

# Experiment 2

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Start by loading functions and data

```
require(latex2exp)
require(bootES)
# load plyr *BEFORE* dplyr (i.e., tidyverse)
require(plyr)
require(tidyverse)
require(assertthat)
df <- read_csv('../data/data.csv')

# create output img directory if not existant
dir.create('../img', showWarning=F)
```

Preprocess the data, that is do the following

- remove trials where participants responded with a third option; that is, if they were presented with morph *ab* and they responded with *c*, that trial will be removed
- binarize the responses to run the model

```
df <- df %>%
  filter(third_option != 1)

# now we need to add a binarized response to compute the psychometric curves
# but this changes depending on morph_type, so we need to make a function that
# checks the conditions
binarize_response <- function(morph_type, response_identity) {
  if (morph_type == 'ab') {
    return(ifelse(response_identity == 'a', 0, 1))
  } else if (morph_type == 'bc') {
    return(ifelse(response_identity == 'b', 0, 1))
  } else if (morph_type == 'ac') {
    return(ifelse(response_identity == 'a', 0, 1))
  }
}

df <-
df %>%
  rowwise() %>%
```

```

mutate(response_bin=binarize_response(morph_type, response_identity))

df$pos <- as.factor(df$pos)
df$session <- as.factor(df$session)

# make sure that binarize_response worked, i.e. we should get 1s and 0s for each
# morph_type
check_ <-
df %>%
  group_by(morph_type) %>%
  summarise(check=length(unique(response_bin)))
assert_that(length(check_$check) == 3)

## [1] TRUE

assert_that(all(check_$check == c(2, 2, 2)))

## [1] TRUE

```

## Procedure and equations

We are going to fit a linear mixed effect model to the data. We will model the data as follows

$$y^k = \text{logit}(g(x)) \quad g(x) = \beta_0 x + \sum_{i=1}^8 (\beta_i + z_i^k) I_i$$

Where  $y^k$  is the response for subject  $k$ ,  $x$  is the (scaled) percentage of morphing,  $\beta_i, i = 1 \dots 4$  are the fixed-effects for each angular location (0 to 315 in 45 deg steps), and  $z_i^k$  are the random-effects (random slopes for location) for each subject, and  $I_i$  is an indicator variable, indicating the angular location for each trial.

In this way for each subject we can find the PSE as the point where  $y^k = 0.5$ , that is the point  $\hat{x}$

$$\text{logit}(g(\hat{x})) = 0.5 \iff g(\hat{x}) = 0 \iff \beta_0 \hat{x} + \sum_{i=1}^4 (\beta_i + z_i^k) I_i = 0 \iff \hat{x} = -\frac{\sum_{i=1}^4 (\beta_i + z_i^k) I_i}{\beta_0}$$

Thus for every angular location  $i$  we have that

$$\hat{x}_i = -\frac{\beta_i}{\beta_0} - \frac{z_i^k}{\beta_0} = \text{PSE}_i^p + \Delta \text{PSE}_i^s$$

with  $\text{PSE}_i^p$  being the population-level PSE at location  $i$ , and  $\Delta \text{PSE}_i^s$  being the change at location  $i$  for subject  $s$ .

We will fit one such model for each of the morph types, and one for each session.

## Model fitting

```

require(lme4)

# define some functions

```

```

extract_morph_session <- function(df, mt, ses) {
  # Extract trials from one particular morph_type and session
  df_ <-
    df %>%
    filter(morph_type == mt, session == ses) %>%
    mutate(morph_resc=(morph - 50)/100)
  return(df_)
}
run_model_session <- function(df) {
  # Run the following logit mixed-effect model for one session
  # response_bin ~ morph_resc + pos - 1 + (pos - 1 | subject)
  m <- glmer(response_bin ~ morph_resc + pos - 1 + (pos - 1 | subject),
    data=df,
    family=binomial(link='logit'),
    control=glmerControl(optimizer='bobyqa', optCtrl=list(maxfun=20000)))
  return(m)
}

```

Run the model separately for every morph

```

morphs <- unique(df$morph_type)
df_ses1 <- sapply(morphs, function(x) extract_morph_session(df, x, '1'),
  simplify=F, USE.NAMES=T)
df_ses2 <- sapply(morphs, function(x) extract_morph_session(df, x, '2'),
  simplify=F, USE.NAMES=T)
# check that we have all the trials
assert_that(sum(sapply(df_ses1, nrow)) + sum(sapply(df_ses2, nrow)) == nrow(df))

## [1] TRUE

# now compute models
model_ses1 <- sapply(df_ses1, run_model_session, simplify=F, USE.NAMES=T)
model_ses2 <- sapply(df_ses2, run_model_session, simplify=F, USE.NAMES=T)

```

The following functions are used to extract both the population  $PSE^p$  and the subject  $\Delta PSE^s$ . Remember that  $PSE^s = PSE^p + \Delta PSE^s$ .

```

population_pse <- function(model) {
  # Computes population-level PSE (see equations above)
  # Note: they are on the scale of morph_resc
  morph_beta <- fixef(model)[1]
  pos_betas <- fixef(model)[-1]
  return(-pos_betas/morph_beta)
}

subjects_pse <- function(model) {
  # Computes subject-level PSE (see equations above)
  morph_pop <- fixef(model)[1]
  position_pop <- fixef(model)[-1]
  ranef_model <- ranef(model)$subject
  # this is the denominator
  morph_subj <- ranef_model[, 1] + morph_pop
  # these are the numerators
  position_subj <- ranef_model[, -1]
  # extend position_beta to get the same shape as ranef_pos
  position_pop <- matrix(rep(position_beta, nrow(ranef_pos)),

```

```

                                byrow=T, nrow=nrow(ranef_pos))
position_subj <- position_subj + position_pop
# extend morph_subj to get the same shape as position_subj
morph_subj <- matrix(rep(morph_subj, ncol(position_subj)), ncol=ncol(position_subj))
return(-position_subj/morph_subj)
}

delta_pse <- function(model) {
  # Computes subject-level PSE (see equations above)
  # Note: they are on the scale of morph_resc
  # IF 'morph_resc' is entered as a random effect, add that for each individual
  # morph
  morph_beta <- fixef(model)[1]
  ranef_model <- ranef(model)$subject
  if ('morph_resc' %in% names(ranef_model)) {
    pse_pop <- population_pse(model)
    pse_subj <- subjects_pse(model)
    pse_pop <- matrix(rep(pse_pop, nrow(pse_subj)), nrow=nrow(pse_subj), byrow=T)
    return(pse_subj - pse_pop)
  } else {
    return(-ranef_model/morph_beta)
  }
}

```

Let's look at the population estimates for the PSEs across the two sessions, as well as the subject-level estimates.

```

psep_ses1 <- sapply(model_ses1, population_pse, simplify=F, USE.NAMES=T)
psep_ses2 <- sapply(model_ses2, population_pse, simplify=F, USE.NAMES=T)
dpse_ses1 <- sapply(model_ses1, delta_pse, simplify=F, USE.NAMES=T)
dpse_ses2 <- sapply(model_ses2, delta_pse, simplify=F, USE.NAMES=T)

```

## Predict psychometric curves

Let's plot the population estimates first

```

df_predict <-
  expand.grid(morph_resc=seq(-0.5, 0.5, 0.01), pos=c('1', '3', '5', '7'))

predict_pop_ses1 <- sapply(model_ses1, predict, newdata=df_predict,
  re.form=NA, type='response', simplify=F, USE.NAMES=T)
predict_pop_ses2 <- sapply(model_ses2, predict, newdata=df_predict,
  re.form=NA, type='response', simplify=F, USE.NAMES=T)

# add df_predict to each of them
predict_pop_ses1 <- lapply(predict_pop_ses1, function(x) cbind(df_predict, pred=x))
predict_pop_ses2 <- lapply(predict_pop_ses2, function(x) cbind(df_predict, pred=x))

predict_pop_ses1 <- ldply(predict_pop_ses1, data.frame)
predict_pop_ses2 <- ldply(predict_pop_ses2, data.frame)
predict_pop_ses1$session <- '1'
predict_pop_ses2$session <- '2'

predict_pop <- rbind(predict_pop_ses1, predict_pop_ses2)

```

```

predict_pop <-
  predict_pop %>%
  mutate(morph=resc*100 + 50,
         pos_num=as.numeric(as.character(pos))*45,
         morph_type=.id)

# modified from https://rpubs.com/Koundy/71792
theme_Publication <- function(base_size=12) {
  library(ggthemes)
  (theme_foundation(base_size=base_size)
   + theme(plot.title = element_text(face = "bold",
                                     size = rel(1.2), hjust = 0.5),

          text = element_text(),
          panel.background = element_rect(colour = NA),
          plot.background = element_rect(colour = NA),
          panel.border = element_rect(colour = NA),
          axis.title = element_text(size = rel(1)),
          axis.title.y = element_text(angle=90,vjust =2),
          axis.title.x = element_text(vjust = -0.2),
          axis.text = element_text(),
          axis.line = element_line(colour="black"),
          axis.ticks = element_line(),
          panel.grid.major = element_blank(), #element_line(colour="#f0f0f0"),
          panel.grid.minor = element_blank(),
          legend.key = element_rect(colour = NA),
          legend.position = "bottom",
          legend.direction = "horizontal",
          #legend.key.size= unit(0.2, "cm"),
          legend.spacing = unit(0, "cm"),
          legend.title = element_text(),
          plot.margin = unit(c(10,5,5,5),"mm"),
          strip.background = element_rect(colour="#f0f0f0",fill="#f0f0f0"),
          strip.text = element_text(face="bold"),
          strip.text.y = element_text(angle = 0)

  ))
}

```

```

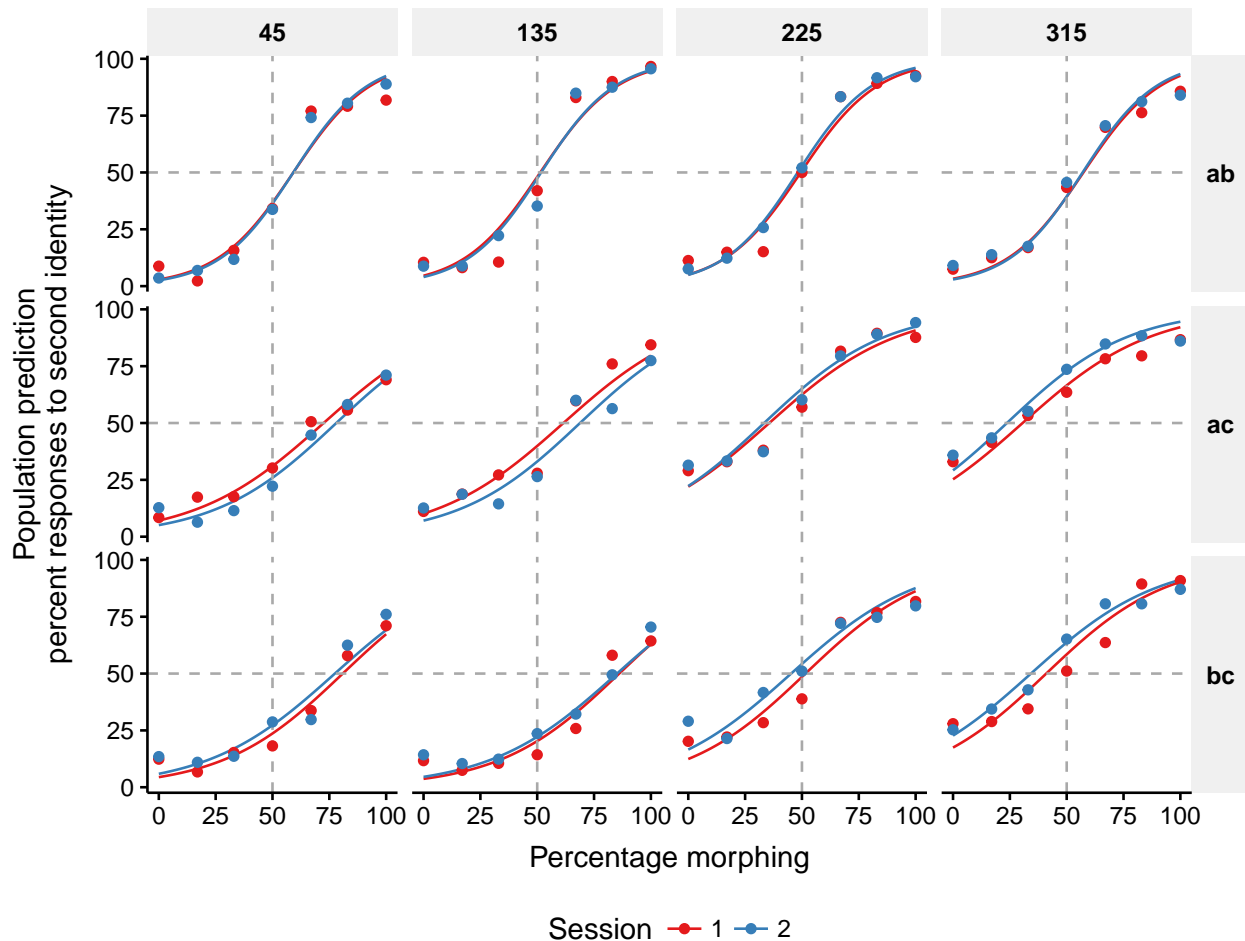
df <-
  df %>%
  mutate(pos_num=as.numeric(as.character(pos))*45)

predict_pop %>%
  ggplot(aes(morph, pred*100, color=session)) +
  geom_line() +
  geom_hline(yintercept=50, size=0.5, color='darkgray', alpha=1, linetype='dashed') +
  geom_vline(xintercept=50, size=0.5, color='darkgray', alpha=1, linetype='dashed') +
  # add individual data
  stat_summary(data=df,
              aes(morph, response_bin),
              fun.y=function(x) sum(x)/length(x)*100, geom='point') +
  facet_grid(morph_type ~ pos_num) +
  labs(x='Percentage morphing', y='Population prediction\npercent responses to second identity', color=
  scale_color_brewer(palette='Set1') +

```

```
theme_Publication() +
coord_equal()
```

```
## Warning: package 'ggthemes' was built under R version 3.2.5
```



```
ggsave('../img/pred_pop_gmm.png', width=8, height=6)
```

Now we can predict for each individual subject

```
df_predict <-
  expand_grid(
    morph_resc=seq(-0.5, 0.5, 0.01),
    pos=c('1', '3', '5', '7'),
    subject=unique(df$subject))

predict_subj_ses1 <- sapply(model_ses1, predict, newdata=df_predict,
                           type='response', simplify=F, USE.NAMES=T)
predict_subj_ses2 <- sapply(model_ses2, predict, newdata=df_predict,
                           type='response', simplify=F, USE.NAMES=T)

# add df_predict to each of them
predict_subj_ses1 <- lapply(predict_subj_ses1, function(x) cbind(df_predict, pred=x))
predict_subj_ses2 <- lapply(predict_subj_ses2, function(x) cbind(df_predict, pred=x))

predict_subj_ses1 <- ldply(predict_subj_ses1, data.frame)
```

```

predict_subj_ses2 <- ldply(predict_subj_ses2, data.frame)
predict_subj_ses1$session <- '1'
predict_subj_ses2$session <- '2'

predict_subj <- rbind(predict_subj_ses1, predict_subj_ses2)
predict_subj <-
  predict_subj %>%
  mutate(morph=resc*morph_resc*100 + 50,
         pos_num=as.numeric(as.character(pos))*45,
         morph_type=.id)

```

Now we can save each individual plot to disk

```

subjects <- unique(df$subject)
for (subj in subjects) {
  out_dir <- file.path('../img', 'pred_gmm')
  fnout <- file.path(out_dir, paste(subj, '_pred_gmm.png', sep=''))
  # setup dataframes for plotting
  this_subject_df <- df %>%
    filter(subject == subj) %>%
    # add position in angles
    mutate(pos_num=as.numeric(as.character(pos))*45)
  this_predict_subj <- predict_subj %>%
    filter(subject == subj) %>%
    # add position in angles
    mutate(pos_num=as.numeric(as.character(pos))*45)

  plot_curve <-
    this_predict_subj %>%
    # add prediction
    ggplot(aes(morph, pred*100, color=session)) +
    geom_hline(yintercept=50, size=0.5, color='darkgray', alpha=1, linetype='dashed') +
    geom_vline(xintercept=50, size=0.5, color='darkgray', alpha=1, linetype='dashed') +
    geom_line() +
    # add individual data
    stat_summary(data=this_subject_df,
                 aes(morph, response_bin),
                 fun.y=function(x) sum(x)/length(x)*100, geom='point') +
    facet_grid(morph_type ~ pos_num) +
    labs(x='Percentage morphing', y='Percent responses to second identity', color='Session') +
    scale_color_brewer(palette='Set1') +
    ggtitle(paste("Subject", subj)) +
    theme_Publication() +
    coord_equal()

  # save
  dir.create(out_dir, recursive=T)
  ggsave(filename=fnout, plot=plot_curve, width=8, height=6)
}

```

```

## Warning in dir.create(out_dir, recursive = T): '../img/pred_gmm' already
## exists

```

```

## Warning in dir.create(out_dir, recursive = T): '../img/pred_gmm' already

```

```
## exists

## Warning in dir.create(out_dir, recursive = T): '../img/pred_gmm' already
## exists

## Warning in dir.create(out_dir, recursive = T): '../img/pred_gmm' already
## exists

## Warning in dir.create(out_dir, recursive = T): '../img/pred_gmm' already
## exists

## Warning in dir.create(out_dir, recursive = T): '../img/pred_gmm' already
## exists

## Warning in dir.create(out_dir, recursive = T): '../img/pred_gmm' already
## exists

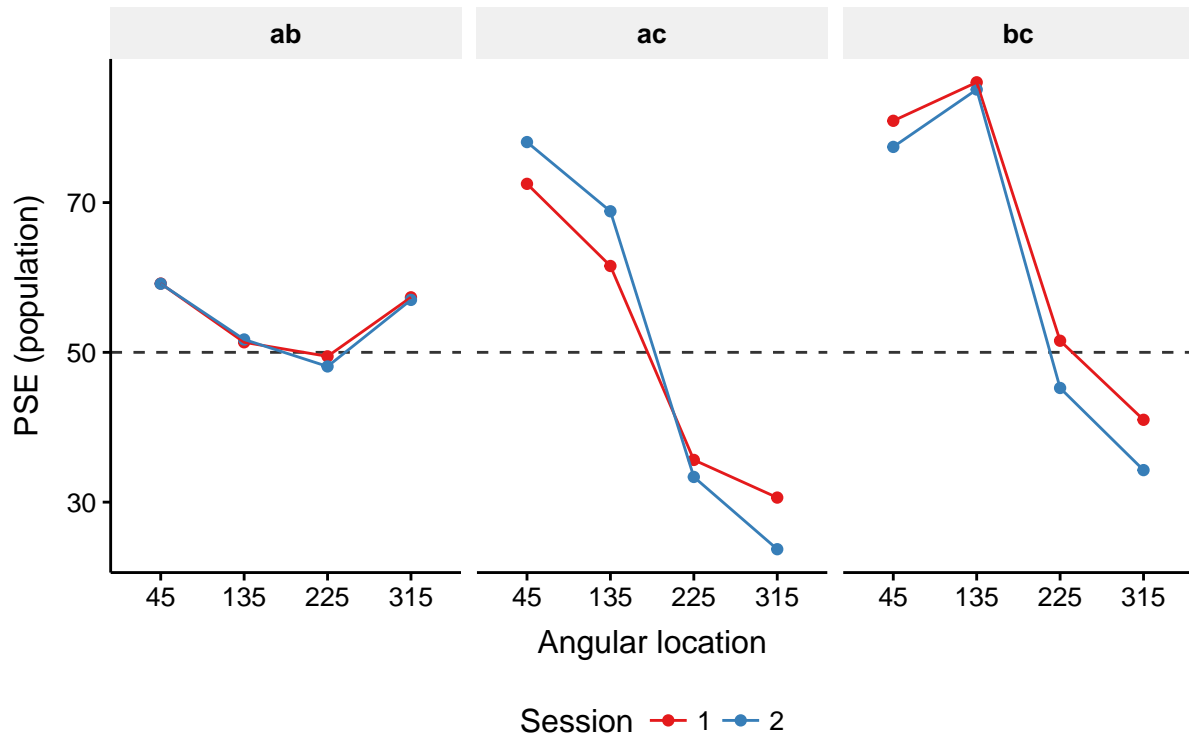
## Warning in dir.create(out_dir, recursive = T): '../img/pred_gmm' already
## exists

## Warning in dir.create(out_dir, recursive = T): '../img/pred_gmm' already
## exists
```

These are the population-level estimates

```
psep %>%
  ggplot(aes(pos_num, pse*100 + 50, color=session, group=session)) +
  geom_hline(yintercept=50, linetype='dashed', alpha=0.8) +
  geom_point() +
  geom_line() +
  labs(x='Angular location', y='PSE (population)', color='Session') +
  #ggtitle('Population-level PSE') +
  scale_color_brewer(palette='Set1') +
  facet_grid(~morph_type) +
  scale_y_continuous(breaks=c(30, 50, 70)) +
  theme_Publication()
```



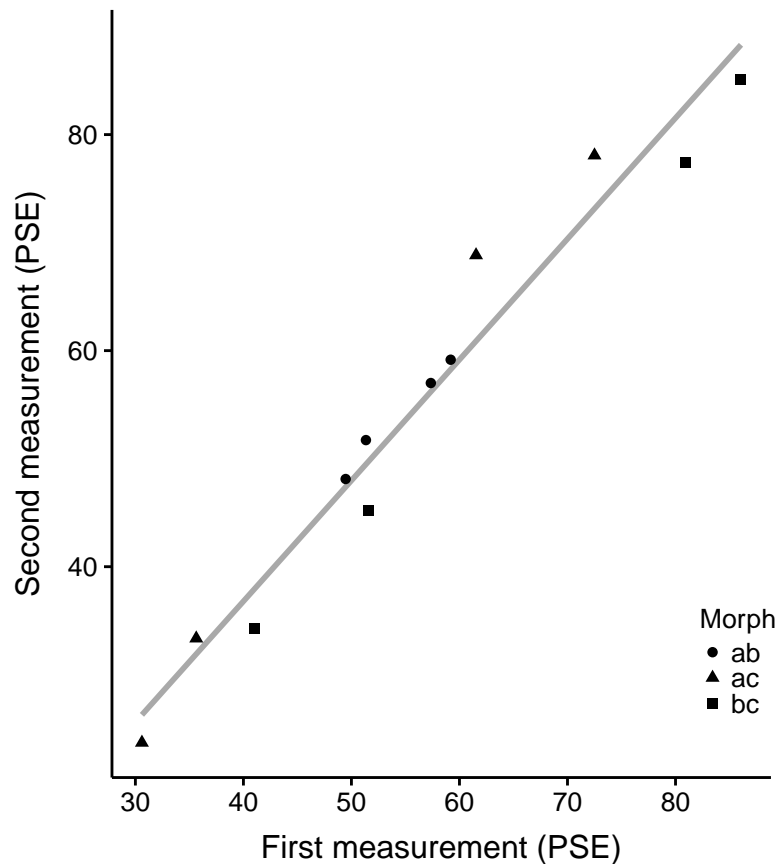


##

Stability of population level estimates across sessions Let's compute a correlation across sessions

```
psep_wide <-
psep %>%
  spread(session, pse) %>%
  mutate(session1=.$'1', session2=.$'2')

ggplot(psep_wide, aes(session1*100 + 50, session2*100 + 50, shape=morph_type, group=1)) +
  geom_smooth(method='lm', color='darkgray', se=F) +
  geom_point() +
  labs(x='First measurement (PSE)', y='Second measurement (PSE)', shape='Morph') +
  theme_Publication() +
  coord_equal() +
  #guides(shape=F)
  theme(legend.position=c(0.95, 0.15),
        legend.direction='vertical',
        legend.key.size=unit(.8, 'picas'),
        legend.title=element_text(size=10))
```



```
#coord_equal(xlim=c(20, 95), ylim=c(20, 95)) +
#scale_x_continuous(breaks=seq(20, 90, 10)) +
#scale_y_continuous(breaks=seq(20, 90, 10))

ggsave('../img/pse_pop_scatter.png', width=5, height=5)
```

And these are the correlations

```
set.seed(3432)
bootES(psep_wide[c('session1', 'session2')], R=10000)

##
## 95.00% bca Confidence Interval, 10000 replicates
## Stat      CI (Low)    CI (High)    bias      SE
## 0.979      0.929      0.992      0.000     0.011

cor.test(psep_wide$session1, psep_wide$session2)

##
## Pearson's product-moment correlation
##
## data:  psep_wide$session1 and psep_wide$session2
## t = 15.219, df = 10, p-value = 3.042e-08
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
##  0.9248693 0.9942948
## sample estimates:
##      cor
```

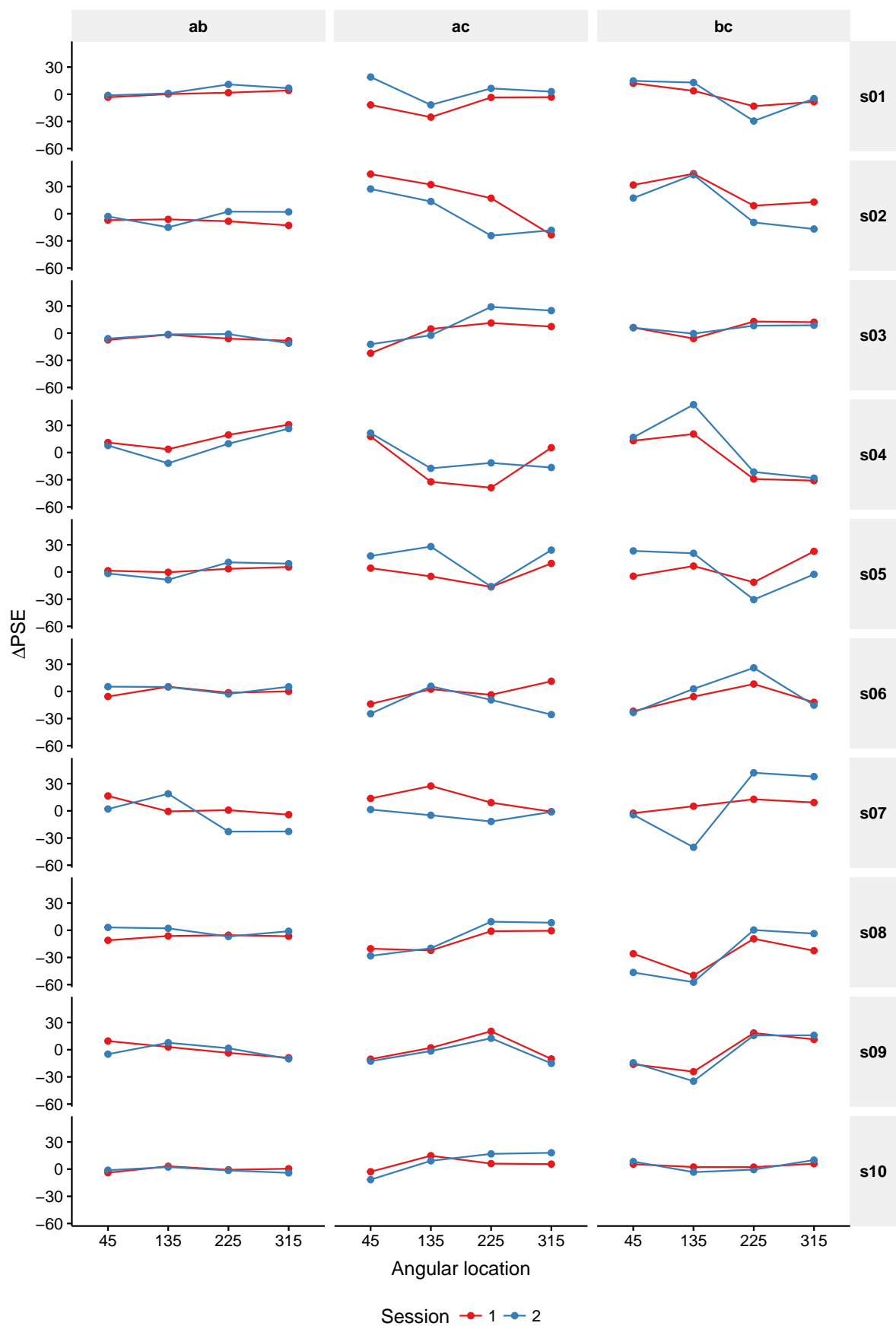
```
## 0.9790871
```

They are very consistent across sessions.

## Stability of subject-level estimates across sessions

```
# add morph type
for (morph in morphs) {
  dpse_ses1[[morph]]$subject <- row.names(dpse_ses1[[morph]])
  dpse_ses2[[morph]]$subject <- row.names(dpse_ses2[[morph]])
}
# get dpse in long format for plotting
dpse_ses1_long <-
  ldply(
    lapply(dpse_ses1, function(x) gather(x, pos, pse, -subject)),
    data.frame) %>%
  mutate(morph_type=.id, session='1')
dpse_ses2_long <-
  ldply(
    lapply(dpse_ses2, function(x) gather(x, pos, pse, -subject)),
    data.frame) %>%
  mutate(morph_type=.id, session='2')

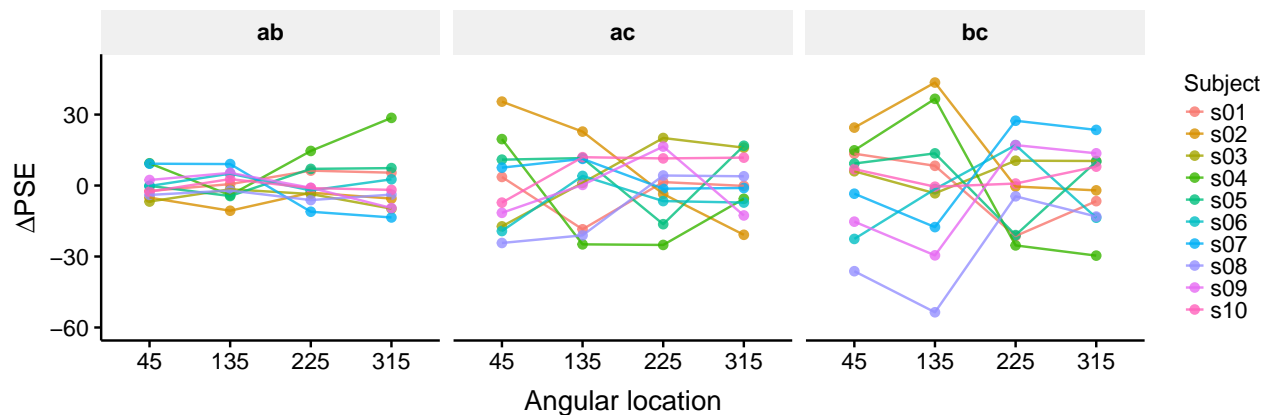
dpse <-
  rbind(dpse_ses1_long, dpse_ses2_long) %>%
  mutate(pos_num=mapvalues(pos,
    paste('pos', c(1, 3, 5, 7), sep=''),
    c(1, 3, 5, 7)*45))
dpse$pos_num <- factor(dpse$pos_num, levels=c(1, 3, 5, 7)*45)
```



Aver-

age the estimates across sessions and plot them

```
dpse %>%
  group_by(subject, morph_type, pos_num) %>%
  summarise(pse=mean(pse)) %>%
  ggplot(aes(pos_num, pse*100, color=subject, group=subject)) +
  geom_point(alpha=0.8) +
  geom_line(alpha=0.8) +
  facet_wrap(~morph_type, nrow=1) +
  theme_Publication() +
  labs(x='Angular location', y=TeX('$\\Delta$PSE'), color='Subject') +
  scale_y_continuous(limits=c(-60, 50)) +
  theme(#legend.position=c(0.95, 0.15),
        legend.direction='vertical',
        legend.position='right',
        legend.key.size=unit(.8, 'picas'),
        legend.title=element_text(size=10))
```



```
ggsave(' ../img/dpse_subjects.png', width=8, height=3)
```

Let's also plot an example fit for subject s10

```
# get the data to plot individual points
df_s10_plot_ab <-
  df %>%
  filter(subject == 's10', session == '1') %>%
  filter(morph_type == 'ab', pos_num %in% c('225', '315'))
df_s10_plot_ac <-
  df %>%
  filter(subject == 's10', session == '1') %>%
  filter(morph_type == 'ac', pos_num %in% c('135', '315'))
df_s10_plot_bc <-
  df %>%
  filter(subject == 's10', session == '1') %>%
  filter(morph_type == 'bc', pos_num %in% c('135', '315'))

df_s10_plot <-
  rbind(df_s10_plot_ab, df_s10_plot_ac, df_s10_plot_bc)

# now get the pse values to plot
psep1_plot <-
psep %>%
```

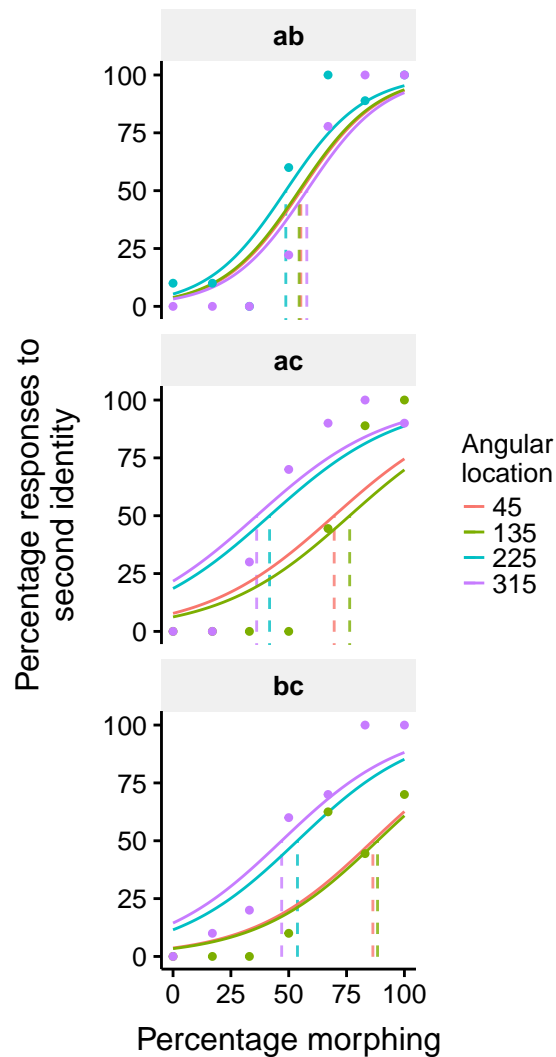
```

  filter(session == '1') %>%
  arrange(pos, morph_type)
dpse1_plot <-
dpse %>%
  filter(subject == 's10', session == '1') %>%
  arrange(pos, morph_type) %>%
  mutate(dpse=pse) %>%
  select(morph_type, pos, pos_num, dpse)

pse_s10_plot <- merge(psep1_plot, dpse1_plot)
pse_s10_plot <-
pse_s10_plot %>%
  mutate(tpse=(pse+dpse)*100+50)

predict_subj %>%
  filter(subject == 's10', session == '1') %>%
  ggplot(aes(morph_resc*100 + 50, pred*100, color=as.factor(pos_num))) +
  geom_segment(aes(x=tpse, xend=tpse, y=-10, yend=50), alpha=0.8, linetype='dashed', data=pse_s10_plot)
  geom_line() +
  facet_wrap(~morph_type, ncol=1) +
  coord_equal(ylim=c(-0.4, 101)) +
  theme_Publication() +
  theme(#legend.position=c(0.95, 0.15),
        legend.direction='vertical',
        legend.position='right',
        legend.key.size=unit(.8, 'picas'),
        legend.title=element_text(size=10)) +
  labs(x='Percentage morphing', y='Percentage responses to\nsecond identity', color='Angular\nlocation')
  stat_summary(aes(morph, response_bin), fun.y=function(x) sum(x)/length(x)*100, geom='point',
               data=df_s10_plot, size=0.9, show.legend=F)

```

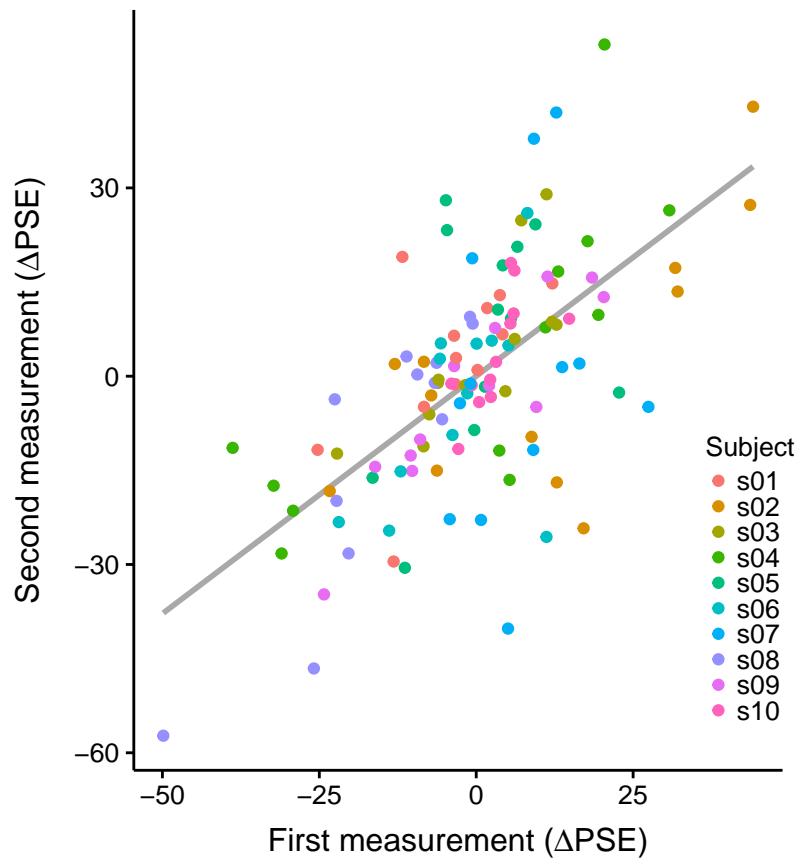


```
ggsave('../img/examplefit_s10.png', width=8, height=6)
```

Correlation of subject-level estimates across sessions

```
dpse_wide <-
dpse %>%
  spread(session, pse) %>%
  mutate(session1=.$'1', session2=.$'2')

ggplot(dpse_wide, aes(session1*100, session2*100, color=subject)) +
  geom_smooth(method='lm', color='darkgray', se=F) +
  geom_point() +
  labs(x=TeX('First measurement ( $\Delta$ PSE)'),
       y=TeX('Second measurement ( $\Delta$ PSE)'),
       color='Subject') +
  theme_Publication() +
  coord_equal() +
  theme(legend.position=c(0.95, 0.25),
        legend.direction='vertical',
        legend.key.size=unit(.8, 'picas'),
        legend.title=element_text(size=10))
```



```
#coord_equal(xlim=c(-60, 45), ylim=c(-60, 45)) +
#scale_x_continuous(breaks=seq(-60, 40, 20)) +
#scale_y_continuous(breaks=seq(-60, 40, 20))

ggsave('../img/pse_subj_scatter.png', width=5, height=5)
```

And these are the correlations.

```
set.seed(23448)
bootES(dpse_wide[c('session1', 'session2')], R=10000)

##
## 95.00% bca Confidence Interval, 10000 replicates
## Stat      CI (Low)    CI (High)    bias      SE
## 0.639      0.499      0.750      -0.003    0.064

cor.test(dpse_wide$session1, dpse_wide$session2)

##
## Pearson's product-moment correlation
##
## data:  dpse_wide$session1 and dpse_wide$session2
## t = 9.0205, df = 118, p-value = 3.997e-15
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
##  0.5190456 0.7340393
## sample estimates:
##      cor
```



```
## 0.6388518
```

## Within- vs. between-subjects correlation

Let's compute the correlation between the first and the second session

```
cor_ses12 <- list()
for (morph in morphs) {
  this_cor <- cor(t(dpse_ses1[[morph]][, 1:4]), t(dpse_ses2[[morph]][, 1:4]))
  # make it symmetric
  this_cor <- (this_cor + t(this_cor))/2.
  cor_ses12[[morph]] = this_cor
}

# make a dataframe in long format
make_cor_long <- function(cor) {
  within <- diag(cor)
  between <- cor[lower.tri(cor)]

  df_within_between <- data.frame(corr=c(within, between),
                                  type=c(rep('within', length(within)),
                                         rep('between', length(between))))
  return(df_within_between)
}

cor_ses12_long <- sapply(cor_ses12, make_cor_long, simplify=F, USE.NAMES=T)
# add everything together for plotting
cor_ses12_long_plot <- ldply(cor_ses12_long, data.frame)
```

We can check whether the estimates are consistent across sessions, and also subject-specific, by comparing the within-subject correlations with the between-subject correlations. We will compute the bootstrapped difference Within – Between.

```
require(bootES)
require(broom)
```

```
## Loading required package: broom
```

```
## Warning: package 'broom' was built under R version 3.2.5
```

```
bootstrap_withinbetween <- function(corr_df) {
  b <- bootES(corr_df,
              data.col='corr', group.col='type',
              contrast=c(within=1, between=-1), R=10000)
  return(b)
}

set.seed(124)
boot_cis <- sapply(cor_ses12_long,
                  bootstrap_withinbetween,
                  simplify=F,
                  USE.NAMES=T)

extract_cis <- function(bootes_out) {
```

```

t0 <- boot_out$t0
bounds <- boot_out$bounds
df <- data.frame(t0=t0, lci=bounds[1], rci=bounds[2])
return(df)
}

extract_distribution <- function(boot_out) {
  df <- data.frame(t=boot_out$t)
  return(df)
}

boot_cis_df <- ldply(boot_cis, extract_cis)
boot_dist_df <- ldply(boot_cis, extract_distribution)

```

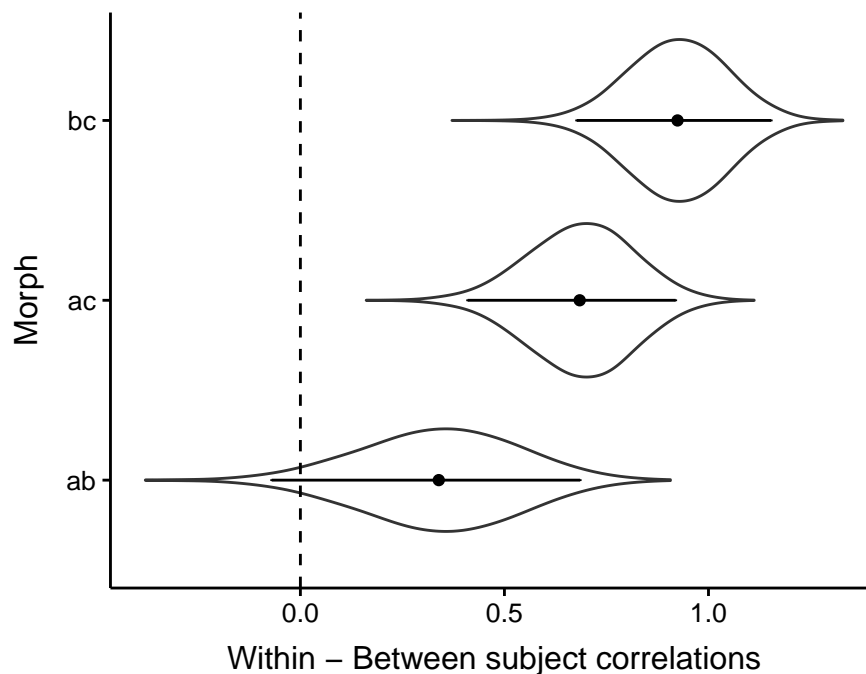
And finally plot them

```

ggplot(data=boot_dist_df, aes(.id, t)) +
  geom_violin(adjust=2) +
  geom_errorbar(data=boot_cis_df, aes(ymin=lci, ymax=rci, y=t0), width=0.01) +
  geom_point(data=boot_cis_df, aes(y=t0)) +
  geom_hline(yintercept=0, linetype='dashed') +
  labs(x='Morph', y='Within - Between subject correlations') +
  theme_Publication() +
  coord_flip() +
  theme(aspect.ratio=3/4)

```

## Warning: Ignoring unknown aesthetics: y



And these are the values

```
boot_cis_df %>% arrange(.id)
```

```

##   .id      t0      lci      rci
## 1  ab 0.3391053 -0.07029715 0.6855457
## 2  ac 0.6843746  0.40988453 0.9196337
## 3  bc 0.9241400  0.67701408 1.1536785

```

Let's also do it separately for within and between

```
bootstrap <- function(corr_df) {  
  b <- bootES(corr_df,  
              data.col='corr', R=10000)  
  return(b)  
}  
  
cor_ses12_long_within <- sapply(cor_ses12_long,  
                                function(x) filter(x, type=='within'),  
                                simplify=F,  
                                USE.NAMES=T)  
cor_ses12_long_between <- sapply(cor_ses12_long,  
                                  function(x) filter(x, type=='between'),  
                                  simplify=F,  
                                  USE.NAMES=T)  
  
set.seed(3243)  
bs_within <- ldply(  
  lapply(cor_ses12_long_within, bootstrap),  
  extract_cis)  
set.seed(23423)  
bs_between <- ldply(  
  lapply(cor_ses12_long_between, bootstrap),  
  extract_cis)
```

```
bs_within %>% arrange(.id)
```

```
##   .id      t0      lci      rci  
## 1  ab 0.3189388 -0.1007527 0.6151277  
## 2  ac 0.6179501  0.3513429 0.7855175  
## 3  bc 0.8490889  0.6115145 0.9463881
```

```
bs_between %>% arrange(.id)
```

```
##   .id      t0      lci      rci  
## 1  ab -0.02016651 -0.1510517 0.10897154  
## 2  ac -0.06642447 -0.2118914 0.08072318  
## 3  bc -0.07505119 -0.2658810 0.12166054
```

## Correlation with familiarity ratings

```
df_quest <- read_csv('../data/questionnaire.csv',  
                     col_types='ccdddddddddd')
```

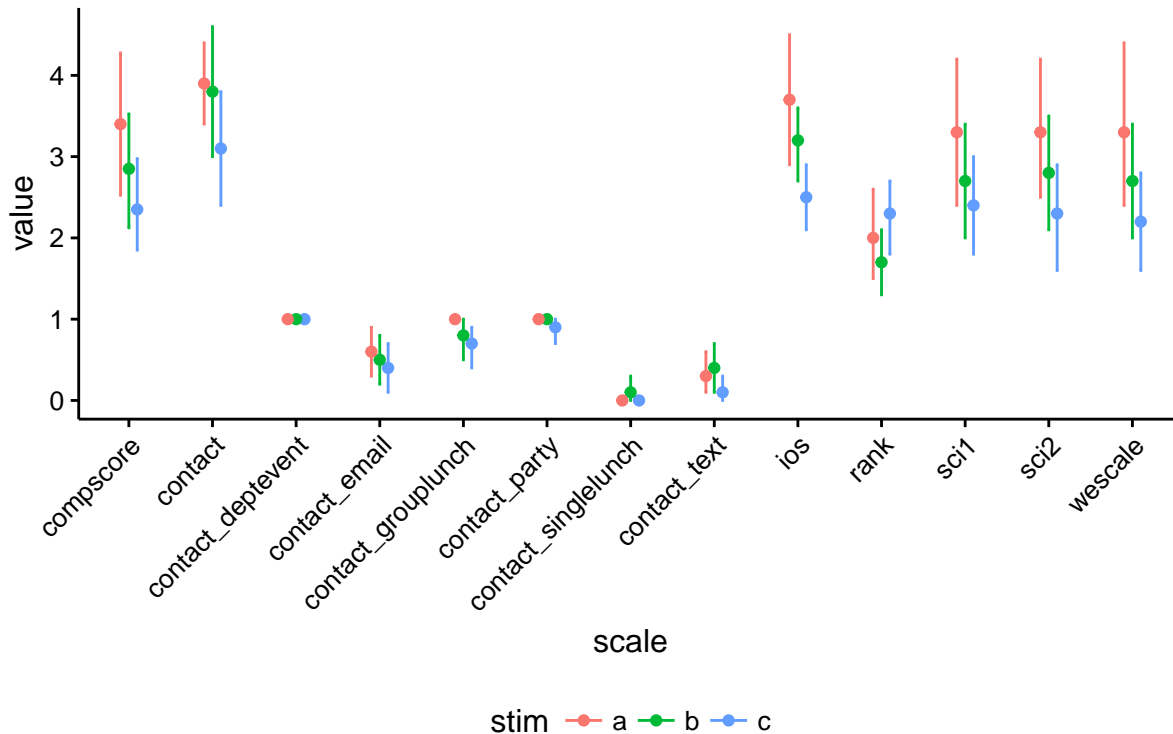
We are going to create a composite score by simply averaging the questions related to closeness.

```
# add composite score  
df_quest <-  
df_quest %>%  
  # everybody got the name right  
  select(-name) %>%  
  rowwise() %>%  
  # Let's make a composite score by averaging ios, wescale, sci1, sci2  
  mutate(compscore=mean(c(ios, wescale, sci1, sci2)))
```

Let's start simply by looking at the average values for each id

```
df_quest_long <-
df_quest %>%
  gather(scale, value, -subject, -stim)

df_quest_long %>%
  ggplot(aes(scale, value, color=stim)) +
  stat_summary(fun.data=mean_cl_boot, geom='errorbar', width=0.1, position=position_dodge(w=0.3)) +
  stat_summary(fun.y=mean, geom='point', position=position_dodge(w=0.3)) +
  theme_Publication() +
  theme(axis.text.x=element_text(angle=45, hjust=1))
```



```
ggsave('..img/scores.png', width=8, height=6)

# compute a score of deviance from 0 for each subject and morph type
# basically the variance
dpse_deviance <-
dpse %>%
  group_by(subject, morph_type, pos_num) %>%
  summarise(pse=mean(pse)) %>%
  group_by(subject, morph_type) %>%
  summarise(psescore=sum(pse^2))

# now for each subject compute the average score for the morphs
df_quest_avg <-
df_quest %>%
  select(subject, stim, compscore) %>%
  spread(stim, compscore) %>%
  rowwise() %>%
  mutate(score_ab=mean(c(a, b)),
         score_bc=mean(c(b, c)),
```

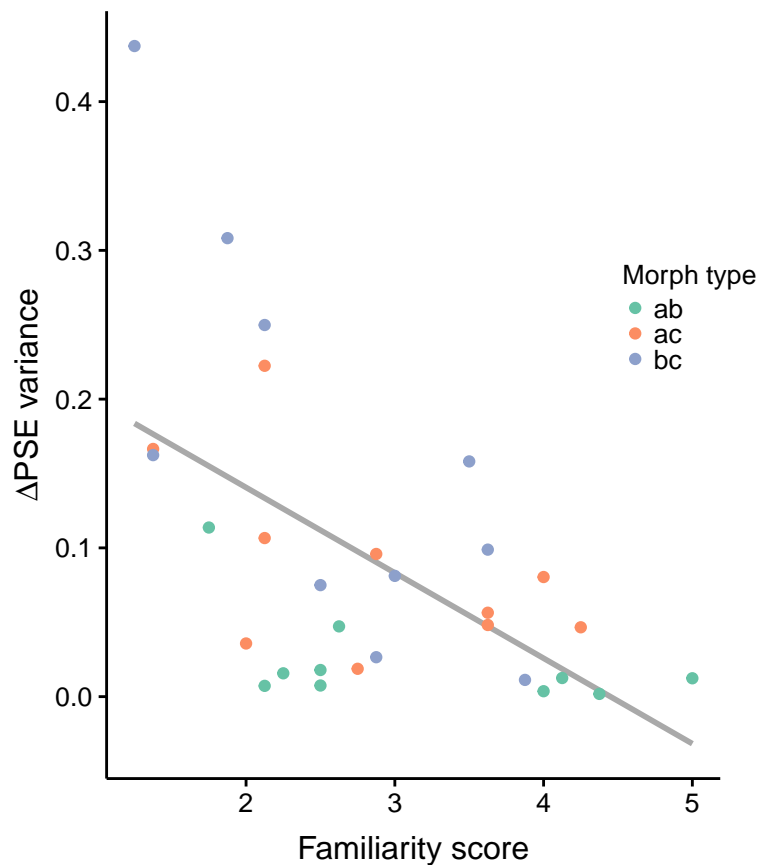
```

    score_ac=mean(c(a, c))) %>%
  select(-a, -b, -c) %>%
  gather(morph_type, score, -subject)

df_quest_avg$morph_type <-
  mapvalues(df_quest_avg$morph_type,
            c('score_ab', 'score_ac', 'score_bc'),
            c('ab', 'ac', 'bc'))
df_quest_deviance <-
df_quest_avg %>%
  arrange(subject, morph_type) %>%
  merge(dpse_deviance)

ggplot(aes(score, psescore, color=morph_type), data=df_quest_deviance) +
  geom_smooth(method='lm', aes(group=1), se=F, color='darkgray') +
  geom_point() +
  theme_Publication() +
  coord_fixed(ratio=10) +
  labs(x='Familiarity score', y=TeX('$\\Delta$PSE variance'), color='Morph type') +
  theme(legend.position=c(0.95, 0.6),
        legend.direction='vertical',
        legend.key.size=unit(.8, 'picas'),
        legend.title=element_text(size=10)) +
  scale_color_brewer(palette='Set2')

```



```
ggsave(' ../img/psequest_scatter.png', width=5, height=5)

cor.test(df_quest_deviance$score, df_quest_deviance$psescore)

##
## Pearson's product-moment correlation
##
## data: df_quest_deviance$score and df_quest_deviance$psescore
## t = -3.5895, df = 28, p-value = 0.001248
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## -0.7666035 -0.2520935
## sample estimates:
## cor
## -0.5613776

set.seed(324)
bootES(df_quest_deviance[c('score', 'psescore')], R=10000)

##
## 95.00% bca Confidence Interval, 10000 replicates
## Stat CI (Low) CI (High) bias SE
## -0.561 -0.708 -0.298 0.006 0.100
```

## Is it personal familiarity or is it contact?

```
df_quest_contact_avg <-
df_quest %>%
  select(subject, stim, contact) %>%
  spread(stim, contact) %>%
  rowwise() %>%
  mutate(ab=mean(c(a, b)),
         bc=mean(c(b, c)),
         ac=mean(c(a, c))) %>%
  select(-a, -b, -c) %>%
  gather(morph_type, contact, -subject)

df_contact_deviance <-
df_quest_deviance %>%
  arrange(subject, morph_type) %>%
  merge(df_quest_contact_avg)
```

Are familiarity score and contact correlated?

```
cor.test(df_contact_deviance$score, df_contact_deviance$contact)

##
## Pearson's product-moment correlation
##
## data: df_contact_deviance$score and df_contact_deviance$contact
## t = 2.6517, df = 28, p-value = 0.01304
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## 0.1046298 0.6959508
```

```
## sample estimates:
##      cor
## 0.4480113

set.seed(324)
bootES(df_contact_deviance[c('score', 'contact')], R=10000)

##
## 95.00% bca Confidence Interval, 10000 replicates
## Stat      CI (Low)    CI (High)    bias      SE
## 0.448      0.168      0.675      -0.002    0.130
```

Run a model that predicts the variance of  $\Delta$ PSE using only the familiarity score and then also the contact score.

```
m1 <- lm(psescore ~ score, data=df_contact_deviance)
m2 <- update(m1, . ~ . + contact)

tidy(m1)
```

	term	estimate	std.error	statistic	p.value
## 1	(Intercept)	0.25543170	0.04848293	5.268487	1.331582e-05
## 2	score	-0.05740843	0.01599341	-3.589506	1.248105e-03

```
tidy(m2)
```

	term	estimate	std.error	statistic	p.value
## 1	(Intercept)	0.36532810	0.06236289	5.858101	3.077435e-06
## 2	score	-0.03894801	0.01640093	-2.374744	2.492620e-02
## 3	contact	-0.04522674	0.01800160	-2.512374	1.827090e-02

```
anova(m1, m2)
```

```
## Analysis of Variance Table
##
## Model 1: psescore ~ score
## Model 2: psescore ~ score + contact
##   Res.Df    RSS Df Sum of Sq   F Pr(>F)
## 1      28 0.20880
## 2      27 0.16924   1  0.039565 6.312 0.01827 *
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

```
require(heplots)
```

```
## Loading required package: heplots
## Warning: package 'heplots' was built under R version 3.2.5
## Loading required package: car
## Warning: package 'car' was built under R version 3.2.5
##
## Attaching package: 'car'
## The following object is masked from 'package:dplyr':
##
##      recode
## The following object is masked from 'package:purrr':
```

```
##
##      some
## The following object is masked from 'package:boot':
##
##      logit
round(etasq(m1), 2)

##          Partial eta^2
## score          0.32
## Residuals      NA
round(etasq(m2), 2)

##          Partial eta^2
## score          0.17
## contact        0.19
## Residuals      NA

# from http://www.statmethods.net/advstats/bootstrapping.html
bs <- function(formula, data, indices) {
  d <- data[indices,] # allows boot to select sample
  fit <- lm(formula, data=d)
  return(coef(fit))
}

set.seed(234)
results <- boot(data=df_contact_deviance, statistic=bs,
  R=10000, formula=psescore ~ score + contact)

t0 <- results$t0
conf <- c()
for (i in 1:length(t0)) {
  b <- boot.ci(results, type='bca', index=i)
  conf <- rbind(conf, b$bca[4:5])
}

df_ci <- data.frame(t0=t0, lci=conf[,1], rci=conf[,2])

df_ci

##          t0          lci          rci
## (Intercept) 0.36532810 0.21465489 0.55147559
## score      -0.03894801 -0.07431666 -0.01631785
## contact     -0.04522674 -0.08609399 -0.01694056
```

Now compute partial correlations

```
partialout_variable <- function(df, x, y, z) {
  mx <- lm(df[[x]] ~ df[[z]])
  my <- lm(df[[y]] ~ df[[z]])
  df_out <- data.frame(mx$residuals, my$residuals)
  names(df_out) <- c(x, y)
  return(df_out)
}

# Make plots of partial correlation with residuals
```



```

partial_contact <- partialout_variable(df_contact_deviance, "pscore", "score", "contact")
partial_score <- partialout_variable(df_contact_deviance, "pscore", "contact", "score")
cor.test(partial_contact$pscore, partial_contact$score)

##
## Pearson's product-moment correlation
##
## data: partial_contact$pscore and partial_contact$score
## t = -2.4183, df = 28, p-value = 0.02235
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## -0.67487275 -0.06515499
## sample estimates:
## cor
## -0.4156671

cor.test(partial_score$pscore, partial_score$contact)

##
## Pearson's product-moment correlation
##
## data: partial_score$pscore and partial_score$contact
## t = -2.5585, df = 28, p-value = 0.01621
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## -0.68771457 -0.08898003
## sample estimates:
## cor
## -0.4352951

set.seed(3242)
bootES(partial_contact[c('pscore', 'score')], R=10000)

##
## 95.00% bca Confidence Interval, 10000 replicates
## Stat CI (Low) CI (High) bias SE
## -0.416 -0.610 -0.158 0.008 0.116

bootES(partial_score[c('pscore', 'contact')], R=10000)

##
## 95.00% bca Confidence Interval, 10000 replicates
## Stat CI (Low) CI (High) bias SE
## -0.435 -0.619 -0.166 -0.001 0.113

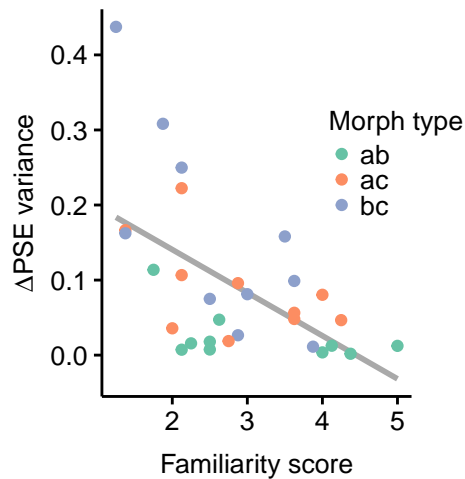
# add the morph types to partial_contact and partial_score
partial_contact$morph_type <- df_contact_deviance$morph_type
partial_score$morph_type <- df_contact_deviance$morph_type

# just a theme to uniform scatters
theme_Publication_scatter <- function(base_size=12) {
  (theme_Publication(base_size=base_size) +
   theme(legend.position=c(0.95, 0.6),
         legend.direction='vertical',
         legend.key.size=unit(.8, 'picas'),
         legend.title=element_text(size=10),
         axis.title=element_text(size=10)))
}

```

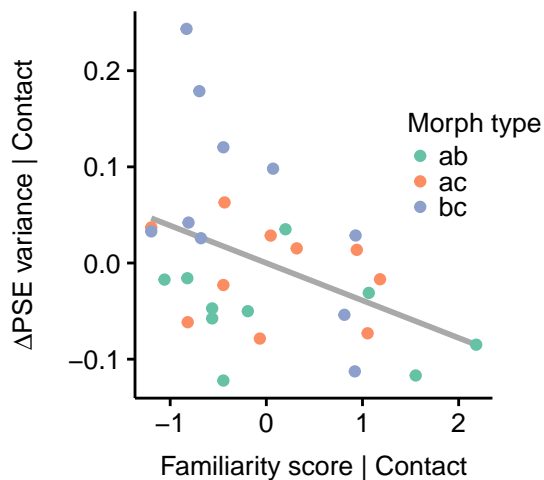
```
}
```

```
ggplot(aes(score, psescore, color=morph_type), data=df_quest_deviance) +
  geom_smooth(method='lm', aes(group=1), se=F, color='darkgray') +
  geom_point() +
  coord_fixed(ratio=10) +
  labs(x='Familiarity score', y=TeX('$\\Delta$PSE variance'), color='Morph type') +
  theme_Publication_scatter() +
  scale_color_brewer(palette='Set2')
```



```
ggsave(' ../img/psequest_scatter_psefam.png', width=3, height=3)
```

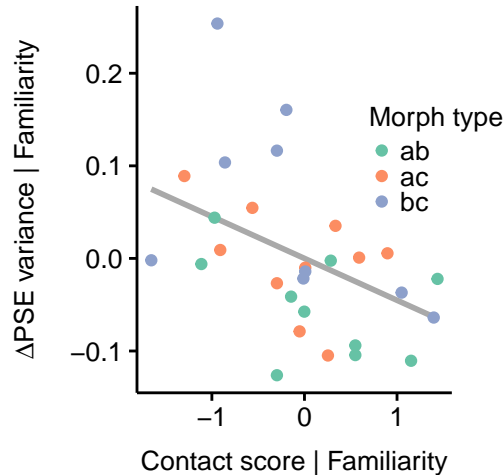
```
ggplot(aes(score, psescore, color=morph_type), data=partial_contact) +
  geom_smooth(method='lm', aes(group=1), se=F, color='darkgray') +
  geom_point() +
  coord_fixed(ratio=10) +
  labs(x='Familiarity score | Contact', y=TeX('$\\Delta$PSE variance | Contact'), color='Morph type') +
  theme_Publication_scatter() +
  scale_color_brewer(palette='Set2')
```



```
ggsave(' ../img/psequest_scatter_psefam_contact.png', width=3, height=3)
```

```
ggplot(aes(contact, psescore, color=morph_type), data=partial_score) +
  geom_smooth(method='lm', aes(group=1), se=F, color='darkgray') +
```

```
geom_point() +
coord_fixed(ratio=10, expand=T) +
labs(x='Contact score | Familiarity', y=TeX('$\\Delta$PSE variance | Familiarity'), color='Morph type') +
theme_Publication_scatter() +
scale_color_brewer(palette='Set2')
```



```
ggsave('../img/psequest_scatter_psecontact_fam.png', width=3, height=3)
```

## PSE for individual identities

We are going to evaluate the bias towards specific identities, instead of pairs of morphs. To do so, we'll fit a linear model predicting the PSE, taking the data of each pair of morphs containing the same identity. For example, to evaluate the bias for identity **a**, we'll consider the PSE estimates for morphs **ab** and **ac** across the two sessions, and fit a linear model with participants, angular location, and their interaction as predictors.

```
require(tidyverse)
require(lme4)
require(car)
require(broom)
```

First we compute the PSE for each participant by adding PSEp and DPSE.

```
cols <- c('pos1', 'pos3', 'pos5', 'pos7')
pse_ses1 = list()
pse_ses2 = list()

# compute the PSE by adding PSEp to DPSE for each subject
for (morph in c('ac', 'ab', 'bc')) {
  psep <- psep_ses1[[morph]]
  dpse <- dpse_ses1[[morph]]
  pse <- (matrix(rep(psep[cols], 10), nrow=10, byrow=T) +
           dpse[cols])*100 + 50
  pse$subject <- row.names(pse)
  pse_ses1[[morph]] <- pse

  psep <- psep_ses2[[morph]]
  dpse <- dpse_ses2[[morph]]
}
```

```

pse <- (matrix(rep(psep[cols], 10), nrow=10, byrow=T) +
        dpse[cols])*100 + 50
pse$subject <- row.names(pse)
pse_ses2[[morph]] <- pse
}

# now store it in long format
pse_ses1_long <-
  ldply(
    lapply(pse_ses1, function(x) mutate(x, subject=row.names(x)) %>% gather(pos, pse, -subject)),
    data.frame) %>%
    mutate(morph_type=.id, session='1')
pse_ses2_long <-
  ldply(
    lapply(pse_ses2, function(x) mutate(x, subject=row.names(x)) %>% gather(pos, pse, -subject)),
    data.frame) %>%
    mutate(morph_type=.id, session='2')

pse <-
  rbind(pse_ses1_long, pse_ses2_long) %>%
  mutate(pos_num=mapvalues(pos,
                           paste('pos', c(1, 3, 5, 7), sep=''),
                           c(1, 3, 5, 7)*45))
pse$pos_num <- factor(pse$pos_num, levels=c(1, 3, 5, 7)*45)

```

Let's save it for later use.

```

# save pse
pse_fn <- '../derivatives/pse.csv'

if (!file.exists(pse_fn)) {
  pse_ <- pse %>%
    select(subject, session, morph_type, pos_num, pse)
  write_csv(pse_, pse_fn)
}

```

Here we load the pre-estimated data

```

pse <- read_csv('../derivatives/pse.csv', col_types='ciccd')
pse$morph_type <- as.factor(pse$morph_type)
pse$pos_num <- factor(pse$pos_num,
                     levels=c('45', '135', '225', '315'))

```

We set up a function to filter the data for each individual identity. Moreover, we make sure that the PSE value indicates consistently the bias with respect to the same identity.

```

filter_dataset <- function(id, df) {
  identity2morphs <- list(a=c('ab', 'ac'),
                          b=c('ab', 'bc'),
                          c=c('bc', 'ac'))

  df_ <-
    df %>%
    filter(morph_type %in% identity2morphs[[id]]) %>%
    droplevels(.$morph_type)

  # put the PSE in the right order: > 50 indicates

```

```

# more towards the identity of interest
if (id == 'b') {
  df_ <-
  df_ %>%
    mutate(pse=ifelse(morph_type == 'ab', 100 - pse, pse))
} else if (id == 'c') {
  df_ <-
  df_ %>%
    mutate(pse=100 - pse)
}

return(df_)
}

```

Finally we can fit the models

```

ids <- c('a', 'b', 'c')
# split the datasets
pse_ids <- map(ids, filter_dataset, pse)

fit_model <- function(df) {
  df$subject <- as.factor(df$subject)
  df$pos_num <- as.factor(df$pos_num)
  contrasts(df$subject) <- contr.sum(length(unique(df$subject)))
  contrasts(df$pos_num) <- contr.sum(length(unique(df$pos_num)))

  return(lm(pse ~ subject * pos_num, data=df))
}

model_ids <-
  map(pse_ids, fit_model) %>%
  set_names(ids)

anova_model <-
  model_ids %>%
  map(Anova, type=3)

tidy_anova <-
  anova_model %>%
  map_df(tidy, .id='id')

```

And here are the results

```
anova_model
```

```

## $a
## Anova Table (Type III tests)
##
## Response: pse
##           Sum Sq Df  F value    Pr(>F)
## (Intercept) 438129  1 1501.3371 < 2.2e-16 ***
## subject      1395   9   0.5311   0.8495
## pos_num      18777  3  21.4481  3.43e-11 ***
## subject:pos_num  9191 27   1.1664   0.2807
## Residuals    35019 120

```

```
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## $b
## Anova Table (Type III tests)
##
## Response: pse
##           Sum Sq Df  F value    Pr(>F)
## (Intercept) 469993  1 1184.2655 < 2.2e-16 ***
## subject      5970   9   1.6713   0.10322
## pos_num      16226  3   13.6287 1.039e-07 ***
## subject:pos_num 18981 27   1.7714   0.01947 *
## Residuals    47624 120
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## $c
## Anova Table (Type III tests)
##
## Response: pse
##           Sum Sq Df  F value    Pr(>F)
## (Intercept) 302383  1 1093.0426 < 2.2e-16 ***
## subject      10701  9   4.2981 7.363e-05 ***
## pos_num      62758  3   75.6183 < 2.2e-16 ***
## subject:pos_num 24919 27   3.3361 3.229e-06 ***
## Residuals    33197 120
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

We find a significant interaction between participants and angular location for identities b and c, but not for identity a, suggesting that for identity a the bias is consistently more homogeneous across participants.

## Use the model to predict the biases

We can also use these models now to predict the biases at each angular location for each individual participant.

```
df_pred <- expand_grid(
  subject=unique(pse$subject),
  pos_num=unique(pse$pos_num)
)

df_pred <-
map_df(model_ids, predict, df_pred) %>%
  gather(identity, pse) %>%
  cbind(df_pred, .)

ggplot(aes(pos_num, pse-50, color=subject, group=subject), data=df_pred) +
  geom_point() +
  geom_line() +
  facet_wrap(~identity, nrow=1)
```

