Cross-cultural relationships between music, emotion, and visual imagery: A comparative study of Iran, Canada, and Japan [Stage 1 Registered Report]

Code and analyses: Simulation-based power analysis

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Descripción

This document contains all code, and step by step explanations for all analyses, figures and tables (including supplementary figures and tables) for:

hadavi, S., Kuroda, J., Shimozono, T., Leongómez, J. D., & Savage, P. E. (in prep). Cross-cultural relationships between music, emotion, and visual imagery: A comparative study of Iran, Canada, and Japan [Stage 1 Registered Report]. https://doi.org/10.31234/osf.io/26yg5

Data available from the Open Science Framework (OSF): https://doi.org/10.XXXXX/OSF.IO/XXXXXX. This document and its underlying code were created in R Markdown by Juan David Leongómez using LATEX.

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	Session info (for reproducibility)

1 Preliminaries

1.1 Load packages

This file was created using knitr (Xie, 2014), mostly using tidyverse (Wickham et al., 2019) syntax. As such, data wrangling was mainly done using packages such as dplyr (Wickham et al., 2022), and most figures were created using ggplot2 (Wickham, 2016).

Cumulative Link Mixed Models (CLMM) were fitted using ordinal (Christensen, 2022), and contrasts and interactions were explored using emmeans (Lenth, 2022).

All packages used in this file can be directly installed from the Comprehensive R Archive Network (CRAN). For a complete list of packages used to create this file, and their versions, see section 5, at the end of the document.

```
library(tidyverse)
library(ggpubr)
library(emmeans)
library(ordinal)
```

1.2 Custom functions

To simulate ordinal data with a specific distribution, we used the function clmm_generate_data, created by Borders et al. (2022) as part of their tutorial (https://osf.io/e6usd/). To run this function, a number of other custom functions need to be defined and implemented as well. The original code can be found in the clmm-power-library.R file (https://osf.io/tjpkf). Here, it is run from source, but in a version oif the file changing the mc.cores parameter to 1, to avoid errors.

```
source("clmm-power-library.R", local = knitr::knit_global())
```

2 Data

2.1 Pilot data

To simulate data, first we looked at the distribution of the pilot data.

```
# Load data
sh1 <- read.csv("../sh1.csv") |>
    mutate(Tempo = str_replace(Tempo, "low", "Low")) |>
    mutate(Tempo = str_replace(Tempo, "high", "High")) |>
    mutate_if(is.character,as.factor) |>
    mutate(Tempo = fct_relevel(Tempo, c("Low", "High")))

# Assign unique id to pairs
uniqueval <- unique(sh1[, c("Participant", "Music.country", "Solo.group")])
sh1$groupid <- 0
for (i in 1:dim(uniqueval)[1]) {
    idx <- sh1$Participant == uniqueval$Participant[i] &
        sh1$Music.country == uniqueval$Music.country[i] &
        sh1$Solo.group == uniqueval$Solo.group[i]
    sh1$groupid[idx] <- i
}</pre>
```

```
data <- sh1 |>
  select(1:9,20)
p1a <- ggplot(data, aes(x = density.tempo, y = after_stat(density),</pre>
                       fill = Tempo, color = Tempo)) +
  scale_fill_hue(direction = -1) + scale_colour_hue(direction = -1) +
  geom_histogram(alpha = 0.3, position = "identity", binwidth = 1) +
  labs(y= "Probability", x = "Rating", title = "Density") +
  geom_text(aes(label = format(after_stat(density), digits = 1), y= after_stat(density)),
            stat= "bin", binwidth = 1,
            vjust = -0.2,
            show.legend = FALSE) +
  facet_wrap(~Solo.group)
p1b <- ggplot(data, aes(x = arousal.tempo, y = after_stat(density),
                       fill = Tempo, color = Tempo)) +
  scale_fill_hue(direction = -1) + scale_colour_hue(direction = -1) +
  geom_histogram(alpha = 0.3, position = "identity", binwidth = 1) +
  labs(y= NULL, x = "Rating", title = "Arousal") +
  geom_text(aes(label = format(after_stat(density), digits = 1), y= after_stat(density)),
            vjust = -0.2,
            show.legend = FALSE) +
  facet_wrap(~Solo.group)
p1 <- ggarrange(p1a, p1b,
                common.legend = TRUE,
                legend = "bottom",
                labels = "AUTO")
p1
```

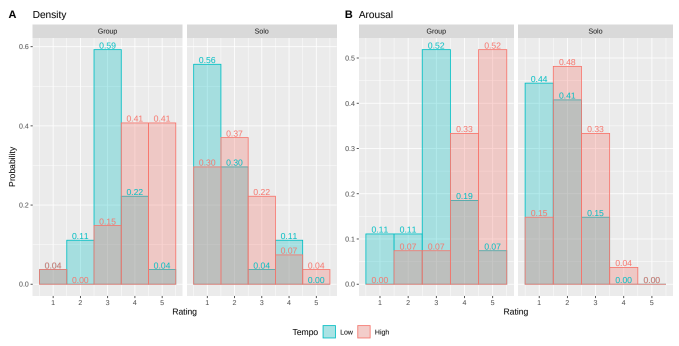


Figure S1. Distribution of results from pilot data for ratings of both Density (**A**) and Arousal (**B**), by Instrumentation (Group, Solo), and Tempo (High, Low).

2.2 Data simulation

While basing a power analysis on pilot data is problematic and can bias the sample size estimation (see e.g., Albers & Lakens, 2018), we used this distribution as a starting point.

The clmm_generate_data function (Borders et al., 2022) allows to simulate data with random and fixed effects with an ordinal outcome. In this case, we simulated data for Tempo, the within-subject variable for which there was an a-priori prediction, replicating the distribution of the first level (Low) using a specified probability of each rating score (using the argument control_distribution). The second level of that within-subject variable (Tempo = High) is generated based on a hypothesized effect (using the argument effect). Other important factors such as participant country and music country were randomly assigned, as there are no a-priori predictions regarding these factors.

2.2.1 Replication of the pilot data distribution

To obtain distributions similar to the probabilities observed in the pilot data, we first simulated individual data for Density and Arousal ratings, separated by Instrumentation type (Group, Solo). In all cases, we simulated data from 10000 participants, and changed the value assigned to the effect argument until the distribution of the second level of the within-subject variable (Tempo = High) resembled the pilot data.

```
select(1:3) |>
  mutate(pair = rep(1:3, times = 20000)) |>
 mutate(Participant.country = rep(rep(c("Iran", "Canada", "Japan"), each = 10000), 2)) |>
 mutate(Music.country = rep(c("Iran", "Canada", "Japan"), times = 20000)) |>
 mutate(Solo.group = rep("Group", TIMES = 60000)) |>
 select(c(1,6,5,2,7,3))
dat.den.so.PILOT <- clmm_generate_data(n_participants = 10000,
                                  n_trials = 3,
                                  control_distribution = c(.55, .30, .04, .10, .01),
                                  effect = 1.2,
                                  participant_variation = 1,
                                  within_subject = TRUE,
                                  control_weight = .5) |>
 mutate(group = ifelse(group == 0, "Low", "High")) |>
  dplyr::rename(Density.tempo = pas) |>
 dplyr::rename(Tempo = group) |>
  dplyr::rename(Participant = id) |>
 mutate(Participant = paste0("p", Participant)) |>
 select(1:3) |>
 mutate(pair = rep(1:3, times = 20000)) |>
 mutate(Participant.country = rep(rep(c("Iran", "Canada", "Japan"), each = 10000), 2)) |>
 mutate(Music.country = rep(c("Iran", "Canada", "Japan"), times = 20000)) |>
 mutate(Solo.group = rep("Solo", TIMES = 60000)) |>
  select(c(1,6,5,2,7,3))
dat.aro.gr.PILOT <- clmm_generate_data(n_participants = 10000,</pre>
                                  n trials = 3,
                                  control_distribution = c(.11, .11, .52, .19, .07),
                                  effect = 3.1,
                                  participant_variation = 1,
                                  within_subject = TRUE,
                                  control weight = .5) |>
 mutate(group = ifelse(group == 0, "Low", "High")) |>
 dplyr::rename(Arousal.tempo = pas) |>
  dplyr::rename(Tempo = group) |>
 dplyr::rename(Participant = id) |>
 mutate(Participant = paste0("p", Participant)) |>
 select(1:3) |>
 mutate(pair = rep(1:3, times = 20000)) |>
 mutate(Participant.country = rep(rep(c("Iran", "Canada", "Japan"), each = 10000), 2)) |>
mutate(Music.country = rep(c("Iran", "Canada", "Japan"), times = 20000)) |>
 mutate(Solo.group = rep("Group", TIMES = 60000)) |>
 select(c(1,6,5,2,7,3))
dat.aro.so.PILOT <- clmm_generate_data(n_participants = 10000,</pre>
                                  n_trials = 3,
                                  control_distribution = c(.43, .41, .14, .01, .01),
                                  effect = 1.6,
                                  participant_variation = 1,
                                  within_subject = TRUE,
                                  control_weight = .5) |>
 mutate(group = ifelse(group == 0, "Low", "High")) |>
```

```
dplyr::rename(Arousal.tempo = pas) |>
dplyr::rename(Tempo = group) |>
dplyr::rename(Participant = id) |>
mutate(Participant = paste0("p", Participant)) |>
select(1:3) |>
mutate(pair = rep(1:3, times = 20000)) |>
mutate(Participant.country = rep(rep(c("Iran", "Canada", "Japan"), each = 10000), 2)) |>
mutate(Music.country = rep(c("Iran", "Canada", "Japan"), times = 20000)) |>
mutate(Solo.group = rep("Solo", TIMES = 60000)) |>
select(c(1,6,5,2,7,3))
```

We then merged these four data frames to obtain a simulated data frame.

The distribution of this replication of the pilot data distribution is represented in S2

```
# Plot distribution of results for density ratings
p2a <- ggplot(dat.sim.PILOT, aes(x = Density.tempo, y = after_stat(density),
                                 fill = Tempo, color = Tempo)) +
  scale_fill_hue(direction = -1) + scale_colour_hue(direction = -1) +
  geom_histogram(alpha = 0.3, position = "identity", binwidth = 1) +
  labs(y= "Probability", x = "Rating") +
  geom_text(aes(label = format(after_stat(density), digits = 1), y= after_stat(density)),
            stat= "bin", binwidth = 1,
            vjust = -0.2,
            show.legend = FALSE) +
  facet_wrap(~Solo.group)
p2b <- ggplot(dat.sim.PILOT, aes(x = Arousal.tempo, y = after_stat(density),
                                 fill = Tempo, color = Tempo)) +
  scale_fill_hue(direction = -1) + scale_colour_hue(direction = -1) +
  geom_histogram(alpha = 0.3, position = "identity", binwidth = 1) +
  labs(y= NULL, x = "Rating") +
  geom_text(aes(label = format(after_stat(density), digits = 1), y= after_stat(density)),
            stat= "bin", binwidth = 1,
            vjust = -0.2,
            show.legend = FALSE) +
  facet_wrap(~Solo.group)
p2 <- ggarrange(p2a +
                  labs(subtitle = "Density"),
                p2b +
                  labs(subtitle = "Arousal"),
                common.legend = TRUE,
                legend = "bottom",
                labels = "AUTO")
p2
```

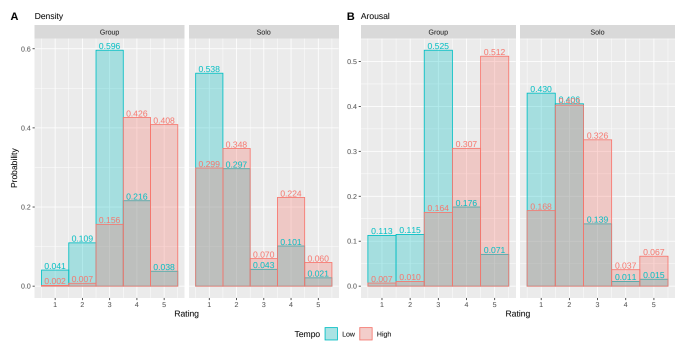


Figure S2. Distribution of results from a simulated replication of the pilot data distribution for ratings of both Density (**A**) and Arousal (**B**), by Instrumentation (Group, Solo), and Tempo (High, Low).

2.2.2 Simulation of data with a more conservative effect

While the data simulated in the previous section resembled the distribution of the pilot data, using these data would be problematic for at least two reasons:

First, the effects of Tempo were extremely large (3.2, 1.2, 3.1, 1.6), particularly for ratings of Group instrumentation (3.2, and 3.1). With such differences, a very high statistical power of about $1 - \beta = 0.95$ can be achieved with 2 or 3 participants. And second, as mentioned before, estimating effects from pilot data is problematic, as it biases the results (Albers & Lakens, 2018).

For this reason, we decided to maintain the distributions of probabilities for the first Tempo level (Low), as it is a good starting point to estimate the distribution of ratings that participants would assign, but simulated the second level (Tempo = High) with a much more conservative effect of 1 in all cases.

```
dat.den.gr <- clmm_generate_data(n_participants = 10000,
                                 n trials = 3,
                                 control_distribution = c(.04, .11, .59, .22, .04),
                                 effect = 1,
                                 participant_variation = 1,
                                 within_subject = TRUE,
                                 control_weight = .5) |>
 mutate(group = ifelse(group == 0, "Low", "High")) |>
 dplyr::rename(Density.tempo = pas) |>
 dplyr::rename(Tempo = group) |>
 dplyr::rename(Participant = id) |>
 mutate(Participant = paste0("p", Participant)) |>
 select(1:3) |>
 mutate(pair = rep(1:3, times = 20000)) |>
 mutate(Participant.country = rep(rep(c("Iran", "Canada", "Japan"), each = 10000), 2)) |>
 mutate(Music.country = rep(c("Iran", "Canada", "Japan"), times = 20000)) |>
 mutate(Solo.group = rep("Group", TIMES = 60000)) |>
  select(c(1,6,5,2,7,3))
```

```
dat.den.so <- clmm generate data(n participants = 10000,
                                  n_trials = 3,
                                  control_distribution = c(.55, .30, .04, .10, .01),
                                  effect = 1,
                                  participant_variation = 1,
                                  within_subject = TRUE,
                                  control_weight = .5) |>
 mutate(group = ifelse(group == 0, "Low", "High")) |>
  dplyr::rename(Density.tempo = pas) |>
 dplyr::rename(Tempo = group) |>
 dplyr::rename(Participant = id) |>
 mutate(Participant = paste0("p", Participant)) |>
  select(1:3) |>
 mutate(pair = rep(1:3, times = 20000)) |>
 mutate(Participant.country = rep(rep(c("Iran", "Canada", "Japan"), each = 10000), 2)) |>
 mutate(Music.country = rep(c("Iran", "Canada", "Japan"), times = 20000)) |>
 mutate(Solo.group = rep("Solo", TIMES = 60000)) |>
 select(c(1,6,5,2,7,3))
dat.aro.gr <- clmm_generate_data(n_participants = 10000,</pre>
                                  n_trials = 3,
                                  control_distribution = c(.11, .11, .52, .19, .07),
                                  effect = 1,
                                  participant_variation = 1,
                                  within_subject = TRUE,
                                  control_weight = .5) |>
 mutate(group = ifelse(group == 0, "Low", "High")) |>
 dplyr::rename(Arousal.tempo = pas) |>
  dplyr::rename(Tempo = group) |>
 dplyr::rename(Participant = id) |>
 mutate(Participant = paste0("p", Participant)) |>
 select(1:3) |>
 mutate(pair = rep(1:3, times = 20000)) |>
 mutate(Participant.country = rep(rep(c("Iran", "Canada", "Japan"), each = 10000), 2)) |>
mutate(Music.country = rep(c("Iran", "Canada", "Japan"), times = 20000)) |>
 mutate(Solo.group = rep("Group", TIMES = 60000)) |>
 select(c(1,6,5,2,7,3))
dat.aro.so <- clmm_generate_data(n_participants = 10000,</pre>
                                  n_trials = 3,
                                  control_distribution = c(.43, .41, .14, .01, .01),
                                  effect = 1,
                                  participant_variation = 1,
                                  within_subject = TRUE,
                                  control_weight = .5) |>
 mutate(group = ifelse(group == 0, "Low", "High")) |>
  dplyr::rename(Arousal.tempo = pas) |>
  dplyr::rename(Tempo = group) |>
 dplyr::rename(Participant = id) |>
 mutate(Participant = paste0("p", Participant)) |>
 select(1:3) |>
 mutate(pair = rep(1:3, times = 20000)) |>
 mutate(Participant.country = rep(rep(c("Iran", "Canada", "Japan"), each = 10000), 2)) |>
```

```
mutate(Music.country = rep(c("Iran", "Canada", "Japan"), times = 20000)) |>
mutate(Solo.group = rep("Solo", TIMES = 60000)) |>
select(c(1,6,5,2,7,3))
```

Again, we then merged these four data frames to obtain a final simulated data frame.

The distribution of these simulated data is represented in S3

```
# Plot distribution of results for density ratings
p3a <- ggplot(dat.sim, aes(x = Density.tempo, y = after_stat(density),
                           fill = Tempo, color = Tempo)) +
  scale_fill_hue(direction = -1) + scale_colour_hue(direction = -1) +
  geom_histogram(alpha = 0.3, position = "identity", binwidth = 1) +
  labs(y= "Probability", x = "Rating") +
  geom_text(aes(label = format(after_stat(density), digits = 1), y= after_stat(density)),
            stat= "bin", binwidth = 1,
            vjust = -0.2,
            show.legend = FALSE) +
  facet_wrap(~Solo.group)
p3b <- ggplot(dat.sim, aes(x = Arousal.tempo, y = after_stat(density),
                           fill = Tempo, color = Tempo)) +
  scale_fill_hue(direction = -1) + scale_colour_hue(direction = -1) +
  geom_histogram(alpha = 0.3, position = "identity", binwidth = 1) +
  labs(y= NULL, x = "Rating") +
  geom_text(aes(label = format(after_stat(density), digits = 1), y= after_stat(density)),
            stat= "bin", binwidth = 1,
            vjust = -0.2,
            show.legend = FALSE) +
  facet_wrap(~Solo.group)
p3 <- ggarrange(p3a +
                  labs(subtitle = "Density"),
                  labs(subtitle = "Arousal"),
                common.legend = TRUE,
                legend = "bottom",
                labels = "AUTO")
рЗ
```

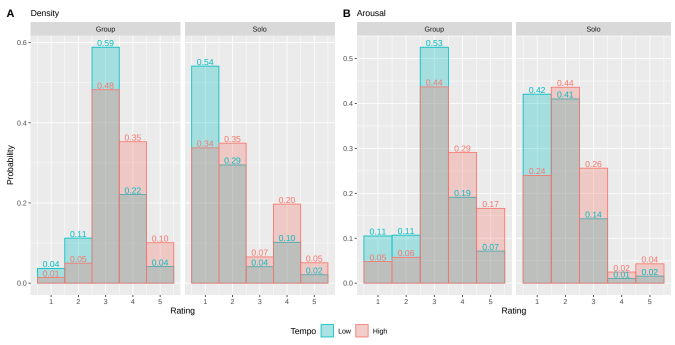


Figure S3. Distribution of results from a conservative simulated distribution for ratings of both Density (**A**) and Arousal (**B**), by Instrumentation (Group, Solo), and Tempo (High, Low).

2.3 Comparisons of the distribuition of results between pilot and simulated data

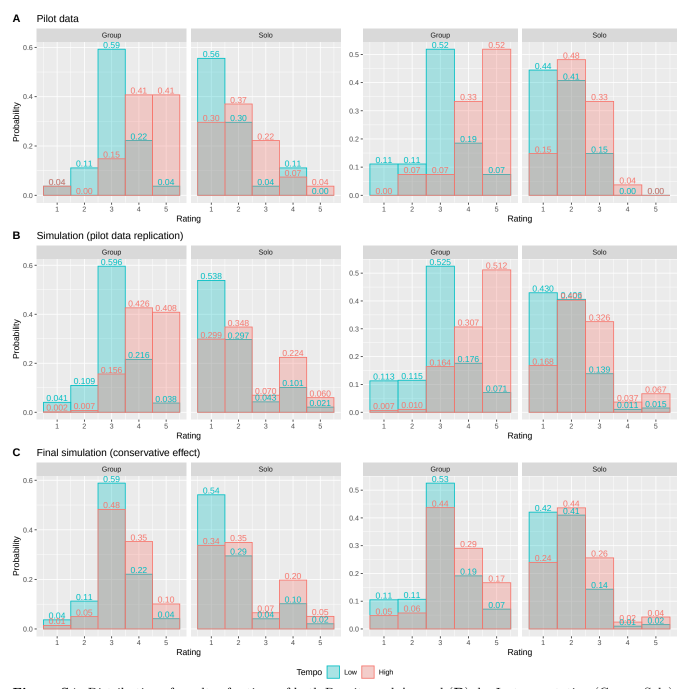


Figure S4. Distribution of results of ratings of both Density and Arousal (**B**), by Instrumentation (Group, Solo), and Tempo (High, Low). **A** pilot data; **B** simulation replicating pilot data distribution; **C** simulation with a more conservative effect.

3 Power analysis

Using the final, simulated population (N=10000 participants, with 6 paired observations per participant), we conducted a simulation-based power analysis. To do this, we first defined the desired number of simulations, the sample size (per country) of each simulation, and the α value (significance level) for all statistical tests:

```
# Number of simulations to run
num_sims.clmm = 100

# Sample size for each simulation (number of participants per country)
```

```
sample_size.clmm = 25

# Significance level
alpha.clmm = 0.05
```

Then, models were fitted from the defined 100 random samples extracted from the simulated population.

From each sample, two Cumulative Link Mixed Models (CLMM) were fitted: one for Density ratings, and one for arousal ratings. All models were fitted with the following call: clmm(DV ~ Tempo * Solo.group * Participant.country * Music.country + (1 + Tempo | Participant).

This is, all models had the same structure, which included the main effects and all possible interactions between Tempo (Low, High), Instrumentation (Group, Solo), Participant country (Iran, Canada, Japan), and Music country (Iran, Canada, Japan) as fixed effects, as well as random intercepts and random slopes between Tempo conditions for each participant.

For each random sample, we adjusted the number of participants per country (in this case, n=25) to achieve the target statistical power of at least $1-\beta=0.95$ for both the Density and Arousal models.

3.1 Select random samples

3.2 Density power analysis

```
message('Error')
             print(e)
           })
  clmm.comps.den$p[i] <- round(data.frame(comp$contrast)$p.value, 3)</pre>
  clmm.comps.den$`Statistical signicance`[i] <- ifelse(clmm.comps.den$p[i] <= alpha.clmm,
## <simpleError in vcov.clm(object, method = "Cholesky"): Cannot compute vcov:
## Hessian is not positive definite>
## <simpleError in vcov.clm(object, method = "Cholesky"): Cannot compute vcov:
## Hessian is not positive definite>
## <simpleError in vcov.clm(object, method = "Cholesky"): Cannot compute vcov:
## Hessian is not positive definite>
## <simpleError in vcov.clm(object, method = "Cholesky"): Cannot compute vcov:
## Hessian is not positive definite>
## <simpleError in vcov.clm(object, method = "Cholesky"): Cannot compute vcov:
## Hessian is not positive definite>
## <simpleError in vcov.clm(object, method = "Cholesky"): Cannot compute vcov:
## Hessian is not positive definite>
## <simpleError in vcov.clm(object, method = "Cholesky"): Cannot compute vcov:
## Hessian is not positive definite>
## <simpleError in vcov.clm(object, method = "Cholesky"): Cannot compute vcov:
## Hessian is not positive definite>
## <simpleError in vcov.clm(object, method = "Cholesky"): Cannot compute vcov:
## Hessian is not positive definite>
power.clmm.den <- sum(clmm.comps.den$`Statistical signicance` == "Significant")/num_sims.clmm</pre>
```

3.3 Arousal power analysis

```
## <simpleError in vcov.clm(object, method = "Cholesky"): Cannot compute vcov:
## Hessian is not positive definite>
```

```
## <simpleError in vcov.clm(object, method = "Cholesky"): Cannot compute vcov:
## Hessian is not positive definite>
## <simpleError in vcov.clm(object, method = "Cholesky"): Cannot compute vcov:
## Hessian is not positive definite>
## <simpleError in vcov.clm(object, method = "Cholesky"): Cannot compute vcov:
## Hessian is not positive definite>
## <simpleError in vcov.clm(object, method = "Cholesky"): Cannot compute vcov:
## Hessian is not positive definite>
## <simpleError in vcov.clm(object, method = "Cholesky"): Cannot compute vcov:
## Hessian is not positive definite>
## <simpleError in vcov.clm(object, method = "Cholesky"): Cannot compute vcov:
## Hessian is not positive definite>
## <clculate power (proportion of simulations in which the Tempo contrast was significant)
power.clmm.aro <- sum(clmm.comps.aro$`Statistical signicance` == "Significant")/num_sims.clmm</pre>
```

3.4 Achieved power

```
p5a <- ggplot(clmm.comps.den, aes(x = p, fill = `Statistical signicance`)) +
  scale fill hue(direction = -1) +
  geom_histogram(bins = 1/alpha.clmm, breaks = seq(0, 1, alpha.clmm)) +
  labs(y = "Count", x = "p-value", title = "Density") +
  annotate("text", x = 0.5, y = sum(clmm.comps.den$`Statistical signicance` == "Significant") * 0.9,
           label = paste0("Power = ", round(power.clmm.den, 3),
                          "\nSample size = ", sample_size.clmm,
                          " participants per country\n(6 paired responses per participant)"))
p5b <- ggplot(clmm.comps.aro, aes(x = p, fill = `Statistical signicance`)) +
  scale_fill_hue(direction = -1) +
  geom_histogram(bins = 1/alpha.clmm, breaks = seq(0, 1, alpha.clmm)) +
  labs(y = "Count", x = "p-value", title = "Arousal") +
  annotate("text", x = 0.5, y = sum(clmm.comps.aro$`Statistical signicance` == "Significant") * 0.9,
           label = paste0("Power = ", round(power.clmm.aro, 3),
                          "\nSample size = ", sample_size.clmm,
                          " participants per country\n(6 paired responses per participant)"))
p5 <- ggarrange(p5a, p5b,
                common.legend = TRUE,
                legend = "bottom",
                labels = "AUTO")
p5
```

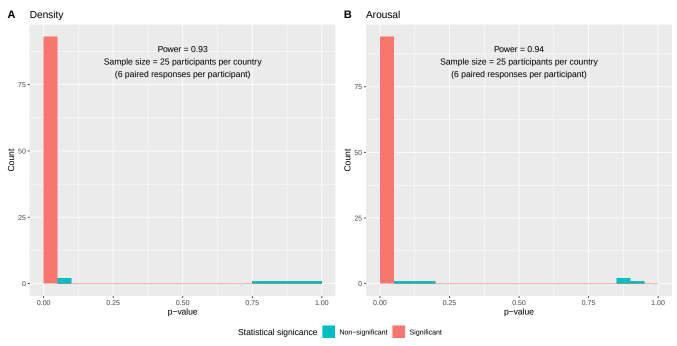


Figure S5. Histogram of p-values for the contrasts between Tempo conditions (Low, High) for all simulations. The estatistical power is the proportion of significant results. Significant results are in red.

4 Analysis script

Here, using a randomly selected sample, we fit one model as if it was the data collected from the experiment. This is not only to make sure that the data are sufficient and the structure is adequate to fit the model, but to plan in advance all the code.

```
# Select a random sample
ex.data <- samples.clmm.long %>%
  filter(sample.clmm == sample(1:num_sims.clmm, 1, replace = TRUE))
```

Because only the effect of Tempo was predicted and simulated a-priori, all other effects and interactions are completely random.

```
p6a <- ggplot(ex.data, aes(x = Tempo, y = Density.tempo, color = Participant.country, shape = Music.countr
  geom_violin(aes(group = Tempo), draw_quantiles = 0.5) +
  geom_point(position = position_dodge(0.3), size = 3) +
  geom_line(aes(group = factor(groupid)), size = 0.5, linetype = "dashed", position = position_dodge(0.3)
  labs(y= "Density", x = "Tempo", title = "Density") +
  scale_x_discrete(limits = c("Low", "High")) +
  scale_size_discrete(breaks = c("Solo", "Group")) +
  theme(plot.title = element_text(face = "bold", hjust = 0.5)) +
  facet_wrap(~Solo.group)
## Warning: Using 'size' aesthetic for lines was deprecated in ggplot2 3.4.0.
## i Please use 'linewidth' instead.
## This warning is displayed once every 8 hours.
```

- ## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
- ## generated.
- ## Warning: Using size for a discrete variable is not advised.

```
p6b <- ggplot(ex.data, aes(x = Tempo, y = Arousal.tempo, color = Participant.country, shape = Music.countr
  geom_violin(aes(group = Tempo), draw_quantiles = 0.5) +
  geom_point(position = position_dodge(0.3), size = 3) +
```

```
geom_line(aes(group = factor(groupid)), size = 0.5, linetype = "dashed", position = position_dodge(0.3),
labs(y= "Arousal", x = "Tempo", title = "Arousal") +
scale_x_discrete(limits = c("Low", "High")) +
scale_size_discrete(breaks = c("Solo", "Group")) +
theme(plot.title = element_text(face = "bold", hjust = 0.5)) +
facet_wrap(~Solo.group)
```

```
## Warning: Using size for a discrete variable is not advised.
p6 <- ggarrange(p6a, p6b,
                common.legend = TRUE,
                legend = "bottom")
## Warning: The following aesthetics were dropped during statistical transformation:
## colour, shape
## i This can happen when ggplot fails to infer the correct grouping structure in
     the data.
## i Did you forget to specify a 'group' aesthetic or to convert a numerical
    variable into a factor?
## Warning: The following aesthetics were dropped during statistical transformation:
## colour, shape
## i This can happen when ggplot fails to infer the correct grouping structure in
    the data.
## i Did you forget to specify a 'group' aesthetic or to convert a numerical
   variable into a factor?
## The following aesthetics were dropped during statistical transformation:
## colour, shape
## i This can happen when ggplot fails to infer the correct grouping structure in
    the data.
## i Did you forget to specify a 'group' aesthetic or to convert a numerical
    variable into a factor?
## The following aesthetics were dropped during statistical transformation:
## colour, shape
## i This can happen when ggplot fails to infer the correct grouping structure in
##
    the data.
## i Did you forget to specify a 'group' aesthetic or to convert a numerical
    variable into a factor?
## The following aesthetics were dropped during statistical transformation:
## colour, shape
## i This can happen when ggplot fails to infer the correct grouping structure in
##
    the data.
## i Did you forget to specify a 'group' aesthetic or to convert a numerical
     variable into a factor?
## The following aesthetics were dropped during statistical transformation:
## colour, shape
## i This can happen when ggplot fails to infer the correct grouping structure in
##
    the data.
## i Did you forget to specify a 'group' aesthetic or to convert a numerical
     variable into a factor?
p6
```

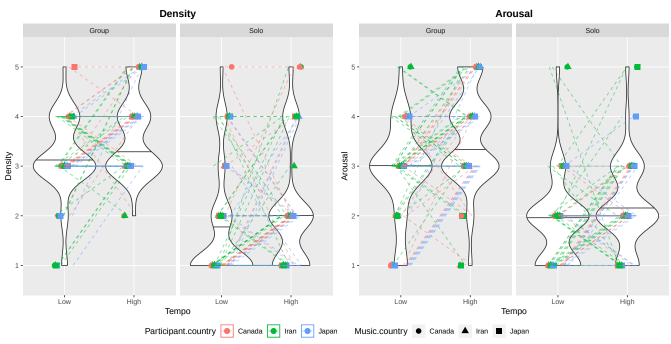


Figure S6. Data for Density and Arousal ratings from a ramdomly selected, simulated sample. Dashed lines represent the within-subject effect of the Tempo manipulation.

We fitted the model with the same structure: clmm(DV ~ Tempo * Solo.group * Participant.country * Music.country + (1 + Tempo | Participant). Estimated marginal means and confidence intervals for this simulation are represented in S7.

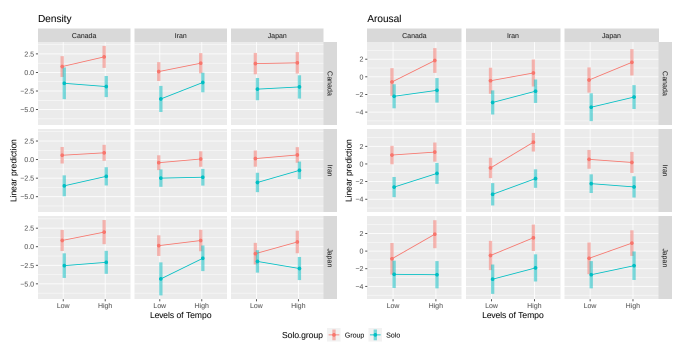


Figure S7. Estimated marginal means and confidence intervals for the within-subject effects of Tempo from a ramdomly selected, simulated sample. Columns represent data from simulated participants from each country, while rows represent the simulated response to music from each country.

5 Session info (for reproducibility)

```
library(pander)
pander(sessionInfo(), locale = FALSE)
```

R version 4.3.1 (2023-06-16 ucrt)

Platform: x86 64-w64-mingw32/x64 (64-bit)

attached base packages: stats, graphics, grDevices, utils, datasets, methods and base

other attached packages: pander(v.0.6.5), ordinal(v.2022.11-16), emmeans(v.1.8.6), ggpubr(v.0.6.0), lubridate(v.1.9.2), forcats(v.1.0.0), stringr(v.1.5.0), dplyr(v.1.1.2), purrr(v.1.0.1), readr(v.2.1.4), tidyr(v.1.3.0), tibble(v.3.2.1), ggplot2(v.3.4.2), tidyverse(v.2.0.0) and knitr(v.1.43)

loaded via a namespace (and not attached): gtable(v.0.3.3), xfun(v.0.39), rstatix(v.0.7.2), lattice(v.0.21-8), tzdb(v.0.4.0), numDeriv(v.2016.8-1.1), vctrs(v.0.6.3), tools(v.4.3.1), generics(v.0.1.3), parallel(v.4.3.1), fansi(v.1.0.4), highr(v.0.10), ucminf(v.1.2.0), pkgconfig(v.2.0.3), Matrix(v.1.5-4.1), lifecycle(v.1.0.3), compiler(v.4.3.1), farver(v.2.1.1), tinytex(v.0.45), munsell(v.0.5.0), codetools(v.0.2-19), carData(v.3.0-5), htmltools(v.0.5.5), yaml(v.2.3.7), crayon(v.1.5.2), pillar(v.1.9.0), car(v.3.1-2), MASS(v.7.3-60), abind(v.1.4-5), nlme(v.3.1-162), tidyselect(v.1.2.0), digest(v.0.6.31), mvtnorm(v.1.2-2), stringi(v.1.7.12), bookdown(v.0.34), labeling(v.0.4.2), cowplot(v.1.1.1), fastmap(v.1.1.1), grid(v.4.3.1), colorspace(v.2.1-0), cli(v.3.6.1), beepr(v.1.3), magrittr(v.2.0.3), utf8(v.1.2.3), broom(v.1.0.5), withr(v.2.5.0), scales(v.1.2.1), backports(v.1.4.1), timechange(v.0.2.0), estimability(v.1.4.1), rmarkdown(v.2.22), audio(v.0.1-10), gridExtra(v.2.3), ggsignif(v.0.6.4), hms(v.1.1.3), evaluate(v.0.21), rlang(v.1.1.1), Rcpp(v.1.0.10), xtable(v.1.8-4), glue(v.1.6.2), rstudioapi(v.0.14) and R6(v.2.5.1)

Supplementary references

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