

Does the strength of women's attraction to male vocal masculinity track changes in steroid hormones?

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Abstract

Recent studies that either used luteinizing hormone tests to confirm the timing of ovulation or measured steroid hormones from saliva have found little evidence that women's preferences for facial or body masculinity track within-subject changes in women's fertility or hormonal status. Fewer studies using these methods have examined women's preferences for vocal masculinity, however, and those that did report mixed results. Consequently, we used a longitudinal design and measured steroid hormones from saliva to test for evidence of hormonal regulation of women's (N=351) preferences for two aspects of male vocal masculinity (low pitch and low formants). Analyses suggested that preferences for masculine pitch, but not masculine formants, may track within-woman changes in estradiol. Although these results present some evidence for the hypothesis that within-subject hormones regulate women's attraction to masculine men, we do not discount the possibility that the effect of estradiol on pitch preferences in the current study is a false positive.

Introduction

The Dual Mating Strategy hypothesis of ovulatory shifts in women's mate preferences proposes that women show stronger preferences for masculine men during high-fertility ovulatory phase of their menstrual cycle (Gangestad & Simpson, 2000; Little et al., 2011; Penton-Voak et al., 1999). Early tests of this hypothesis suggested women do indeed show stronger preferences for masculine characteristics in men's faces (Little & Jones, 2012; Penton-Voak et al., 1999), voices (Puts, 2005, 2006; Feinberg et al., 2006), bodies (Little et al., 2007), behavioral displays (Gangestad et al., 2004), and body odors (Havlicek et al., 2005). However, these studies have recently been criticized for employing small sample sizes, using weak methods to assess women's position in the menstrual cycle and/or hormonal status, and being overly-reliant on between-subject designs (Blake et al., 2016; Gangestad et al., 2016; Jones et al., 2018a).

These criticisms have triggered a concerted effort from many research groups to test for fertility- and/or hormone-linked changes in women's masculinity preferences using methods that address the methodological limitations of earlier work. Recent large-scale studies of women's preferences for masculinity in men's faces (Jones et al., 2018a; Marcinkowska et al., 2018a; Marcinkowska et al., 2018b) and bodies (Jünger et al., 2018a; Marcinkowska et al., 2018a; Marcinkowska et al., 2018b) that used luteinizing hormone tests and/or measured women's hormone levels found no clear evidence that face or body preferences tracked within-subject changes in women's fertility or hormonal status. One exception to this pattern was Dixson et al. (2018), who found that women's preferences for facial masculinity and femininity tracked changes in estradiol and progesterone, respectively. Dixson et al. (2018) did not find significant correlations between face preferences and fertility, however.

Recent studies of women's preferences for vocal masculinity using similar designs to those described above have reported more mixed results, however. One study found that women's preferences for vocal masculinity were slightly stronger in test sessions where women had higher salivary estradiol, although this effect was not significant¹ (Pisanski et al., 2014). By contrast, two other recent studies found no significant effects of fertility on women's vocal masculinity preferences and found no evidence that women's vocal masculinity preferences were reliably with steroid hormone levels (Jünger et al., 2018b).

In light of the mixed results for vocal masculinity described above, here we report new data on the hormonal correlates of within-women changes in preferences for vocal masculinity. Following Pisanski et al. (2014), we used two different types of masculinity preference test. One test assessed women's preferences for masculinized versus feminized voice pitch. The other test assessed women's preferences for voices that had been masculinized and

¹ p values for the effect of estradiol were .055 and .050, depending on the model.

feminized in formant frequencies. Both pitch and format frequencies are reliably sexually dimorphic in young adults (Pisanski et al., 2016).

Methods

Participants

We tested 371 heterosexual women (mean age=21.6 years, SD=3.31 years) who reported that they were not using any form of hormonal contraceptive (i.e., all women reported having natural menstrual cycles). Women participated as part of a larger study investigating possible effects of steroid hormones on different aspects of women's behavior (Jones et al., 2018a, 2018b, 2018c). In this larger study, participants completed up to three blocks of test sessions. Each of the three blocks of test sessions consisted of five weekly test sessions. The data reported here are all responses from blocks of test sessions in the larger study where women provided voice preference data in at least one test session. Following these restrictions, 328 women had completed five or more test sessions and 96 of these women completed ten test sessions. Forty-three women completed fewer than five test sessions.

Sixty-two of our participants' first block of test sessions was previously reported in Pisanski et al. (2014). Analyses are reported with and without these data points.

Stimuli

Voice stimuli were identical to those used in Kandrik et al. (2016) and Pisanski et al. (2014). Recordings of 6 men between the ages of 18 and 25 speaking the English monophthong vowels, "ah"/a/, "ee"/i/, "e"/ɛ/, "oh"/o/, and "oo"/u/, were made in an anechoic sound-controlled booth using a Sennheiser MKH 800 cardioid condenser microphone (MKH-P48), at an approximate distance of 5-10 cm. Voice recordings were digitally encoded using an M-Audio Fast Track Ultra interface at a sampling rate of 96 kHz and 32-bit amplitude quantization, and transferred to a computer as PCM WAV files using Adobe Soundbooth CS5 version 3.0.

Following other recent work on perceptions of sexually dimorphic vocal characteristics (e.g., Kandrik et al., 2016; Pisanski et al., 2014), we created two feminized and two masculinized versions of each original voice recording by independently manipulating voice pitch or formants using the Pitch-Synchronous Overlap Add (PSOLA) algorithm in Praat version 5.2.15 (Boersma & Weenink, 2013). Pitch was lowered (masculinized) or raised (feminized) by 10% from baseline while holding formants constant. Likewise, formants were lowered (masculinized) or raised (feminized) by 10% from baseline while holding pitch constant. This process created 6 pairs of voices that differed in pitch and 6 pairs of voices that differed in formants. Following these manipulations, we amplitude-normalized the sound pressure level of all voices to 70 decibels using the root mean squared method. Voice pitch and formant measures for the feminized and masculinized voice stimuli are given in Kandrik et al. (2016). We have previously shown that these voice manipulations reliably alter masculinity and dominance perceptions in the predicted way (i.e., lowered pitch or formants increase relative judgments of masculinity and dominance; Kandrik et al., 2016; Pisanski et al., 2014).

Vocal masculinity preference test

In each test session, participants listened to 12 pairs of voices (each pair consisting of a masculinized and a feminized version of the same voice) through headphones (Sennheiser HD202). Women were instructed to select the more attractive voice in each pair and to indicate the strength of that preference by choosing from the options “slightly more attractive”, “somewhat more attractive”, more attractive”, and “much more attractive”. Trial order and the order in which participants listened to the masculinized and feminized versions in each pair were fully randomized.

Responses on the masculinity preference test were coded using the following scale (higher scores indicate stronger masculinity preferences and the scale is centered on chance, i.e., zero):

0.5 to 3.5: masculinized voice rated ‘slightly more attractive’ (=0.5), ‘somewhat more attractive’ (=1.5), ‘more attractive’ (=2.5) or ‘much more attractive’ (=3.5) than feminized voice.

-0.5 to -3.5: feminized voice rated ‘slightly more attractive’ (=−0.5), ‘somewhat more attractive’ (=−1.5), ‘more attractive’ (=−2.5) or ‘much more attractive’ (=−3.5) than masculinized voice.

Each woman’s average masculinity preference score was calculated separately for the pitch-manipulated and formant-manipulated trials for each test session. This method for assessing and coding masculinity preferences has been used in many previous studies (e.g., Jones et al., *in press a*; Zietsch et al., 2015). Higher scores indicate stronger masculinity preferences.

Saliva samples

Participants provided a saliva sample via passive drool (Papacosta & Nassis, 2011) in each test session. Participants were instructed to avoid consuming alcohol and coffee in the 12 hours prior to participation and avoid eating, smoking, drinking, chewing gum, or brushing their teeth in the 60 minutes prior to participation. Each woman’s test sessions took place at approximately the same time of day to minimize effects of diurnal changes in hormone levels (Veldhuis et al., 1988; Bao et al., 2003).

Saliva samples were frozen immediately and stored at -32°C until being shipped, on dry ice, to the Salimetrics Lab (Suffolk, UK) for analysis, where they were assayed using the Salivary 17 β -Estradiol Enzyme Immunoassay Kit 1-3702 ($M=3.30$ pg/mL, $SD=1.27$ pg/mL, sensitivity=0.1 pg/mL, intra-assay CV=7.13%, inter-assay CV=7.45%), Salivary Progesterone Enzyme Immunoassay Kit 1-1502 ($M=148.45$ pg/mL, $SD=95.95$ pg/mL, sensitivity=5 pg/mL, intra-assay CV=6.20%, inter-assay CV=7.55%), Salivary Testosterone Enzyme Immunoassay Kit 1-2402 ($M=87.58$ pg/mL, $SD=27.18$ pg/mL, sensitivity<1.0 pg/mL, intra-assay CV=4.60%, inter-assay CV=9.83%), and Salivary Cortisol Enzyme Immunoassay Kit 1-3002 ($M=0.23$ μ g/dL, $SD=0.16$

µg/dL, sensitivity<0.003 µg/dL, intra-assay CV=3.50%, inter-assay CV=5.08%).

Hormone levels more than three standard deviations from the sample mean for that hormone, or where Salimetrics indicated levels were outside the sensitivity range of their relevant ELISA, were excluded from the dataset (~1% of hormone measures were excluded for these reasons). The descriptive statistics given above do not include these excluded values. Values for each hormone were centered on their subject-specific means to isolate effects of within-subject changes in hormones. They were then scaled so the majority of the distribution for each hormone varied from -.5 to .5 to facilitate calculations in the linear mixed models. As hormone levels were centered on their subject-specific means, women with only one value for a hormone could not be included in the analyses.

Analyses

Linear mixed models were used to test for possible effects of hormonal status on masculinity preferences. Analyses were conducted using R version 3.3.2 (R Core Team, 2016), with lme4 version 1.1-13 (Bates et al., 2014) and lmerTest version 2.0-33 (Kuznetsova et al., 2013). The dependent variable was the masculinity preference score for each test session (separate models were run for the pitch and formant manipulations). Predictors were the scaled and centered hormone levels. Random slopes were specified maximally following Barr et al. (2013) and Barr (2013). Full model specifications and full results for each analysis are given in our Supplemental Materials. Data files and analysis scripts are publicly available at <https://osf.io/byu3h/>. 95% confidence intervals are given in our Supplemental Materials.

Results

Masculinity preferences for pitch- or formant-manipulated voice stimuli were analyzed separately. For each type of masculinity manipulation (pitch manipulation, formant manipulation) we ran three models. Our first model (Model 1) included estradiol, progesterone, and their interaction as predictors. Our second model (Model 2) included estradiol, progesterone, and estradiol-

to-progesterone ratio as predictors. Our third model (Model 3) included testosterone and cortisol as predictors, but did not consider possible effects of estradiol or progesterone. This analysis strategy is identical to that used in Jones et al. (2018a), Jones et al. (2018b), and Jones et al. (2018c) to investigate the hormonal correlates of women's facial masculinity preferences, sexual desire, and disgust sensitivity, respectively.

Preferences for masculinized pitch

The intercept was significant in all three models (all estimates >0.70 , all $ts>26.60$, all $ps<.001$, $M=0.70$, $SEM=0.03$), indicating that women generally judged pitch-masculinized versions of male voices to be more attractive than pitch-feminized versions. Model 1 revealed a significant positive effect of estradiol (estimate=0.19, $t=2.38$, $p=.017$), but neither the effect of progesterone (estimate=-0.04, $t=-0.60$, $p=.55$) nor the interaction between estradiol and progesterone (estimate=-0.04, $t=-0.10$, $p=.92$) were significant. Model 2 also revealed a significant positive effect of estradiol (estimate=0.21, $t=2.58$, $p=.010$), but neither the effects of progesterone (estimate=-0.09, $t=-1.15$, $p=.25$) nor estradiol-to-progesterone ratio (estimate=-0.05, $t=-1.23$, $p=.22$) were significant. In Model 3, neither testosterone (estimate=0.15, $t=1.77$, $p=.08$) nor cortisol (estimate=-0.06, $t=-0.86$, $p=.39$) had a significant effect on women's pitch preferences.

Preferences for masculinized formants

The intercept was significant in all three models (all estimates >0.44 , all $ts>13.08$, all $ps<.001$, $M=0.45$, $SEM=0.03$), indicating that women generally judged formant-masculinized versions of male voices to be more attractive than formant-feminized versions. None of the three models testing for effects of hormones on formant-masculinized voices revealed significant effects of any hormones (all absolute estimates <0.18 , all absolute $ts<0.46$, all $ps>.50$).

Repeating the analyses of pitch and formant preferences described above including women's partnership status (partnered versus unpartnered) as an additional effect-coded factor revealed no significant interactions involving partnership status. The significant effects of estradiol on pitch preferences

described above were also significant in these analyses. These analyses are described in full in our Supplemental Materials.

Additional within-subject analyses

An analysis of voice preference data from 62 of our participants' first block of test sessions was previously reported in Pisanski et al. (2014). Excluding these data from our analyses did not alter the pattern of significant results. Full results of these additional analyses are reported in our Supplemental Materials.

Between-subject analyses

Recent work on links between between-subject (i.e., average) progesterone levels and face (Marcinkowska et al., 2018a; DeBruine et al., 2018) and body (Marcinkowska et al., 2018a) preferences suggest that masculinity preferences are predicted by the combined effects of relationship status and average progesterone. The corresponding effects for pitch and formant preferences in this data set (all participants included) were not significant. These results are described in full in our Supplemental Materials.

Discussion

Consistent with previous research (e.g., Feinberg et al., 2005; Pisanski et al., 2014), women in our study showed strong preferences for male voices with masculinized pitch and formants over those with feminized pitch and formants, respectively. We also found that women's preferences for masculinized pitch, but not masculinized formants, were stronger in test sessions with higher estradiol. These results differ from those reported in other recent work on fertility- and hormone-regulated voice preferences in important ways.

First, and despite using identical stimuli to Pisanski et al. (2014), we found that only pitch preferences tracked changes in women's estradiol. This contrasts with Pisanski et al. (2014), who reported that vocal masculinity preferences tended to track changes in estradiol regardless of whether they had been manipulated in pitch or formants. Since masculine formants are

reliably associated with masculine body shapes (Pisanski et al., 2016), our null results for formant preferences are arguably consistent with recent work finding no within-subject effects of hormonal status on women's preferences for masculine body characteristics (Jünger et al., 2018a).

Second, our results differ from those reported in Jünger et al. (2018b). Across two studies, Jünger et al. (2018b) found no evidence that women's preferences for vocal masculinity tracked changes in either fertility or estradiol. Jünger et al.'s (2018b) results suggest that the within-subject effect of estradiol observed in the current study is not robust.

In conclusion, the current study saw some evidence that women's preferences for one component of vocal masculinity (pitch), but not another (formants), track changes in estradiol. This is surprising since there is no a priori reason to expect hormonal effects for only one component of vocal masculinity. Although these results partially replicate those of a previous study using identical methods (Pisanski et al., 2014), the evidence is not particularly strong and they conflict with other recent work finding no evidence for hormonal regulation of vocal masculinity preferences (Jünger et al., 2018b). Thus, we do not rule out the possibility that the effect of estradiol on pitch preferences is a false positive. Indeed, the effects of estradiol on pitch preferences would not be significant if corrected for multiple comparisons. Arguably, more work is needed to investigate the reliability and robustness of hormonal effects on women's vocal masculinity preferences.

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Code ▾

Women's preferences for masculine pitch in men's voices may track changes in estradiol

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- Setup and data processing
 - Custom Functions
 - Load full dataset
 - Age
 - The number of sessions completed per woman
 - Exclude observations with missing estradiol, progesterone or testosterone
 - Exclude subjects with only a single session in a block
 - Remove outlier hormone values
 - Subject-mean-centre hormones
- Descriptive stats
 - Voice Masculinity preferences
 - Hormones
- Analyses of all data
 - Masculinity preference (formants)
 - Model 1: formants ~ E + P + E x P
 - Model 2: formants ~ E + P + EP_ratio
 - Model 3: formants ~ T + C
 - Masculinity preference (pitch)
 - Model 1: Pitch ~ E + P + E x P
 - Model 2: Pitch ~ E + P + EP_ratio
 - Model 3: Pitch ~ T + C
- Analyses excluding responses from Pisanski et al (2014)
 - Masculinity preference (formants)
 - Model 1: formants ~ E + P + E x P
 - Model 2: formants ~ E + P + EP_ratio
 - Model 3: formants ~ T + C
 - Masculinity preference (pitch)
 - Model 1: Pitch ~ E + P + E x P
 - Model 2: Pitch ~ E + P + EP_ratio
 - Model 3: Pitch ~ T + C
- Data subset: data_hormones_partner (with partnership status)
- Masculinity preference (formants)
 - Model 1: formants ~ E + P + E x P (with partnership status)
 - Model 2: formants ~ E + P + EP_ratio (with partnership status)
 - Model 3: formants ~ T + C (with partnership status)
- Masculinity preference (Pitch)

- Model 1: Pitch ~ E + P + E x P (with partnership status)
- Model 2: Pitch ~ E + P + EP_ratio (with partnership status)
- Model 3: Pitch ~ T + C (with partnership status)
- Average Progesterone by Partnership Status

Setup and data processing

```
R version 3.5.1 (2018-07-02)
Platform: x86_64-apple-darwin15.6.0 (64-bit)
Running under: macOS High Sierra 10.13.6

Matrix products: default
BLAS: /System/Library/Frameworks/Accelerate.framework/Versions/A/Frameworks/vecLib.framework/Versions/A/libBLAS.dylib
LAPACK: /Library/Frameworks/R.framework/Versions/3.5/Resources/lib/libRlapack.dylib

locale:
[1] en_GB.UTF-8/en_GB.UTF-8/en_GB.UTF-8/C/en_GB.UTF-8/en_GB.UTF-8

attached base packages:
[1] stats      graphics    grDevices   utils      datasets    methods     base

other attached packages:
[1] lubridate_1.7.4 car_3.0-0       carData_3.0-1    lmerTest_3.0-1
[5] lme4_1.1-17   Matrix_1.2-14   forcats_0.3.0   stringr_1.3.1
[9] dplyr_0.7.6    purrr_0.2.5    readr_1.1.1    tidyverse_1.2.1
[13] tibble_1.4.2   ggplot2_3.0.0   tidyverse_1.2.1

loaded via a namespace (and not attached):
[1] tidyselect_0.2.4  splines_3.5.1    haven_1.1.2    lattice_0.20-35
[5] colorspace_1.3-2 yaml_2.2.0      rlang_0.2.1    pillar_1.3.0
[9] nloptr_1.0.4     foreign_0.8-71   glue_1.3.0    withr_2.1.2
[13] modelr_0.1.2    readxl_1.1.0    bindrcpp_0.2.2 bindr_0.1.1
[17] plyr_1.8.4      munsell_0.5.0   gtable_0.2.0   cellranger_1.1.0
[21] zip_1.0.0       rvest_0.3.2     rio_0.5.10   knitr_1.20
[25] curl_3.2        broom_0.5.0    Rcpp_0.12.18  scales_1.0.0
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[33] openxlsx_4.1.0  stringi_1.2.4   numDeriv_2016.8-1 grid_3.5.1
[37] cli_1.0.0       tools_3.5.1    magrittr_1.5   lazyeval_0.2.1
[41] crayon_1.3.4    pkgconfig_2.0.1  MASS_7.3-50   data.table_1.11.4
[45] xml2_1.2.0      assertthat_0.2.0 minqa_1.2.4   httr_1.3.1
[49] rstudioapi_0.7   R6_2.2.2      nlme_3.1-137 compiler_3.5.1
```

Custom Functions

[Hide](#)

```
# calculate standard errors
se <- function(x, na.rm = FALSE) {
  if (na.rm) {
    the.SE <- sqrt(var(x,na.rm=TRUE)/length(na.omit(x)))
  } else {
    the.SE <- sqrt(var(x,na.rm=FALSE)/length(x))
  }

  return(the.SE)
}
```

Load full dataset

Data entered from all white, heterosexual women not using hormonal contraceptives.

Each row is all data from a single session (i.e. oc_id:date)

- “oc_id” = ID of the subject
- “age” = age (in years) of subject on day of testing
- “date” = date of testing session
- “block” = testing block (1, 2 or 3)
- “block_partner” = partnership status (0 = no partner, 1 = partner)
- “manip” = type of masculinity manipulation (formant or pitch)
- “rating.c” = mean voice rating (centered on chance)
- “prog” = salivary progesterone for that session
- “estr” = salivary estradiol for that session
- “test” = salivary testosterone for that session
- “cort” = salivary cortisol for that session
- “partner.e” = effect-coded partnership status (-0.5 = no partner, 0.5 = partner)

[Hide](#)

```
# read in the raw data
data_all <- read.csv("OCMATE_voicepref_anon.csv")
```

Age

[Hide](#)

```
data_all %>%
  group_by(oc_id) %>%
  summarise(age = min(age)) %>%
  ungroup() %>%
  group_by() %>%
  summarise(
    n = n(),
    mean_age = mean(age, na.rm = TRUE),
    sd_age = sd(age, na.rm = TRUE),
    se_age = se(age, na.rm = TRUE)
  ) %>% t() %>%
  knitr::kable()
```

The table should have a header (column names)

```
n      371.0000000
mean_age 21.5602703
sd_age    3.3136844
se_age    0.1722702
```

The number of sessions completed per woman

[Hide](#)

```
data_all %>%
  group_by(oc_id) %>%
  summarise(
    sessions = n_distinct(date)
  ) %>%
  group_by(sessions) %>%
  summarise(
    n = n()
  ) %>%
  spread(sessions, n) %>% t()
```

	[,1]
1	11
2	9
3	6
4	17
5	224
6	1
7	1
8	2
9	4
10	96

Exclude observations with missing estradiol, progesterone or testosterone

[Hide](#)

```
## Exclude observations with EPT missing hormone values
sub_nopill_hormones<-data_all %>%
  filter(!is.na(prog) |
         !is.na(est) |
         !is.na(test))
```

Exclude subjects with only a single session in a block

This is necessary because you can't calculate subject-centered means with only one data point.

[Hide](#)

```
## exclude oc_id with only a single session in a block
## (can't calculate subject-centered means)
check_single_session <- sub_nopill_hormones %>%
  group_by(oc_id, block) %>%
  summarise(sessions = n_distinct(date)) %>%
  ungroup() %>%
  filter(sessions == 1)
sub_hormones_full <- sub_nopill_hormones %>%
  anti_join(check_single_session, by=c('oc_id', 'block'))
```

Remove outlier hormone values

Remove below bottom sensitivity thresholds for assays (prog < 5, estr < 0.1) and remove outlier values (+/- 3SD from the mean).

[Hide](#)

```
sub_hormones_avg_no_outliers <- sub_hormones_full %>%
  mutate(
    prog = ifelse(prog >= 5, prog, NA),
    estr = ifelse(estr >= 0.1, estr, NA),
    prog = ifelse(prog>mean(prog, na.rm=TRUE)+3*sd(prog, na.rm=TRUE) | 
                  prog<mean(prog, na.rm=TRUE)-3*sd(prog, na.rm=TRUE), NA, prog),
    estr = ifelse(estr>mean(estr, na.rm=TRUE)+3*sd(estr, na.rm=TRUE) | 
                  estr<mean(estr, na.rm=TRUE)-3*sd(estr, na.rm=TRUE), NA, estr),
    test = ifelse(test>mean(test, na.rm=TRUE)+3*sd(test, na.rm=TRUE) | 
                  test<mean(test, na.rm=TRUE)-3*sd(test, na.rm=TRUE), NA, test),
    cort = ifelse(cort>mean(cort, na.rm=TRUE)+3*sd(cort, na.rm=TRUE) | 
                  cort<mean(cort, na.rm=TRUE)-3*sd(cort, na.rm=TRUE), NA, cort)
  )
```

Hide

```
# how many included?
sub_hormones_avg_no_outliers %>%
  group_by(oc_id, date, block) %>%
  summarise(
    e = is.na(mean(estr)),
    p = is.na(mean(prog)),
    t = is.na(mean(test)),
    c = is.na(mean(cort))
  ) %>%
  ungroup() %>%
  select(block, e:c) %>%
  gather('hormone', 'na', e:c) %>%
  group_by(hormone, block) %>%
  summarise(
    'valid' = n() - sum(na),
    'excluded' = sum(na)
  ) %>%
  arrange(block, hormone) %>%
  group_by() %>%
  summarise(
    total_hormone_samples_valid = sum(valid),
    total_hormone_samples_excluded = sum(excluded)
  ) %>% gather("stat", "value", !! 1:ncol(.)) %>%
  knitr::kable()
```

stat	value
total_hormone_samples_valid	8529
total_hormone_samples_excluded	95

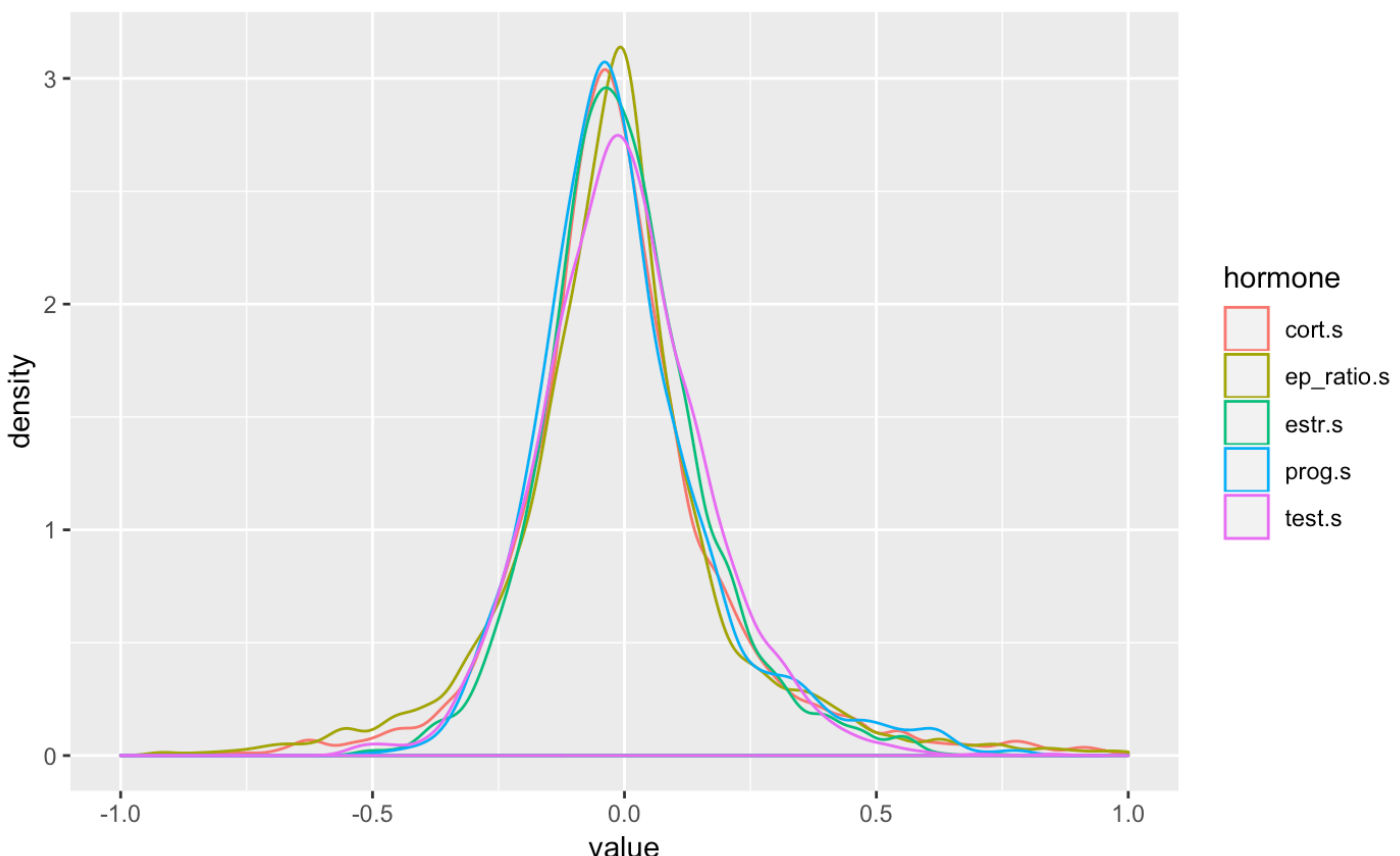
Hide

Subject-mean-centre hormones

Divide results by a constant to put all hormones on ~ -0.5 to +0.5 scale

[Hide](#)

```
data_hormones <- sub_hormones_avg_no_outliers %>%
  group_by(oc_id) %>%
  mutate(prog.s=(prog-mean(prog, na.rm=TRUE))/400,
        estr.s=(estr-mean(estr, na.rm=TRUE))/5,
        test.s=(test-mean(test, na.rm=TRUE))/100,
        cort.s=(cort-mean(cort, na.rm=TRUE))/0.5,
        ep_ratio.s = ((estr/prog)-mean(estr/prog, na.rm=TRUE))/0.075 ) %>%
  ungroup() %>%
  as.data.frame()
data_hormones %>%
  group_by(oc_id, date, prog.s, estr.s, test.s, cort.s, ep_ratio.s) %>%
  summarise(n = n()) %>%
  ungroup() %>%
  gather("hormone", "value", prog.s:ep_ratio.s) %>%
  ggplot(aes(value, colour=hormone)) +
  geom_density(alpha=.5) +
  scale_x_continuous(limits = c(-1,1))
```



NA

Hide

Descriptive stats

Voice Masculinity preferences

Hide

```
# create mean DV for all ratings by oc_id
data_hormones %>%
  group_by(oc_id, manip) %>%
  summarise(
    overall_rating.c = mean(rating.c)
  ) %>%
  ungroup() %>%
  group_by(manip) %>%
  summarise(
    n= n_distinct(oc_id),
    mean_dv = mean(overall_rating.c),
    sd_dv = sd(overall_rating.c),
    se_dv = se(overall_rating.c)
  ) %>%
  knitr::kable()
```

manip	n	mean_dv	sd_dv	se_dv
formant	352	0.4494538	0.6299405	0.0335759
pitch	352	0.7021594	0.4849671	0.0258488

Hormones

Hide

```
data_hormones %>%
  group_by() %>%
  summarise(
    mean_prog = mean(prog, na.rm = TRUE),
    sd_prog =sd(prog, na.rm = TRUE),
    se_prog =se(prog, na.rm = TRUE),
    mean_estr = mean(estr, na.rm = TRUE),
    sd_estr =sd(estr, na.rm = TRUE),
    se_estr =se(estr, na.rm = TRUE),
    mean_test = mean(test, na.rm = TRUE),
    sd_test =sd(test, na.rm = TRUE),
    se_test =se(test, na.rm = TRUE),
    mean_cort = mean(cort, na.rm = TRUE),
    sd_cort =sd(cort, na.rm = TRUE),
    se_cort =se(cort, na.rm = TRUE)
  ) %>% gather("stat", "value", !! 1:ncol(.)) %>%
  mutate(value = round(value, 4)) %>%
  separate(stat, c("stat", "hormone")) %>%
  spread(stat, value) %>%
  knitr::kable()
```

hormone	mean	sd	se
cort	0.2292	0.1646	0.0025
estr	3.2983	1.2707	0.0194
prog	148.4495	95.9503	1.4753
test	87.5753	27.1783	0.4161

[Hide](#)

Analyses of all data

Masculinity preference (formants)

Model 1: formants ~ E + P + E x P

[Hide](#)

```
model.formants.horm.EP <- lmer(rating.c ~ estr.s * prog.s +  
                                (1 | oc_id) +  
                                (0 + estr.s:prog.s || oc_id) +  
                                (1 | block:oc_id) +  
                                (0 + estr.s:prog.s || block:oc_id),  
                                data = filter(data_hormones, manip=="formant"),  
                                REML = FALSE)  
summary(model.formants.horm.EP)
```

Linear mixed model fit by maximum likelihood . t-tests use
Satterthwaite's method [lmerModLmerTest]
Formula: rating.c ~ estr.s * prog.s + (1 | oc_id) + (0 + estr.s:prog.s ||
oc_id) + (1 | block:oc_id) + (0 + estr.s:prog.s || block:oc_id)
Data: filter(data_hormones, manip == "formant")

AIC	BIC	logLik	deviance	df.resid
4038.7	4089.5	-2010.3	4020.7	2087

Scaled residuals:

Min	1Q	Median	3Q	Max
-4.3809	-0.5561	-0.0255	0.5404	3.3650

Random effects:

Groups	Name	Variance	Std.Dev.
block.oc_id	estr.s:prog.s	5.693e-12	2.386e-06
block.oc_id.1	(Intercept)	3.144e-02	1.773e-01
oc_id	estr.s:prog.s	2.199e-12	1.483e-06
oc_id.1	(Intercept)	3.281e-01	5.728e-01
Residual		2.745e-01	5.240e-01

Number of obs: 2096, groups: block:oc_id, 445; oc_id, 352

Fixed effects:

	Estimate	Std. Error	df	t value	Pr(> t)
(Intercept)	0.44743	0.03420	369.88550	13.084	<2e-16 ***
estr.s	0.03030	0.07941	1582.74293	0.382	0.703
prog.s	-0.01237	0.06903	1762.02840	-0.179	0.858
estr.s:prog.s	0.17017	0.40079	1837.64647	0.425	0.671

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

Correlation of Fixed Effects:

	(Intr)	estr.s	prog.s
estr.s	0.007		
prog.s	0.015	-0.321	
estr.s:prg.	-0.103	-0.023	-0.150

Hide

```
confint(model.formants.horm.EP, method = "Wald") %>%
  as.data.frame() %>%
  rownames_to_column() %>%
  filter(!is.na(`2.5 %`))
```

	rowname	2.5 %	97.5 %
1	(Intercept)	0.3804082	0.5144524
2	estr.s	-0.1253377	0.1859282
3	prog.s	-0.1476730	0.1229362
4	estr.s:prog.s	-0.6153611	0.9557027

Model 2: formants ~ E + P + EP_ratio

[Hide](#)

```
model.formants.horm.EPratio <- lmer(rating.c ~ estr.s + prog.s + ep_ratio.s +
  (1 | oc_id) +
  (0 + estr.s + prog.s + ep_ratio.s || oc_id) +
  (1 | block:oc_id) +
  (0 + estr.s + prog.s + ep_ratio.s || block:oc_id),
  data = filter(data_hormones, manip=="formant" ),
  REML = FALSE)
summary(model.formants.horm.EPratio)
```

```
Linear mixed model fit by maximum likelihood . t-tests use
Satterthwaite's method [lmerModLmerTest]
Formula: rating.c ~ estr.s + prog.s + ep_ratio.s + (1 | oc_id) + (0 +
estr.s + prog.s + ep_ratio.s || oc_id) + (1 | block:oc_id) +
(0 + estr.s + prog.s + ep_ratio.s || block:oc_id)
Data: filter(data_hormones, manip == "formant")

      AIC      BIC logLik deviance df.resid
4043.5   4116.9 -2008.8    4017.5     2083

Scaled residuals:
    Min      1Q Median      3Q      Max
-4.2737 -0.5421 -0.0221  0.5247  3.3182

Random effects:
Groups        Name        Variance Std.Dev.
block.oc_id   ep_ratio.s 0.000e+00 0.000e+00
block.oc_id.1  prog.s     3.752e-11 6.125e-06
block.oc_id.2  estr.s     2.453e-01 4.953e-01
block.oc_id.3  (Intercept) 2.991e-02 1.729e-01
oc_id         ep_ratio.s 5.282e-03 7.268e-02
oc_id.1       prog.s     3.217e-02 1.794e-01
oc_id.2       estr.s     0.000e+00 0.000e+00
oc_id.3       (Intercept) 3.289e-01 5.735e-01
Residual      2.667e-01 5.164e-01
Number of obs: 2096, groups: block:oc_id, 445; oc_id, 352

Fixed effects:
            Estimate Std. Error      df t value Pr(>|t|)    
(Intercept)  0.44879  0.03395 360.73315 13.219 <2e-16 ***
estr.s       0.03986  0.08746 187.84165  0.456  0.649    
prog.s      -0.02204  0.08203 180.27461 -0.269  0.788    
ep_ratio.s  -0.01810  0.04620 14.15097 -0.392  0.701    
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Correlation of Fixed Effects:
            (Intr) estr.s prog.s ep_ratio.s
estr.s      0.006
prog.s     -0.001 -0.368
ep_ratio.s -0.001 -0.209  0.524
```

Hide

```
confint(model.formants.horm.EPratio, method = "Wald") %>%
  as.data.frame() %>%
  rownames_to_column() %>%
  filter(!is.na(`2.5 %`))
```

rowname	2.5 %	97.5 %
1 (Intercept)	0.3822532	0.51533423
2 estr.s	-0.1315672	0.21128823
3 prog.s	-0.1828287	0.13874135
4 ep_ratio.s	-0.1086468	0.07245605

Model 3: formants ~ T + C

[Hide](#)

```
model.formants.horm.TC <- lmer(rating.c ~ test.s + cort.s +
  (1 | oc_id) +
  (0 + test.s + cort.s || oc_id) +
  (1 | block:oc_id) +
  (0 + test.s + cort.s|| block:oc_id),
  data = filter(data_hormones, manip=="formant"),
  REML = FALSE)
summary(model.formants.horm.TC)
```

```
Linear mixed model fit by maximum likelihood . t-tests use
Satterthwaite's method [lmerModLmerTest]

Formula:
rating.c ~ test.s + cort.s + (1 | oc_id) + (0 + test.s + cort.s ||
oc_id) + (1 | block:oc_id) + (0 + test.s + cort.s || block:oc_id)
Data: filter(data_hormones, manip == "formant")

      AIC      BIC logLik deviance df.resid
4101.3   4157.9 -2040.6    4081.3     2112

Scaled residuals:
    Min      1Q Median      3Q      Max
-4.4011 -0.5477 -0.0277  0.5289  3.2725

Random effects:
Groups        Name        Variance Std.Dev.
block.oc_id   cort.s     0.000e+00 0.000e+00
block.oc_id.1 test.s     0.000e+00 0.000e+00
block.oc_id.2 (Intercept) 3.227e-02 1.796e-01
oc_id         cort.s     0.000e+00 0.000e+00
oc_id.1       test.s     2.519e-15 5.019e-08
oc_id.2       (Intercept) 3.279e-01 5.726e-01
Residual      2.770e-01 5.263e-01
Number of obs: 2122, groups: block:oc_id, 444; oc_id, 351

Fixed effects:
            Estimate Std. Error          df t value Pr(>|t|)    
(Intercept)  0.45060   0.03407  359.83939 13.227 <2e-16 ***
test.s      -0.02124   0.07989 1750.14278 -0.266    0.790  
cort.s      -0.03980   0.05968 1698.58852 -0.667    0.505  
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Correlation of Fixed Effects:

```
(Intr) test.s
test.s -0.003
cort.s  0.007 -0.441
```

Hide

```
confint(model.formants.horm.TC, method = "Wald") %>%
  as.data.frame() %>%
  rownames_to_column() %>%
  filter(!is.na(`^2.5 %`))
```

```
rownames      2.5 %     97.5 %
1 (Intercept) 0.3838292 0.51737328
2      test.s -0.1778115 0.13533200
3      cort.s -0.1567730 0.07716821
```

Masculinity preference (pitch)

Model 1: Pitch ~ E + P + E x P

[Hide](#)

```
model.pitch.horm.EP <- lmer(rating.c ~ estr.s * prog.s +
  (1 | oc_id) +
  (0 + estr.s:prog.s || oc_id) +
  (1 | block:oc_id) +
  (0 + estr.s:prog.s|| block:oc_id),
  data = filter(data_hormones, manip=="pitch"),
  REML = FALSE)
summary(model.pitch.horm.EP)
```

```
Linear mixed model fit by maximum likelihood . t-tests use
Satterthwaite's method [lmerModLmerTest]
Formula: rating.c ~ estr.s * prog.s + (1 | oc_id) + (0 + estr.s:prog.s || 
oc_id) + (1 | block:oc_id) + (0 + estr.s:prog.s || block:oc_id)
Data: filter(data_hormones, manip == "pitch")

      AIC      BIC      logLik deviance df.resid
3935.4   3986.2   -1958.7    3917.4      2087

Scaled residuals:
    Min     1Q Median     3Q    Max
-4.6078 -0.5446 -0.0350  0.5303  4.4362

Random effects:
Groups        Name        Variance Std.Dev.
block.oc_id  estr.s:prog.s 1.615e-14 1.271e-07
block.oc_id.1 (Intercept) 2.957e-02 1.720e-01
oc_id         estr.s:prog.s 0.000e+00 0.000e+00
oc_id.1       (Intercept) 1.603e-01 4.004e-01
Residual                 2.884e-01 5.370e-01
Number of obs: 2096, groups: block:oc_id, 445; oc_id, 352

Fixed effects:
            Estimate Std. Error      df t value Pr(>|t|)    
(Intercept)  0.70121  0.02631  366.12574 26.650 <2e-16 ***
estr.s       0.19333  0.08115 1533.86813  2.382  0.0173 *  
prog.s      -0.04266  0.07065 1760.08934 -0.604  0.5460  
estr.s:prog.s -0.04159  0.40606 1902.36782 -0.102  0.9184  
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Correlation of Fixed Effects:
            (Intr) estr.s prog.s
estr.s      0.009
prog.s      0.019 -0.320
estr.s:prg. -0.136 -0.023 -0.148
```

Hide

```
confint(model.pitch.horm.EP, method = "Wald") %>%
as.data.frame() %>%
rownames_to_column() %>%
filter(!is.na(`2.5 %`))
```

	rowname	2.5 %	97.5 %
1	(Intercept)	0.64964273	0.7527854
2	estr.s	0.03427162	0.3523814
3	prog.s	-0.18113085	0.0958038
4	estr.s:prog.s	-0.83745519	0.7542755

Model 2: Pitch ~ E + P + EP_ratio

[Hide](#)

```
model.pitch.horm.EPratio <- lmer(rating.c ~ estr.s + prog.s + ep_ratio.s +
(1 | oc_id) +
(0 + estr.s + prog.s + ep_ratio.s || oc_id) +
(1 | block:oc_id) +
(0 + estr.s + prog.s + ep_ratio.s || block:oc_id),
data = filter(data_hormones, manip=="pitch" ),
REML = FALSE)
summary(model.pitch.horm.EPratio)
```

```
Linear mixed model fit by maximum likelihood . t-tests use
Satterthwaite's method [lmerModLmerTest]
Formula: rating.c ~ estr.s + prog.s + ep_ratio.s + (1 | oc_id) + (0 +
estr.s + prog.s + ep_ratio.s || oc_id) + (1 | block:oc_id) +
(0 + estr.s + prog.s + ep_ratio.s || block:oc_id)
Data: filter(data_hormones, manip == "pitch")

      AIC      BIC logLik deviance df.resid
3941.9   4015.3 -1957.9    3915.9     2083

Scaled residuals:
    Min      1Q Median      3Q      Max
-4.6089 -0.5403 -0.0377  0.5364  4.4322

Random effects:
Groups        Name        Variance Std.Dev.
block.oc_id   ep_ratio.s 0.000e+00 0.000e+00
block.oc_id.1  prog.s     5.482e-14 2.341e-07
block.oc_id.2  estr.s     0.000e+00 0.000e+00
block.oc_id.3  (Intercept) 2.982e-02 1.727e-01
oc_id         ep_ratio.s 0.000e+00 0.000e+00
oc_id.1       prog.s     0.000e+00 0.000e+00
oc_id.2       estr.s     0.000e+00 0.000e+00
oc_id.3       (Intercept) 1.601e-01 4.002e-01
Residual      2.881e-01 5.368e-01
Number of obs: 2096, groups: block:oc_id, 445; oc_id, 352

Fixed effects:
            Estimate Std. Error          df t value Pr(>|t|)    
(Intercept)  0.70086   0.02607  352.76221 26.888 < 2e-16 ***
estr.s       0.21420   0.08290 1572.63621  2.584  0.00986 ** 
prog.s      -0.09287   0.08049 1754.50021 -1.154  0.24876  
ep_ratio.s  -0.05094   0.04149 1732.83289 -1.228  0.21968  
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Correlation of Fixed Effects:
            (Intr) estr.s prog.s ep_ratio.s
estr.s      0.006
prog.s     -0.001 -0.381
ep_ratio.s  0.000 -0.207  0.497
```

Hide

```
confint(model.pitch.horm.EPratio, method = "Wald") %>%
  as.data.frame() %>%
  rownames_to_column() %>%
  filter(!is.na(`2.5 %`))
```

rowname	2.5 %	97.5 %
1 (Intercept)	0.64977557	0.75195191
2 estr.s	0.05171203	0.37669013
3 prog.s	-0.25063000	0.06489325
4 ep_ratio.s	-0.13225536	0.03037491

Model 3: Pitch ~ T + C

[Hide](#)

```
model.pitch.horm.TC <- lmer(rating.c ~ test.s + cort.s +
  (1 | oc_id) +
  (0 + test.s + cort.s || oc_id) +
  (1 | block:oc_id) +
  (0 + test.s + cort.s|| block:oc_id),
  data = filter(data_hormones, manip=="pitch"),
  REML = FALSE)
summary(model.pitch.horm.TC)
```

```
Linear mixed model fit by maximum likelihood . t-tests use
Satterthwaite's method [lmerModLmerTest]

Formula:
rating.c ~ test.s + cort.s + (1 | oc_id) + (0 + test.s + cort.s ||
oc_id) + (1 | block:oc_id) + (0 + test.s + cort.s || block:oc_id)
Data: filter(data_hormones, manip == "pitch")

      AIC      BIC      logLik deviance df.resid
3992.1   4048.7   -1986.1    3972.1     2112

Scaled residuals:
    Min      1Q  Median      3Q      Max
-4.6443 -0.5491 -0.0480  0.5332  4.3182

Random effects:
Groups        Name        Variance Std.Dev.
block.oc_id   cort.s     0.06619  0.2573
block.oc_id.1 test.s     0.00000  0.0000
block.oc_id.2 (Intercept) 0.03282  0.1812
oc_id         cort.s     0.00000  0.0000
oc_id.1       test.s     0.00000  0.0000
oc_id.2       (Intercept) 0.15542  0.3942
Residual      0.28692  0.5357
Number of obs: 2122, groups: block:oc_id, 444; oc_id, 351

Fixed effects:
            Estimate Std. Error          df t value Pr(>|t|)    
(Intercept)  0.70151   0.02593  350.33503 27.049 <2e-16 ***
test.s       0.14612   0.08245 1573.69929  1.772  0.0766 .
cort.s      -0.05728   0.06676  131.30995 -0.858  0.3924 
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Correlation of Fixed Effects:
  (Intr) test.s
test.s -0.005
cort.s  0.011 -0.442
```

[Hide](#)

```
confint(model.pitch.horm.TC, method = "Wald") %>%
  as.data.frame() %>%
  rownames_to_column() %>%
  filter(!is.na(`^2.5 %`))
```

```
rownames      2.5 %     97.5 %
1 (Intercept) 0.6506820 0.75234485
2      test.s -0.0154798 0.30771562
3      cort.s -0.1881335 0.07356425
```

Analyses excluding responses from Pisanski et al (2014)

[Hide](#)

```
data_hormones_post_2013 <- data_hormones %>%
  filter(dmy(date) > ymd('2013-09-06'))
```

Masculinity preference (formants)

Model 1: formants ~ E + P + E x P

[Hide](#)

```
model.formants.horm.EP <- lmer(rating.c ~ estr.s * prog.s +
  (1 | oc_id) +
  (0 + estr.s:prog.s || oc_id) +
  (1 | block:oc_id) +
  (0 + estr.s:prog.s|| block:oc_id),
  data = filter(data_hormones_post_2013, manip=="formant"),
  REML = FALSE)
summary(model.formants.horm.EP)
```

```
Linear mixed model fit by maximum likelihood . t-tests use
Satterthwaite's method [lmerModLmerTest]
Formula: rating.c ~ estr.s * prog.s + (1 | oc_id) + (0 + estr.s:prog.s || 
oc_id) + (1 | block:oc_id) + (0 + estr.s:prog.s || block:oc_id)
Data: filter(data_hormones_post_2013, manip == "formant")

      AIC      BIC      logLik deviance df.resid
3458.4   3507.8   -1720.2    3440.4     1777

Scaled residuals:
    Min      1Q Median      3Q      Max
-4.3285 -0.5618 -0.0258  0.5329  3.3328

Random effects:
Groups      Name        Variance Std.Dev.
block.oc_id  estr.s:prog.s 0.000e+00 0.000e+00
block.oc_id.1 (Intercept) 2.538e-02 1.593e-01
oc_id        estr.s:prog.s 3.533e-11 5.944e-06
oc_id.1      (Intercept) 3.277e-01 5.724e-01
Residual                 2.789e-01 5.281e-01
Number of obs: 1786, groups: block:oc_id, 378; oc_id, 300

Fixed effects:
            Estimate Std. Error      df t value Pr(>|t|)    
(Intercept) 4.311e-01 3.681e-02 3.151e+02 11.712 <2e-16 ***
estr.s      2.769e-02 8.999e-02 1.407e+03  0.308    0.758  
prog.s      4.012e-03 7.501e-02 1.505e+03  0.053    0.957  
estr.s:prog.s 1.206e-01 4.451e-01 1.567e+03  0.271    0.786  
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Correlation of Fixed Effects:
            (Intr) estr.s prog.s
estr.s      0.022
prog.s      0.004 -0.320
estr.s:prg. -0.102 -0.027 -0.110
```

Hide

```
confint(model.formants.horm.EP, method = "Wald") %>%
as.data.frame() %>%
rownames_to_column() %>%
filter(!is.na(`2.5 %`))
```

```
rownames      2.5 %    97.5 %
1  (Intercept) 0.3589187 0.5031937
2      estr.s -0.1486796 0.2040572
3      prog.s -0.1429994 0.1510230
4 estr.s:prog.s -0.7517601 0.9929421
```

Model 2: formants ~ E + P + EP_ratio

[Hide](#)

```
model.formants.horm.EPratio <- lmer(rating.c ~ estr.s + prog.s + ep_ratio.s +
(1 | oc_id) +
(0 + estr.s + prog.s + ep_ratio.s || oc_id) +
(1 | block:oc_id) +
(0 + estr.s + prog.s + ep_ratio.s || block:oc_id),
data = filter(data_hormones_post_2013, manip=="formant" ),
REML = FALSE)
```

Model failed to converge with 1 negative eigenvalue: -1.4e+01

[Hide](#)

```
summary(model.formants.horm.EPratio)
```

```
Linear mixed model fit by maximum likelihood . t-tests use
Satterthwaite's method [lmerModLmerTest]
Formula: rating.c ~ estr.s + prog.s + ep_ratio.s + (1 | oc_id) + (0 +
estr.s + prog.s + ep_ratio.s || oc_id) + (1 | block:oc_id) +
(0 + estr.s + prog.s + ep_ratio.s || block:oc_id)
Data: filter(data_hormones_post_2013, manip == "formant")

      AIC      BIC logLik deviance df.resid
3459.2   3530.6 -1716.6    3433.2     1773

Scaled residuals:
    Min      1Q Median      3Q      Max
-4.1107 -0.5491 -0.0226  0.5074  3.2600

Random effects:
Groups        Name        Variance Std.Dev.
block.oc_id  ep_ratio.s 3.275e-10 1.810e-05
block.oc_id.1 prog.s     0.000e+00 0.000e+00
block.oc_id.2 estr.s     4.659e-01 6.826e-01
block.oc_id.3 (Intercept) 2.303e-02 1.518e-01
oc_id        ep_ratio.s 3.361e-02 1.833e-01
oc_id.1      prog.s     6.434e-03 8.021e-02
oc_id.2      estr.s     5.816e-14 2.412e-07
oc_id.3      (Intercept) 3.277e-01 5.724e-01
Residual            2.655e-01 5.153e-01
Number of obs: 1786, groups: block:oc_id, 378; oc_id, 300

Fixed effects:
            Estimate Std. Error       df t value Pr(>|t|)    
(Intercept)  0.432497  0.036463 306.172049 11.861 <2e-16 ***
estr.s       0.034974  0.104133 197.207932  0.336    0.737  
prog.s       0.006005  0.093339 214.912755  0.064    0.949  
ep_ratio.s   -0.010397  0.067076 21.391213 -0.155    0.878  
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Correlation of Fixed Effects:
            (Intr) estr.s prog.s ep_ratio.s
estr.s      0.020
prog.s      -0.004 -0.385
ep_ratio.s   0.004 -0.248  0.584
```

Hide

```
confint(model.formants.horm.EPratio, method = "Wald") %>%
  as.data.frame() %>%
  rownames_to_column() %>%
  filter(!is.na(`^2.5 %`))
```

```
rownome      2.5 %    97.5 %
1 (Intercept) 0.3610304 0.5039634
2      estr.s -0.1691240 0.2390716
3      prog.s -0.1769370 0.1889462
4   ep_ratio.s -0.1418634 0.1210699
```

Model 3: formants ~ T + C

[Hide](#)

```
model.formants.horm.TC <- lmer(rating.c ~ test.s + cort.s +
(1 | oc_id) +
(0 + test.s + cort.s || oc_id) +
(1 | block:oc_id) +
(0 + test.s + cort.s|| block:oc_id),
data = filter(data_hormones_post_2013, manip=="formant"),
REML = FALSE)
summary(model.formants.horm.TC)
```

Linear mixed model fit by maximum likelihood . t-tests use Satterthwaite's method [lmerModLmerTest]

Formula:

```
rating.c ~ test.s + cort.s + (1 | oc_id) + (0 + test.s + cort.s || oc_id) + (1 | block:oc_id) + (0 + test.s + cort.s || block:oc_id)
```

Data: filter(data_hormones_post_2013, manip == "formant")

AIC	BIC	logLik	deviance	df.resid
3516.6	3571.6	-1748.3	3496.6	1797

Scaled residuals:

Min	1Q	Median	3Q	Max
-4.3367	-0.5452	-0.0278	0.5222	3.2407

Random effects:

Groups	Name	Variance	Std.Dev.
block.oc_id	cort.s	0.00000	0.0000
block.oc_id.1	test.s	0.00000	0.0000
block.oc_id.2	(Intercept)	0.02687	0.1639
oc_id	cort.s	0.00000	0.0000
oc_id.1	test.s	0.00000	0.0000
oc_id.2	(Intercept)	0.32752	0.5723
Residual		0.28231	0.5313

Number of obs: 1807, groups: block:oc_id, 377; oc_id, 299

Fixed effects:

	Estimate	Std. Error	df	t value	Pr(> t)
(Intercept)	4.334e-01	3.671e-02	3.062e+02	11.805	<2e-16 ***
test.s	-9.419e-03	8.452e-02	1.465e+03	-0.111	0.911
cort.s	-3.268e-02	6.299e-02	1.426e+03	-0.519	0.604

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

Correlation of Fixed Effects:

(Intr)	test.s
test.s	-0.003
cort.s	0.005 -0.444

Hide

```
confint(model.formants.horm.TC, method = "Wald") %>%  
  as.data.frame() %>%  
  rownames_to_column() %>%  
  filter(!is.na(`^2.5 %`))
```

```
rownames      2.5 %     97.5 %
1 (Intercept) 0.3614060 0.50531018
2      test.s -0.1750803 0.15624146
3      cort.s -0.1561409 0.09077523
```

Masculinity preference (pitch)

Model 1: Pitch ~ E + P + E x P

[Hide](#)

```
model.pitch.horm.EP <- lmer(rating.c ~ estr.s * prog.s +
  (1 | oc_id) +
  (0 + estr.s:prog.s || oc_id) +
  (1 | block:oc_id) +
  (0 + estr.s:prog.s || block:oc_id),
  data = filter(data_hormones_post_2013, manip=="pitch"),
  REML = FALSE)
summary(model.pitch.horm.EP)
```

```
Linear mixed model fit by maximum likelihood . t-tests use
Satterthwaite's method [lmerModLmerTest]
Formula: rating.c ~ estr.s * prog.s + (1 | oc_id) + (0 + estr.s:prog.s || 
oc_id) + (1 | block:oc_id) + (0 + estr.s:prog.s || block:oc_id)
Data: filter(data_hormones_post_2013, manip == "pitch")

      AIC      BIC      logLik deviance df.resid
3358.0  3407.3   -1670.0    3340.0     1777

Scaled residuals:
    Min      1Q Median      3Q      Max
-4.6041 -0.5556 -0.0318  0.5546  4.4898

Random effects:
Groups        Name        Variance Std.Dev.
block.oc_id  estr.s:prog.s 0.00000  0.0000
block.oc_id.1 (Intercept) 0.02196  0.1482
oc_id        estr.s:prog.s 0.00000  0.0000
oc_id.1      (Intercept)  0.16705  0.4087
Residual                 0.28995  0.5385
Number of obs: 1786, groups: block:oc_id, 378; oc_id, 300

Fixed effects:
            Estimate Std. Error       df t value Pr(>|t|)    
(Intercept)  0.70537  0.02852 308.86686  24.730 <2e-16 ***
estr.s       0.19477  0.09122 1357.43580   2.135  0.0329 *  
prog.s      -0.05148  0.07629 1501.99430  -0.675  0.4999    
estr.s:prog.s -0.19055  0.44859 1619.44124  -0.425  0.6711    
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Correlation of Fixed Effects:
            (Intr) estr.s prog.s
estr.s      0.029
prog.s      0.005 -0.319
estr.s:prg. -0.134 -0.026 -0.108
```

Hide

```
confint(model.pitch.horm.EP, method = "Wald") %>%
as.data.frame() %>%
rownames_to_column() %>%
filter(!is.na(`2.5 %`))
```

	rowname	2.5 %	97.5 %
1	(Intercept)	0.6494692	0.76127542
2	estr.s	0.0159860	0.37356189
3	prog.s	-0.2010074	0.09805076
4	estr.s:prog.s	-1.0697710	0.68867938

Model 2: Pitch ~ E + P + EP_ratio

[Hide](#)

```
model.pitch.horm.EPratio <- lmer(rating.c ~ estr.s + prog.s + ep_ratio.s +
(1 | oc_id) +
(0 + estr.s + prog.s + ep_ratio.s || oc_id) +
(1 | block:oc_id) +
(0 + estr.s + prog.s + ep_ratio.s || block:oc_id),
data = filter(data_hormones_post_2013, manip=="pitch" ),
REML = FALSE)
summary(model.pitch.horm.EPratio)
```

```
Linear mixed model fit by maximum likelihood . t-tests use
Satterthwaite's method [lmerModLmerTest]
Formula: rating.c ~ estr.s + prog.s + ep_ratio.s + (1 | oc_id) + (0 +
estr.s + prog.s + ep_ratio.s || oc_id) + (1 | block:oc_id) +
(0 + estr.s + prog.s + ep_ratio.s || block:oc_id)
Data: filter(data_hormones_post_2013, manip == "pitch")

      AIC      BIC logLik deviance df.resid
3364.5  3435.8 -1669.3    3338.5     1773

Scaled residuals:
    Min      1Q Median      3Q      Max
-4.6042 -0.5502 -0.0313  0.5497  4.4850

Random effects:
Groups        Name        Variance Std.Dev.
block.oc_id   ep_ratio.s 1.301e-17 3.607e-09
block.oc_id.1  prog.s     1.142e-14 1.069e-07
block.oc_id.2  estr.s     0.000e+00 0.000e+00
block.oc_id.3  (Intercept) 2.278e-02 1.509e-01
oc_id         ep_ratio.s 0.000e+00 0.000e+00
oc_id.1       prog.s     0.000e+00 0.000e+00
oc_id.2       estr.s     0.000e+00 0.000e+00
oc_id.3       (Intercept) 1.665e-01 4.081e-01
Residual      2.895e-01 5.380e-01
Number of obs: 1786, groups: block:oc_id, 378; oc_id, 300

Fixed effects:
            Estimate Std. Error      df t value Pr(>|t|)    
(Intercept)  0.70363   0.02827 298.11748 24.887 <2e-16 ***
estr.s       0.22119   0.09371 1387.61739  2.360  0.0184 *  
prog.s      -0.11147   0.08777 1497.56504 -1.270  0.2043    
ep_ratio.s  -0.06129   0.04793 1479.08990 -1.279  0.2011    
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Correlation of Fixed Effects:
            (Intr) estr.s prog.s ep_ratio.s
estr.s      0.024
prog.s     -0.007 -0.389
ep_ratio.s  0.003 -0.231  0.504
```

Hide

```
confint(model.pitch.horm.EPratio, method = "Wald") %>%
  as.data.frame() %>%
  rownames_to_column() %>%
  filter(!is.na(`2.5 %`))
```

```
rownome      2.5 %    97.5 %
1 (Intercept) 0.64821931 0.75904632
2      estr.s  0.03752448 0.40486529
3      prog.s -0.28348577 0.06055463
4   ep_ratio.s -0.15522511 0.03264313
```

Model 3: Pitch ~ T + C

[Hide](#)

```
model.pitch.horm.TC <- lmer(rating.c ~ test.s + cort.s +
                           (1 | oc_id) +
                           (0 + test.s + cort.s || oc_id) +
                           (1 | block:oc_id) +
                           (0 + test.s + cort.s|| block:oc_id),
                           data = filter(data_hormones_post_2013, manip=="pitch"),
                           REML = FALSE)
summary(model.pitch.horm.TC)
```

```
Linear mixed model fit by maximum likelihood . t-tests use
Satterthwaite's method [lmerModLmerTest]

Formula:
rating.c ~ test.s + cort.s + (1 | oc_id) + (0 + test.s + cort.s ||
oc_id) + (1 | block:oc_id) + (0 + test.s + cort.s || block:oc_id)
Data: filter(data_hormones_post_2013, manip == "pitch")

      AIC      BIC logLik deviance df.resid
3400.4   3455.4 -1690.2    3380.4     1797

Scaled residuals:
    Min      1Q Median      3Q      Max
-4.6481 -0.5516 -0.0444  0.5353  4.3748

Random effects:
Groups        Name        Variance Std.Dev.
block.oc_id   cort.s     7.511e-02 2.741e-01
block.oc_id.1 test.s     0.000e+00 0.000e+00
block.oc_id.2 (Intercept) 2.692e-02 1.641e-01
oc_id         cort.s     2.051e-14 1.432e-07
oc_id.1       test.s     0.000e+00 0.000e+00
oc_id.2       (Intercept) 1.604e-01 4.005e-01
Residual      2.868e-01 5.356e-01
Number of obs: 1807, groups: block:oc_id, 377; oc_id, 299

Fixed effects:
            Estimate Std. Error          df t value Pr(>|t|)    
(Intercept)  0.70210   0.02809 294.85214 24.991 <2e-16 ***
test.s       0.14427   0.08662 1305.82025  1.666   0.096 .
cort.s      -0.11320   0.07089  113.69237 -1.597   0.113  
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Correlation of Fixed Effects:
  (Intr) test.s
test.s -0.005
cort.s  0.010 -0.446
```

Hide

```
confint(model.pitch.horm.TC, method = "Wald") %>%
  as.data.frame() %>%
  rownames_to_column() %>%
  filter(!is.na(`^2.5 %`))
```

```
rownames      2.5 %    97.5 %
1 (Intercept) 0.64703490 0.75716014
2 test.s     -0.02550343 0.31404172
3 cort.s     -0.25214760 0.02573793
```

Data subset: data_hormones_partner (with partnership status)

Only women with consistently reported partnership status for tests of effects of endogenous hormones considering effects of partnership status. Because women who reported their partnership status inconsistently are not included in partnership analyses, Ns for all partnership analyses will be slightly smaller.

[Hide](#)

```
data_hormones_partner <- data_hormones %>%
  filter(block_partner == "0" | block_partner == "1") %>%
  group_by(oc_id) %>%
  mutate(pchange = mean(partner.e)) %>%
  ungroup() %>%
  filter(pchange %in% c(-.5, .5)) %>%
  select(-pchange)
```

[Hide](#)

```
data_hormones_partner %>%
  group_by(manip, oc_id, partner.e) %>%
  summarise(overall_rating.c = mean(rating.c)) %>%
  group_by(manip, partner.e) %>%
  summarise(
    n = n_distinct(oc_id),
    mean_dv = mean(overall_rating.c),
    sd_dv = sd(overall_rating.c),
    se_dv = se(overall_rating.c)
  )
```

manip	partner.e	n	mean_dv	sd_dv	se_dv
<fctr>	<dbl>	<int>	<dbl>	<dbl>	<dbl>
formant	-0.5	211	0.4213841	0.6505473	0.04478552
formant	0.5	111	0.5355802	0.6038185	0.05731191
pitch	-0.5	211	0.7098084	0.5162464	0.03553987
pitch	0.5	111	0.6884867	0.4461727	0.04234884
4 rows					

Masculinity preference (formants)

Model 1: formants ~ E + P + E x P (with partnership status)

Hide

```
model.formants.horm.EP.partner <- summary(lmer(rating.c ~ estr.s * prog.s * partner.e +
                                                (1 | oc_id) +
                                                (0 + estr.s:prog.s || oc_id) +
                                                (1 | block:oc_id) +
                                                (0 + estr.s:prog.s || block:oc_id),
                                                data = filter(data_hormones_partner, manip =
                                                "formant"),
                                                REML = FALSE))
summary(model.formants.horm.EP.partner)
```

Linear mixed model fit by maximum likelihood . t-tests use
Satterthwaite's method [lmerModLmerTest]
Formula:
rating.c ~ estr.s * prog.s * partner.e + (1 | oc_id) + (0 + estr.s:prog.s ||
oc_id) + (1 | block:oc_id) + (0 + estr.s:prog.s || block:oc_id)
Data: filter(data_hormones_partner, manip == "formant")

AIC	BIC	logLik	deviance	df.resid
3515.0	3586.6	-1744.5	3489.0	1820

Scaled residuals:

Min	1Q	Median	3Q	Max
-4.3911	-0.5376	-0.0129	0.5277	3.1144

Random effects:

Groups	Name	Variance	Std.Dev.
block.oc_id	estr.s:prog.s	3.122e-12	1.767e-06
block.oc_id.1	(Intercept)	3.536e-02	1.880e-01
oc_id	estr.s:prog.s	2.398e-14	1.548e-07
oc_id.1	(Intercept)	3.210e-01	5.666e-01
Residual		2.673e-01	5.170e-01

Number of obs: 1833, groups: block:oc_id, 390; oc_id, 322

Fixed effects:

	Estimate	Std. Error	df	t value	Pr(> t)
(Intercept)	0.47626	0.03754	335.10252	12.687	<2e-16
estr.s	0.02603	0.09129	1436.68182	0.285	0.776
prog.s	-0.03960	0.07438	1526.87566	-0.532	0.595
partner.e	0.11291	0.07508	335.10252	1.504	0.134

```
estr.s:prog.s          0.43914   0.44309 1591.72875   0.991   0.322
estr.s:partner.e       0.03182   0.18257 1436.68182   0.174   0.862
prog.s:partner.e      -0.14026   0.14876 1526.87566  -0.943   0.346
estr.s:prog.s:partner.e 0.05811   0.88619 1591.72875   0.066   0.948

(Intercept) ***

estr.s
prog.s
partner.e
estr.s:prog.s
estr.s:partner.e
prog.s:partner.e
estr.s:prog.s:partner.e
---
Signif. codes:  0 '****' 0.001 '***' 0.01 '**' 0.05 '*' 0.1 '.' 1

Correlation of Fixed Effects:
              (Intr) estr.s prog.s prtnr. estr.s:prg. estr.s:prt. prg.::
estr.s        0.014
prog.s        0.007 -0.315
partner.e     0.312  0.009 -0.009
estr.s:prg.  -0.101 -0.007 -0.129 -0.020
estr.s:prt.   0.009  0.335 -0.056  0.014  0.014
prg.s:prtn.  -0.009 -0.056  0.167  0.007  0.033      -0.315
estr.s:p...   -0.020  0.014  0.033 -0.101  0.243      -0.007      -0.129
```

Model 2: formants ~ E + P + EP_ratio (with partnership status)

[Hide](#)

```
model.formants.horm.EPratio.partner <- summary(lmer(rating.c ~ estr.s * partner.e +
+                                         prog.s * partner.e +
+                                         ep_ratio.s * partne
r.e +
+                                         (1 | oc_id) +
+                                         (0 + estr.s + prog.s + ep_ratio.s || oc
_id) +
+                                         (1 | block:oc_id) +
+                                         (0 + estr.s + prog.s + ep_ratio.s || bl
ock:oc_id),
+                                         data = filter(data_hormones_partner,
manip=="formant" ),
+                                         REML = FALSE))
```

Model failed to converge with 1 negative eigenvalue: -4.2e+01

Hide

summary(model.formants.horm.EPratio.partner)

Linear mixed model fit by maximum likelihood . t-tests use Satterthwaite's method [lmerModLmerTest]
Formula: rating.c ~ estr.s * partner.e + prog.s * partner.e + ep_ratio.s * partner.e + (1 | oc_id) + (0 + estr.s + prog.s + ep_ratio.s || oc_id) + (1 | block:oc_id) + (0 + estr.s + prog.s + ep_ratio.s || block:oc_id)
Data: filter(data_hormones_partner, manip == "formant")

AIC	BIC	logLik	deviance	df.resid
3519.2	3613.0	-1742.6	3485.2	1816

Scaled residuals:

Min	1Q	Median	3Q	Max
-4.2753	-0.5256	-0.0169	0.5215	3.0747

Random effects:

Groups	Name	Variance	Std.Dev.
block.oc_id	ep_ratio.s	2.570e-10	1.603e-05
block.oc_id.1	prog.s	1.204e-10	1.097e-05
block.oc_id.2	estr.s	2.683e-01	5.179e-01
block.oc_id.3	(Intercept)	3.318e-02	1.821e-01
oc_id	ep_ratio.s	6.859e-03	8.282e-02
oc_id.1	prog.s	8.857e-02	2.976e-01
oc_id.2	estr.s	0.000e+00	0.000e+00
oc_id.3	(Intercept)	3.224e-01	5.678e-01
Residual		2.568e-01	5.067e-01

Number of obs: 1833, groups: block:oc_id, 390; oc_id, 322

Fixed effects:

	Estimate	Std. Error	df	t value	Pr(> t)
(Intercept)	0.480203	0.037265	327.251341	12.886	<2e-16 ***
estr.s	0.031984	0.101006	175.990404	0.317	0.752
partner.e	0.113653	0.074529	327.251341	1.525	0.128
prog.s	-0.033168	0.090597	198.515487	-0.366	0.715
ep_ratio.s	-0.009857	0.048937	27.009921	-0.201	0.842
estr.s:partner.e	0.070518	0.202012	175.990404	0.349	0.727
partner.e:prog.s	-0.170465	0.181194	198.515487	-0.941	0.348
partner.e:ep_ratio.s	0.009344	0.097873	27.009921	0.095	0.925

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

Correlation of Fixed Effects:

(Intr) estr.s prtnr. prog.s ep_rt. est... prt...

```
estr.s      0.015
partner.e   0.314  0.011
prog.s     -0.007 -0.349 -0.006
ep_ratio.s -0.002 -0.187 -0.001  0.510
estr.s:prt. 0.011  0.336  0.015 -0.050  0.011
prtnr.:prg. -0.006 -0.050 -0.007  0.136 -0.061 -0.349
prtnr.:p_r. -0.001  0.011 -0.002 -0.061 -0.214 -0.187  0.510
```

Model 3: formants ~ T + C (with partnership status)

[Hide](#)

```
model.formants.horm.TC.partner <- summary(lmer(rating.c ~ test.s * partner.e + cor
t.s * partner.e +
                               (1 | oc_id) +
                               (0 + test.s + cort.s || oc_id) +
                               (1 | block:oc_id) +
                               (0 + test.s + cort.s|| block:oc_id),
                               data = filter(data_hormones_partner, manip
== "formant"),
                               REML = FALSE))
summary(model.formants.horm.TC.partner)
```

```
Linear mixed model fit by maximum likelihood . t-tests use
Satterthwaite's method [lmerModLmerTest]

Formula:
rating.c ~ test.s * partner.e + cort.s * partner.e + (1 | oc_id) +
(0 + test.s + cort.s || oc_id) + (1 | block:oc_id) + (0 +
test.s + cort.s || block:oc_id)
Data: filter(data_hormones_partner, manip == "formant")

      AIC      BIC logLik deviance df.resid
3562.3  3634.1 -1768.2    3536.3     1838

Scaled residuals:
    Min      1Q Median      3Q      Max
-4.4801 -0.5378 -0.0130  0.5259  3.0331

Random effects:
Groups        Name        Variance Std.Dev.
block.oc_id   cort.s     1.099e-15 3.315e-08
block.oc_id.1 test.s     3.975e-16 1.994e-08
block.oc_id.2 (Intercept) 3.560e-02 1.887e-01
oc_id         cort.s     0.000e+00 0.000e+00
oc_id.1       test.s     0.000e+00 0.000e+00
oc_id.2       (Intercept) 3.229e-01 5.682e-01
Residual                  2.701e-01 5.197e-01
Number of obs: 1851, groups: block:oc_id, 389; oc_id, 321

Fixed effects:
            Estimate Std. Error          df t value Pr(>|t|)    
(Intercept)  0.48104   0.03743  325.30470 12.852 <2e-16 ***
test.s      -0.07201   0.08752 1525.96801 -0.823  0.411  
partner.e    0.11504   0.07486  325.30470  1.537  0.125  
cort.s      -0.04674   0.06797 1500.30638 -0.688  0.492  
test.s:partner.e -0.19119   0.17504 1525.96801 -1.092  0.275  
partner.e:cort.s  0.08494   0.13595 1500.30638  0.625  0.532  
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Correlation of Fixed Effects:
            (Intr) test.s prtnr. cort.s tst.::
test.s      -0.004
partner.e    0.309 -0.002
cort.s      0.009 -0.402  0.010
tst.s:prtn. -0.002  0.223 -0.004 -0.044
prtnr.:crt.  0.010 -0.044  0.009  0.288 -0.402
```

Masculinity preference (Pitch)

Model 1: Pitch ~ E + P + E x P (with partnership status)

Hide

```
model.pitch.horm.EP.partner <- summary(lmer(rating.c ~ estr.s * prog.s * partner.e
+
(1 | oc_id) +
(0 + estr.s:prog.s || oc_id) +
(1 | block:oc_id) +
(0 + estr.s:prog.s || block:oc_id),
data = filter(data_hormones_partner, manip=="pitch"),
REML = FALSE))
summary(model.pitch.horm.EP.partner)
```

Linear mixed model fit by maximum likelihood . t-tests use Satterthwaite's method [lmerModLmerTest]

Formula:

rating.c ~ estr.s * prog.s * partner.e + (1 | oc_id) + (0 + estr.s:prog.s || oc_id) + (1 | block:oc_id) + (0 + estr.s:prog.s || block:oc_id)

Data: filter(data_hormones_partner, manip == "pitch")

AIC	BIC	logLik	deviance	df.resid
3430.9	3502.5	-1702.4	3404.9	1820

Scaled residuals:

Min	1Q	Median	3Q	Max
-4.6732	-0.5421	-0.0373	0.5358	3.5651

Random effects:

Groups	Name	Variance	Std.Dev.
block.oc_id	estr.s:prog.s	0.00000	0.0000
block.oc_id.1	(Intercept)	0.03069	0.1752
oc_id	estr.s:prog.s	0.00000	0.0000
oc_id.1	(Intercept)	0.16671	0.4083
Residual		0.28131	0.5304

Number of obs: 1833, groups: block:oc_id, 390; oc_id, 322

Fixed effects:

	Estimate	Std. Error	df	t value	Pr(> t)
(Intercept)	0.70095	0.02952	337.55096	23.748	<2e-16
estr.s	0.19899	0.09302	1422.08017	2.139	0.0326
prog.s	-0.01293	0.07610	1534.45034	-0.170	0.8651
partner.e	-0.01197	0.05903	337.55096	-0.203	0.8394
estr.s:prog.s	-0.03205	0.44922	1649.07416	-0.071	0.9431
estr.s:partner.e	-0.03168	0.18604	1422.08017	-0.170	0.8648
prog.s:partner.e	0.13569	0.15221	1534.45034	0.891	0.3728

```
estr.s:prog.s:partner.e -0.17530 0.89844 1649.07416 -0.195 0.8453
```

(Intercept) ***
estr.s *

prog.s
partner.e
estr.s:prog.s
estr.s:partner.e
prog.s:partner.e
estr.s:prog.s:partner.e

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

Correlation of Fixed Effects:

	(Intr)	estr.s	prog.s	prtnr.	estr.s:prg.	estr.s:prt.	prg... estr.s	0.018	prog.s	0.009 -0.314	partner.e	0.313 0.011 -0.012	estr.s:prg.	-0.132 -0.005 -0.127 -0.026	estr.s:prt.	0.011 0.332 -0.056 0.018 0.015	prg.s:prtn.	-0.012 -0.056 0.168 0.009 0.034	estr.s:p... estr.s	-0.026 0.015 0.034 -0.132 0.242	estr.s	-0.005 -0.314 -0.005 -0.127
--	--------	--------	--------	--------	-------------	-------------	------------------	-------	--------	--------------	-----------	--------------------	-------------	-----------------------------	-------------	--------------------------------	-------------	---------------------------------	-----------------------	---------------------------------	--------	-----------------------------

Model 2: Pitch ~ E + P + EP_ratio (with partnership status)

Hide

```
model.pitch.horm.EPratio.partner <- summary(lmer(rating.c ~ estr.s * partner.e +  
                                                 prog.s * partner.e +  
                                                 ep_ratio.s * partner.e  
+  
                                                 (1 | oc_id) +  
                                                 (0 + estr.s + prog.s + ep_ratio.s || o  
c_id) +  
                                                 (1 | block:oc_id) +  
                                                 (0 + estr.s + prog.s + ep_ratio.s || b  
lock:oc_id),  
                                                 data = filter(data_hormones_partner, man  
ip=="pitch" ),  
                                                 REML = FALSE))  
summary(model.pitch.horm.EPratio.partner)
```

```
Linear mixed model fit by maximum likelihood . t-tests use  
Satterthwaite's method [lmerModLmerTest]  
Formula: rating.c ~ estr.s * partner.e + prog.s * partner.e + ep_ratio.s *  
partner.e + (1 | oc_id) + (0 + estr.s + prog.s + ep_ratio.s ||  
oc_id) + (1 | block:oc_id) + (0 + estr.s + prog.s + ep_ratio.s ||
```

```
block:oc_id)
Data: filter(data_hormones_partner, manip == "pitch")

      AIC      BIC    logLik deviance df.resid
3437.5  3531.2 -1701.8    3403.5     1816

Scaled residuals:
    Min      1Q  Median      3Q     Max
-4.6684 -0.5427 -0.0400  0.5369  3.5917

Random effects:
Groups        Name        Variance Std.Dev.
block:oc_id   ep_ratio.s 0.000e+00 0.000e+00
block:oc_id.1  prog.s    2.363e-14 1.537e-07
block:oc_id.2  estr.s    8.355e-03 9.141e-02
block:oc_id.3  (Intercept) 3.099e-02 1.760e-01
oc_id         ep_ratio.s 0.000e+00 0.000e+00
oc_id.1       prog.s    0.000e+00 0.000e+00
oc_id.2       estr.s    1.056e-15 3.249e-08
oc_id.3       (Intercept) 1.665e-01 4.080e-01
Residual      2.808e-01 5.299e-01
Number of obs: 1833, groups: block:oc_id, 390; oc_id, 322

Fixed effects:
            Estimate Std. Error      df t value Pr(>|t|)    
(Intercept) 7.008e-01 2.925e-02 3.261e+02 23.955 <2e-16 ***
estr.s      2.208e-01 9.506e-02 1.550e+02  2.323  0.0215 *  
partner.e   -1.328e-02 5.851e-02 3.261e+02 -0.227  0.8205  
prog.s      -6.236e-02 8.685e-02 1.505e+03 -0.718  0.4729  
ep_ratio.s  -4.933e-02 4.486e-02 1.340e+03 -1.100  0.2717  
estr.s:partner.e -2.000e-02 1.901e-01 1.550e+02 -0.105  0.9164  
partner.e:prog.s  1.199e-01 1.737e-01 1.505e+03  0.690  0.4900  
partner.e:ep_ratio.s 1.614e-03 8.972e-02 1.340e+03  0.018  0.9856  
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Correlation of Fixed Effects:
              (Intr) estr.s prtnr. prog.s ep_rt. est.. prt..
estr.s      0.017
partner.e   0.316  0.014
prog.s     -0.008 -0.367 -0.007
ep_ratio.s -0.002 -0.186 -0.002  0.494
estr.s:prt.  0.014  0.327  0.017 -0.050  0.018
prtnr.:prg. -0.007 -0.050 -0.008  0.112 -0.121 -0.367
prtnr.:p_r. -0.002  0.018 -0.002 -0.121 -0.370 -0.186  0.494
```

Model 3: Pitch ~ T + C (with partnership status)

Hide

```
model.pitch.horm.TC.partner <- summary(lmer(rating.c ~ test.s * partner.e + cort.s  
* partner.e +  
          (1 | oc_id) +  
          (0 + test.s + cort.s || oc_id) +  
          (1 | block:oc_id) +  
          (0 + test.s + cort.s|| block:oc_id),  
          data = filter(data_hormones_partner, manip=="  
pitch"),  
          REML = FALSE))  
summary(model.pitch.horm.TC.partner)
```

```
Linear mixed model fit by maximum likelihood . t-tests use
Satterthwaite's method [lmerModLmerTest]

Formula:
rating.c ~ test.s * partner.e + cort.s * partner.e + (1 | oc_id) +
(0 + test.s + cort.s || oc_id) + (1 | block:oc_id) + (0 +
test.s + cort.s || block:oc_id)
Data: filter(data_hormones_partner, manip == "pitch")

      AIC      BIC    logLik deviance df.resid
3468.9   3540.7   -1721.4     3442.9      1838

Scaled residuals:
    Min      1Q  Median      3Q      Max
-4.7166 -0.5505 -0.0500  0.5316  3.4218

Random effects:
Groups        Name        Variance Std.Dev.
block.oc_id   cort.s     0.09798  0.3130
block.oc_id.1 test.s     0.00000  0.0000
block.oc_id.2 (Intercept) 0.03513  0.1874
oc_id         cort.s     0.00000  0.0000
oc_id.1       test.s     0.00000  0.0000
oc_id.2       (Intercept) 0.16031  0.4004
Residual      0.27846  0.5277
Number of obs: 1851, groups: block:oc_id, 389; oc_id, 321

Fixed effects:
            Estimate Std. Error          df t value Pr(>|t|)    
(Intercept)  0.69932   0.02904  323.53575  24.083 <2e-16 ***
test.s       0.11534   0.09064 1377.87798   1.273   0.203  
partner.e    -0.02552   0.05807  323.53575  -0.439   0.661  
cort.s       -0.06863   0.08001  101.67030  -0.858   0.393  
test.s:partner.e  0.01896   0.18129 1377.87798   0.105   0.917  
partner.e:cort.s -0.02299   0.16003  101.67030  -0.144   0.886  
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Correlation of Fixed Effects:
            (Intr) test.s prtnr. cort.s tst.::
test.s     -0.006
partner.e    0.310 -0.004
cort.s      0.015 -0.410  0.015
tst.s:prtn. -0.004  0.237 -0.006 -0.085
prtnr.:crt.  0.015 -0.085  0.015  0.371 -0.410
```

Average Progesterone by Partnership Status

Testing whether the combined effects of partnership status and average progesterone predict masculinity preference.

[Hide](#)

```
data_avg <- data_hormones_partner %>%
  group_by(oc_id, manip, partner.e) %>%
  summarise(AP = mean(prog, na.rm = T),
            mean_rating = mean(rating.c)) %>%
  ungroup()
```

[Hide](#)

```
model.ap.formant <- lm(mean_rating ~ AP * partner.e,
                        data = filter(data_avg, manip == "formant"))
summary(model.ap.formant)
```

Call:

```
lm(formula = mean_rating ~ AP * partner.e, data = filter(data_avg,
  manip == "formant"))
```

Residuals:

Min	1Q	Median	3Q	Max
-2.26681	-0.41929	-0.05476	0.44126	2.13349

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	0.5165353	0.0899226	5.744	2.16e-08 ***
AP	-0.0002487	0.0005366	-0.464	0.643
partner.e	0.1494925	0.1798452	0.831	0.406
AP:partner.e	-0.0002262	0.0010731	-0.211	0.833

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

Residual standard error: 0.6366 on 318 degrees of freedom

Multiple R-squared: 0.00803, Adjusted R-squared: -0.001328

F-statistic: 0.8581 on 3 and 318 DF, p-value: 0.4632

[Hide](#)

```
model.ap.pitch <- lm(mean_rating ~ AP * partner.e,
                      data = filter(data_avg, manip == "pitch"))
summary(model.ap.pitch)
```

Call:

```
lm(formula = mean_rating ~ AP * partner.e, data = filter(data_avg,  
manip == "pitch"))
```

Residuals:

Min	1Q	Median	3Q	Max
-1.19762	-0.35902	-0.06594	0.29976	1.66121

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	0.7682743	0.0697113	11.021	<2e-16 ***
AP	-0.0004566	0.0004160	-1.098	0.273
partner.e	-0.0820151	0.1394227	-0.588	0.557
AP:partner.e	0.0004095	0.0008319	0.492	0.623

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

Residual standard error: 0.4935 on 318 degrees of freedom

Multiple R-squared: 0.005633, Adjusted R-squared: -0.003747

F-statistic: 0.6005 on 3 and 318 DF, p-value: 0.6151