, whereas the factorized version creates a regression line per scenario.

For this dataset, scenario should be a factor, since the scenarios are not a continuous variable and depending on the prescribed scenario, the participants may have a different f0 (mean fundamental frequency of speech), and we are interested in following the trajectory of f0 across scenarios not as a variable that consistently decreases or increases, but a separate regression line showing the changes in f0 between the 7 different scenarios. And in order, to be able to see that crucial difference we need to consider the 'scenario' variable as a factor when we run a model predicting f0 across scenarios.

### Exercise 1, part 3 – (EH)

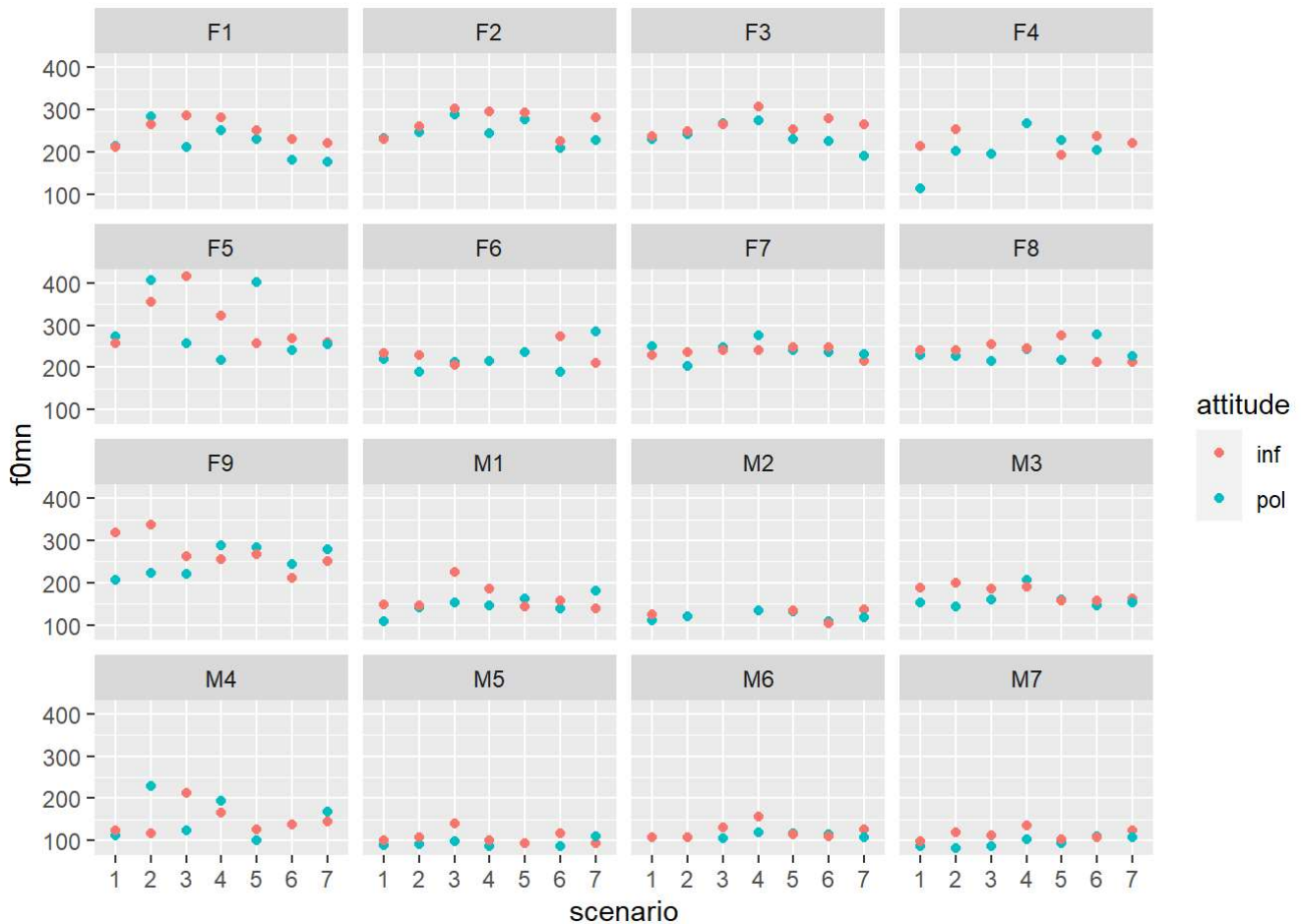
3. Make a plot that includes a subplot for each subject that has *scenario* on the x-axis and *f0mn* on the y-axis and where points are colour coded according to *attitude*
  - i. Describe the differences between subjects

```

politeness %>% ggplot(aes(scenario, f0mn, color = attitude)) +
  geom_point() +
  facet_wrap(vars(subject))

```

```
## Warning: Removed 12 rows containing missing values (geom_point).
```



...

We can visually observe that there are baseline differences between the male and female subjects' mean fundamental frequency of speech, where the males' f0 is consistently lower, across scenario and attitude. Between the different scenarios there is variability in the f0 values for both male and female subjects depending on both the scenario type and the attitude (informal or formal). There is a consistent tendency across scenario type and gender for the mean fundamental frequency of speech to be slightly higher when the attitude is informal as opposed to formal. This visual information interpreted from this plot is consistent with the results of Winter and Grawunder (2012)

##Exercise 2 - comparison of models

```
mixed.model <- lmer(formula=..., data=...)
example.formula <- formula(dep.variable ~ first.level.variable + (1 | second.level.variable))
```

###Part 1 – (VK) 1) Build four different models and do some comparisons

```
# the single level model  
m3 <- lm(formula = f0mn ~ gender, data = politeness)  
  
# a two-level model where each scenario has a unique intercept  
m4 <- lmer(formula = f0mn ~ gender + (1 | scenario), data = politeness)  
# a two-level model that has models subject as intercept  
m5 <- lmer(formula = f0mn ~ gender + (1 | subject), data = politeness)  
  
# a two-level model that incorporate intercepts for both subject and scenario  
m6 <- lmer(formula = f0mn ~ gender +  
           (1 | subject) + (1 | scenario), data = politeness)  
  
#comparing AIC and Deviance values for all the models  
AIC(m3)
```

```
## [1] 2163.971
```

```
AIC(m4)
```

```
## [1] 2152.314
```

```
AIC(m5)
```

```
## [1] 2099.626
```

```
AIC(m6)
```

```
## [1] 2092.482
```

```
deviance(m3)
```

```
## [1] 327033.6
```

```
deviance(m4)
```

```
## Warning in deviance.merMod(m4): deviance() is deprecated for REML fits;  
## use REMLcrit for the REML criterion or deviance(.,REML=FALSE) for deviance  
## calculated at the REML fit
```

```
## [1] 2144.314
```

```
deviance(m5)
```

```
## Warning in deviance.merMod(m5): deviance() is deprecated for REML fits;  
## use REMLcrit for the REML criterion or deviance(.,REML=FALSE) for deviance  
## calculated at the REML fit
```

```
## [1] 2091.626
```

```
deviance(m6)
```

```
## Warning in deviance.merMod(m6): deviance() is deprecated for REML fits;
## use REMLcrit for the REML criterion or deviance(.,REML=FALSE) for deviance
## calculated at the REML fit
```

```
## [1] 2082.482
```

```
anova(m4, m5, m6)
```

```
## refitting model(s) with ML (instead of REML)
```

```
## Data: politeness
## Models:
## m4: f0mn ~ gender + (1 | scenario)
## m5: f0mn ~ gender + (1 | subject)
## m6: f0mn ~ gender + (1 | subject) + (1 | scenario)
##   npar    AIC    BIC  logLik deviance Chisq Df Pr(>Chisq)
## m4     4 2162.3 2175.7 -1077.1   2154.3
## m5     4 2112.1 2125.5 -1052.0   2104.1 50.2095  0
## m6     5 2105.2 2122.0 -1047.6   2095.2  8.8725  1   0.002895 **
```

## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

```
piecewiseSEM:::rsquared(c(m4, m5, m6))
```

```
##   Response family link method Marginal Conditional
## 1      f0mn gaussian identity   none 0.6779555  0.6967788
## 2      f0mn gaussian identity   none 0.6681651  0.7899229
## 3      f0mn gaussian identity   none 0.6677206  0.8077964
```

The single level model performs the worst and this makes sense as we do not expect all participants to have the same f0 as their voices have naturally occurring differences (not just ones predicted by gender). There are differences that might be explained by the scenario or individual subject (as we observed in the plot above), however, neither of those are taken into account when using a single level model.

So then using two-level models that explain variance by either taking scenario or subject as random intercepts, is definitely an improvement to help explain more of the scenario/attitude based differences, not just the gender differences.

Consequently, Of the three multi-level models,it is model m6, which includes random intercepts for both subject and scenario,that has the most explained variance with for the entire model  $R^2 \approx 0.81$  or 81%.

Additionally, we see that model m6 has the lowest AIC and deviance.

## Exercise 2, part 2 and 3 – (LR)

2. Why is our single-level model bad?

- i. create a new data frame that has three variables, *subject*, *gender* and *f0mn*, where *f0mn* is the average of all responses of each subject, i.e. averaging across *attitude* and *\_scenario*
- ii. build a single-level model that models *f0mn* as dependent on *gender* using this new dataset

- iii. make Quantile-Quantile plots, comparing theoretical quantiles to the sample quantiles) using `qqnorm` and `qqline` for the new single-level model and compare it to the old single-level model (from 1).i). Which model's residuals ( $\epsilon$ ) fulfil the assumptions of the General Linear Model better?)
- iv. Also make a quantile-quantile plot for the residuals of the multilevel model with two intercepts. Does it look alright?

### 3. Plotting the two-intercepts model

- i. Create a plot for each subject, (similar to part 3 in Exercise 1), this time also indicating the fitted value for each of the subjects for each for the scenarios (hint use `fixef` to get the “grand effects” for each gender and `ranef` to get the subject- and scenario-specific effects)

```
# scenario x f0mn y, attitude = color
ff <- fixef(m6)
ff
```

```
## (Intercept)      genderM
##   246.7650    -115.1746
```

```
rf <- ranef(m6)
rf <- as.data.frame(rf)
rf
```

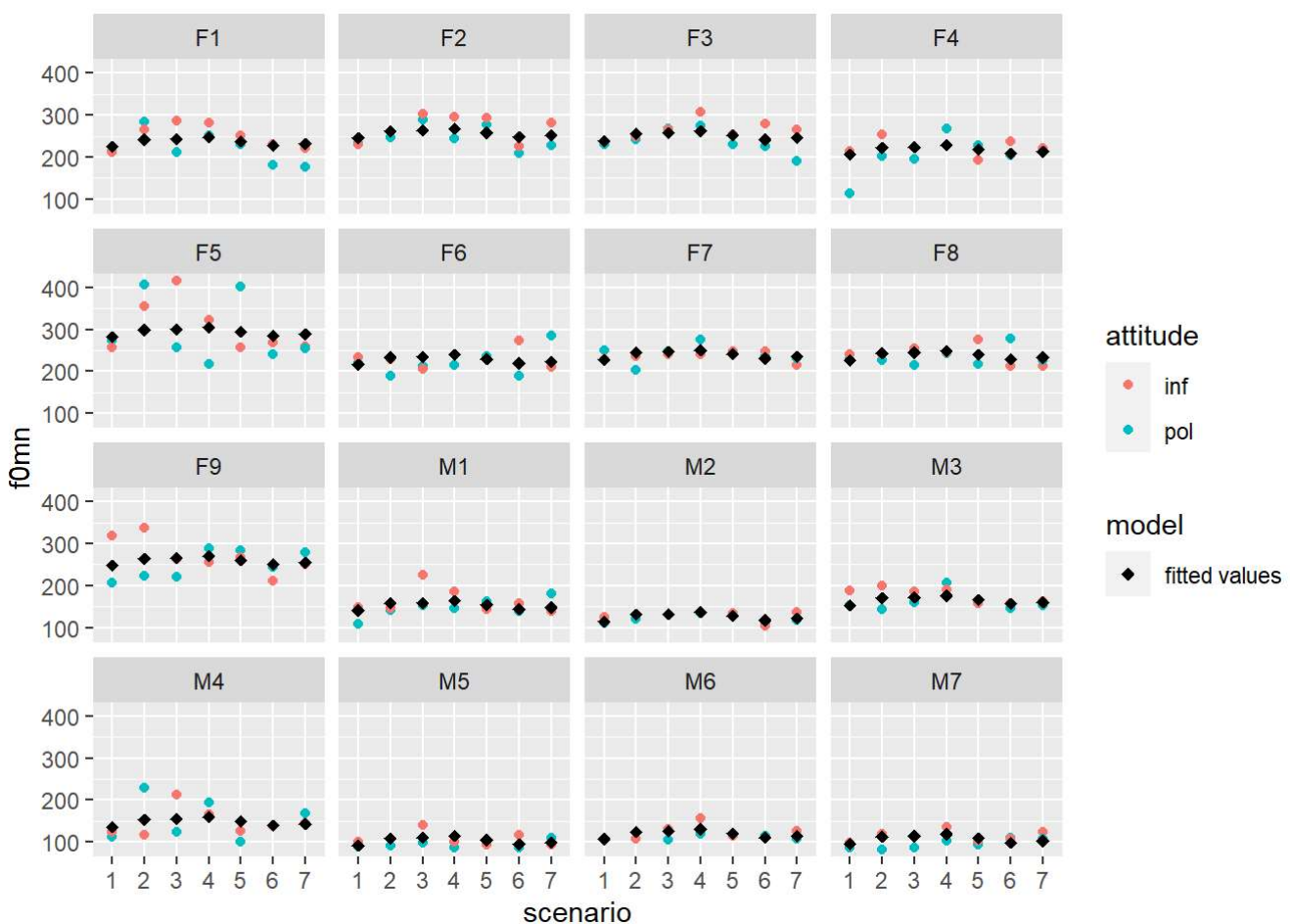
	grpvar	term	grp	condval	condsd
## 1	subject	(Intercept)	F1	-10.490356	8.280794
## 2	subject	(Intercept)	F2	10.251809	8.280794
## 3	subject	(Intercept)	F3	3.795129	8.280794
## 4	subject	(Intercept)	F4	-29.495270	9.095403
## 5	subject	(Intercept)	F5	47.093999	8.280794
## 6	subject	(Intercept)	F6	-18.396273	8.794359
## 7	subject	(Intercept)	F7	-6.976691	8.280794
## 8	subject	(Intercept)	F8	-8.521934	8.280794
## 9	subject	(Intercept)	F9	12.739587	8.280794
## 10	subject	(Intercept)	M1	21.052117	8.280794
## 11	subject	(Intercept)	M2	-5.462358	9.453009
## 12	subject	(Intercept)	M3	33.561535	8.280794
## 13	subject	(Intercept)	M4	16.093337	8.524543
## 14	subject	(Intercept)	M5	-28.267430	8.280794
## 15	subject	(Intercept)	M6	-12.640202	8.794541
## 16	subject	(Intercept)	M7	-24.336998	8.280794
## 17	scenario	(Intercept)	1	-11.595496	5.488728
## 18	scenario	(Intercept)	2	5.321218	5.532205
## 19	scenario	(Intercept)	3	6.795658	5.586194
## 20	scenario	(Intercept)	4	11.348815	5.578013
## 21	scenario	(Intercept)	5	1.411037	5.488705
## 22	scenario	(Intercept)	6	-8.622136	5.489258
## 23	scenario	(Intercept)	7	-4.659096	5.488058

```

politeness$effect_gender <- 0.0
politeness[politeness$gender == "F", ]$effect_gender <- ff[1]
politeness[politeness$gender == "M", ]$effect_gender <- ff[1] + ff[2]
politeness$intercept_subject <- left_join(politeness, rf, by = c("subject" = "grp"), copy
  = TRUE, keep = FALSE)$condval
politeness$intercept_scenario <- left_join(politeness, rf, by = c("scenario" = "grp"), cop
y = TRUE, keep = FALSE)$condval
politeness$predicted <- politeness$effect_gender + politeness$intercept_subject + politene
ss$intercept_scenario
politeness %>% ggplot(aes(scenario, f0mn, color = attitude)) +
  geom_point() +
  geom_point(aes(y = predicted, shape = "fitted values"), color = "black", size = 2) +
  scale_shape_manual(name = "model", values = c(18)) +
  facet_wrap(vars(subject))

```

## Warning: Removed 12 rows containing missing values (geom\_point).



`deviance(m3)`

## [1] 327033.6

`deviance(m4, REML = FALSE)`

## [1] 2154.33

`deviance(m5, REML = FALSE)`

```
## [1] 2104.175
```

```
deviance(m6, REML = FALSE)
```

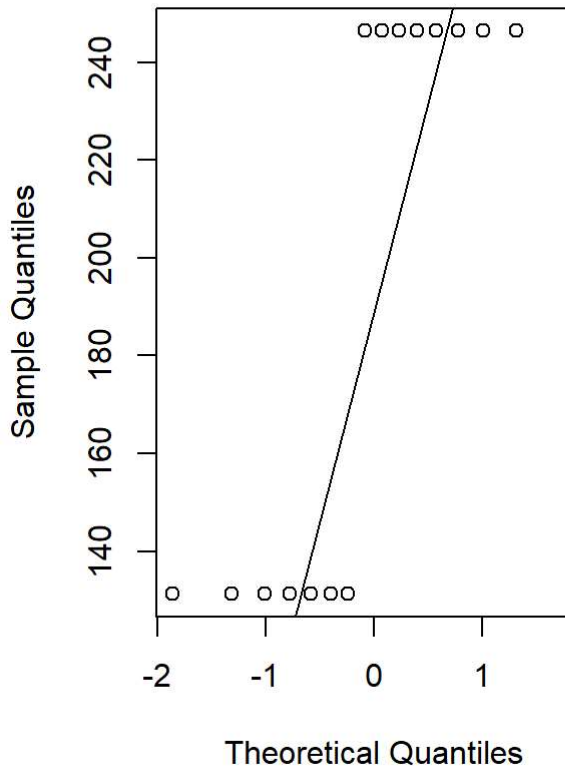
```
## [1] 2095.279
```

```
politeness_aggregated <- politeness[!is.na(politeness$f0mn), ] %>% group_by(subject) %>% s
ummarize(subject = subject[1], gender = gender[1], f0mn = mean(f0mn))
politeness_aggregated
```

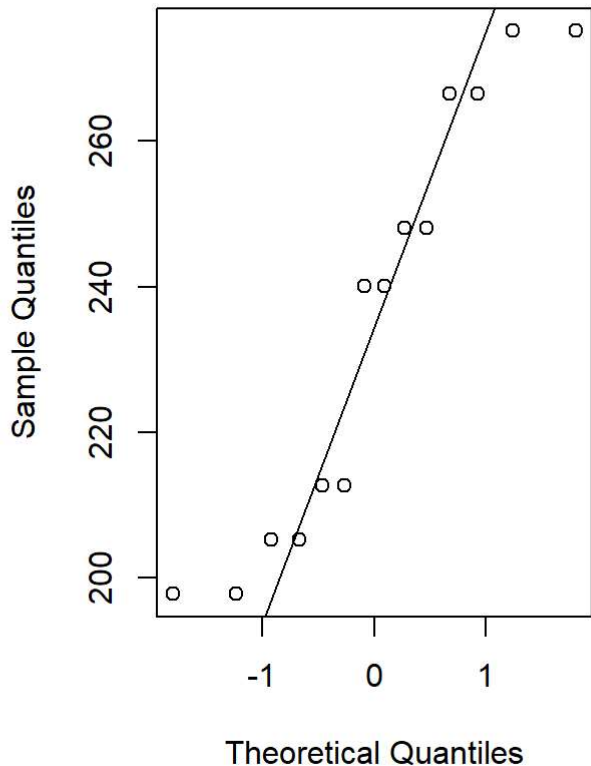
```
## # A tibble: 16 x 3
##   subject gender   f0mn
##   <chr>    <fct>   <dbl>
## 1 F1       F      235.
## 2 F2       F      258.
## 3 F3       F      251.
## 4 F4       F      212.
## 5 F5       F      299.
## 6 F6       F      225.
## 7 F7       F      239.
## 8 F8       F      237.
## 9 F9       F      261.
## 10 M1      M      155.
## 11 M2      M      122.
## 12 M3      M      169.
## 13 M4      M      150.
## 14 M5      M      100.
## 15 M6      M      118.
## 16 M7      M      104.
```

```
m7 <- lm(f0mn ~ gender, data = politeness_aggregated)
par(mfrow=c(1,2))
qqnorm(fitted.values(m7))
qqline(fitted.values(m7))
qqnorm(fitted.values(m2))
qqline(fitted.values(m2))
```

### Normal Q-Q Plot

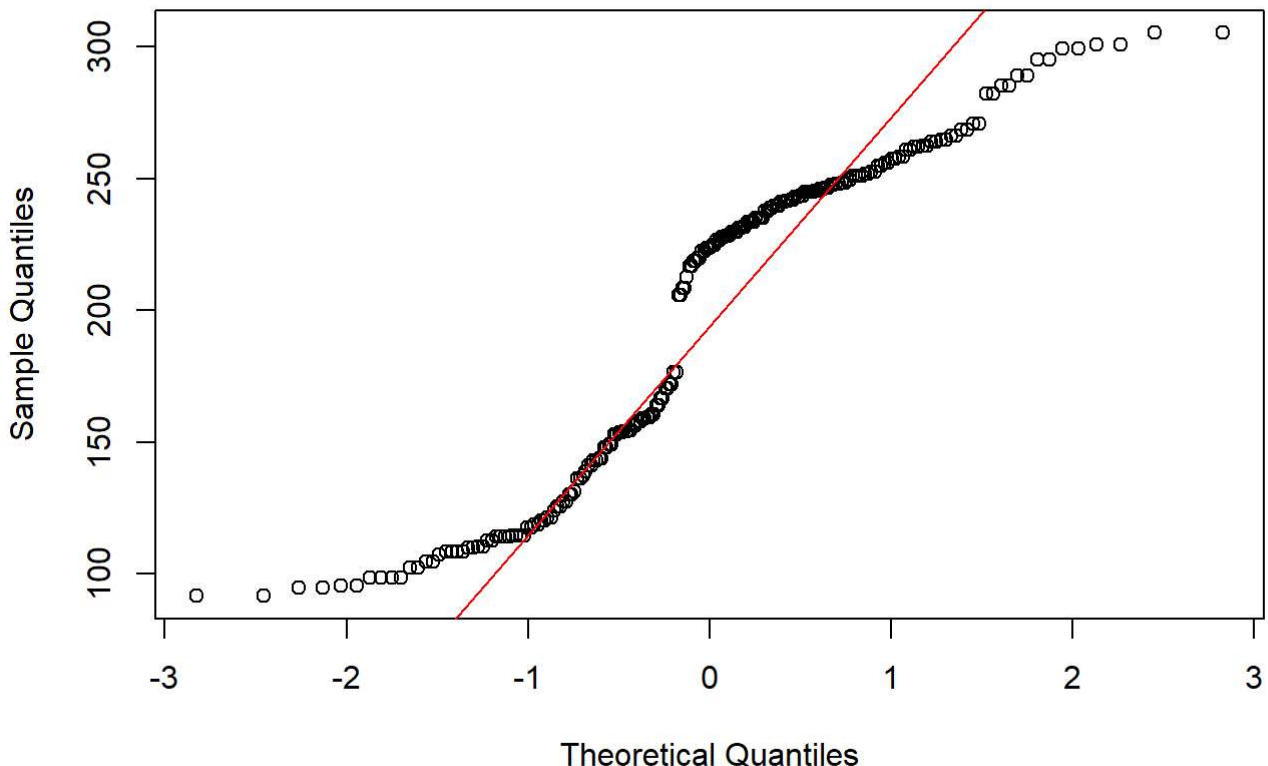


### Normal Q-Q Plot



```
par(mfrow=c(1,1))
qqnorm(fitted.values(m6))
qqline(fitted.values(m6), col = "red")
```

## Normal Q-Q Plot



Assessing the QQ-plots of the single-level models it seems that the aggregated model m7's residuals are worse off than those of model m2. The residuals of model m2 are better dispersed along the line - however it still doesn't look fantastic. The QQ-plot of the multilevel model m6 looks better than any of the single level ones, with data points closer to the line and more evenly dispersed on both sides of the line.

Looking at the plot for the observed and the fitted values it looks as if the model m6 performs reasonably as well.

## Exercise 3 - now with attitude

**Exercise 3, part 1–(VK) 1)** Carry on with the model with the two unique intercepts fitted (*scenario* and *subject*) but now build a new model that has *attitude* as a main effect besides *gender*. After create a separate model that besides the main effects of *attitude* and *gender* also include their interaction

```
#making a model with two unigue intercepts and having attitude and gender as main effects
m8 <- lmer(formula = f0mn ~ gender + attitude + (1 | subject) + (1 | scenario), data = politeness)
summary(m8)
```

```
## Linear mixed model fit by REML ['lmerMod']
## Formula: f0mn ~ gender + attitude + (1 | subject) + (1 | scenario)
##   Data: politeness
##
## REML criterion at convergence: 2065.1
##
## Scaled residuals:
##     Min      1Q  Median      3Q     Max 
## -2.8511 -0.6081 -0.0602  0.4329  3.8745 
##
## Random effects:
##   Groups   Name        Variance Std.Dev. 
##   subject (Intercept) 585.6    24.20  
##   scenario (Intercept) 106.7    10.33  
##   Residual           882.7    29.71  
## Number of obs: 212, groups: subject, 16; scenario, 7
##
## Fixed effects:
##             Estimate Std. Error t value
## (Intercept) 254.398    9.597  26.507
## genderM     -115.437   12.881  -8.962
## attitudepol -14.819    4.096  -3.618
##
## Correlation of Fixed Effects:
##          (Intr) gendrM
## genderM -0.587
## attitudepol -0.220  0.006
```

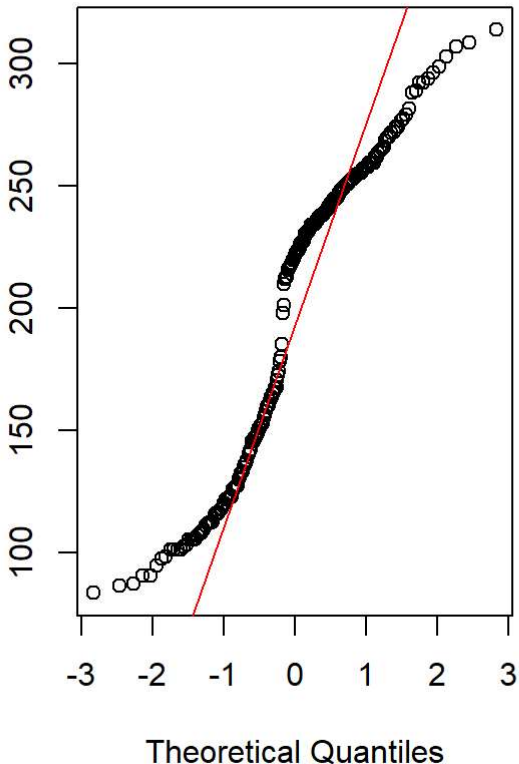
```
fitted.values(m8)
```

	1	2	3	4	5	6	7	8
##	216.47929	231.29853	234.27048	249.08973	236.09087	250.91011	241.21606	256.03530
##	9	10	11	12	13	14	15	16
##	230.41934	245.23858	219.58594	234.40518	238.54395	223.72471	237.34031	252.15956
##	17	18	19	20	21	22	23	24
##	255.13151	269.95075	256.95189	271.77114	262.07708	276.89632	251.28036	266.09960
##	25	26	27	28	29	30	31	32
##	240.44696	255.26621	259.40498	244.58573	230.84663	245.66588	248.63783	263.45707
##	33	34	35	36	37	38	39	40
##	250.45822	265.27746	255.58340	270.40264	244.78668	259.60593	233.95328	248.77253
##	41	42	43	44	45	46	47	49
##	252.91130	238.09206	198.00782	212.82706	215.79901	230.61826	217.61940	222.74458
##	51	52	53	54	55	57	58	59
##	211.94787	226.76711	201.11447	215.93371	220.07248	274.39362	289.21286	292.18481
##	60	61	62	63	64	65	66	67
##	307.00406	294.00520	308.82444	299.13038	313.94963	288.33367	303.15291	277.50027
##	68	69	70	71	72	73	74	75
##	292.31951	296.45828	281.63904	209.70329	224.52253	227.49448	242.31372	229.31487
##	76	77	79	81	82	83	84	85
##	244.13411	234.44005	223.64333	212.80994	227.62918	231.76795	216.94871	220.01309
##	86	87	88	89	90	91	92	93
##	234.83233	237.80428	252.62353	239.62467	254.44391	244.74985	259.56910	233.95314
##	94	95	96	97	98	99	100	101
##	248.77238	223.11974	237.93898	242.07775	227.25851	218.45899	233.27823	236.25019
##	102	103	104	105	106	107	108	109
##	251.06943	238.07057	252.88982	243.19576	258.01500	232.39904	247.21828	221.56564
##	110	111	112	113	114	115	116	117
##	236.38488	240.52366	225.70441	239.84235	254.66159	257.63354	272.45278	259.45393
##	118	119	120	121	122	123	124	125
##	274.27317	264.57911	279.39835	253.78240	268.60164	242.94900	257.76824	261.90701
##	126	127	128	129	130	131	132	133
##	247.08777	133.00242	147.82166	150.79361	165.61285	152.61400	167.43324	157.73918
##	134	135	136	137	138	139	140	141
##	172.55843	146.94247	161.76171	136.10907	150.92831	155.06708	140.24784	107.79414
##	142	143	147	149	150	151	152	153
##	122.61338	125.58533	132.53090	121.73419	136.55343	110.90079	125.72003	129.85880
##	154	155	156	157	158	159	160	161
##	115.03956	145.58352	160.40276	163.37471	178.19395	165.19510	180.01434	170.32028
##	162	163	164	165	166	167	168	169
##	185.13952	159.52357	174.34281	148.69017	163.50941	167.64818	152.82894	127.46793
##	170	171	172	173	174	175	176	177
##	142.28717	145.25913	160.07837	147.07951	161.89876	152.20470	167.02394	141.40798
##	178	180	181	182	183	184	185	186
##	156.22722	145.39382	149.53260	134.71335	83.40025	98.21950	101.19145	116.01069
##	187	188	189	190	191	192	193	194
##	103.01184	117.83108	108.13702	122.95626	97.34030	112.15955	86.50690	101.32615
##	195	196	198	200	201	202	203	204
##	105.46492	90.64568	112.78128	130.57248	117.57362	132.39287	122.69881	137.51805
##	205	206	207	208	209	210	211	212
##	111.90209	126.72133	101.06869	115.88793	120.02671	105.20746	87.35321	102.17245
##	213	214	215	216	217	218	219	220
##	105.14440	119.96365	106.96479	121.78403	112.08998	126.90922	101.29326	116.11250
##	221	222	223	224				
##	90.45986	105.27910	109.41787	94.59863				

```
par(mfrow=c(1,2))
qqnorm(fitted.values(m8))
qqline(fitted.values(m8), col = "red")
qqnorm(politeness$f0mn)
qqline(politeness$f0mn, col = "red")
```

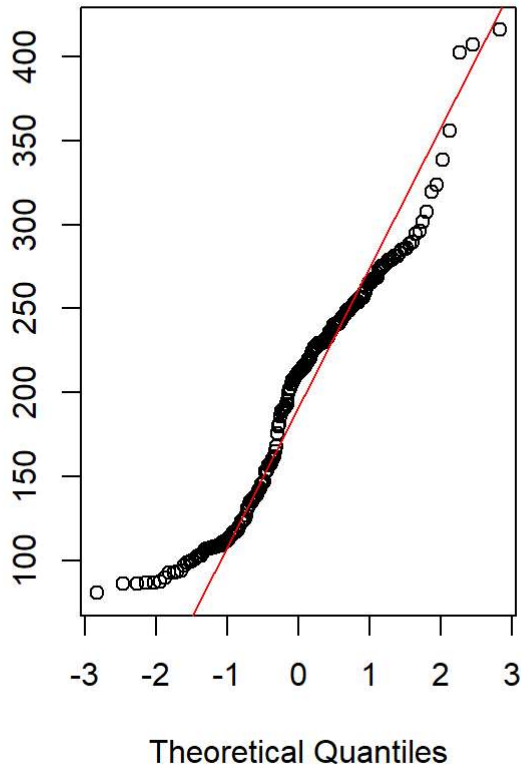
Normal Q-Q Plot

Sample Quantiles

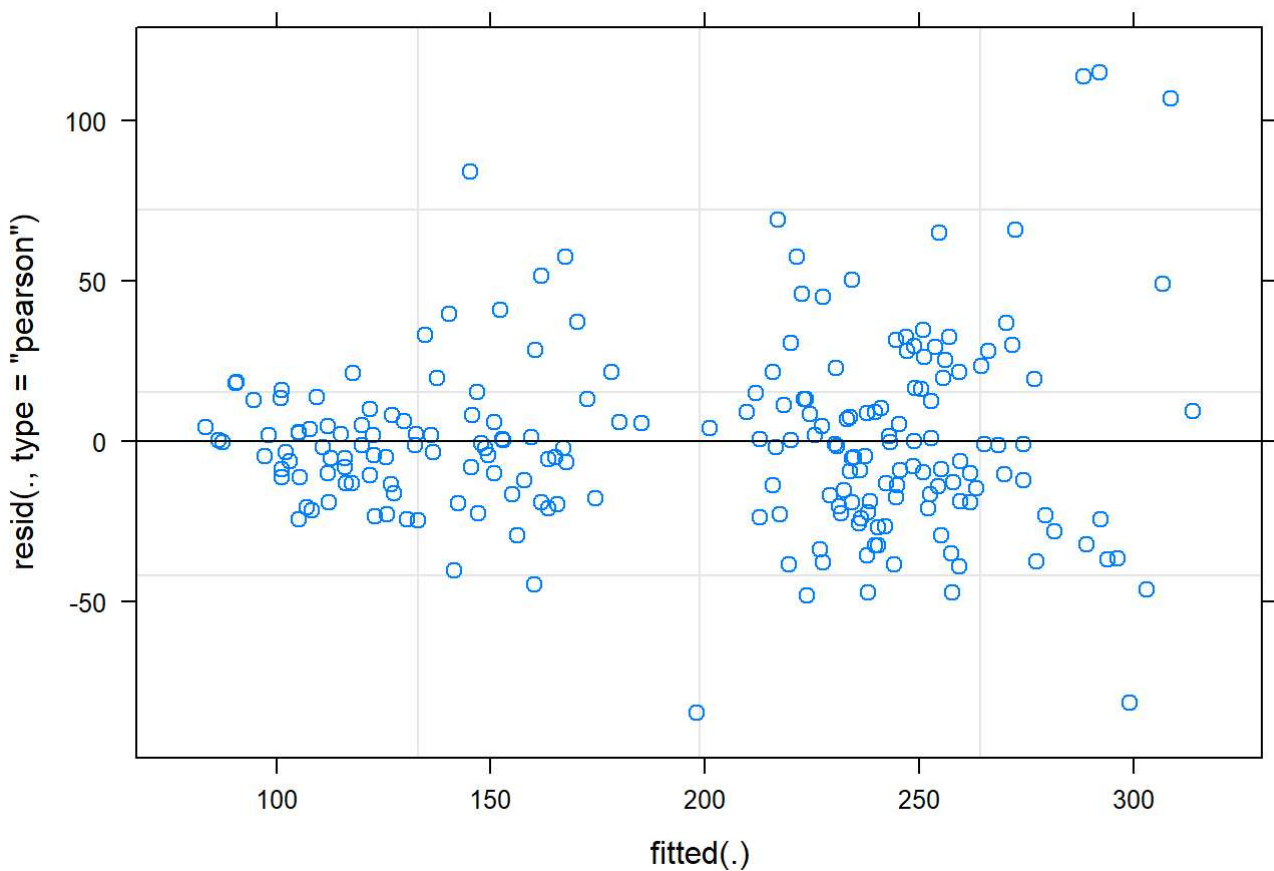


Normal Q-Q Plot

Sample Quantiles



```
plot(m8)
```



```
#a model that additionally includes the interaction between the main effects (attitude and gender)
m9 <- lmer(formula = f0mn ~ gender + attitude + gender:attitude + (1 | subject) + (1 | scenario), data = politeness)
summary(m9)
```

```

## Linear mixed model fit by REML ['lmerMod']
## Formula: f0mn ~ gender + attitude + gender:attitude + (1 | subject) +
##           (1 | scenario)
## Data: politeness
##
## REML criterion at convergence: 2058.6
##
## Scaled residuals:
##     Min      1Q  Median      3Q     Max
## -2.8120 -0.5884 -0.0645  0.4014  3.9100
##
## Random effects:
##   Groups   Name        Variance Std.Dev.
##   subject (Intercept) 584.4    24.17
##   scenario (Intercept) 106.4    10.32
##   Residual            885.5    29.76
## Number of obs: 212, groups: subject, 16; scenario, 7
##
## Fixed effects:
##                   Estimate Std. Error t value
## (Intercept)      255.618    9.761  26.186
## genderM         -118.232   13.531 -8.738
## attitudepol     -17.192    5.423 -3.170
## genderM:attitudepol  5.544    8.284  0.669
##
## Correlation of Fixed Effects:
##          (Intr) gendrM attt dp
## genderM     -0.606
## attitudepol -0.286  0.206
## gndrM:tttdp  0.187 -0.309 -0.654

```

The model m9 can be read as following: The intercept for women/inf are \$ \approx 256 \text{ Hz}\$, when we look at men with the same attitude their pitch drops by \$ \approx 118 \text{ Hz} \$. Overall a polite attitude will result in a drop in pitch by \$ \approx 17 \text{ Hz} \$, however for men it will only be \$ -17.2+5.5 \approx 11.6 \text{ Hz} \$. Korean women's relative drop in pitch is therefore larger than male's in a polite situation according to this sample.

### Exercise 3, part 2–(KV)

2. Compare the three models (1. gender as a main effect; 2. gender and attitude as main effects; 3. gender and attitude as main effects and the interaction between them.

```
#comparing the models
anova(m6, m8, m9)
```

```
## refitting model(s) with ML (instead of REML)
```

```

## Data: politeness
## Models:
## m6: f0mn ~ gender + (1 | subject) + (1 | scenario)
## m8: f0mn ~ gender + attitude + (1 | subject) + (1 | scenario)
## m9: f0mn ~ gender + attitude + gender:attitude + (1 | subject) +
## m9:      (1 | scenario)
##   npar   AIC   BIC logLik deviance Chisq Df Pr(>Chisq)
## m6     5 2105.2 2122.0 -1047.6   2095.2
## m8     6 2094.5 2114.6 -1041.2   2082.5 12.6868  1 0.0003683 ***
## m9     7 2096.0 2119.5 -1041.0   2082.0  0.4551  1 0.4998998
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

```

```
piecewiseSEM:::rsquared(c(m6, m8, m9))
```

```

## Response family link method Marginal Conditional
## 1   f0mn gaussian identity none 0.6677206  0.8077964
## 2   f0mn gaussian identity none 0.6782542  0.8196777
## 3   f0mn gaussian identity none 0.6782490  0.8192531

```

**Exercise 3, part 3–(KV) 3** Choose the model that you think describe the data the best - and write a short report on the main findings based on this model.

```
summary(m8)
```

```

## Linear mixed model fit by REML ['lmerMod']
## Formula: f0mn ~ gender + attitude + (1 | subject) + (1 | scenario)
##   Data: politeness
##
## REML criterion at convergence: 2065.1
##
## Scaled residuals:
##   Min     1Q Median     3Q    Max
## -2.8511 -0.6081 -0.0602  0.4329  3.8745
##
## Random effects:
##   Groups   Name        Variance Std.Dev.
##   subject (Intercept) 585.6    24.20
##   scenario (Intercept) 106.7    10.33
##   Residual           882.7    29.71
## Number of obs: 212, groups: subject, 16; scenario, 7
##
## Fixed effects:
##             Estimate Std. Error t value
## (Intercept) 254.398    9.597  26.507
## genderM     -115.437   12.881 -8.962
## attitudepol -14.819    4.096 -3.618
##
## Correlation of Fixed Effects:
##          (Intr) gendrM
## genderM   -0.587
## attitudepol -0.220  0.006

```

This dataset consists of the basic demographic information of 16 Korean participants, and their observed pitch in different situations that require either an informal or polite(formal) attitude.

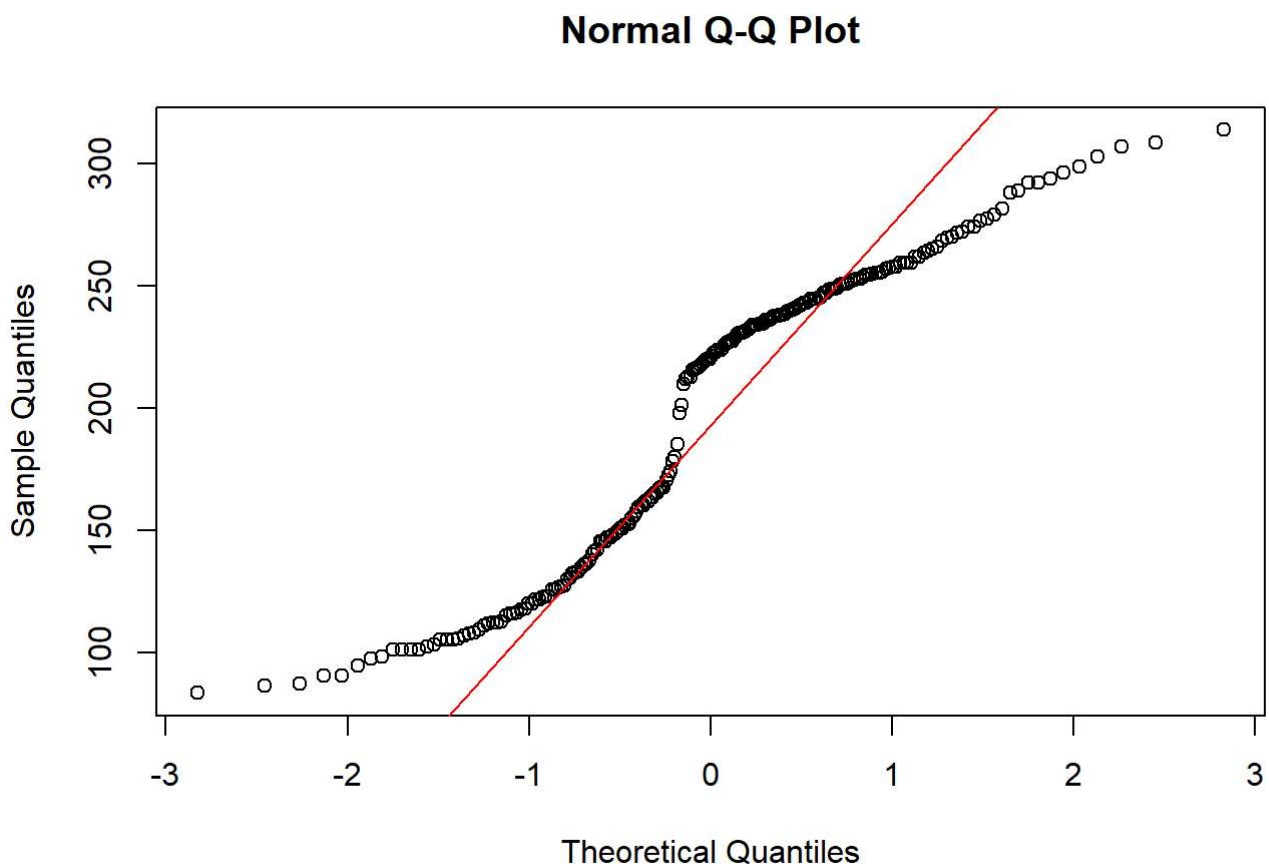
The model we chose as the best is m8. Consistent with the conclusions of the original oauthor's study, this model showed that women on average have a higher pitch than men BUT it also suggested a negative relationship between *attitude* and pitch with p-values<0.001. It would seem that both Korean men's and women's frequency of voice drops when having a polite attitude.

Subjects and scenarios should have different intercepts because it would be assumed they would all have different baselines, (and therefore need different intercepts to account for this). Different subjects will naturally already speak at a different pitch level, so separate intercepts allows us to account for these differences. The different scenarios may also need separate intercepts as certain scenarios may result in participants lowering or raising their pitch to meet the appropriate ambiance of the scenario. Once again, by including separate intercepts for scenarios then we should be accounting for these differences in our models.

Furthermore the output of the summary function shows that more variance is explained by the random effect of the subject than that of the scenario, further strengthening the choice of multilevel modelling.

And here's a QQ-plot of our chosen model

```
qqnorm(fitted.values(m8))
qqline(fitted.values(m8), col = "red")
```



# Portfolio Assignment 2 part 1, Methods 3, 2021, autumn semester

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2021-10-02

## Exercises and objectives

The objectives of the exercises of this assignment are:

- 1) Download and organise the data and model and plot staircase responses based on fits of logistic functions
- 2) Fit multilevel models for response times
- 3) Fit multilevel models for count data

REMEMBER: In your report, make sure to include code that can reproduce the answers requested in the exercises below (**MAKE A KNITTED VERSION**)

REMEMBER: This assignment will be part of your final portfolio

## Exercise 1

Go to <https://osf.io/ecxsj/files/> (<https://osf.io/ecxsj/files/>) and download the files associated with Experiment 2 (there should be 29).

The data is associated with Experiment 2 of the article at the following DOI

<https://doi.org/10.1016/j.concog.2019.03.007>

(<https://doi.org/10.1016/j.concog.2019.03.007>)

1. Put the data from all subjects into a single data frame
2. Describe the data and construct extra variables from the existing variables
  - i. add a variable to the data frame and call it *correct* (have it be a *logical* variable). Assign a 1 to each row where the subject indicated the correct answer and a 0 to each row where the subject indicated the incorrect answer (**Hint:** the variable *obj.resp* indicates whether the subject answered “even”, e or “odd”, o, and the variable *target\_type* indicates what was actually presented).
  - ii. describe what the following variables in the data frame contain, *trial.type*, *pas*, *trial*, *target.contrast*, *cue*, *task*, *target\_type*, *rt.subj*, *rt.obj*, *obj.resp*, *subject* and *correct*. (That means you can ignore the rest of the variables in your description). For each of them, indicate and argue for what *class* they should be classified into, e.g. *factor*, *numeric* etc.
  - iii. for the staircasing part **only**, create a plot for each subject where you plot the estimated function (on the *target.contrast* range from 0-1) based on the fitted

- values of a model (use `glm`) that models *correct* as dependent on *target.contrast*. These plots will be our *no-pooling* model. Comment on the fits  
- do we have enough data to plot the logistic functions?
- iv. on top of those plots, add the estimated functions (on the *target.contrast* range from 0-1) for each subject based on partial pooling model (use `glmer` from the package `lme4`) where unique intercepts and slopes for *target.contrast* are modelled for each *subject*
- v. in your own words, describe how the partial pooling model allows for a better fit for each subject

## Answers

### Exercise 1, part 1 - EH

```
# get list of all CSV files
temp <- list.files(path = "./experiment_2/",
  pattern = "*.csv", full.names = TRUE)
# load into single data frame
samples <- map_df(temp, read_csv, trim_ws = TRUE, na = c("", "NA")),
  col_types = cols(
    trial.type = col_factor(),
    pas = col_factor(),
    trial = col_factor(),
    jitter.x = col_double(),
    jitter.y = col_double(),
    odd.digit = col_integer(),
    target.contrast = col_double(),
    target.frames = col_double(),
    cue = col_factor(),
    task = col_factor(),
    target.type = col_factor(),
    rt.subj = col_double(),
    rt.obj = col_double(),
    even.digit = col_integer(),
    seed = col_double(),
    obj.resp = col_factor(),
    subject = col_factor()
  )
rm(temp)
# peek data
head(samples)
```

```

## # A tibble: 6 x 17
##   trial.type pas    trial jitter.x jitter.y odd.digit target.contrast
##   <fct>     <dbl> <dbl>      <dbl>      <dbl>      <int>       <dbl>
## 1 staircase  4     0     -0.343     0.449      9        1
## 2 staircase  4     1      0.0623    0.0291     9        1
## 3 staircase  4     2     -0.406     0.500      7        0.9
## 4 staircase  4     3     -0.362     -0.222     7        0.9
## 5 staircase  4     4      0.289     0.413      7        0.8
## 6 staircase  4     5      0.0824    -0.0934    7        0.8
## # ... with 10 more variables: target.frames <dbl>, cue <fct>, task <fct>,
## #   target.type <fct>, rt.subj <dbl>, rt.obj <dbl>, even.digit <int>,
## #   seed <dbl>, obj.resp <fct>, subject <fct>

```

```

# total number of samples
nrow(samples)

```

```

## [1] 18131

```

## Exercise 1, part 2 - KV

```

# Add column to indicate if participant response was correct
samples <- mutate(samples,
  correct = as.logical(
    ifelse(substr(target.type, 1, 1) == obj.resp, 1, 0)
  )
)

```

The dataset contains the following variables:

Variable	Description	Class
trial.type	either staircase(practice) or experiment	factor: categorical, reused
pas	any number from 1-4, indicating the reported experience on the Perceptual Awareness Scale	factor: categorical, not continuous
trial	trial number zero-indexed for the practice and experiment blocks	factor: categorical, not continuous
target.contrast	the grey-scale proportion of the target digit	double: numeric, continuous variable from 0-1
cue	number code for pre-stimulus cue	factor: categorical, not continuous

Variable	Description	Class
task	how many numbers could be shown; has the levels quadruplet: (all numbers); pairs: (2 even and 2 odd numbers); singles: (1 even and 1 odd number)	factor: categorical, no continuous relationship
target_type	whether the target shown was the chosen even.digit or the chosen odd.digit	factor: categorical, no continuous relationship
rt.subj	reaction time (seconds) on the PAS response	double: continuous variable with decimal places
rt.obj	reaction time (seconds) on the target digit	double: continuous variable with decimal places
obj.resp	the key actually pressed e for even and o for odd	factor: categorical, no continuous relationship
subject	subject number	factor: categorical
correct	Whether answer was correct(1) or incorret (0)	logical, boolean either true or false

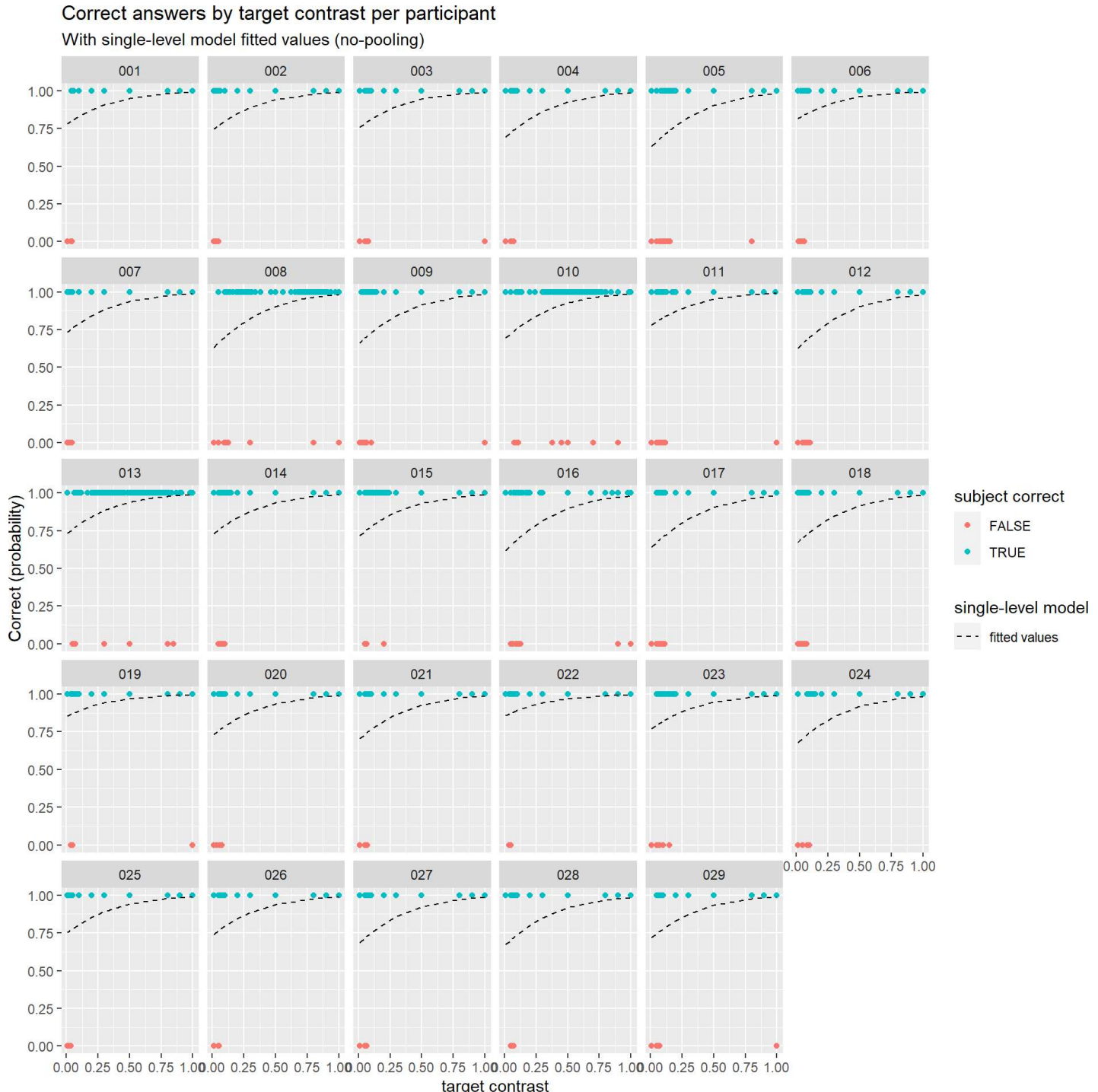
```
# only use staircase trial types.
samples_staircase <- samples %>% filter(trial.type == "staircase")
#making a no-pooling model
m1 <- glm(correct ~ target.contrast + subject,
           data = samples_staircase,
           family = binomial(link = "logit"))
# We should see an intercept per subject
summary(m1)
```

```

## 
## Call:
## glm(formula = correct ~ target.contrast + subject, family = binomial(link = "logit"),
##      data = samples_staircase)
##
## Deviance Residuals:
##    Min      1Q  Median      3Q     Max
## -3.2225  0.1836  0.7262  0.7907  0.9772
##
## Coefficients:
##                               Estimate Std. Error z value Pr(>|z| )
## (Intercept)           1.237950  0.310228  3.990 6.59e-05 ***
## target.contrast   3.446129  0.439005  7.850 4.17e-15 ***
## subject002        -0.200038  0.396544 -0.504  0.6139
## subject003        -0.126797  0.367239 -0.345  0.7299
## subject004        -0.447583  0.341680 -1.310  0.1902
## subject005        -0.729034  0.342889 -2.126  0.0335 *
## subject006         0.204637  0.412265  0.496  0.6196
## subject007        -0.257607  0.397907 -0.647  0.5174
## subject008        -0.728765  0.345702 -2.108  0.0350 *
## subject009        -0.600476  0.339050 -1.771  0.0766 .
## subject010        -0.450820  0.353083 -1.277  0.2017
## subject011        -0.003327  0.353925 -0.009  0.9925
## subject012        -0.751795  0.339166 -2.217  0.0267 *
## subject013        -0.251208  0.354408 -0.709  0.4784
## subject014        -0.279840  0.345979 -0.809  0.4186
## subject015        -0.330359  0.349908 -0.944  0.3451
## subject016        -0.781364  0.340545 -2.294  0.0218 *
## subject017        -0.691345  0.339823 -2.034  0.0419 *
## subject018        -0.546979  0.340267 -1.607  0.1079
## subject019         0.502524  0.488717  1.028  0.3038
## subject020        -0.271168  0.356758 -0.760  0.4472
## subject021        -0.399562  0.346301 -1.154  0.2486
## subject022         0.523698  0.488385  1.072  0.2836
## subject023        -0.065056  0.359321 -0.181  0.8563
## subject024        -0.531046  0.341990 -1.553  0.1205
## subject025        -0.160067  0.426526 -0.375  0.7075
## subject026        -0.222237  0.374445 -0.594  0.5528
## subject027        -0.487602  0.342189 -1.425  0.1542
## subject028        -0.547571  0.339972 -1.611  0.1073
## subject029        -0.335918  0.355318 -0.945  0.3445
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
## Null deviance: 6127.2 on 5602 degrees of freedom
## Residual deviance: 5936.1 on 5573 degrees of freedom
## AIC: 5996.1
##
## Number of Fisher Scoring iterations: 6

```

```
# plot the fitted values per subject
samples_staircase %>%
  ggplot(aes(target.contrast, as.integer(correct), color = correct)) +
  geom_point() +
  geom_line(aes(target.contrast, fitted(m1),
    linetype = "fitted values"), inherit.aes = FALSE) +
  scale_linetype_manual(name = "single-level model", values = c("dashed")) +
  facet_wrap(~subject) +
  labs(title = "Correct answers by target contrast per participant",
    subtitle = "With single-level model fitted values (no-pooling)",
    color = "subject correct") +
  ylab("Correct (probability)") +
  xlab("target contrast")
```



Ex1.p2.iii Comment on the fits - do we have enough data to plot the logistic functions?

The fits are not great. We can see that the lower contrast seems to result in more incorrect answers but even so, subjects still get correct answer at low contrast. As this is a logical outcome variable, the probabilities for most subjects start around 75%. If the contrast is 0.00, we would expect that the subject would not be able to correctly identify the target stimulus, and should start at 50% (chance level). It does seem as though there is sufficient data, the minimum number of samples for a subject is 70, although there are not the same number of samples for each contrast value.

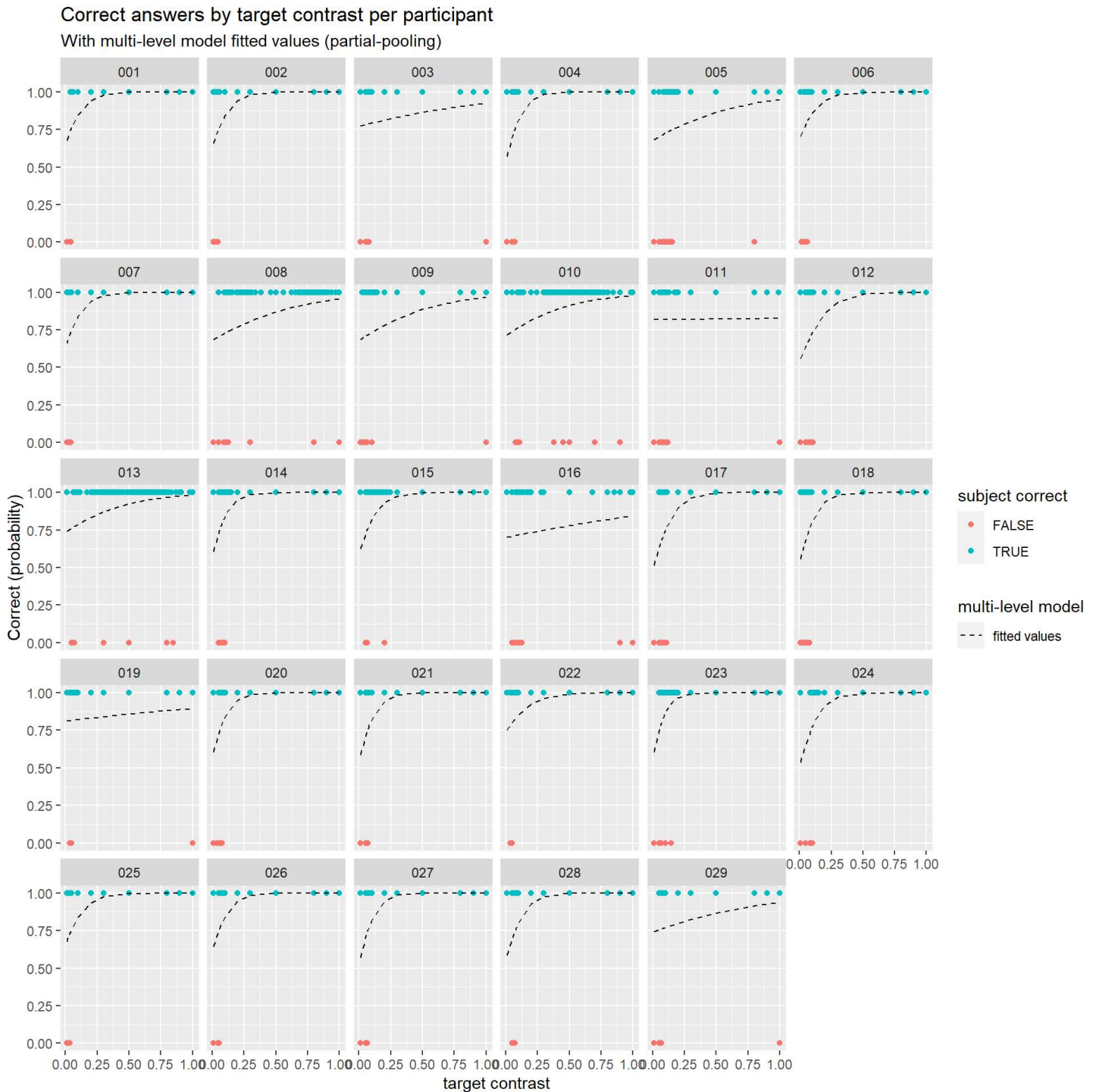
```
m2 <- glmer(correct ~ target.contrast + (1 + target.contrast | subject),
  data = samples_staircase,
  family = binomial(link = "logit"))
summary(m2)
```

```
## Generalized linear mixed model fit by maximum likelihood (Laplace
## Approximation) [glmerMod]
## Family: binomial ( logit )
## Formula: correct ~ target.contrast + (1 + target.contrast | subject)
## Data: samples_staircase
##
##      AIC      BIC  logLik deviance df.resid
##  5988.5   6021.6  -2989.2    5978.5     5598
##
## Scaled residuals:
##      Min      1Q  Median      3Q      Max
## -5.7671  0.0068  0.5532  0.5915  0.9264
##
## Random effects:
## Groups   Name        Variance Std.Dev. Corr
## subject (Intercept)  0.2717   0.5213
##           target.contrast 42.7577  6.5389  -0.84
## Number of obs: 5603, groups: subject, 29
##
## Fixed effects:
##             Estimate Std. Error z value Pr(>|z|)
## (Intercept)  0.5619    0.1580   3.557 0.000376 ***
## target.contrast 8.7132    2.3604   3.691 0.000223 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##          (Intr)
## trgt.cntrst -0.907
```

```

samples_staircase %>%
  ggplot(aes(target.contrast, as.integer(correct)), color = correct) +
  geom_point() +
  geom_line(aes(target.contrast, fitted(m2),
    linetype = "fitted values"), inherit.aes = FALSE) +
  scale_linetype_manual(name = "multi-level model", values = c("dashed")) +
  facet_wrap(~subject) +
  labs(title = "Correct answers by target contrast per participant",
      subtitle = "With multi-level model fitted values (partial-pooling)",
      color = "subject correct") +
  ylab("Correct (probability)") +
  xlab("target contrast")

```



Ex1.p2. v. in your own words, describe how the partial pooling model allows for a better fit for each subject

It allows for per-subject differences in their response to the target contrast. If some subjects have more difficulty seeing differences in low contrast images, we would expect their accuracy to decrease more than other subjects as the contrast values becomes lower. We would conceptually expect no-pooling models to create a better fit for each subject(?), as the subjective models are modelled solely on a given subject. However we would of course always choose the partial pooling models as these would give us great subject-fits but also a model generalizable for the population.

## Exercise 2

Now we **only** look at the *experiment* trials (*trial.type*)

1. Pick four subjects and plot their Quantile-Quantile (Q-Q) plots for the residuals of their objective response times (*rt.obj*) based on a model where only intercept is modelled
  - i. comment on these
  - ii. does a log-transformation of the response time data improve the Q-Q-plots?
2. Now do a partial pooling model modelling objective response times as dependent on *task*? (set *REML=FALSE* in your *lmer* -specification)
  - i. which would you include among your random effects and why? (support your choices with relevant measures, taking into account variance explained and number of parameters going into the modelling)
  - ii. explain in your own words what your chosen models says about response times between the different tasks
3. Now add *pas* and its interaction with *task* to the fixed effects
  - i. how many types of group intercepts (random effects) can you add without ending up with convergence issues or singular fits?
  - ii. create a model by adding random intercepts (without modelling slopes) that results in a singular fit - then use  
`print(VarCorr(<your.model>), comp='Variance')` to inspect the variance vector - explain why the fit is singular (Hint: read the first paragraph under details in the help for *isSingular* )
  - iii. in your own words - how could you explain why your model would result in a singular fit?

## Answers

### Exercise 2, part 1 - VK

We will use subjects 002, 009, 014 and 015.

```

samples_experiment <- samples %>% filter(
  trial.type == "experiment" &
  (
    subject == "002" |
    subject == "009" |
    subject == "014" |
    subject == "015"
  ))
m3 <- lmer(rt.obj ~ (1 | subject),
  data = samples_experiment)
summary(m3)

```

```

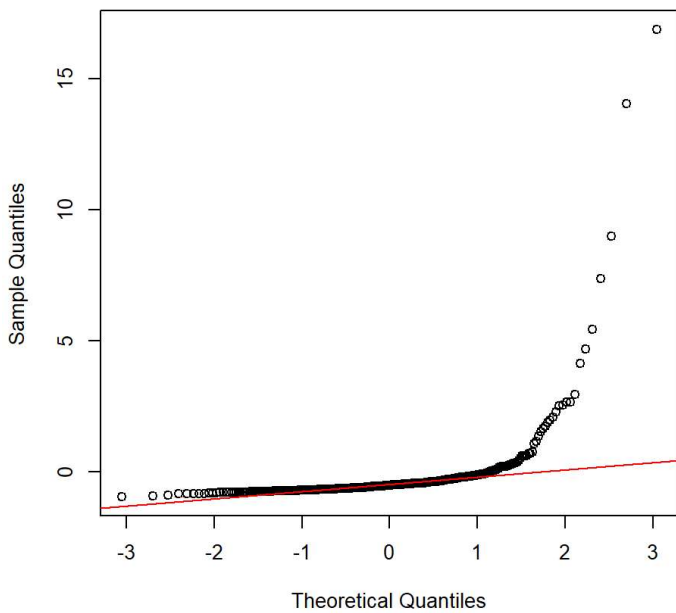
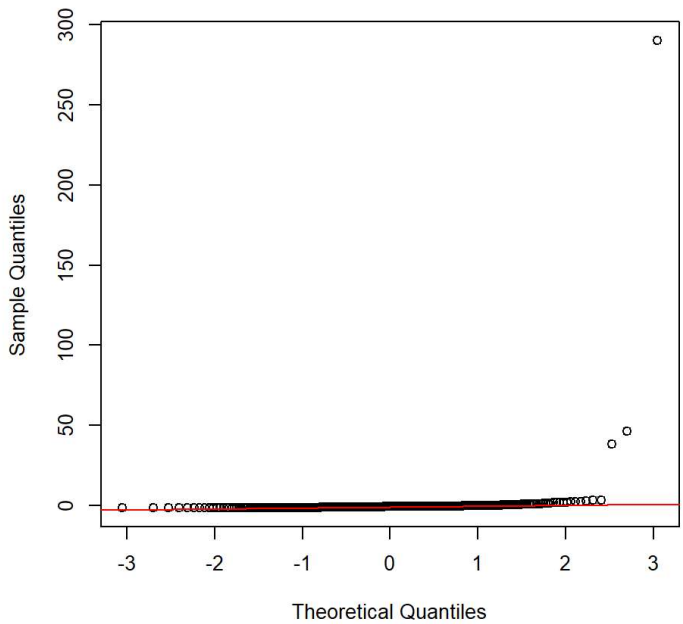
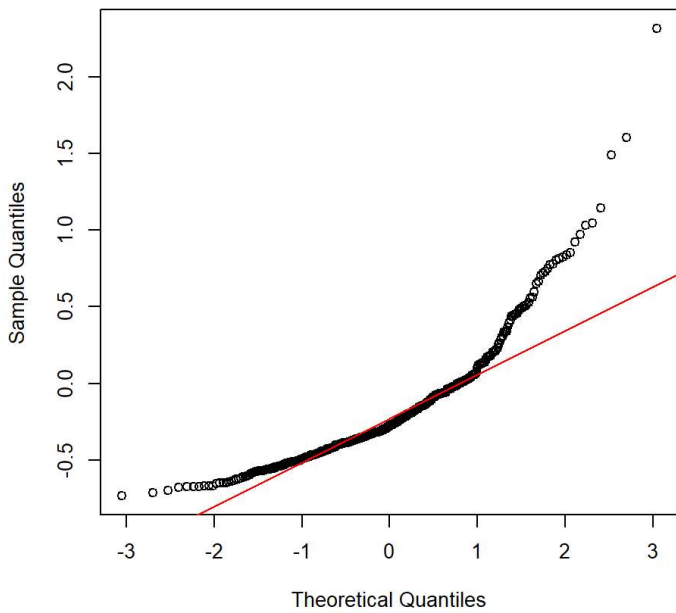
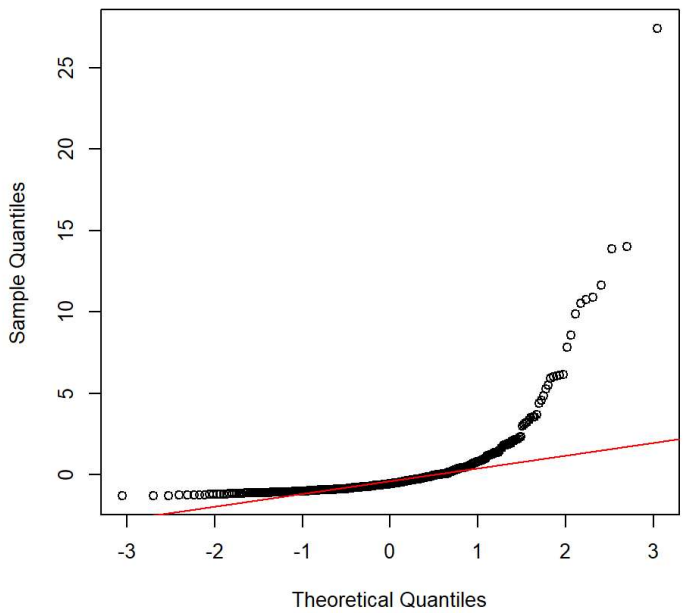
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [ 
## lmerModLmerTest]
## Formula: rt.obj ~ (1 | subject)
##   Data: samples_experiment
##
## REML criterion at convergence: 11774.5
##
## Scaled residuals:
##   Min     1Q Median     3Q    Max
## -0.214 -0.098 -0.063 -0.019 39.793
##
## Random effects:
##   Groups   Name        Variance Std.Dev.
##   subject (Intercept) 0.1621  0.4026
##   Residual           53.2091  7.2945
## Number of obs: 1728, groups: subject, 4
##
## Fixed effects:
##             Estimate Std. Error   df t value Pr(>|t|)    
## (Intercept) 1.239      0.267 3.000  4.64   0.0189 *
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

```

```

samples_experiment$m3_resid <- residuals(m3)
# make 2x2 plot grid
par(mfrow = c(2, 2))
# plot qq plots for objective response time of subjects:
# 002
qqnorm(samples_experiment[samples_experiment$subject == "002", ]$m3_resid)
qqline(samples_experiment[samples_experiment$subject == "002", ]$m3_resid,
  col = "red")
# 009
qqnorm(samples_experiment[samples_experiment$subject == "009", ]$m3_resid)
qqline(samples_experiment[samples_experiment$subject == "009", ]$m3_resid,
  col = "red")
# 014
qqnorm(samples_experiment[samples_experiment$subject == "014", ]$m3_resid)
qqline(samples_experiment[samples_experiment$subject == "014", ]$m3_resid,
  col = "red")
# 015
qqnorm(samples_experiment[samples_experiment$subject == "015", ]$m3_resid)
qqline(samples_experiment[samples_experiment$subject == "015", ]$m3_resid,
  col = "red")

```

**Normal Q-Q Plot****Normal Q-Q Plot****Normal Q-Q Plot****Normal Q-Q Plot**

## Ex2.p1.i

These data show a significant tail with some reaction times going far beyond the mean (e.g. 15 seconds +).

```
m3l <- lmer(log(rt.obj) ~ (1 | subject),  
           data = samples_experiment)  
summary(m3l)
```

```

## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: log(rt.obj) ~ (1 | subject)
##   Data: samples_experiment
##
## REML criterion at convergence: 4775.9
##
## Scaled residuals:
##      Min     1Q Median     3Q    Max 
## -8.4491 -0.4344  0.0168  0.4933  6.4941 
##
## Random effects:
## Groups   Name        Variance Std.Dev. 
## subject (Intercept) 0.08853  0.2975 
## Residual           0.92008  0.9592 
## Number of obs: 1728, groups: subject, 4 
##
## Fixed effects:
##             Estimate Std. Error   df t value Pr(>|t|)    
## (Intercept) -0.3916    0.1505 3.0000 -2.601   0.0803 .  
## --- 
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

```

```

samples_experiment$m3l_resid <- residuals(m3l)
# plot qq plots for log transformed objective response time of subjects:
# 002
qqnorm(log(samples_experiment[samples_experiment$subject == "002", ]$m3l_resid))

```

```

## Warning in log(samples_experiment[samples_experiment$subject == "002", ]
## $m3l_resid): NaNs produced

```

```

qqline(log(samples_experiment[samples_experiment$subject == "002", ]$m3l_resid),
       col = "red")

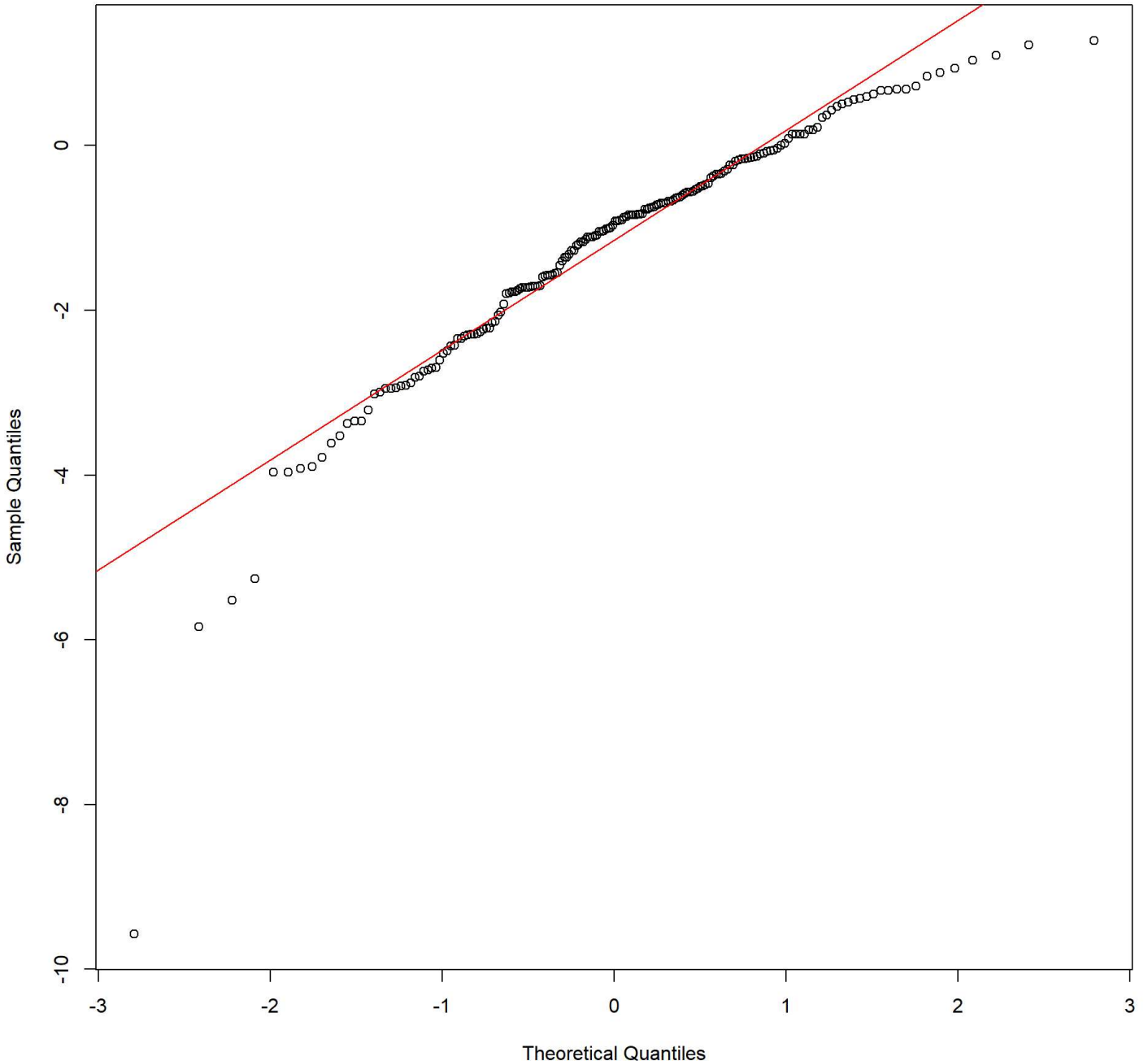
```

```

## Warning in log(samples_experiment[samples_experiment$subject == "002", ]
## $m3l_resid): NaNs produced

```

### Normal Q-Q Plot



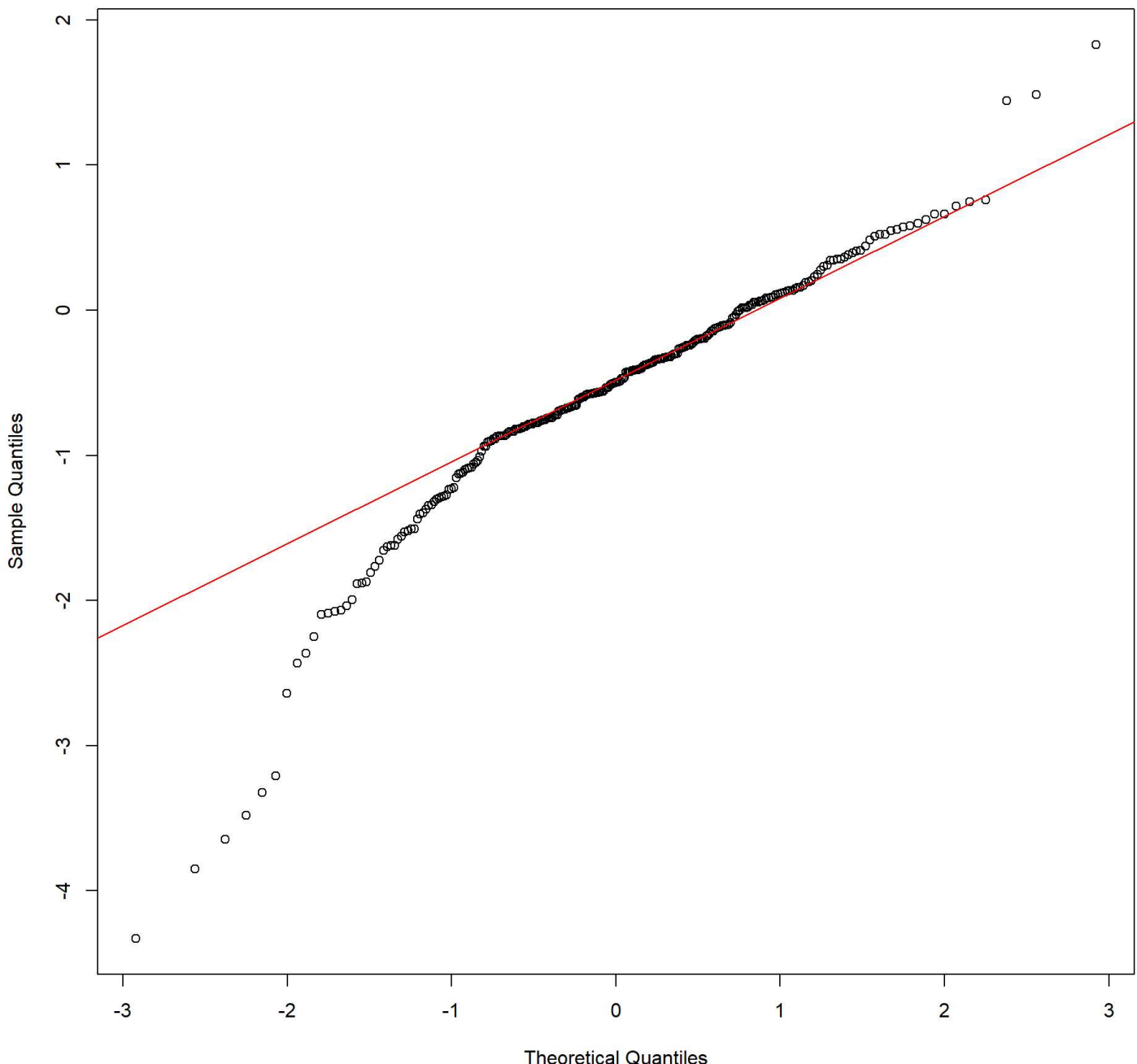
```
# 009
qnorm(log(samples_experiment[samples_experiment$subject == "009", ]$m3l_resid))
```

```
## Warning in log(samples_experiment[samples_experiment$subject == "009", ]
## $m3l_resid): NaNs produced
```

```
qqline(log(samples_experiment[samples_experiment$subject == "009", ]$m3l_resid),
       col = "red")
```

```
## Warning in log(samples_experiment[samples_experiment$subject == "009", ]
## $m3l_resid): NaNs produced
```

### Normal Q-Q Plot



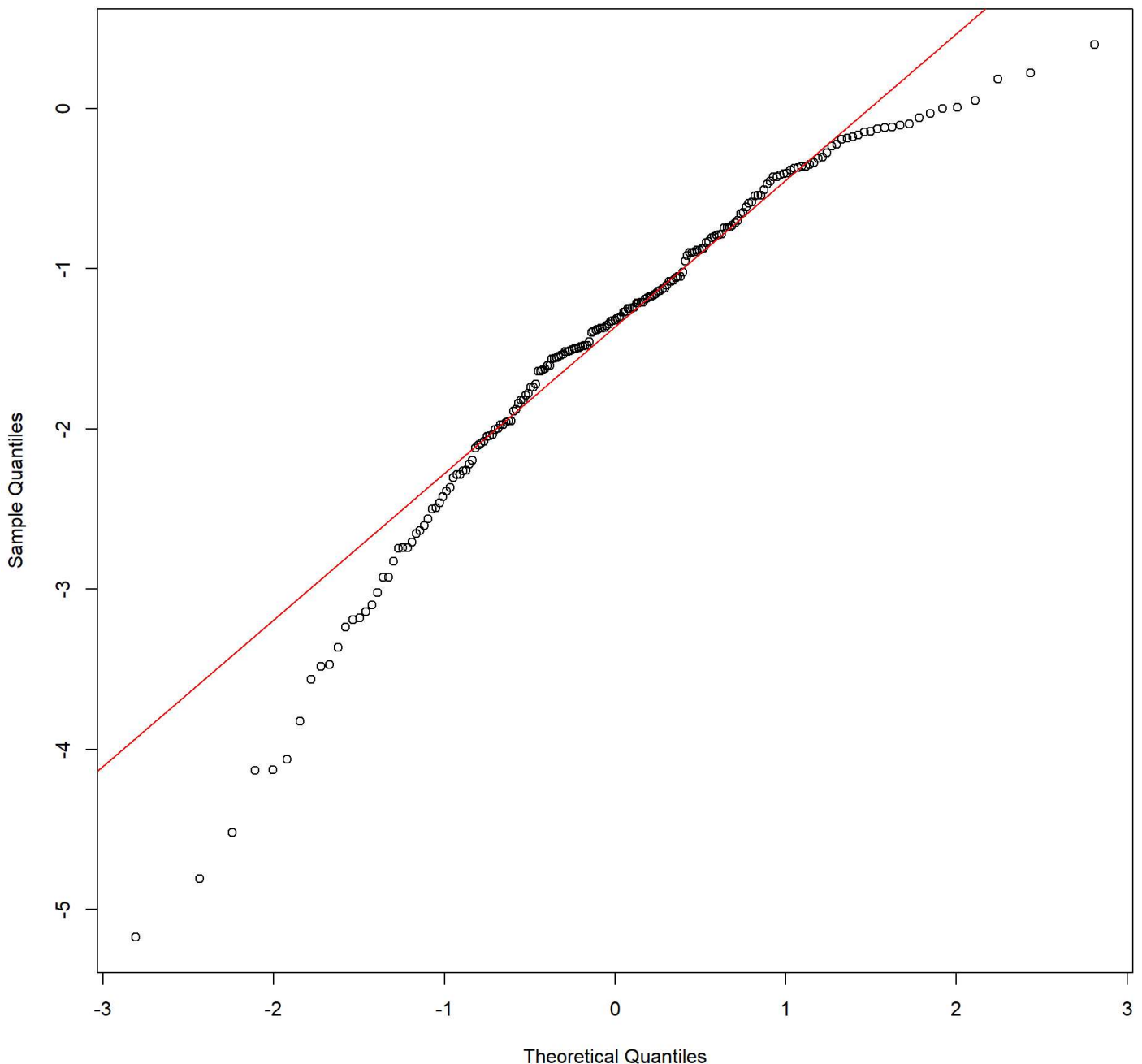
```
# 014
qnorm(log(samples_experiment[samples_experiment$subject == "014", ]$m3l_resid))
```

```
## Warning in log(samples_experiment[samples_experiment$subject == "014", ]
## $m3l_resid): NaNs produced
```

```
qqline(log(samples_experiment[samples_experiment$subject == "014", ]$m3l_resid),
       col = "red")
```

```
## Warning in log(samples_experiment[samples_experiment$subject == "014", ]
## $m3l_resid): NaNs produced
```

### Normal Q-Q Plot



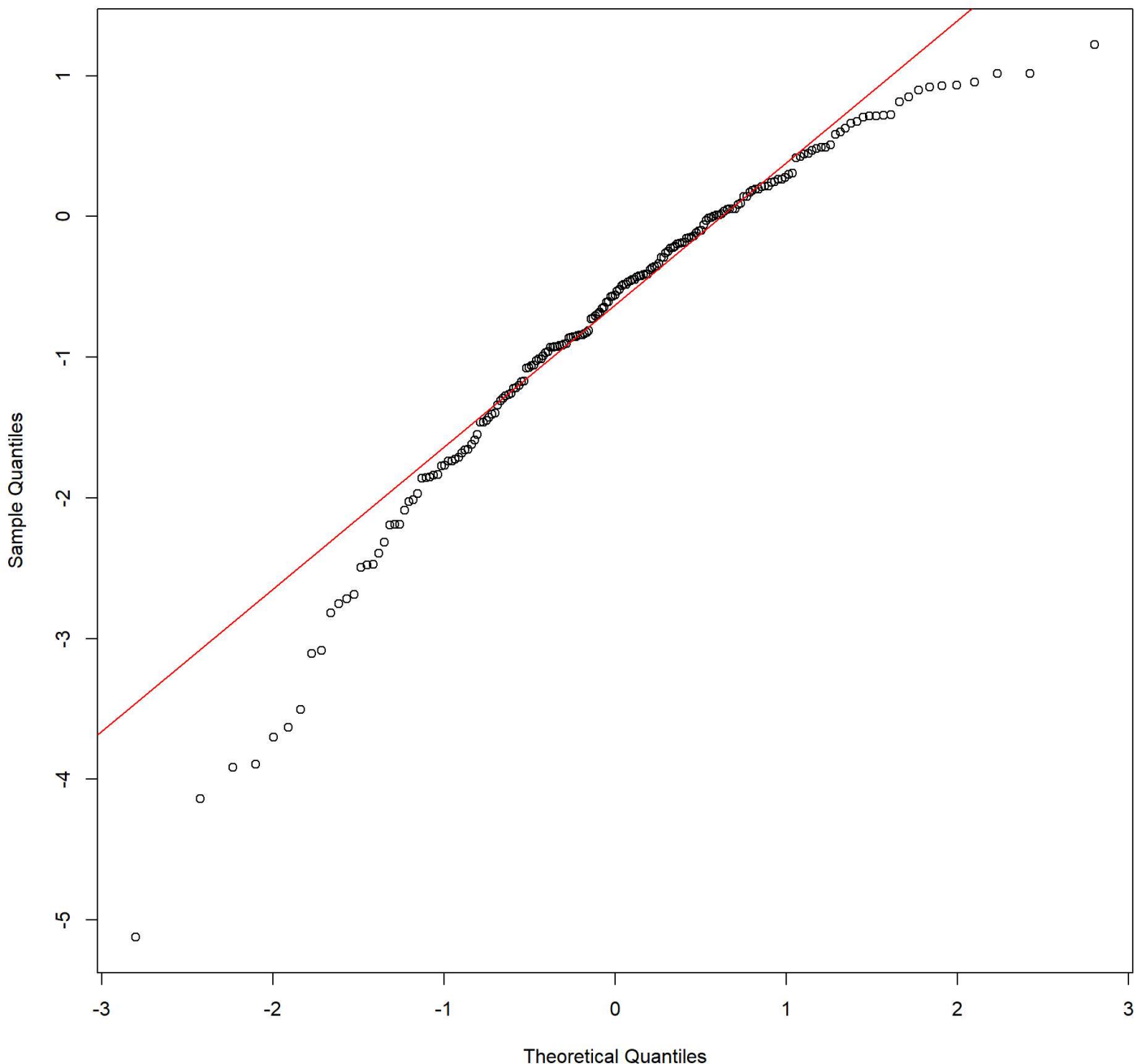
```
# 015
qqnorm(log(samples_experiment[samples_experiment$subject == "015", ]$m3l_resid))
```

```
## Warning in log(samples_experiment[samples_experiment$subject == "015", ]
## $m3l_resid): NaNs produced
```

```
qqline(log(samples_experiment[samples_experiment$subject == "015", ]$m3l_resid),
       col = "red")
```

```
## Warning in log(samples_experiment[samples_experiment$subject == "015", ]
## $m3l_resid): NaNs produced
```

Normal Q-Q Plot



```
# reset plot grid
par(mfrow = c(1, 1))
```

## Ex2.p1.ii

Log-transforming the data definitely creates a better fit, as can be seen in the qq-plots.

## Exercise 2, part 2 - VK

```
m4 <- lmer(rt.obj ~ task + (1 | subject),
            REML = FALSE, data = samples_experiment)
summary(m4)
```

```

## Linear mixed model fit by maximum likelihood . t-tests use Satterthwaite's
##   method [lmerModLmerTest]
## Formula: rt.obj ~ task + (1 | subject)
## Data: samples_experiment
##
##      AIC      BIC  logLik deviance df.resid
## 11780.8 11808.0 -5885.4 11770.8     1723
##
## Scaled residuals:
##    Min     1Q Median     3Q    Max
## -0.258 -0.111 -0.058 -0.010 39.780
##
## Random effects:
## Groups   Name        Variance Std.Dev.
## subject (Intercept) 0.09096 0.3016
## Residual           53.12390 7.2886
## Number of obs: 1728, groups: subject, 4
##
## Fixed effects:
##             Estimate Std. Error      df t value Pr(>|t|)    
## (Intercept) 1.6487    0.3391 18.4270 4.863 0.000117 ***
## taskquadruplet -0.6554    0.4295 1724.0000 -1.526 0.127187  
## tasksingles   -0.5736    0.4295 1724.0000 -1.336 0.181883  
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##          (Intr) tskqdr
## taskquadrpl -0.633
## tasksingles -0.633  0.500

```

```
piecewiseSEM:::rsquared(m4)
```

```

## Response family link method Marginal Conditional
## 1 rt.obj gaussian identity none 0.001596197 0.00330281

```

The above model uses random intercepts for subject. We considered that task and trial might be good additional random effects, but these result in a singular fit. We considered task to be a relevant random effect, as it was expected that the different tasks would influence the reaction time. We considered trial to be a relevant random effect as participants might perform better throughout trials.

Overall this model does not explain much of the variance (0.33% about 0.16% fixed effects and 0.17% random effects). The model shows no statistically significant difference between response times per task.

## Exercise 2, part 3 - LR

```

# model with PAS-task interaction.
m5 <- lmer(rt.obj ~ task + pas:task + (1 | subject),
            REML = FALSE, data = samples_experiment)
summary(m5)

```

```

## Linear mixed model fit by maximum likelihood . t-tests use Satterthwaite's
##   method [lmerModLmerTest]
## Formula: rt.obj ~ task + pas:task + (1 | subject)
## Data: samples_experiment
##
##      AIC      BIC  logLik deviance df.resid
## 11796.6 11872.9 -5884.3 11768.6     1714
##
## Scaled residuals:
##    Min     1Q Median     3Q    Max
## -0.310 -0.100 -0.054 -0.005 39.752
##
## Random effects:
## Groups   Name        Variance Std.Dev.
## subject (Intercept) 0.08247 0.2872
## Residual           53.06137 7.2843
## Number of obs: 1728, groups: subject, 4
##
## Fixed effects:
##             Estimate Std. Error      df t value Pr(>|t|) 
## (Intercept)  0.8696   1.7248 1487.2418  0.504   0.614
## taskquadruplet -0.2514   2.5048 1727.9918 -0.100   0.920
## tasksingles   -0.2318   2.1467 1726.1882 -0.108   0.914
## taskpairs:pas3  0.4921   1.9054 1726.6724  0.258   0.796
## taskquadruplet:pas3 0.3666   2.0816 1720.4025  0.176   0.860
## tasksingles:pas3  0.3560   1.5592 1727.9856  0.228   0.819
## taskpairs:pas2   0.4343   1.7897 1726.8105  0.243   0.808
## taskquadruplet:pas2 0.5845   1.8898 1663.2112  0.309   0.757
## tasksingles:pas2   0.3804   1.3794 1675.4298  0.276   0.783
## taskpairs:pas1    1.1935   1.7775 1717.3364  0.671   0.502
## taskquadruplet:pas1 0.2221   1.8775 1711.0815  0.118   0.906
## tasksingles:pas1   0.5702   1.3781 1486.2328  0.414   0.679
##
## Correlation of Fixed Effects:
##          (Intr) tskqdr tsksng tskp:3 tskq:3 tsks:3 tskp:2 tskq:2 tsks:2
## taskquadruplt -0.682
## tasksingles  -0.796  0.548
## taskprs:ps3  -0.898  0.617  0.720
## tskqdrpl:3   0.000 -0.638  0.000  0.001
## tsksngls:p3 -0.001  0.001 -0.496  0.000 -0.001
## taskprs:ps2  -0.957  0.657  0.767  0.865  0.001  0.000
## tskqdrpl:2   -0.003 -0.703  0.000  0.001  0.847 -0.001  0.003
## tsksngls:p2 -0.004  0.000 -0.561  0.001  0.001  0.772  0.003  0.005
## taskprs:ps1  -0.964  0.662  0.772  0.871  0.000  0.002  0.929  0.003  0.005
## tskqdrpl:1   -0.002 -0.707  0.000  0.001  0.852  0.000  0.002  0.940  0.004
## tsksngls:p1 -0.005  0.001 -0.563  0.002 -0.001  0.776  0.003  0.003  0.880
##                  tskp:1 tskq:1
## taskquadruplt
## tasksingles
## taskprs:ps3
## tskqdrpl:3
## tsksngls:p3
## taskprs:ps2
## tskqdrpl:2
## tsksngls:p2
## taskprs:ps1

```

```
## tskqdrplt:1 0.002
## tsksngls:p1 0.007 0.004
```

## Ex2.p3.i

Any additional random effects we added besides subject resulted in singular fits.

```
# here already it becomes a singular fit
m6 <- lmer(rt.obj ~ task + pas:task + (1 | subject) + (1 | task),
    REML = FALSE, data = samples_experiment)
```

```
## boundary (singular) fit: see ?isSingular
```

```
summary(m6)
```

```

## Linear mixed model fit by maximum likelihood . t-tests use Satterthwaite's
##   method [lmerModLmerTest]
## Formula: rt.obj ~ task + pas:task + (1 | subject) + (1 | task)
## Data: samples_experiment
##
##      AIC      BIC logLik deviance df.resid
## 11798.6 11880.4 -5884.3 11768.6     1713
##
## Scaled residuals:
##    Min     1Q Median     3Q    Max
## -0.310 -0.100 -0.054 -0.005 39.752
##
## Random effects:
## Groups   Name        Variance Std.Dev.
## subject  (Intercept) 8.247e-02 2.872e-01
## task     (Intercept) 1.268e-14 1.126e-07
## Residual            5.306e+01 7.284e+00
## Number of obs: 1728, groups: subject, 4; task, 3
##
## Fixed effects:
##             Estimate Std. Error      df t value Pr(>|t|) 
## (Intercept)  0.8696   1.7248 1487.2418  0.504  0.614
## taskquadruplet -0.2514   2.5048 1727.9918 -0.100  0.920
## tasksingles   -0.2318   2.1467 1726.1882 -0.108  0.914
## taskpairs:pas3  0.4921   1.9054 1726.6724  0.258  0.796
## taskquadruplet:pas3 0.3666   2.0816 1720.4025  0.176  0.860
## tasksingles:pas3  0.3560   1.5592 1727.9856  0.228  0.819
## taskpairs:pas2   0.4343   1.7897 1726.8105  0.243  0.808
## taskquadruplet:pas2 0.5845   1.8898 1663.2112  0.309  0.757
## tasksingles:pas2   0.3804   1.3794 1675.4298  0.276  0.783
## taskpairs:pas1    1.1935   1.7775 1717.3364  0.671  0.502
## taskquadruplet:pas1 0.2221   1.8775 1711.0815  0.118  0.906
## tasksingles:pas1    0.5702   1.3781 1486.2328  0.414  0.679
##
## Correlation of Fixed Effects:
##          (Intr) tskqdr tsksng tskp:3 tskq:3 tsks:3 tskp:2 tskq:2 tsks:2
## taskquadruplet -0.682
## tasksingles   -0.796  0.548
## taskprs:ps3   -0.898  0.617  0.720
## tskqdrplt:3   0.000 -0.638  0.000  0.001
## tsksngls:p3   -0.001  0.001 -0.496  0.000 -0.001
## taskprs:ps2   -0.957  0.657  0.767  0.865  0.001  0.000
## tskqdrplt:2   -0.003 -0.703  0.000  0.001  0.847 -0.001  0.003
## tsksngls:p2   -0.004  0.000 -0.561  0.001  0.001  0.772  0.003  0.005
## taskprs:ps1   -0.964  0.662  0.772  0.871  0.000  0.002  0.929  0.003  0.005
## tskqdrplt:1   -0.002 -0.707  0.000  0.001  0.852  0.000  0.002  0.940  0.004
## tsksngls:p1   -0.005  0.001 -0.563  0.002 -0.001  0.776  0.003  0.003  0.880
##          tskp:1 tskq:1
## taskquadruplet
## tasksingles
## taskprs:ps3
## tskqdrplt:3
## tsksngls:p3
## taskprs:ps2
## tskqdrplt:2
## tsksngls:p2
## taskprs:ps1

```

```
## tskqdrplt:1 0.002
## tsksngls:p1 0.007 0.004
## optimizer (nloptwrap) convergence code: 0 (OK)
## boundary (singular) fit: see ?isSingular
```

```
print(VarCorr(m6), comp = "Variance")
```

```
## Groups   Name        Variance
## subject (Intercept) 8.2468e-02
## task    (Intercept) 1.2681e-14
## Residual                      5.3061e+01
```

Ex2.p3.iii. in your own words - how could you explain why your model would result in a singular fit?

In the example above, model m6, it results in a singular fit. Singular fits occur when the variances of some of the linear combinations of the effects are either zero or close to zero, and it is indicating that the model may be overfitting for the data. In model m6, we get the variance for task as 0, which results in a singular fit - this indicates to us that this model is overfitted to the data.

We expect that the reason that the model results in a singular fit is because we have too many parameters, and is therefore too complex. As mentioned above, adding any additional random effects besides subject resulted in getting singular fits, so in order for the example above, m6, to not result in a singular fit, we would have to remove (1|Task).

## Exercise 3

1. Initialise a new data frame, `data.count`. `count` should indicate the number of times they categorized their experience as `pas` 1-4 for each `task`. I.e. the data frame would have for subject 1: for task:singles, `pas1` was used # times, `pas2` was used # times, `pas3` was used # times and `pas4` was used # times. You would then do the same for task:pairs and task:quadruplet
2. Now fit a multilevel model that models a unique “slope” for `pas` for each `subject` with the interaction between `pas` and `task` and their main effects being modelled
  - i. which family should be used?
  - ii. why is a slope for `pas` not really being modelled?
  - iii. if you get a convergence error, try another algorithm (the default is the `Nelder_Mead`) - try (`bobyqa`) for which the `dfoptim` package is needed. In `glmer`, you can add the following for the `control` argument:  
`glmerControl(optimizer="bobyqa")` (if you are interested, also have a look at the function `allFit`)
  - iv. when you have a converging fit - fit a model with only the main effects of `pas` and `task`. Compare this with the model that also includes the interaction

- v. indicate which of the two models, you would choose and why
  - vi. based on your chosen model - write a short report on what this says about the distribution of ratings as dependent on *pas* and *task*
  - vii. include a plot that shows the estimated amount of ratings for four subjects of your choosing
3. Finally, fit a multilevel model that models *correct* as dependent on *task* with a unique intercept for each *subject*
- i. does *task* explain performance?
  - ii. add *pas* as a main effect on top of *task* - what are the consequences of that?
  - iii. now fit a multilevel model that models *correct* as dependent on *pas* with a unique intercept for each *subject*
  - iv. finally, fit a model that models the interaction between *task* and *pas* and their main effects
  - v. describe in your words which model is the best in explaining the variance in accuracy

## Answers

### Exercise 3, part 1 - KV

```
data.count <- samples %>% select(subject, pas, task, correct) %>%
  group_by(subject, task, pas) %>%
  summarize(
    subject = subject[1],
    task = task[1],
    pas = pas[1],
    count = n(),
    accuracy = sum(correct) / n(),
    .groups = "drop")
head(data.count)
```

```
## # A tibble: 6 x 5
##   subject task      pas  count accuracy
##   <fct>   <fct>  <fct> <int>    <dbl>
## 1 001     pairs    4      12     1
## 2 001     pairs    3       4     1
## 3 001     pairs    2      45    0.756
## 4 001     pairs    1     109    0.532
## 5 001     quadruplet 4       1     1
## 6 001     quadruplet 3       3     1
```

### Exercise 3, part 2 EH

```
# multi-level with slope and interaction
m7 <- glmer(count ~
  pas + task + pas:task +
  (1 + pas | subject),
  data = data.count,
  family = poisson,
  control = glmerControl(optimizer = "bobyqa")) #added control, ex3.p2.iii
summary(m7)
```

```

## Generalized linear mixed model fit by maximum likelihood (Laplace
## Approximation) [glmerMod]
## Family: poisson  ( log )
## Formula: count ~ pas + task + pas:task + (1 + pas | subject)
## Data: data.count
## Control: glmerControl(optimizer = "bobyqa")
##
##      AIC      BIC  logLik deviance df.resid
##  3148.4  3232.7 -1552.2   3104.4     318
##
## Scaled residuals:
##      Min    1Q Median    3Q   Max
## -4.3871 -0.7853 -0.0469  0.7550  6.5438
##
## Random effects:
## Groups   Name        Variance Std.Dev. Corr
## subject (Intercept) 1.172    1.083
##         pas3        1.146    1.071   -0.80
##         pas2        1.956    1.398   -0.97  0.79
##         pas1        2.374    1.541   -0.96  0.70  0.92
## Number of obs: 340, groups: subject, 29
##
## Fixed effects:
##                               Estimate Std. Error z value Pr(>|z| )
## (Intercept)            3.262772  0.206184 15.825 < 2e-16 ***
## pas3                  0.259281  0.206702  1.254  0.20971
## pas2                  0.749155  0.264869  2.828  0.00468 **
## pas1                  0.772932  0.290752  2.658  0.00785 **
## taskquadruplet       -0.100104  0.041920 -2.388  0.01694 *
## tasksingles           0.332516  0.037868  8.781 < 2e-16 ***
## pas3:taskquadruplet  0.005989  0.059782  0.100  0.92020
## pas2:taskquadruplet  0.101249  0.053853  1.880  0.06010 .
## pas1:taskquadruplet  0.215004  0.052299  4.111 3.94e-05 ***
## pas3:tasksingles     -0.320468  0.056111 -5.711 1.12e-08 ***
## pas2:tasksingles     -0.368099  0.050973 -7.221 5.15e-13 ***
## pas1:tasksingles     -0.563462  0.051014 -11.045 < 2e-16 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
## (Intr) pas3  pas2  pas1  tskqdr tsksng ps3:tskq ps2:tskq
## pas3   -0.797
## pas2   -0.969  0.786
## pas1   -0.959  0.702  0.911
## taskquadruplet -0.105  0.104  0.081  0.074
## tasksingles  -0.107  0.107  0.083  0.076  0.526
## ps3:tskqdrp   0.073 -0.142 -0.057 -0.052 -0.701 -0.369
## ps2:tskqdrp   0.081 -0.081 -0.103 -0.058 -0.778 -0.410  0.546
## ps1:tskqdrp   0.084 -0.084 -0.065 -0.093 -0.802 -0.422  0.562   0.624
## ps3:tsksngl   0.072 -0.146 -0.056 -0.051 -0.355 -0.675  0.506   0.276
## ps2:tsksngl   0.079 -0.079 -0.104 -0.056 -0.391 -0.743  0.274   0.513
## ps1:tsksngl   0.079 -0.079 -0.062 -0.091 -0.391 -0.742  0.274   0.304
##                  ps1:tskq ps3:tsks ps2:tsks
## pas3
## pas2
## pas1
## taskquadruplet

```

```
## tasksingles
## ps3:tskqdrp
## ps2:tskqdrp
## ps1:tskqdrp
## ps3:tsksngl  0.285
## ps2:tsksngl  0.313    0.501
## ps1:tsksngl  0.507    0.501    0.551
```

### Ex3.p2.i

For this model (m7) we used the poisson family because the poisson distribution is specifically used to describe probabilities of event frequencies.

### Ex3.p2.ii

There is not really a slope being modelled for PAS as PAS is not continuous, it is a factor, so we only get estimates for each PAS level. As a factor, PAS is treated categorically (PAS1, PAS2, PAS3, PAS4), there is no such thing as PAS 1.3 etc and as such, it can not really be modelled as a slope.

```
m7.1 <- glmer(count ~
  pas + task + (1 + pas | subject),
  data = data.count,
  family = poisson,
  control = glmerControl(optimizer = "bobyqa"))
summary(m7.1)
```

```

## Generalized linear mixed model fit by maximum likelihood (Laplace
## Approximation) [glmerMod]
## Family: poisson  ( log )
## Formula: count ~ pas + task + (1 + pas | subject)
## Data: data.count
## Control: glmerControl(optimizer = "bobyqa")
##
##      AIC      BIC  logLik deviance df.resid
##  3398.5  3459.8 -1683.3   3366.5     324
##
## Scaled residuals:
##      Min      1Q  Median      3Q     Max
## -5.5885 -0.9001 -0.0477  0.8253  6.5100
##
## Random effects:
## Groups   Name        Variance Std.Dev. Corr
## subject (Intercept) 1.203    1.097
##           pas3        1.122    1.059   -0.80
##           pas2        2.002    1.415   -0.97  0.80
##           pas1        2.422    1.556   -0.96  0.71  0.92
## Number of obs: 340, groups: subject, 29
##
## Fixed effects:
##             Estimate Std. Error z value Pr(>|z|)
## (Intercept) 3.340760  0.207576 16.094 <2e-16 ***
## pas3        0.153914  0.201722  0.763  0.4455
## pas2        0.657300  0.266004  2.471  0.0135 *
## pas1        0.663828  0.291951  2.274  0.0230 *
## taskquadruplet 0.003294  0.018188  0.181  0.8563
## tasksingles  0.004307  0.018177  0.237  0.8127
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##            (Intr) pas3  pas2  pas1  tskqdr
## pas3     -0.802
## pas2     -0.971  0.799
## pas1     -0.961  0.714  0.917
## taskquadruplet -0.047  0.004  0.003  0.002
## tasksingles -0.044  0.000  0.000  0.000  0.501

```

AIC(m7, m7.1)

```

##      df      AIC
## m7    22  3148.441
## m7.1 16  3398.549

```

Ex3.p2.v Indicate which of the two models, you would choose and why?

Between m7 and m7.1, we believe that the m7 is the best model. Out of the two, m7 is the only one that's predictors have any significance on the outcome, in particular, the interactions in m7 have significant effect on the count variable. When comparing the

AIC output, it is also shown that model m7 has a better fit, with an AIC of 3148.441, compared to m7.1 of 3398.549. Taking all this into consideration, we are inclined to use this model m7 out of the two as it is the better model.

Ex3.p2.vi based on your chosen model - write a short report on what this says about the distribution of ratings as dependent on *pas* and *task* ?

m7 tells us that there is an important interaction between *pas* and *task* when it comes to the distribution of ratings, as all interactions reach significance,  $p < 0.05$ , except for *pas3:taskquadruplet*,  $p = 0.92020$  and *pas2:taskquadruplet*,  $p = 0.06010$

We can see from m7 that there is a decrease of all the interactions between pass's and quadruplet tasks (*pas:quadruplet*), est. -0.0719. We see the opposite in the case all counts of interactions between pass and single tasks, where there is an increase of est. ~0.169.

Looking at *pas* in m7, there is an increase of count in all *pas* when going from *pas4* to *pas1* (0.772932), 2 (est. 0.749155) and 3 (est. 0.259281). Only the predictions for *pas1* and *pas2* are significant,  $p < 0.05$ .

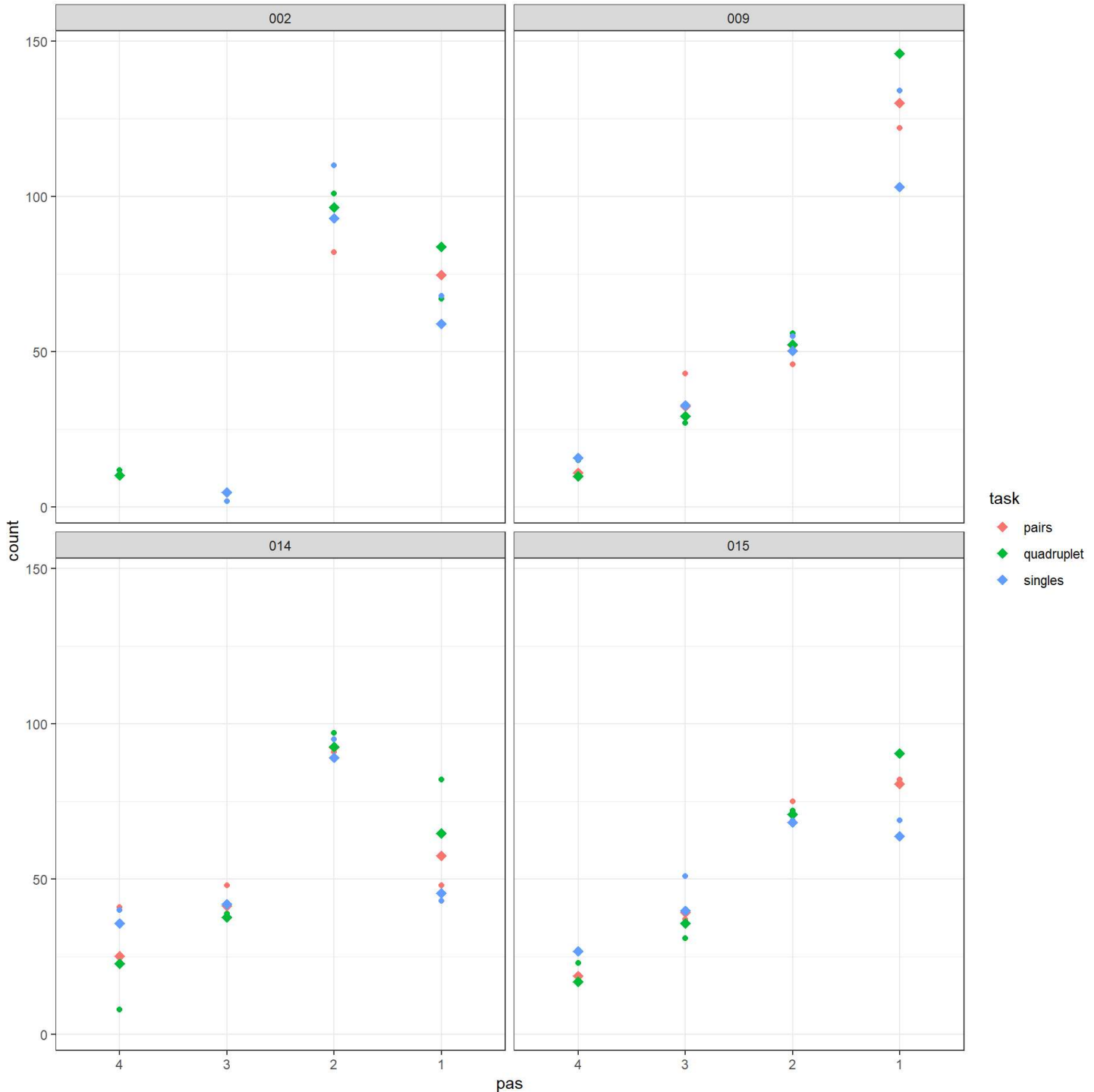
Regarding *task* in our model, count decreases going from task pairs to task quadruplets, est. -0.100, and count increases going from task pairs to task singles, ~ est. 0.333. Both of these predictions are significant,  $p < 0.05$ .

```
# get only specified subjects
data.count_subset <- data.count %>%
  filter(
    subject == "002" |
    subject == "009" |
    subject == "014" |
    subject == "015")
# predict using model for only selected subjects
m7_fitted <- predict(m7, newdata = data.count_subset)
data.count_subset$count_estimate <- expm1(m7_fitted)

print(m7_fitted)
```

```
##      1      2      3      4      5      6      7      8
## 1.746384 4.579301 4.326593 2.422614 4.580446 4.441493 1.758433 4.543718
##      9     10     11     12     13     14     15     16
## 4.095647 2.489337 3.508646 3.974888 4.875774 2.389234 3.414531 3.976033
##     17     18     19     20     21     22     23     24
## 4.990674 2.821854 3.520694 3.939305 4.644828 3.270353 3.749821 4.537376
##     25     26     27     28     29     30     31     32
## 4.070358 3.170249 3.655707 4.538521 4.185258 3.602869 3.761869 4.501793
##     33     34     35     36     37     38     39     40
## 3.839412 2.990216 3.697020 4.274627 4.402163 2.890113 3.602906 4.275772
##     41     42     43     44     45
## 4.517063 3.322733 3.709068 4.239044 4.171217
```

```
# plot
data.count_subset %>% ggplot(aes(pas, count, color = task)) + #smaller dots to see the original ratings
  geom_point() +
  theme_bw() +
  geom_point(aes(x = pas, y = count_estimate, color = task, ), shape = 18, size = 3) + #larger dots are to see the estimated ratings over top
  facet_wrap(data.count_subset$subject)
```



## Exercise 3, part 3 - LR

```
m8 <- glmer(correct ~ task +
  (1 | subject),
  data = samples,
  family = binomial(link = "logit"))
summary(m8)
```

```

## Generalized linear mixed model fit by maximum likelihood (Laplace
## Approximation) [glmerMod]
## Family: binomial ( logit )
## Formula: correct ~ task + (1 | subject)
## Data: samples
##
##      AIC      BIC  logLik deviance df.resid
## 19927.2 19958.4 -9959.6 19919.2     18127
##
## Scaled residuals:
##    Min     1Q Median     3Q    Max
## -2.7426 -1.0976  0.5098  0.6101  0.9111
##
## Random effects:
##   Groups   Name        Variance Std.Dev.
##   subject (Intercept) 0.1775   0.4214
## Number of obs: 18131, groups: subject, 29
##
## Fixed effects:
##             Estimate Std. Error z value Pr(>|z|)
## (Intercept) 1.10071   0.08386 13.125 < 2e-16 ***
## taskquadruplet -0.09825   0.04190 -2.345   0.019 *
## tasksingles    0.18542   0.04337  4.276 1.91e-05 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##          (Intr) tskqdr
## taskquadruplet -0.256
## tasksingles    -0.247  0.495

```

```

# seems task only explains a tiny amount of performance
piecewiseSEM:::rsquared(m8)

```

```

## Response family link method Marginal Conditional
## 1 correct binomial logit delta 0.002526141  0.03489787

```

### Ex3.p3.i

Differences in the task seem to explain performance (number of correct answers) to a statistically significant level.

```

m9 <- glmer(correct ~ task + pas +
  (1 | subject),
  data = samples,
  family = binomial(link = "logit"))
summary(m9)

```

```

## Generalized linear mixed model fit by maximum likelihood (Laplace
## Approximation) [glmerMod]
## Family: binomial ( logit )
## Formula: correct ~ task + pas + (1 | subject)
## Data: samples
##
##      AIC      BIC  logLik deviance df.resid
## 17424.9 17479.5 -8705.5 17410.9     18124
##
## Scaled residuals:
##    Min     1Q Median     3Q    Max
## -8.4872 -0.6225  0.3240  0.5767  1.6144
##
## Random effects:
##   Groups   Name        Variance Std.Dev.
##   subject (Intercept) 0.1979   0.4449
## Number of obs: 18131, groups: subject, 29
##
## Fixed effects:
##             Estimate Std. Error z value Pr(>|z|)
## (Intercept)  3.21262  0.11772 27.290 <2e-16 ***
## taskquadruplet -0.03055  0.04497 -0.679  0.497
## tasksingles   -0.01059  0.04687 -0.226  0.821
## pas3          -1.15022  0.09368 -12.279 <2e-16 ***
## pas2          -2.17254  0.08623 -25.195 <2e-16 ***
## pas1          -3.12732  0.08628 -36.248 <2e-16 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##            (Intr) tskqdr tsksng pas3   pas2
## taskquadruplet -0.190
## tasksingles   -0.233  0.489
## pas3          -0.557  0.004  0.043
## pas2          -0.625 -0.006  0.059  0.764
## pas1          -0.637 -0.017  0.079  0.764  0.869

```

```

# better
piecewiseSEM:::rsquared(m9)

```

```

## Response family link method Marginal Conditional
## 1 correct binomial logit delta 0.2001362  0.2289711

```

## Ex3.p3.ii

When we add PAS as a main effect on top of task, we can see that PAS actually has a significant effect on performance, whereas now task no longer displays a significant effect. This means that by including PAS as a main effect in the model, it altered the significance of task on performance.

```

m10 <- glmer(correct ~ pas +
  (1 | subject),
  data = samples,
  family = binomial(link = "logit"))
summary(m10)

```

```

## Generalized linear mixed model fit by maximum likelihood (Laplace
## Approximation) [glmerMod]
## Family: binomial ( logit )
## Formula: correct ~ pas + (1 | subject)
## Data: samples
##
##      AIC      BIC  logLik deviance df.resid
## 17421.4 17460.4 -8705.7 17411.4     18126
##
## Scaled residuals:
##    Min     1Q Median     3Q    Max
## -8.5665 -0.6243  0.3244  0.5754  1.6017
##
## Random effects:
## Groups   Name        Variance Std.Dev.
## subject (Intercept) 0.1981   0.4451
## Number of obs: 18131, groups: subject, 29
##
## Fixed effects:
##             Estimate Std. Error z value Pr(>|z|)
## (Intercept) 3.19985  0.11404  28.06  <2e-16 ***
## pas3       -1.15049  0.09358 -12.29  <2e-16 ***
## pas2       -2.17365  0.08601 -25.27  <2e-16 ***
## pas1       -3.12940  0.08582 -36.46  <2e-16 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
## (Intr) pas3   pas2
## pas3 -0.567
## pas2 -0.637  0.764
## pas1 -0.647  0.764  0.868

```

```

# task makes negligible diff
piecewiseSEM:::rsquared(m10)

```

```

## Response family link method Marginal Conditional
## 1 correct binomial logit delta 0.2001568  0.2290084

```

```

m11 <- glm(correct ~ pas + task +
  pas:task,
  data = samples,
  family = binomial(link = "logit"))
summary(m11)

```

```

## 
## Call:
## glm(formula = correct ~ pas + task + pas:task, family = binomial(link = "logit"),
##      data = samples)
##
## Deviance Residuals:
##    Min      1Q  Median      3Q     Max
## -2.5349 -1.2539  0.5089  0.7922  1.1028
##
## Coefficients:
##                               Estimate Std. Error z value Pr(>|z|)
## (Intercept)                2.97483   0.13461 22.100 < 2e-16 ***
## pas3                     -1.04172   0.16112 -6.465 1.01e-10 ***
## pas2                     -1.97686   0.14502 -13.631 < 2e-16 ***
## pas1                     -2.76473   0.14218 -19.445 < 2e-16 ***
## taskquadruplet            -0.12316   0.18907 -0.651   0.515
## tasksingles               0.19699   0.18347  1.074   0.283
## pas3:taskquadruplet       0.01682   0.22714  0.074   0.941
## pas2:taskquadruplet       0.07579   0.20373  0.372   0.710
## pas1:taskquadruplet       0.12540   0.19928  0.629   0.529
## pas3:tasksingles          -0.15153   0.22253 -0.681   0.496
## pas2:tasksingles          -0.15565   0.19913 -0.782   0.434
## pas1:tasksingles          -0.22907   0.19593 -1.169   0.242
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
## Null deviance: 20418  on 18130  degrees of freedom
## Residual deviance: 17853  on 18119  degrees of freedom
## AIC: 17877
##
## Number of Fisher Scoring iterations: 5

```

```
piecewiseSEM:::rsquared(m11)
```

```

## Response  family link      method R.squared
## 1 correct  binomial logit nagelkerke      NaN

```

```

m12 <- glmer(correct ~ pas + task +
  pas:task + (1 | subject),
  data = samples,
  family = binomial(link = "logit"))

```

```

## Warning in checkConv(attr(opt, "derivs"), opt$par, ctrl = control$checkConv, :
## Model failed to converge with max|grad| = 0.0036472 (tol = 0.002, component 1)

```

```
summary(m12)
```

```

## Generalized linear mixed model fit by maximum likelihood (Laplace
## Approximation) [glmerMod]
## Family: binomial ( logit )
## Formula: correct ~ pas + task + pas:task + (1 | subject)
## Data: samples
##
##      AIC      BIC  logLik deviance df.resid
## 17431.0 17532.4 -8702.5 17405.0     18118
##
## Scaled residuals:
##    Min     1Q Median     3Q    Max
## -7.9339 -0.6276  0.3186  0.5750  1.6411
##
## Random effects:
##   Groups   Name        Variance Std.Dev.
##   subject (Intercept) 0.1987   0.4458
## Number of obs: 18131, groups: subject, 29
##
## Fixed effects:
##             Estimate Std. Error z value Pr(>|z|)
## (Intercept)  3.13206  0.15977 19.604 < 2e-16 ***
## pas3       -1.04839  0.16303 -6.430 1.27e-10 ***
## pas2       -2.08499  0.14809 -14.080 < 2e-16 ***
## pas1       -3.04828  0.14599 -20.881 < 2e-16 ***
## taskquadruplet -0.08775  0.18953 -0.463   0.643
## tasksingles   0.23912  0.18410  1.299   0.194
## pas3:taskquadruplet -0.03995  0.22823 -0.175   0.861
## pas2:taskquadruplet  0.04490  0.20484  0.219   0.827
## pas1:taskquadruplet  0.09419  0.20023  0.470   0.638
## pas3:tasksingles   -0.21805  0.22376 -0.974   0.330
## pas2:tasksingles   -0.25772  0.20058 -1.285   0.199
## pas1:tasksingles   -0.29812  0.19720 -1.512   0.131
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##            (Intr) pas3   pas2   pas1   tskqdr tksng ps3:tskq ps2:tskq
## pas3      -0.712
## pas2      -0.792  0.773
## pas1      -0.808  0.783  0.874
## taskquadruplet -0.601  0.590  0.650  0.658
## tasksingles -0.619  0.608  0.670  0.678  0.522
## ps3:tskqdrp  0.499 -0.705 -0.540 -0.547 -0.831 -0.433
## ps2:tskqdrp  0.557 -0.546 -0.702 -0.610 -0.926 -0.483  0.769
## ps1:tskqdrp  0.569 -0.559 -0.616 -0.699 -0.947 -0.494  0.786   0.876
## ps3:tsksngl  0.507 -0.718 -0.548 -0.555 -0.429 -0.823  0.512   0.397
## ps2:tsksngl  0.566 -0.558 -0.715 -0.618 -0.479 -0.919  0.398   0.517
## ps1:tsksngl  0.576 -0.567 -0.623 -0.707 -0.487 -0.934  0.404   0.451
##            ps1:tskq ps3:tsks ps2:tsks
## pas3
## pas2
## pas1
## taskquadruplet
## tasksingles
## ps3:tskqdrp
## ps2:tskqdrp
## ps1:tskqdrp

```

```
## ps3:tsksngl 0.406
## ps2:tsksngl 0.454    0.756
## ps1:tsksngl 0.517    0.768    0.858
## optimizer (Nelder_Mead) convergence code: 0 (OK)
## Model failed to converge with max|grad| = 0.0036472 (tol = 0.002, component 1)
```

```
piecewiseSEM:::rsquared(m12)
```

```
## Response family link method Marginal Conditional
## 1 correct binomial logit delta 0.201629  0.2305216
```

```
AIC(m8, m9, m10, m12)
```

```
##      df      AIC
## m8     4 19927.21
## m9     7 17424.91
## m10   5 17421.39
## m12  13 17430.97
```

Ex3.p3.v Describe in your words which model is the best in explaining the variance in accuracy?

Of the above models, m12 is the best at explaining the variance in accuracy. But we think that model 10 is a better model to use because the difference in explained variance is negligible and it is a simpler model which means it is less prone to overfitting. Adding the random intercepts per subject allows for differences in subjects ability to correctly classify the stimuli. These differences in subjects may come from situational stress. Model m10 also has the lowest AIC score (17421.39), even though it has slightly more residual deviance than model 12, but this makes sense as more complex models are punished in this area.

# Portfolio Assignment 2, part 2, Methods 3, 2021, autumn semester

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2021-10-27

## EXERCISE 4 - Download and organise the data from experiment 1

Go to <https://osf.io/ecxsj/files/> (<https://osf.io/ecxsj/files/>) and download the files associated with Experiment 1 (there should be 29).

The data is associated with Experiment 1 of the article at the following DOI

<https://doi.org/10.1016/j.concog.2019.03.007> (<https://doi.org/10.1016/j.concog.2019.03.007>)

1. Put the data from all subjects into a single data frame - note that some of the subjects do not have the *seed* variable. For these subjects, add this variable and make it *NA* for all observations. (The *seed* variable will not be part of the analysis and is not an experimental variable)
  - i. Factorise the variables that need factorising
  - ii. Remove the practice trials from the dataset (see the *trial.type* variable)
  - iii. Create a *correct* variable
  - . iv. Describe how the *target.contrast* and *target.frames* variables differ compared to the data from part 1 of this assignment

## Answers

### Exercise 4, part 1 - EH

```
#loading in the data
dat <- list.files(path = "./experiment_1", full.names = TRUE)

#Changing all the variables classes to the ones we want them to be
samples <- map_df(dat, read_csv,
  trim_ws = TRUE, na = c("", "NA"), # i
  col_types = cols(
    trial.type = col_factor(),
    pas = col_factor(),
    trial = col_factor(),
    jitter.x = col_double(),
    jitter.y = col_double(),
    odd.digit = col_integer(),
    target.contrast = col_double(),
    target.frames = col_double(),
    cue = col_factor(),
    task = col_factor(),
    target.type = col_factor(),
    rt.subj = col_double(),
    rt.obj = col_double(),
    even.digit = col_integer(),
    seed = col_double(),
    obj.resp = col_factor(),
    subject = col_factor()
  )
)
```

```
## Warning: The following named parsers don't match the column names: seed
## Warning: The following named parsers don't match the column names: seed
## Warning: The following named parsers don't match the column names: seed
```

```
# removing all the practice trials #ii
exp1 <- samples[samples$trial.type == "experiment", ]
exp1 <- mutate(exp1, # iii
  correct = as.logical(
    ifelse(substr(target.type, 1, 1) == obj.resp, 1, 0)
  )
)
```

- iv. In the previous experiment, the number of target frames was fixed at 3, whereas this time it the target frames was not fixed (they were integer numbers from 1-6), and in the last experiment where *target.contrast* was not fixed but for this experiment *target.contrast* was fixed at 0.1

# EXERCISE 5 - Use log-likelihood ratio tests to evaluate logistic regression models

1. Do logistic regression - *correct* as the dependent variable and *target.frames* as the independent variable. (Make sure that you understand what *target.frames* encode). Create two models - a pooled model and a partial-pooling model. The partial-pooling model should include a subject-specific intercept.

i. the likelihood-function for logistic regression is:  $L(p) = \prod_{i=1}^N p^{y_i} (1 - p)^{(1-y_i)}$  (Remember the probability mass function for the Bernoulli Distribution).

Create a function that calculates the likelihood.

ii. the log-likelihood-function for logistic regression is:  $l(p) = \sum_{i=1}^N [y_i \ln p + (1 - y_i) \ln (1 - p)]$ .

Create a function that calculates the log-likelihood

iii. apply both functions to the pooling model you just created. Make sure that the log-likelihood matches what is returned from the *logLik* function for the pooled model. Does the likelihood-function return a value that is surprising? Why is the log-likelihood preferable when working with computers with limited precision?

iv. now show that the log-likelihood is a little off when applied to the partial pooling model - (the likelihood function is different for the multilevel function - see section 2.1 of [https://www.researchgate.net/profile/Douglas-Bates/publication/2753537\\_Computational\\_Methods\\_for\\_Multilevel\\_Modelling/links/00b4953b4108d73427000000-Methods-for-Multilevel-Modelling.pdf](https://www.researchgate.net/profile/Douglas-Bates/publication/2753537_Computational_Methods_for_Multilevel_Modelling/links/00b4953b4108d73427000000-Methods-for-Multilevel-Modelling.pdf) if you are interested)

2. Use log-likelihood ratio tests to argue for the addition of predictor variables, start from the null model, `glm(correct ~ 1, 'binomial', data)`, then add subject-level intercepts, then add a group-level effect of *target.frames* and finally add subject-level slopes for *target.frames*. Also assess whether or not a correlation between the subject-level slopes and the subject-level intercepts should be included.

i. write a short methods section and a results section where you indicate which model you chose and the statistics relevant for that choice. Include a plot of the estimated group-level function with `xlim=c(0, 8)` that includes the estimated subject-specific functions.

ii. also include in the results section whether the fit didn't look good for any of the subjects. If so, identify those subjects in the report, and judge (no statistical test) whether their performance (accuracy) differed from that of the other subjects. Was their performance better than chance? (Use a statistical test this time) (50 %)

3. Now add *pas* to the group-level effects - if a log-likelihood ratio test justifies this, also add the interaction between *pas* and *target.frames* and check whether a log-likelihood ratio test justifies this

i. if your model doesn't converge, try a different optimizer

ii. plot the estimated group-level functions over `xlim=c(0, 8)` for each of the four PAS-ratings - add this plot to your report (see: 5.2.i) and add a description of your chosen model. Describe how *pas* affects accuracy together with target duration if at all. Also comment on the estimated functions' behaviour at *target.frame=0* - is that behaviour reasonable?

## Answers

### Exercise 5, part 1 - KV

```
#creating a pooled model
m1 <- glm(correct ~ target.frames, data = exp1,
           family = binomial(link = "logit"))
summary(m1)
```

```

## 
## Call:
##   glm(formula = correct ~ target.frames, family = binomial(link = "logit"),
##       data = exp1)
##
## Deviance Residuals:
##   Min      1Q  Median      3Q     Max
## -2.6452  0.2478  0.3546  0.7039  1.2621
##
## Coefficients:
##             Estimate Std. Error z value Pr(>|z|)
## (Intercept) -0.92992    0.03485 -26.69 <2e-16 ***
## target.frames  0.73296    0.01219   60.11 <2e-16 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
## Null deviance: 26683 on 25043 degrees of freedom
## Residual deviance: 21730 on 25042 degrees of freedom
## AIC: 21734
##
## Number of Fisher Scoring iterations: 5

```

```

#creating a partial-pooled model
m2 <- glmer(correct ~ target.frames + (1 | subject),
             data = exp1, family = binomial(link = "logit"))
summary(m2)

```

```

## Generalized linear mixed model fit by maximum likelihood (Laplace
## Approximation) [glmerMod]
## Family: binomial ( logit )
## Formula: correct ~ target.frames + (1 | subject)
## Data: exp1
##
##      AIC      BIC  logLik deviance df.resid
## 21250.1 21274.4 -10622.0 21244.1   25041
##
## Scaled residuals:
##   Min      1Q  Median      3Q     Max
## -7.7520  0.1436  0.2604  0.4816  2.0730
##
## Random effects:
##   Groups Name        Variance Std.Dev.
##   subject (Intercept) 0.1549   0.3936
##   Number of obs: 25044, groups: subject, 29
##
## Fixed effects:
##             Estimate Std. Error z value Pr(>|z|)
## (Intercept) -0.95968    0.08125 -11.81 <2e-16 ***
## target.frames  0.75493    0.01251   60.37 <2e-16 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##          (Intr)
## target.frms -0.382

```

```

#5.i creating our own likelihood function
likelihood <- function(p, y) {
  prod((p^y) * (1 - p)^(1 - y))
}

#5.ii creating our own log-likelihood function
log_likelihood <- function(p, y) {
  sum(y * log(p) + (1 - y) * log(1 - p))
}

#5.iii Getting some likelihood and log-likelihood values and double-checking that our function spits out the right
#t values for m1
p <- fitted.values(m1)
likelihood(p, exp1$correct)

```

```

## [1] 0

```

```

log_likelihood(p, exp1$correct)

```

```

## [1] -10865.25

```

```

(m1_ll <- logLik(m1))

```

```

## 'log Lik.' -10865.25 (df=2)

# 5.iv Showing that the log-likelihood is slightly different when using _LogLik_ in comparison to our own function for m2
p <- fitted.values(m2)
likelihood(p, exp1$correct)

## [1] 0

log_likelihood(p, exp1$correct)

## [1] -10565.53

(m2_ll <- logLik(m2))

## 'log Lik.' -10622.03 (df=3)

```

5.iii a likelihood value of 0 is returned for both m1 and m2, which is due to the lack of precision of the computer. This is why it is preferable to use the log-likelihood function instead.

5.iv The log-likelihood value returned from our function is -10565.53 whereas the value for the *LogLik* function is 10622.03, which can probably be explained by the fact that our function does not take multilevel modelling into consideration.

## Exercise 5, part 2 - LR

```

#creating the null-model
m0 <- glm(correct ~ 1, data = exp1, family = binomial(link = "logit"))

#adding subject-level intercepts
m3 <- glmer(correct ~ 1 + (1 | subject),
             data = exp1, family = binomial(link = "logit"))

#adding subject-level intercepts and slopes
m4 <- glmer(correct ~ target.frames + (target.frames | subject),
             data = exp1, family = binomial(link = "logit"))

#checking the log-likelihood of the different models
(m0_ll <- logLik(m0))

## 'log Lik.' -13341.54 (df=1)

(m1_ll <- logLik(m1))

## 'log Lik.' -10865.25 (df=2)

(m2_ll <- logLik(m2))

## 'log Lik.' -10622.03 (df=3)

(m3_ll <- logLik(m3))

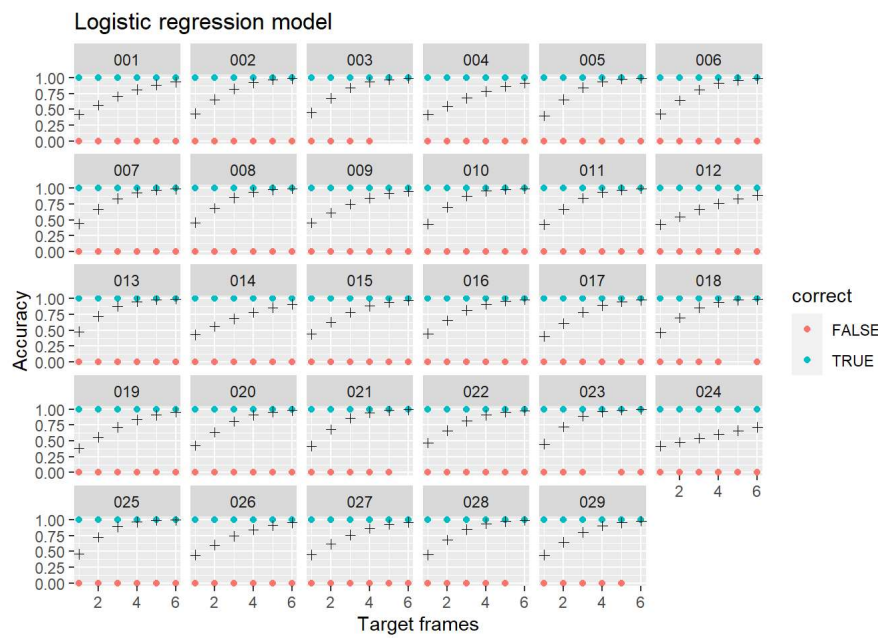
## 'log Lik.' -13157.55 (df=2)

(m4_ll <- logLik(m4))

## 'log Lik.' -10448.83 (df=5)

#creating some great plots to investigate the data
ggplot(exp1, aes(target.frames, as.numeric(correct), color = correct)) +
  geom_point() +
  geom_point(aes(target.frames, fitted.values(m4)),
             shape = 3, color = "black", inherit.aes = FALSE) +
  facet_wrap(~subject) +
  labs(
    x = "Target frames",
    y = "Accuracy",
    title = "Logistic regression model")

```



```
#zooming in on subject 024
(subj24_accuracy <- exp1 %>%
  filter(subject == "024") %>%
  summarise(accuracy = sum(correct) / n()))
```

```
## # A tibble: 1 × 1
##   accuracy
##     <dbl>
## 1 0.568
```

2.i+ii We have run binomial regression models, in which we tested the relationship between the variable *correct* predicted by *target.frames*. Different models using fixed and random effects were created for comparison. The quality of the models Was compared by using the log-likelihood function.

Out of the 5 models, we choose m4 as our best model, as it has the highest LogLik value of -10676.02, with 5 degrees of freedom. After observing the plots, we have identified subject number 024 as having a bad fit. As the fit looks more linear than sigmoid, and has values all along the axis x = 0. Judging from this graph we predict that their performance is different to the other participants possibly having a more equal ratio of correct and incorrect, as they are the only subject that had a graph that resulted in looking more linear. We then took subject 024 and took the number of correct divided by trials to retrieve their accuracy score and compare it to chance, 50%. They have a accuracy percentage of 56.9%, which is only slightly higher than chance, which could explain why their graph looks poor.

### Exercise 5, part 3 - VK

```
#Creating some models
m5 <- glmer(correct ~ target.frames + pas + (target.frames | subject),
  data = exp1, family = binomial(link = "logit"))
(m5_ll <- logLik(m5))
```

```
## 'log Lik.' -9931.828 (df=8)
```

```
m6 <- glmer(correct ~ target.frames + pas +
  (pas * target.frames) + (target.frames | subject),
  data = exp1, family = binomial(link = "logit"), control = glmerControl(optimizer = "bobyqa"))
(m6_ll <- logLik(m6))
```

```
## 'log Lik.' -9742.039 (df=11)
```

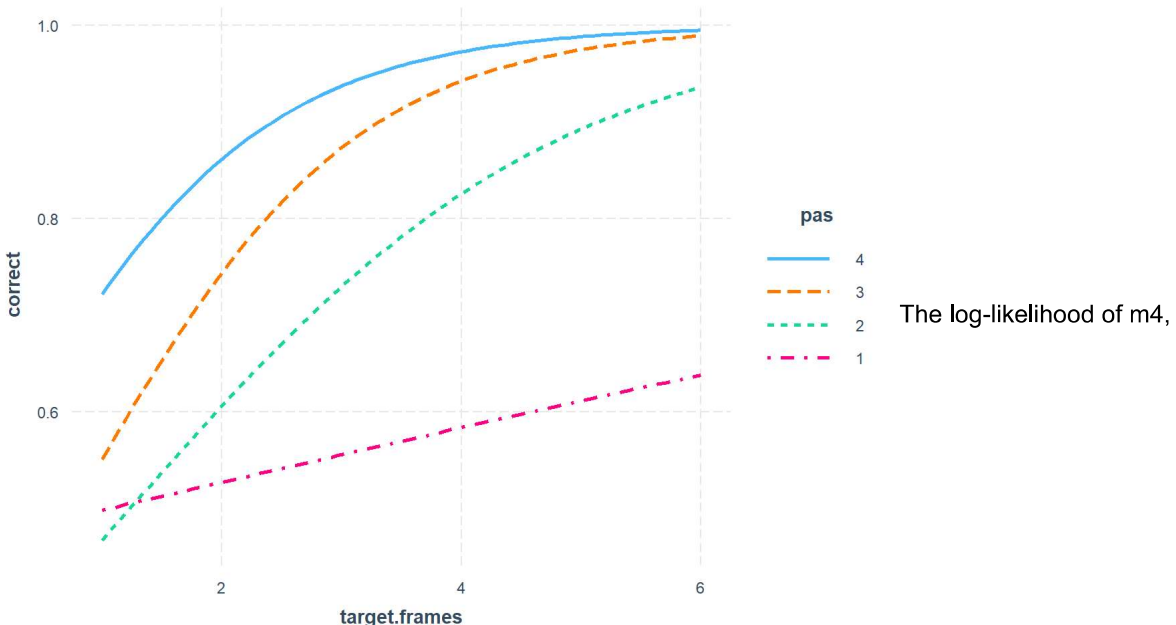
```
summary(m6)
```

```

## Generalized linear mixed model fit by maximum likelihood (Laplace
## Approximation) [glmerMod]
## Family: binomial ( logit )
## Formula:
## correct ~ target.frames + pas + (pas * target.frames) + (target.frames |
##     subject)
## Data: exp1
## Control: glmerControl(optimizer = "bobyqa")
##
##      AIC      BIC  logLik deviance df.resid
## 19506.1 19595.5 -9742.0 19484.1    25033
##
## Scaled residuals:
##    Min     1Q  Median     3Q    Max 
## -19.0101  0.0537  0.1606  0.4849  1.4465
##
## Random effects:
## Groups   Name        Variance Std.Dev. Corr
## subject (Intercept) 0.03698  0.1923
##         target.frames 0.02057  0.1434  -0.76
## Number of obs: 25044, groups: subject, 29
##
## Fixed effects:
##             Estimate Std. Error z value Pr(>|z|)    
## (Intercept) 0.08009  0.24320  0.329  0.74193    
## target.frames 0.87404  0.06792 12.869 < 2e-16 ***  
## pas3       -0.74022  0.27065 -2.735  0.00624 **  
## pas2       -0.77311  0.25133 -3.076  0.00210 **  
## pas1       -0.20172  0.24613 -0.820  0.41246    
## target.frames:pas3 -0.01055  0.07335 -0.144  0.88565    
## target.frames:pas2 -0.31206  0.06850 -4.555 5.23e-06 *** 
## target.frames:pas1 -0.75924  0.06749 -11.250 < 2e-16 *** 
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##          (Intr) trgt.f pas3   pas2   pas1   trg.:3 trg.:2
## target.frms -0.896
## pas3        -0.869  0.759
## pas2        -0.945  0.826  0.847
## pas1        -0.966  0.842  0.859  0.936
## trgt.frms:3  0.775 -0.776 -0.926 -0.761 -0.769
## trgt.frms:2  0.841 -0.841 -0.762 -0.921 -0.839  0.790
## trgt.frms:1  0.853 -0.850 -0.767 -0.838 -0.916  0.791  0.870

```

```
interactions::interact_plot(model = m6, pred = "target.frames", modx = "pas")
```



m5 and m6 are these:

$m4\_ll = -10676.02$ ,  $df=(5)$   $m5\_ll = -9931.828$ ,  $df=(8)$   $m6\_ll = -9742.039$ ,  $df=(11)$

Thereby justifying the choice of model 6 as our preferred model

5.3 ii The chosen model, m6, is a binomial model in which the variable *correct* is predicted by *target.frames*, *PAS* and their interaction.

The estimate of *target.frames* shows that an increase in this variable “generally” increases accuracy - however, let's look at the interactions\_

We use this interaction between PAS and target duration because the lesser amount of frames (target duration) the less confident participants are (PAS).

Together, PAS with time duration (target.frames), the correctness gets worse as target.frames:PAS decreases (from PAS\_4 to PAS\_1). It can be read from the model summary that *target.frames* influences correctness less with lower PAS level (e.g. target.frames:PAS\_1 = -0.75924 etc.)

The intercept of 0.08009, does sound strange as one might incorrectly expect it to be around 50%, but naturally this is because it is not possible to show a figure at a duration of 0 frames <3 The intercept of 0.08 is thus only meaningful within the model but not in reality.

## EXERCISE 6 - Test linear hypotheses

In this section we are going to test different hypotheses. We assume that we have already proved that more objective evidence (longer duration of stimuli) is sufficient to increase accuracy in and of itself and that more subjective evidence (higher PAS ratings) is also sufficient to increase accuracy in and of itself.

We want to test a hypothesis for each of the three neighbouring differences in PAS, i.e. the difference between 2 and 1, the difference between 3 and 2 and the difference between 4 and 3. More specifically, we want to test the hypothesis that accuracy increases faster with objective evidence if subjective evidence is higher at the same time, i.e. we want to test for an interaction.

1. Fit a model based on the following formula:

```
correct ~ pas * target.frames + (target.frames | subject))
```

i. First, use `summary` (yes, you are allowed to!) to argue that accuracy increases faster with objective evidence for PAS 2 than for PAS 1.

2. `summary` won't allow you to test whether accuracy increases faster with objective evidence for PAS 3 than for PAS 2 (unless you use `relevel`, which you are not allowed to in this exercise). Instead, we'll be using the function `glht` from the `multcomp` package

i. To redo the test in 6.1.i, you can create a *contrast* vector. This vector will have the length of the number of estimated group-level effects and any specific contrast you can think of can be specified using this. For redoing the test from 6.1.i, the code snippet below will do

ii. Now test the hypothesis that accuracy increases faster with objective evidence for PAS 3 than for PAS 2.

iii. Also test the hypothesis that accuracy increases faster with objective evidence for PAS 4 than for PAS 3

3. Finally, test that whether the difference between PAS 2 and 1 (tested in 6.1.i) is greater than the difference between PAS 4 and 3 (tested in 6.2.iii)

## Answers

### Exercise 6, part 1 - EH

```
#creating the desired model
m7 <- glmer(correct ~ pas * target.frames +
  (target.frames | subject), data = exp1,
  family = binomial(link = "logit"),
  control = glmerControl(optimizer = "bobyqa"))
#getting that summary
summary(m7)
```

```

## Generalized linear mixed model fit by maximum likelihood (Laplace
## Approximation) [glmerMod]
## Family: binomial ( logit )
## Formula: correct ~ pas * target.frames + (target.frames | subject)
## Data: exp1
## Control: glmerControl(optimizer = "bobyqa")
##
##      AIC      BIC  logLik deviance df.resid
## 19506.1 19595.5 -9742.0 19484.1    25033
##
## Scaled residuals:
##      Min     1Q  Median     3Q    Max 
## -19.0102  0.0537  0.1606  0.4849  1.4465
##
## Random effects:
## Groups   Name        Variance Std.Dev. Corr
## subject (Intercept) 0.03698  0.1923
##          target.frames 0.02057  0.1434  -0.76
## Number of obs: 25044, groups: subject, 29
##
## Fixed effects:
##             Estimate Std. Error z value Pr(>|z|)    
## (Intercept)  0.08010  0.24575  0.326  0.74446  
## pas3       -0.74025  0.27355 -2.706  0.00681 ** 
## pas2       -0.77313  0.25370 -3.047  0.00231 ** 
## pas1       -0.20174  0.24871 -0.811  0.41729  
## target.frames 0.87404  0.06850 12.760 < 2e-16 *** 
## pas3:target.frames -0.01054  0.07403 -0.142  0.88678  
## pas2:target.frames -0.31205  0.06904 -4.520 6.19e-06 *** 
## pas1:target.frames -0.75924  0.06809 -11.151 < 2e-16 *** 
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##            (Intr) pas3  pas2  pas1  trgt.f ps3:t. ps2:t.
## pas3      -0.871
## pas2      -0.946  0.850
## pas1      -0.966  0.862  0.937
## target.frms -0.898  0.763  0.829  0.845
## ps3:trgt.fr  0.780 -0.927 -0.766 -0.773 -0.780
## ps2:trgt.fr  0.843 -0.766 -0.923 -0.841 -0.844  0.794
## ps1:trgt.fr  0.856 -0.771 -0.841 -0.917 -0.853  0.794  0.872

```

6.1.i we see that the interaction-effect between *pas* and *target.frames* is clearly higher for PAS\_2 vs PAS\_1, thus indicating that accuracy increases faster with objective evidence for PAS\_2 than for PAS\_1.

## Exercise 6, part 2 - KV

```

#trying out glht on m7
glht(m7)

```

```

## General Linear Hypotheses
##
## Linear Hypotheses:
##             Estimate
## (Intercept) == 0    0.08010
## pas3 == 0       -0.74025
## pas2 == 0       -0.77313
## pas1 == 0       -0.20174
## target.frames == 0 0.87404
## pas3:target.frames == 0 -0.01054
## pas2:target.frames == 0 -0.31205
## pas1:target.frames == 0 -0.75924

```

## Snippet for 6.2.i+ii+iii

```

## testing whether PAS 2 is different from PAS 1
contrast.vector <- matrix(c(0, 0, -1, 1, 0, 0, 0, 0), nrow = 1)
gh <- glht(m7, contrast.vector)
print(summary(gh))
## as another example, we could also test whether there is a difference in
## intercepts between PAS 2 and PAS 3
contrast.vector <- matrix(c(0, -1, 1, 0, 0, 0, 0, 0), nrow = 1)
gh <- glht(m7, contrast.vector)
print(summary(gh))

## PAS 4 and 3
contrast.vector <- matrix(c(-1, 1, 0, 0, 0, 0, 0, 0), nrow = 1)
gh <- glht(m7, contrast.vector)
print(summary(gh))

```

## Exercise 6, part 3 - LR

```
#creating the contrast_vector
K <- rbind(
  c(0, 0, -1, 1),
  c(-1, 1, 0, 0)
)

gh <- glht(m7, mcp(pas = K))
print(summary(gh))
```

6.3.i - as seen in the output there is a greater difference between PAS 2 and 1 than between 4 and 3.

## EXERCISE 7 - Estimate psychometric functions for the Perceptual Awareness Scale and evaluate them

We saw in 5.3 that the estimated functions went below chance at a target duration of 0 frames (0 ms). This does not seem reasonable, so we will be trying a different approach for fitting here.

We will fit the following function that results in a sigmoid,  $f(x) = a + \frac{b-a}{1+e^{\frac{c-x}{d}}}$

It has four parameters:  $a$ , which can be interpreted as the minimum accuracy level,  $b$ , which can be interpreted as the maximum accuracy level,  $c$ , which can be interpreted as the so-called inflection point, i.e. where the derivative of the sigmoid reaches its maximum and  $d$ , which can be interpreted as the steepness at the inflection point. (When  $d$  goes towards infinity, the slope goes towards a straight line, and when it goes towards 0, the slope goes towards a step function).

We can define a function of a residual sum of squares as below

```
sigfit <- function(a, b, c, d, x) {
  a + (b - a) / (1 + exp((c - x) / d))
}

RSS <- function(dataset, par) {
  x <- dataset$x
  y <- dataset$y
  y.hat <- sigfit(par[1], par[2], par[3], par[4], x)
  RSS <- sum((y - y.hat)^2)
  return(RSS)
}
```

1. Now, we will fit the sigmoid for the four PAS ratings for Subject 7

- i. use the function `optim`. It returns a list that among other things contains the four estimated parameters. You should set the following arguments:
    - par : you can set  $c$  and  $d$  as 1. Find good choices for  $a$  and  $b$  yourself (and argue why they are appropriate)
    - fn : which function to minimise?
    - data : the data frame with  $x$ , `target.frames`, and  $y$ , `correct` in it
    - method : 'L-BFGS-B'
    - lower : lower bounds for the four parameters, (the lowest value they can take), you can set  $c$  and  $d$  as  $-\text{Inf}$ . Find good choices for  $a$  and  $b$  yourself (and argue why they are appropriate)
    - upper : upper bounds for the four parameters, (the highest value they can take) can set  $c$  and  $d$  as  $\text{Inf}$ . Find good choices for  $a$  and  $b$  yourself (and argue why they are appropriate)
  - ii. Plot the fits for the PAS ratings on a single plot (for subject 7) `xlim=c(0, 8)`
  - iii. Create a similar plot for the PAS ratings on a single plot (for subject 7), but this time based on the model from 6.1 `xlim=c(0, 8)`
  - iv. Comment on the differences between the fits - mention some advantages and disadvantages of each way
2. Finally, estimate the parameters for all subjects and each of their four PAS ratings. Then plot the estimated function at the group-level by taking the mean for each of the four parameters,  $a$ ,  $b$ ,  $c$  and  $d$  across subjects. A function should be estimated for each PAS-rating (it should look somewhat similar to Fig. 3 from the article: <https://doi.org/10.1016/j.concog.2019.03.007> (<https://doi.org/10.1016/j.concog.2019.03.007>))
- i. compare with the figure you made in 5.3.ii and comment on the differences between the fits - mention some advantages and disadvantages of both.

## Answers

## Exercise 7, part 1 - VK

```

#filtering out subject 7
subj7 <- exp1[exp1$subject == "007", ]

#choosing our desired variables
subj7xy <- data.frame(
  x = subj7$target.frames,
  y = subj7$correct,
  pas = subj7$pas)

#Setting up the function and using it on the four pas_levels

#PAS 1
s7p1 <- optim(
  par = c(0.5, 1, 1, 1), # we set a to be 0.5 as we expect chance level to be the appropriate minimum accuracy
  fn = RSS,
  data = subj7xy[subj7xy$pas == "1", ],
  method = "L-BFGS-B",
  lower = c(0.5, 0.5, -Inf, -Inf),
  upper = c(1, 1, Inf, Inf)
)
p1fit <- sigfit(
  a = s7p1$par[1],
  b = s7p1$par[2],
  c = s7p1$par[3],
  d = s7p1$par[4],
  x = subj7xy[subj7xy$pas == "1", ]$x
)

#PAS 2
s7p2 <- optim(
  par = c(0.5, 1, 1, 1),
  fn = RSS,
  data = subj7xy[subj7xy$pas == "2", ],
  method = "L-BFGS-B",
  lower = c(0.5, 0.5, -Inf, -Inf),
  upper = c(1, 1, Inf, Inf)
)
p2fit <- sigfit(
  a = s7p2$par[1],
  b = s7p2$par[2],
  c = s7p2$par[3],
  d = s7p2$par[4],
  x = subj7xy[subj7xy$pas == "2", ]$x
)

#PAS 3
s7p3 <- optim(
  par = c(0.5, 1, 1, 1),
  fn = RSS,
  data = subj7xy[subj7xy$pas == "3", ],
  method = "L-BFGS-B",
  lower = c(0.5, 0.5, -Inf, -Inf),
  upper = c(1, 1, Inf, Inf)
)
p3fit <- sigfit(
  a = s7p3$par[1],
  b = s7p3$par[2],
  c = s7p3$par[3],
  d = s7p3$par[4],
  x = subj7xy[subj7xy$pas == "3", ]$x
)

#PAS 4
s7p4 <- optim(
  par = c(0.5, 1, 1, 1),
  fn = RSS,
  data = subj7xy[subj7xy$pas == "4", ],
  method = "L-BFGS-B",
  lower = c(0.5, 0.5, -Inf, -Inf),
  upper = c(1, 1, Inf, Inf)
)
p4fit <- sigfit(
  a = s7p4$par[1],
  b = s7p4$par[2],
  c = s7p4$par[3],
  d = s7p4$par[4],
  x = subj7xy[subj7xy$pas == "4", ]$x
)

length(subj7xy[subj7xy$pas == "1", ]$x)

```

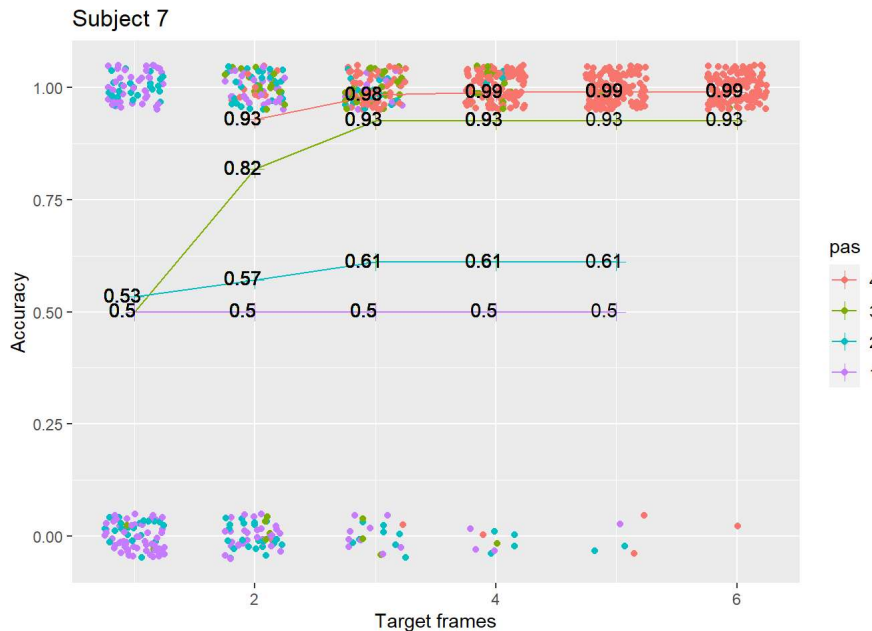
```
## [1] 183
```

```
length(p1fit)
```

```
## [1] 183
```

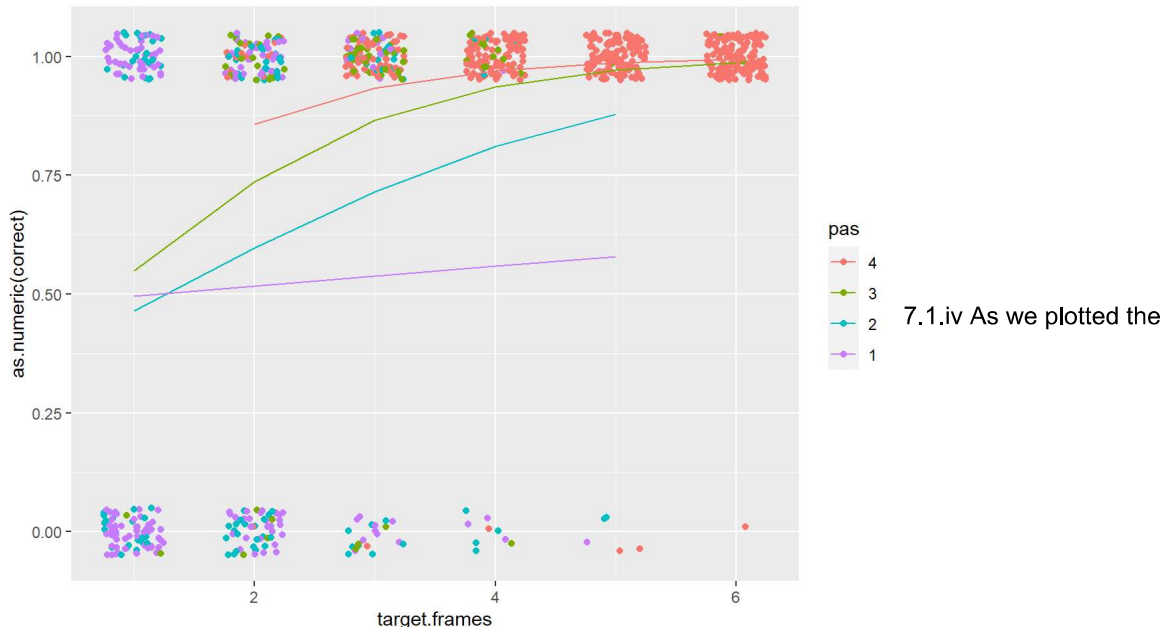
```
subj7xy$yhat <- NA
subj7xy[subj7xy$pas == "1", ]$yhat <- p1fit
subj7xy[subj7xy$pas == "2", ]$yhat <- p2fit
subj7xy[subj7xy$pas == "3", ]$yhat <- p3fit
subj7xy[subj7xy$pas == "4", ]$yhat <- p4fit

#plotting the fits of the sigmoid
ggplot(aes(x = x, y = as.numeric(y), color = pas), data = subj7xy) +
  geom_jitter(width = 0.25, height = 0.05) +
  geom_line(aes(y = yhat)) +
  geom_point(aes(y = yhat), shape = 3, size = 3) +
  geom_text(aes(y = yhat, label = round(yhat, 2)),
            color = "black", nudge_x = -0.1, nudge_y = 0.005
  ) +
  labs(x = "Target frames", y = "Accuracy", title = "Subject 7")
```



```
s7m7fit <- predict(m7, newdata = subj7, type = "response")
```

```
#plotting m7
ggplot(aes(x = target.frames, y = as.numeric(correct), color = pas),
       data = subj7) +
  geom_jitter(width = 0.25, height = 0.05) +
  geom_line(aes(y = s7m7fit))
```



predicted values of the fits, the intercepts at *target.frames*=0 is not shown, though this would have been a crucial difference between the plots. The visible difference between the plots must primarily stem from the fact that we manually set the lowest possible value to 0.5 in our optim function. The advantage of manually creating the function is that we can control for the fact that below chance level accuracy is highly unlikely and thus avoid an uninterpretable intercept, however we might be prone to bias as we set the parameters ourselves. On the other hand, using the binomial model might prone to less bias as we do not decide that any parameters, but might return (as in this case) values which are uninterpretable (as the intercept in this). It should however be noticed that the plot of m7 is based upon all the subject data whereas the "optim"-plot is only based on subject 7, thus making them not entirely comparable.

### Exercise 7, part 2 - EH

```

sigmodel_pas <- function(pas_lev, subjx) {
  dat <- data.frame(
    x = subjx[subjx$pas == pas_lev, ]$target.frames,
    y = subjx[subjx$pas == pas_lev, ]$correct)
  subjpxpx_opt <- optim(
    par = c(0.5, 1, 1, 1),
    fn = RSS,
    data = dat,
    method = "L-BFGS-B",
    lower = c(0.5, 0.5, -Inf, -Inf),
    upper = c(1, 1, Inf, Inf)
  )
  list(
    subjpxpx_opt$par[1],
    subjpxpx_opt$par[2],
    subjpxpx_opt$par[3],
    subjpxpx_opt$par[4]
  )
}

sigmodel <- function(x, dat, paslevs) {
  subjx <- dat[dat$subject == x, ]
  sapply(paslevs, sigmodel_pas, subjx = subjx,
         USE.NAMES = FALSE, simplify = FALSE)
}

fit_sigmodel <- function(pas, dat, params) {
  sigfit(
    a = params[, pas][1],
    b = params[, pas][2],
    c = params[, pas][3],
    d = params[, pas][4],
    x = dat[dat$pas == pas, ]$target.frames
  )
}

pars <- c("a", "b", "c", "d")
subjects <- levels(exp1$subject)
paslevs <- levels(exp1$pas)
N <- length(subjects)

sigmodel_params <- array(
  unlist(sapply(
    subjects,
    sigmodel,
    dat = exp1,
    paslevs = paslevs,
    USE.NAMES = FALSE,
    simplify = FALSE
  )),
  dim = c(4, 4, N), dimnames = list(pars, paslevs, subjects)
)

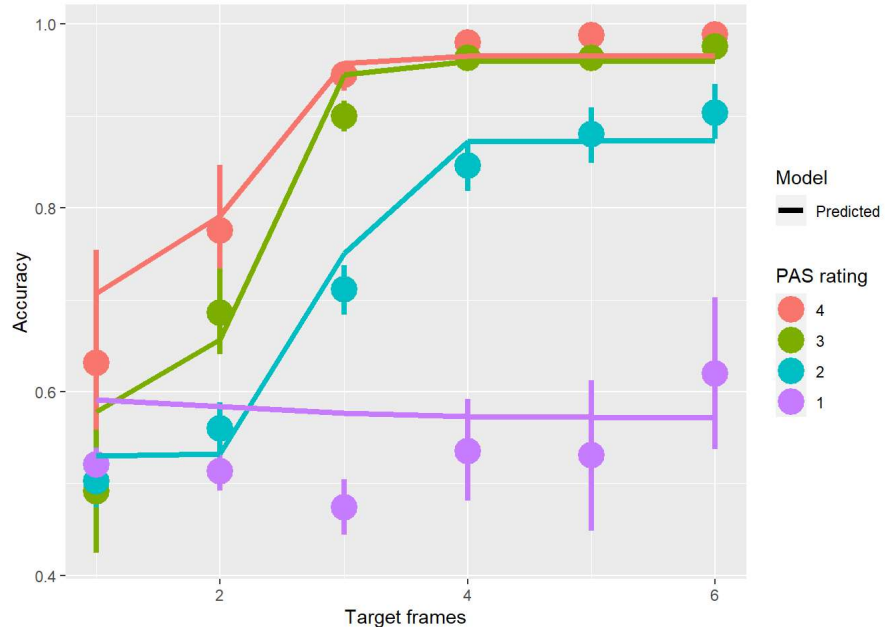
inter_subj_par_means_by_pas <- rowMeans(sigmodel_params, dims = 2)

sig_fitted <- sapply(
  paslevs,
  fit_sigmodel,
  dat = exp1,
  params = inter_subj_par_means_by_pas
)

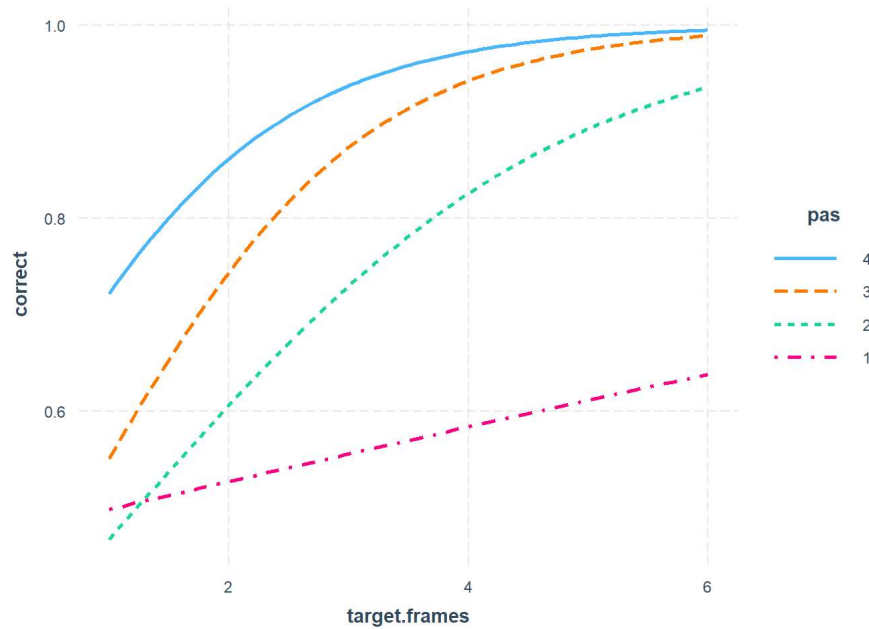
exp1$y_hat <- NA
exp1[exp1$pas == 1, ]$y_hat <- sig_fitted`1`
exp1[exp1$pas == 2, ]$y_hat <- sig_fitted`2`
exp1[exp1$pas == 3, ]$y_hat <- sig_fitted`3`
exp1[exp1$pas == 4, ]$y_hat <- sig_fitted`4`

ggplot(exp1, aes(target.frames, as.numeric(correct), color = pas)) +
  stat_summary(fun.data = "mean_cl_boot", size = 1.5) +
  geom_line(
    aes(x = target.frames, y = y_hat, linetype = "Predicted"),
    size = 1.5
  ) +
  labs(
    x = "Target frames",
    y = "Accuracy",
    color = "PAS rating",
    linetype = "Model"
  )
}

```



```
interactions::interact_plot(model = m6, pred = "target.frames", modx = "pas")
```



```
summary(m6)
```

```

## Generalized linear mixed model fit by maximum likelihood (Laplace
## Approximation) [glmerMod]
## Family: binomial ( logit )
## Formula:
## correct ~ target.frames + pas + (pas * target.frames) + (target.frames |
##     subject)
## Data: exp1
## Control: glmerControl(optimizer = "bobyqa")
##
##      AIC      BIC  logLik deviance df.resid
## 19506.1 19595.5 -9742.0 19484.1    25033
##
## Scaled residuals:
##      Min      1Q  Median      3Q      Max
## -19.0101  0.0537  0.1606  0.4849  1.4465
##
## Random effects:
## Groups   Name        Variance Std.Dev. Corr
## subject (Intercept) 0.03698  0.1923
##         target.frames 0.02057  0.1434  -0.76
## Number of obs: 25044, groups: subject, 29
##
## Fixed effects:
##             Estimate Std. Error z value Pr(>|z|)
## (Intercept) 0.08009  0.24320  0.329  0.74193
## target.frames 0.87404  0.06792 12.869 < 2e-16 ***
## pas3       -0.74022  0.27065 -2.735  0.00624 **
## pas2       -0.77311  0.25133 -3.076  0.00210 **
## pas1       -0.20172  0.24613 -0.820  0.41246
## target.frames:pas3 -0.01055  0.07335 -0.144  0.88565
## target.frames:pas2 -0.31206  0.06850 -4.555 5.23e-06 ***
## target.frames:pas1 -0.75924  0.06749 -11.250 < 2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##          (Intr) trgt.f pas3   pas2   pas1   trg.:3 trg.:2
## target.frms -0.896
## pas3        -0.869  0.759
## pas2        -0.945  0.826  0.847
## pas1        -0.966  0.842  0.859  0.936
## trgt.frms:3 0.775 -0.776 -0.926 -0.761 -0.769
## trgt.frms:2 0.841 -0.841 -0.762 -0.921 -0.839  0.790
## trgt.frms:1 0.853 -0.850 -0.767 -0.838 -0.916  0.791  0.870

```

7.2.i Of course the plot for the optim function looks a bit wonky (as it is based on the predicted values), but the shape of the slope can still be assessed. A crucial difference between the plots is the slope of PAS 1, which is positive for m6 and negative for the plot of the optim function. As in the previous exercise, the advantage of the optim function is that you can control the parameters resulting in avoidance of below chance level estimates - but as we decide the parameters it might be prone to bias. For m6, we use the inbuilt binomial function making us less prone to bias, but giving us estimates which are uninterpretable.

# Portfolio Assignment 3, Methods 3, 2021, autumn semester

Study Group 8, Emma Rose Hahn (EH), Viara Krasteva (VK), Kristian Nøhr Villebro (KV), Luke Ring (LR)

```
In [ ]: #!/usr/bin/env python3
# -*- coding: utf-8 -*-
from multiprocessing.connection import wait
import sys
from time import sleep
import numpy as np
import matplotlib.pyplot as plt
from sklearn.preprocessing import StandardScaler
from sklearn.linear_model import LogisticRegression
from sklearn.model_selection import cross_val_score, StratifiedKFold
from sklearn.svm import SVC
from sklearn.model_selection import train_test_split
from sklearn.metrics import accuracy_score, ConfusionMatrixDisplay
from sklearn import preprocessing
import seaborn as sns
```

## Exercise 1

### 1.1 Describe data structure (EH)

```
In [ ]: data = np.load('megmag_data.npy')
dshape = data.shape
print('Megmag data shape: {}'.format(dshape))
```

Megmag data shape: (682, 102, 251)

1.1.i There are 682 repetitions, 102 sensors and 251 timestamps

### Add time offset (ms) (EH)

```
In [ ]: times = np.arange(-200, 801, 4)
print('Time offset array shape: {}'.format(times.shape))
```

Time offset array shape: (251,)

### Create covariance matrix (VK)

```
In [ ]: output = []
for i in range(len(data)):
    X = data[i]
    Xt = X.T
    output.append(np.matrix(np.dot(X, Xt)))
cov_mat = 1 / dshape[0] * np.sum(i for i in output)

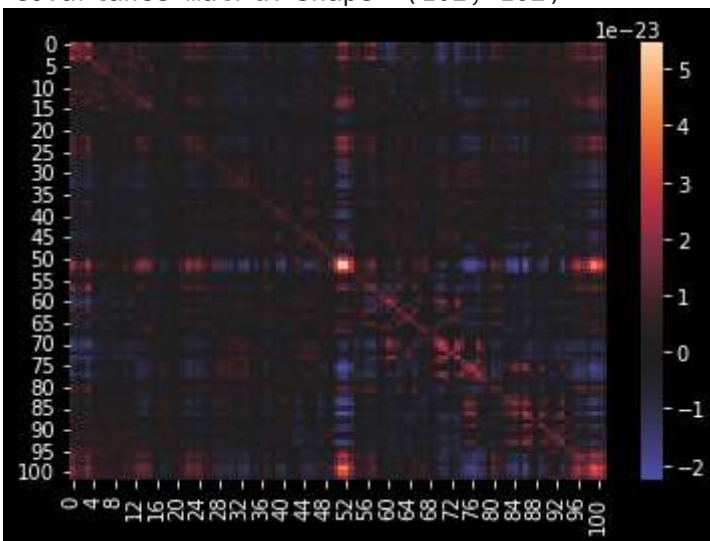
print('Covariance matrix shape: {}'.format(cov_mat.shape))
sns.heatmap(cov_mat, cmap='icefire', center = 0)
#plt.imshow(cov_mat)
plt.show()
```

C:\Users\webma\AppData\Local\Temp\ipykernel\_24288/1960884245.py:6: DeprecationWarn

```
ing: Calling np.sum(generator) is deprecated, and in the future will give a different result. Use np.sum(np.fromiter(generator)) or the python sum builtin instead.
```

```
cov_mat = 1 / dshape[0] * np.sum(i for i in output)
```

```
Covariance matrix shape: (102, 102)
```



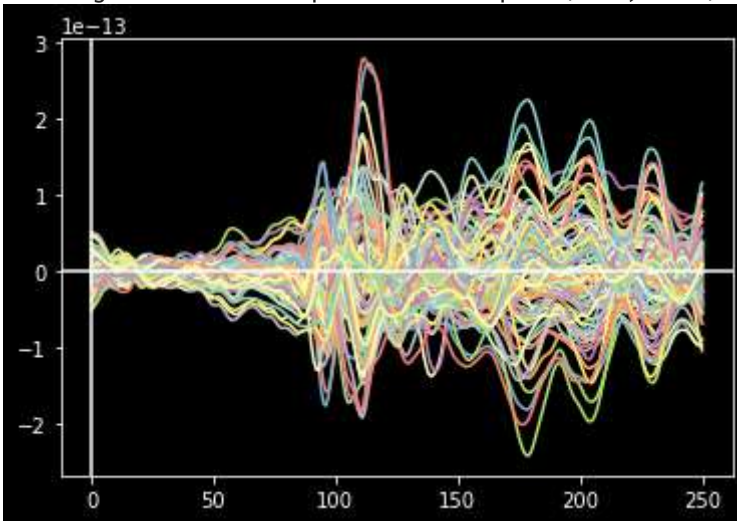
1.1.iii It looks like there is a lot of covariance (which would make sense), however it is quite difficult to get a clear impression with such big covariance matrix.

## Create signal averages across repetitions (KV)

```
In [ ]:
```

```
avg_for_reps = np.mean(data, axis=0).T
print('Average for each repetition shape: {}'.format(avg_for_reps.shape))
plt.plot(avg_for_reps)
plt.axvline()
plt.axhline()
plt.show()
```

```
Average for each repetition shape: (251, 102)
```



## Find and plot maximum response channel (KV)

```
In [ ]:
```

```
max_resp = np.unravel_index(np.argmax(avg_for_reps), avg_for_reps.shape)
print('Max response index: {}, data shape: {}'.format(max_resp, data.shape))
#plot average for all repetitions
plt.plot(times, avg_for_reps[:, max_resp[1]])
plt.axvline()
plt.axvline(x = times[max_resp[0]], color='r', label='max response')
plt.axhline()
plt.show()

# plot all repetitions for max response channel
```

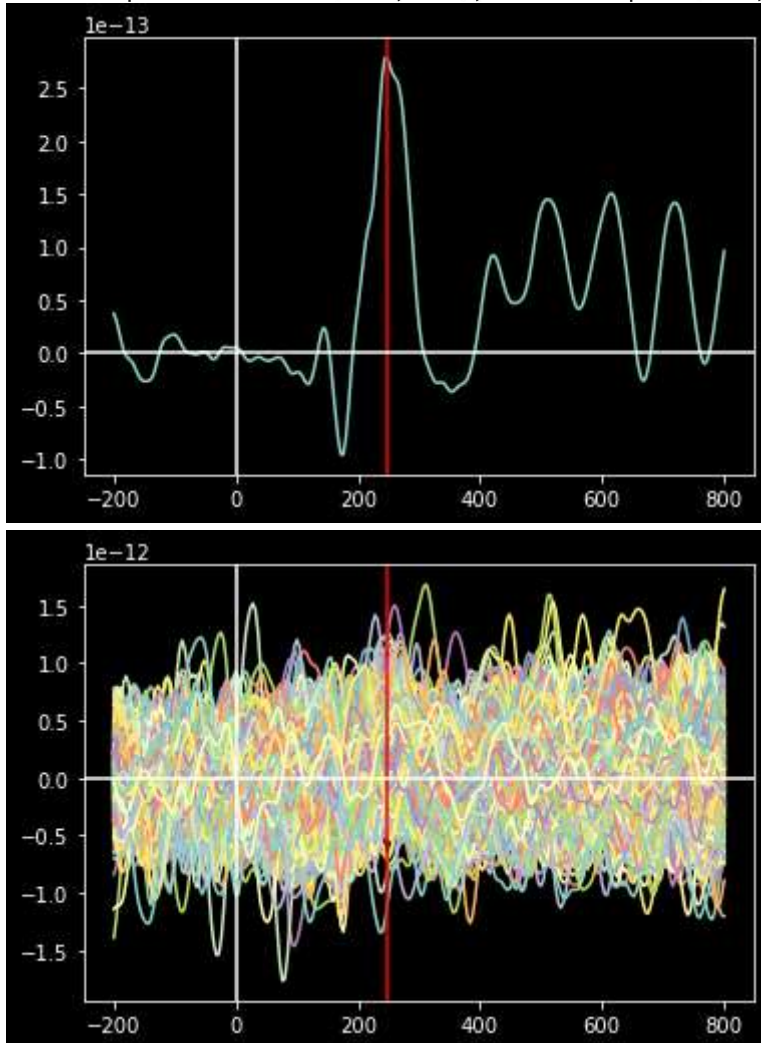
```

plt.plot(times, data[:, max_resp[1], :].squeeze().T)
plt.axvline()
plt.axvline(x = times[max_resp[0]], color='r', label='max response')
plt.axhline()

plt.show()

```

Max response index: (112, 73), data shape: (682, 102, 251)



1.1.vi We found that the maximum response signal was received by sensor 74 (as python index starts at 0)

1.1.viii The messy plot showing the single repetitions includes a lot of fluctuation around the y-axis, thus including lots of noise, making it difficult to assess anything from. Even though the messy plot looks as if the activity is more or less the same across the whole time period, the averaged plot reveals that activation is quite different over time. As the signal noise goes in both negative and positive direction, they cancel each other out thus resulting in the averaged plot we have seen.

## load pas\_vector.npy (LR)

```

In [ ]:
y = np.load('pas_vector.npy')
# same as number of repetitions
print('Pas vector shape: {}'.format(y.shape))
pas_data = []
pas_index = {}
for i in np.unique(y):
    print('Creating pas slice for pas {}'.format(i))
    pas_slice = np.argwhere(y == i)
    data_slice = data[pas_slice].squeeze()
    pas_data.append({
        'paslevel': i,

```

```

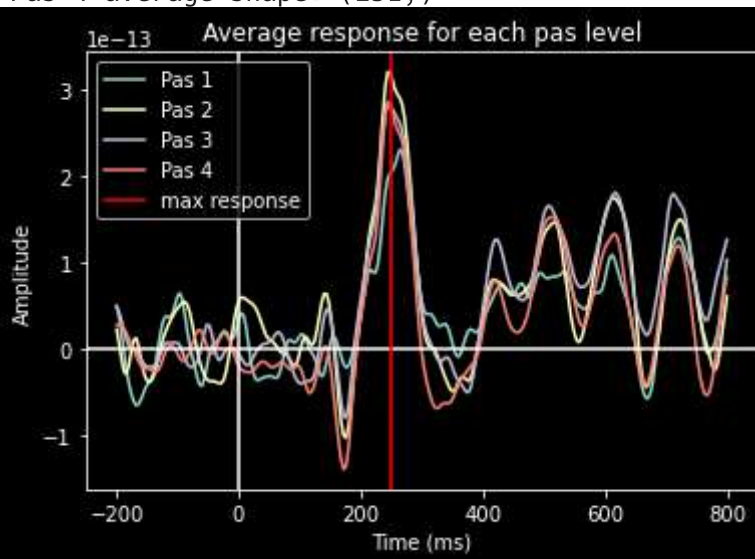
        'data': data_slice,
    })
pas_index[i] = len(pas_data) - 1
for i in pas_data:
    print('Pas {} shape: {}'.format(i['paslevel'], i['data'].shape))
    avg_for_pas = np.mean(i['data'][:,max_resp[1],:], axis=0).T
    print('Pas {} average shape: {}'.format(i['paslevel'], avg_for_pas.shape))
    plt.plot(times, avg_for_pas, label='Pas {}'.format(i['paslevel']))
plt.axvline()
plt.axhline()
plt.axvline(x = times[max_resp[0]], color='r', label='max response')
plt.legend(loc="upper left")
plt.title('Average response for each pas level')
plt.xlabel('Time (ms)')
plt.ylabel('Amplitude')
plt.show()

```

```

Pas vector shape: (682,)
Creating pas slice for pas 1
Creating pas slice for pas 2
Creating pas slice for pas 3
Creating pas slice for pas 4
Pas 1 shape: (99, 102, 251)
Pas 1 average shape: (251,)
Pas 2 shape: (115, 102, 251)
Pas 2 average shape: (251,)
Pas 3 shape: (208, 102, 251)
Pas 3 average shape: (251,)
Pas 4 shape: (260, 102, 251)
Pas 4 average shape: (251,)

```



1.2.i The pas\_vector has length 682, thus having the same length as repetitions.

1.2.iii One would expect that the lower the PAS level the lower the response signal – e.g., PAS 1, not perceiving anything, would result in a low signal, and PAS 4 would result in the highest signal. This seems to correlate with what we see in plot. However, surprisingly PAS 2 has a higher max at 250ms than PAS 4 (which we would expect would have the highest).

## Exercise 2

### 2.1 Logistic regression (EH)

```

In [ ]: # create a new array called data_1_2 that only contains PAS responses 1 and 2
data_1_2 = np.concatenate((pas_data[pas_index[1]]['data'], pas_data[pas_index[2]]])
print('Data for pas 1 and 2 shape: {}'.format(data_1_2.shape))
# Similarly, create a y_1_2 for the target vector

```

```

y_1_2 = y[np.where((y == 1) | (y == 2))]
print('y for pas 1 and 2 shape: {}'.format(y_1_2.shape))
#Our data_1_2 is a three-dimensional array. Our strategy will be to collapse our 1
X_1_2 = np.reshape(data_1_2, (data_1_2.shape[0], data_1_2.shape[1] * data_1_2.shape[2]))
print('Reshaped data for pas 1 and 2 shape: {}'.format(X_1_2.shape))
# and scale X_1_2
sc = StandardScaler()
sc.fit(X_1_2)
X_1_2_std = sc.transform(X_1_2)
# Do a standard LogisticRegression - make sure there is no penalty applied
lr = LogisticRegression(penalty='none', solver='lbfgs', multi_class='ovr', n_jobs=-1)
lr.fit(X_1_2_std, y_1_2)
# we get a score of 1.0...that'd definitely an overfit, right? haha
print('Logistic regression score with no penalty: {}'.format(lr.score(X_1_2_std, y_1_2)))
#found classes
print('Classes found in regression: {}'.format(lr.classes_))
# total coefs
coefs_orig = lr.coef_
coefs = len(lr.coef_[0])
print('Total coefs: {}'.format(coefs))
# non-zero coefs
coefs_nonzero = len(lr.coef_[0].nonzero()[0])
print('Nonzero coefs: {}'.format(coefs_nonzero))
# zero coefs
coefs_zero = coefs - coefs_nonzero
print('Zero coefs: {}'.format(coefs_zero))
# apply l1 penalty
lr = LogisticRegression(penalty='l1', solver='liblinear', multi_class='ovr')
lr.fit(X_1_2_std, y_1_2)
print('Logistic regression score with l1 penalty: {}'.format(lr.score(X_1_2_std, y_1_2)))
# default_threshold = np.get_printoptions()['threshold']
# np.set_printoptions(threshold=sys.maxsize)
# print(lr.coef_)
# np.set_printoptions(threshold=default_threshold)
#found classes
print('Classes found in regression: {}'.format(lr.classes_))
# total coefs
print('Total coefs: {}'.format(lr.coef_.shape))
coefs_orig = lr.coef_[0]
coefs = len(lr.coef_[0])
print('Total coefs: {}'.format(coefs))
# non-zero coefs
coefs_nonzero = len(lr.coef_[0].nonzero()[0])
print('Nonzero coefs: {}'.format(coefs_nonzero))
# zero coefs
coefs_zero = coefs - coefs_nonzero
print('Zero coefs: {}'.format(coefs_zero))
# part 2.1.3: something is wrong with the above code because there is only one non-zero coefficient
cov_nz_index = lr.coef_[0].nonzero()[0]
X_reduced = X_1_2_std[:, cov_nz_index]

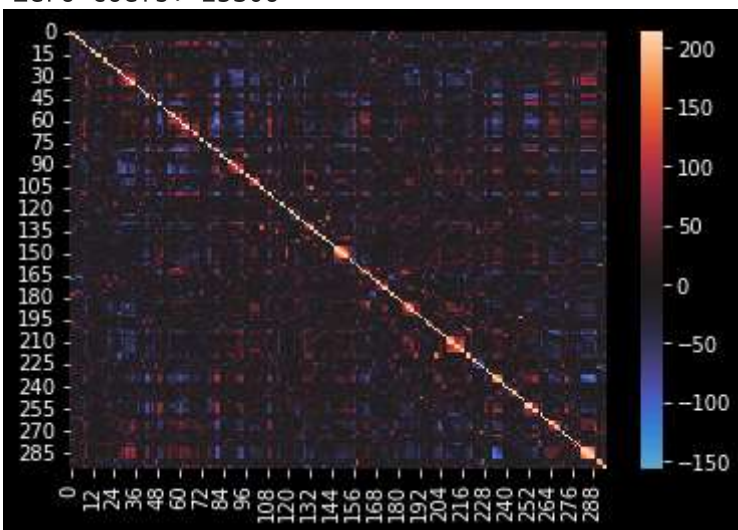
cmat = X_reduced.T @ X_reduced

sns.heatmap(cmat, cmap='icefire', center = 0)
plt.show()

```

Data for pas 1 and 2 shape: (214, 102, 251)  
 y for pas 1 and 2 shape: (214,)  
 Reshaped data for pas 1 and 2 shape: (214, 25602)  
 Logistic regression score with no penalty: 1.0  
 Classes found in regression: [1 2]  
 Total coefs: 25602  
 Nonzero coefs: 25602  
 Zero coefs: 0  
 Logistic regression score with l1 penalty: 1.0

```
Classes found in regression: [1 2]
Total coefs: (1, 25602)
Total coefs: 25602
Nonzero coefs: 296
Zero coefs: 25306
```



2.1.v As we get a score of 1.0, it seems like an overfitting situation – besides the score of 1.0 we also suspect overfitting based on the lack of model-penalties.

2.1.vi We found there to be 215 non-zero coefficients.

2.1.vii It could seem as there is less covariance, it is however super difficult to assess this from these complex plots still. <3

## Exercise 2.2 cross validation (VK)

```
In [ ]: def equalize_targets_class(data, y):
    np.random.seed(7)
    targets = np.unique(y) ## find the number of targets
    if len(targets) < 2:
        raise NameError("at least 2 targets required")
    counts = list()
    indices = list()
    for target in targets:
        counts.append(np.sum(y == target)) ## find the number of each target
        indices.append(np.where(y == target)[0]) ## find their indices
    min_count = np.min(counts)
    # randomly choose trials
    choices = []
    for i in range(len(targets)):
        choices.append(np.random.choice(indices[i], size=min_count, replace=False))

    # create the new data sets
    new_indices = np.concatenate(tuple(choices))
    new_y = y[new_indices]
    new_data = data[new_indices, :, :]

    return new_data, new_y
```

## Stratified 5-fold cross validation logistic model (KV)

```
In [ ]: X_1_2_equal, y_1_2_equal = equalize_targets_class(data_1_2, y_1_2)
X_1_2_equal = np.reshape(X_1_2_equal, (X_1_2_equal.shape[0], X_1_2_equal.shape[1]))
sc.fit(X_1_2_equal)
X_1_2_equal_std = sc.transform(X_1_2_equal)
cv = StratifiedKFold(n_splits=5)
```

```

lr = LogisticRegression(penalty='none', solver='lbfgs', multi_class='ovr', n_jobs=-1)
print('Getting cross validation score (Stratified 5-fold)...')
scores = cross_val_score(lr, X_1_2_equal_std, y_1_2_equal, cv=cv)
print('Cross validation score (Stratified 5-fold): {}'.format(np.mean(scores)))

```

Getting cross validation score (Stratified 5-fold)...  
Cross validation score (Stratified 5-fold): 0.4746153846153846

## Stratified 5-fold cross validation logistic model with L2 (LR)

```

In [ ]: best_score = 0.0
for c in [1e5, 1e1, 1e-5]:
    print('Getting cross validation score (Stratified 5-fold, C={})...'.format(c))
    lr = LogisticRegression(penalty='l2', solver='lbfgs', multi_class='ovr', C=c, n_jobs=-1)
    scores = cross_val_score(lr, X_1_2_equal_std, y_1_2_equal, cv=cv)
    score = np.mean(scores)
    print('Cross validation score (Stratified 5-fold, C={}): {}'.format(c, score))
    if score >= best_score:
        best_score = score
        best_c = c
print('Best C: {}'.format(best_c))

```

Getting cross validation score (Stratified 5-fold, C=100000.0)...  
Cross validation score (Stratified 5-fold, C=100000.0): 0.4746153846153846  
Getting cross validation score (Stratified 5-fold, C=10.0)...  
Cross validation score (Stratified 5-fold, C=10.0): 0.4746153846153846  
Getting cross validation score (Stratified 5-fold, C=1e-05)...  
Cross validation score (Stratified 5-fold, C=1e-05): 0.45487179487179485  
Best C: 10.0

2.2.iv It seems as if C=1e-05 gives the worst score at 0.455 and C=10.0 gives the best score at 0.475. The difference between them is thus 0.02. It is however surprising that C=1e5 seems to give almost the exact same number as C=1e1. This indicates that regularization is not making a large difference for these data.

## now fit a model for each time sample using best c (EH)

```

In [ ]: # 2.2.v
scores = []
lr = LogisticRegression(penalty='l2', solver='newton-cg', multi_class='ovr', C=best_c)
X_1_2_equal, y_1_2_equal = equalize_targets_class(data_1_2, y_1_2)
print('Fitting l2 model (c={}) for time sample: {}'.format(best_c), end='', flush=True)
for time_ofs in range(data_1_2.shape[2]):
    X_1_2_equal_time = X_1_2_equal[:, :, time_ofs].squeeze()
    print('{}.'.format(time_ofs), end='..', flush=True)
    # print('Shape of X: {}'.format(X_1_2_equal_time.shape))
    sc.fit(X_1_2_equal_time)
    X_1_2_equal_std = sc.transform(X_1_2_equal_time)
    # print('Getting Cross validation score (Stratified 5-fold, C={}, time={}ms)'.format(best_c, time_ofs))
    time_scores = cross_val_score(lr, X_1_2_equal_std, y_1_2_equal, cv=cv)
    score = np.mean(time_scores)
    # print('Cross validation score (Stratified 5-fold, C={}, time={}ms): {}'.format(best_c, time_ofs, score))
    scores.append(score)
print('\nCompleted fitting time-based models')
scores = np.array(scores)
max_score = np.unravel_index(np.argmax(scores), scores.shape)
print('Best score: {} at timestamp {}ms'.format(scores[max_score], times[max_score]))
plt.scatter(times, scores)
plt.axvline(x = times[max_score[0]], color='r', label='max model score')
plt.axhline(y = 0.5, label = 'random chance', color='w')
plt.legend(loc="upper left")
plt.show()

```

```
Fitting l2 model (c=10.0) for time sample: 0..1..2..3..4..5..6..7..8..9..10..11..1
2..13..14..15..16..17..18..19..20..21..22..23..24..25..26..27..28..29..30..31..3
2..33..34..35..36..37..38..39..40..41..42..43..44..45..46..47..48..49..50..51..5
2..53..54..55..56..57..58..59..60..61..62..63..64..65..66..67..68..69..70..71..7
2..73..74..75..76..77..78..79..80..81..82..83..84..85..86..87..88..89..90..91..9
2..93..94..95..96..97..98..99..100..101..102..103..104..105..106..107..108..109..1
10..111..112..113..114..115..116..117..118..119..120..121..122..123..124..125..12
6..127..128..129..130..131..132..133..134..135..136..137..138..139..140..141..14
2..143..144..145..146..147..148..149..150..151..152..153..154..155..156..157..15
8..159..160..161..162..163..164..165..166..167..168..169..170..171..172..173..17
```

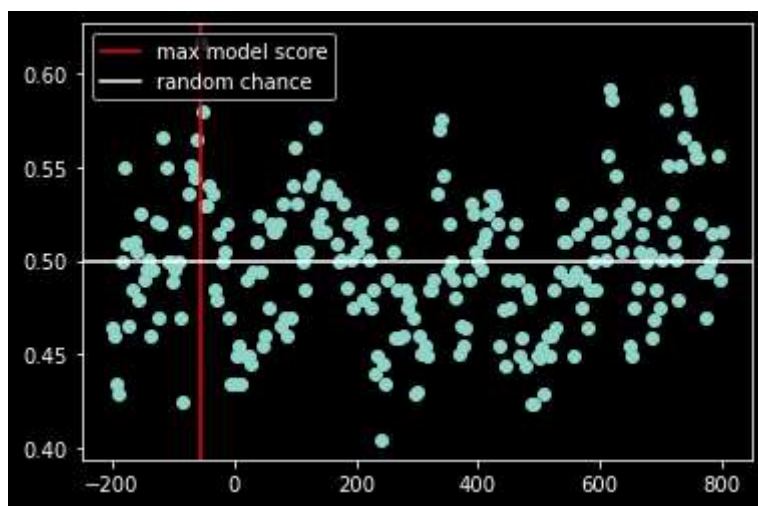
2.2.v The best score 61.1% is at timestamp = 176 ms. Chance level should be at 50% as we are predicting PAS 1 and PAS 2

## now fit a model for each time sample using c=1e-1 and l1 (VK)

In [ ]:

```
scores = []
lr = LogisticRegression(penalty='l1', solver='liblinear', multi_class='ovr', C=1e-1)
X_1_2_equal, y_1_2_equal = equalize_targets_class(data_1_2, y_1_2)
print('Fitting l1 model (c=1e-1) for time sample: ', end='', flush=True)
for time_ofs in range(data_1_2.shape[2]):
    X_1_2_equal_time = X_1_2_equal[:, :, time_ofs].squeeze()
    print('{}.'.format(time_ofs), end='.', flush=True)
    # print('Shape of X: {}'.format(X_1_2_equal_time.shape))
    sc.fit(X_1_2_equal_time)
    X_1_2_equal_std = sc.transform(X_1_2_equal_time)
    # print('Getting Cross validation score (Stratified 5-fold, C={}, time={}ms)'.format(C, time))
    time_scores = cross_val_score(lr, X_1_2_equal_std, y_1_2_equal, cv=cv)
    score = np.mean(time_scores)
    # print('Cross validation score (Stratified 5-fold, C={}, time={}ms): {}'.format(C, time))
    scores.append(score)
print('\nCompleted fitting time-based models')
scores = np.array(scores)
max_score = np.unravel_index(np.argmax(scores), scores.shape)
print('Best score: {} at timestamp {}ms'.format(scores[max_score], times[max_score]))
plt.scatter(times, scores)
plt.axvline(x = times[max_score[0]], color='r', label='max model score')
plt.axhline(y = 0.5, label = 'random chance', color='w')
plt.legend(loc="upper left")
plt.show()
```

```
Fitting l1 model (c=1e-1) for time sample: 0..1..2..3..4..5..6..7..8..9..10..11..1
2..13..14..15..16..17..18..19..20..21..22..23..24..25..26..27..28..29..30..31..3
2..33..34..35..36..37..38..39..40..41..42..43..44..45..46..47..48..49..50..51..5
2..53..54..55..56..57..58..59..60..61..62..63..64..65..66..67..68..69..70..71..7
2..73..74..75..76..77..78..79..80..81..82..83..84..85..86..87..88..89..90..91..9
2..93..94..95..96..97..98..99..100..101..102..103..104..105..106..107..108..109..1
10..111..112..113..114..115..116..117..118..119..120..121..122..123..124..125..12
6..127..128..129..130..131..132..133..134..135..136..137..138..139..140..141..14
2..143..144..145..146..147..148..149..150..151..152..153..154..155..156..157..15
8..159..160..161..162..163..164..165..166..167..168..169..170..171..172..173..17
4..175..176..177..178..179..180..181..182..183..184..185..186..187..188..189..19
0..191..192..193..194..195..196..197..198..199..200..201..202..203..204..205..20
6..207..208..209..210..211..212..213..214..215..216..217..218..219..220..221..22
2..223..224..225..226..227..228..229..230..231..232..233..234..235..236..237..23
8..239..240..241..242..243..244..245..246..247..248..249..250..
Completed fitting time-based models
Best score: 0.6161538461538462 at timestamp -56ms
```



2.2.vi classification accuracy was best (61.62%) at time point -56ms.

**fit again but for PAS 1 and 4 (KV)**

```
In [ ]: data_1_4 = np.concatenate((pas_data[pas_index[1]]['data'], pas_data[pas_index[4]]) print('Data for pas 1 and 4 shape: {}'.format(data_1_4.shape)) y_1_4 = y[np.where((y == 1) | (y == 2))] print('y for pas 1 and 4 shape: {}'.format(y_1_4.shape)) #Our data_1_2 is a three-dimensional array. Our strategy will be to collapse our i X_1_4 = np.reshape(data_1_4, (data_1_4.shape[0], data_1_4.shape[1] * data_1_4.shape[2])) scores = [] lr = LogisticRegression(penalty='l1', solver='liblinear', multi_class='ovr', C=1e-10) X_1_4_equal, y_1_4_equal = equalize_targets_class(data_1_4, y_1_4) print('Fitting l1 model (c=1e-1) for time sample: ', end='', flush=True) for time_ofs in range(data_1_4.shape[2]): X_1_4_equal_time = X_1_4_equal[:, :, time_ofs].squeeze() print('{}'.format(time_ofs), end='..', flush=True) # print('Shape of X: {}'.format(X_1_4_equal_time.shape)) sc.fit(X_1_4_equal_time) X_1_4_equal_std = sc.transform(X_1_4_equal_time) # print('Getting Cross validation score (Stratified 5-fold, C={}, time={}ms)'.format(C, time)) time_scores = cross_val_score(lr, X_1_4_equal_std, y_1_4_equal, cv=cv) score = np.mean(time_scores) # print('Cross validation score (Stratified 5-fold, C={}, time={}ms): {}'.format(C, time), score) scores.append(score) print('\nCompleted fitting time-based models') scores = np.array(scores) max_score = np.unravel_index(np.argmax(scores), scores.shape) print('Best score: {} at timestamp {}ms'.format(scores[max_score], times[max_score])) plt.scatter(times, scores) plt.axvline(x = times[max_score[0]], color='r', label='max model score') plt.axhline(y = 0.5, label = 'random chance', color='w') plt.legend(loc="upper left") plt.show()
```

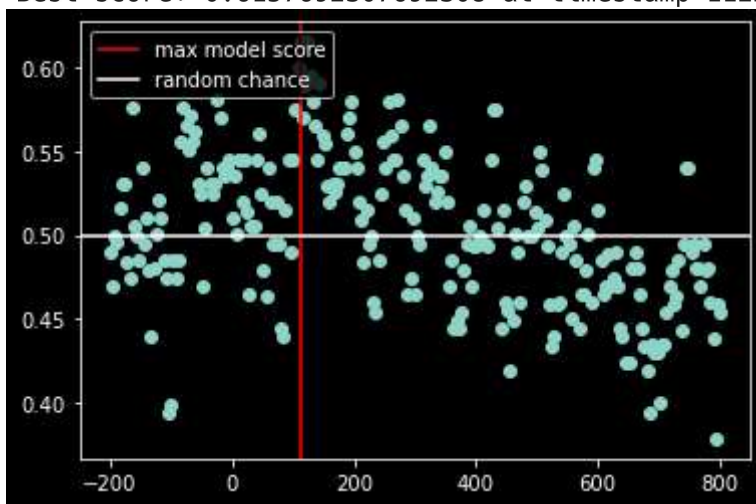
Data for pas 1 and 4 shape: (359, 102, 251)

y for pas 1 and 4 shape: (214, )

```

6..207..208..209..210..211..212..213..214..215..216..217..218..219..220..221..22
2..223..224..225..226..227..228..229..230..231..232..233..234..235..236..237..23
8..239..240..241..242..243..244..245..246..247..248..249..250..
Completed fitting time-based models
Best score: 0.6157692307692308 at timestamp 112ms

```



2.2.vii Best classification (61.58%) is at time point 112ms. Chance level for this analysis should be 50% as we are working with PAS 1 and PAS 4.

2.3 As our scores is all above chance level (however not a lot) it seems to be possible to do pairwise classification. It is quite surprising that the accuracy level of PAS 1 vs 2 and PAS 1 vs 4 are almost identical, though at very different time points.

## EXERCISE 3 - Do a Support Vector Machine Classification on all four PAS-ratings

### 3.1 Support Vector Machine Classification (LR)

In [ ]:

```

# 3.1.i
X_equal, y_equal = equalize_targets_class(data, y)
print('X shape: {}'.format(X_equal.shape))
print('y shape: {}'.format(y_equal.shape))

X_equal_f = np.reshape(X_equal, (X_equal.shape[0], X_equal.shape[1] * X_equal.shape[2]))
sc.fit(X_equal_f)
X_equal_std = sc.transform(X_equal_f)

print('X reshaped shape: {}'.format(X_equal_std.shape))

X shape: (396, 102, 251)
y shape: (396,)
X reshaped shape: (396, 25602)

```

### Linear kernel classifier (LR)

In [ ]:

```

cv = StratifiedKFold()
svm = SVC(kernel='linear')
scores_svm = cross_val_score(svm, X_equal_std, y_equal, cv=cv)
print(np.mean(scores_svm))

```

0.2928164556962025

### Radial basis (EH)

```
In [ ]: cv = StratifiedKFold()
svm = SVC(kernel='rbf')
scores_svm = cross_val_score(svm, X_equal_std, y_equal, cv=cv)
print(np.mean(scores_svm))
```

0.3333544303797468

3.1.iv) Which one is better predicting the category?

Based on the cross-validation results, the linear kernel classifier achieved 29.9% accuracy, and the radial basis kernel classifier achieved 33.4% accuracy, so for these data the radial basis kernel classifier is the best predictor.

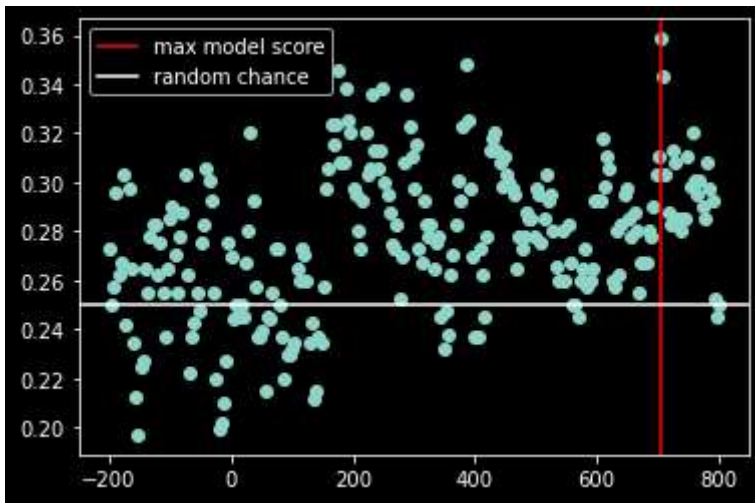
## Sample-by-sample analysis (VK)

```
In [ ]: # radial basis score better than linear
cv = StratifiedKFold()
svm = SVC(kernel='rbf')

scores = []
print(X_equal.shape)
print('Fitting rbv svm model time sample: ', end='', flush=True)
for time_ofs in range(X_equal.shape[2]):
    X_equal_time = X_equal[:, :, time_ofs].squeeze()
    print('{}'.format(time_ofs), end='..', flush=True)
    # print('Shape of X: {}'.format(X_1_2_equal_time.shape))
    sc.fit(X_equal_time)
    X_equal_std = sc.transform(X_equal_time)
    # print('Getting Cross validation score (Stratified 5-fold, C={}, time={}ms)'.format(
    time_scores = cross_val_score(svm, X_equal_std, y_equal, cv=cv)
    score = np.mean(time_scores)
    # print('Cross validation score (Stratified 5-fold, C={}, time={}ms): {}'.format(score))
    scores.append(score)
print('\nCompleted fitting time-based models')
scores = np.array(scores)
max_score = np.unravel_index(np.argmax(scores), scores.shape)
print('Best score: {} at timestamp {}ms'.format(scores[max_score], times[max_score]))
plt.scatter(times, scores)
# random chance is 1/4
plt.axvline(x = times[max_score[0]], color='r', label='max model score')
plt.axhline(y = 0.25, label = 'random chance', color='w')
plt.legend(loc="upper left")
plt.show()
```

(396, 102, 251)  
Fitting rbv svm model time sample: 0..1..2..3..4..5..6..7..8..9..10..11..12..13..1  
4..15..16..17..18..19..20..21..22..23..24..25..26..27..28..29..30..31..32..33..3  
4..35..36..37..38..39..40..41..42..43..44..45..46..47..48..49..50..51..52..53..5  
4..55..56..57..58..59..60..61..62..63..64..65..66..67..68..69..70..71..72..73..7  
4..75..76..77..78..79..80..81..82..83..84..85..86..87..88..89..90..91..92..93..9  
4..95..96..97..98..99..100..101..102..103..104..105..106..107..108..109..110..11  
1..112..113..114..115..116..117..118..119..120..121..122..123..124..125..126..12  
7..128..129..130..131..132..133..134..135..136..137..138..139..140..141..142..14  
3..144..145..146..147..148..149..150..151..152..153..154..155..156..157..158..15  
9..160..161..162..163..164..165..166..167..168..169..170..171..172..173..174..17  
5..176..177..178..179..180..181..182..183..184..185..186..187..188..189..190..19  
1..192..193..194..195..196..197..198..199..200..201..202..203..204..205..206..20  
7..208..209..210..211..212..213..214..215..216..217..218..219..220..221..222..22  
3..224..225..226..227..228..229..230..231..232..233..234..235..236..237..238..23  
9..240..241..242..243..244..245..246..247..248..249..250..

Completed fitting time-based models  
Best score: 0.3586075949367088 at timestamp 704ms



3.1.iv) Is classification of subjective experience possible at around 200-250 ms?

It seems that the model would perform above random chance at classification of subjective experience for 200-250ms although the accuracy is still not high, ranging from around 0.27-0.34.

## Exercises 3.2 Test/train (KV)

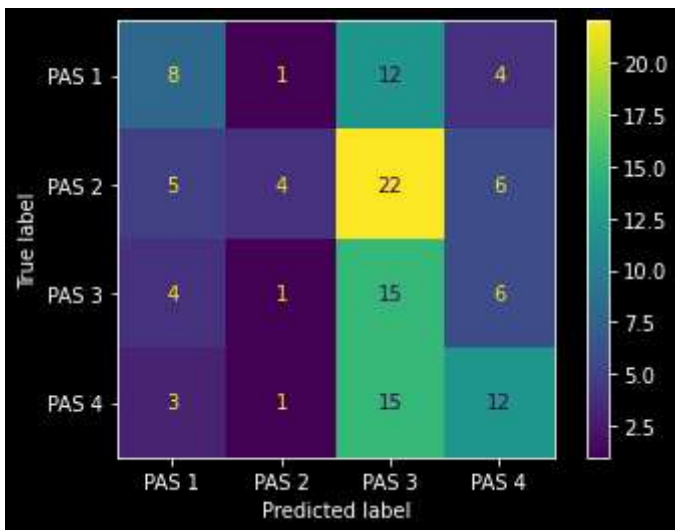
```
In [ ]: X_train, X_test, y_train, y_test = train_test_split(X_equal, y_equal, test_size=0.2)
print(X_train.shape)
print(y_train.shape)
# assignment says: This time your features are the number of sensors multiplied by
# So, we need to reshape the data to be (n_samples, n_features)
X_train_f = np.reshape(X_train, (X_train.shape[0], X_train.shape[1] * X_train.shape[2]))
X_test_f = np.reshape(X_test, (X_test.shape[0], X_test.shape[1] * X_test.shape[2]))

sc.fit(X_train_f)
X_train_f_std = sc.transform(X_train_f)
X_test_f_std = sc.transform(X_test_f)

svm = SVC(kernel='rbf')

svm.fit(X_train_f_std, y_train)
predictions = svm.predict(X_test_f_std)
print('Accuracy: {}'.format(accuracy_score(y_test, predictions)))
# confusion matrix
# print(confusion_matrix(y_test, predictions))
ConfusionMatrixDisplay.from_predictions(y_test, predictions, display_labels=['PAS', 'NPA'])
plt.show()

(277, 102, 251)
(277,)
Accuracy: 0.3277310924369748
```



3.2.iii) Based on the confusion matrix, describe how ratings are misclassified and if that makes sense given that ratings should measure the strength/quality of the subjective experience. Is the classifier biased towards specific ratings?

The confusion matrix shows a bias towards predicting PAS 3, which can be seen from the PAS 3 predicted label on the x axis, this should not come from PAS 3 being the most frequent rating, because the samples were stratified. Of the predicted labels classified as PAS 1, 2 and 4, the prediction was correct most frequently, which fits with subjects indicating that they either not aware of the stimulus (PAS 1) or did not doubt their experience of the stimulus (PAS 4). The PAS 3 label was most frequently misclassified, especially when the true label was PAS 2, but both 2 and 3 ratings that relate to uncertainty and ambiguity.

# Portfolio Assignment 4, Methods 3, 2021, autumn semester

## Exercise 1 - Use principal component analysis to improve the classification of subjective experience

### 1.1 - Create a covariance matrix, find the eigenvectors and the eigenvalues (EH)

```
In [ ]: # methods3 conda environment
# imports
import numpy as np
from sklearn.preprocessing import StandardScaler
from sklearn.linear_model import LogisticRegression
from sklearn.model_selection import cross_val_score, StratifiedKFold
from sklearn.decomposition import PCA
import matplotlib.pyplot as plt
import seaborn as sns
```

```
In [ ]: # common functions
def equalize_targets(data, y):
    np.random.seed(7)
    targets = np.unique(y)
    counts = list()
    indices = list()
    for target in targets:
        counts.append(np.sum(y == target))
        indices.append(np.where(y == target)[0])
    min_count = np.min(counts)
    first_choice = np.random.choice(indices[0], size=min_count, replace=False)
    second_choice = np.random.choice(indices[1], size=min_count, replace=False)
    third_choice = np.random.choice(indices[2], size=min_count, replace=False)
    fourth_choice = np.random.choice(indices[3], size=min_count, replace=False)

    new_indices = np.concatenate((first_choice, second_choice,
                                 third_choice, fourth_choice))
    new_y = y[new_indices]
    new_data = data[new_indices, :, :]

    return new_data, new_y
```

```
In [ ]: data = np.load('megmag_data.npy')
pas_data = np.load('pas_vector.npy')
```

```
In [ ]: # 1.1.ii) Equalize the number of targets in y and data using equalize_targets
data, y = equalize_targets(data, pas_data)
```

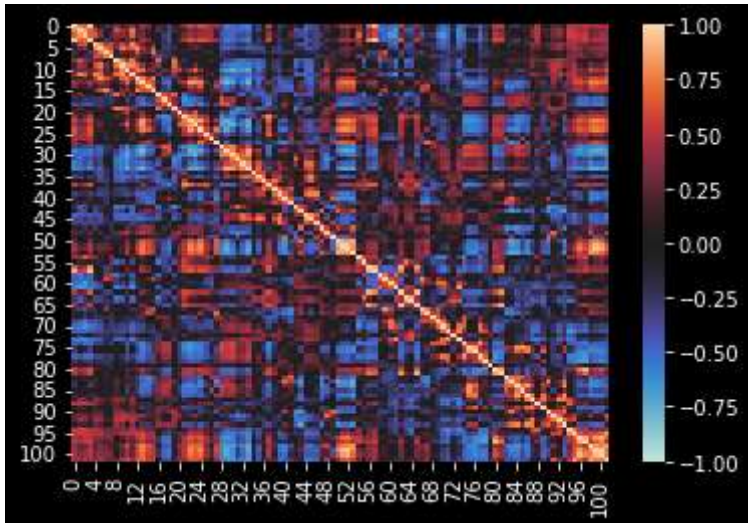
```
In [ ]: # 1.1.iii) times
times=np.arange(-200, 804, 4)
# 1.1.iii) reduce to two dimensions only using 248ms
data_reduced = data[:, :, np.where(times == 248)[0][0]]
```

```
In [ ]: # 1.1.iv) Scale the data using StandardScaler
```

```
sc = StandardScaler()
data_scaled = sc.fit_transform(data_reduced, y)
```

In [ ]:

```
# 1.1.v) Calculate the sample covariance matrix using np.cov
cov_matrix = np.cov(data_scaled, rowvar=False)
# plot using seaborn heatmap
sns.heatmap(cov_matrix, cmap='icefire', vmin=-1, vmax=1)
plt.show()
```



### 1.1.vi) What does the off-diagonal activation imply about the independence of the signals measured by the 102 sensors? (KV)

We see that the off-diagonal activation implies that the signals measured by the sensors are not independent. This makes sense as activity in the brain can involve multiple regions, and we would expect to see some correlation and covariance between sensors.

In [ ]:

```
# 1.1.vii) Run np.linalg.matrix_rank on the covariance matrix - what integer value
rank = np.linalg.matrix_rank(cov_matrix)
print("Matrix rank: {}".format(rank))
```

Matrix rank: 97

In [ ]:

```
# 1.1.viii) Find the eigenvalues and eigenvectors of the covariance matrix using r
eigenvalues, eigenvectors = np.linalg.eig(cov_matrix)
# Use np.real to retrieve only the real parts of the eigenvalues and eigenvectors
eigenvalues = np.abs(np.real(eigenvalues))
eigenvectors = np.real(eigenvectors)
```

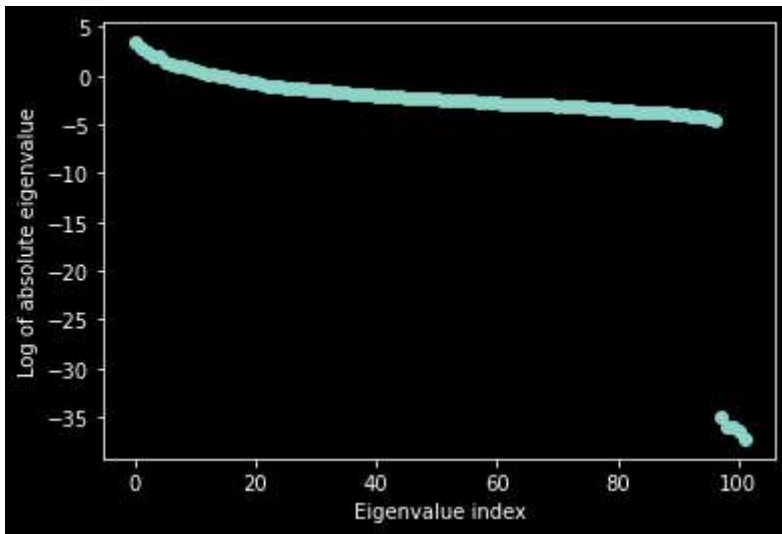
### 1.2 - Create the weighting matrix W and the projected data, Z

In [ ]:

```
# 1.2.i to 1.2.iii) sort the eigenvectors and eigenvalues according to the absolute
# value of the eigenvalues
sorted_indices = np.argsort(eigenvalues)[::-1]
# sorted eigenvalues
eigenvalues = eigenvalues[sorted_indices]
# sorted eigenvectors
eigenvectors = eigenvectors[:, sorted_indices]
```

In [ ]:

```
# 1.2.iv) Plot the log of the eigenvalues
plt.plot(np.log(eigenvalues), 'o')
plt.xlabel('Eigenvalue index')
plt.ylabel('Log of absolute eigenvalue')
plt.show()
```



### 1.2.iv) are there some values that stand out from the rest? (LR)

The last 5 of the sorted log eigenvalues are much smaller than the previous 97. This matches the result of our matrix rank test which gives values above a threshold.

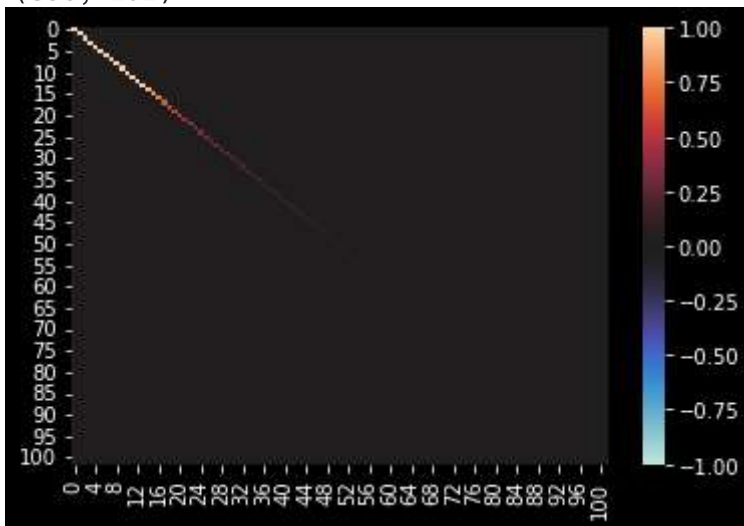
```
In [ ]: # 1.2.v - 1.2.vi) eigenvectors are the weighting matrix, create projected data Z =
Z = data_scaled @ eigenvectors
X = Z @ eigenvectors.T

# check if calculations are correct
closeness_check = np.isclose(data_scaled, X)
number_of_false_values = np.sum(closeness_check == False)
print("Number values that are not close: {}".format(number_of_false_values))
```

Number values that are not close: 0

```
In [ ]: # 1.2.vii) Create a new covariance matrix of the principal components (n=102)
cov_matrix_pca = np.cov(Z, rowvar=False)
print(Z.shape)
# plot using seaborn heatmap
sns.heatmap(cov_matrix_pca, cmap='icefire', vmin=-1, vmax=1)
plt.show()
```

(396, 102)



### 1.2.vii What has happened to the off-diagonals and why? (LR)

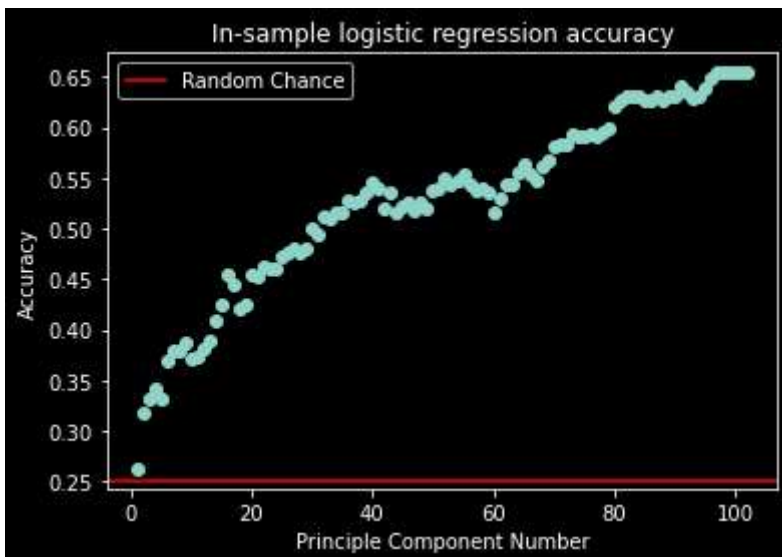
The weighting matrix is the orthogonal projection of the data onto the eigenvectors, this results in values that are not correlated with each other, giving a 0 value for the covariance matrix.

## 2 - Use logistic regression with cross-validation to find the optimal number of principal components (VK)

In [ ]:

```
# 2.1.i) run standard logistic regression (no regularization) based on Zdxk and y
lr = LogisticRegression(fit_intercept=False, solver='newton-cg')
# fit 102 models based on: k=[1,2,...,101,102] and d=102. For each fit get the class
scores = list()
d = Z.shape[1]
for k in range(1, d+1):
    Z_kxd = Z[:, :k]
    lr.fit(Z_kxd, y)
    scores.append(lr.score(Z_kxd, y))

plt.scatter(np.arange(1, d+1), scores)
plt.axhline(0.25, color='red', label='Random Chance')
plt.xlabel('Principle Component Number')
plt.ylabel('Accuracy')
plt.title('In-sample logistic regression accuracy')
plt.legend()
plt.show()
```



2.1.ii) what is the general trend and why is this so? (EH)

Based on the above plot we could say the general trend is that adding more principle components results in a higher classification accuracy. This is because more variance is accounted for, but it also could mean that the model is overfitting.

2.1.iii) In terms of classification accuracy, what is the effect of adding the five last components? Why do you think this is so? (KV)

The last 5 components have a negligible effect, this is because they are the principle components that have the least variance, adding them to our models doesn't provide much additional information.

In [ ]:

```
# 2.2) cross validate using cross_val_score and StratifiedKFold (LR)
cv = StratifiedKFold()
scores = []
for k in range(1, d+1):
    Z_k = Z[:, :k]
    scores.append(cross_val_score(lr, Z_k, y, cv=cv))

scores = np.asarray(scores)
# get mean score for each k
mean_scores = np.mean(scores, axis=1)
```

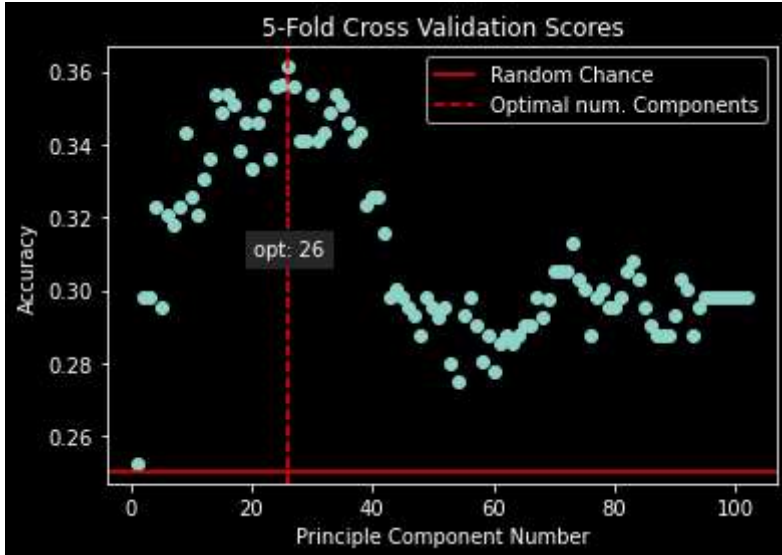
```

# get index of max
max_index = np.argmax(mean_scores)
grand_mean = np.mean(mean_scores)

print(mean_scores.shape)
plt.scatter(np.arange(1, d+1), mean_scores)
plt.axhline(0.25, color='red', label='Random Chance')
plt.axvline(max_index+1, color='red', label='Optimal num. Components', linestyle='--')
plt.text(max_index+1, grand_mean, 'opt: {}'.format(max_index+1), horizontalalignment='center')
plt.xlabel('Principle Component Number')
plt.ylabel('Accuracy')
plt.title('5-Fold Cross Validation Scores')
plt.legend()
plt.show()

```

(102, )



2.2.ii) how is this plot different from the one in Exercise 2.1.ii? (VK)

This shows a different trend, rather than each additional component increasing the model accuracy, we see a peak at 26 components and after that a decline in accuracy which then levels out at around 70 components. Based on the first model we might assume that adding all the components would result in a higher accuracy, whereas using cross-validation we see that the results of the previous model were likely due to overfitting.

2.2.iii) What is the number of principal components,  $k_{max\_accuracy}$ , that results in the greatest classification accuracy when cross-validated?

The peak model performance is 26 components, at around 36% accuracy.

```
In [ ]: print("Maximum classification accuracy: {:.2%}, classification accuracy with full dataset: {:.2%}, improvement: {:.2%}"
```

Maximum classification accuracy: 36.13%, classification accuracy with full dataset: 29.79%, improvement: 6.33%

2.2.iv) How many percentage points is the classification accuracy increased with relative to the full-dimensional,  $d$ , dataset? (EH)

using all of the components, the 5-fold cross-validation accuracy is increased by about 6.3%.

2.2.v) How do the analyses in Exercises 2.1 and 2.2 differ from one another? Make sure to comment on the differences in optimization criteria. (KV)

Both use a logistic regression model, the key difference here is that the first analysis is adding components and fitting the model to the entire dataset. 2.2 uses stratified 5-fold cross-validation which tries to ensure balanced classes across the folds, as well as training and testing on different subsets of the data. This is a more robust method of testing the model, as it is less prone to overfitting and while the classification accuracy may seem lower, the accuracy in 2.1 is likely overinflated.

In [ ]:

```
# 2.3.i) For each of the 251 time samples, use the same estimator and cross-validate
scores_pca = []
scores_all = []
time_steps = len(times)
print("Running analyses for {} time samples from {}ms to {}ms:".format(time_steps,
for t in range(time_steps):
    print("{}ms".format(times[t]), end="", flush=True)
    # get data for current timestamp
    data_t = data[:, :, t]
    data_t_scaled = sc.transform(data_t)
    # use pca to reduce to k_max
    pca_kmax = PCA(n_components=max_index+1)
    pcs = pca_kmax.fit_transform(data_t_scaled)
    # calculate cross validation scores for reduced data
    scores_pca.append(cross_val_score(lr, pcs, y, cv=cv))
    print(".", end="", flush=True)
    # calculate cross validation scores for all data
    scores_all.append(cross_val_score(lr, data_t_scaled, y, cv=cv))
    print(".", end="", flush=True)
print("done\nPreparing plot...", flush=True)
```

Running analyses for 251 time samples from -200ms to 800ms:  
-200ms..-196ms..-192ms..-188ms..-184ms..-180ms..-176ms..-172ms..-168ms..-164ms..-160ms..-156ms..-152ms..-148ms..-144ms..-140ms..-136ms..-132ms..-128ms..-124ms..-120ms..-116ms..-112ms..-108ms..-104ms..-100ms..-96ms..-92ms..-88ms..-84ms..-80ms..-76ms..-72ms..-68ms..-64ms..-60ms..-56ms..-52ms..-48ms..-44ms..-40ms..-36ms..-32ms..-28ms..-24ms..-20ms..-16ms..-12ms..-8ms..-4ms..0ms..4ms..8ms..12ms..16ms..20ms..24ms..28ms..32ms..36ms..40ms..44ms..48ms..52ms..56ms..60ms..64ms..68ms..72ms..76ms..80ms..84ms..88ms..92ms..96ms..100ms..104ms..108ms..112ms..116ms..120ms..124ms..128ms..132ms..136ms..140ms..144ms..148ms..152ms..156ms..160ms..164ms..168ms..172ms..176ms..180ms..184ms..188ms..192ms..196ms..200ms..204ms..208ms..212ms..216ms..220ms..224ms..228ms..232ms..236ms..240ms..244ms..248ms..252ms..256ms..260ms..264ms..268ms..272ms..276ms..280ms..284ms..288ms..292ms..296ms..300ms..304ms..308ms..312ms..316ms..320ms..324ms..328ms..332ms..336ms..340ms..344ms..348ms..352ms..356ms..360ms..364ms..368ms..372ms..376ms..380ms..384ms..388ms..392ms..396ms..400ms..404ms..408ms..412ms..416ms..420ms..424ms..428ms..432ms..436ms..440ms..444ms..448ms..452ms..456ms..460ms..464ms..468ms..472ms..476ms..480ms..484ms..488ms..492ms..496ms..500ms..504ms..508ms..512ms..516ms..520ms..524ms..528ms..532ms..536ms..540ms..544ms..548ms..552ms..556ms..560ms..564ms..568ms..572ms..576ms..580ms..584ms..588ms..592ms..596ms..600ms..604ms..608ms..612ms..616ms..620ms..624ms..628ms..632ms..636ms..640ms..644ms..648ms..652ms..656ms..660ms..664ms..668ms..672ms..676ms..680ms..684ms..688ms..692ms..696ms..700ms..704ms..708ms..712ms..716ms..720ms..724ms..728ms..732ms..736ms..740ms..744ms..748ms..752ms..756ms..760ms..764ms..768ms..772ms..776ms..780ms..784ms..788ms..792ms..796ms..800ms..done  
Preparing plot...

In [ ]:

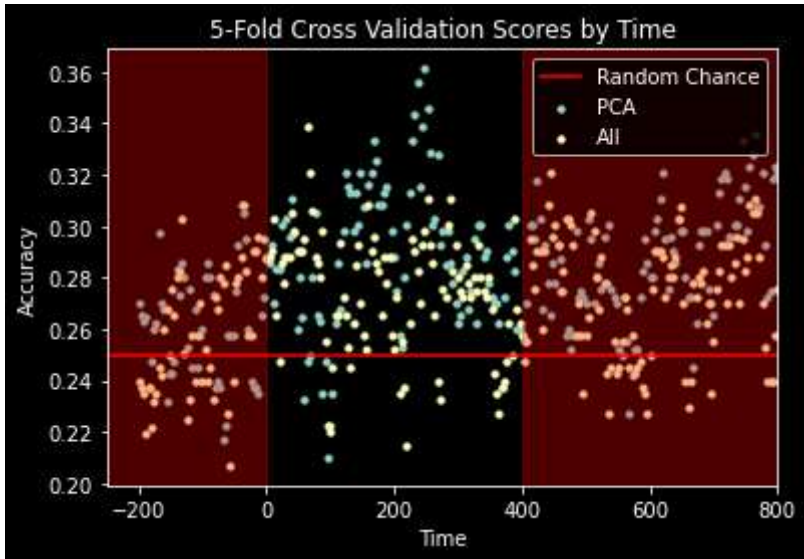
```
# plot scores by time (VK)
mean_pca_scores = np.mean(scores_pca, axis=1)
mean_all_scores = np.mean(scores_all, axis=1)

fig, ax = plt.subplots()
ax.axhline(0.25, color='red', label='Random Chance')
ax.scatter(times, mean_pca_scores, label='PCA', s=10)
ax.scatter(times, mean_all_scores, label='All', s=10)
ax.axvspan(-250, 0, color="red", alpha=0.3)
ax.axvspan(400, 800, color="red", alpha=0.3)
```

```

plt.xlim([-250, 800])
plt.xlabel('Time')
plt.ylabel('Accuracy')
plt.title('5-Fold Cross Validation Scores by Time')
plt.legend(loc='upper right')
plt.show()

```



2.3.iii) Describe the differences between the two analyses - focus on the time interval between 0 ms and 400 ms - describe in your own words why the logistic regression performs better on the PCA-reduced dataset around the peak magnetic activity (EH)

The PCA reduced dataset performance tends to be higher than the full dataset, especially around peak activity. The full dataset contains the signal from all the sensors, but some of these may be less relevant to the response being investigated thereby introducing noise into the model. As hypothesized in 2.2.v, the overall accuracy of the full dataset was inflated due to overfitting which reduced its ability to be generalized to other time intervals. Unsurprisingly, as the PCA model underwent more robust validation, the accuracy at the previously modelled timestamp (248ms) remained the same (~36%), and seems to be able to generalize to time intervals around the peak activity, where the signal is likely similar.