- Tutorial for Customized Power Simulations and Data Analyses for Human-AI Interaction
- Experiments using Generalized Linear Mixed Models
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Abstract

19 Understanding how humans interact with articifical intelligence through experiments

becomes increasingly relevant. However, human-AI interaction researchers lack the

21 appropriate tools to conduct power and data analyses for the required complex study

designs. In this work, we provide a tutorial on how to run customized power analyses using

<sup>23</sup> data simulation based on Generalized Linear Mixed Models (GLMMs). By providing code

4 in a case study, we equip human-AI interaction researchers with the tools to simulate their

own data and run analyses based on GLMMs. We discuss implications and give an outlook.

26 Keywords: generalized linear mixed model, data simulation, sample size, power

27 analysis, human-AI interaction

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Introduction

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As the integration of artificial intelligence (AI) into our daily lives continues to grow, it has become increasingly important to study how humans interact with these intelligent 33 systems. This is particularly important as AI becomes more sophisticated and is used in decision-making processes, for instance in medicine, that have significant impacts on 35 individuals and society as a whole. Therefore, thorough experiments are needed to study human-AI interaction to understand how people perceive and respond to these novel technologies. When conducting experimental research in this emerging field, it is essential to determine an appropriate sample size to ensure that the results obtained are both statistically significant and reliable. Therefore, running sample size estimations is crucial for designing experiments that can provide meaningful insights into human-AI interaction. In empirical research relying on hypothesis testing, the most common strategy to 42 determine appropriate samples size is based on statistical power. Statistical power is defined as the probability that a hypothesis test of interest turns out significant, when analyzing repeated samples from a population with a true effect of some pre-specified size. Less formally, power is also described as the probability that a hypothesis test accepts the alternative hypothesis  $H_1$  if  $H_1$  is indeed true. Power analysis (i.e. calculating the statistical power for a given true effect and sample size) plays a crucial role in addressing the replication crisis and meeting the requirements set forth by journals and funding agencies in human-AI interaction research. The replication crisis has underscored the problem of high false discovery rates (FDR) coupled with low-powered studies. Power 51 analysis offers a valuable solution by allowing researchers to estimate the appropriate sample sizes needed to achieve sufficient statistical power for reliable results. Moreover, many journals and funding agencies now mandate the inclusion of power analysis as part of study protocols and grant proposals, recognizing their significance in ensuring robust and meaningful findings. By incorporating power analysis into research planning, researchers can enhance the replicability and credibility of their work, ultimately contributing to the advancement of psychological science.

## <sup>59</sup> Customized power simulations are hard

Power analysis can be challenging, particularly when analyzing complex study designs 60 using (generalized) linear mixed models (GLMMs; also called multilevel models). While user-friendly software is available for simple statistical models like t-tests, ANOVA, and linear regression, the increasing use of GLMMs in human-AI interaction research introduces greater complexity to power analysis. For GLMMs, analytic options to derive sample sizes are not available for general research designs. In contrast, simulations are often required, necessitating assumptions for all parameters in the model. An increasing number of tutorials describe how to perform power simulation for multilevel models. However, most 67 tutorials discuss only linear mixed models (LMMs) and focus on the most common designs, limiting the guidance available for researchers facing more complex study designs [REF]. Moreover, existing tutorials often rely on heuristics for specifying variance components (e.g. standard deviation of random intercepts) or assume that meta-analytic results or data 71 from pilot studies is available to determine plausible values for all model parameters. 72 When planning a human-AI interaction experiment with complexities such as the use 73 of GLMMs, non-standard designs, and missing by design conditions, the process of power simulations becomes more challenging. Currently available software packages for power simulations do not include common designs in human-AI interaction research, making it necessary to build data simulations tailored specifically to the study design. Unfortunately, there are no general recommendations that can be applied universally in such cases. Instead, researchers must acquire the necessary skills based on the specific application at 79 hand. In this paper, we present a case study that serves as a practical demonstration of

how to perform a data simulation for a concrete study for which no meta-analytic results or conclusive data from pilot studies is available. Through this case study, we aim to provide guidance and insights into the complexities involved in addressing complex design considerations and obtaining accurate power estimates.

# 85 LMMs, GLMMs and the lme4 package in R

LMMs and GLMMs are powerful statistical frameworks that handle complex data structures by incorporating both fixed and random effects.

LMMs are extensions of linear regression models that account for correlated data and hierarchical structures. They are used when the outcome variable is continuous and follow a normal distribution. LMMs allow for the modeling of fixed effects, which capture the relationships between predictors and the outcome, as well as random effects, which account for the correlation and variability within groups or subjects. Random effects are typically assumed to follow a normal distribution with a mean of zero and a variance that quantifies the heterogeneity across the groups or subjects.

GLMMs extend the LMM framework to accommodate non-normal and categorical outcome variables. They are used when the outcome variable does not follow a normal distribution, but instead belongs to a different distribution family, such as binomial, Poisson, or gamma. GLMMs incorporate both fixed and random effects, similar to LMMs, but also involve a link function that connects the linear predictor to the expected value of the outcome variable. The link function allows for modeling the relationship between predictors and the outcome in a way that is appropriate for the specific distribution family of the response variable.

GLMMs are gaining increasing popularity in the field of Human-AI interaction research. As the complexity of studying human interactions with artificial intelligence systems grows, researchers require more sophisticated statistical models to capture the

nuanced relationships and hierarchical structures within the data. GLMMs offer a flexible 106 framework for analyzing data with non-normal and categorical outcomes, accounting for 107 both fixed and random effects. This versatility makes GLMMs particularly suitable for 108 investigating various aspects of Human-AI interaction, such as user preferences, trust, 109 engagement, and performance. By incorporating GLMMs into their analyses, researchers in 110 the field of Human-AI interaction can obtain more robust and comprehensive insights into 111 the intricate dynamics between humans and AI systems, leading to a deeper understanding 112 of the psychological and behavioral aspects involved. 113

The lme4 R package is a state-of-the-art tool for fitting frequentist GLMMs (for 114 Bayesian GLMMs, the brms R package is currently the most prominent option). The lme4 package includes a useful function called "simulate" that allows researchers to simulate the 116 dependent variable based on the same model formula used for model fitting, enabling power simulations and other related analyses. However, the model parameterization used 118 by the lme4 package is quite technical, making it more difficult for applied researchers to 119 determine whether their specified population model implies plausible associations in their 120 simulated data. Therefore in this tutorial, we simulate data for GLMMs from first 121 principles to assist applied researchers in better understanding all model assumptions, and 122 then use lme4 to analyze the simulated datasets. 123

124 Methods

In this section, we describe the steps to conduct a data simulation and a power analysis for GLMMs. In doing so, we provide a concrete example in the form of a case study.

We used R (Version 4.2.3; R Core Team, 2023) and the R-packages *papaja* (Version 0.1.1; Aust & Barth, 2022), and *tinylabels* (Version 0.2.3; Barth, 2022) for all our analyses.

## The present case study

In the case study, we simulate the data for a human-AI Interaction experiment where
a health care AI is to be evaluated. Participants view head CT cases and evaluate if a
bleeding is present. Also, there can be AI advice that is either correct or incorrect. This
medical diagnosis can be correct or incorrect.

In this case, it would be more challenging to recruit task expert as there is only a limited amount of such people. Non-experts would be easier to recruit. The open question is how many task experts and non-experts to recruit to achieve sufficient power.

describe study design and sample size restrictions - describe research question
(i.e. the coefficients targeted by power simulation)

Power simulations in GLMMs are essential for estimating the statistical power of 140 complex psychological study designs. The model equation for a GLMM combines fixed effects, random effects, and an appropriate link function to model the relationship between predictors and the outcome variable. The necessary assumptions for power simulations in GLMMs include assumptions about the distributional form of the outcome variable, the random effects, and the error structure. The distributional assumption specifies the family of distributions for the outcome variable, such as Gaussian, Poisson, or binomial. Assumptions about the random effects include the assumption of normality and the covariance structure among the random effects. Additionally, assumptions about the error 148 structure, such as independence or correlation, must be specified. Interpreting these 149 assumptions entails understanding the underlying assumptions of the model and ensuring 150 they align with the characteristics of the data being analyzed. 151

# 2 TODO

153

points not discussed yet:

- discuss the specific missing by design complexity
- discuss different possible parametrizations of the fixed effects
- find the appropriate lme4 formula (simple models first, more complex models later)
- determine the correct labels for model coefficients in the lme4 output
- excursus: estimate expected with of confidence intervals for different sample sizes and number of stimuli

# 160 Generalized linear mixed models

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In a GLMM, the expected value of the dependent variable Y conditioned on the vector of predictor variables  $\mathbf{X}$  and random effects  $\mathbf{U}$ , transformed by a link function g() is modeled as a linear combination  $\eta$  of the predictor variables  $\mathbf{X}$ , the random effects  $\mathbf{U}$  and the model parameters  $\beta$ .

$$q(E(Y|\mathbf{X} = \mathbf{x}, \mathbf{U} = \mathbf{u})) = \eta$$

Equivalently, the conditional expected value is modeled as the linear combination  $\eta$ , transformed by the inverse link function  $g^{-1}()$ .

$$E(Y|\mathbf{X} = \mathbf{x}, \mathbf{U} = \mathbf{u})) = g^{-1}(\eta)$$

If the dependent variable Y is a binary variable with values 0 or 1, the conditional expected value is equivalent to the probability:

$$P_{si} = P(Y = 1 | \mathbf{X} = \mathbf{x}, \mathbf{U} = \mathbf{u})$$

In our case study,  $P_{si}$  is the conditional probability that subject s gives the correct response to item i.

In such a setting, we model the probability as

$$P_{si} = inverse\_logit(\eta_{si})$$

with the inverse-logit link  $g^{-1}(\eta_{si}) = inverse\_logit(\eta_{si}) = \frac{exp(\eta_{si})}{1 + exp(\eta_{si})}$  or equivalently

$$logit(P_{si}) = \eta_{si}$$

- with the logit link  $g(P_{si}) = logit(P_{si}) = ln(\frac{P_{si}}{1 P_{si}})$ .
- In our case study, the probability to give a correct response is assumed to depend on the predictors:
- $advice\_present_{si}$ : whether subject s was presented with algorithmic advice (1) or not

  (0) when asked to asses item i
- $advice\_correct_{si}$ : whether this advice was correct (1) or not (0)
- $expert_s$ : whether subject s was a professional neurologist (1) or not (0)
- and the random effects:
- $u_{0s}$ : the deviation of suject s from the average ability to solve an item with average difficulty; assumed to be distributed as  $u_{0s} \sim N(0, \sigma_S^2)$
- $u_{0i}$ : the deviation of item i from the average difficulty to be solved by a person with average ability; assumed to be distributed as  $u_{0i} \sim N(0, \sigma_I^2)$ 
  - In total, we assume the model

$$logit[P_{si}] = (\beta_0 + u_{0s} + u_{0i}) +$$

$$\beta_a \cdot advice\_present_{si} + \beta_c \cdot advice\_correct_{si} + \beta_e \cdot expert_s +$$

$$\beta_{ea} \cdot expert_s \cdot advice\_present_{si} + \beta_{ec} \cdot expert_s \cdot advice\_correct_{si}$$

or equivalently

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$$P_{si} = inverse\_logit[(\beta_0 + u_{0s} + u_{0i}) + \\ \beta_a \cdot advice\_present_{si} + \beta_c \cdot advice\_correct_{si} + \beta_e \cdot expert_s + \\ \beta_{ea} \cdot expert_s \cdot advice\_present_{si} + \beta_{ec} \cdot expert_s \cdot advice\_correct_{si}]$$

with model parameters  $\beta_0$ ,  $\beta_e$ ,  $\beta_a$ ,  $\beta_c$ ,  $\beta_{ea}$ ,  $\beta_{ec}$ ,  $\sigma_S$ , and  $\sigma_I$ .

In the GLMM literature, this would be called a binomial GLMM with two random intercepts (for subjects and items), two level-1 predictors (advice\_present,

```
advice\_correct), one level-2 predictor (expert) and two cross-level interactions (expert \cdot advice\_present, expert \cdot advice\_correct).
```

To limit complexity, we do not consider random slopes, additional predictors or higher-level interactions here.

## Data Simulation

The following R function simulates a full dataset structured according to our case study. The faux package contains useful functions when simulating factorial designs including random effects.

```
simulate <- function(n subjects = 100, n items = 50,
 b = 0.847, b_e = 1.350, b_a = -1.253, b_c = 2.603,
 b_{ea} = 0.790, b_{ec} = -1.393,
 sd\ u0s = 0.5, sd\ u0i = 0.5, ...){
 require(dplyr)
 require(faux)
  # simulate design
 dat <- add random(subject = n subjects, item = n items) %>%
    add_between("subject", expert = c(1, 0), .prob = c(0.25, 0.75)) %>%
    mutate(advice present = rbinom(n(), 1, prob = 2/3)) %>%
    mutate(advice_correct = if_else(advice_present == 1,
                                    rbinom(n(), 1, prob = 0.8), 0)) %>%
    # add random effects
    add ranef("subject", u0s = sd u0s) %>%
    add ranef("item", u0i = sd u0i) %>%
    # compute dependent variable
    mutate(linpred = b_0 + u0i + u0s +
```

```
b_e * expert + b_a * advice_present + b_c * advice_correct +
    b_ea * expert * advice_present + b_ec * expert * advice_correct) %>%
    mutate(y_prob = plogis(linpred)) %>%
    mutate(y_bin = rbinom(n = n(), size = 1, prob = y_prob))
    dat
}
```

In our case study, each subject (n subjects in total) is assumed to respond to each 198 item (n items in total). Thus the add random command creates a fully-crossed 199 data.frame with n subjects × n items rows. We add a between subject effect with the 200 add between command, simulating that about 25 of subjects are experts. The next two 201 lines simulate that in about  $\frac{2}{3}$  of trials, subjects will be presented with AI advice and if 202 advice is presented, the advice will be correct in about 80 of cases (the variable 203 advice correct is always 0 when no advice is presented). Next we simulate one random 204 effect each subject (u0s) and for each item (u0i). As assumed by standard GLMMs, the 205 add ranef function draws the random effects from a normal distribution with mean 0 and 206 a standard deviation specified by the user. With all design variables done, we are ready to 207 simulate our model equation as outlined in equation X. The linear predictor variable 208 lingred ( $\eta$  in the GLMM model equations), combines the predictor variables, random effects and model parameters as assumed by our model. We then transform the linear 210 predictor with the inverse-link function to compute y\_prob, the probability that the subject 211 correctly solved the item (in R the inverse-logit link is computed with plogis and the logit 212 link with qlogis). In the final step, we simulate the binary dependent variable y\_bin by -213 for each trial – drawing from a bernoulli distribution with success probability y\_prob. 214

## Model fitting

In this section, we show how to fit a GLMM with lme4, interpret the model and test hypotheses derived from a research question.

We simulate data according to our model, in which 100 subjects respond to 50 items
(we use set.seed to make the simulation reproducible). However, for the sake of the
exercise, we can imagine that this would be real data resulting from our future experiment
and think about how we would analyse this data.

```
library(tidyverse)
set.seed(1)
dat <- simulate(n_subjects = 100, n_items = 50)</pre>
```

The lme4 package uses a special syntax for model specification. Our assumed GLMM is represented by the formula:

```
library(lme4)

f <- y_bin ~ 1 + expert + advice_present + advice_correct +
    expert:advice_present + expert:advice_correct +
    (1|subject) + (1|item)</pre>
```

The first two lines looks similar to any linear model in R (general intercept indicated by 1; main effects indicated by variable names in the dataset; interactions indicated by variable1:variable2). The third line specifies a random intercept for each subject (1|subject) and for each item (1|item). The complete set of rules for the syntax are outlined in (CITE LME4 PAPER) and in the documentation of the lme4 package.

In lme4, a GLMM is fitted with the glmer function. By setting family =

"binomial" we request a binomial GLMM, appropriate for our binary dependent variable y\_bin (the binomial GLMM used the canonical logit link by default).

```
fit <- glmer(f, data = dat, family = "binomial")</pre>
```

# 232 Model interpretation

We can inspect the estimated model parameters with the summary command:

```
summary(fit)
```

```
## Generalized linear mixed model fit by maximum likelihood (Laplace
         Approximation) [glmerMod]
   ##
235
       Family: binomial (logit)
236
   ## Formula:
237
   ## y_bin ~ 1 + expert + advice_present + advice_correct + expert:advice_present +
238
           expert:advice correct + (1 | subject) + (1 | item)
   ##
239
          Data: dat
   ##
240
   ##
241
                            logLik deviance df.resid
   ##
            AIC
                      BIC
242
         4149.4
   ##
                  4201.6
                           -2066.7
                                      4133.4
                                                  4992
243
   ##
244
   ## Scaled residuals:
245
   ##
           Min
                     1Q
                         Median
                                      3Q
                                              Max
246
   ## -5.7669 0.2125
                        0.3046 0.4317
                                          2.1057
247
   ##
248
   ## Random effects:
249
   ##
       Groups Name
                             Variance Std.Dev.
        subject (Intercept) 0.3148
                                       0.5611
   ##
251
                (Intercept) 0.1624
                                       0.4029
252
   ##
        item
   ## Number of obs: 5000, groups: subject, 100; item, 50
253
```

```
##
254
   ## Fixed effects:
255
   ##
                              Estimate Std. Error z value Pr(>|z|)
256
   ## (Intercept)
                                                             < 2e-16 ***
                                1.0339
                                            0.1103
                                                      9.374
257
   ## expert
                                1.1849
                                            0.2096
                                                      5.654 1.56e-08 ***
258
   ## advice_present
                                            0.1206 -11.143
                               -1.3436
                                                             < 2e-16 ***
259
   ## advice_correct
                                2.6154
                                            0.1273
                                                     20.541
                                                             < 2e-16 ***
260
   ## expert:advice_present
                                            0.2940
                                                      3.601 0.000317 ***
                                1.0589
261
   ## expert:advice_correct
                               -1.8104
                                            0.2915
                                                    -6.211 5.27e-10 ***
262
   ## ---
263
                       0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
   ## Signif. codes:
   ##
265
   ## Correlation of Fixed Effects:
   ##
                   (Intr) expert advc_p advc_c exprt:dvc_p
   ## expert
                   -0.377
268
   ## advic prsnt -0.349
                            0.176
269
   ## advic crrct 0.023
                           0.001 -0.668
   ## exprt:dvc p 0.143 -0.448 -0.412
271
   ## exprt:dvc c -0.008 0.004 0.292 -0.435 -0.686
272
```

The output shows the estimates for all model parameters: the Estimate column in the Fixed effects table contains the estimates for the  $\beta$  parameters, while the Std.Dev. column in the Random effects table contains the estimates for  $\sigma_S$  and  $\sigma_I$ .

Unfortunately, the model parameters in a binomial GLMM are hard to interpret, because 1) the  $\beta$  parameters are connected to the modeled probability via the non-linear inverse-logit link, and 2) we also have to consider the random effects. The most simple interpretation works by imagining a subject with average ability (u0s = 0) responding to an item with average difficulty (u0i = 0). Then the model implied probability that such a person solves such an item is given by:

$$P(Y = 1 | \mathbf{X} = \mathbf{x}, \mathbf{U} = \mathbf{0}) =$$

$$= inverse\_logit[\beta_0 + \beta_a \cdot advice\_present_{si} + \beta_c \cdot advice\_correct_{si} + \beta_e \cdot expert_s +$$

$$\beta_{ea} \cdot expert_s \cdot advice\_present_{si} + \beta_{ec} \cdot expert_s \cdot advice\_correct_{si}]$$

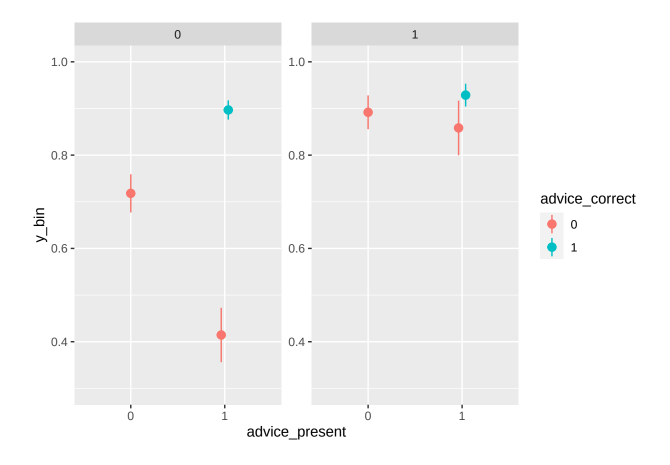
In fact, we would only need the full equation if the subject is an expert and correct advice is presented. In all other experimental conditions, some terms drop from the equation because they are multiplied by 0. The other extreme case would be the probability that a non-expert with average ability solves and item with average difficulty when presented without any advice:

$$P(Y = 1 | expert = 0, advice\_present = 0, advice\_correct = 0, u_{0s} = 0, u_{0i} = 0) = inverse \ logit[\beta_0]$$

Due to this complicated relationship, many experts argue not to focus too much on interpreting single model parameters when working with GLMMs. Instead, it can be more intuitive to consider the implied predicted distribution of the dependent variable for each experimental conditions across all subjects and items.

With the marginal effects package, we can easily compute predictions for all observations in the dataset based on the fitted GLMM (including all fixed **and** random effects), and plot the average probability with confidence intervals for each experimental condition:

```
library(marginaleffects)
plot_predictions(fit, by = c("advice_present", "advice_correct", "expert"),
    type = "response") + ylim(c(0.3, 1))
```



### Hypothesis testing

However, we need to think about the model parameters again when wanting to test hypotheses which we have theoretically derived from some research question. Because the inverse-logit link is still a continuously increasing function, positive parameter values always correspond to increases in probability and vice versa.

The Fixed effects table in Figure X also includes p-values for hypothesis tests with null hypotheses of the style  $H_0: \beta = 0$ . However, for many research questions of interest, we are not interested in these two-sided tests referring to only one parameter.

For our case study, imagine the following combined hypothesis: We expect that for both experts and non-experts, correct advice leads to a higher probability to solve an item compared to no advice presented, AND, we expect that for both experts and non-experts,

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incorrect advice leads to a lower probability to solve an item compared to no advice presented.

This combined hypothesis leads to the following four separate null hypotheses to be tested:

$$H_{01}: \beta_a + \beta_c + \beta_{ea} + \beta_{ec} \le 0$$

$$H_{02}: \beta_a + \beta_c \le 0$$

$$H_{03}: \beta_a + \beta_{ea} \ge 0$$

$$H_{04}: \beta_a > 0$$

We arrive at these inequalities based on the following logic, exemplified here only for  $H_{01}$ : The first null hypothesis states that an expert responding to an item while presented with correct advice has lower or equal probability to solve the item compared to the same expert facing the same item without any advice. This implies the following inequality for each subject s and item i

```
inverse\_logit[(\beta_0 + u_{0s} + u_{0i}) + \beta_e + \beta_a + \beta_c + \beta_{ea} + \beta_{ec}] \leq inverse\_logit[(\beta_0 + u_{0s} + u_{0i}) + \beta_e]
which simplifies to \beta_a + \beta_c + \beta_{ea} + \beta_{ec} \leq 0.
```

We can specify and test hypotheses like these with the multcomp package as follows:

```
library(multcomp)
null_hypotheses <- c(
    "advice_present + advice_correct + expert:advice_present +
    expert:advice_correct <= 0",
    "advice_present + advice_correct <= 0",
    "-1 * (advice_present + expert:advice_present) <= 0",
    "-1 * (advice_present) <= 0")
glht <- glht(fit, linfct = null_hypotheses)
summary(glht, test = univariate())$test$pvalues</pre>
```

```
## advice present + advice correct + expert:advice present + expert:advice correct
318
   ##
                                                                                  0.006407541
319
   ##
                                                            advice present + advice correct
320
                                                                                  0.00000000
   ##
321
                                            -1 * (advice_present + expert:advice_present)
   ##
322
                                                                                  0.143933522
   ##
323
   ##
                                                                       -1 * (advice_present)
324
   ##
                                                                                  0.00000000
325
```

Because all hypotheses tested simultaneously with the glht function must have the 326 same direction, we flip the sign of inequalities three and four by multiplying them with -1. 327 The multcomp package automatically adjusts p-values when multiple hypotheses are tested 328 simultaneously. However, the combined null hypothesis in our exemplary research question 329 should only be rejected if all individual null hypotheses are rejected. In such cases, the 330 error probabilities do not accumulate and we would waste power when correcting for 331 multiple testing. Thus, we request unadjusted p-values by setting test = univariate() in the summary command. With a standard significance level of  $\alpha = 0.05$ , we would reject all four null hypotheses and therefore also reject the combined null hypothesis for this simulated dataset.

### Specification of plausible parameter values

When introducing our simulation function and simulating data for the above example, we have used theoretically plausible values as defaults for all model parameters ( $\beta_0$ ,  $\beta_e$ ,  $\beta_a$ ,  $\beta_c$ ,  $\beta_{ea}$ ,  $\beta_{ec}$ ,  $\sigma_S$ , and  $\sigma_I$ ), but have not talked about where the numbers came from. All parameter values have been determined in repeated exchanges with our affiliated domain experts and we here outline a few strategies on how to determine plausible parameter values.

We have already seen in our discussion of model interpretation, how we can derive
the model implied probability for each experimental condition, that a subject with average
ability solves an item with average difficulty. We can revert this perspective by choosing
plausible probability values and deriving the parameter values implied by these
probabilities (for an average subject and an average item).

Table X shows our set of assumptions concerning the probability that an average subject solves an average item for each experimental condition, as well as the corresponding equations implied by the model:

Experimental condition	$P(Y=1 \mathbf{X}=\mathbf{x},\mathbf{U}=0)$	$Implied\ equation$
no advice, no expert	0.70	$logit(0.70) = \beta_0$
no advice, expert	0.90	$logit(0.90) = \beta_0 + \beta_e$
false advice, no expert	0.40	$logit(0.40) = \beta_0 + \beta_a$
false advice, expert	0.85	logit(0.85) =
		$\beta_0 + \beta_e + \beta_a + \beta_{ea}$
correct advice, no	0.90	logit(0.90) =
expert		$\beta_0 + \beta_a + \beta_c$
correct advice, expert	0.95	logit(0.95) =
		$\beta_0 + \beta_e + \beta_a + \beta_c + \beta_{ea} + \beta_{ec}$

This table can be used to compute the implied values for the  $\beta$  parameters, starting with the first equation and reinserting the computed  $\beta$  values in all following equations:

```
b_0 <- qlogis(0.7)

b_e <- qlogis(0.9) - b_0

b_a <- qlogis(0.4) - b_0

b_ea <- qlogis(0.85) - b_0 - b_e - b_a

b_c <- qlogis(0.9) - b_0 - b_a</pre>
```

$$b_{ec} \leftarrow qlogis(0.95) - b_{e} - b_{e} - b_{a} - b_{c} - b_{ea}$$

$$c(b_{e} = b_{e}, b_{e} = b_{e}, b_{a} = b_{a}, b_{c} = b_{c}, b_{ea} = b_{ea}, b_{ec} = b_{ec})$$

It is always possible to double-check these computations by transforming the parameter values back to probabilities, e.g.

$$P(Y = 1 | expert = 1, advice\_present = 1, advice\_correct = 1, u_{0s} = 0, u_{0i} = 0) = inverse\_logit[\beta_0 + \beta_e + \beta_a + \beta_c + \beta_{ea} + \beta_{ec}]$$

357 ## [1] 0.95

Although the derivations above are straightforward, it is important not to 358 misinterpret their implications: In binomial GLMMs, the average probability to solve an 350 item (averaged across persons of varying ability and items of varying difficulty) is **not** 360 equal to the probability that a person with average ability solves an item with average 361 difficulty. For example, we determined the  $\beta$  parameters in a way that correspond to a 362 desired probability, that an expert with average ability solves an item with average 363 difficulty when presented with a correct advice. However even if the model were true, we would not observe this probability if we estimated the probability in a group of expert 365 responding to items presented with correct advice from a big sample of subjects drawn from their natural distribution of ability and items drawn from their natural distribution of difficulty. This implies that one must be careful when specifying parameter values based on 368 previous studies or pilot data. 369

The well known inequality of conditional and marginal effects in GLMMs makes their interpretation more difficult, however, this does not mean that we cannot use the marginal

interpretation (average probability across persons and items) to inform plausible parameter values: When parameter values have selected, we can compute the implied marginal distributions and compare this information to our domain knowledge. Then we can iteratively adjust the parameter values until we are satisfied with the implied distributions.

Earlier, we have already encountered one way to visualize the implied marginal
distributions: We can fit our model to a simulated dataset and use the convenience
functions from the marginaleffects package to compute averaged predictions that
correspond to our quantities of interest. However, the model predictions will only be close
to the true distribution if the simulated dataset is very large, but then the model fitting
consumes a lot of time and memory. A more sophisticated strategy is to simulate a large
dataset and directly compute the averages, contrasts and distributions we are interested in.

```
library(tidyverse)
library(ggdist)
dat <- simulate(n_subjects = 2000, n_items = 2000, sd_u0s = 0.5, sd_u0i = 0.5)
dat %>%
    mutate(condition = fct_cross(
        factor(expert), factor(advice_present), factor(advice_correct))) %>%
    mutate(condition = fct_recode(condition,
        "no expert, no advice" = "0:0:0", "expert, no advice" = "1:0:0",
        "no expert, wrong advice" = "0:1:0", "expert, wrong advice" = "1:1:1")) %>%
    ggplot(aes(x = y_prob, y = condition)) +
    stat_histinterval(point_interval = "mean_qi", slab_color = "gray45") +
    scale_x_continuous(breaks = seq(0, 1, 0.1), limits = c(0, 1))
```

383

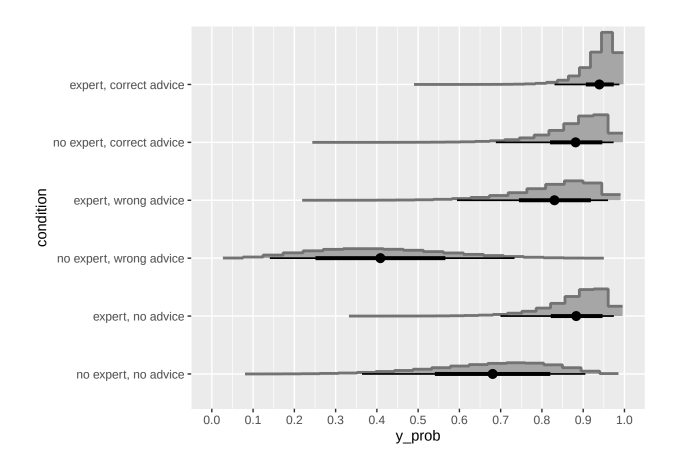


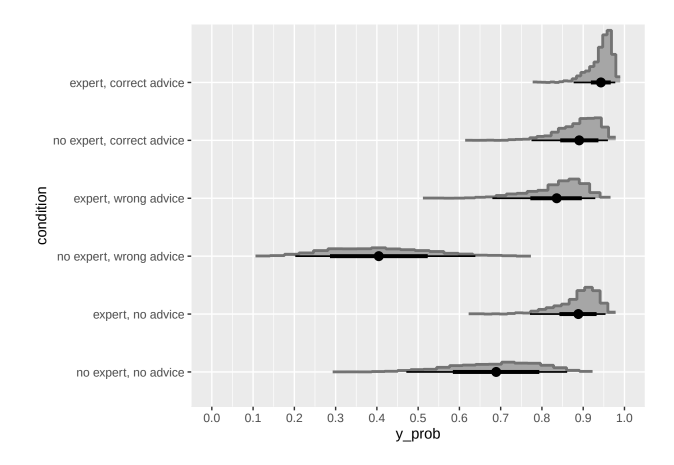
Figure X shows the model implied marginal distributions, including the mean, 66% and 95% intervals. We can see that indeed, the average probabilities (block dots) differ from the probabilities of average subjects and items considered in the previous section.

This difference increases with the variability of the random effects.

In fact, up to this point we have not talked about plausible values for the standard deviations of the subject and item random intercepts ( $\sigma_S$  and  $\sigma_I$ ). Plots like the one above are a useful tool to decide whether the specified standard deviations are reasonable, by comparing the ranges and overlap between conditions to domain knowledge.

In the next plot, we have set the item standard deviation to almost zero ( $\sigma_I = 0.01$ ).

This gives us a better way to see the variability between persons.

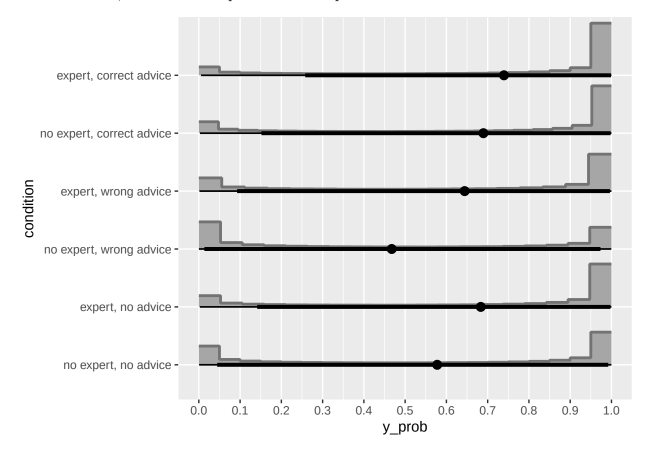


As an example, this presentation reveals a number of implicit assumptions about the comparison between experts and non-experts: With wrong advice, virtually all experts have a higher probability to make a correct diagnosis compared to non-experts when considering only items with average difficulty. In contrast, there is considerable overlap in probability between experts and non-experts with no advice and even higher overlap with correct advice. Patterns like these should be considered carefully, discussed with the domain experts and parameter values ( $\beta$  parameters, and  $\sigma_s$ ) should be adjusted if the implications do not seem reasonable.

We could also have a closer look at variability between items, by setting the subject standard deviation to almost zero ( $\sigma_S = 0.01$ , see Figure X in Appendix X).

The final plot demonstrates, that these plots are also useful to spot standard deviations specified to high. For example, if we set  $\sigma_S = 3$  and  $\sigma_I = 3$  this implies that in

each experimental condition, the probabilities that a subject solves an item are always close to either 0 or 1, which is not a plausible assumption.



410 Results

#### 411 Power simulation

409

With all these considerations out of the way, we are finally ready to perform a power simulation. Wrapping the simulate function already introduced earlier, the helper function sim\_and\_analyse performs all previous steps (simulate a dataset, fit a GLMM, compute p-values) in a single command.

```
sim_and_analyse <- function(
  formula_chr = "y_bin ~ 1 + expert + advice_present + advice_correct +
      expert:advice_present + expert:advice_correct + (1|subject) + (1|item)",</pre>
```

```
null hypotheses = c("advice present + advice correct +
    expert:advice present + expert:advice correct <= 0",</pre>
    "advice present + advice correct <= 0",</pre>
    "-1 * (advice present + expert:advice present) <= 0",
    "-1 * (advice present) <= 0"), ...){
  require(lme4)
  require(multcomp)
  # simulate data
  dat <- simulate(...)</pre>
  # fit model
  model <- glmer(as.formula(formula chr), data = dat, family = "binomial")</pre>
  # compute p-values
  glht <- glht(model, linfct = null hypotheses)</pre>
  pvalues <- summary(glht, test = univariate())$test$pvalues</pre>
  setNames(pvalues, paste0("p_H0", 1:length(null_hypotheses)))
}
```

Power analysis can quickly become computationally intensive, when we repeatedly
simulate data and fit model for different parameter combinations or sample sizes. Here, we
use the future and furr packages to perform computations in parallel. First, we enable
parallelization and specify how many parallel cores of our computer to use (users can find
out the maximum number of cores on their computer with the command
parallel::detectCores()), and set a seed to make the simulation reproducible.

```
library(future)
plan("multisession", workers = 6)
set.seed(2)
```

The next code chunk, specifies a simulation design with different settings for both the number of subjects (n\_subjects) and the number of items (n\_items), each combination being repeated rep times.

The result of the computation is a data frame that contains the p-values of all tested hypotheses for each simulated dataset.

For our exemplary combined hypothesis, power is defined as the (long-run)

percentage of simulations, in which all four p-values of our component hypotheses are

significant at the  $\alpha = 0.05$  level. Based on our simulation outcomes, we compute a power

estimate for each combination of n\_subjects  $\times$  n\_items (including 95% confidence

intervals) and visualize the results with the following code (which is heavily inspired by the

"Mixed Design Simulation" vignette of the faux package at

https://debruine.github.io/faux/articles/sim\_mixed.html).

```
library(binom)
alpha <- 0.05

power <- sim_design %>%

group_by(n_subjects, n_items) %>%
```

```
summarise(power = mean(p HO1 < alpha & p HO2 < alpha &</pre>
                            p H03 < alpha & p H04 < alpha),
    n_sig = sum(p_H01 < alpha & p_H02 < alpha &</pre>
                  p H03 < alpha & p <math>H04 < alpha),
    n = n(),
    ci.lwr = binom.confint(n_sig, n, method = "wilson")$lower,
    ci.upr = binom.confint(n sig, n, method = "wilson")$upper,
    .groups = "drop")
power %>%
  mutate(across(c(n subjects, n items), factor)) %>%
  ggplot(aes(n subjects, n items, fill = power)) +
  geom tile() +
  geom text(aes(label = sprintf("%.2f \n [%.2f; %.2f]",
                                 power, ci.lwr, ci.upr)),
    color = "white", size = 6) +
  scale fill viridis c(limits = c(0, 1))
```

As should be the case, power estimates increase with both the number ob subjects
and the number of items. The confidence intervals indicate how precisely power was
estimated by our simulation. Higher precision (which would be reflected in narrower
confidence intervals) could be obtained by increasing the number of repetitions (rep) in the
simulation. In practice, power simulations are often run multiple times with adjusted
combinations of sample sizes. When run for the first time, it might be revealed that power
is way too low (or too high) for some combinations of n\_subjects and n\_items. When
narrowing down the best combination that achieves sufficient power, while at the same
time strikes a good balance of how many subjects and items are practically feasible, later
rounds of power simulations will typically include a smaller grid of sample sizes combined

with a higher number of repetitions. This will assure high precision for the final power estimates which are then used for sample size justification of the future study.

Much has been written on the optimal amount of power to target in empirical 446 research. The most prominent heuristic is to target a power of 0.8 (when combined with a 447 type I error rate of  $\alpha = 0.05$ ), but depending on the research goals of the study, there are 448 often good reasons to move away from this standard. When target power has been specified, the number of subjects and the number of items in our study design can be traded against each other, based on practical considerations. For the sake of the example, let the targeted power be indeed about 0.8 to detect an effect of the expected size. This 452 could be achieved by collecting data of 200 subjects (about 25% of which will be experts) each completing the same 50 items (with advice present in about 67% of cases which is correct in about 80% of cases with present advice). If collecting 200 subjects is not 455 practically feasible, it would also be possible to recruit 150 subjects but increase the length 456 of the experiment to over 70 items. However, 70 items might take to long to complete 457 (especially for the expert radiologists who might have a busy schedule). The simulation 458 suggests that it might also be possible to plan an even shorter experiment with only 30 459 items, if it is feasible to recruit an even higher number of subjects (> 250, to be determined 460 by additional rounds of power analysis). Design parameters which also affect power and 461 which could be investigated in the simulation to find a more optimal trade-off are the ratio 462 of experts, the frequency of whether advice is presented and whether it is correct. 463

Discussion

Human-AI interaction research requires meticulous planning and consideration of
statistical power to ensure reliable and meaningful results. While heuristics and helper
programs can be useful for simple designs and models, they often fall short when more
complex and customized simulations are required. In this paper, we discuss the necessity of
bespoke power simulations in human-AI interaction research, their implications for research

- design, and the importance of teaching these skills to Human-AI interaction researchers.
- Furthermore, we highlight the discrepancy between the perceived effort involved in
- 472 performing bespoke power simulations and the actual effort required. Finally, we
- emphasize the value of power simulations as research contributions and their alignment
- with open science and preregistration practices.
- 1. The Need for Power Simulations in Human-AI Interaction Research: Human-AI interaction research often involves intricate designs and complex models that cannot be adequately addressed by heuristics or simple helper programs. Bespoke power simulations offer a solution by providing Human-AI interaction researchers with a tailored approach to estimating statistical power. These simulations take into account the specific study design, account for the underlying assumptions, and offer more accurate power estimates.
- 2. Managing Simulations with Discrete Predictor Variables: Bespoke power simulations become more manageable when all predictor variables are discrete and fixed by the study design. This allows Human-AI interaction researchers to focus on simulating outcome variables while avoiding the need to simulate predictor values, which would introduce additional assumptions. By simplifying the simulation process, researchers can obtain reliable power estimates without compromising accuracy.
- 3. Teaching Bespoke Power Simulation Skills to Human-AI Interaction Researchers: 488 The ability to conduct bespoke power simulations is a valuable skill that should be 489 taught to Human-AI interaction researchers. By incorporating this training into 490 research methods courses and workshops, researchers can gain a deeper 491 understanding of statistical power and improve the quality of their experimental 492 designs. Equipping Human-AI interaction researchers with the knowledge and tools 493 to perform be power simulations empowers them to make informed decisions 494 and enhance the rigor of their studies. 495

- 4. Addressing the Mismatch in Effort Perception: There is often a significant disconnect between the perceived effort required to perform bespoke power simulations and the actual effort estimated by researchers and collaborators in Human-AI interaction research. Many content researchers and collaborators request power simulations from statisticians or methodological experts without fully comprehending the complexity and time-consuming nature of these simulations. It is crucial to raise awareness about the effort involved in bespoke power simulations to ensure realistic expectations and effective collaboration between researchers and methodological experts.
  - 5. Recognizing the Value of Bespoke Power Simulations: Bespoke power simulations are not mere technicalities; they are valuable research contributions that deserve recognition in Human-AI interaction research. They offer insights into the reliability and sensitivity of experimental designs, helping researchers make informed decisions about sample sizes, effect sizes, and statistical power. The importance of bespoke power simulations can be reflected by allocating them a separate publication or incorporating them as a significant component of stage 1 preregistered reports.
  - 6. Integration with Open Science and Preregistration Practices: Bespoke power simulations align well with the principles of open science and preregistration in Human-AI interaction research. When researchers have access to simulated data based on their prespecified model, analyzing the collected dataset becomes straightforward and unambiguous. By preregistering their power simulations, researchers enhance transparency and accountability in their experimental procedures, contributing to the credibility and reproducibility of Human-AI interaction research.

### Expected effect size vs. smallest effect size of interest: sensitivity power analysis

In our case study, we have performed power simulations based on a single set of parameter values that reflect our assumptions of an expected effect size. Instead of

extracting this expected effect size from meta-analyses or pilot data which has been the 522 main focus of previous tutorials, we have demonstrated some strategies to determine 523 plausible parameter values in GLMMs based on domain knowledge which can be considered 524 a vague theoretical model about the data-generating process. When sample sizes are 525 chosen based on the results of our power simulations, a future study will be informative to 526 reject the null hypothesis if an effect of our expected size is present. However, if the true 527 effect is indeed smaller, power will be lower the study might not be sufficiently informative. 528 A common, more conservative strategy for sample size justification is to perform power 529 analysis for a smallest effect size of interest (SESOI). An effect smaller than the SESOI 530 would be considered too small to be interesting or practically meaningful, even if the effect 531 is not actually zero. For strategies on the even more difficult task of specifying a plausible 532 SESOI as well as a thorough discussion of various topics concerning power analysis, see (lakensImprovingYourStatistical2022?). When domain knowledge or formal theories about the research topic of interest are too vague to specify a meaningful SESOI, it is still 535 recommended to demonstrate power for different effect sizes in what is called sensitivity 536 power analysis. By simulating power for different effect sizes (in addition to the different 537 number of subjects and items), one can make sure that power would still be sufficient to 538 detect smaller effect sizes than our expected effect, or at least get an impression how 539 strongly power depends on the size of the true effect. In simple study designs, it is possible 540 to perform sensitivity power analysis based on a single standardized effect size (e.g. analyse 541 power in a two-sample t-test for a standardized mean difference varying between 0.1 and 542 0.8). However, for our case study that investigates combined hypotheses in a GLMM 543 modeling framework, the effect size is implicitly represented by the complex distribution of 544 probabilities within and between experimental conditions. In this setting, sensitivity power 545 analysis would require to manually specify additional sets of plausible parameter values 546 that reflect scenarios with smaller or larger differences between groups with respect to our 547 specific research question. Power could then be simulated for several of these scenarios

(across different number of subjects and items as considered earlier).

550 Conclusion

Bespoke power simulations play a critical role in human-AI interaction research,
allowing researchers to tailor power estimation to the unique aspects of their experiments.
The skills required to perform these simulations should be taught to Human-AI interaction
researchers, fostering a deeper understanding of statistical power and enhancing research
design. It is essential to bridge the gap between perceived and actual effort associated with
power simulations and recognize their value as research contributions. By integrating
bespoke power simulations with open science and preregistration practices, researchers can
improve the robustness and transparency of their findings, advancing the field.

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